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PROTEOMICS OF *BACTEROIDES*
FRAGILIS AND *ENTEROBACTER*
CANCEROGENUS

A thesis submitted in partial fulfilment of
the requirements of the University of
Northumbria at Newcastle for the degree
of Doctor of Philosophy

LAKSHMY MANICKAN

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Abstract

Bacteroides fragilis NCTC 9343 is a Gram-negative anaerobic bacterium with genomic DNA of 5205 Kb and a GC ratio of 43%. It is a commensal organism that can act as an opportunistic pathogen and is commonly present on the mucous membranes. It causes a variety of infections including intra abdominal infections, perirectal abscesses and decubitus ulcers. Enterotoxigenic forms are capable of causing diarrhoea in children and animals.

Enterobacter cancerogenus ATCC 35316 is also a Gram-negative facultatively anaerobic bacterium with genomic DNA of 4602 Kb and a GC ratio of 55%. It is a naturally occurring human gut symbiont known to exhibit resistance to antibiotics like aminopenicillins. It has also been reported in cases of severe osteomyelitis and infections of bones and joints.

This study aims to analyse the differential expression of proteins in the presence of mucin since it serves as the first site of adherence for the bacteria. The *E. cancerogenus* and *B. fragilis* proteins were extracted and separated by two dimensional electrophoresis from logarithmic phase cultures grown in semi-defined media enriched with or without porcine gastric mucin Types II and III. The gel images were analysed using Bio-Rad PDQuest, Ludesi Redfin and Nonlinear Dynamics SameSpots softwares. It was observed that the presence of mucin in the media affected the expression of a number of proteins in *E. cancerogenus* and *B. fragilis* cells. The protein spots of interest were excised, hydrolysed using trypsin and subjected to electrospray ionisation based LC-MS analysis in order to determine the identity of the digested proteins and obtain a better understanding of the interactions of *B. fragilis* and *E. cancerogenus* with mucin.

The outer membrane protein surface antigen X was found to be up-regulated in both mucin Type II and III enriched media in *E. cancerogenus*. Some of the other proteins that were differentially regulated in both *E. cancerogenus* and *B. fragilis* included the elongation factor Ts, malate dehydrogenase, triose phosphate isomerase and thiol peroxidase proteins indicating that these proteins may be associated with the ability of bacteria to grow in mucin and may be potential virulence factors.

Genes encoding the proteins CAH06598 and CAH09443 from the glycoside hydrolase families 95 and 97 in *B. fragilis* strain NCTC9343 were cloned, overexpressed and purified using nickel affinity and gel filtration chromatography. The enzymes were found to be active by performing fluorimetric assays using methyl-umbelliferyl sugar substrates. Diffracting crystals of CAH09443 were obtained from the PACT ANION screens containing polyethylene glycol and sodium malonate as a precipitant. Structure determination was achieved via molecular replacement using the glycoside hydrolase Family 97 α -galactosidase, *BtGH97b*, from *Bacteroides thetaiotaomicron* as a starting model. The structure of CAH09443 was shown to be composed of a N-terminal β -super-sandwich domain and a canonical $(\beta/\alpha)_8$ barrel, similar to the two other glycoside hydrolase family 97 enzyme structures reported.

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I would like to dedicate this thesis to my loving parents and sister, Lavanya without whose unconditional support, help and encouragement, this work would have been impossible. Finally, I thank Northumbria University for making this experience a special and memorable one.

Abbreviation list

APS	Ammonium persulphate
bp	Base pair
BSA	Bovine serum albumin
CBM	Carbohydrate binding module
CFE	Cell free extract
Da	Dalton
dNTP	deoxynucleotide triphosphate
DTT	Dithiothreitol
EDTA	Ethylene diamine tetraacetic acid
GH	Glycoside hydrolase
HEPES	N-[2-hydroxyethyl] piperazine-N'-[2-ethanosulphonic acid]
IMAC	Immobilised metal affinity chromatography
IPTG	Isopropyl- β -D-thiogalactopyranoside
kb	Kilobase pair
LB	Luria Bertani
PAGE	Polyacrylamide gel electrophoresis
PCR	Polymerase chain reaction
SDS	Sodium dodecyl sulphate
TAE	Tris-acetate-EDTA
TEMED	N,N,N',N'-tetramethylethylene diamine
TRIS	tris(hydroxymethyl)aminomethane
v/v	volume per volume

w/v	weight per volume
ACN	Acetonitrile
ATP	Adenosine tri phosphate
BHI	Brain Heart Infusion
BPB	Bromophenol blue
BPC	Base peak chromatogram
CHES	N-cyclohexyl-2-aminoethane sulphonic acid
DHAP	Dihydroxy acetone phosphate
ESI	Electrospray ionisation
EF	Elongation factor
IAA	Iodoacetamide
MES	2-(N-morpholino)-ethanesulphonic acid
MPD	2-Methyl-1,3 propanediol
MS	Mass spectrometry
FA	Formic acid
GDP	Guanosine di phosphate
G-3-P	Glyceraldehyde-3-phosphate
GTP	Guanosine tri phosphate
IEF	Isoelectric focussing
LC	Liquid chromatography
CAZY	Carbohydrate active enzymes
dCMP	deoxy cytidine mono phosphate
MPD	2-methyl-1,3 propanediol

NAD	Nicotinamide adenine dinucleotide
NADP	Nicotinamide adenine dinucleotide phosphate
NCBI	National Centre for Biotechnology Information
Omp	Outer membrane protein
PEG	Polyethylene glycol
PEG MME	Polyethylene glycol monomethyl ether
PEPCK	Phosphoenol pyruvate carboxykinase
PBS	Phosphate buffered saline
PFL	Pyruvate formate lyase
SOB	Super optimal broth
SOC	Super optimal culture
SSP	Standard spot numbers
TIC	Total ion current
TPI	Triose phosphate isomerase

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NB: * Tables 5 to 10 have been included in the Appendix section E.

DECLARATION

I declare that the work contained in this thesis has not been submitted for any other award and that it is all my own work.

NAME: LAKSHMY MANICKAN

SIGNATURE:

DATE: 26th April 2010

4. General introduction

The study aims at understanding the interaction between gut commensal bacteria and mucin. The GI tract is colonised by a number of bacteria which are opportunistic pathogens that can cause infection when the host immune system is impaired. This work mainly focuses on the gut infection causing bacteria, *Enterobacter cancerogenus* strain ATCC 35316 and *Bacteroides fragilis* strain NCTC 9343.

4.1 Bacteroidaceae

Based on the comparative analysis of the 16S rRNA sequences, *Bacteroides* species have been classified under the family 'Bacteroidaceae' and phylum 'Bacteroidetes'. They play an important role in the maintenance of human and animal health by colonising the intestine and forming an indigenous flora of the colon. They have been known to colonise the distal regions of the stomach where the gastric secretions are diluted and the hindgut beginning from the distal regions of the small intestine with increasing numbers in the terminal ileum and colon. They help in the fermentation of carbohydrates in the colon where the fermentation activity is very high at the caecum. As the carbohydrate sources become limited, the activity is slowed down in the transverse colon and directed towards protein fermentation (Smith *et al.*, 2006).

4.2 Introduction to *Bacteroides fragilis*

Bacteroides fragilis is classified as a Gram-negative anaerobe which is included in the family of Bacteroidaceae. They commonly occur on mucous membranes and are responsible for endogenous infections. *Bacteroides* species cause more than 50% of anaerobic infections. They normally range from 0.5 - 0.6 μm in diameter to 1.5 - 4.5 μm in length and are found in nature as round ended bacilli (Fig. 1). They have a genome size of 5205 Kb with a guanine+cytosine (GC) ratio of 42% (Brook, 2006). They are opportunistic pathogens present in the gastrointestinal flora and cause a number of intra-abdominal infections.

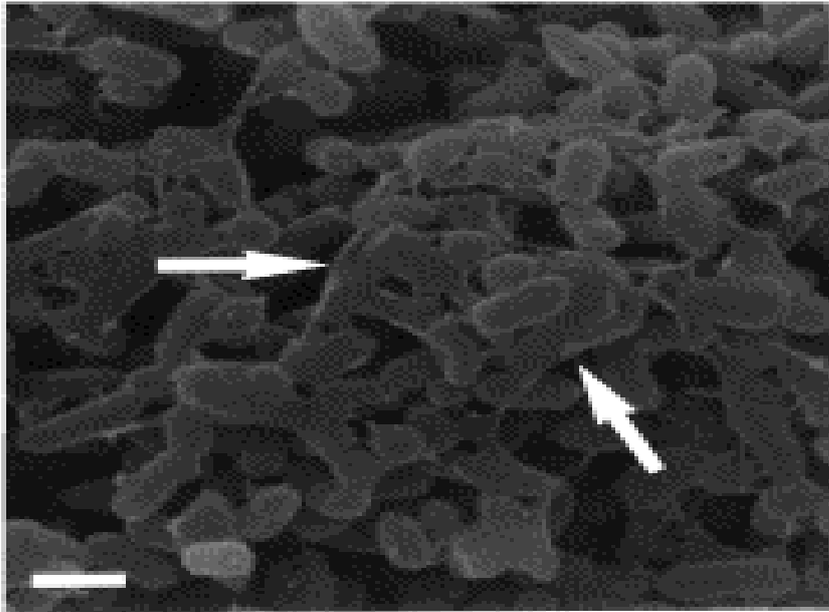


Figure 1: An electron micrograph showing aggregates of *B. fragilis*. The bar refers to a size of 1 μ m (Ferreiraa *et al.*, 2002).

They have also been found to exhibit resistance to a variety of antibiotics including penicillin and the β -lactam antibiotics because they have the ability to produce the enzyme, β -lactamase which attacks the β -lactam ring within the antibiotic. They have also shown multidrug resistance to other antibiotics like clindamycin, tetracycline and metronidazole (Su & Honek, 2007).

4.2.1 Types of *Bacteroides fragilis* infections

B. fragilis is a Gram-negative bacterium that is capable of causing a variety of infections within the host. These include central nervous system infections, intra-abdominal and pleuropulmonary infections, skin and soft tissue infections, female genital tract infections, bacteraemia, osteomyelitis, abscesses and septic arthritis (Pumbwe *et al.*, 2006). *B. fragilis* is commonly present on the mucous membranes and causes a variety of infections including intra abdominal infections, perirectal abscesses and decubitus ulcers. The enterotoxigenic forms are capable of causing diarrhoea in children and animals (Nakano *et al.*, 2007).

4.2.2 Uses of *Bacteroides fragilis* to man

Bacteroides spp. are also beneficial to man in a number of ways. They are capable of producing certain vitamins like Vitamin K2 and menaquinone which are utilised by the human host.

Anaerobic growth of *Bacteroides* species results in the production of a number of end products such as butyrate, acetate and propionate and this contributes to nearly 70% of the energy source for enterocytes, a distinct type of epithelial cell that constitutes the innermost layer of the large and small intestine and helps to transport molecules into the tissues after breaking them down, present in the colon.

They play a very valuable role in the enterohepatic cycles which include the bile acid recirculation and the bile acid transformation by producing the enzyme, bile salt hydrolase (BSH) (Stellwag & Hylemon, 1976).

Bile acid recirculation involves the absorption of bile acids by the hepatocytes and this process is repeated several times within the intestine to make sure that the maximum amounts of nutrients are absorbed. The presence of anaerobic bacteria help to enhance this process (Hopley & Schalkwyk, 2006).

Bile acid transformation also plays an important role in the regulation of bile and cholesterol in the system. The terminal ileum and large bowel are largely characterised by the presence of those anaerobes which enhance the deconjugation process (Hopley & Schalkwyk, 2006). They compete with other pathogens in the colon thereby developing a competition at food supply and receptor sites by producing acidic end products like acetic acid and lactic acid that lowers the pH of the environment. This regulates the bacterial growth in the intestine.

4.2.3 General characteristics of *Bacteroides fragilis*

The selective medium which can be used to distinguish *B. fragilis* is Bile esculin agar which has the presence of oxgall, ferric ammonium sulphate (colour change indicator) and an antibiotic, gentamycin. Gentamycin and oxgall act as inhibitors of

other facultative anaerobes and Gram-negative anaerobes. Esculin is hydrolysed by *B. fragilis* to produce dextrose and esculetin and this reacts with the salt to produce black complexes which get accumulated around the colony (Livingston *et al.*, 1978).

B. fragilis have the ability to exhibit haemolytic activity. Studies showed that the haemolytic activity diminished in the presence of oxygen or hydrogen peroxide and was controlled by the *hlyA* and *B* genes (Robertson *et al.*, 2006). This was confirmed when haemolytic activity was observed in blood agar plates containing *E. coli* transformed with the *hlyA* and *B* genes.

4.2.4 Cell wall structure of *Bacteroides fragilis*

B. fragilis has a distinct outer capsular layer made up of polysaccharides A and B (Fig. 2) (Pruzzo *et al.*, 1989). The capsular layer confers resistance to phagocytosis, assists in adherence to the host and also evades complement mediated lysis (Pumbwe *et al.*, 2006).

The second layer, termed as the lipopolysaccharide layer, projects out from the outer membrane of the lipid layer. This can be of varying thickness exhibiting a considerable level of toxicity and also functioning as an adhesive when contacted by host cells.

They have the presence of fibrils which act as short fine appendages whose role in virulence is not yet determined even though it is suspected to be related to adhesion and biofilm formation. The next important component of the cell wall includes the outer membrane vesicles. They are known to produce endotoxins like neuraminidase, sialidase and fibrinogenolytic enzymes (Pumbwe *et al.*, 2006). The fibrinogenolytic enzymes, chondroitin sulphatases and hyaluronidases are histolytic in nature and *B. fragilis* is also capable of producing antimicrobial chemicals, namely bacteriocins. Neuraminidase exhibits a hydrolytic activity on the mucin polysaccharide. One of the previous studies showed that neuraminidase activity plays a very important role in the successful survival of *Bacteroides* in mammalian

tissues thereby acting as one of the major virulence conferring antigens (Godoy *et al.*, 1993).

Studies suggest that *B. fragilis* produces outer membrane proteins which play an important role in maintaining the structure of the cell and the production of these proteins are controlled by a set of 4 *omp* genes (Wexler, 2002).

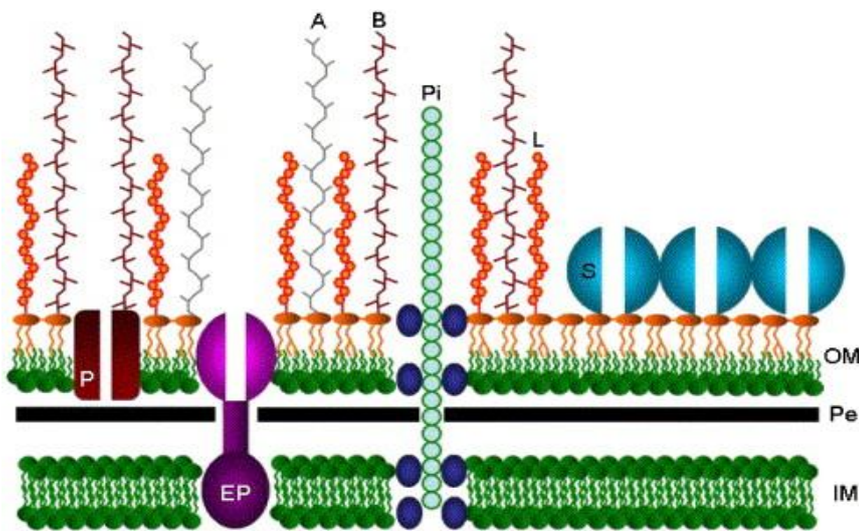


Figure 2:Cell wall structure of *B. fragilis* (Pumbwe *et al.*, 2006)

Pe- periplasmic space, OM- outer membrane, L- side chains, A and B- Capsular polysaccharides, Pi- Pilus, P- outer membrane porin, EP- efflux pump, IM- inner membrane.

4.2.5 Types of *B. fragilis* toxins

B. fragilis produces two main types of toxins,

- The endotoxin from the outer membrane vesicles and
- The enterotoxin whose production is controlled by the regulation of three *bft* genes, *I*, *II* and *III*. They can cause diarrhoea within the host by damaging the tight junctions of the intestine. Actin cytoskeleton rearrangement of the

epithelial cells causes a leakage in the internal contents of the intestine (Nakano *et al.*, 2007).

- The *bft* genes also induce a fluid secretion in the intestinal epithelia (Pumbwe *et al.*, 2006).
- *B. fragilis* also produces proteins termed as bacteriocins which inhibit the growth of other bacteria by targeting their RNA polymerase thereby providing the bacteria a better chance of survival during competition from other pathogens (Stein, 1998).

4.2.6 *Bacteroides fragilis* virulence factors

Some of the virulence factors produced by *B. fragilis* include capsules where capsulated strains have been shown to be more virulent in nature (Pruzzo *et al.*, 1989). They produce several extracellular or membrane bound enzymes like collagenases, fibrinolysin, proteases, lipases, ribonucleases and deoxyribonucleases involved in tissue degradation and destruction. The other virulence factors are the oxygen scavenging enzymes namely superoxide dismutases, catalases and peroxidases which enable the bacteria to survive in the presence of oxygen or in aerobic environments (Namavar *et al.*, 1991). They produce endotoxins that are transported to the target host sites and induce complement activation. The capsular polysaccharides inhibit phagocytosis and induce abscess formation, the charged surface ligands help in adherence and bacterial interaggregation sometimes promoting the formation of biofilms of the same or different bacterial communities. The IgA protease causes impairment of secretory and mucosal immunity and heparinase promotes coagulation with tissue ischemia. Some strains of bacteria are capable of producing β -lactamases which confer resistance to β lactam antibiotics. Bacteriocins and other secondary metabolites including fatty acids, hydrogen sulphide and ammonia competitively inhibit the growth of normal flora (Stein, 1998).

Studies in *Bacteroides forsythus* have revealed the presence of a S-layer which is a unique surface structure that has been known to be involved in virulence. They help in the adherence of the bacterial cells to the host, surface recognition and resistance to phagocytosis. The S-layer ranges in size from 40-200 kDa in size and is composed of regularly aligned glycoprotein subunits. These proteins could contribute

to the ability of the bacteria to adhere and form biofilms on the surface of teeth resulting in dental plaque (Sabet *et al.*, 2003).

B. fragilis can exist in both capsulated and non-capsulated forms but the capsulated forms tend to have a higher degree of virulence conferring capacity. Their capsules contain two different types of polysaccharides called A and B which are linked together by oppositely charged groups and they confer resistance to the organism against the host defences (Pantosti *et al.*, 1993).

They also produce a number of virulence conferring enzymes which include neuraminidase, hyaluronate lyase, DNase, phosphatase and rare fibrinolysins. They produce the enzyme heparinase which may result in vascular thrombosis (Hopley & Schalkwyk, 2006). They also produce a toxin called fragilysin that is a zinc metalloprotease, 20 kDa in molecular weight. This damages the tight junctions of the intestine resulting in the destruction of enterocytes due to its cytotoxic effects and fluid secretion (Fasano & Nataro, 2004).

4.2.7 Treatment of *Bacteroides fragilis* infections

A number of normal antibiotics can be used for the treatment of *B. fragilis* infections like metronidazole, chloramphenicol, moxalactam, ceftriaxone, clarithromycin, cefoperazone, cefotaxime, ceftazidime, sparfloxacin, carbapenems and clindamycin (Katsandri *et al.*, 2006). However, recent studies showed that certain strains of *B. fragilis* exhibit resistance to carbapenem by producing a β -lactamase enzyme, carbapenemase. This was found to be induced by the activation of the *cfiA* gene with or without the insertion sequence elements indicating that this gene may be involved in the development of antibiotic resistance in *Bacteroides* (Edwards & Read, 2000). They also exhibited a resistance percentage of 0.8 and 1.3 for antibiotics imipenem and meropenem, respectively (Edwards & Read, 2000).

4.3 Enterobacteriaceae

Enterobacter species are capable of causing a number of infections ranging from bacteraemia, lower respiratory tract infections, skin and soft tissue infections to endocarditis, urinary tract infections, intra abdominal infections and post surgical wound infections. One of the major characteristic of Gram negative bacteria is their ability to produce lipopolysaccharides. Enterobacteriaceae produce lipopolysaccharides which are composed of three parts namely the oligomeric repeating O polysaccharide units, a core and lipid A. The core links the O polysaccharide units to the lipid A and this causes the lipopolysaccharide to remain anchored to the cell envelope. O polysaccharides exhibit structural variation which induces inflammatory responses in the host. The toxin is also capable of causing sepsis within the host which later leads to the development of *Enterobacter* bacteraemia characterised by cyanosis (Janda & Abbott, 2006). Biochemical characteristics reveal that these bacteria are straight rods, Gram negative, facultatively anaerobic and exhibit both respiratory and fermentative type of metabolism. They are citrate positive, indole negative and are chemoorganotrophic in nature (Bergey & Holt, 1994).

4.3.1 General virulence factors of *Enterobacter* species

Some of the major virulence factors associated with *Enterobacter* species are the mannose sensitive haemagglutinins (MSHA) type 1 or 3, siderophores, toxins and outer membrane proteins. The MSHA are putative fimbriae of 35 kDa size and have a receptor site that is recognised as a high mannose oligosaccharide. Siderophores have been known to be associated with invasive systemic infections. *E. cloacae* produces hydroxamate siderophore aerobactins that may be associated with invasive diseases whereas *E. coli* has been known to produce a 72 kDa aerobactin associated protein. Certain non-aerobactin hydroxamate compounds have also been found to be infrequently expressed in *Enterobacter* species. Details about their role in pathogenicity remain unknown (Keller *et al.*, 1998). *Enterobacter* species also produce toxins like α - haemolysin which have a glycine rich motif and exhibit cytotoxic activity against human erythrocytes and leukocytes. Molecular characterisation of invasive infections caused by *E. cloacae* showed an up-regulation

of the outer membrane protein OmpX suggesting that this could be a potential virulence factor, 17 kDa in size. It was observed that an up-regulation of *ompX* was accompanied by a down-regulation of other outer membrane porins OmpF and OmpC and may be associated with the ability of the organism to exhibit resistance to β -lactam compounds. Bacterial strains lacking the *omp X* gene were constructed to study their role in pathogenicity and it was observed that these proteins increased the infection rate by 10 fold in non mutant strains (Janda & Abbott, 2006).

4.3.2 *Enterobacter cancerogenus*

Enterobacter cancerogenus is also a Gram-negative facultatively anaerobic bacterium that is a naturally occurring human gut symbiont. They have a GC ratio of 55% with a size of 4602 Kb and are known to exhibit resistance to antibiotics like aminopenicillins. They have also been reported in cases of severe osteomyelitis and infection of bones and joints and belong to the CDC Enteric group 19.

This bacterium was formerly known as *Erwinia cancerogena* and several morphological, physiological and biochemical studies revealed that *E. cancerogena* was not a member of the genus *Erwinia* but was included in the genus *Enterobacter*. *E. cancerogenus* are capsulated, motile straight rod shaped bacteria with peritrichous flagella. *E. cancerogena* was found to exhibit 75% phenotypic similarity to *Erwinia carotovora* and 92% phenotypic similarity to *Enterobacter nimipressuralis*. Hence it was proposed that *E. cancerogena* could be moved to the genus *Enterobacter* based on phenotypic characterisation.(Dickey & Zumoff, 1988).

They are commonly found in nature and occur as commensals but when the host immune system is impaired they are capable of causing infections which make them opportunistic pathogens. Five cases of infection were reported in 1997. *E. cancerogenus* was identified from blood culture tests where wound infections and septicaemia occurred following environmental exposure of wounds from traumatic events or injuries in adults (Abbott & Janda, 1997).

Studies revealed that *E. cancerogenus* was formerly known as *Enterobacter taylorae* after being moved from the genus *Erwinia*. Based on the evidence obtained from the study of the biochemical and physiological properties of *E. taylorae*, it was proposed

that the organism needed to be included in the *Enterobacteriaceae* family. They were able to ferment glucose to produce gas and showed 84- 95% similarity to ATCC 35317 strain (Farmer *et al.*, 1985). There was a very strong relation between these two bacteria where they had a high degree of DNA relatedness. Since no difference could be ascertained between these two species, it was accepted that *E. cancerogenus* was a senior synonym of *E. taylorae* (Grimont & Ageron, 1989).

Even though it is known that these bacteria are capable of causing infections and occur predominantly as gut microbiota, details regarding their virulence factors or mode of action are not available. Several cases of infection have been reported since 1987 and the bacteria have been included in the Proteobacteria division and isolated mainly from human faeces. Infections occurred following a traumatic injury and the bacterium was found to exhibit resistance to aminopenicillins (Garazzino *et al.*, 2005). Four cases of nosocomial infections were reported in 1989 where all the cases described localised infection following open fractures, wounds or abrasions. The cases reported were bacteraemia, pneumonia and urinary tract infections. The bacteria were also found to produce β -lactamases showing resistance to antibiotics like penicillin and cephalosporins. Administration of cephalosporin prophylactically during cardiac surgery induced a drastic increase in the number of *Enterobacter* species in the intestine of the patient (Rubinstien *et al.*, 1993). Another reported case of *E. taylorae* infection was osteomyelitis in a patient following an open fracture wound being infected (Westblom & Coggins, 1987). The most recent occurrence of *E. cancerogenus* infection was reported in 2005 where the infection occurred following a multiple bone fracture with abundant environmental exposure (Garazzino *et al.*, 2005).

4.3.3 *Enterobacter* species attachment to intestinal cells

The outer membrane protein A has been known to play an important role in the attachment of *E. sakazakii* to the epithelial cells of the intestine of human hosts according to previous studies (Nair & Venkitanarayanan, 2007). Even though no reports are available regarding the attachment of *E. cancerogenus* to intestinal epithelial cells, *E. cloacae* which belongs to the same genus has been known to be associated with host intestinal tissues and mucin (Schierack *et al.*, 2007).

4.3.4 Antibiotic treatment

The commonly used antibiotic treatment for *Enterobacter* infections includes carbapenems and cefepime. Cross transmission contributes to some of the major factors that result in the spread of the bacteria. They have been predominantly found in intensive care units especially if infection control is poor.

4.4 Epithelial cells of the intestine and their role in protection against antigens

There are three types of cells that are produced to confer resistance to bacteria. This includes the goblet cells that produce proteoglycans and glycoproteins in the form of viscous mucin that covers the entire surface of the wet intestinal epithelial layer; the M cells that help to transport antigens from the exterior into the lymphoid tissue. This assists the host in identifying bacterial antigens and preparing the immune system. The third type of cells includes the Paneth cells that produce intra epithelial lymphocytes and antibacterial proteins that act against exogenous bacteria (Wilson, 2002). Intestinal cells also protect themselves against bacteria by shedding off the surface epithelia that fill up with keratin. This sloughing off of the outer layers results in the removal of adhered bacteria which get flushed away. Mucosal epithelia also have the ability to produce proteins that recognise lipopolysaccharides present exogenously. This may be produced due to continual exposure to certain specific surface components of bacteria or for identification of potential antigens (Laux *et al.*, 2005).

The entire length of the gastrointestinal tract consists of the intestinal epithelial cells with the outer protective mucous membrane. The upper part of the tract has been known to be colonised by aerobic bacterial flora since the oxygen concentration of the intestine decreases from the duodenum to the colon. The lower part of the intestinal tract is colonised by the anaerobic flora. Variations in pH and nutrition can affect the microflora balance and also determines the general metabolism of the body. Changes in microflora of the gastrointestinal tract have been known to result in metabolic disorders (Serino *et al.*, 2009).

4.5 Mucin

Mucin is a heavily glycosylated protein with a high molecular weight. Mucin forms a major component of the body secretions which include the mucous and secretions of the mucous gland, the saliva and secretions of the salivary gland, fibres of the tendons and the connective tissue. It is albuminoid in nature with a ropy appearance and possesses heavy glycosylation (Dekker *et al.*, 2002). Mucins contain hydrophobic membrane spanning domains that enable them to be retained within the plasma membrane making them membrane bound.

They generally form aggregates of proteins which are about 1-10 million Daltons and are known to contain neuraminic acids in their side chains (Dekker *et al.*, 2002). They have the ability to resist proteolysis and act as the first line of defence against infection.

4.5.1 Structure of mucin

Mucin consists of a number of monomeric units that are linked by O and N linked oligosaccharides. Fig. 3 represents the mucin monomers linked together in an oligomeric gel and Figs. 4 and 5 represent the N and C termini of individual mucin monomers with N and O linked oligosaccharide units, cysteine knot and disulphide rich domains containing disulphide bonds. The disulphide rich domain is also termed as the D domain (Refer to Fig. 5) (Wilson, 2002). Porcine gastric mucin is predominantly made up of O-linked oligosaccharides and mucin side chains contain a number of groups attached to them including sulphates, neuraminic acids, sialic acids and so on.

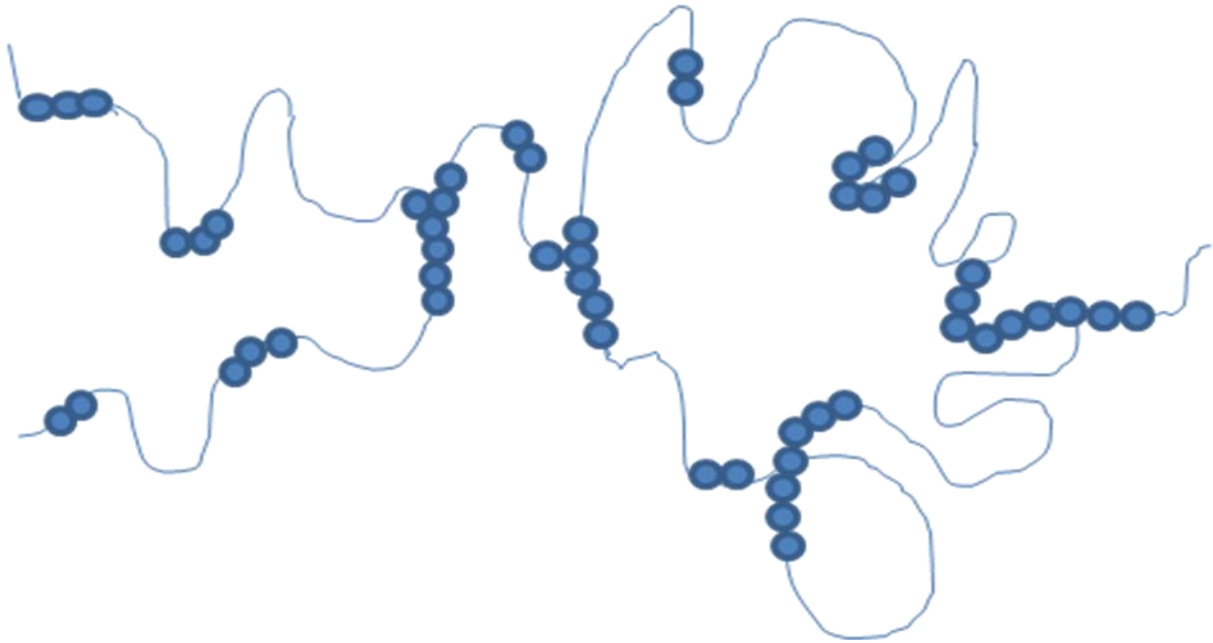


Figure 3: A number of mucin monomers linked together in an oligomeric gel

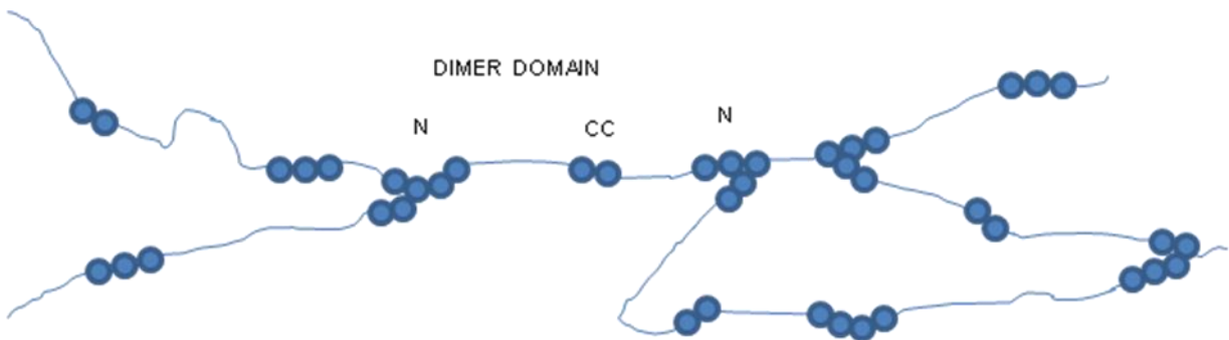


Figure 4: Structure of mucin showing the dimer domain

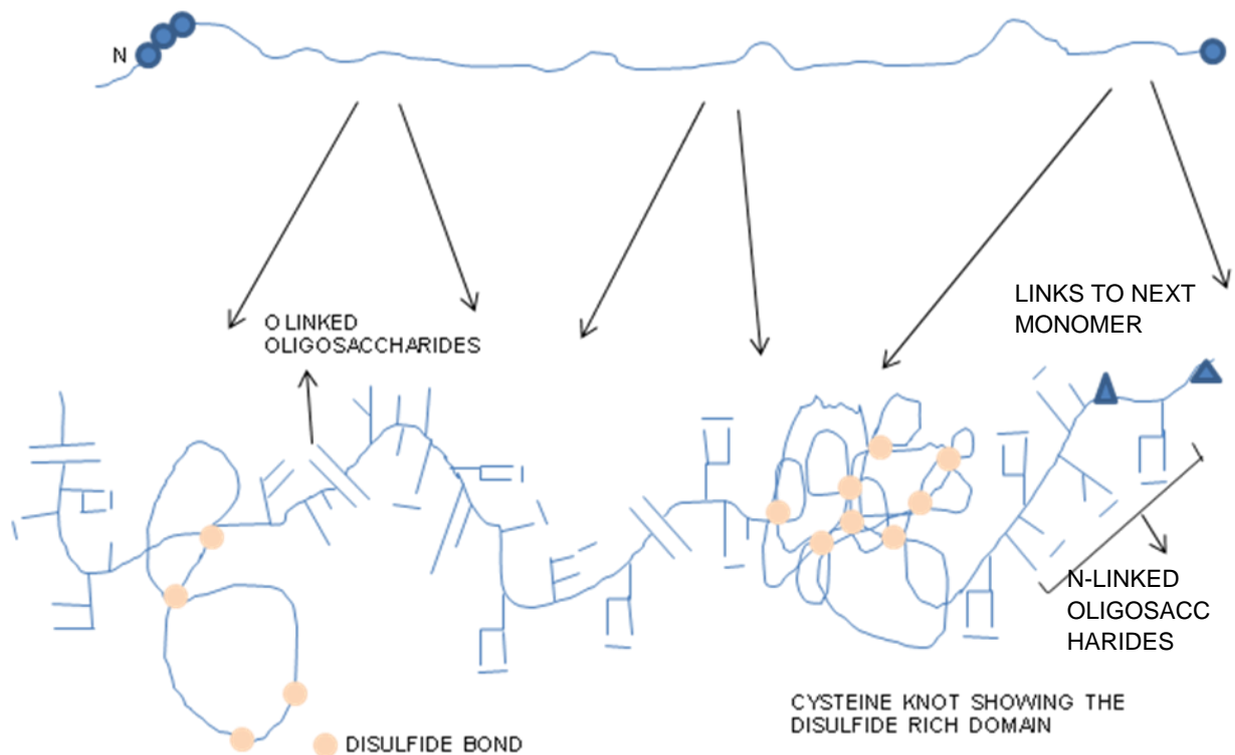


Figure 5: Oligomeric structure of mucin focussing on the monomer linkages and bonds (Wilson, 2002).

Mucins are of two main types namely soluble mucins and membrane mucins. Soluble mucin secretion associated with the respiratory tract is known to be encoded by a cluster of 18 genes that results in the formation of heavily glycosylated disulphide bond rich glycoproteins in the mucous in man (Rose & Voynow, 2006). Membrane mucins are highly diverse structurally and functionally and contain a transmembrane domain that enables them to associate with the cell surface thereby acting as the first line of defense and can mediate transport of solutes. Examples of membrane mucin include the mucin encoded by the *muc1* and *muc4* genes. They function as anti-adhesive agents that prevent cell-cell, cell-molecular interactions. They also act as modulators of cell signalling pathways where *muc1* encodes for mucin that act as docking protein and *muc4* encodes mucin that acts as a modulator of receptor tyrosine kinase ErbB2 (Zhang *et al.*, 2006). Sialomucin complexes have the ability to act as anti-adhesive agents by disrupting cell-matrix interactions and have been overexpressed in cancer cells (Komatsu *et al.*, 1997). This demonstrates the role of mucin and its protective functions within the host.

A study was conducted to study the variation of mucin distribution in the gut. The four main parts that were studied were the small intestine, caecum, colon and luminal contents and three types of mucin were found to predominantly occur in them. These were classified as neutral mucins, sialomucins and sulphomucins of which sialo and sulphomucins were found to be acidic in nature and sulphomucins were sulphated. It was found that conventional rats produced higher amounts to mucin with higher villi and deeper crypts in the small intestine when compared to germ free rats indicating that exposure to bacteria may be one of the main reasons for the production of mucin. It was also observed that conventional rats had decreased amounts of neutral and sulpho mucins in the intestine indicating that bacteria could have a higher hydrolytic activity on them when compared to sialomucins that exhibit higher resistance (Meslin *et al.*, 1999). This could explain the presence of bound sialic acids in the partially purified and crude extracts of mucin from the porcine intestine.

The synthesis of mucin is controlled by a set of mucin genes termed *muc*. Mucin monomers are linked to form aggregates. They have specific non-covalent and intermolecular disulfide bonds to link them. The central region of the gene encodes the synthesis of hundreds of O-linked oligosaccharides of varying lengths whereas the N and C terminals have a much lesser level of glycosylation. The central region also possesses 10-80 repeating units of serine or threonine amino acid residues as shown in the figure below (Salyers *et al.*, 1977).



Figure 6: Structure of the mucin gene '*muc*'

4.5.2 Sugar composition of mucin

Some of the oligosaccharides which have been released upon breakdown of mucin include 2-acetamido-2-deoxy- α -D-galactopyranosyl (1 \rightarrow 3)-[α -L-fucopyranosyl-(1 \rightarrow 2)]-deoxy- β -D-galactopyranosyl-(1 \rightarrow 3)-[N-glycoylneuraminy-(2 \rightarrow 6)]-2-acetamido-2-deoxy-D-galacitol, N-glycolylneuraminy \rightarrow N-acetylgalactosaminitol, N-acetylgalactosaminitol. Other oligosaccharides include GlcNAc β (1-3)GalNAc-ol, Gal β (1-4)GlcNAc β (1-3)GalNAc-ol, Gal β (1-4)GlcNAc β (1-3)Gal β (1-4)GlcNAc β (1-3)GalNAc-ol, Gal β (1-4)GlcNAc β (1-3)GalNAc-ol, GalNAc α (1-3)Gal β (1-4)GlcNAc β (1-6), and GalNAc α (1-3)Gal β (1-4)GlcNAc β (1-3)Gal β (1-4)GlcNAc β (1-3)Gal α (1-4)GlcNAc β (1-3)GalNAc-ol irrespective of the source of mucin (Podolsky, 1985).

The saccharide composition of mucin depends upon the part of the body where it might be produced, nutrition and several other factors including the environmental conditions. Studies were performed in porcine submaxillary mucins based on their ability to inhibit human A- anti A haemagglutination. The breakdown of the complex polysaccharide into oligosaccharides was studied and the resulting oligosaccharides were grouped from I to V. The study aimed at determining the type of mucin oligosaccharides that was found to be active in inhibiting haemagglutination (Carlson, 1968). Breakdown of mucin resulted in the release of the monosaccharide, 2-acetamido-2-deoxy-D-galacitol (N-acetylgalactosaminitol), a pentasaccharide, 2-acetamido-2-deoxy- α -D-galactopyranosyl (1 \rightarrow 3)-[α -L-fucopyranosyl-(1 \rightarrow 2)]-deoxy- β -D-galactopyranosyl-(1 \rightarrow 3)-[N-glycoylneuraminy-(2 \rightarrow 6)]-2-acetamido-2-deoxy-D-galacitol. The other reduced oligosaccharides released include N-acetylglucosamine, N-glycolylneuraminy \rightarrow N-acetylgalactosaminitol and N-acetylgalactosaminitol (Carlson, 1968).

Similar studies were performed in cervical mucin and it was found to contain neutral, sialylated and sulphated oligosaccharides including fucose, galactose, N-acetylgalactosamine, N-acetylglucosamine, N-acetyl-galactosamine and so on. A neutral tetrasaccharide and an acidic trisaccharide were also isolated. It has been observed that there is a high degree of heterogeneity in mucins derived from various parts of the body like bronchial mucin, colonic mucin, cervical mucin, gastric mucin and so on (Yurewicz & Moghissi, 1981).

Previous studies performed in colonic mucin showed the release of sugars like galactosamine, galactose, glucosamine, sialic acids, fucose, mannose and glucose (Podolsky & Isselbacher, 1983).

4.5.3 Functions of mucin

Mucins are present in various parts of the body and play a very important role by acting as protective barriers of the host against pathogenic bacteria. Intestinal mucin helps the system internally by protecting the walls against the activity of strong digestive enzymes. They lubricate the walls of the intestine, exhibit surface hydrophobicity and can reduce the absorption of toxins (Lugea *et al.*, 2000). Innate mucosal defense involves two main components; mechanisms that are involved in preventing the invasion of bacteria and their toxins and the second mechanism involved the rapid regeneration of the mucosal layer by repairing damages and defects. They work in conjunction with the tight junctions of the intestine controlling the passive diffusion of solutes beneficial to the host while regulating the entry of bacterial toxins and antigens (Blikslager *et al.*, 2007). Even though it is known that the epithelial cell layer of the intestine secretes the mucin glycoproteins that form a viscoelastic gel, functional evidence of their protective functions are limited and not available in detail. Recent studies have revealed the presence of a distinctive and highly conserved structural motif containing 3 small proteins called the trefoil peptides. The goblet cells secrete these proteins and they were found to span the entire length of the gastrointestinal tract. The three proteins were designated as PS2, SP and the Intestinal trefoil factor (ITF) of which the first two were found to occur predominantly in the mucosa of the stomach, mouth, hepatobiliary duct and the pancreatic duct whereas the intestinal trefoil factor was found to be predominant in the small and large intestinal mucosa. These proteins contain a highly conserved 6 cysteine residue motif that form three intrachain loops with disulphide bond interactions. It has also been suggested that the occurrence of secondary structures of these proteins on the mucosal layer could confer resistance to the digestive activity of a variety of protease enzymes. This helps to maintain the intestinal walls structurally and functionally intact from the activity of proteases (Podolsky, 1999).

4.5.4 Mucin degradation by bacteria

Bacteria have the ability to produce a variety of enzymes that degrade the mucin glycoproteins. This enables them to adhere efficiently to the host and thereby cause infection within the host. The glycoside hydrolases produced by bacteria, cleave the carbohydrate chains in mucin and hydrolyse them to produce monosaccharide units. These monosaccharides are then used by the organism as a source of energy and nutrition (Robertson & Stanley, 1982).

Mucin, being the main component of mucosal surfaces acts as a point of adherence for the bacteria where they grow upon the glycoside residues of the complex glycoprotein oligomer by producing glycoside hydrolases and this leads to further damage of the mucosal surface and impairment of the membrane function (Salyers *et al.*, 1977). *B. fragilis* has also been known to ferment mucin to release sulphates which favour the growth of other sulphate reducing bacteria. They break down sulphates into sulphides which can be toxic to the human system and this suggests that *B. fragilis* may possess sulfatase activity (Willis *et al.*, 1996).

Mucin, a high molecular weight glycoprotein that is heavily glycosylated and phosphorylated acts as a potential target for commensal bacteria like sulphate reducing bacteria which metabolise the sulphates and phosphates to produce sulphide ions and toxic hydrogen sulphide. This alters the environment in the gut and encourages the growth of other intestinal infection causing bacteria. According to previous studies, the sulphation of the gastrointestinal tract increases from 1-3% in the stomach to 3-5% in the colon (Willis *et al.*, 1996). The intestinal microflora depend upon the type of nutrition and it has been observed that changes in nutrition can affect metabolism and cause an imbalance or change in the microflora of the gut. The other factor that contributes to the type of microflora present in the gut is the oxygen concentration. The upper part of the gastrointestinal tract is colonised by aerobic flora whereas the lower part of the gut is colonised by anaerobes. This may be due to the decrease in concentration of oxygen from the duodenum to the colon (Serino *et al.*, 2009). Evasion of the host immune system results in the development of an infection. Previous studies have shown that micro-organisms have the ability to colonise the reproductive tract by producing enzymes like mucinases and sialidases which degrade the protective mucin layers in the cervix (Wiggins *et al.*, 2001).

Some of the reasons for the colonisation of mucosal surfaces by bacteria may be due to chemotaxis and motility. Previous studies have shown that mutants lacking motility and chemotactic abilities exhibited reduced virulence (Laux *et al.*, 2005). Bacterial flagella can function as adhesins too. Adherence of bacteria to the mucous layer receptors prevents their adherence to the cell receptors of the intestine. But the process behind how this adherence enhances colonisation is not known. Hence the cell surface structures that are expressed in bacteria during colonisation play an important part in understanding the mechanism of pathogenesis (Laux *et al.*, 2005). The *B. fragilis* genes BF0855 and BF3763 which were cloned and expressed in *E. coli* encoded the putative expression of the enzymes α -fucosidase and α -glucosidase respectively. These enzymes have the ability to hydrolyse oligosaccharides into simple sugars like glucose or galactose and may play an important role in the degradation of complex polysaccharides like mucins. These enzymes may be associated with the differential expression of proteins in mucin enriched media and was one of the main reasons for their purification and characterisation in this study.

The study of the genome of organisms is termed as 'genomics' and the sequencing of genomes forms the basic blue print behind the study of proteins. There may be several modifications, interactions and other complex structures formed by proteins but all of these may be linked to the genome through various factors like the environment (Zivy & de Vienne, 2000).

4.6 Proteomics

Proteomics is a field of science that merges the study of genes and proteins employing technologies such as two dimensional gel electrophoresis and mass spectrometry. It is basically a genomic study of the functions and expression of proteins where the two dimensional gel electrophoresis is used to measure the expression of a protein and mass spectrometry is used in protein identification and characterisation. The advancements in the field of genomics has led to the more frequent employment of proteomics to study proteins/enzymes, and has led to advancements in antibacterial drug discovery, identification of drug targets and the

mechanisms of resistance exhibited by bacteria and various other characteristics related to it (Brotz-Oesterhelt *et al.*, 2005).

The term 'proteome' has been derived from the terms 'protein' and 'genome'. There are several factors that can influence the proteome of an organism, namely stress, culture conditions, environment, metabolism, temperature, drugs, illness and so on. The genome encodes the cell specific expression of proteins based on the effect of these factors. Protein modifications also act as important factors that contribute to the functions of a cell and the study of all the proteins produced by a particular species in an environment is collectively termed as proteomics. The five major aspects that contribute to research in proteomics include mass spectrometry based proteomics, proteome-wide biochemical assays, systematic structural biology and imaging techniques, proteome informatics and clinical applications of proteomics (Tyers & Mann, 2003). Proteomics serves as a tool in studying post translational modifications that alter physical and chemical properties of proteins including folding, conformation, stability and functions. Proteomic techniques like stable isotope labelling and novel mass spectrometric peptide sequencing can be used to characterise modifications in proteins like phosphorylation or in membrane proteins. Proteomics has also been used as a valuable tool in the identification of protein isoforms in eukaryotes (Blakeley *et al.*, 2010). Alternate splicing of mRNAs result in a major difference between the sequences of genes present and the translated protein isoforms that can be of varying complexities. Protease associated degradation of proteins acts as an important factor in regulating signalling pathways and other physiological processes where selective affinity purification and tandem mass spectrometric analysis of resulting peptides can be used to study the processing events that may be taking place during cell signalling (Xu *et al.*, 2009).

4.7 Transcriptomics

Analysing or studying proteins that are expressed in a cell through the mRNA present in the cell is called transcriptomics. Transcriptomics forms the link between proteomics and genomics since messenger RNA is the link between genes and proteins (Hegde *et al.*, 2003). Transcriptomics works based on the principle that genes showing similarities in their expression patterns may be functionally related too and hence it may be of importance to study the genes that control these expression profiles. Mutations in gene sequences can be used to study their potential as prospective drug targets, determining the functions of DNA sequences with no known activity based on their similarity to other conserved genes that may be homologous in nature (Twyman, 2003).

4.8 Two dimensional polyacrylamide gel electrophoresis (2D-PAGE)

This technique plays a major role in the separation of mixtures of proteins. It is mainly composed of two techniques namely isoelectric focussing (IEF) and sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS PAGE).

IEF is the first dimensional separation of the protein based on the net charge that it possesses. A high voltage is applied to the proteins of interest which are to be separated and an immobilised pH gradient technique is used in the separation of proteins using specific reagents. Proteins usually possess a charge and tend to migrate towards the oppositely charged electrode in the presence of an electric field. When a positively charged protein migrates towards the cathode, it passes through an increasing pH gradient which causes an overall decrease in the charge of the protein. Eventually, the protein reaches a pH region that corresponds to its isoelectric point where the net charge becomes zero and migration stops. This results in the formation of a protein band at that particular point. Following the separation based on charges, they are then run on a second dimension SDS-PAGE where the separation is based on the molecular weight and size. This technique employs the use of a denaturing anionic detergent, sodium dodecyl sulphate, in the effective separation of proteins (Gorg *et al.*, 2004). The anionic detergent confers a negative charge to the protein in proportion to its length by wrapping itself around the

polypeptide backbone. When an electric field is applied, the proteins migrate towards the positive electrode. The size of the pores in the polyacrylamide gel are user-defined and differentially retard the migration of proteins based on their size. Smaller protein molecules tend to travel farther down the gel when compared to larger protein molecules that remain closer to the point of origin resulting in effective separation of proteins. Therefore, by combining the separation of proteins based on their charge, or isoelectric point, and size, or relative molecular mass, differential expression studies can be performed.

Previous proteomic studies have been done in analysing the oxidative stress response in *B. fragilis* since they are known to be strict anaerobes and it was found that these bacteria are capable of inducing OxyR regulon genes that respond to the presence of oxygen or hydrogen peroxide in the environment and this might play an important role in their ability to survive within the tissues in the colon (Rocha *et al.*, 2003).

4.9 Mass spectrometry

Separation of molecules on the basis of their mass to charge ratio following ionisation is called mass spectrometry where a mass spectrum is generated based on the intensity of ionisation. The detected spectrum is then compared to a standard database to determine the identity of the complex peptide mixture. The sample is first passed through the ioniser that charges the molecules and the ions that are generated, enter the analyser where the ions are separated in vacuum. The individual ions from the analyser are then passed on to the detector where a spectrum based on the intensity of the ions is generated. The height of the peak in a spectrum is referred to as intensity. Mass spectrometry acts as a major tool in genomics and proteomics where complex molecules and protein mixtures can be identified and analysed (Liebler, 2002). Identification of proteins from a database is called peptide mass fingerprinting (PMF) or peptide mass mapping. Some methods of analysis also involve the comparison of a theoretical calculated spectrum of a database to an experimental spectrum generated from the mass spectrometer. One of the common methods used for the identification of proteins is the tandem mass spectrometry which consists of two phases. The first phase constitutes the ionisation

of peptides to produce a spectrum that is used for peptide mass fingerprinting. Some of the peaks from this spectrum are selected based on the intensity and subjected to the second phase of dissociation. The ions are transmitted through a high pressure region of the tandem mass spectrometer containing gas molecules. The gas molecules collide with the ions and further fragment them into ions of varying masses called collision induced dissociation (CID) (Liptak, 2005). Since fragmentation can occur between any two residues along the peptide, differing masses of ions are generated and can be termed as b and y ions (Refer to Figs. 7 and 8). These ions are then mapped to their respective proteins using a database like MASCOT (Johnson, 1987).

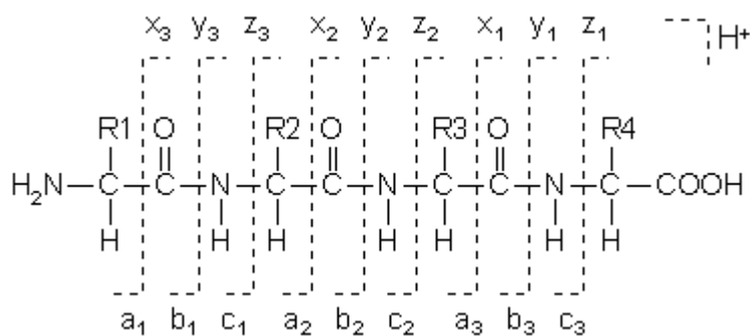


Figure 7 represents the fragment ions in an MS/MS spectrum (Anon, 2010)

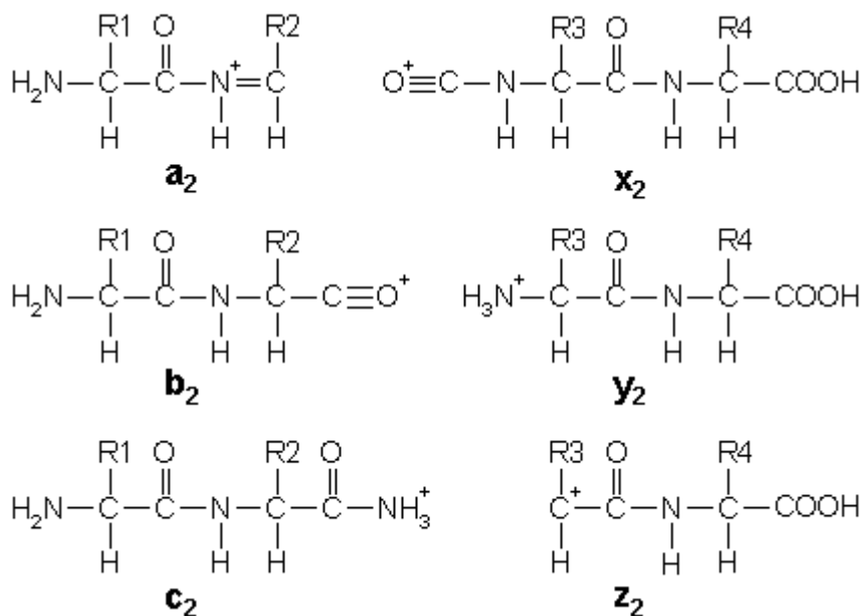


Figure 8 shows the structures of six singly charged sequence ions (Anon, 2010).

4.10 Glycoside hydrolases

Glycoside hydrolases or glycosidases are enzymes that catalyse the hydrolysis of sugars and other complex polysaccharides into smaller and simpler sugar units by hydrolysing their glycosidic linkages. These simple sugars or end products help in carbohydrate metabolism and are used as a source of carbon and energy in bacteria and higher animals. Glycosidases in association with glycosyltransferases, form the main machinery for the synthesis and breakdown of glycosidic bonds. They are produced as extra and intracellular enzymes in bacteria that help to acquire nutrients by breaking down complex compounds into simple sugars. Within the intestinal tract, they are capable of degrading carbohydrates like lactose, starch, sucrose and trehalose.

Glycoside hydrolases can be classified based on the stereochemical outcome of their hydrolysis reactions as inverting or retaining mechanisms. They act on oligo and polysaccharide chains by hydrolysing the glycosidic linkages in the non-reducing end or the middle of the chain. Hence this can be used to classify them as exo-acting or endo-acting glycoside hydrolases. The exo-acting enzymes are capable of cleaving residues from the ends of the polysaccharide chains whereas endo-acting enzymes remove residues from the middle or internal positions in a random manner resulting in several oligosaccharides of different sizes. They have also been classified on sequence based methods where sequence similarity acts as a major factor. This series of sequence based classification of enzymes allows the prediction of their mechanism of hydrolysis, active site residues and possible substrates that can confer this activity. The CAZy website enzyme database is regularly updated and classifies enzymes based on their sequence and three dimensional structural similarities. The glycosidase sequence analysis and three dimensional structure comparison has led to the generation of a hierarchical classification of these glycoside hydrolases (http://www.cazy.org/fam/acc_GH.html).

4.10.1 Inverting mechanism in glycoside hydrolases

Reactions involving the inverting mechanism of hydrolysis of carbohydrates employ the use of two enzymatic residues where one acts as an acid and the other acts as a base. The reaction occurs in the presence of carboxylic acid at the active site. This type of catalysis can be termed as a general acid base assisted catalysis method (Sinnott, 1990).

Figure 9: Figure showing the catalytic mechanism in glycoside hydrolases

(http://www.cazy.org/fam/ghf_INV_RET.html)

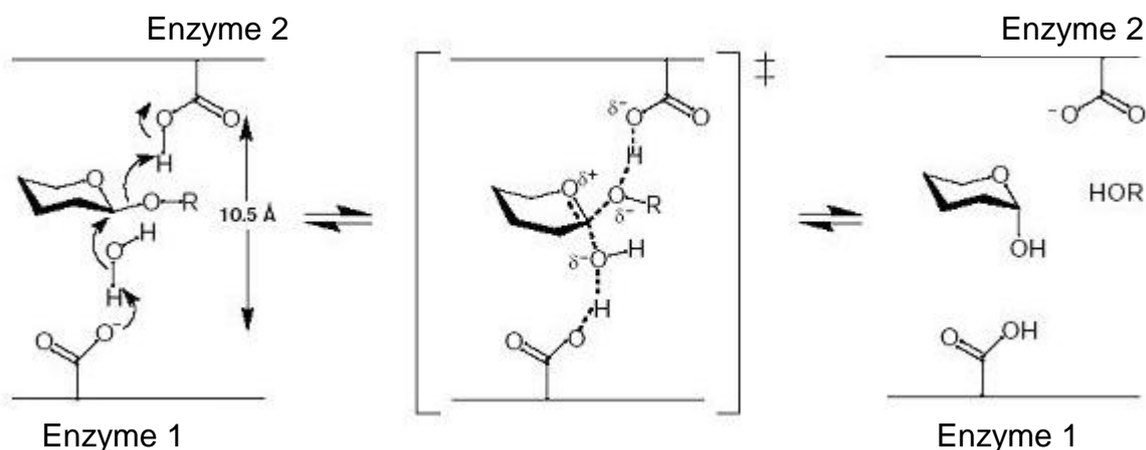


Figure 9.1: Figure showing the inverting mechanism of hydrolysis. Enzyme 1 acts as a base and the enzyme 2 acts as an acid by donating protons.

4.10.2 Retaining mechanism in glycoside hydrolases

The retaining mechanism involves a 2 step reaction and each step results in an inversion of compounds. There are two enzyme carboxylic acid residues present where one acts as a nucleophile and the other acts as an acid or base by donating or accepting protons respectively. When the nucleophile attacks the anomeric centre in the first step, a glycosyl intermediate is formed. The acidic carboxylate acts as a proton donor and assists the reaction. The second step involves the hydrolysis of the

glycosyl enzyme intermediate by the deprotonated acid carboxylate which acts as a base to produce the hydrolysed product. The reaction also involves the use of a nucleophilic water molecule. The following figure represents the inverting and retaining mechanism of glycoside hydrolase enzymes.

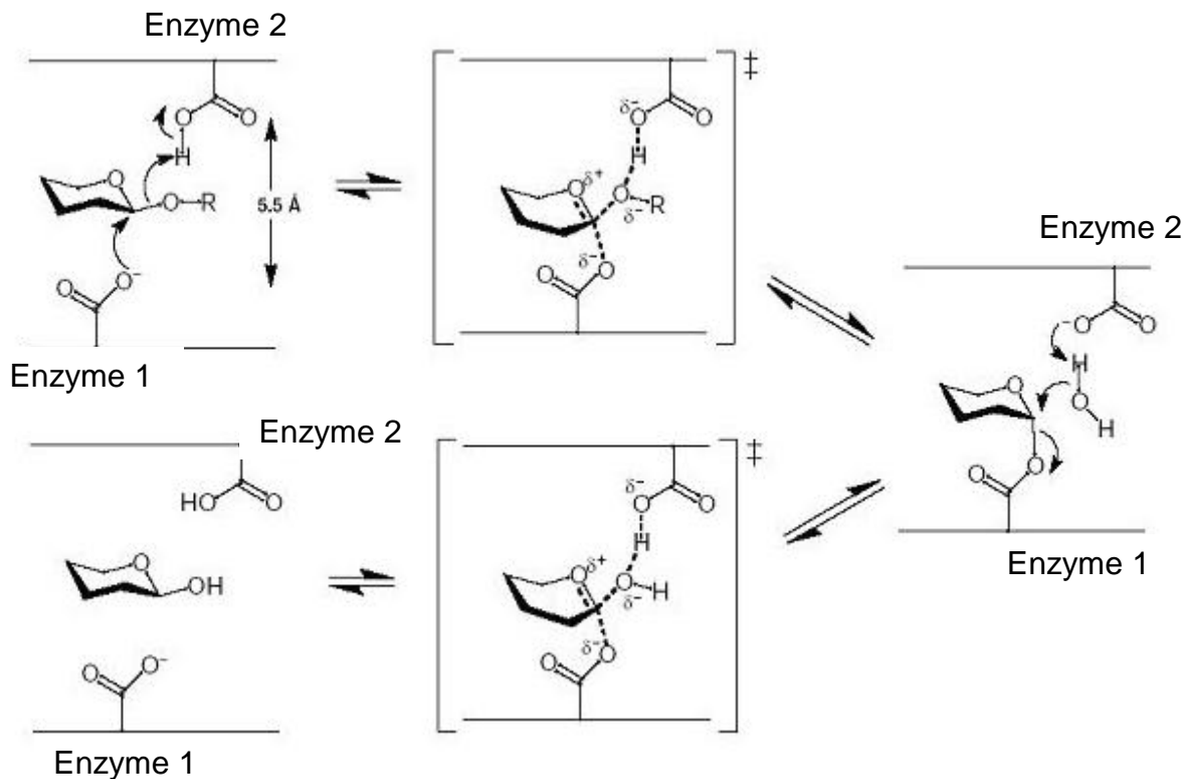


Figure 9.2: Figure showing the retaining mechanism of hydrolysis. The enzyme 1 acts as a nucleophile and the enzyme two acts as an acid or a base. A glycosyl enzyme intermediate is formed before the substrate is converted into the hydrolysed product.

4.10.3 Carbohydrate active enzymes (CAZy) based classification of enzymes

The CAZY website classifies enzymes into five main categories namely glycoside hydrolases (GHs), polysaccharide lyases (PLs), glycosyltransferases (GTs) and carbohydrate esterases (CEs) and carbohydrate binding modules (CBMs). Substrate specificity and molecular mechanisms contribute to some of the main factors considered for classification. The relationship between sequences and their folding similarities led to the consideration of amino acid sequence similarities as a major

factor. The active sites remain highly conserved and are maintained with integrity even though the structures of enzymes may differ due to evolutionary divergence. The families have also been grouped into 'clans' based on the strong conservation of the folds of proteins even though sequences were not found to be as conserved as the folds of proteins. This grouping of families into clans would help to overcome structural and sequential resemblances or relatedness of enzymes to more than one family. The other factor that supported the grouping of families into clans was the improved sensitivity of sequence comparison methods. As of May 2009, 115 families of enzymes have been identified and classified in the CAZY website (http://www.cazy.org/fam/acc_GH.html).

4.10.4 Active site topology

Glycoside hydrolases have the presence of specific active sites based on the position of hydrolysis of the polypeptide chain. These may be of three types namely, the pocket site which is characteristic of exo enzymes, the cleft structure that occurs in endo enzyme hydrolysis and the tunnel topology in exo enzyme hydrolysis (refer to the Fig. 10). While the pocket topology binds to the ends of the polysaccharide chains, the cleft topology binds chains randomly within a cleft. The tunnel topology being very similar to the cleft topology has additional polypeptide loops which are present on the top of the cleft giving it a tunnel like appearance. The end of the polypeptide chain enters the tunnel and the digested peptides are released out at the other end. In the case of the cleft active site, random hydrolysis of the polypeptide chain results in the release of oligosaccharides or various lengths (Davies & Henrissat, 1995).

Figure 10: Figure showing the active site topology in glycoside hydrolases (Davies & Henrissat, 1995)

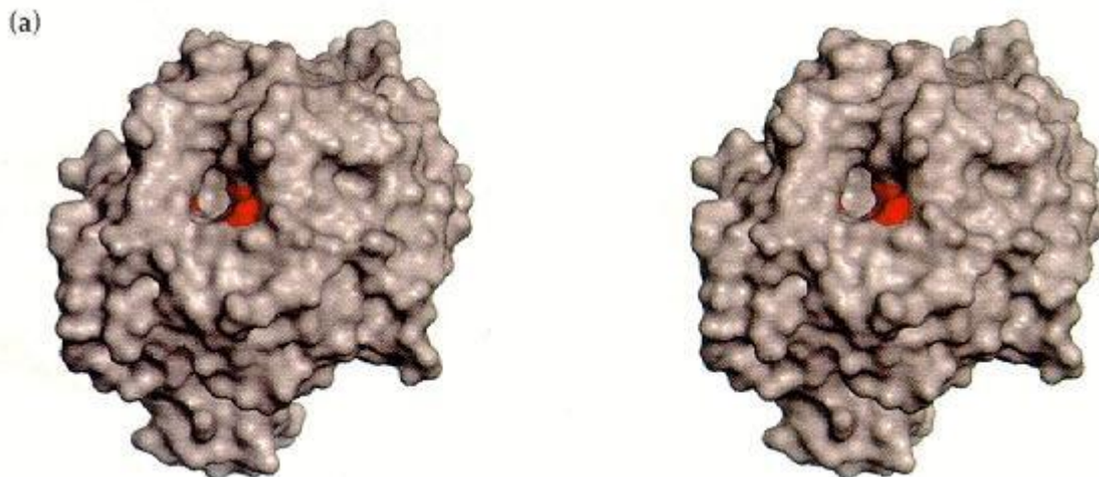


Figure 10.1: Pocket topology. Coloured area indicates the substrate binding site of the enzyme.

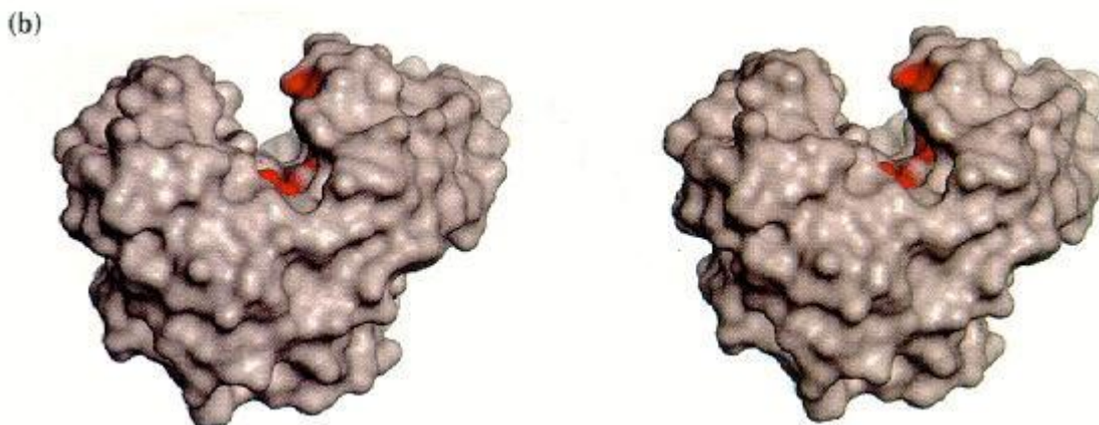


Figure 10.2: Cleft topology. Coloured area indicates the substrate binding site of the enzyme.

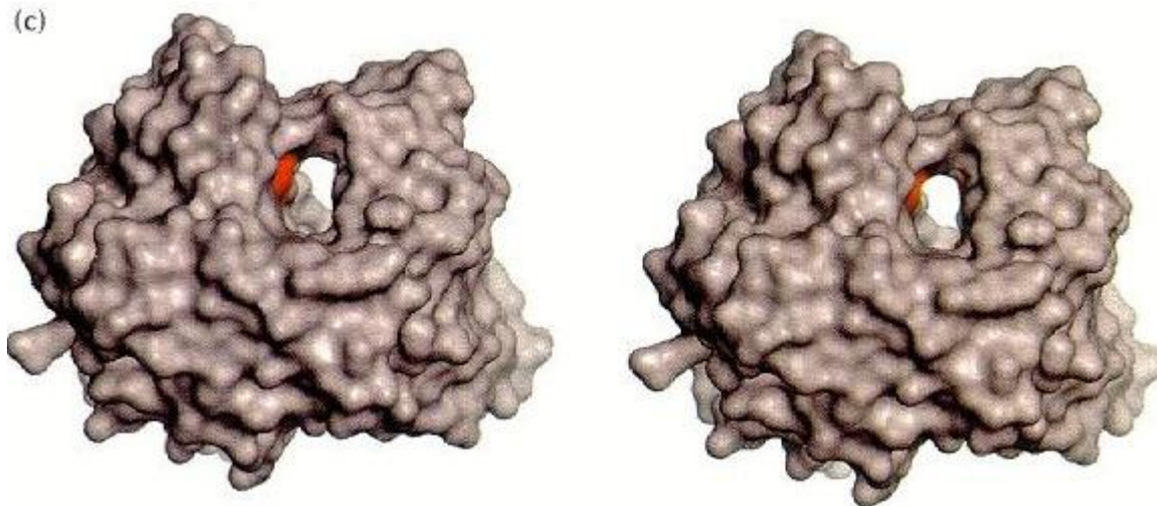


Figure 10.3: Tunnel topology. Coloured area indicates the substrate binding site of the enzyme.

4.10.5 Genomes of *B. fragilis* and *E. cancerogenus*

Completed published sequences of the genome of a number of different bacteria are readily available online. As of 11th May, 2009, 992 genomes have been published and are available via the Gold genome database. The online database reports 2523 ongoing bacterial genome sequencing projects along with 1029 eukaryotic genomes and 96 archael ongoing projects. The total number of ongoing genome projects has been reported to be 4807 when compared to 2006 when it was 2172 (<http://www.genomesonline.org/gold.cgi>).

The sequencing of genomes provides important information regarding the function of proteins. Once a genome is sequenced, the DNA is annotated as putative open reading frames (ORFs) containing information about the coded regions including exons, introns, promoters and so on (Zhang, 2006). Based on the translation of these ORFs, proteins or amino acid sequences can be obtained and searched against a sequence database to determine a list of proteins with the highest scores. Protein hits that show a very high degree of similarity or homology indicate a functional similarity between the two proteins. This can also be used to study the similarities and relationships between different bacterial species. Some of the problems that may be associated with this approach are that one particular gene

could encode for more than one protein and this could create confusion during analysis (Pennington & Dunn, 2001).

The genome of *B. fragilis* NCTC 9343 has been sequenced and published by the Sanger Institute (http://www.sanger.ac.uk/Projects/B_fragilis/). The genome of *E. cancerogenus* ATCC 35316 is still being sequenced and is available through the NCBI website but work is still ongoing. The sequencing information is available from the Washington University or through the Gold genomes online database (<http://www.genomesonline.org/gold.cgi?want=Bacterial+Ongoing+Genomes>).

Genomes of various species of *Bacteroides* namely *B. vulgatus*, *B. diastonis* were compared with *B. fragilis* and *B. thetaiotomicron* in a study. The conserved gene sets were grouped into Clusters of orthologous groups. Gut associated Bacteroidetes proteomes showed the predominant expression of conserved genes in carbohydrate transport and metabolism followed by cell wall or membrane or envelope biogenesis. The shared proteome of Bacteroidetes showed the expression of genes involved in amino acid transport and metabolism followed by translation, ribosomal structure and biogenesis genes (Xu *et al.*, 2007).

4.10.6 Glycoside hydrolase families GH95 and 97 and the putative activity of their enzymes

The GH95 BF0855 gene encodes for a conserved hypothetical protein containing 755 amino acids with a molecular weight of 86 kDa. The CAH06598.1 protein exhibits putative α 1-2, L-fucosidase and α -L-fucosidase activity and the catalytic nucleophile is asparagine that is activated by aspartic acid residues (Cerdeno-Tarraga *et al.*, 2005). The catalytic proton donor has been identified to be glutamic acid residues. Previous structural studies of these enzymes in *Bifidobacterium bifidum* have shown that the enzyme has a high degree of similarity to the immunoglobulin Ig domains with similar folding patterns. They possess α/α six barrel or catalytic TIM type barrel domain and are active against substrates like 2'-fucosyllactose and lacto-N-fucopentaose I (Katayama, 2004).

The GH97 BF3763 gene encodes for a putative export protein containing 649 amino acids with a molecular weight of 74 kDa. The CAH09443.1 protein is known to be a

putative α -glucosidase or α -galactosidase. Enzymes that exhibit inverting mechanisms of catalysis contain glutamic acid as nucleophiles and those that exhibit retaining mechanisms of catalysis possess aspartic acid residues as nucleophiles (Gloster *et al.*, 2008). Previous studies of the starch utilisation system in *Bacteroides thetaiotomicron* showed an inversion mechanism of catalysis liberating a β -anomer of glucose (Kitamura *et al.*, 2008). They possess α/β eight barrel or catalytic TIM barrel type domains and the amino acids of the active site are located at the C-termini of the β -strands. These proteins exhibit a high degree of similarity to proteins of other families that exhibit a retaining mechanism of glycoside bond hydrolysis (Naumoff, 2005).

The BF0855 gene from *B. fragilis* NCTC 9343 which was cloned and expressed in *E. coli* with a putative activity of α -L-fucosidase or 1, 2 α -L- fucosidase catalyses the reaction between α -L- fucoside and water resulting in the formation of L- fucose as an end product. These enzymes belong to the family of glycoside hydrolases that hydrolyse O and S glycosyl compounds and are also known as α -L- fucoside fucosidase. It is involved in n- glycan and glycan structure degradation and may be involved in the degradation of mucin by bacteria. 1, 2- α -L- fucosidase catalyses the following reaction:

Methyl- 2- para- L- fucopyranosyl- β - D- galactoside + H₂O \leftrightarrow L- fucose + methyl β - D- galactoside.

5 Aims of the study

This study aims at understanding the differences in protein expression in *E. cancerogenus* ATCC 35316 and *B. fragilis* strain NCTC9343 in mucin-supplemented media in comparison to basic semi-defined media containing glucose as the carbohydrate source. These are commensal organisms that act as opportunistic pathogens and can cause infections within the host when the immune system is impaired.

Two dimensional gel electrophoresis technologies would be employed, where the proteins were first separated in the first dimension based on their charge and pH using isoelectric focussing and then separated using the second dimension polyacrylamide gel electrophoresis based on their molecular weight. Protein expression profiles would be generated for both bacteria grown in the presence and absence of mucin and compared. Comparative analysis of gels would be used to determine the protein spots that show a differential expression in the presence of mucin.

The protein spots of interest will be excised and subjected to in-gel trypsin digestion where the peptides produced from the degradation of proteins would be analysed using a nano flow liquid chromatography coupled mass spectrometer. This would help to identify the protein based on the mass spectrum (m/z ratio) generated from the MS-MS fragmentation of peptide ions. The identification of the differentially expressed proteins may provide an insight into the indirect association of bacteria and their colonisation of mucin. It may also be interesting to use mucin-coated cell culture plates to study the interaction between bacteria and mucin in the presence of a host immune response.

This study also aims to express, purify and crystallise the proteins CAH06598 and CAH09443 from the glycoside hydrolase families 97 and 95 in *B. fragilis* strain NCTC 9343 which produce the enzymes with the putative functions of α -fucosidase and α -glucosidase respectively. The recombinant enzyme activity was analysed against different chromophoric and fluorogenic substrates. Crystallisation of proteins would be useful in determining its three dimensional structure and obtain a detailed understanding of its active sites.

6. Materials for the Proteomics experiment

6.1 Culture Conditions

Bacteroides fragilis strain NCTC 9343 and *Enterobacter cancerogenus* ATCC 35316 were cultured in Columbia agar plates (CBA). A single colony from the plate was used to inoculate 20 mL of sterile anaerobic broth media (Bac T/ ALERT SN). The medium was incubated overnight in an anaerobic chamber at 37 °C. A loopful of the liquid culture was used to streak a CBA plate (Figure 3). The purity of the subculture was confirmed by Gram staining which showed Gram-negative rods.

6.2 Anaerobic chamber

The bacteria was cultured and maintained in an anaerobic chamber that was flushed with 80% nitrogen gas, 10% carbon dioxide and 10% hydrogen (Cox & Mangels, 1976).

6.3 Minimal media used for the growth of *B. fragilis*

Minimal media used for the growth of *B. fragilis* according to the method of Varel and Bryant (1974)

Per 100 ml

Glucose	0.5 g
Mineral solution ¹	5.0 mL
Hemin solution ²	0.1 µL
Resazurin solution ³	0.1 µL
Volatile fatty acid solution ⁴	0.45 mL
Vitamin B solution ⁵	0.5 mL
FeSO ₄ -7H ₂ O	0.4 mg
6 mM (NH ₄) ₂ SO ₄	5.0 mL
Casitone (Difco)	0.2 g
Amino acid mixture ⁶	20 mL
2.5% (w/v) Cysteine-HCl solution	2.0 g
8% (w/v) Na ₂ CO ₃ solution	5.0 mL

The medium was made up with distilled water and autoclaved before use. The pH of the medium was adjusted to 7.0 and all components were added before autoclaving except sodium carbonate and cysteine. The medium was incubated overnight in the anaerobic chamber at 37°C in the absence of oxygen before inoculation of starter cultures.

6.3.1 Stock solutions

6.3.1.1 Mineral solution¹

Per litre

KH ₂ PO ₄	18 g
NaCl	18 g
CaCl ₂ -2H ₂ O	0.53 g
MgCl ₂ -6H ₂ O	0.4 g
MnCl ₂ -4H ₂ O	0.2 g
CoCl ₂ -6H ₂ O	0.02 g

6.3.1.2 Hemin stock solution²

A stock concentration of 0.1% (w/v) of hemin solution was obtained by dissolving 0.1 g of hemin in 1 mL of 0.1 M sodium hydroxide and then diluted to 100 mL with distilled water and appropriate dilutions of the stock were added to obtain a final concentration of 0.0001% (w/v) of hemin solution.

6.3.1.3 Resazurin solution³

A stock concentration of 0.1% (w/v) of resazurin solution was prepared by dissolving 0.1 g of resazurin in 100 mL of sterile 18.2 MΩ/cm and appropriate dilutions of the stock were added to obtain a final concentration of 0.001% (w/v) of resazurin solution.

6.3.1.4 Volatile fatty acid solution⁴

Per 100 mL

Acetic acid	36 mL
Isobutyric acid	1.8 mL

<i>n</i> -Valeric acid	2.0 mL
DL-2 Methylbutyric acid	2.0 mL
Isovaleric acid	2.0 mL

6.3.1.5 Vitamin B solution⁵

Per 100 mL

Thiamin-hydrochloride	20 mg
Calcium-D-pantothenate	20 mg
Nicotinamide	20 mg
Riboflavin	20 mg
Pyridoxine-HCl	20 mg
p-aminobenzoic acid	1 mg
Biotin	0.25 mg
Folic acid	0.25 mg
Vitamin B12	0.1 mg

6.3.1.6 Amino acid mixture⁶

Per 50 mL

L-histidine-hydrochloride	25 mg
L-tryptophan	25 mg
Glycine	25 mg
L-tyrosine	25 mg
L-arginine-hydrochloride	50 mg
L-phenylalanine	50 mg
L-methionine	50 mg
L-threonine	50 mg
L-alanine	50 mg
L-lysine	75 mg
L-serine	75 mg
L-valine	75 mg
L-isoleucine	75 mg
L-proline	75 mg
L-aspartic acid	75 mg
L-leucine	100 mg

L-glutamic acid 224 mg

6.4 Semi defined growth media

Per 100 mL

Casitone	0.5 g
Yeast extract	0.5 g
Ammonium sulphate	0.08 g
Salts solution A*	0.04 mL
Salts solution B *	0.005 mL
Hemin solution	0.01 mL
Glucose	0.5 g
Sodium carbonate	0.8 g
Cysteine-HCl	0.05 g

The medium was adjusted to a pH of 7 and all components were added before sterilisation except glucose and Na₂SO₄. Sterile glucose and sodium carbonate solutions were prepared separately and added aseptically to the medium. The medium was incubated overnight at 37 °C in the anaerobic chamber before inoculation with the starter cultures. The hemin solution was prepared by dissolving 50 mg of hemin in 1 ml of 0.1 M NaOH which was diluted with distilled water to 100 mL. In 100 mL of the mucin supplemented semi-defined media, 0.5 g of mucin Type II or III were added to the media in addition to 0.5 g of glucose and autoclaved. The 0.5 g refers to the total amount weighed out and added rather than the concentration of sugars in mucin.

Sigma-Aldrich porcine gastric mucin Type II and Type III were used in the proteomics experiments. A quantity of 0.5 per of the glycoprotein was weighed out and added to 100 mL of the semi-defined medium and dissolved by vigorous shaking. Mucin was added to the semi-defined medium just before autoclaving and care was taken to avoid inhaling the chemical dust.

6.4.1 Stock solutions

6.4.1.1 Salts solution A

Per litre

CaCl ₂	0.2 g
MgSO ₄	0.2 g
K ₂ HPO ₄	1.0 g
KH ₂ PO ₄	1.0 g
NaCl	2.0 g
NaHCO ₃	10.0 g

6.4.1.2 Salts solution B

Per litre

FeSO ₄ ·7H ₂ O	2.0 g
CoCl	0.08 g

6.5 Gram staining reagents

Crystal violet	5 mL
Gram's iodine	5 mL
Acetone	5 mL
Safranin solution	5 mL

6.6 SDS-PAGE Loading buffer

Per 10 mL

60 mM Tris Base pH 6.8	0.6 mL
50% (w/v) glycerol	5 mL
10% (w/v) sodium dodecyl sulphate (SDS)	2 mL
14.4 mM β-mercaptoethanol	0.5 mL
1% (w/v) Bromophenol blue (BPB)	1 mL

Stored as 1 mL aliquots at -20 °C

6.7 Protein size standards

High molecular weight standard (M.W. 36, 45, 55, 66, 84, 97, 116 and 205 kDa)

Low molecular weight standard (M.W. 20, 24, 29, 36, 45 and 66 kDa)

To obtain a final concentration of about 2.0-3.5 mg/mL, the lyophilised standards were reconstituted in 100 μ L of 18.2 M Ω /cm water and aliquoted out in 4 μ L quantities into 1.5 mL microcentrifuge tubes and stored at -20°C. Refer to appendix D for the list of proteins used as a source for producing these size standards.

6.8 Buffers and solutions used for the Proteomics experiment

6.8.1 Cell resuspension buffer

The cell resuspension buffer has the same recipe as that of the phosphate buffered saline (6.8.2)

6.8.2 Phosphate buffered saline (PBS)

Per litre (pH 7.0)

NaCl	8 g
KCl	0.2 g
KH ₂ PO ₄	0.2 g
Na ₂ HPO ₄ -12H ₂ O	1.44 g

6.8.3 Lysis solution

Per 10 mL

Urea	4.8 g
3-[(3-Cholamidopropyl) dimethylammonio]-2-hydroxy-1-propane sulphonate (CHAPS)	0.4 g

The solution was made up in a sterile universal and aliquoted out into microcentrifuge tubes in 1 mL quantities. This was then stored at -20°C and 2 μ L of IPG buffer was added to every 100 μ L before use.

6.8.4 Rehydration solution (for IPG strips pH 3-10 or 4-7)

Per 10 ml

Urea	4.8 g
CHAPS	0.4 g
1% (v/v) BPB	20 μ L

1% (w/v) bromophenol blue solution was prepared by dissolving 1 g of bromophenol blue dye in 100 mL of 18.2 M Ω /cm water.

The rehydration solution was divided into 700 μ L aliquots and stored at -20°C. Prior to use, 14 μ L IPG buffer (pH 3-10 or 4-7) and 1.4 mg of dithiothreitol (DTT) was added to each tube to give a final concentration of 2% (v/v) and 0.2% (w/v) respectively.

Isoelectric focusing or the first dimension separation of proteins was carried out using a Multiphor II electrophoresis system involving the use of other components listed in the Appendix section B.

Main parts include the anode and cathode electrodes, tray and electrode holder, dry strip aligners, IEF electrode strips.

6.8.5 Equilibration solution

Stock solution:

Per 200 mL

1.5 M Tris-HCl (pH 8.8)	10 mL
Urea	72 g
Glycerol	69 mL
SDS	2 g
1% (v/v) BPB	200 μ L

The stock solution was stored at -20°C in 20 mL aliquots.

6.8.5.1 Equilibration buffer with DTT

Per 20 mL

Equilibration stock solution	20 mL
DTT	0.2 g

6.8.5.2 Equilibration buffer with Iodoacetamide (IAA)

Per 20 mL

Equilibration stock solution	20 mL
Iodoacetamide (IAA)	0.9 g

6.9 Second dimension-SDS-PAGE

The SDS-PAGE was performed using a Protean II XL 2-D cells (Bio-Rad).

4 mm xi clamp notch vs 13 mm XL clamp notch

19 mm xi spacer vs. 8 mm XL spacer

181 mm xi core gasket vs. 198 mm XL core gasket

153 mm xi comb vs. 184 mm XL comb

The main parts include tank and lid, central cooling core, casting stand, sandwich clamps, alignment card and combs.

Small gels- glass plate sizes were 10.1 X 8.2 cm

Large gels- glass plate sizes were 20 X 20 and 20 X 22 cm separated by spacers of 1.5 mm

6.9.1 14% (w/v) Resolving gel components

18.2 MΩ/cm water	58 mL
1.5 M TRIS-HCl, pH 8.8	37.5 mL
10% (w/v) SDS stock	1.5 mL
40% (v/v) solution (37.5:1 acrylamide: bisacrylamide)	52.2 mL
10% (w/v) APS	750 μL
N, N,N',N'-Tetramethylethylenediamine (TEMED)	75 μL

6.9.2 4% (w/v) Stacking gel components

18.2 MΩ/cm water	12.85 mL
0.5 M TRIS-HCl, pH 6.8	5 mL
10% (w/v) SDS stock	200 μL
40% (v/v) solution (37.5:1 acrylamide: bisacrylamide)	1.95 mL
10% (w/v) ammonium persulphate (APS)	100 μL
TEMED	20 μL

1% (w/v) BPB	20 μ L
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6.9.3 Agarose sealing solution

Per 20 mL

Running buffer (1 X)	19 mL
Agarose	0.1 g
1% (w/v) BPB	40 μ L

Protein size standards were the same as those mentioned in the above sections (Refer to Appendix D for details).

6.9.4 Staining and destaining solutions

6.9.4.1 Colloidal Coomassie blue stock

Per litre

Ammonium sulphate	100 g
Phosphoric acid	20 mL
Coomassie blue G 250	1 g

20 mL of 18.2 M Ω /cm was used to dissolve 1 g of Coomassie blue G 250. A 20 mL solution of phosphoric acid was used to dissolve 100 g of ammonium sulphate and 18.2 M Ω /cm water was added if necessary. The two solutions were then mixed together and made up to 1 litre with 18.2 M Ω /cm water.

The stock solution was stored at room temperature until use.

6.9.4.2 Fixing solution (Colloidal Coomassie blue staining)

Per litre

Methanol	500 mL
Glacial acetic acid	120 mL

6.9.4.3 Staining solution (Colloidal Coomassie blue staining)

The stock solution was shaken well to suspend the Coomassie blue stain and four parts of the Coomassie stock was diluted with one part methanol before use.

6.9.4.4 Fixing solution (Fluorescent staining)

Per litre

Ethanol	400 mL
Acetic acid	100 mL

6.9.4.5 Staining solution (Fluorescent staining)

A 10 X concentration of the Bio-Rad Flamingo™ fluorescent gel stain was diluted to a 1 X working concentration of the solution with 18.2 MΩ/cm.

6.9.5 Visualisation of the stained gels

The Bio Rad Chemi doc XRS (Quantity One™ software, Resolution: 1392 X 1040 pixels) was used to capture images of the gel using UV transillumination for fluorescence stained gels and the GS-800 densitometer (Resolution: 1360 X 1024 pixels) was used to scan Coomassie blue stained gels. Hard copies of the gel were printed out using the Mitsubishi Video Copy Processor attached to it.

Image acquisition for Coomassie blue stained gels was carried out using Bio-Rad GS-800 image densitometer and further analysis was done using the PDQuest™ Advanced v 8.0 software. Hard copies of the gel were produced using the Mitsubishi Video Copy Processor (K65HM-CE / High density type, 110 cm X 21 m). The raw image data which was obtained from scanning the gels using the GS-800 densitometer (Bio-Rad Quantity One™ software) was uploaded into the other two softwares, Ludesi Redfin and Non-linear Dynamics Progenesis SameSpots and used in analysis.

6.10 Digestion of proteins for Mass Spectrometric analysis

6.10.1 Protein digestion in solution

6.10.1.1 Dissolve solution

Per 100 mL

SDS	0.1 g
Tris base	6.06 g
Dithiothreitol (DTT)	77.1 mg

The solution was adjusted to a pH of 8 using HCl and stored at room temperature.

6.10.1.2 Trypsin stock (1 µg/µL) (Promega and NEB)

Vial containing the lyophilised powder was stored at -20°C

Trypsin lyophilised powder 20 µg/mL	100 µg
Glacial acetic acid (50 mM)	100 µL

The trypsin powder was dissolved in 50 mM glacial acetic acid and stored at -20°C for upto 1 month or at -80°C for a long term.

6.11 In-gel protein digestion reagents

6.11.1 100 mM NH₄HCO₃

Per 10 mL

NH ₄ HCO ₃	79 mg
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6.11.2 50 mM NH₄HCO₃

Per 10 mL

100 mM NH ₄ HCO ₃	5 mL
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6.11.3 Trypsin solution (20 µg/mL)

Per 100 µL

1 µg /µL trypsin stock	2 µL
50 mM NH ₄ HCO ₃	98 µL

6.12 Buffers used in LC-MS analysis

6.12.1 Start buffer A

Per 100 mL

LC-MS Grade Water	95 mL
LC-MS Grade Acetonitrile	5 mL
Formic acid	0.1 mL

6.12.2 Start buffer B

Per 100 mL

LC-MS Grade Acetonitrile 95 mL

LC-MS Grade Water 5 mL

Formic acid 0.1 mL

The solution was made up fresh before use.

7 Methodology used for the proteomics experiment

7.1 Growth study in *B. fragilis*

Ampules of *B. fragilis* were ordered from the National Collection of Type Cultures (NCTC) and used to inoculate the anaerobic basal broth media and incubated in the anaerobic chamber at 37°C overnight. The bacteria were sub cultured in Columbia blood agar media or brain heart infusion broth media.

The initial growth curve experiment was carried out by growing the bacteria in minimal medium whose components have been mentioned in the Materials section 6.3. But since the growth in the medium was not found to be reasonable enough for analysis, a semi defined medium was used. The components of the media varied with respect to yeast extract being used as the source of vitamin B. The medium was autoclaved before use and incubated overnight in the anaerobic chamber before inoculation. Mucin and glucose were weighed out and added at 0.5 g per 100 mL to the semi-defined growth media.

The semi defined media supported quicker growth of bacteria. Experiments were performed by using different carbon sources.

The culture was incubated at 37°C in the anaerobic chamber. The anaerobic condition within the chamber was monitored by using resazurin indicator strips that turned pink in the presence of oxygen in the chamber environment. Refer to Appendix E1 for details.

7.2 Growth curve studies

Small volume (20 mL) overnight starter cultures were used to inoculate large volumes (450 mL) of the media and the optical density of the media were monitored at various time intervals by measuring the absorbance at 600 nm and plating out serial dilutions of the culture. The time periods that were used to monitor the growth in *E. cancerogenus* were 0 h, 2 h, 3 h, 4 h, 5h, 6 h, 7 h, and 8 h whereas the time periods in *B. fragilis* were 0 h, 4 h, 6 h, 21 h, 22h, 23 h, 24 h, 25 h, 26 h, 27 h, 28 h, 29 h, 30 h, 31h, 46 h, 47h. The growth of *B. fragilis* in the medium was found to be slower and hence more time points were monitored. For each and every time point, the culture was mixed well and 1 mL was transferred into a plastic cuvette with a sterile pipette tip. The OD was measured at 600 nm using a Cecil Spectrophotometer. 100 µL of the culture was spread out onto a sterile CBA agar

plate and incubated overnight at 37°C. 1 mL of the culture was used to perform appropriate serial dilutions into test tubes containing 9 mL of sterile distilled water. Care was taken to maintain aseptic conditions within the chamber while performing the experiment. All experiments were performed in triplicates and the average values were calculated. Growth curves were plotted out to determine the phase of growth of the bacterium.

7.3 Two dimensional polyacrylamide gel electrophoresis (2D-PAGE) methodology

Two dimensional gel electrophoresis comprised of a series of steps from the growth of bacteria, protein extraction and one dimensional iso-electric focussing to the two dimensional separation of proteins using SDS-PAGE. The methodology has been explained in detail in the following sections.

7.4 Growth of *E. cancerogenus* and *B. fragilis* in semi defined media

A loopful of the glycerol stock was used to streak out a plate of Columbia blood agar medium and incubated overnight at 37°C in an anaerobic chamber.

A single colony of the bacteria from the culture plate was used to inoculate 450 mL of the semi-defined medium supplemented with and without mucin the following day. The semi defined medium was incubated overnight in the anaerobic chamber before inoculation. The culture was incubated in the anaerobic chamber till the optical density reached a value of 0.7 at 600 nm.

7.5 Protein extraction

A volume of 100 mL of the culture was then centrifuged at 4000 x g for 30 min at 4°C. Following this, the pellets were resuspended in 5 mL of PBS and washed at 4000 x g for 10 min at 4°C. The washing step was repeated 3 times and the final resuspension step was performed in 1 mL of PBS buffer. To 2 µL of IPG buffer pH 4-7 or 3-10, 100 µL of lysis solution stock was added to prepare the lysis solution and 100 µL of this was added to each of the samples. The pellets were then ultra-sonicated on ice for 1 min at 10 s intervals and then centrifuged at 14000 x g for 30 min at 4°C. The pellets were then discarded and the supernatant was transferred into clean 1.5 mL microcentrifuge tubes.

The cell free extract was subjected to removal of contaminants using the 2D clean up kit. It also helped to get rid of residual polysaccharides. To 100 μ L of the cell free extract, 300 μ L of precipitant solution was added and incubated on ice for 15 min.

To this, 300 μ L of the co-precipitant was added and vortexed well before centrifuging it for 5 min at 14000 x g. The supernatant was discarded and 40 μ L of the co-precipitant was added to the pellets and incubated on ice for 5 min. This was then centrifuged at 14000 x g for 5 min and the supernatant was discarded. 25 μ L of 18.2 M Ω /cm water was added and vortexed well. To this, 1 mL of chilled wash buffer and 5 μ L of wash additive were added before incubating the tubes at -20°C for at least 30 min. The tubes containing the proteins could be stored at -20°C for a maximum of one week with minimal protein loss. The solution was centrifuged at 14000 x g for 5 min after which the supernatant was removed and the pellets were allowed to dry for half a min.

The pellets were re-suspended in 350 μ L of rehydration solution containing 0.001 g of DTT, 4 μ L of BPB, 7 μ L of IPG buffer and 350 μ L of the rehydration stock solution. The solution was vortexed and centrifuged at 14000 x g for half a min to remove the insoluble residue and the solution was transferred into the wells of the reswelling tray.

IPG strips of 11 cm length, 3.3 mm width and 0.5 mm thickness (pH 3-10 or 4-7) which were stored in the freezer at -20°C were taken and the plastic was removed from its surface. The strips were layered in the wells of the reswelling tray in such a way that the gel end of the strip was facing the rehydration solution and the positive end of the strip was located at the lower end of the reswelling tray. Once the strips were layered in the wells, the surfaces of the strips were covered with Dry strip cover fluid and left to rehydrate overnight at room temperature.

7.6 Iso-electric focussing

The temperature was maintained at 20°C

pH intervals	Step	Voltage (V)	Time (h)	kVh	Current (mA)	Power (W)
4-7	1	500	0:01	-	2	5
	2#	500	6:00	3.0	2	5
	3	3500	1:30	3.0	2	5
	4	3500	8:00	30	2	5

Total: 15:31

This step was included only when the Multiphor II Electrophoresis system was left running overnight since this is an optional step.

Table 1: Table showing the parameters used in isoelectric focussing, pH 4-7

pH intervals	Step	Voltage (V)	Time (h)	kVh	Current (mA)	Power (W)
3-10	1	500	0:01	-	2	5
	2	3500	1:30	3.0	2	5
	3	3500	6:00	22.0	2	5

Total: 7:31

Table 2: Table showing the parameters used in isoelectric focussing, pH 3-10

The Multiphor II from Amersham Biosciences was used to run the 1st dimension isoelectric focussing.

Two electrode contact strips of 11 cm were cut out and moistened with 1 mL of 18.2 MΩ/cm water. The Dry strip cover fluid was poured on to the cooling plate and the Drystrip tray with its anode at the upper end [red] and the cathode at the lower end [black] was placed on it. The cathode and anode ends were connected to the Multiphor II isoelectric focussing unit. About 15 mL of the Dry strip cover fluid was poured on the tray and the Dry strip tray aligner was placed on it with the groove side facing upwards. Care was taken to avoid formation of air bubbles while aligning the dry strip tray on the cover fluid since it could affect the thermal contact. The

rehydrated Dry strip gels from the reswelling tray were removed and placed parallel to each other along the lanes of the tray with their gel side facing upwards. The cathodic and anodic ends of the strips along with the gel surface were kept in contact by the moistened electrode strip. The electrodes were placed and pressed down on these electrode strips. The cover fluid was poured over the strips to immerse them completely and a programmed EPS 3501 XL power supply (Amersham Biosciences) was switched on. The Techne Circulator C-100 with a TECAM Heat exchanger was used to maintain a temperature of 20°C throughout the process of separation. The gels were left to run for at least 8 h and the strips were stored at -80°C in plastic petridishes until use. The parameters used in running the IEF strips have been detailed in Tables 1 and 2.

7.7 Casting of SDS-PAGE gel

Two glass plates of size 20 X 20 cm and 20 X 22 cm were wiped clean with 50% ethanol. The glass plates were aligned parallel to each other with a spacer of 1 mm thickness placed in between them around the edges. Single screw clamps were used to clamp the plates together tightly and the whole set up was mounted on a rubber gasket placed on a casting stand.

A 12% (w/v) resolving gel was prepared as mentioned in the materials section and the solution was degassed for 30 min after which it was pipetted out into the space between the glass plates upto about 4 cm from the top of the smaller plate. The surface of the solution was overlaid with 18.2 MΩ/cm water and allowed to polymerise for 45 min at least. After the gel was set, the overlaid water was poured out and a clean filter paper was used to remove any residual liquid.

A 4% (w/v) stacking gel acrylamide solution was prepared and layered on top of the resolving gel solution. A comb containing a long 17 cm well and a small size standard well measuring 1.5 mm in thickness was immediately inserted into the gap between the plates. Care was taken to avoid formation of air bubbles by inserting the combs at an angle. This was allowed to polymerise for 30 min at least. Once set, the combs were removed and the surface of the gel was rinsed with copious amounts of 18.2 MΩ/cm water to get rid of any gel debris or unpolymerised acrylamide.

The frozen IPG strips were equilibrated with 0.1 g of dithiothreitol (DTT) in 10 mL of the equilibration stock solution by gentle rocking on a shaker for 15 min. The equilibration step was repeated again with 0.45 g of iodoacetamide (IAA) instead of DTT. IAA was dissolved in 10 mL of the equilibration stock solution and the strips containing the gel were gently rocked on a shaker for 15 min.

The strips were placed on a filter paper moistened with 18.2 MΩ/cm water to drain all the excess liquid. Approximately 1 cm of plastic from the negative edge of the strip was cut off in order to obtain a suitable length. The strips were layered on the top of the gel and liquid agarose sealing solution was poured in between the glass plates allowing it to cool down. Care was taken to ensure that the strips were placed with their gel side facing the smaller glass plate.

The gels containing the strips were loaded onto the SDS PAGE electrophoresis unit by assembling them on a cooling core which was gently immersed at an angle into a gel tank containing about 2.0 L of running buffer. Protein size standards of 8 µl were loaded into the respective wells of each gel present on either side of the cooling core. Running buffer was also used to fill up the upper compartment containing the electrodes.

The second dimension was run in the electrophoresis unit where the lower and upper parts of the tank were filled with the running buffer. A current of 8 mA was applied to two gels and the process was normally complete in about 4 h which was indicated by the bromophenol blue dye that ran down the gel and started to come off. Throughout the separation process, the cooling core was maintained at a temperature of 10°C using the Techne Circulator C-100 with TECAM Heat Exchanger 1000.

On completion of the process, the clamps were removed and the gels were transferred to large storage containers and 250 mL of the fixing solution was added to it. The gels were fixed for about 1 h with gentle rocking.

The fixing solution was replaced by 250 mL of the Coomassie blue staining solution which was left rocking on a shaker overnight at room temperature.

Following this, the gels were destained using 18.2 MΩ/cm water for 4-6 h and the images were scanned and analysed using the PDQuest software. The raw image data obtained from scanning the gels using GS-800 Bio Rad densitometer was uploaded and used in analysis in the Redfin and SameSpots softwares.

7.8 Fluorescent staining

Attempts were also made to use fluorescent staining for visualising gels containing protein spots. Once the second dimension run was complete, the gels were left overnight in the fluorescent stain fixing solution with gentle rocking on a laboratory rocker. The gels were transferred to the staining solution the following day and allowed to stain from 3 to 8 h. The gels were stored in special storage boxes wrapped with aluminium foil to avoid contact with sunlight.

7.9 Visualisation of the stained gel

The Bio Rad Chemi doc XRS (Quantity One™ software) was used to capture images of the gel using UV transillumination. Hard copies of the gel were printed out using the Mitsubishi Video Copy Processor attached to it.

7.10 Gram staining procedure

7.10.1 Heat fixed film preparation

A droplet of water was spread on to the surface of a clean glass slide with the help of a sterile plastic loop. A single colony of the bacteria was picked from a fresh culture plate with another sterile loop and mixed gently with the droplet of water and allowed to dry at room temperature. The slide was heat fixed by passing it through a Bunsen flame. Care was taken to avoid heating the slide too much since it could result in the charring of proteins. The slide was allowed to cool after heat fixation and subjected to Gram staining.

7.10.2 Gram staining

The heat fixed slide was flooded with methyl violet stain for 30 s. The stain was poured off after 30 s and the slide was flooded with Gram's iodine using a Pasteur pipette. The slide was left for 30 s with Gram's iodine on it and then gently rinsed off with tap water. Following this, the decolourising agent, acetone was used to flood the slide for just a few seconds (3-4 s) and washed off instantly in running tap water. The slide was finally flooded with Safranin stain for 30 s. The slide was rinsed off with tap water and dried with a blotting paper. Once the slide was completely dry, it was observed under the microscope using the oil immersion lens (X 100).

7.10.3 Expected Result

The presence of violet or deep purple stained cells indicate that the bacteria may be Gram positive and the presence of red, pink or orange cells indicate that they are Gram negative in nature.

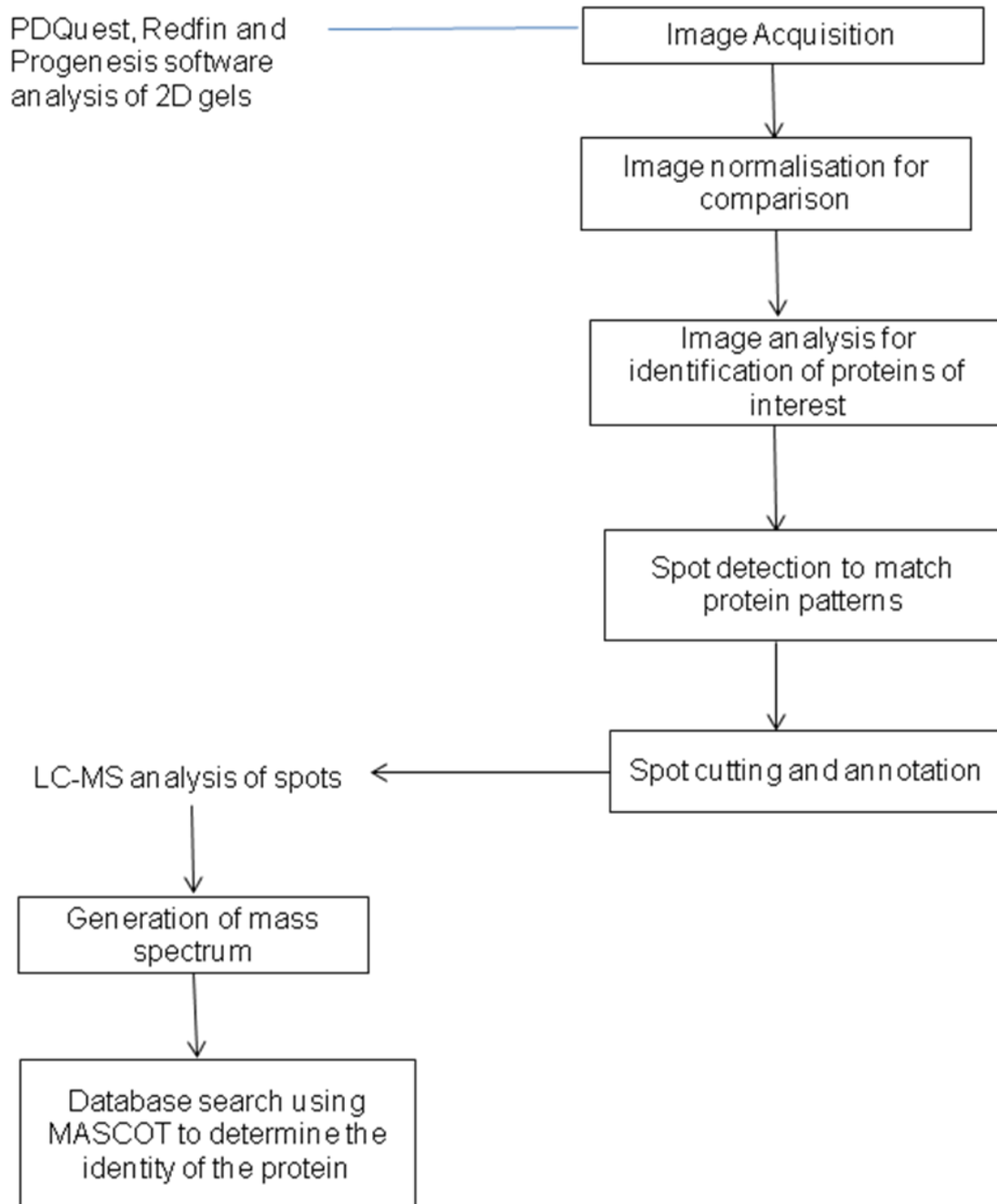


Figure 11: Methodology used for the analysis of spot data

7.11 PDQuest analysis

7.11.1 Acquiring images

The Bio-Rad GS-800 image densitometer was used for scanning and obtaining images of the 2D gels. At least 6 gel images were generated from every condition in order to ensure reproducibility of results. The cultures were grown in 450 mL volumes and separated out into 4 samples (Figs. 20.1-20.4, 21.1-21.4, 22.1-22.4, 23.1-23.4, 24.1-24.4, 25.1-25.4) of 100 mL volume each when they reached the appropriate optical density. A second lot of 450 mL cultures were grown independently to process the rest of the 2 or 3 samples (Figs. 20.5-20.7, 21.5-21.6, 22.5-22.8, 23.5-23.6, 24.5-24.6, 25.5-25.6)). Proteins were extracted from these biological replicates and 100 μ L of the cell free extract from each sample was processed separately using the 2D Clean up kit. Hence 4 samples of the cell free extracts used in sequential optimisation experiments were from the same pool of material whereas the other 2 were from a separate culture. The PDQuestTM software uses Bio-Rad imaging services to convert the signals from biological samples into digital data. The gel was initially 'Preview' scanned and specific sections of the gel which contained all the protein spots were framed and acquired. Fig. 9 shows the steps involved in the processing of data.

All the gels were scanned at the same time and care was taken to keep the frame dimension the same in all the cases. Once scanned, the images were saved and copies were printed out using the Mitsubishi Video Copy Processor. The gels can be cropped or rotated according to the crop and rotation tools on the Image menu. PDQuest was used as the default software for analysis based on the Student t-test.

7.11.2 Spot detection

The spot detection wizard was used to start up an experiment and step wise analysis of the gels was carried out. The spot detection wizard was used to detect spots on the gels. A template gel was selected and parameters were generated based on the smallest, faintest and the largest spot identified in it. The set parameters were used to scan every gel accordingly since gels show a high degree of variability with respect to streaking, intensity or clarity. The spot detection wizard was set to identify Gaussian, salt and pepper noise characteristics and filter them using a 7 X 7 pixel size filter dimension based on the individual noise pixels in the gel. Other features

including subtracting backgrounds and speckles detection were also enabled to make the detection process more efficient.

Once the gels were filtered and smoothed to clarify all the spots including fuzzy, streaked or overlapping spots, three dimensional spots called Gaussian spots were created. All types of analysis were then performed on the Gaussian image which contained the Gaussian spots that were precisely identified and quantified. Three types of images were created during spot detection namely the original 2D scan, the filtered image and the synthetic Gaussian image. The Gaussian spots were indicated on the gels using crosshairs that was located at spot centres and spot ellipses that encircled the spot shape.

7.11.3 Editing spots

Editing of spots involved several features including adding, removing, combining or changing spot boundaries. All the editing was performed on the Gaussian image. Since the same parameters were used to identify spots in all the gels, some spots seemed to have been missed out. In certain cases, air bubbles or stain marks were identified as spots. Hence all the gels were reviewed manually to minimise errors. The spot adding tool was used to add spots to the Gaussian image and the spot removal tool was used to remove incorrectly identified or positioned spots.

Other tools include the spot combining tool which is used to combine spots that have been identified to be different and the changing boundary tool which helps with the altering of spot boundary either by removing or moving them around.

7.11.4 Matching spots

Setting up an experiment automatically results in the generation of a matchset once the spots on the gels have been detected. Further grouping of the gels based on the various media conditions in which they were grown was also performed. The three main groups were the Media without mucin, Media with Type II mucin and media with Type III mucin. The gels could be matched manually or automatically. Automated matching was performed with a 50% restriction where the master image was matched to all the other images. The master image contained all the spots from all the gels mapped to it. However, a classic manual match was also performed after the automated match in order to minimise errors during matching. Landmarking was the first step performed which involved choosing reference spots to align and

position gels for matching. Once the landmarks are selected, an automatic match of all the spots is performed. The remaining spots can be reviewed individually and either matched or simply added to the master.

7.11.5 Data Analysis

Upon completion of the matching, an experiment summary report can be obtained from the analysis menu. The report shows the number of spots identified in each gel including its group details, match rates 1 and 2. Match rate 1 shows the percentage of matched spots relative to the total number of spots on the gel. Match rate 2 shows the percentage of matched spots on the gel relative to the total number of spots on the master gel.

Standard spot numbers (SSP) are assigned automatically to all the spots. These are generated by the software when the standard gel is divided into rows and columns with each section having the same number of spots. The spots in a section are numbered sequentially and the first two digits of the SSP number indicate the X and Y coordinates of that section. The spot review tool was used to study the histograms of all the spots in a matchset. The spot quantities, intensities, location and the number of gels showing expression of a protein can be identified using this tool. The highlighted spot can be compared between all the gels and match modifications can also be made. The histograms of interest can be selected and used to create analysis sets.

The scatter plot analysis shows the relatedness between two gels or two groups in a matchset. The quantity of each spot in the first gel is plotted along the X axis against the quantity of spots in the second gel. The slope of the scatter plot shows the degree of similarity. The closer the value is to 1.00, the stronger the relatedness between the gels.

From the reports menu, quantity and quality tables can be obtained where the spot quality report shows a range of values from 0 to 100 based on the Gaussian fit, X and Y streaking, overlap and linear range of the scanner.

7.11.6 Normalization

Normalization refers to the process where quantitative data from different gels are adjusted in order to compare different samples to one another. Variations in gels occur due to a number of factors which include handling and pipetting errors, loss of sample during loading, low cell density or inconsistent staining. Hence this process helps to compensate for non expression related variations in spot intensity.

The normalization formula:

Normalized spot quantity = Raw spot quantity X Scaling factor X Pipetting error compensation factor / Normalization factor.

Where raw spot quantity refers to the intensity of each spot, scaling factor refers to a constant value defaulted as 106 parts per million (ppm) and the normalized quantity is multiplied to it.

The normalization factor was calculated for each gel based on 'total quantity in valid spots' method where the raw quantity of a spot in a gel was divided by the total quantity of all the spots in each gel included in the master.

All the spots which showed variations in expression in at least 6 gels were selected for analysis using LC-MS. Further, spots that showed expression in all three conditions were also selected for analysis.

7.11.7 Consensus

Group consensus was used to find potential spot detection errors and use the spot matching information uniformly across a set of gels in replicate group. The consensus tool helps to edit and match spots across specific gels, replicate groups or all the gels in a matchset. The ambiguous spots could be identified from the consensus analysis and matched to one or more spots on other gels. Once updated, the results are added to the spot review analysis report.

7.11.8 Gel analysis software

Apart from the PDQuest™ Advanced software, the Ludesi Redfin software and the Non linear Dynamics Progenesis SameSpots software were also used to compare and analyse the spot data. Spot data and match result tables have been attached to the Appendix F. Redfin software uses the Analysis of variance (ANOVA) and principal component statistical analysis (PCA) for summarising the spot data in E.

cancerogenus and *B. fragilis* whereas a cluster analysis is used to describe the same data in Progenesis SameSpots software. Study of gels using three different software packages was performed to determine the reliability and reproducibility of analyses data from PDQuest, Redfin and SameSpots rather than depend just on the data obtained from PDQuest analysis. The differentially expressed spot identification criteria included a significant p value of 0.05 or less and a fold change of 1.0 or more.

7.11.8.1 Redfin analysis of 2DE gels

The six main steps used in the analysis of data using the Redfin software included the pre-processing of images, choosing of a warp reference image, warping of all the gel images, creation of a fusion image, detection of spots and generation of spot borders. Completion of all the above mentioned steps provided a series of results showing the experiment overview, statistics used, details of filtered spots including details on its p value (p values < 0.05 are considered to be significant), presence (set as 100%), volume and fold change with respect to all the three different growth conditions.

The pre-processing of images involved an initial quality control check on the gels based on a number of factors to determine its suitability for analysis. The image editor was used to further crop, flip or rotate images as necessary. The step involving the creation of a fusion image is automated and involves transferring of the detected spots and their borders on to every individual image for quantification. This step ensures 100% matching and helps to get rid of cracks or other damages to the image. The “less-more” slider helps to optimise the number of spots detected on a gel and the “loose-tight” slider helps to determine the spot boundaries.

Redfin analysis of spots also supports the categorisation of data into multiple groups to facilitate comparison. Once the reference image is selected and warped, global and local spot editing tools can be used to edit the spots on the gel. Once the spot borders are detected, an analysis report is generated containing the entire list of spots detected. The spots of interest are then filtered out by setting a p value filter of <0.05, a fold change of 1.0 and an expression volume of 100%. The filtered spots

could then be exported as a .pdf or .xls document. Spots of interest can be viewed using a 2D and 3D montage or statistically analysed.

7.11.8.2 SameSpots analysis of 2DE gels

The steps used in the analysis of data using the Progenesis SameSpots included the initial quality assessment followed by the alignment of images. The initial quality assessment step also helps to prevent positional errors with tools for cropping, flipping or rotating images. This software uses an automated image alignment system in order to minimise the repeated editing and re-matching of images.

The pre-filtering step of analysis involved the filtering out of damaged, noisy and spots from the edges of the images. This was followed by appropriate grouping of gels based on their growth conditions to study differential expression and an automated gel analysis involving spot detection, background subtraction, normalisation and matching was performed.

The analysis provides a review of results containing a list of spots from which the spots with relevant p values (<0.05) were selected. Fold change of 1.0 or more can also be used to select relevant spots since fold change is calculated by subtracting the highest and the lowest mean values and this is added on to the mean value of the third group. They provide a 2D and 3D montage view of spots to be able to study them better.

Apart from analysis of variance, this software also uses other statistical applications like PCA analysis, correlation analysis and power analysis for describing the data obtained from the experiment. PCA analysis helps to determine the variation in expression based on the levels of spot expression spread out across all the gels.

The correlation analysis helps to determine the degree of relatedness between two sets of data. Normalised spot expression volumes with a value closer to 1 indicate a higher correlation when compared to lower correlation spots which have a value closer to -1. The results of the correlation analysis are visually presented using dendrograms. Based on the variations in expression, difference between mean groups and sample size, the power value is calculated. The larger the difference between the group means, the more significant the power value of the spot. Usually a power value of 0.8 or more is categorised as a relevant spot. The data is presented

as average values based on the differential expression of spots in three different conditions of growth. The spots of interest could then be selected and used to generate a report for future use.

7.12 Preparation of protein spots for analysis using the Mass spectrometer

The spot outlines of interest were cut out using a scalpel and the gel bits were stored in silanised 0.5 mL tubes at -20°C until use.

The Coomassie blue stain was removed from the gel slice by adding 100 µL of 100 mM NH_4HCO_3 and 60 µL of acetonitrile and shaken for 30 min. The same washing process was repeated at least 3 times in order to get rid of the stain completely. Following this, the gel slices were dehydrated by adding 50 µL of ACN and incubated in room temperature for 5 min. The same process was repeated again. The gel slices containing the proteins were dried using the centrifugal evaporator for 15 min at room temperature. In the meanwhile a 20 µg/mL concentration of trypsin solution was prepared in 50 mM NH_4HCO_3 solution and 25 µL of the trypsin solution was added to each of the tubes. The tubes were incubated on ice for 30 min and 30 µL of 50 mM NH_4HCO_3 was added to cover the solution before incubating it at 37°C in a waterbath overnight.

The following day, 30 µL of a 50% (v/v) ACN and 5% (v/v) formic acid solution was added to stop the reaction and the tubes were shaken for 30 min. The solution containing the digested peptides were removed and stored in a clean microcentrifuge tube. A 30 µL solution containing 83% (v/v) ACN and 0.1% (v/v) formic acid was added to the tubes containing the gel slices and shaken for 30 min to remove all the remaining digested peptides which were then pooled into tubes containing the initial collection of digested peptides.

7.13 Liquid chromatography

Liquid chromatography experiments were performed using the Dionex Ultimate 3000 nano LC system which consists of a pump that regulates the flow of buffers, temperature compartment that is maintained at 60°C while running samples, the autosampler that is attached to the syringe facilitating injection of the samples from

the multi-well plate and the UV detector that shows the absorbance values of samples at 254 nm. Separation of samples was carried out using a 200 μm X 5 cm long monolithic column made up of polystyrene divinyl benzene polymers. The peptide elution gradient was set in such a way that the first 0-0.5 min was used to perform equilibration and injection at a 100% Buffer A concentration. The elution of peptides occurred between 0.5 and 16 min of the gradient where the concentration of acetonitrile was gradually increased by pumping Buffer B to a 100%. The column was washed and prepared for the loading of the next sample during the last 6 min of the gradient (Total 22 min gradient).

The nano LC system is coupled to a HystarTM Bruker Daltonics ESI Ion trap MS system which carries out the MS/MS fragmentation of peptides.

7.14 Mass spectrometric analysis

The microcentrifuge tubes containing the digested peptides were freeze dried completely and then resuspended in 10 μL of the start buffer A. The solution was then pipetted into the respective microtitre well for performing an LC-MS analysis.

The HCTUltra was used as the ion trap system with the ESI source. In order to create a stable spray with high ionisation yields, optimised ESI conditions were used. Identification experiments were performed at a flow rate of 3 $\mu\text{L}/\text{min}$ with a nebulizer pressure of 15 psi and a drying gas flow rate of 5 L/min at a drying gas temperature of 300°C.

7.14.1 Tuning

Optimising the voltages that guide the flow of ions along the CapExit, Skimmer, Octopoles, Lenses and Tap Drive help to achieve maximum signal intensity called tuning. The Smart parameter setting was used for qualitative analysis and this was done using a standard compound. The target mass was set to 622.03 m/z. Tuning solution that was used in our experiments was 20% (w/v) acetonitrile.

7.14.2 Parameter settings for peptide mapping

The Standard enhanced scan mode was used to obtain MS spectra in the order of singly, doubly and triply charged precursor ions. The Ion Charge Control (ICC) target was set to 200.000. Scan range was 300-2000 m/z for MS and the average was set to 5 (Low concentration samples required higher average numbers whereas low average numbers were set to complex samples).

The Octopole 1 DC was set to 8.0 V. All ions above the intensity threshold were selected in the Auto MS (n) precursor selection analysis. Singly and doubly charged ions were detected whereas other mass ranges were excluded as being contaminants.

The compound spectra generated from the Auto MS (n) were exported to the Biotoools software as *.mgf files and then database searched through Mascot.

Ion chromatograms from data processing included the UV chromatogram, BPC All MS and the TIC All MSn. The MS/MS ion search was used to connect to the Mascot search where the query file peptides were compared to the peptides present in the database to determine the identity of the protein.

The username, email and an appropriate search title were filled in the search box. The taxonomy used for carrying out the searches was 'Eubacteria' since the protein spots were cut out from gels expressing Gram negative bacterial culture proteins. The enzyme used was trypsin and the database normally used to conduct the searches was NCBI. The global modifications were set to a default of Carbamidomethyl (C) and the variable modifications were set to Oxidation (M). The missed cleavages were set to 2 and the charge state to doubly and triply charged ions. The mass tolerance was set to 1.7 Da, MS/MS tolerance to 0.5 Da and the output was set to 50 proteins. The results were saved by clicking on the 'get hits' option. (Refer to section G1 of the Appendix for more details) Random matching results could also provide false positive results and hence certain criteria were set up to segregate false positives from correct identifications. This included selection of results where the protein score was more than 40 and at least 2 peptides matched to the query sequence.

8. Materials used for the cloning, purification and structural studies of glycoside hydrolase enzymes from families 95 and 97 in *B. fragilis*

8.1 Liquid media

Unless stated otherwise, all media were prepared using distilled water, and sterilised by autoclaving. In cases where solutions could not be autoclaved, they were filter-sterilised using 0.2 µm Ministart[®] filter units (Sartorius) and stored in sterile plastic 30 mL containers.

Liquid media were stored at room temperature, and solid media at 4°C. All adjustments to the pH of solutions are stated under the list of components, and were achieved using HCl or NaOH.

8.2 Solid media

Columbia blood agar containing 5% (v/v) horse blood

Per 100 mL

Columbia agar powder	3.7 g
Horse blood	5 mL

To 95 mL of distilled water, the agar powder was added and mixed well before autoclaving. The medium was then allowed to cool to about 50°C and 5 mL of horse blood was added to it and mixed gently. This was aseptically poured into labelled Petri dishes and allowed to set for 30 min. The horse blood was aliquoted out in 5 mL quantities, stored in the freezer at -20°C and thawed before use.

8.3 Plasmid vectors and competent cells used for cloning

The plasmid vector used was pET-YSBLIC which was a modified form of the pET-28a vector developed in the University of York to allow ligation independent cloning (LIC) with over expression of proteins in BL21 strains. Refer to Appendix C for details.

The vector has the presence of a hexa-histidine tag at the *N*-terminal end that helps in easy purification of proteins using the immobilised metal affinity chromatography column.

The chemically competent cells used for the transformation were TOP10 *E. coli* cells and *E. coli* strain BL21 respectively.

8.4 Luria-Bertani (LB) broth

Per 1000 mL

Tryptone	10 g
Yeast extract	5 g
NaCl	10 g
pH 7.0	

8.5 LB agar

Per 100 mL

LB broth	100 mL
Agar (bacteriological agar N° 1)	2 g

The medium was autoclaved to make the agar soluble and then poured into Petri dishes when cooled to about 55°C.

8.6 Selective media

The antibiotic used in the selective medium was kanamycin where a stock concentration of 10 mg/mL was prepared in 18.2 MΩ/cm water and stored at -20°C in the freezer. A 1 in 100 dilution was used as the working concentration making it 50 µg/mL.

Isopropylthio-β-D-galactoside (IPTG) was used as an inducer in LB medium at a final concentration of 240 µg/mL and this was prepared from a stock concentration of 24 mg/mL dissolved in sterile 18.2 MΩ/cm water.

8.7 Cryogenic storage of bacterial stocks

The *B. fragilis* stock was stored at -80°C after mixing 0.5 mL of an overnight anaerobic basal broth culture with 0.5 mL of sterile 50% (v/v) glycerol.

Per 100 mL

100% (v/v) glycerol (Fisher)	50.0 mL
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8.8 SOB broth

Per 100 mL

Tryptone	2.0 g
Yeast extract	0.5 g
5 M NaCl	0.2 mL
1 M KCl	0.25 mL
4 M MgCl ₂ ·6H ₂ O	0.25 mL
4 M MgSO ₄	0.25 mL

The solution was adjusted to a pH of 7.0 and autoclaved before use. MgCl₂·6H₂O and MgSO₄ were prepared and autoclaved separately and then aseptically added to the solution.

8.9 SOC broth

Per 100 mL of the SOB broth

1 M D (+) glucose (filter sterilised)	2.0 mL
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8.10 NZY Enrichment broth

Per litre

NZ Amine	10 g
Yeast extract	5 g
NaCl	5 g
MgSO ₄ ·7H ₂ O	2 g

The solution was adjusted to a pH of 7.5 before autoclaving.

8.11 NZY supplement

Per 45 mL

1 M MgCl ₂ ·6H ₂ O	12.5 mL
1 M MgSO ₄	12.5 mL
20% (w/v) D (+) glucose	20 mL

The solution was filter sterilised and 0.45 mL of the NZY supplement was added to 10 mL of the NZY broth to make up the NZY⁺ enrichment broth aseptically.

8.12 TE buffer

Per litre

0.5 M Tris-base pH 7.5 20 mL

0.5 M EDTA pH 8.0 2 mL

The buffer was adjusted to a pH of 7.5

8.13 Starter cultures for auto-induction media

Per 10 mL

Water 9.55 mL

1M MgSO₄ 20 µL

1000 x metals 2 µL

40% (w/v) glucose 125 µL

25% (w/v) aspartate 100 µL

50 X M 200 µL

kanamycin 100 µL

8.13.1 ZYM-5052 auto-induction media

Per 500 mL

ZY 478.5 mL

1M MgSO₄ 1 mL

1000 X metals 0.5 mL

50 X 5052 10 mL

50 X M 10 mL

kanamycin 5 mL

8.13.2 Stock solutions

8.13.2.1 50 X M

Per 100 mL

Na₂SO₄ (anhydrous) 3.6 g

NH₄Cl (anhydrous) 13.4 g

KH₂PO₄ (anhydrous) 17.0 g

Na₂HPO₄ (anhydrous) 17.7 g

The salts were sequentially dissolved in distilled water and autoclaved before use.

8.13.2.2 1000 X metals

Per 100 mL

Sterile 18.2 M Ω /cm water 36 mL

0.1 M $\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$ 50 mL

(Dissolved in 100-fold dilution of HCl, approximately 0.12 M HCl and filter sterilised using 0.2 μm filters)

1 M CaCl_2 (anhydrous) 2 mL

1 M $\text{ZnSO}_4 \cdot 7\text{H}_2\text{O}$ 1 mL

0.2 M $\text{CoCl}_2 \cdot 6\text{H}_2\text{O}$ 1 mL

0.1 M $\text{CuCl}_2 \cdot 2\text{H}_2\text{O}$ 2 mL

0.2 M $\text{NiCl}_2 \cdot 6\text{H}_2\text{O}$ 1 mL

0.1 M $\text{Na}_2\text{MoO}_4 \cdot 5\text{H}_2\text{O}$ 2 mL

0.1 M $\text{Na}_2\text{SeO}_3 \cdot 5\text{H}_2\text{O}$ 2 mL

0.1 M H_3BO_3 (anhydrous) 2 mL

All the solutions were made separately and autoclaved unless stated otherwise.

20 mL of a 25% (w/v) aspartate solution was prepared and the pH was adjusted to 7.0 by neutralisation with NaOH and autoclaved separately.

8.13.2.3 50 X 5052

Per 100 mL (The compounds were sequentially dissolved in distilled water)

Glycerol 25 g

Glucose 2.5 g

α -lactose 10 g

8.13.2.4 ZY

Per litre (sequentially dissolved in distilled water)

Tryptone	10 g
Yeast extract	5 g

8.13.3 PASM-5052 SeMet-labelling auto-induction media

Per 500 mL

Water	450.3 mL
1 M MgSO ₄	1 mL
1000 X metals	0.5 mL
50 X 5052	10 mL
20 X P	25 mL
17 amino acid mix	10 mL
25 mg/mL methionine	0.2 mL
25 mg/mL selenomethionine	2.5 mL
100 µM vitamin B12	0.5 mL
kanamycin	5.0 mL

8.13.4 Stock solutions

8.13.4.1 20 X P

Per 100 mL (sequentially dissolved in water)

(NH ₄) ₂ SO ₄ (anhydrous)	6.6 g
KH ₂ PO ₄ (anhydrous)	13.6 g
Na ₂ HPO ₄ (anhydrous)	14.4 g

8.13.4.2 17 amino acid mix (17 aa mix)

Per 100 mL (sequentially dissolved in water)

Alanine	1 g
Arginine-HCl	1 g
Asparagine-H ₂ O	1 g
Aspartate	1 g

Na glutamate	1 g
Glutamine	1 g
Glycine	1 g
Histidine-HCl-H ₂ O	1 g
Leucine	1 g
Isoleucine	1 g
Lysine-HCl	1 g
Phenylalanine	1 g
Proline	1 g
Serine	1 g
Threonine	1 g
Tryptophan	1 g
Valine	1 g

The amino acids were dissolved by continuous stirring and also heating with the Bunsen when necessary. The solution was made up to a pH of 7.0 by neutralisation with NaOH, filter sterilised using 0.2 µm filters and stored at 4°C until use.

8.13.4.3 Methionine and Selenomethionine solutions

20 mL of a 25 mg/mL solution of methionine solution was prepared and autoclaved before use.

5 mL of a 25 mg/mL L (+)-selenomethionine solution was made up just prior to use and filter sterilised using a 0.2 µm filter.

20 mL of a 100 µM solution of Vitamin B12 was prepared by filter sterilisation using a 0.2 µm filter and stored at 4°C until use.

8.14 Protein extraction and purification solutions and buffers

8.14.1 Cell resuspension buffer in protein extraction

Per litre (pH 7.4)

HEPES	4.76 g
NaCl	29.22 g
imidazole	0.68 g

8.14.2 Nickel column purification resin

Fast flow chelating sepharose from GE Healthcare

8.14.3 Gel filtration purification resin

HiLoad 16/60 Superdex 200 prep grade (All the resins were stored in 20% (v/v) ethanol)

8.14.4 Buffers used for nickel affinity purification

8.14.4.1 Start buffer

Per litre (pH 7.4)

HEPES	4.76 g
NaCl	29.22 g
imidazole	0.68 g

8.14.4.2 Elution buffer

Per litre (pH 7.4)

HEPES	4.76 g
NaCl	29.22 g
imidazole	34 g

8.14.5 Buffers used for gel filtration purification

Per litre (pH 7.4)

HEPES	4.76 g
NaCl	11.9 g

8.14.6 Buffer exchange buffers

The proteins were concentrated using 30 kDa concentrators.

For CAH06598 and CAH09443

Per litre (pH 7.4)

HEPES (5 mM)	0.595 g
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8.15 Crystallisation screens and reagents

The screen compositions for the various conditions used for the preparation of crystal trays have been included in the Appendix H.

8.16 Chemicals and solutions used for SDS-PAGE

8.16.1 Resolving gel (12% (w/v) acrylamide)

40% (w/v) solution (37.5:1 acrylamide: bisacrylamide)	3 mL
Buffer B	2.5 mL
18.2 MΩ/cm water	4.5 mL
10% (w/v) ammonium persulphate (APS)	50 µL
Tetramethylethylenediamine (TEMED)	10 µL

8.16.2 Buffer B

Per 100 mL	
2 M Tris-base pH 8.8	75 mL
10% (w/v) sodium dodecyl sulphate (SDS)	4.0 mL

8.16.3 Stacking gel (4% (w/v) acrylamide)

40% (w/v) solution (37.5:1 acrylamide: bisacrylamide)	0.5 mL
Buffer C	1 mL
18.2 MΩ/cm	2.5 mL
10% (w/v) APS	30 µL
TEMED	10 µL

8.16.4 Buffer C

Per 100 mL	
1 M Tris-base pH 6.8	50 mL
10% (w/v) SDS	4 mL

8.16.5 Running buffer for SDS-PAGE

Per litre

Tris-base	30.3 g
Glycine	144 g
SDS	10 g

The pH of the solution was adjusted to 8.8 before the addition of SDS and the running buffer was made up at a 10 X stock concentration which was diluted at 1:10 before use with distilled water.

8.16.6 SDS-PAGE Loading buffer

Per 10 mL

60 mM Tris-base pH 6.8	0.6 mL
50% (w/v) glycerol	5 mL
10% (w/v) SDS	2 mL
14.4 mM β -mercaptoethanol	0.5 mL
1% (w/v) bromophenol blue	1 mL

Stored at 1 ml aliquots at -20°C

8.16.7 SDS-PAGE solubilisation buffer

Per 10 mL

SDS-PAGE loading buffer	7.6 mL
urea	2.4 g

Stored at 4°C

8.16.8 Coomassie Blue staining solution

Per litre

Coomassie Blue R-250	1 g
glacial acetic acid	100 mL
methanol	450 mL

Coomassie blue gel staining solution can be filtered through a filter paper and re-used.

8.16.9 Destaining solution for gels

Per litre

glacial acetic acid	100 mL
methanol	100 mL

The Coomassie blue gel destaining solution can be filtered through a filter paper funnel containing activated charcoal and re-used once the stain is removed.

8.16.10 Bradford assay

Bradford's reagent solution	200 μ L
Bovine serum albumin solution at various concentrations ranging from 0.1 mg/mL to 1 mg/mL.	
Protein solution to be assayed	20 μ L

8.17 Solutions used for enzyme assay

8.18 Stock solutions

20 mM solution of HEPES pH 7.4	25 mL
10 mg/mL solution of BSA	20 mL
10 mM 4-methylumbelliferyl α -D-glucoside	10 mL
10 mM 4-methylumbelliferyl α -L-fucoside	10 mL
50 mM 4-nitrophenyl α -D-glucoside	10 mL
50 mM 4-nitrophenyl α -L-fucoside	1 mL
50 mM 4-nitrophenyl β -D-maltoside	1 mL
10 mg/mL α -glucosidase (Recombinant protein purified from the study)	100 μ L
10 mg/mL α -fucosidase (Recombinant protein purified from the study)	100 μ L

8.19 Reagents for agarose gel electrophoresis

8.19.1 TAE Running buffer (50 X stock)

Per litre

Tris-base (ultra-pure)	242 g
17.51 M glacial acetic acid	57.1 g
0.5 M EDTA pH 8.0	100 mL

A 1: 50 dilution of this buffer is used for making agarose gels and for running gels in the electrophoresis tanks.

8.19.2 Bromophenol blue (6 X) sample loading buffer

Per 10 mL

Bromophenol blue	0.025 g
Glycerol	3.0 g

A 1 X dilution of the buffer was added to the sample before loading it onto a gel.

8.19.3 Size standards for agarose gel electrophoresis

The Bioline hyperladder I or NEB 1 kb ladder was used as a size standard after diluting to a concentration of 1 µg/12 µL. Bioline hyperladder I size standard was added at a concentration of 1 µg/5 µL before loading it on to the gel and stored at 4°C for up to 12 months. Refer to Appendix D for details.

8.20 Kits, reagents and enzymes for DNA extraction and purification

Quiagen miniprep kit was used for the extraction of genomic DNA.

8.21 Polymerase chain reaction

1 µL of the forward primer was mixed with 1 µL of the reverse primer and made up to 5 µL with sterile 18.2 MΩ/cm water.

The other reagents include KOD DNA polymerase enzyme, 1M MgSO₄, deoxy nucleotide tri phosphate (dNTP), KOD DNA polymerase buffer and DNA to be amplified.

9. Methodology used for the cloning, purification and structural studies of glycoside hydrolase enzymes from families 95 and 97 in *B. fragilis*

9.1 Bioinformatics analysis

The gene of interest was identified from GenPept in the NCBI sequence viewer v2.0 and the FASTA format of the sequence was obtained. The conserved and semi conserved residues of the sequence was determined using the ClustalW software (www.ebi.ac.uk/Tools/clustalw).

The expert protein analysis system (EXPASY) identification and characterisation tool was initially used for the determination of the molecular weight and pI of the proteins of interest. The presence of signal peptides and transmembrane helices in BF3763 and BF0855 was identified using the SignalP 3.0 server (<http://www.cbs.dtu.dk/services/SignalP/>), and the TMHMM server (<http://www.cbs.dtu.dk/services/TMHMM/>). The 'Translate tool' in Expasy was used to translate the gene sequence into amino acid sequence thereby determining the amino acids that are present in the protein (<http://www.expasy.org/tools/dna.html>)

The coding sequence (CDS) for the BF0855 protein from GH95 is

```
1 ttaagggttc aactcgatcc attcaccggc ttgagtatct atgttcaaga tatccgtgcc
61 ggataaagta tagttcttat tccgcttcaa tgtaccggga cgggccattt tcaacatgcc
121 gcctttctcc gaatagatgc gtacccgatt gattttctcca ccttccatcc gggccgatac
181 caggaatgca cccatggcgc gtaagttttc gaaggataca tctttccatt ctttcggtat
241 tgccgggaag atgcgtatta cgctgtgtg actttgcagc aacatttctt gtatgcctgc
301 tgcaaaagcg aagttacctt cgagcgtgaa ggggcggtaa gtgaagcggg atttgccgct
361 ttgggtctgg tcaccgttgg catggaaggt gttcttcagg cagaagcatt ccgcaaaggt
421 tttcagggct tgtgcggcac cttcaccatc gaatgcacgg gctttcatat tggcgagcca
481 actgtatgaa tatcctgtcc agtagtccgg ccctacttta tcgagtctct tcaaggtggc
541 gcggtatgat tgttgtgact tttctccgtc gctccagtct atcagtccca gcggtatgat
601 agccatggca tgtgaaaaat gacgggtgta ttctttgtag ggatatacctt tggcgaaagt
661 gaggcaacct tcttcgtcta tgtcgtaatc cggtagttgg gcttccaggg atgccagtg
721 tccggcttcg tccggccagg tcagttcgtg tgccagttcg gaagtagcct taaagaggaa
781 gtgcatcatt gccaggtcat aattggatcat gtcgctgaac cacgcttgca gggagttgtc
841 gaatatttcg ggacttgaac tgaattccag tttacgtact ccttcggggg taacttctga
901 aatttgctcc agatagattg ccacatcctt gatgaatgga taagcacgct ctttgaggaa
961 agtgccggtc gccgaatatt tccattgcag atagaagtgt tgtgccagcc atgcagctac
1021 ggtttgcgac atggagtatt gtatccatcc tcccatgggt tctcctgtca gggtaaaaac
1081 ccccggtata ttcacccctt cggtagcggaa atagcggcgg gtgtagcgtt tgtatgcatc
1141 ccggttggtc cacaacgtgt tcagatagcc cattccttcg gtcaggtgat tgctgtata
1201 ggccggccag taacttagct gggatattaa atcgtgatga tagtctccct tccagggcgg
1261 aagtttgccg ttatcggctg tccatacagc ctgcaggag atggggtatg agtgctcggc
1321 ggtagtggaa ccgaatttat acatttcggt ctgatactgt ttttgagta cagaatcggg
1381 tagagtgatc gatgattgtg cccagtactt gtcccaatat tccaggtgtg cctgataatc
1441 atgctttaga ccgcgttgca gggcggcttc tgccttttcg gaagcttgcg cgccggataa
1501 ggaagagggt acgctccagg taccgtaaaa ggtttcgcct tcttgcttcc aacagactgt
1561 tacgtcatac gaaaagtctc cgtatccctt ttgatggtag gttatttggg ttccttcacg
1621 aaccactttt ccttgttgat agcctaactg gtgcagatct tgtcccgaac ccggtatcag
1681 gctgccatcc ggcttcggtt tattgtatac aggagtgatg atggagggtt ctatcggggg
1741 tttcagggta ccgaatacaa accatccgat cggttcggta gcatgaacga aagtctgcat
1801 ttgtgttccg tctgcccagt ctgcttcaca taaggcattg ttcaggtaaa gtcttacctg
1861 agtcggagtg ccgatctgct ctaatggaaa ttcgatggca gctccgggaa tcttgagcgg
1921 cgcgggattc atgtcataag gccagtccag ctttttctgt acgggcagat agttcttttg
1981 acgaatatgt tcttttacc aggagaagcg gaagtatct cccgacagac tgtctaccgg
2041 acgtaaatcc cataaatcgg ttctgtccag tgacaggcgt aggggtggag cacggtgcca
2101 tacciaaggc ccgacggtag cttttcccaa gggcattgct tcgtcccacg accgggcaag
2161 tccggtgaat acaagatcac tctctgaagg agtgaccgga caatagtcac acgagttgca
2221 actgctccac agaccacaac agagtagtaa tagtagtttg attttcat
```

The CDS for the BF3763 protein from GH97 is

```
1 atgaaaagaa aaatgatgtc cttattactc gcattggcgg taataagcgg aagtagcgtg
61 tacgctaaag tgattgacgt aatgtctccc aacggagcca ttaaagtatc ggtagacatc
121 aaggaccgga tttattattc ggtgtcctat gataatgacc agttattaaa agattgctat
181 ctcaacctgc aactgcagaa tgagacgtta ggtacgaatc cccacttacg gagcaccaaa
241 cgtggaacca ttgacgaaag tgtaaaacgt gaaatacctt tcaagaatgc gatcgtaaaga
301 aatcactgta ataccctgag aatgaatttc agcggaaatt atgccgttga attccgcgta
361 ttcgacaatg gtatcgctta ccgttttgtg acagataaaa aaggagataa catcgtaatg
421 ggggaagact tcgcaattaa ctttccaacc aattataaag ctcatctctc ccaaccgat
481 ggcttttaaaa cctcatacga atgcccatac actcatgtag ataccgaaaa gtatgctgct
541 accgaccgca tgagttacct gcctgtattg atagaaacgg ataaagcata taaaatactg
601 atatctgaag ccgacttatc cgattatccc tgtatgttcc ttaaaagtac cggtaagaac
661 ggaatgcagt ctatTTTTTcc caaagcacct ttagccttcg gagaagatgg tgaccgtagc
721 ctcaagatta ccgaagaagc cgattacatt gccaaagcgg acggcaaacg ttcattcccc
781 tggcgcgatga tgggtgatttc gaaagaagac aaagaactga ttgaaaacga aatgggtgat
841 aaactgtctg ctcttTgtgt tcttgaagac tacagttgga tcaaaccggg acaagtgagt
901 tgggaatggg ggcacgacgc acgcctctat ggggtagatt tccgttcggg tttcaatatg
961 gattcctata aatactacat tgacttcgca tccaagttcg gtattcctta tatcatcatg
1021 gacgaaggat gggcgaaaaa cacacgtgat ccgtttacc ccaatcccac catcaatctt
1081 accgaactga taaaatacgg aaaggaccgc aacgtaaaaa tcgtactttg gctgccatgg
1141 ctgactgtcg agaatcattt cgacctcttt aaaacatttg ccgattgggg catcgcagga
1201 gtgaagatcg acttcatgga ccgcagtgac cagtggatgg taaactatta tgaacgtgta
1261 gccaaagaag ccgccaaagc taaactgttt gtagattttc atgggtgctt taaaccagcc
1321 ggacttgaac gcaaatatcc gaatgtgctt tcctatgaag gcgtattggg catggaacaa
1381 ggtggtaatt gcaaacctga aaacagcatt tatctgccct ttatgcgtaa tgccgtggga
1441 ccgatggatt tcaactccggg ttcaatgatc tctgcacagc cggaagacaa ccgttccacc
1501 cgggccaatg ccatgggctc aggaacacgt gctttccaaa tggctctttt catcatcttc
1561 gaaagtggtc tgcaaatggt agccgacaat ccggtttact attacagaga acttccctgt
1621 accgaattta tcacaagtgt tcccgtcacc tgggatgaaa ccaaggtcct ctatgccaaa
1681 gtaggtgaag cagtcgtcgt agccaaacga aaaggagaac agtggttcat cggaggtatc
1741 accggcaatc aaccacaaaa catcgagatc gacctcggat tcattccggc aggacaatca
1801 ttcacattaa cctcatttga agatggcatt aacgctgacc gtcaagcaat ggattacaag
1861 aaaaaggagt ctaccgtgaa caatcaaacc cgcattgacat tgaaaatggt acgcaacggg
1921 ggatgggccg gaacaattaa aatgaaatag
```

9.2 BLAST analysis

The similarity of the genes and proteins of interest with other sequences was determined using the BLASTP (for proteins) and BLASTN (for DNA) links in the National Centre for Biotechnology Information site (www.ncbi.nlm.nih.gov/BLAST).

9.3 Primer Design

The primers were designed using the Web Primer server (<http://genome-www2.stanford.edu/cgi-bin/SGD/web-primer>) with a default value of 18 for the minimum length of the primer.

PCR primers for the amplification of the glycoside hydrolase family GH97 BF3763 gene in *B. fragilis* NCTC 9343.

Forward primer sequence:

5'- CACCACCACCAC ATGAAAGTGATTGACGTAATGTCTCCCAACGG-3' (44 bp)

Reverse primer sequence:

5'- GAGGAGAAGGCGCGTTATTTCAATTTAATTGTTCCGGCCCATCCCC- 3' (47 bp)

Refer to Appendix I for tables showing the conditions used for the amplification of the BF3763 gene from *B. fragilis*

PCR of the glycoside hydrolase gene GH95_BF0855 from *B. fragilis* NCTC 9343.

Primer sequences used for amplification

Forward primer

5'-CACCACCACCACATGTATGACTATTGTCCGGTCACTCCTTCAGAGAG-3'

Reverse primer

5'-GAGGAGAAGGCGCGTTAAGGGTTCAACTCGATCCATTCACCGGCTTGAGT-3'

The primer sequences were ordered from MWG Biotech, Germany and the Ligation independent cloning (LIC) specific ends were added to the respective ends of the primer sequences.

9.4 Genomic DNA extraction and purification in *B. fragilis* NCTC 9343

The DNeasy tissue kit was used for the extraction of DNA as per a standard protocol. A volume of 10 mL of LB medium was inoculated with a single colony of *B. fragilis* from a pure culture CBA plate. The medium was incubated overnight at 37°C anaerobically.

0.5 mL of the culture was centrifuged at 5000 x g for 10 min. The supernatant was discarded and the pellets were resuspended in 180 µL of Buffer ATL (Recipe patented). To this, 20 µL of proteinase K was added and vortexed well. The mixture was incubated overnight at 56°C in a water bath. The lysate was vortexed for 15 s and 200 µL of Buffer AL (Recipe patented) was added to it. The sample was mixed thoroughly and 200 µL of absolute ethanol (96-100%) was added to it. The sample was vortexed again vigorously and the mixture was pipette out into a DNeasy Mini spin column placed in a 2 mL collection tube. This was centrifuged at 14000 x g for 1 min. The flow through was discarded and the collection tube was replaced. 500 µL of Buffer AW1 (Recipe patented) was added to the Spin column and centrifuged at 14000 x g for 1 min. The flow through was discarded again and the 2 mL collection tube was replaced. 500 µL of Buffer AW2 (Recipe patented) was added to the spin column and centrifuged at 20000 x g for 3 min. The flow through and the collection tube were discarded. The DNeasy mini spin column was placed in a clean 2.0 mL microcentrifuge tube and 200 µL of Buffer AE (10 mM Tris-Cl, 0.5 mM EDTA pH 9.0) was pipetted out onto the column. The sample was incubated in room temperature for 1 min and the DNA was eluted out by centrifuging at 14000 x g for 1 min. The eluted DNA of interest can be stored at -20°C until use.

Agarose gel electrophoresis was performed to confirm the presence of the purified *B. fragilis* genomic DNA.

9.5 Agarose gel electrophoresis

A 1% (w/v) agarose gel was prepared by adding 1 g of agarose in 100 mL of 1 X TAE buffer and bringing it to boil to dissolve the solid bits. This was allowed to cool to about 60°C and then poured into a mini gel casting tray. Care must be taken to avoid air bubbles. A 12 toothed comb was then inserted into the solution. The gel was then left to set on the bench for 20 to 30 min.

Following this, the gel tray was placed horizontally in the electrophoresis tank and 1 x TAE buffer was poured to submerge the gel completely.

The comb was removed without damaging the gel and the samples were loaded into the wells. The samples were prepared by adding 5 µL of amplified DNA to 1 µL of the loading dye (bromophenol blue 6 X) along with 5 µL of the Bioline hyperladder standard (Refer to Appendix D). The electrophoresis was carried out at 100 mA, 200 V for 35 min.

The gels were then visualised using the Bio-Rad Gel doc system (using Quantity One™ software) and hard copies of the gel were printed out using the Mitsubishi Video Copy Processor attached to it.

9.6 PCR amplification of the genes of interest

KOD Hot start DNA polymerase was used to amplify the genes of interest in the PCR protocol using the Eppendorf MasterCycler™ machine.

9.7 Reaction components

PCR components added to each tube	GH95 (BF0855)	GH97 (BF3763)
Sterile 18.2 MΩ/cm water	34 µL	33.6 µL
12.5 µM Forward primer	1 µL	1.2 µL
12.5 µM Reverse primer	1 µL	1.2 µL
KOD reaction buffer 10 X	5 µL	5 µL
(Refer to Appendix for details of the KOD reaction buffer recipe)		
<i>B. fragilis</i> genomic DNA (50 ng/ml)	1 µL	1 µL
MgSO ₄ (25 mM)	2 µL	2 µL
dNTPs (5 mM)	5 µL	5 µL
KOD DNA polymerase (1U/µL)	1 µL	1 µL

The PCR conditions used to amplify the BF3763 and BF0855 genes in *B. fragilis* has been detailed in tables in the Appendix section I.

The amplified PCR products of BF3763 and BF0855 were analysed using agarose gel electrophoresis and purified using the MinElute PCR Purification Kit, QIAGEN.

To 90 μL of the PCR product, 450 μL of Buffer PB (Recipe patented) was added and mixed so that they were in the ratio of 1:5 respectively. The mixture was applied onto a MinElute column that was placed in a 2 mL collection tube and centrifuged at 14000 x g for 1 min. The flow through was discarded and the collection tube replaced. 750 μL of Buffer PE (Recipe patented) was used to wash the column and then centrifuged at 14000 x g for 1 min. The washing step was repeated again. The flow through was discarded and the column was placed in a clean 1.5 mL microcentrifuge tube. 10 μL of buffer BE (Recipe patented) was pipetted onto the centre of the membrane in the column, left for 1 min at room temperature and centrifuged at 14000 x g for 1 min. This step eluted the purified PCR product which was stored at -20°C until use.

9.8 Insertion of LIC T4 polymerase (T4 pol) reaction

The purified PCR product (10 μL) was mixed with 2 μL of 10 X T4 pol buffer (Refer to Appendix A2 for recipe details), 2 μL of 25 mM dATP, 1 μL of 100 mM dithiothreitol (DTT), 0.4 μL of 2.5U/ μL of LIC T4 DNA pol (Novagen) and made up to 20 μL by adding 5.6 μL of sterile 18.2 M Ω /cm water. The mixture was incubated at 22°C for 30 min. To stop the reaction of T4 DNA polymerase, the mixture was incubated at 75°C for 20 min and centrifuged at 14000 x g for 10 s.

For effective annealing of the genes, BF3763 and BF0855 to the LIC vector pET-YSBLIC, 2 μL of the reaction mixture resulting from the above was added to 1 μL of the prepared pET-YSBLIC vector (50 ng/ μL) and incubated at $20-22^{\circ}\text{C}$ for 10 min. This was then made up to 4 μL by adding 1 μL of 100 mM EDTA and incubated at room temperature for 10 min.

9.9 Preparation of electrocompetent TOP10 *E.coli* cells

The glycerol stock of the cells was used to inoculate a plate that was incubated overnight at 37°C . A single colony from this was used to inoculate 50 mL of low salt LB which was incubated at 37°C in a shaker at 200 rpm until it reached an OD of 0.5 to 0.7 at 600 nm. The culture was incubated on ice for 30 min and centrifuged at 4000 x g for 15 min at 4°C . A volume of 25 mL of cold filter sterilised 18.2 M Ω /cm water was used to resuspend the pellets which were then centrifuged again and the supernatant was discarded. The pellets were resuspended in 12 mL of cold 18.2

MΩ/cm water. The resuspension and centrifugation steps were repeated again reducing the volume of water to 5 mL and then to 2 mL. The steps were repeated once final time reducing the volume of water used to resuspend the pellets to 0.2 mL. The cells were aliquoted out into 40 μL quantities and stored at -80°C.

9.10 Transformation

A volume of 2 μL of the pET-YSBLIC vector containing the gene of interest was added to 40 μL of TOP10 *E. coli* cells, gently mixed with a pipette tip and incubated on ice for 1 min. This was transferred to a 2 mm electroporation cuvette which was kept ice cold and sterile. The cuvette was tapped gently in order to make sure that the solution covered the bottom of it. The cuvette was placed in a Bio-Rad ShockPod after drying the outside thoroughly and the lid was closed. The cells were electroporated using the BioRad Gene Pulser Xcell at 2.5 V for 20 ms. 1 mL of the SOC medium was added to the electroporated cells immediately and transferred into a clean sterile micro centrifuge tube using a sterile glass pipette. This was incubated at 37 °C for 60 min at 200 rpm and then plated out onto an LB agar plate containing kanamycin at a concentration of 100 μg/mL. The transformed cells were allowed to grow by incubating the plates at 37°C overnight.

9.11 Screening of colonies for successful inserts

The number of colonies present in the LB agar plates containing kanamycin was counted the following day. The colonies of interest were transferred onto a fresh 24 grid LB agar plate supplemented with an appropriate concentration of kanamycin. The first colony was streaked across the first grid on the agar plate. The cells remaining on the pipette were gently and thoroughly mixed into 50 μL of sterile 18.2 MΩ/cm water contained in a 0.2 mL PCR tube. The PCR tubes containing the cell-water suspensions were incubated in the PCR machine at 95°C for 5 min to disrupt the cells and centrifuged at 14000 x g for 2 min. The pellet was discarded and the supernatant was transferred into a clean sterile 0.2 mL PCR tube. This was stored at -20°C until use. The same process was repeated for each and every colony of interest on the agar plate.

9.12 PCR screening to identify colonies containing gene inserts

A volume of 10 μL of the supernatant from above was mixed with 1 μL of the forward primer (0.4 μM), 1 μL of the reverse primer (0.4 μM), 5 μL of dNTPs (2 mM), 1 μL of (5 U/ μL) Taq polymerase, 5 μL of Taq buffer (10 X) (Refer to Appendix A2 for recipe details) and made up to 50 μL with sterile 18.2 M Ω /cm water.

PCR conditions for the amplification of the BF3763 gene from transformed *E. coli* to screen for colonies containing successful inserts has been included in the Appendix section I.

The presence of successful inserts was confirmed by agarose gel electrophoresis. Refer to Methods section 9.5 for details on how to perform agarose gel electrophoresis.

Colonies containing successful inserts were used to inoculate LB broth media supplemented with an appropriate quantity of kanamycin and incubated overnight at 37°C in a 200 rpm shaking incubator.

9.13 Purification of the plasmid DNA from the colonies containing inserts

The Bio/Spin-PTM Plasmid Miniprep Purification protocol was used to purify the plasmid DNA from the *E. coli* TOP10 cell colonies containing the gene of interest. A volume of 1.5 mL of the cell culture from overnight incubation was centrifuged at 14000 x g for 1 min and the supernatant was discarded. The same step was repeated twice and the supernatant was discarded with a sterile pipette tip. A volume of 200 μL of Buffer I (Recipe patented) was added and vortexed to resuspend the cells. To this, Buffer II (Recipe patented) of a volume of 200 μL was added and mixed by inverting 5 to 6 times. To this, 200 μL of Buffer III (Recipe patented) was added and mixed by inverting 5 to 6 times and centrifuged at 14000 x g for 5 min. A spin column was placed in a collection tube. The mixture was pipetted into the column and centrifuged at 14000 x g for 30 s. The flow through was discarded and 700 μL of the washing solution was added to the column and centrifuged at 14000 x g for 1min. The flow through was discarded and the column was centrifuged at 14000 x g for 3 min to remove any residual ethanol from the washing step. The flow through was discarded and the column was placed in a clean sterile 1.5 mL microcentrifuge tube. A volume of 50 μL of TE solution was added to the column and

centrifuged at 14000 x g for 1 min to elute the plasmid DNA which was stored at -20°C until use.

9.14 Method used to prepare chemically competent cells

The glycerol stock of the cells was used to inoculate a plate that was incubated overnight at 37°C. A single colony from this was used to inoculate 50 mL of low salt LB which was incubated at 37°C in a shaker at 200 rpm until it reached an OD of 0.35 to 0.4 at 600 nm. The cells were incubated on ice for 30 min and then centrifuged at 4000 x g for 10 min at 4°C. After discarding the supernatant, the pellets were resuspended in 15 mL of cold 80 mM MgCl₂, 20 mM CaCl₂ solution and then centrifuged at 4000 x g for 10 min at 4°C. The pellets were then resuspended in 1 mL of 100 mM CaCl₂. The cells were incubated on ice for 1 h. The cells were added to 50 % (v/v) glycerol making it up to a final concentration of 15 % (v/v) before freezing it in liquid nitrogen and storing at -80°C until use.

9.15 Restriction digests of the recombinant BF3763 and BF0855 genes from pET-YSBLIC

The restriction enzymes used to carry out the digestion was *Eco* RI and *Xba* I according to the analysis of the sequence in Webcutter 2.0 server (<http://www.firstmarket.com/cutter/cut2.html>). The restriction enzyme digestion was carried out to remove the recombinant genes from the vector. The restriction enzymes were kept on ice and centrifuged for 3 s before use. To 5 µL of the plasmid DNA, 0.5 µL of *Xba* I, 0.5 µL of *Eco* RI, 1 µL of 10 X NE Buffer 2, (Refer to Appendix A2 for recipe details) 1 µL of BSA (50 µg/ml (w/v)) and 2 µL of sterile 18.2 MΩ/ cm water was added to make it up to 10 µL. The mixture was incubated at 37°C for 2 h and the presence of the recombinant genes was analysed using agarose gel electrophoresis (Refer to section 9.5 in the Methods section).

9.16 Transformation of the recombinant genes into BL21 cells

Chemically competent BL21 cells were prepared and 50 µl of the cells were treated with 2.5 µL of the purified plasmids by gentle mixing on ice for 5 min with a sterile tip. The cells were subjected to heat shock at 42°C in a water bath for 90 s and then immediately transferred back onto ice. A volume of 200 µL of NZY⁺ medium was

added to the mixture aseptically. The cell wall recovery was induced by incubating the mixture in a water bath at 37°C for 45 min. The transformed cells were plated out onto LB agar plates containing an appropriate amount of kanamycin and incubated overnight at 37°C. The number of colonies showing successful inserts was 2 for GH95 and 3 for GH97.

The genes for GH95 and 97 have been cloned by previous students. All the details are available in the dissertations ('Isolation and transformation of GH95_BF0855 gene from *Bacteroides fragilis* NCTC 9343 in competent cells to produce soluble protein' by Cheun Hong Yeap, 2006 and 'Cloning and expression of glycoside hydrolase family 97 BF3763 (putative exported protein) gene from the *Bacteroides fragilis* NCTC 9343 ATCC 25285 in *Escherichia coli*' by Lee Ling, Bong in 2006). Only experiments pertaining to the purification and characterisation of these enzymes have been performed by me.

9.17 Solubilisation buffer experiment

9.17.1 Growth of bacteria in starter cultures

E. coli cells were grown in 30 mL sterile plastic containers containing 10 mL of LB media or 10 mL of autoinduction media starter cultures supplemented with an appropriate amount of kanamycin. The cultures were inoculated from an agar plate or glycerol stock using a sterile inoculation loop and grown overnight in a shaking incubator at 37°C and 200 rpm.

9.17.2 Growth of bacteria for expression and purification using IPTG induced media

An initial study was done to compare the expression of protein in LB and auto-induction media. In both cases, 0.5 mL of the starter culture of BL21 *E. coli* cells was used to inoculate 50 mL of the starter culture. Following the inoculation of the media, the cultures were grown at 37°C in a shaking incubator at 200 rpm.

When the culture reached an OD of 0.6, the flask containing LB medium was induced for the expression of the proteins of interest by the addition of 100 mM IPTG

(Isopropyl β -D-1-thiogalactopyranoside) to make the final concentration 1 mM and then the temperature was reduced to 30°C and the shaking reduced to 100 rpm. However, for auto-induction media cultures, the temperature was reduced to 30°C and the shaking to 100 rpm when it reached an OD of 0.6 without the addition of IPTG. The cultures were then incubated overnight and processed the following morning.

9.17.3 Small scale protein extraction for the solubilisation buffer experiment

The overnight culture was taken the following day and 3.0 mL of it was transferred into two clean microcentrifuge tubes of 1.5 mL each. The tubes were centrifuges at 14000 x g for 1 min. Following centrifugation, the supernatant was discarded and the pellets were dissolved in 300 μ L of the start buffer. The mixture was ultra-sonicated for 5 s at amplitude of 15 and maintained on ice while doing so. The cells were centrifuged at 14000 x g for 10 min and 200 μ L of the supernatant also known as the cell free extract was transferred into a clean microcentrifuge tube. The remaining supernatant was discarded and the pellets were resuspended in 300 μ L of the solubilisation buffer. An SDS-PAGE gel was run with 20 μ L of the sample and 5 μ L of the loading buffer except the solubilisation mixture which was placed in a boiling water bath for 1 min to dissolve the pellets and 20 μ L of it was loaded on to the SDS-PAGE gel. Molecular weight size standards were prepared with 8 μ L of the loading buffer and run along with the samples in order to determine the size of the expressed protein.

9.17.4 SDS-PAGE electrophoresis

The BioRad Mini Protean III kit was used to perform 2D SDS-PAGE. Two clean glass plates of dimensions 10.1 X 7.2 cm and 10.1 X 8.2 cm were aligned parallel to each other and clamped. The larger glass plate had a spacer ridge of 0.75 mm attached to it. The two glass plates along with their casting clamp stand were mounted on the surface of a rubber gasket by applying vertical downward pressure. The resolving gel solution was made up in a plastic container and pipetted into the space between the two plates. A 2 cm mark was made from the top of the smallest plate and the solution was filled up to it. For efficient polymerisation, the surface of the gel was layered with 18.2 M Ω /cm water and the gel was allowed to polymerise

for 20 to 30 min. The water was removed from the surface of the set gel and blotting paper was used to remove any residual unset gel or water. Care was taken to avoid touching the surface of the gel with the blotting paper. The stacking gel solution was then made up and pipetted on to the surface of the resolving gel. This was filled up to the top of the small glass plate and a 10 toothed comb was inserted into the gap between the plates immediately (Size of the comb was 1.1 X 0.75 cm). The gel was then allowed to polymerise for 20 min and the combs were slowly removed to reveal the wells. Care was taken to avoid the formation of air bubbles while doing so and the surface of the gel was rinsed with 18.2 MΩ/cm water. The gels were placed in the electrophoresis module in such a way that the smaller glass plates faced towards the centre. The module was then lowered into the electrophoresis tank and SDS-PAGE (1 X) running buffer was poured into it to fill both the central and the outer compartments. Care was taken to ensure that the electrodes were completely immersed in the buffer and no air bubbles formed. The buffer in the central compartment was allowed to flow into the wells and formed a layer on the top of the gel.

The prepared samples were placed in a boiling water bath for 3 min and then centrifuged at 14000 x g for 1 min before loading them onto the gel with the help of a Hamilton syringe. The electrophoresis was performed at 120 mA and 200 V for about 50 min. The completion of the process was indicated by the migration of the bromophenol blue dye to the bottom of the gel.

Upon completion of electrophoresis, the gels were removed from the set up and placed in a plastic container containing Coomassie blue staining solution and rocked gently on a shaker for 10 min at room temperature. Following this, the gels were immersed in the destaining solution for atleast 6 to 8 h. Once the bands were clearly visible, the gels were rinsed with 18.2 MΩ/cm water and then visualised using the BioRad gel documentation system. Photographs of the gel were obtained using the Mitsubishi Video Copy Processor containing the Mitsubishi thermal paper (Refer to Appendix B for details).

9.18 Large scale growth of bacteria

A large scale growth of bacteria was carried out using the auto-induction medium where 500 mL of the medium was inoculated with 5 mL of the starter culture.

The growth procedure was followed exactly the same way as mentioned above.

9.18.1 Protein extraction

The induced cells were centrifuged at 4000 x g for 15 min and were resuspended in 5 mL of re-suspension buffer. The cells were ultra-sonicated on ice at amplitude of 14 for 2 min in 10 s bursts with 10 s intervals. The lysate was then transferred into centrifuge tubes and then spun at 24000 x g for 30 min at 4°C. The liquid supernatant or the cell free extract was transferred into a sterile plastic universal and stored in ice until purification.

9.18.2 Protein Purification

The recombinant proteins were purified using immobilised metal affinity chromatography (IMAC). Proteins have the presence of hexa-histidine tags at the N terminal that could be used to separate it out from other *E. coli* proteins using the SepharoseTM chelating fast flow resin which acts as the metal affinity matrix. The column was first stripped and charged using 1M NiSO₄ followed by equilibration with 200 mL of start buffer. After equilibration, the cell free extract (10-20 mL) was loaded onto the column and then eluted out with 100 mL of the elution buffer at a flow rate of 4 mL/min. The presence of the protein was monitored using a UV spectrophotometer and the eluted fractions were collected in 4 mL quantities in plastic tubes. A volume of 20 µL of each sample was run on an SDS-PAGE gel to confirm and detect the fractions containing the pure protein. The pure protein fractions were then concentrated using a Vivaspin protein concentrator and then resuspended in the gel filtration buffer.

Further purification of the proteins was carried out using gel filtration chromatography. The column was first equilibrated using 120 mL of the gel filtration buffer at a rate of 1 mL/min. A volume of 0.5 mL of the protein sample was loaded onto the column by injection and then eluted out at a rate of 1 mL/min. The presence of the pure protein was detected using the UV spectrophotometer and the eluted fractions were collected in 5 mL quantities in plastic tubes. A 20 µL sample from every fraction was run on an SDS PAGE gel to confirm the presence of the pure protein. The fractions of interest were pooled and concentrated.

9.18.3 Concentration of proteins

The pure proteins were concentrated using the 30 kDa Vivaspin 6 mL cut off concentrator units (Viva Science). The pooled fractions were concentrated by centrifuging at 4000 x g and were buffer exchanged with 5 mM HEPES (pH 7.4) within the concentrator units.

9.19 Concentration of proteins determined by Bradford's assay

Standard dilutions of BSA were prepared within a range of 1-10 µg/mL and the total volume was made up to 500 µL. The absorbance of the solutions was recorded at 595 nm using a visible spectrophotometer and a standard curve was drawn based on the values obtained. A blank was set up containing 18.2 MΩ/cm instead of BSA and added to the Bradford reagent. The protein solution was then diluted appropriately and 500 µL volumes were transferred into cuvettes. The absorbance values were measured and plotted against the standard to determine the protein concentration.

From the standard curve, the value of c was found to be 0.6 which was the point on the y axis where the line intercepted. Substituting the rest of the values in the equation below, the value of m was determined to be 0.52.

Unknown concentrations of protein samples and their dilutions 1:10, 1:50 and 1:100 were prepared and mixed with the Bradford reagent. The absorbance values at 545 nm was measured and applied to the equation $y = mx + c$ where y = Absorbance at 545 nm, x = the concentration of the unknown sample, m = the gradient of the straight line and c = the point of intercept of the line with the y axis.

Once the concentration of the unknown protein was determined, it was multiplied by the dilution factor in order to obtain the concentration in mg/ mL.

Protein conc. mg/ mL = Conc. obtained from the curve X dilution of the sample/1000.

20 µL of known concentrations of BSA were prepared and mixed with 200 µL of the Bradford reagent and the absorbance was measured at 545 nm. The concentration range of BSA was 0.1 to 1.0 mg/mL to generate a standard curve and the standard curve has been included in the Appendix section J.

The concentration of the GH95 protein was determined to be 43 mg/mL from the equation with an absorbance of 0.823 and a dilution factor of 100. The concentration of the GH97 protein was determined to be 29.2 mg/mL from the equation with an absorbance of 0.752 and a dilution factor of 100.

The proteins were diluted to obtain a final concentration of 15 mg/mL with 5 mM HEPES buffer pH 7.4 before crystallisation.

9.20 Crystallisation

The protein concentration was determined using Bradford assay and then crystallisation trays were set up.

9.20.1 Crystallogenesis

1 ml of liquid Aquasil was mixed with 999 mL of 18.2 M Ω /cm water in a 2000 mL beaker and the cover slips were added to the solution.

The liquid solution was stirred every 15 min for about 2 h. The cover slips were taken out and allowed to dry on a tissue paper. The cover slips were wiped down with a clean cloth before use. The crystal tray wells were filled with the appropriate buffer solutions from various screen aliquots and the cover slips containing drops of protein and buffer in three different concentrations of 1:1, 1:2, 2:1 respectively were inverted onto the surface of the wells which were greased beforehand. Once the tray of 24 wells was set up, they were incubated at 22.4°C in the crystallisation incubator and monitored for the presence of crystals every week. The various crystal screens and their compositions have been included in Appendix H. Crystals were harvested in rayon fiber loops, bathed in cryo-protectant solution prior to flash freezing in liquid nitrogen.

9.20.2 Data collection

Diffraction data were collected on the European Synchrotron Radiation Facility (ESRF) from single crystals at 100 K with an oscillation range of 0.5 °. Data were collected at a wavelength of 0.933 Å on beamline ID14-2 using an ADSC Q4R CCD detector. All diffraction data were processed with MOSFLM and reduced and scaled with SCALA (Leslie, 1992). All other computing was undertaken using the CCP4

suite (Anon, 1994) unless otherwise specified. The structure of *BfGH97* was solved by molecular replacement using the program PHASER using the *Bacteroides thetaiotaomicron BtGH97b* (UniProtKB/TrEMBL entry Q8A6L0) structure (Okuyama *et al.*, 2009) (PDB code 3A24) as a search model. 5 % of the total reflections were set aside for cross validation (“R_{free}”) purposes and for the weighting of geometrical and temperature factor restraints. The structure was manually built with cycles of REFMAC (Murshudov *et al.*, 1997). Solvent molecules were added using COOT (Emsley & Cowtan, 2004) and checked manually. The structure was validated using MOLPROBITY (Davis *et al.*, 2007).

9.21 Enzyme assay

9.21.1 Preparation of substrates

A volume of 1 mL of 6 mM substrates were prepared by dissolving the substrate in 18.2 MΩ/ cm. Since the methylumbelliferone substrates were found to be difficult to dissolve, they were ultra-sonicated (15 amplitude) at 1 min intervals until the substrates dissolved completely. The appropriate volumes required for the assay were aliquoted out into clean microcentrifuge tubes. Care was taken to perform the assays with freshly prepared substrates.

9.21.2 Enzyme assay protocol

Two different assay methods were used to characterise and determine the specific activity of the enzymes expressed from GH95 and 97 namely fluorogenic and chromophoric assays. The putative enzymes expressed from GH95 and 97 were α-fucosidase and α- glucosidase respectively.

9.21.3 Chromophoric assay

p-Nitrophenyl α-D-Glucoside → α-D-Glucose + p-Nitrophenol

CONDITIONS: T = 37°C, pH = 6.8, A 400nm, Light path = 1 cm

The substrates used were p-nitrophenyl α-D-glucoside, p-nitrophenyl α-D- fucoside, p-nitrophenyl β-D-maltoside and p-nitrophenyl α-D-galactoside. The activities of the enzymes were measured by their action against the substrates resulting in the release of 4- nitrophenol indicated by a change of colour to yellow. The assay

mixture contained 20 mM of the substrate, 5 mM HEPES pH 7.4, 10 mg/mL BSA and 15 μ L of the enzyme at a final concentration of 10 mg/mL. The volume of the assay mixture was made up to a total volume of 500 μ L. The mixture was transferred to a clean plastic cuvette, mixed well with a sterile pipette tip and incubated at 37°C. The change in absorbance values at a wavelength of 420 nm was noted at 1 min intervals.

9.21.4 Fluorimetric assay

The substrates used to carry out the fluorimetric assays include 4-methylumbelliferyl α -D glucoside, 4-methylumbelliferyl α -D-fucoside and 4-methylumbelliferyl β -D-maltoside. These are synthetic substrates that react with enzymes to release a product called 4-methylumbelliferone (4-MU). 4-MU is a fluorescent dye whose fluorescence was measured using the Bio-Tek FL600 fluorescence microplate reader. The excitation wavelength used was 360 nm and the emission wavelength was 460 nm. Standard assays were performed at 37°C in a total volume of 500 μ L of 5 mM Hepes buffer, pH 7.4, containing 10 mg/mL BSA and 0.15 μ g of the enzymes, α -D-glucosidase and α -D-fucosidase. The reaction mixture without the enzyme was transferred into a clean glass cuvette and placed into the cuvette holder in the fluorescence densitometer. Once the excitation and emission wavelengths were set, the enzyme was quickly added and mixed well with a sterile tip and the photometric values were measured at 10 min intervals for 60 min. The cell containing the reaction mixture was maintained at 37°C by a temperature controlled circulator. The photometric recordings were traced out onto chart paper using a chart recorder with a recorder pen attached to it. Positive activity was detected in the case of both enzymes.

Two different types of substrates were used in the study to determine the activity of the recombinant enzyme, α -glucosidase purified from our experiment. Para nitrophenyl- α -D- glucopyranoside was used as a substrate where α -D-glucose and p-nitrophenol were expected to be produced as end products as a result of the hydrolysis of the α 1 \rightarrow 6/ α 1 \rightarrow 4 linkages in the substrate. The release of para nitrophenol into the reaction mixture would result in the development of a yellow colour. The intensity of the yellow colour indicated the amount of product released and was measured spectrophotometrically at 420 nm in the colorimetric assay

(Anon, 1996). The other substrate that was used in the study was 4-methylumbelliferol α -D-glucoside. The α -glucosidase carried out an enzymatic hydrolysis of the fluorescent sugar resulting in the release of 4-methylumbelliferone as a byproduct of the reaction. The presence of the product in the reaction mixture was analysed using a fluorescent spectrophotometer where the excitation and emission wavelengths were used to measure the amount of product released in the fluorimetric assay (Sheldon *et al.*, 2006).

The BF0855 gene from *B. fragilis* NCTC 9343 was cloned and expressed in *E. coli* and the putative activity of the enzyme was predicted to be α -L-fucosidase or 1, 2 α -L-fucosidase. α -L-fucosidase catalyses the reaction between α -L-fucoside and water resulting in the formation of L-fucose as an end product. They belong to the family of glycoside hydrolases that hydrolyse O and S glycosyl compounds. They are also known as α -L-fucoside fucosidase. It is involved in n-glycan and glycan structure degradation. In the presence of 4-methylumbelliferol α -L-fucoside, 1, 2- α -L-fucosidase catalyses the release of 4-methylumbelliferone as a byproduct of the reaction and this can be used in its fluorimetric assay. The substrate used in the colorimetric assay of α -L-fucosidase was para-nitrophenyl- α -D-fucoside.

10. Results and discussion of the proteomic analysis of *Bacteroides fragilis* NCTC 9343 and *Enterobacter cancerogenus* ATCC 35316

Enterobacter cancerogenus and *Bacteroides fragilis* are some of the predominant flora of the gut and are capable of acting as opportunistic pathogens by causing gut associated infections. *B. fragilis* originate in the endogenous flora of the mucous membranes and are capable of causing a number of infections including intra abdominal abscesses, soft skin and tissue infections, bacteraemia, gangrene, necrotizing cellulitis and so on. They are also capable of causing severe secondary infections that include central nervous system infections, pleuropulmonary infections and septic arthritis (Pumbwe *et al.*, 2006). Since *Bacteroides* spp. are strictly anaerobic in nature, it hard to detect them during specimen culture and analysis. *E. cancerogenus* has been known to occur as a secondary wound infection causing bacteria. They have been isolated from human faeces, found to be associated with the human gut and form a part of the normal human gut microbiota.

A number of studies have been performed on the differential expression of proteins in *B. fragilis* and *Enterobacter* sp. in response to oxidative stress, antibiotic and so on. In order to determine the proteins that may be differentially expressed in metronidazole-resistant strains of *B. fragilis*, mutants lacking the flavodoxin and pyruvate-ferredoxin oxidoreductase genes were generated and studied (Diniz *et al.*, 2004). Other proteomic studies in *B. fragilis* were performed to determine the oxidative stress response (Rocha *et al.*, 2003). A top down proteomics approach has been used to study the differential expression of heat resistant strains (Williams *et al.*, 2005) and osmotic stress response (Riedel & Lehner, 2007) in *E. sakazakii* and selenite resistance in *E. cloacae* (Barasa, 2008). Since *E. cancerogenus* infections are not as commonly occurring as *B. fragilis* infections, no proteomics experiments have been performed to determine its virulence factors.

10.1 Growth analysis of *E. cancerogenus* and *B. fragilis*

The two strains, *Bacteroides fragilis* NCTC 9343 and *Enterobacter cancerogenus* ATCC 35316, were grown anaerobically at 37°C in an anaerobic chamber. The presence of Gram-negative bacteria was confirmed by Gram staining (Refer to the Figs.12 and 13).

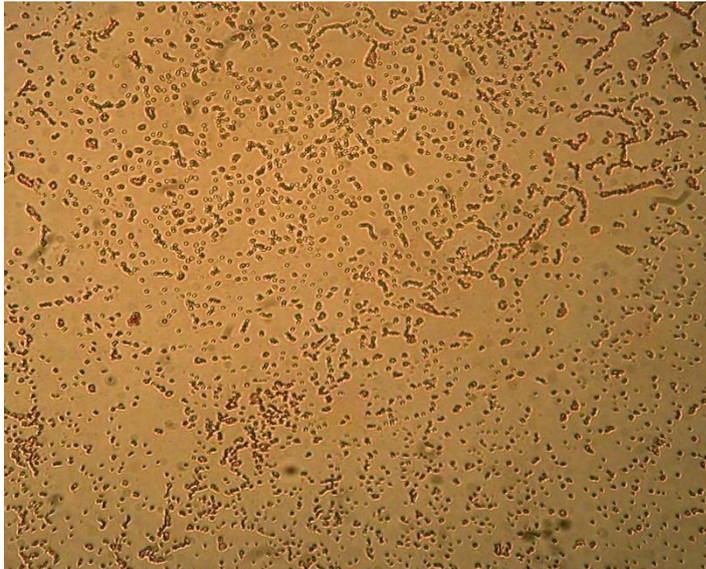


Figure 12: Gram staining results for *E. cancerogenus* showing the presence of small rod shaped bacilli that stained pink.

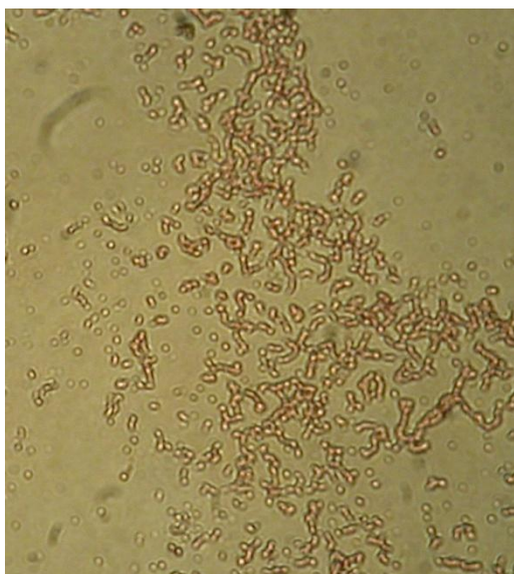


Figure 13: Gram staining results for *B. fragilis* showing the presence of small rod shaped bacilli that stained pink.

Minimal medium was prepared as described in the methodology section and the bacteria were grown in the media. The growth rate of both *B. fragilis* and *E. cancerogenus* was found to be very slow taking about 160 to 170 h to reach an OD of 1.0 at 600 nm. Hence modifications were made in the minimal media where yeast extract was used as a source of vitamin B and the concentration of casitone was increased from 0.2 g to 0.5 g per 100 mL of medium. The growth in this medium (semi- defined media) was found to be much quicker with the culture attaining an overnight stationary phase in case of *E. cancerogenus* (12-16 h) and 40 h for *B. fragilis*. Table 3 showing the growth of *E. cancerogenus* and *B. fragilis* in minimal growth media have been included below. Table 4 included below details the growth of bacteria (*E. cancerogenus* and *B. fragilis*) in semi-defined media containing various sources of carbon.

Time in h	Average absorbance at 600 nm	
	<i>E. cancerogenus</i>	<i>B. fragilis</i>
0	0.000	0.000
12	0.008	0.015
24	0.023	0.044
48	0.158	0.102
72	0.278	0.304
96	0.488	0.417
120	0.821	0.632
144	1.025	0.954
168	1.238	1.189
192	1.143	1.178

Table 3: Growth of *E. cancerogenus* and *B. fragilis* in minimal media

Type of carbon source used in the semi-defined media	Average OD at 4 h after inoculation		Average OD at 8 h after inoculation		Average OD at 16 h after inoculation (overnight incubation)	
	<i>E. cancerogenus</i>	<i>B. fragilis</i>	<i>E. cancerogenus</i>	<i>B. fragilis</i>	<i>E. cancerogenus</i>	<i>B. fragilis</i>
Glucose	0.839±0.0008	0.223±0.0009	1.036± 0.002	0.539±0.0012	1.304± 0.0025	0.728±0.0008
Mucin Type II	0.378±0.001	0.035±0.0008	0.367± 0.156	0.156±0.0008	0.358± 0.002	0.462±0.0017
Mucin Type III	0.355 ±0.0005	0.038±0.0012	0.353± 0.093	0.201±0.0008	0.325± 0.0008	0.512±0.0008
Glucose and Type II mucin (0.5 g/100 mL of glucose and mucin Type II)	0.993±0.0012	0.324±0.0012	1.148 ±0.001	0.498±0.0012	1.316±0.0016	0.819±0.0017
Glucose and Type III media (0.5 g/100 mL of glucose and mucin Type III)	1.090±0.0124	0.287±0.0009	1.220±0.025	0.562±0.0016	1.414 ±0.0017	0.753±0.0016
Type II and III mucin (0.5 g/100 mL of mucin Type II and Type III)	0.956±0.0008	0.045±0.001	1.134±0.002	0.198±0.0008	0.614±0.0012	0.485±0.0012
Glucose and Type II and III mucin (0.5 g/100 mL of glucose, mucin Type II and Type III)	0.068±0.0017	0.015±0.0017	0.881±0.0017	0.367±0.0008	0.821± 0.0012	0.328±0.0008

Table 4: Growth of bacteria in the semi-defined media with different carbon sources

All experiments were performed in duplicate and the values represented in the tables indicate average values of absorbance (600 nm) at the respective time points. All the carbon sources were added at a concentration of 0.5 g per 100 mL of the media. A quantity of 0.5 g of mucin was weighed out from the bottle and added to the medium rather than measuring the concentration of sugar in it.

10.1.1 Growth analysis of *E. cancerogenus*

Media (450 mL) inoculated with a 1 mL volume from 10 mL overnight starter cultures showed the following growth results over a period of 8 h (Fig. 14). The optical density values were measured at 600 nm at various time periods. An average of three results is represented in the graph below for both OD values against time and three different media have been used.

All the growth curve experiments were performed in triplicate and vertical error bars representing the standard deviation from the mean can be observed in the graph.

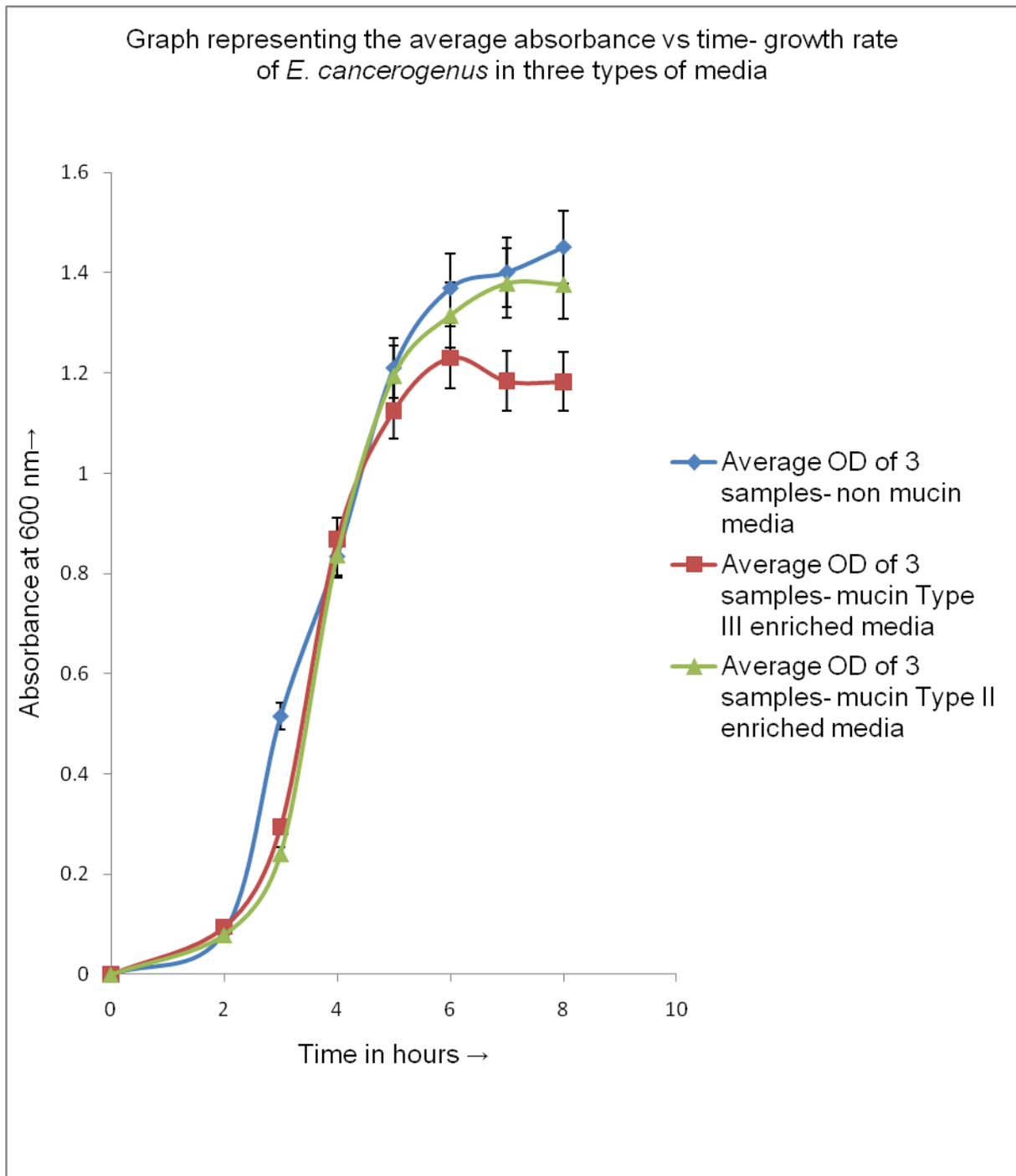


Figure 14: Graph representing the average absorbance vs time- growth rate of *E. cancerogenus* in three types of media

10.1.2 Growth analysis of *B. fragilis*

Media (450 mL) inoculated with a 1 mL volume from 10 mL overnight starter cultures showed the following growth results over a period of 48 h (Fig. 15). The optical density values were measured at 600 nm at various time periods. An average of three results is represented in the graph below for both OD values and colonies forming units against time and three different media have been used.

All the growth curve experiments were performed in triplicate and vertical error bars representing the standard deviation from the mean can be observed in the graph.

The average absorbances for the growth of *E. cancerogenus* and *B. fragilis* in semi-defined media have been included in the Appendix section E in the following tables. Tables 5, 6 and 7 included in the Appendix E show the growth of *E. cancerogenus* in non mucin, mucin Type II and Type III enriched media, respectively. Tables 8, 9 and 10 included in the Appendix E show the data for the growth of *B. fragilis* in non mucin, mucin Type II and Type III enriched media, respectively.

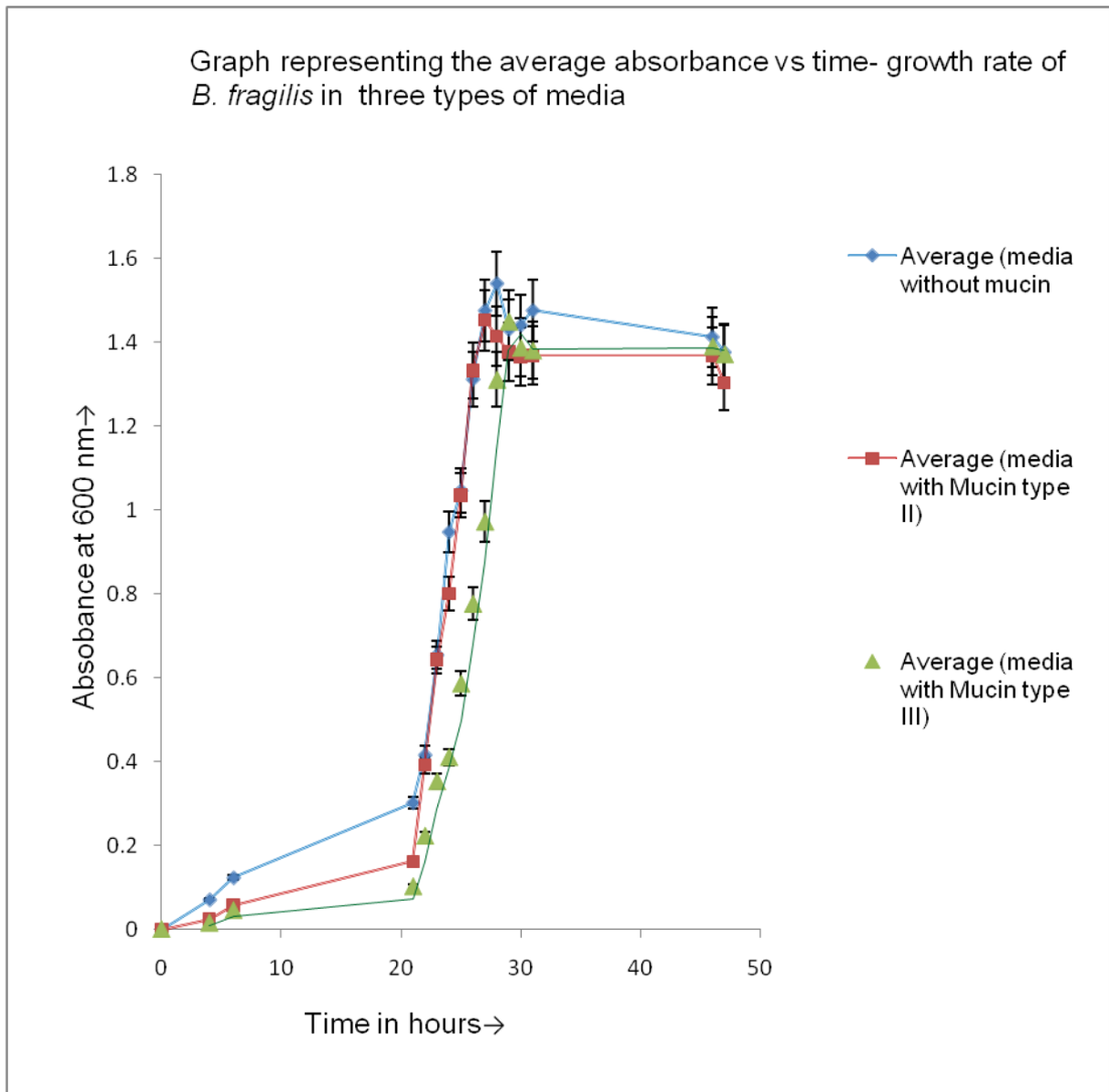


Figure 15: Graph representing the average absorbance vs time- growth rate of *B. fragilis* in three types of media

The harvesting time point for the cultures of *E. cancerogenus* and *B. fragilis* were determined based on halving the maximum OD value achieved during its growth in the semi-defined media as 0.6 and 0.7, respectively. This would also ensure that the cells were in their mid log phase.

10.2 Growth of *E. cancerogenus* ATCC 35316 and *B. fragilis* NCTC 9343 in semi-defined media

Both bacteria were grown in three types of semi-defined media which included media without mucin, media enriched with porcine intestinal mucin Type II and media enriched with porcine intestinal mucin Type III. Even though initial attempts were aimed at growing the bacteria in minimal media with and without mucin, it was found that the extremely slow rate of growth of bacteria hindered the progress of experiments and hence the medium was slightly modified by the addition of yeast extract, replacing the vitamin solution and increasing the concentration of casitone from 0.2 to 0.5 g to enable quicker growth of the bacteria.

Growth of the bacteria was studied in the logarithmic phase rather than the stationary phase because most of the virulence factors and other important proteins are normally expressed in the logarithmic phase of growth (Ernst *et al.*, 1990).

Several carbon sources were used to study the growth of *E. cancerogenus* and *B. fragilis* and it was found that the semi-defined medium containing glucose on its own (or in combination with mucin) supported efficient growth. In a semi-defined medium containing only mucin as a source of carbon, the growth rate was found to be low, with the optical density not increasing beyond 1.0, when compared to a semi-defined medium containing glucose and mucin. This may be due to the general tendency of bacteria to use glucose as the first source of carbon and other carbohydrate sources are utilised only when glucose is completely depleted in the media (Studier, 2005).

10.3 Mucin used in the study

The two types of mucin used were porcine gastric/intestinal mucin Type II and III. The main difference between the Type II and Type III medium is that mucin Type III is a partially purified powder that has the presence of 0.5 to 1.5% bound sialic acid. The sugar and amino acid composition of mucin has not been determined by the Sigma-Aldrich company from where it was purchased. The Type II mucin is a crude extract of mucin glycoproteins from the hog stomach using the enzyme pepsin and it has about 1% of bound sialic acid. Since the study aimed at analysing the differential protein expression in opportunistic pathogens when they come in contact with the

first line of cell defence, mucin, the closest possible system to the human body was necessary. Previous studies show that the core structures found in porcine gastric mucins are the same as that in human gastric mucins (Karlsson *et al.*, 1997). One of the most easily accessible types of mucin was the porcine gastric mucin without the involvement of any ethical issues thereby encouraging the incorporation of mucin as a potential source of carbohydrates in the semi-defined medium.

Out of several mucin types used in a study, ovomucoid, porcine gastric mucin and beef submaxillary mucin were found exhibit high similarity to human colonic mucin (Salyers *et al.*, 1977). The variations mainly occur in a relatively small number of basic structures. Some of these variations include the presence of acidic structures derived from neutral oligosaccharides in human colonic mucin, variations in the degree of sialylation, the biantennary structures of mucin oligosaccharides and also the branched chain oligosaccharides which appear to be linear chain derivatives of the same core structure (Podolsky, 1985). High levels of sialylation have been observed in both human and porcine colonic mucin. The hydroxyl groups present at the end of the sialic acids that contain acetyl esters inhibit bacterial degradation of mucin (Corfield *et al.*, 1992). Disulphide rich domains have been identified in both human intestinal and porcine submaxillary mucins but there may be a low degree of sequence identity (PerezVilar *et al.*, 1996).

It is possible that the activity of *B. fragilis* on colonic mucin may be different to that of gastric mucin since gastric mucin has been used in our studies due to the unavailability of colonic mucin.

My results show the expression of a number of proteins related to oxidative stress and heat shock which indicate the presence of stress during growth. Previous proteomics studies in *E. faecalis* have revealed the expression of a number of stress associated proteins in exponentially grown cells, viable but non culturable cells and proteins expressed during starvation conditions (Heim *et al.*, 2002). These stress proteins are generally expressed in response to bacterial growth and adaptability to various environmental conditions.

Proteome profiles obtained from the growth of bacteria in mucin enriched media suggest that the differentially expressed proteins may be associated with colonisation where bacteria adapt themselves to bind to the mucous layers of the

gut. Bacterial proteins associated with the ability to induce an immune response within the host could act as potential virulence factors and eventually result in infection. In the proteomics studies, first dimension separation of proteins was limited to a pH range of 4-7 rather than a variety of different pH gradients. Performing experiments with different second dimension separations may also provide a better insight into understanding differential protein expression. There are a large number of bacteria that occur commonly on mucous membranes forming a part of the normal gut flora by adhering and colonising mucin. But certain environmental adaptations within them can result in their ability to produce enzymes and compounds that lead to virulence.

10.4 Optimisation of two dimensional polyacrylamide gel electrophoresis (2DE-PAGE) gels

10.4.1 Comparison of staining methods

An attempt was used to compare the Coomassie blue staining with fluorescent staining, but due to the inconvenience of the requirement to irradiate with UV light when excising protein spots and poorer resolution, the Coomassie blue staining technique was chosen to analyse the spots of interest. On comparison of the gels represented in Figs 16.1 and 16.2 it is evident that the spots have been expressed with better clarity and resolution when colloidal Coomassie blue stain was used. From the figure, it was evident that fluorescent staining failed to achieve the required sensitivity levels and seemed inconsistent when compared to Coomassie blue staining. The process of cutting out spots of interest would be a more tedious process if fluorescent staining was used since this would involve working in the presence of UV light. The gel looks smudgy and the presence of a dark background may obscure the number of low abundance proteins observed. The improper background staining may be attributed to the trapping of the fluorescent dye in detergent micelles and this has been observed in previous studies (Malone *et al.*, 2001).

Figure 16: Figure showing the difference between fluorescent staining and Coomassie blue staining.

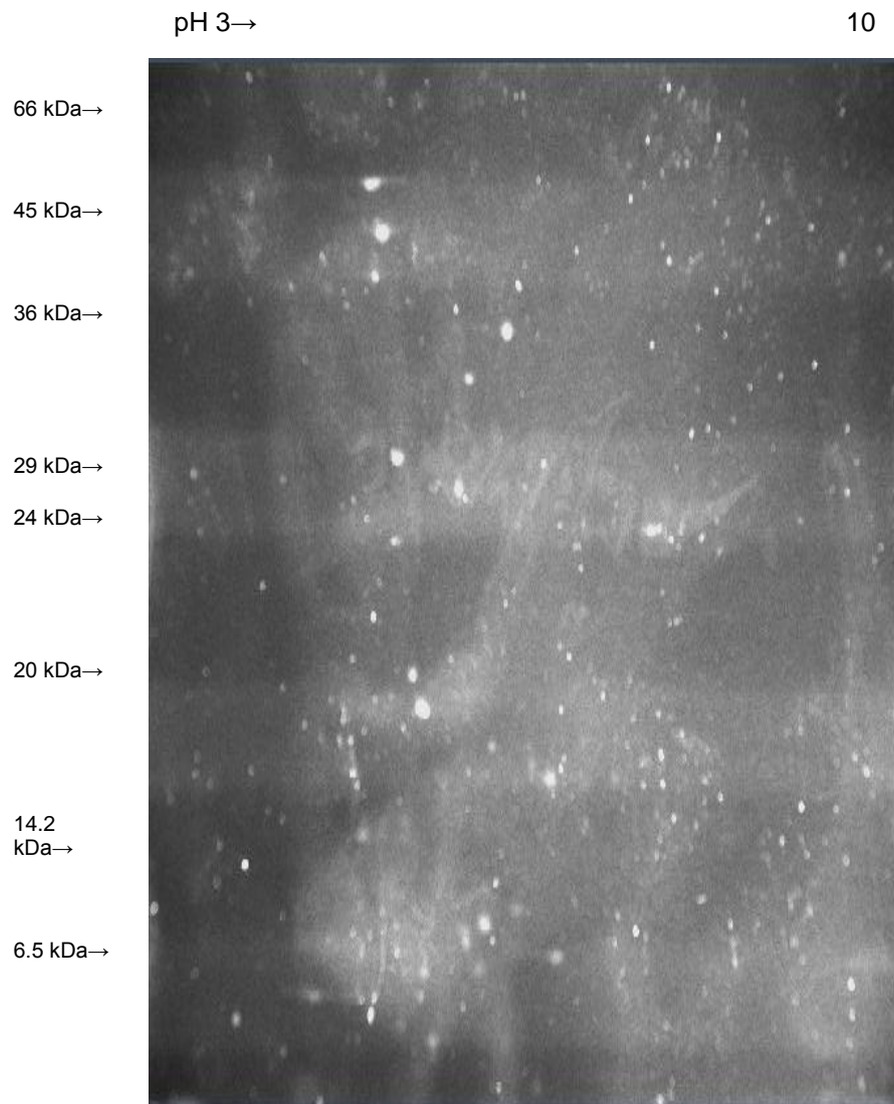


Figure 16.1: Proteome map of *E. cancerogenus* grown in semi-defined media without mucin. Separation was achieved using pH range 3-10 IEF strips. Gel was stained using fluorescent staining.

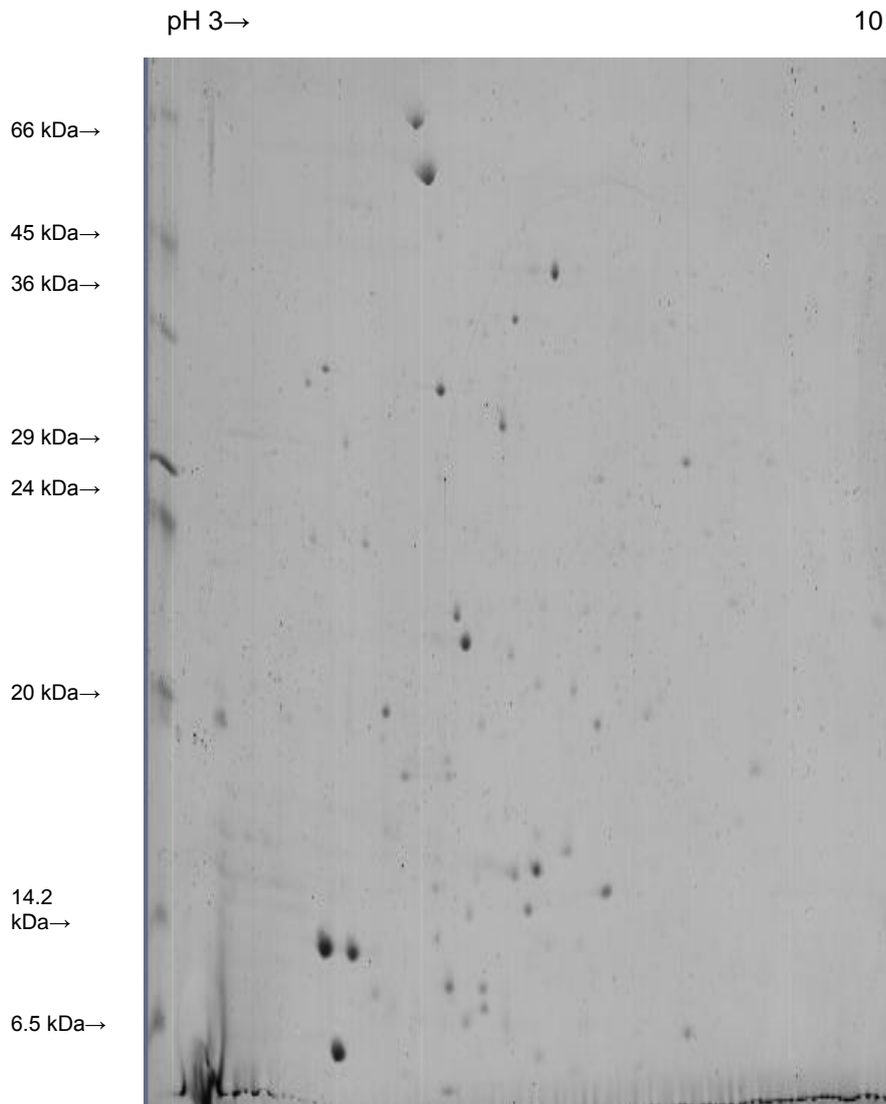


Figure 16.2: Proteome map of *E. cancerogenus* grown in semi-defined media without mucin. Separation was achieved using pH range 3-10 IEF strips. Gel was stained using colloidal Coomassie blue staining.

10.4.2 Comparison of pH ranges of IEF strips

The proteins from the culture supernatant were initially analysed using pH range 3-10 IEF strips. However the location of proteins seemed to be predominantly within the 4-7 pH range (Figures 17.1, 17.2 and 17.3) and hence the IPG strip pH range was changed to 4-7.

Figure 17: Isolation of *E. cancerogenus* proteins from 450 mL mid log phase cultures at an OD of 0.6 separated using pH range of 3-10 IEF strips

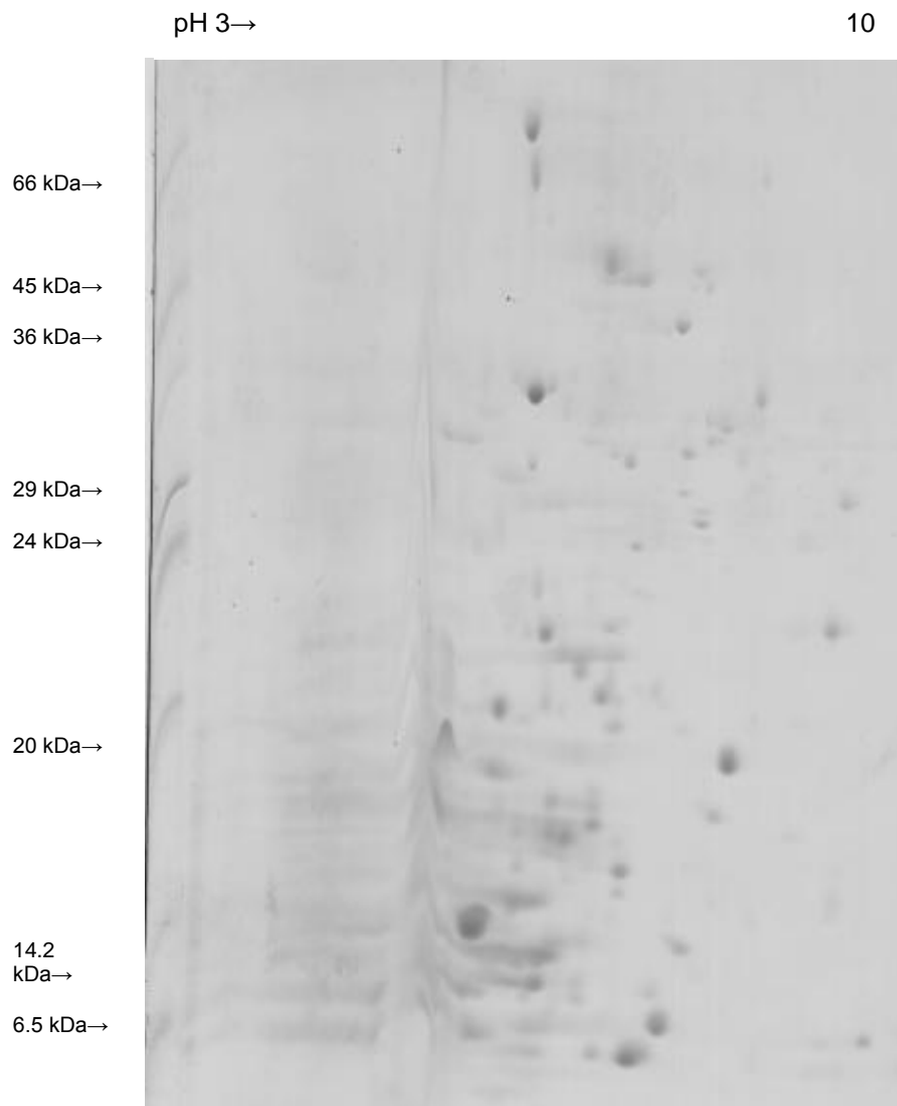


Figure 17.1: Proteome map of *E. cancerogenus* grown in semi-defined media without mucin. Separation was achieved using pH range 3-10 IEF strips.

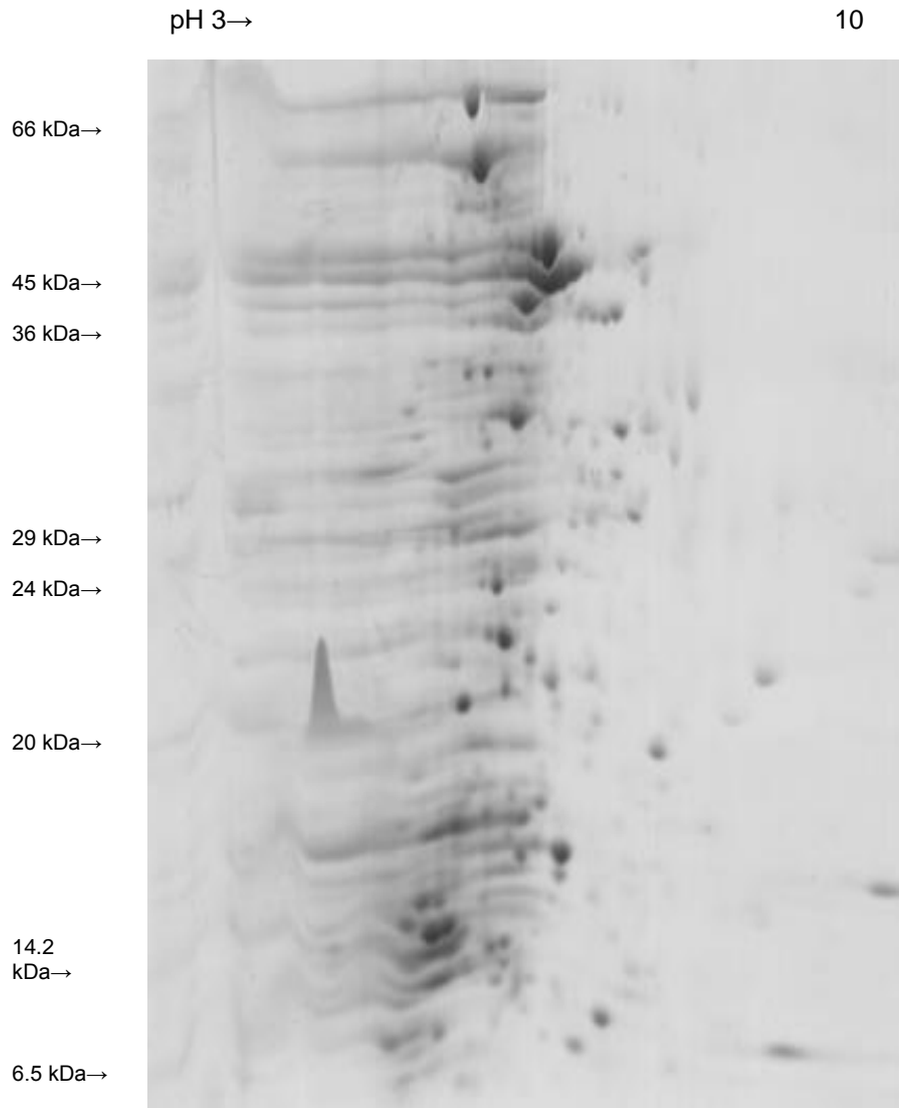


Figure 17.2: Proteome map of *E. cancerogenus* grown in semi-defined media enriched with mucin Type II. Separation was achieved using pH 3-10 IEF strips.

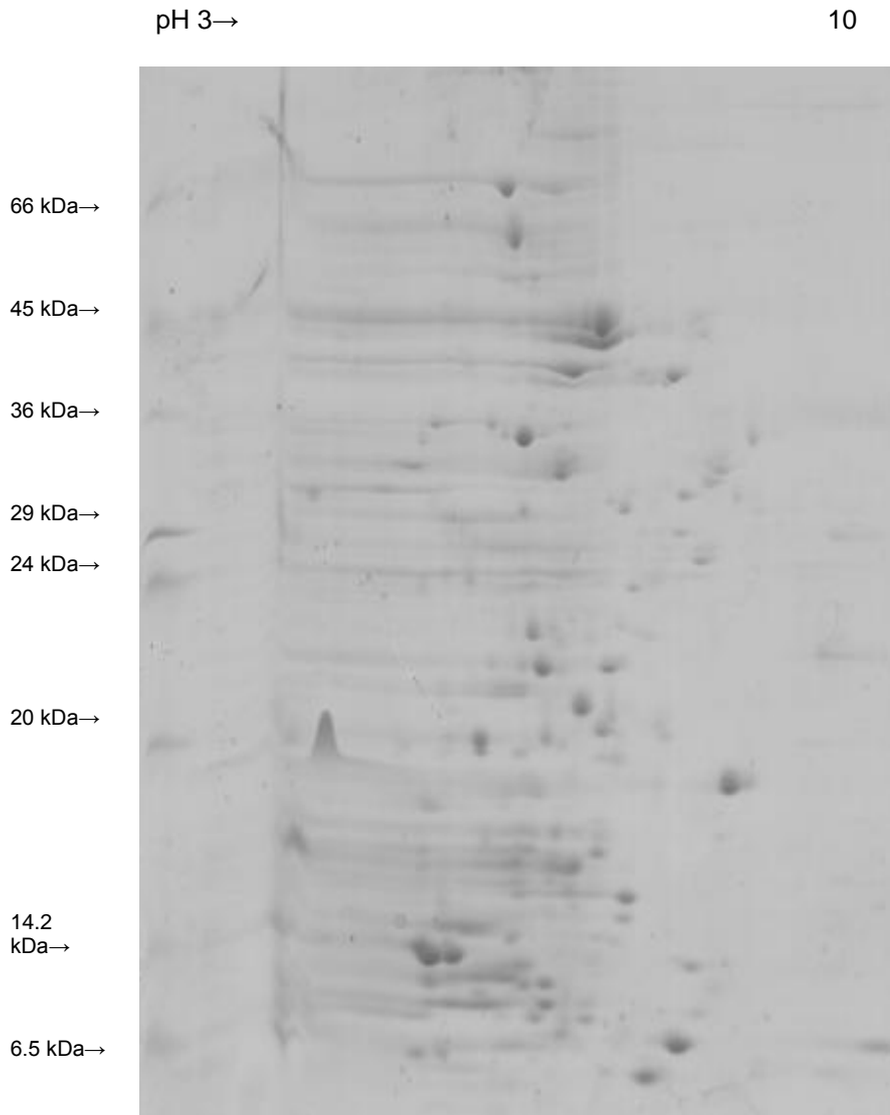


Figure 17.3: Proteome map of *E. cancerogenus* grown in semi-defined media enriched with mucin Type III. Separation was achieved using pH 3-10 IEF strips.

10.4.3 Comparison of protein loadings

The expression of proteins was studied in the mid logarithmic phase of growth of the culture and several optimisation experiments were performed to determine the appropriate volume of the culture that could provide a good separation and resolution of spots. Initial experiments were performed using 450 mL of the log phase cultures and were later reduced to a volume of 100 mL in order to obtain the best possible resolution (Figs. 18.1 and 18.2).

Figure 18. Proteins isolated from 100 and 450 mL of *B.fragilis* and *E. cancerogenus* grown in semi-defined media separated by pH 4-7 IEF strips

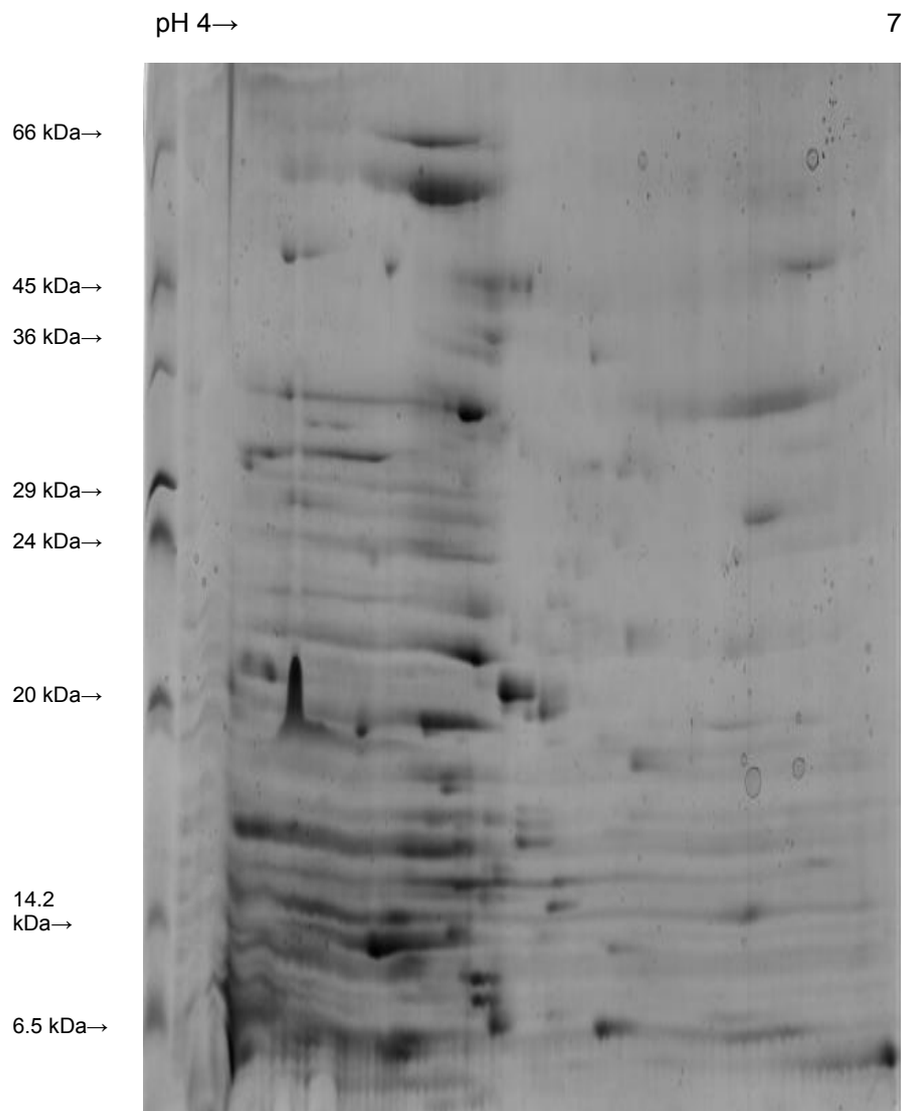


Figure 18.1: Proteins isolated from 450 mL of *E. cancerogenus* grown in semi-defined media without mucin separated by pH 4-7 IEF strips.

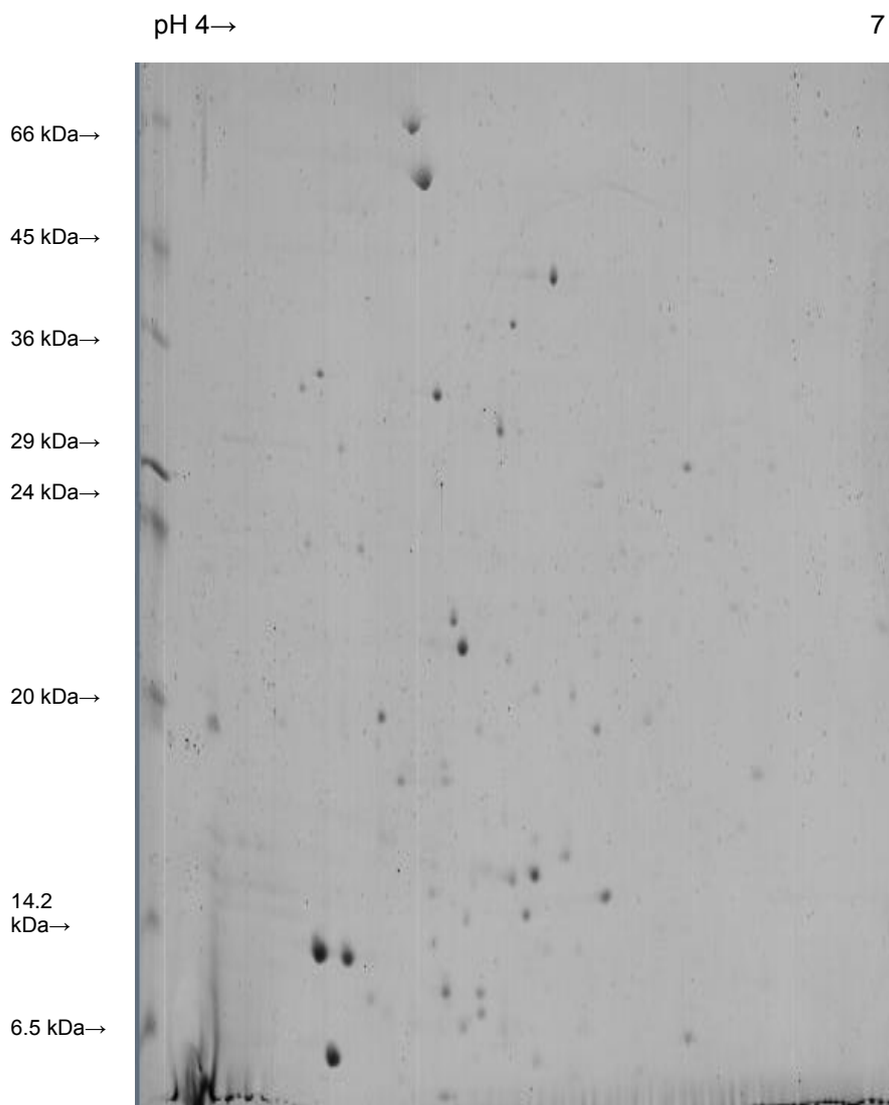


Figure 18.2: Proteins isolated from 100 mL of *E.cancerogenus* grown in semi-defined media without mucin separated by pH 4-7 IEF strips.

10.4.4 Comparison of polyacrylamide gel concentrations

The protein expression patterns were found to be more distinct and better resolved in 12 % (w/v) rather than 14% (w/v) gels and hence 12 % (w/v) gels were used during the study (Figures 19.1 and 19.2).

Figure 19: Isolation of *E. cancerogenus* proteins from 100 mL log phase cultures at an OD of 0.6 separated using pH range of 4-7 IEF strips

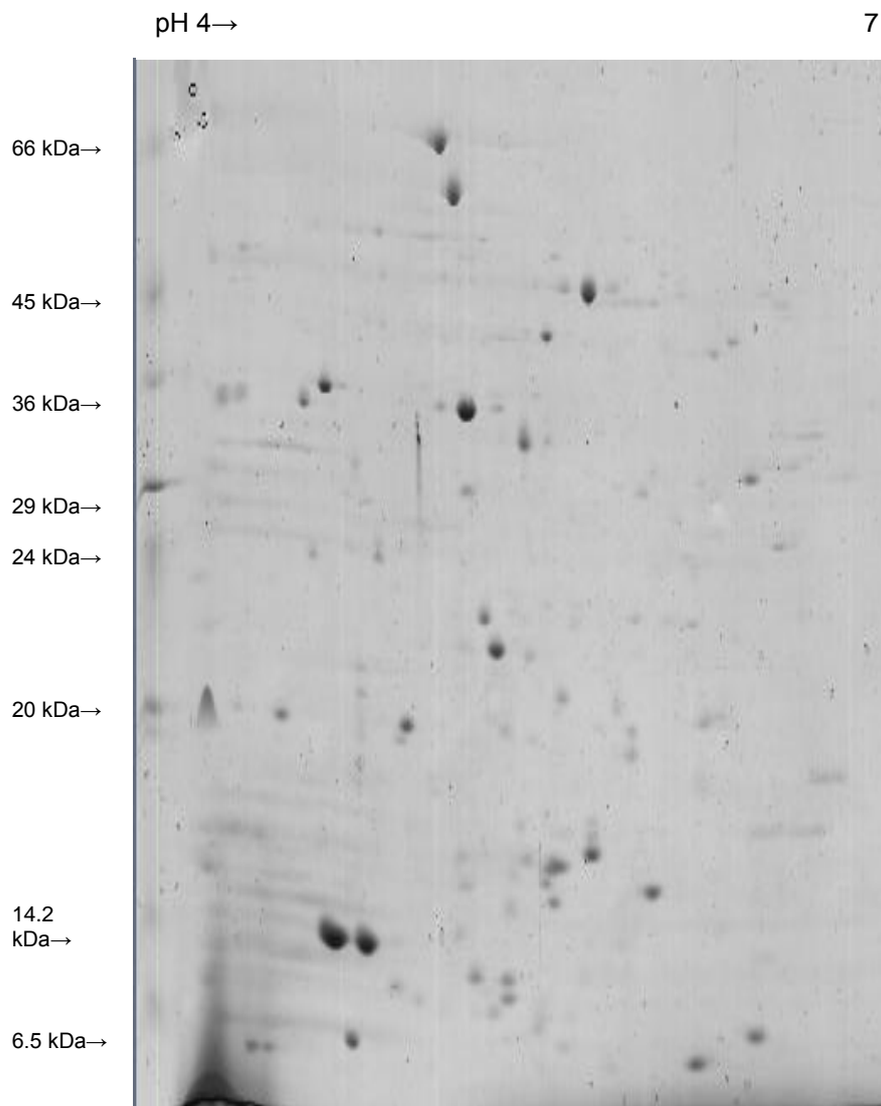


Figure 19.1: Proteome map of *E. cancerogenus* grown in semi-defined media without mucin. Separation was achieved using a 12% (w/v) gel

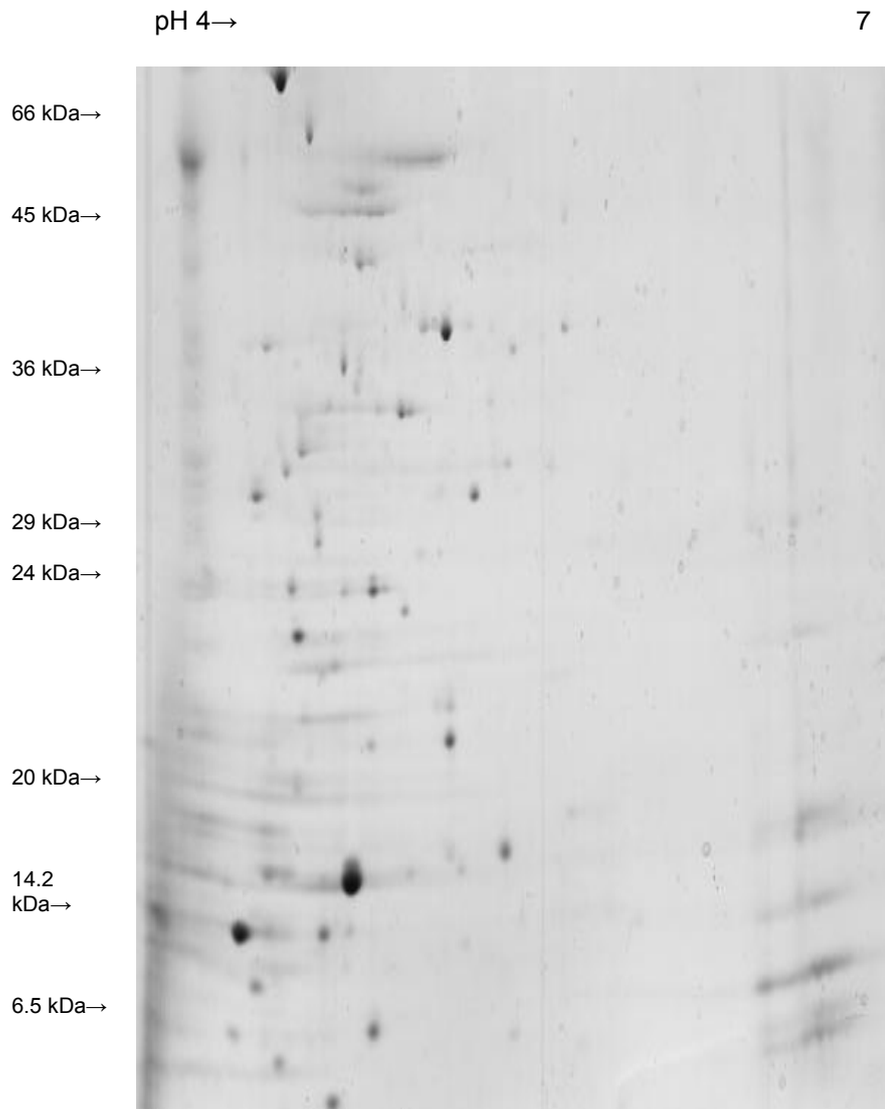


Figure 19.2: Proteome map of *E. cancerogenus* grown in semi-defined media without mucin. Separation was achieved using a 14% (w/v) gel

10.5 Two dimensional gel electrophoresis

One of the initial problems was the culture volume required to obtain a protein expression profile with good resolution and minimising horizontal streaks. Hence different volumes of cultures were used to obtain expression patterns in the log phase and a 100 mL culture volume at an OD_{600nm} of 0.6 or 0.7 in *E. cancerogenus* or *B. fragilis*, respectively, was found to be the most appropriate. Based on the results obtained, 100 mL culture volumes were used in the study (Refer to Figs. 18.1, 18.2 with different volumes in results section).

Sample preparation is one of the most important parts of the two dimensional gel electrophoresis (2DE) technique since even small variations in methodology could

result in massive changes in protein expression profiles. Reproducibility of gels was another major issue faced in the 2DE method. Care was taken to ensure that proteins studied from different conditions were extracted and analysed in a similar fashion aiming at minimising variation as much as possible (Choe & Lee, 2003). At least six gels were generated from each condition in order to confirm that they were reproducible. During Mann-Whitney U analysis using the PDQuest and Redfin softwares, spots that were not up- or down-regulated in all the replicate gels compared to all the replicated control gels were excluded. During Student t-test analysis using the PDQuest, Redfin and SameSpots software, an average expression in each condition was considered rather than the expression levels in each replicate gel.

Initial experiments resulted in a high amount of horizontal and vertical streaks on the gels with poor resolution. One of the main reasons for this was the incomplete resolubilisation of pellets following their treatment with the 2D Clean up kit. The protein pellets were resolubilised in the rehydration buffer by ultra-sonication on ice for a few seconds and the insoluble pellets were removed by centrifugation for 1 min. This helped to reduce protein overload on to the IPG strips and also resulted in efficient separation of proteins.

The composition of the solutions used in the 2D Clean up kit are not freely available. Previous studies have shown that they help to remove contaminating polysaccharides from the protein samples (Zhang, 2006) (Zhang *et al.*, 2007). The 2D Clean up kit consists of the precipitant, co-precipitant, wash buffer and wash additive solutions whose recipe has been patented. Alternatively, the acetone precipitation method could be used to prepare protein samples from *B. fragilis* cultures where acetone is the main compound precipitating out the proteins present in the cell free extract.

Similar studies were performed to determine the pH range for the IPG strips used in the first dimension electrophoresis. The pH range of 3-10 was used initially but since most of the proteome expression was concentrated around the acidic pH range of 4-7, further experiments were performed using IPG strips of pH 4-7 (refer to Figs. 17).

An attempt was used to compare the Coomassie blue staining with fluorescent staining but due to the inconvenience of cutting out spots and improper resolution,

the Coomassie blue staining technique was chosen to analyse the spots of interest. The inconvenience of cutting out spots was that UV light was required to locate the exact spot boundaries and this would make the cutting process a tedious task and visualisation in visible light impossible. From Figs. 16.1 and 16.2, it is evident that the spots have been expressed with better clarity and resolution when colloidal Coomassie blue stain was used. The increased convenience of using colloidal Coomassie blue staining method when compared to fluorescent staining supported its use in the study.

Since reproducibility and uniform dimensions of scan area on gels were some of the most important factors to be considered, the scanning of all the gels were performed together at the same time ensuring that the scan area was exactly the same for all of them. Further adjustments were possible using the PDQuest, Redfin and Samespots software tools where flipping of gels, cropping and background subtractions were carried out to optimise and compensate for differences in intensities of staining and other experimental variations. The spots on the gels were reviewed using the spot review tool and the consensus tool was used to edit any errors that may have occurred during spot analysis.

10.6 Separation of proteins from *E. cancerogenus* and *B. fragilis* grown in semi-defined media with and without mucin using 2DE gels.

10.6.1 Sets of replicate 2DE gels of proteins from *E. cancerogenus*

Multiple sets of gels of biological replicates of proteins from *E. cancerogenus* grown in media without mucin, media with mucin Type II and media with mucin Type III (IEF strip pH range 4-7, processing volume: 100 mL, harvesting OD at 600 nm: 0.6) (Figs. 20-22).

Figure 20: Proteome maps of *E. cancerogenus* grown in semi-defined media without mucin (control media; using pH 4-7 IEF strips).

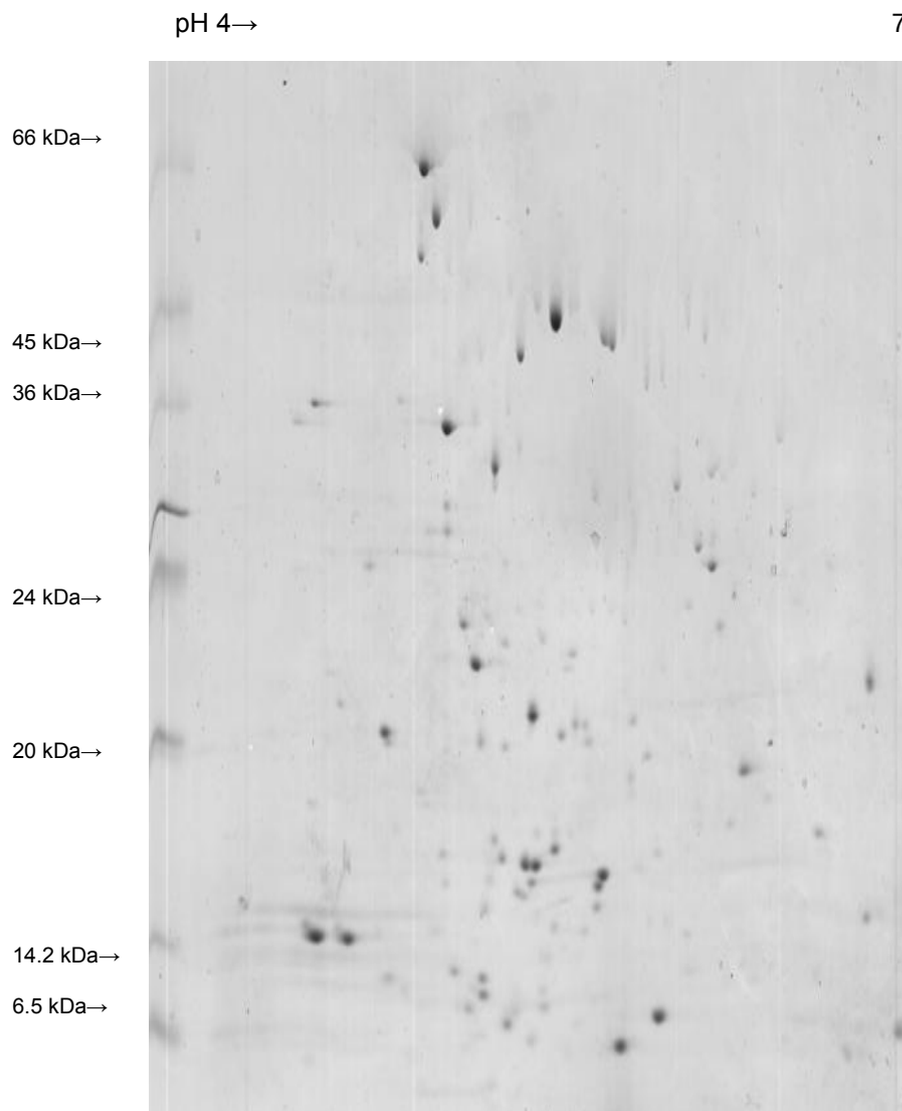


Figure 20.1: Proteome map no.1 of *E. cancerogenus* grown in semi-defined media without mucin

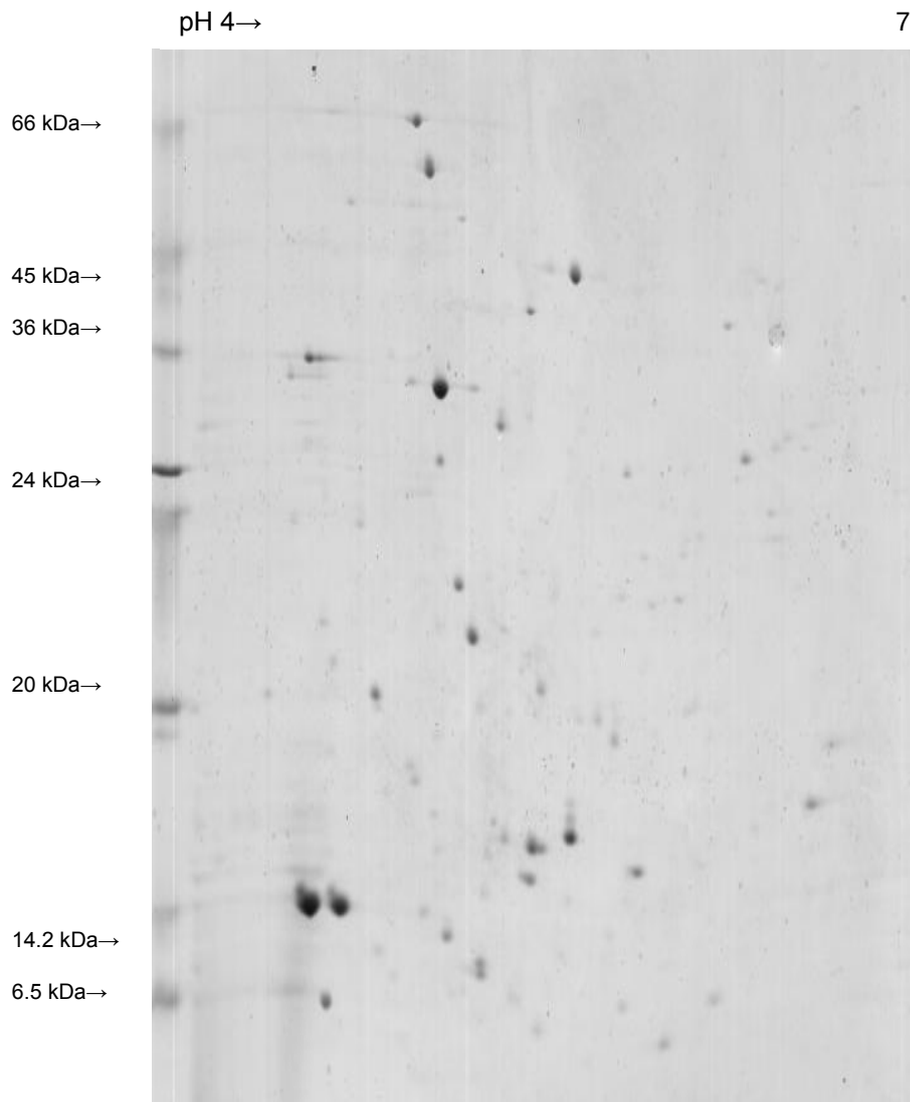


Figure 20.2: Proteome map no.2 of *E. cancerogenus* grown in semi-defined media without mucin.

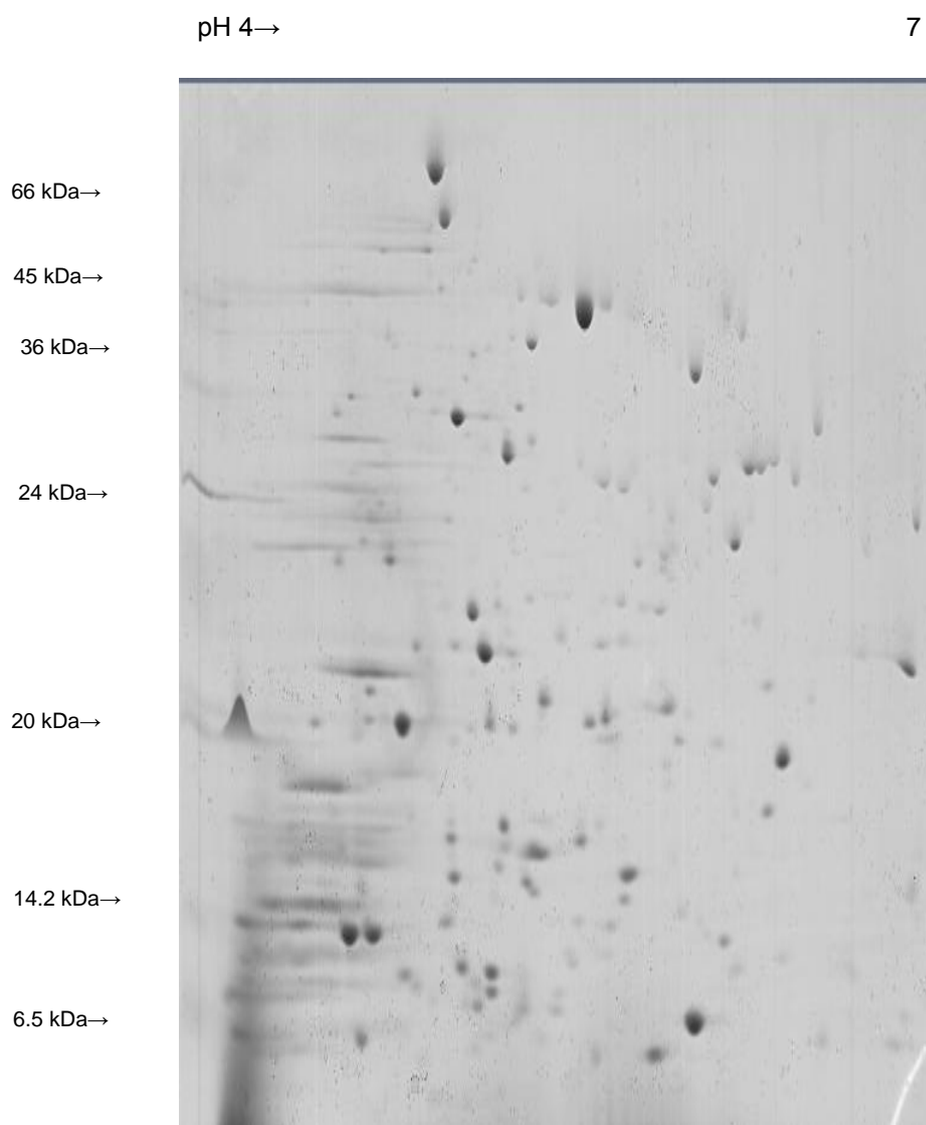


Figure 20.3: Proteome map no.3 of *E. cancerogenus* grown in semi-defined media without mucin.

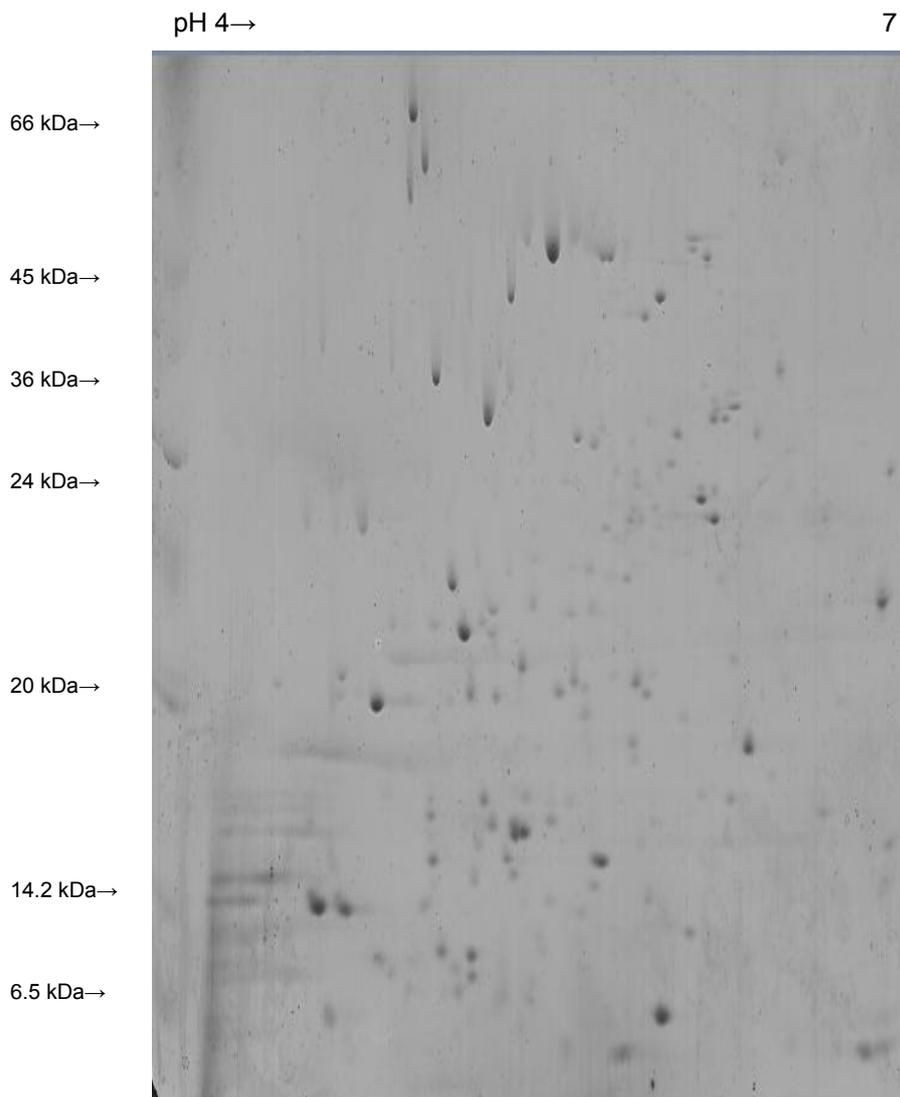


Figure 20.4: Proteome map no.4 of *E. cancerogenus* grown in semi-defined media without mucin.

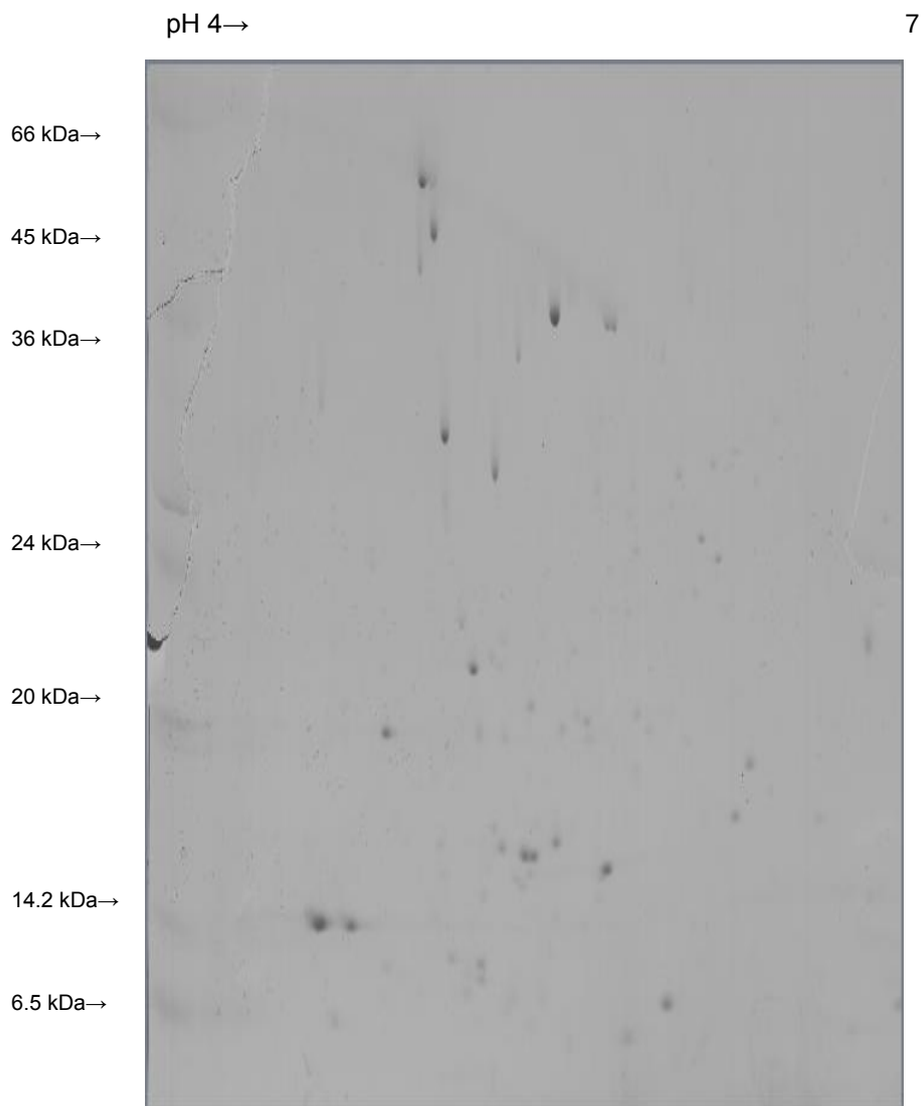


Figure 20.5: Proteome map no.5 of *E. cancerogenus* grown in semi-defined media without mucin.

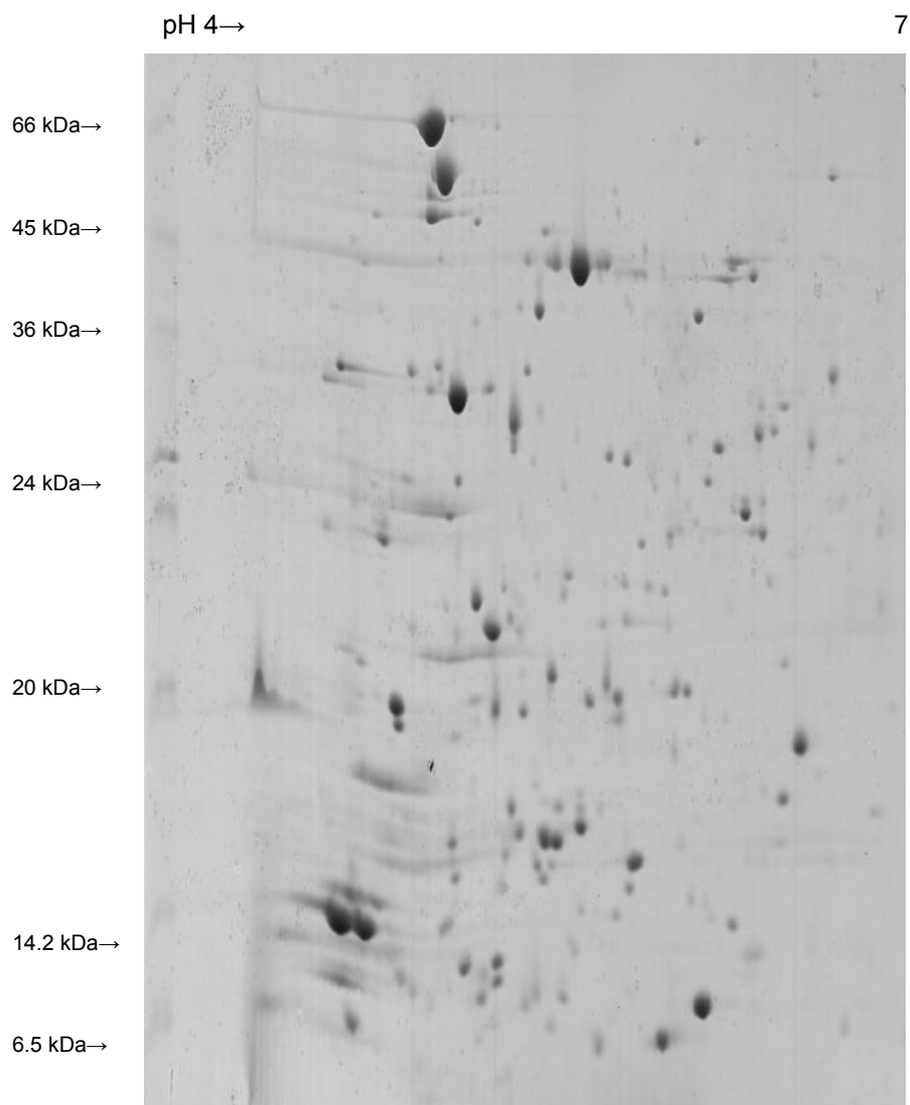


Figure 20.6: Proteome map no.6 of *E. cancerogenus* grown in semi-defined media without mucin.

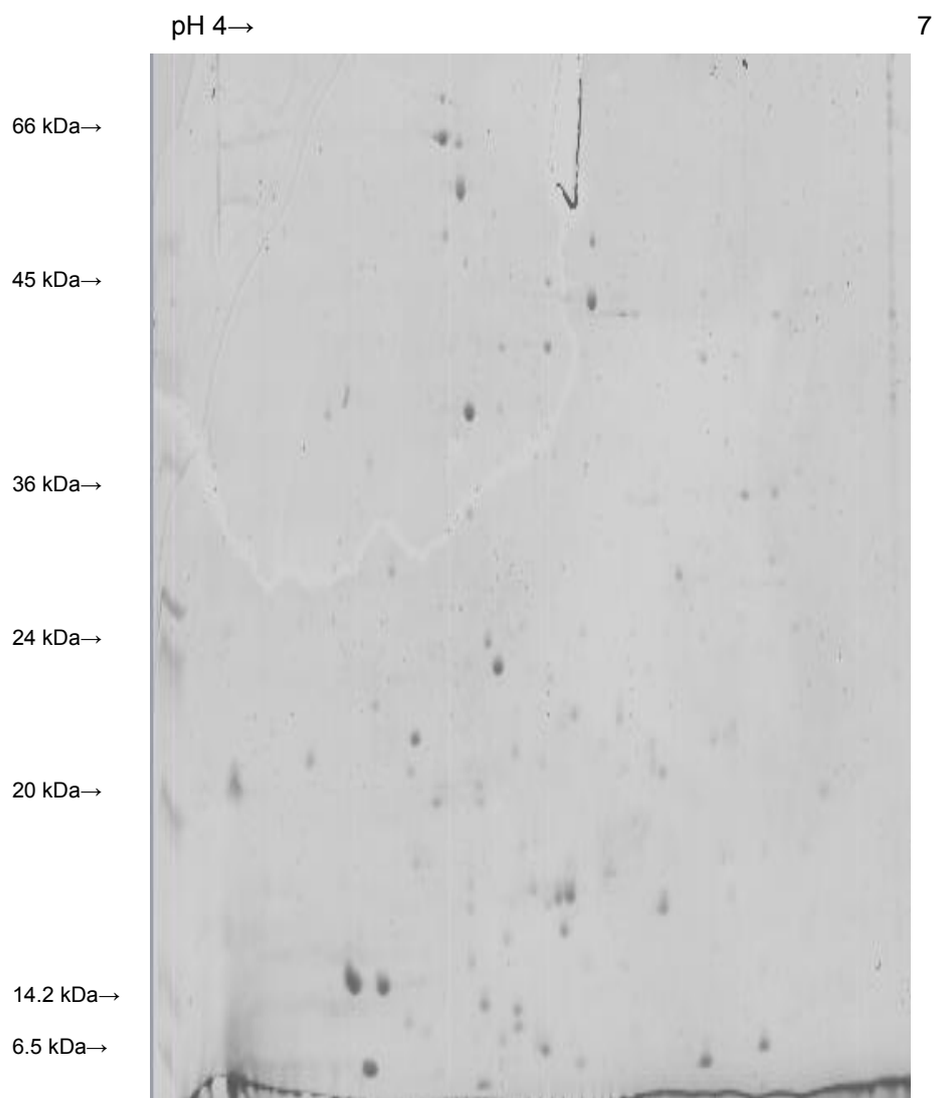


Figure 20.7: Proteome map no.7 of *E. cancerogenus* grown in semi-defined media without mucin.

Figure 21: Proteome maps of *E. cancerogenus* grown in semi-defined media enriched with mucin Type II (using pH 4-7 IEF strips)

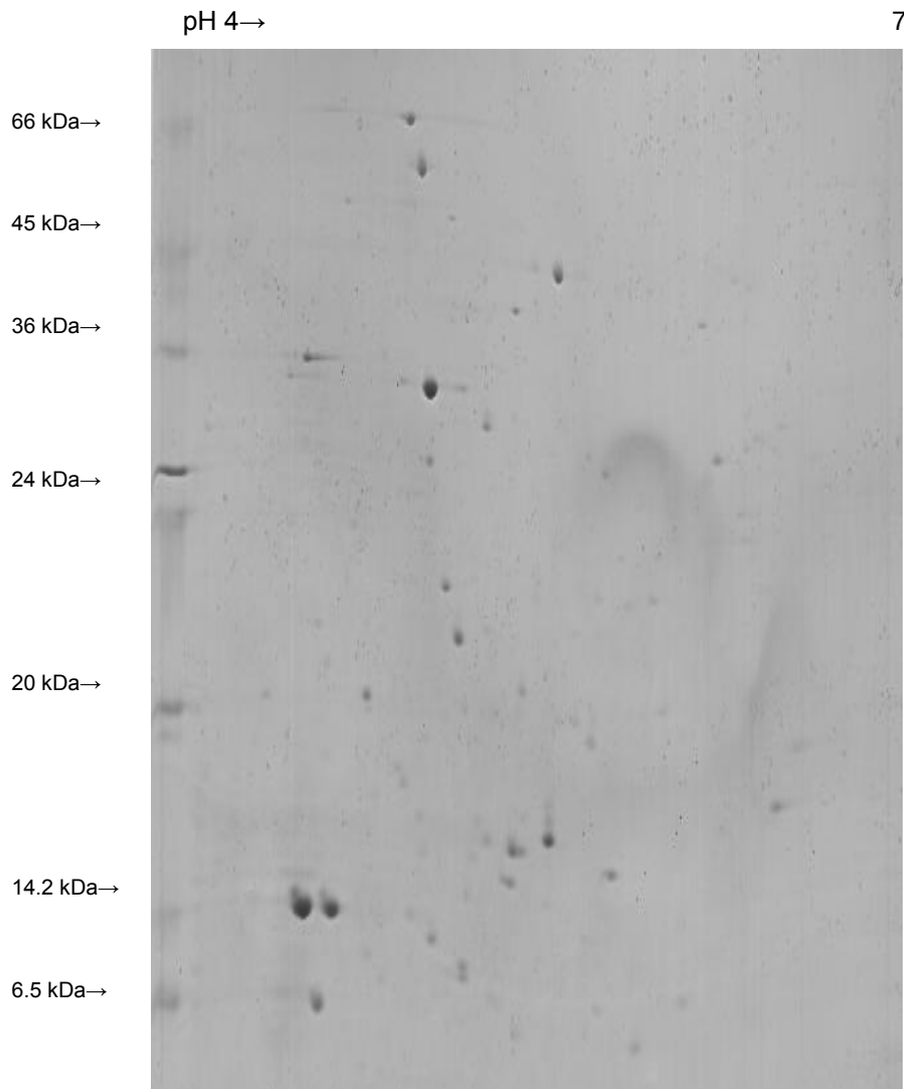


Figure 21.1: Proteome map no.1 of *E. cancerogenus* grown in mucin Type II enriched media

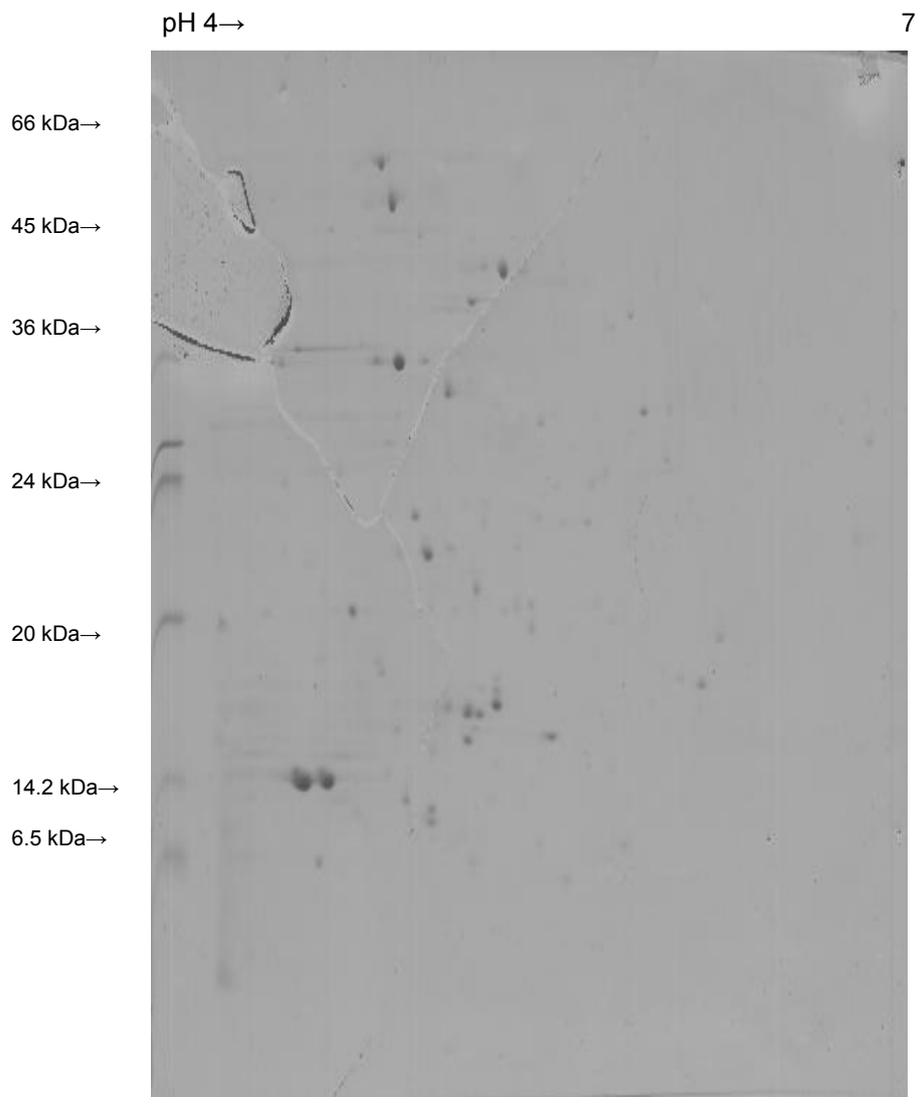


Figure 21.2: Proteome map no.2 of *E. cancerogenus* grown in mucin Type II enriched media

pH 4→

7

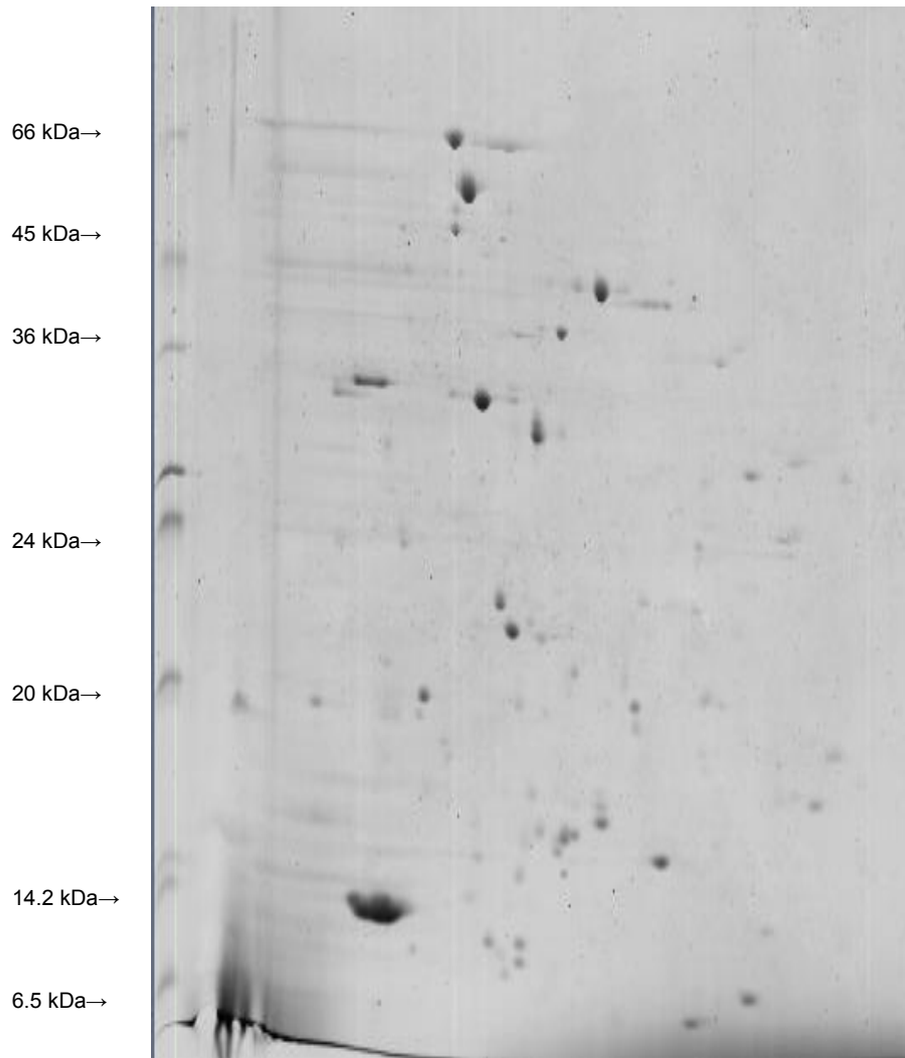


Figure 21.3: Proteome map no.3 of *E. cancerogenus* grown in mucin Type II enriched media

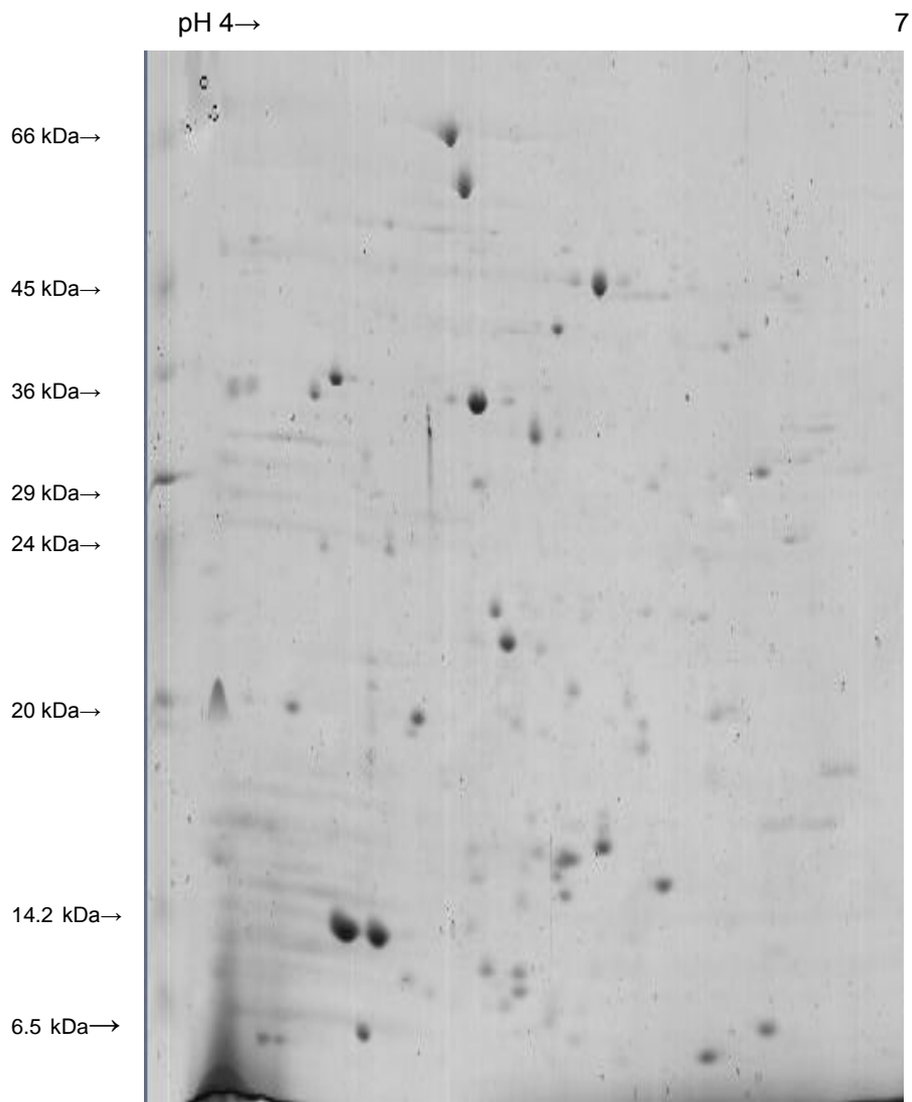


Figure 21.4: Proteome map no.4 of *E. cancerogenus* grown in mucin Type II enriched media

pH 4→

7

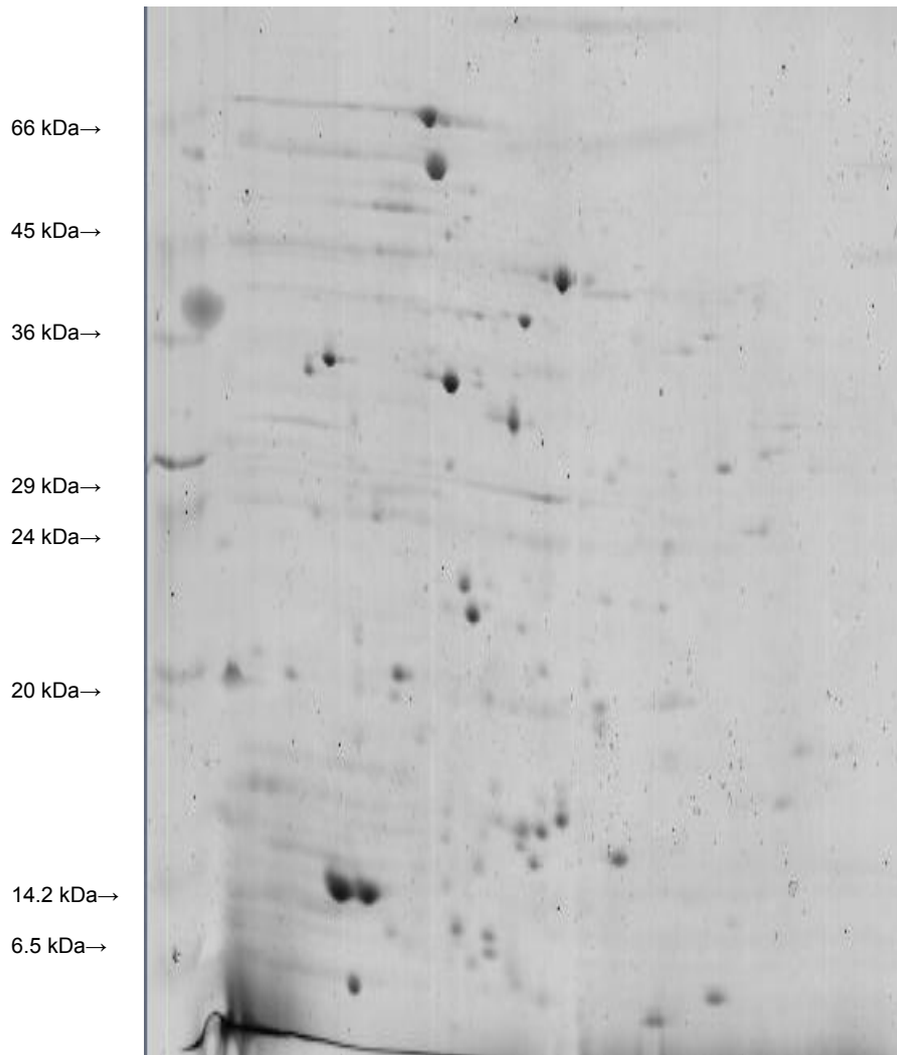


Figure 21.5: Proteome map no.5 of *E. cancerogenus* grown in mucin Type II enriched media

pH 4→

7

66 kDa→

45 kDa→

36 kDa→

29 kDa→

24 kDa→

20 kDa→

14.2 kDa→

6.5 kDa→

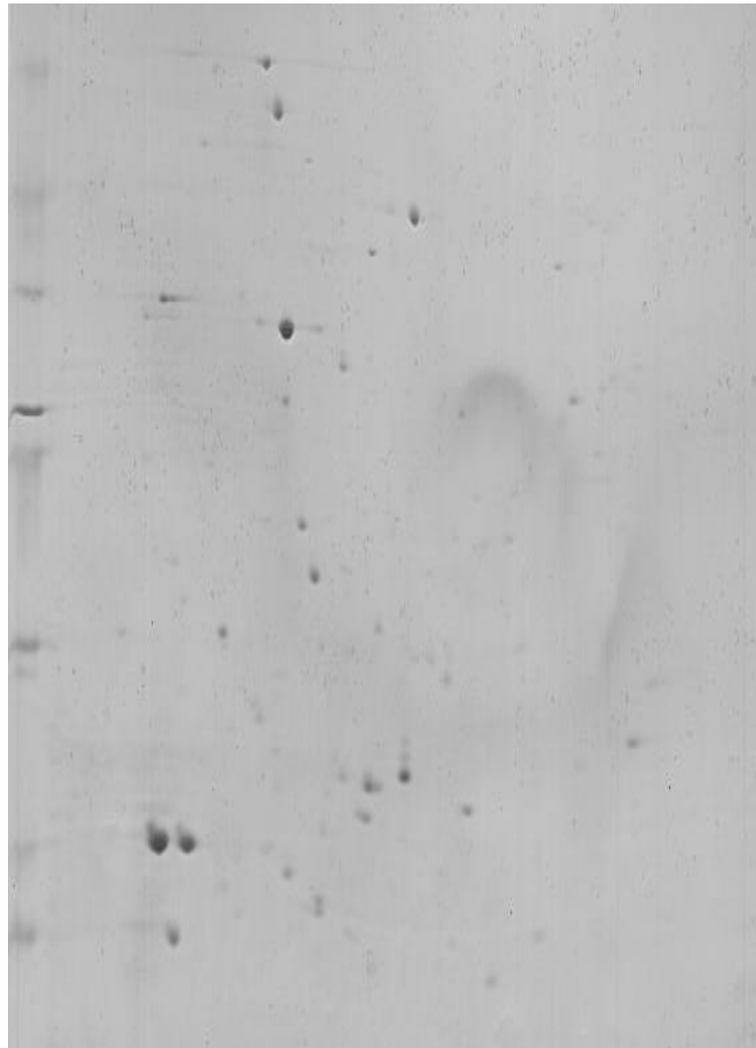


Figure 21.6: Proteome map no.6 of *E. cancerogenus* grown in mucin Type II enriched media

Figure 22: Proteome maps of *E. cancerogenus* grown in semi-defined media enriched with mucin Type III (using pH 4-7 IEF strips)

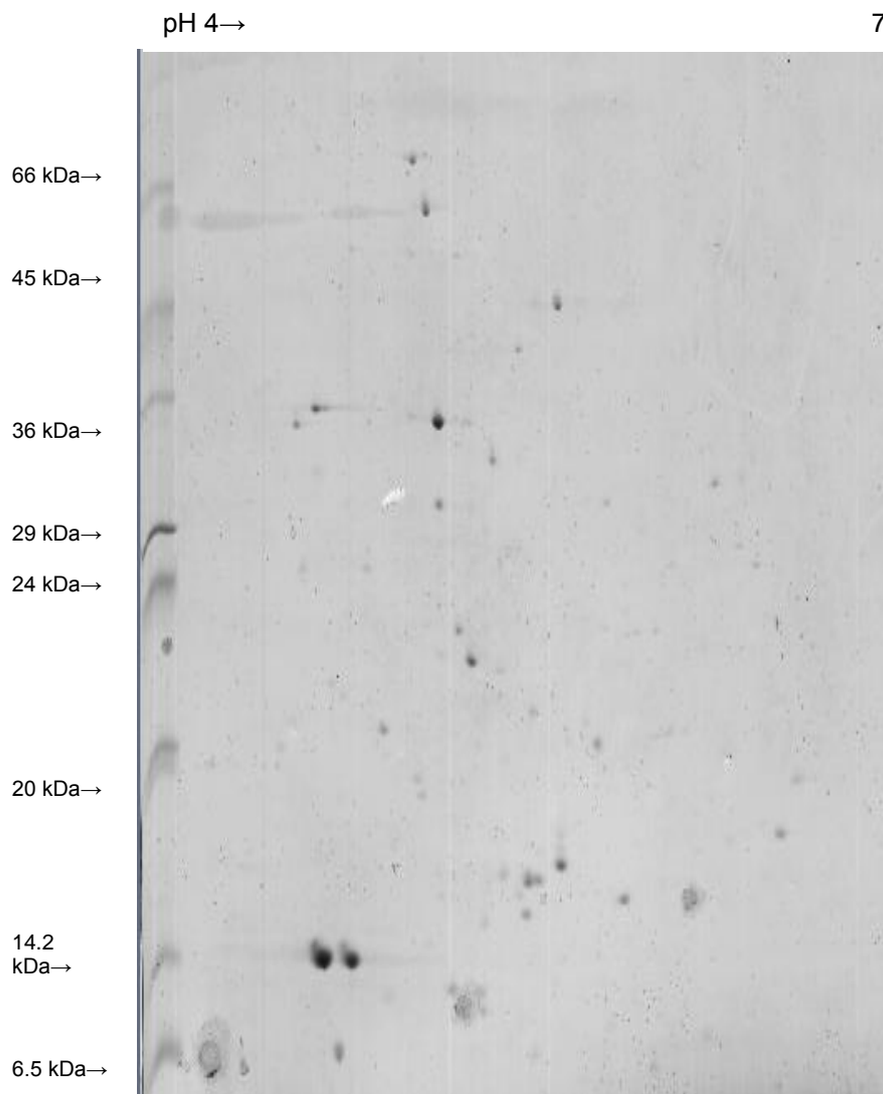


Figure 22.1: Proteome map no.1 of *E. cancerogenus* grown in mucin Type III enriched media

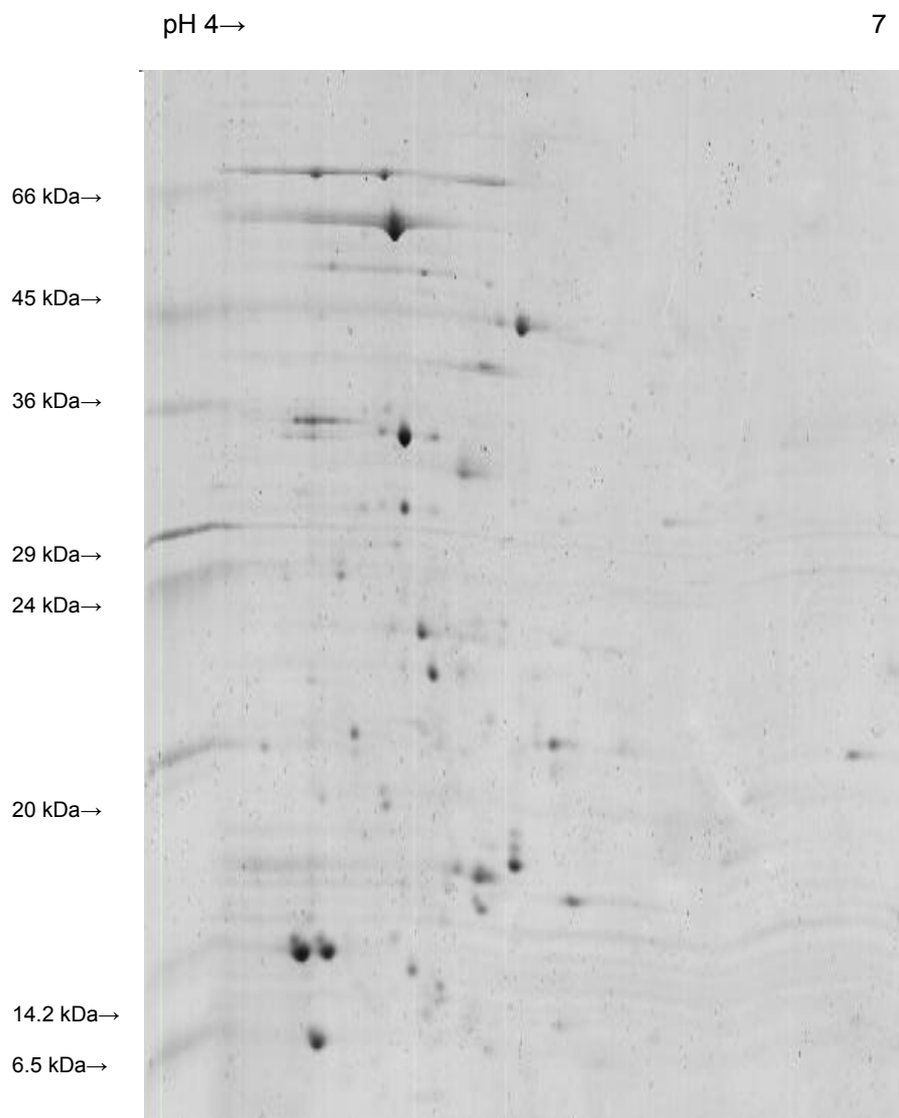


Figure 22.2: Proteome map no.2 of *E. cancerogenus* grown in mucin Type III enriched media

pH 4→

7

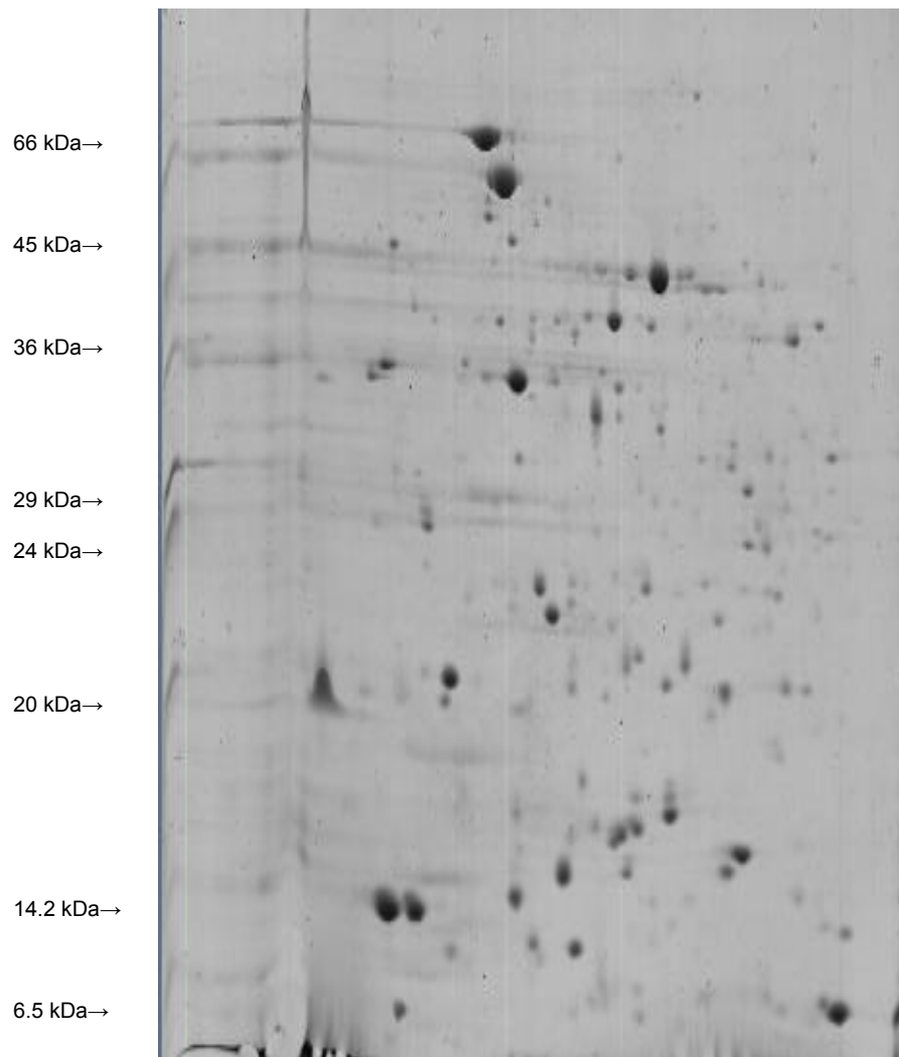


Figure 22.3: Proteome map no.3 of *E. cancerogenus* grown in mucin Type III enriched media

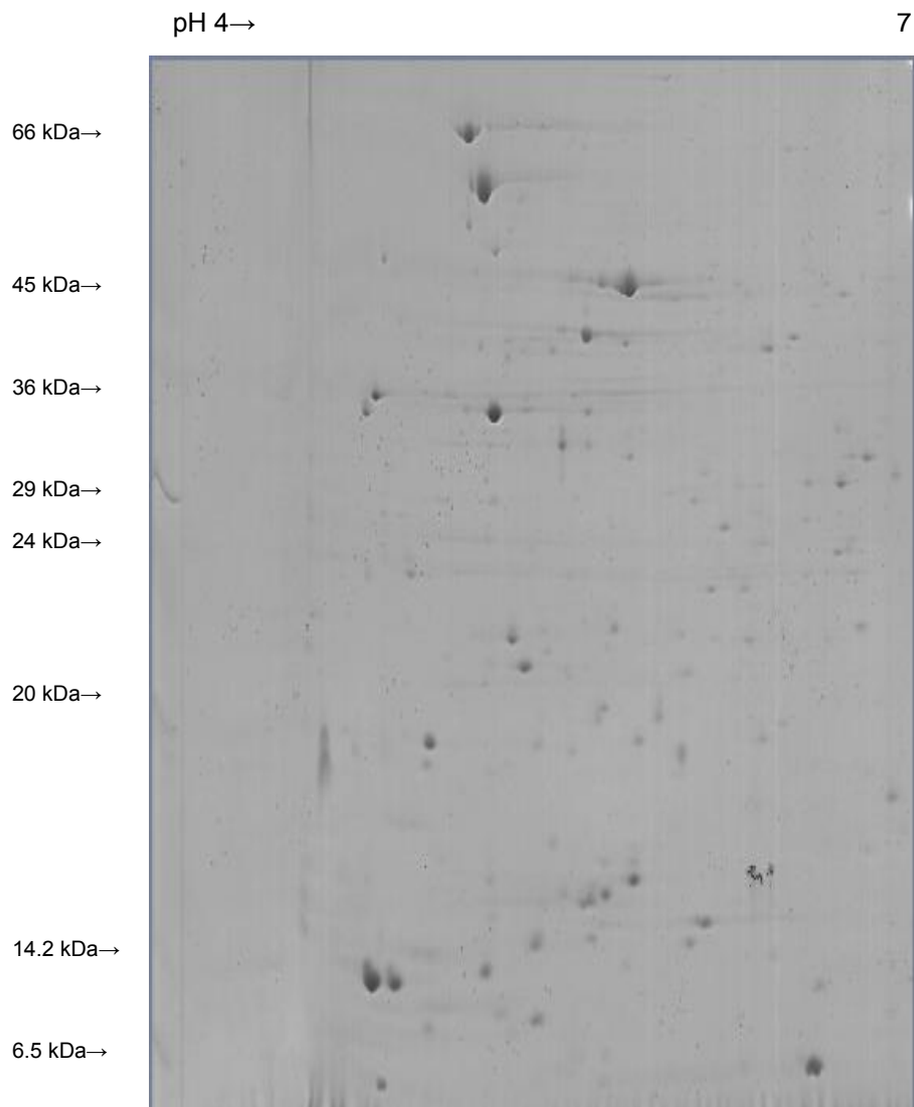


Figure 22.4: Proteome map no.4 of *E. cancerogenus* grown in mucin Type III enriched media

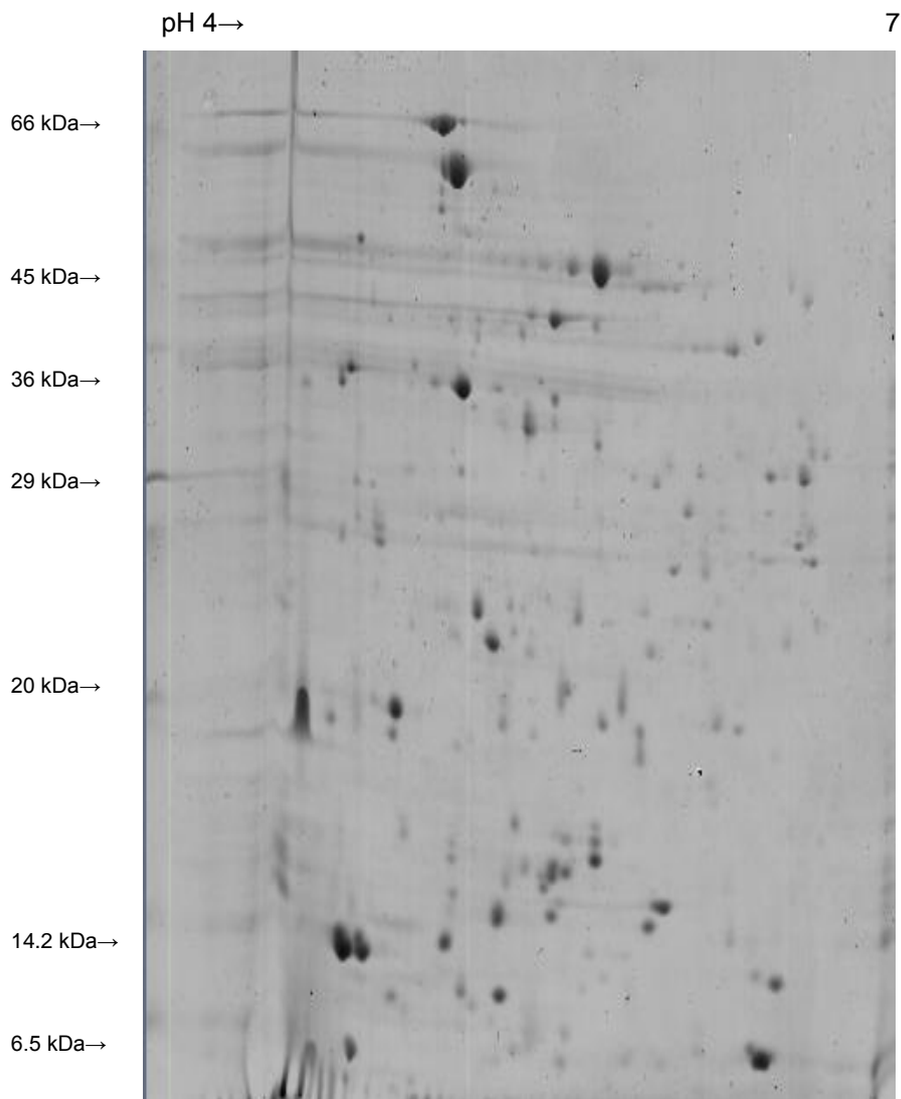


Figure 22.5: Proteome map no.5 of *E. cancerogenus* grown in mucin Type III enriched media

pH 4→

7

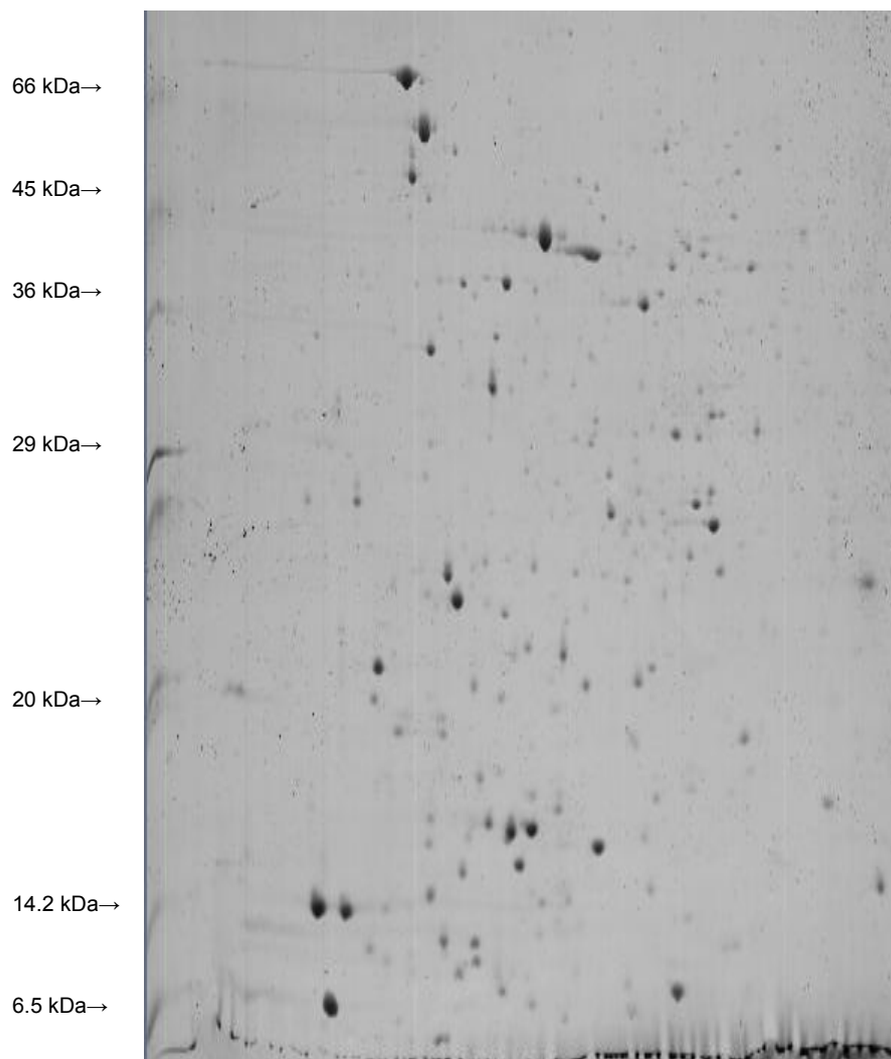


Figure 22.6: Proteome map no.6 of *E. cancerogenus* grown in mucin Type III enriched media

pH 4→

7

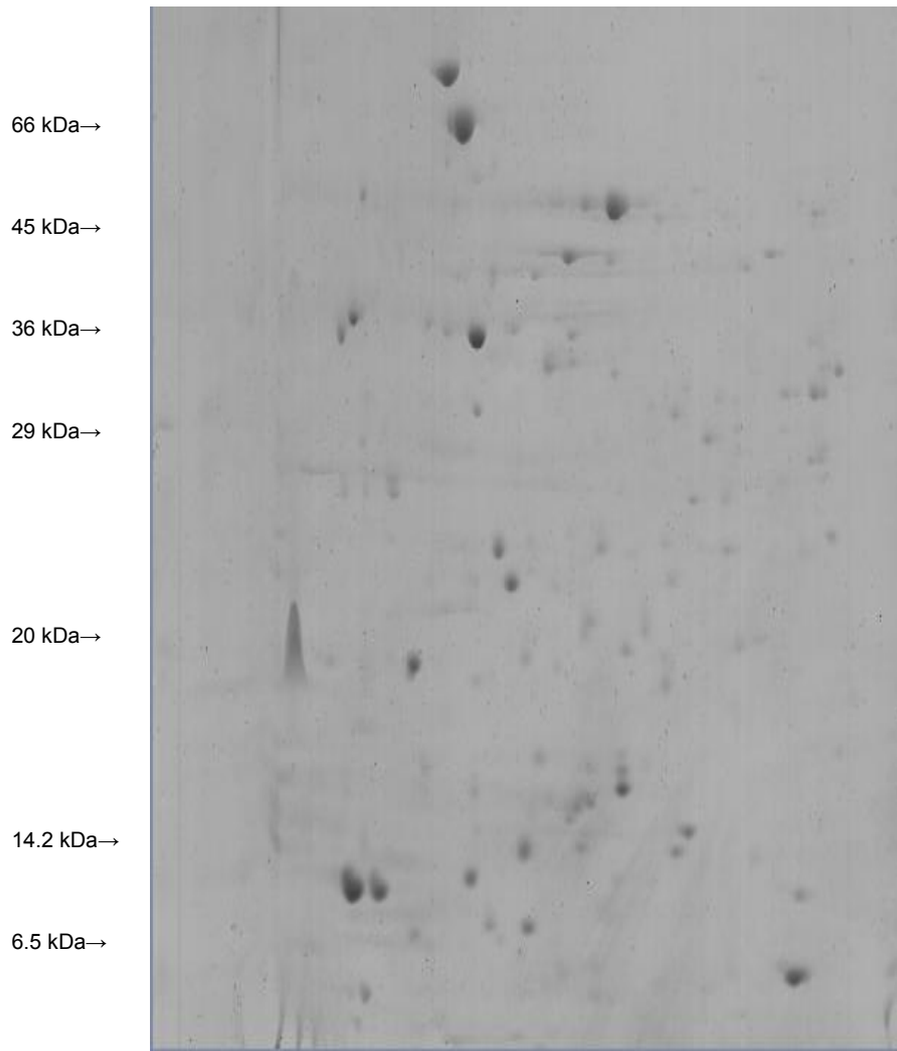


Figure 22.7: Proteome map no.7 of *E. cancerogenus* grown in mucin Type III enriched media

pH 4→

7

66 kDa→

45 kDa→

36 kDa→

29 kDa→

24 kDa→

20 kDa→

14.2 kDa→

6.5 kDa→



Figure 22.8: Proteome map no.8 of *E. cancerogenus* grown in mucin Type III enriched media

10.6.2 Sets of replicate 2DE gels of proteins from *B. fragilis*

Multiple sets of gels of proteins from *B. fragilis* grown in media without mucin, media with mucin Type II and media with mucin Type III (IEF strip pH range 4-7, processing volume: 100 mL, harvesting OD at 600 nm: 0.7) (Figs. 23-25).

Figure 23: Proteome maps of *B. fragilis* grown in semi-defined media without mucin (control media; using pH 4-7 IEF strips)

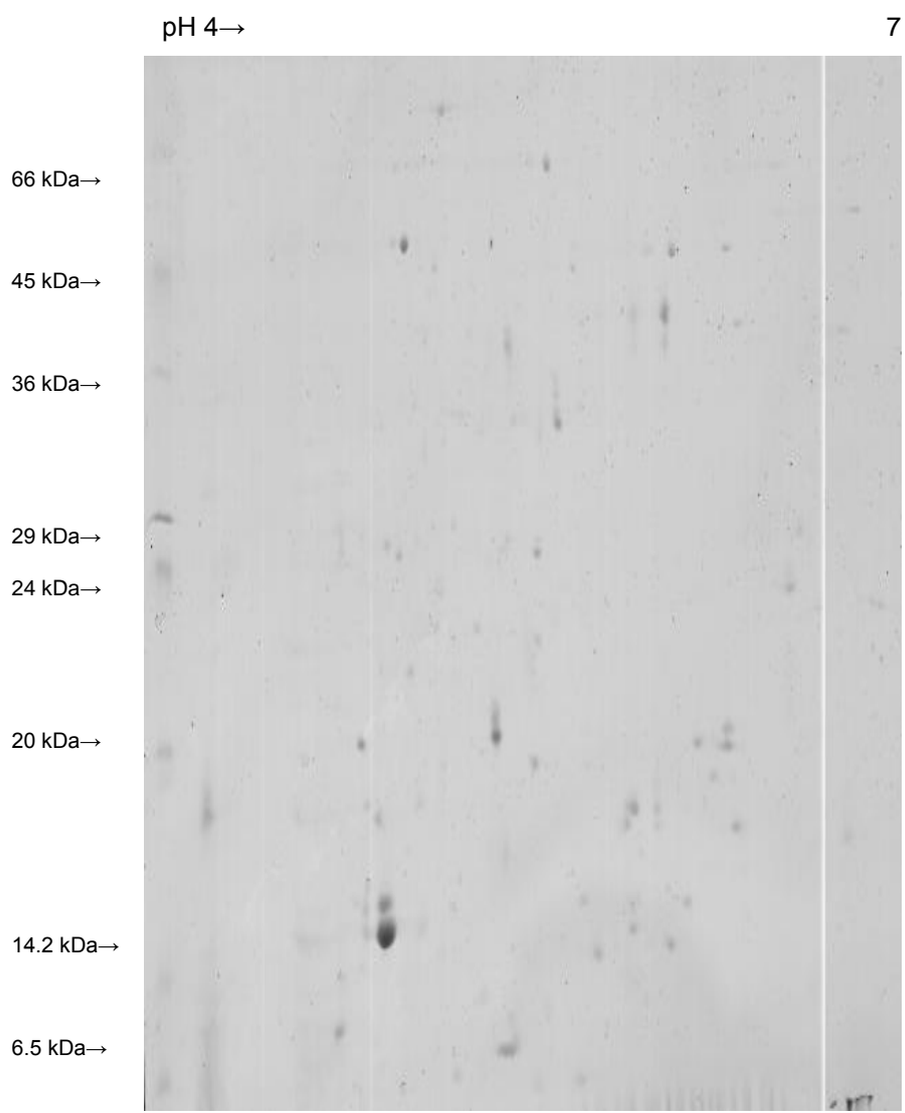


Figure 23.1 Proteome map no.1 of *B. fragilis* grown in semi-defined media without mucin

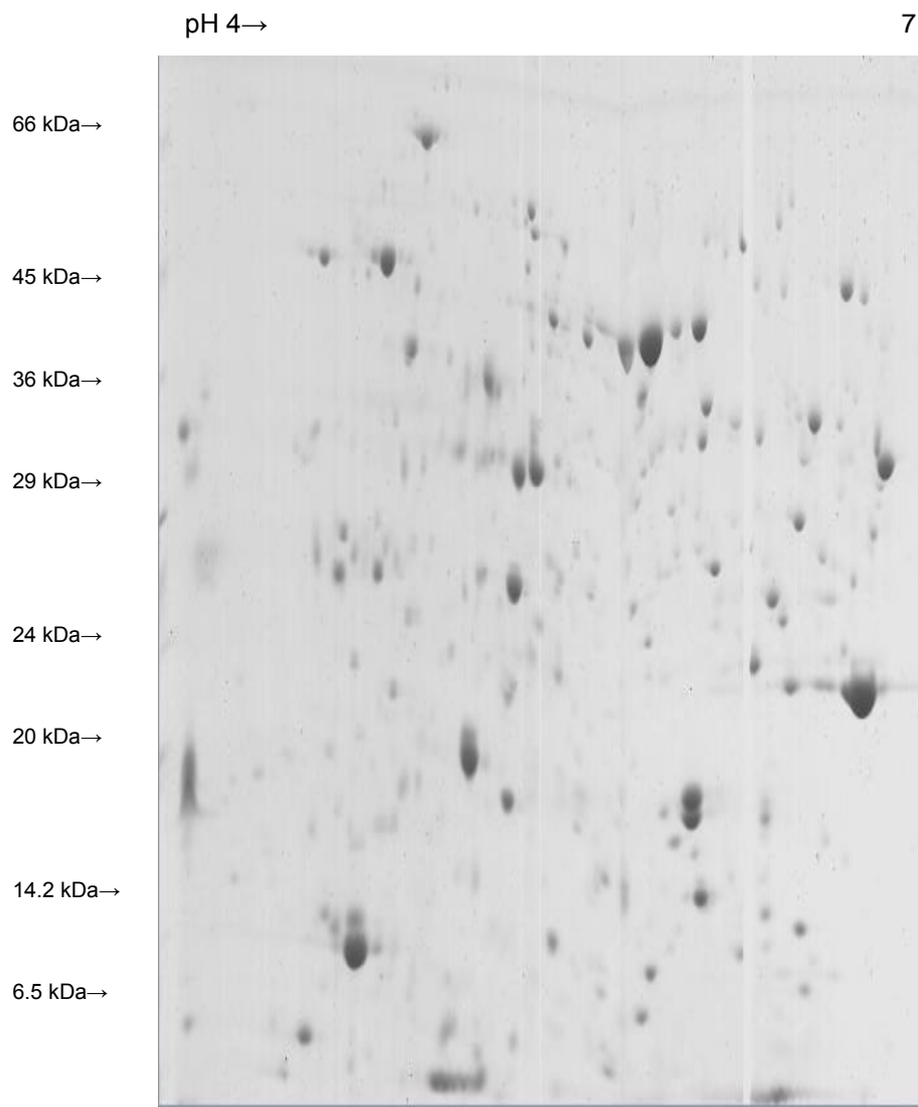


Figure 23.2: Proteome map no.2 of *B. fragilis* grown in semi-defined media without mucin

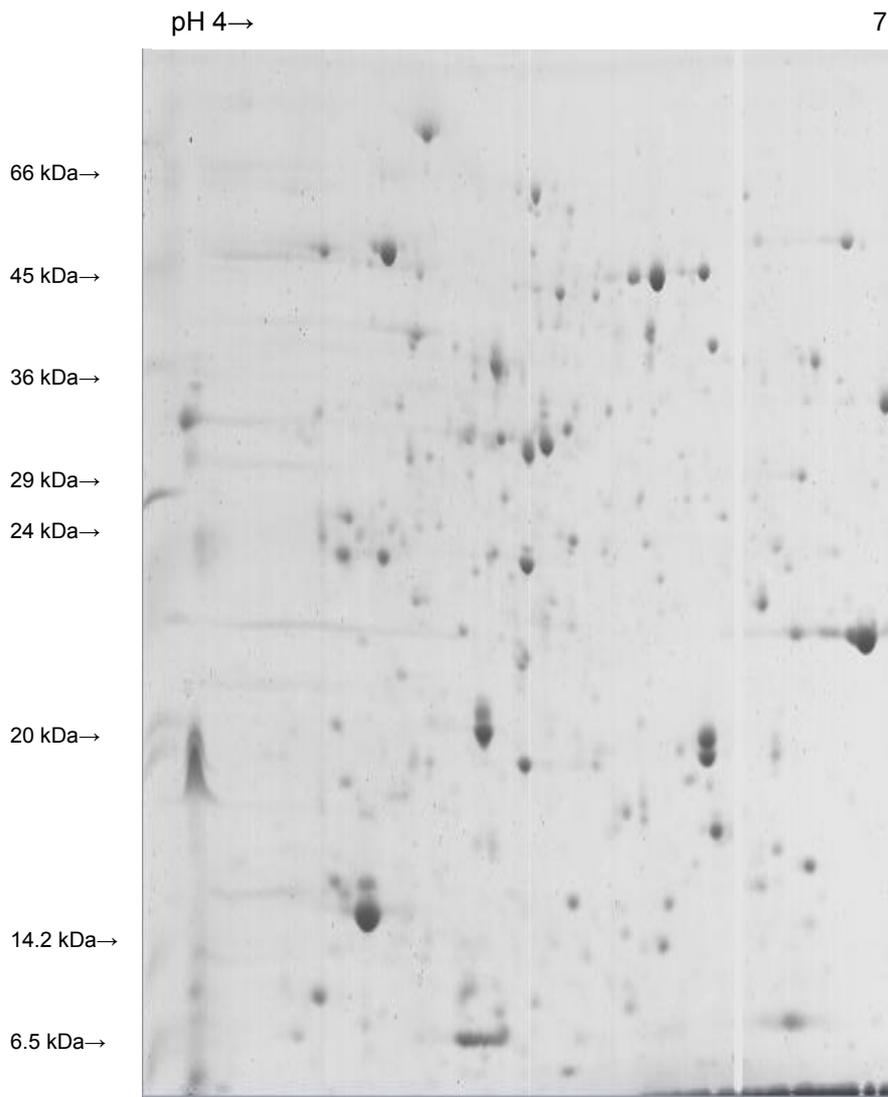


Figure 23.3: Proteome map no.3 of *B. fragilis* grown in semi-defined media without mucin

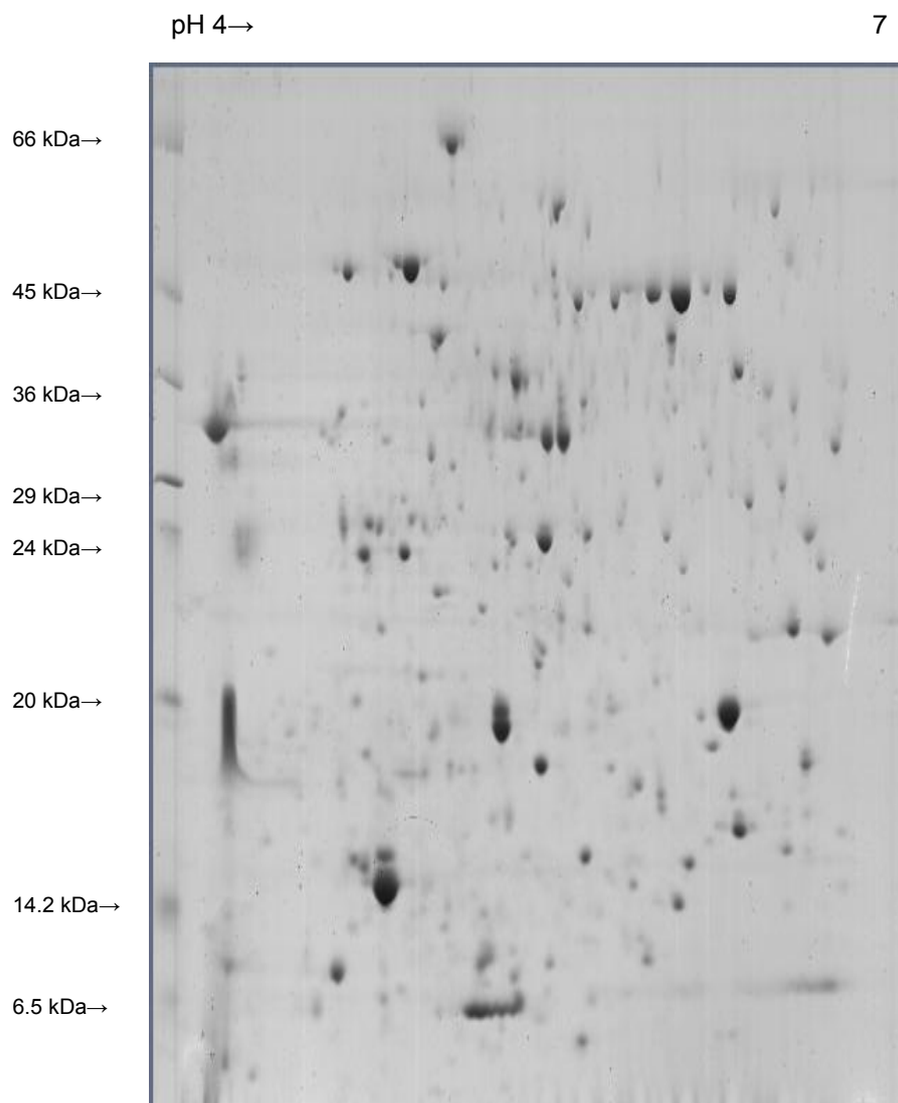


Figure 23.4: Proteome map no.4 of *B. fragilis* grown in semi-defined media without mucin

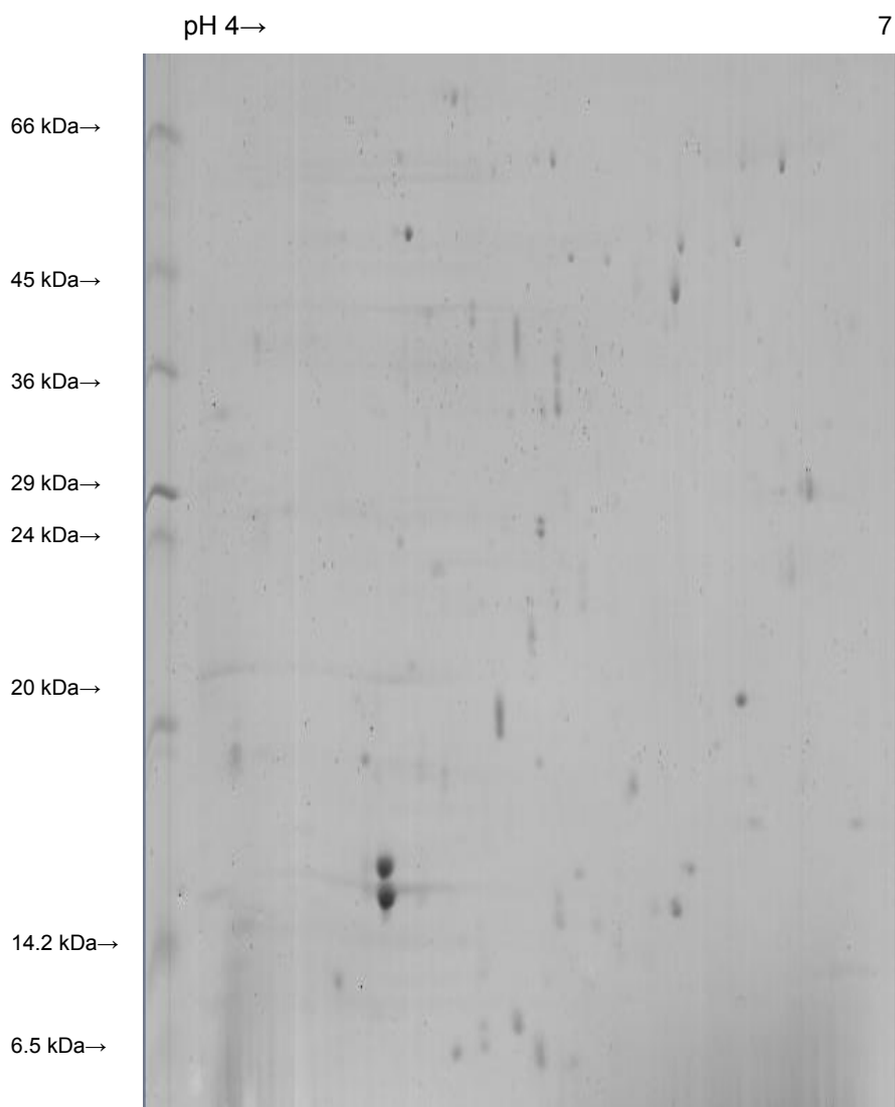


Figure 23.5: Proteome map no.5 of *B. fragilis* grown on semi-defined media without mucin

pH 4→

7

66 kDa→
45 kDa→
36 kDa→
29 kDa→
24 kDa→
20 kDa→
14.2 kDa→
6.5 kDa→



Figure 23.6: Proteome map no.6 of *B. fragilis* grown in semi-defined media without mucin

Figure 24: Proteome maps of *B. fragilis* grown in media containing mucin Type II (using pH 4-7 IEF strips).

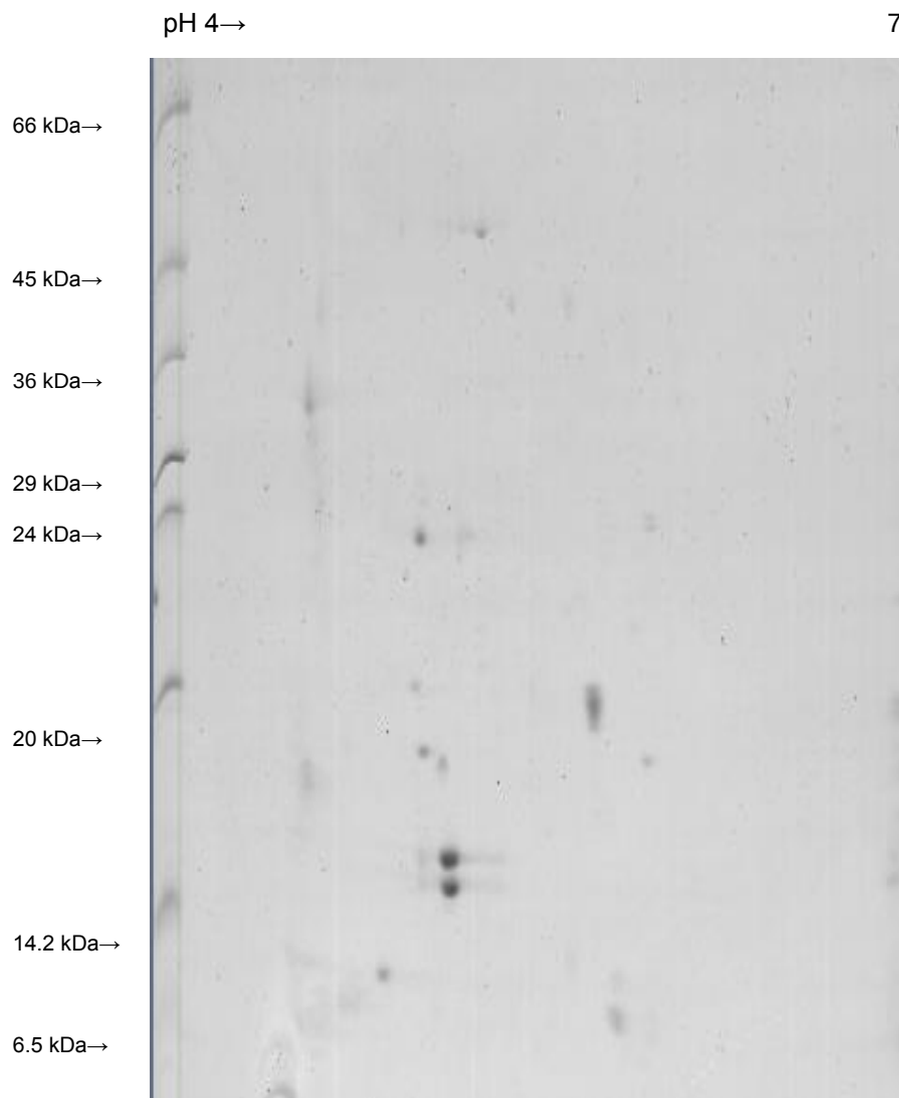


Figure 24.1: Proteome map no.1 of *B. fragilis* grown in mucin Type II enriched media

pH 4→

7

66 kDa→

45 kDa→

36 kDa→

29 kDa→

24 kDa→

20 kDa→

14.2 kDa→

6.5 kDa→

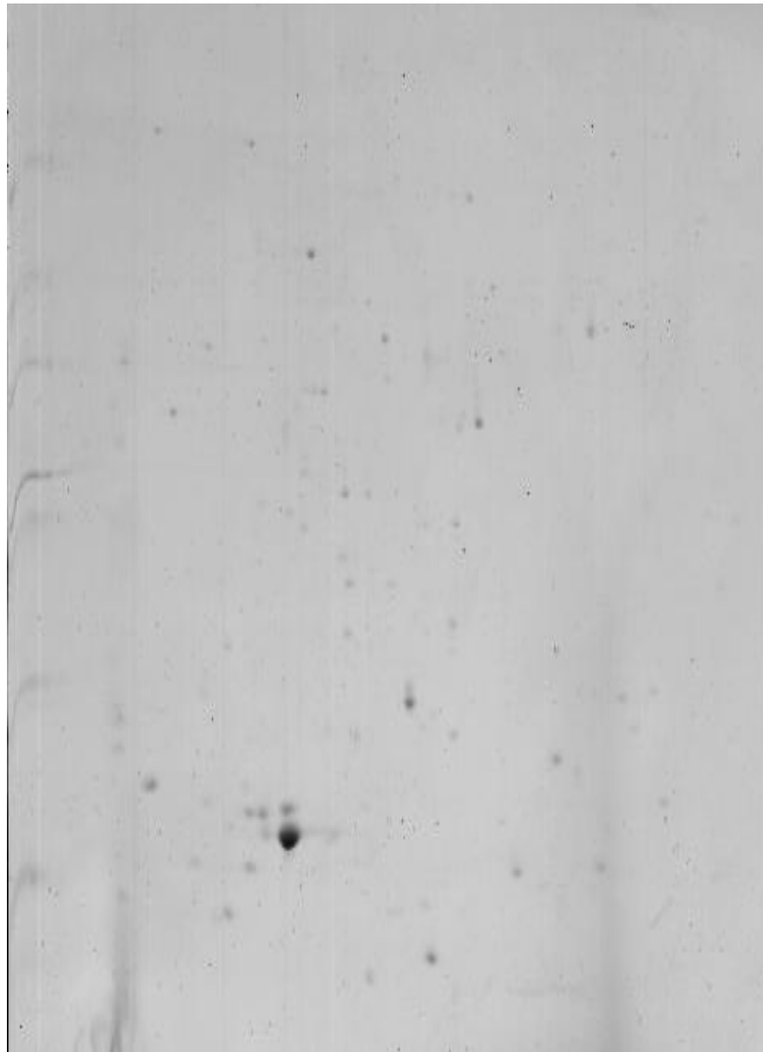


Figure 24.2: Proteome map no.2 of *B. fragilis* grown in mucin Type II enriched media

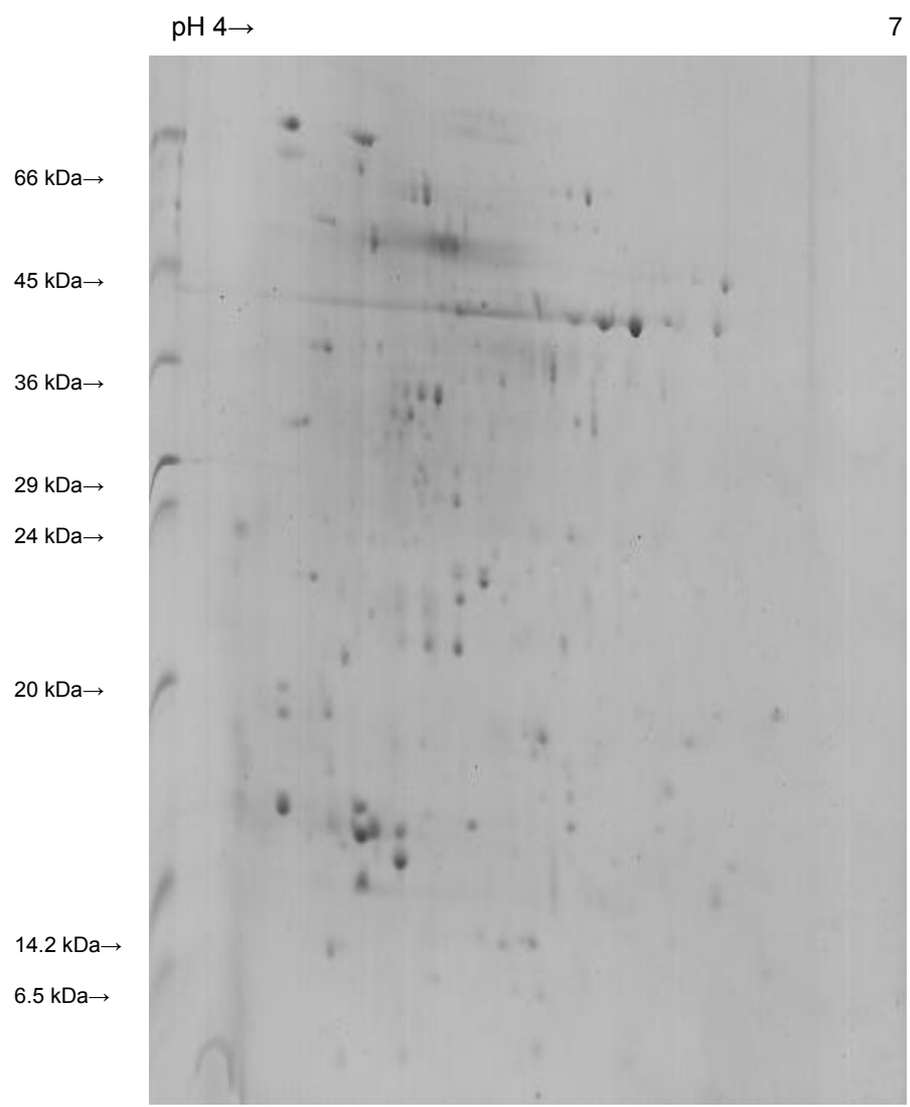


Figure 24.3: Proteome map no.3 of *B. fragilis* grown in mucin Type II enriched media

pH 4 →

7

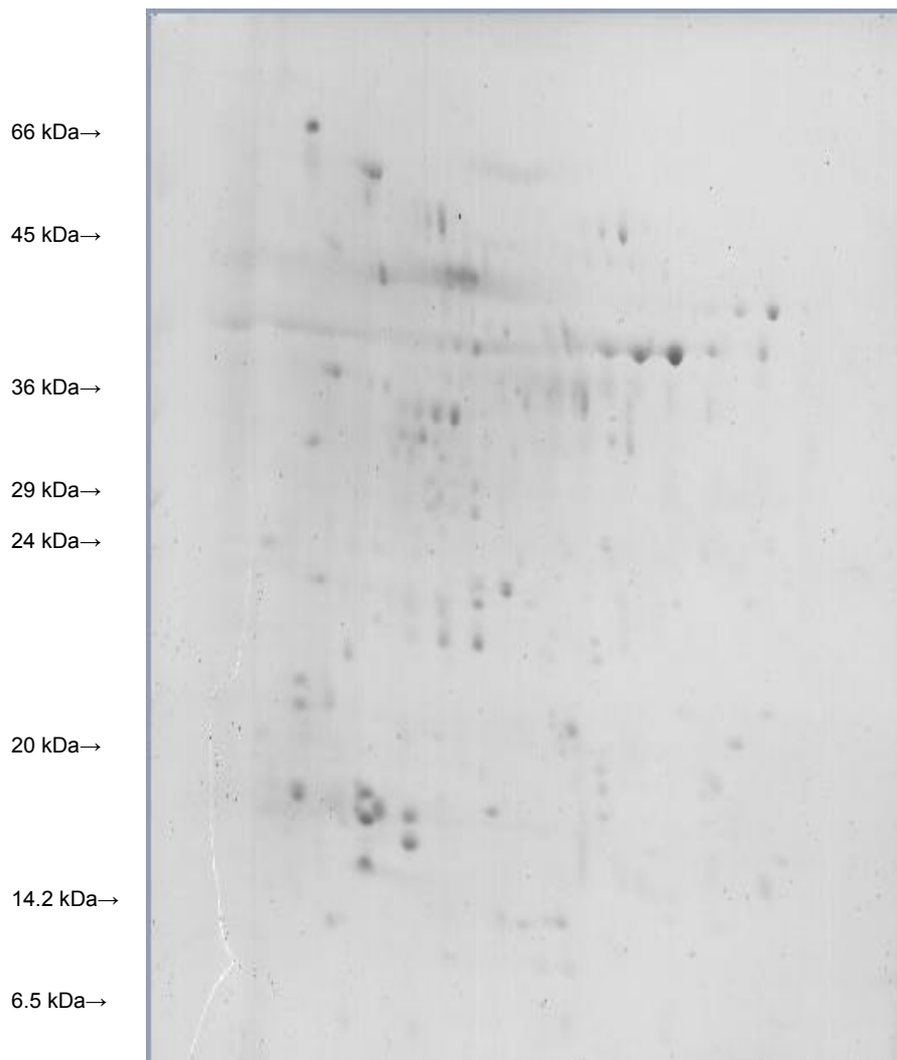


Figure 24.4: Proteome map no.4 of *B. fragilis* grown in mucin Type II enriched media

pH 4→

7

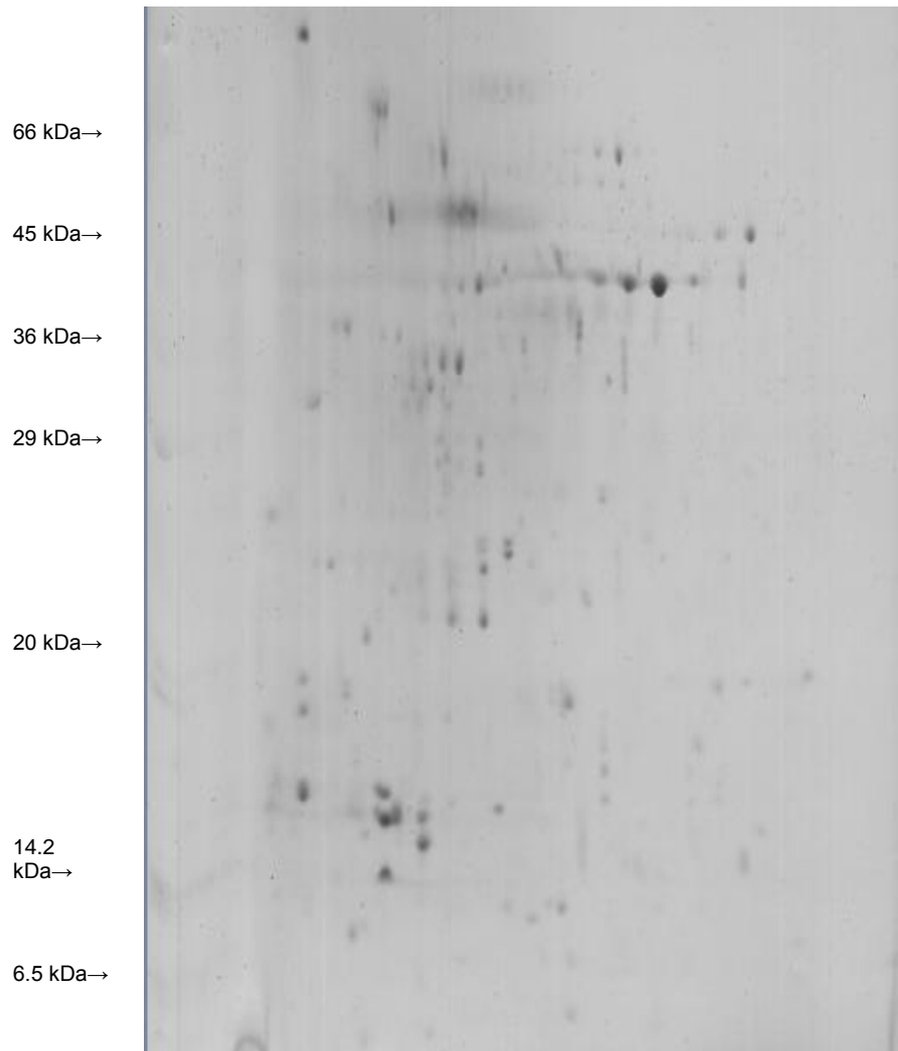


Figure 24.5: Proteome map no.5 of *B. fragilis* grown in mucin Type II enriched media

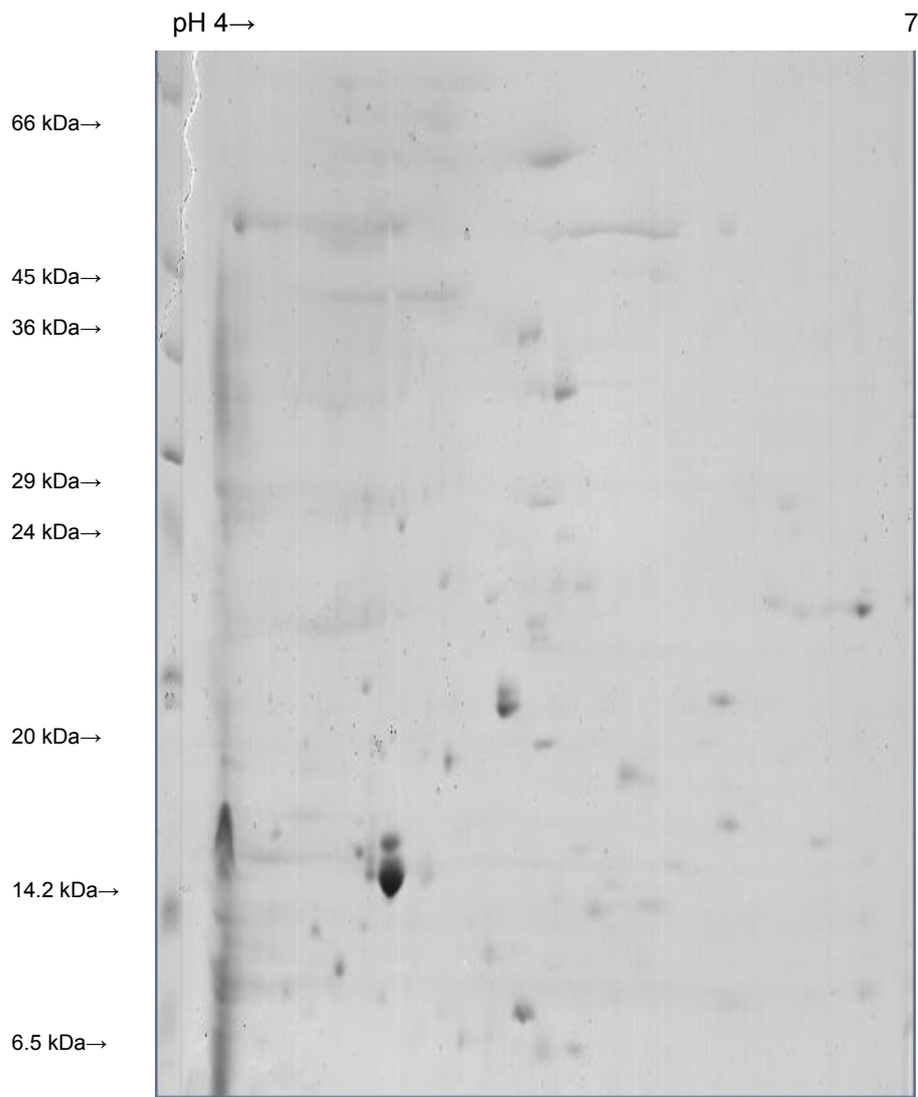


Figure 24.6: Proteome map no.6 of *B. fragilis* grown in mucin Type II enriched media

Figure 25: Proteome maps of *B. fragilis* grown in semi-defined media enriched with mucin Type III (pH 4-7 IEF strips)

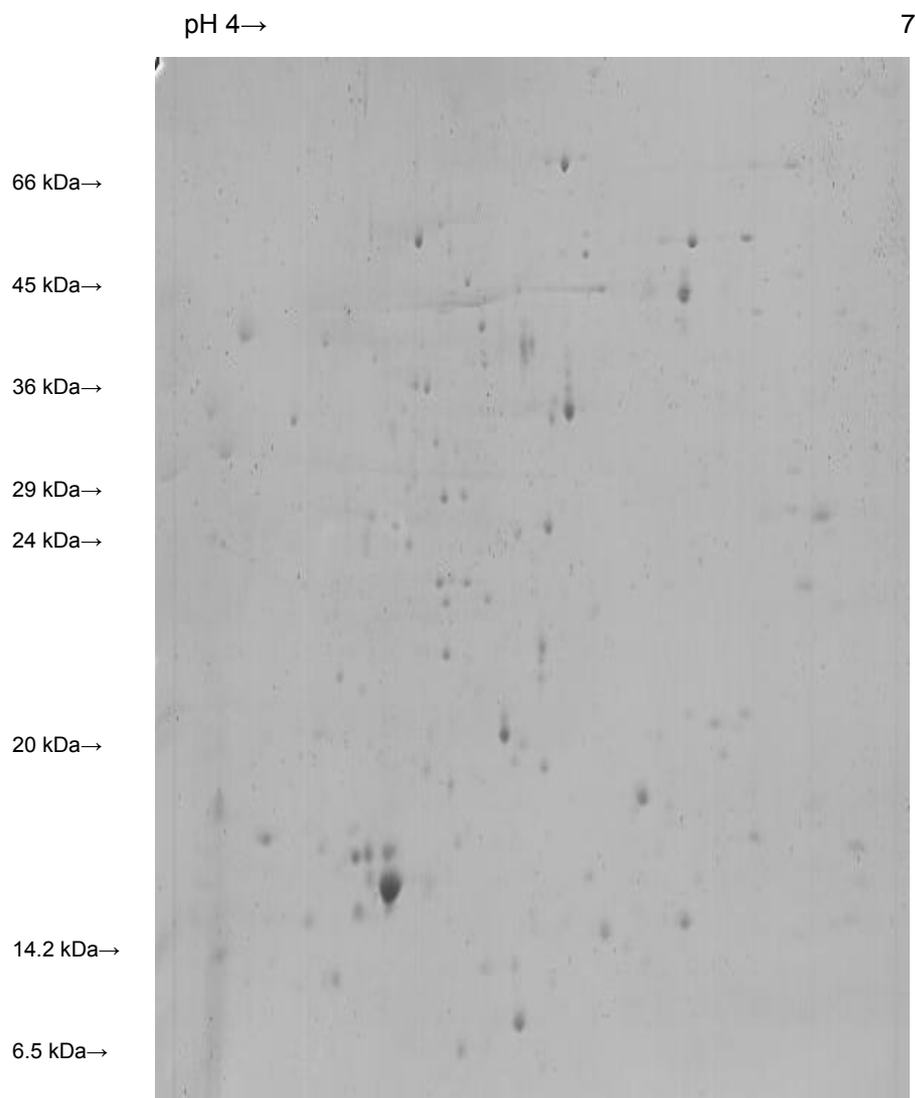


Figure 25.1: Proteome map no.1 of *B. fragilis* grown in mucin Type III enriched media

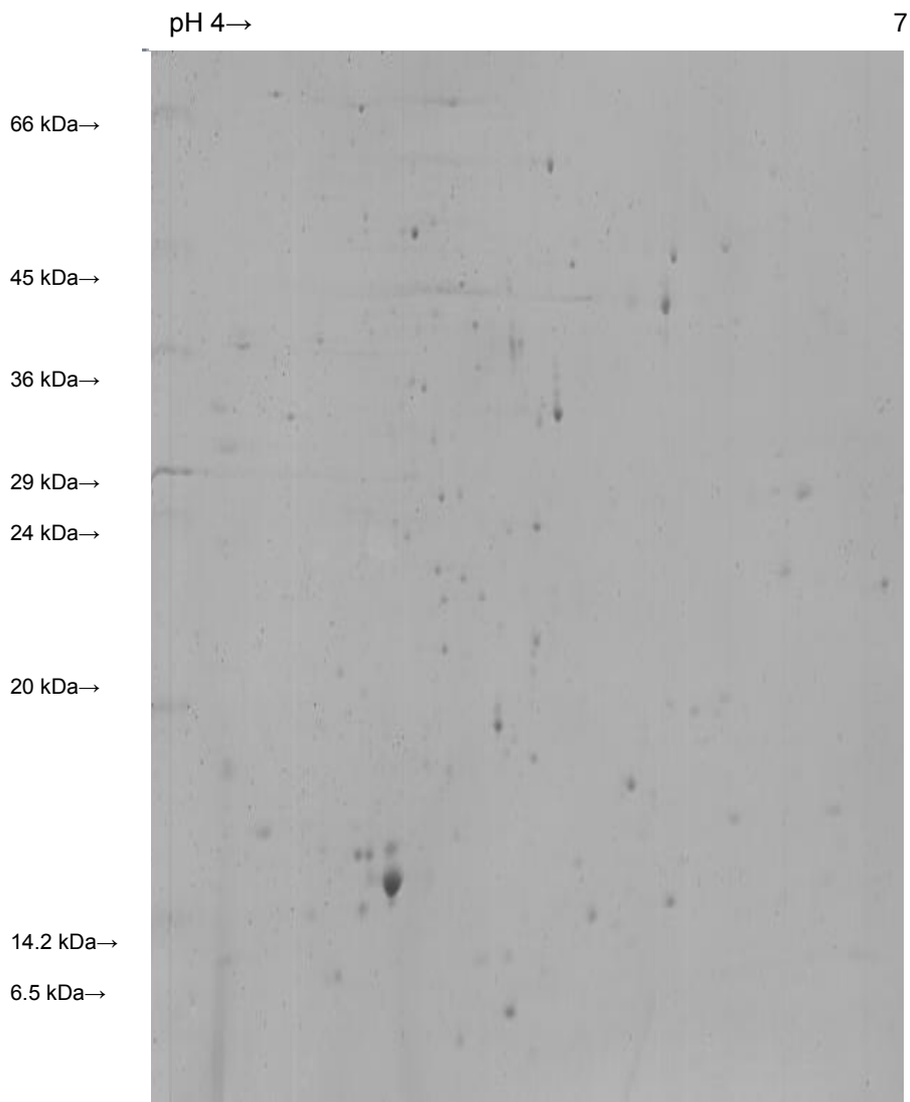


Figure 25.2: Proteome map no.2 of *B. fragilis* grown in mucin Type III enriched media

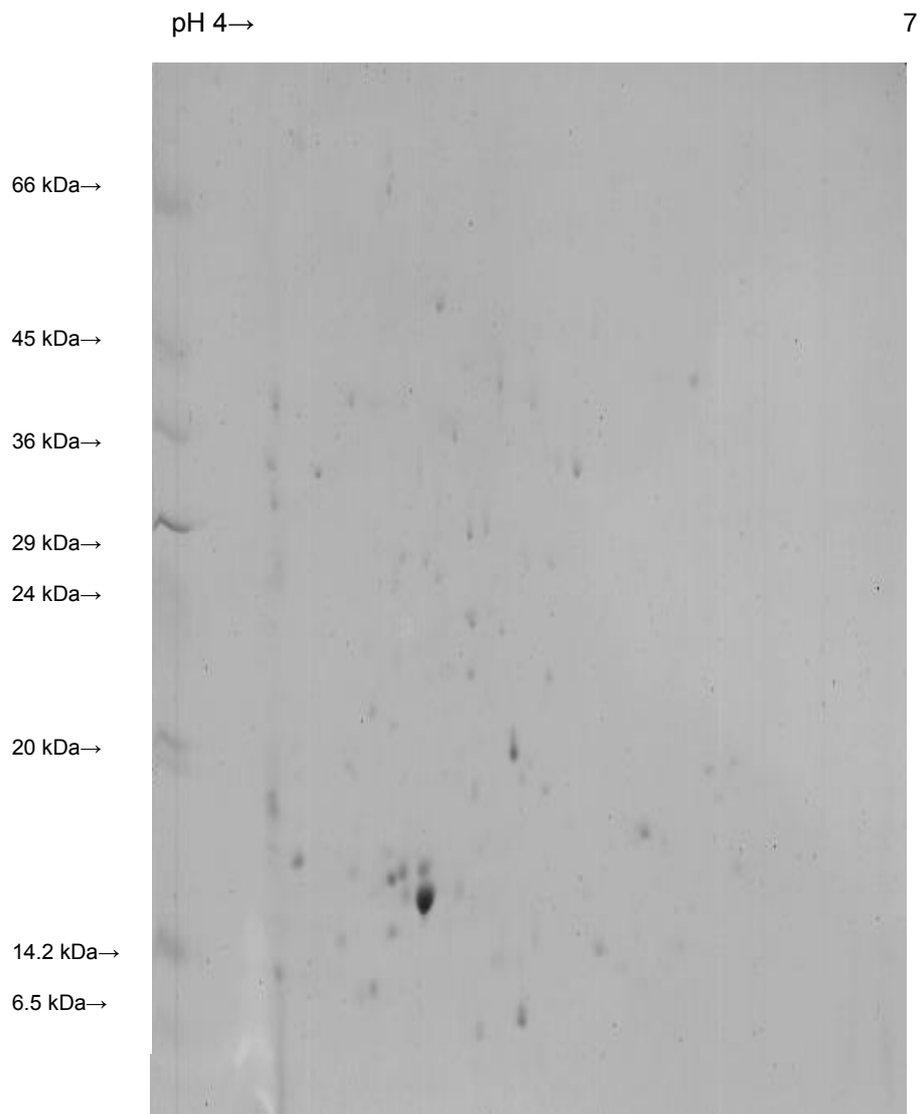


Figure 25.3: Proteome map no.3 of *B. fragilis* grown in mucin Type III enriched media

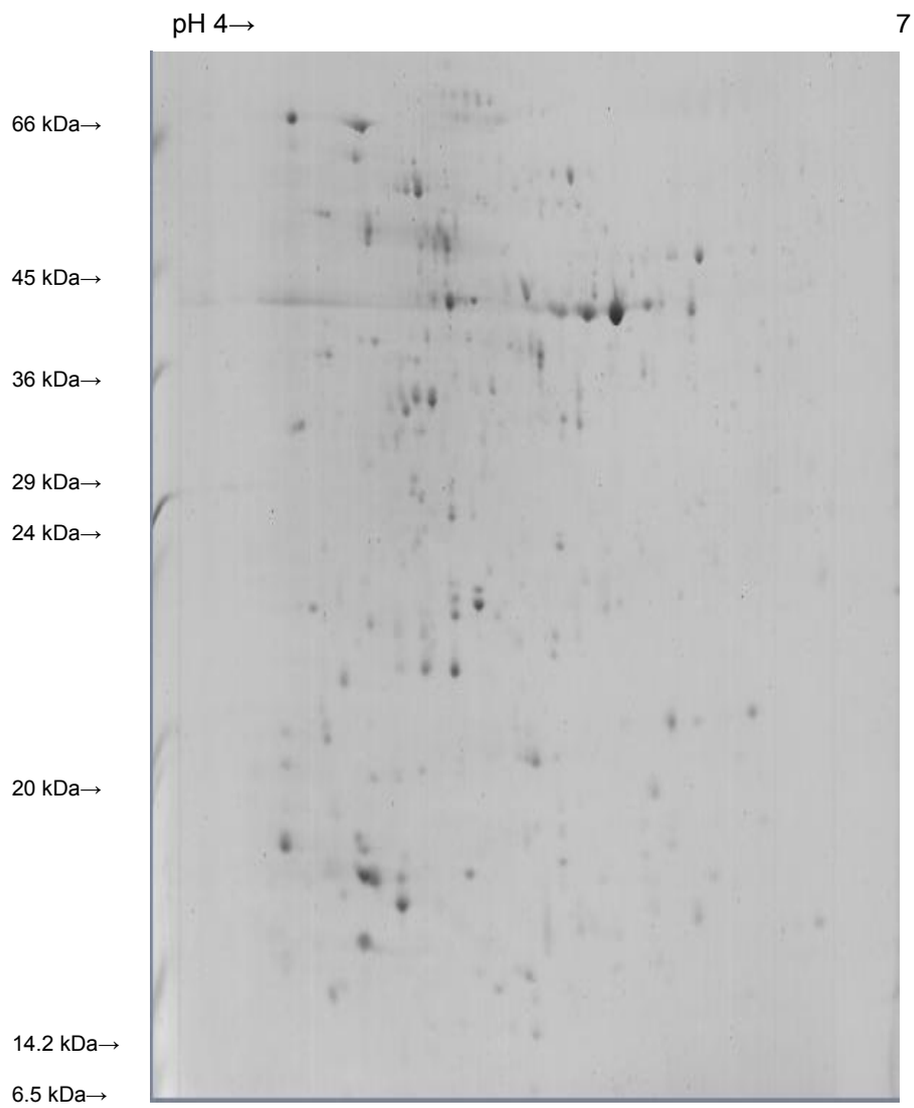


Figure 25.4: Proteome map no.4 of *B. fragilis* grown in mucin Type III enriched media

pH 4→

7

66 kDa→
45 kDa→
36 kDa→
29 kDa→
24 kDa→
20 kDa→
14.2 kDa→
6.5 kDa→

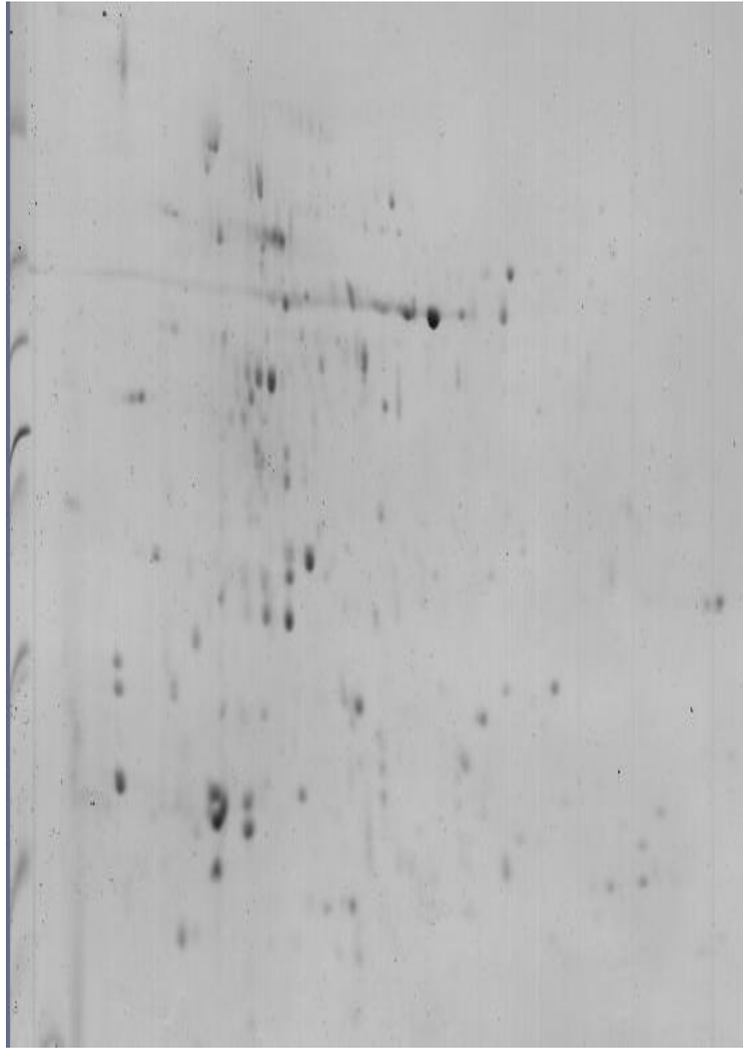


Figure 25.5: Proteome map no.5 of *B. fragilis* grown in mucin Type III enriched media

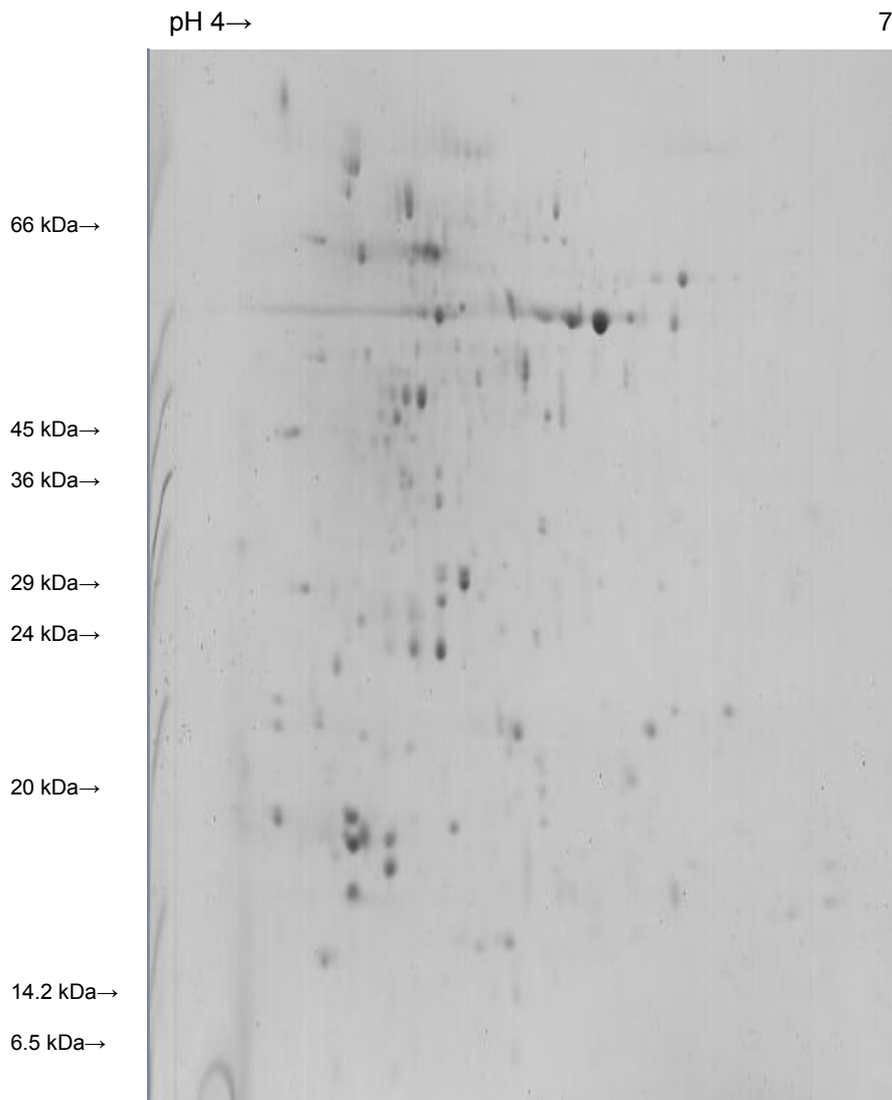


Figure 25.6: Proteome map no.6 of *B. fragilis* grown in mucin Type III enriched media

10.7 Analysis of 2DE gels

10.7.1 Analysis of 2DE gels using Bio-Rad PDQuest software

PDQuest software was used to analyse the spots present in the three different conditions. A reference master image was created which contained all the spots from all the three different conditions. Further landmarking and matching of gels was performed using the reference master gel as the template (Fig. 26.1, 26.3). The filename 'PROTBF 1' refers to the 2D gel analysis in *E. cancerogenus* and the filename 'PROTEOMICS 3' refers to the 2D gel analysis in *B. fragilis*. Refer to Appendix F for details.

The spot review tool was used to generate bar charts showing the expression of all the spots included in the reference master gel (Appendix F1). The spots are assigned SSP numbers and the graphs show the level of expression of a spot in all the individual gels or can be adjusted to show average values within replicate groups. Differentially expressed proteins were identified based on these expression patterns and cut out from the gel for further analysis. The analysis manager was used to perform statistical analysis on the spot data and the two main statistical tests performed were the Student t-test and the Mann-Whitney U test (Table 11).

Replicate groups used in analysis	Student t-test	Mann-Whitney U test	Organism
Non mucin v. mucin Type II enriched media	85/127	42/127	<i>E. cancerogenus</i>
Non mucin v. mucin Type III enriched media	32/65	33/65	<i>E. cancerogenus</i>
Non mucin v. mucin Type II enriched media	45/84	39/84	<i>B. fragilis</i>
Non mucin v. mucin Type III enriched media	66/94	28/94	<i>B. fragilis</i>

Table 11: Table represents the statistical test results for spot data in replicate groups using the Bio-Rad PDQuest software. The values represented in the table as x/y indicate that 'x' refers to the differentially expressed proteins and 'y' refers to the total proteins detected in each statistical test.

Figure 26: Reference reference master gels used in the PDQuest analysis of spot data

pH 4→

7

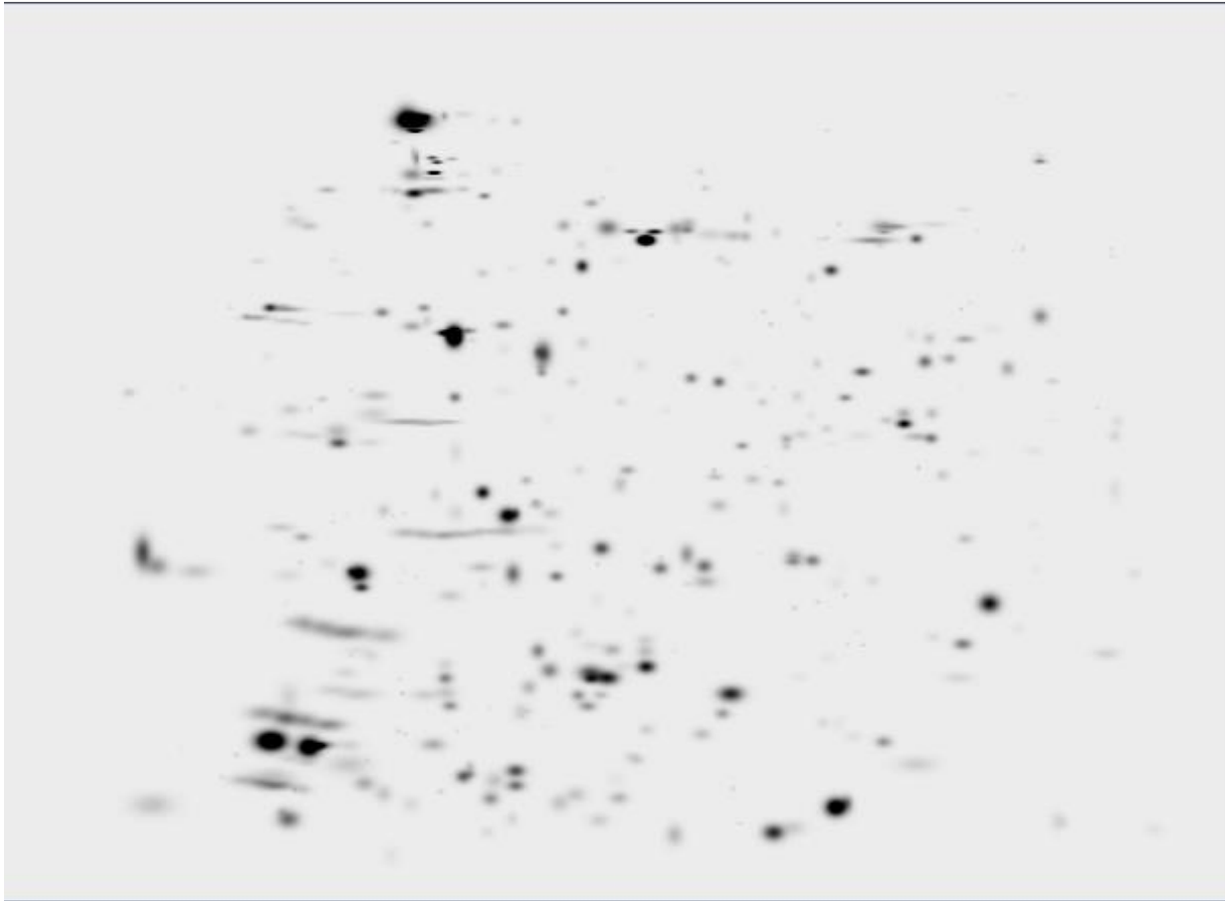


Figure 26.1: Reference master gel obtained from PDQuest analysis of *E. cancerogenus* spot data.

10.7.2 Differential expression of proteins in *E. cancerogenus*

Out of a total of 132 spots that were analysed, 98 proteins were identified using ESI LC MS/MS in *E. cancerogenus*. From the 98 proteins that were identified, 23 protein spots were differentially expressed in mucin enriched media. Seven proteins were shown to be up-regulated in both types of mucin enriched media (SSP numbers- 4008, 5306, 5601, 5607, 5701, 5702, 6203, 6605), three proteins were up-regulated in mucin Type II enriched media (SSP numbers- 5102, 7602, 8205) and 5 proteins in mucin Type III enriched media (SSP numbers- 6306, 6401, 6408, 8707). Four

proteins were found to be down-regulated in both types of mucin media (SSP numbers 3002, 5003, 7001, 8206), six proteins were found to be down-regulated in mucin Type II enriched media (SSP numbers 3601, 5302, 6303, 6408, 8707, 9601) and five proteins were down-regulated in mucin Type III enriched media (SSP numbers 2302, 5003, 5601, 6711, 8205) See Fig. 26.2 for graphical summary of differentially expressed proteins.

Proteins involved in translation namely the elongation factors Tu and greA were found to be down-regulated in both mucin enriched media but elongation factor Ts was found to be up-regulated in mucin Type II enriched media (refer to Fig. 26.2, Table 20) Proteins involved in carbohydrate metabolism namely phosphoglycerate kinase and enolase were found to be up-regulated in both mucin enriched media. Immune response proteins and toxins, ecotin was found to be down-regulated in both mucin enriched media except the surface antigen outer membrane protein X and outer membrane protein A which was shown to be up-regulated in both mucin enriched media. The amino acid biosynthesis protein domains for carbamate kinase, DHAP synthetase I and cytidylate kinase were also found to be up-regulated in Type III mucin enriched media. A DNA synthesis associated glutaredoxin related protein region was also identified as up-regulated in mucin Type II enriched media. The oxidative stress associated protein regions namely universal stress protein, ribonucleotide reductase region; pyruvate formate lyase region and autonomous glycy radical cofactor were down-regulated in mucin Type III enriched media except the universal stress protein which was up-regulated in both mucin enriched media. The other up-regulated proteins include the protein transport associated periplasmic, translocation associated tolB proteins. These were found to be up-regulated in mucin Type III enriched media. The protein folding associated thiodisulphide oxidoreductase and protein metabolism associated acetoin reductase were also up-regulated in mucin Type II enriched media. Four hypothetical proteins were detected (similar to Yba B and Yfa Z protein, Hypothetical protein 20877, 17757) of which the first two (similar to Yba B and Yfa Z protein) were found to be up-regulated in mucin Type II enriched media and the hypothetical protein 20877 was up-regulated in mucin Type III enriched media. The hypothetical protein 17757 was found to be

down-regulated in mucin Type III enriched media. Refer to Appendix G for details on the putative identifications obtained using BLAST analysis.

Analyses performed using the Redfin and Samespots softwares confirmed the up-regulation of the cell wall, membrane and envelope biogenesis associated protein outer membrane surface antigen X and carbohydrate metabolism associated phosphoglycerate kinase in both mucin enriched media in *E. cancerogenus*. Transcription associated elongation factor Ts was found to be up-regulated in mucin Type III enriched media in all the software analyses in *E. cancerogenus*. Refer to Tables 16 and 17 for further details.

Percentage regulation of spots in mucin enriched media with respect to the control non mucin media in *E. cancerogenus*

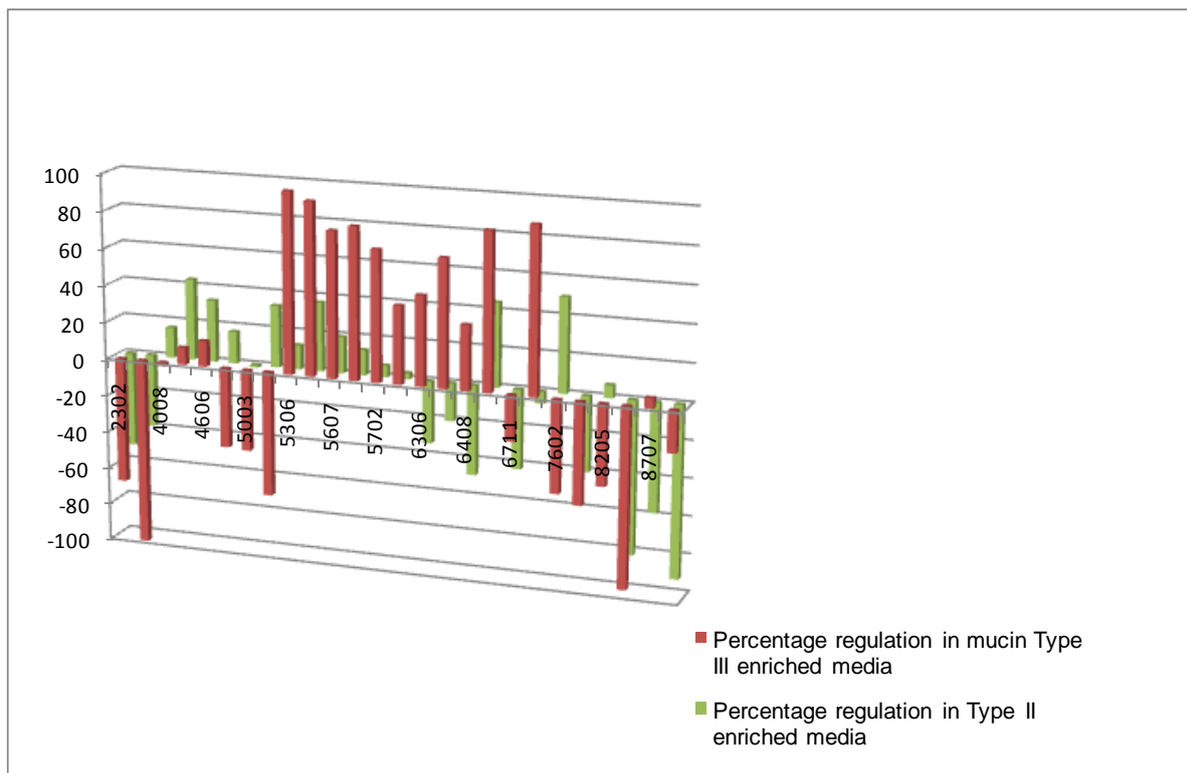


Figure 26.2: Percentage regulation of spots in mucin enriched media with respect to the control non-mucin media in *E. cancerogenus*

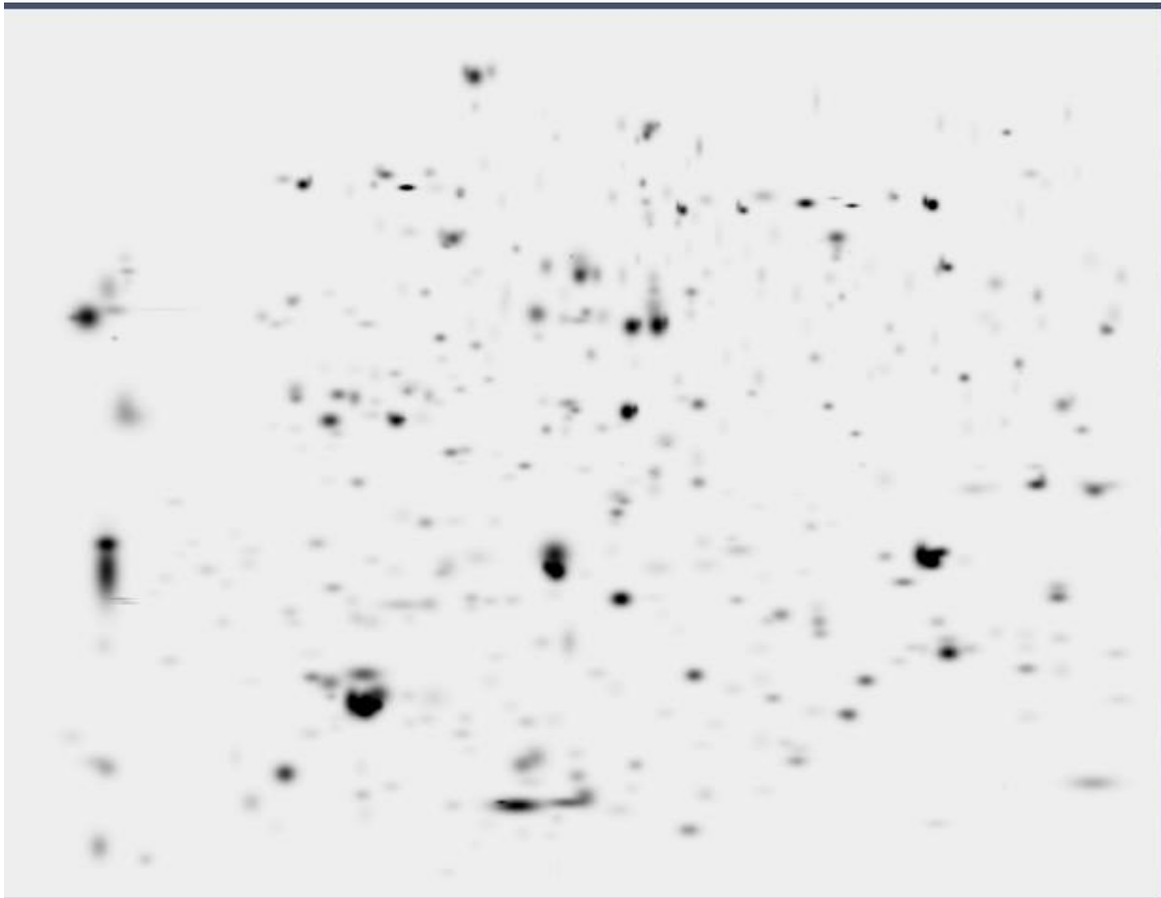


Figure 26.3: Reference master gel obtained from PDQuest analysis of *B. fragilis* spot data.

10.7.3 Differential expression of proteins in *B. fragilis*

Out of a total of 106 proteins that were analysed, 73 were identified using LC MS/MS in *B. fragilis* and 45 were found to be differentially expressed. A total of 31 spots (SSP numbers- 1104, 2101, 2102, 2401, 2701, 2704, 2804, 2806, 3303, 3404, 4502, 4601, 4607, 4608, 4702, 4804, 5106, 5201, 5205, 5402, 6102, 6103, 6202, 6206, 6301, 6304, 6802, 6807, 6901, 7301, 7401) were differentially expressed in mucin enriched media but only 5 proteins (SSP numbers- 1104, 2701, 2704, 2806 and 4804) were found to be up-regulated in mucin enriched media. SSP number 1104 was up-regulated in both types of mucin media but SSP 2701 and 2704 were up-regulated in mucin Type III enriched media, absent in mucin Type II enriched media. SSP 2806 was found to be up-regulated in both mucin enriched media. Twenty-five proteins were found to be down-regulated in mucin enriched media. Eight proteins

were found to be down-regulated in both mucin enriched media (SSP numbers 1803, 2102, 3001, 3303, 4601, 5106, 5205, 6103) whereas 12 proteins were down-regulated in mucin Type II enriched media and 5 proteins were down-regulated in mucin Type III enriched media. The presence of a host immune response may trigger the up-regulation of more proteins in the bacteria (Tran *et al.*, 2003). The predominant proteins showing regulation in mucin enriched media have been indicated in Fig. 26.4 and Table 20.

The 4 protein spots that were found to be up-regulated were the hypothetical protein BF2494 (SSP 1104), proton transport and ATP synthesis associated ATP synthase subunit E (SSP 2701, 2704) and translation associated 50S ribosomal protein L7/L12 (SSP 2806).

The presence of protein isoforms may have been indicated by the identification of the same protein in more than one spot that occurred adjacent to each other on the gels but not much information is available regarding the existence of isoforms in prokaryotic bacteria. Proteins that showed this phenomenon were the citric acid cycle associated malate dehydrogenase in *B. fragilis*, oxidative stress associated thiol peroxidase in *B. fragilis* and the immune response associated outer membrane proteins in *E. cancerogenus* and *B. fragilis* (refer to Figs. 30, 31, 33 and 34 for zoomed in images in the Discussion section).

The proteins that showed down-regulation included the carbohydrate metabolism associated phosphoenol pyruvate carboxylase, triose phosphate isomerase, oxidative stress associated thiol peroxidase, immune response associated putative outer membrane protein, glutathione detoxification system associated lactoylglutathione lyase, RNA protection associated putative RNA binding protein, translation associated ribosomal protein L7/L12 and elongation factor Ts, citric acid cycle associated malate dehydrogenase and chaperone associated GrpE protein. Refer to Table 21 for details. Two down-regulated hypothetical proteins were identified namely BF1203 and BF0301 and one hypothetical protein BF2494 was up-regulated. Refer to Appendix G4 for details on the putative identifications obtained using BLAST analysis. Hypothetical proteins BF2494 were found to be up-regulated in mucin Type II enriched media when analysed using both PDQuest and SameSpots softwares but not in Redfin. Similarly, BF1203 hypothetical proteins were

found to be down-regulated in both mucin Type II and III enriched media when analysed using both PDQuest and SameSpots softwares (tests were based on the Student t-test).

Analyses performed using the Redfin and Samespots softwares in *B. fragilis* confirmed the down-regulation of the carbohydrate metabolism associated triose phosphate isomerase in mucin Type II enriched media. Energy production and conversion associated malate dehydrogenase proteins were found to be down-regulated in mucin Type II enriched media but up-regulated in mucin Type III enriched media in *B. fragilis*. All analyses also showed the up-regulation of the transcription protein elongation factor Ts in mucin Type II enriched media. Student t-test was used to perform the analyses using all the three softwares where expression levels were considered based on average values in each condition.

Percentage regulation of spots in mucin enriched media with respect to the control non mucin media in *B. fragilis*

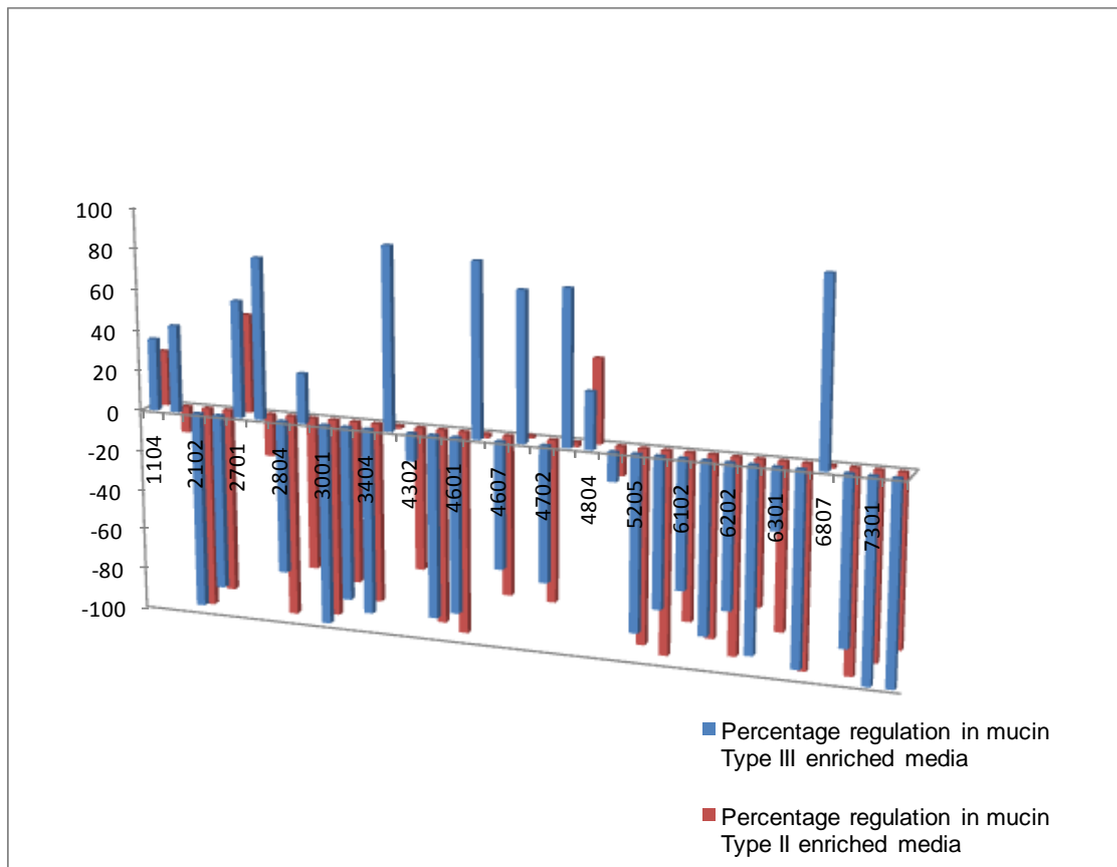


Figure 26.4: Percentage regulation of spots in mucin enriched media with respect to the control non-mucin media in *B. fragilis*

The general growth rate of both *B. fragilis* and *E. cancerogenus* were not profoundly influenced by the presence of mucin in the media especially with reference to culture density.

10.7.4 Analysis of 2DE gels using Ludesi Redfin software

Reports have been generated for the 2DE gel analysis using the Ludesi Redfin software for studying the differential expression of proteins in control vs mucin enriched media in *E. cancerogenus* and *B. fragilis*. Refer to Appendix F2 for results, including p values and expression levels of *E. cancerogenus* and *B. fragilis* proteins in non-mucin, mucin Type II and Type III enriched media. Spots selected for analysis were based on significant p values of <0.05 and a fold change of 1.5 or more. Expression profile filters were also used to detect up or down regulation of proteins.

10.7.5 Analysis of 2DE gels using Nonlinear Dynamics SameSpots software

The SameSpots analysis software from Nonlinear Dynamics was also used to study the differential expression of proteins in both bacteria. The spots of interest were selected based on the analysis of variance where significant p values of <0.05 and fold changes of 1.0 or more were considered. Refer to Appendix F3 for the analysis report of *E. cancerogenus* and *B. fragilis* spot data, respectively.

10.7.6 Comparative analysis of spot data

The spots with relevant p values from each analysis were manually compared in all the three softwares (Fig 27; Tables 12-19). The proteins highlighted in 'bold' font in the tables indicate differential expression in all software analyses under those specific growth conditions and statistical tests. SSP number refers to the Standard spot number which has been used to identify the protein spots using the default Bio-Rad PDQuest software. Each circle of the Venn diagram indicates the differentially expressed proteins in one particular software package and the overlap regions

indicate the proteins identified commonly in two of the software packages. The central region where all the three circles overlap shows the proteins that have been differentially expressed in all the three software packages (Fig. 27). SSP numbers have been indicated for spots that remained unidentified using LC-MS analysis.

Since SameSpots works only on the basis of average expression in each growth condition, the Mann-Whitney U test, where expression of a protein in each individual gel can be considered, could not be performed using data from SameSpots. Therefore the Student t-test was used to compare SameSpots, PDQuest and Redfin data, and the Mann-Whitney U test was used to compare PDQuest and Redfin data.

The Student t-test was used to identify proteins with a significant p value (<0.05) from all three softwares and the average values were used to compare expression. The Mann-Whitney U test was used to identify proteins with a significant p value (<0.05) from all three softwares but the expression in individual gels was considered unlike the Student t-test where the average values were compared.

Figure 27: Comparative analyses of spot data using PDQuest, Redfin and SameSpots softwares

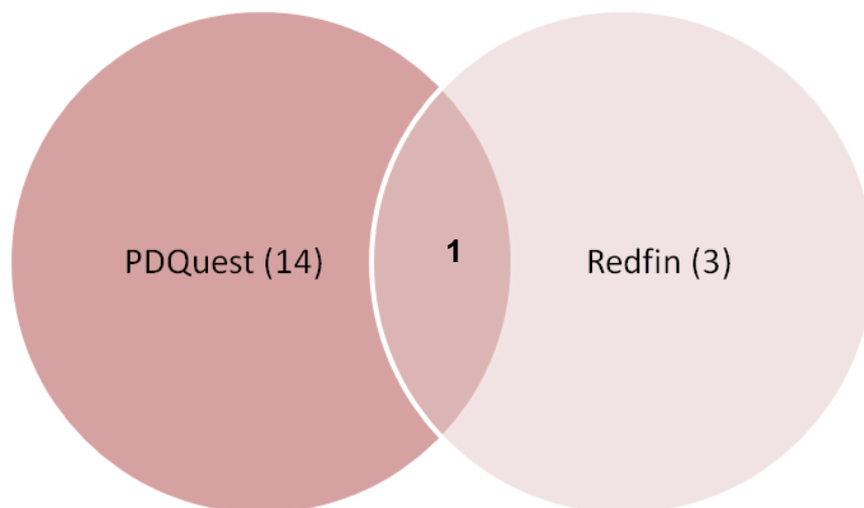


Figure 27.1: Comparative analysis of proteins differentially expressed by *E. cancerogenus* in semi-defined media without mucin vs mucin Type II enriched media by PDQuest and Redfin softwares using Mann-Whitney U test. Refer to Table 12 for details on the differentially expressed proteins in media without mucin vs mucin Type II enriched media by PDQuest and Redfin softwares using the Mann-Whitney U test.

Protein identity- <i>E. cancerogenus</i>	PDQuest analysis	Redfin analysis
SSP 7001 (Not identified)	↑	↑ (4.57, 0.0144)
SSP 6102 (Not identified)		↑ (2.55, 0.0247)
Peptidoglycan binding domain (5305)		↑ (2.51, 0.0089)
Ecotin (8206)	↓	
Glutaredoxin related protein (3002)	↓	
SSP 2802 (Not identified)	↑	
Autonomous glycy radical cofactor (4008)	↑	
Outer membrane protein surface antigen X (6203)	↑	
Thiol disulphide oxidoreductase protein (6408)	↓	
Acetoin reductase (7602)	↑	
Tol B protein (8707)	↓	
Outer membrane protein A (4606)	↑	
Phosphoglycerate kinase (5701)	↑	
SSP 1602 (Not identified)	↑	
SSP 5302 (Not identified)	↓	
SSP 6303 (Not identified)	↓	

Table 12: Differentially expressed *E. cancerogenus* proteins in non mucin vs mucin Type II enriched media by PDQuest and Redfin softwares using the Mann-Whitney U test. ↑ indicates up-regulation of the protein, ↓ indicates down-regulation of the protein. (Fold change, p value) for differentially expressed spots has been indicated in brackets for Redfin and SameSpots softwares. SSP numbers of protein spots are indicated in the first column.

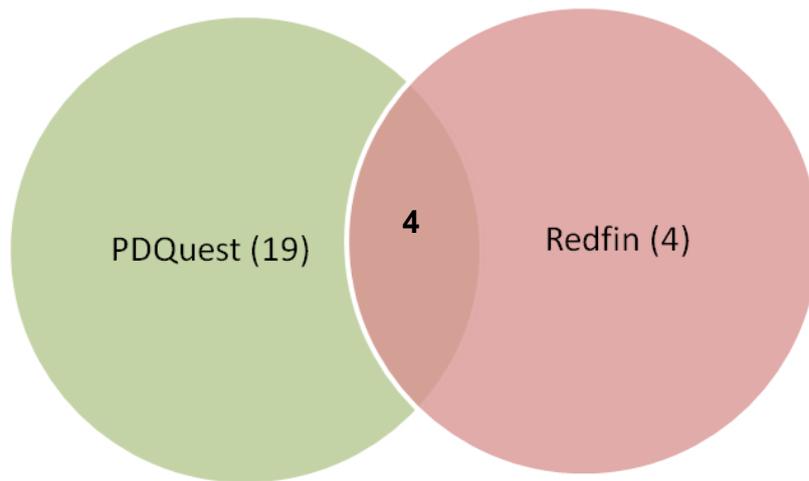


Figure 27.2: Comparative analysis of proteins differentially expressed by *E. cancerogenus* in semi-defined media without mucin vs mucin Type III enriched media by PDQuest and Redfin softwares using the Mann-Whitney U test. Refer to Table 13 for details on the differentially expressed *E. cancerogenus* proteins in media without mucin vs. mucin Type III enriched media by PDQuest and Redfin softwares using the Mann-Whitney U test.

Protein identity- <i>E. cancerogenus</i>	PDQuest analysis	Redfin analysis
Elongation factor Ts (5601)	↓	↓ (2.17, 0.0026)
Carbamate kinase (5607)	↑	↑ (2.58, 0.0296)
Periplasmic protein (5306)	↑	↑ (2.93, 0.0262)
Ecotin (8206)	↓	
Autonomous glycy radical cofactor (4008)	↓	
Phosphoglycerate kinase (5701)	↑	
Outer membrane protein surface antigen X (6203)	↑	↑ (2.85, 0.0143)
Elongation factor greA (2302)	↓	
Glutaredoxin related protein (3002)	↓	
Hypothetical protein 17757(5003)	↓	
Universal stress protein A (5102)	↑	
Enolase (5705)	↑	
Hypothetical protein 20877 (6306)	↑	
Cytidylate kinase (6401)	↑	
2-dehydro-3-deoxy phosphooctonate aldolase (6605)	↑	
Elongation factor Tu (6711)	↓	
SSP 2402 (Not identified)	↑	
SSP 3602 (Not identified)	↑	
Tol B (8707)	↑	

Table 13: Differentially expressed *E. cancerogenus* proteins in non mucin vs mucin Type III enriched media by PDQuest and Redfin softwares using the Mann-Whitney U test

↑ indicates up-regulation of the protein

↓ indicates down-regulation of the protein. (Fold change, p value) for differentially expressed spots has been indicated in brackets for Redfin and SameSpots softwares. SSP numbers of protein spots are indicated in the first column.

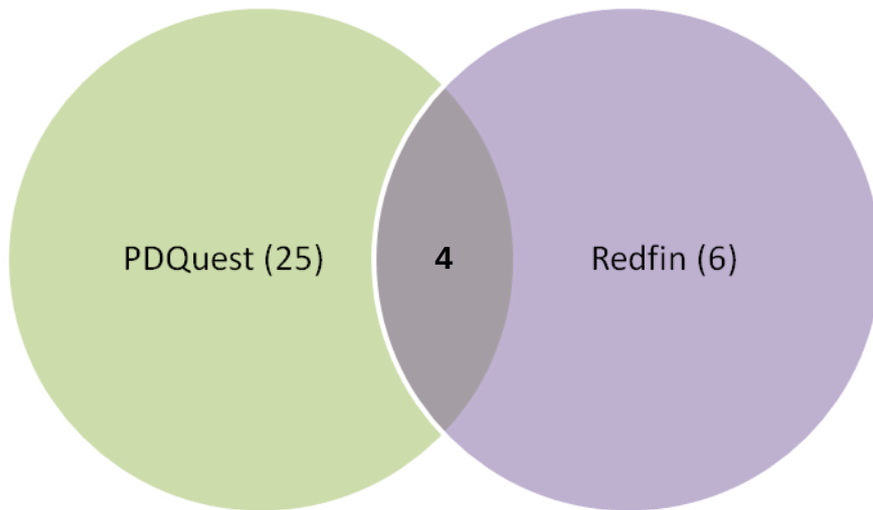


Figure 27.3: Comparative analysis of proteins differentially expressed by *B. fragilis* in semi-defined media without mucin v mucin Type II enriched media by PDQuest and Redfin software using Mann-Whitney U test. Refer to Table 14 for details on the differentially expressed *B. fragilis* proteins in media without mucin vs. mucin Type II enriched media by PDQuest and Redfin softwares using the Mann-Whitney U test based.

Protein identity- <i>B. fragilis</i>	PDQuest analysis	Redfin analysis
Putative thiol peroxidase (6304)		↓ (2.22, 0.0264)
Outer membrane protein H (6202)	↓	
Putative RNA binding protein (6102)	↓	↓ (3.66, 0.0139)
SSP 3206 (Not identified)	↓	↓ (3.11, 7.293e-4)
Putative thiol peroxidase (4601)	↓	↓ (3.07, 0.0163)
Malate dehydrogenase (4607)	↓	↓ (3.07, 3.5558e-4)
Hypothetical protein BF2494 (1104)	↑	
50S Ribosomal protein L7/L12 (2102)	↓	
Hypothetical protein BF2494 (2804)	↓	
50S Ribosomal protein L7/L12 (3001)	↓	
Triose phosphate isomerase (4502)	↓	↓ (2.69, 0.0136)
Elongation factor Ts (4702)	↓	
50S Ribosomal protein L7/L12 (5106)	↓	
Hypothetical protein BF1203 (5205)	↓	
Phosphoenol pyruvate carboxykinase (6901)	↓	
Lactoylglutathione lyase (6103)	↓	
Putative outer membrane protein H (5402)	↓	
Hypothetical protein BF 0301 (6206)	↓	
Fructose bisphosphate aldolase (6802)	↓	
Superoxide dismutase (7301)	↓	
SSP 1103 (Not identified)	↑	
SSP 1211 (Not identified)	↑	
SSP 5401(Not identified)	↓	
SSP 1107(Not identified)	↑	
SSP 1803 (Not identified)	↓	
Putative thiol peroxidase (6301)	↑	

Table 14: Differentially expressed *B. fragilis* proteins in non mucin vs mucin Type II enriched media by PDQuest and Redfin softwares using the Mann-Whitney U test. ↑ indicates up-regulation of the protein; ↓ indicates down-regulation of the protein. (Fold change, p value) for differentially expressed spots has been indicated in brackets for Redfin and SameSpots softwares. SSP numbers of protein spots are indicated in the first column.

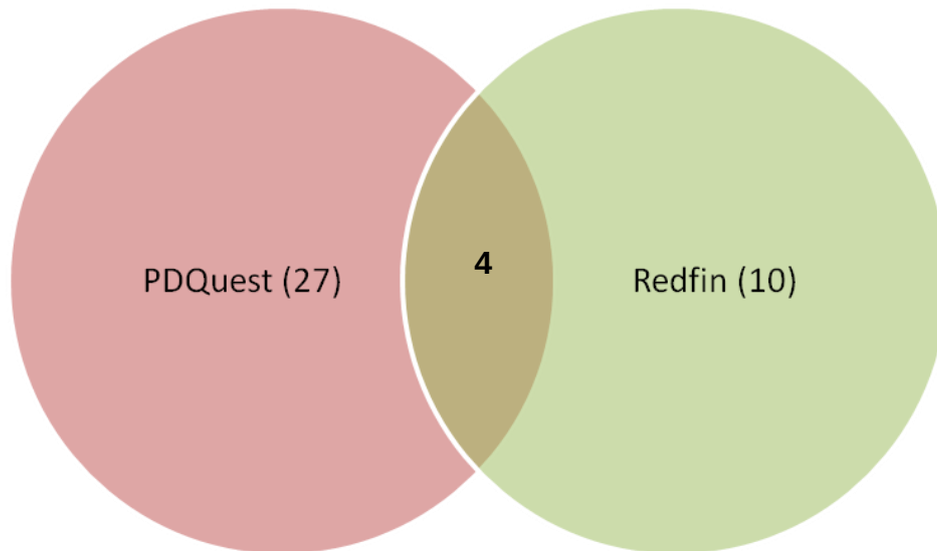


Figure 27.4: Comparative analysis of proteins differentially expressed by *B. fragilis* in semi-defined media without mucin vs mucin Type III enriched media by PDQuest and Redfin software using the Mann-Whitney U test. Refer to Table 15 for details on the differentially expressed *B. fragilis* proteins in media without mucin vs. mucin Type III enriched media by PDQuest and Redfin softwares using the Mann-Whitney U test.

Protein identity- <i>B. fragilis</i>	PDQuest analysis	Redfin analysis
Putative thiol peroxidase (6304)		↓ (2.22, 0.0264)
Outer membrane protein H (6202)	↓	↓ (4.69, 0.0176)
SSP 3206 (Not identified)		↑ (3.11, 7.293e-4)
Putative thiol peroxidase (4601)	↓	↓ (3.07, 0.0163)
SSP 1207 (Not identified)		↓ (2.65, 0.0101)
SSP 3305 (Not identified)		↓ (2.47, 0.0479)
Malate dehydrogenase (4607)	↓	↓ (2.62, 0.0211)
Malate dehydrogenase (4608)	↑	↑ (3.07, 3.5558e-4)
SSP 5002 (Not identified)		↓ (2.89, 0.0037)
50S Ribosomal protein L7/L12 (2102)	↓	
50S Ribosomal protein L7/L12 (5106)	↓	
Lactoylglutathione lyase (6103)	↓	
Triose phosphate isomerase (6302)	↓	↓ (2.69, 0.0136)
Hypothetical protein BF2494 (2804)	↓	
SSP 1103 (Not identified)	↑	
Hypothetical protein BF2494 (1104)	↑	
GrpE protein (2401)	↓	
ATP synthase subunit E (2701)	↑	
SSP 4101 (Not identified)	↑	
Malate dehydrogenase (4804)	↑	
Hypothetical protein BF1203 (5201)	↓	
Superoxide dismutase (3404)	↑	
Thioredoxin (6301)	↑	
Hypothetical protein BF1203 (5205)	↓	
Hypothetical protein BF 0301(6807)	↓	
SSP 1211 (Not identified)	↑	
SSP 1402 (Not identified)	↓	
SSP 1803 (Not identified)	↓	
SSP 2403 (Not identified)	↑	
SSP 2504 (Not identified)	↑	
ATP synthase subunit E (2704)	↑	
50S Ribosomal protein L7/L12 (2806)	↑	

Table 15: Differentially expressed *B. fragilis* proteins in non mucin vs mucin Type III enriched media by PDQuest and Redfin softwares using the Mann-Whitney U test. ↑ indicates up-regulation of the protein; ↓ indicates down-regulation of the protein. (Fold change, p value) for differentially expressed spots has been indicated in brackets for Redfin and SameSpots softwares.

SSP numbers of protein spots are indicated in the first column.

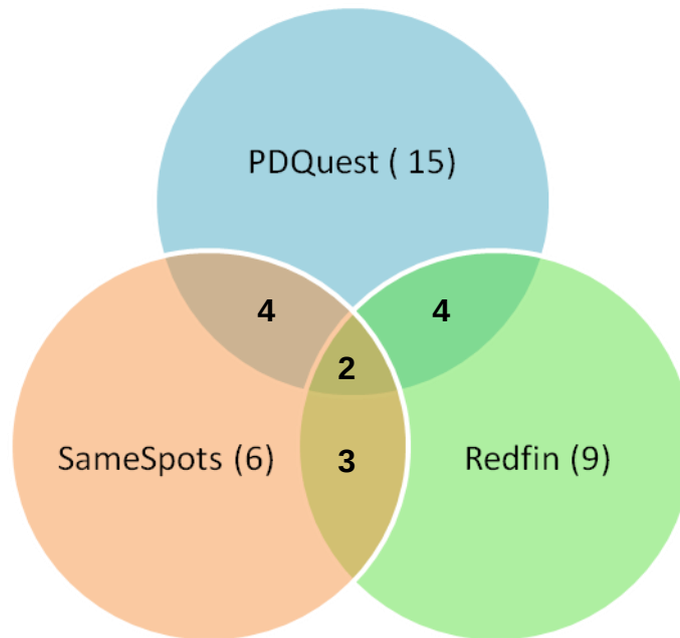


Figure 27.5: Comparative analysis of proteins differentially expressed by *E. cancerogenus* in semi-defined media without mucin vs mucin Type II enriched media by PDQuest, Redfin and Same spots software using the Student t-test. Refer to Table 16 for details on the differentially expressed *E.cancerogenus* proteins in media without mucin vs. mucin Type II enriched media by PDQuest, SameSpots and Redfin softwares using the Student t-test.

Protein identity- <i>E. cancerogenus</i>	PDQuest analysis	Redfin analysis	Samespots analysis
SSP 7001 (Not identified by mass spectrometry)	↑	↑ (4.57, 0.0144)	
SSP 6102 (Not identified)		↑ (2.55, 0.0247)	
SSP 1602 (Not identified)	↓		↓ (3.4, 0.024)
Peptidoglycan binding domain (5305)		↑ (2.51, 0.0089)	
Ecotin (8206)	↓		
Glutaredoxin related protein (3002)	↓		
Autonomous glycy radical cofactor (4008)	↓		
Outer membrane protein X (6203)	↑	↑ (2.85, 0.0143)	↑ (2.3, 0.006)
Acetoin reductase (7602)	↑		
Tol B protein (8707)	↓		
Outer membrane protein II (4505)	↓		
Enolase (5705)			↑ (6.5, 0.029)
Phosphoglycerate kinase (5701)	↑	↑(2.51, 0.0089)	↑ (2.3, 0.032)
SSP 7003 (Not identified)		↑ (2.55, 0.0116)	↑ (2.3, 0.006)
Inorganic pyrophosphatase (5405)	↑	↑ (1.73, 0.0033)	
Thiol disulphide oxidoreductase protein (6408)	↓		↓ (2.2, 0.003)
SSP 2509 (Not identified)	↑		
SSP 3602 (Not identified)	↓		
SSP 5302 (Not identified)	↓		
Putative glutathione peroxidase (4701)		↑ (1.81, 0.0459)	
ABC type sugar transport system (6001)		↑ (4.41, 0.045)	

Table 16: Differentially expressed *E. cancerogenus* proteins in non mucin vs mucin Type II enriched media by PDQuest and Redfin and Samespots softwares using the Student t-test (i.e. based on averages) ↑ indicates up-regulation of the protein. ↓ indicates down-regulation of the protein. (Fold change, p value) for differentially expressed spots has been indicated in brackets for Redfin and SameSpots softwares. SSP numbers of protein spots are indicated in the first column.

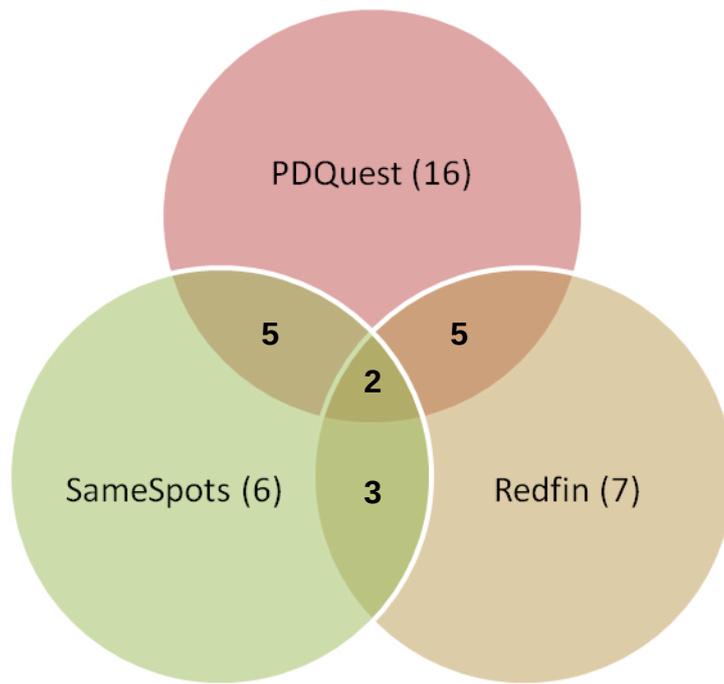


Figure 27.6: Comparative analysis of proteins differentially expressed by *E. cancerogenus* in semi-defined media without mucin v mucin Type III enriched media by PDQuest, Redfin and SameSpots softwares using the Student t-test (i.e. based on averages). Refer to Table 17 for details on the differentially expressed *E.cancerogenus* proteins in media without mucin vs. mucin Type III enriched media by PDQuest, SameSpots and Redfin softwares using the Student t-test.

Protein identity- <i>E. cancerogenus</i>	PDQuest analysis	Redfin analysis	Samespots software
Elongation factor Ts (5601)	↓	↓ (2.17, 0.0026)	↓ (2.5, 0.041)
Carbamate kinase (5607)	↓	↓ (2.58, 0.0296)	
Periplasmic protein (5306)	↑	↑ (2.93, 0.0262)	
Ecotin (8206)	↓		↓ (6.0, 3.043e-004)
Autonomous glyceryl radical cofactor (4008)	↓		
Phosphoglycerate kinase (5701)	↑		
Outer membrane protein surface antigen X (6203)	↑	↑ (2.85, 0.0143)	↑ (2.3, 0.006)
Elongation factor greA (2302)	↓		
Glutaredoxin related protein (3002)	↓		
Hypothetical protein 17757 (5003)	↓		↓ (2.1, 0.054)
Universal stress protein (5102)	↑		
Enolase (5705)	↑		
Outer membrane protein A (4606)	↑		
Cytidylate kinase (6401)	↑		↑ (2.5, 0.045)
DAHPh synthetase (6605)	↑		
Elongation factor Tu (6711)	↑	↑ (2.55, 0.0247)	
SSP 7003 (Not identified)		↑ (2.55, 0.0116)	↑ (2.3, 0.006)
ABC type sugar transport system (6001)		↑ (4.41, 0.045)	

Table 17: Differentially expressed *E. cancerogenus* proteins in non mucin vs mucin Type III enriched media by PDQuest and Redfin and SameSpots softwares using the Student t-test (i.e. based on averages). ↑ indicates up-regulation of the protein

↓ indicates down-regulation of the protein. (Fold change, p value) for differentially expressed spots has been indicated in brackets for Redfin and SameSpots softwares. SSP numbers of protein spots are indicated in the first column.

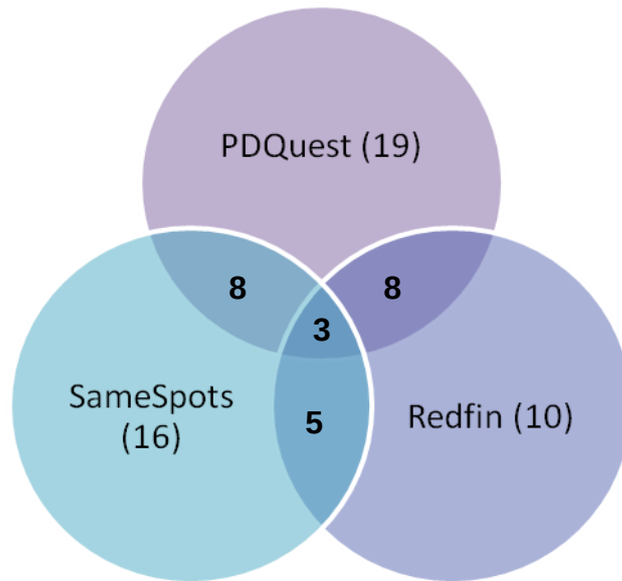


Figure 27.7: Comparative analysis of proteins differentially expressed by *B. fragilis* in semi-defined media without mucin vs mucin Type II enriched media by PDQuest, Redfin and Same Spots softwares using the Student t-test (i.e. based on averages).

Refer to Table 18 for details on the differentially expressed *B. fragilis* proteins in media without mucin vs. mucin Type II enriched media by PDQuest, SameSpots and Redfin softwares using the Student t-test.

Protein identity- <i>B. fragilis</i>	PDQuest analysis	Redfin analysis	Samespots software
Putative thiol peroxidase (6304)		↓ (2.22, 0.0264)	↑(4.2, 0.024)
Outer membrane protein H (6202)	↓	↓ (4.69, 0.0176)	
Putative RNA binding domain (6102)	↓	↓ (3.66, 0.0139)	
SSP 3206 (Not identified)		↓ (3.11, 7.293e-4)	↓ (1.9, 0.003)
Putative thiol peroxidase (4601)	↓	↓ (3.07, 0.0163)	
Malate dehydrogenase (4607)	↓	↓ (3.07, 3.5558e-4)	↓ (3.2, 0.002)
Hypothetical protein BF2494 (1104)	↑		↑ (5.1, 6.672e-004)
50S Ribosomal protein L7/L12 (2102)	↓		
Hypothetical protein BF2494 (2804)	↓		
50S Ribosomal protein L7/L12 (3001)	↓		↓ (2.4, 0.043)
Triose phosphate isomerase (4502)	↓	↓ (2.69, 0.0136)	↓ 4.5, 0.052)
Elongation factor Ts (4702)	↑	↑ (2.34, 0.0250)	↑ (2.4, 0.043)
50S Ribosomal protein L7/L12 (5106)	↓		
Hypothetical protein BF1203 (5205)	↓		↓ (1.5, 0.033)
Lactoylglutathione lyase (6103)	↓	↓ (2.68, 0.0211)	
Hypothetical protein 1203 (5402)	↓		
Hypothetical protein 0301 (6206)	↓		
Phosphoenol pyruvate carboxykinase (6901)	↓		
Fructose biphosphate aldolase (6802)	↓		↓ (3.0, 0.002)
SSP 3904 (Not identified)			↓ (3.7, 0.003)
50S Ribosomal protein L7/L12 (3106)			↓ (1.7, 0.009)
Superoxide dismutase (6206)	↓		↓ (6.9, 0.004)
SSP 5806 (Not identified)			↓ (4.3, 0.028)
SSP 1101 (Not identified)			↓ (6.1, 5.017e-004)
Triose phosphate isomerase (6302)	↓	↓ (6.78, 0.0156)	
SSP 1302 (Not identified)			↓(2.0, 6.287e-004)
Adenylate kinase (4302)			↓(2.6, 0.046)

Table 18: Differentially expressed *B. fragilis* proteins in non mucin vs mucin Type II enriched media by PDQuest and Redfin and SameSpots softwares using the Student t-test (i.e. based on averages) ↑ indicates up-regulation of the protein ↓ indicates down-regulation of the protein. (Fold change, p value) for differentially expressed spots has been indicated in brackets for Redfin and SameSpots softwares. SSP numbers of protein spots are indicated in the first column.

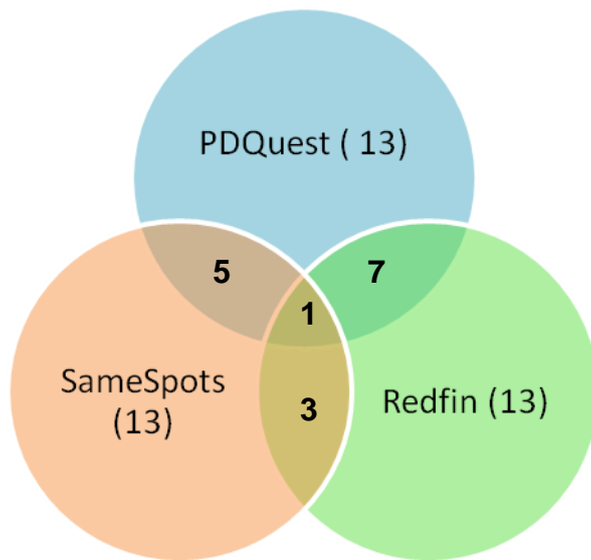


Figure 27.8: Comparative analysis of proteins differentially expressed by *B. fragilis* in semi-defined media without mucin vs mucin Type III enriched media by PDQuest, Redfin and SameSpots softwares using the Student t-test (i.e. based on averages). Refer to the Table 19 for details on the differentially expressed *B. fragilis* proteins in media without mucin vs. mucin Type III enriched media by PDQuest, SameSpots and Redfin softwares using the Student t-test.

Protein identity- <i>B. fragilis</i>	PDQuest analysis	Redfin analysis	Samespots software
Putative thiol peroxidase (6304)	↓	↓ (2.22, 0.0264)	
Outer membrane protein H (6202)	↓	↓ (4.69, 0.0176)	
SSP 3206 (Not identified)		↑ (3.11, 7.293e-4)	↑ (1.9, 0.003)
SSP 1207 (Not identified)		↓ (2.65, 0.0101)	
SSP 3305 (Not identified)		↓ (2.47, 0.0479)	
Malate dehydrogenase (4607)		↓ (3.07, 3.5558e-4)	
Malate dehydrogenase (4608)	↑	↑ (3.24, 0.0431)	↑ (1.3, 0.053)
SSP 5002 (Not identified)		↓ (2.89, 0.0037)	
50S Ribosomal protein L7/L12 (2102)	↓		↓ (2.4, 0.043)
50S Ribosomal protein L7/L12 (5106)	↓		
Lactoylglutathione lyase (6103)	↓	↓ (2.68, 0.0211)	
SSP 3904 (Not identified)			↓ (3.7, 0.003)
Adenylate kinase (4302)			↓ (2.6, 0.046)
SSP 7001 (Not identified)			↓ (3.8, 0.001)
Hypothetical protein BF2494 (1104)	↑		
Hypothetical protein BF1203 (5205)	↓		↓ (1.5, 0.033)
Hypothetical protein BF1203 (5201)	↓		↓ (2.8, 0.024)
Malate dehydrogenase (4705)			↓ (2.1, 0.047)
Putative thiol peroxidase (4701)			↑(4.2, 0.024)
SSP 3702 (Not identified)			↑ (3.6, 0.012)
SSP 1803 (Not identified)	↓		↓ (1.5, 0.049)
SSP 5103 (Not identified)		↓ (2.46, 0.0271)	↓ (1.8, 0.043)
Outer membrane protein precursor (3303)	↓	↓ (5.07, 2.848e-6)	
SSP 2403 (Not identified)	↑	↑ (2.70, 0.0381)	
GrpE protein (2401)	↓	↓ (2.66, 0.0034)	

Table 19: Differentially expressed *B. fragilis* proteins in non mucin vs mucin Type III enriched media by PDQuest and Redfin and SameSpots softwares using the Student t-test (i.e. based on averages) ↑ indicates up-regulation of the protein; ↓ indicates down-regulation of the protein. (Fold change, p value) for differentially expressed spots has been indicated in brackets for Redfin and SameSpots softwares. SSP numbers of protein spots are indicated in the first column.

10.7.7 Analysis using Redfin and SameSpots softwares

Initial analysis of results was performed using the PDQuest software using the Student t-test and hence this has been used as the default method for discussing the results. Following further analysis using the Redfin and SameSpots softwares, the differential expression data from all the three softwares were compared. PDQuest and Redfin software analyses were based on the Mann Whitney U test performed to determine the proteins showing a regulation based on the levels of expression in each individual gel in every growth condition. Analyses using PDQuest, SameSpots and Redfin softwares were based on the Student t-test performed based on the average level of expression of proteins in every growth condition.

The Mann Whitney U test based analysis of proteins showed a down-regulation of elongation factor Ts in mucin Type III enriched media and an up-regulation of the outer membrane surface antigen protein in mucin Type III enriched media in *E. cancerogenus*. Apart from these two proteins, the periplasmic proteins and carbamate kinase proteins were up-regulated in mucin Type III enriched media whereas SSP number 7001 protein which was unidentifiable from mass spectrometry was up-regulated in mucin Type II enriched media in *E. cancerogenus*. The Student t-test based analyses of proteins from all the 3 softwares showed an up-regulation of outer membrane surface antigen protein in both mucin Type II and III enriched media and an up-regulation of phosphoglycerate kinase in mucin Type II enriched media in *E. cancerogenus*. In mucin Type III enriched media in *E. cancerogenus*, the elongation factor Ts protein was found to be down-regulated.

From both the tests, it could be identified that the outer membrane surface antigen was up-regulated in mucin Type III enriched media in *E. cancerogenus* indicating that this protein may be a potential virulence factor whose expression may be associated with the presence of mucin in the growth media.

The Mann Whitney U test based analysis of proteins in *B. fragilis* showed a down-regulation of the RNA binding protein, thiol peroxidase, malate dehydrogenase and triose phosphate isomerase proteins in mucin Type II enriched media. In mucin Type III enriched media, down-regulation of the outer membrane protein H, thiol peroxidase and triose phosphate isomerase proteins was observed in *B. fragilis*.

The Student t-test based analyses of proteins from all the 3 softwares showed a down-regulation of malate dehydrogenase, triose phosphate isomerase and an up-regulation of elongation factor EFTs in mucin Type II enriched media in *B. fragilis*. In mucin Type III enriched media only malate dehydrogenase was found to be up-regulated in all the 3 softwares.

From both the tests, it could be identified that malate dehydrogenase was found to be up-regulated in mucin Type III enriched media and down-regulated in mucin Type II enriched media suggesting that they may be existing as isoforms. Triose phosphate isomerase was also found to be down-regulated in mucin Type II enriched media suggesting that these proteins may be associated with the pathogenicity of *B. fragilis* when grown in the presence of mucin.

10.8 Liquid chromatography - Mass spectrometric analysis of protein spots

A total of 132 *E. cancerogenus* protein spots were analysed using LC-MS of which 98 spots were identified and 34 spots were found to be differentially expressed, according to PDQuest software analysis using Student t-test. Of the 34 spots that showed a regulation of expression, 23 spots were identifiable and the remaining 11 were unidentifiable. For *B. fragilis*, a total of 106 protein spots were analysed of which 73 spots were identified and 45 were found to be differentially expressed. Of the 45 spots that showed differential expression, 31 spots were identifiable and the remaining 14 were unidentifiable. Figure 28 shows the differentially expressed proteins in the reference master gel for *E. cancerogenus* and Figure 29 shows the differentially expressed proteins in the reference master gel for *B. fragilis*. Refer to Appendix sections G2 and G3 for details on the peptide match scores and identities of proteins using the Mascot search for both *E. cancerogenus* and *B. fragilis*.

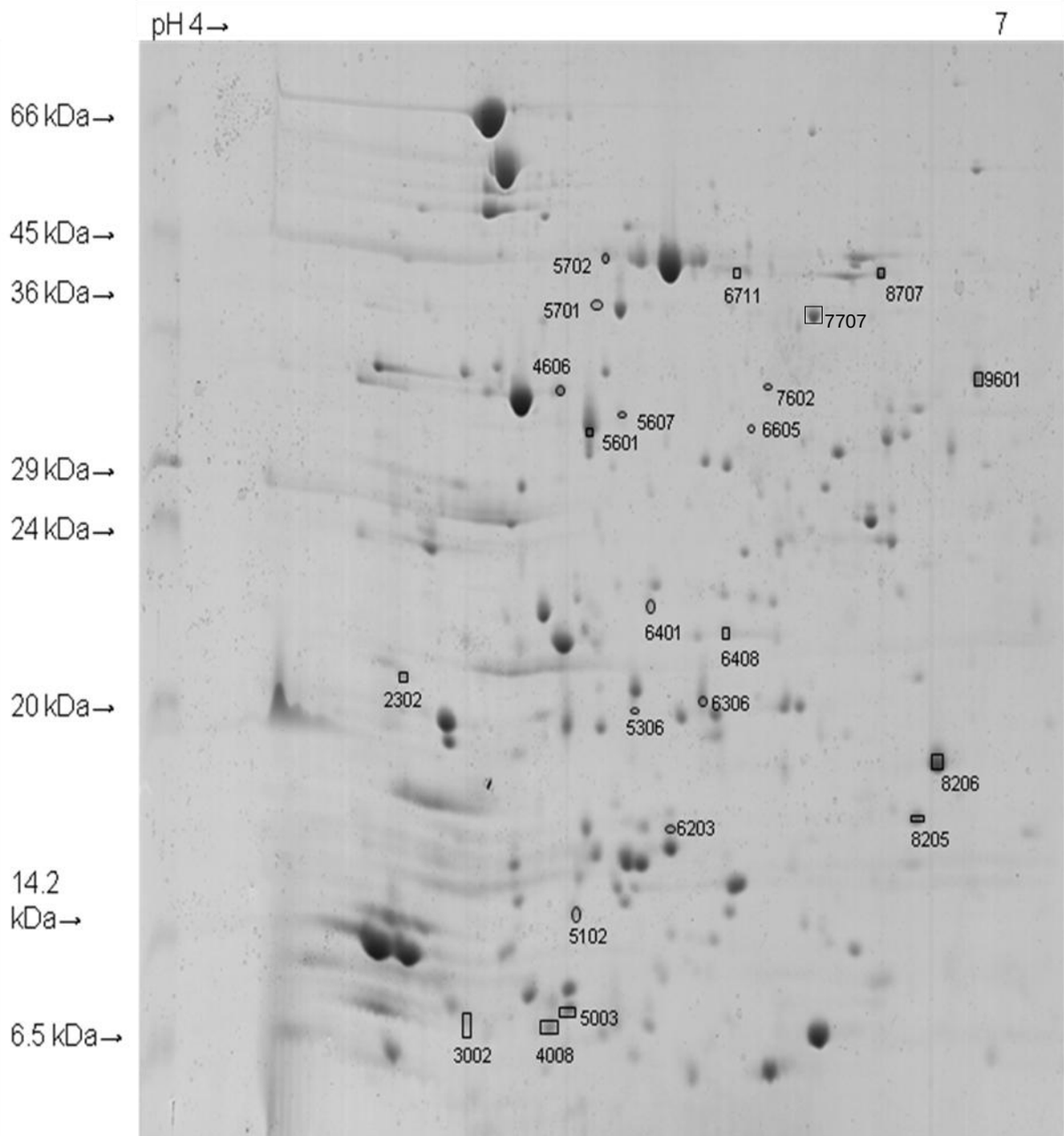


Figure 28: The SSP numbers of differentially expressed protein spots on the reference master gel of *E. cancerogenus* based on the PDQuest software analysis. ○ Indicates up-regulation and □ indicates down-regulation in mucin enriched media

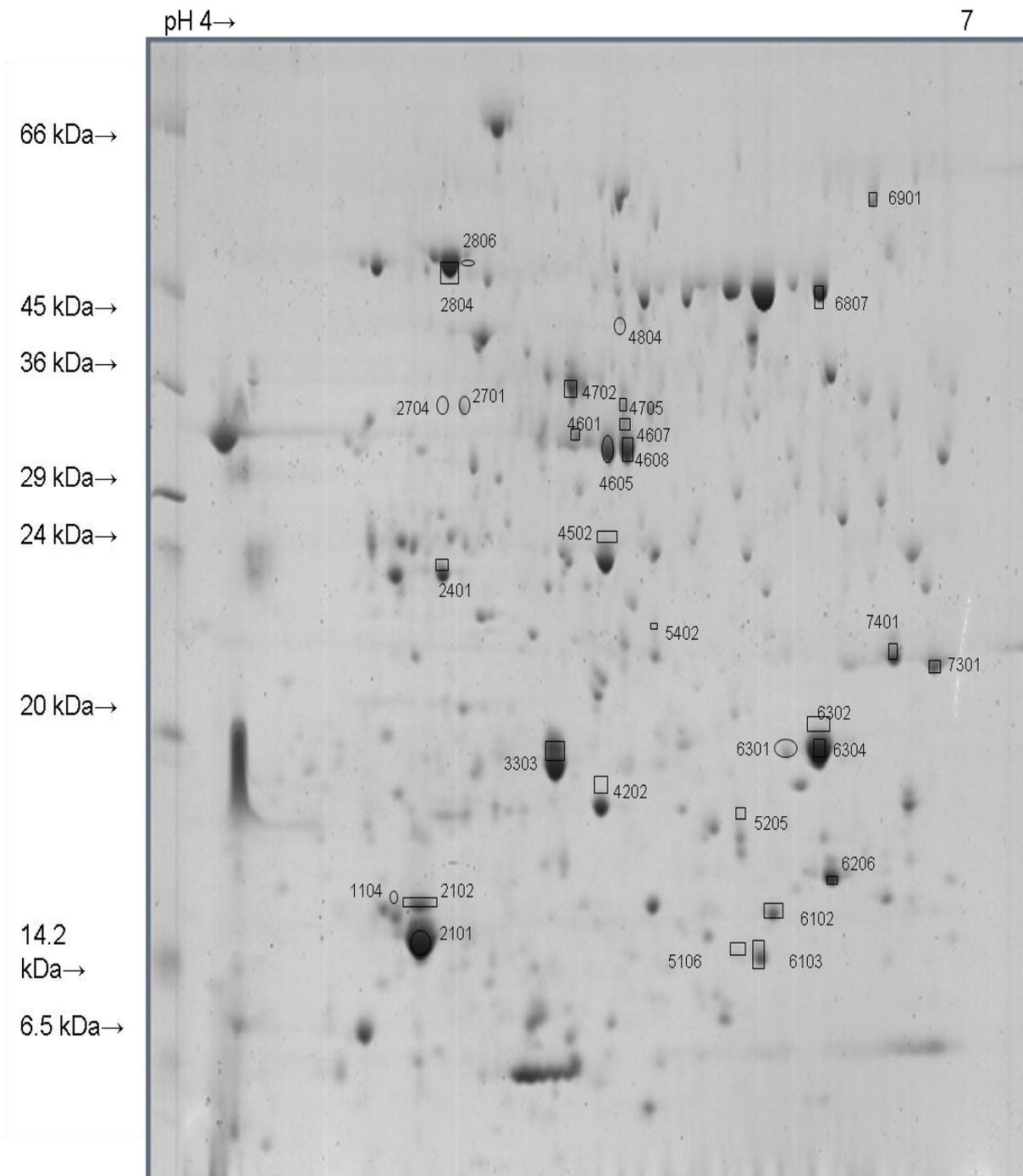


Figure 29: The SSP numbers of differentially expressed protein spots on the reference master gel of *B. fragilis* based on the PDQuest software analysis.

○ Indicates up-regulation and □ indicates down-regulation in mucin enriched media

Tables 20 and 22 represent the list of proteins identified, including the spots that were found to be differentially expressed in *E. cancerogenus* and *B. fragilis* respectively. Tables 21 and 23 represent the list of proteins identified, excluding the spots that were found to be differentially expressed in *E. cancerogenus* and *B. fragilis*.

S.No	SSP number	Identity of the protein	Function	Regulation	Peptide match score/ number of peptides	Fold change (PDQuest)	Accession number
1	2302	Elongation factor gre A	Unknown function	↓ mucin Type III media	125/3	5.9	gi 261342624
2	3002	Hypothetical protein 03947	Post translational modifications, protein turnover and chaperones	↓ mucin media	101/2	6.9	gi 261339586
3	4008	Autonomous glycy radical cofactor GrcA	General function prediction only	↓ mucin Type III media	150/4	1.7	gi 261340878
4	4606	Omp A	Cell wall, membrane, envelope biogenesis	↑ mucin media	84/2	2.4	gi 261339284
5	5003	Hypothetical protein 17757	Protein function unknown	↓ mucin Type III media	126/2	2.2	gi 261342318
6	5102	Universal stress protein A	Signal transduction mechanisms	↑ mucin Type III media	195/7	7.2	gi 261341988
7	5306	Periplasmic protein	Function unknown	↑ mucin Type III media	130/4	5.5	gi 261339031
8	5601	Elongation factor Ts	Translation, ribosomal structure and biogenesis	↓ mucin Type III media	376/8	6.5	gi 261338832
9	5607	Carbamate kinase	Amino acid transport	↑ mucin Type III	355/6	4.9	gi 261339130

			and metabolism	media			
10	5701	Phosphoglycerate kinase	Carbohydrate transport and metabolism	↑ mucin Type II media	684/13	4.7	gi 261342323
11	5702	Phosphopyruvate hydratase	Carbohydrate transport and metabolism	↑ mucin Type III media	238/4	3.8	gi 261342208
12	6203	Outer membrane protein surface antigen X	Cell wall, membrane, envelope biogenesis	↑ mucin media	129/2	2.3	gi 261341400
13	6306	Hypothetical protein 20877	General function prediction only	↑ mucin Type III media	91/3	7.5	gi 261342932
14	6401	Cytidylate kinase	Nucleotide transport and metabolism	↑ mucin Type III media	115/3	2.4	gi 261339244
15	6408	Periplasmic protein disulphide isomerase I	Energy production and conversion	↓ mucin Type II media	189/5	6.6	gi 261342934
16	6605	2-dehydro-3-deoxyphosphooctonate aldolase	Amino acid transport and metabolism	↑ mucin Type III media	144/3	6.5	gi 261340146
17	6711	Elongation factor Tu	Translation, ribosomal structure and biogenesis	↓ mucin Type III media	246/7	3.3	gi 261341842
18	7602	Acetoin reductase	Secondary metabolites	↑ mucin Type II media	298/5	1.5	gi 261341164

			biosynthesis, transport and catabolism, signal transduction mechanisms				
19	7707	Gamma glutamyl phosphate reductase	Energy production and conversion	↓ mucin Type II media	97/2	4.7	gi 261340945
20	8205	yfaZ family protein	Unknown function	↓ mucin Type III media	195/3	1.8	gi 261339072
21	8206	Ecotin	Cell wall, membrane, envelope biogenesis	↓ mucin media	147/5	8.9	gi 261340622
22	8707	tolB translocation protein	Intracellular trafficking, secretion and vesicular transport	↓ mucin Type II media	271/7	2.6	gi 261341330
23	9601	Omp A	Cell wall, membrane, envelope biogenesis	↓ mucin Type II media	152/3	5.7	gi 261339284

Table 20: Table represents *E. cancerogenus* proteins that have been identified to be expressed in all three conditions with any up or down-regulation using PDQuest software. Regulation in mucin media in the table refers to both mucin Type II and Type III enriched media.

S.No	SSP number	Identity of the protein	Function	Regulation	Peptide match score/ number of peptides
1	1108	50S ribosomal protein L7/L12	Translation, ribosomal structure and biogenesis	None	155/5
2	1703	Porin 02457	Cell wall, membrane and envelope biogenesis	None	55/2
3	2001	Hypothetical protein 03856	Unknown function	None	116/8
4	2104	50S ribosomal protein L7/L12	Translation, ribosomal structure and biogenesis	None	134/6
5	2305	PTS system II A subunit	Carbohydrate transport and metabolism	None	237/7
6	2403	Heat shock protein grpE	Post translational modifications, protein turnover and chaperones	None	77/2
7	2506	Heat shock protein grpE	Post translational modifications, protein turnover and chaperones	None	60/2
8	3808	Trigger factor protein	Post translational modifications, protein turnover and chaperones	None	573/12
9	3809	Trigger factor protein	Post translational modifications, protein	None	71/2

			turnover and chaperones		
10	3901	Hypothetical protein 00599, Dnak chaperone protein	Heat shock proteins	None	560/9
11	3902	Hypothetical protein 00599, Dnak chaperone protein	Heat shock proteins	None	561/12
12	4101	Hypothetical protein 03459, Autonomous glycy radical cofactor	General function prediction only	None	185/3
13	4202	Hypothetical protein 03416, 5-methylaminomethyl-2-thiouridylate methyltransferase	Translation, ribosomal structure and biogenesis	None	126/3
14	4401	Peroxiredoxin 01092	Post translational modifications, protein turnover and chaperones	None	191/4
15	4402	00804 inorganic pyrophosphatase	Energy production and conversion	None	71/2
16	4505	Omp II, 01735	Cell wall, membrane, envelope biogenesis	None	278/5
17	4602	Omp A	Cell wall, membrane, envelope biogenesis	None	468/16
18	4701	Putative glutathione peroxidase	Post translational modification, protein turnover,	None	137/4

			chaperones		
19	4812	Porin 02457	Cell envelope biogenesis	None	84/2
20	4814	Tol C protein	Intracellular trafficking, secretion and vesicular transport	None	175/4
21	4901	Molecular chaperone DnaK	Post translational modification, protein turnover, chaperones	None	60/2
22	5002	01181, YbaB family, hypothetical protein	Protein function unknown	None	166/3
23	5107	Riboflavin synthase subunit β , 01103 hypothetical protein, 6,7, dimethylribityllumazine synthase	Biosynthesis of riboflavin	None	265/4
24	5207	DNA binding protein, 02636	General function prediction only	None	218/8
25	5305	Peptidoglycan binding domain, 01554	Cell wall degradation	None	111/3
26	5401	Inorganic pyrophosphatase, 00804	Energy production and conversion	None	160/5
27	5403	Inorganic pyrophosphatase, 00804	Energy production and conversion	None	185/4
28	5405	Inorganic pyrophosphatase	Energy production and conversion	None	235/6
29	5406	Inorganic	Energy production and	None	163/5

		pyrophosphatase	conversion		
30	5604	Elongation factor Ts, 00441	Translation, ribosomal structure and biogenesis	None	259/5
31	5704	Phosphoglycerate kinase	Carbohydrate transport and metabolism	None	638/15
32	5705	Enolase, 04083	Carbohydrate transport and metabolism	None	483/14
33	6001	ABC type sugar transport system	Carbohydrate transport and metabolism	None	85/2
34	6202	OmpX, 01529	Cell wall, membrane, envelope biogenesis	None	247/3
35	6307	F0F1 ATP synthase subunit B	Energy production and conversion	None	186/4
36	6603	01449, hypothetical protein (DNA uptake, outer membrane assembly)	Function unknown	None	328/6
37	6702	Phosphopyruvate hydratase, 04083	Carbohydrate transport and metabolism	None	479/9
38	7301	Single stranded DNA binding protein	General function prediction only	None	219/4
39	7401	03228, hypothetical protein, TRX like ferredoxin family, NADH: ubiquinone oxidoreductase subunit E	Energy production and conversion	None	74/1

40	7403	02508, hypothetical protein, phage shock protein psp A	Lipid transport and metabolism	None	76/2
41	7501	Hypothetical protein, 2,5 diketo D gluconate reductase A	Biosynthesis of vitamin C, carbohydrate metabolism	None	152/4
42	7704	01034, Aldehyde dehydrogenase	Energy production and conversion	None	87/2
43	7710	03735, Flavodoxin	Energy production and conversion	None	247/7
44	8602	04209, protein of unknown function	Unknown function	None	126/3
45	8604	Phosphopyruvate hydratase	Carbohydrate transport and metabolism	None	635/16
46	8610	01936, Aldose-1-epimerase	Carbohydrate transport and metabolism	None	115/2
47	8611	03358, Dihydrodipicolinate synthase	Amino acid transport and metabolism	None	78/2
48	8704	00565, Peptidyl-prolyl cis trans isomerase SurA	Post translational modification, protein turnover and chaperones	None	223/4

Table 21: Table represents *E. cancerogenus* proteins that have been identified to be expressed in all three conditions without regulation using PDQuest software

S. No	SSP number	Identity of the protein	Function	Regulation	Peptide match score/ number of peptides	Fold change (PDQuest)	Accession number
1	1104	BF2494, Hypothetical protein	Mediates protein-protein interactions	↑ mucin media	366/8	3.1	gi 60681974
2	2101	50S ribosomal protein L7/L12	Translation, ribosomal structure and biogenesis	↓ mucin Type II media	265/4	1.5	gi 60683451
3	2102	50S ribosomal protein L7/L12	Translation, ribosomal structure and biogenesis	↓ mucin media	347/30	9.9	gi 60683451
4	2401	GrpE protein	Post translational modification, protein turnover and chaperones	↓ mucin Type III media	83/2	8.9	gi 60681311
5	2701	ATP synthase subunit E	Energy production and conversion	↑ mucin Type III media	197/4	5.3	gi 60682222
6	2704	ATP synthase subunit E	Energy production and	↑mucin Type III media	175/5	2.9	gi 60682222

			conversion				
7	2804	TRP repeat containing protein, 2494	Mediates protein-protein interactions	↓ mucin Type II media	393/7,366/8	8.8	gi 60681974
8	2806	50S ribosomal protein L7/L12	Translation, ribosomal structure and biogenesis	↑mucin Type III media	165/6	2.6	gi 60683451
9	3303	Omp precursor	Cell wall, membrane, envelope biogenesis	↓ mucin media	184/8	8.3	gi 60680025
10	4202	Omp precursor	Cell wall, membrane, envelope biogenesis	↓ mucin media	473/13	4.4	gi 60680025
11	4502	Triose phosphate isomerase	Carbohydrate transport and metabolism	↓ mucin Type II media	183/4	9.3	gi 60683174
12	4601	Putative thiol peroxidase	Post translational modification, protein turnover and chaperones	↓mucin media	112/2	9.3	gi 60682249

13	4605	Thioredoxin	Post translational modification, protein turnover and chaperones	↓mucin Type II media	140/5	4.2	gi 60680203
14	4607	Malate dehydrogenase	Amino acid transport and metabolism	↓ mucin Type II media	159/4	7	gi 60683199
15	4608	Malate dehydrogenase	Amino acid transport and metabolism	↑mucin Type III media	85/2	3.7	gi 60683199
16	4702	Elongation factor Ts	Translation, ribosomal structure and biogenesis	↓ mucin Type II media	331/7	7.3	gi 60683229
17	4705	Malate dehydrogenase	Amino acid transport and metabolism	↓mucin media	159/4	3.7	gi 60683199
18	4804	Malate dehydrogenase	Amino acid transport and metabolism	↑mucin Type III media	171/5	3.5	gi 60683199
19	5106	50S ribosomal protein L7/L12, Hypothetical	Translation, ribosomal structure and	↓ mucin media	93/1, 60/3	1.4	gi 60683451

		protein 2161	biogenesis				
20	5205	Hypothetical protein 1203, putative anti-sigma factor	Signal transduction mechanisms	↓mucin media	403/7	9.1	gi 60680695
21	5402	Hypothetical protein 1203	Signal transduction mechanisms	↓ mucin Type II media	361/9	8.7	gi 60680695
22	6102	Putative RNA binding protein	General function prediction only	↓ mucin Type II media	159/4	7.3	gi 60682207
23	6103	Lactoylglutathione lyase	Amino acid transport and metabolism	↓mucin media	195/3	8.7	gi 60682596
24	6206	Superoxide dismutase (Fe), hypothetical protein 0301	General function prediction only	↓mucin Type II media	100/3, 136/2	8.1	gi 60679842
25	6301	Thioredoxin	Post translational modification, protein turnover, chaperones	↑ in mucin Type III media	87/2	5.6	gi 60680203
26	6302	Triose phosphate isomerase	Carbohydrate transport and	↓ mucin media	87/2	9.8	gi 60683174, gi 60682249

			metabolism				
27	6802	Fructose-bisphosphate aldolase	Carbohydrate transport and metabolism	↓mucin media	196/6	8.3	gi 60682607
28	6807	Conserved hypothetical protein	Unknown function	↓ mucin media	610/14	4.1	gi 60491969
29	6901	Phosphoenol pyruvate carboxykinase	Carbohydrate transport and metabolism.	↓ mucin Type II media	663/14	9.1	gi 53715725
30	7301	FeS Superoxide dismutase	Inorganic ion transport and metabolism	↓ mucin media	466/37	9.6	gi 60682036
31	7401	Ribosome recycling factor	Translation, ribosomal structure and biogenesis	↓mucin media	235/5	9.2	gi 60680182

Table 22: *B. fragilis* (Table represents proteins that have been identified to be expressed in all three conditions with any up or down regulation using PDQuest software). Regulation in mucin media in the table refers to both mucin Type II and Type III enriched media.

S. No	SSP number	Identity of the protein	Function	Regulation	Peptide match score/ number of peptides
1	0605	BF2494, TRP repeat containing protein similar to tetratricopeptide repeat family	Mediates protein-protein interactions	None	45/2
2	1208	50S ribosomal protein L7/L12	Translation, ribosomal structure and biogenesis	None	141/4
3	2106	50S ribosomal protein L7/L12	Translation, ribosomal structure and biogenesis	None	121/2
4	3001	50S ribosomal protein L7/L12	Translation, ribosomal structure and biogenesis	None	93/1
5	3105	50S ribosomal protein L7/L12	Translation, ribosomal structure and biogenesis	None	160/2
6	3302	Co-chaperonin GroES	Post translational modification, protein turnover and chaperones	None	48/1
7	3401	FeS superoxide dismutase	Inorganic ion transport and metabolism	None	211/5
8	3403	FeS superoxide dismutase	Inorganic ion transport and metabolism	None	466/38
9	3902	Chaperone dnaK	Post translational modification,	None	345/10

			protein turnover and chaperones		
10	4302	Adenylate kinase	Nucleotide transport and metabolism	None	105/2
11	4408	GrpE protein	Post translational modification, protein turnover and chaperones	None	98/3
12	4701	Putative thiol peroxidase	Post translational modification, protein turnover and chaperones	None	172/5
13	5201	Hypothetical protein 1203, putative anti-sigma factor	Signal transduction mechanisms	None	133/3
14	5204	Hypothetical protein 1203	Signal transduction mechanisms	None	361/9
15	5808	Enolase	Carbohydrate transport and metabolism	None	469/8
16	5810	Phosphopyruvate hydratase	Carbohydrate transport and metabolism	None	354/6
17	6202	Putative OmpH	Cell wall, membrane, envelope biogenesis	None	60/1
18	6304	Thiol peroxidase	Carbohydrate transport and metabolism	None	484/18
19	6709	Hypothetical protein 1004	Protein function unknown	None	358/9
20	6801	Chaperonin groEL	Post translational	None	703/12

			modification, protein turnover, chaperones		
21	6803	Enolase	Carbohydrate transport and metabolism	None	179/4

Table 23: *B. fragilis* (Table represents proteins that have been identified to be expressed in all three conditions without any up or down regulation using PDQuest software)

MASCOT search results with the peptide scores and identity of the protein spots can be seen in Appendix G.

This study was able to identify 62% of the differentially expressed proteins using LC-MS analysis in *E. cancerogenus* and 67% of the differentially expressed proteins were identified in *B. fragilis*.

10.9 Mass spectrometric analysis of proteins

Formic acid (FA) was used in the preparation of buffers for LC-MS analysis instead of trifluoroacetic acid (TFA). This is because TFA has known negative effects which include suppression of sensitivity and ionisation of protein samples by binding and pairing to the basic groups of peptides when ionised in the gaseous phase. At a concentration of 0.1% (w/v) or more, TFA is capable of preventing spray formation or nebulisation due to the generation of a high surface tension during the ionisation process (Anon, 2002). Also, better resolution of peaks has been observed during the separation of proteins using liquid chromatography (Anon, 2002). Hence this buffer was used in our study.

Sample carry over was another potential problem that was found to influence the results obtained from the MS data. This was overcome by performing water runs in between the sample runs to remove any of the remaining peptides present in the monolith column.

Concentration of protein present in a spot was another important issue for determining its identity. Determining the concentration of the protein to be loaded to identify it from the LC-MS analysis was essential since the differential protein expression profile contained several different proteins of varying intensities. Since at least six replicates of each gel condition was generated, each of the spot was processed separately and pooled together for analysis. Initial trial experiments were performed by cutting out six sets of faint, medium and dark spots. It was observed that three or four faint spots pooled together gave a positive identification for the majority protein on MS analysis. On average, 4 spots were pooled together to detect fainter spots whereas 2 or 3 replicate spot samples were pooled together to detect dark and medium spots respectively.

Some of the spots remained undetectable because of the very low concentrations or due to sample loss during processing. Pressure variations in LC set up and needle blockages in the MS set up resulted in loss of some of the samples too. The autosampler that transferred the sample on to the column used an injection needle to pick up the sample from the multi-well microtitre plate and care had to be taken to avoid the pipetting of air bubbles along with the sample into the wells since this sometimes resulted in the blockage of the system. Frequent ultra-sonication and cleaning of the needles and the shield in the MS system helped to prevent blockages.

The possibility of adjacent spots overlapping each other on the gel suggests that the cutting out of spots need to be very accurate. It was also a possibility that some of the adjacent overlapping spots were isoforms resolving poorly. Some of the adjacent overlapping spots were identified to be the same protein. The inability to obtain a positive identity for some protein spots may be attributed to the fact that proteins were lost during processing, freeze drying or experimental errors. Several steps involved the solubilisation, freeze drying and resuspension of proteins which added to the loss of proteins.

In order to avoid random matching of measured masses to peptides in the MASCOT sequence database, the searches were restricted to the grouping of 'Eubacteria.' Random matching results could also provide false positive results and hence certain criteria were set up to segregate false positives from correct identifications. This included selection of results where the protein score was more than 40 and at least 2 peptides matched to the query sequence.

A total of 23 differentially expressed proteins were identified out of 34 in *E. cancerogenus* and 31 proteins were identified out of 45 in *B. fragilis*. The default statistical analysis used in PDQuest was the Student t-test. More than 60% of the differentially expressed proteins were identified in both cases but a substantial amount of the protein samples were lost due to processing or experimental errors.

The LC-MS spectra generated from the protein spots were searched through the MASCOT server in the NCBI and SWISS PROT databases. It was found that the NCBI prioritised the *B. fragilis* strain YCH76 over the NCTC 9343 strain. The exact reason for this is unknown. The significant hits in the YCH76 strain prompted us to

perform NCBI blast searches with its NCTC 9343 counterparts and it was observed that the matched peptide sequences were identical in both cases. There is a possibility that the unique classification, categorisation and annotation of sequences in the NCBI database is responsible for the results obtained. Similar searches in the SWISS PROT database produced matching peptide sequences in the NCTC 9343 strain but the ion scores were found to be lower. This may be because SWISS PROT database has a manually annotated protein sequence record (Mizrachi, 2002). Previous work by other scientists have suggested the presence of similar problems in data analysis (Ingrell, 2009). Refer to the second section of the discussion for further details.

Differential expression of virulence associated proteins like neuraminidases, sialidases, mucinases and glycoside hydrolases were expected to be evident from the proteomic experiments but surprisingly, none of these proteins were identified. *B. fragilis*, known to exhibit a well established catabolic pathway in its activity against sialic acids by producing sialidases did not express the protein in the presence of mucin, unless it was expressed in undetectable amounts. The reason behind repression of these enzymes may be due to the presence of glucose in the semi-defined growth media in addition to mucin which may be inducing a catabolite repression.

It may be interesting to perform specific assays to determine the presence of these enzymes in the cell free extracts since they have not been detected from the proteomics experiments.

10.10 Expression of proteins involved in translation, ribosomal structure and biogenesis

The ribosome recycling factor (RRF) protein was found to be down-regulated in mucin enriched media in *B. fragilis* but no regulation was observed in *E. cancerogenus*. The protein plays a vital role in the recycling of ribosomes after completion of protein synthesis. The two main signal transduction associated proteins required for the release of ribosomes from the mRNA are elongation factor (EF-G) and the ribosome recycling factor (originally known as the ribosome recycling factor (RRF)); (Wilson *et al.*, 2005). RRF dissociates ribosomes from mRNA after termination of translation and is essential for bacterial growth (Janosi *et al.*, 1994). The activity of this protein is important for the availability of free ribosomes for further continuation of protein synthesis. These proteins play an important role in inducing immune responses within the host during infection by activating the production of IgG antibodies (Cassataro *et al.*, 2007). This suggests that these proteins may be up-regulated during adverse growth conditions but has been found to be down-regulated in mucin enriched media in *B. fragilis*. Previous studies have shown that the growth of *Streptococcus oralis* in acidic conditions resulted in an up-regulation of these surface-associated proteins (Wilkins *et al.*, 2003).

The 50S ribosomal protein L7/12 was found to be down-regulated in mucin enriched media in *B. fragilis* even though no regulation was observed in *E. cancerogenus*. The main functions of the 50S ribosomal subunit L7/L12 is mRNA directed protein synthesis, increase in polypeptide synthesis and reduction of missense error rate. Variable expression levels of 50S ribosomal subunit L7/L12 were observed in mucin media in *B. fragilis* which could be related to the possible existence of isoforms even though this phenomenon has not been observed in bacteria. But a general down-regulation of the protein was observed in mucin enriched media except SSP number 2806 which shows an up-regulation in expression.

10.11 Expression of proteins involved in transcription

Elongation factor Ts was found to be down-regulated in mucin enriched media in *B. fragilis* and *E. cancerogenus*. Interestingly, it has been known from previous studies in *S. pneumoniae* that opaque invasive variants of the bacteria were found to exhibit a down regulation of elongation factor Ts when compared to the less invasive variants that expressed higher amounts of the elongation factor (Overweg *et al.*, 2000). The elongation factor Ts plays an important role in protein synthesis by acting as a nucleotide exchange factor that is required for the regeneration of elongation factor Tu (EF-Tu) from its inactive EF-Tu-GDP form to the active form EF-Tu-GTP. This enables EF-Tu to interact with the next incoming amino acid (Hwang & Miller, 1985).

Elongation factor EF-Tu was found to be down-regulated in mucin enriched media in *E. cancerogenus* even though the protein showed no differential expression in *B. fragilis*. EF-Tu is an intracellular protein that contains a number of other proteins associated with it and is involved in translation elongation factor activity and GTP binding. It plays an important role in the translation process by initiating the selection and binding of the cognate amino-acyl t-RNA to the acceptor A site (Pape *et al.*, 1998). They may be associated with the translation and biogenesis of ribosomal structures apart from being involved in transcription. Differential expression studies in mycobacteria showed the down-regulation of these proteins until they came in contact with the host macrophages which indicates that the expression of these proteins may be triggered by the presence of a host immune response (Monahan *et al.*, 2001).

Putative RNA binding protein domains were down-regulated in mucin enriched media in *B. fragilis*. These proteins play an important role in regulating transcription termination; protect the mRNA from degradation during unfavourable growth conditions like cold shock and nucleic acid recognition (Stulke, 2002). They are capable of stopping translation by preventing the binding of mRNA to ribosomes and can control the secondary structure formation of mRNA by promoting or preventing

translation initiation thereby modulating metabolic processes. They induce antitermination of carbon and nitrogen catabolic genes to suit the survival of bacteria in the growth media and the CRM domain forms the conserved region of RNA binding protein with 100 amino acids involved in nucleic acid recognition (Barkan *et al.*, 2007). RNA binding attenuation protein *trp*, was found to be associated with the production of a putative efflux protein in *B. subtilis* suggesting that it may be a potential virulence factor (Yakhnin *et al.*, 2006). Down-regulation of these proteins in mucin enriched media suggests that the adaptability of *B. fragilis* in this media is much better when compared to the control semi-defined media.

10.12 Expression of proteins involved in secondary metabolite biosynthesis, transport and catabolism

Acetoin reductase, a very essential enzyme involved in acetoin catabolism and acetoin dehydrogenation activity has been found to be up-regulated in media enriched with mucin Type II but down-regulated in media enriched with mucin Type III in *E. cancerogenus*. This protein was not identified in *B. fragilis*. The enzyme catalyses the conversion of acetoin and NADH to 2, 3 butanediol and NAD (+) which is a characteristic of anaerobic fermentation processes. This is also used in the commercial production of 2,3 butanediol (Xiao & Xu, 2007). A study in *B. subtilis* has revealed that this cytoplasmic protein is expressed under anaerobic conditions when the cells enter the late log phase where the production of butanediol can be observed whereas the early log phase shows accumulation of acetoin (Nicholson, 2008). The production of acetoin is essential for maintaining the pH of the media where it compensates for the acidic end products that are produced when glucose is present as a source of carbon in the media and this has been observed in *V. cholerae* (Kovacikova *et al.*, 2005). This suggests that the growth stage of the bacteria during media harvest play a very important role in their protein expression.

10.13 Inorganic ion transport and metabolism associated protein expression

The enzyme superoxide dismutase has been identified to be down-regulated in mucin enriched media in *B. fragilis*. This protein has not been identified in *E. cancerogenus*. They are housekeeping enzymes that respond or confer resistance to oxidative stress in bacteria. Their location is periplasmic and catalyses the dismutation of superoxide into oxygen and hydrogen peroxide thus functioning as an antioxidant in cells exposed to oxygen. They are produced to allow the host to outcompete any aerobic flora or reduce reactive oxygen species. They destroy radicals that are commonly produced within cells and are toxic to biological systems. They occur as homodimers or homotetramers (Messerschmidt *et al.*, 2001). Previous studies have shown that the expressions of these enzymes are encoded by the *sodA* and *sodB* genes. The *sodA* gene was activated when the growth conditions shifted from anaerobic to aerobic conditions and the *sodB* gene was switched off under aerobic conditions (Matsumura *et al.*, 1993). Growth of bacteria in media containing glucose and phosphate has been known to induce autoxidation (Carlsson *et al.*, 1978) but a down-regulation of the enzyme results in a reduction in control of autoxidation and formation of superoxide radicals indicating that *B. fragilis* has a poorer survival rate in the presence of mucin.

10.14 Expression of proteins involved in cell wall, membrane and envelope biogenesis

Several outer membrane proteins have been differentially expressed in mucin enriched media with both *B. fragilis* and *E. cancerogenus*. Outer membrane protein A was up-regulated while outer membrane protein II was found to be down-regulated in *E. cancerogenus*. The outer membrane surface antigen X was found to be up-regulated in both mucin Type II and III enriched media in *E. cancerogenus*. In *B. fragilis*, two different spots were found to contain the same outer membrane protein precursors which were down-regulated in both cases in mucin enriched media. This could possibly indicate that the protein is present as isoforms in *B. fragilis* but no information is available in literature regarding the existence of isoforms in outer membrane proteins of bacteria. Refer to Figs. 30, 31 and 32 which indicate spots that expressed the same protein in *B. fragilis* and *E. cancerogenus*.

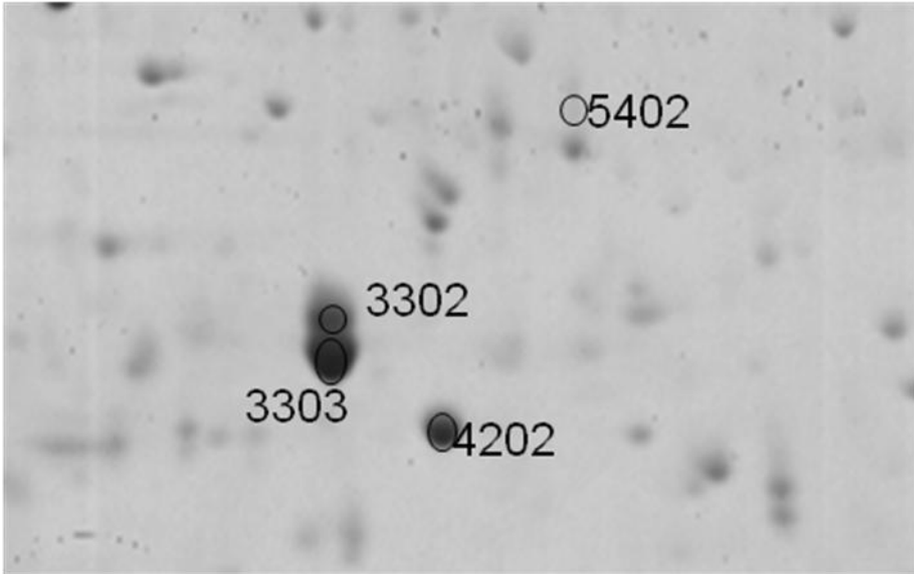


Figure 30: Figure showing the differentially expressed spots of outer membrane proteins in *B. fragilis*. SSP numbers 3302 and 5402 were identified to be outer membrane protein H and SSP numbers 3303 and 4202 were identified to be outer membrane protein precursors.

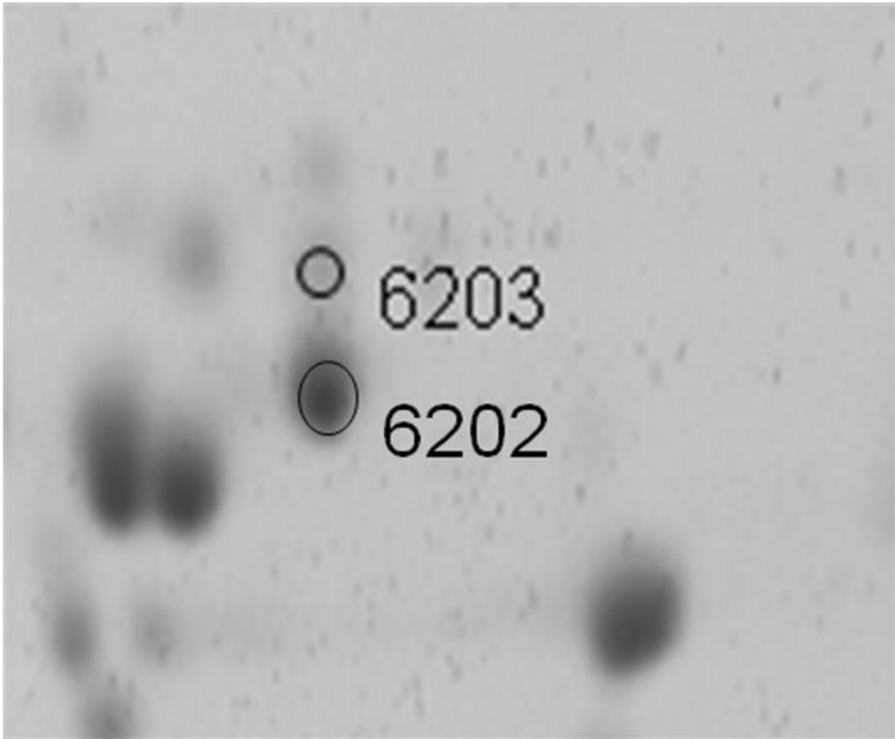


Figure 31: Figure showing the differentially expressed spots of outer membrane protein H in *B. Fragilis*

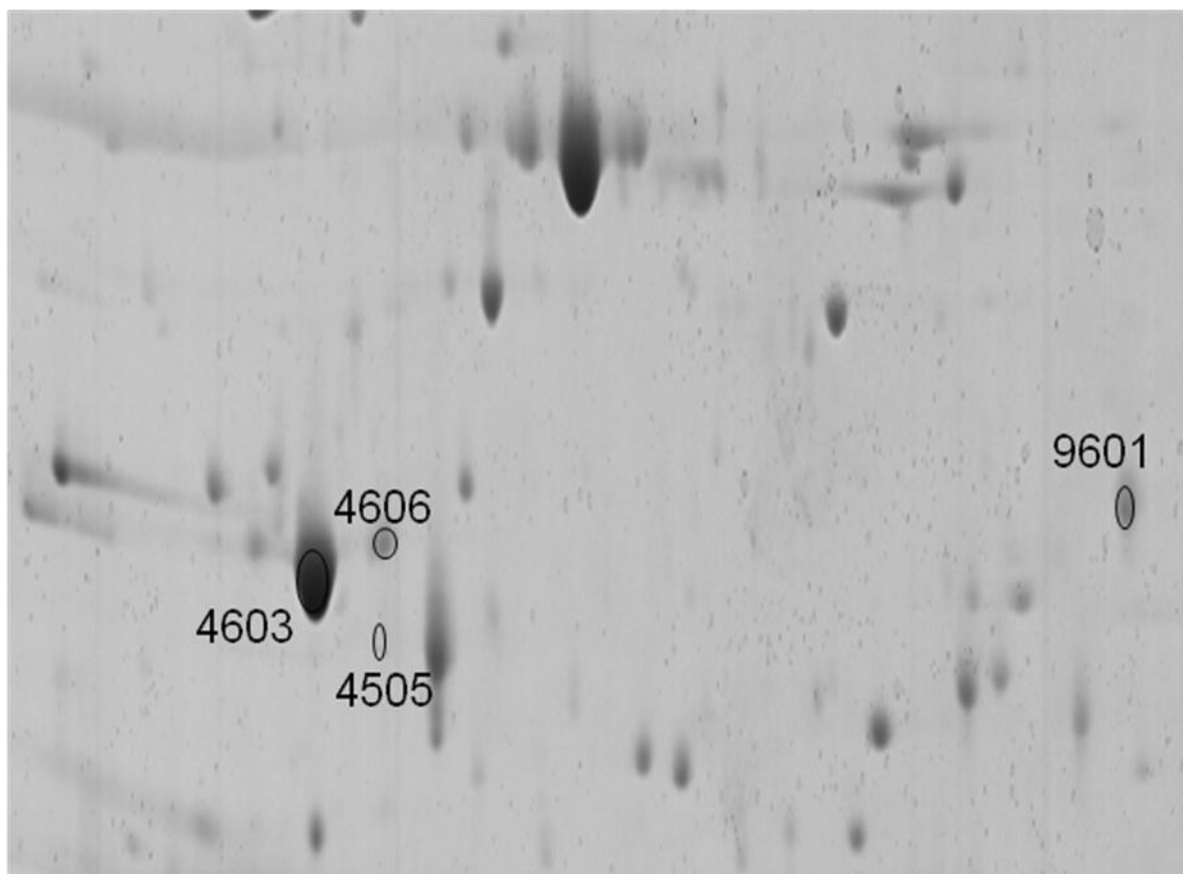


Figure 32: Figure showing the differentially expressed spots of outer membrane proteins in *E. cancerogenus*. SSP numbers 4606, 4505 and 9601 were identified to be outer membrane protein II and SSP numbers 4602 and 4603 were identified to be outer membrane protein A.

Outer membrane proteins are porins involved in transport of solutes, are immunogenic in nature and occur as epitopes on the bacterial surface exhibiting antigenic properties (Luo *et al.*, 1997). Outer membrane protein H have been known to be up-regulated during periods of iron starvation where lack of iron activates the *fur* gene which then activates the *ompH* gene leading to the increased production of the protein (Bosch *et al.*, 2001). These proteins may also be associated with intracellular trafficking, transport and secretion of vesicles apart from cell envelope biogenesis. They enable the adaptation of bacteria to various niches by obtaining iron from host-iron complexes. Increased amounts of glucose could also result in the suppression of the outer membrane protein H expression (Wheeler, 2009). Previous studies have shown that depletion of carbon sources in the growth media can result

in the activation of the gene encoding the synthesis of the protein since general expression in the presence of sugars is prevented by catabolite repression as studied in *Photobacterium* sp. (Bartlett & Welch, 1995). Outer membrane proteins A and II have been known to play an important role in conjugation, invasion of host tissues and detection of viruses apart from inducing immunogenic responses in the host (Singamsetty *et al.*, 2008). Deletion of the gene encoding the synthesis of the outer membrane proteins caused a reduction in the biomass, reduced survival rate at high temperatures, loss of viability in stationary phase (Behr *et al.*, 1980) and accumulation of low molecular weight solutes (Barrios *et al.*, 2006). Previous studies in *B. fragilis* (Ko *et al.*, 2009) and *Chlamydia pneumonia* (Hogan *et al.*, 2003) have shown the outer membrane protein encoding genes to be associated with virulence and seem to be up-regulated in severe cases of infection.

10.15 Intracellular trafficking, secretion and vesicular transport associated protein expression

Translocation associated TolB proteins were found to be differentially expressed in mucin enriched media in *E. cancerogenus* where up-regulation was observed in mucin Type III enriched media and down-regulation was observed in the mucin Type II enriched media. This protein was not identified in *B. fragilis*. TolB proteins are known to be involved in a number of functions that include translocation of colicins of group E and A including the delivery of the toxins to their respective target sites, assembly of porins in the outer membrane, recycling of peptidoglycan and linking the peptidoglycan layer to the outer membrane of the cell (Abergel *et al.*, 1999). They have been known to interact with trimeric outer membrane complexes including ompF and ompC but do not interact with monomeric forms like ompA (Rigal *et al.*, 1997). They are a part of the Tol dependent translocation system where they form complexes with porins and can initiate the import or export of colicin but increased expression of TolA proteins in the form of TolA II His-porin complexes can result in the down-regulation of the TolB proteins (Lazzaroni *et al.*, 2002). They may also be essential for maintenance of the outer membrane stability (Lazzaroni *et al.*, 1999).

10.16 Expression of proteins involved in post translational modifications, protein turnover and chaperones

Putative thiol peroxidase was found to be differentially expressed in *B. fragilis* in mucin Type II and III enriched media but was not identified in *E. cancerogenus*. The protein was found to be up-regulated in SSP number 6301 and down-regulated in SSP numbers 4601 and 4605 in mucin enriched media in *B. fragilis* (Figure 33). . They play an important role as antioxidants and may be activated under conditions of oxidative stress and protect cells against the damaging effects of reactive oxygen species produced during metabolism (Missall *et al.*, 2004). In the presence of hydrogen peroxide, they convert the reduced form of thioredoxin to the oxidised form and are also known as peroxiredoxins. The presence of varying concentrations of oxidants in the growth media produced as a result of the accumulation of toxic metabolic end products may be responsible for the differential expression of these enzymes in mucin enriched media. Under anaerobic growth conditions, the expression of these proteins is controlled by the binding of designated proteins that prevent the activation of the *tpx* promoter (Kim *et al.*, 1999). The regulation of these proteins may be attributed to the adaptation capabilities of bacteria to various ecological niches even under conditions of stress.

These proteins protect DNA and detoxify peroxides. The activation of the genes controlling the expression of these proteins is controlled by the *tpx* promoter (Kim *et al.*, 1999). Considering the lack of oxygen in anaerobic growth conditions, the down-regulation of these proteins suggests the possible lowering of oxidative stress in mucin enriched media (Kim *et al.*, 1999).

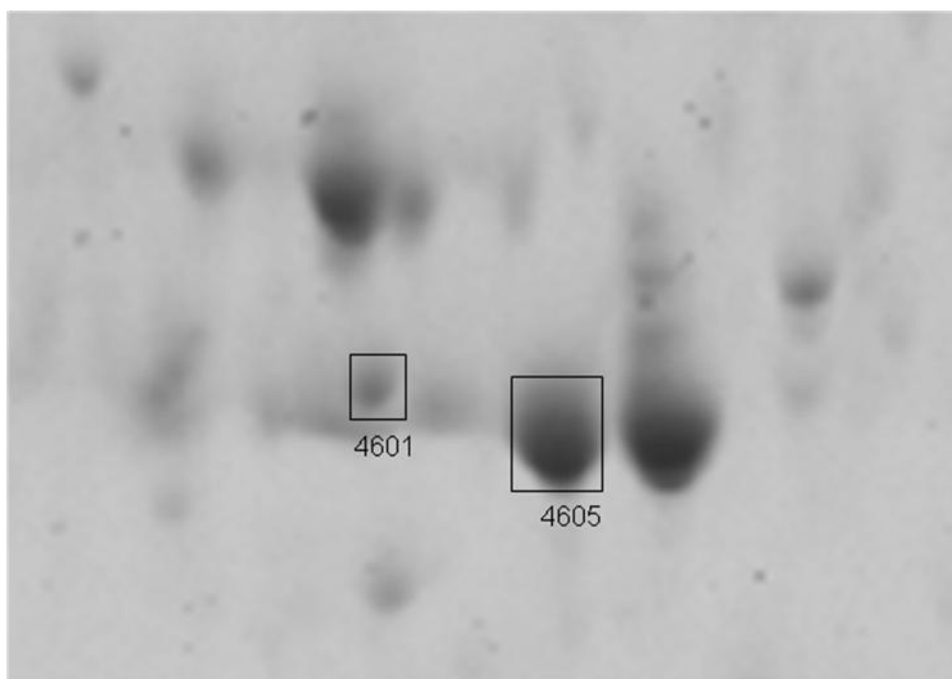


Figure 33: Figure showing the differentially expressed spots of putative thiol peroxidase in *B. fragilis*

E. cancerogenus showed a down-regulation of a protein from the glutaredoxin family in mucin Type III enriched media but was not identified in *B. fragilis*. Glutaredoxin related proteins are disulphide reductases that contain glutaredoxin and NADPH as cofactors. They have the presence of an intramolecular disulphide bond, act as electron carriers in the synthesis of deoxyribonucleotides and contribute to cellular functions like cell proliferation, viability and stabilisation during stress conditions (Holmgren *et al.*, 2005). Thiol disulphide oxidoreductases play an important role in disulphide bond formation in cytochrome biogenesis (Holmgren *et al.*, 2005). Previous studies in *E. coli* reveal the ability of these proteins to adhere to tissue culture cells and may contribute to the formation of pili (Donnenberg *et al.*, 1997). They have been known to contribute to bacterial virulence by acting as a folding catalyst suggesting that the down-regulation of these proteins in mucin media may be associated with the absence of a host immune response that induced expression (Yu & Kroll, 1999). This explains the reason behind the observation of a down-regulation of the glutaredoxin family related protein like thiol disulphide oxidoreductase in mucin Type III enriched media in *E. cancerogenus*. Other factors such as general metabolic requirements or toxic by product build up could also be responsible.

The co-chaperonin protein GroES was found to be up-regulated in mucin Type III enriched media in *B. fragilis*. This protein was not identified in *E. cancerogenus*. This is an oligomeric molecular chaperone which functions in protein folding and intracellular signals. It is a surface protein which is released from cells when required. It captures the substrate with the help of co-chaperone GroES and ATP which help with the binding. The substrate is then discharged into the microenvironment inside the chaperone promoting productive folding. The polypeptide is later released into the solution. GroES works as a co-chaperone with GroEL during protein folding. They have metal binding enzyme sites and can play an important role in host immunogenicity (Ranford *et al.*, 2004). These chaperonins are known to act as intercellular signals interacting with a variety of cell types, including leukocytes, vascular endothelial cells and epithelial cells and may be responsible for key cellular activities such as synthesis of cytokines and adhesion proteins (Landry *et al.*, 1996). Co-chaperonin GroES is a known virulence factor and has been known to be associated with inducing inflammation and promoting cell proliferation (Lin *et al.*, 2006). Up-regulation of these virulence associated proteins in mucin Type III enriched media indicate that they may be expressed during infection of the host by bacteria.

A heat shock protein, GrpE was found to be differentially down-regulated in mucin enriched media in *B. fragilis* but no regulation was observed in *E. cancerogenus*. This is a stress related protein that works in conjunction with two other proteins namely DnaK and DnaJ forming complexes and is activated to repair damaged proteins or to initiate folding of proteins (Packschies *et al.*, 1997). Apart from being involved in protein folding and disaggregation of heat shock proteins, these proteins also carry out the degradation of misfolded proteins. Unfolded proteins bind to DnaJ protein to form an unstable complex which is then stabilised by the hydrolysis action of DnaK. The movement of the folded protein is initiated by the removal of an ADP molecule which is converted to ATP coupled to the DnaK protein caused by the GrpE protein (Straus *et al.*, 1990). Inactivation of the gene encoding the GrpE protein in *E. coli* has known to enhance the production of heat shock proteins and the continual production of the heat shock protein could not be stopped even though the growth conditions were returned to normal (Lipinska *et al.*, 1988). Down-regulation of

the protein indicates the lowering of heat shock or oxidative stress when grown in mucin enriched media when compared to the control media.

ATP synthase subunit E was up-regulated in mucin Type II and III enriched media in *B. fragilis* but no regulation was observed in *E. cancerogenus*. ATP synthases form proton channels that cause an influx of ions across the membrane thereby generating a proton gradient formed by the hydrolysis of ATP or oxidative phosphorylation of ADP and resulting in the generation of energy for metabolic processes (Feniouk & Junge, 2005). Differential expression studies in *Mycobacterium avium* showed the up-regulation of the the ATP synthase subunit C called the *AtpC* component of the ATP synthase complex (Radosevich *et al.*, 2007). The exact function of the subunit E is not known but the up-regulation of the protein may be associated with energy generation for efficient growth of the bacteria. Up-regulation of the ATPase subunit E may be associated with the availability of a large amount of sugar substrates for growth thereby supporting an increased generation of energy in *B. fragilis*. It is quite interesting that only the subunit E of the ATP synthase complex appeared to be up-regulated in mucin enriched media instead of the entire protein complex. ATP synthase subunit B, another component of the ATP synthase complex was identified during MS analysis of protein spots but did not show any differential expression in mucin enriched media. The exact reasons for the differential expression of the ATP synthase subunit E in mucin enriched media remain unknown.

10.17 Expression of proteins involved in energy production and conversion

Phosphoenol pyruvate carboxykinase (PEPCK), an important enzyme in the gluconeogenic pathway catalysing the decarboxylation of oxaloacetate into phosphoenol pyruvate and carbon-di-oxide was shown to be down-regulated in mucin Type II and III enriched media in *B. fragilis*. This protein was not identified in *E. cancerogenus*. They utilise ATP or GTP as a source of phosphates and exhibit catalytic activity by binding to metal containing sites. The activation of these enzymes occurs when non fermentable sources of carbon are present in the media

and have been known to occur in ruminal and other anaerobic bacteria (Schocke & Weimer, 1997). Over expression of these proteins could affect the growth balance since they are known to have an influence over the oxygen concentrations in the media and this alteration in expression has been studied in *E. coli* (Chao & Liao, 1993). Since mucin acts as a rich source of carbon substrates, the fermentation of these substrates probably depend on the energy conservation and nutritional specialisation of *B. fragilis*. Studies have shown the up-regulation of the *acuF* gene which encodes the synthesis of PEPCK when non fermentable sources of carbon are present in the growth media (Hynes *et al.*, 2002). Hence the down-regulation of the protein in mucin enriched media can be attributed to the lack of requirement of more sugars for growth as mucin itself acts as a major source of carbon.

One of the interesting results from my study showed the differential regulation of malate dehydrogenase enzymes in *B. fragilis* but was not identified in *E. cancerogenus*. These enzymes were found to be down-regulated in mucin Type II and III enriched media in SSP number 4607 but the same enzyme was found to be up-regulated in mucin Type II and III enriched media in SSP number 4804 (Fig. 34). This could indicate that the enzyme exists in isoforms that may be differentially expressed based on their function but has been observed only in eukaryotes and is not known to exist in *B. fragilis*. Malate dehydrogenases catalyse the conversion of malate to oxaloacetate in the citric acid cycle and their expression has been known to vary with the carbon substrates and growth conditions used for culturing bacteria (Minarik *et al.*, 2002). Apart from being involved in carbohydrate metabolism, they may also be associated with amino acid transport and metabolism. Previous studies in malate dehydrogenase enzymes have shown that they exist as isoenzymes and also exhibit allelic polymorphisms in eukaryotes (Gietl, 1992) and in phototropic bacteria *Rhodobacter sphaeroides* and *Rhodopseudomonas palustris* (Eprintsev *et al.*, 2008) . There were considerable amounts of distinctive amino acids associated with the allelic polymorphisms of the *mdh* gene indicating that this could have been caused by the horizontal transfer of genes in *E. coli* and *S. enterica* (Boyd *et al.*, 1994). The *mdh* gene in *E. coli* encodes for the ArcA protein whose expression is down-regulated under anaerobic conditions and regulation of expression was studied in response to varying oxygen, carbon and haeme availability in the growth media

(Park *et al.*, 1995). The differential expression of these proteins may be associated with the presence of mucin in the semi-defined growth media which acts as a rich source of carbon providing a better survival rate for the bacteria.

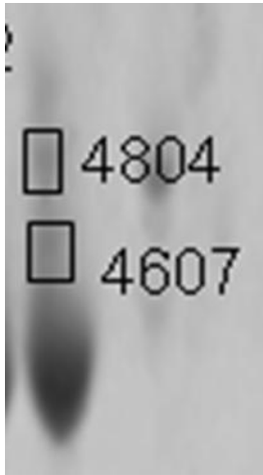


Figure 34: Figure showing the differentially expressed malate dehydrogenase spots in *B. fragilis*

NAD dependent aldehyde dehydrogenase enzyme was found to be down-regulated in media containing mucin Type II in *E. cancerogenus*. These proteins were not identified in *B. fragilis*. The aldehyde dehydrogenase enzymes are very closely associated with the gamma glutamyl phosphate reductase enzymes. They catalyse the dehydrogenation or oxidation of aldehydes to produce acidic end products in the presence of NADP as a co-factor. One of the important features of this enzyme is that it can act as an allergen that induces IgE responses in humans and mammals. The enzyme may be acting as a potential virulence factor by triggering immune responses within the host by activating the IgG antibodies. Under anaerobic conditions, aldehyde dehydrogenases metabolise endogenous acetaldehyde and ferment nutrients to produce energy (Mikulskis *et al.*, 1997). Previous studies showed that the expression of these proteins in germ free mice were modulated when challenged by bacteria and were found to be down-regulated (Fukushima *et al.*, 2003). Down-regulation of aldehyde dehydrogenase in mucin Type II enriched media indicates that the presence of mucin may be modulating the expression of this enzyme.

10.18 Expression of carbohydrate transport and metabolism associated proteins

Triose phosphate isomerase (TPI) which is involved in the interconversion of the TPI isomers dihydroxy acetone phosphate (DHAP) and glyceraldehyde-3-phosphate was found to be down-regulated in mucin Type II and III enriched media in *B. fragilis*. The enzyme plays an important role in carbohydrate metabolism through glycolysis and energy production. Regulation of the *tpi* gene which encodes the enzyme that can result in the accumulation of sugar moieties which becomes impossible to metabolise (Solem *et al.*, 2008). They are also known to be immunogenic in nature where these enzymes were up-regulated in convalescent phase serum obtained from patients suffering from pneumococcal infections caused by *S. pneumoniae* (Zysk *et al.*, 2000). Growth of *B. fragilis* in cell culture could trigger an up-regulation of these proteins since they are associated with the host immune system. Differential expression studies in planktonic and biofilm cultures showed the up-regulation of these enzymes in biofilm conditions (Becker *et al.*, 2001). Previous studies have shown that these enzymes can be over expressed during stress conditions like deprivation of oxygen and this suggests that the down-regulation of these proteins may be due to the ability of bacteria to grow well in the presence of mucin (Yamaji *et al.*, 2004). Regulation of expression of these proteins indicates that they play a vital role in carbohydrate metabolism and immune response suggesting that they could act as potential virulence factors when grown in host cells.

Up-regulation of the conserved domains of enolase or phosphopyruvate hydratase was observed in *E. cancerogenus* media enriched with mucin Type II and III. The enzyme forms an integral part of the glycolytic pathway catalysing the conversion of 2-phosphoglycerate to phosphoenol pyruvate. The expression of these enzymes has been found to be enhanced under anaerobic conditions and mutants lacking the presence of the enolase gene have been unable to utilise gluconeogenic and glycolytic carbon sources (Lemaire & Wesolowski-Louvel, 2004). Enolase has also been known to play an important role in tissue invasion and pathogenesis in Gram-positive bacteria and may be over expressed in infection conditions (Liu & Shih,

2007). The up-regulation of these proteins in *E. cancerogenus* suggests the possibility that they may be indirectly involved in pathogenicity and may act as potential virulence factors and are therefore worthy of further study.

Phosphoglycerate kinase; a key enzyme involved in the glycolytic pathway was up-regulated in mucin Type II and III enriched media in *E. cancerogenus*. This protein was not identified in *B. fragilis*. These enzymes are involved in the anaerobic fermentation and generation of energy from carbon sources by catalysing the conversion of 1, 3-bisphosphoglycerate into 3-phosphoglycerate in the glycolytic cycle. Inactivation of the *pgk* gene encoding the synthesis of phosphoglycerate kinase resulted in the accumulation of 1, 3-bisphosphoglycerate thereby leading to the inability to generate energy for metabolism (Nakano *et al.*, 1999). Up-regulation of these enzymes indicates the efficient metabolisation of sugars which is probably enhanced in the presence of mucin which acts as a rich source of carbon.

Fructose 1,6-bisphosphate aldolase was found to be down-regulated in mucin Type II and III enriched media in *B. fragilis* following analysis using the PDQuest gel analysis software. This protein was not identified in *E. cancerogenus*. This enzyme catalyses the conversion of fructose 1,6 bisphosphate to dihydroxyacetone phosphate and glyceraldehyde-3-phosphate in the glycolytic pathway involving the binding of a divalent metal ion (Zinc) to the active site (Anon, 2009). This enzyme has been known to be expressed when *B. fragilis* cell suspensions were grown in media containing glucose and lactate as carbon sources (Macy *et al.*, 1978). Previous studies in *Trichomonas vaginalis* revealed the up-regulation of malate dehydrogenases and fructose 1,6 bisphosphate aldolases and were found to be associated with virulence (Cuervo *et al.*, 2008). Hence these enzymes may be potential virulence factors and play a vital role in the adherence of bacteria to host cells. They were found to be over expressed in the viable but non culturable state of bacteria suggesting that they play a vital role in the alternate metabolic pathways involved in the generation of energy (Heim *et al.*, 2002) . In our experiments, fructose 1,6 bisphosphate aldolase was found to be down-regulated in mucin enriched media indicating better energy generation and viability of cells in mucin enriched media.

10.19 Amino acid transport and metabolism associated protein expression

Proteins from the carbamate kinase family involved in ATP and amino acid synthesis pathways was found to be up-regulated in mucin enriched media in *E. cancerogenus*. This protein was not identified in *B. fragilis*. Carbamate kinases catalyse the conversion of carbamoyl phosphate and ADP to carbamate and ATP. Hence this acts as a major source of ATP for energy metabolism. Previous studies have shown the over expression of this enzyme induced by growth of bacteria in anaerobic conditions, moderate expression induced by the presence of arginine and repression caused by the presence of ammonia (Abdelal *et al.*, 1982). Expression of carbamate kinase was found to increase when *S. aureus* cultures were grown under biofilm conditions and seem to play an important role in the growth of bacteria (Resch *et al.*, 2005). This indicates that the enzyme is involved in energy production reactions for the better growth of bacteria and its up-regulation may be attributed to the availability of rich sources of carbon under strict anaerobic conditions. Up-regulation of these proteins in mucin enriched media indicates an increase in energy metabolism in the presence of mucin in *E. cancerogenus*.

One of the other proteins that were differentially under expressed in mucin Type II and III enriched media was the lactoylglutathione lyase enzyme in *B. fragilis*. This protein was not identified in *E. cancerogenus*. This enzyme has been known to play an important role in the detoxification of the bacterial system and may be produced in response to oxidative stress (MacLean *et al.*, 1998). Only a general prediction of the function of these proteins is available associating them with amino acid transport and metabolism. This enzyme is also known as glyoxalase I and it catalyses the isomerisation of hemithioacetal adducts which are produced as a result of the reaction between glutathione (containing a glutathionyl group) and methylglyoxal (containing an aldehyde group). The end product of the reaction was lactoylglutathione which protected the system from the toxic effects of methylglyoxal (Korithoski *et al.*, 2007). The glutathione conjugates formed from the glyoxalase I-II pathway can activate KefB and KefC potassium channels that cause a lowering of the pH of the bacterial cell intracellularly thereby protecting them against the toxic effects of electrophiles (Ferguson *et al.*, 1998). Studies have shown that the genes encoding the expression of these enzymes when over expressed, resulted in an increased tolerance of the organism to methylglyoxal and osmotic stress exhibiting a

protective function (Takatsume *et al.*, 2005). They may be over expressed in cases of oxidative stress especially when acidic conditions prevail in the growth media (Wilkins *et al.*, 2002). Similar results have been observed in *S. mutans* where the *Igl* gene was over expressed during growth in acidic media (Korithoski *et al.*, 2007). But the down-regulation of the enzyme in mucin enriched media indicates a decrease in toxic metabolites or oxidative stress which suggests better viability and survival rates in mucin media rather than the control semi-defined media.

Proteins from the 3-deoxy-D-arabino-heptulosonate 7-phosphate synthetase I (DAHP) family was found to be up-regulated in mucin enriched media in *E. cancerogenus*. This protein was identified to be 2-dehydro-3-deoxy phosphooctonate aldolase and was not expressed in *B. fragilis*. 2-dehydro-3-deoxy phosphooctonate aldolase belongs to the transferase enzyme family and has been known to be up-regulated at transition stages from logarithmic to stationary phase in growth cultures (Sowell *et al.*, 2008). The enzyme, 3-deoxy-D-arabino-heptulosonate 7-phosphate synthetase I plays an important role in the aromatic amino acid biosynthesis and shikimate pathway (Panina *et al.*, 2001). Two of the main amino acids synthesised using the shikimate pathway are tyrosine and phenylalanine and the feedback inhibition caused by the end products of the shikimate pathway does not affect the activity of DAHP synthase enzyme in *Bacillus sp.* (Kim, 2001). It catalyses the condensation reaction that converts phosphoenol pyruvate and D-erythrose-4-phosphate to 3-deoxy-D-arabino-heptulosonate 7-phosphate and this step plays an important role in the synthesis of amino acids like tyrosine and phenyl alanine (Herrmann, 1995). Since the substrates used in the condensation reaction are mainly sugar phosphates and mucin acts as a rich source of carbohydrates and complex polysaccharides, the up-regulation of these enzymes in *B. fragilis* can be understood.

10.20 Expression of poorly characterized proteins with a general function prediction only

The periplasmic protein, ecotin was found to be down-regulated in *E. cancerogenus* but not identified in *B. fragilis*. The protein acts as a broad range serine protease inhibitor that is capable of acting against enzymes like trypsin, chymotrypsin, blood peptidase factor Xa, thrombin and urokinase type plasminogen activator (Mcgrath *et al.*, 1995). They exhibit a high degree of variation and develop adaptive interactions that suit the survival of the pathogen in host tissue and studies show that its location in the cell is periplasmic (Mcgrath *et al.*, 1991). They have a protective function and may be responsible for the bacteriostatic activity of the bacteria against neutrophil elastases (Eggers *et al.*, 2004). Since the presence of a host immune response triggers the expression of ecotin, its down-regulation suggests the lack of host digestive enzymes. This indicates that ecotin may be used as a protective system against proteolytic enzymes or other antibacterial agents and the lack of any components that challenge the growth of bacteria may have caused its down-regulation in mucin enriched media. In fact, media containing mucin support better growth of bacteria.

Differential expression of ribonucleotide reductase, pyruvate formate lyase (PFL) and autonomous glycyl radical cofactor were observed in *E. cancerogenus*. This protein was not identified in *B. fragilis*. Proteomic analysis of the spots revealed up-regulation of the protein in mucin Type II enriched media whereas down-regulation was observed in mucin Type III media. The autonomous glycyl radical cofactor plays an important role in the regulation of transcription and may be induced under conditions of oxidative stress since it acts as an antioxidant (Wyborn *et al.*, 2002). The enzyme ribonucleotide reductase is involved in the aerobic glycolysis reaction whereas pyruvate formate lyase takes part in the anaerobic glycolysis reaction (Leppanen *et al.*, 1999). Under oxidative stress conditions, the fragmentation of pyruvate formate lyase takes place and this induces the activation of the autonomous glycyl radical cofactor gene which then acts as an independent glycyl radical carrier and replaces the C terminal glycyl radical in PFL (Wagner *et al.*, 2001). Previous studies have shown that the enzyme pyruvate formate lyase is over expressed in the presence of pyruvate or milk and can yield mixed acid end products following anaerobic fermentation (Derzelle *et al.*, 2005). Since the exact composition

of mucin remains unknown, the end products of glycolysis may be inducing the expression of this protein in mucin Type II enriched media when compared to the mucin Type III enriched media as revealed in previous work (Sawers & Bock, 1988).

A transcription elongation associated protein, elongation factor GreA was identified to be down-regulated in mucin Type II and III enriched media in *E. cancerogenus*. They are involved in the process of transcription regulation where the elongation factors cleave 2 or 3 nucleotides from the N terminal end of the nascent transcripts of RNA polymerases which are trapped by the arresting sites in the DNA forming locked DNA/RNA polymerase tertiary complexes as they pass through them. Once cleaved, resumption of the process of elongation takes place from the 3' end (Lu *et al.*, 1997). Their functions include DNA dependent regulation of transcription, DNA binding and transcription elongation regulator activity (Stepanova *et al.*, 2007). These are cell surface associated proteins that are over expressed during stress conditions. Induced stress conditions like growth in lower pH (5.0) in *Streptococcus mutans* (Len *et al.*, 2004) and mutation of the *pgi* gene that encodes the synthesis of the phosphoglycerate kinase enzyme trigger the increased expression of these proteins (Kabir & Shimizu, 2003). The lack of any stress conditions and the ability to survive effectively in the media could be responsible for the down-regulation of this protein in mucin enriched media when compared to the control non mucin media in *E. cancerogenus*.

10.21 Expression of signal transduction mechanism associated proteins

The universal stress proteins were found to be up-regulated in mucin Type II and mucin Type III enriched media in *E. cancerogenus*. This protein has not been identified in *B. fragilis*. This result is similar to the autonomous glycy radical cofactor differential expression suggesting reduced accumulation of toxic end products or oxidative stress in mucin Type III media when compared to mucin Type II media. These cytoplasmic proteins play a major role in protecting DNA against damaging agents and they have been known to be over expressed when cells enter a preparatory phase before transition from the log phase to the stationary phase (Nystrom & Neidhardt, 1994). Expression may also be induced by depletion of carbon sources in growth media (Persson *et al.*, 2007) or in the presence of stress

inducing agents like salt or toxic by-products of metabolism (Weber & Jung, 2002). In some cases, these proteins have been known to be expressed even before complete depletion of carbon sources in the media (Chang *et al.*, 2002). Previous studies in the universal stress proteins expressed by *E.coli* showed that 6 different proteins which were encoded by 6 different genes *uspA, B, C, D* and *E* were produced under different conditions of stress. The genes *usp C* and *E* were found to produce proteins that were involved in cell adhesion and motility and this could be related to the up-regulation of these proteins in mucin enriched media (Nachin *et al.*, 2005).

10.22 Nucleotide transport and metabolism associated protein expression

Three other proteins found to be differentially expressed were cytidylate kinase in *E. cancerogenus*, phosphoglycerate kinase in *E. cancerogenus* and the 50S ribosomal protein L7/12 in *B. fragilis*. Cytidylate kinase proteins were found to be up-regulated in mucin Type III enriched media and down-regulated in mucin Type II enriched media in *E. cancerogenus*. These are cytoplasmic proteins that play a vital role in pyrimidine and nucleic acid metabolism by catalysing the transfer of a phosphate group from ATP thereby converting dCMP to dCDP and ADP. The two main functions of the protein include cytidylate kinase activity and ATP binding and they also require the binding of metal ions like magnesium (Mg^{2+}). When mutants lacking the conserved serine residues were generated, there was a dramatic decrease in the phosphorylation of dCMP indicating that these residues were highly conserved (Bertrand *et al.*, 2002). Not much information is available regarding the differential expression of these proteins but over expression in mucin Type III media suggests the possible availability of more energy in the form of ATP for nucleic acid synthesis unlike the mucin Type II or non mucin media.

E. cancerogenus showed the up-regulation of a periplasmic protein with a phospholipid binding domain in mucin enriched media. SSP number 6408 also showed a differential expression of a periplasmic protein disulphide isomerase I. This protein showed a down-regulation in the mucin Type II enriched media in *E. cancerogenus*. This protein was not identified in *B. fragilis*. The periplasmic protein disulphide isomerase belongs to the thiol disulphide oxidoreductase protein family

and is known to play an important role in the oxidative protein folding pathway. Gram-negative cell walls have the presence of phospholipids associated with their peptidoglycan and periplasmic layers. Periplasmic proteins form a part of bacterial transport systems with a high affinity and solute binding extracellular site (Ito *et al.*, 1981). They may also be involved in the formation and aggregation of porins that transport nutrients across the membrane but the exact function of these proteins remain unknown (Lazar & Kolter, 1996). Previous studies have shown that increased concentrations of sucrose is capable of inducing the over expression of these genes and mucin being a rich source of carbohydrates like sucrose could be responsible for the up-regulation of these proteins (Costerto *et al.*, 1974).

10.23 Summary

Some of the differentially expressed proteins identified in *E. cancerogenus* and *B. fragilis* were found to be hypothetical proteins that showed the presence of unknown domains of function. These include the SSP numbers 5003 (hypothetical protein 03947), 6306 (hypothetical protein 20877) and 8205 (Yfaz family protein) in *E. cancerogenus* and SSP numbers 1104 (BF2494), 2804 (BF2494), 5205 (BF1203) and 6206 (BF0301) in *B. fragilis*. Two of the proteins that were found to be differentially expressed in both *E. cancerogenus* and *B. fragilis* were elongation factor Ts and outer membrane proteins.

Apart from the differentially expressed proteins, several other proteins were found to be expressed in *B. fragilis* and *E. cancerogenus*. These can be classified as proteins required for the general metabolism and growth of bacteria in both control and mucin enriched media. A list of these proteins is available in the 'Result section Table numbers 20 and 22 '.

10.24 Future work

Cell culture coated multiwell plates can be used to study the interactions between the bacteria and eukaryotic cells. This may help in the better understanding of the virulence factors involved in the host infections caused by the bacteria since the presence of a host immune response may trigger the expression of virulence associated proteins.

The database used for searching and identifying the proteins of interest from the mass spectra was predominantly NCBI. It may be better to use other protein identification databases like PRIDE for determining the identity of proteins since they provide an identity based on searches in several databases including NCBI and UniProt.

A non gel based approach may also be used in studying differential expression in mucin media in both log and stationary phase. Even though the log phase is known to be associated more with virulence, it may be interesting to find out the differences in regulation that occurs in the stationary phase. One of the modern proteomics technologies includes Shotgun proteomics or Multidimensional protein identification technology (MUDPIT) where the proteins are subjected to a 2D-LC separation through strong cation exchange columns in the first dimension. The second dimension separation employs reverse phase chromatography. The cell lysates are tryptic digested before separation and pre-fractionation is carried out with iso-electric focussing. In order to obtain a better resolution of proteins, affinity chromatography using non-specific dyes could also be performed. The digestion and separation of proteins is followed by mass spectrometric analysis. The peptides are subjected to MS/MS by using electrospray ionisation (ESI) or matrix assisted laser desorption/ionisation (MALDI). The spectrum generated from the mass spectrometric analysis is compared to a database to determine the identity of the protein of interest. This is considered an efficient method because it helps to identify proteins with high molecular masses, extreme pI, hydrophobic or when they occur in low abundance. This approach could also help in the identification of integral membrane proteins and overcome some of the drawbacks of two dimensional gel electrophoresis. Analysis of proteins using this technique could be used as a future method of studying differential expression of proteins.

In the proteomics technique, the sample processing step plays a very important role since slight variations can cause drastic changes in expression. In my methodology, the use of the clean up kit seems to help in obtaining a good resolution of proteins but the components of this kit remains unknown. It would be interesting to compare the expression of proteins when the acetone precipitation method is used instead of the 2D clean up kit.

Further work could also be done in trying to generate mutants that lack the genes associated with the expression of proteins like the outer membrane surface antigens which are known for their virulence and appear to be differentially expressed in both *B. fragilis* and *E. cancerogenus*. This could be useful in providing more information regarding the pathogenicity of these bacteria.

11. Results and discussion of the expression and structural analysis of *B. fragilis* proteins from glycoside hydrolase families 97 and 95.

11.1 Construction of plasmid vector expression constructs

NB. Expression vector constructs carrying the genes encoding proteins CAH09443 and CAH06598 from glycoside hydrolase families 97 and 95 respectively, were provided by Lee Ling Bong and Cheun Hong Yeap from Northumbria University.

FASTA format of the putative α -fucosidase (CAH06598) of glycoside hydrolase family 95 from *B. fragilis* NCTC 9343:

>gi|60491840|emb|CAH06598.1| conserved hypothetical protein [*Bacteroides fragilis* NCTC 9343]

```
1 mkikllllllc cglwsscnsy dycpvtpses dlvtftglars wdeamplgna tvgalvwqrd
61 stlrslsldrt dlwdlrpvds lsgdnfrfsw vkehrqkny lpvqkkldwp ydmnpapski
121 pgaaielfple qigtptqvrl ylinalcead wadgtqmqtf vhatpigwf vfrnlktpie
181 psiitpvynk tkpdgsldpv sqqdlhrlgy qqgkvvregn qityhqkgyg dfsydvttvcw
241 kqegetlygt wsvtsslsge qasekaaal qrglkhdyqa hleywdkywa qssitlpdsv
301 lqkqyqnemy kfgsttrehs ypislqavwt adngklppwk gdyhhdntq lsywpaytgn
361 hltegmgyln tlwnqrdayk rytrryfgte gmnipgvctl tgepmggwiq ysmsqtvaaw
421 laqhfylqwk ysadrtflke raypfikdva iyleqisevt pegvrklefs sspeifdnsl
481 qawfsdmtny dlammhflfk atselaheln ladeaghwas leaqlpdydi deegcltfak
541 gypykeshrh fshamaihpl glidwsdgek sqhiiratlk rldkvgpdyw tgysyswlan
601 mkarafdgeg aaqalktfae cfclknfha ngdqtqsgks rftyrypftle gnfafaagiq
661 emllqshtgv irifpaipke wkdvsvfenlr amgaflvsar meggeinrvr iysekggmlk
721 marpgtltkpn knytlsgtdi lnidtqagew ielnp
```

The protein was found to contain a signal peptide that was cleaved between the 19th and 20th position of the amino acid sequence (Appendix I). The protein parameters were determined from the Protparam tool (Refer to Appendix I for details).

FASTA format of the putative α -glucosidase (CAH09443) of the glycoside hydrolase family 97 from *B. fragilis* NCTC 9343:

>gi|60494642|emb|CAH09443.1| putative exported protein [*Bacteroides fragilis* NCTC 9343]

```
1 mkrkmmssl11 alavisgssv yakvidvmsp ngaikvsvdi kdriyysvsy dndqllkdcy
61 ln1qlqnetl gtnphlrstk rgtidesvkr eipfknaivr nhcntlrnmf sgnvavefrv
121 fdngiayrfv tdkkgdnivm gedfainfpt nykahlsqpd gfktsyecpy thvdtekyaa
181 tdrmsylpvl ietdkaykil iseadlsdyp cmflkstgkn gmqsfpkap lafgedgdrs
241 lkiteeadyi aktdgkrsfp wrmmvsked kelienemvy nlsapcvled yswikpgqvs
301 wewwhdarly gvdfrrsgfnm dsykyidfa skfgipyim degwakntrd pftpnptinl
361 telikygkdr nvkivlwlpw ltvenhfdlf ktfadwgiag vkidfmdrsd qwmvnyyerv
421 akeaakhklf vdfhgafkpa glerkypnvl syegvlgmeq ggnckpensi ylpfmrnavg
481 pmdftpgsmi saqpednrst ranamsgtr afqmalfiif esglqmladn pvyvyrelpc
541 tefitsvpvt wdetkvlyak vgeavvvakr kgeqwfiggi tgnqpniei dlgfipagqs
601 ftltsfedgi nadrqamdyk kkestvnnqt rmtlkmvrng gwagtikmk
```

The SignalP server predicted the presence of a signal peptide with probable cleavage positions between the 22nd and the 23rd residues of the amino acid sequence (Appendix I). The protein parameters were determined from the Protparam tool (Appendix I).

The genes were amplified using the following primers:

Forward primer for BF3763

5'- ATGAAAGTGATTGACGTAATGTCTCCCAACGG-3'

Reverse primer for BF3763

5'- ATTGTTCCGGCCCATCCCC-3' and

Forward primer for BF0855

5'- TATGACTATTGTCCGGTCACTCCTTCAGAGAG-3'

Reverse primer for BF0855

5'- AGGGTTCAACTCGATCCATTCACCGGCTTGAGT-3'

The amplified genes which encoded the mature CAH09443 and CAH06598 proteins were cloned into the pETYSBLIC vector (see Appendix C)

The BF3763 and BF0855 genes were cloned and expressed in *E. coli* using the pET-YSBLIC vector system. *E. coli* was used as a host for the expression of recombinant proteins because of the ease of promoter control, high amount of product yields, high densities of growth in cultures, easy to grow with regards to metabolic carbon requirements (Brown, 1953). The pET-YSBLIC vector is a modification of the pET28a vector developed at York University where the gene of interest can be inserted into the vector through ligation independent cloning. The LIC vector is modified to encode a hexa-histidine tag that is incorporated at the *N* terminus of the protein of the encoded gene thereby making its purification easy. These factors make the LIC vector the most suitable candidate for cloning and expression experiments.

The SignalP and TMHMM websites were used to determine the presence of signal peptides and transmembrane helices in the proteins of interest so that genes sequences encoding these protein elements could be excluded during cloning. This was done to prevent the accumulation of secreted and transmembrane proteins in the inner membrane of the host which would inhibit growth.

11.2 Protein expression comparison using IPTG induced media and auto-induction media

Figs. 35 and 36 show SDS gels of CAH06598 and CAH09443 proteins, respectively, isolated from recombinant *E.coli* cells grown in IPTG induced media and auto-induction media.

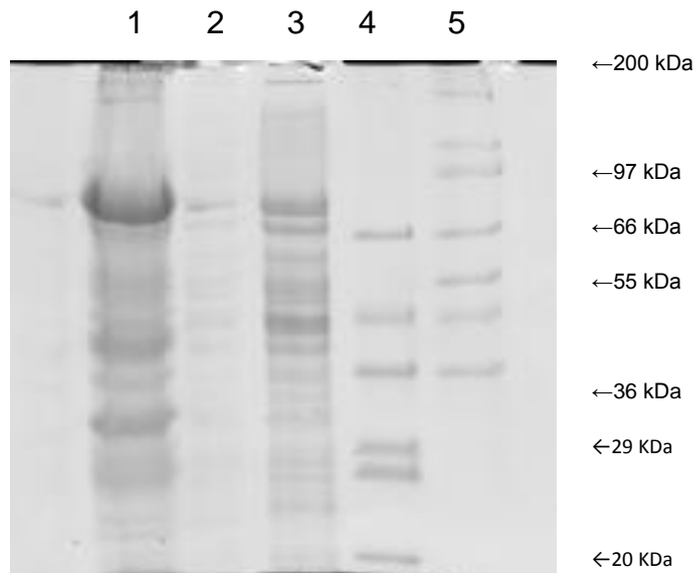


Figure 35: Analysis of CAH06598 expression (86 kDa) using a 12% (w/v) SDS gel obtained from growing the recombinant *E. coli* in IPTG induced media. Lane 1- Solubilisation buffer sample, Lane 2- 1 in 10 dilution of CFE, Lane 3- Cell free extract (CFE), Lane 4- Low molecular weight marker, Lane 5- High molecular weight marker.

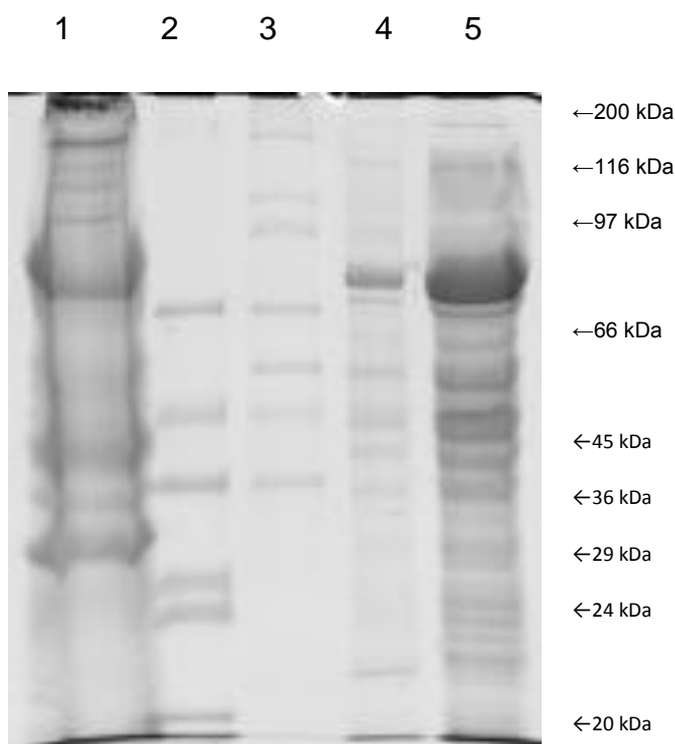


Figure 36: Analysis of CAH09443 expression (74 kDa) using a 12% (w/v) SDS gel obtained from growing recombinant *E. coli* in auto-induction media. Lane 1- Solubilisation buffer sample, 2- Low molecular weight marker, 3- High molecular weight marker, 4- 1 in 10 dilution of CFE, 5- CFE

Since the cell free extract in the auto-induction media showed better expression than the IPTG induced media, the former was used in the protein purification experiment.

The analysis of protein production was performed by comparing the expression levels in auto-induction and IPTG induced media. The auto-induction media which had the presence of glucose and α -lactose in it supported the growth of the recombinant *E. coli* and glucose was initially used as the source of carbon. Once the glucose present in the media was completely depleted, the cells started metabolising the lactose present. According to previous studies, this stage was normally achieved when the cells entered their mid or late log phase (Studier, 2005). The cultures were incubated overnight at 30°C with constant aeration and harvested at their stationary phase of growth (OD >4.0). However, the differences in level of expression of the protein of interest was not studied in lower temperatures or at various time points post induction even though lower temperatures (20°C) have been known to support

better growth due to increased solubility of oxygen (Donovan *et al.*, 1996). But since the expression of proteins was found to be very significant in the culture conditions used in the experiment and no other contaminating proteins were detected, the same system was used to obtain stable protein expression.

11.3 Protein expression and purification of *B. fragilis* proteins CAH06598 and CAH09443

CAH06598 and CAH09443 were extracted, purified using immobilised nickel affinity chromatography (Figs 37 and 41, respectively) and run on 12% (w/v) SDS-gels (Figs. 38 and 42, respectively) to confirm the proteins of the right molecular weight. Further purification of the proteins was performed using gel filtration and fractions observed on SDS gels.

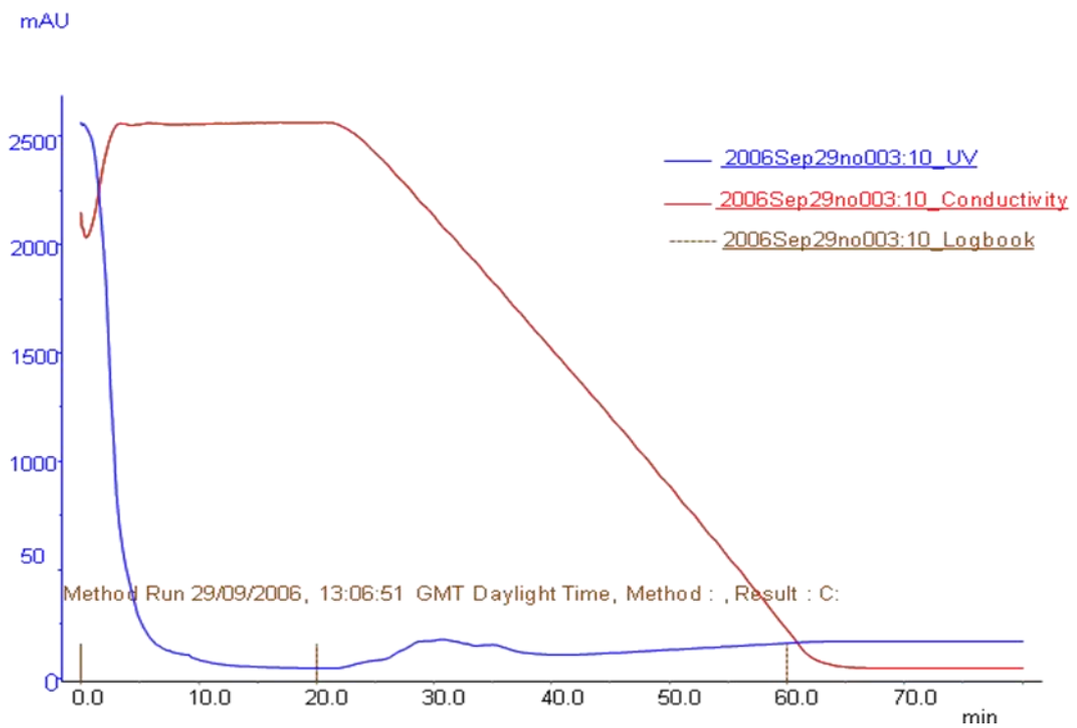


Figure 37: Nickel column purification of CAH06598 protein. The blue line represents the UV absorbance and the peak that appears between 20 and 40 min confirms the results from the SDS gels.

Figures 38: Nickel column purification results for CAH06598 protein

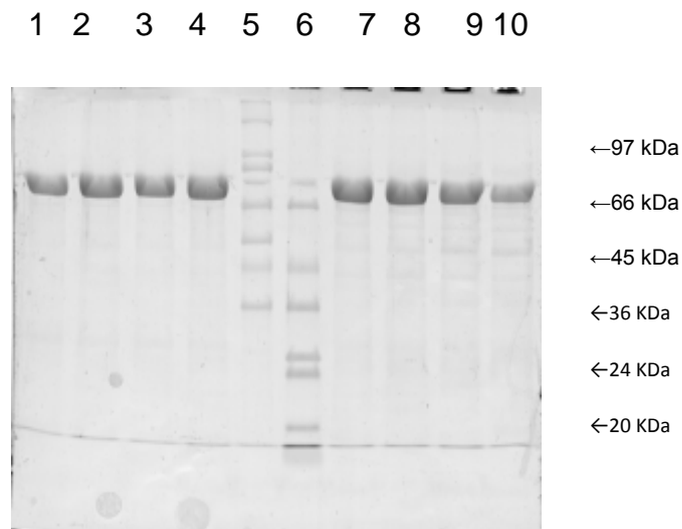


Figure 38.1: 12% (w/v) SDS gel showing purified fractions. Lanes 1, 2, 3, 4, 7, 8, 9 and 10 shows the presence of a band at 86 kDa indicating the presence of the CAH06598 protein which corresponded to the eluted fraction numbers of 26, 27, 28, 29, 30, 31, 32 and 33 respectively. Lanes 5 and 6 represent the high and low molecular weight standards respectively.

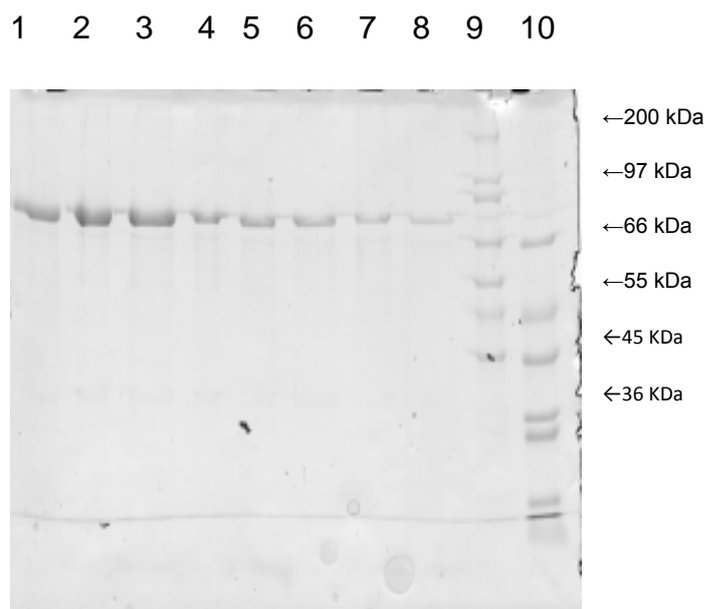


Figure 38.2: 12% (w/v) SDS gel showing purified fractions. Lanes 1 to 8 shows the decrease in intensity of the band at 86 kDa, indicating the presence of the CAH06598 protein, as elution continued and represent fraction numbers 34 to 41 respectively. Lanes 9 and 10 show the high and low molecular weight standards.

Fractions 34-41 were pooled and concentrated in a 30 kDa cut-off centrifugal concentrator and washed into 20 mM HEPES buffer pH 7.4. The concentrated protein was then purified using a HiLoad 16/60 Superdex 200 prep grade gel filtration column (Fig. 39) and the fractions containing CAH09443 analysed via SDS-PAGE (Fig. 40).

Figure 39: Gel filtration graph of CAH06598 protein showing the UV absorbance peak. Refer to Appendix section I for details.

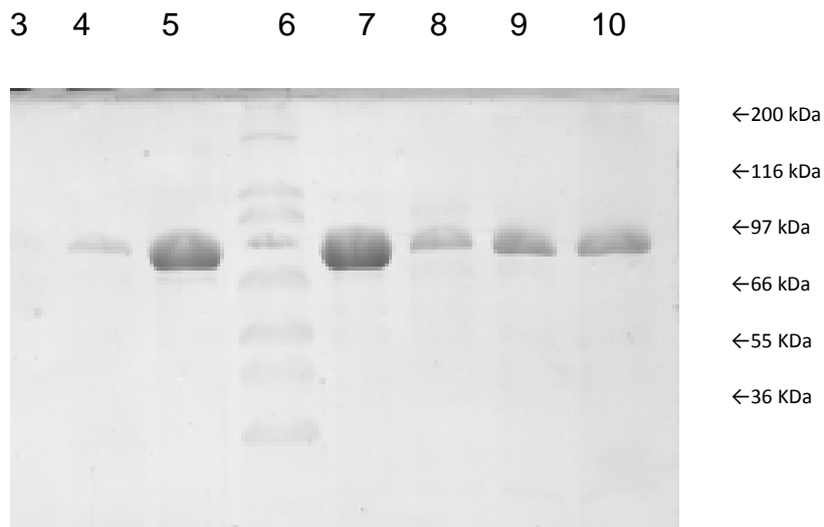


Figure 40: 12% (w/v) SDS gel showing the gel filtration fractions containing CAH06598 protein. Lanes 3, 4, 5, 7, 8, 9 and 10 represent the fractions 14, 15, 16, 17, 18, 19 and 20 respectively indicating the presence of protein bands at 86 kDa. Lane 6 shows the high molecular weight standard.

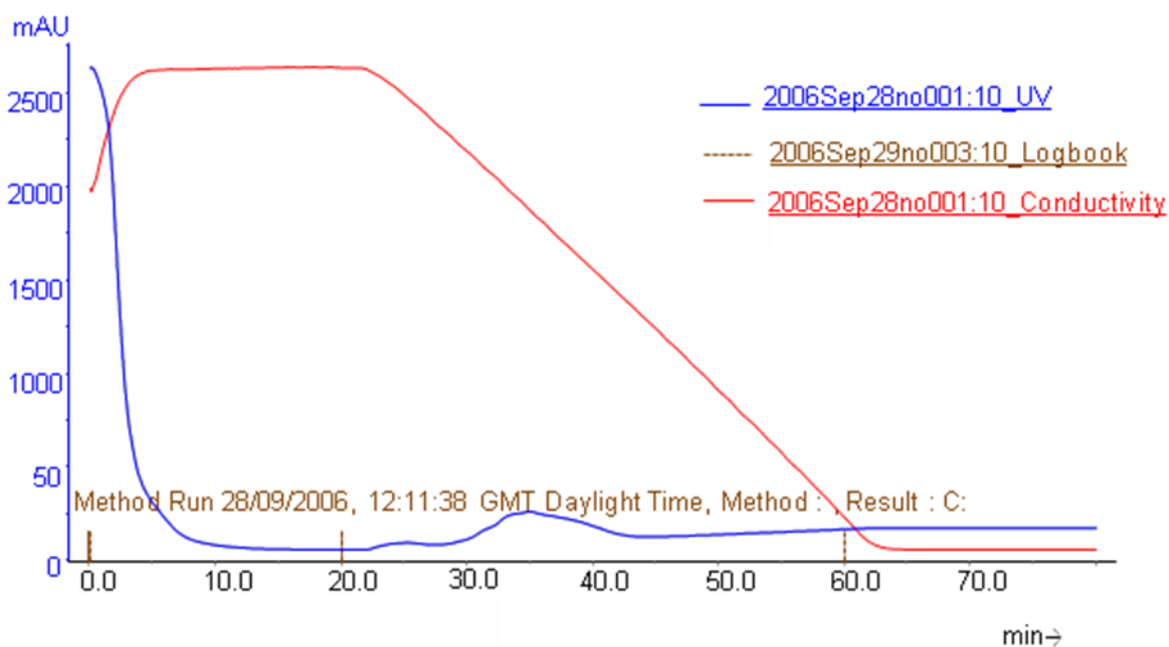


Figure 41: Nickel column purification results for CAH09443 protein. The blue line represents the UV absorbance and the peak that appears between 30 and 40 min indicates the elution of the protein at these corresponding fractions.

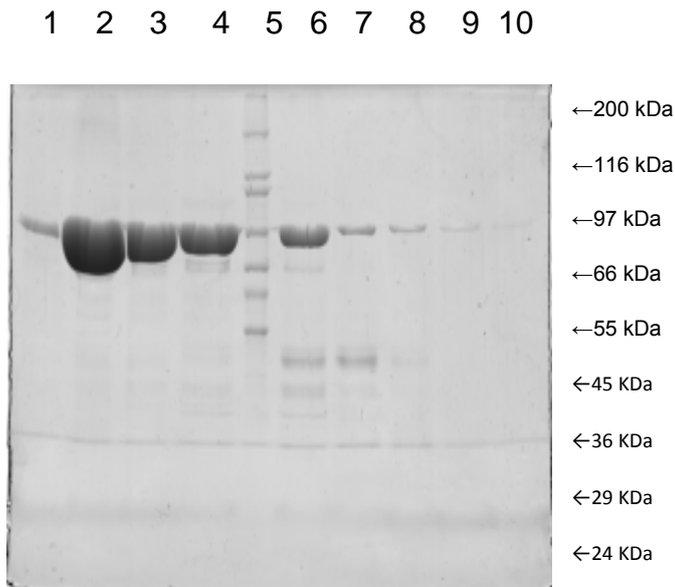


Figure 42: Nickel column purification results for CAH09443 protein (12% (w/v) SDS-gel). Lanes 2, 3, 4 and 6 represent fractions 34, 36, 38 and 40 respectively showing the presence of bands at 74 kDa. Lanes 1, 7, 8, 9 and 10 represent the fractions 31, 41, 43, 45 and 47 and Lane 5 represents the high molecular weight standard.

Fractions 34-40 were pooled and concentrated in a 30 kDa cut-off centrifugal concentrator and washed into 20 mM HEPES buffer pH 7.4. The concentrated protein was then purified using a HiLoad 16/60 Superdex 200 prep grade gel filtration column (Fig. 43) and the fractions containing CAH09443 analysed via SDS-PAGE (Fig. 44).

Figure 43: Gel filtration graph of CAH09443 protein showing the UV absorbance peak. Refer to Appendix section I for details.

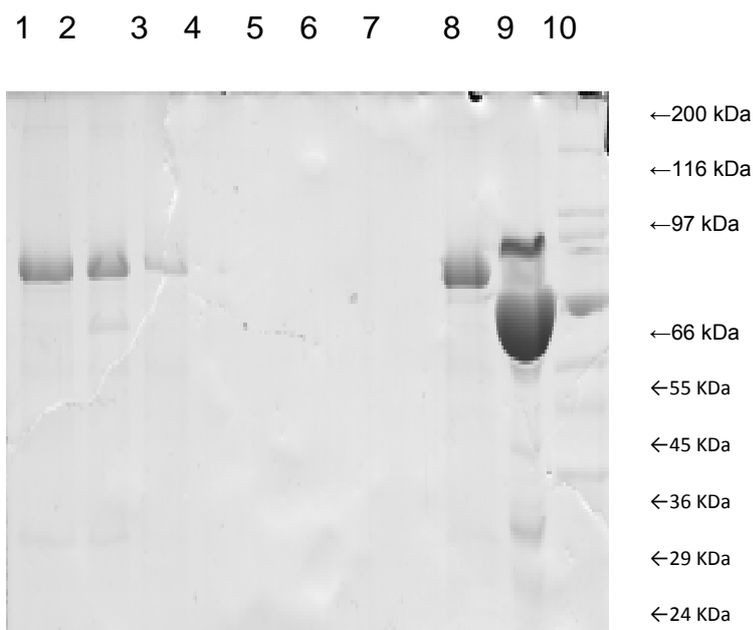


Figure 44: 12% (w/v) SDS gel showing the gel filtration results for CAH09443 protein. Lanes 1, 2, 3 and 8 show the fractions 10, 12, 13 and 15 respectively and Lane 9 and 10 represent the low and high molecular weight standards. Lanes 4 to 7 represent fractions 17, 19, 21 and 23 which do not show the presence of bands. Fractions 10 to 15 were pooled for further concentration.

The purified proteins were concentrated to a volume of 0.5 mL and the concentration of protein was estimated using Bradford's assay. They were diluted down to a concentration of 15 mg/mL from 43 mg/mL (CAH06598) and 29 mg/mL (CAH09443) respectively before crystallogenesis.

The proteins expressed and purified from CAH09443 showed tendencies towards precipitation at pH 9 and above. Hence the proteins were maintained in 5 mM HEPES buffer at a pH of 7.4. The nickel column purification of recombinant proteins from *E. coli* did not yield good peaks of absorbance at 280 nm in the UV chromatograms but strong and well detectable bands were observed on performing the SDS-PAGE analysis. The UV chromatogram peaks for GH95 proteins were found to be better resolved when compared to CAH09443.

11.4 Protein crystallogenes and diffraction results for *B. fragilis* CAH06598 and CAH09443.

A number of different crystallisation screens were used in the attempt to crystallise *B. fragilis* CAH06598 and CAH09443 proteins. These include PEG/Ion, Hampton Screen (HS) I and II, Clear Strategy Screen (CSS) I and II, PEG/Anion, PEG/Cation, PEG/pH, SaltRx, Newcastle and Index screens.

Crystals of CAH06598 proteins were observed in the Peg/Ion screen 3 containing 0.2 M ammonium fluoride and 20% (w/v) polyethylene glycol (PEG) 3350, CSS II screen condition 35 containing 10% PEG 8000, 10% PEG 1000 and 0.2 M Calcium acetate, HS I screen condition 20 containing 0.2 M ammonium sulphate, 0.1 M sodium acetate pH 4.6 and 25% (w/v) PEG 4000, Newcastle screen condition 1 containing 50% PEG 400, 0.2 M lithium sulphate and 0.1 M sodium acetate pH 5.1 and PEG/Anion screen condition 8 containing 0.2 M sodium sulphate and 20% PEG 3350. But no diffraction was obtained from the crystals of the above mentioned conditions.

The PACT ANION screen number 12 for CAH09443 produced crystals that diffracted and the screen was composed of 20% (v/v) PEG 3350 (Polyethylene Glycol) and 200 mM of sodium malonate. The proteins were buffered in 5 mM HEPES at a pH of 7.4. Other conditions that produced crystals include Peg/Ion condition 40 containing 0.2 M potassium thiocyanate, 0.1 M Bis-tris propane pH 8.5 and 20% PEG 3350, PEG/Anion condition 34 containing 0.2 M sodium potassium phosphate, 0.1 M Bis-tris propane pH 7.5 and 20% PEG 3350 and PEG/Anion condition 22 containing 0.2 M sodium potassium phosphate, 0.1 M Bis-tris propane pH 6.5 and 20% PEG 3350. But unfortunately these crystals did not diffract.

Crystallisation studies were performed on both the CAH06598 and CAH09443 protein families but no crystals were detected in CAH06598 that gave a positive diffraction pattern. CAH09443, yielded crystals with a positive diffraction pattern in the PACT/ANION screen number 12 but when the selenomethionine proteins were produced in minimal media, crystals were obtained but did not show a diffraction pattern. Attempts were made at growing selenomethionine protein crystals by seeding existing crystals into new screen conditions but no detectable crystals were observed.

11.5 Production, purification and crystallogenesis of a selenomethionine derivative for *B. fragilis* CAH09443

A selenomethionine derivative of CAH09443 was expressed, purified (Fig. 45) and crystallised but no positive diffraction patterns were obtained from the crystals.

Figure 45: Purification results for selenomethionine incorporated CAH09443 derivative.

12% (w/v) SDS gels showing the nickel column purification and gel filtration purification results for the selenomethionine preparations of CAH09443 respectively.

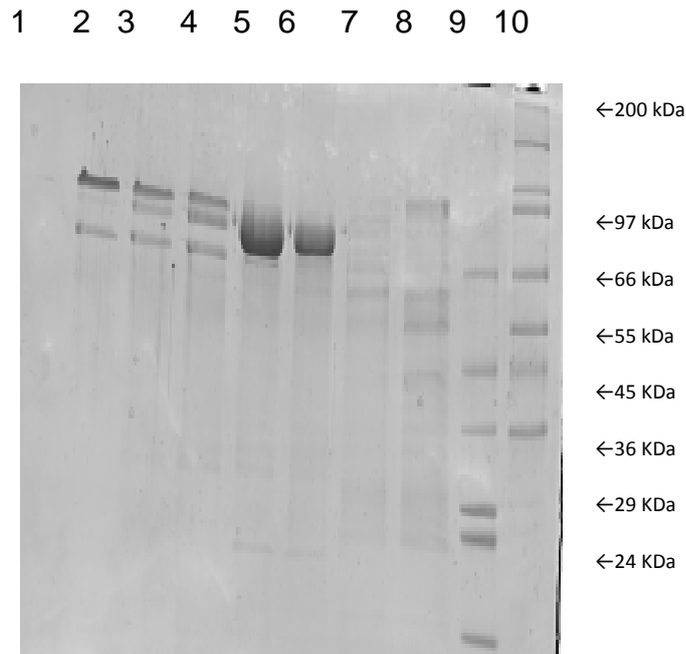


Figure 45.1: 12% (w/v) SDS gel showing the nickel column purification results for selenomethionine preparation of CAH09443 protein. Lanes 5 and 6 shows the presence of bands at 74 kDa and these were from fractions 41 and 43 and Lanes 1, 2, 3, 4, 7 and 8 represent fractions 31, 33, 36, 38, 46 respectively. Lanes 9 and 10 represent low and high molecular weight standard.

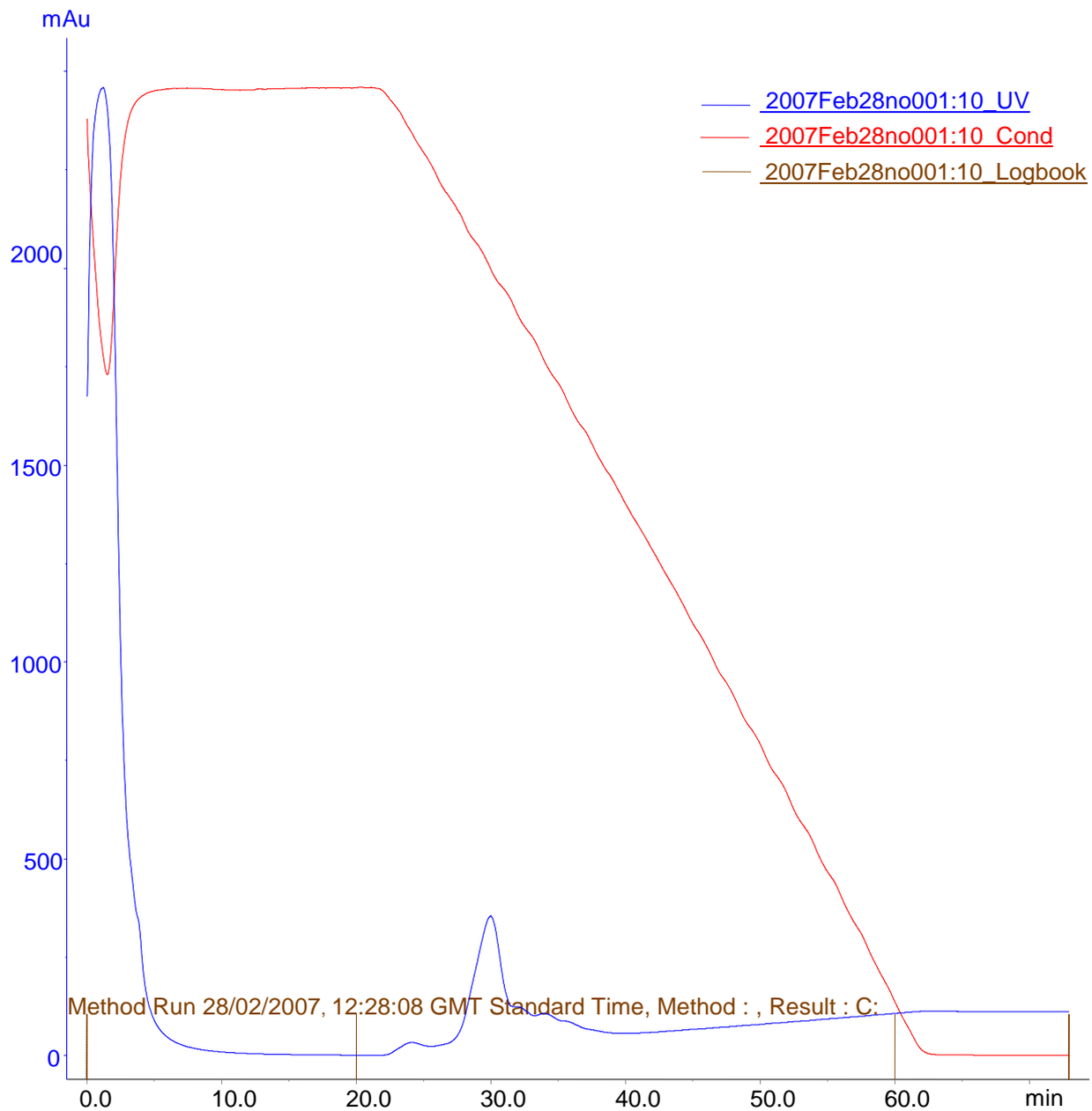


Figure 45.2: Nickel column purification results for the selenomethionine derivative of CAH09443 protein.

Figure 45.3: Gel filtration graph of CA09443 protein showing the UV absorbance peak for the selenomethionine derivative. Refer to Appendix section I for details.

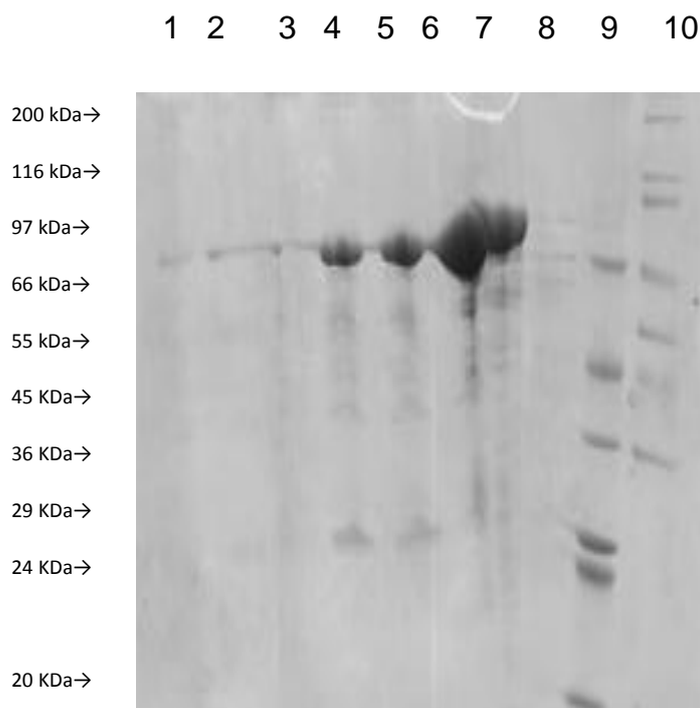


Figure 45.4: 12% (w/v) SDS gel showing gel filtration purification results for selenomethionine preparation of CAH09443 protein.

Lanes 4 to 7 shows the presence of purified fractions and the fraction numbers were 12, 13, 14 and 15. Lanes 9 and 10 were the low and high molecular weight standards and lanes 1 to 3 and 8 represent fractions 9, 10, 11 and 16 respectively.

The gels indicate the presence of bands at the correct molecular weight of 74 kDa.

Crystals were produced on screening using the Newcastle crystal screen condition 17 containing 40% MPD, 5% PEG 8000 and 0.1 M sodium cacodylate pH 7.0 but did not show diffraction.

11.6 Structure solution of CAH09443 via molecular replacement

NB. Structure solution of CAH09443 was performed by Dr Edward Taylor at York Structural Biology Laboratory.

The crystals were found to belong to space group P 2 21 21 with two molecule in the asymmetric unit with the approximate cell dimensions of $a = 60.33 \text{ \AA}$ $b = 132.82 \text{ \AA}$ $c = 163.65 \text{ \AA}$ (Table 24). The structure was determined via molecular replacement using *Bacteroides thetaiotaomicron* BtGH97b (UniProtKB/TrEMBL entry Q8A6L0) structure (PDB code 3A24) as a search model and refined to a resolution of 2.70 \AA with an R-factor of 0.225 and R-free of 0.296 (Table 24).

Data Processing	<i>BfGH97</i>
Space Group (No.)	P 2 21 21 (18)
Unit Cell lengths (Å)	a = 60.33 b = 132.82 c = 163.65
Unit Cell angles (°)	$\alpha = \beta = \gamma = 90$
Molecules in asymmetric unit	2
Resolution Range (outer shell) ¹	81.82- 2.70 (2.85-2.70)
R _{merge} *	0.11 (0.464)
$\langle I/\sigma_I \rangle^*$	6.3 (15.6)
Completeness*	99.8 (99.7)
Redundancy*	6.6 (5.7)
Refinement Statistics	
Resolution Range (Å)	103.13- 2.70
R _{cryst}	0.225
R _{free} ²	0.296
No. protein atoms	9953
Mean <i>B</i> value protein atoms (Å ²)	28.0
Mean <i>B</i> value solvent atoms (Å ²)	29.4
Ramachandran Statistics³	88.2% preferred regions, 7.5% allowed regions, 4.2% outliers

Table 24: Data collection and refinement statistics for *BfGH97* (*B. fragilis* CAH09443 protein). Structure figures were drawn with PyMOL (DeLano Scientific www.pymol.org).

1 Numbers in parenthesis correspond to the high resolution outer shell

2 Estimated Standard Uncertainty, based upon R_{free}, calculated using REFMAC

3 Calculated using Validation options in COOT

11.7 Three dimensional structure of CAH09443.

The tertiary structure of CAH09443 reveals two domains (Fig. 46); an N-terminal β -super-sandwich domain (in yellow) and a canonical $(\beta/\alpha)_8$ barrel (in red). This is a similar architecture to other GH97 enzymes, specifically the *Bacteroides thetaiotaomicron* α -glucosidase, *BtGH97a* (Gloster *et al.*, 2008; Kitamura *et al.*, 2008) and α -galactosidase, *BtGH97b* (Okuyama *et al.*, 2009). Previous studies in the *Bacteroides thetaiotomicron* GH97 showed that one of the general characteristics of GH97 enzymes may be their ability to possess two catalytic residues that act as bases at the end of β strands 3 and 5 or a conserved nucleophilic residue at the end of the β strand 4 (Refer to Fig. 47 for calcium binding sites of the *BtGH97* enzymes) (Okuyama *et al.*, 2009). Our enzyme, BfGH97 seems to show the presence of a conserved nucleophilic aspartic acid residue at the position 405 which corresponds to the position 415 in *BtGH97b* (Refer to Fig. 48 for details of the ClustalW alignment).

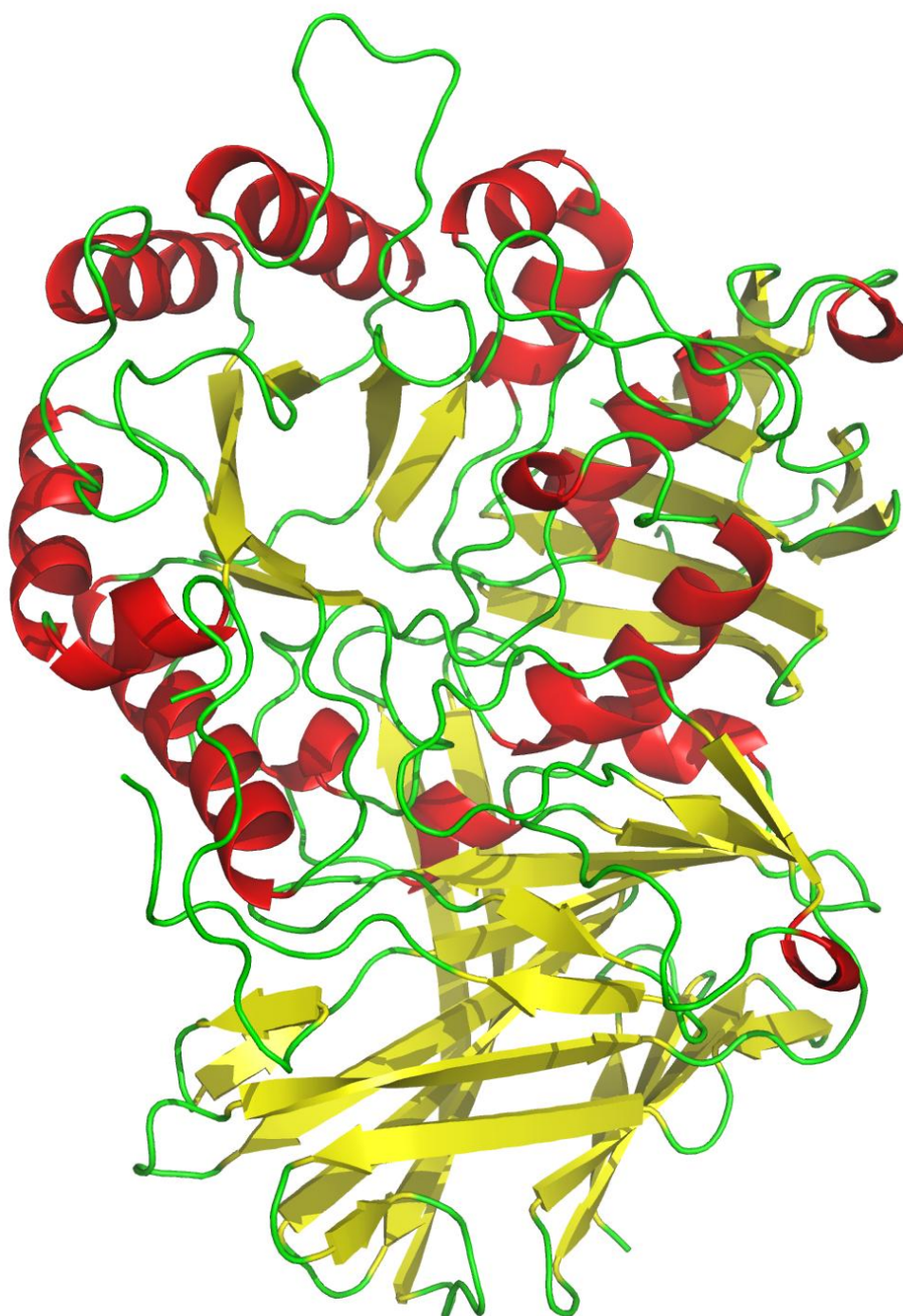


Figure 46: Structure of CAH09443. Crystal structure of CAH09443 coloured according to domains. The N-terminal β -super-sandwich domain is in yellow and a canonical $(\beta/\alpha)_8$ barrel is in red.

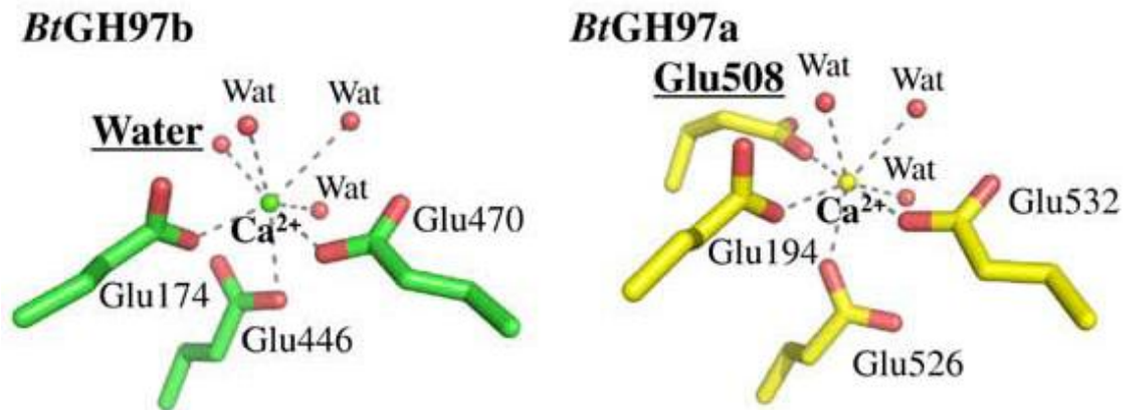


Figure 47: Figure shows the calcium binding sites in *BtGH97a* and *BtGH97b* (Okuyama *et al.*, 2009).

11.8 Biochemical assay for CAH06598 and CAH09443.

CAH06598 was predicted to be a putative uncharacterized protein and CAH09443 was predicted to be α -glucosidase.

Table 25 shows the presence or absence of activity in fluorescent and chromogenic assays for CAH06598 and CAH09443 proteins against methylumbelliferyl and p-nitrophenyl glycoside substrates, respectively.

Substrates used in enzyme assay	<i>B. fragilis</i> CAH06598 (GH95)	<i>B. fragilis</i> CAH09443 (GH97)
Para nitrophenyl α -D-glucoside	-	+ (18 h incubation)
Para nitrophenyl α -D-fucoside	+ (18 h incubation)	-
Para nitrophenyl α -D-galactoside	-	+ (18 h incubation)
Para nitrophenyl β -D-maltoside	-	-
4-methylumbelliferyl α -D-glucoside	-	+ (2 min incubation)
4-methylumbelliferyl α -D-fucoside	+ (2 min incubation)	+ (2 min incubation)
4-methylumbelliferyl β -D-maltoside	-	-

Table 25: Determination of the activity of CAH06598 and CAH09443 against a variety of para nitrophenyl and methylumbelliferyl substrates at 37°C.

+ indicates presence of activity

- indicates absence of activity

Two types of enzyme assays were performed to determine the putative activity of the enzymes from CAH06598 and CAH09443 which was α -fucosidase and α -glucosidase respectively. The fluorimetric assays showed activity towards their respective substrates but the chromophoric assays did not show positive activity. As mentioned in the results section Table number 23, the pNP assay showed a change in colour of the reaction mixture when incubated overnight at 37°C but no instant reactions or change of colour to yellow was observed. This suggests that the reaction occurred at a slower rate in the pNP assay which could probably be optimised by changing the concentration of the substrates, enzymes or by modifying the reaction conditions. The enzymes were found to be very specific in their reactions towards the substrates even though a wide range of substrates were used. This also indicates that the fluorescent assays were more sensitive when compared to the chromophoric assays. The CAH06598 putative α -fucosidase enzyme exhibited activity towards 4-methylumbelliferyl α -D-fucoside and CAH09443 putative α -glucosidase exhibited activity towards 4-methylumbelliferyl α -D-glucoside indicating the specificity of the enzyme in its activity against the substrates used. No assay experiments were performed to determine the optimum temperature and pH for the activity of the enzymes since positive activity was observed at 37°C and the pH was maintained at 7.4 since the CAH09443 enzyme precipitated at higher pH.

α -Glucosidase enzymes are also known as acid maltases, glucoinvertases, glucosidosucrases, lysosomal α -glucosidases, maltases and maltase-glucoamylases. They are capable of hydrolysing terminal, non reducing 1 \rightarrow 4 linked D-glucose residues resulting in the release of α -D-glucose as an end product. They are also known as exo-enzymes because they carry out exohydrolysis of α 1 \rightarrow 4 glucosidic linkages and hydrolyse oligosaccharides rapidly. The hydrolysis of polysaccharides is comparatively slower and industrial enzymes are capable of hydrolysing 1, 6 α -D-glucose linkages in polysaccharides (Hughes *et al.*, 2003).

These enzymes play an important role in the hydrolysis of complex polysaccharides into glucose and act as one of the major factors that contribute to type 2 diabetes. Attempts are being made to use acarbose, which is a competitive inhibitor of α -glucosidase to lower the blood glucose levels in diabetic patients following the consumption of food containing carbohydrates (Van de Laar *et al.*, 2005). The

enzymes also take part in glycogen metabolism and nutrient uptake of bacteria. They catalyse the transglycosylation of α -D-glucose moieties and are industrially useful in the biosynthesis of oligosaccharides or glycoconjugates (Shirai *et al.*, 2008).

11.9 Future work

Further experiments could be performed to generate a catalytic mutant of CAH09443 that lacks the putative acid catalyst, Asp404. This amino acid residue has been identified to be the acid catalyst as it aligns with the acid catalyst, Asp415 of the α -galactosidase, BtGH97b, from *Bacteroides thetaiotaomicron* (Fig. 47). Once the mutant is generated, its structure can be solved in the presence of substrate, therefore providing additional structural data with respect to enzyme-substrate interactions.

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gi | 60494642 | emb | CAH09443.1 | MKRKMSLLALLAVISGSSVYAKVIDVMSPNGAIKVSVDIKDRIYYSVSY 50
gi | 29339180 | gb | AAO76978.1 | MKKLTFLLLCVLCTLS--LQAQKQFTLASPDGNLKTITIGDRLTYDITC 48
** : ** *..* * : ** * : ** : * ** : *..:

gi | 60494642 | emb | CAH09443.1 | DNDQLLKDCYLNLQLQN-ETLGTNPHLRSTKRGTIDESVKREIPFKNAIV 99
gi | 29339180 | gb | AAO76978.1 | NGRQILTPSPISMTLDNGTVWGENAKLSGTSRKSVDEMIPSPF-YRASEL 97
:. *:* . :. : *:* . * *:* . ** * : ** : : : :

gi | 60494642 | emb | CAH09443.1 | RHNCLTRMNFSGNYAVEFRVFDNGIAYRFVTDKKGDNIVMGEDFAINF 149
gi | 29339180 | gb | AAO76978.1 | RNHYGLTLRFKKDWNVEFRAYNDGIAYRFVQGGKPFVTVSDYCF 147
*** * * :.* : : *****:*****: * * : * **

gi | 60494642 | emb | CAH09443.1 | TNYKAHL----SQPDG-----FKTSYECPYTHVDTEKYAATDRMSYLPVL 190
gi | 29339180 | gb | AAO76978.1 | SDMTASVPYVKSGKGDYNSQFNFSEFNTYTTDKLSKLNK-QRLMFLPLV 196
: . * : * ** * . ** * . * . * : * ** :

gi | 60494642 | emb | CAH09443.1 | IETDKAYKILISEADLSDYPCMFLLKSTG---KNGMQSIFPKAPLAFGED 236
gi | 29339180 | gb | AAO76978.1 | VDAGDGVKVCITESDLENYPLGLYSASEGANRLSSMHAPYKRTVQGGHN 246
: : . . . * : * : ** : ** : * : * : * : * : * : * : * :

gi | 60494642 | emb | CAH09443.1 | GDRSLKITEEADYIAKTGKRSPWRMMVISKEDKELIENEMVYNLSAPC 286
gi | 29339180 | gb | AAO76978.1 | -QLQMLVKEHEDYIAKVDKPRNFPWRIAVVTTDKDLAATNLSYLLGAPS 295
: . : : . * . * ** * * * . * : * : * : * : * : * : * :

gi | 60494642 | emb | CAH09443.1 | VLEDYSWIKPGQVSWEWHHDARLYGVDFRSGFNMSYKYIIDFASKFGIP 336
gi | 29339180 | gb | AAO76978.1 | RMSDLSWIKPGKVAWDWNLDGVDFVTGVNNTYKAYIDFASANGIE 345
: . * * ** * : * : * : * : * : * : * : * : * : * : * :

gi | 60494642 | emb | CAH09443.1 | YIIMDEGWAKNTR-DPFTPNPTINLTELIKYKDRNVKIVLWLPWLTVEN 385
gi | 29339180 | gb | AAO76978.1 | YVILDEGWAVNLQADLMQVVKEDLKLVDYAASKNVGIIWAGYHAFER 395
* : * : * ** * * : * : * : * : * : * : * : * : * : * :

gi | 60494642 | emb | CAH09443.1 | HFD-LFKTFADWGIAGVKIDFMDRSDQWMVNYERVAKAAKHKLFVDFH 434
gi | 29339180 | gb | AAO76978.1 | DMENVCRHYAEMGVKGFVDFMDRDDQEMTAFNYRAAEMCAKYKLILDH 445
. : : : : * : * : * : * : * : * : * : * : * : * : * : * :

gi | 60494642 | emb | CAH09443.1 | GAFKPAGLERKYPNVLSEYGLGMEQGGNCKP-----ENSIYLPFMRNAV 479
gi | 29339180 | gb | AAO76978.1 | GTHKPAGLNRTYPNVLNFEVNGLEQMKWSSPSVDQVKYDVMIPFIRQVS 495
* : * : * : * : * : * : * : * : * : * : * : * : * : * : * :

gi | 60494642 | emb | CAH09443.1 | GPMDFTPGSMISAQPEDNRSTRANAMGSGTRAFQMALFIIFESGLQMLAD 529
gi | 29339180 | gb | AAO76978.1 | GPMDYTQAMRNASKGNYPYCYSEPMQGTTRCRLALYVVFESPFNMLCD 545
* : * : * : * : * : * : * : * : * : * : * : * : * : * : * :

gi | 60494642 | emb | CAH09443.1 | NPVYYYRELPCTEFITSVPTWDETKVLYAKVGEAVVAKRKGQWFI 579
gi | 29339180 | gb | AAO76978.1 | TPSNYMREPESTAFIAEIPVWDESIVLDGKMGEYIVTARRKGDVWVYVGG 595
. * * * * . * : * : * : * : * : * : * : * : * : * : * : * :

gi | 60494642 | emb | CAH09443.1 | ITGNQPONIEIDLGFIPAGQSFTLTSFEDGINADRQAMDYKKKESTVNNQ 629
gi | 29339180 | gb | AAO76978.1 | ITDWSARDIEVDCSFLGD-KSYHATLFKDGVNHRAGRDKCESFPIK 644
* . . : : * : * : * : * : * : * : * : * : * : * : * : * :

gi | 60494642 | emb | CAH09443.1 | TRMTLKMVRNGWAGTIKMK 649
gi | 29339180 | gb | AAO76978.1 | GKLKVHLAPGGGFALKIK-- 662
: : : : . . * : * . *

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Figure 48: Sequence alignment (of the CAH09443 amino acid residues and the AA076978 residues of *BtGH97b* showing the conserved aspartic acid residue (emboldened and underlined) positioned at 415 and 404 respectively using ClustalW). (<http://www.ebi.ac.uk/Tools/clustalw2/index.html>).

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APPENDICES

Appendix A- Chemicals, media and enzymes

This section of the Appendix contains a list of the chemicals, enzymes and media used in the study.

A1 Chemicals used in the study

Acros

Lactose

p-aminobenzoic acid

Cysteine

Imidazole

Iodoacetamide

Monosodium phosphate

Duchefa

Ethylenediaminetetraacetic acid disodium salt

Fisher BioReagents

Acrylamide/Bisacrylamide 37.5:1 40% solution

Brilliant blue G-250

Coomassie Blue R-250

Methanol

Fisher chemicals

Acetic acid, glacial

Acetone

Acetonitrile

Ammonium bicarbonate

Ammonium sulphate

Ethanol

Glycerol

D-(+)-Glucose

Ammonium per sulphate

N,N,N',N'-tetra methylethylene diamine (TEMED)

Phosphoric acid

Potassium dihydrogen phosphate

Propan-2-ol

Dipotassium hydrogen phosphate

Sodium carbonate

Sodium hydroxide

Sodium tetraborate

Melford

Agarose (High gel strength)

Dithiothreitol

Glycine

N-[2-hydroxyethylpiperazine-N']-[2-ethanesulphonic acid] (HEPES)

Isopropyl- β -D-thiogalactopyranoside

Kanamycin

Sodium chloride

Sodium dodecyl sulphate

Tris [Hydroxymethyl] aminomethane (Tris-HCl)

Urea

Promega

Trypsin Gold

Riedel-deHaen

Hydrochloric acid

Sulphuric acid

Sigma

L-Aspartic acid

Ammonium sulphate

Bovine serum albumin (BSA)

Bromophenol blue

CHAPS

Citric acid

Dipotassium phosphate

Ethidium bromide

Ferrous sulphate

Magnesium sulphate

Magnesium chloride

β -mercaptoethanol

Pantothenic acid

Potassium chloride

Xylene cyanol

Potassium monophosphate monobasic

Phenol

Resazurin

Riboflavin

Sodium acetate

Sodium chloride

Sodium hydroxide

Sodium sulphite

Calcium chloride

Cobalt chloride

Cupric chloride

Manganese chloride

Nickel (II) chloride

Barium chloride

Strontium chloride

Zinc chloride

Bradford's reagent

Porcine gastric Mucin Type II

Porcine gastric mucin Type III

Media:

Oxoid

Agar (Bacteriological agar N° 1)

NZ amine (casein hydrolysate)

Tryptone

Yeast extract

Columbia Agar base

Anaerobic CO₂ generation compact [CO₂ Gen™ (Atmosphere Generation System)]

Difco

Casitone (Bactopeptone)

BactAlert™ Anaerobic Basal media

Hampton Research

Aquasil

Amino acids and vitamins used in media:

Acros	Sigma	Fisher Scientific	BDH	Aldrich
L-alanine	L-arginine	L-lysine	4-aminobenzoic acid	Isoleucine
L-aspartic acid	Ascorbic acid		L(-)- Proline	L-tryptophan
L-glutamic acid	L-asparagine			
Pyridoxine	Folic acid			
Riboflavin	Glutamine			
L-serine	D(+) Biotin			
L-threonine	L-histidine			
L-valine	L-leucine			
	Niacinamide			
	Nicotinic acid			
	L-phenylalanine			
	L-tyrosine			

Crystallisation chemicals:

Ammonium acetate- Fisher chemicals

Ammonium chloride- Fisher

Tri ammonium citrate- Fisher

Ammonium citrate dibasic- Sigma

Ammonium formate- Acros

Ammonium fluoride- Sigma

Ammonium dihydrogen orthophosphate- Fisher

Ammonium nitrate- Fisher

Ammonium sulphate- Sigma

Ammonium tartrate dibasic- Sigma

Bicine- Acros

Bis-Tris- Melford

Bis-Tris propane- Sigma

1, 4- Butanediol- Acros

Caesium chloride- Fisher

Calcium acetate (dried) - Fisher

Calcium chloride hexahydrate- Fisher

CAPS free acid- Melford

CHES- Melford

Citric acid- Fisher

Cobalt (II) chloride- Fisher

1, 4- Dioxane- Fisher
Diammonium hydrogen citrate- Fisher
HEPES, free acid- Melford
Hexadecyl trimethyl ammonium bromide- Sigma
1, 6- Hexanediol- Acros, Fisher
Jeffamine- Fluka
Imidazole (GPG) – Fisher
Iron (II) chloride- Fisher
DL-Malic acid- Fisher
Tertiary butyl alcohol- Fisher
Barbitone sodium- BDH
Barium chloride- Sigma
Calcium sulphate- Fisher
Cadmium chloride- Acros
Ethylene glycol- Fisher
Formaldehyde (40%) – BDH
Formamide- BDH
Lithium acetate dehydrate- Acros
Lithium chloride anhydrous- Fisher
Lithium sulphate- Fisher
Magnesium acetate- Sigma
Magnesium chloride- Fisher
Magnesium formate dehydrate- Fluka

Magnesium nitrate- Fisher
Magnesium sulphate- Fisher
MES- Fisher
2-methyl-1, 2- pentanediol- Acros
Nickel (II) chloride- Fisone
Nickel chloride hexahydrate- sigma
Nickel sulphate- BDH
Nickel (II) sulphate hexahydrate- Sigma
Pentaerythritol ethoxylate- Aldrich
Phenylmethyl sulfonyl fluoride- Sigma
Poly (acrylic acid) – Acros
Polyethylene glycol- Fluka
Polyethylene glycol 4000 grade- Fisher
Polyethylene glycol 6000 grade- Fisher
Potassium dichromate-Melford
Potassium hydrogen tartrate- Fisher
Potassium iodide- Fisone
Sodium cacodylate trihydrate- Fisher
Sodium cyanoborohydride- Sigma
Thiomersal- BDH
Tri-lithium citrate tetrahydrate- Fluka
Sodium chloride- Fisher
Potassium thiocyanate- Fisher

Sodium acetate trihydrate- Fisher

Potassium-dihydrogen orthophosphate- Fisher

Potassium nitrate- Fisher

Sodium formate- Fisher

Potassium chloride- Fisher

Potassium iodide- Fisher

Sodium sulphate anhydrous- Fisher

Zinc acetate- Fisher

Sodium thiocyanate dehydrate- Fisher

Tri-sodium citrate- Fisher

Potassium sulphate- Fisher

Sodium selenite- Sigma

Polyethylenimine- Sigma

Zinc sulphate- Fisher

di- sodium tetraborate- Fisher

Succinic acid- Fisher

Potassium bromide- Fisher

Di-sodium hydrogen orthophosphate dodecahydrate- Fisher

Sodium nitrate- Fisher

Sodium bromide- Fisher

Sodium fluoride- Fisher

Zinc chloride- Fisher

Sodium succinate- Fisher

Tri-sodium citrate- Fisher

Potassium sodium tartrate- Fisher

Trimethylamine N-oxide dehydrate- Acros

Tri potassium citrate- Fisher

Tris Base Ultrapure- Melford

Sodium propionate- Fisher

Polyvinyl pyrrolidone K15- Fluka

Sodium dihydrogen orthophosphate dehydrate- Fisher

Sodium tartrate- Fisher

Potassium formate- Fluka

Potassium fluoride anhydrous- Fluka

Potassium acetate- Fisher

Sodium selenite- Fisher

Sodium acetate trihydrate- Fisher

Di-potassium hydrogen orthophosphate- Fisher

Polyethylene glycol 20000- Fluka

Polyethylene glycol 5000 monomethyl ether- Fluka

Propan-2-ol- BDH

Polypropylene glycol 2000 grade- Fisher

Propane-1, 2- diol- Fisher

Propylene glycol 400- Fluka

Pentaerythritol propoxylate- Aldrich

Polyethylene glycol 8000- Fisher

Polyethylene glycol 400 grade- Fisher

Polyethylene glycol 600 grade- Fisher

Polyethylene glycol 1000 grade- Fisher

Polyethylene glycol- Sigma

Polyethylene glycol 3000- Fluka

2-Ethoxyethanol- Fisher

Polyethylene glycol 200 grade- Fisher

Polyethylene glycol 300- Sigma

Glycerol- Fisher

Polyethylene glycol 2000 monomethyl ether- Fluka.

A2 Recipe for buffers used in the cloning experiment

Taq reaction buffer (10 X)

Tris pH 8.4 200 mM

KCl 500 mM

MgCl₂ 50 mM

T4 DNA polymerase reaction buffer (10 X)

Tris- HCl pH 8.8 67 mM

MgCl₂ 6.7 mM

(NH₄)₂SO₄ 16.6 mM

β mercaptoethanol 10 mM

EDTA 6.7 μM

BSA 170 µg/mL

KOD polymerase reaction buffer (10 X)

Tris HCl pH 7.5 at 25°C 20 mM

MgCl₂ 8 mM

DTT 7.5 mM

BSA 50 µg/mL

NE Buffer 2 (1 X)

NaCl 50 mM

Tris-HCl pH 7.9 at 25°C 10 mM

MgCl₂ 10 mM

DTT 1 mM

The buffer is supplied at a 10 X concentration.

Appendix B

Lab Equipment

Autoclaving

A benchtop Prestige® Medical 2100 Classic autoclave was used to sterilise solutions at 121°C, 32 lb/inch² pressure for 20 min.

pH meter

pH of solutions were adjusted to the required values using a Jenway Ion Meter 3340 calibrated with the standard buffer values of pH 4.0, 7.0 and 9.2.

Spectrophotometer

The Cecil 2000 Series Spectrophotometer was used to determine the cell density of cultures of bacteria at various stages of growth. The wavelength was set to 660 nm for bacterial cultures and 420 nm for determining the optical density of Bradford assay reaction mixtures.

The Hitach Fluorescence Spectrophotometer was used to determine the enzyme activity against methylumbelliferone substrates. The excitation wavelength was set at 360 nm and the emission wavelength was set at 460 nm.

Incubators

A Gallenkamp orbital shaker was used to grow liquid bacterial cultures and a Gallenkamp static incubator was used to grow bacteria on solid media. The..... anaerobic chamber was used to grow bacteria anaerobically at 37°C. The static incubator maintained a stable anaerobic environment by being flushed with a mixture of 10% carbon-di-oxide gas, 80% nitrogen gas and 10% hydrogen gas.

Freeze drier

The trypsinised protein samples containing digested peptides were lyophilised using the Christ® Alpha 1-2 freeze drier.

Centrifugation

The Sigma* 1-15 bench top microcentrifuge was used to centrifuge samples of upto 1.5 ml in volume. The Sigma 3K18C refrigerated bench top centrifuge was used to centrifuge larger volumes of cultures or solutions (>1.5 ml). The rotors used have been detailed as follows:

The rotor number 11133 was used to harvest cells at 4000 xg in clay universal holders (part No. 17049, 25 ml plastic universals were used), concentrate proteins at 4000 xg in 30 kDa cut off protein concentrators which were inserted into clay universal holders (part No. 17049). The rotor number 12158 was used to separate soluble cell extract from lysed cells at 14000 xg in 30 ml plastic centrifuge tubes (part No. 31190030). The rotor number 12131 was used to isolate DNA from the Qiagen kit and to extract proteins from lysed cells by centrifuging at 14000 xg in 1.5 ml microcentrifuge tubes.

Sonication

The MSE Soniprep 150-ultrasonication machine was used to lyse cells.

Protein purification

Large scale purification of proteins was carried out using an AKTA prime and the FPLC system which was controlled by Prime View Software.

PCR machine

Eppendorf MasterCycler™ PCR machine was used to perform all the PCR reactions.

Electroporator

A Bio-Rad Gene Pulser Xcell™ with an attached Bio-Rad Pc and CE module was used for the transformation of electrocompetent cells. The Bio-Rad ShockPod™ was used to place the cuvettes.

Agarose gel kits

Owl Scientific EASY-CAST electrophoresis system was used for the electrophoresis of DNA and the power supply was an E-C 570-90 E-C Apparatus Corporation power pack.

2D Clean up kit from Amersham Biosciences

The solutions used in the Clean up kit include the precipitant solution which was used to in solubilise proteins, the co-precipitant solution which worked with the precipitant solution in enhancing the precipitation of proteins, the wash buffer to get rid of non protein contaminants and the wash additive to facilitate rapid and complete resuspension of the proteins in the solution. Due to trade reasons, the company have not revealed the exact composition of these solutions.

SDS-PAGE gel kits for small gels

A Bio-Rad Mini-PROTEAN 3 Cell kit was used for the electrophoresis of proteins and the same E-C 570-90 E-C Apparatus corporation power pack was used.

SDS-PAGE kit for 2D analysis of large gels

Protean II XL 2-D cells and accessories were used to perform the electrophoresis. The kit included a tank and a lid, a central cooling core unit, casting stand, sandwich clamps, alignment card and combs. The accessories provided include a 4 mm xi clamp notch vs. 13 mm XL clamp notch, 19 mm xi spacer vs. 8 mm XL spacer, 181 mm xi core gasket vs. 198 mm XL core gasket and a 153 mm xi comb vs. 184 mm XL comb.

The molecular biology kits used in the study include the Qiagen Miniprep kit for the purification of recombinant plasmids, the Qiagen Maxiprep kit for the large scale preparation of cloning vectors, the Qiagen Qiaquick gel extraction kit for the purification of DNA from agarose gels and the Qiagen DNeasy kit for the purification of genomic DNA.

Immobiline DryStrip Kit

Immobiline™ DryStrip gels of pH range 4-7 and 3-10 were ordered as a part of the electrophoresis kit. The other parts include the electrodes (anode and cathode), tray and electrode holder, DryStrip aligners, IEF electrode strips and the Bio-Rad power pack 200 and Universal power supply pack for protein and nucleic acid electrophoresis.

Gel Documentation

The Quantity One™ software in the Bio-Rad Gel Doc 2000 system was used to visualise the agarose gels. The Bio-Rad GS-800 densitometer was used to visualise SDS-PAGE gels. Hard copies of the gel were obtained using the Mitsubishi Video Copy Processor (Model P91) with Mitsubishi thermal paper (K65HM-CE/High density type, 110 mm X 21 m).

Gel spot analysis

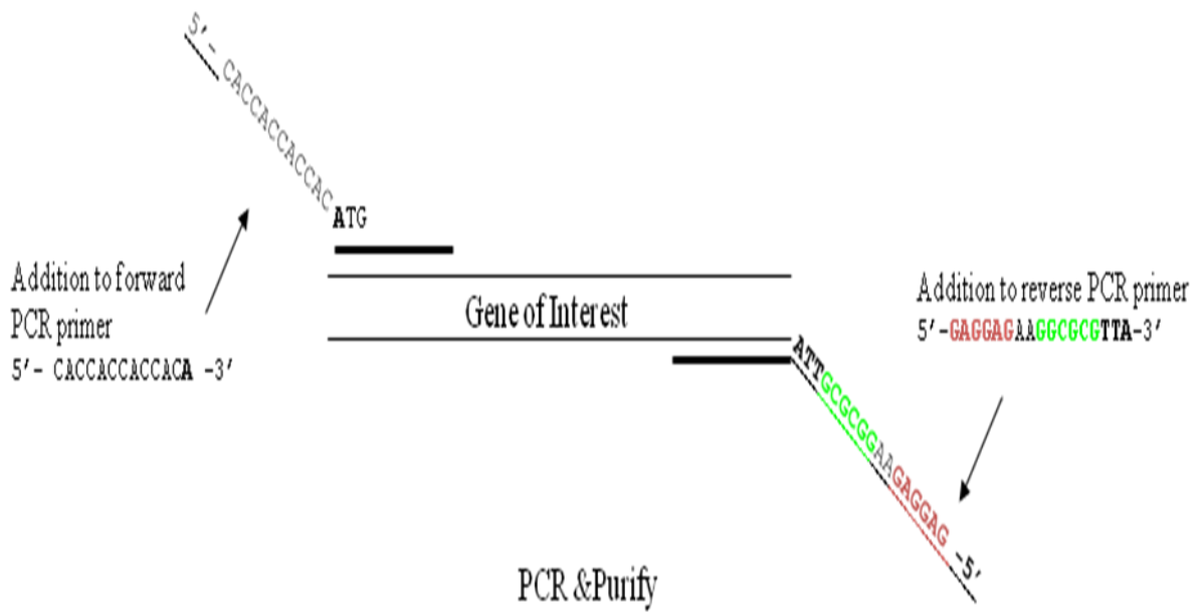
The spots on the gels were compared and analysed using the Advanced PDQuest™ 8.0 software in the Bio-Rad Gel Doc 2000 system.

LC-MS analysis

The Ultimate 3000 Dionex LC system was used to separate the peptide fragments before Mass spectrometric analysis. The LC system was equipped with a vacuum degasser, a UV cell compartment, flow manager, column compartment fitted with an oven maintained at 60°C and the autosampler system with a multiwell sample microtitre plate.

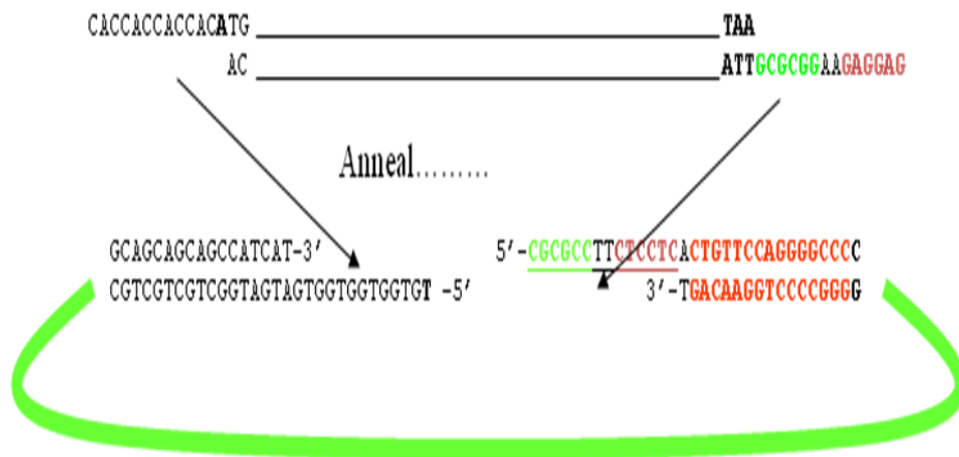
Mass spectrometric analysis

The peptides of interest were identified using the Bruker Daltonics ESI-ion trap Mass spectrometer equipped with the Esquire series, Hystar sample processing, Chromeleon LC system and the Biotools software for Mascot database searches.



CACCACCACCACATG _____ TAA CGCGCCTTCTCCTC
 GTGGTGGTGGTGTAC _____ ATT CGCGGAAAGAGGAG

Treat with T4 pol and dATP



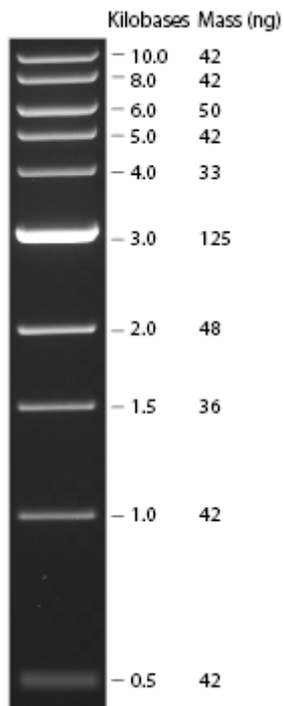
...then transform

Appendix D

Protein and DNA size standards

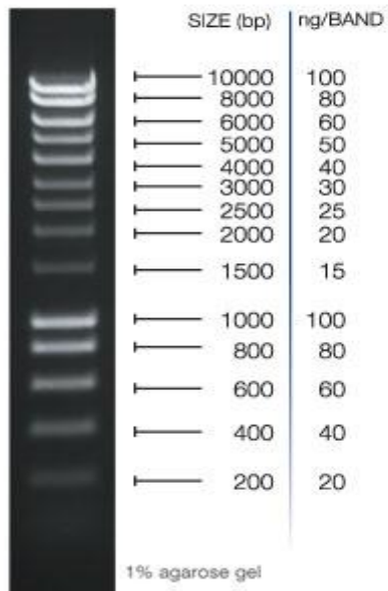
Proteins	Molecular weight	High range (S8320)	Wide range (S8445)	Low range (M3913)
Myosin from rabbit muscle	200000	X	X	
B-galactosidase from <i>E. coli</i>	116000	X	X	
Phosphorylase b from rabbit muscle	97000	X	X	
Albumin, bovine serum	66000	X	X	X
Glutamic dehydrogenase from bovine liver	55000	X	X	
Ovalbumin from chicken egg	45000	X	X	X
Glyceraldehyde-3-phosphate dehydrogenase from rabbit muscle	36000	X	X	X
Carbonic anhydrase from bovine erythrocytes	29000		X	X
Trypsinogen from bovine pancreas	24000		X	X
Trypsin inhibitor from soyabean	20000		X	X
α -lactalbumin from bovine milk	14200		X	X
Aprotinin from bovine lung	6500		X	X

High and low molecular weight markers for SDS PAGE gels



NEB 1 kb Ladder used to determine the mass of the DNA fragments.

HyperLadder I



- Higher intensity bands:
1000bp and 10000bp
- Supplied in a ready-to-use format
- Each lane (5 μ l) provides
720ng of DNA

Bioline hyperladder I used as a size standard for agarose gel electrophoresis.

Appendix E

Growth curve data for *E. cancerogenus* and *B. fragilis*

Time in h	Absorbance at 600 nm- Experiment 1	Absorbance at 600 nm- Experiment 2	Absorbance at 600 nm- Experiment 3	Average absorbance at 600 nm
0	0	0	0	0
2	0.081	0.083	0.087	0.083667
3	0.416	0.58	0.55	0.515333
4	0.789	0.882	0.831	0.834
5	1.156	1.288	1.186	1.21
6	1.386	1.415	1.306	1.369
7	1.412	1.426	1.364	1.400667
8	1.434	1.445	1.474	1.451

Table 5: Growth of *E. cancerogenus* in control media without mucin

Time in h	Absorbance at 600 nm- Experiment 4	Absorbance at 600 nm- Experiment 5	Absorbance at 600 nm- Experiment 6	Average absorbance at 600 nm
0	0	0	0	0
2	0.053	0.076	0.108	0.079
3	0.21	0.256	0.257	0.241
4	0.931	0.794	0.786	0.837
5	1.264	1.132	1.188	1.194667
6	1.38	1.31	1.256	1.315333
7	1.392	1.370	1.376	1.379333
8	1.396	1.350	1.386	1.377333

Table 6: Growth of *E. cancerogenus* in media enriched with mucin Type II

Time in h	Absorbance at 600 nm- Experiment 7	Absorbance at 600 nm- Experiment 8	Absorbance at 600 nm- Experiment 9	Average absorbance at 600 nm
0	0	0	0	0
2	0.104	0.107	0.072	0.094333
3	0.31	0.292	0.280	0.294
4	0.901	0.897	0.807	0.868333
5	1.106	1.116	1.152	1.124667
6	1.186	1.242	1.264	1.230667
7	1.208	1.144	1.2	1.184
8	1.188	1.156	1.204	1.182667

Table 7: Growth of *E. cancerogenus* in media with mucin Type III

Time in h	Absorbance at 600 nm- Experiment 10	Absorbance at 600 nm- Experiment 11	Absorbance at 600 nm- Experiment 12	Average absorbance at 600 nm
0	0	0	0	0
4	0.067	0.087	0.059	0.071
6	0.112	0.1	0.158	0.123333
21	0.313	0.286	0.307	0.302
22	0.439	0.383	0.426	0.416
23	0.748	0.596	0.619	0.654333
24	1.004	0.909	0.93	0.947667
25	1.018	1.052	1.069	1.046333
26	1.394	1.298	1.243	1.311667
27	1.526	1.483	1.418	1.475667
28	1.556	1.552	1.512	1.54
29	1.488	1.418	1.385	1.430333
30	1.452	1.448	1.422	1.440667
31	1.475	1.466	1.486	1.475667
46	1.402	1.412	1.423	1.412333
47	1.328	1.396	1.404	1.376

Table 8: Growth of *B. fragilis* in control media without mucin

Time in h	Absorbance at 600 nm- Experiment 13	Absorbance at 600 nm- Experiment 14	Absorbance at 600 nm- Experiment 15	Average absorbance at 600 nm
0	0	0	0	0
4	0.012	0.018	0.015	0.015
6	0.05	0.041	0.045	0.045333
21	0.098	0.106	0.102	0.102
22	0.206	0.236	0.226	0.222667
23	0.34	0.358	0.36	0.352667
24	0.397	0.412	0.422	0.410333
25	0.532	0.622	0.603	0.585667
26	0.718	0.814	0.798	0.776667
27	0.922	1.039	0.956	0.972333
28	1.294	1.334	1.306	1.311333
29	1.433	1.461	1.457	1.450333
30	1.359	1.398	1.405	1.387333
31	1.382	1.368	1.393	1.381
46	1.401	1.387	1.384	1.390667
47	1.385	1.355	1.376	1.372

Table 9: Growth of *B. fragilis* in media with mucin Type II

Time in h	Absorbance at 600 nm- Experiment 16	Absorbance at 600 nm- Experiment 17	Absorbance at 600 nm- Experiment 18	Average absorbance at 600 nm
0	0	0	0	0
4	0.029	0.023	0.019	0.023667
6	0.063	0.052	0.058	0.057667
21	0.165	0.158	0.162	0.161667
22	0.384	0.4	0.391	0.391667
23	0.598	0.675	0.656	0.643
24	0.738	0.842	0.823	0.801
25	1.012	1.058	1.035	1.035
26	1.328	1.338	1.332	1.332667
27	1.438	1.463	1.457	1.452667
28	1.402	1.417	1.423	1.414
29	1.334	1.393	1.404	1.377
30	1.374	1.355	1.369	1.366
31	1.348	1.382	1.375	1.368333
46	1.368	1.376	1.36	1.368
47	1.284	1.318	1.307	1.303

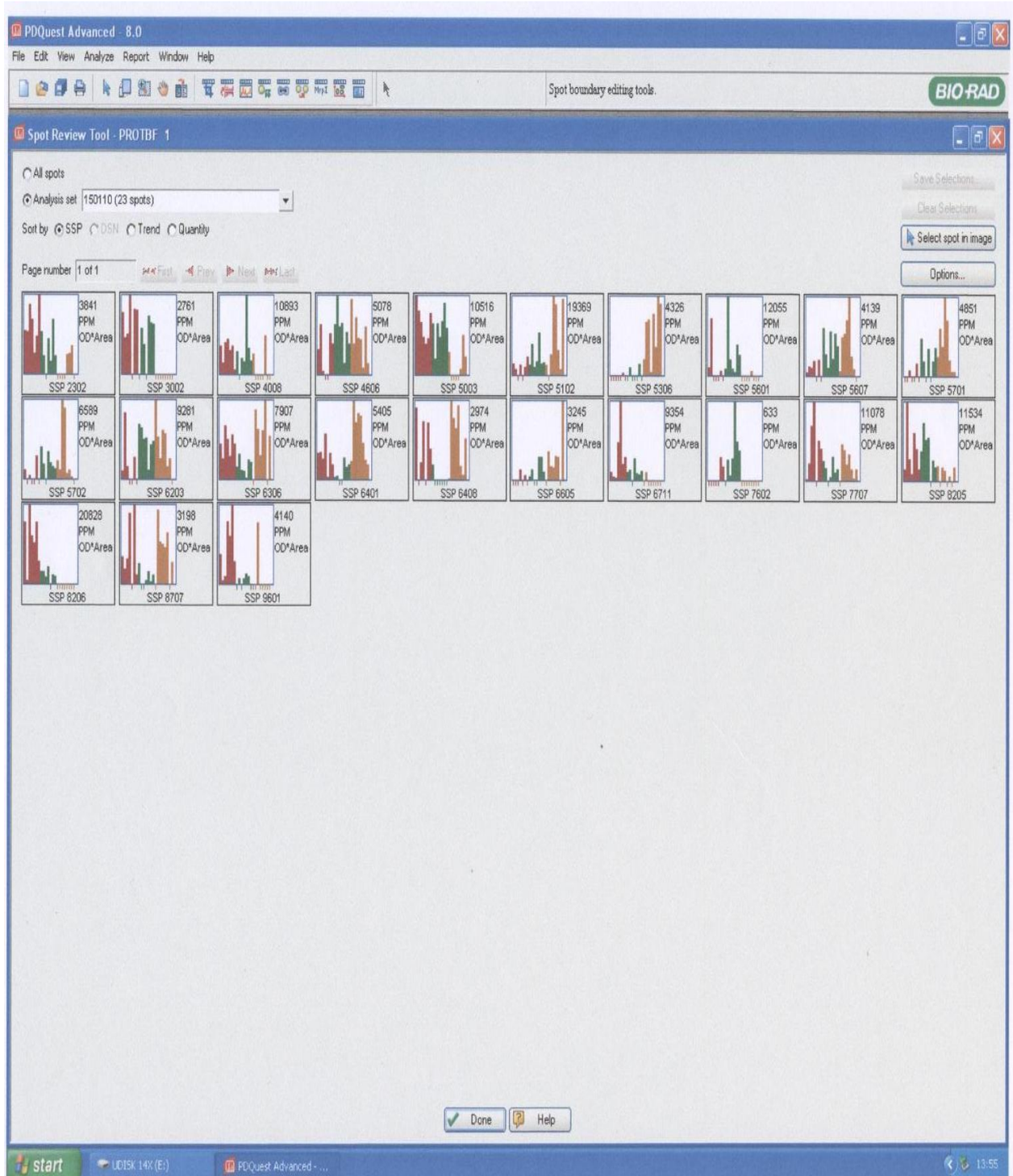
Table 10: Growth of *B. fragilis* in media with mucin Type III

E1 Resazurin test method

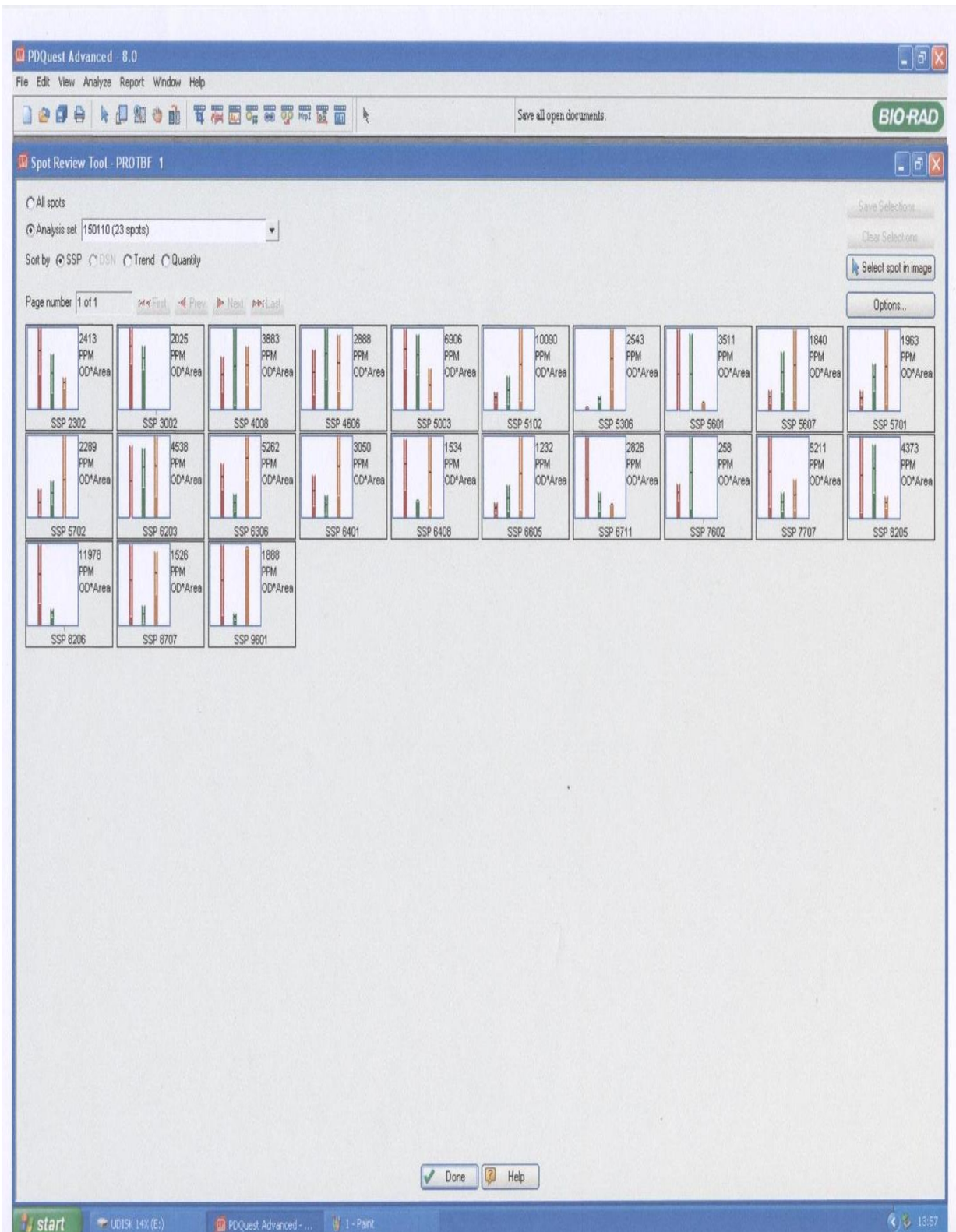
A 0.001% resazurin solution was prepared by dissolving 1 mg of resazurin and 2 g of cysteine-HCl in sterile 18.2 MΩ/cm making it up to 100 ml. The solution was aliquoted into 5 ml quantities and used as indicators. The resazurin strips or solutions were pink to purple in the presence of oxygen and gradually turn colourless when left in an environment lacking oxygen. This was normally used as a test to confirm the absence of oxygen in the bacterial growth environment.

Appendix F- Analysis data for 2DE gels

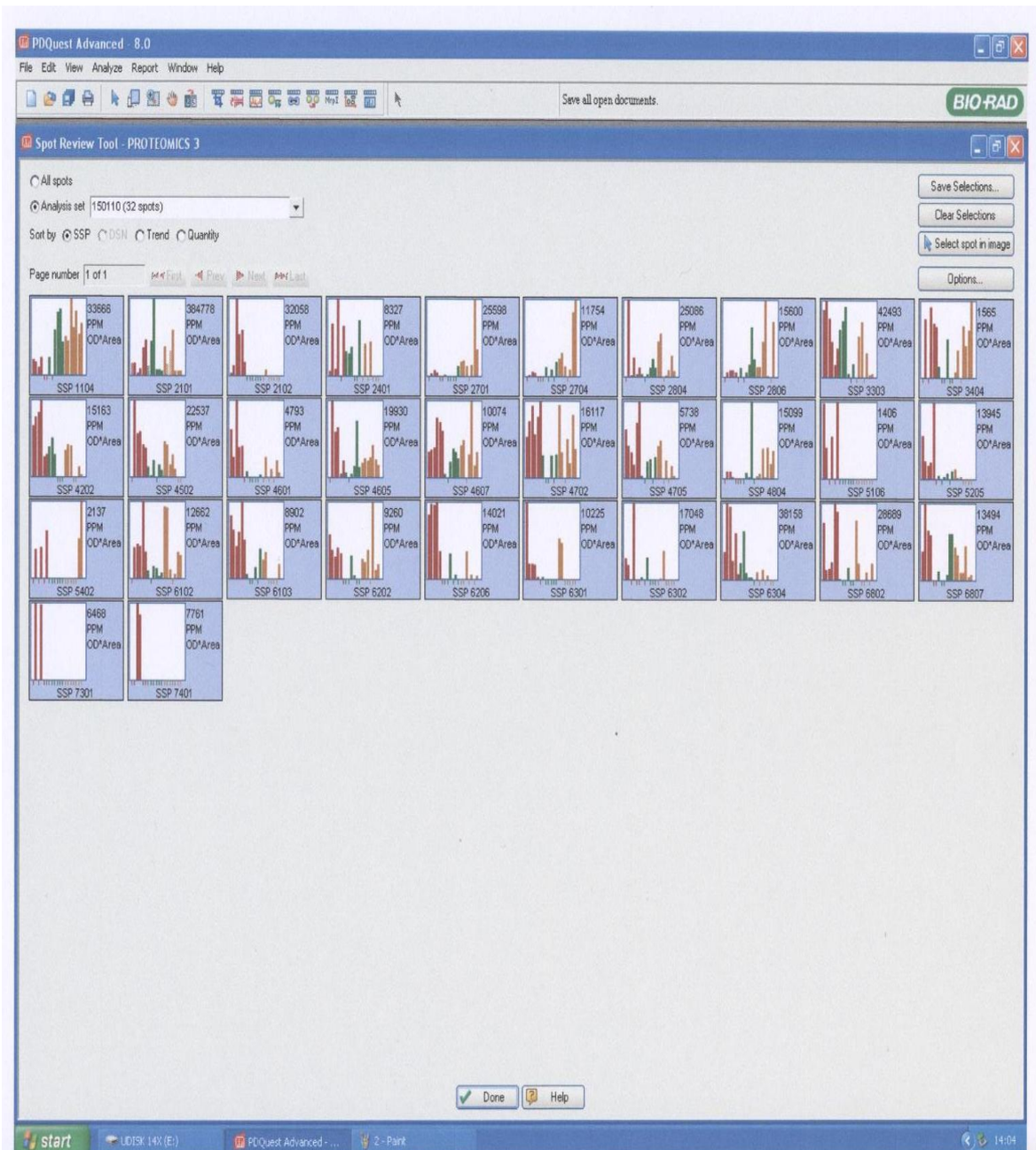
F1 Gel analysis data for *E. cancerogenus* and *B. fragilis* using Bio-Rad PDQuest software



'Protbf 1' Protein quantities in individual gels of differentially expressed spots in three different growth conditions. Red bars indicate non-mucin media, Green bars- mucin Type II enriched media, Brown bars- mucin Type III enriched media.



'Proteomics 3' Average quantities of differentially expressed spots in three different growth conditions. Red bars indicate non-mucin media, Green bars- mucin Type II enriched media, Brown bars- mucin Type III enriched media.



'Proteomics 3' Protein quantities in individual gels of differentially expressed spots in three different growth conditions. Red bars indicate non-mucin media, Green bars- mucin Type II enriched media, Brown bars- mucin Type III enriched media.

F2 Gel analysis data for *E. cancerogenus* and *B. fragilis* using Ludesi Redfin software

Experiment overview

Experiment	PROTBF 1
Comparison	NON MUCIN MEDIA RD EC
Groups	MUCIN TYPE II MEDIA RD EC, NON MUCIN MEDIA RD EC, MUCIN TYPE III MEDIA RD EC

Experiment statistics

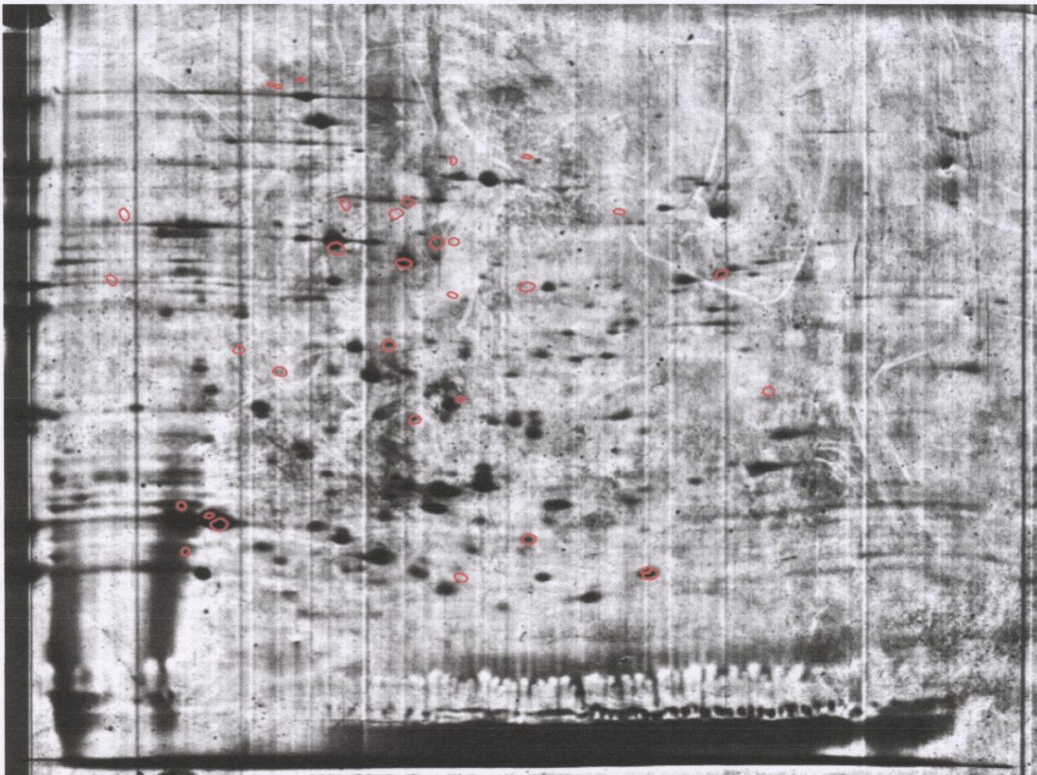
Statistical test	Anova
Applied filters	Anova < 0.05
	Presence \geq 100
	Fold change > 1.5
	Location filter (top-left: 203, 238 to lower-right: 2323, 2853)
Number filtered spots	32

Group color legend

 MUCIN TYPE II MEDIA RD EC	 NON MUCIN MEDIA RD EC	 MUCIN TYPE III MEDIA RD EC
---	---	--

Protein overview

Overview gel name: Scanned gel 10 (2) RD EC



Overview

Filtered spots

ID	619						
Anova	0.0470						
Presence	100 %						
Fold	2.01						
Volume	323						
ID	733						
Anova	0.0459						
Presence	100 %						
Fold	1.81						
Volume	468						
ID	127						★
Anova	0.0450						
Presence	100 %						
Fold	4.41						
Volume	2140						
ID	320						
Anova	0.0434						
Presence	100 %						
Fold	1.79						
Volume	1608						
ID	681						
Anova	0.0433						
Presence	100 %						
Fold	2.67						
Volume	655						
ID	583						★
Anova	0.0430						
Presence	100 %						
Fold	3.03						
Volume	1182						
ID	695						
Anova	0.0425						
Presence	100 %						
Fold	2.32						
Volume	650						
ID	636						
Anova	0.0418						
Presence	100 %						
Fold	2.01						
Volume	2436						
ID	24						
Anova	0.0354						
Presence	100 %						
Fold	1.54						
Volume	27676						
ID	445						
Anova	0.0345						
Presence	100 %						
Fold	2.46						
Volume	1025						

ID	672						★
Anova	0.0324						
Presence	100 %						
Fold	3.44						
Volume	366						
ID	87						
Anova	0.0296						
Presence	100 %						
Fold	2.58						
Volume	1795						
ID	483						
Anova	0.0262						
Presence	100 %						
Fold	2.93						
Volume	609						
ID	551						★
Anova	0.0250						
Presence	100 %						
Fold	4.36						
Volume	667						
ID	210						
Anova	0.0247						
Presence	100 %						
Fold	2.55						
Volume	1400						
ID	565						
Anova	0.0242						
Presence	100 %						
Fold	2.45						
Volume	635						
ID	684						
Anova	0.0204						
Presence	100 %						
Fold	1.64						
Volume	610						
ID	58						★
Anova	0.0202						
Presence	100 %						
Fold	3.32						
Volume	7216						
ID	131						High fold increase Type III ★
Anova	0.0144						
Presence	100 %						
Fold	4.57						
Volume	1871						
ID	307						
Anova	0.0143						
Presence	100 %						
Fold	2.40						
Volume	1247						
ID	548						
Anova	0.0143						
Presence	100 %						
Fold	2.85						
Volume	871						

ID	631					
Anova	0.0116					
Presence	100 %					
Fold	2.55					
Volume	245					
ID	130					
Anova	0.0100					
Presence	100 %					
Fold	6.67					
Volume	862					
						High fold increase in Mucin III
ID	297					
Anova	0.0089					
Presence	100 %					
Fold	2.51					
Volume	1325					
ID	384					
Anova	0.0085					
Presence	100 %					
Fold	2.18					
Volume	1205					
ID	137					
Anova	0.0050					
Presence	100 %					
Fold	2.71					
Volume	1208					
ID	351					
Anova	0.0047					
Presence	100 %					
Fold	2.75					
Volume	1312					
ID	1					
Anova	0.0033					
Presence	100 %					
Fold	1.73					
Volume	24393					
ID	48					
Anova	0.0026					
Presence	100 %					
Fold	2.17					
Volume	9679					
ID	437					
Anova	0.0020					
Presence	100 %					
Fold	2.25					
Volume	1262					
ID	497					
Anova	0.0020					
Presence	100 %					
Fold	2.13					
Volume	705					
ID	238					
Anova	0.0016					
Presence	100 %					
Fold	2.33					
Volume	1956					

Experiment overview

Experiment	PROTEOMICS 3
Comparison	PROTEOMICS 3
Groups	NON MUCIN MEDIA RD, MUCIN TYPE II MEDIA RD, MUCIN TYPE III MEDIA RD

Experiment statistics

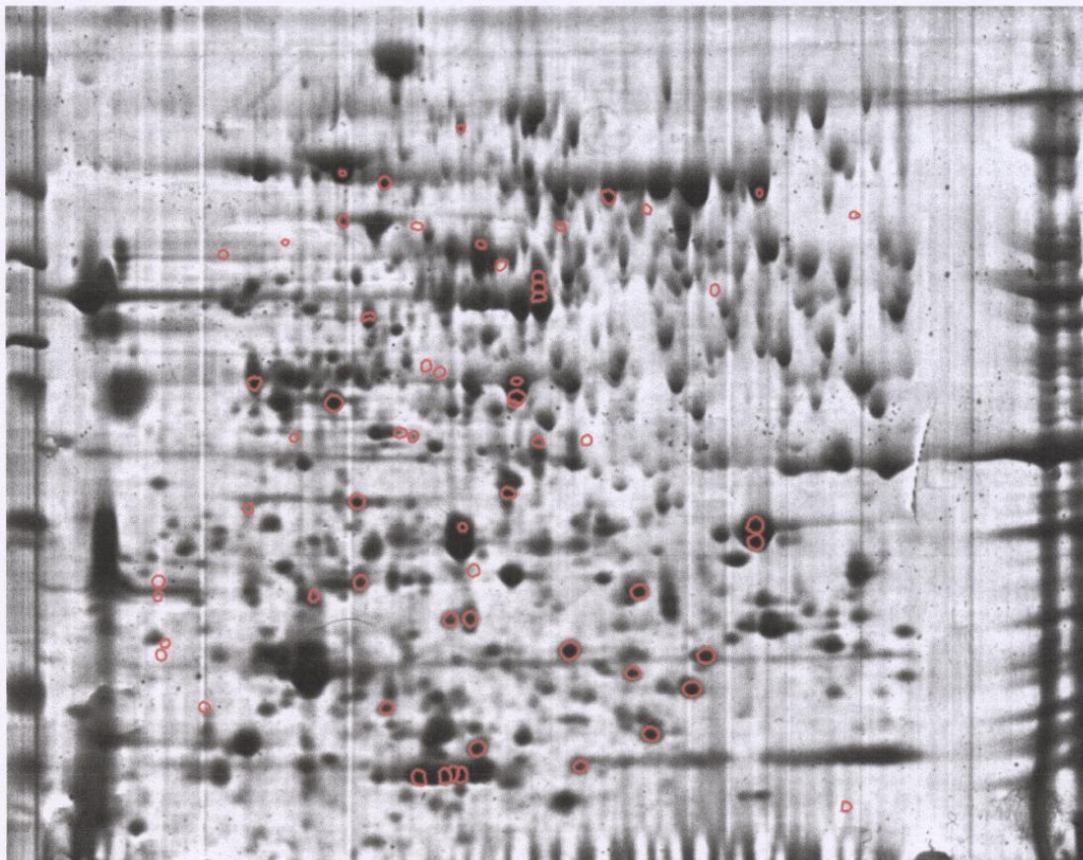
Statistical test	Anova
Applied filters	Anova < 0.05 Presence >= 100 Fold change > 1.5 Location filter (top-left: 87, 195 to lower-right: 2272, 2755)
Number filtered spots	61

Group color legend

 NON MUCIN MEDIA RD	 MUCIN TYPE II MEDIA RD	 MUCIN TYPE III MEDIA RD
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Protein overview

Overview gel name: Staff 2008-10-24 14hr 21min (10) 4 RD



Overview

Filtered spots

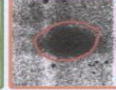
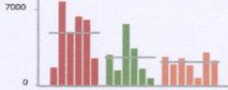
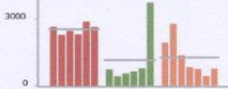
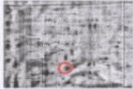
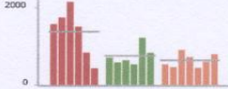
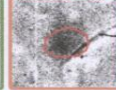
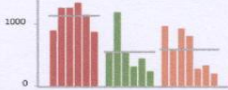
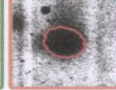
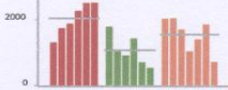
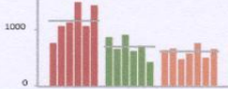
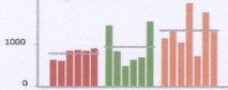
ID	10						★
Anova	0.0028						
Presence	100 %						
Fold	7.04						
Volume	4270						
ID	17						★
Anova	0.0156						
Presence	100 %						
Fold	6.78						
Volume	4047						
ID	138						★
Anova	0.0204						
Presence	100 %						
Fold	6.56						
Volume	1230						
ID	8						
Anova	0.0375						
Presence	100 %						
Fold	5.16						
Volume	636						
ID	141						★
Anova	2.848e-6						
Presence	100 %						
Fold	5.07						
Volume	1206						
ID	300						
Anova	0.0233						
Presence	100 %						
Fold	4.99						
Volume	1999						
ID	7						★
Anova	0.0176						
Presence	100 %						
Fold	4.69						
Volume	4147						
ID	104						★
Anova	1.734e-4						
Presence	100 %						
Fold	4.62						
Volume	4452						
ID	3						
Anova	0.0365						
Presence	100 %						
Fold	4.32						
Volume	879						
ID	253						
Anova	0.0138						
Presence	100 %						
Fold	4.30						
Volume	541						

ID	893					
Anova	0.0074					
Presence	100 %					
Fold	4.00					
Volume	237					
ID	558					
Anova	0.0072					
Presence	100 %					
Fold	3.88					
Volume	1378					
ID	709					
Anova	0.0091					
Presence	100 %					
Fold	3.85					
Volume	508					
ID	126					
Anova	0.0162					
Presence	100 %					
Fold	3.78					
Volume	2211					
ID	267					
Anova	0.0025					
Presence	100 %					
Fold	3.76					
Volume	2064					
ID	24					
Anova	0.0242					
Presence	100 %					
Fold	3.74					
Volume	1818					
ID	149					
Anova	0.0139					
Presence	100 %					
Fold	3.66					
Volume	5225					
ID	645					
Anova	0.0185					
Presence	100 %					
Fold	3.55					
Volume	731					
ID	415					
Anova	0.0055					
Presence	100 %					
Fold	3.43					
Volume	874					
ID	402					
Anova	0.0220					
Presence	100 %					
Fold	3.43					
Volume	587					
ID	776					
Anova	0.0422					
Presence	100 %					
Fold	3.26					
Volume	605					

ID	578						★
Anova	0.0101						
Presence	100 %						
Fold	3.26						
Volume	711						
ID	80						
Anova	8.341e-4						
Presence	100 %						
Fold	3.26						
Volume	3433						
ID	983						★
Anova	0.0431						
Presence	100 %						
Fold	3.24						
Volume	460						
ID	573						
Anova	0.0156						
Presence	100 %						
Fold	3.24						
Volume	581						
ID	342						
Anova	0.0239						
Presence	100 %						
Fold	3.23						
Volume	629						
ID	512						
Anova	0.0171						
Presence	100 %						
Fold	3.11						
Volume	829						
ID	547						
Anova	7.293e-4						
Presence	100 %						
Fold	3.11						
Volume	1365						
ID	276						
Anova	0.0205						
Presence	100 %						
Fold	3.07						
Volume	507						
ID	1074						★
Anova	0.0163						
Presence	100 %						
Fold	3.07						
Volume	557						
ID	603						
Anova	0.0239						
Presence	100 %						
Fold	3.07						
Volume	581						
ID	399						★
Anova	3.558e-4						
Presence	100 %						
Fold	3.07						
Volume	1869						

ID	768					
Anova	0.0021					
Presence	100 %					
Fold	2.96					
Volume	754					
ID	610					
Anova	0.0449					
Presence	100 %					
Fold	2.95					
Volume	399					
ID	991					
Anova	0.0413					
Presence	100 %					
Fold	2.95					
Volume	600					
ID	960					
Anova	0.0165					
Presence	100 %					
Fold	2.92					
Volume	706					
ID	270					
Anova	0.0037					
Presence	100 %					
Fold	2.89					
Volume	1649					
ID	753					
Anova	0.0383					
Presence	100 %					
Fold	2.88					
Volume	283					
ID	6					
Anova	0.0164					
Presence	100 %					
Fold	2.84					
Volume	1385					
ID	1008					
Anova	0.0378					
Presence	100 %					
Fold	2.75					
Volume	727					
ID	218					
Anova	0.0126					
Presence	100 %					
Fold	2.71					
Volume	1895					
ID	524					
Anova	0.0096					
Presence	100 %					
Fold	2.70					
Volume	364					
ID	1004					
Anova	0.0381					
Presence	100 %					
Fold	2.70					
Volume	364					

ID	815					
Anova	0.0136					
Presence	100 %					
Fold	2.69					
Volume	580					
ID	305					
Anova	0.0211					
Presence	100 %					
Fold	2.68					
Volume	1346					
ID	891					
Anova	0.0252					
Presence	100 %					
Fold	2.67					
Volume	379					
ID	244					
Anova	0.0034					
Presence	100 %					
Fold	2.66					
Volume	1905					
ID	287					
Anova	0.0101					
Presence	100 %					
Fold	2.65					
Volume	1357					
ID	708					
Anova	0.0211					
Presence	100 %					
Fold	2.62					
Volume	1976					
ID	819					
Anova	0.0475					
Presence	100 %					
Fold	2.50					
Volume	348					
ID	34					
Anova	0.0479					
Presence	100 %					★
Fold	2.47					
Volume	4074					
ID	238					
Anova	0.0271					
Presence	100 %					
Fold	2.46					
Volume	1498					
ID	140					
Anova	0.0226					
Presence	100 %					
Fold	2.41					
Volume	1211					
ID	1063					
Anova	0.0250					
Presence	100 %					
Fold	2.34					
Volume	746					

ID	87						
Anova	0.0301						
Presence	100 %						
Fold	2.28						
Volume	3128						
ID	466						
Anova	0.0264						
Presence	100 %						
Fold	2.22						
Volume	1636						
ID	539						
Anova	0.0096						
Presence	100 %						
Fold	2.14						
Volume	907						
ID	712						
Anova	0.0052						
Presence	100 %						
Fold	2.06						
Volume	748						
ID	563						
Anova	0.0120						
Presence	100 %						
Fold	1.91						
Volume	1553						
ID	531						
Anova	2.280e-4						
Presence	100 %						
Fold	1.90						
Volume	810						
ID	877						
Anova	0.0441						
Presence	100 %						
Fold	1.70						
Volume	1036						

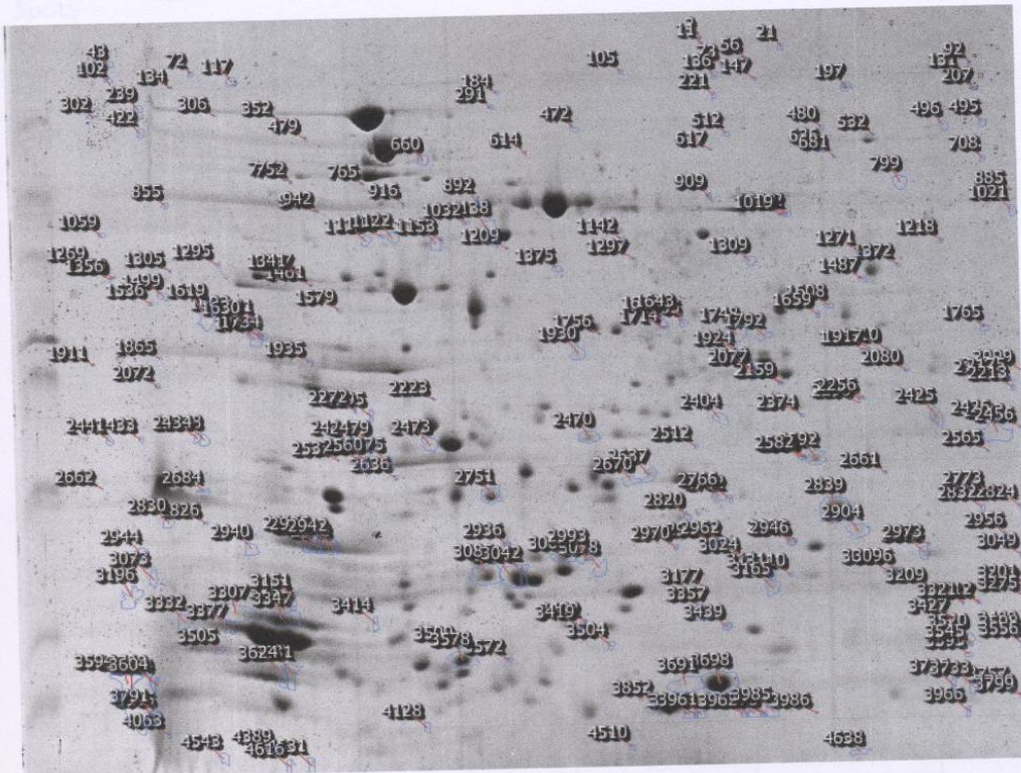
F3 Gel analysis data for *E. cancerogenus* and *B. fragilis* using NonLinear Dynamics SameSpots software

PROTBF 1

Experiment: PROTBF 1

Report created: 16/01/2010 22:18:43

Reference image



Spot ID	Intensity	pI	pK	MW (kDa)	pI (theoretical)	pK (theoretical)	MW (theoretical)
11	0.001	4.5	5.5	1.25	4.5	5.5	1.25
21	0.001	4.5	5.5	1.25	4.5	5.5	1.25
31	0.001	4.5	5.5	1.25	4.5	5.5	1.25
41	0.001	4.5	5.5	1.25	4.5	5.5	1.25
51	0.001	4.5	5.5	1.25	4.5	5.5	1.25
61	0.001	4.5	5.5	1.25	4.5	5.5	1.25
71	0.001	4.5	5.5	1.25	4.5	5.5	1.25
81	0.001	4.5	5.5	1.25	4.5	5.5	1.25
91	0.001	4.5	5.5	1.25	4.5	5.5	1.25
101	0.001	4.5	5.5	1.25	4.5	5.5	1.25
111	0.001	4.5	5.5	1.25	4.5	5.5	1.25
121	0.001	4.5	5.5	1.25	4.5	5.5	1.25
131	0.001	4.5	5.5	1.25	4.5	5.5	1.25
141	0.001	4.5	5.5	1.25	4.5	5.5	1.25
151	0.001	4.5	5.5	1.25	4.5	5.5	1.25
161	0.001	4.5	5.5	1.25	4.5	5.5	1.25
171	0.001	4.5	5.5	1.25	4.5	5.5	1.25
181	0.001	4.5	5.5	1.25	4.5	5.5	1.25
191	0.001	4.5	5.5	1.25	4.5	5.5	1.25
201	0.001	4.5	5.5	1.25	4.5	5.5	1.25
211	0.001	4.5	5.5	1.25	4.5	5.5	1.25
221	0.001	4.5	5.5	1.25	4.5	5.5	1.25
231	0.001	4.5	5.5	1.25	4.5	5.5	1.25
241	0.001	4.5	5.5	1.25	4.5	5.5	1.25
251	0.001	4.5	5.5	1.25	4.5	5.5	1.25
261	0.001	4.5	5.5	1.25	4.5	5.5	1.25
271	0.001	4.5	5.5	1.25	4.5	5.5	1.25
281	0.001	4.5	5.5	1.25	4.5	5.5	1.25
291	0.001	4.5	5.5	1.25	4.5	5.5	1.25
301	0.001	4.5	5.5	1.25	4.5	5.5	1.25
311	0.001	4.5	5.5	1.25	4.5	5.5	1.25
321	0.001	4.5	5.5	1.25	4.5	5.5	1.25
331	0.001	4.5	5.5	1.25	4.5	5.5	1.25
341	0.001	4.5	5.5	1.25	4.5	5.5	1.25
351	0.001	4.5	5.5	1.25	4.5	5.5	1.25
361	0.001	4.5	5.5	1.25	4.5	5.5	1.25
371	0.001	4.5	5.5	1.25	4.5	5.5	1.25
381	0.001	4.5	5.5	1.25	4.5	5.5	1.25
391	0.001	4.5	5.5	1.25	4.5	5.5	1.25
401	0.001	4.5	5.5	1.25	4.5	5.5	1.25
411	0.001	4.5	5.5	1.25	4.5	5.5	1.25
421	0.001	4.5	5.5	1.25	4.5	5.5	1.25
431	0.001	4.5	5.5	1.25	4.5	5.5	1.25
441	0.001	4.5	5.5	1.25	4.5	5.5	1.25
451	0.001	4.5	5.5	1.25	4.5	5.5	1.25
461	0.001	4.5	5.5	1.25	4.5	5.5	1.25
471	0.001	4.5	5.5	1.25	4.5	5.5	1.25
481	0.001	4.5	5.5	1.25	4.5	5.5	1.25
491	0.001	4.5	5.5	1.25	4.5	5.5	1.25
501	0.001	4.5	5.5	1.25	4.5	5.5	1.25
511	0.001	4.5	5.5	1.25	4.5	5.5	1.25
521	0.001	4.5	5.5	1.25	4.5	5.5	1.25
531	0.001	4.5	5.5	1.25	4.5	5.5	1.25
541	0.001	4.5	5.5	1.25	4.5	5.5	1.25
551	0.001	4.5	5.5	1.25	4.5	5.5	1.25
561	0.001	4.5	5.5	1.25	4.5	5.5	1.25
571	0.001	4.5	5.5	1.25	4.5	5.5	1.25
581	0.001	4.5	5.5	1.25	4.5	5.5	1.25
591	0.001	4.5	5.5	1.25	4.5	5.5	1.25
601	0.001	4.5	5.5	1.25	4.5	5.5	1.25
611	0.001	4.5	5.5	1.25	4.5	5.5	1.25
621	0.001	4.5	5.5	1.25	4.5	5.5	1.25
631	0.001	4.5	5.5	1.25	4.5	5.5	1.25
641	0.001	4.5	5.5	1.25	4.5	5.5	1.25
651	0.001	4.5	5.5	1.25	4.5	5.5	1.25
661	0.001	4.5	5.5	1.25	4.5	5.5	1.25
671	0.001	4.5	5.5	1.25	4.5	5.5	1.25
681	0.001	4.5	5.5	1.25	4.5	5.5	1.25
691	0.001	4.5	5.5	1.25	4.5	5.5	1.25
701	0.001	4.5	5.5	1.25	4.5	5.5	1.25
711	0.001	4.5	5.5	1.25	4.5	5.5	1.25
721	0.001	4.5	5.5	1.25	4.5	5.5	1.25
731	0.001	4.5	5.5	1.25	4.5	5.5	1.25
741	0.001	4.5	5.5	1.25	4.5	5.5	1.25
751	0.001	4.5	5.5	1.25	4.5	5.5	1.25
761	0.001	4.5	5.5	1.25	4.5	5.5	1.25
771	0.001	4.5	5.5	1.25	4.5	5.5	1.25
781	0.001	4.5	5.5	1.25	4.5	5.5	1.25
791	0.001	4.5	5.5	1.25	4.5	5.5	1.25
801	0.001	4.5	5.5	1.25	4.5	5.5	1.25
811	0.001	4.5	5.5	1.25	4.5	5.5	1.25
821	0.001	4.5	5.5	1.25	4.5	5.5	1.25
831	0.001	4.5	5.5	1.25	4.5	5.5	1.25
841	0.001	4.5	5.5	1.25	4.5	5.5	1.25
851	0.001	4.5	5.5	1.25	4.5	5.5	1.25
861	0.001	4.5	5.5	1.25	4.5	5.5	1.25
871	0.001	4.5	5.5	1.25	4.5	5.5	1.25
881	0.001	4.5	5.5	1.25	4.5	5.5	1.25
891	0.001	4.5	5.5	1.25	4.5	5.5	1.25
901	0.001	4.5	5.5	1.25	4.5	5.5	1.25
911	0.001	4.5	5.5	1.25	4.5	5.5	1.25
921	0.001	4.5	5.5	1.25	4.5	5.5	1.25
931	0.001	4.5	5.5	1.25	4.5	5.5	1.25
941	0.001	4.5	5.5	1.25	4.5	5.5	1.25
951	0.001	4.5	5.5	1.25	4.5	5.5	1.25
961	0.001	4.5	5.5	1.25	4.5	5.5	1.25
971	0.001	4.5	5.5	1.25	4.5	5.5	1.25
981	0.001	4.5	5.5	1.25	4.5	5.5	1.25
991	0.001	4.5	5.5	1.25	4.5	5.5	1.25
1001	0.001	4.5	5.5	1.25	4.5	5.5	1.25

PROTBF 1

Experiment: PROTBF 1

Report created: 20/01/2010 23:53:23

Spots

#	Anova (p)	Fold	Tags	Notes	Average Normalised Volumes		
					Non mucin media	Mucin Type II media	Mucin Type III media
3698	3.043e-004	6.0			3.148e+006	7.924e+005	5.241e+005
2484	0.002	3.2			1.516e+005	4.788e+004	6.999e+004
2470	0.003	2.2			2.941e+005	2.068e+005	1.350e+005
3006	0.006	2.3			2.202e+005	5.007e+005	2.233e+005
1608	0.011	2.7			1.558e+004	2.609e+004	4.131e+004
2592	0.017	2.1			3.104e+005	1.456e+005	1.927e+005
1138	0.024	3.4			2.021e+005	6.919e+005	4.931e+005
2584	0.025	1.6			4950.707	3996.665	3040.097
3691	0.028	2.8			6.981e+005	9.814e+005	3.530e+005
1295	0.029	6.5			3192.412	2.091e+004	5940.371
3572	0.030	1.8			1.101e+004	1.267e+004	6902.432
4128	0.031	3.0			3.009e+004	9.073e+004	6.460e+004
4319	0.033	6.0			8.717e+004	3.030e+005	5.204e+005
3986	0.054	7.7			1361.911	1.043e+004	2036.838

Tags	
	Power spots
	p value spots
	fold change of 3.0 or more
	p value spots

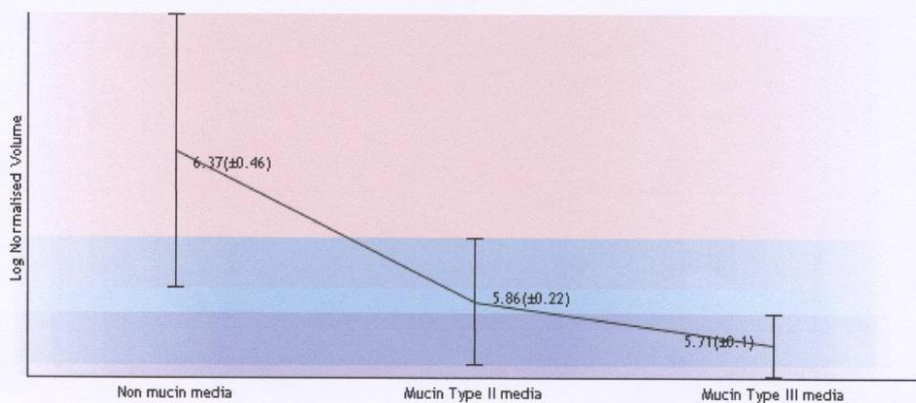
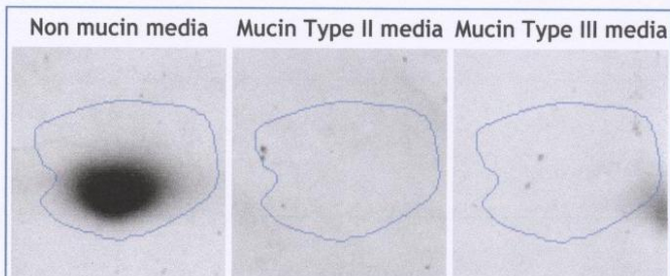
Identifier 3698

Position (2138, 2189)

Notes

- fold change of 3.0 or more
- p value spots
- p value spots

 Power spots

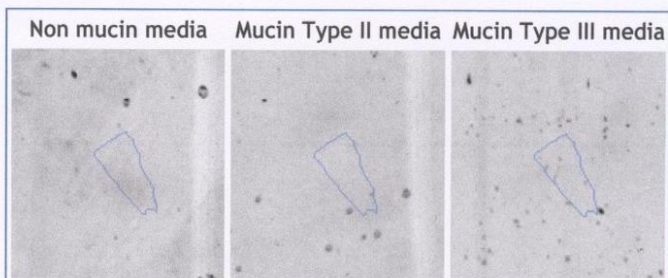


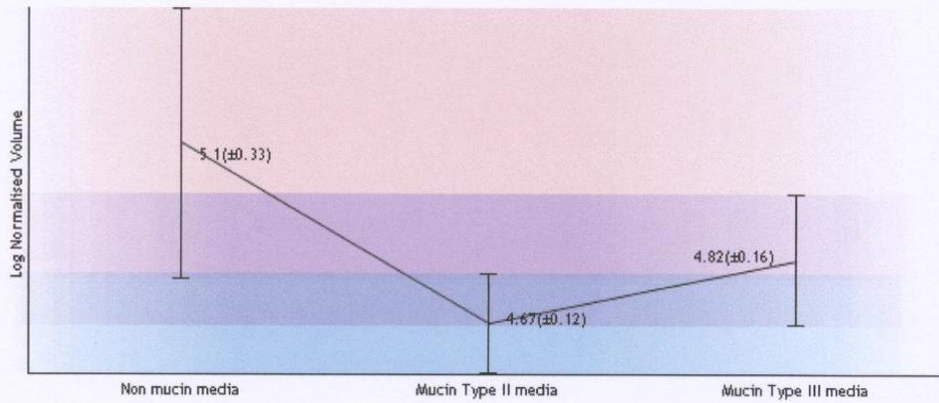
Identifier 2484

Position (1039, 1385)

Notes

-  fold change of 3.0 or more
-  p value spots
-  p value spots
-  Power spots



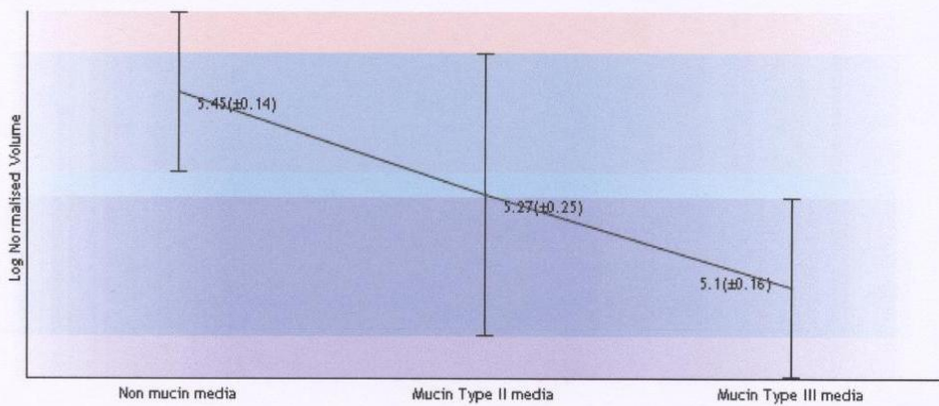
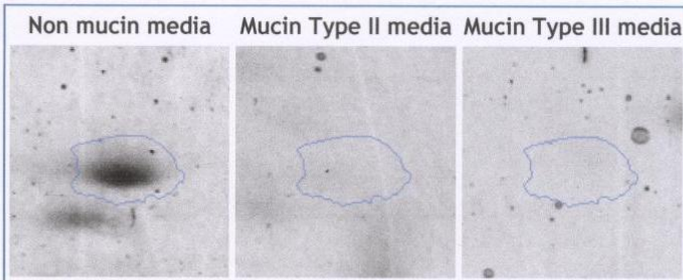


Identifier 2470

Position (1760, 1373)

Notes

- p value spots
- p value spots
- Power spots

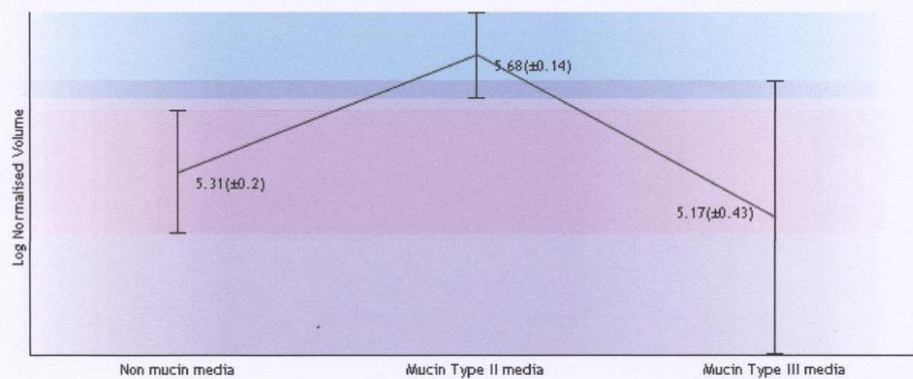
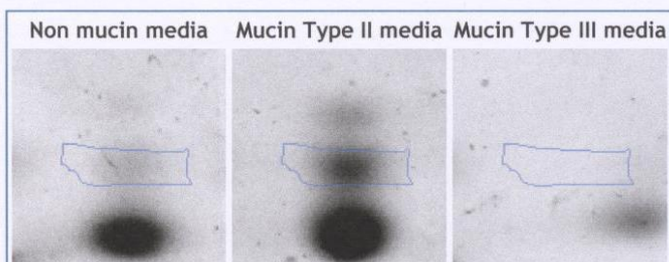


Identifier 3006

Position (1674, 1761)

Notes

- p value spots
- p value spots
- Power spots

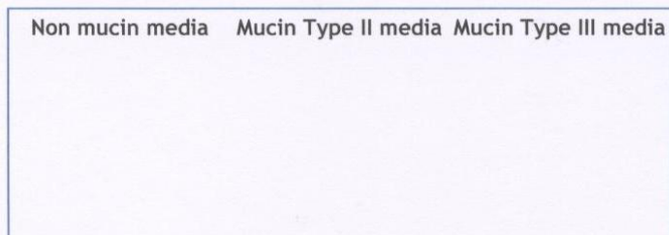


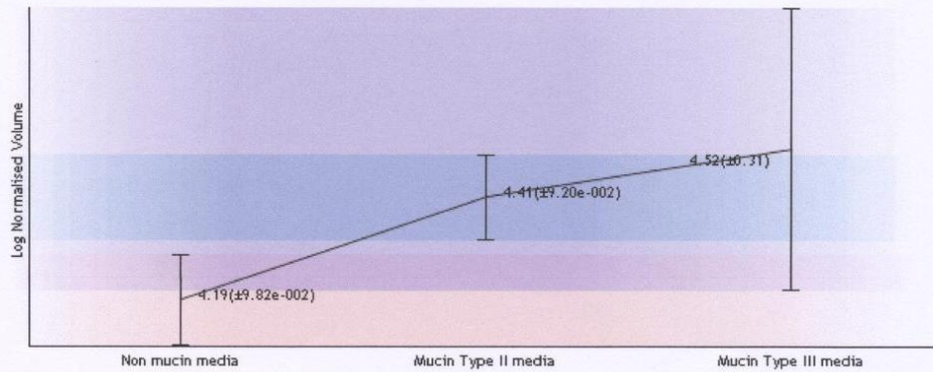
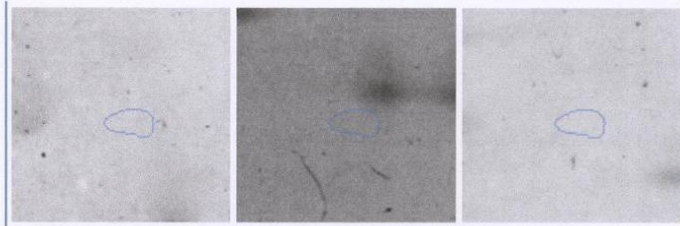
Identifier 1608

Position (2509, 958)

Notes

- p value spots
- p value spots
- Power spots



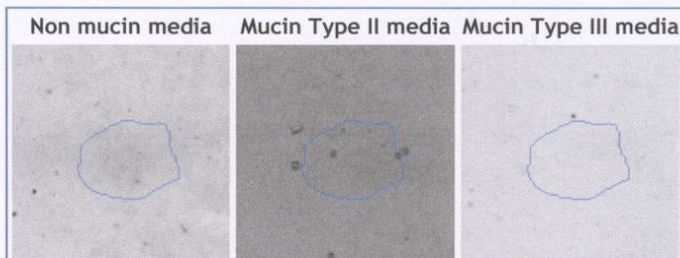


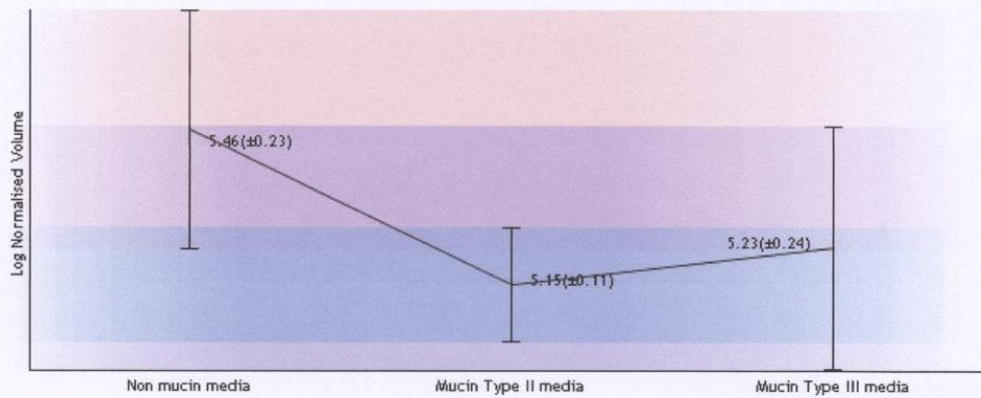
Identifier 2592

Position (2452, 1455)

Notes

-  p value spots
-  p value spots



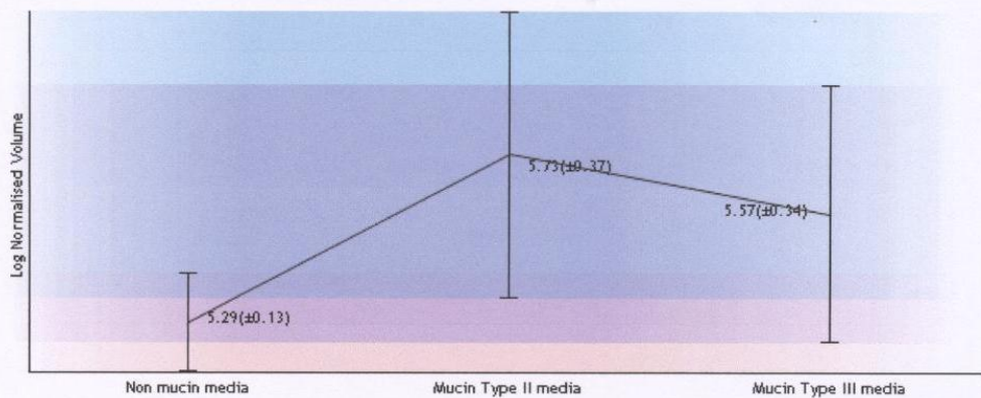
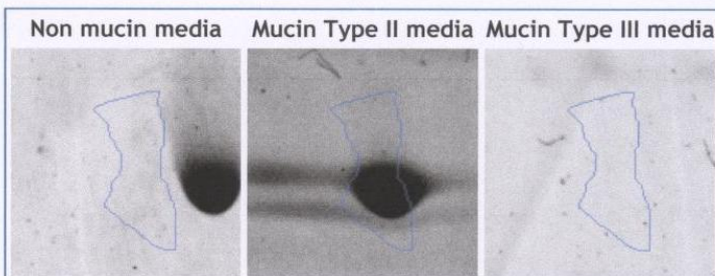


Identifier 1138

Position (1489, 721)

Notes

- fold change of 3.0 or more
- p value spots
- p value spots

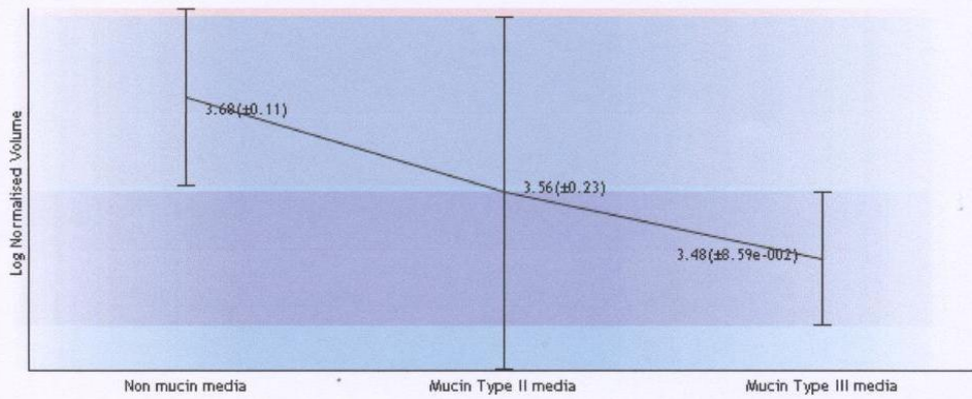
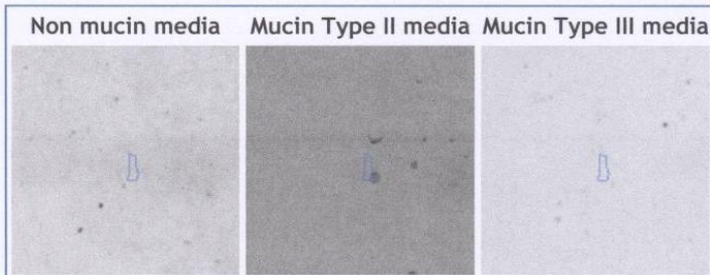


Identifier 2584

Position (2415, 1450)

Notes

- p value spots
- p value spots

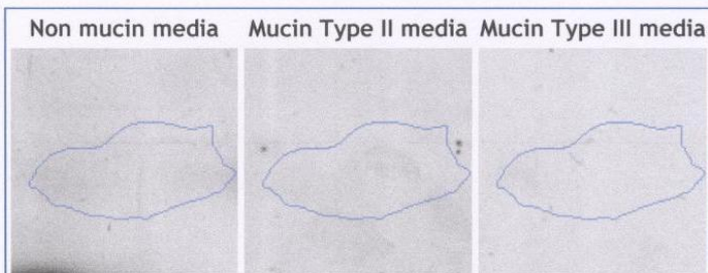


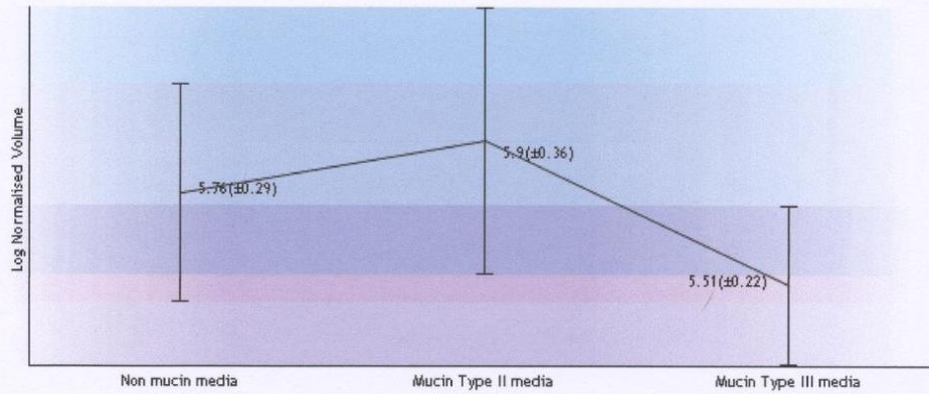
Identifier 3691

Position (2047, 2183)

Notes

- p value spots
- p value spots



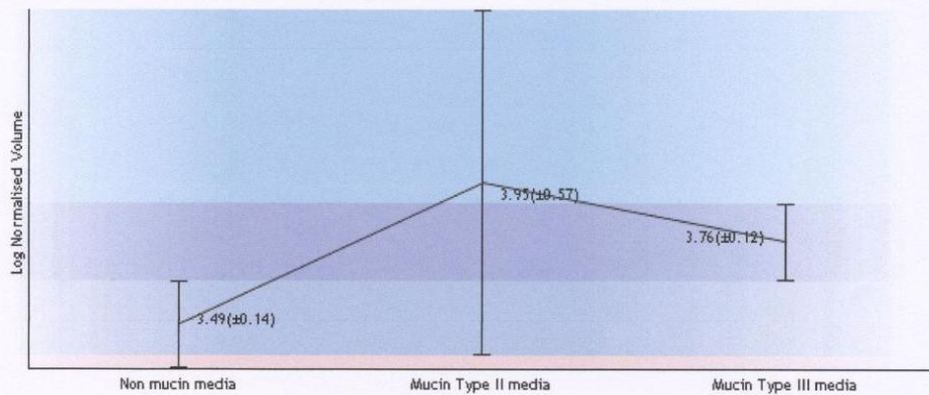
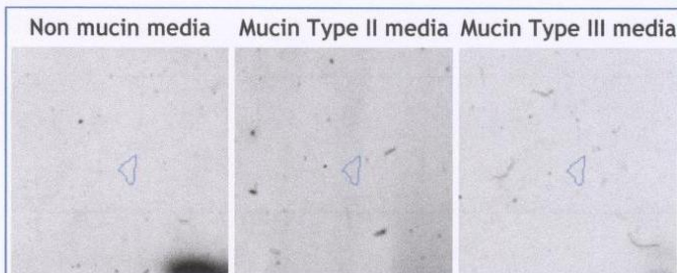


Identifier 1295

Position (639, 795)

Notes

- fold change of 3.0 or more
- p value spots
- p value spots



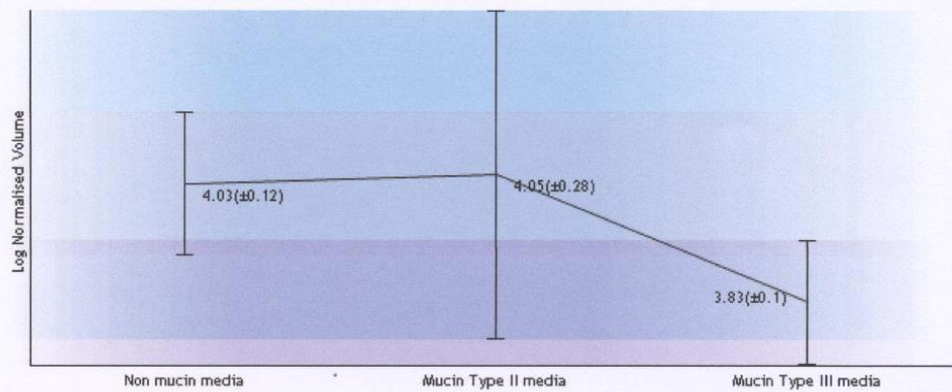
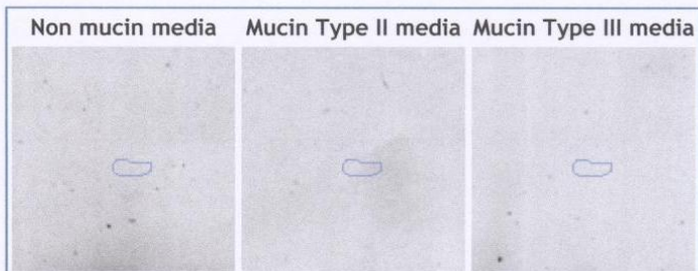
Identifier 3572

Position (1501, 2083)

Notes

p value spots

p value spots



Identifier 4128

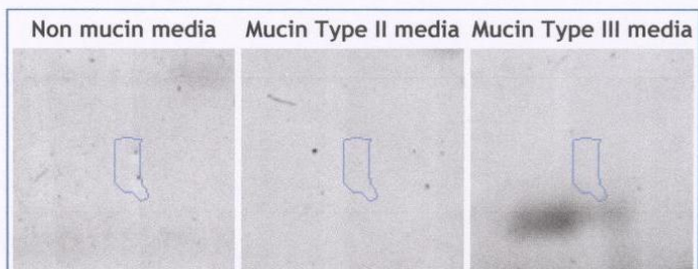
Position (1241, 2305)

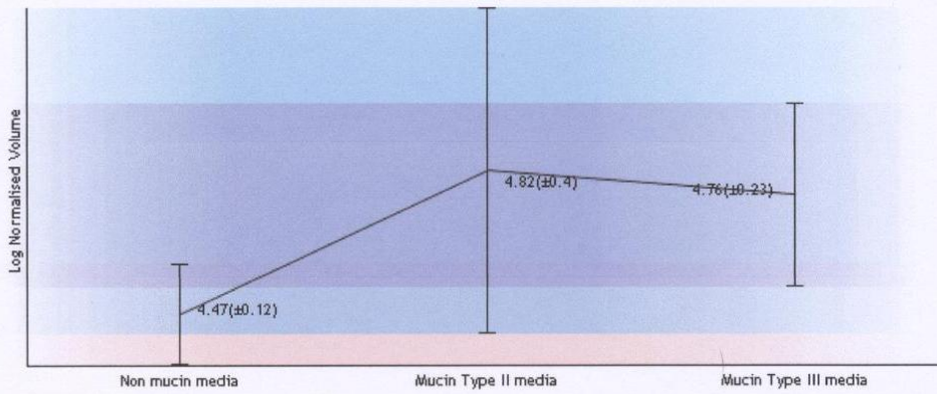
Notes

fold change of 3.0 or more

p value spots

p value spots



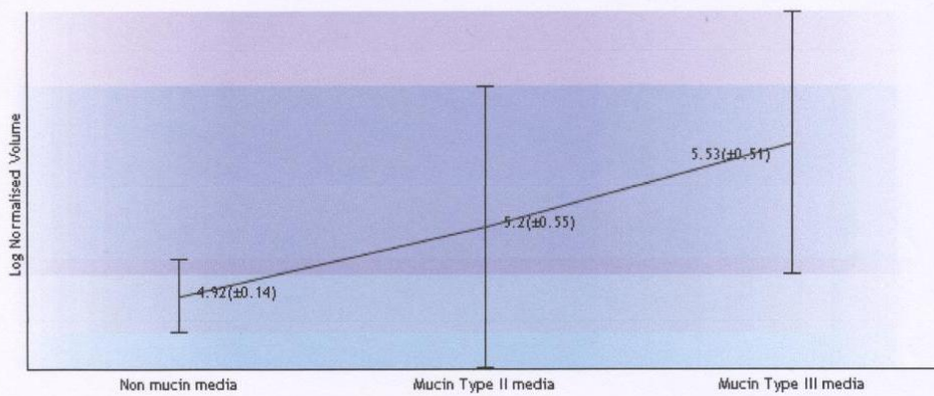
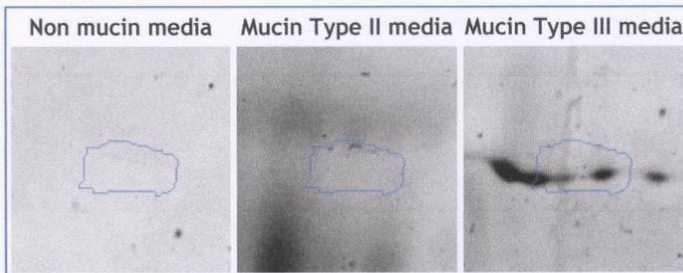


Identifier 4319

Position (565, 2343)

Notes

- fold change of 3.0 or more
- p value spots
- p value spots

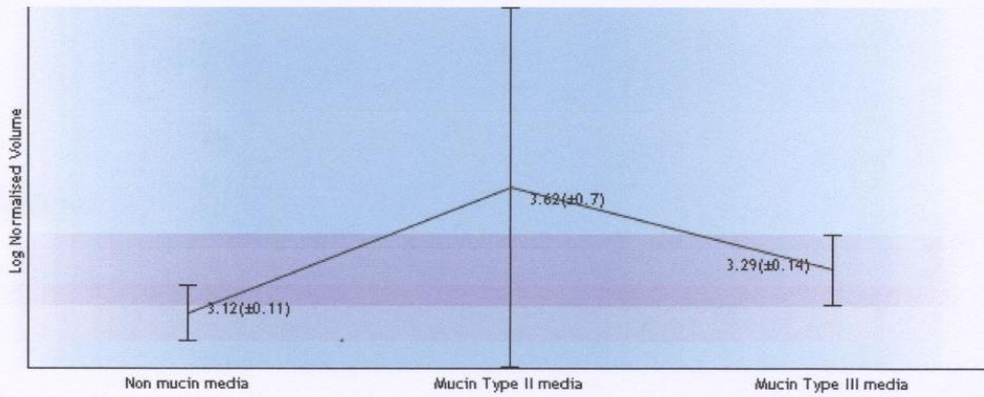
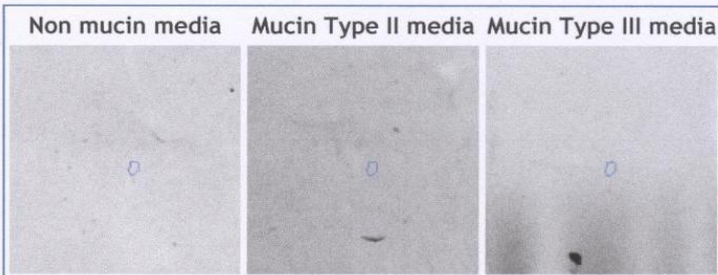


Identifier 3986

Position (2438, 2281)

Notes

- fold change of 3.0 or more
- p value spots
- p value spots

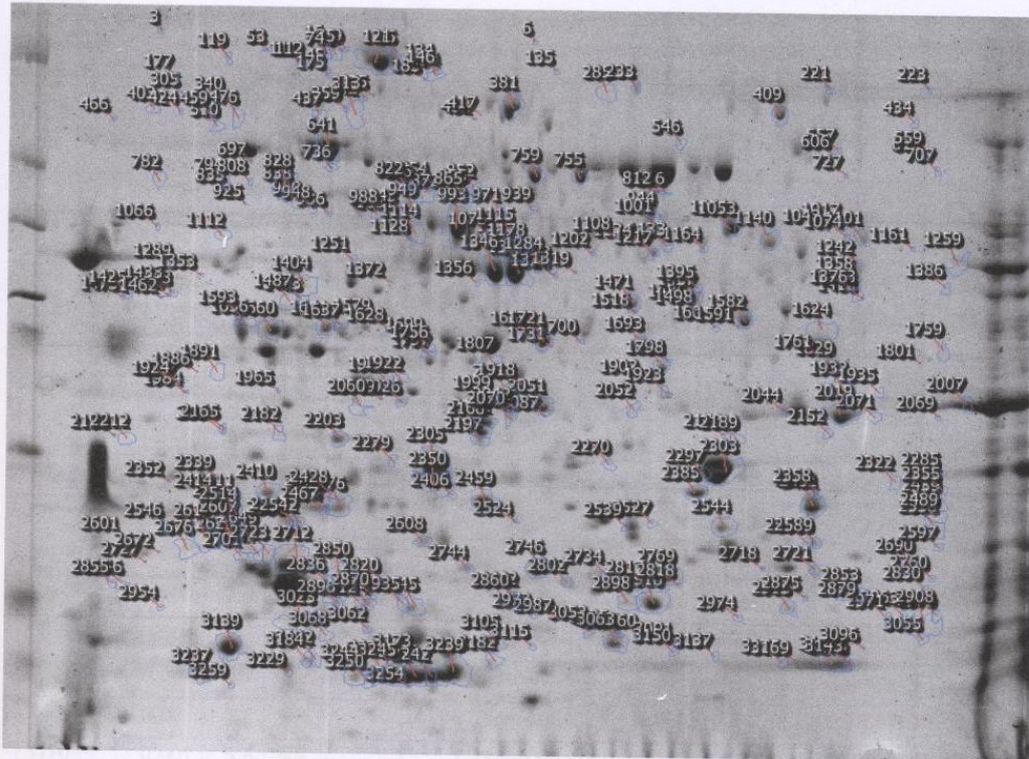


PROTEOMICS 3

Experiment: PROTEOMICS 3

Report created: 16/01/2010 22:22:41

Reference image



171	0.002	3.0	1.58e+005	1.47e+007	1.87e+005
189	0.002	3.3	1.78e+006	5.44e+003	1.89e+005
201	0.002	3.1	1.13e+005	2.44e+003	1.02e+005
224	0.002	2.0	2.19e+004	5.21e+004	1.41e+004
239	0.002	1.4	6.21e+004	1.77e+002	6.21e+004
246	0.002	2.0	4.25e+003	1.23e+004	1.10e+003
193	0.002	3.8	1.52e+005	9.34e+004	8.27e+004
124	0.001	3.0	4.19e+005	1.29e+005	1.37e+005
128	0.002	2.5	1.01e+005	8.17e+004	1.19e+005
205	0.002	2.3	1.25e+004	7.91e+004	1.12e+004
204	0.002	3.3	1.02e+004	3.51e+004	1.08e+004
241	0.002	3.1	8.75e+004	2.49e+005	1.41e+004

PROTEOMICS 3

Experiment: PROTEOMICS 3

Report created: 20/01/2010 23:57:30

Spots

#	Anova (p)	Fold	Tags	Notes	Average Normalised Volumes		
					Non mucin media	Mucin type II media	Mucin type III media
95	1.039e-004	3.0			3.023e+005	1.031e+005	1.012e+005
2914	1.114e-004	2.3			1.123e+005	4.946e+004	5.120e+004
2026	1.362e-004	1.9			6.443e+004	3.867e+004	3.409e+004
2132	3.050e-004	4.8			4.420e+005	1.284e+005	9.248e+004
755	3.761e-004	6.7			5.050e+005	7.543e+004	8.154e+004
3176	5.017e-004	6.1			1.166e+006	2.160e+005	1.925e+005
2484	6.287e-004	2.0			2.108e+004	1.329e+004	1.052e+004
2717	6.672e-004	5.1			6.585e+004	3.377e+005	8.572e+004
2945	6.762e-004	2.3			2.858e+005	1.318e+005	1.254e+005
2365	8.818e-004	2.9			2.652e+004	9878.909	9019.675
2608	0.001	2.7			1.663e+005	6.131e+004	7.626e+004
2746	0.001	3.8			9.243e+005	2.420e+005	2.470e+005
1171	0.002	3.0			3.900e+005	1.467e+005	1.303e+005
3139	0.002	3.3			1.783e+006	5.443e+005	5.895e+005
854	0.002	3.1			9.330e+004	2.854e+005	2.040e+005
1234	0.002	3.2			4.140e+005	1.294e+005	1.517e+005
3234	0.002	2.5			2.018e+005	8.117e+004	1.190e+005
2352	0.003	1.9			3754.338	1944.626	3165.534
2410	0.003	3.7			5.802e+005	5.349e+005	1.571e+005
1579	0.004	1.9			1433.976	2319.226	2752.321
1342	0.004	4.9			2.388e+006	4.888e+005	7.282e+005
3060	0.009	1.7			2.947e+004	1.909e+004	3.283e+004
813	0.010	3.0			4591.714	1.368e+004	9587.014
1922	0.022	3.5			4.540e+004	1.605e+005	9.859e+004
872	0.024	4.1			8057.272	2.099e+004	3.320e+004
3244	0.026	1.7			2.928e+005	2.396e+005	1.745e+005
2083	0.028	2.0			8.762e+004	4.398e+004	7.334e+004

file://C:\Documents and Settings\larcje1\My Documents\d.htm

21/01/2010

1902	0.028	1.7		1753.296	3052.482	2859.892
285	0.037	3.6		9.781e+004	3.564e+005	2.645e+005
1807	0.038	3.2		2.874e+005	2.417e+005	9.005e+004
2712	0.043	2.4		3.925e+006	3.254e+006	1.648e+006
835	0.049	1.5		8994.086	8357.012	6113.549

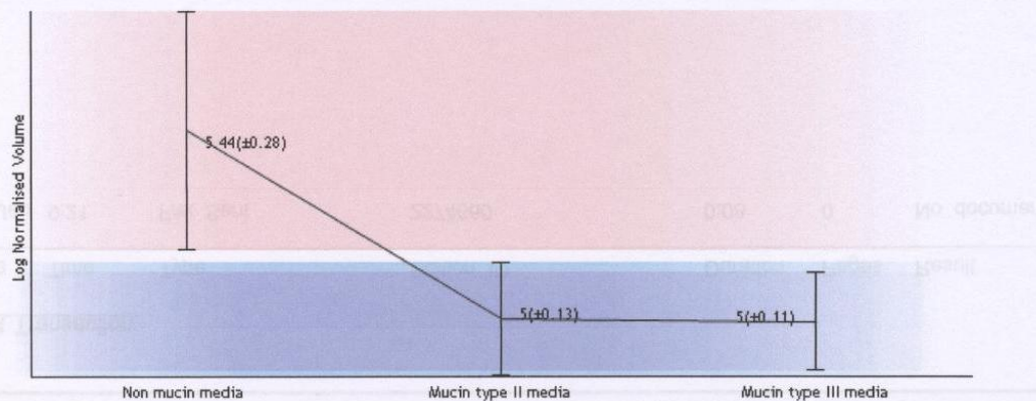
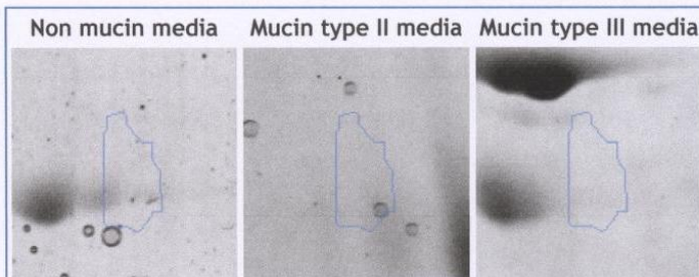
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	p value spots
	fold change of 3.0 or more
	spots of interest

Identifier 95

Position (1149, 163)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest

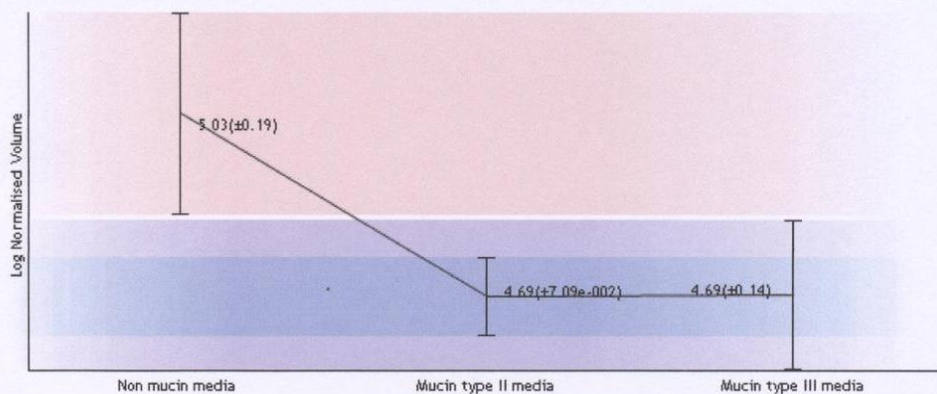
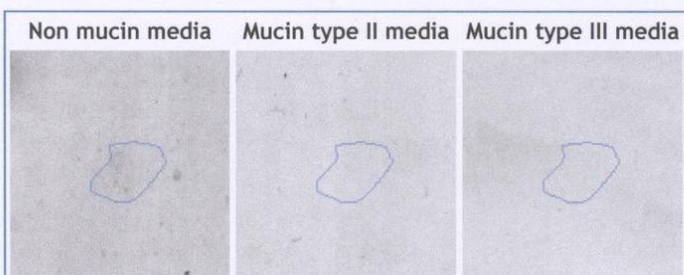


Identifier 2914

Position (1101, 1843)

Notes

- p value spots
- Power spots
- spots of interest

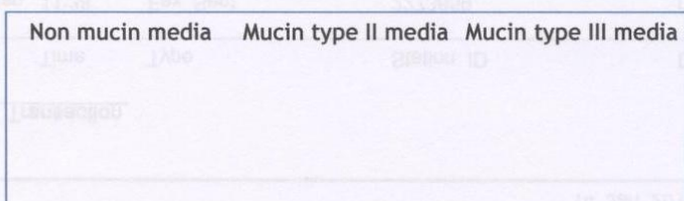


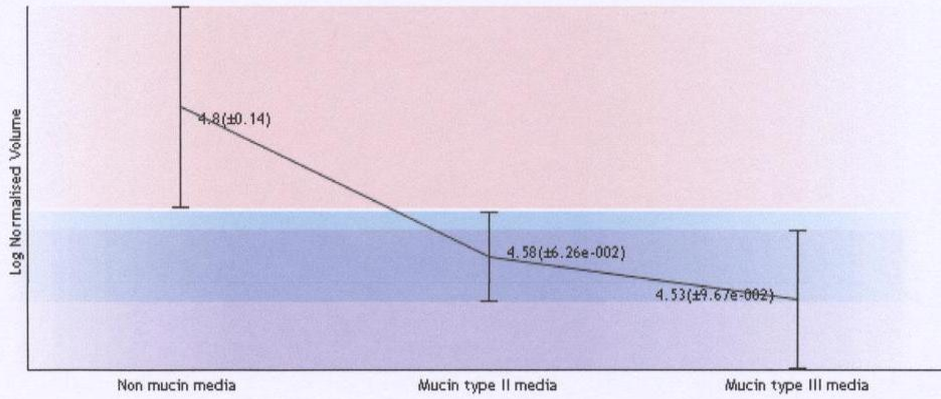
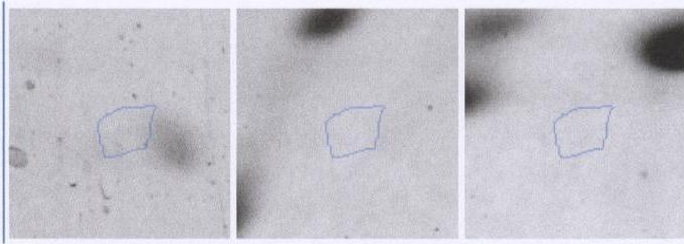
Identifier 2026

Position (1193, 1219)

Notes

- p value spots
- Power spots
- spots of interest



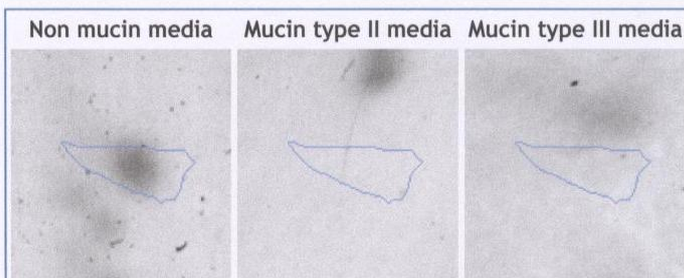


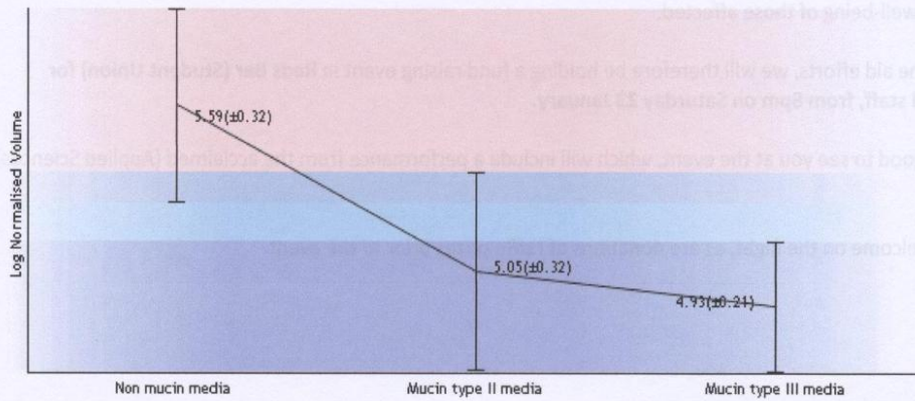
Identifier 2132

Position (1447, 1279)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest



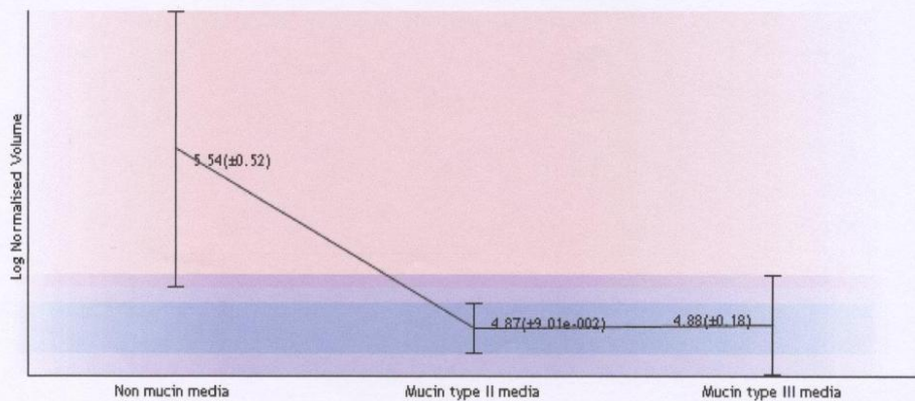
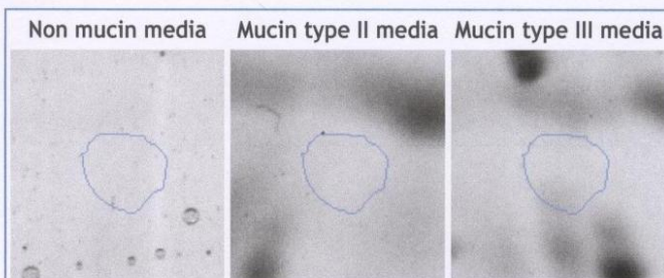


Identifier 755

Position (1721, 521)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest

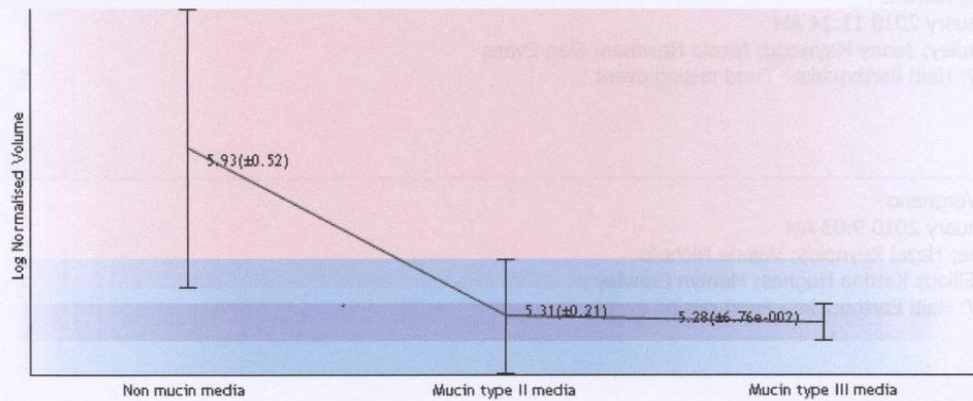
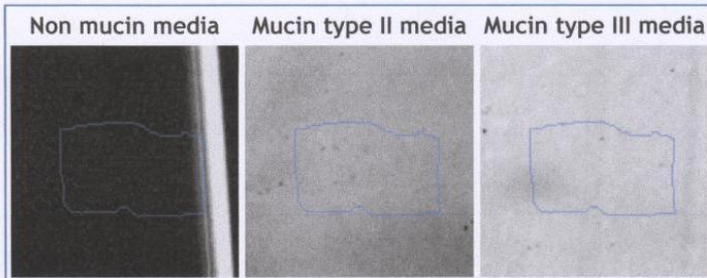


Identifier 3176

Position (2461, 2018)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest

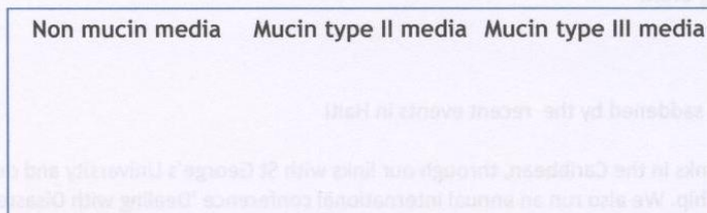


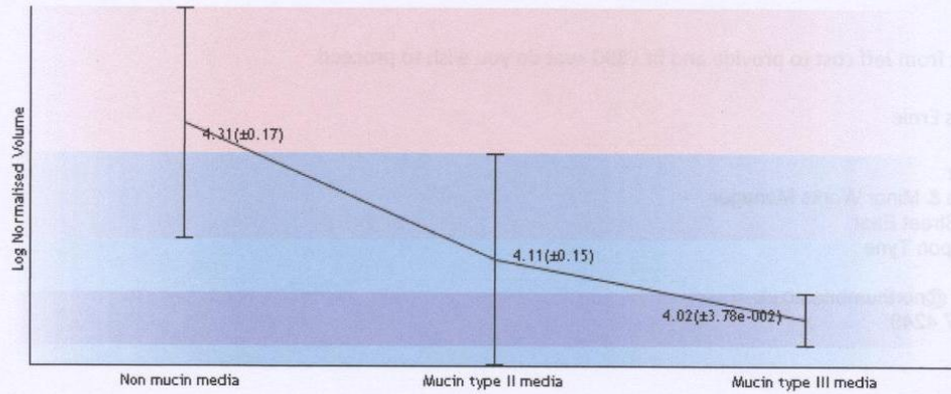
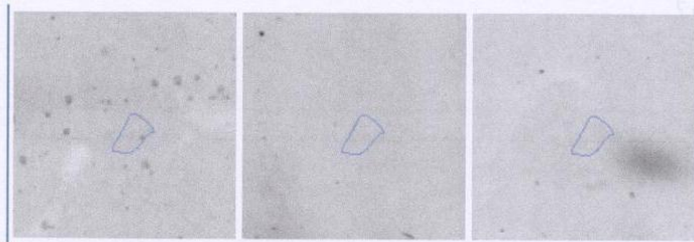
Identifier 2484

Position (696, 1555)

Notes

- p value spots
- Power spots
- spots of interest



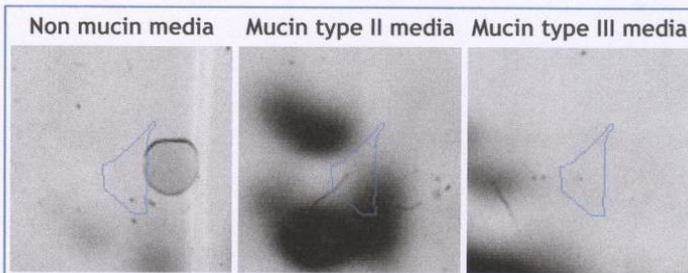


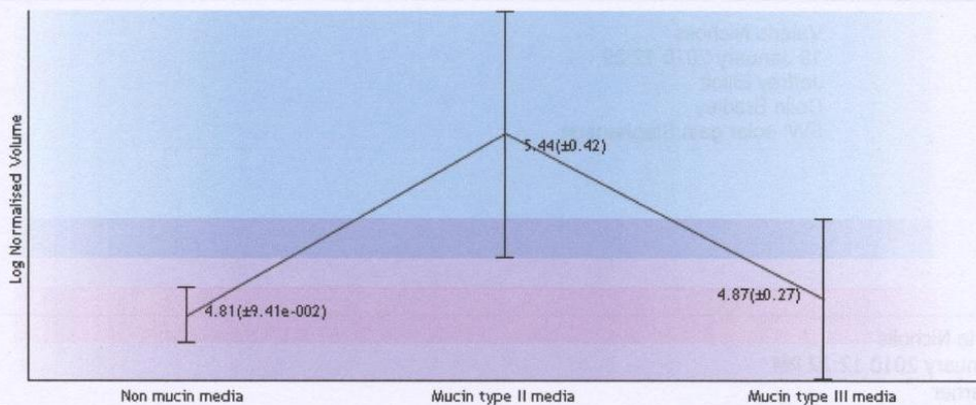
Identifier 2717

Position (778, 1719)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest



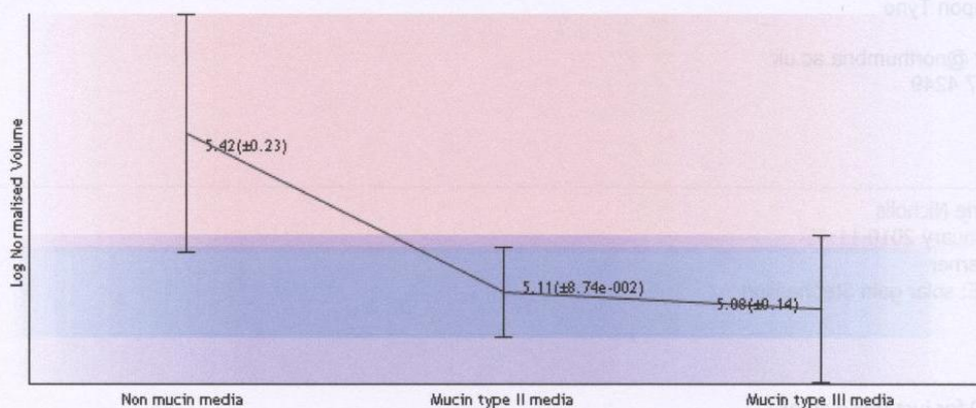
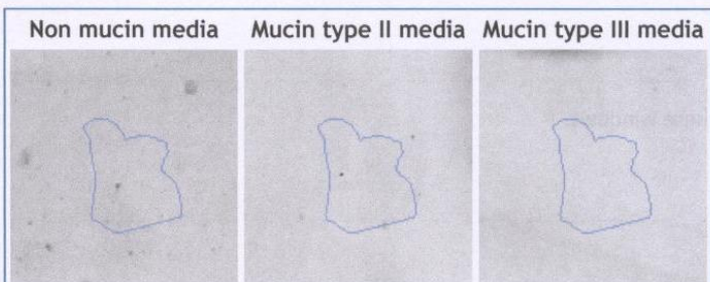


Identifier 2945

Position (1232, 1861)

Notes

- p value spots
- Power spots
- spots of interest



Identifier 2365

Jeffrey Elliott

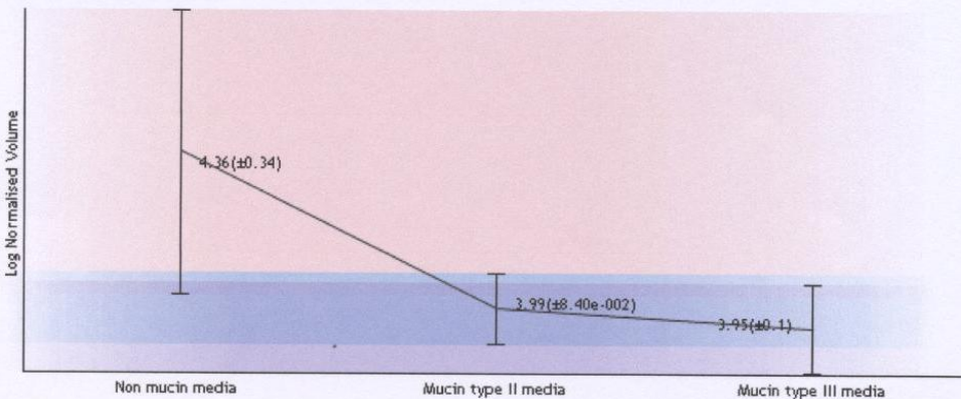
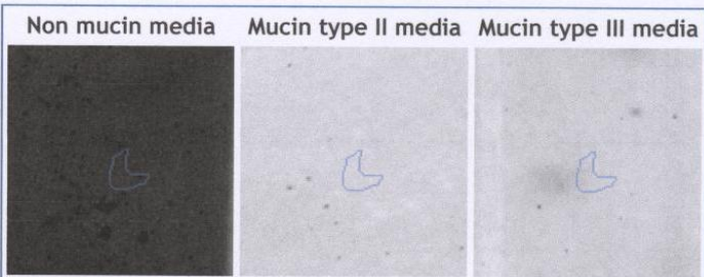
Emile Garner (emile.garner@northumbria.ac.uk)
19 January 2010 18:14

From:
Sent:

Position (2450, 1480)

Notes

- p value spots
- Power spots
- spots of interest

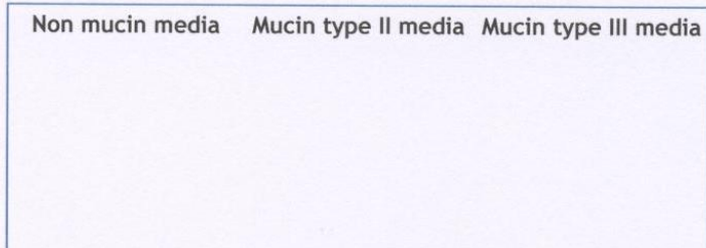


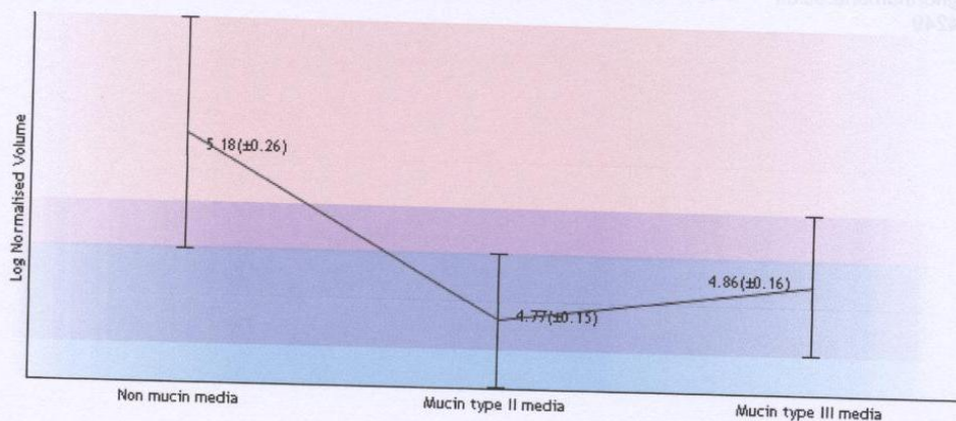
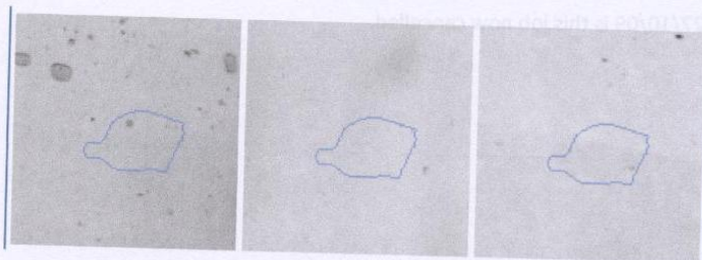
Identifier 2608

Position (1264, 1650)

Notes

- p value spots
- Power spots
- spots of interest



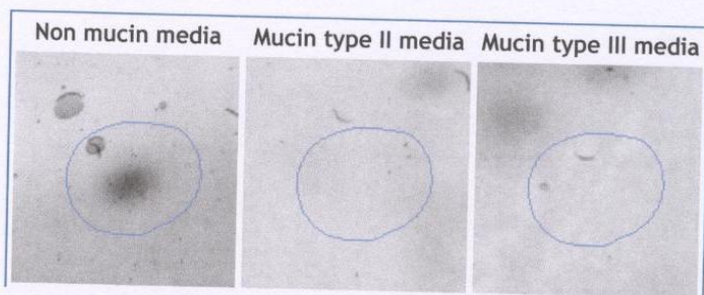


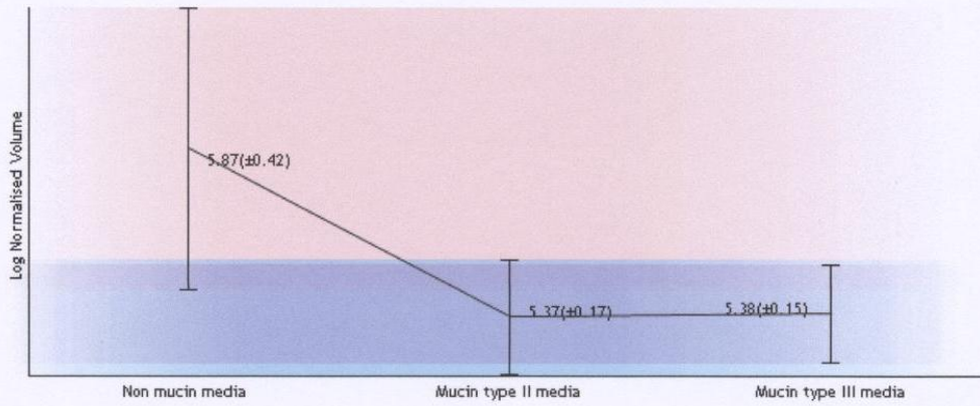
Identifier 2746

Position (1613, 1735)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest



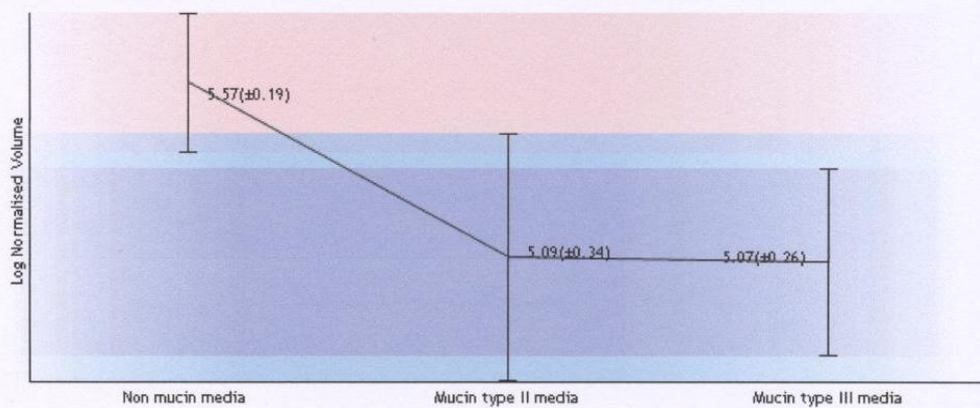
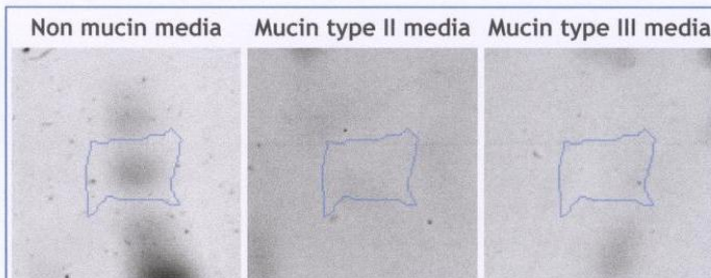


Identifier 1171

Position (1520, 729)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest

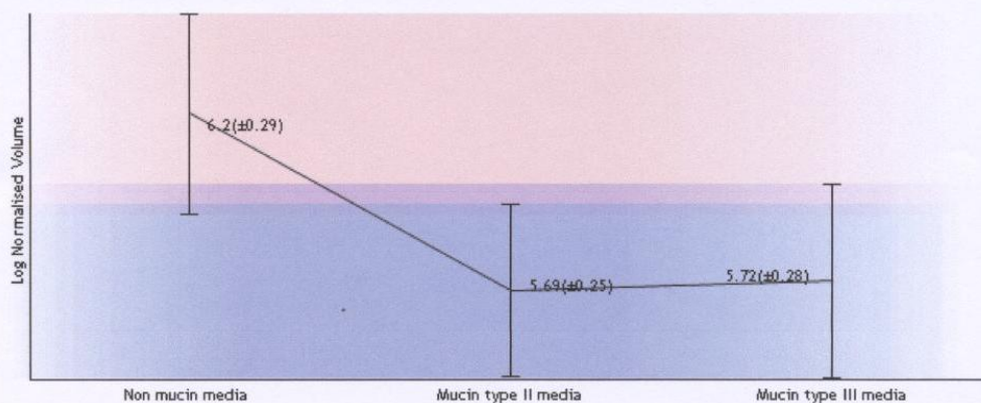
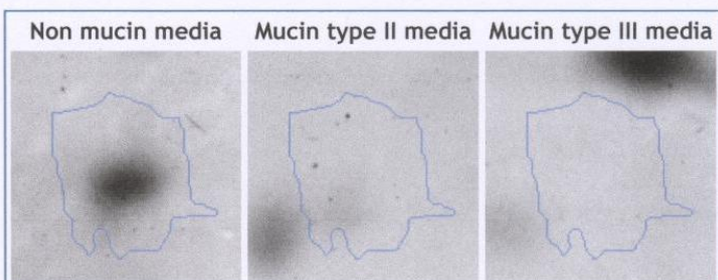


Identifier 3139

Position (685, 1993)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest

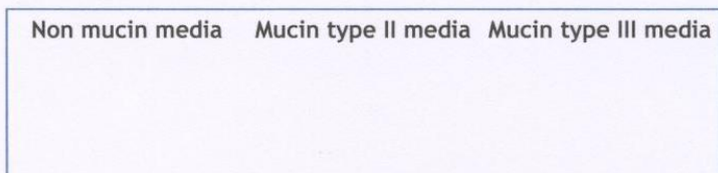


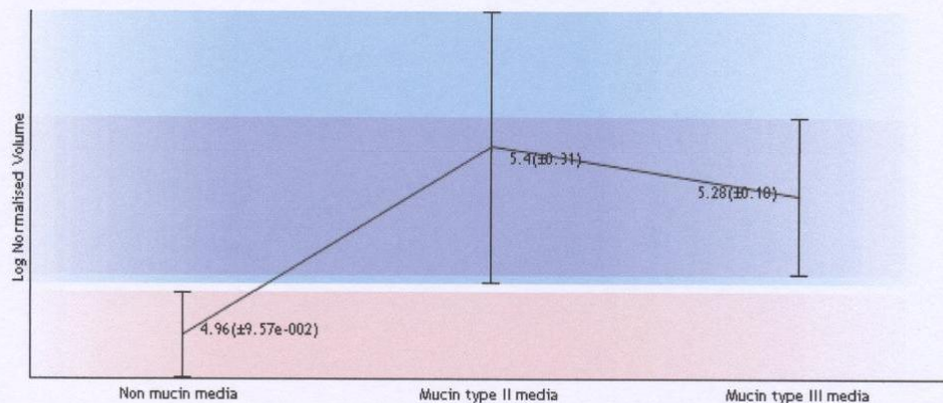
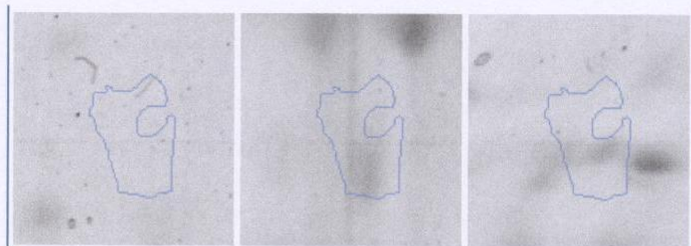
Identifier 854

Position (1255, 574)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest



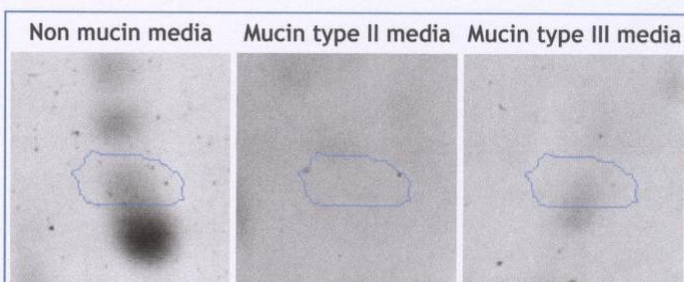


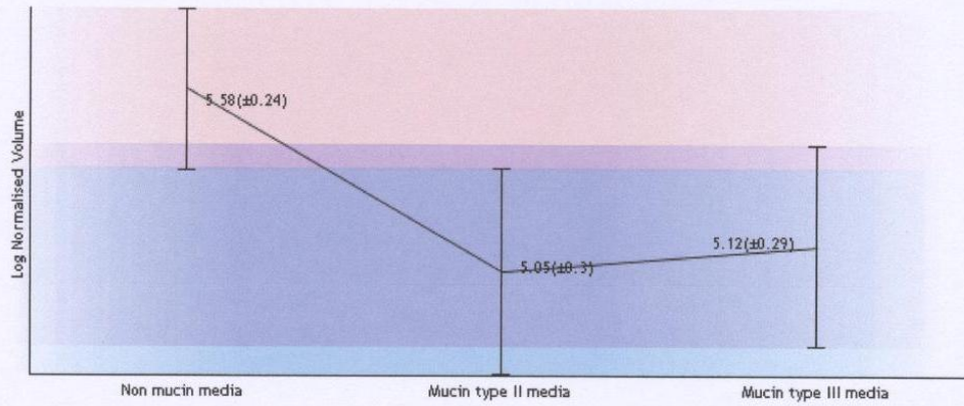
Identifier 1234

Position (1523, 765)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest



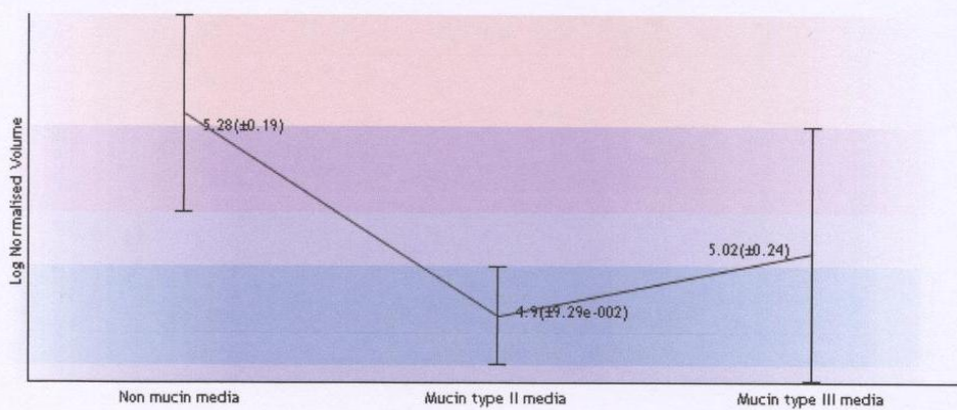
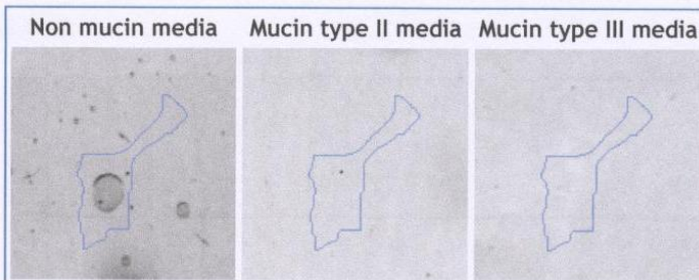


Identifier 3234

Position (1143, 2065)

Notes

- p value spots
- Power spots
- spots of interest

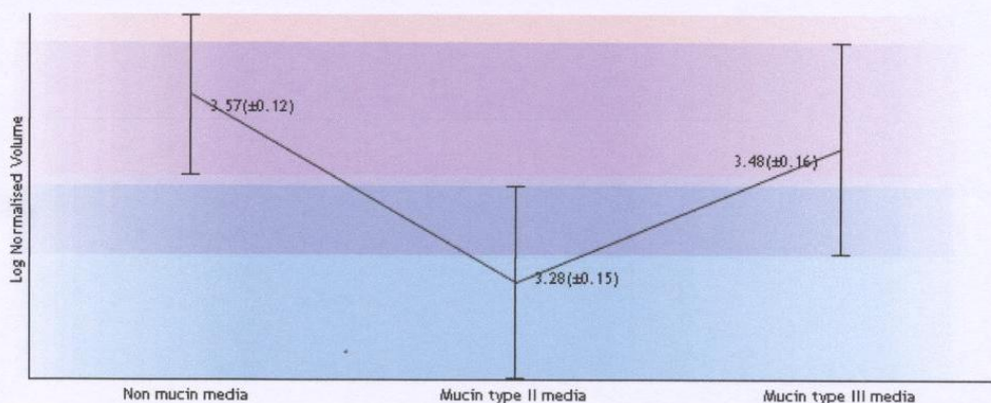
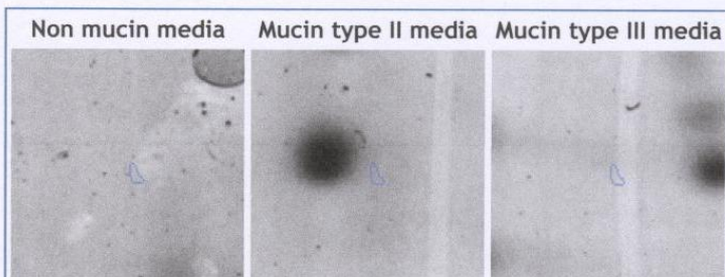


Identifier 2352

Position (496, 1470)

Notes

- p value spots
- Power spots
- spots of interest

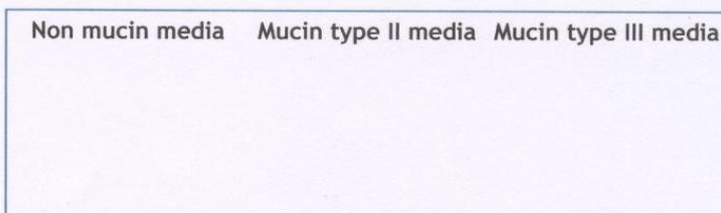


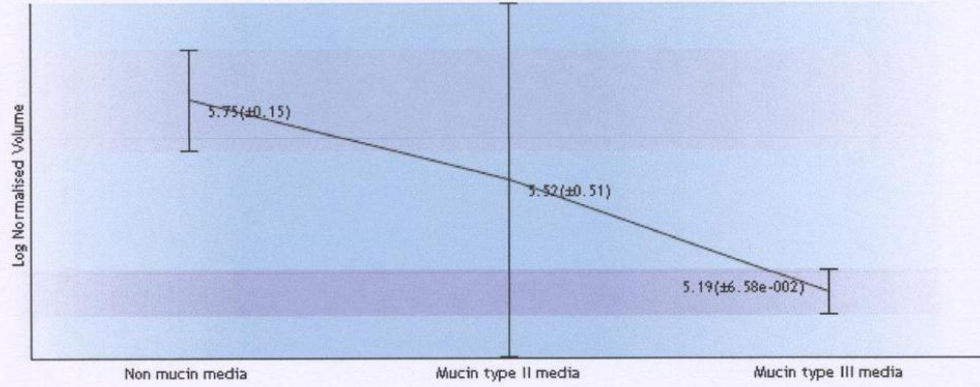
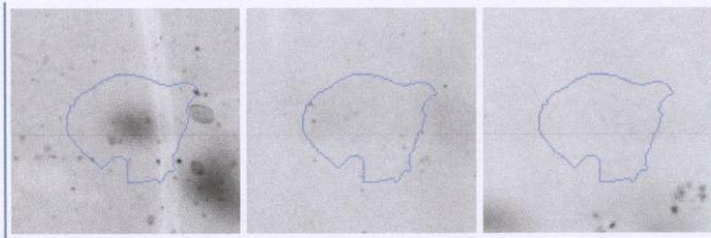
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Position (795, 1507)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest



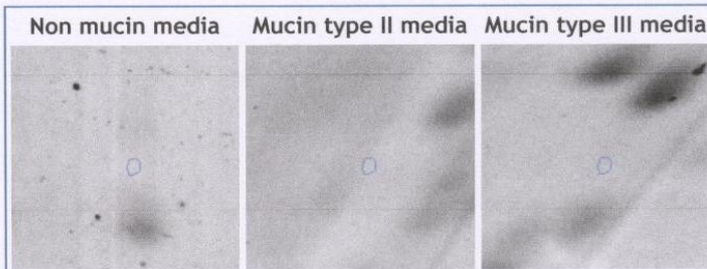


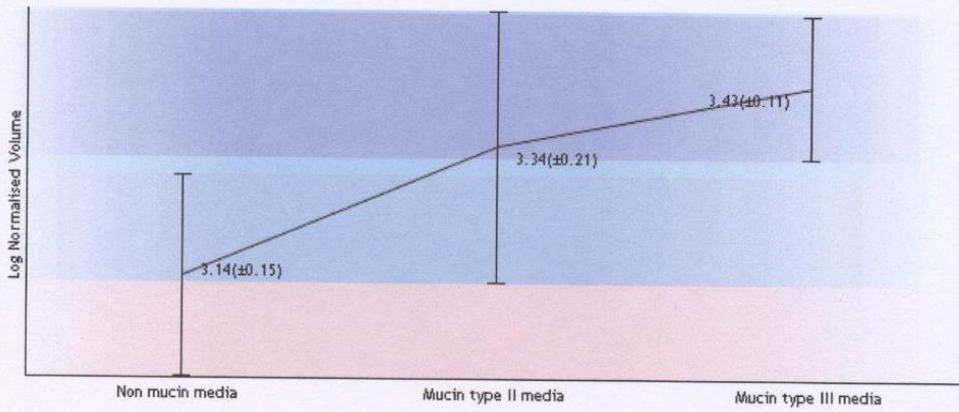
Identifier 1579

Position (1121, 960)

Notes

- p value spots
- Power spots
- spots of interest



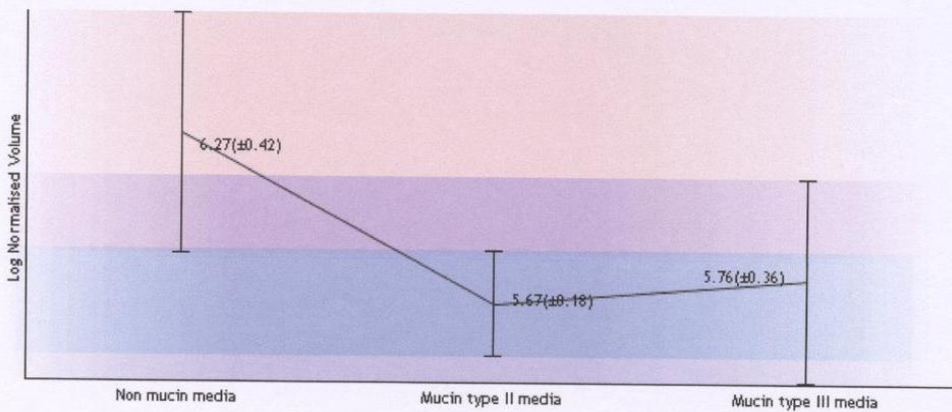
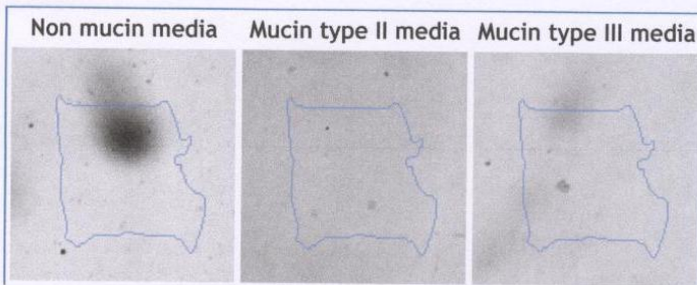


Identifier 1342

Position (1533, 827)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest

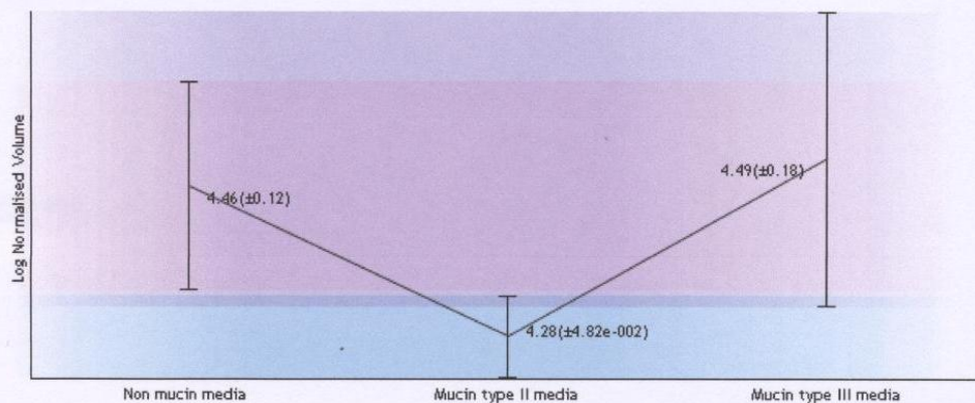
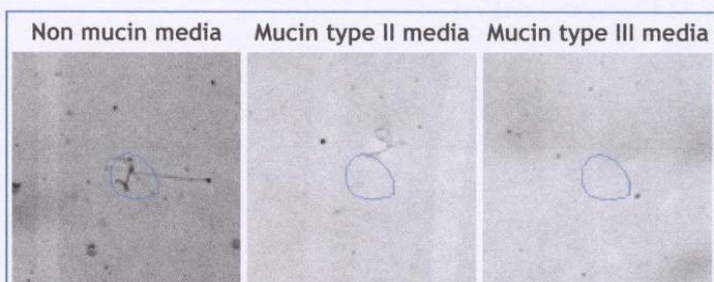


Identifier 3060

Position (1918, 1933)

Notes

- p value spots
- Power spots
- spots of interest

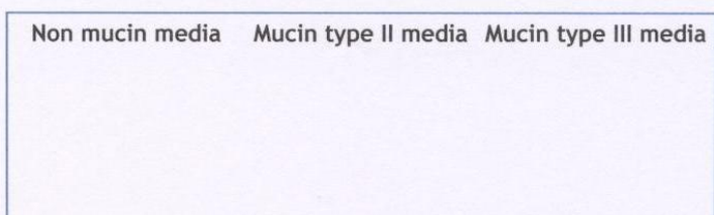


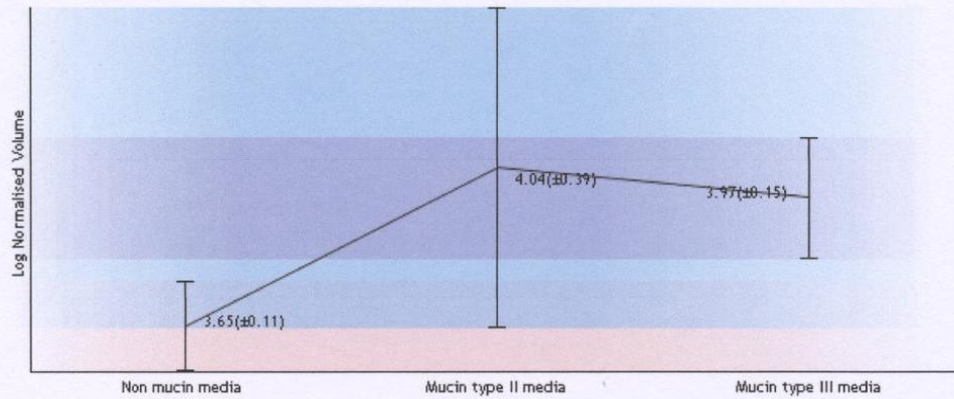
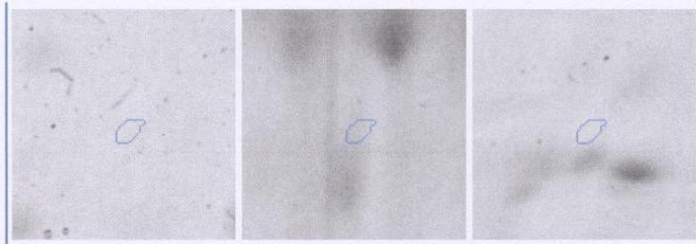
Identifier 813

Position (1270, 552)

Notes

- fold change of 3.0 or more
- p value spots
- Power spots
- spots of interest



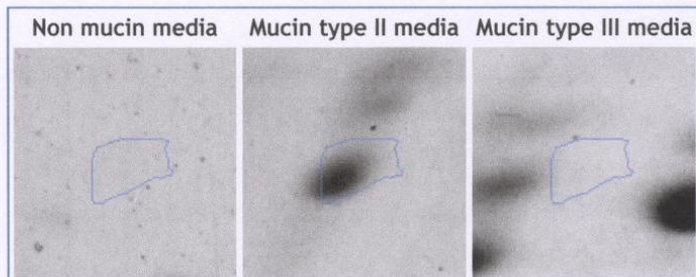


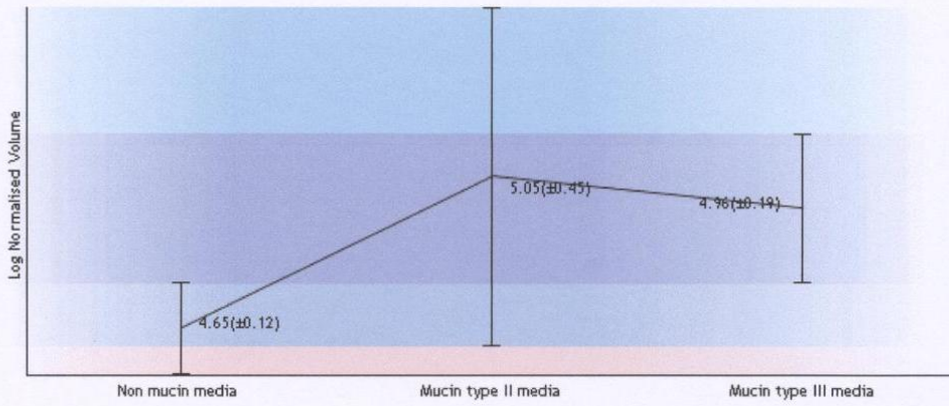
Identifier 1922

Position (1171, 1158)

Notes

- fold change of 3.0 or more
- p value spots
- spots of interest



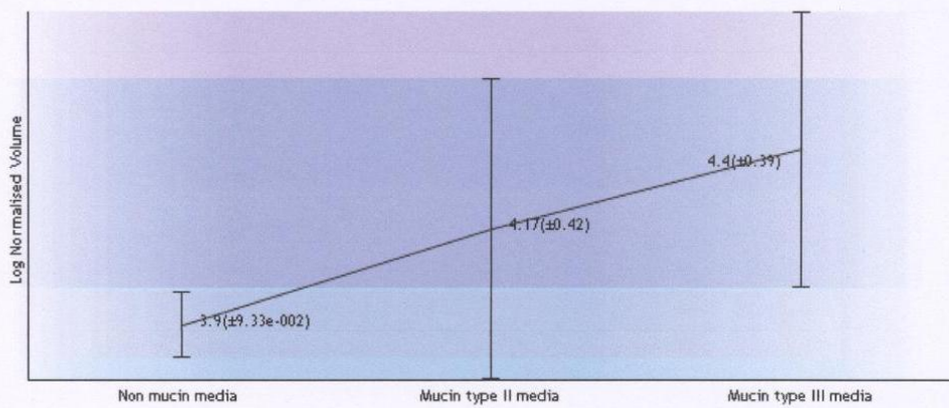
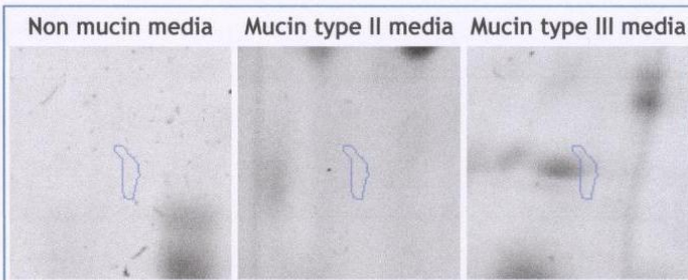


Identifier 872

Position (1314, 579)

Notes

- fold change of 3.0 or more
- p value spots
- spots of interest

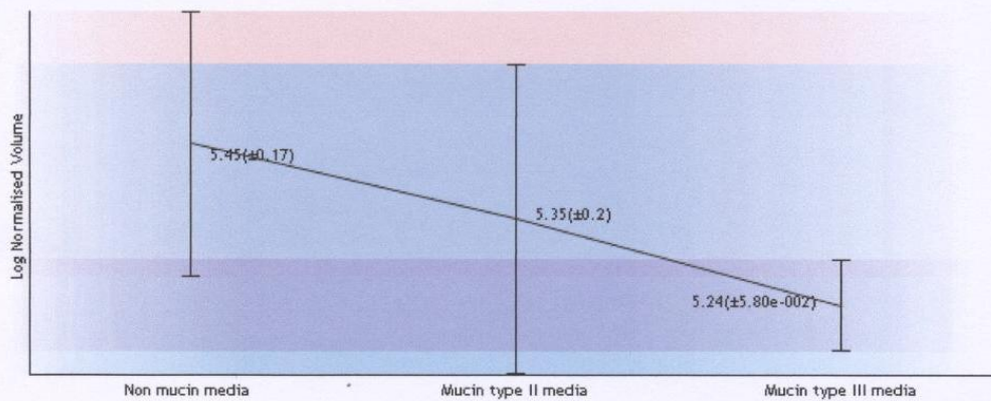
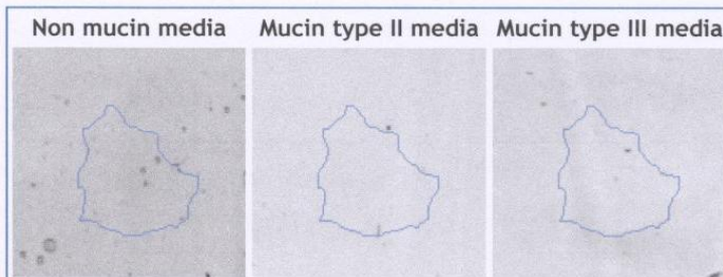


Identifier 3244

Position (1067, 2075)

Notes

- p value spots
- spots of interest

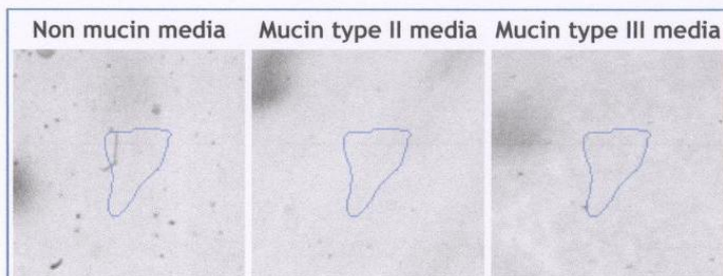


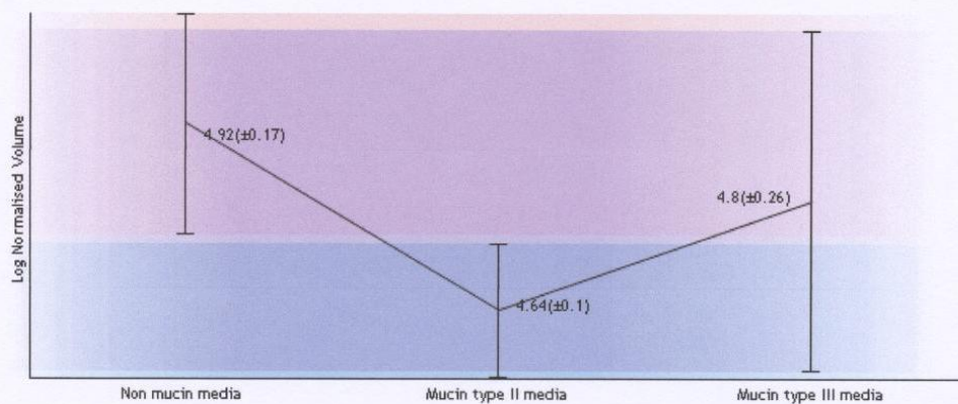
Identifier 2083

Position (1523, 1251)

Notes

- p value spots
- spots of interest



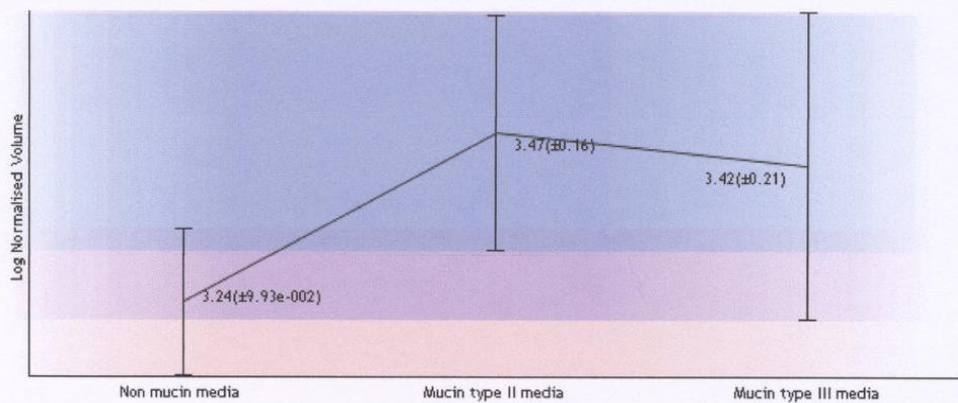
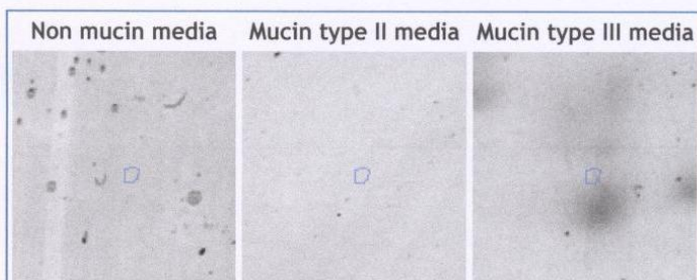


Identifier 1902

Position (1923, 1144)

Notes

- p value spots
- spots of interest

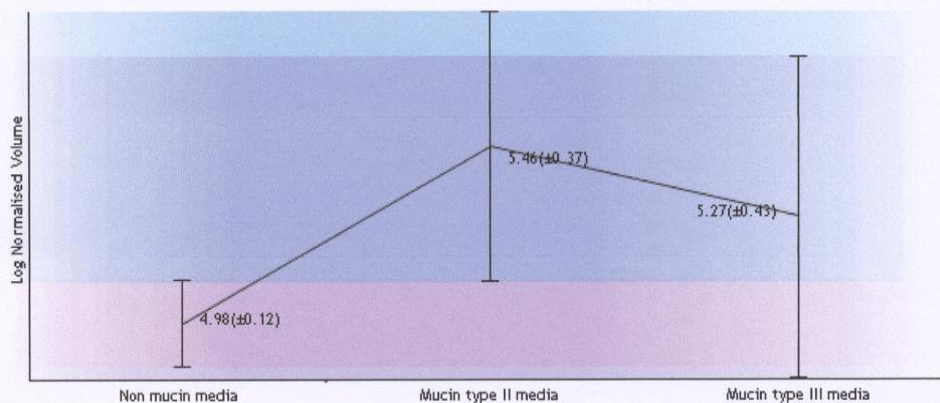
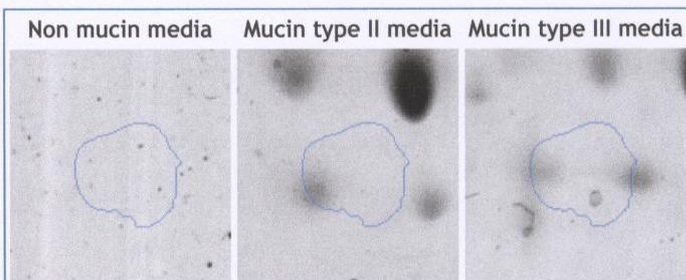


Identifier 285

Position (1797, 259)

Notes

- fold change of 3.0 or more
- p value spots
- spots of interest

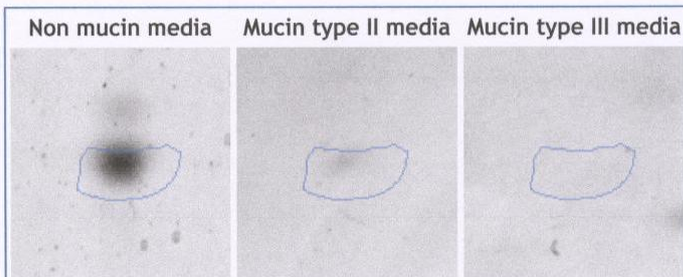


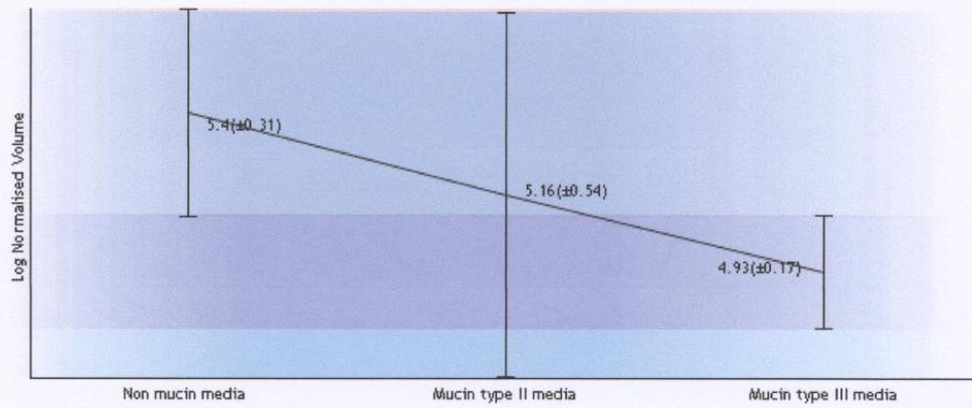
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Position (1461, 1079)

Notes

- fold change of 3.0 or more
- p value spots
- spots of interest



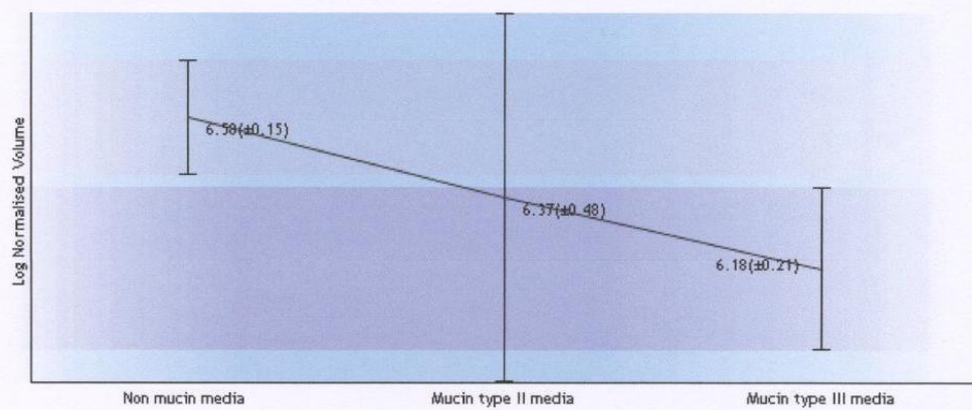
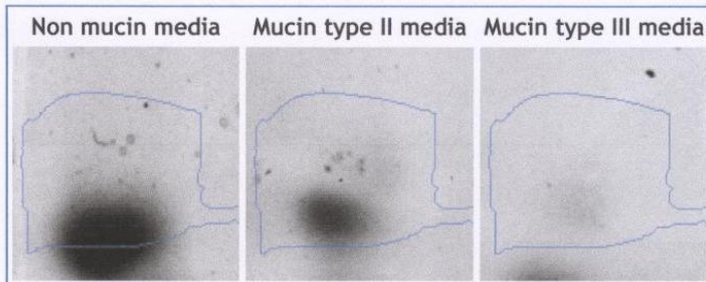


Identifier 2712

Position (923, 1717)

Notes

- p value spots
- spots of interest

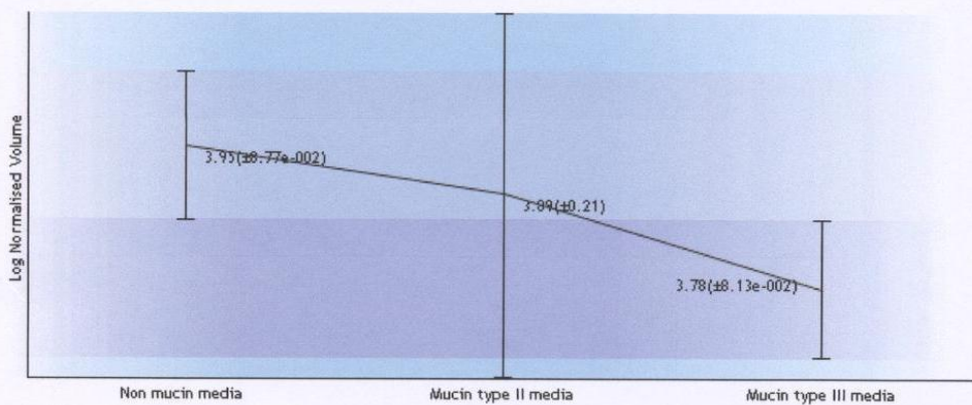
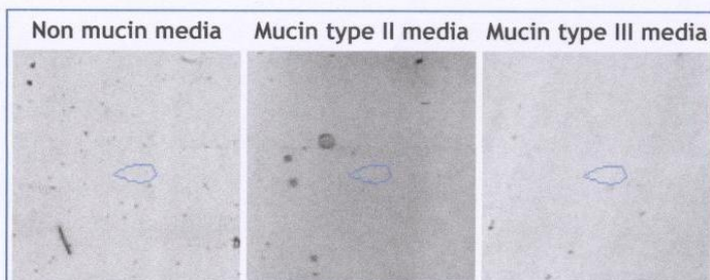


Identifier 835

Position (666, 565)

Notes

- p value spots
- spots of interest



Appendix G

G1 Details on MS analysis

Bruker Daltonics Mass spectrometer that worked based on the principle of electrospray ionisation was used for fragmenting, ionising and identifying peptides from the protein spots of interest. The liquid sample which passed through an emitter was ionised to form a liquid jet under high voltage electricity. The liquid then radially dispersed to form small and highly charged liquid droplets due to Coulomb repulsion. Following nebulisation, the ions were dried and desorbed under the influence of the same high potential electrostatic field.

The generation of ions

The three important steps in the generation of ions were nebulisation, desolvation and ion evaporation. The sample containing the trypsinised proteins was first separated using the LC column in the Dionex Ultimate 3000 system and the droplets were fed into the mass spectrometer through the needle at a flow rate of 3 $\mu\text{L}/\text{min}$. The droplets enter the spray chamber from the tip of the needle where the force of Coulomb repulsion exceeds the surface tension causing the droplets to explode into smaller droplets that undergo evaporation. During the process of evaporation, the droplets become highly charged and get ionised. When the surface tension on the surface of the droplets is exceeded by the ionic field, ions are emitted into the gas phase. Peptides containing amino acids either accept or donate protons based on their charge. A similar effect can be observed in the buffers and solvents that also ionise in the presence of an electric field. pH of the buffers and solvents can have a great effect on the peptide signals detected. The ionisation process can also be affected by the conductivity of solutions and their flow rate.

Parts of the Ion trap mass spectrometer

The spray shield forms a protective covering that prevents the deposition of contaminants, buffer salts and other impurities. The transfer capillary plays an important role in the declustering and desolvation of ions. It acts as the first barrier in the vacuum system. The skimmer forms the main part of the next vacuum stage where it helps to remove dried gas and other solvent molecules. The ions that pass

through the transfer capillary and the skimmer enter the dual octopoles that focus, slow down or accelerate them for efficient ion trapping.

Ion trap system

The first step in the trapping of ions involves the generation of a potential well which acts as the site for accumulation of ions. The end cap generates a repelling potential that enhances the trapping efficiency. The ions collide with helium thereby slowing them down and oscillating around the centre of the trap. The ions focussed in the centre of the trap are ejected by applying an increased RF amplitude and auxiliary RF which cause each m/z to come into resonance. The ions accumulated in the 3D multipolar field were ejected by increasing the RF amplitudes and the response was measured at the detector.

The conversion dynode emitted electrons when hit by ions. These electrons were accelerated onto an electron multiplier that converted the electrons into a resulting current that was measured as the MS spectrum.

Gases used in Ion traps

One rough pump and two turbomolecular pumps were used to create four stages of vacuum within the system (Refer to drawing). Helium gas was used for cooling the ions during trapping and also as a collision gas for fragmentation. Nitrogen gas was used generally for keeping the system clean, nebulising samples and drying liquid droplets.

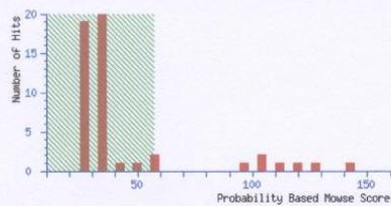
G2 Mass spectrometric analysis data for differentially expressed spots *E. cancerogenus*

MASCOT (SCIENCE) **Mascot Search Results**

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 2302
 MS data file : D:\Data\Lakshmy\120808\140808\SSP2302_RE13_01_281.d\SSP2302_RE13_01_281.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 11:11:44 GMT
 Protein hits :
 gi|126496 RecName: Full=Lysostaphin; AltName: Full=Glycyl-glycine endopeptidase; Flags: Precursor
 gi|261342624 transcription elongation factor GreA [Enterobacter cancerogenus ATCC 35316]
 gi|16762061 transcription elongation factor GreA [Salmonella enterica subsp. enterica serovar Typhi str. CT18]
 gi|37528357 transcription elongation factor GreA [Photobacterium luminescens subsp. laumondii TT01]
 gi|226329694 hypothetical protein PROPEN_03606 [Proteus penneri ATCC 35198]
 gi|238750117 Transcription elongation factor greA [Yersinia rohdei ATCC 43380]
 gi|269137754 transcription elongation factor [Edwardsiella tarda EIB202]
 gi|238757567 Transcription elongation factor greA [Yersinia aldovae ATCC 35236]

Probability Based Mowse Score

Ions score is $-10 * \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: Help

Significance threshold p<: Max. number of hits:

Standard scoring: MudPIT scoring Ions score or expect cut-off: Show sub-sets:

Show pop-ups: Suppress pop-ups Sort unassigned: Require bold red:

Select All Select None Search Selected Error tolerant

1. [gi|126496](#) Mass: 51656 Score: 143 Queries matched: 2 emPAI: 0.13
 RecName: Full=Lysostaphin; AltName: Full=Glycyl-glycine endopeptidase; Flags: Precursor
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 102	819.8730	1637.7314	1637.7897	-0.0582	0	42	1.7	1	K.SESASFTPTNDIITR.T
<input checked="" type="checkbox"/> 182	1007.4620	2012.9094	2012.9336	-0.0241	0	101	2e-06	1	R.MVNSFSNSTAQDPMPLK.S

Proteins matching the same set of peptides:
[gi|153047](#) Mass: 42213 Score: 143 Queries matched: 2
 Lysostaphin (ttg start codon) [Staphylococcus simulans]
[gi|3287967](#) Mass: 53058 Score: 143 Queries matched: 2
 RecName: Full=Lysostaphin; AltName: Full=Glycyl-glycine endopeptidase; Flags: Precursor

2. [gi|261342624](#) Mass: 17799 Score: 125 Queries matched: 3 emPAI: 0.42
 transcription elongation factor GreA [Enterobacter cancerogenus ATCC 35316]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 72	656.3350	1310.6554	1310.7306	-0.0752	0	61	0.022	1	K.QNLISVNSPIAR.G
<input checked="" type="checkbox"/> 74	667.8120	1333.6094	1333.6765	-0.0671	0	40	2.9	1	R.TPGGEVEYEIIK.V
<input checked="" type="checkbox"/> 145	919.9400	1837.8654	1837.9349	-0.0695	1	24	1.1e+02	1	R.TPGGEVEYEIIRVEYL.-

3. [gi|16762061](#) Mass: 17703 Score: 120 Queries matched: 3 emPAI: 0.69
 transcription elongation factor GreA [Salmonella enterica subsp. enterica serovar Typhi str. CT18]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 70	655.3790	1308.7434	1308.7513	-0.0079	0	(57)	0.059	1	R.RPEIIAALAEAR.E
<input checked="" type="checkbox"/> 71	437.2600	1308.7582	1308.7513	0.0068	0	58	0.039	1	R.RPEIIAALAEAR.E
<input checked="" type="checkbox"/> 72	656.3350	1310.6554	1310.7306	-0.0752	0	61	0.022	1	K.QNLISVNSPIAR.G

Proteins matching the same set of peptides:
[gi|24114470](#) Mass: 17688 Score: 120 Queries matched: 3
 transcription elongation factor GreA [Shigella flexneri 2a str. 301]

Mascot Search Results

Protein View

Match to: **gi|261342624** Score: **125**
transcription elongation factor GreA [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\140808\SSP2302_RE13_01_281.d\SSP2302_RE13_01_281.mgf

Nominal mass (M_r): **17799**; Calculated pI value: **4.82**
 NCBI BLAST search of **gi|261342624** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **17%**

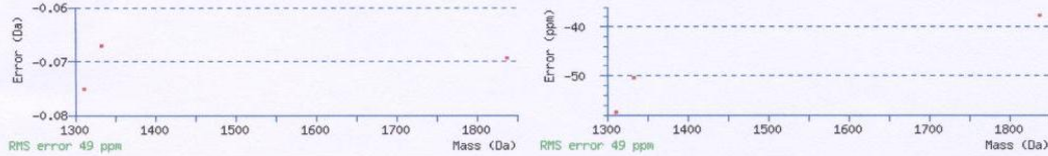
Matched peptides shown in **Bold Red**

- 1** MQAIPMTLRG AEKLRREELDF LKSVRRPEII AAIADAREHG DLKENAEYHA
- 51** AREQQGFCEG RIKDIEAKLS NAQVIDITKM PNNGRVIFGA TTVLNLNDND
- 101** EEQRYRIVGD DEADFK**QNL**I **SVNSPIAR**GL IGREQDDVVV I**RTPGGEVEY**
- 151** **EIKRVEYL**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
117 - 128	656.3350	1310.6554	1310.7306	-0.0752	0 K.QNLISVNSPIAR.G (Ions_score 61)
143 - 154	667.8120	1333.6094	1333.6765	-0.0671	0 R.TPGGEVEYEIIK.V (Ions_score 40)
143 - 158	919.9400	1837.8654	1837.9349	-0.0695	1 R.TPGGEVEYEIIKRVEYL.- (Ions_score 24)



LOCUS ZP_05970482 158 aa linear BCT 15-OCT-2009
 DEFINITION transcription elongation factor GreA [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05970482
 VERSION ZP_05970482.1 GI:261342624
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000035.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 158)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B., Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C., Delahaanty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C., Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M., Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 REFERENCE 2 (residues 1 to 158)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H., Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 COMMENT WGS [REFSEQ](#): This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from [ABWM02000035](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no

http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100105/FtmmSGSS... 21/01/2010

manual curation.

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project (<http://nihroadmap.nih.gov/hmp/>).

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7%. Sequencing Technology: 454.

Method: conceptual translation.

```

FEATURES
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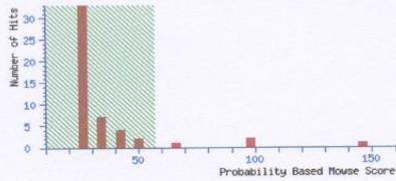
Mascot: <http://www.matrixscience.com/>

MASCOT Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 3002
 MS data file : D:\Data\Lakshmy\120808\260808\SSP 3002_RG16_01_396.d\SSP 3002_RG16_01_396.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 11:13:53 GMT
 Protein hits : [gi|42543180](#) Chain A, Ige Fv Spe7 Complexed With A Recombinant Thioredoxin
[gi|15802068](#) hypothetical protein Z2676 [Escherichia coli O157:H7 EDL933]
[gi|238920001](#) hypothetical protein NT01EI_2105 [Edwardsiella ictaluri 93-146]
[gi|154310594](#) putative thioredoxin [Photobacterium profundum SS9]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|42543180](#) Mass: 13560 Score: 146 Queries matched: 4 emPAI: 0.96
 Chain A, Ige Fv Spe7 Complexed With A Recombinant Thioredoxin
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 128	607.9250	1820.7532	1820.8866	-0.1334	0	(21) 1.8e+02	1		K.MIAPILDEIADEYQGG.L + Oxidation (M)
<input checked="" type="checkbox"/> 129	607.9690	1820.8852	1820.8866	-0.0014	0	(61) 0.022	1		K.MIAPILDEIADEYQGG.L + Oxidation (M)
<input checked="" type="checkbox"/> 130	911.4580	1820.9014	1820.8866	0.0149	0	113 1.3e-07	1		K.MIAPILDEIADEYQGG.L + Oxidation (M)
<input checked="" type="checkbox"/> 131	913.7260	2738.1562	2738.3183	-0.1621	1	35	11	1	R.YDLVGGCRMIAPILDEIADEYQGG.L

2. [gi|15802068](#) Mass: 13044 Score: 101 Queries matched: 2 emPAI: 0.26
 hypothetical protein Z2676 [Escherichia coli O157:H7 EDL933]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 81	724.9020	1447.7894	1447.7744	0.0150	0	32	18	1	R.QIAENPILLYMK.G + Oxidation (M)
<input checked="" type="checkbox"/> 91	782.4400	1562.8654	1562.8093	0.0562	0	69	0.0033	1	R.FAYVDILGNPDIR.A

Proteins matching the same set of peptides:

- [gi|16760480](#) Mass: 13074 Score: 101 Queries matched: 2
hypothetical protein STY1689 [Salmonella enterica subsp. enterica serovar Typhi str. CT18]
- [gi|150120857](#) Mass: 13245 Score: 101 Queries matched: 2
hypothetical protein ECA1927 [Pectobacterium atrosepticum SCRI1043]
- [gi|174312019](#) Mass: 13044 Score: 101 Queries matched: 2
hypothetical protein SSON_1502 [Shigella sonnei Ss046]
- [gi|1123442416](#) Mass: 13060 Score: 101 Queries matched: 2
hypothetical protein YE2155 [Yersinia enterocolitica subsp. enterocolitica 8081]
- [gi|1152970544](#) Mass: 13068 Score: 101 Queries matched: 2
hypothetical protein KFN_01992 [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
- [gi|1156934186](#) Mass: 12933 Score: 101 Queries matched: 2
hypothetical protein ESA_02013 [Enterobacter sakazakii ATCC BAA-894]
- [gi|1157145914](#) Mass: 13074 Score: 101 Queries matched: 2
hypothetical protein CKO_01665 [Citrobacter koseri ATCC BAA-895]
- [gi|1161503474](#) Mass: 13002 Score: 101 Queries matched: 2
hypothetical protein SARI_01548 [Salmonella enterica subsp. arizonae serovar 62:z4,z23:--]
- [gi|1170768703](#) Mass: 13030 Score: 101 Queries matched: 2
putative monothiol glutaredoxin [Escherichia albertii TW07627]
- [gi|108533942](#) Mass: 12734 Score: 101 Queries matched: 2
hypothetical protein ETA_18040 [Erwinia tasmaniensis ET1/99]
- [gi|1227111796](#) Mass: 13212 Score: 101 Queries matched: 2
hypothetical protein PcarbP_02467 [Pectobacterium carotovorum subsp. brasiliensis PBR1692]
- [gi|237731383](#) Mass: 13089 Score: 101 Queries matched: 2

conserved hypothetical protein [Citrobacter sp. 30_2]
 gi|238749425 Mass: 13318 Score: 101 Queries matched: 2
 Glutaredoxin-4 [Yersinia rohdei ATCC 43380]
 gi|238782581 Mass: 13289 Score: 101 Queries matched: 2
 Glutaredoxin-4 [Yersinia bercovieri ATCC 43970]
 gi|238787142 Mass: 13288 Score: 101 Queries matched: 2
 Glutaredoxin-4 [Yersinia frederiksenii ATCC 33641]
 gi|238792292 Mass: 13262 Score: 101 Queries matched: 2
 Glutaredoxin-4 [Yersinia intermedia ATCC 29909]
 gi|241913123 Mass: 15206 Score: 101 Queries matched: 2
 Chain A, Structure Of E. Coli Monothiol Glutaredoxin Grx4 Homodimer
 gi|242239124 Mass: 12992 Score: 101 Queries matched: 2
 glutaredoxin-like protein [Dickeya dadantii Ech703]
 gi|251789329 Mass: 13316 Score: 101 Queries matched: 2
 glutaredoxin-like protein [Dickeya zeae Ech1591]
 gi|253688763 Mass: 13228 Score: 101 Queries matched: 2
 glutaredoxin-like protein [Pectobacterium carotovorum subsp. carotovorum PC1]
 gi|258635113 Mass: 12512 Score: 101 Queries matched: 2
 glutaredoxin-like protein [Pantoea sp. At-9b]
 gi|259908543 Mass: 12733 Score: 101 Queries matched: 2
 Putative glutaredoxin protein Ydh [Erwinia pyrifoliae Epl/96]
 gi|260597772 Mass: 12927 Score: 101 Queries matched: 2
 Glutaredoxin-4 [Cronobacter turicensis]
 gi|261339586 Mass: 13045 Score: 101 Queries matched: 2
 hypothetical protein EcanA3_03947 [Enterobacter cancerogenus ATCC 35316]
 gi|261821927 Mass: 13259 Score: 101 Queries matched: 2
 glutaredoxin-like protein [Pectobacterium wasabiae WPP163]
 gi|269139028 Mass: 12773 Score: 101 Queries matched: 2
 hypothetical protein ETAE_1679 [Edwardsiella tarda EIB202]
 gi|271500215 Mass: 13334 Score: 101 Queries matched: 2
 glutaredoxin-like protein [Dickeya dadantii Ech586]

3. gi|236920001 Mass: 12760 Score: 101 Queries matched: 2 emPAI: 0.27
 hypothetical protein NT01EI_2105 [Edwardsiella ictaluri 93-146]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
81	724.9020	1447.7894	1447.7744	0.0150	0	32	18	1	R.QIAENPILYMK.G + Oxidation (M)
91	782.4400	1562.8654	1562.8093	0.0562	0	69	0.0033	1	R.FAYVDILQNPDIR.A

4. gi|54310594 Score: 65 Queries matched: 3
 putative thioredoxin [Photobacterium profundum SS9]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
128	607.9250	1820.7532	1820.8866	-0.1334	0	(19)	3.2e+02	2	K.MIAPILDEIANEYEGK.V + Oxidation (M)
129	607.9690	1820.8852	1820.8866	-0.0014	0	(58)	0.044	2	K.MIAPILDEIANEYEGK.V + Oxidation (M)
130	911.4580	1820.9014	1820.8866	0.0149	0	65	0.0079	2	K.MIAPILDEIANEYEGK.V + Oxidation (M)

Proteins matching the same set of peptides:
 gi|84393642 Score: 65 Queries matched: 3
 gi|90414228 Score: 65 Queries matched: 3

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
70	679.3270	1356.6394	1356.7976	-0.1582	2	45	1	1	LNDISILKAKK
55	515.2470	1028.4794	1028.5866	-0.1071	0	33	18	1	VLDPELLTAR
65	632.3680	1262.7214	1263.6320	-0.9105	1	31	23	1	REQVQNTAYR
71	681.1670	2040.4792	2040.1367	0.3424	1	29	43	1	NIVSSTELHQNVKAFIK
52	492.3030	982.5914	983.5916	-1.0002	1	29	32	1	KIATWLFPR
58	582.1380	1743.3922	1743.8540	-0.4618	0	29	50	1	NLPLPLYSQFNASASSR
198	737.6880	2210.0422	2211.0598	-1.0177	0	28	33	1	EHPRFPVMSSTASQMLVER + Oxidation (M)
136	615.8910	1844.6512	1845.0472	-0.3960	1	26	44	1	ILNDSIAYIRSLAQLR
225	1164.4980	3490.4722	3489.7207	0.7515	2	26	77	1	DYGFILVHDASPVKMGMPVLPMTGSRQESVR + Oxidation (M)
77	701.4550	2101.3432	2101.0361	0.3070	1	26	1.1e+02	1	CAGLADGPRSSVLLIDEVDK
202	742.3760	2224.1062	2223.1681	0.9380	0	26	62	1	MLTGTATLPSTAAQPPAQIQR
44	465.7930	1394.3572	1393.6403	0.7169	0	26	1.5e+02	1	AAVIEAMAQCMK
121	900.0780	2697.2122	2698.3021	-1.0899	0	25	1.2e+02	1	QIGSQEAAAFSPSTLSMQAPSPAR + Oxidation (M)
193	1084.4090	3250.2052	3250.3690	-0.1639	1	25	92	1	MMFRGFLMPSASVNDWMDGVYQMAK + 2 Oxidation (M)
122	904.2980	1806.5814	1805.8180	0.7634	2	25	1e+02	1	VVGDFLDDNDRKGN
132	609.5150	1825.5232	1824.8135	0.7097	0	24	66	1	SMGNDTFVAVGFAMGIER + Oxidation (M)
106	847.9540	2540.8402	2540.3737	0.4664	1	24	1.6e+02	1	IELIKDYLIHPEGSVLITGDTK
241	905.5410	2713.6012	2714.4391	-0.8380	1	24	69	1	RIGLSAHDFAQLLGADEVILTANK
109	577.9310	1730.7712	1729.8999	0.8713	0	24	1.1e+02	1	NDSAIVQALEALFPSR
144	951.8300	2852.4682	2851.4279	1.0402	2	24	1.6e+02	1	KPGIVPEFEDEREIEIQEPIK
73	692.3460	1382.6774	1383.6704	-0.9929	0	23	1.3e+02	1	MSQSSALASFLDK
218	767.1490	2298.4252	2299.2001	-0.7749	1	23	84	1	FRFADTLGILDFFATFEAIR
152	974.8230	2921.4472	2920.3735	1.0737	2	23	1.8e+02	1	KEPMIADLARYENQISITGFCCASQK + Oxidation (M)
45	931.6700	930.6627	931.4611	-0.7983	0	23	1.7e+02	1	AETSQTPAK
89	771.6520	2311.9342	2311.1466	0.7876	2	23	1.9e+02	1	MHAQSLFCFRAARIAFASR
124	906.2870	2715.8392	2716.2477	-0.4086	2	23	1.7e+02	1	DPFSQGARAGDNFPYTGQTNATTR
53	499.3350	996.6554	996.4624	0.1930	0	23	1.4e+02	1	SQAALDAAAR
161	991.6440	2971.9102	2972.5351	-0.6249	2	23	1.8e+02	1	LKFDIVDVTAGRLHVEAMMDEIK + Oxidation (M)
221	1158.1100	3471.3082	3470.7359	0.5723	1	22	1.9e+02	1	HNGVDVAVMTAVGQALVNGVVGSGASTLTMQTAR + 2 Oxidation (M)

Mascot Search Results

Protein View

Match to: **gi|261339586** Score: **101**
hypothetical protein EcanA3_03947 [**Enterobacter cancerogenus ATCC 35316**]
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Nominal mass (M_r): **13045**; Calculated pI value: **4.87**
 NCBI BLAST search of [gi|261339586](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **21%**

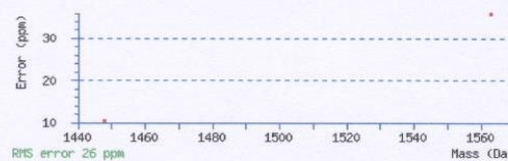
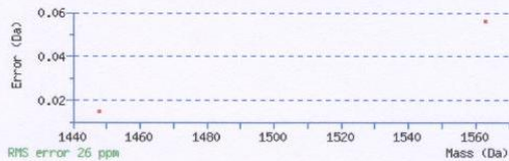
Matched peptides shown in **Bold Red**

- 1** MSTTIEKIQR **QIAENPILLY** MKGSPKLPSC GFSQAQVQAL SACGER**FAYV**
- 51** **DILQNPDIRA** ELPKYANWPT FPQLWIDGEL VGGCDILIEM YQRGELQQLI
- 101** KETAARYKTE EPGAE

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
11 - 22	724.9020	1447.7894	1447.7744	0.0150	0	R.QIAENPILLYMK.G Oxidation (M) (Ions score 32)
47 - 59	782.4400	1562.8654	1562.8093	0.0562	0	R.FAYVDILQNPDIR.A (Ions score 69)



LOCUS ZP_05967444 115 aa linear BCT 15-OCT-2009
 DEFINITION hypothetical protein EcanA3_03947 [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05967444
 VERSION ZP_05967444.1 GI:261339586
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000005.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 115)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B., Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C., Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C., Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M., Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 REFERENCE 2 (residues 1 to 115)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H., Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 COMMENT WGS [REFSEQ](#): This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from [ABWM02000005](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter

cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454.

Method: conceptual translation.

```

FEATURES             Location/Qualifiers
    source            1..115
                     /organism="Enterobacter cancerogenus ATCC 35316"
                     /strain="ATCC 35316"
                     /db_xref="taxon:500639"
    Protein           1..115
                     /product="hypothetical protein"
                     /calculated_mol_wt=12748
    Region            8..97
                     /region_name="GRX_PICOT like"
                     /note="Glutaredoxin (GRX) family, PKC-interacting cousin
                     of TRX (PICOT)-like subfamily; composed of PICOT and
                     GRX-PICOT-like proteins. The non-PICOT members of this
                     family contain only the GRX-like domain, whereas PICOT
                     contains an N-terminal TRX-like domain...; cd03028"
                     /db_xref="CDD:48577"
    Site              order(22,30..32,70..71)
                     /site_type="other"
                     /note="putative GSH binding site"
                     /db_xref="CDD:48577"
    Site              order(30,33)
                     /site_type="other"
                     /note="catalytic residues"
                     /db_xref="CDD:48577"
    CDS               1..115
                     /locus_tag="EcanA3_020100003947"
                     /coded_by="NZ_ABWM02000005.1:254505..254852"
                     /note="COG0278 Glutaredoxin-related protein"
                     /transl_table=11
                     /db_xref="CDD:138199"
    
```

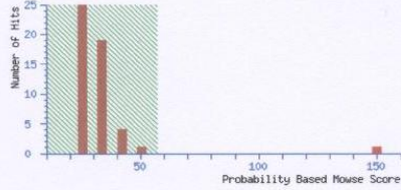
Mascot: <http://www.matrixscience.com/>

MASCOT Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4008
 MS data file : D:\Data\meng\20080728\SSP4008_RB7_01_222.d\SSP4008_RB7_01_222.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 11:17:21 GMT
 Protein hits : [gi|15803106](#) autonomous glycy radical cofactor GrcA [Escherichia coli O157:H7 EDL933]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: **Peptide Summary** [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|15803106](#) Mass: 14283 Score: 150 Queries matched: 4 emPAI: 0.53
 autonomous glycy radical cofactor GrcA [Escherichia coli O157:H7 EDL933]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 47	445.7760	889.5374	889.5021	0.0353	0	38	5.7	1	K.YFQLTIR.V
<input checked="" type="checkbox"/> 110	478.9540	1433.8402	1433.7739	0.0663	0	(37)	6.2	1	R.VEGGQHILNVNVL.R
<input checked="" type="checkbox"/> 111	717.9630	1433.9114	1433.7739	0.1375	0	39	3.3	1	R.VEGGQHILNVNVL.R
<input checked="" type="checkbox"/> 184	976.0110	1950.0074	1949.9370	0.0704	0	75	0.00082	1	K.AANDLLNSFWLDSEK.G

Proteins matching the same set of peptides:

- [gi|15832699](#) Mass: 14333 Score: 150 Queries matched: 4
autonomous glycy radical cofactor GrcA [Escherichia coli O157:H7 str. Sakai]
- [gi|16761504](#) Mass: 14393 Score: 150 Queries matched: 4
autonomous glycy radical cofactor GrcA [Salmonella enterica subsp. enterica serovar Typhi str. CT18]
- [gi|82778007](#) Mass: 14375 Score: 150 Queries matched: 4
autonomous glycy radical cofactor GrcA [Shigella dysenteriae Sd197]
- [gi|110642741](#) Mass: 14317 Score: 150 Queries matched: 4
autonomous glycy radical cofactor GrcA [Escherichia coli 536]
- [gi|146312705](#) Mass: 14385 Score: 150 Queries matched: 4
autonomous glycy radical cofactor GrcA [Enterobacter sp. 638]
- [gi|152971433](#) Mass: 14350 Score: 150 Queries matched: 4
autonomous glycy radical cofactor GrcA [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
- [gi|157144488](#) Mass: 14376 Score: 150 Queries matched: 4
autonomous glycy radical cofactor GrcA [Citrobacter koseri ATCC BAA-895]
- [gi|198244654](#) Mass: 14403 Score: 150 Queries matched: 4
autonomous glycy radical cofactor [Salmonella enterica subsp. enterica serovar Dublin str. CT_02021853]
- [gi|237705089](#) Mass: 15033 Score: 150 Queries matched: 4
yfiD [Escherichia sp. 3_2_53FAA]
- [gi|237732565](#) Mass: 15009 Score: 150 Queries matched: 4
protein yfiD [Citrobacter sp. 30_2]
- [gi|261340878](#) Mass: 14490 Score: 150 Queries matched: 4
autonomous glycy radical cofactor GrcA [Enterobacter cancerogenus ATCC 35316]
- [gi|262040312](#) Mass: 14378 Score: 150 Queries matched: 4
autonomous glycy radical cofactor [Klebsiella pneumoniae subsp. rhinoscleromatis ATCC 13884]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 96	681.3590	1360.7034	1360.6470	0.0564	2	49	0.36	1	KSGGRSPEDEAEK
<input checked="" type="checkbox"/> 95	679.3320	1356.6494	1356.7976	-0.1482	2	37	6.7	1	LNDIDSILKAKK
<input checked="" type="checkbox"/> 220	1113.0770	2224.1394	2224.0035	0.1360	2	36	5.9	1	MCAEVARRGGELVDGGLMR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 83	629.3340	1256.6534	1256.7816	-0.1281	2	33	15	1	IKASLQRTIQK
<input checked="" type="checkbox"/> 137	553.9480	1658.8222	1658.8410	-0.0188	2	32	16	1	THARGRELSDPEGK

mhtml:file://C:\Documents and Settings\larcje1\Local Settings\Temp\Peptide Summar... 21/01/2010

{MATRIX} Mascot Search Results
{SCIENCE}

Protein View

Match to: **gi|261340878** Score: **150**
autonomous glycy radical cofactor GrcA [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\meng\20080728\SSP4008_RB7_01_222.d\SSP4008_RB7_01_222.mgf

Nominal mass (M_r): **14490**; Calculated pI value: **4.85**
 NCBI BLAST search of **gi|261340878** against nr
 Unformatted [sequence string](#) for pasting into other applications
 Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)
 Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **29%**

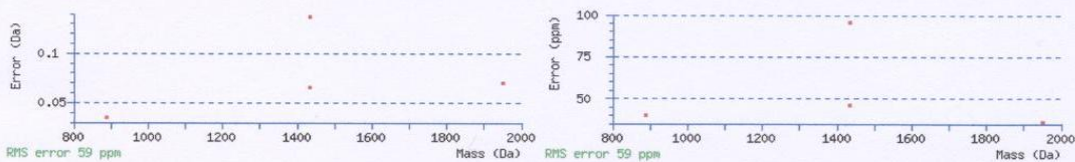
Matched peptides shown in **Bold Red**

- 1** MITGIIQITKA **ANDDLINSEFW LLDSEK**NEAR CVVAKAGFAE DEIVPVNKLIG
- 51** EIEYREIPME VQPEVR**VEGG QHLNVNVLRR** ETLMDAVEHP EKYPQLTIRV
- 101** SGYAVRFNSL TPEQQRDVIA RTFTESL

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
10 - 26	976.0110	1950.0074	1949.9370	0.0704	0	K.AANDDLINSEFWLLDSEK.N (Ions score 75)
67 - 79	478.9540	1433.8402	1433.7739	0.0663	0	R.VEGGQHLNVNVLRR (Ions score 37)
67 - 79	717.9630	1433.9114	1433.7739	0.1375	0	R.VEGGQHLNVNVLRR (Ions score 39)
93 - 99	445.7760	889.5374	889.5021	0.0353	0	K.YPQLTIR.V (Ions score 38)



LOCUS ZP_05968736 127 aa linear BCT 15-OCT-2009
 DEFINITION autonomous glycy radical cofactor GrcA [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05968736
 VERSION ZP_05968736.1 GI:261340878
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000010.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM Enterobacter cancerogenus ATCC 35316
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 127)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B., Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C., Delahaanty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C., Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M., Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 REFERENCE 2 (residues 1 to 127)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H., Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from [ABWM02000010](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no

manual curation.
Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x
Sequencing Technology: 454.
Method: conceptual translation.

```
FEATURES             Location/Qualifiers
    source             1..127
                     /organism="Enterobacter cancerogenus ATCC 35316"
                     /strain="ATCC 35316"
                     /db_xref="taxon:500639"
    Protein            1..127
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                     /calculated_mol_wt=14310
    Region             <69..123
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                     /note="RNR_PFL. Ribonucleotide reductase (RNR) and
                     pyruvate formate lyase (PFL) have a structurally similar
                     ten-stranded alpha-beta barrel active site domain and are
                     believed to have diverged from a common ancestor. RNRs
                     are found in all organisms and...; c109939"
    CDS                /db_xref="CDD:142336"
                     1..127
                     /locus_tag="EcanA3_020100010438"
                     /coded_by="complement(NZ_ABWM02000010.1:359251..359634)"
                     /note="COG3445 Acid-induced glycyI radical enzyme"
                     /transl_table=11
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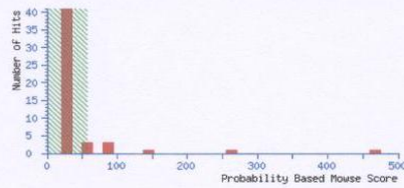
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 4602
MS data file : D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf
Database : NCBItr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 11:19:04 GMT
Protein hits : [gi|261339284](#) outer membrane protein A [Enterobacter cancerogenus ATCC 35316]
[gi|62901665](#) OmpA [Cronobacter sakazakii]
[gi|157146352](#) outer membrane protein A [Citrobacter koseri ATCC BAA-895]
[gi|262039986](#) outer membrane protein A [Klebsiella pneumoniae subsp. rhinoscleromatis ATCC 13884]
[gi|14578722](#) major outer membrane protein [Pectobacterium cypripedi]
[gi|129136](#) RecName: Full=Outer membrane protein A; Flags: Precursor

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p < 0.05$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|261339284](#) Mass: 37561 Score: 468 Queries matched: 16 emPAI: 0.53
 outer membrane protein A [Enterobacter cancerogenus ATCC 35316]
 Check to include this hit in error tolerant search

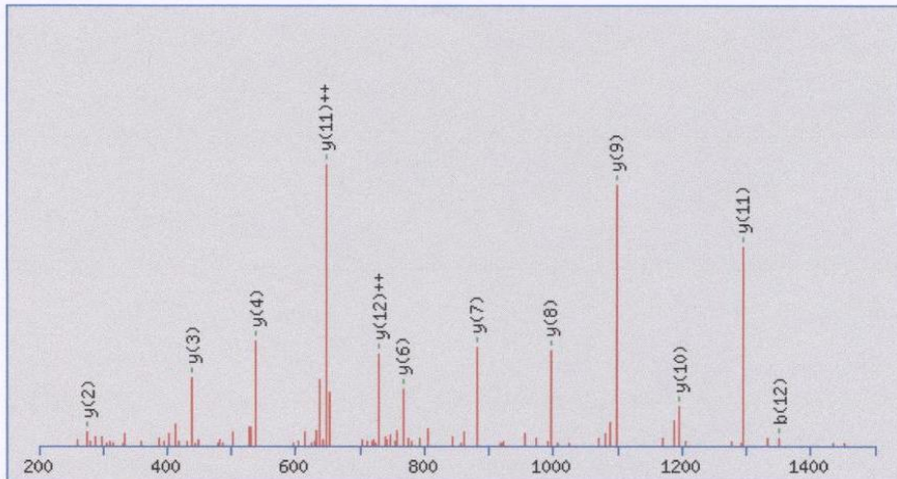
Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 84	542.2320	1082.4494	1082.5397	-0.0902	0	39	3.3	1	K.SDVLFNFK.A
<input checked="" type="checkbox"/> 94	604.8540	1207.6934	1207.6449	0.0486	0	44	1.1	1	R.AQSVVDYLVSK.G
<input checked="" type="checkbox"/> 99	632.8000	1263.5854	1263.6460	-0.0605	0	65	0.01	1	K.DGSSVVLGFTDR.I
<input checked="" type="checkbox"/> 104	455.2530	1362.7372	1363.7460	-1.0088	1	(50)	0.26	1	K.RAQSVVDYLVSK.G
<input checked="" type="checkbox"/> 105	682.3840	1362.7534	1363.7460	-0.9925	1	51	0.25	1	K.RAQSVVDYLVSK.G
<input checked="" type="checkbox"/> 127	813.6590	1625.3034	1625.7937	-0.4903	0	(22)	2.3e+02	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 128	813.8930	1625.7714	1625.7937	-0.0223	0	90	2.7e-05	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 129	813.9310	1625.8474	1625.7937	0.0537	0	(62)	0.016	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 130	813.9930	1625.9714	1625.7937	0.1777	0	(58)	0.039	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 131	814.4100	1626.8054	1625.7937	1.0117	0	(69)	0.0038	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 228	783.0930	2346.2572	2346.1856	0.0716	0	57	0.046	1	R.FGQGEDAPVVPAPAPAPEVQTR.H
<input checked="" type="checkbox"/> 244	890.8220	2669.4442	2670.3864	-0.9423	0	(33)	9.2	1	K.ATLKEGQALDQLYTQLSNLDPK.D
<input checked="" type="checkbox"/> 243	891.1400	2670.3982	2670.3864	0.0117	0	52	0.14	1	K.ATLKEGQALDQLYTQLSNLDPK.D
<input checked="" type="checkbox"/> 245	891.4750	2671.4032	2670.3864	1.0167	0	(21)	1.7e+02	8	K.ATLKEGQALDQLYTQLSNLDPK.D
<input checked="" type="checkbox"/> 192	1017.1370	3048.3892	3048.3520	0.0372	0	71	0.003	1	R.ADSSNFIAGDDHDTGVSPVFAGGVEWAMTR.D
<input checked="" type="checkbox"/> 195	1022.5150	3064.5232	3064.3469	0.1763	0	(24)	1.6e+02	1	R.ADSSNFIAGDDHDTGVSPVFAGGVEWAMTR.D + Oxidation (M)

2. [gi|62901665](#) Mass: 37147 Score: 258 Queries matched: 9 emPAI: 0.29
 OmpA [Cronobacter sakazakii]
 Check to include this hit in error tolerant search

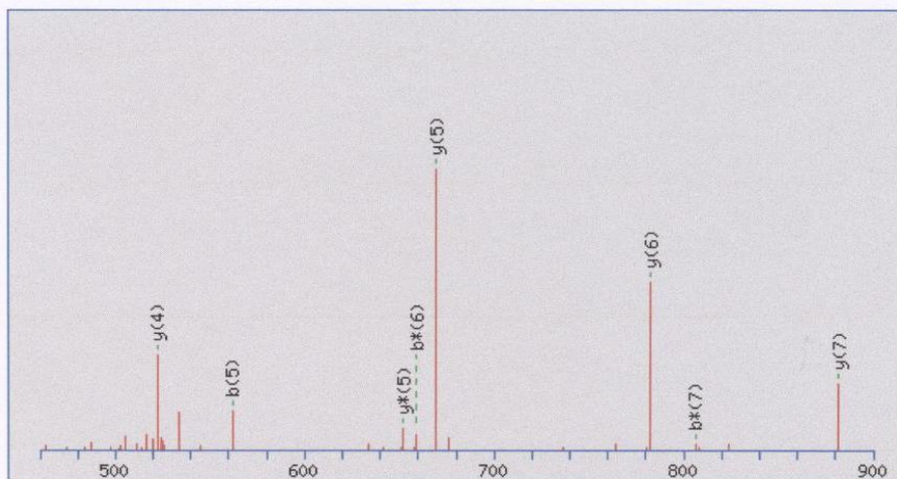
Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 84	542.2320	1082.4494	1082.5397	-0.0902	0	39	3.3	1	K.SDVLFNFK.A
<input checked="" type="checkbox"/> 99	632.8000	1263.5854	1263.6460	-0.0605	0	65	0.01	1	K.DGSSVVLGFTDR.I
<input checked="" type="checkbox"/> 127	813.6590	1625.3034	1625.7937	-0.4903	0	(22)	2.3e+02	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 128	813.8930	1625.7714	1625.7937	-0.0223	0	90	2.7e-05	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 129	813.9310	1625.8474	1625.7937	0.0537	0	(62)	0.016	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 130	813.9930	1625.9714	1625.7937	0.1777	0	(58)	0.039	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 131	814.4100	1626.8054	1625.7937	1.0117	0	(69)	0.0038	1	K.LGYFVTDDLQVYTR.L
<input checked="" type="checkbox"/> 192	1017.1370	3048.3892	3048.3520	0.0372	0	65	0.012	2	R.ADSSNFIAGDDHDTGVSPVFAGGVEWAMTR.D
<input checked="" type="checkbox"/> 195	1022.5150	3064.5232	3064.3469	0.1763	0	(24)	1.6e+02	1	R.ADSSNFIAGDDHDTGVSPVFAGGVEWAMTR.D + Oxidation (M)

Enterobacter cancerogenus SSP 4602 peptide sequences

LGYPVTDDLVDVYTR – Ion score 90



SDVLFNFNK- Ion score 39



Mascot Search Results

Protein View

Match to: gi|261339284 Score: 468
outer membrane protein A [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

Nominal mass (M_n): 37561; Calculated pI value: 5.19
 NCBI BLAST search of gi|261339284 against nr
 Unformatted sequence string for pasting into other applications

Taxonomy: Enterobacter cancerogenus ATCC 35316

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 35%

Matched peptides shown in **Bold Red**

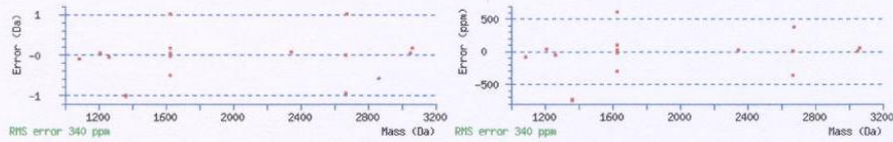
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1 MKKTAIAIIV ALAGFATVAQ AAKPKDNTWYA GKGKLGWSQFH DTGWYNSSLIAN
51 NDGPTHESQL GAGAFGGYQV NPYVGFEMGY DWLGRMPEYK DNUNGAFAKQ
101 GVOLTAKLGY EYTDLDLVYT RLGGWVWRAD SSNSIAGDDH DTGVSFVFAAG
151 GVENAMTRDI ATRLEYQWVN NIGDGTATGVV RPDNGLSVG VSYRFGQGED
201 APVVAPAPAP APEVQTKHFT LKSDVLFNFN KATLKPEGQQ ALDQLYQLS
251 NLDPKDGSVV VLGFTDRIGS DAYNQLSEK RAQSVVDYLV SKGIPANKIS
301 PRMGESPNV TGMTCDNVKP RAALDCLAP DRRVEIEVKG IKDVTTPAA
351
    
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Show predicted peptides also

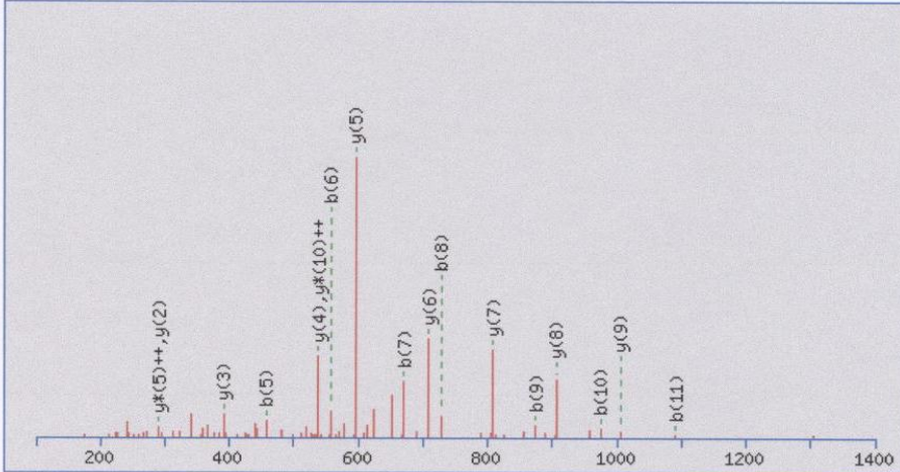
Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
108 - 121	813.6590	1625.3034	1625.7937	-0.4903	0 K.LGYFPTDLDLVYTR.L (Ions score 22)
108 - 121	813.8930	1625.7714	1625.7937	-0.0223	0 K.LGYFPTDLDLVYTR.L (Ions score 30)
108 - 121	813.9310	1625.8474	1625.7937	0.0537	0 K.LGYFPTDLDLVYTR.L (Ions score 62)
108 - 121	813.9930	1625.9714	1625.7937	0.1777	0 K.LGYFPTDLDLVYTR.L (Ions score 58)
108 - 121	814.4100	1626.8054	1625.7937	1.0117	0 K.LGYFPTDLDLVYTR.L (Ions score 59)
129 - 158	1017.1370	3048.3892	3048.3520	0.0372	0 R.ADSSNSIAGDDHDTGVSFVFAAGVWAMTR.D (Ions score 71)
129 - 158	1022.5150	3064.5232	3064.3469	0.1763	0 R.ADSSNSIAGDDHDTGVSFVFAAGVWAMTR.D Oxidation (M) (Ions score 24)
195 - 217	783.0930	2346.2572	2346.1856	0.0716	0 R.FGQQEDAPVVAPAPAPEVQTK.H (Ions score 57)
223 - 231	542.2320	1082.4494	1082.5397	-0.0902	0 K.SDVLFNFK.A (Ions score 39)
232 - 255	890.8220	2669.4442	2670.3864	-0.9423	0 K.ATLKPEGQQALDQLYQLSNLDPK.D (Ions score 33)
232 - 255	891.1400	2670.3982	2670.3864	0.0117	0 K.ATLKPEGQQALDQLYQLSNLDPK.D (Ions score 52)
232 - 255	891.4750	2671.4032	2670.3864	1.0167	0 K.ATLKPEGQQALDQLYQLSNLDPK.D (Ions score 21)
256 - 267	632.8000	1263.5854	1263.6460	-0.0605	0 K.DGSVVVLGFTDR.I (Ions score 65)
281 - 292	455.2530	1362.7372	1363.7460	-1.0088	1 K.RAQSVVDYLVSK.G (Ions score 50)
281 - 292	682.3840	1362.7534	1363.7460	-0.9925	1 K.RAQSVVDYLVSK.G (Ions score 51)
282 - 292	604.8540	1207.6934	1207.6449	0.0486	0 R.AQSVVDYLVSK.G (Ions score 44)



LOCUS ZP_05967142 350 aa linear BCT 15-OCT-2009
 DEFINITION outer membrane protein A [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05967142
 VERSION ZP_05967142.1 GI:261339284
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession NZ_ABWM02000004.1
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM Enterobacter cancerogenus ATCC 35316
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 350)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 350)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from ABWM02000004.
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be
 aware that the annotation is done automatically with little or no

DGSVVVLGFTDR- Ion score 65



manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

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                     /note="outer membrane protein A; Reviewed; PRK10808"
                     /db_xref="CDD:138188"
    Region            23..196
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                     /note="Surface antigen; c101155"
                     /db_xref="CDD:141128"
    Region            224..338
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                     /note="Peptidoglycan binding domains similar to the
                     C-terminal domain of outer-membrane protein OmpA; cd07185"
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                     peptidoglycan-associated (lipo)proteins"
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                     /db_xref="CDD:138188"
    
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Mascot: <http://www.matrixscience.com/>

(MATRIX) **Mascot Search Results**
(SCIENCES)

Peptide View

MS/MS Fragmentation of **LGYPVTDDLVDYTR**

Found in gi|261339284, outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

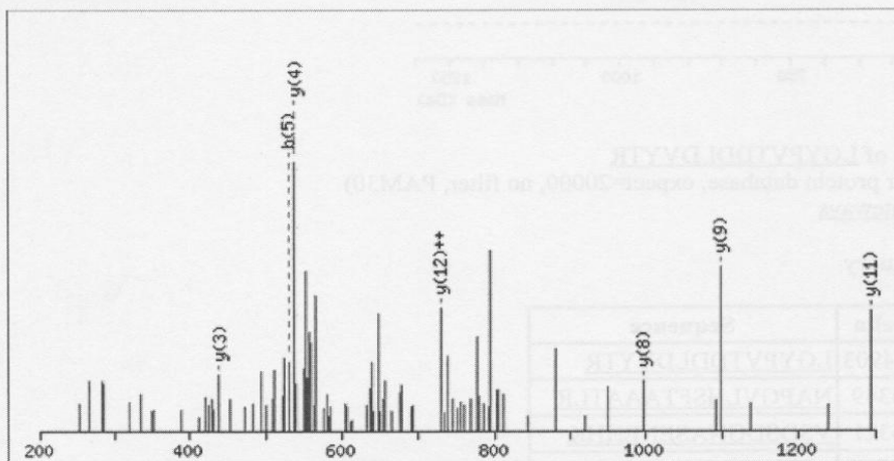
Match to Query 127: 1625.303448 from(813.659000,2+) intensity(2804959.0000)

Title: Cmpd 235, +MSn(814.24), 19.7 min

Data file D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 1625.7937

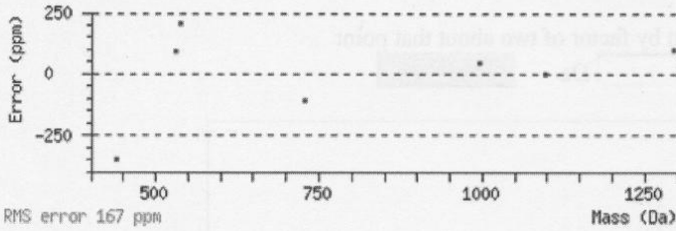
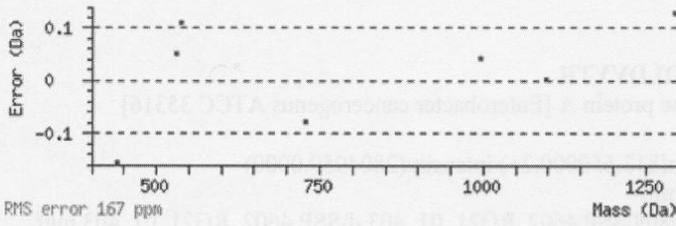
Fixed modifications: Carboxymethyl (C)

Ions Score: 22 Expect: 2.3e+02

Matches (Bold Red): 7/104 fragment ions using 18 most intense peaks

#	a	a ⁺⁺	b	b ⁺⁺	Seq.	y	y ⁺⁺	y*	y ^{*++}	#
1	86.0964	43.5519	114.0913	57.5493	L					14
2	143.1179	72.0626	171.1128	86.0600	G	1513.7169	757.3621	1496.6904	748.8488	13
3	306.1812	153.5942	334.1761	167.5917	Y	1456.6955	728.8514	1439.6689	720.3381	12
4	403.2340	202.1206	431.2289	216.1181	P	1293.6321	647.3197	1276.6056	638.8064	11
5	502.3024	251.6548	530.2973	265.6523	V	1196.5794	598.7933	1179.5528	590.2800	10
6	603.3501	302.1787	631.3450	316.1761	T	1097.5109	549.2591	1080.4844	540.7458	9
7	718.3770	359.6921	746.3719	373.6896	D	996.4633	498.7353	979.4367	490.2220	8
8	833.4040	417.2056	861.3989	431.2031	D	881.4363	441.2218	864.4098	432.7085	7
9	946.4880	473.7477	974.4829	487.7451	L	766.4094	383.7083	749.3828	375.1951	6
10	1061.5150	531.2611	1089.5099	545.2586	D	653.3253	327.1663	636.2988	318.6530	5
11	1160.5834	580.7953	1188.5783	594.7928	V	538.2984	269.6528	521.2718	261.1396	4
12	1323.6467	662.3270	1351.6416	676.3245	Y	439.2300	220.1186	422.2034	211.6053	3
13	1424.6944	712.8508	1452.6893	726.8483	T	276.1666	138.5870	259.1401	130.0737	2

14 | | | | | R | 175.1190 | 88.0631 | 158.0924 | 79.5498 | 1 |



NCBI **BLAST** search of LGYPVTDDLVDVYTR
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST web gateways

All matches to this query

Score	Mr(cal):	Delta	Sequence
21.8	1625.7937	-0.4903	<u>LGYPVTDDLVDVYTR</u>
11.7	1624.8685	0.4349	<u>NAPGVLHSFTAAATLR</u>
10.9	1624.6713	0.6321	<u>VSDSDGNASENEHHK</u>
10.6	1624.7468	0.5567	<u>AETINTDEVDVYEK</u>
10.6	1624.7986	0.5048	<u>MRVKVILACTECK</u>
10.6	1625.7607	-0.4573	<u>FMTGLDDDLTQVGAK</u>
9.6	1625.8049	-0.5015	<u>EHGDLEGPLKAFADK</u>
9.3	1625.9902	-0.6868	<u>GIVSPILQVVKAMKK</u>
9.1	1624.7072	0.5962	<u>KENCPVASSSTMDAK</u>
8.5	1625.7257	-0.4222	<u>WFGSGGDGVSVOGAMVR</u>

Mascot: <http://www.matrixscience.com/>

(MATRIX)
(SCIENCES) Mascot Search Results

Peptide View

MS/MS Fragmentation of LGYPVTDDLDVYTR

Found in gi|261339284, outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

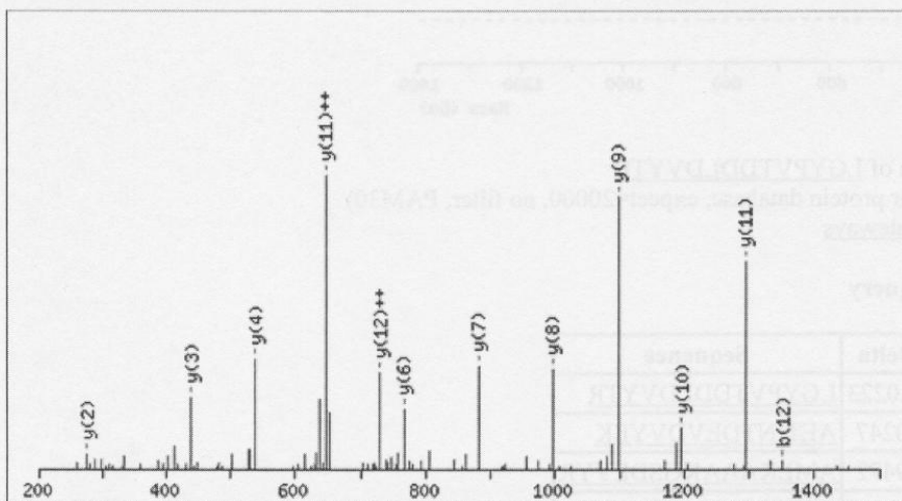
Match to Query 128: 1625.771448 from(813.893000,2+) intensity(85683688.0000)

Title: Cmpd 37, +MSn(814.35), 7.3 min

Data file D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 1625.7937

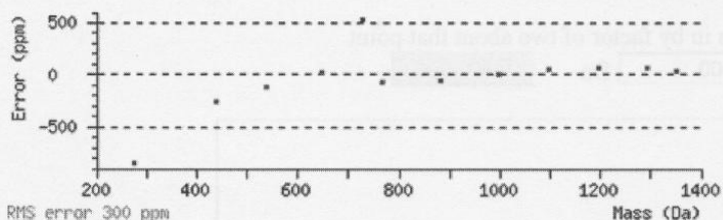
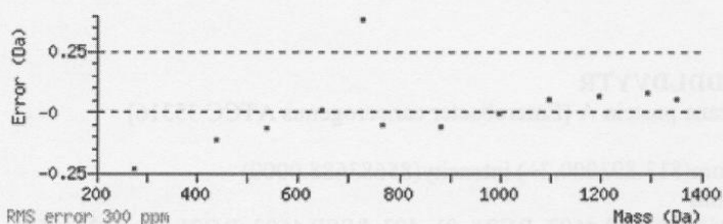
Fixed modifications: Carboxymethyl (C)

Ions Score: 90 Expect: 2.7e-05

Matches (Bold Red): 12/104 fragment ions using 15 most intense peaks

#	a	a ⁺⁺	b	b ⁺⁺	Seq.	y	y ⁺⁺	y*	y ⁺⁺⁺	#
1	86.0964	43.5519	114.0913	57.5493	L					14
2	143.1179	72.0626	171.1128	86.0600	G	1513.7169	757.3621	1496.6904	748.8488	13
3	306.1812	153.5942	334.1761	167.5917	Y	1456.6955	728.8514	1439.6689	720.3381	12
4	403.2340	202.1206	431.2289	216.1181	P	1293.6321	647.3197	1276.6056	638.8064	11
5	502.3024	251.6548	530.2973	265.6523	V	1196.5794	598.7933	1179.5528	590.2800	10
6	603.3501	302.1787	631.3450	316.1761	T	1097.5109	549.2591	1080.4844	540.7458	9
7	718.3770	359.6921	746.3719	373.6896	D	996.4633	498.7353	979.4367	490.2220	8
8	833.4040	417.2056	861.3989	431.2031	D	881.4363	441.2218	864.4098	432.7085	7
9	946.4880	473.7477	974.4829	487.7451	L	766.4094	383.7083	749.3828	375.1951	6
10	1061.5150	531.2611	1089.5099	545.2586	D	653.3253	327.1663	636.2988	318.6530	5
11	1160.5834	580.7953	1188.5783	594.7928	V	538.2984	269.6528	521.2718	261.1396	4
12	1323.6467	662.3270	1351.6416	676.3245	Y	439.2300	220.1186	422.2034	211.6053	3
13	1424.6944	712.8508	1452.6893	726.8483	T	276.1666	138.5870	259.1401	130.0737	2

14 | | | | | R | 175.1190 | 88.0631 | 158.0924 | 79.5498 | 1



NCBI **BLAST** search of LGYPVTDDLVDVYTR
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
90.0	1625.7937	-0.0223	<u>LGYPVTDDLVDVYTR</u>
33.0	1624.7468	1.0247	<u>AETINTDEVDVYEK</u>
28.1	1624.8242	0.9472	<u>AMLKAAAKGSDEVYR</u>
23.7	1625.8672	-0.0957	<u>MTTPVIVRSAAHSTR</u>
23.2	1624.7733	0.9982	<u>KFYDDKGNEVPGEK</u>
20.3	1625.6748	0.0966	<u>MECACARAHVYTR</u>
19.9	1625.9002	-0.1287	<u>RPGRVXFVDLSEVR</u>
18.5	1625.6553	0.1161	<u>DDHAEEPSDEAQQR</u>
18.4	1625.7798	-0.0083	<u>DLORDTPFNTYTR</u>
17.6	1625.8157	-0.0442	<u>GMSYLAEMLSLNGIK</u>

Mascot: <http://www.matrixscience.com/>

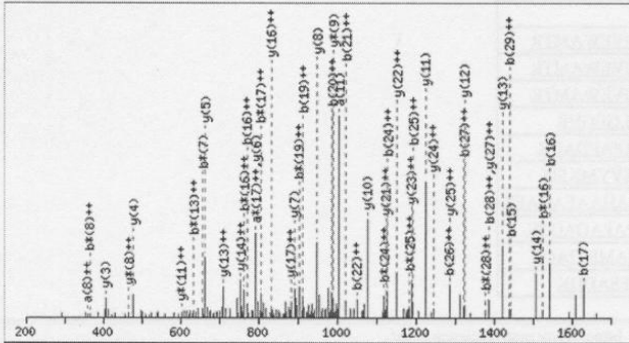
Mascot Search Results

Peptide View

MS/MS Fragmentation of ADSSNSIAGDDHDTGVSPVFAGGVEWAMTR
 Found in gi|261339284, outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

Match to Query 192: 3048.389172 from(1017.137000,3+) intensity(4976399.0000)
 Title: Cmpd 86, +MSn(1017.75), 9.2 min
 Data file D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

Click mouse within plot area to zoom in by factor of two about that point
 Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 3048.3520
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 71 Expect: 0.003
 Matches (Bold Red): 51/332 fragment ions using 88 most intense peaks

#	a	a ⁺⁺	a [*]	a ⁺⁺⁺	b	b ⁺⁺	b [*]	b ⁺⁺⁺	Seq.	y	y ⁺⁺	y [*]	y ⁺⁺⁺	#
1	44.0495	22.5284			72.0444	36.5258			A					30
2	159.0764	80.0418			187.0713	94.0393			D	2978.3221	1489.6647	2961.2956	1481.1514	29
3	246.1084	123.5579			274.1034	137.5553			S	2863.2952	1432.1512	2846.2686	1423.6379	28
4	333.1405	167.0739			361.1354	181.0713			S	2776.2631	1388.6352	2759.2366	1380.1219	27
5	447.1834	224.0953	430.1569	215.5821	475.1783	238.0928	458.1518	229.5795	N	2689.2311	1345.1192	2672.2046	1336.6059	26
6	534.2154	267.6114	517.1889	259.0981	562.2103	281.6088	545.1838	273.0955	S	2575.1882	1288.0977	2558.1616	1279.5845	25
7	647.2995	324.1534	630.2729	315.6401	675.2944	338.1508	658.2679	329.6376	I	2488.1562	1244.5817	2471.1296	1236.0684	24
8	718.3366	359.6719	701.3101	351.1587	746.3315	373.6694	729.3050	365.1561	A	2375.0721	1188.0397	2358.0455	1179.5264	23
9	775.3581	388.1827	758.3315	379.6694	803.3530	402.1801	786.3264	393.6669	G	2304.0350	1152.5211	2287.0084	1144.0079	22
10	890.3850	445.6961	873.3585	437.1829	918.3799	459.6936	901.3534	451.1803	D	2247.0135	1124.0104	2229.9870	1115.4971	21
11	1005.4120	503.2096	988.3854	494.6963	1033.4069	517.2071	1016.3803	508.6938	D	2131.9866	1066.4969	2114.9600	1057.9836	20
12	1142.4709	571.7391	1125.4443	563.2258	1170.4658	585.7365	1153.4392	577.2233	H	2016.9596	1008.9835	1999.9331	1000.4702	19
13	1257.4978	629.2525	1240.4713	620.7393	1285.4927	643.2500	1268.4662	634.7367	D	1879.9007	940.4540	1862.8742	931.9407	18
14	1358.5455	679.7764	1341.5189	671.2631	1386.5404	693.7738	1369.5139	685.2606	T	1764.8738	882.9405	1747.8472	874.4272	17
15	1415.5670	708.2871	1398.5404	699.7738	1443.5619	722.2846	1426.5353	713.7713	G	1663.8261	832.4167	1646.7995	823.9034	16
16	1514.6354	757.8213	1497.6088	749.3080	1542.6303	771.8188	1525.6037	763.3055	V	1606.8046	803.9060	1589.7781	795.3927	15
17	1601.6674	801.3373	1584.6408	792.8241	1629.6623	815.3348	1612.6358	806.8215	S	1507.7362	754.3717	1490.7097	745.8585	14
18	1698.7202	849.8637	1681.6936	841.3504	1726.7151	863.8612	1709.6885	855.3479	P	1420.7042	710.8557	1403.6776	702.3425	13
19	1797.7886	899.3979	1780.7620	890.8847	1825.7835	913.3954	1808.7569	904.8821	V	1323.6514	662.3293	1306.6249	653.8161	12
20	1944.8570	972.9321	1927.8304	964.4189	1972.8519	986.9296	1955.8254	978.4163	F	1224.5830	612.7951	1207.5565	604.2819	11
21	2015.8941	1008.4507	1998.8676	999.9374	2043.8890	1022.4481	2026.8625	1013.9349	A	1077.5146	539.2609	1060.4880	530.7477	10
22	2072.9156	1036.9614	2055.8890	1028.4481	2100.9105	1050.9589	2083.8839	1042.4456	G	1006.4775	503.7424	989.4509	495.2291	9
23	2129.9370	1065.4722	2112.9105	1056.9589	2157.9319	1079.4696	2140.9054	1070.9563	G	949.4560	475.2316	932.4295	466.7184	8
24	2229.0054	1115.0064	2211.9789	1106.4931	2257.0004	1129.0038	2239.9738	1120.4905	V	892.4346	446.7209	875.4080	438.2076	7
25	2358.0480	1179.5277	2341.0215	1171.0144	2386.0430	1193.5251	2369.0164	1185.0118	E	793.3661	397.1867	776.3396	388.6734	6
26	2544.1274	1272.5673	2527.1008	1264.0540	2572.1223	1286.5648	2555.0957	1278.0515	W	664.3235	332.6654	647.2970	324.1521	5
27	2615.1645	1308.0859	2598.1379	1299.5726	2643.1594	1322.0833	2626.1328	1313.5701	A	478.2442	239.6258	461.2177	231.1125	4
28	2746.2050	1373.6061	2729.1784	1365.0928	2774.1999	1387.6036	2757.1733	1379.0903	M	407.2071	204.1072	390.1806	195.5939	3
29	2847.2526	1424.1300	2830.2261	1415.6167	2875.2475	1438.1274	2858.2210	1429.6141	T	276.1666	138.5870	259.1401	130.0737	2
30									R	175.1190	88.0631	158.0924	79.5498	1

MASCOT Mascot Search Results

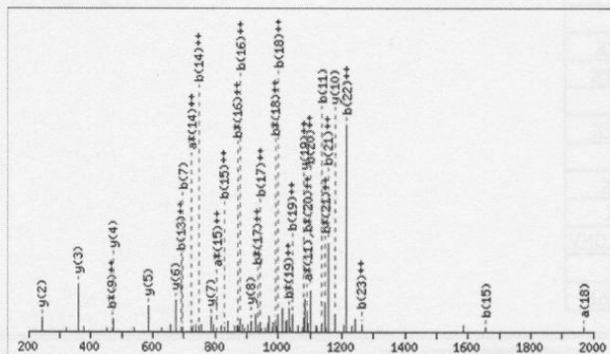
Peptide View

MS/MS Fragmentation of ATLKPEGQQALDQLYTQLSNLDPK
 Found in gi|261339284, outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

Match to Query 244: 2670.398172 from(891.140000,3+) intensity(46648868.0000)
 Title: Cmpd 82, +MSn(891.82), 9.1 min
 Data file D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

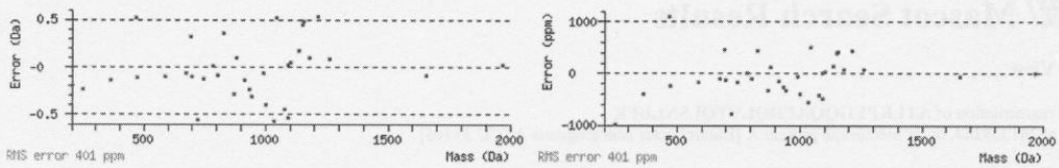
Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 2670.3864
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 52 Expect: 0.14
 Matches (Bold Red): 34/264 fragment ions using 62 most intense peaks

#	a	a ⁺⁺	a*	a ⁺⁺⁺	b	b ⁺⁺	b*	b ⁺⁺⁺	Seq.	y	y ⁺⁺	y*	y ⁺⁺⁺	#
1	44.0495	22.5284			72.0444	36.5258			A					24
2	145.0972	73.0522			173.0921	87.0497			T	2600.3566	1300.6820	2583.3301	1292.1687	23
3	258.1812	129.5942			286.1761	143.5917			L	2499.3089	1250.1581	2482.2824	1241.6448	22
4	386.2762	193.6417	369.2496	185.1285	414.2711	207.6392	397.2445	199.1259	K	2386.2249	1193.6161	2369.1983	1185.1028	21
5	483.3289	242.1681	466.3024	233.6548	511.3239	256.1656	494.2973	247.6523	P	2258.1299	1129.5686	2241.1034	1121.0553	20
6	612.3715	306.6894	595.3450	298.1761	640.3665	320.6869	623.3399	312.1736	E	2161.0772	1081.0422	2144.0506	1072.5289	19
7	669.3930	335.2001	652.3665	326.6869	697.3879	349.1976	680.3614	340.6843	G	2032.0346	1016.5209	2015.0080	1008.0076	18
8	797.4516	399.2294	780.4250	390.7162	825.4465	413.2269	808.4199	404.7136	Q	1975.0131	988.0102	1957.9866	979.4969	17
9	925.5102	463.2587	908.4836	454.7454	953.5051	477.2562	936.4785	468.7429	Q	1846.9545	923.9809	1829.9280	915.4676	16
10	996.5473	498.7773	979.5207	490.2640	1024.5422	512.7747	1007.5156	504.2615	A	1718.8959	859.9516	1701.8694	851.4383	15
11	1109.6313	555.3193	1092.6048	546.8060	1137.6263	569.3168	1120.5997	560.8035	L	1647.8588	824.4331	1630.8323	815.9198	14
12	1224.6583	612.8328	1207.6317	604.3195	1252.6532	626.8302	1235.6266	618.3170	D	1534.7748	767.8910	1517.7482	759.3777	13
13	1352.7169	676.8621	1335.6903	668.3488	1380.7118	690.8595	1363.6852	682.3462	Q	1419.7478	710.3775	1402.7213	701.8643	12
14	1465.8009	733.4041	1448.7744	724.8908	1493.7958	747.4016	1476.7693	738.8883	L	1291.6892	646.3483	1274.6627	637.8350	11
15	1628.8643	814.9358	1611.8377	806.4225	1656.8592	828.9332	1639.8326	820.4199	Y	1178.6052	589.8062	1161.5786	581.2930	10
16	1729.9119	865.4596	1712.8854	856.9463	1757.9068	879.4571	1740.8803	870.9438	T	1015.5419	508.2746	998.5153	499.7613	9
17	1857.9705	929.4889	1840.9440	920.9756	1885.9654	943.4863	1868.9389	934.9731	Q	914.4942	457.7507	897.4676	449.2374	8
18	1971.0546	986.0309	1954.0280	977.5176	1999.0495	1000.0284	1982.0229	991.5151	L	786.4356	393.7214	769.4090	385.2082	7
19	2058.0866	1029.5469	2041.0600	1021.0337	2086.0815	1043.5444	2069.0550	1035.0311	S	673.3515	337.1794	656.3250	328.6661	6
20	2172.1295	1086.5684	2155.1030	1078.0551	2200.1244	1100.5659	2183.0979	1092.0526	N	586.3195	293.6634	569.2930	285.1501	5
21	2285.2136	1143.1104	2268.1870	1134.5972	2313.2085	1157.1079	2296.1820	1148.5946	L	472.2766	236.6419	455.2500	228.1287	4
22	2400.2405	1200.6239	2383.2140	1192.1106	2428.2354	1214.6214	2411.2089	1206.1081	D	359.1925	180.0999	342.1660	171.5866	3
23	2497.2933	1249.1503	2480.2667	1240.6370	2525.2882	1263.1477	2508.2617	1254.6345	P	244.1656	122.5864	227.1390	114.0731	2
24									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of [ATLKPEGQOALDOLYTOLSNLDPK](#)
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc)	Delta	Sequence
51.9	2670.3864	0.0117	ATLKPEGQOALDOLYTOLSNLDPK
51.9	2670.3864	0.0117	ATLKPEGQOALDOLYTOLSNLDPK
34.7	2669.4891	0.9091	IIFEIDQALGIENVTLDIKDGLLK
34.4	2671.4068	-1.0087	ATLKPEGQOALDOLYTOLTTLDPK
32.8	2671.2118	-0.8137	NREIHEQECFNLNPATNVMNPK
30.7	2670.2806	0.1175	DNQMRPLTDEELLQEQATALPDK
29.4	2669.4170	0.9812	QINDEMTVONKALEIARQOOLSK
28.8	2670.3646	0.0335	AGNKGSEAAOGLLEMISLNOOILGNV
28.0	2671.2662	-0.8680	MVEPAEEQFSGFIFKIQANMDPK
27.9	2671.4908	-1.0927	FVIEEGIKESLLGGRTINATLVGGAK

Mascot: <http://www.matrixscience.com/>

Rank	Score	Protein	Accession	Length	Mass	Charge	Delta	Ion	Mod	Ident	Score	Protein	Accession	Length	Mass	Charge	Delta	Ion	Mod	Ident
1	51.9	ATLKPEGQOALDOLYTOLSNLDPK		16	2670.3864	1	0.0117			100	51.9	ATLKPEGQOALDOLYTOLSNLDPK		16	2670.3864	1	0.0117		100	
2	51.9	ATLKPEGQOALDOLYTOLSNLDPK		16	2670.3864	1	0.0117			100	51.9	ATLKPEGQOALDOLYTOLSNLDPK		16	2670.3864	1	0.0117		100	
3	34.7	IIFEIDQALGIENVTLDIKDGLLK		16	2669.4891	1	0.9091			100	34.7	IIFEIDQALGIENVTLDIKDGLLK		16	2669.4891	1	0.9091		100	
4	34.4	ATLKPEGQOALDOLYTOLTTLDPK		16	2671.4068	1	-1.0087			100	34.4	ATLKPEGQOALDOLYTOLTTLDPK		16	2671.4068	1	-1.0087		100	
5	32.8	NREIHEQECFNLNPATNVMNPK		16	2671.2118	1	-0.8137			100	32.8	NREIHEQECFNLNPATNVMNPK		16	2671.2118	1	-0.8137		100	
6	30.7	DNQMRPLTDEELLQEQATALPDK		16	2670.2806	1	0.1175			100	30.7	DNQMRPLTDEELLQEQATALPDK		16	2670.2806	1	0.1175		100	
7	29.4	QINDEMTVONKALEIARQOOLSK		16	2669.4170	1	0.9812			100	29.4	QINDEMTVONKALEIARQOOLSK		16	2669.4170	1	0.9812		100	
8	28.8	AGNKGSEAAOGLLEMISLNOOILGNV		16	2670.3646	1	0.0335			100	28.8	AGNKGSEAAOGLLEMISLNOOILGNV		16	2670.3646	1	0.0335		100	
9	28.0	MVEPAEEQFSGFIFKIQANMDPK		16	2671.2662	1	-0.8680			100	28.0	MVEPAEEQFSGFIFKIQANMDPK		16	2671.2662	1	-0.8680		100	
10	27.9	FVIEEGIKESLLGGRTINATLVGGAK		16	2671.4908	1	-1.0927			100	27.9	FVIEEGIKESLLGGRTINATLVGGAK		16	2671.4908	1	-1.0927		100	

MASCOT Mascot Search Results

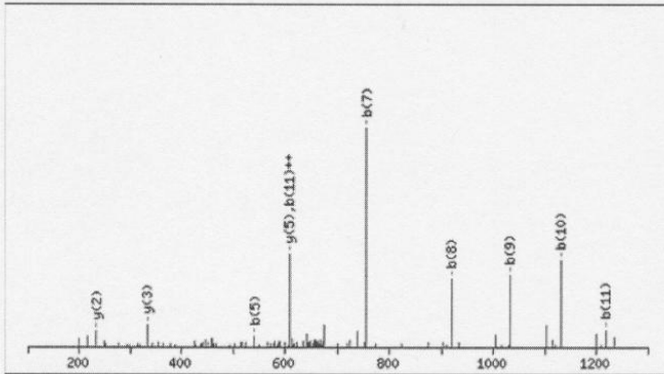
Peptide View

MS/MS Fragmentation of **RAQSVVDYLVSK**
 Found in gi|261339284, outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

Match to Query 105: 1362.753448 from(682.384000,2+) intensity(2643055.0000)
 Title: Cmpd 2, +MSn(683.26), 5.4 min
 Data file D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

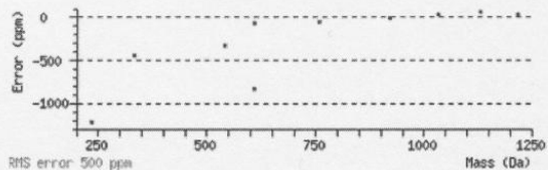
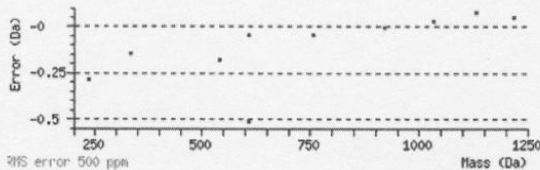
Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from 100 to 1300 Da Full range



Monoisotopic mass of neutral peptide Mr(calc): 1363.7460
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 51 Expect: 0.25
 Matches (Bold Red): 10/132 fragment ions using 11 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{*++}	b	b ⁺⁺	b [*]	b ^{*++}	Seq.	y	y ⁺⁺	y [*]	y ^{*++}	#
1	129.1135	65.0604	112.0869	56.5471	157.1084	79.0578	140.0818	70.5446	R					12
2	200.1506	100.5789	183.1240	92.0657	228.1455	114.5764	211.1190	106.0631	A	1208.6521	604.8297	1191.6256	596.3164	11
3	328.2092	164.6082	311.1826	156.0949	356.2041	178.6057	339.1775	170.0924	Q	1137.6150	569.3111	1120.5885	560.7979	10
4	415.2412	208.1242	398.2146	199.6110	443.2361	222.1217	426.2096	213.6084	S	1009.5564	505.2819	992.5299	496.7686	9
5	514.3096	257.6584	497.2831	249.1452	542.3045	271.6559	525.2780	263.1426	V	922.5244	461.7658	905.4979	453.2526	8
6	613.3780	307.1926	596.3515	298.6794	641.3729	321.1901	624.3464	312.6768	V	823.4560	412.2316	806.4294	403.7184	7
7	728.4050	364.7061	711.3784	356.1928	756.3999	378.7036	739.3733	370.1903	D	724.3876	362.6974	707.3610	354.1842	6
8	891.4683	446.2378	874.4417	437.7245	919.4632	460.2352	902.4367	451.7220	Y	609.3606	305.1840	592.3341	296.6707	5
9	1004.5524	502.7798	987.5258	494.2665	1032.5473	516.7773	1015.5207	508.2640	L	446.2973	223.6523	429.2708	215.1390	4
10	1103.6208	552.3140	1086.5942	543.8007	1131.6157	566.3115	1114.5891	557.7982	V	333.2132	167.1103	316.1867	158.5970	3
11	1190.6528	595.8300	1173.6262	587.3168	1218.6477	609.8275	1201.6212	601.3142	S	234.1448	117.5761	217.1183	109.0628	2
12									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of **RAQSVVDYLVSK**
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
50.6	1363.7460	-0.9925	RAQSVVDYLVSK
36.3	1363.7248	-0.9714	HFSKANAYLVSK

(MATRIX)
(SCIENCE) **Mascot Search Results**

Peptide View

MS/MS Fragmentation of **DGSVVVLGFTDR**

Found in **gi|261339284**, outer membrane protein A [*Enterobacter cancerogenus* ATCC 35316]

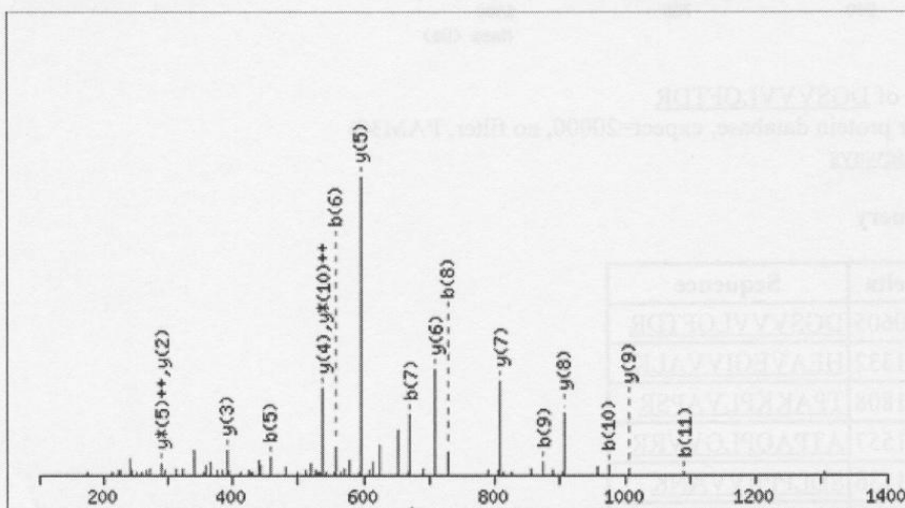
Match to Query 99: 1263.585448 from(632.800000,2+) intensity(34117664.0000)

Title: Cmpd 23, +MSn(633.16), 6.6 min

Data file D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from to Da



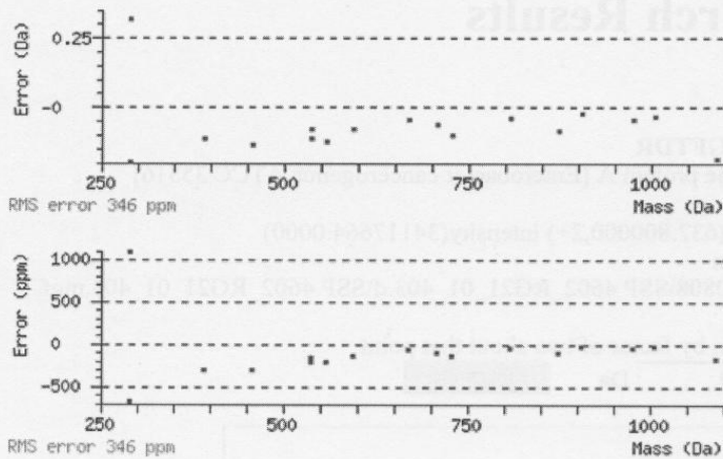
Monoisotopic mass of neutral peptide $M_r(\text{calc})$: 1263.6460

Fixed modifications: Carboxymethyl (C)

Ions Score: 65 Expect: 0.01

Matches (Bold Red): 17/88 fragment ions using 23 most intense peaks

#	a	a ⁺⁺	b	b ⁺⁺	Seq.	y	y ⁺⁺	y*	y ^{*++}	#
1	88.0393	44.5233	116.0342	58.5207	D					12
2	145.0608	73.0340	173.0557	87.0315	G	1149.6263	575.3168	1132.5997	566.8035	11
3	232.0928	116.5500	260.0877	130.5475	S	1092.6048	546.8060	1075.5782	538.2928	10
4	331.1612	166.0842	359.1561	180.0817	V	1005.5728	503.2900	988.5462	494.7767	9
5	430.2296	215.6184	458.2245	229.6159	V	906.5043	453.7558	889.4778	445.2425	8
6	529.2980	265.1527	557.2930	279.1501	V	807.4359	404.2216	790.4094	395.7083	7
7	642.3821	321.6947	670.3770	335.6921	L	708.3675	354.6874	691.3410	346.1741	6
8	699.4036	350.2054	727.3985	364.2029	G	595.2835	298.1454	578.2569	289.6321	5
9	846.4720	423.7396	874.4669	437.7371	F	538.2620	269.6346	521.2354	261.1214	4
10	947.5197	474.2635	975.5146	488.2609	T	391.1936	196.1004	374.1670	187.5872	3
11	1062.5466	531.7769	1090.5415	545.7744	D	290.1459	145.5766	273.1193	137.0633	2
12					R	175.1190	88.0631	158.0924	79.5498	1



NCBI **BLAST** search of DGSVVVLGFTDR
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

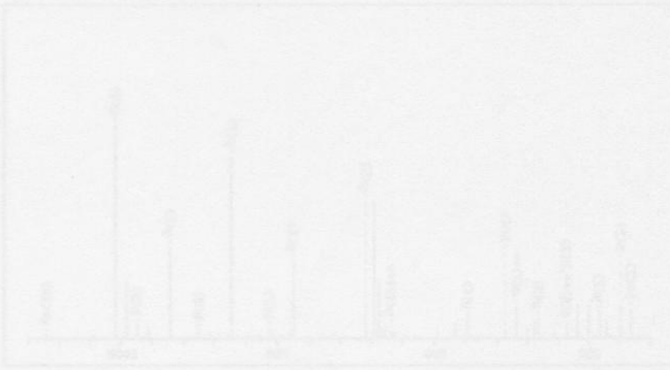
All matches to this query

Score	Mr(calc):	Delta	Sequence
64.5	1263.6460	-0.0605	<u>DGSVVVLGFTDR</u>
43.3	1263.7187	-0.1332	<u>HEAVEGIVVALK</u>
35.8	1263.7663	-0.1808	<u>TPAKKPLVAPSR</u>
34.0	1263.7412	-0.1557	<u>ATPAQPLGVVRR</u>
33.2	1264.7391	-1.1536	<u>SDLPPIVVANK</u>
31.6	1263.8166	-0.2311	<u>LAALLILVVPDK</u>
29.7	1263.8642	-0.2788	<u>VVGLLILVVIAR</u>
29.2	1263.8027	-0.2172	<u>KIPQKIPTLAR</u>
29.0	1262.6143	0.9712	<u>GDIVNGIGFDEK</u>
28.8	1263.7299	-0.1445	<u>GLAAPKGLPEGVRR</u>

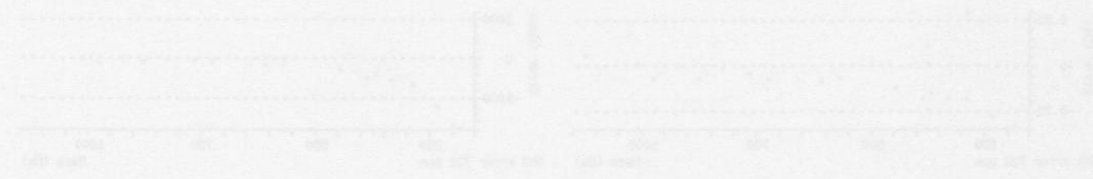
Mascot: <http://www.matrixscience.com/>

21.4	1208.6877	-0.9943	TVSVVEAVHIR
19.9	1208.6513	-0.9579	DODQHKVVIK
19.8	1207.6019	0.0915	ASQVVQFCLR
19.7	1206.5993	1.0942	REVVEQEYR
19.2	1206.6179	1.0755	REVMPPVYASR
19.1	1206.6608	1.0326	AQSVVNYLVSK
18.9	1208.5707	-0.8772	SISVVSNDMNK

Mascot: <http://www.matrixscience.com/>



Rank	Score	Protein	Accession	Length	Start	End	Score	Protein	Accession	Length	Start	End	Score
1	1208.6877	TVSVVEAVHIR	Q9VYD1	100	1	100	1208.6877	TVSVVEAVHIR	Q9VYD1	100	1	100	1208.6877
2	1208.6513	DODQHKVVIK	Q9VYD1	100	1	100	1208.6513	DODQHKVVIK	Q9VYD1	100	1	100	1208.6513
3	1207.6019	ASQVVQFCLR	Q9VYD1	100	1	100	1207.6019	ASQVVQFCLR	Q9VYD1	100	1	100	1207.6019
4	1206.5993	REVVEQEYR	Q9VYD1	100	1	100	1206.5993	REVVEQEYR	Q9VYD1	100	1	100	1206.5993
5	1206.6179	REVMPPVYASR	Q9VYD1	100	1	100	1206.6179	REVMPPVYASR	Q9VYD1	100	1	100	1206.6179
6	1206.6608	AQSVVNYLVSK	Q9VYD1	100	1	100	1206.6608	AQSVVNYLVSK	Q9VYD1	100	1	100	1206.6608
7	1208.5707	SISVVSNDMNK	Q9VYD1	100	1	100	1208.5707	SISVVSNDMNK	Q9VYD1	100	1	100	1208.5707



Rank	Score	Protein	Accession	Length	Start	End	Score
1	1208.6877	TVSVVEAVHIR	Q9VYD1	100	1	100	1208.6877
2	1208.6513	DODQHKVVIK	Q9VYD1	100	1	100	1208.6513
3	1207.6019	ASQVVQFCLR	Q9VYD1	100	1	100	1207.6019
4	1206.5993	REVVEQEYR	Q9VYD1	100	1	100	1206.5993
5	1206.6179	REVMPPVYASR	Q9VYD1	100	1	100	1206.6179
6	1206.6608	AQSVVNYLVSK	Q9VYD1	100	1	100	1206.6608
7	1208.5707	SISVVSNDMNK	Q9VYD1	100	1	100	1208.5707

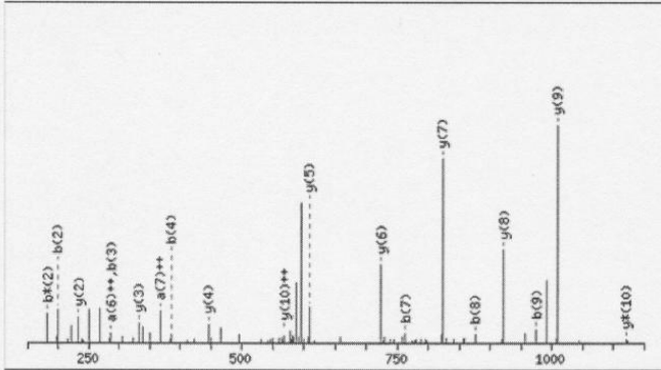
MASCOT Mascot Search Results

Peptide View

MS/MS Fragmentation of AQSVDYLVS**K**
 Found in gi|261339284, outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

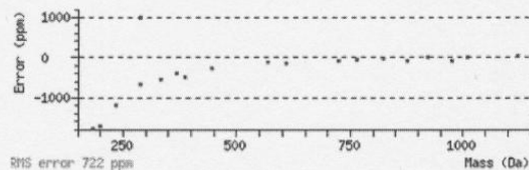
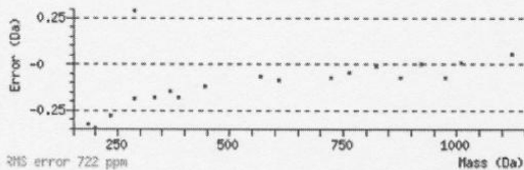
Match to Query 94: 1207.693448 from(604.854000,2+) intensity(17956968.0000)
 Title: Cmpd 7, +MSn(605.23), 5.7 min
 Data file D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

Click mouse within plot area to zoom in by factor of two about that point
 Or, Plot from 150 to 1150 Da Full range



Monoisotopic mass of neutral peptide Mr(calc): 1207.6449
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 44 Expect: 1.1
 Matches (Bold Red): 19/116 Fragment ions using 47 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{*++}	b	b ⁺⁺	b [*]	b ^{*++}	Seq.	y	y ⁺⁺	y [*]	y ^{*++}	#
1	44.0495	22.5284			72.0444	36.5258			A					11
2	172.1081	86.5577	155.0815	78.0444	200.1030	100.5551	183.0764	92.0418	Q	1137.6150	569.3111	1120.5885	560.7979	10
3	259.1401	130.0737	242.1135	121.5604	287.1350	144.0711	270.1084	135.5579	S	1009.5564	505.2819	992.5299	496.7686	9
4	358.2085	179.6079	341.1819	171.0946	386.2034	193.6053	369.1769	185.0921	V	922.5244	461.7658	905.4979	453.2526	8
5	457.2769	229.1421	440.2504	220.6288	485.2718	243.1396	468.2453	234.6263	V	823.4560	412.2316	806.4294	403.7184	7
6	572.3039	286.6556	555.2773	278.1423	600.2988	300.6530	583.2722	292.1397	D	724.3876	362.6974	707.3610	354.1842	6
7	735.3672	368.1872	718.3406	359.6740	763.3621	382.1847	746.3355	373.6714	Y	609.3606	305.1840	592.3341	296.6707	5
8	848.4512	424.7293	831.4247	416.2160	876.4462	438.7267	859.4196	430.2134	L	446.2973	223.6523	429.2708	215.1390	4
9	947.5197	474.2635	930.4931	465.7502	975.5146	488.2609	958.4880	479.7477	V	333.2132	167.1103	316.1867	158.5970	3
10	1034.5517	517.7795	1017.5251	509.2662	1062.5466	531.7769	1045.5201	523.2637	S	234.1448	117.5761	217.1183	109.0628	2
11									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of AQSVDYLVS**K**
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
44.0	1207.6449	0.0486	<u>AQSVDYLVSK</u>
29.4	1206.7085	0.9850	<u>VVSVVQAGVPPR</u>
24.0	1207.6673	0.0261	<u>GHGQVIGNLVSK</u>

MASCOT Mascot Search Results

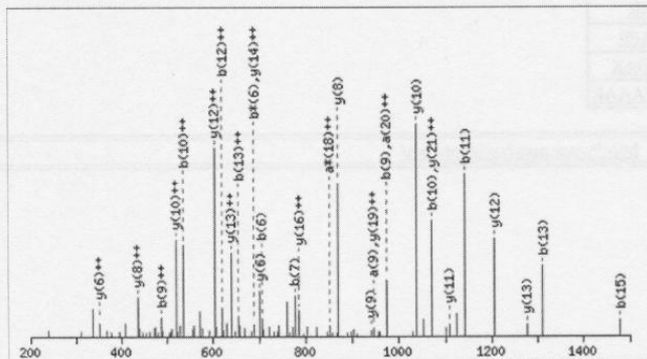
Peptide View

MS/MS Fragmentation of FGQQEDAPVVAPAPAPEVQTK
 Found in gi|261339284, outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

Match to Query 228: 2346.257172 from(783.093000,3+) intensity(68730400.0000)
 Title: Cmpd 39, +MSn(783.69), 7.3 min
 Data file D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

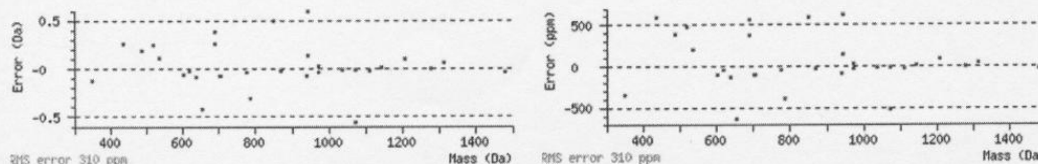
Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 2346.1856
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 57 Expect: 0.046
 Matches (Bold Red): 31/256 fragment ions using 40 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{*++}	b	b ⁺⁺	b [*]	b ^{*++}	Seq.	y	y ⁺⁺	y [*]	y ^{*++}	#
1	120.0808	60.5440			148.0757	74.5415			F					23
2	177.1022	89.0548			205.0972	103.0522			G	2200.1244	1100.5659	2183.0979	1092.0526	22
3	305.1608	153.0840	288.1343	144.5708	333.1557	167.0815	316.1292	158.5682	Q	2143.1030	1072.0551	2126.0764	1063.5419	21
4	433.2194	217.1133	416.1928	208.6001	461.2143	231.1108	444.1878	222.5975	Q	2015.0444	1008.0258	1998.0179	999.5126	20
5	562.2620	281.6346	545.2354	273.1214	590.2569	295.6321	573.2304	287.1188	E	1886.9858	943.9965	1869.9593	935.4833	19
6	677.2889	339.1481	660.2624	330.6348	705.2838	353.1456	688.2573	344.6323	D	1757.9432	879.4753	1740.9167	870.9620	18
7	748.3260	374.6667	731.2995	366.1534	776.3210	388.6641	759.2944	380.1508	A	1642.9163	821.9618	1625.8897	813.4485	17
8	845.3788	423.1930	828.3523	414.6798	873.3737	437.1905	856.3472	428.6772	P	1571.8792	786.4432	1554.8526	777.9299	16
9	944.4472	472.7272	927.4207	464.2140	972.4421	486.7247	955.4156	478.2114	V	1474.8264	737.9168	1457.7999	729.4036	15
10	1043.5156	522.2615	1026.4891	513.7482	1071.5106	536.2589	1054.4840	527.7456	V	1375.7580	688.3826	1358.7314	679.8694	14
11	1114.5528	557.7800	1097.5262	549.2667	1142.5477	571.7775	1125.5211	563.2642	A	1276.6896	638.8484	1259.6630	630.3352	13
12	1211.6055	606.3064	1194.5790	597.7931	1239.6004	620.3039	1222.5739	611.7906	P	1205.6525	603.3299	1188.6259	594.8166	12
13	1282.6426	641.8250	1265.6161	633.3117	1310.6375	655.8224	1293.6110	647.3091	A	1108.5997	554.8035	1091.5732	546.2902	11
14	1379.6954	690.3513	1362.6688	681.8381	1407.6903	704.3488	1390.6638	695.8355	P	1037.5626	519.2849	1020.5360	510.7717	10
15	1450.7325	725.8699	1433.7060	717.3566	1478.7274	739.8673	1461.7009	731.3541	A	940.5098	470.7586	923.4833	462.2453	9
16	1547.7853	774.3963	1530.7587	765.8830	1575.7802	788.3937	1558.7536	779.8805	P	869.4727	435.2400	852.4462	426.7267	8
17	1618.8224	809.9148	1601.7958	801.4016	1646.8173	823.9123	1629.7908	815.3990	A	772.4199	386.7136	755.3934	378.2003	7
18	1715.8751	858.4412	1698.8486	849.9279	1743.8701	872.4387	1726.8435	863.9254	P	701.3828	351.1951	684.3563	342.6818	6
19	1844.9177	922.9625	1827.8912	914.4492	1872.9127	936.9600	1855.8861	928.4467	E	604.3301	302.6687	587.3035	294.1554	5
20	1943.9862	972.4967	1926.9596	963.9834	1971.9811	986.4942	1954.9545	977.9809	V	475.2875	238.1474	458.2609	229.6341	4
21	2072.0447	1036.5260	2055.0182	1028.0127	2100.0396	1050.5235	2083.0131	1042.0102	Q	376.2191	188.6132	359.1925	180.0999	3
22	2173.0924	1087.0498	2156.0659	1078.5366	2201.0873	1101.0473	2184.0608	1092.5340	T	248.1605	124.5839	231.1339	116.0706	2
23									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of FGQOEDAPVVAPAPAPAPEVOTK
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
56.9	2346.1856	0.0716	<u>FGQOEDAPVVAPAPAPAPEVOTK</u>
35.8	2346.1855	0.0716	<u>FGQOEEAAPIAPAPAPAPEVOTK</u>
23.3	2346.1420	0.1152	<u>MDEDERMARAALSGLLEANLVR</u>
22.7	2346.2042	0.0530	<u>VLMLVSGGLGOALWYSHDEKK</u>
21.8	2345.2267	1.0304	<u>FGODDAVAPVVAPAPAPVETK</u>
20.1	2346.2154	0.0417	<u>QVYGTVKGKLEMAWDPVAOEAR</u>
19.8	2345.1838	1.0734	<u>EQGIDTVWFGAAAPLGLMAGVAR</u>
19.7	2345.3008	0.9564	<u>FNAPKAAPAPAPVVAAPAPAPAPR</u>
19.5	2347.2318	-0.9746	<u>VAIISGAAOQMGGAATARLFAAEGAK</u>
18.3	2346.2880	-0.0308	<u>SARADLLGPVGAPASGRAAAAIVAAAR</u>

Mascot: <http://www.matrixscience.com/>



Mascot Search Results

Peptide View

MS/MS Fragmentation of SDVLFNFNK

Found in gi|261339284, outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

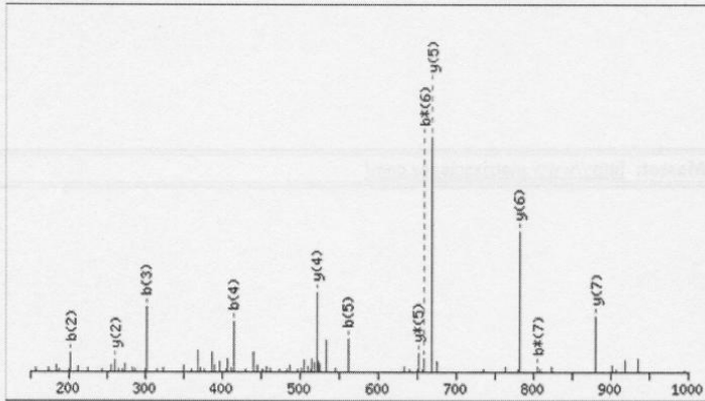
Match to Query 84: 1082.449448 from(542.232000,2+) intensity(14554407.0000)

Title: Cmpd 14, +MSn(543.11), 6.2 min

Data file D:\Data\Lakshmy\120808\260808\SSP 4602_RG21_01_403.d\SSP 4602_RG21_01_403.mgf

Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from to Da



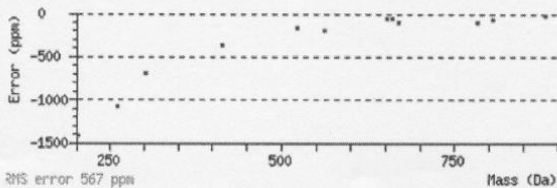
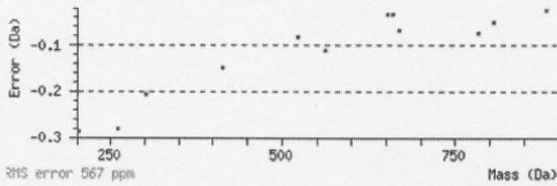
Monoisotopic mass of neutral peptide Mr(calc): 1082.5397

Fixed modifications: Carboxymethyl (C)

Ions Score: 39 Expect: 3.3

Matches (Bold Red): 12/76 fragment ions using 17 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{*++}	b	b ⁺⁺	b [*]	b ^{*++}	Seq.	y	y ⁺⁺	y [*]	y ^{*++}	#
1	60.0444	30.5258			88.0393	44.5233			S					9
2	175.0713	88.0393			203.0662	102.0368			D	996.5149	498.7611	979.4884	490.2478	8
3	274.1397	137.5735			302.1347	151.5710			V	881.4880	441.2476	864.4614	432.7343	7
4	387.2238	194.1155			415.2187	208.1130			L	782.4196	391.7134	765.3930	383.2001	6
5	534.2922	267.6498			562.2871	281.6472			F	669.3355	335.1714	652.3089	326.6581	5
6	648.3352	324.6712	631.3086	316.1579	676.3301	338.6687	659.3035	330.1554	N	522.2671	261.6372	505.2405	253.1239	4
7	795.4036	398.2054	778.3770	389.6921	823.3985	412.2029	806.3719	403.6896	F	408.2241	204.6157	391.1976	196.1024	3
8	909.4465	455.2269	892.4199	446.7136	937.4414	469.2243	920.4149	460.7111	N	261.1557	131.0815	244.1292	122.5682	2
9									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of SDVLFNFNK
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
38.9	1082.5397	-0.0902	<u>SDVLFNFNK</u>
37.4	1082.5318	-0.0824	<u>SDVIFPMIK</u>
37.4	1082.5318	-0.0824	<u>SDVIFLMDK</u>
32.4	1082.5396	-0.0902	<u>AEELFGYVR</u>
30.3	1082.6448	-0.1953	<u>LVALODGLVR</u>
29.7	1082.5397	-0.0902	<u>DEGLFTFVR</u>
24.5	1082.6059	-0.1565	<u>MFRVFIVR</u>
24.5	1082.5253	-0.0758	<u>MFTPGMIVR</u>
23.8	1082.5760	-0.1266	<u>FLSSDFIVR</u>
22.9	1082.5720	-0.1226	<u>ADPLDGQIVR</u>

Mascot: <http://www.matrixscience.com/>



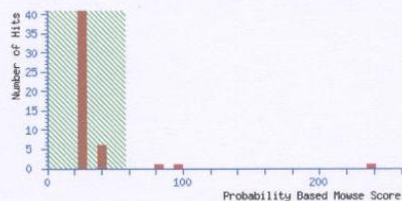
Protein	Score	Mr(calc)	Delta	Sequence
SDVLFNFNK	38.9	1082.5397	-0.0902	SDVLFNFNK
SDVIFPMIK	37.4	1082.5318	-0.0824	SDVIFPMIK
SDVIFLMDK	37.4	1082.5318	-0.0824	SDVIFLMDK
AEELFGYVR	32.4	1082.5396	-0.0902	AEELFGYVR
LVALODGLVR	30.3	1082.6448	-0.1953	LVALODGLVR
DEGLFTFVR	29.7	1082.5397	-0.0902	DEGLFTFVR
MFRVFIVR	24.5	1082.6059	-0.1565	MFRVFIVR
MFTPGMIVR	24.5	1082.5253	-0.0758	MFTPGMIVR
FLSSDFIVR	23.8	1082.5760	-0.1266	FLSSDFIVR
ADPLDGQIVR	22.9	1082.5720	-0.1226	ADPLDGQIVR

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 4606
MS data file : D:\Data\Lakshmy\120808\241108\SSP 4606_RB21_01_773.d\SSP 4606_RB21_01_773.mgf
Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 11:20:40 GMT
Protein hits : [gi|50582463](#) oxidoreductase [Exiguobacterium acetylicum]
[gi|146275769](#) short-chain dehydrogenase/reductase SDR [Novosphingobium aromaticivorans DSM 12444]
[gi|148368](#) outer membrane protein II [Enterobacter aerogenes]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|50582463](#) Mass: 27360 Score: 238 Queries matched: 5 emPAI: 0.41
 oxidoreductase [Exiguobacterium acetylicum]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 125	497.5760	1489.7062	1489.7413	-0.0351	0	53	0.15	1	K.VLAPATETEFADR.S
<input checked="" type="checkbox"/> 155	573.2590	1716.7552	1716.8570	-0.1019	0	(25)	87	1	K.YTVITGASSGIGYETAK.L
<input checked="" type="checkbox"/> 156	859.3900	1716.7654	1716.8570	-0.0916	0	91	2.3e-05	1	K.YTVITGASSGIGYETAK.L
<input checked="" type="checkbox"/> 197	676.9390	2027.7952	2028.9752	-1.1801	0	(37)	3.6	1	K.SVDLADNQNVDLYEGLK.E
<input checked="" type="checkbox"/> 198	1014.9250	2027.8354	2028.9752	-1.1398	0	94	7.5e-06	1	K.SVDLADNQNVDLYEGLK.E

2. [gi|146275769](#) Mass: 27018 Score: 97 Queries matched: 2 emPAI: 0.26
 short-chain dehydrogenase/reductase SDR [Novosphingobium aromaticivorans DSM 12444]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 107	455.1670	1362.4792	1362.6204	-0.1413	0	52	0.14	1	K.FEDTFLSDFHR.V
<input checked="" type="checkbox"/> 211	700.5920	2098.7542	2099.0027	-0.2486	0	44	0.63	1	R.VNSVHPGGIDTPMLGSIMDK.Y + 2 Oxidation (M)

3. [gi|148368](#) Mass: 25654 Score: 84 Queries matched: 2 emPAI: 0.13
 outer membrane protein II [Enterobacter aerogenes]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 141	813.8050	1625.5954	1625.7937	-0.1983	0	48	0.32	1	K.LGYPVTDLDVYTR.L
<input checked="" type="checkbox"/> 228	783.0500	2346.1282	2346.1856	-0.0574	0	36	5.2	1	R.FGQQEDAPVAPAPAPAEVQTK.H

Proteins matching the same set of peptides:

[gi|261339284](#) Mass: 37561 Score: 84 Queries matched: 2
 outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 101	651.8070	1301.5994	1302.5914	-0.9920	0	40	2.9	1	GYCSLLAAR
<input checked="" type="checkbox"/> 108	692.3100	1382.6054	1382.8245	-0.2191	2	39	3.1	1	LSIALGKGRNPAK
<input checked="" type="checkbox"/> 120	738.3480	1474.6814	1475.8209	-1.1394	1	36	7.1	1	TRTFVQQLINR
<input checked="" type="checkbox"/> 229	789.7300	2366.1682	2366.3533	-0.1851	2	34	9.9	1	GDNVGRKTLKILAGLLQPSGK

Mascot Search Results

Protein View

Match to: **gi|261339284** Score: **84**
outer membrane protein A [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\241108\SSP 4606_RB21_01_773.d\SSP 4606_RB21_01_773.mgf

Nominal mass (M_r): **37561**; Calculated pI value: **5.19**
 NCBI BLAST search of **gi|261339284** against nr
 Unformatted **sequence_string** for pasting into other applications

Taxonomy: **Enterobacter cancerogenus ATCC 35316**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **10%**

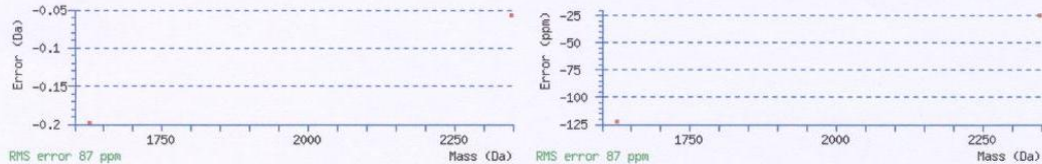
Matched peptides shown in **Bold Red**

1 MKKTAIAIAV ALAGFATVAQ AAPKDNWYA GKKLGWSQFH DTGWYNSSLN
 51 NDGPETHESQL GAGAFGGYQV NPYVGFEMGY DWLGRMPYKG DNVNGAFKAQ
101 GVQLTAKLGY FVTDDLVDVYR RLGGMVWRAD SSNSIAGDDH DTGVSVPFAG
151 GVENAMTRDI ATRLEYQWVN NIGDGATVGV RPDNGMLSVG VSYRFGQED
201 AEPVAPAPAP AEPVQTRHFT LKSDVLFNFN KATLKPEGQQ ALDQLYTQLS
251 NLDPKDGSVV VLGFTDRIGS DAYNQLSEK RAQSVVDYLV SKGPIANKIS
301 PRGMGESNPV TGNCTDNVKP RAALIDCLAP DRRVEIEVKG IKDVTQPAA
351

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
108 - 121	813.8050	1625.5954	1625.7937	-0.1983	0	K.LGYFVTDDLVDVYR.L (Ions score 48)
195 - 217	783.0500	2346.1282	2346.1856	-0.0574	0	R.FGQEDAPVVAPAPAPAPEVQTR.H (Ions score 36)



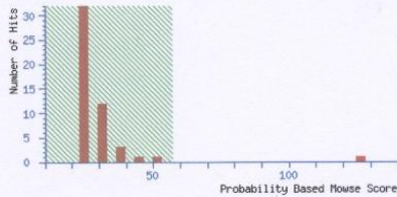
LOCUS ZP_05967142 350 aa linear BCT 15-OCT-2009
 DEFINITION outer membrane protein A [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05967142
 VERSION ZP_05967142.1 GI:261339284
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000004.1](#)
 KEYWORDS
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM **Enterobacter cancerogenus ATCC 35316**
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 350)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 350)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM02000004](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 5003
MS data file : D:\Data\Lakshmy\120808\280808\SSP 5003_RJ6_01_513.d\SSP 5003_RJ6_01_513.mgf
Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 11:22:03 GMT
Protein hits : [gi|261342318](#) hypothetical protein EcanA3_17757 [Enterobacter cancerogenus ATCC 35316]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event.
 Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$).
 Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)
 Significance threshold $p <$ Max. number of hits
 Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets
 Show pop-ups Suppress pop-ups Sort unassigned Require bold red
 Error tolerant

1. [gi|261342318](#) Mass: 26485 Score: 126 Queries matched: 2 **emPAI**: 0.27
 hypothetical protein EcanA3_17757 [Enterobacter cancerogenus ATCC 35316]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
<input checked="" type="checkbox"/> 112	556.2900	1665.8482	1665.8726	-0.0244	0 36	7.1 1 R.VAQYLSFLEQNGVAK.K
<input checked="" type="checkbox"/> 166	628.6650	1882.9732	1882.9537	0.0195	0 90	2.4e-05 1 K.AAIDDAIHQAQLASGFK.G

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
<input checked="" type="checkbox"/> 67	679.3770	1356.7394	1356.7976	-0.0582	2 48	0.45 1 LNDIDSILKAKK
<input checked="" type="checkbox"/> 233	749.4760	2245.4062	2245.1512	0.2550	0 42	1.1 1 TLVLA MSPDPLNTVDQVVAK + Oxidation (M)
<input checked="" type="checkbox"/> 61	516.3400	1030.6654	1030.5546	0.1108	0 38	5.5 1 VLDELDSLK
<input checked="" type="checkbox"/> 76	736.1480	1470.2814	1469.6126	0.6688	0 31	29 1 MGESGDSGAMLLDR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 69	692.3760	1382.7374	1382.6388	0.0987	0 28	48 1 EGDLA AAFSMIDK + Oxidation (M)
<input checked="" type="checkbox"/> 232	747.0540	2238.1402	2237.0569	1.0833	1 28	40 1 MRMAAEFGAAVTFGGVDDVVR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 226	738.0380	2211.0922	2210.0968	0.9954	1 28	42 1 KYVTTVAQDATGNQFANSPK
<input checked="" type="checkbox"/> 199	1011.1660	3030.4762	3029.5433	0.9329	1 28	63 1 GLVEKGWQALPDHAA PTTSTMVFLVR + Oxidation (M)
<input checked="" type="checkbox"/> 64	633.3340	1264.6534	1265.5267	-0.8733	0 27	50 1 QEMDMENEIK
<input checked="" type="checkbox"/> 229	742.4370	2224.2892	2223.1456	1.1435	1 27	39 1 YLKTMGALIEGEETGITIR
<input checked="" type="checkbox"/> 129	876.6700	2626.9882	2627.3894	-0.4012	2 27	62 1 MYRVVVSQFNDGIRTPLAQIYIK + Oxidation (M)
<input checked="" type="checkbox"/> 190	987.5410	2959.6012	2960.5650	-0.9638	1 27	86 1 VAARLFGGVVTVAVVPAVMCVLTVVACR
<input checked="" type="checkbox"/> 244	892.6220	2674.8442	2675.2757	-0.4315	2 27	29 1 MYGIRGTAKASDDGIYPMMAQNGIK + Oxidation (M)
<input checked="" type="checkbox"/> 188	982.2840	2943.8302	2943.4470	0.3832	1 27	63 1 GGVHFLLEVDTPAALQRMEVASARMK + Oxidation (M)
<input checked="" type="checkbox"/> 98	805.0860	2412.2362	2411.0984	1.1377	1 27	90 1 DLTLLSDYERHEMIEEQMK + 2 Oxidation (M)
<input checked="" type="checkbox"/> 155	914.2040	2739.5902	2740.4653	-0.8751	2 26	91 1 AALTLAQIEDLNKRLANAAAAAQR + Oxidation (M)
<input checked="" type="checkbox"/> 118	853.0970	2556.2692	2557.2119	-0.9427	0 25	1.2e+02 1 SGLVDVGFVAVCEEIGVPLSNHR
<input checked="" type="checkbox"/> 72	701.8480	1401.6814	1402.6690	-0.9875	0 25	1.5e+02 1 LEELSMYDLYK
<input checked="" type="checkbox"/> 235	757.8950	2270.6632	2271.1318	-0.4686	0 25	53 1 VLAAPTAIAGFDNGHVMETETTR
<input checked="" type="checkbox"/> 228	1111.6590	3331.9552	3331.5602	0.3950	2 25	1.1e+02 1 SYFFVLDLQGHSDCDDVVMREGLEKIR + Oxidation (M)
<input checked="" type="checkbox"/> 196	1006.5340	2011.0534	2009.9782	1.0753	2 25	1.5e+02 1 GRTGLYFCGRGQSFRLR
<input checked="" type="checkbox"/> 187	979.3990	1956.7834	1957.0190	-0.2355	0 24	79 1 ALVLQVLDIIDNEAMR + Oxidation (M)
<input checked="" type="checkbox"/> 130	876.9690	2627.8852	2627.3317	0.5535	2 24	1.6e+02 1 HGDVNTFLKTAHVSLYARGASR
<input checked="" type="checkbox"/> 214	695.0820	2082.2242	2082.9536	-0.7295	0 24	92 1 MNLYGNAGSAGAVMIMLEK + 3 Oxidation (M)
<input checked="" type="checkbox"/> 206	1029.2490	2056.4834	2055.9944	0.4891	0 24	1.4e+02 1 MFMGLLWMLAASVQAAER + 2 Oxidation (M)
<input checked="" type="checkbox"/> 75	734.9050	2201.6932	2201.1362	0.5570	1 24	1.9e+02 1 DTALAIEGMLKATGPEVVSER + Oxidation (M)
<input checked="" type="checkbox"/> 74	731.2890	2190.8452	2191.1307	-0.2855	1 24	1.9e+02 1 SVVFGQDPAIDALASAIKMSR + Oxidation (M)
<input checked="" type="checkbox"/> 126	867.5370	2599.5892	2599.3581	0.2311	2 23	1.8e+02 1 SRIGVCKVDITPPGIDFVGYHR
<input checked="" type="checkbox"/> 200	1012.5140	3034.5202	3033.3538	1.1664	0 23	2.1e+02 1 DLHALMTDSQDWPADFHYGGIMVR + Oxidation (M)

<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project (<http://nihroadmap.nih.gov/hmp/>).

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

FEATURES	Location/Qualifiers
source	1..350 /organism="Enterobacter cancerogenus ATCC 35316" /strain="ATCC 35316" /db_xref="taxon:500639"
Protein	1..350 /product="outer membrane protein A" /calculated_mol_wt=37337
Region	1..346 /region_name="PRK10808" /note="outer membrane protein A; Reviewed; PRK10808" /db_xref="CDD:138188"
Region	23..196 /region_name="Surface Ag 2" /note="Surface antigen; cl01155" /db_xref="CDD:141128"
Region	224..338 /region_name="OmpA_C-like" /note="Peptidoglycan binding domains similar to the C-terminal domain of outer-membrane protein OmpA; cd07185" /db_xref="CDD:143586"
Site	order(229..230,265..266,269,273..274,277,328,332) /site_type="other" /note="ligand binding site" /db_xref="CDD:143586"
CDS	1..350 /locus_tag="EcanA3_020100002425" /coded_by="complement(NZ_ABWM02000004.1:545400..546452)" /note="COG2885 Outer membrane protein and related peptidoglycan-associated (lipo)proteins" /transl_table=11 /db_xref="CDD:138188"

Mascot: <http://www.matrixscience.com/>

Mascot Search Results

Protein View

Match to: **gi|261342318** Score: **126**
hypothetical protein EcanA3_17757 [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\280808\SSP 5003_RJ6_01_513.d\SSP 5003_RJ6_01_513.mgf

Nominal mass (M_r): **26485**; Calculated pI value: **5.95**
 NCBI BLAST search of **gi|261342318** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **13%**

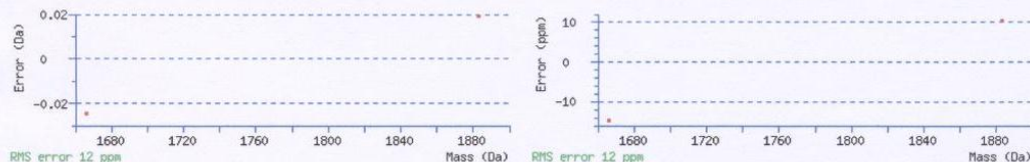
Matched peptides shown in **Bold Red**

1 MKNFVIALAA LVSLGAVSAQ ANELPAGPHI VTSGTASVDA VPDVATLAI E
51 VNVAAKDAAS AKQADDRVA **QYLSFLEQNG VAK**DINSAN LRTQPDYDQ
101 NGKSILKGYR AVRTVEVTLR QLDKLNLLD GALKAGLNEI RSVSLGVAQP
151 EKYKDEARKA **AIDDAIHQAQ QLASGFK**GKL GPVYSVRYHV SNYQSPMVR
201 MMKADAAVPS AQETYEQPTI QFDDQVVVF QLEPTQTQT EAAKAQ

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
69 - 83	556.2900	1665.8482	1665.8726	-0.0244	0	R.VAQLSFLAQNGVAK.K (Ions score 36)
160 - 177	628.6650	1882.9732	1882.9537	0.0195	0	K.AAIDDAIHQAQQLASGFK.G (Ions score 90)



LOCUS ZP_05970176 246 aa linear BCT 15-OCT-2009
 DEFINITION hypothetical protein EcanA3_17757 [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05970176
 VERSION ZP_05970176.1 GI:261342318
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000033.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 246)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B., Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C., Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C., Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M., Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 REFERENCE 2 (residues 1 to 246)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H., Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from [ABWM02000033](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no

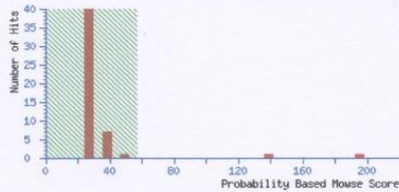
http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100105/FtmmSGSa... 21/01/2010

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 5102
 MS data file : D:\Data\Lakshmy\120808\260808\SSP 5102_RG17_01_398.d\SSP 5102_RG17_01_398.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:03:53 GMT
 Protein hits : [gi|152972368](#) universal stress protein A [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
 [gi|50119012](#) universal stress protein A [Pectobacterium atrosepticum SCRI1043]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: **Peptide Summary** [Help](#)

Significance threshold $p < 0.05$ Max. number of hits **AUTO**

Standard scoring MudPIT scoring Ions score or expect cut-off **0** Show sub-sets **0**

Show pop-ups Suppress pop-ups Sort unassigned **Decreasing Score** Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|152972368](#) Mass: 16229 Score: 195 Queries matched: 7 emPAI: 0.46
 universal stress protein A [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 96	711.3850	1420.7554	1420.7926	-0.0371	0	(93)	1.6e-05	1	K.HILIAVDLSPESK.V
<input checked="" type="checkbox"/> 97	711.4060	1420.7974	1420.7926	0.0049	0	(82)	0.00018	1	K.HILIAVDLSPESK.V
<input checked="" type="checkbox"/> 98	711.4080	1420.8014	1420.7926	0.0089	0	94	1.2e-05	1	K.HILIAVDLSPESK.V
<input type="checkbox"/> 212	718.3550	2152.0432	2152.8983	-0.8551	0	25		86	K.YDMDLVVCGHHQDFWSK.L + Oxidation (M)
<input checked="" type="checkbox"/> 111	761.3180	2280.9322	2280.9932	-0.0611	1	20	4.2e+02	1	K.KYDMDLVVCGHHQDFWSK.L + Oxidation (M)
<input checked="" type="checkbox"/> 237	826.7860	2477.3362	2477.2472	0.0890	1	56	0.051	1	R.QLINTVHVDMLIVPLRDEEDE.-
<input checked="" type="checkbox"/> 240	832.0960	2493.2662	2493.2421	0.0241	1	(51)	0.19	1	R.QLINTVHVDMLIVPLRDEEDE.- + Oxidation (M)

Proteins matching the same set of peptides:
[gi|261341988](#) Mass: 16201 Score: 195 Queries matched: 7
 universal stress protein A [Enterobacter cancerogenus ATCC 35316]

2. [gi|50119012](#) Mass: 16177 Score: 134 Queries matched: 5 emPAI: 0.21
 universal stress protein A [Pectobacterium atrosepticum SCRI1043]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 96	711.3850	1420.7554	1420.7926	-0.0371	0	(93)	1.6e-05	1	K.HILIAVDLSPESK.V
<input type="checkbox"/> 97	711.4060	1420.7974	1420.7926	0.0049	0	(82)	0.00018	1	K.HILIAVDLSPESK.V
<input type="checkbox"/> 98	711.4080	1420.8014	1420.7926	0.0089	0	94	1.2e-05	1	K.HILIAVDLSPESK.V
<input type="checkbox"/> 237	826.7860	2477.3362	2476.2632	1.0730	1	(34)	8.2	2	R.QLINTVHVDMLIVPLRDEEDE.-
<input type="checkbox"/> 240	832.0960	2493.2662	2492.2581	1.0081	1	40	2.1	2	R.QLINTVHVDMLIVPLRDEEDE.- + Oxidation (M)

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 128	836.9540	1671.8934	1671.9858	-0.0924	1	37	5.7	1	MKVVSIVGARPFQIK
<input checked="" type="checkbox"/> 230	767.0360	2298.0862	2299.1961	-1.1099	1	36	5.7	1	YLITAAFTDRGGGFNNQTILK
<input checked="" type="checkbox"/> 239	831.7950	2492.3632	2492.3097	0.0535	2	35	7.6	1	YPFLLVDRILEMDRENQTIK
<input checked="" type="checkbox"/> 249	983.8360	2948.4862	2949.4001	-0.9139	0	34	7.6	1	NSIIYYVGCPCPKPGVVGSCGPTTSGR
<input checked="" type="checkbox"/> 87	651.8450	1301.6754	1301.6914	-0.0160	1	34	13	1	RAFMHTLAEVK
<input checked="" type="checkbox"/> 238	826.8270	2477.4592	2476.3471	1.1120	2	33	8.9	1	MLLAEKNVSVLNTAIVERAARY + Oxidation (M)
<input checked="" type="checkbox"/> 233	802.1120	2403.3142	2403.3413	-0.0272	1	32	13	1	KGYVLTGVPSLSPTQIEFLK
<input checked="" type="checkbox"/> 218	737.7020	2210.0842	2211.1787	-1.0945	1	31	17	1	VNVTYEVLDSNVTFKVSVAK
<input checked="" type="checkbox"/> 221	742.3240	2223.9502	2223.0841	0.8660	2	30	20	1	SAGVDDIQMEIDEKFINRK + Oxidation (M)
<input checked="" type="checkbox"/> 200	702.0780	2103.2122	2103.0406	0.1716	0	29	26	1	ADVVLMLVIDATEGVTEQDAK
<input checked="" type="checkbox"/> 126	831.3840	2491.1302	2490.2875	0.8426	2	29	60	1	MKQFQCAIFDMLLGAAVTNWR + Oxidation (M)
<input checked="" type="checkbox"/> 214	1091.3100	3270.9082	3271.6078	-0.6996	2	28	44	1	GGHCLRAADVGEATVRQLYMPDTAVLVTR

manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

FEATURES	Location/Qualifiers
source	1..246 /organism="Enterobacter cancerogenus ATCC 35316" /strain="ATCC 35316" /db_xref="taxon:500639"
Protein	1..246 /product="hypothetical protein" /calculated_mol_wt=26370
Region	22..234 /region_name="DUF541" /note="Protein of unknown function (DUF541); c101077" /db_xref="CDD:141087"
CDS	1..246 /locus_tag="EcanA3_020100017757" /coded_by="complement(NZ_ABWM02000033.1:268250..268990)" /note="COG2968 Uncharacterized conserved protein" /transl_table=11 /db_xref="CDD:138394"

Mascot: <http://www.matrixscience.com/>

Mascot Search Results

Protein View

Match to: **gi|261341988** Score: 195
universal stress protein A [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\260808\SSP 5102_RG17_01_398.d\SSP 5102_RG17_01_398.mgf

Nominal mass (M_r): **16201**; Calculated pI value: **4.98**
 NCBI BLAST search of **gi|261341988** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **35%**

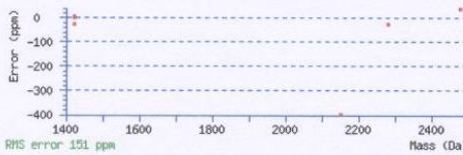
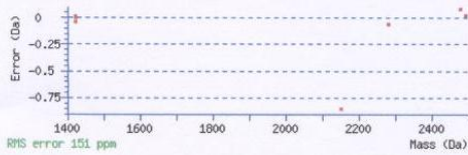
Matched peptides shown in **Bold Red**

- 1 **MAYKHILIAV DLSPE**SKVLV DKAVSMARPY NAKVSLIHVD VNYS DLYTGL
- 51 IDVNLGDMQK RISEETHAL SELSTNAGYP ITETLGGSGD LGQVLVDAIK
- 101 **KYDMDLVVCG HHQDFWSKLM** SSARQLINTV **HVDMLIVPLR DEEDE**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
5 - 17	711.3850	1420.7554	1420.7926	-0.0371	0	K.HILLIAVDLSPEK.V (Ions_score 93)
5 - 17	711.4060	1420.7974	1420.7926	0.0049	0	K.HILLIAVDLSPEK.V (Ions_score 82)
5 - 17	711.4080	1420.8014	1420.7926	0.0089	0	K.HILLIAVDLSPEK.V (Ions_score 94)
101 - 118	761.3180	2280.9322	2280.9932	-0.0611	1	K.KYDMDLVVCGHHQDFWSK.L Oxidation (M) (Ions_score 20)
102 - 118	718.3550	2152.0432	2152.8983	-0.8551	0	K.YDMDLVVCGHHQDFWSK.L Oxidation (M) (Ions_score 25)
125 - 145	826.7860	2477.3362	2477.2472	0.0890	1	R.QLINTVHVDMLIVPLRDEEDE.- (Ions_score 56)
125 - 145	832.0960	2493.2662	2493.2421	0.0241	1	R.QLINTVHVDMLIVPLRDEEDE.- Oxidation (M) (Ions_score 51)



LOCUS ZP_05969846 145 aa linear BCT 15-OCT-2009
 DEFINITION universal stress protein A [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05969846
 VERSION ZP_05969846.1 GI:261341988
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession NZ_ABWM02000029.1
 KEYWORDS
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM Enterobacter cancerogenus ATCC 35316
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 145)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 145)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from ABWM02000029.
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be
 aware that the annotation is done automatically with little or no
 manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA
 gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter
 cancerogenus, not the sequenced strain), is a member of the
 Proteobacteria division of the domain bacteria and has been
 isolated from human feces. The sequenced strain was obtained from

http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100106/FtmmSxStt... 21/01/2010

ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

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This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x
Sequencing Technology: 454.
Method: conceptual translation.

FEATURES	Location/Qualifiers
source	1..145 /organism="Enterobacter cancerogenus ATCC 35316" /strain="ATCC 35316" /db_xref="taxon:500639"
Protein	1..145 /product="universal stress protein A" /calculated_mol_wt=16022
Region	5..137 /region_name="USP_Like" /note="Usp: Universal stress protein family. The universal stress protein Usp is a small cytoplasmic bacterial protein whose expression is enhanced when the cell is exposed to stress agents. Usp enhances the rate of cell survival during prolonged exposure to...; cd00293" /db_xref="CDD:30165"
Site	order(9..11,39,109..110,111..112,121..123) /site_type="other" /note="Ligand Binding Site" /db_xref="CDD:30165"
CDS	1..145 /locus_tag="EcanA3_020100016092" /coded_by="NZ_ABWM02000029.1:151914..152351" /note="COG0589 Universal stress protein UspA and related nucleotide-binding proteins" /transl_table=11 /db_xref="CDD:137695"

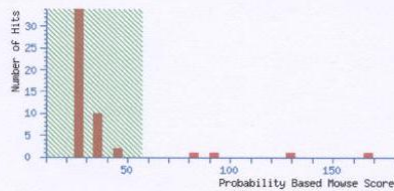
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy_manickan@unn.ac.uk
Search title : SSP 5306
MS data file : D:\Data\Lakshmy\120808\260808\SSP 5306_RG13_01_389.d\SSP 5306_RG13_01_389.mgf
Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 11:29:43 GMT
Protein hits : [gi|157155787](#) periplasmic protein [Escherichia coli E24377A]
[gi|146310199](#) periplasmic protein [Enterobacter sp. 638]
[gi|238794784](#) Osmotically-inducible protein Y [Yersinia intermedia ATCC 29909]
[gi|16763356](#) periplasmic protein [Salmonella enterica subsp. enterica serovar Typhi str. CT18]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: **Peptide Summary** [Help](#)

Significance threshold p<: Max. number of hits:

Standard scoring: MudPIT scoring Ions score or expect cut-off: Show sub-sets:

Show pop-ups: Suppress pop-ups Sort unassigned: Require bold red:

Select All Select None Search Selected Error tolerant

1. [gi|157155787](#) Mass: 21075 Score: 167 Queries matched: 6 emPAI: 0.56
 periplasmic protein [Escherichia coli E24377A]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 78	549.7840	1097.5534	1097.6081	-0.0546	0	(62)	0.016	1	K.LLADDIVPSR.H
<input checked="" type="checkbox"/> 79	549.8330	1097.6514	1097.6081	0.0434	0	65	0.0095	1	K.LLADDIVPSR.H
<input checked="" type="checkbox"/> 80	550.3010	1098.5874	1097.6081	0.9794	0	(46)	1.5	1	K.LLADDIVPSR.H
<input checked="" type="checkbox"/> 96	692.8440	1383.6734	1383.6340	0.0394	0	20	3.2e+02	3	K.VGNFMDDSAITAK.V + Oxidation (M)
<input checked="" type="checkbox"/> 188	1011.5000	2020.9854	2021.0317	-0.0463	0	83	0.00013	1	K.VVTLGPFVESQTQAEAVK.V
<input checked="" type="checkbox"/> 189	674.6710	2020.9912	2021.0317	-0.0405	0	(80)	0.00025	1	K.VVTLGPFVESQTQAEAVK.V

2. [gi|146310199](#) Mass: 21390 Score: 130 Queries matched: 4 emPAI: 0.34
 periplasmic protein [Enterobacter sp. 638]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 78	549.7840	1097.5534	1097.6081	-0.0546	0	(62)	0.016	1	K.LLADDIVPSR.K
<input type="checkbox"/> 79	549.8330	1097.6514	1097.6081	0.0434	0	65	0.0095	1	K.LLADDIVPSR.K
<input type="checkbox"/> 80	550.3010	1098.5874	1097.6081	0.9794	0	(46)	1.5	1	K.LLADDIVPSR.K
<input checked="" type="checkbox"/> 96	692.8440	1383.6734	1383.6340	0.0394	0	65	0.0098	1	K.VGNFMDDSSITAK.V

Proteins matching the same set of peptides:
[gi|261339031](#) Mass: 21366 Score: 130 Queries matched: 4
 periplasmic protein [Enterobacter cancerogenus ATCC 35316]

3. [gi|238794784](#) Mass: 21354 Score: 94 Queries matched: 4 emPAI: 0.16
 Osmotically-inducible protein Y [Yersinia intermedia ATCC 29909]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 78	549.7840	1097.5534	1097.6081	-0.0546	0	(62)	0.016	1	K.LLADDIVPSR.K
<input type="checkbox"/> 79	549.8330	1097.6514	1097.6081	0.0434	0	65	0.0095	1	K.LLADDIVPSR.K
<input type="checkbox"/> 80	550.3010	1098.5874	1097.6081	0.9794	0	(46)	1.5	1	K.LLADDIVPSR.K
<input checked="" type="checkbox"/> 118	781.5810	2341.7212	2341.1510	0.5702	1	29	45	1	K.VETTDGVVQLSGHVESKAQSDR.A

4. [gi|16763356](#) Score: 84 Queries matched: 4
 periplasmic protein [Salmonella enterica subsp. enterica serovar Typhi str. CT18]
 Check to include this hit in error tolerant search

mhtml:file://C:\Documents and Settings\larcje1\Local Settings\Temp\Peptide Summar... 21/01/2010

Mascot Search Results

Protein View

Match to: **gi|261339031** Score: **130**
periplasmic protein [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\260808\SSP 5306_RG13_01_389.d\SSP 5306_RG13_01_389.mgf

Nominal mass (M_r): **21366**; Calculated pI value: **7.79**
 NCBI BLAST search of **gi|261339031** against nr
 Unformatted **sequence_string** for pasting into other applications

Taxonomy: **Enterobacter cancerogenus ATCC 35316**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **11%**

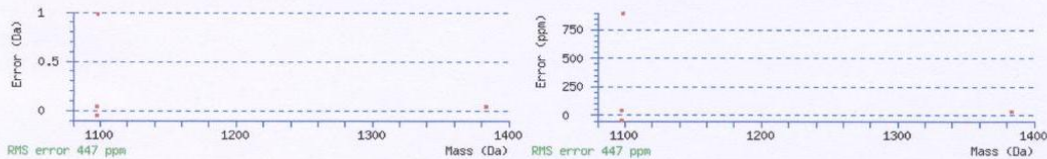
Matched peptides shown in **Bold Red**

1 MNMFLRLKISK TLLAVTLGVS LVSGSALAEV TTMDKAQSTA DTAGDKIDSS
51 MNK**VGNFMDD** **SSITAK**VKAA LVDDEAIKST DISVKTDKV VTLSGFVESQ
101 AQAEQAVKVA KGVGGVTSVS DKLHVRDSKN ASVKGYAGDA ATTSEIKAKL
151 **LADDIVPSRK** VKVETDGVV QLSGTVDSQA QSDRAESIAK AIDGKSVKN
201 DLRTK

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
54 - 66	692.8440	1383.6734	1383.6340	0.0394	0	K.VGNFMDDSSITAK.V (Ions score 65)
150 - 159	549.7840	1097.5534	1097.6081	-0.0546	0	K.LLADDIVPSR.K (Ions score 62)
150 - 159	549.8330	1097.6514	1097.6081	0.0434	0	K.LLADDIVPSR.K (Ions score 65)
150 - 159	550.3010	1098.5874	1097.6081	0.9794	0	K.LLADDIVPSR.K (Ions score 46)



LOCUS ZP_05966889 205 aa linear BCT 15-OCT-2009
 DEFINITION periplasmic protein [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05966889
 VERSION ZP_05966889.1 GI:261339031
 DBLINK Project:28863
 DBSOURCE REFSEQ: accession [NZ_AEWM02000004.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 205)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 205)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [AEWM02000004](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be

http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100105/FtmmSrcL... 21/01/2010

aware that the annotation is done automatically with little or no manual curation.

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

FEATURES	Location/Qualifiers
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Protein	1..205 /product="periplasmic protein" /calculated_mol_wt=21248
Region	3..205 /region_name="PRK10568" /note="periplasmic protein; Provisional; PRK10568" /db_xref="CDD:137994"
Region	64..127 /region_name="BON" /note="Putative phospholipid-binding domain; c102771" /db_xref="CDD:141694"
Region	143..203 /region_name="BON" /note="Putative phospholipid-binding domain; c102771" /db_xref="CDD:141694"
CDS	1..205 /locus_tag="EcanA3_020100001160" /coded_by="complement(NZ_ABWM02000004.1:262747..263364)" /note="COG2823 Predicted periplasmic or secreted lipoprotein" /transl_table=11 /db_xref="CDD:137994"

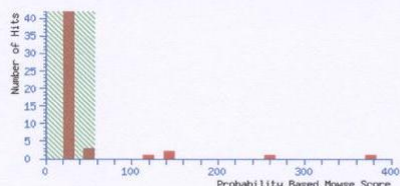
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 5601
MS data file : D:\Data\Lakshmy\120808\260808\SSP 5601_RG24_01_409.d\SSP 5601_RG24_01_409.mgf
Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 11:30:58 GMT
Protein hits : [gi|261338832](#) elongation factor Ts [Enterobacter cancerogenus ATCC 35316]
[gi|1942722](#) Chain B, Elongation Factor Complex Ef-TuEF-Ts From Escherichia Coli
[gi|259907543](#) Elongation factor Ts [Erwinia pyrifoliae Ep1/96]
[gi|50119972](#) elongation factor Ts [Pectobacterium atrosepticum SCRI1043]
[gi|156935315](#) elongation factor Ts [Enterobacter sakazakii ATCC BAA-894]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p < 0.05$ Max. number of hits

Standard scoring MUDPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|261338832](#) Mass: 30472 Score: 376 Queries matched: 8 emPAI: 0.86
 elongation factor Ts [Enterobacter cancerogenus ATCC 35316]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 91	742.8650	1483.7154	1483.6864	0.0290	0	108	4.8e-07	1	K.VETDFAAEVAAMSK.Q + Oxidation (M)
<input checked="" type="checkbox"/> 133	887.9130	1773.8114	1773.8646	-0.0531	0	87	4.9e-05	1	R.VSSLEGDVLSYQHGAR.I
<input checked="" type="checkbox"/> 134	592.2800	1773.8182	1773.8646	-0.0464	0	(76)	0.00064	1	R.VSSLEGDVLSYQHGAR.I
<input checked="" type="checkbox"/> 151	617.6220	1849.8442	1849.9244	-0.0802	0	32	14	5	K.EYQVQLDIAMQSGKPK.E + Oxidation (M)
<input checked="" type="checkbox"/> 155	938.4440	1874.8734	1874.9043	-0.0309	0	91	2e-05	1	K.ALTEANGDIELAIENMR.K + Oxidation (M)
<input checked="" type="checkbox"/> 156	626.3090	1875.9052	1874.9043	1.0008	0	(33)	14	1	K.ALTEANGDIELAIENMR.K + Oxidation (M)
<input checked="" type="checkbox"/> 167	652.6170	1954.8292	1954.9347	-0.1055	0	(41)	1.9	1	K.FTGEVSLTGQPFVMDPSK.S + Oxidation (M)
<input checked="" type="checkbox"/> 168	978.4730	1954.9314	1954.9347	-0.0032	0	57	0.048	1	K.FTGEVSLTGQPFVMDPSK.S + Oxidation (M)

2. [gi|1942722](#) Mass: 30389 Score: 267 Queries matched: 6 emPAI: 0.23
 Chain B, Elongation Factor Complex Ef-TuEF-Ts From Escherichia Coli
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 91	742.8650	1483.7154	1483.6864	0.0290	0	108	4.8e-07	1	K.VETDFAAEVAAMSK.Q + Oxidation (M)
<input checked="" type="checkbox"/> 151	617.6220	1849.8442	1849.9244	-0.0802	0	32	14	5	K.EYQVQLDIAMQSGKPK.E + Oxidation (M)
<input checked="" type="checkbox"/> 155	938.4440	1874.8734	1874.9043	-0.0309	0	91	2e-05	1	K.ALTEANGDIELAIENMR.K + Oxidation (M)
<input checked="" type="checkbox"/> 156	626.3090	1875.9052	1874.9043	1.0008	0	(33)	14	1	K.ALTEANGDIELAIENMR.K + Oxidation (M)
<input checked="" type="checkbox"/> 221	757.0520	2268.1342	2269.0936	-0.9595	0	36	6	1	K.IDGNYGIILEVNCQTFVAK.D
<input checked="" type="checkbox"/> 224	757.7650	2270.2732	2269.0936	1.1795	0	(21)	1.9e+02	3	K.IDGNYGIILEVNCQTFVAK.D

Proteins matching the same set of peptides:
[gi|15799852](#) Mass: 30520 Score: 267 Queries matched: 6
 elongation factor Ts [Escherichia coli O157:H7 EDL933]
[gi|157147400](#) Mass: 32239 Score: 267 Queries matched: 6
 elongation factor Ts [Citrobacter koseri ATCC BAA-895]
[gi|170768398](#) Mass: 30504 Score: 267 Queries matched: 6
 translation elongation factor Ts [Escherichia albertii TW07627]
[gi|172045630](#) Mass: 30471 Score: 267 Queries matched: 6
 RecName: Full=Elongation factor Ts; Short=EF-Ts
[gi|194433429](#) Mass: 30538 Score: 267 Queries matched: 6
 translation elongation factor Ts [Shigella dysenteriae 1012]

Mascot Search Results

Protein View

Match to: **gi|261338832** Score: **376**
elongation factor Ts [**Enterobacter cancerogenus ATCC 35316**]
 Found in search of D:\Data\Lakshmy\120808\260808\SSP 5601_RG24_01_409.d\SSP 5601_RG24_01_409.mgf

Nominal mass (M_n): **30472**; Calculated pI value: **5.13**
 NCBI BLAST search of **gi|261338832** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: **Enterobacter cancerogenus ATCC 35316**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **28%**

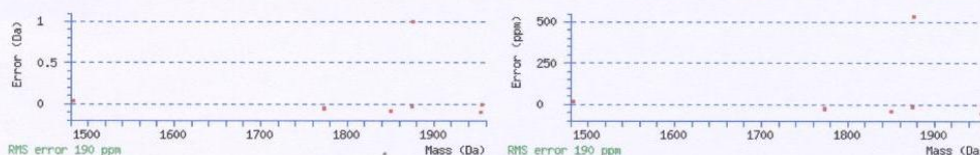
Matched peptides shown in **Bold Red**

1 MAEITASLVK ELRERTGAGM MDCK**KALTEA** NGDI**LAIEN** MRKSGAIKAA
 51 KKAGNVAADC VIITKIDGTY GIILEVNCQT DFLVAKDGGFQ AFANKVLDA
 101 VAGKITDVEV LKAQFEERY ALVAKIGENI NRRV**SLEG** DVLGSY**Q**RGA
 151 RIGVLVAARK ADEELVRQLA MHIAASKPEF VKPEDVSAEV VEKEY**Q**QLD
 201 **IAMQSGKPK**E TAEKMEVGRM KK**FTGEVSLT** **QPFVMDPSK** SVAQLLKEHN
 251 ADVTGFIREF VEGEIER**VET** **DFAAEVAAMS** KQS

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
26 - 42	938.4440	1874.8734	1874.9043	-0.0309	0	K.ALTEANGDI LAIEN MR.K Oxidation (M) (Ions score 91)
26 - 42	626.3090	1875.9052	1874.9043	1.0008	0	K.ALTEANGDI LAIEN MR.K Oxidation (M) (Ions score 33)
135 - 151	887.9130	1773.8114	1773.8646	-0.0531	0	R.VSSLEGDVLGSY Q HGAR.I (Ions score 87)
135 - 151	592.2800	1773.8182	1773.8646	-0.0464	0	R.VSSLEGDVLGSY Q HGAR.I (Ions score 76)
194 - 209	617.6220	1849.8442	1849.9244	-0.0802	0	K.EYQVQLD IAMQSGKPK .E Oxidation (M) (Ions score 32)
223 - 240	652.6170	1954.8292	1954.9347	-0.1055	0	K.FTGEVSLT QPFVMDPSK .S Oxidation (M) (Ions score 41)
223 - 240	978.4730	1954.9314	1954.9347	-0.0032	0	K.FTGEVSLT QPFVMDPSK .S Oxidation (M) (Ions score 57)
268 - 281	742.8650	1483.7154	1483.6864	0.0290	0	K.VET DFAAEVAAMS K.Q Oxidation (M) (Ions score 108)



LOCUS ZP_05966690 283 aa linear BCT 15-OCT-2009
 DEFINITION elongation factor Ts [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05966690
 VERSION ZP_05966690.1 GI:261338832
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM0200004.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM **Enterobacter cancerogenus ATCC 35316**
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 283)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 283)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS [REFSEQ](#): This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM0200004](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be

aware that the annotation is done automatically with little or no manual curation.

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7%
Sequencing Technology: 454.
Method: conceptual translation.

```

FEATURES
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                    certain protein kinases. Although its specific role is so
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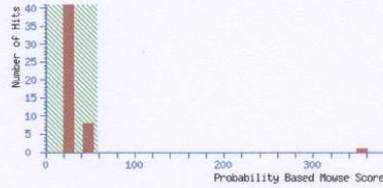
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

(MATRIX) SCIENCE
 User : Lakshmy Manickan
 Email : lakshmy_manickan@unn.ac.uk
 Search title : SSP 5607
 MS data file : D:\Data\Lakshmy\120808\280808\SSP 5607_RH20_01_448.d\SSP 5607_RH20_01_448.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 11:32:24 GMT
 Protein hits : gi|261339130 carbamate kinase [Enterobacter cancerogenus ATCC 35316]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event.
 Individual ions scores > 57 indicate identity or extensive homology (p<0.05).
 Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: **Peptide Summary** [Help](#)

Significance threshold p<: Max. number of hits:

Standard scoring: MudPIT scoring Ions score or expect cut-off: Show sub-sets:

Show pop-ups: Suppress pop-ups Sort unassigned: Require bold red:

Error tolerant

1. **gi|261339130** Mass: 32346 Score: 355 Queries matched: 6 emPAI: 0.48
 carbamate kinase [Enterobacter cancerogenus ATCC 35316]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 33	465.2430	928.4714	928.5341	-0.0627	0	50	0.4	1	K.DLSAALLAR.Q
<input checked="" type="checkbox"/> 63	714.8890	1427.7634	1427.7984	-0.0349	0	86	6.9e-05	1	R.IVESDAITALIQR.D
<input checked="" type="checkbox"/> 65	498.6310	1492.8712	1492.9341	-0.0629	0	27	55	5	R.KPTLVVALGGNALLK.R
<input checked="" type="checkbox"/> 98	878.0740	1754.1334	1753.9938	0.1397	1	80	0.0002	1	R.GIEAVIDRDLAALLAR.Q
<input checked="" type="checkbox"/> 181	766.7240	2297.1502	2297.0859	0.0643	0	55	0.075	1	R.DHLVICNGGGGPPVVENANGYR.G
<input checked="" type="checkbox"/> 188	798.1100	2391.3082	2391.2910	0.0171	0	57	0.047	1	R.VVLVHGNGPQVGLLALQNSAYDK.V

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 39	476.8730	951.7314	952.5818	-0.8503	1	43	0.96	1	AAPGGAILKR
<input checked="" type="checkbox"/> 51	587.8290	1173.6434	1173.7332	-0.0898	2	36	7.2	1	TLKLTKETIK
<input checked="" type="checkbox"/> 65	498.6310	1492.8712	1491.7351	1.1360	1	33	12	1	RGASATGLGEDLMK + Oxidation (M)
<input checked="" type="checkbox"/> 104	608.9710	1823.8912	1823.9377	-0.0465	1	33	12	1	QLADEAEIVLPRGR
<input checked="" type="checkbox"/> 88	563.1710	1686.4912	1685.9147	0.5764	2	30	18	1	ADRRVLLWLANQR + Oxidation (M)
<input checked="" type="checkbox"/> 49	577.7330	1730.1772	1730.8951	-0.7179	1	30	43	1	LPNIDYRVALAEDSR
<input checked="" type="checkbox"/> 55	618.9030	1235.7914	1235.6067	0.1847	1	30	27	1	VEEAMDTVKAK + Oxidation (M)
<input checked="" type="checkbox"/> 140	1034.4450	3100.3132	3099.5196	0.7936	1	29	49	1	GHAEEALNIGTAVASSGLPDMAAAFQR + Oxidation (M)
<input checked="" type="checkbox"/> 199	873.4630	2617.3672	2617.2417	0.1255	2	28	31	1	MRMTFRWYGENNDSVTLQIK
<input checked="" type="checkbox"/> 122	976.7320	2927.1742	2926.4827	0.6915	2	28	52	1	NPDAEILVHGSGRIVGCLTALMITMK + Oxidation (M)
<input checked="" type="checkbox"/> 90	565.5930	1693.7572	1692.8366	0.9206	2	27	47	1	SQMRSAEPRSLPSFR + Oxidation (M)
<input checked="" type="checkbox"/> 109	927.1790	2778.5152	2779.4691	-0.9539	2	27	71	1	QFGKKTAVSVTFDIPQQMVGIIGR + Oxidation (M)
<input checked="" type="checkbox"/> 139	1034.1020	3099.2842	3098.6652	0.6190	2	27	73	1	ITKLRLYVSGNLLTFSSLYQLDPEGK
<input checked="" type="checkbox"/> 182	771.7530	2312.2372	2312.1107	0.1265	1	26	54	1	CRSFDAGADGTAFAEGIVVALK
<input checked="" type="checkbox"/> 166	742.3730	2224.0972	2224.0549	0.0423	0	25	70	1	FGPPGASSNGFNENLALFNSGK
<input checked="" type="checkbox"/> 124	985.8260	2954.4562	2954.4194	0.0368	0	25	1.1e+02	1	ETGAWASHVFPFVADAVMEAAASSGK + Oxidation (M)
<input checked="" type="checkbox"/> 113	944.0300	2829.0682	2829.4669	-0.3988	2	25	1.3e+02	1	GIRHLKLDLCTIPVSYAWAEMR + Oxidation (M)
<input checked="" type="checkbox"/> 115	947.1350	2838.3832	2838.4182	-0.0350	1	25	1.3e+02	1	TTSDAALSGGPPAPAPRALILMSGLQSGK + Oxidation (M)
<input checked="" type="checkbox"/> 46	563.4510	1687.3312	1687.8199	-0.4887	0	24	1.3e+02	1	SSMAQAGKPIEPATER + Oxidation (M)
<input checked="" type="checkbox"/> 202	883.8770	2648.6092	2648.3129	0.2963	2	24	57	1	LQAVRFMEGLVAGETGSPGRSER + Oxidation (M)
<input checked="" type="checkbox"/> 99	881.5150	2641.5232	2640.3619	1.1612	0	24	1.6e+02	1	FLSQALSLQAEQINTANAGPNIVR
<input checked="" type="checkbox"/> 149	1059.9540	3176.8402	3176.6713	0.1689	1	24	1.3e+02	1	DLLKEIMESALFPTMDVLLVFANVSGVK + 2 Oxidation (M)
<input checked="" type="checkbox"/> 151	1067.5050	3199.4932	3198.5734	0.9198	2	24	1.5e+02	1	KTEELSSSYRVFNFRPVASTADYFR
<input checked="" type="checkbox"/> 94	573.3000	1716.8782	1716.9860	-0.1078	2	24	1.2e+02	1	GRHLHDAGVAAVVGRR
<input checked="" type="checkbox"/> 101	599.0420	1794.1042	1794.8788	-0.7746	2	24	99	1	DWIELKADKNSYDK
<input checked="" type="checkbox"/> 54	615.6450	1843.9132	1843.8332	0.0800	0	23	2.2e+02	1	VDVYGTMEASAAMGLAK + Oxidation (M)
<input checked="" type="checkbox"/> 152	1067.5820	2133.1494	2133.0815	0.0680	0	23	1.9e+02	1	ANFVTQQPPGSLGTSPAAATR
<input checked="" type="checkbox"/> 117	952.4810	1902.9474	1904.0441	-1.0967	1	23	1.2e+02	1	MTPEQLQGVLTGVLYK
<input checked="" type="checkbox"/> 89	847.3550	2539.0432	2538.4866	0.5566	2	23	2e+02	1	MALVNVKVLIIQGGIGMSAAILLER + Oxidation (M)

MASCOT **SCIENCE** Mascot Search Results

Protein View

Match to: **gi|261339130** Score: **355**
carbamate kinase [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\280808\SSP 5607_RH20_01_448.d\SSP 5607_RH20_01_448.mgf

Nominal mass (M_r): **32346**; Calculated pI value: **5.13**
 NCBI BLAST search of **gi|261339130** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **29%**

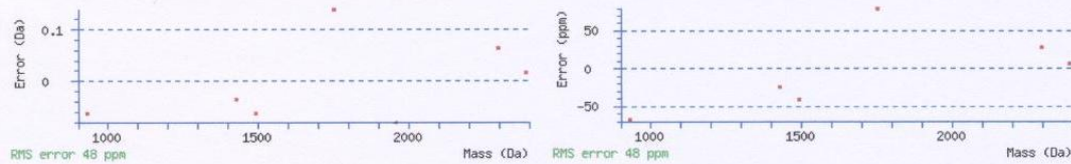
Matched peptides shown in **Bold Red**

1 MERKPTLVVA LGGNALLKRG EPLEAEIQRQ NIELAARTIA GLTAQWRVVL
51 VHGNGPQVGL LALQNSAYDK VTPYPLDILG AESQGMIGYM LQQALKNNLP
101 QREVSLLTQ VEVDPADPAF TNPTKYIGPI YTDAAKALA AERGWIFKAD
151 GSAFRRVVPS PPKRIVESD AITALIQRDH LVICNGGGGV **PVVENANGYR**
201 GIEAVIDKDL SAALLARQIE ADALLILTDA DAVYLDWGPQ TQRPLAQTVP
251 ALLNDMQFDA GSMGPRVAAC REFVEACGGI AGIGALADGA EILAGEKGTLL
301 IRN

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
4 - 18	498.6310	1492.8712	1492.9341	-0.0629	0	R.KPTLVVALGGNALLK.R (Ions score 27)
48 - 70	798.1100	2391.3082	2391.2910	0.0171	0	R.VVLVHGNGPQVGLLALQNSAYDK.V (Ions score 57)
166 - 178	714.8890	1427.7634	1427.7984	-0.0349	0	R.IVESDAITALIQR.D (Ions score 86)
179 - 200	766.7240	2297.1502	2297.0859	0.0643	0	R.DELVICNGGGGVVVVENANGYR.G (Ions score 55)
201 - 217	878.0740	1754.1334	1753.9938	0.1397	1	R.GIEAVIDKDL SAALLAR .Q (Ions score 80)
209 - 217	465.2430	928.4714	928.5341	-0.0627	0	K.DLSAALLAR.Q (Ions score 50)



LOCUS ZP_05966988 303 aa linear BCT 15-OCT-2009
 DEFINITION carbamate kinase [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05966988
 VERSION ZP_05966988.1 GI:261339130
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000004.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM Enterobacter cancerogenus ATCC 35316
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 303)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 303)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM02000004](#).

Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be found here: <http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x
 Sequencing Technology: 454.
 Method: conceptual translation.

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FEATURES
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                  ATP-phosphorylation of carbamate and carbamoyl phosphate
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Mascot: <http://www.matrixscience.com/>

<input checked="" type="checkbox"/>	81	754.9170	1507.8194	1507.7518	0.0676	1	67	0.006	1	K.SLYEADLVDEAKR.L	
<input type="checkbox"/>	111	580.6830	1739.0272	1738.9763	0.0508	1	24	1.1e+02	3	K.ALKEPARFMVAIVGGSK.V + Oxidation (M)	
<input type="checkbox"/>	135	938.0120	1874.0094	1873.9462	0.0633	0	97	5.8e-06	1	K.ISYISTGGGAFLEFVEGK.V	
<input type="checkbox"/>	147	665.0230	1992.0472	1992.0258	0.0214	0	55	0.084	1	K.TILWNGPVGVFFFNFR.K	
<input type="checkbox"/>	162	687.3070	2058.8992	2059.9132	-1.0140	0	49	0.31	1	K.YAALCDVFMDFGTAAHR.A + Oxidation (M)	
<input checked="" type="checkbox"/>	163	1030.5570	2059.0994	2060.0136	-0.9141	0	(32)		16	1	K.FADVACAGPLLALELALGK.A
<input checked="" type="checkbox"/>	164	687.3740	2059.1002	2060.0136	-0.9134	0	50	0.22	1	K.FADVACAGPLLALELALGK.A	
<input type="checkbox"/>	233	817.4700	2449.3882	2449.3078	0.0804	0	80	0.00022	1	K.IADQLIVGGGIANTFVAAQGHNVGK.S	

3. [gi|1156932627](#) Mass: 41426 Score: 613 Queries matched: 11 emPAI: 1.00
 phosphoglycerate kinase [Enterobacter sakazakii ATCC BAA-894]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
56	578.3620	1154.7094	1154.6910	0.0184	0	47	0.62	1	R.ASLPTIEIALK.Q
58	622.3450	1242.6754	1242.6642	0.0113	0	53	0.17	1	K.VLPAVAMLEER.A + Oxidation (M)
65	676.8330	1351.6514	1351.6507	0.0008	0	58	0.042	1	K.SLYEADLVDEAKR.R
81	754.9170	1507.8194	1507.7518	0.0676	1	67	0.006	1	K.SLYEADLVDEAKR.L
111	580.6830	1739.0272	1738.9763	0.0508	1	24	1.1e+02	3	K.ALKEPARFMVAIVGGSK.V + Oxidation (M)
135	938.0120	1874.0094	1873.9462	0.0633	0	97	5.8e-06	1	K.ISYISTGGGAFLEFVEGK.V
147	665.0230	1992.0472	1992.0258	0.0214	0	55	0.084	1	K.TILWNGPVGVFFFNFR.K
163	1030.5570	2059.0994	2060.0136	-0.9141	0	(19)	3.5e+02	7	K.FADVACAGPLLAELDALGK.A
164	687.3740	2059.1002	2060.0136	-0.9134	0	34	9.3	2	K.FADVACAGPLLAELDALGK.A
200	1109.6410	2217.2674	2217.1165	0.1510	0	102	1.5e-06	1	K.DYLDGVVAVAGELVLENVR.F
233	817.4700	2449.3882	2449.3078	0.0804	0	80	0.00022	1	K.IADQLIVGGGIANTFVAAQGHNVGK.S

Proteins matching the same set of peptides:
[gi|260599261](#) Mass: 41470 Score: 613 Queries matched: 11
 Phosphoglycerate Kinase [Cronobacter turicensis]

4. [gi|242238006](#) Mass: 41399 Score: 435 Queries matched: 7 emPAI: 0.54
 Phosphoglycerate Kinase [Dickeya dadantii Ech703]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
56	578.3620	1154.7094	1154.6910	0.0184	0	47	0.62	1	R.ASLPTIEIALK.Q
72	734.9040	1467.7934	1467.7457	0.0478	0	112	1.9e-07	1	R.VATEFSETATATLK.S
73	490.2880	1467.8422	1467.7457	0.0965	0	(28)	40	1	R.VATEFSETATATLK.S
135	938.0120	1874.0094	1873.9462	0.0633	0	97	5.8e-06	1	K.ISYISTGGGAFLEFVEGK.K
147	665.0230	1992.0472	1992.0258	0.0214	0	55	0.084	1	K.TILWNGPVGVFFFNFR.K
162	687.3070	2058.8992	2059.9132	-1.0140	0	49	0.31	1	K.YAALCDVFMDFGTAAHR.A + Oxidation (M)
233	817.4700	2449.3882	2449.3078	0.0804	0	80	0.00022	1	R.IADQLIVGGGIANTFVAAQGHNVGK.S

5. [gi|227114292](#) Mass: 41338 Score: 362 Queries matched: 6 emPAI: 0.54
 phosphoglycerate kinase [Pectobacterium carotovorum subsp. brasiliensis FBR1692]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide	
56	578.3620	1154.7094	1154.6910	0.0184	0	47	0.62	1	R.ASLPTIEIALK.Q	
135	938.0120	1874.0094	1873.9462	0.0633	0	97	5.8e-06	1	K.ISYISTGGGAFLEFVEGK.K	
147	665.0230	1992.0472	1992.0258	0.0214	0	55	0.084	1	K.TILWNGPVGVFFFNFR.K	
162	687.3070	2058.8992	2059.9132	-1.0140	0	49	0.31	1	K.YAALCDVFMDFGTAAHR.A + Oxidation (M)	
233	817.4700	2449.3882	2449.3078	0.0804	0	80	0.00022	1	K.IADQLIVGGGIANTFVAAQGHNVGK.S	
<input checked="" type="checkbox"/>	242	903.9790	2708.9152	2708.3545	0.5607	0	38	2.3	1	R.AIAESDAFSIAGGGDTLAAIDLEGIADK.I

Proteins matching the same set of peptides:
[gi|253690062](#) Mass: 41334 Score: 362 Queries matched: 6
 Phosphoglycerate Kinase [Pectobacterium carotovorum subsp. carotovorum PC1]
[gi|261823130](#) Mass: 41340 Score: 362 Queries matched: 6
 Phosphoglycerate kinase [Pectobacterium wasabiae WPP163]

6. [gi|157372179](#) Mass: 41484 Score: 319 Queries matched: 5 emPAI: 0.30
 phosphoglycerate kinase [Serratia proteamaculans 568]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
58	622.3450	1242.6754	1242.6642	0.0113	0	28	45	2	K.ALPAVVMLEER.A + Oxidation (M)
135	938.0120	1874.0094	1873.9462	0.0633	0	97	5.8e-06	1	K.ISYISTGGGAFLEFVEGK.A
147	665.0230	1992.0472	1992.0258	0.0214	0	55	0.084	1	K.TILWNGPVGVFFFNFR.K
162	687.3070	2058.8992	2059.9132	-1.0140	0	38	4	2	K.YAALCDVFMDFGTAAHR.A
200	1109.6410	2217.2674	2217.1165	0.1510	0	102	1.5e-06	1	K.DYLDGVVAVAGELVLENVR.F

7. [gi|238759311](#) Mass: 39537 Score: 306 Queries matched: 5 emPAI: 0.44
 Phosphoglycerate kinase [Yersinia aldovae ATCC 35236]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
135	938.0120	1874.0094	1873.9462	0.0633	0	97	5.8e-06	1	K.ISYISTGGGAFLEFVEGK.K
147	665.0230	1992.0472	1992.0258	0.0214	0	55	0.084	1	K.TILWNGPVGVFFFNFR.K
162	687.3070	2058.8992	2059.9132	-1.0140	0	38	4	2	K.YAALCDVFMDFGTAAHR.A
233	817.4700	2449.3882	2449.3078	0.0804	0	80	0.00022	1	K.IADQLIVGGGIANTFVAAQGHNVGK.S
242	903.9790	2708.9152	2708.3545	0.5607	0	38	2.3	1	R.AIAESDAFSIAGGGDTLAAIDLEGIADK.I

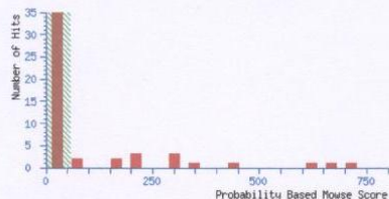
Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 5701
 MS data file : D:\Data\Lakshmy\120808\190808\SSP 5701_RF24_01_363.d\SSP 5701_RF24_01_363.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:02:00 GMT

- Protein hits
- gi|146312978 phosphoglycerate kinase [Enterobacter sp. 638]
 - gi|261342323 phosphoglycerate kinase [Enterobacter cancerogenus ATCC 35316]
 - gi|156932627 phosphoglycerate kinase [Enterobacter sakazakii ATCC BAA-894]
 - gi|242238006 Phosphoglycerate kinase [Dickeya dadantii Ech703]
 - gi|227114292 phosphoglycerate kinase [Pectobacterium carotovorum subsp. brasiliensis PBR1692]
 - gi|157372179 phosphoglycerate kinase [Serratia proteamaculans 568]
 - gi|238759311 Phosphoglycerate kinase [Yersinia aldovae ATCC 35236]
 - gi|261346331 phosphoglycerate kinase [Providencia rustigianii DSM 4541]
 - gi|268592003 phosphoglycerate kinase [Providencia rettgeri DSM 1131]
 - gi|53733312 COG0126: 3-phosphoglycerate kinase [Haemophilus influenzae R2866]
 - gi|183601149 hypothetical protein PROSTU_04779 [Providencia stuartii ATCC 25827]
 - gi|22127184 phosphoglycerate kinase [Yersinia pestis KIM]
 - gi|220936052 Phosphoglycerate kinase [Thioalkalivibrio sp. HL-EbGR7]
 - gi|15640504 phosphoglycerate kinase [Vibrio cholerae O1 biovar El Tor str. N16961]
 - gi|261210056 phosphoglycerate kinase [Vibrio sp. RC341]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|146312978](#) Mass: 41414 Score: 713 Queries matched: 13 emPAI: 1.00
 phosphoglycerate kinase [Enterobacter sp. 638]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
<input checked="" type="checkbox"/> 56	578.3620	1154.7094	1154.6910	0.0184	0 47 0.62	1 R.ASLPTIELALK.Q
<input checked="" type="checkbox"/> 58	622.3450	1242.6754	1242.6642	0.0113	0 53 0.17	1 K.VLPAVAMLEER.A + Oxidation (M)
<input checked="" type="checkbox"/> 72	734.9040	1467.7934	1467.7457	0.0478	0 112 1.9e-07	1 R.VATEFSETATATLK.S
<input checked="" type="checkbox"/> 73	490.2880	1467.8422	1467.7457	0.0965	0 (28) 40	1 R.VATEFSETATATLK.S
<input checked="" type="checkbox"/> 111	580.6830	1739.0272	1738.9763	0.0508	1 24 1.1e+02	3 K.ALKEPARPMVAIVGSK.V + Oxidation (M)
<input checked="" type="checkbox"/> 135	938.0120	1874.0094	1873.9462	0.0633	0 97 5.8e-06	1 K.ISYISTGGGAFLEFVEGK.V
<input checked="" type="checkbox"/> 147	665.0230	1992.0472	1992.0258	0.0214	0 55 0.084	1 K.TILWNGPVGVFFPNFR.K
<input checked="" type="checkbox"/> 162	687.3070	2058.8992	2059.9132	-1.0140	0 49 0.31	1 K.YAALCDVFMDFGTAR.A + Oxidation (M)
<input checked="" type="checkbox"/> 163	1030.5570	2059.0994	2060.0136	-0.9141	0 (19) 3.5e+02	7 K.FADVACGPLLAELDALK.A
<input checked="" type="checkbox"/> 164	687.3740	2059.1002	2060.0136	-0.9134	0 34 9.3	2 K.FADVACGPLLAELDALK.A
<input checked="" type="checkbox"/> 200	1109.6410	2217.2674	2217.1165	0.1510	0 102 1.5e-06	1 K.DYLDGVEVARGELVVLNVR.F
<input checked="" type="checkbox"/> 220	772.7840	2315.3302	2315.1492	0.1809	1 68 0.0031	1 K.SVNDIRDEEQILDLDVSAQK.L
<input checked="" type="checkbox"/> 233	817.4700	2449.3882	2449.3078	0.0804	0 80 0.00022	1 K.IADQLIVGGGIANTFVAAQGHNVGK.S

2. [gi|261342323](#) Mass: 41343 Score: 684 Queries matched: 13 emPAI: 1.19
 phosphoglycerate kinase [Enterobacter cancerogenus ATCC 35316]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
56	578.3620	1154.7094	1154.6910	0.0184	0 47 0.62	1 R.ASLPTIELALK.Q
58	622.3450	1242.6754	1242.6642	0.0113	0 53 0.17	1 K.VLPAVAMLEER.A + Oxidation (M)
<input checked="" type="checkbox"/> 65	676.8330	1351.6514	1351.6507	0.0008	0 58 0.042	1 K.SLYEADLVDEAK.R
72	734.9040	1467.7934	1467.7457	0.0478	0 112 1.9e-07	1 R.VATEFSETATATLK.S
73	490.2880	1467.8422	1467.7457	0.0965	0 (28) 40	1 R.VATEFSETATATLK.S

MASCOT Mascot Search Results

Protein View

Match to: **gi|261342323** Score: **684**
phosphoglycerate kinase [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120800\190800\SSP 5701_RF24_01_363.d\SSP 5701_RF24_01_363.mgf

Nominal mass (M_r): **41343**; Calculated pI value: **5.08**

NCBI BLAST search of **gi|261342323** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **42%**

Matched peptides shown in **Bold Red**

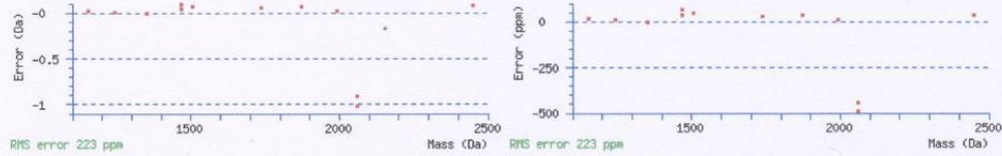
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101 VARGELVVLE NVRFNKGEKK DDETLSSKYYA ALCDVFMVDA FGTAAHQAS
151 THGIGKPADV ACAGPLLADE LEALGKALKE PARPMVAIVG GSKVSTKLTV
201 LDLSLKIADQ LIVGGGIANT FVAAQGHNVG KSLYEADLVD EAKRLITTC
251 IPVPTDVRVA TEFSETATAT LKSVNDIKDD EQILDLDVDS AQKLAELTKN
301 AKTILWNGFV GVPEFPNFRK GTEIVANAIA DSEAFSIAGG GDTLAAIDL
351 GIADKISYIS TGGGAFLEFV EGRVLEPAM LEERAKQ
    
```

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
39 - 49	578.3620	1154.7094	1154.6910	0.0184	0	R.ASLEPTIELALK.Q (Ions score 47)
129 - 146	687.3070	2058.8992	2059.9132	-1.0140	0	K.YAALCDVFMVDAFGTAHR.A Oxidation (M) (Ions score 49)
157 - 176	1030.5570	2059.0994	2060.0136	-0.9141	0	K.FADVACAGPLLADELEALGK.A (Ions score 32)
157 - 176	687.3740	2059.1002	2060.0136	-0.9134	0	K.FADVACAGPLLADELEALGK.A (Ions score 50)
177 - 193	580.6830	1739.0272	1738.9763	0.0508	1	K.ALKEPARPMVAIVGGSK.V Oxidation (M) (Ions score 24)
207 - 231	817.4700	2449.3882	2449.3078	0.0804	0	K.IADQLIVGGGIANTFVAAQGHNVGK.S (Ions score 80)
232 - 243	676.8330	1351.6514	1351.6507	0.0008	0	K.SLYEADLVDEAK.R (Ions score 58)
232 - 244	754.9170	1507.8194	1507.7518	0.0676	1	K.SLYEADLVDEAKR.L (Ions score 67)
259 - 272	734.9040	1467.7934	1467.7457	0.0478	0	R.VATEFSETATATLK.S (Ions score 112)
259 - 272	490.2880	1467.8422	1467.7457	0.0965	0	R.VATEFSETATATLK.S (Ions score 28)
303 - 319	665.0230	1992.0472	1992.0258	0.0214	0	K.TILWNGPVGVEFPNFR.K (Ions score 55)
356 - 373	938.0120	1874.0094	1873.9462	0.0633	0	K.ISYITGGGAFLEFVEGK.V (Ions score 97)
374 - 384	622.3450	1242.6754	1242.6642	0.0113	0	K.VLPAVAMLEER.A Oxidation (M) (Ions score 53)



LOCUS ZP_05970181 387 aa linear BCT 15-OCT-2009
 DEFINITION phosphoglycerate kinase [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05970181
 VERSION ZP_05970181.1 GI:261342323
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABW02000033.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM Enterobacter cancerogenus ATCC 35316
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 387)
 AUTHORS Weinstein,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 387)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA

COMMENT WGS REFSEQ: This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from [ABWM02000033](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be found here: <http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).
 We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.
 This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project (<http://nihroadmap.nih.gov/hmp/>).

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

FEATURES Location/Qualifiers

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 - /strain="ATCC 35316"
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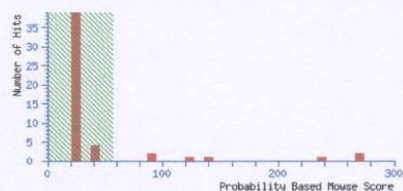
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

Matrix Science
 User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 5702
 MS data file : D:\Data\Lakshmy\120808\280808\SSP 5702_RI12_01_475.d\SSP 5702_RI12_01_475.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 11:34:50 GMT
 Protein hits :
 gi|146312874 phosphopyruvate hydratase [Enterobacter sp. 638]
 gi|156932734 phosphopyruvate hydratase [Enterobacter sakazakii ATCC BAA-894]
 gi|157148317 phosphopyruvate hydratase [Citrobacter koseri ATCC BAA-895]
 gi|260771907 enolase [Vibrio metschnikovii CIP 69.14]
 gi|198241971 phosphopyruvate hydratase [Salmonella enterica subsp. enterica serovar Dublin str. CT_02021853]
 gi|15642443 phosphopyruvate hydratase [Vibrio cholerae O1 biovar El Tor str. N16961]
 gi|54310177 phosphopyruvate hydratase [Photobacterium profundum SS9]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)
 Significance threshold p < 0.05 Max. number of hits
 Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets
 Show pop-ups Suppress pop-ups Sort unassigned Require bold red
 Error tolerant

1. [gi|146312874](#) Mass: 45497 Score: 270 Queries matched: 4 emPAI: 0.15
 phosphopyruvate hydratase [Enterobacter sp. 638]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 68	478.8270	955.6394	955.6066	0.0328	0	45	0.95	1	K.GIVNSILIK.F
<input checked="" type="checkbox"/> 105	733.3860	1464.7574	1463.8460	0.9114	0	55	0.091	2	K.AVGAVNGPIAQAVLQK.D
<input checked="" type="checkbox"/> 117	781.9560	1561.8974	1561.8352	0.0623	0	92	2e-05	1	K.IQLVGDLLFVNTNK.I
<input checked="" type="checkbox"/> 177	957.9100	1913.8054	1913.9047	-0.0993	0	79	0.00031	1	K.APTSEEPHFLEDLTK.Q

2. [gi|156932734](#) Mass: 45573 Score: 262 Queries matched: 4 emPAI: 0.23
 phosphopyruvate hydratase [Enterobacter sakazakii ATCC BAA-894]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 105	733.3860	1464.7574	1463.8460	0.9114	0	69	0.0032	1	K.AVGAVNGPIAQAVLQK.D
<input checked="" type="checkbox"/> 117	781.9560	1561.8974	1561.8352	0.0623	0	92	2e-05	1	K.IQLVGDLLFVNTNK.I
<input checked="" type="checkbox"/> 148	875.8420	1749.6694	1750.7430	-1.0735	0	22	1.2e+02	5	K.DITLAMDCAASEFYK.D + Oxidation (M)
<input checked="" type="checkbox"/> 177	957.9100	1913.8054	1913.9047	-0.0993	0	79	0.00031	1	K.APTSEEPHFLEDLTK.Q

Proteins matching the same set of peptides:
[gi|260599132](#) Mass: 45516 Score: 262 Queries matched: 4
 Enolase [Cronobacter turicensis]

3. [gi|157148317](#) Mass: 45646 Score: 238 Queries matched: 4 emPAI: 0.15
 phosphopyruvate hydratase [Citrobacter koseri ATCC BAA-895]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 68	478.8270	955.6394	955.6066	0.0328	0	45	0.95	1	K.GIVNSILIK.F
<input checked="" type="checkbox"/> 117	781.9560	1561.8974	1561.8352	0.0623	0	92	2e-05	1	K.IQLVGDLLFVNTNK.I
<input checked="" type="checkbox"/> 148	875.8420	1749.6694	1750.7430	-1.0735	0	22	1.2e+02	5	K.DITLAMDCAASEFYK.D + Oxidation (M)
<input checked="" type="checkbox"/> 177	957.9100	1913.8054	1913.9047	-0.0993	0	79	0.00031	1	K.APTSEEPHFLEDLTK.Q

Proteins matching the same set of peptides:
[gi|261342208](#) Mass: 45644 Score: 238 Queries matched: 4
 phosphopyruvate hydratase [Enterobacter cancerogenus ATCC 35316]

MASCOT Mascot Search Results

Protein View

Match to: **gi|261342208** Score: 238
phosphopyruvate hydratase [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\280808\SSP 5702_RI12_01_475.d\SSP 5702_RI12_01_475.mgf

Nominal mass (M_r): **45644**; Calculated pI value: **5.19**
 NCBI BLAST search of **gi|261342208** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **12%**

Matched peptides shown in **Bold Red**

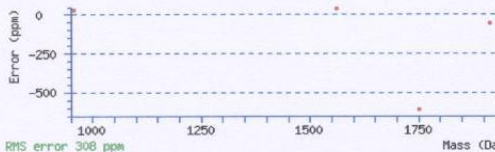
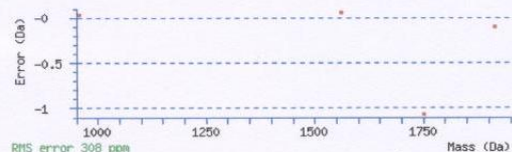
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151 MNIINGGEHA DNNVDIQEPM IQPVGAKTLK EAVRMGSEVF HNLAKVLKAK
201 GMNTAVGDEG GYAVNLGSNA EALAVIAEAV KAAGYELGKD ITLAMDCAAS
251 EFYKDGKIVL AGEGNKAFTS EEFTHFLEDL TKQYPIVSIE DGLDESDWDG
301 FAYQTKVLGD KIQLVGGDLF VTNTHKILKEG IEKGIVNSIL IKFNQIGSLT
351 ETLAAIKMAK DAGYTAIVISH RSGETEDATI ADLAVGTAAG QIKTGSMRSR
401 DRVAKYNQLI RIEEALGEKA PYNGRKEIKG QA
    
```

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
240 - 254	875.8420	1749.6694	1750.7430	-1.0735	0	K.DITLAMDCAASEFYK.D Oxidation (M) (Ions score 22)
267 - 282	957.9100	1913.8054	1913.9047	-0.0993	0	K.AFTSEEFTHFLEDLTK.Q (Ions score 79)
312 - 325	781.9560	1561.8974	1561.8352	0.0623	0	K.IQLVGGDLFVNTK.I (Ions score 92)
334 - 342	478.8270	955.6394	955.6066	0.0328	0	K.GIVNSILIK.F (Ions score 45)



LOCUS ZP_05970066 432 aa linear BCT 15-OCT-2009
 DEFINITION phosphopyruvate hydratase [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05970066
 VERSION ZP_05970066.1 GI:261342208
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000033.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 432)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahanty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 432)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM02000033](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be

4. [gi|260771907](#) Mass: 45789 Score: 136 Queries matched: 2 emPAI: 0.07
 enolase [Vibrio metschnikovii CIP 69.14]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
68	478.8270	955.6394	955.6066	0.0328	0	45	0.95	1	K.GIVNSILIK.F
117	781.9560	1561.8974	1561.8352	0.0623	0	92	2e-05	1	K.LQLVGGDDLFTVNTK.I

5. [gi|198241971](#) Mass: 45627 Score: 126 Queries matched: 2 emPAI: 0.07
 phosphopyruvate hydratase [Salmonella enterica subsp. enterica serovar Dublin str. CT_02021853]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
117	781.9560	1561.8974	1561.8352	0.0623	0	92	2e-05	1	K.IQLVGGDDLFTVNTK.I
<input checked="" type="checkbox"/> 148	875.8420	1749.6694	1749.7589	-0.0895	0	34	7.8	1	K.NITLAMDCAASEFYK.D + Oxidation (M)

6. [gi|15642443](#) Mass: 45836 Score: 92 Queries matched: 1 emPAI: 0.07
 phosphopyruvate hydratase [Vibrio cholerae O1 biovar El Tor str. N16961]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
117	781.9560	1561.8974	1561.8352	0.0623	0	92	2e-05	1	K.IQLVGGDDLFTVNTK.I

Proteins matching the same set of peptides:

- [gi|29839249](#) Mass: 45848 Score: 92 Queries matched: 1
 RecName: Full=Enolase; AltName: Full=2-phosphoglycerate dehydratase; AltName: Full=2-phospho-D-glycerate hydro-lyase
- [gi|77918834](#) Mass: 46350 Score: 92 Queries matched: 1
 enolase [Pelobacter carbinolicus DSM 2380]
- [gi|90412096](#) Mass: 45596 Score: 92 Queries matched: 1
 phosphopyruvate hydratase [Photobacterium profundum 3TCK]
- [gi|1117617509](#) Mass: 45750 Score: 92 Queries matched: 1
 phosphopyruvate hydratase [Aeromonas hydrophila subsp. hydrophila ATCC 7966]
- [gi|121591783](#) Mass: 33004 Score: 92 Queries matched: 1
 enolase [Vibrio cholerae 2740-80]
- [gi|145300350](#) Mass: 45766 Score: 92 Queries matched: 1
 phosphopyruvate hydratase [Aeromonas salmonicida subsp. salmonicida A449]
- [gi|153803630](#) Mass: 17966 Score: 92 Queries matched: 1
 enolase, C- TIM barrel domain [Vibrio cholerae M20-3]
- [gi|153827653](#) Mass: 40048 Score: 92 Queries matched: 1
 enolase [Vibrio cholerae M20-2]
- [gi|229527053](#) Mass: 45850 Score: 92 Queries matched: 1
 enolase [Vibrio cholerae 12129(1)]
- [gi|237809481](#) Mass: 45583 Score: 92 Queries matched: 1
 enolase [Tolomonas auensis DSM 9187]
- [gi|258626906](#) Mass: 45864 Score: 92 Queries matched: 1
 Enolase [Vibrio mimicus VM603]
- [gi|261211542](#) Mass: 45820 Score: 92 Queries matched: 1
 enolase [Vibrio sp. RC341]
- [gi|269123733](#) Mass: 46259 Score: 92 Queries matched: 1
 enolase [Streptococcus moniliformis DSM 12112]

7. [gi|54310177](#) Score: 92 Queries matched: 1
 phosphopyruvate hydratase [Photobacterium profundum SS9]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
117	781.9560	1561.8974	1561.8352	0.0623	0	92	2e-05	1	K.LQLVGGDDLFTVNTK.I

Proteins matching the same set of peptides:

- [gi|184390114](#) Score: 92 Queries matched: 1
- [gi|86146349](#) Score: 92 Queries matched: 1
- [gi|218710558](#) Score: 92 Queries matched: 1
- [gi|260888598](#) Score: 92 Queries matched: 1

Peptide matches not assigned to protein hits: (no details means no match)

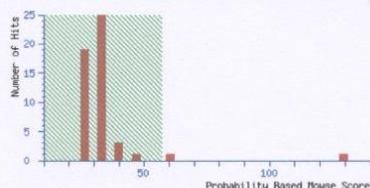
Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 84	647.3270	1292.6394	1291.7136	-0.9259	1	47	0.56	1	KTFADIGQTIAK
<input checked="" type="checkbox"/> 233	767.0460	2298.1162	2298.2947	-0.1785	1	39	3.2	1	QKPNQAQALEYLDTIKFLK
<input checked="" type="checkbox"/> 226	747.0360	2238.0862	2238.1314	-0.0453	1	29	27	1	SGLDGGGMITGSAGAFVLESRK + Oxidation (M)
<input checked="" type="checkbox"/> 130	816.4800	2446.4182	2447.1421	-0.7239	2	29	55	1	DLTSDHSTCEGRTLPRIK
<input checked="" type="checkbox"/> 209	1067.4960	3199.4662	3200.4945	-1.0284	2	27	73	1	TTTLQQRSTANVDRFCEWITSTENR
<input checked="" type="checkbox"/> 100	714.6410	2140.9012	2140.1496	0.7515	2	27	75	1	LTMGRAPDVLAMSATPIPR + Oxidation (M)
<input checked="" type="checkbox"/> 222	742.7000	2225.0782	2224.0206	-1.0576	0	27	46	1	MTDVTATPASADPVAAYDPTSK + Oxidation (M)
<input checked="" type="checkbox"/> 158	902.4180	2704.2322	2704.3234	-0.0913	0	27	96	1	SSPDVGGQVLQMLPAGATLSLTGCMIR + Oxidation (M)
<input checked="" type="checkbox"/> 140	846.6630	2536.9672	2537.1793	-0.2121	2	27	71	1	TFVFLKAMFYGMEYSRGTK
<input checked="" type="checkbox"/> 51	428.2160	854.4174	855.5066	-1.0891	0	27	54	1	IATVLPDK
<input checked="" type="checkbox"/> 218	737.7080	2210.1022	2210.1219	-0.0197	1	26	54	1	VEAEKAFSTINTEFQNLAAK
<input checked="" type="checkbox"/> 93	686.3130	2055.9172	2057.0542	-1.1370	0	26	1.3e+02	1	VFTGNPALDAAAAPVTSSNR
<input checked="" type="checkbox"/> 107	738.8360	2213.4862	2214.2115	-0.7254	1	26	1.3e+02	1	MSAVELLGINREVALIMIK
<input checked="" type="checkbox"/> 214	729.3010	2184.8812	2184.1397	0.7415	0	25	54	1	FILGFINGAVLMCAIPSLAK + 2 Oxidation (M)
<input checked="" type="checkbox"/> 92	683.7950	1365.5754	1365.6268	-0.0514	0	25	1.4e+02	1	MTGSLVDDEMR
<input checked="" type="checkbox"/> 49	835.7180	834.7107	834.3905	0.3202	0	25	67	1	SLEGLCR

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6203
 MS data file : D:\Data\Lakshmy\120808\241108\SSP 6203 RD22_01_943.d\SSP 6203_RD22_01_943.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 11:47:36 GMT
 Protein hits : [gi|129164](#) RecName: Full=Outer membrane protein X; Flags: Precursor
 [gi|1440183](#) similar to outer membrane protein X from Enterobacter cloacae, Swiss-Prot Accession Number P25253; ORF2 [Es.

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|129164](#) Mass: 18700 Score: 129 Queries matched: 2 emPAI: 0.39
 RecName: Full=Outer membrane protein X; Flags: Precursor
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 78	708.7790	1415.5434	1415.6834	-0.1399	0	66	0.0056	1	K.GGYGITAGPAYR.L
<input checked="" type="checkbox"/> 118	898.9310	1795.8474	1796.9097	-1.0623	0	63	0.014	1	R.LNDWASIVGVVGVGYGK.F

Proteins matching the same set of peptides:

- [gi|16759751](#) Mass: 18541 Score: 129 Queries matched: 2
outer membrane protein X [Salmonella enterica subsp. enterica serovar Typhi str. CT18]
- [gi|152969400](#) Mass: 18452 Score: 129 Queries matched: 2
outer membrane protein X [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
- [gi|161504000](#) Mass: 18554 Score: 129 Queries matched: 2
outer membrane protein X [Salmonella enterica subsp. arizonae serovar 62:z4,z23:--]
- [gi|198245355](#) Mass: 18640 Score: 129 Queries matched: 2
outer membrane protein X [Salmonella enterica subsp. enterica serovar Dublin str. CT_02021853]
- [gi|205352088](#) Mass: 18540 Score: 129 Queries matched: 2
outer membrane protein X [Salmonella enterica subsp. enterica serovar Gallinarum str. 287/91]
- [gi|206578835](#) Mass: 18528 Score: 129 Queries matched: 2
outer membrane protein X [Klebsiella pneumoniae 342]
- [gi|213583694](#) Mass: 14338 Score: 129 Queries matched: 2
outer membrane protein X [Salmonella enterica subsp. enterica serovar Typhi str. E98-0664]
- [gi|224582645](#) Mass: 18771 Score: 129 Queries matched: 2
outer membrane protein X [Salmonella enterica subsp. enterica serovar Paratyphi C strain RKS4594]
- [gi|261341400](#) Mass: 15248 Score: 129 Queries matched: 2
outer membrane protein X [Enterobacter cancerogenus ATCC 35316]

2. [gi|1440183](#) Mass: 14762 Score: 63 Queries matched: 1 emPAI: 0.23
 similar to outer membrane protein X from Enterobacter cloacae, Swiss-Prot Accession Number P25253; ORF2 [Escherichia coli]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 118	898.9310	1795.8474	1796.9097	-1.0623	0	63	0.014	1	R.LNDWASIVGVVGVGYGK.F

Proteins matching the same set of peptides:

- [gi|16435772](#) Mass: 16350 Score: 63 Queries matched: 1
Chain A, Crystal Structure Of The Outer Membrane Protein OmpX From Escherichia Coli
- [gi|15800566](#) Mass: 18649 Score: 63 Queries matched: 1
outer membrane protein X [Escherichia coli O157:H7 EDL933]
- [gi|26246790](#) Mass: 18879 Score: 63 Queries matched: 1
outer membrane protein X [Escherichia coli CFT073]
- [gi|30749947](#) Mass: 16390 Score: 63 Queries matched: 1
Chain A, Nmr Fold Of The Outer Membrane Protein OmpX In Dhpc Micelles

aware that the annotation is done automatically with little or no manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

```

FEATURES
  source              1..432
                    /organism="Enterobacter cancerogenus ATCC 35316"
                    /strain="ATCC 35316"
                    /db_xref="taxon:500639"
  Protein            1..432
                    /product="phosphopyruvate hydratase"
                    /EC_number="4.2.1.11"
                    /calculated_mol_wt=45484
  Region            1..432
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                    /note="phosphopyruvate hydratase; Provisional; PRK00077"
                    /db_xref="CDD:134085"
  Region            6..414
                    /region_name="enolase"
                    /note="Enolase: Enolases are homodimeric enzymes that
                    catalyse the reversible dehydration of
                    2-phospho-D-glycerate to phosphoenolpyruvate as part of
                    the glycolytic and gluconeogenesis pathways. The reaction
                    is facilitated by the presence of metal ions; cd03313"
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  Site              order(8,10..18,22,34,160..162,183..184,187..188,191..192,
                    205..206,213,215,374..376,399..401,403..404,407,410..411,
                    414)
                    /site_type="other"
                    /note="dimer interface"
                    /db_xref="CDD:48188"
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                    /site_type="metal-binding"
                    /db_xref="CDD:48188"
  Site              order(159,209,342,370..372,393)
                    /site_type="other"
                    /note="substrate binding pocket"
                    /db_xref="CDD:48188"
  CDS               1..432
                    /gene="eno"
                    /locus_tag="EcanA3_020100017202"
                    /coded_by="complement(NZ_ABWM02000033.1:143450..144748)"
                    /note="COG0148 Enolase"
                    /transl_table=11
                    /db_xref="CDD:134085"
    
```

Mascot: <http://www.matrixscience.com/>

MASCOT **SCIENCE** Mascot Search Results

Protein View

Match to: **gi|261341400** Score: **129**
outer membrane protein X [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\241108\SSP 6203_RD22_01_943.d\SSP 6203_RD22_01_943.mgf

Nominal mass (M_r): **15248**; Calculated pI value: **5.38**
 NCBI BLAST search of **gi|261341400** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **22%**

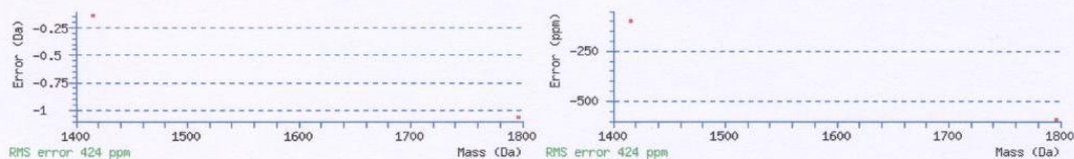
Matched peptides shown in **Bold Red**

1 MQGVNKNKING FNLKYRYEQD NNPLGVIGSF TYTEKDRTEN GAYNKG**QYYG**
51 **ITAGPAYRLN DWASIVGVVG VGYGK**FQOTE NEGLNRTASN SDYGFSYGAG
101 MQFNPIENVA LDFSIEQSRI RNVDVGTWIA GVGYRF

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
46 - 58	708.7790	1415.5434	1415.6834	-0.1399	0 K.GQYYGITAGPAYR.L (Ions score 66)
59 - 75	898.9310	1795.8474	1796.9097	-1.0623	0 R.LNDWASIVGVVGVGK.F (Ions score 63)



LOCUS ZP_05969258 136 aa linear BCT 15-OCT-2009
 DEFINITION outer membrane protein X [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05969258
 VERSION ZP_05969258.1 GI:261341400
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000020.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE
 AUTHORS 1 (residues 1 to 136)
 Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE
 AUTHORS 2 (residues 1 to 136)
 Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS [REFSEQ](#): This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM02000020](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be
 aware that the annotation is done automatically with little or no
 manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA
 gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter

cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project (<http://nihroadmap.nih.gov/hmp/>).

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454.

Method: conceptual translation.

FEATURES	Location/Qualifiers
source	1..136 /organism="Enterobacter cancerogenus ATCC 35316" /strain="ATCC 35316" /db_xref="taxon:500639"
Protein	1..136 /product="outer membrane protein X" /calculated_mol_wt=15127
Region	<1..136 /region_name="Surface Ag 2" /note="Surface antigen; cl01155" /db_xref="CDD:141128"
CDS	1..136 /gene="ompX" /locus_tag="EcanA3_020100013096" /coded_by="NZ_ABWMD2000020.1:323158..323568" /note="COG3637 Opacity protein and related surface antigens" /transl_table=11 /db_xref="CDD:137256"

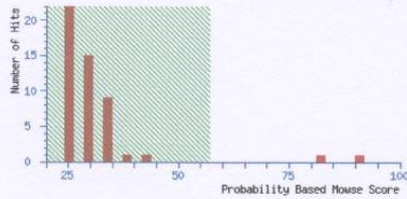
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6306
 MS data file : D:\Data\Lakshmy\120808\SSP 6306_RB6_01_719.d\SSP 6306_RB6_01_719.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 11:53:30 GMT
 Protein hits : [gi|152972668](#) hypothetical protein KPN_04168 [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
 [gi|50582463](#) oxidoreductase [Exiguobacterium acetylicum]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|152972668](#) Mass: 10606 Score: 91 Queries matched: 3 emPAI: 0.75
 hypothetical protein KPN_04168 [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 104	753.3170	1504.6194	1504.7595	-0.1401	0	48	0.43	1	K.EPELNLKQFLQK.L + Oxidation (M)
<input checked="" type="checkbox"/> 123	840.9220	1679.8294	1679.9246	-0.0952	0	43	1.3	1	R.LNEVIELLQPAWQK.E
<input checked="" type="checkbox"/> 124	560.9530	1679.8372	1679.9246	-0.0874	0	(29)	36	3	R.LNEVIELLQPAWQK.E

Proteins matching the same set of peptides:
[gi|206579500](#) Mass: 10388 Score: 91 Queries matched: 3
 hypothetical protein KPK_5514 [Klebsiella pneumoniae 342]
[gi|261342932](#) Mass: 10363 Score: 91 Queries matched: 3
 hypothetical protein EcanA3_20877 [Enterobacter cancerogenus ATCC 35316]

2. [gi|50582463](#) Mass: 27360 Score: 83 Queries matched: 3 emPAI: 0.12
 oxidoreductase [Exiguobacterium acetylicum]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 174	677.2520	2028.7342	2028.9752	-0.2411	0	24	67	9	K.SVDLADNQNVDLYEGLK.E
<input checked="" type="checkbox"/> 250	984.4460	2950.3162	2950.4236	-0.1075	0	58	0.025	1	K.ELDIETWINNAGFGDFLVQDIELGK.I
<input checked="" type="checkbox"/> 162	984.7600	2951.2582	2950.4236	0.8345	0	(55)	0.096	1	K.ELDIETWINNAGFGDFLVQDIELGK.I

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 85	610.2780	1218.5414	1218.6972	-0.1558	0	41	2.7	1	LTDLQLFLLR
<input checked="" type="checkbox"/> 149	948.4010	1894.7874	1894.9312	-0.1438	0	39	2.9	1	IGDTAPAQYQTPEYTIK
<input checked="" type="checkbox"/> 220	761.6840	2282.0302	2282.2867	-0.2565	0	35	6.5	1	LLEYGLGIGYLLLPFPWR
<input checked="" type="checkbox"/> 124	560.9530	1679.8372	1680.0371	-0.2000	2	35	8.2	1	KLELVPMIKLIIDR
<input checked="" type="checkbox"/> 75	563.2460	1686.7162	1686.9212	-0.2051	2	34	21	1	VHHGVARALLAACRR
<input checked="" type="checkbox"/> 234	837.2810	2508.8212	2509.1802	-0.3590	2	33	8	1	DGARIGTTGNIGIGPCYADLAARMR + Oxidation (M)
<input checked="" type="checkbox"/> 82	602.2490	1202.4834	1202.6441	-0.1607	1	32	19	1	LTDQVKAAMR
<input checked="" type="checkbox"/> 88	629.2630	1256.5114	1255.6996	0.8118	1	32	18	1	AEALRQGATLAR
<input checked="" type="checkbox"/> 248	972.1250	2913.3532	2912.4425	0.9107	1	30	16	1	VYRNHINNVEMIMAGHIALPEYQK + Oxidation (M)
<input checked="" type="checkbox"/> 73	562.4190	1122.8234	1121.6346	1.1889	2	30	20	1	FQRFQGRV
<input checked="" type="checkbox"/> 157	647.5700	1939.6882	1939.8689	-0.1808	0	29	20	1	MGLAEGDMIGTAAGLATCGK + Oxidation (M)
<input checked="" type="checkbox"/> 175	677.2520	2028.7342	2027.9881	0.7461	1	29	23	1	AVADEGARGVLPAGPMQAMR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 174	677.2520	2028.7342	2028.8881	-0.1540	0	29	24	1	GGSMTTMDHSLFVGSSTALGNK + 2 Oxidation (M)
<input checked="" type="checkbox"/> 222	767.0440	2298.1102	2298.2107	-0.1006	1	28	34	1	FYDVSSIGLGLDGGQTLTKLSK
<input checked="" type="checkbox"/> 118	822.2250	2463.6532	2464.2420	-0.5888	0	28	49	1	YAAESFTPNRPVLEALELMAR + Oxidation (M)

Mascot Search Results

Protein View

Match to: **gi|261342932** Score: **91**
hypothetical protein EcanA3_20877 [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\SSP 6306_RB6_01_719.d\SSP 6306_RB6_01_719.mgf

Nominal mass (M_r): **10363**; Calculated pI value: **5.02**
 NCBI BLAST search of **gi|261342932** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **29%**

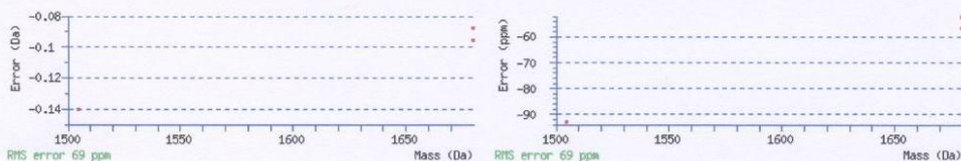
Matched peptides shown in **Bold Red**

1 MKCKRLNEVI **ELLQPAWQRE** **PELNLMQFLQ** KLAKESGFDG ELTDLSDIIL
51 IYQLKMRDSA KDAVIPGIQK DYEEFKTAL LRARGVIKE

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss Sequence
6 - 19	840.9220	1679.8294	1679.9246	-0.0952	0 R.LNEVI ELLQPAWQK .E (Ions score 43)
6 - 19	560.9530	1679.8372	1679.9246	-0.0874	0 R.LNEVI ELLQPAWQK .E (Ions score 29)
20 - 31	753.3170	1504.6194	1504.7595	-0.1401	0 K.EPELNLMQFLQK .L Oxidation (M) (Ions score 48)



LOCUS ZP_05970790 89 aa linear BCT 15-OCT-2009
 DEFINITION hypothetical protein EcanA3_20877 [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05970790
 VERSION ZP_05970790.1 GI:261342932
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000045.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM Enterobacter cancerogenus ATCC 35316
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 89)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B., Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C., Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C., Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M., Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 REFERENCE 2 (residues 1 to 89)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H., Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from [ABWM02000045](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: 296078, the 16S rDNA gene of a related strain of Enterobacter

cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

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FEATURES             Location/Qualifiers
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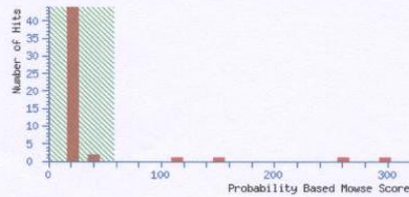
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 6401
MS data file : D:\Data\Lakshmy\120808\280808\SSP 6401_RK4_01_560.d\SSP 6401_RK4_01_560.mgf
Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 11:56:40 GMT
Protein hits : [gi|195940647](#) cytidylate kinase [Escherichia coli O157:H7 str. EC4024]
[gi|6137462](#) Chain A, Cmp Kinase From Escherichia Coli Free Enzyme Structure
[gi|152969494](#) cytidylate kinase [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
[gi|261339244](#) cytidylate kinase [Enterobacter cancerogenus ATCC 35316]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 58 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from mowse scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|195940647](#) Mass: 24783 Score: 297 Queries matched: 5 emPAI: 0.46
 cytidylate kinase [Escherichia coli O157:H7 str. EC4024]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 130	696.3070	1390.5994	1390.6803	-0.0808	0	68	0.0041	1	R.DMSTVVFEDAPVK.I + Oxidation (M)
<input checked="" type="checkbox"/> 150	784.4830	1566.9514	1566.8617	0.0897	0	79	0.00031	1	M.TAIAPVITIDGPGGAGK.G
<input checked="" type="checkbox"/> 161	830.4350	1658.8554	1658.8376	0.0178	0	90	2.9e-05	1	R.TQEVANAASQVAAPFR.V
<input checked="" type="checkbox"/> 247	793.4150	2377.2232	2377.1649	0.0583	0	40	2.4	1	R.FVSTGNGNLEVILEGEDVSGEIR.T
<input type="checkbox"/> 250	940.8840	2819.6302	2820.5372	-0.9070	0	20	1.5e+02	3	R.AVAPLVPALVDLSTSLTIEQVIEK.A

2. [gi|6137462](#) Mass: 24759 Score: 264 Queries matched: 4 emPAI: 0.46
 Chain A, Cmp Kinase From Escherichia Coli Free Enzyme Structure
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 130	696.3070	1390.5994	1390.6803	-0.0808	0	68	0.0041	1	R.DMSTVVFEDAPVK.I + Oxidation (M)
<input type="checkbox"/> 150	784.4830	1566.9514	1566.8617	0.0897	0	79	0.00031	1	M.TAIAPVITIDGPGGAGK.G
<input type="checkbox"/> 161	830.4350	1658.8554	1658.8376	0.0178	0	90	2.9e-05	1	R.TQEVANAASQVAAPFR.V
<input type="checkbox"/> 247	793.4150	2377.2232	2376.1809	1.0423	0	27	49	3	R.FVSTGNGNLEVILEGEDVSGEIR.T

Proteins matching the same set of peptides:

[gi|15800771](#) Mass: 24789 Score: 264 Queries matched: 4
 cytidylate kinase [Escherichia coli O157:H7 EDL933]
[gi|24112319](#) Mass: 24803 Score: 264 Queries matched: 4
 cytidylate kinase [Shigella flexneri 2a str. 301]
[gi|74311467](#) Mass: 24805 Score: 264 Queries matched: 4
 cytidylate kinase [Shigella sonnei Ss046]
[gi|82777567](#) Mass: 24729 Score: 264 Queries matched: 4
 cytidylate kinase [Shigella dysenteriae Sd197]
[gi|88192972](#) Mass: 24764 Score: 264 Queries matched: 4
 Chain A, Mutant R188m Of The Cytidine Monophosphate Kinase From E. Coli
[gi|157146404](#) Mass: 24818 Score: 264 Queries matched: 4
 cytidylate kinase [Citrobacter koseri ATCC BAA-895]

3. [gi|152969494](#) Mass: 24917 Score: 158 Queries matched: 2 emPAI: 0.29
 cytidylate kinase [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
130	696.3070	1390.5994	1390.6803	-0.0808	0	68	0.0041	1	R.DMGTVVFPDAPVK.I + Oxidation (M)
161	830.4350	1658.8554	1658.8740	-0.0185	1	90	2.9e-05	1	R.TQEVANAASKVAAPFR.V

Proteins matching the same set of peptides:
 qi|206579846 Mass: 24874 Score: 158 Queries matched: 2
 cytidylate kinase [Klebsiella pneumoniae 342]
 qi|238893966 Mass: 24931 Score: 158 Queries matched: 2
 cytidylate kinase [Klebsiella pneumoniae NTUH-K2044]
 qi|262040991 Mass: 27105 Score: 156 Queries matched: 2
 cytidylate kinase [Klebsiella pneumoniae subsp. rhinoscleromatis ATCC 13884]

4. qi|261339244 Mass: 24768 Score: 115 Queries matched: 3 emPAI: 0.14
 cytidylate kinase [Enterobacter cancerogenus ATCC 35316]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
130	696.3070	1390.5994	1390.6803	-0.0808	0	68	0.0041	1	R.DMGTVVFPDAPVK.I + Oxidation (M)
247	793.4150	2377.2232	2376.1809	1.0423	0	27	49	3	R.FVSTNGNLEVLGEDVSGEIR.T
250	940.8840	2819.6302	2820.5372	-0.9070	0	20	1.5e+02	3	R.AVAPLVPADALVLDSTSLTIEQVIEK.A

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
198	971.0240	1940.0334	1939.8266	0.2068	1	46	0.71	1	MHFKEGEMTLQSCR
238	737.7190	2210.1352	2210.1439	-0.0087	1	30	25	1	DAYLIIGGGTIISMQAGMKGAK + Oxidation (M)
223	1043.4990	2084.9834	2084.0432	0.9402	1	30	45	1	SERNAGAGAVIMHALESAIR + Oxidation (M)
106	565.3480	1693.0222	1693.8471	-0.8249	2	30	63	1	EMTRSFQLGRWQR
243	747.0480	2238.1222	2238.0409	0.0813	1	28	36	1	LEQMTREASQWLEMQGAK + Oxidation (M)
189	623.0330	1866.0772	1865.7148	0.3624	0	28	40	1	QDGAGEDSGSDGASGSDGAK
142	759.4010	2275.1812	2276.0295	-0.8484	0	28	90	1	MNNYVQAWQLGLSFSGR + Oxidation (M)
241	742.3940	2224.1602	2223.0240	1.1362	2	26	61	1	MFRGGAFWGGSRQTDAEGR + Oxidation (M)
144	771.9940	2312.9602	2313.0954	-0.1352	2	26	1.3e+02	1	GGREVMARLHQELVACAER + Oxidation (M)
197	647.6310	1939.8712	1939.9356	-0.0644	1	25	84	1	MLEQAHQDAGACIAR
120	642.3800	1924.1182	1924.0088	0.1094	2	24	2e+02	1	VKGTMTVAQTGGDKYAIK
250	940.8840	2819.6302	2819.5197	0.1105	1	24	64	1	TRASTIMHGFWILLFAVPPFAGLVEK + Oxidation (M)
199	972.5850	2914.7332	2914.4297	0.3035	2	24	1.6e+02	1	VHPYRLNFAFAGEAGPLGWMEEGRVR + Oxidation (M)
183	910.0500	2727.1282	2726.3407	0.7875	1	24	1.8e+02	1	LMLQKAATLYDAGQFAAAEANNKAK
205	1002.0500	2002.0854	2000.9480	1.1375	0	23	1.2e+02	1	LITWSSDYDDGFSVALGR
93	491.3570	1471.0492	1470.8042	0.2450	1	23	2.1e+02	1	AVIAREGTEASVEK
173	879.1510	2634.4312	2634.4315	-0.0004	2	23	1.9e+02	1	YAAMPFLKKAELLEGVVHLSAVR + Oxidation (M)
230	1072.9460	2143.8774	2143.1830	0.6944	2	23	1e+02	1	KLYVSLVKNYFPFVVDOR
195	645.3600	1933.0582	1932.9615	0.0967	0	23	1.4e+02	1	DITTLAAMAAIAPDFDAR
111	587.8760	1760.6062	1760.9156	-0.3094	1	23	2.7e+02	1	VTVNESDLAALGKSTK
126	676.0180	2025.0322	2025.8846	-0.8524	0	23	2.4e+02	1	SEYSGVHEVMAMPGLIEK + 2 Oxidation (M)
129	691.9220	1381.8294	1380.7222	1.1072	1	22	1.6e+02	1	GVRGGAGGPAEAGR
118	632.3890	1894.1452	1894.9981	-0.8530	2	22	3e+02	1	QAKGVAAALMLLAEHR
196	970.7840	2909.3302	2910.4592	-1.1291	2	22	2e+02	1	LNQGMRLQDTPFVIYGMGSAYNGRIR
202	982.2250	2943.6532	2944.4569	-0.8037	1	22	2.1e+02	1	MSAMDAGIPAGTVAAVMGRAPVAQLEAR + 2 Oxidation (M)
48	375.1880	1122.5422	1123.6349	-1.0928	0	22	3.6e+02	1	VQALLEAAGR
190	936.7680	2807.2822	2806.3068	0.9754	2	22	2.3e+02	1	FGRHYGGCGSPRPVPGYGDVPAVCR
240	1110.5770	3328.7092	3327.7615	0.9477	2	22	2.5e+02	1	GEVRFYQAVGSEEAFFKAYRQGLSLVLEK
137	735.8300	2204.4682	2203.9700	0.4981	1	22	2.8e+02	1	GQHSICCVHTEAFLRYK + Oxidation (M)
248	853.9570	2558.8492	2559.3526	-0.5034	2	22	98	1	VGGATYQVFEVRRPARRMALAMR + 2 Oxidation (M)
231	1075.6540	3223.9402	3223.6005	0.3396	2	22	2.3e+02	1	VLWKQNIYASGMCTGPIIRVSDANLSK + Oxidation (M)
222	1042.1070	2082.1994	2083.0334	-0.8340	2	22	1.6e+02	1	SGYERVDIVENKGEFSVR
155	811.2030	2430.5872	2431.2529	-0.6657	2	21	2.5e+02	1	NFLMIIGGFARSLKNSDENDK
128	680.9010	1359.7874	1359.6895	0.0980	0	21	3.4e+02	1	EALBAEGHIVR
184	911.8940	2732.6602	2732.3922	0.2680	2	21	2.9e+02	1	LREFFSSRSYPTIIGYTNELPR
162	838.7720	2513.2942	2514.1155	-0.8213	1	21	3.2e+02	1	SFMFNGNIQSTGPRYCPISIEDK + Oxidation (M)
53	391.1710	1170.4912	1171.5330	-1.0418	1	21	4.2e+02	1	AQQGNGPSSRGN
194	964.0440	2889.1102	2889.3744	-0.2643	1	21	3.2e+02	1	MHLTQLFASIMVAGALTACAQNGADMK + 3 Oxidation (M)
166	844.0780	2529.2122	2530.3465	-1.1344	1	21	3.4e+02	1	LGAAGVHVIQDNLQVVFQPKSDSIK + Oxidation (M)
98	514.3690	1540.0852	1539.7682	0.3170	0	21	3.9e+02	1	VPADVASPWLSDAGR
171	860.3590	2578.0552	2577.3360	0.7192	1	21	3.4e+02	1	QELVEVLDTHLLPQDKTINK + Oxidation (M)
233	1082.6550	3244.9432	3245.6179	-0.6747	2	21	2.8e+02	1	EHQEFKQGFIFTQILLADEINRCSPK
102	553.9120	1105.8094	1106.5179	-0.7084	0	21	2e+02	1	GYNIVACGPR
104	557.3730	1669.0972	1668.7930	0.3042	1	21	4.4e+02	1	HSQVMVGGTSPFGTKK + Oxidation (M)
179	902.2060	2703.5962	2703.3438	0.2523	0	20	3.1e+02	1	LAALMQLNGEIVYQLEEIHHTHK + Oxidation (M)
168	849.9520	2546.8342	2546.3162	0.5179	1	20	3.8e+02	1	ARSASPEVVLITAYGTAAAVEAMR + Oxidation (M)
145	772.8310	2315.4712	2316.1420	-0.6708	1	20	3.9e+02	1	DEETFWNTIFIDAMFSPIRR + Oxidation (M)
226	1047.7250	3140.1532	3140.6401	-0.4869	1	20	2.6e+02	1	RASSAGVLHVGSLFMAGITIAKANGAGALTR
135	479.2060	1434.5962	1435.7459	-1.1498	0	20	2.4e+02	1	AWLFIDPATFAER
151	790.3940	2368.1602	2369.1222	-0.9621	0	20	4.8e+02	1	NVNLLMPEAYAPHNSYIDR + Oxidation (M)
224	1046.0650	3135.1732	3135.6434	-0.4702	1	20	3.8e+02	1	GGHASPHEAADPVFAAEIVLALQAMVTRR
158	824.1480	2469.4222	2470.1917	-0.7695	2	20	4e+02	1	DRDGGFLPMDGPEFAEAMKR
175	895.7880	2684.3422	2683.2661	1.0761	1	20	4.1e+02	1	GQVCGLRKWSVDLDAQITVHDNR

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project (<http://nihroadmap.nih.gov/hmp/>).

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

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FEATURES
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Mascot: <http://www.matrixscience.com/>

MASCOT **SCIENCE** Mascot Search Results

Protein View

Match to: **gi|261339244** Score: **115**
cytidylate kinase [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\280808\SSP 6401_RK4_01_560.d\SSP 6401_RK4_01_560.mgf

Nominal mass (M_r): **24768**; Calculated pI value: **5.32**
 NCBI BLAST search of **gi|261339244** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **27%**

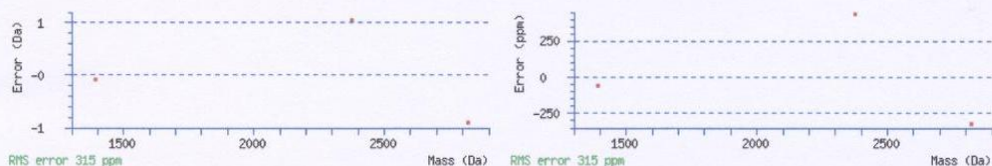
Matched peptides shown in **Bold Red**

1 MTAVAPVITI DGPSGAGKGT LCKAMAEALQ WHLLDSGAIY RVLALALHH
51 HVDVASEEAL VPLAAHLVDR FVSTNGNLEV ILEGEDVSGE IRTQDVANAA
101 SQVAAPFPRV EALLRRQRGF REAPGLIADG RDMGTVVFDP APVKIFLDAS
151 SEERAQRML QLQEKGFVFN FERLLSEIKE RDDRDRNRV APLVPAEDAL
201 VLDSTSLTIE QVIEKALQYA RQKLALA

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
71 - 92	793.4150	2377.2232	2376.1809	1.0423	0	R.FVSTNGNLEVILEGEDVSGEIR.T (Ions score 27)
132 - 144	696.3070	1390.5994	1390.6803	-0.0808	0	R.DMGTVVFDPAPVK.I Oxidation (M) (Ions score 68)
189 - 215	940.8840	2819.6302	2820.5372	-0.9070	0	R.AVALVPAEDALVLDSTSLTIEQVIEK.A (Ions score 20)



LOCUS ZP_05967102 227 aa linear BCT 15-OCT-2009
 DEFINITION cytidylate kinase [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05967102
 VERSION ZP_05967102.1 GI:261339244
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABNM02000004.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 227)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 227)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS [REFSEQ](#): This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABNM02000004](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be
 aware that the annotation is done automatically with little or no
 manual curation.

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

FEATURES	Location/Qualifiers
source	1..227 /organism="Enterobacter cancerogenus ATCC 35316" /strain="ATCC 35316" /db_xref="taxon:500639"
Protein	1..227 /product="cytidylate kinase" /calculated_mol_wt=24594
Region	5..225 /region_name="cmk" /note="Cytidylate kinase; Provisional; PRK00023" /db_xref="CDD:134038"
Region	7..205 /region_name="CMPK" /note="Cytidine monophosphate kinase (CMPK) catalyzes the reversible phosphorylation of cytidine monophosphate (CMP) to produce cytidine diphosphate (CDP), using ATP as the preferred phosphoryl donor; cd02020" /db_xref="CDD:73296"
Site	order(110,131..132,188) /site_type="other" /note="CMP-binding site" /db_xref="CDD:73296"
Site	order(181,185) /site_type="other" /note="The sites determining sugar specificity" /db_xref="CDD:73296"
CDS	1..227 /locus_tag="EcanA3_020100002225" /coded_by="NZ_ABWM02000004.1:495167..495850" /note="COG0283 Cytidylate kinase" /transl_table=11 /db_xref="CDD:134038"

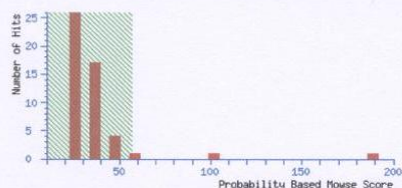
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy_manickan@unn.ac.uk
 Search title : SSP 6408
 MS data file : D:\Data\Lakshmy\120808\280808\SSP 6408_RI7_01_467.d\SSP 6408_RI7_01_467.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 11:58:13 GMT
 Protein hits : [gi|261342934](#) periplasmic protein disulfide isomerase I [Enterobacter cancerogenus ATCC 35316]
 [gi|2558846](#) disulfide oxidoreductase [Salmonella enterica subsp. enterica serovar Typhimurium]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: Peptide Summary [Help](#)

Significance threshold p<: Max. number of hits:

Standard scoring: MudPIT scoring Ions score or expect cut-off: Show sub-sets:

Show pop-ups: Suppress pop-ups Sort unassigned: Require bold red:

Select All Select None Search Selected Error tolerant

1. [gi|261342934](#) Mass: 23134 Score: 189 Queries matched: 5 emPAI: 0.50
 periplasmic protein disulfide isomerase I [Enterobacter cancerogenus ATCC 35316]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
73	630.3790	1258.7434	1258.6710	0.0724	0	42	2.1	5	K.YHVEFLGGLGK.D
<input checked="" type="checkbox"/> 114	800.8990	1599.7834	1599.7205	0.0629	0	67	0.006	1	K.GEDYDAWNISFVVK.S
<input checked="" type="checkbox"/> 155	600.6850	1799.0332	1798.9465	0.0867	0	(44)	1.1	1	K.DLTQAMAVAIALGVEDK.V
<input checked="" type="checkbox"/> 156	900.5290	1799.0434	1798.9465	0.0969	0	80	0.0024	1	K.DLTQAMAVAIALGVEDK.V
<input checked="" type="checkbox"/> 157	900.5730	1799.1314	1798.9465	0.1849	0	(50)	0.21	1	K.DLTQAMAVAIALGVEDK.V

Proteins matching the same set of peptides:

[gi|152972670](#) Mass: 23047 Score: 187 Queries matched: 5
 periplasmic protein disulfide isomerase I [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
[gi|206577959](#) Mass: 23034 Score: 187 Queries matched: 5
 thiol:disulfide interchange protein DsbA [Klebsiella pneumoniae 342]

2. [gi|2558846](#) Score: 102 Queries matched: 2
 disulfide oxidoreductase [Salmonella enterica subsp. enterica serovar Typhimurium]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
73	630.3790	1258.7434	1258.6710	0.0724	0	42	2.1	5	K.YHVEFLGGLGK.E
114	800.8990	1599.7834	1598.7365	1.0469	0	61	0.025	2	K.GENYDAWNISFVVK.S

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 70	608.8640	1215.7134	1215.6870	0.0265	2	53	0.16	1	L.TAPLMRRSR + Oxidation (M)
<input checked="" type="checkbox"/> 73	630.3790	1258.7434	1258.6128	0.1306	1	51	0.25	1	RAWGGLDVPK
<input checked="" type="checkbox"/> 72	420.5380	1258.5922	1257.6717	0.9204	0	35	11	1	ALLDGLFAGPER
<input checked="" type="checkbox"/> 219	747.0790	2238.2152	2238.1202	0.0950	1	34	8.8	1	THNVTKSYVLGEGEIVSSVAK + Oxidation (M)
<input checked="" type="checkbox"/> 197	1042.0290	2082.0434	2082.0456	-0.0021	1	32	14	1	VQNKFDVIELVDGIGDFR + Oxidation (M)
<input checked="" type="checkbox"/> 226	767.4120	2299.2142	2298.2947	0.9195	1	32	14	1	QKPNQALETLDTIKFLK
<input checked="" type="checkbox"/> 223	762.1160	2283.3262	2284.0971	-0.7710	1	31	18	1	KWTEKQGSFLNSDATASTSK
<input checked="" type="checkbox"/> 212	1106.1540	2210.2934	2210.0598	0.2336	2	31	19	1	YNQAKMPEIELVDMEEK + Oxidation (M)
<input checked="" type="checkbox"/> 76	685.7070	2054.0992	2054.1850	-0.0858	1	29	59	1	VFHLSSAVLIVSLYKQK + Oxidation (M)
<input checked="" type="checkbox"/> 154	897.9300	1793.8454	1793.8692	-0.0237	1	28	38	1	MCILTSYAYGDIETR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 125	828.9490	2483.8252	2484.3019	-0.4768	2	28	74	1	MQRSPFFRADEVGSLRPEK + Oxidation (M)
<input checked="" type="checkbox"/> 189	1007.7040	3020.0902	3019.4855	0.6047	2	27	55	1	AGLADRVTGLDLAIEAGADFDRLASAPR + Oxidation (M)
<input checked="" type="checkbox"/> 48	435.7510	869.4874	870.4923	-1.0049	1	27	53	1	KDGVTVPR

MASCOT **SCIENCE** Mascot Search Results

Protein View

Match to: **gi|261342934** Score: **189**
periplasmic protein disulfide isomerase I [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\280808\SSP 6408_RI7_01_467.d\SSP 6408_RI7_01_467.mgf

Nominal mass (M_r): **23134**; Calculated pI value: **5.65**
 NCBI BLAST search of **gi|261342934** against nr
 Unformatted **sequence string** for pasting into other applications

Taxonomy: **Enterobacter cancerogenus ATCC 35316**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **20%**

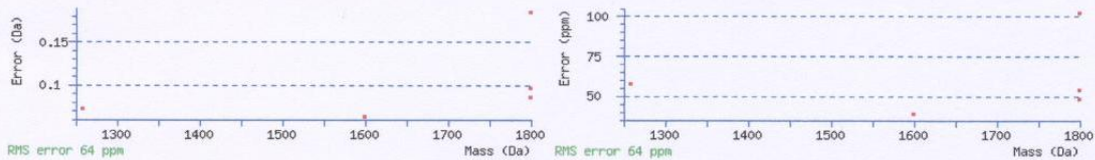
Matched peptides shown in **Bold Red**

1 MKKIWLALAG MILAFSASAA QFTDGKQYIT LDKPVAGEFQ VLEFFSFYCP
 51 HCYQFEVVLH VSDNVKKKLP EGTRKMTKYHV **EFLGPLGKDL TQAWAVAIAL**
 101 **GVEDKVTIAPL** FEAVKQTQTV QNTADIRKVF VDAGVK**GEDY DA**WNSFVVK
 151 SLVAQQEKAA ADLQLQGVPA MFVNGRYQLN MQGMDTSSMD IPVQQYADTV
 201 KYLVEKK

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
78 - 88	630.3790	1258.7434	1258.6710	0.0724	0 K.YHVEFLG PLGK.D (Ions score 42)
89 - 105	600.6850	1799.0332	1798.9465	0.0867	0 K.DLTQAWAVAIALG VEDK.V (Ions score 44)
89 - 105	900.5290	1799.0434	1798.9465	0.0969	0 K.DLTQAWAVAIALG VEDK.V (Ions score 80)
89 - 105	900.5730	1799.1314	1798.9465	0.1849	0 K.DLTQAWAVAIALG VEDK.V (Ions score 50)
137 - 150	800.8990	1599.7834	1599.7205	0.0629	0 K.GEDY DAWNSFVVK.S (Ions score 67)



LOCUS ZP_05970792 207 aa linear BCT 15-OCT-2009
 DEFINITION periplasmic protein disulfide isomerase I [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05970792
 VERSION ZP_05970792.1 GI:261342934
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWMO2000045.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM **Enterobacter cancerogenus ATCC 35316**
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 207)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B., Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C., Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C., Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M., Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 REFERENCE 2 (residues 1 to 207)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H., Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from [ABWMO2000045](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be

http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100105/FtmmSranE... 21/01/2010

found here:

<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

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This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x
Sequencing Technology: 454.
Method: conceptual translation.

FEATURES	Location/Qualifiers
source	1..207 /organism="Enterobacter cancerogenus ATCC 35316" /strain="ATCC 35316" /db_xref="taxon:500639"
Protein	1..207 /product="periplasmic protein disulfide isomerase I" /calculated_mol_wt=22901
Region	24..202 /region_name="DsbA_DsbA" /note="DsbA family, DsbA subfamily; DsbA is a monomeric thiol disulfide oxidoreductase protein containing a redox active CXXC motif imbedded in a TRX fold. It is involved in the oxidative protein folding pathway in prokaryotes, and is the strongest thiol...; cd03019" /db_xref="CDD:48568"
Region	40..197 /region_name="DSBA" /note="DSBA-like thioredoxin domain; pfam01323" /db_xref="CDD:110334"
Site	order(49,51..52,169) /site_type="other" /note="catalytic residues" /db_xref="CDD:48568"
Site	82..84 /site_type="other" /note="hinge region" /db_xref="CDD:48568"
Site	order(89..97,102..121,124..132,138..147) /site_type="other" /note="alpha helical domain" /db_xref="CDD:48568"
CDS	1..207 /locus_tag="EcanA3_020100020887" /coded_by="NZ_ABWM02000045.1:3023..3646" /note="COG0526 Thiol-disulfide isomerase and thioredoxins" /transl_table=11 /db_xref="CDD:138304"

Mascot: <http://www.matrixscience.com/>

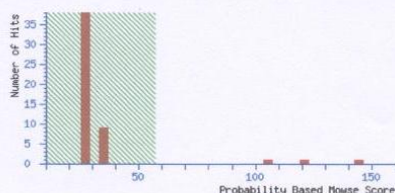
http://www.matrixscience.com/cgi/protein_view.pl?file=./data/20100105/FtmmSranE... 21/01/2010

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6605
 MS data file : D:\Data\Lakshmy\120808\241108\SSP 6605_RC3_01_783.d\SSP 6605_RC3_01_783.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 12:01:23 GMT
 Protein hits : [gi|261340146](#) 2-dehydro-3-deoxyphosphooctonate aldolase [Enterobacter cancerogenus ATCC 35316]
[gi|50582463](#) oxidoreductase [Exiguobacterium acetylicum]
[gi|146311979](#) 2-dehydro-3-deoxyphosphooctonate aldolase [Enterobacter sp. 638]

Probability Based Mowse Score

Ions score is $-10 * \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold p<: Max. number of hits:

Standard scoring: MudPIT scoring Ions score or expect cut-off: Show sub-sets:

Show pop-ups: Suppress pop-ups Sort unassigned: Require bold red:

Select All Select None Search Selected Error tolerant

1.	gi 261340146	Mass: 30917	Score: 144	Queries matched: 3	emPAI: 0.23		
	2-dehydro-3-deoxyphosphooctonate aldolase [Enterobacter cancerogenus ATCC 35316]						
	<input type="checkbox"/> Check to include this hit in error tolerant search						
	Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
	<input checked="" type="checkbox"/> 194	723.3040	2166.8902	2167.0732	-0.1830 0 48	0.32 1	R.AGMATGLAGLFIEARHPDPAK.C + Oxidation (M)
	<input checked="" type="checkbox"/> 248	964.5330	2890.5772	2890.4899	0.0873 0 59	0.023 1	K.VVSIQDINVANDLPPVLPFGMNVLESR.D + Oxidation (M)
	<input checked="" type="checkbox"/> 174	1034.5570	3100.6492	3100.6557	-0.0065 0 39	5.1 1	K.VITDVHEASQAQPVADVVDVIQLPAFLAR.Q
2.	gi 50582463	Mass: 27360	Score: 118	Queries matched: 2	emPAI: 0.12		
	oxidoreductase [Exiguobacterium acetylicum]						
	<input type="checkbox"/> Check to include this hit in error tolerant search						
	Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
	<input checked="" type="checkbox"/> 133	859.3740	1716.7334	1716.8570	-0.1236 0 86	6.4e-05 1	K.YTVITGASSGIGYETAK.L
	<input checked="" type="checkbox"/> 89	677.2430	2028.7072	2028.9752	-0.2681 0 32	25 1	K.SVDLADNQVNDHLYEGLK.E
3.	gi 146311979	Mass: 31019	Score: 107	Queries matched: 2	emPAI: 0.23		
	2-dehydro-3-deoxyphosphooctonate aldolase [Enterobacter sp. 638]						
	<input type="checkbox"/> Check to include this hit in error tolerant search						
	Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
	<input checked="" type="checkbox"/> 194	723.3040	2166.8902	2167.0732	-0.1830 0 48	0.32 1	R.AGMATGLAGLFIEARHPDPAK.C + Oxidation (M)
	<input checked="" type="checkbox"/> 248	964.5330	2890.5772	2890.4899	0.0873 0 59	0.023 1	K.VVSIQDINVANDLPPVLPFGMNVLESR.D + Oxidation (M)
Peptide matches not assigned to protein hits: (no details means no match)							
	Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
	<input checked="" type="checkbox"/> 169	676.8940	2027.6602	2027.0510	0.6092 0 38	2.8 1	IALDQGMDSVFEELLAR
	<input checked="" type="checkbox"/> 85	651.8640	1301.7134	1301.6728	-0.0407 1 34	14 1	VTDLDKAGAWAR
	<input checked="" type="checkbox"/> 221	761.6660	2281.9762	2281.1028	0.8733 2 33	9.5 1	TNFLAGYRLEHRFNDWK
	<input checked="" type="checkbox"/> 138	592.3340	1773.9802	1773.9036	0.0765 0 33	14 1	VSAAEPLIADLFEVDK
	<input checked="" type="checkbox"/> 128	846.1080	2535.3022	2535.1866	0.1156 1 30	37 1	IEAFFRAGYAMHRDITYAIGR + Oxidation (M)
	<input checked="" type="checkbox"/> 246	953.0700	2856.1882	2856.2075	-0.0193 2 30	15 1	VAKCDQCDGDPACVKVCPGTALEFR
	<input checked="" type="checkbox"/> 44	436.2640	870.5134	871.4763	-0.9628 0 29	42 1	EAALAEIR
	<input checked="" type="checkbox"/> 202	740.9910	2219.9512	2220.1678	-0.2166 0 29	28 1	IIDLADYALTLVEPGVTFR
	<input checked="" type="checkbox"/> 206	742.2930	2223.8572	2223.0049	0.8523 0 29	24 1	QMPAGYETQIGEGGMLSGQR
	<input checked="" type="checkbox"/> 161	665.2980	1992.8722	1993.9745	-1.1023 0 28	33 1	ELIGAEDQFFEIGNSLR

MASCOT **SCIENCE** Mascot Search Results

Protein View

Match to: **gi|261340146** Score: **144**
2-dehydro-3-deoxyphosphoconate aldolase [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\241108\SSP 6605_RC3_01_783.d\SSP 6605_RC3_01_783.mgf

Nominal mass (M_r): **30917**; Calculated pI value: **5.63**
 NCBI BLAST search of **gi|261340146** against nr
 Unformatted **sequence string** for pasting into other applications

Taxonomy: **Enterobacter cancerogenus ATCC 35316**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **27%**

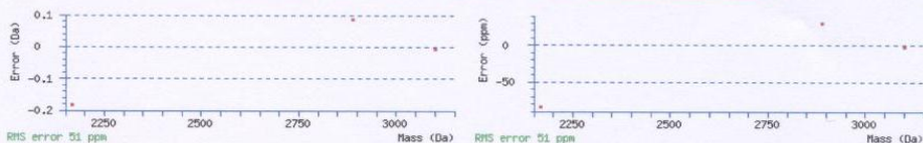
Matched peptides shown in **Bold Red**

1 MKQKVV**SIGD INVANDLPPV LFGGMVLES** RDLAMRICEH YVTVTQKLG**I**
51 PVVFKASFDK ANRSSINSYR GPGLEE**G**MKI FQELKQTFGV KVITD**V**HEAS
101 QAGQVADVVD VIQL**PAFLAR** QTDLVAMAK TGAVINVKRP QFVSPGQMG**N**
151 IVDKFL**EGSN** DRVILDRGA NPYDNLVVD MLGFSVMK**NV** SQGSPV**IFDV**
201 THALQCRD**FF** GASGGR**RAQ** VTELAR**AGMA** TGLAGL**FIEA** HFD**PANAKCD**
251 GPSALPLDKL E**PF**LKQIKAI DDLVRS**PDEL** DTS**N**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
5 - 31	964.5330	2890.5772	2890.4899	0.0873	0 K.VV SIGD INVANDLPPV LFGGMVLES .D Oxidation (M) (Ions_score 59)
92 - 120	1034.5570	3100.6492	3100.6557	-0.0065	0 K.VITD V HEASQAPVADVVDVIQL PAFLAR .Q (Ions_score 39)
227 - 248	723.3040	2166.8902	2167.0732	-0.1830	0 R.AGMATGLAGL FIEA HFD PANAK .C Oxidation (M) (Ions_score 48)



LOCUS ZP_05968004 284 aa linear BCT 15-OCT-2009
 DEFINITION 2-dehydro-3-deoxyphosphoconate aldolase [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05968004
 VERSION ZP_05968004.1 GI:261340146
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession **NZ_ABNM02000008.1**
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM **Enterobacter cancerogenus ATCC 35316**
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 284)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B., Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C., Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C., Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M., Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 REFERENCE 2 (residues 1 to 284)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H., Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 COMMENT WGS **REFSEQ**: This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from **ABNM02000008**.
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).
 We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where

http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100105/FtmmSratE.... 21/01/2010

possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project (<http://nihroadmap.nih.gov/hmp/>).

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

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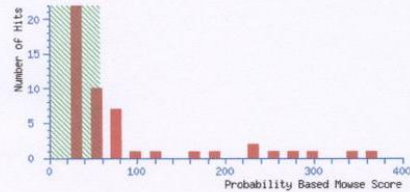
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 6711
MS data file : D:\Data\Lakshmy\120808\SSP 6711_RB5_01_717.d\SSP 6711_RB5_01_717.mgf
Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 12:02:31 GMT
Protein hits :
 gi|157147228 elongation factor Tu [Citrobacter koseri ATCC BAA-895]
 gi|261819559 translation elongation factor Tu [Pectobacterium wasabiae WPP163]
 gi|110807172 elongation factor Tu [Shigella flexneri 5 str. 8401]
 gi|16272522 elongation factor Tu [Haemophilus influenzae Rd KW20]
 gi|261341842 elongation factor Tu [Enterobacter cancerogenus ATCC 35316]
 gi|259906922 Elongation factor Tu-A [Erwinia pyrifoliae Epl/96]
 gi|37524441 elongation factor Tu [Photobacterium luminescens subsp. laumondii TT01]
 gi|237809513 translation elongation factor Tu [Tolomonas auensis DSM 9187]
 gi|270341211 elongation factor Tu [Pectobacterium carotovorum subsp. carotovorum]
 gi|78358029 elongation factor Tu [Desulfovibrio desulfuricans subsp. desulfuricans str. G20]
 gi|24371815 elongation factor Tu [Shewanella oneidensis MR-1]
 gi|78189809 elongation factor Tu [Chlorobium chlorochromatii CaD3]
 gi|119468170 protein chain elongation factor EF-Tu; GTP-binding factor [Alteromonadales bacterium TW-7]
 gi|2369692 elongation factor Ef-Tu [Buchnera aphidicola]
 gi|88861465 protein chain elongation factor EF-Tu; GTP-binding factor [Pseudoalteromonas tunicata D2]
 gi|19551739 elongation factor Tu [Corynebacterium glutamicum ATCC 13032]
 gi|76786395 elongation factor Tu [endosymbiont of Haematomyzus elephantis]
 gi|270341139 elongation factor Tu [Aeromonas molluscorum]
 gi|57238897 elongation factor Tu [Ehrlichia ruminantium str. Welgevonden]
 gi|15839217 elongation factor Tu [Xylella fastidiosa 9a5c]
 gi|119947040 elongation factor Tu [Psychromonas ingrahamii 37]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: **Peptide Summary** Help

Significance threshold $p < 0.05$ Max. number of hits **AUTO**

Standard scoring MudPIT scoring Ions score or expect cut-off **0** Show sub-sets **0**

Show pop-ups Suppress pop-ups Sort unassigned **Decreasing Score** Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|157147228](#) Mass: 43389 Score: 365 Queries matched: 10 emPAI: 0.67
 elongation factor Tu [Citrobacter koseri ATCC BAA-895]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 78	652.2320	1302.4494	1302.7759	-0.3264	0	62	0.014	1	K.TLTLAAITTVLAK.T
<input checked="" type="checkbox"/> 136	604.9370	1811.7892	1811.9274	-0.1382	0	39	2.8	1	K.MVVTLIHPIAMDGLR.F + 2 Oxidation (M)
<input checked="" type="checkbox"/> 164	982.9230	1963.8314	1963.9527	-0.1213	0	61	0.019	1	R.ELLSQYDFPGDDTFIVR.G
<input checked="" type="checkbox"/> 166	989.4440	1976.8734	1977.0095	-0.1360	0	40	2.6	1	K.IIELAGYLDYIPEPER.A
<input checked="" type="checkbox"/> 167	989.7760	1977.5374	1977.0095	0.5280	0	(25)	99	1	K.IIELAGYLDYIPEPER.A
<input checked="" type="checkbox"/> 194	706.3230	2115.9472	2116.1568	-0.2097	0	(33)	11	1	R.AIDKPFLLPIEDVFSISGR.G
<input checked="" type="checkbox"/> 195	706.3510	2116.0312	2116.1568	-0.1257	0	(48)	0.42	1	R.AIDKPFLLPIEDVFSISGR.G
<input checked="" type="checkbox"/> 196	1059.0410	2116.0674	2116.1568	-0.0894	0	52	0.15	1	R.AIDKPFLLPIEDVFSISGR.G
<input checked="" type="checkbox"/> 241	860.0200	2577.0382	2577.2190	-0.1808	0	54	0.066	1	R.TTDVGTIPELPEGVEMVMPGDNK.M + 2 Oxidation (M)
<input checked="" type="checkbox"/> 246	926.6500	2776.9282	2776.3194	0.6088	0	57	0.026	1	K.NMITGAAGDGLLVVAATDGPMPQTR.E + 3 Oxidation (M)

Proteins matching the same set of peptides:
[gi|157148907](#) Mass: 45003 Score: 365 Queries matched: 10
 elongation factor Tu [Citrobacter koseri ATCC BAA-895]

2. [gi|261819559](#) Mass: 43374 Score: 338 Queries matched: 9 emPAI: 0.55
 translation elongation factor Tu [Pectobacterium wasabiae WPP163]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
78	652.2320	1302.4494	1302.7759	-0.3264	0	62	0.014	1	K.TTLTAIITTVLAK.T
99	737.4140	1472.8134	1472.8755	-0.0621	0	14	1.2e+03	6	R.QVGVFFIIVFLNK.C
136	604.9370	1811.7892	1811.9274	-0.1382	0	39	2.8	1	K.MVVTLIHPIAMDDGLR.F + 2 Oxidation (M)
164	982.9230	1963.8314	1963.9527	-0.1213	0	61	0.019	1	R.ELLSQYDFPGDDTFIVR.G
194	706.3230	2115.9472	2116.1568	-0.2097	0	(33)	11	1	R.AIDKPFLLPIEDVFSISGR.G
195	706.3510	2116.0312	2116.1568	-0.1257	0	(48)	0.42	1	R.AIDKPFLLPIEDVFSISGR.G
196	1059.0410	2116.0674	2116.1568	-0.0894	0	52	0.15	1	R.AIDKPFLLPIEDVFSISGR.G
241	860.0200	2577.0382	2577.2190	-0.1808	0	54	0.066	1	R.TTDVTGTIELPEGVEMVMPGDNK.M + 2 Oxidation (M)
246	926.6500	2776.9282	2776.3194	0.6088	0	57	0.026	1	K.NMITGAAQMDGAILVVAATDGPMPQTR.E + 3 Oxidation (M)

3. **gi|110807172** Mass: 43429 Score: 304 Queries matched: 8 emPAI: 0.44
 elongation factor Tu [Shigella flexneri 5 str. 8401]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
78	652.2320	1302.4494	1302.7759	-0.3264	0	62	0.014	1	K.TTLTAIITTVLAK.T
136	604.9370	1811.7892	1811.9274	-0.1382	0	39	2.8	1	K.MVVTLIHPIAMDDGLR.F + 2 Oxidation (M)
164	982.9230	1963.8314	1963.9527	-0.1213	0	61	0.019	1	R.ELLSQYDFPGDDTFIVR.G
194	706.3230	2115.9472	2116.1568	-0.2097	0	(33)	11	1	R.AIDKPFLLPIEDVFSISGR.G
195	706.3510	2116.0312	2116.1568	-0.1257	0	(48)	0.42	1	R.AIDKPFLLPIEDVFSISGR.G
196	1059.0410	2116.0674	2116.1568	-0.0894	0	52	0.15	1	R.AIDKPFLLPIEDVFSISGR.G
241	860.0200	2577.0382	2576.2350	0.8032	0	32	11	2	R.TTDVTGTIELPEGVEMVMPGDNK.M + 2 Oxidation (M)
246	926.6500	2776.9282	2776.3194	0.6088	0	57	0.026	1	K.NMITGAAQMDGAILVVAATDGPMPQTR.E + 3 Oxidation (M)

4. **gi|16272522** Mass: 43443 Score: 286 Queries matched: 7 emPAI: 0.55
 elongation factor Tu [Haemophilus influenzae Rd KW20]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
78	652.2320	1302.4494	1302.7759	-0.3264	0	62	0.014	1	K.TTLTAIITTVLAK.H
164	982.9230	1963.8314	1963.9527	-0.1213	0	61	0.019	1	R.ELLSQYDFPGDDTFIVR.G
194	706.3230	2115.9472	2116.1205	-0.1733	0	(33)	11	1	R.AIDKPFLLPIEDVFSISGR.G
195	706.3510	2116.0312	2116.1205	-0.0893	0	(48)	0.42	1	R.AIDKPFLLPIEDVFSISGR.G
196	1059.0410	2116.0674	2116.1205	-0.0530	0	52	0.15	1	R.AIDKPFLLPIEDVFSISGR.G
241	860.0200	2577.0382	2577.2190	-0.1808	0	54	0.066	1	R.TTDVTGTIELPEGVEMVMPGDNK.M + 2 Oxidation (M)
246	926.6500	2776.9282	2776.3194	0.6088	0	57	0.026	1	K.NMITGAAQMDGAILVVAATDGPMPQTR.E + 3 Oxidation (M)

Proteins matching the same set of peptides:
gi|53733337 Mass: 43469 Score: 286 Queries matched: 7
 COG0050: GTPases - translation elongation factors [Haemophilus influenzae R2866]
gi|145629756 Mass: 43442 Score: 286 Queries matched: 7
 tRNA-dihydrouridine synthase A [Haemophilus influenzae 22.1-21]
gi|145640555 Mass: 43471 Score: 286 Queries matched: 7
 hypothetical protein CGSHIR3021_10860 [Haemophilus influenzae R3021]
gi|15603222 Mass: 43442 Score: 284 Queries matched: 7
 elongation factor Tu [Pasteurella multocida subsp. multocida str. Pm70]
gi|42630895 Mass: 43386 Score: 284 Queries matched: 7
 COG0050: GTPases - translation elongation factors [Haemophilus influenzae R2866]
gi|251793286 Mass: 43438 Score: 284 Queries matched: 7
 translation elongation factor Tu [Aggregatibacter aphrophilus NJ8700]

5. **gi|261341842** Mass: 38502 Score: 246 Queries matched: 7 emPAI: 0.51
 elongation factor Tu [Enterobacter cancerogenus ATCC 35316]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
78	652.2320	1302.4494	1302.7759	-0.3264	0	62	0.014	1	K.TTLTAIITTVLAK.T
99	737.4140	1472.8134	1472.8755	-0.0621	0	14	1.2e+03	6	R.QVGVFFIIVFLNK.C
164	982.9230	1963.8314	1963.9527	-0.1213	0	61	0.019	1	R.ELLSQYDFPGDDTFIVR.G
194	706.3230	2115.9472	2116.1568	-0.2097	0	(33)	11	1	R.AIDKPFLLPIEDVFSISGR.G
195	706.3510	2116.0312	2116.1568	-0.1257	0	(48)	0.42	1	R.AIDKPFLLPIEDVFSISGR.G
196	1059.0410	2116.0674	2116.1568	-0.0894	0	52	0.15	1	R.AIDKPFLLPIEDVFSISGR.G
246	926.6500	2776.9282	2776.3194	0.6088	0	57	0.026	1	K.NMITGAAQMDGAILVVAATDGPMPQTR.E + 3 Oxidation (M)

6. **gi|259906922** Mass: 43465 Score: 233 Queries matched: 7 emPAI: 0.34
 Elongation factor Tu-A [Erwinia pyrifoliae Epl/96]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
78	652.2320	1302.4494	1302.7759	-0.3264	0	62	0.014	1	K.TTLTAIITTVLAK.T
99	737.4140	1472.8134	1472.8755	-0.0621	0	14	1.2e+03	6	R.QVGVFFIIVFLNK.C
164	982.9230	1963.8314	1963.9527	-0.1213	0	47	0.45	4	R.DLLTQYDFPGDDTFIVR.G
194	706.3230	2115.9472	2116.1568	-0.2097	0	(33)	11	1	R.AIDKPFLLPIEDVFSISGR.G
195	706.3510	2116.0312	2116.1568	-0.1257	0	(48)	0.42	1	R.AIDKPFLLPIEDVFSISGR.G
196	1059.0410	2116.0674	2116.1568	-0.0894	0	52	0.15	1	R.AIDKPFLLPIEDVFSISGR.G
246	926.6500	2776.9282	2776.3194	0.6088	0	57	0.026	1	K.NMITGAAQMDGAILVVAATDGPMPQTR.E + 3 Oxidation (M)

7. **gi|37524441** Mass: 43321 Score: 232 Queries matched: 6 emPAI: 0.44
 elongation factor Tu [Photobacterium luminescens subsp. laumondii T101]
 Check to include this hit in error tolerant search

Mascot Search Results

Protein View

Match to: **gi|261341842** Score: **246**
elongation factor Tu [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\SSP 6711_RB5_01_717.d\SSP 6711_RB5_01_717.mgf

Nominal mass (M_n): **38502**; Calculated pI value: **5.17**
 NCBI BLAST search of **gi|261341842** against nr
 Unformatted **sequence_string** for pasting into other applications

Taxonomy: **Enterobacter cancerogenus ATCC 35316**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **25%**

Matched peptides shown in **Bold Red**

1 MSKEKFERTK PHVNVGTIGH VDHGK**TTLTA AITTVLAKTY** GGAARAFDQI
51 DNAPSEKARG ITINTSHVEY DTPTRHYAHV DCPGHADYVK **NMITGAAQMD**
101 **GAILVVAATD GMPFQTR**EHI LLGRQVGVVF IIVFLNKCDM VDDELELLELV
151 EMEVRELLSQ **YDFPGDDTPI VRG**SALKALE GEAEWEKII ELAGFLDSYI
201 PEPERAI**DKP FLLPIEDVFS ISGR**RTVVTG RVERGLIKVG EEVEIVGIKE
251 TAKSTCTGVE MFRKLLDEGR AGENVGVLLR GIKREIERG QVLAKPGSIK
301 PHTKFESEVY ILSKDEGGRH TPFKGYRQP FYFRTDVTG TIELPEG

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
26 - 38	652.2320	1302.4494	1302.7759	-0.3264	0	K.TTLTA AITTVLAK.T (Ions_score 62)
91 - 117	926.6500	2776.9282	2776.3194	0.6088	0	K.NMITGAAQMD GAILVVAATD GMPQTR.E 3 Oxidation (M) (Ions_score 57)
125 - 137	737.4140	1472.8134	1472.8755	-0.0621	0	R.QVGVFFIIVFLNK.C (Ions_score 14)
156 - 172	982.9230	1963.8314	1963.9527	-0.1213	0	R.ELLSQYDFPGDDTPIVR.G (Ions_score 61)
206 - 224	706.3230	2115.9472	2116.1568	-0.2097	0	R.AIDKPFLLPIEDVFSISGR.G (Ions_score 33)
206 - 224	706.3510	2116.0312	2116.1568	-0.1257	0	R.AIDKPFLLPIEDVFSISGR.G (Ions_score 48)
206 - 224	1059.0410	2116.0674	2116.1568	-0.0894	0	R.AIDKPFLLPIEDVFSISGR.G (Ions_score 52)

Error: try setting browser cache to automatic.

Error: try setting browser cache to automatic.

LOCUS ZP_05969700 347 aa linear BCT 15-OCT-2009
 DEFINITION elongation factor Tu [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05969700
 VERSION ZP_05969700.1 GI:261341842
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000029.1](#)
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 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM Enterobacter cancerogenus ATCC 35316
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 347)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 347)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM02000029](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be
 aware that the annotation is done automatically with little or no
 manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA
 gene: Z36078, the 16S rDNA gene of a related strain of Enterobacter
 cancerogenus, not the sequenced strain), is a member of the
 Proteobacteria division of the domain bacteria and has been
 isolated from human feces. The sequenced strain was obtained from
 ATCC (ATCC 35316).

We have performed one round of automated sequence improvement

http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100105/FtmmSratR... 21/01/2010

(pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project (<http://nihroadmap.nih.gov/hmp/>).

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. COMPLETENESS: incomplete on the carboxy end. Method: conceptual translation.

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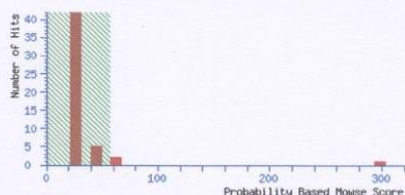
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 7602
 MS data file : D:\Data\Lakshmy\120808\260808\SSP 7602_RH1_01_413.d\SSP 7602_RH1_01_413.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 12:05:52 GMT
 Protein hits : [gi|146311681](#) acetoin reductase [Enterobacter sp. 638]
 [gi|2285882](#) Aalpha subunit of acetyl-CoA carboxylase [Escherichia coli W3110]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)
 Significance threshold p< Max. number of hits
 Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets
 Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|146311681](#) Mass: 26942 Score: 298 Queries matched: 5 emPAI: 0.42
 acetoin reductase [Enterobacter sp. 638]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 104	759.0050	1515.9954	1515.8198	0.1757	0	97	4.7e-06	1	K.GVIWGIQAIDAIFR.K
<input checked="" type="checkbox"/> 128	564.6070	1690.7992	1689.8686	0.9306	1	41	2.1	1	K.VDVSNREQVFAAVEK.A
<input checked="" type="checkbox"/> 132	571.8780	1712.6122	1712.7893	-0.1772	0	(53)	0.089	1	K.DGFAVALADYNEETAK.A
<input checked="" type="checkbox"/> 133	857.4090	1712.8034	1712.7893	0.0141	0	88	4e-05	1	K.DGFAVALADYNEETAK.A
<input checked="" type="checkbox"/> 234	802.0650	2403.1732	2404.1329	-0.9597	0	72	0.0014	1	K.IINACSQAGHTGNPELAVYSSK.F

Proteins matching the same set of peptides:
[gi|261341164](#) Mass: 26884 Score: 298 Queries matched: 5
 acetoin reductase [Enterobacter cancerogenus ATCC 35316]

2. [gi|2285882](#) Score: 59 Queries matched: 2
 Aalpha subunit of acetyl-CoA carboxylase [Escherichia coli W3110]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 111	529.6000	1585.7782	1586.8780	-1.0998	0	32	18	3	K.LIDTSIPEPLGGAHR.N
<input type="checkbox"/> 143	599.9910	1796.9512	1796.9454	0.0058	1	27	59	3	K.SADKAPLAAEAMGIAPR.L + Oxidation (M)

Proteins matching the same set of peptides:
[gi|15799867](#) Score: 59 Queries matched: 2
[gi|16759222](#) Score: 59 Queries matched: 2
[gi|26246132](#) Score: 59 Queries matched: 2
[gi|74310805](#) Score: 59 Queries matched: 2
[gi|82542784](#) Score: 59 Queries matched: 2
[gi|91209255](#) Score: 59 Queries matched: 2
[gi|161504649](#) Score: 59 Queries matched: 2
[gi|170769662](#) Score: 59 Queries matched: 2
[gi|187731373](#) Score: 59 Queries matched: 2
[gi|213029802](#) Score: 59 Queries matched: 2
[gi|213416583](#) Score: 59 Queries matched: 2
[gi|213586740](#) Score: 59 Queries matched: 2
[gi|224582080](#) Score: 59 Queries matched: 2

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 111	529.6000	1585.7782	1586.8780	-1.0999	0	40	3	1	LIDTSIPEPLGGAHR

MASCOT **SCIENCE** Mascot Search Results

Protein View

Match to: **gi|261341164** Score: **298**
acetoin reductase [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\260808\SSP 7602_RH1_01_413.d\SSP 7602_RH1_01_413.mgf

Nominal mass (M_r): **26884**; Calculated pI value: **6.17**
 NCBI BLAST search of **gi|261341164** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **26%**

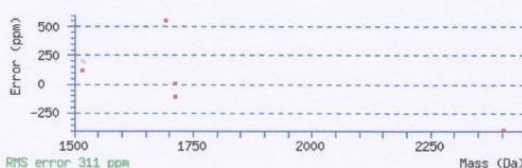
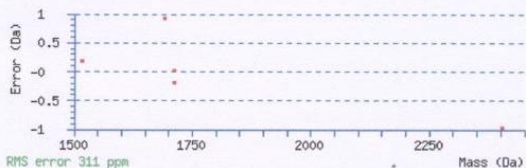
Matched peptides shown in **Bold Red**

- 1 MQKVALVTGS GQGIGKAIAL RLVK**DGFAVA** **LADYNEETAK** AVAEIIVRSQ
- 51 GKAVAVKVDV **SNREQVFAAV** EKARTALGGF NVIVNAGVA PSTPIESITP
- 101 DIVDKVYNIN **VKGVIWGIIQA** **AIDAFRKEGH** GGK**IINACSQ** **AGHTGNPELA**
- 151 **VYSSSK**FAVR GLTQTAARDL APLGITVNAY CPGIVKTPMW AIEDRQVSEA
- 201 AGKPLGYGTE TFAKRITLGR LSEPEDVAAC VSYLAGPDSQ YMTQSQLLID
- 251 GGMVFN

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
25 - 40	571.8780	1712.6122	1712.7893	-0.1772	0	K.DGFAVALADYNEETAK.A (Ions score 53)
25 - 40	857.4090	1712.8034	1712.7893	0.0141	0	K.DGFAVALADYNEETAK.A (Ions score 88)
58 - 72	564.6070	1690.7992	1689.8686	0.9306	1	K.VDVSNREQVFAAVEK.A (Ions score 41)
113 - 126	759.0050	1515.9954	1515.8198	0.1757	0	K.GVIWGIIQAIDAIFR.K (Ions score 97)
134 - 156	802.0650	2403.1732	2404.1329	-0.9597	0	K.IINACSQAGHTGNPELAVYSSK.F (Ions score 72)



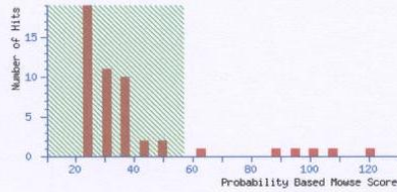
LOCUS ZP_05969022 256 aa linear BCT 15-OCT-2009
 DEFINITION acetoin reductase [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05969022
 VERSION ZP_05969022.1 GI:261341164
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABW02000020.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 256)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B., Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C., Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C., Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M., Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 REFERENCE 2 (residues 1 to 256)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H., Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 COMMENT WGS [REFSEQ](#): This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from [ABW02000020](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 7707
MS data file : D:\Data\Lakshmy\120808\280808\SSP 7707_RI2_01_457.d\SSP 7707_RI2_01_457.mgf
Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 12:11:18 GMT
Protein hits : [gi|157368709](#) inorganic pyrophosphatase [Serratia proteamaculans 568]
[gi|269137738](#) Inorganic pyrophosphatase [Edwardsiella tarda EIB202]
[gi|6453807](#) ProA [Shigella flexneri]
[gi|1943414](#) Chain A, Inorganic Pyrophosphatase
[gi|258634603](#) gamma-glutamyl phosphate reductase [Pantoea sp. At-9b]
[gi|2781250](#) Chain A, Structure Of Inorganic Pyrophosphatase Mutant D42n

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As **Peptide Summary** [Help](#)

Significance threshold $p < 0.05$ Max. number of hits **AUTO**

Standard scoring MudPIT scoring Ions score or expect cut-off **0** Show sub-sets **0**

Show pop-ups Suppress pop-ups Sort unassigned **Decreasing Score** Require bold red

Error tolerant

1. [gi|157368709](#) Mass: 19679 Score: 120 Queries matched: 4
 inorganic pyrophosphatase [Serratia proteamaculans 568]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 68	518.2850	1034.5554	1034.5396	0.0158	0	47	0.62	1	K.AEIIASFER.A
<input checked="" type="checkbox"/> 73	578.3370	1154.6594	1154.6183	0.0412	0	33	13	3	K.DVNDLELLK.A
<input checked="" type="checkbox"/> 130	547.9490	1640.8252	1640.8046	0.0206	1	40	2.8	1	K.YEIDKDTGALFVDR.F
<input checked="" type="checkbox"/> 131	821.4700	1640.9254	1640.8046	0.1209	1	(15) 8.4e+02	1	1	K.YEIDKDTGALFVDR.F

2. [gi|269137738](#) Mass: 19766 Score: 107 Queries matched: 4
 Inorganic pyrophosphatase [Edwardsiella tarda EIB202]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 68	518.2850	1034.5554	1034.5396	0.0158	0	47	0.62	1	K.AEIIASFER.A
<input checked="" type="checkbox"/> 85	464.2630	1389.7672	1389.6830	0.0842	0	20	3.1e+02	3	R.AQIAHFFHYK.D
<input checked="" type="checkbox"/> 130	547.9490	1640.8252	1640.8046	0.0206	1	40	2.8	1	K.YEIDKDTGALFVDR.F
<input checked="" type="checkbox"/> 131	821.4700	1640.9254	1640.8046	0.1209	1	(15) 8.4e+02	1	1	K.YEIDKDTGALFVDR.F

3. [gi|6453807](#) Mass: 22056 Score: 99 Queries matched: 2 emPAI: 0.15
 ProA [Shigella flexneri]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 77	648.8340	1295.6534	1295.6431	0.0103	0	70	0.003	1	R.GPMGLEALTYK.W + Oxidation (M)
<input checked="" type="checkbox"/> 183	619.9010	1856.6812	1856.8905	-0.2093	0	29	24	1	R.FVNEVDSSAVTVNASTR.F

Proteins matching the same set of peptides:
[gi|15799948](#) Mass: 45102 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli O157:H7 EDL933]
[gi|16128229](#) Mass: 45008 Score: 99 Queries matched: 2
 gamma-glutamylphosphate reductase [Escherichia coli str. K-12 substr. MG1655]
[gi|24111735](#) Mass: 45010 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Shigella flexneri 2a str. 301]
[gi|26246288](#) Mass: 45012 Score: 99 Queries matched: 2

found here:

<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x
Sequencing Technology: 454.
Method: conceptual translation.

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FEATURES             Location/Qualifiers
    source            1..256
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                     /strain="ATCC 35316"
                     /db_xref="taxon:500639"
    Protein            1..256
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                     /calculated_mol_wt=26595
    Region            1..256
                     /region_name="PRK08643"
                     /note="acetoin reductase; Validated; PRK08643"
                     /db_xref="CDD:136931"
    Region            1..253
                     /region_name="NADB_Rossmann"
                     /note="Rossmann-fold NAD(P)(+)-binding proteins; c109931"
                     /db_xref="CDD:143725"
    CDS               1..256
                     /locus_tag="EcanA3_020100011911"
                     /coded_by="NZ_ABWM02000020.1:70258..71028"
                     /note="COG1028 Dehydrogenases with different specificities
                     (related to short-chain alcohol dehydrogenases)"
                     /transl_table=11
                     /db_xref="CDD:136931"

```

Mascot: <http://www.matrixscience.com/>

gamma-glutamyl phosphate reductase [Escherichia coli CFT073]
 gi|82542846 Mass: 45013 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Shigella boydii Sb227]
 gi|82775922 Mass: 44980 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Shigella dysenteriae Sd197]
 gi|91209332 Mass: 44954 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli UTI89]
 gi|110640479 Mass: 44959 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli 536]
 gi|1157147174 Mass: 44879 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Citrobacter koseri ATCC BAA-895]
 gi|1157157629 Mass: 45008 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli E24377A]
 gi|170682591 Mass: 45038 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli SMS-3-5]
 gi|170767876 Mass: 45086 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia albertii TW07627]
 gi|188534718 Mass: 45397 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Erwinia tasmaniensis Et1/99]
 gi|194432663 Mass: 44978 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Shigella dysenteriae 1012]
 gi|194438160 Mass: 45038 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli 101-1]
 gi|215485391 Mass: 45012 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli O127:H6 str. E2348/69]
 gi|218550050 Mass: 44924 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia fergusonii ATCC 35469]
 gi|218688135 Mass: 45012 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli ED1a]
 gi|218693726 Mass: 45039 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli 55989]
 gi|218698813 Mass: 45038 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli IAI39]
 gi|218703559 Mass: 44996 Score: 99 Queries matched: 2
 gamma-glutamyl phosphate reductase [Escherichia coli UMNO26]
 gi|237730224 Mass: 45044 Score: 99 Queries matched: 2
 gamma-glutamylphosphate reductase [Citrobacter sp. 30_2]
 gi|260866441 Mass: 44994 Score: 99 Queries matched: 2
 gamma-glutamylphosphate reductase [Escherichia coli O111:H- str. 11128]
 gi|281177456 Mass: 45000 Score: 99 Queries matched: 2
 gamma-glutamylphosphate reductase [Escherichia coli SE15]
 gi|156935257 Mass: 45461 Score: 97 Queries matched: 2
 gamma-glutamyl phosphate reductase [Enterobacter sakazakii ATCC BAA-894]
 gi|259909344 Mass: 45425 Score: 97 Queries matched: 2
 Gamma-glutamyl phosphate reductase [Erwinia pyrifoliae Epl/96]
 gi|260596647 Mass: 45456 Score: 97 Queries matched: 2
 Gamma-glutamyl phosphate reductase [Cronobacter turicensis]
 gi|261340945 Mass: 45230 Score: 97 Queries matched: 2
 gamma-glutamyl phosphate reductase [Enterobacter cancerogenus ATCC 35316]

4. gi|1943414 Score: 93 Queries matched: 3
 Chain A, Inorganic Pyrophosphatase
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
73	578.3370	1154.6594	1154.6183	0.0412	0	33	13	3	K.DVNDLPELLK.A
85	464.2630	1389.7672	1389.6830	0.0842	0	20	3.1e+02	3	K.AQIAHFFEHYK.D
130	547.9490	1640.8252	1640.8046	0.0206	1	40	2.8	1	K.YEIDKESGALFVDR.F

Proteins matching the same set of peptides:
 gi|2554846 Score: 93 Queries matched: 3
 gi|2781252 Score: 93 Queries matched: 3
 gi|2781254 Score: 93 Queries matched: 3
 gi|15804817 Score: 93 Queries matched: 3
 gi|26251130 Score: 93 Queries matched: 3
 gi|130065484 Score: 93 Queries matched: 3
 gi|157831975 Score: 93 Queries matched: 3
 gi|170766732 Score: 93 Queries matched: 3
 gi|215489572 Score: 93 Queries matched: 3
 gi|218551521 Score: 93 Queries matched: 3
 gi|256019879 Score: 93 Queries matched: 3

5. gi|258634603 Mass: 45492 Score: 88 Queries matched: 2 empAI: 0.07
 gamma-glutamyl phosphate reductase [Pantoea sp. At-9b]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
77	648.8340	1295.6534	1295.6431	0.0103	0	70	0.003	1	R.GPMLGALTTYK.W + Oxidation (M)
183	619.9010	1856.6812	1855.9064	0.7747	0	18	3.1e+02	3	R.FVNQVDSAVVYVNASTR.F

6. gi|2781250 Score: 61 Queries matched: 3
 Chain A, Structure Of Inorganic Pyrophosphatase Mutant D42n
 Check to include this hit in error tolerant search

Mascot Search Results

Protein View

Match to: **gi|261340945** Score: **97**
gamma-glutamyl phosphate reductase [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\280808\SSP 7707_RI2_01_457.d\SSP 7707_RI2_01_457.mgf

Nominal mass (M_n): **45230**; Calculated pI value: **5.60**
 NCBI BLAST search of **gi|261340945** against nr
 Unformatted [sequence string](#) for pasting into other applications
 Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)
 Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **6%**

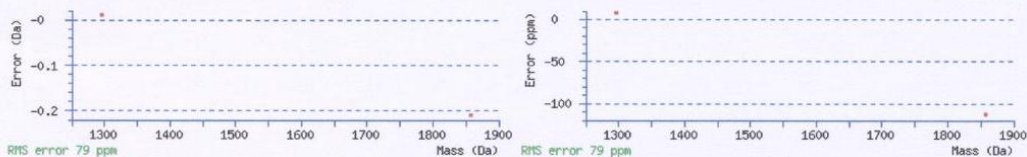
Matched peptides shown in **Bold Red**

1 MLEQMGAAAK AASYKLALLS SREKNRVLEK IADYLESQAS EILLANEQDL
51 LEARRNGLSE AMLDRLALNP ARKLSIADDV RQVCNLADPV GQVIDGGLLD
101 SGLRLERRRRV PLGVIGVIYE ARPNTVDVA SLCLKTGNAI IIRGGKETWR
151 TNAATVVKVIQ QALEECGLPA GAVQAIESP RALVNEMLRM DKYIDMLIPR
201 GGAGLHKLCR EQSTIPVITG GIGVCHILVD DTAEDIDPALK IIVNAKTQRP
251 STCNTVETLL VHQGIARTFL PALSQQAQS GVTLHADASA LALLKDGPPAN
301 VVPVKAEQYD DEFSLDLNV KVVADLDDAI AHIREHGTQH SDALLTRTLR
351 NANRFVNEVD **SSAVYVNASTR** RFTDGGQFGL GAIEVAVSTQK LHARG**PMGLE**
401 ALTTYKWIGF GDDTIRA

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
355 - 371	619.9010	1856.6812	1856.8905	-0.2093	0 R.FVNEVDSSAVYVNASTR.F (Ions score 29)
395 - 406	648.8340	1295.6534	1295.6431	0.0103	0 R.GPMGLEALTTYK.W Oxidation (M) (Ions score 70)



LOCUS ZP_05968803 417 aa linear BCT 15-OCT-2009
 DEFINITION gamma-glutamyl phosphate reductase [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05968803
 VERSION ZP_05968803.1 GI:261340945
 DBLINK Project:20663
 DBSOURCE REFSEQ: accession NZ_ABWM02000017.1
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM Enterobacter cancerogenus ATCC 35316
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales; Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 417)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B., Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C., Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C., Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M., Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 REFERENCE 2 (residues 1 to 417)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H., Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R. and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington University School of Medicine, 4444 Forest Park, St. Louis, MO 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of whole genome shotgun sequences. The reference sequence was derived from ABWM02000017.
 Annotation was added by the NCBI Prokaryotic Genomes Automatic Annotation Pipeline Group. Information about the Pipeline can be

http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100105/FtmmSruT... 21/01/2010

found here:

<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

FEATURES	Location/Qualifiers
source	1..417 /organism="Enterobacter cancerogenus ATCC 35316" /strain="ATCC 35316" /db_xref="taxon:500639"
Protein	1..417 /product="gamma-glutamyl phosphate reductase" /EC_number="1.2.1.41" /calculated_mol_wt=44778
Region	22..412 /region_name="ALDH F18-19 ProA-GPR" /note="Gamma-glutamyl phosphate reductase (GPR), aldehyde dehydrogenase families 18 and 19; cd07079" /db_xref="CDD:143398"
Region	22..406 /region_name="proA" /note="gamma-glutamyl phosphate reductase; TIGR00407" /db_xref="CDD:129501"
Site	253 /site_type="other" /note="putative catalytic cysteine" /db_xref="CDD:143398"
CDS	1..417 /gene="proA" /locus_tag="EcanA3_020100010808" /coded_by="NZ_ABWM02000017.1:31515..32768" /note="COG0014 Gamma-glutamyl phosphate reductase" /transl_table=11 /db_xref="CDD:134165"

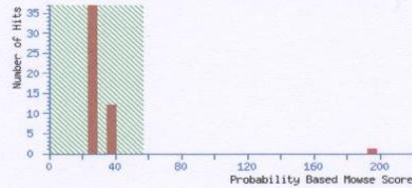
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@sunn.ac.uk
 Search title : SSP 8205
 MS data file : D:\Data\Lakshmy\120808\280808\SSP 8205_RJ23_01_550.d\SSP 8205_RJ23_01_550.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 12:12:48 GMT
 Protein hits : [gi|261339072](#) YfaZ family protein [Enterobacter cancerogenus ATCC 35316]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|261339072](#) Mass: 18559 Score: 195 Queries matched: 3 **emPAI**: 0.65
 YfaZ family protein [Enterobacter cancerogenus ATCC 35316]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 95	648.8060	1295.5974	1295.6180	-0.0205	0	55	0.084	1	K.NFTNLNLEMK.S + Oxidation (M)
<input checked="" type="checkbox"/> 106	721.3580	1440.7014	1440.6885	0.0130	0	61	0.022	1	K.SSSGLYAESHWLK.N
<input checked="" type="checkbox"/> 185	678.2780	2031.8122	2030.9810	0.8311	0	79	0.00025	1	K.DGRFNHTLIDGAYVGGVSP.-

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 82	554.8700	1107.7254	1106.5720	1.1535	1	38	8.2	1	LAAGEFFSKR
<input checked="" type="checkbox"/> 74	516.2960	1030.5774	1030.5546	0.0228	0	37	6.5	1	VLELDLSKR
<input checked="" type="checkbox"/> 88	590.3090	1178.6034	1178.6546	-0.0512	2	33	14	1	YEELLKRRK
<input checked="" type="checkbox"/> 216	737.6970	2210.0692	2210.0062	0.0630	2	32	14	1	NGDLMKEWTFEVRANAQA + Oxidation (M)
<input checked="" type="checkbox"/> 225	747.0420	2238.1042	2237.0646	1.0396	1	32	15	1	GLGPAAGRRGGGGGAGSGGTGGGGAAGR
<input checked="" type="checkbox"/> 229	762.0610	2283.1612	2283.0664	0.0948	0	31	20	1	MESPGAVSLWFGNGTSEMVLIR + Oxidation (M)
<input checked="" type="checkbox"/> 232	767.3850	2299.1332	2300.0896	-0.9564	1	30	23	1	FLRNYHTDFEVSIVCAISK
<input checked="" type="checkbox"/> 81	553.5800	1105.1454	1105.5437	-0.3983	0	30	50	1	DLNFMLDLR + Oxidation (M)
<input checked="" type="checkbox"/> 112	738.3990	1474.7834	1475.8347	-1.0513	2	29	36	1	FNKQQLDIKTK
<input checked="" type="checkbox"/> 92	621.9790	1241.9434	1242.7295	-0.7861	2	28	71	1	ILETNKKLR
<input checked="" type="checkbox"/> 217	737.7010	2210.0812	2211.0598	-0.9787	0	28	39	1	ERPRPVMSTASQMLVER + Oxidation (M)
<input checked="" type="checkbox"/> 201	1065.5450	2129.0754	2128.9657	0.1098	0	27	77	1	TFDNGMICASEQSVIVVDK + Oxidation (M)
<input checked="" type="checkbox"/> 78	536.2960	1070.5774	1069.4862	1.0912	0	27	1.1e+02	1	MIDNYSTAR
<input checked="" type="checkbox"/> 42	435.7910	869.5674	870.5902	-1.0228	1	27	55	1	KATVLVLR
<input checked="" type="checkbox"/> 94	633.2510	1264.4874	1265.5267	-1.0393	0	27	89	1	QEMDMENEIK
<input checked="" type="checkbox"/> 17	386.0810	770.1474	769.4195	0.7280	1	26	45	1	AGVDPRR
<input checked="" type="checkbox"/> 139	876.3290	2625.9652	2625.2964	0.6687	2	26	84	1	LAGMILDPSSELISMSDFLRGCR + Oxidation (M)
<input checked="" type="checkbox"/> 38	428.7700	855.5254	855.4814	0.0440	0	26	75	1	LATLHSSK
<input checked="" type="checkbox"/> 220	742.3550	2224.0432	2223.0240	1.0192	2	26	60	1	MPRGGAFWGGSRRTQDAEGR + Oxidation (M)
<input checked="" type="checkbox"/> 37	428.6860	855.3574	855.5793	-0.2219	0	26	81	1	IATVLVLR
<input checked="" type="checkbox"/> 156	934.9720	2801.8942	2802.2524	-0.3582	2	25	1.3e+02	1	RIHHIMVECSDFNDVGLAYDRCK + Oxidation (M)
<input checked="" type="checkbox"/> 207	720.4810	2158.4212	2158.1780	0.2432	1	25	56	1	GINQLDLRQLMATSLSAVTK
<input checked="" type="checkbox"/> 103	704.6680	2110.9822	2112.0851	-1.1030	1	25	1.4e+02	1	VTDLNGFGFDVPADALDRLK
<input checked="" type="checkbox"/> 248	927.6190	2779.8352	2779.3466	0.4885	1	25	49	1	EIHNSNLDFFPSMHSFRVEHYLK
<input checked="" type="checkbox"/> 238	788.8070	2363.3992	2364.1743	-0.7752	1	24	69	1	KQGPTSVAYVEVNNNSMLNVRK + Oxidation (M)
<input checked="" type="checkbox"/> 233	1155.8640	3464.5702	3465.6441	-1.0739	1	24	1.1e+02	1	QTMNMRKYLHQVMPFFIPADVDDIETR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 69	496.0800	1485.2182	1485.8766	-0.6584	2	24	1.6e+02	1	SLETIGQLKLRK
<input checked="" type="checkbox"/> 108	727.7670	2180.2792	2180.1048	0.1744	1	24	1.7e+02	1	MANDTALANNLLPYRNFK + Oxidation (M)

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MATRIX
SCIENCE Mascot Search Results

Protein View

Match to: **gi|261339072** Score: **195****YfaZ family protein [Enterobacter cancerogenus ATCC 35316]**

Found in search of D:\Data\Lakshmy\120808\280808\SSP 8205_RJ23_01_550.d\SSP 8205_RJ23_01_550.mgf

Nominal mass (M_r): **18559**; Calculated pI value: **6.82**NCBI BLAST search of **gi|261339072** against nrUnformatted [sequence string](#) for pasting into other applicationsTaxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)

Variable modifications: Oxidation (M)

Cleavage by Trypsin: cuts C-term side of KR unless next residue is P

Sequence Coverage: **24%**Matched peptides shown in **Bold Red**

1 MKKLNLLLL ALTVVSGSAL AMGGSIEQ GK **NFTNLNLEMG KSSGLYAES**
 51 **NWLKNTDDGT** QTGGVGAGYN LEVGPVLMNA GAKAIYIGPK KGDNGVAFPI
 101 GGGVNVALTD SIHVFGEGYV APDGLNNSVK NYVEANGGVS WTPIKEVTLK
 151 VGYRHVSVDG **KDGRPNHTLI DGAYVGGGVS F**

 Residue Number
 Increasing Mass
 Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
31 - 41	648.8060	1295.5974	1295.6180	-0.0205	0	K.NFTNLNLEMGK.S Oxidation (M) (Ions score 55)
42 - 54	721.3580	1440.7014	1440.6885	0.0130	0	K.SSSGLYAESNWLK.N (Ions score 61)
162 - 181	678.2780	2031.8122	2030.9810	0.8311	0	K.DGRPNHTLIDGAYVGGGVSF.- (Ions score 79)

Error: try setting browser cache to automatic.

Error: try setting browser cache to automatic.

LOCUS ZP_05966930 181 aa linear BCT 15-OCT-2009
 DEFINITION YfaZ family protein [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05966930
 VERSION ZP_05966930.1 GI:261339072
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000004.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 181)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaanty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 181)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM02000004](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be
 aware that the annotation is done automatically with little or no
 manual curation.

http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100105/FtmmSruT... 21/01/2010

Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

```

FEATURES             Location/Qualifiers
    source            1..181
                     /organism="Enterobacter cancerogenus ATCC 35316"
                     /strain="ATCC 35316"
                     /db_xref="taxon:500639"
    Protein           1..181
                     /product="YfaZ family protein"
                     /calculated_mol_wt=18440
    Region            1..181
                     /region_name="YfaZ"
                     /note="YfaZ precursor; pfam07437"
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    CDS               1..181
                     /locus_tag="EcanA3_020100001365"
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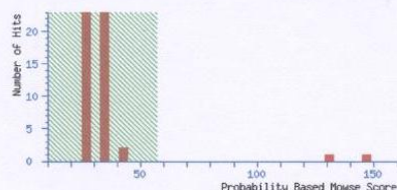
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy_manickan@unn.ac.uk
 Search title : SSP 8206
 MS data file : D:\Data\Lakshmy\120808\241108\SSP 8206_RB19_01_769.d\SSP 8206_RB19_01_769.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 5 Jan 2010 at 12:14:59 GMT
 Protein hits : [gi|237732258](#) ecotin [Citrobacter sp. 30_2]
 [gi|50582463](#) oxidoreductase [Exiguobacterium acetylicum]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: **Peptide Summary** [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|237732258](#) Mass: 19051 Score: 147 Queries matched: 5 emPAI: 0.63
 ecotin [Citrobacter sp. 30_2]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 111	529.2380	1584.6922	1584.8763	-0.1841	0	(33)	12	1	K.LPIVVVTPENVVVK.Y
<input checked="" type="checkbox"/> 113	793.3830	1584.7514	1584.8763	-0.1249	0	60	0.026	1	K.LPIVVVTPENVVVK.Y
<input checked="" type="checkbox"/> 114	793.3980	1584.7814	1584.8763	-0.0949	0	(57)	0.057	1	K.LPIVVVTPENVVVK.Y
<input checked="" type="checkbox"/> 115	793.4280	1584.8414	1584.8763	-0.0349	0	(39)	3.3	1	K.LPIVVVTPENVVVK.Y
<input checked="" type="checkbox"/> 147	878.2990	1754.5834	1754.7828	-0.1994	0	86	4.4e-05	1	K.TLEGWGYDYVFDK.V

Proteins matching the same set of peptides:
[gi|261340622](#) Mass: 18823 Score: 147 Queries matched: 5
 ecotin [Enterobacter cancerogenus ATCC 35316]

2. [gi|50582463](#) Mass: 27360 Score: 129 Queries matched: 2 emPAI: 0.26
 oxidoreductase [Exiguobacterium acetylicum]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 138	859.3720	1716.7294	1716.8570	-0.1276	0	87	5.1e-05	1	K.YTVITGASSGIYETAK.L
<input checked="" type="checkbox"/> 198	1015.4730	2028.9314	2028.9752	-0.0438	0	42	1.5	1	K.SVDLADNQNVHDLVEGLK.E

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 68	455.0960	1362.2662	1362.6204	-0.3543	0	43	1.7	1	FEDTFLSDFHR
<input checked="" type="checkbox"/> 194	675.2970	2022.8692	2023.0520	-0.1829	1	40	2.1	1	VELLIGTQLKVDNQR
<input checked="" type="checkbox"/> 178	644.9340	1931.7802	1932.0720	-0.2919	0	39	2.6	1	IDLLLATPEGWVLDHK
<input checked="" type="checkbox"/> 117	800.8320	1599.6494	1599.7927	-0.1432	0	36	6	1	GDPALPATMREGIGVR + Oxidation (M)
<input checked="" type="checkbox"/> 93	681.7720	1361.5294	1362.6568	-1.1274	0	36	13	1	HGYAFNLDLQK
<input checked="" type="checkbox"/> 200	680.1840	2037.5302	2037.0565	0.4737	1	35	4.8	1	GASVRDIMGGLDGLSISVK + Oxidation (M)
<input checked="" type="checkbox"/> 67	453.2220	904.4294	905.4389	-1.0094	1	34	27	1	MQEAAGRK + Oxidation (M)
<input checked="" type="checkbox"/> 124	546.8830	1637.6272	1637.7937	-0.1665	1	33	9.9	1	KADEYPIITFGDTPK
<input checked="" type="checkbox"/> 101	714.5440	1427.0734	1427.7191	-0.6457	0	33	20	1	ADVPPVQLSGMQR
<input checked="" type="checkbox"/> 140	864.8670	1727.7194	1727.9604	-0.2409	1	32	16	1	INLKIPAGIESGHTLR + Oxidation (M)
<input checked="" type="checkbox"/> 225	747.0320	2238.0742	2239.1174	-1.0432	1	31	20	1	QGPNLVGYETQNWFLKQR
<input checked="" type="checkbox"/> 81	570.6700	1139.3254	1139.5459	-0.2204	1	30	41	1	DSSKGDVTGPK
<input checked="" type="checkbox"/> 208	1037.5310	3109.5712	3108.5332	1.0380	1	30	45	1	DRITAVLMPGSVVSILANDSERPQYNATR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 95	692.2590	1382.5034	1382.8245	-0.3211	2	30	25	1	LSIALGKSGKNPAK
<input checked="" type="checkbox"/> 219	737.6540	2209.9402	2209.1062	0.8340	2	28	36	1	NSSNIFQIKKFTNTQTR + Oxidation (M)

mhtml:file://C:\Documents and Settings\larcje\Local Settings\Temp\Peptide Summar... 21/01/2010

aware that the annotation is done automatically with little or no manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

FEATURES	Location/Qualifiers
source	1..167 /organism="Enterobacter cancerogenus ATCC 35316" /strain="ATCC 35316" /db_xref="taxon:500639"
Protein	1..167 /product="ecotin" /calculated_mol_wt=18529
Region	30..165 /region_name="Ecotin" /note="Protease Inhibitor Ecotin; homodimeric protease inhibitor which binds two chymotrypsin-like serine proteases to form a heterotetramer. Found in bacterial periplasm. Inhibits a broad range of serine proteases including collagenase, trypsin, chymotrypsin...; cd00242"
Site	/db_xref="CDD:29318" order(91..95,133..138) /site_type="other" /note="secondary substrate binding site" /db_xref="CDD:29318"
Site	107..110 /site_type="other" /note="primary substrate binding site" /db_xref="CDD:29318"
Site	109..110 /site_type="inhibition" /note="inhibition loop" /db_xref="CDD:29318"
Site	155..165 /site_type="other" /note="dimerization interface" /db_xref="CDD:29318"
CDS	1..167 /locus_tag="EcanA3_020100009153" /coded_by="NZ_ABWM02000010.1:67048..67551" /note="COG4574 Serine protease inhibitor ecotin" /transl_table=11 /db_xref="CDD:134928"

Mascot: <http://www.matrixscience.com/>

MASCOT SCIENCE Mascot Search Results

Protein View

Match to: **gi|261340622** Score: **147**
ecotin [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\241108\SSP 8206_RB19_01_769.d\SSP 8206_RB19_01_769.mgf

Nominal mass (M_n): **18823**; Calculated pI value: **8.33**
 NCBI BLAST search of [gi|261340622](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **16%**

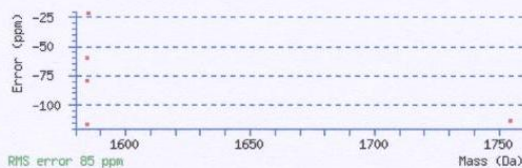
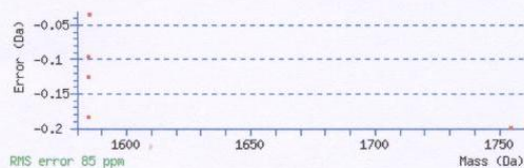
Matched peptides shown in **Bold Red**

- 1 MKNAPKFAIA LLAACVSTVS AFATDTKNDQ PLEKVPYYPQ AEKGMKRQVI
- 51 QVPAQENEAN FKVELLIGQT LEVDCNHHRL GGKLESK**LE** **GWGYDYVFD**
- 101 **KVTSFPVSTMM** ACPDGKKEKK FVTAYLGDNS LLRYNSK**LPI** **VVYTPENVDV**
- 151 **KYRIWKADEN** VGQAVIR

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
88 - 101	878.2990	1754.5834	1754.7828	-0.1994	0	K.TLEGMGYDYVFDK.V (Ions score 86)
138 - 151	529.2380	1584.6922	1584.8763	-0.1841	0	K.LPIVVYTPENVDK.Y (Ions score 33)
138 - 151	793.3830	1584.7514	1584.8763	-0.1249	0	K.LPIVVYTPENVDK.Y (Ions score 60)
138 - 151	793.3980	1584.7814	1584.8763	-0.0949	0	K.LPIVVYTPENVDK.Y (Ions score 57)
138 - 151	793.4280	1584.8414	1584.8763	-0.0349	0	K.LPIVVYTPENVDK.Y (Ions score 39)



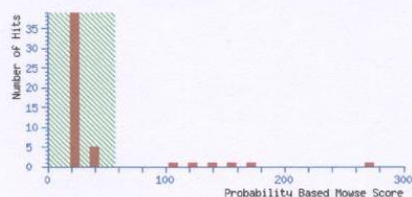
LOCUS ZP_05968480 167 aa linear BCT 15-OCT-2009
 DEFINITION ecotin [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05968480
 VERSION ZP_05968480.1 GI:261340622
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000010.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM Enterobacter cancerogenus ATCC 35316
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 167)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 167)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM02000010](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be
 found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 8707
MS data file : D:\Data\Lakshmy\120808\280808\SSP 8707_RK17_01_587.d\SSP 8707_RK17_01_587.mgf
Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 12:23:26 GMT
Protein hits : [gi|261341330](#) translocation protein TolB [Enterobacter cancerogenus ATCC 35316]
[gi|10120890](#) Chain A, Crystal Structure Of The E. Coli TolB Protein
[gi|62179322](#) translocation protein TolB [Salmonella enterica subsp. enterica serovar Choleraesuis str. SC-B67]
[gi|157369522](#) translocation protein TolB [Serratia proteamaculans 568]
[gi|152969308](#) translocation protein TolB [Klebsiella pneumoniae subsp. pneumoniae MGH 78578]
[gi|242238602](#) Tol-Pal system beta propeller repeat protein TolB [Dickeya dadantii Ech703]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As **Peptide Summary** [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|261341330](#) Mass: 45999 Score: 271 Queries matched: 7 emPAI: 0.32
 translocation protein TolB [Enterobacter cancerogenus ATCC 35316]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 50	571.7910	1141.5674	1141.5768	-0.0093	0	70	0.0028	1	K.LAYVTFESGR.S
<input checked="" type="checkbox"/> 51	571.7940	1141.5734	1141.5768	-0.0033	0	(70)	0.0035	1	K.LAYVTFESGR.S
<input checked="" type="checkbox"/> 62	714.9110	1427.8074	1427.8096	-0.0022	0	40	3.1	1	R.SALVIQTLNGLAVR.Q
<input checked="" type="checkbox"/> 49	567.5860	1699.7362	1699.7876	-0.0514	0	(34)	18	1	R.SPQPLMSPAWSPDGSK.L + Oxidation (M)
<input checked="" type="checkbox"/> 89	850.9190	1699.8234	1699.7591	0.0644	0	37	5.5	1	R.VSDYDGNQFTVHR.S
<input checked="" type="checkbox"/> 90	850.9250	1699.8354	1699.7876	0.0479	0	69	0.0033	1	R.SPQPLMSPAWSPDGSK.L + Oxidation (M)
<input checked="" type="checkbox"/> 105	912.4500	1822.8854	1822.9247	-0.0393	0	59	0.036	1	K.TGSLNLTVMIDIGSGQIR.Q

2. [gi|10120890](#) Mass: 42745 Score: 170 Queries matched: 4 emPAI: 0.16
 Chain A, Crystal Structure Of The E. Coli TolB Protein
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
50	571.7910	1141.5674	1141.5768	-0.0093	0	70	0.0028	1	K.LAYVTFESGR.S
51	571.7940	1141.5734	1141.5768	-0.0033	0	(70)	0.0035	1	K.LAYVTFESGR.S
49	567.5860	1699.7362	1699.8206	-0.0844	0	34	18	1	R.SPQPLMSPAWSPDGSK.L
90	850.9250	1699.8354	1699.7876	0.0479	0	69	0.0033	1	R.SPQPLMSPAWSPDGSK.L + Oxidation (M)

3. [gi|62179322](#) Mass: 46150 Score: 164 Queries matched: 5 emPAI: 0.15
 translocation protein TolB [Salmonella enterica subsp. enterica serovar Choleraesuis str. SC-B67]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
50	571.7910	1141.5674	1141.5768	-0.0093	0	70	0.0028	1	K.LAYVTFESGR.S
51	571.7940	1141.5734	1141.5768	-0.0033	0	(70)	0.0035	1	K.LAYVTFESGR.S
62	714.9110	1427.8074	1427.8096	-0.0022	0	24	1.2e+02	3	R.SALVIQTLNGLAVR.Q
49	567.5860	1699.7362	1699.7876	-0.0514	0	(34)	18	1	R.SPQPLMSPAWSPDGSK.L + Oxidation (M)
90	850.9250	1699.8354	1699.7876	0.0479	0	69	0.0033	1	R.SPQPLMSPAWSPDGSK.L + Oxidation (M)

Proteins matching the same set of peptides:
[gi|224582567](#) Mass: 46584 Score: 164 Queries matched: 5
 translocation protein TolB precursor [Salmonella enterica subsp. enterica serovar Paratyphi C strain RKS4594]

Mascot Search Results

Protein View

Match to: **gi|261341330** Score: 271
translocation protein TolB [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\280808\SSP 8707_RK17_01_587.d\SSP 8707_RK17_01_587.mgf

Nominal mass (M_r): **45999**; Calculated pI value: **6.98**
 NCBI BLAST search of [gi|261341330](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **16%**

Matched peptides shown in **Bold Red**

```

1 MMKQALRVAF GFIMLWAAVL HAEVRIEITQ GVDSARPIGV VPFQWAGPGA
51 APEDIGGIVA ADLRNSGKFN PLDRSRLPQQ PGTAQEVQPA AWSALGIDAV
101 VVGQVTPAPD GGYNVAYQLV DTGGAPGTVL AQNTYKVNKQ WLRVAGHTAS
151 DEVFEKLTGI KGAFRTRIAV VVQTNGGQFP YELRVSDYDG YNQFTVHRSP
201 QPLMSPAWSF DGSKLAVTFE ESGRSALVIQ TLSNGAVRQV ASFPRHNGAP
251 AFSPDGTKLA FALSKTGSLN LYVMDIGSGQ IRQITDGRSN NTEPTWFPDS
301 QNLAFSTDQA GRFOYKVINI NNGAAQRITW EGSGNQDADI SADGKTMVMV
351 STAGGQQHIA KQDLVTTGGVQ VLSSTFLEDT PSLAPNGTMV IYSSSQGMGS
401 VLNLVSTDGR FKARIPATDG QVKSPAWSPY L
    
```

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
185 - 198	850.9190	1699.8234	1699.7591	0.0644	0	R.VSDYDGYNQFTVHR.S (Ions score 37)
199 - 214	567.5860	1699.7362	1699.7876	-0.0514	0	R.SPQPLMSPAWSFDGSK.L Oxidation (M) (Ions score 34)
199 - 214	850.9250	1699.8354	1699.7876	0.0479	0	R.SPQPLMSPAWSFDGSK.L Oxidation (M) (Ions score 69)
215 - 224	571.7910	1141.5674	1141.5768	-0.0093	0	K.LAYVTFESGR.S (Ions score 70)
215 - 224	571.7940	1141.5734	1141.5768	-0.0033	0	K.LAYVTFESGR.S (Ions score 70)
225 - 238	714.9110	1427.8074	1427.8096	-0.0022	0	R.SALVIQTLNSGAVR.Q (Ions score 40)
266 - 282	912.4500	1822.8854	1822.9247	-0.0393	0	K.TGSLNLYVMDIGSGQIR.Q (Ions score 59)

Error: try setting browser cache to automatic.

Error: try setting browser cache to automatic.

LOCUS ZP_05969188 431 aa linear BCT 15-OCT-2009
 DEFINITION translocation protein TolB [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05969188
 VERSION ZP_05969188.1 GI:261341330
 DELINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000020.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 431)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 431)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM02000020](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic

Annotation Pipeline Group. Information about the Pipeline can be found here: <http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x
 Sequencing Technology: 454.
 Method: conceptual translation.

```

FEATURES             Location/Qualifiers
    source            1..431
                     /organism="Enterobacter cancerogenus ATCC 35316"
                     /strain="ATCC 35316"
                     /db_xref="taxon:500639"
    Protein           1..431
                     /product="translocation protein TolB"
                     /calculated_mol_wt=45897
    Region            2..431
                     /region_name="tolB"
                     /note="Translocation protein TolB; Provisional; PRK03629"
                     /db_xref="CDD:134902"
    Region            31..174
                     /region_name="TolB_N"
                     /note="TolB amino-terminal domain; pfam04052"
                     /db_xref="CDD:112850"
    Region            238..273
                     /region_name="PD40"
                     /note="WD40-like Beta Propeller Repeat; pfam07676"
                     /db_xref="CDD:116290"
    Region            282..317
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                     /note="WD40-like Beta Propeller Repeat; pfam07676"
                     /db_xref="CDD:116290"
    CDS                1..431
                     /gene="tolB"
                     /locus_tag="EcanA3_020100012746"
                     /coded_by="NZ_ABWM02000020.1:251385..252680"
                     /note="COG0823 Periplasmic component of the Tol biopolymer
                     transport system"
                     /transl_table=11
                     /db_xref="CDD:134902"
    
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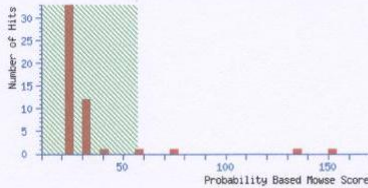
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 9601
MS data file : D:\Data\Lakshmy\120808\140808\SSP9601_RF13_01_334.d\SSP9601_RF13_01_334.mgf
Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 5 Jan 2010 at 12:24:38 GMT
Protein hits : [gi|148368](#) outer membrane protein II [Enterobacter aerogenes]
[gi|146311681](#) acetoin reductase [Enterobacter sp. 638]
[gi|157146352](#) outer membrane protein A [Citrobacter koseri ATCC BAA-895]
[gi|146312800](#) glycine betaine transporter periplasmic subunit [Enterobacter sp. 638]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)
 Significance threshold p<: Max. number of hits:
 Standard scoring MudPIT scoring Ions score or expect cut-off: Show sub-sets:
 Show pop-ups Suppress pop-ups Sort unassigned: Require bold red:

Select All Select None Search Selected Error tolerant

1. [gi|148368](#) Mass: 25654 Score: 152 Queries matched: 3 emPAI: 0.28
 outer membrane protein II [Enterobacter aerogenes]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
90	632.8000	1263.5854	1263.6460	-0.0605	0	41	3.7	2	K.DGSVVVLGFTDR.I
<input checked="" type="checkbox"/> 112	813.9580	1625.9014	1625.7937	0.1077	0	42	1.7	1	K.LGYVPTDDLQVYTR.L
<input checked="" type="checkbox"/> 235	783.3450	2347.0132	2346.1856	0.8276	0	69	0.0027	1	R.FGQEDAPVVAAPAPAPEVQTK.H

Proteins matching the same set of peptides:
[gi|261339284](#) Mass: 37561 Score: 152 Queries matched: 3
 outer membrane protein A [Enterobacter cancerogenus ATCC 35316]

2. [gi|146311681](#) Mass: 26942 Score: 137 Queries matched: 3 emPAI: 0.26
 acetoin reductase [Enterobacter sp. 638]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
126	564.6150	1690.8232	1689.8686	0.9546	1	20	2.5e+02	4	K.VDVSNREQVFAAVEK.A
<input checked="" type="checkbox"/> 132	857.3840	1712.7534	1712.7893	-0.0359	0	55	0.085	1	K.DGFAVAIADYNEETAK.A
<input checked="" type="checkbox"/> 238	802.0750	2403.2032	2404.1329	-0.9297	0	61	0.017	1	K.IINACSQAGHTGNFELAVYSSSK.F

Proteins matching the same set of peptides:
[gi|261341164](#) Mass: 26884 Score: 137 Queries matched: 3
 acetoin reductase [Enterobacter cancerogenus ATCC 35316]

3. [gi|157146352](#) Score: 77 Queries matched: 2
 outer membrane protein A [Citrobacter koseri ATCC BAA-895]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
90	632.8000	1263.5854	1263.6460	-0.0605	0	41	3.7	2	K.DGSVVVLGFTDR.I
235	783.3450	2347.0132	2346.1855	0.8276	0	36	5.2	2	R.FGQEEAAPIAPAPAPEVQTK.H

4. [gi|146312800](#) Mass: 36495 Score: 58 Queries matched: 1 emPAI: 0.09
 glycine betaine transporter periplasmic subunit [Enterobacter sp. 638]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 119	835.3920	1668.7694	1668.8359	-0.0665	0	58	0.043	1	R.EGTFVTGAAGYGLIDK.K

MASCOT SCIENCE Mascot Search Results

Protein View

Match to: **gi|261339284** Score: **152**
outer membrane protein A [Enterobacter cancerogenus ATCC 35316]
 Found in search of D:\Data\Lakshmy\120808\140808\SSP9601_RF13_01_334.d\SSP9601_RF13_01_334.mgf

Nominal mass (M_r): **37561**; Calculated pI value: **5.19**
 NCBI BLAST search of **gi|261339284** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Enterobacter cancerogenus ATCC 35316](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **14%**

Matched peptides shown in **Bold Red**

```

1 MKKTAIAIAV ALAGFATVAQ AAPKDNTWYA GKGWLSQFQ DTGWYNSSLN
51 NDGPETHESQL GAGAFGGYQV NPYVGFEMGY DWLGRMPYKG DNVNGAFKAQ
101 GVQLTAKLGY FVTDLDLVYT RLGGMVWRAD SSNSIAGDDH DTGVSFVFAG
151 GVEWAMTRDI ATRLEYQWVN NIGDGTWGV RPDNGMLSVG VSYRFGQQED
201 APVVAPAPAP APEVQTKHFT LKSDVLEFNF KATLKPEGQQ ALDQLYTQLS
251 NLDPKDGSVV VLGFTDRIGS DAYNQLSEK RAQSVVDYLV SKGIPANKIS
301 PRGMGESNPV TGNTCDNVKP RAALIDCLAP DRRVEIEVKG IKDVVTQPAA
351
    
```

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
108 - 121	813.9580	1625.9014	1625.7937	0.1077	0	K.LGYFVTDDLVDVYTR.L (Ions score 42)
195 - 217	783.3450	2347.0132	2346.1856	0.8276	0	R.FGQQEDAPVVAPAPAPAPEVQTK.H (Ions score 69)
256 - 267	632.8000	1263.5854	1263.6460	-0.0605	0	K.DGSVVVLGFTDR.I (Ions score 41)

Error: try setting browser cache to automatic.

Error: try setting browser cache to automatic.

LOCUS ZP_05967142 350 aa linear BCT 15-OCT-2009
 DEFINITION outer membrane protein A [Enterobacter cancerogenus ATCC 35316].
 ACCESSION ZP_05967142
 VERSION ZP_05967142.1 GI:261339284
 DBLINK Project:28663
 DBSOURCE REFSEQ: accession [NZ_ABWM02000004.1](#)
 KEYWORDS .
 SOURCE Enterobacter cancerogenus ATCC 35316
 ORGANISM [Enterobacter cancerogenus ATCC 35316](#)
 Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
 Enterobacteriaceae; Enterobacter.
 REFERENCE 1 (residues 1 to 350)
 AUTHORS Weinstock,G., Sodergren,E., Clifton,S., Fulton,L., Fulton,B.,
 Courtney,L., Fronick,C., Harrison,M., Strong,C., Farmer,C.,
 Delahaunty,K., Markovic,C., Hall,O., Minx,P., Tomlinson,C.,
 Mitreva,M., Nelson,J., Hou,S., Wollam,A., Pepin,K.H., Johnson,M.,
 Bhonagiri,V., Nash,W.E., Warren,W., Chinwalla,A., Mardis,E.R. and
 Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (06-OCT-2009) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 REFERENCE 2 (residues 1 to 350)
 AUTHORS Fulton,L., Clifton,S., Fulton,B., Xu,J., Minx,P., Pepin,K.H.,
 Johnson,M., Thiruvilangam,P., Bhonagiri,V., Nash,W.E., Mardis,E.R.
 and Wilson,R.K.
 TITLE Direct Submission
 JOURNAL Submitted (17-SEP-2008) Genome Sequencing Center, Washington
 University School of Medicine, 4444 Forest Park, St. Louis, MO
 63108, USA
 COMMENT WGS REFSEQ: This record is provided to represent a collection of
 whole genome shotgun sequences. The reference sequence was derived
 from [ABWM02000004](#).
 Annotation was added by the NCBI Prokaryotic Genomes Automatic
 Annotation Pipeline Group. Information about the Pipeline can be

http://www.matrixscience.com/cgi/protein_view.pl?file=../data/20100105/FtmmSrsST... 21/01/2010

found here:
<http://www.ncbi.nlm.nih.gov/genomes/static/Pipeline.html>. Please be aware that the annotation is done automatically with little or no manual curation.
 Enterobacter cancerogenus (GenBank Accession Number for 16S rDNA gene: Z96078, the 16S rDNA gene of a related strain of Enterobacter cancerogenus, not the sequenced strain), is a member of the Proteobacteria division of the domain bacteria and has been isolated from human feces. The sequenced strain was obtained from ATCC (ATCC 35316).

We have performed one round of automated sequence improvement (pre-finishing), along with manual improvement that includes breaking apart any mis-assembly, and making manual joins where possible. Manual edits also are made where the consensus appears to be incorrect. All low quality data on the ends of contigs is removed. Contigs are ordered and oriented where possible.

This work was supported by the National Human Genome Research Institute and the NIH Roadmap Human Microbiome Project [<http://nihroadmap.nih.gov/hmp/>].

This is a reference genome for the Human Microbiome Project. This project is co-owned with the Human Microbiome Project DACC. Genome Coverage: 29.7x Sequencing Technology: 454. Method: conceptual translation.

FEATURES	Location/Qualifiers
source	1..350 /organism="Enterobacter cancerogenus ATCC 35316" /strain="ATCC 35316" /db_xref="taxon:500639"
Protein	1..350 /product="outer membrane protein A" /calculated_mol_wt=37337
Region	1..346 /region_name="PRK10808" /note="outer membrane protein A; Reviewed; PRK10808" /db_xref="CDD:138188"
Region	23..196 /region_name="Surface_Ag_2" /note="Surface antigen; cl01155" /db_xref="CDD:141128"
Region	224..338 /region_name="OmpA_C-like" /note="Peptidoglycan binding domains similar to the C-terminal domain of outer-membrane protein OmpA; cd07185" /db_xref="CDD:143586"
Site	order(229..230,265..266,269,273..274,277,328,332) /site_type="other" /note="ligand binding site" /db_xref="CDD:143586"
CDS	1..350 /locus_tag="EcanA3_020100002425" /coded_by="complement(NZ_ABWM02000004.1:545400..546452)" /note="COG2885 Outer membrane protein and related peptidoglycan-associated (lipo)proteins" /transl_table=11 /db_xref="CDD:138188"

Mascot: <http://www.matrixscience.com/>

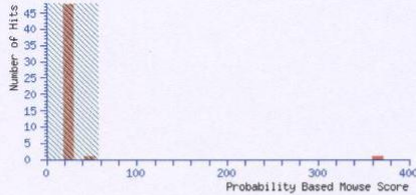
G3 Mass spectrometric analysis data for differentially expressed spots *B. fragilis*

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 1104
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (1104) 10_RJ17_01_1396.d\SSP (1104) 10_RJ17_01_1396.mgf
 Database : NCBI nr 20100108 (10291680 sequences; 3511877860 residues)
 Taxonomy : Bacteria (Subacteria) (5718488 sequences)
 Timestamp : 13 Jan 2010 at 10:39:58 GMT
 Protein hits : [gi|53713703](#) TPR repeat-containing protein [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 * \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As **Peptide Summary** [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|53713703](#) Mass: 46634 Score: 366 Queries matched: 8 emPAI: 0.51
 TPR repeat-containing protein [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 125	776.2740	1550.5334	1550.7365	-0.2031	0	64	0.0073	1	K.DNAATWDVAGYIQK.R
<input checked="" type="checkbox"/> 140	835.8360	1669.6574	1669.8522	-0.1948	0	74	0.00086	1	K.AEQLINBALTNPETK.D
<input checked="" type="checkbox"/> 185	663.5640	1987.6702	1986.9357	* 0.7345	1	51	0.13	1	K.DKENGQFAMQLLTDAYK.A + Oxidation (M)
<input checked="" type="checkbox"/> 186	994.8670	1987.7194	1986.9357	0.7838	1	(26)	97	1	K.DKENGQFAMQLLTDAYK.A + Oxidation (M)
<input checked="" type="checkbox"/> 187	995.8420	1989.6694	1989.9030	-0.2335	0	68	0.0026	1	R.VVYNLNMGPFFREIEK.M + Oxidation (M)
<input checked="" type="checkbox"/> 233	762.6100	2284.8082	2283.9738	0.8344	1	19	2.1e+02	2	R.VVYNLNMGPFFREIEK.M - + 3 Oxidation (M)
<input checked="" type="checkbox"/> 238	797.0170	2388.0292	2387.2961	0.7331	0	32	12	1	K.TILAEKFNLIINGIYQPNLNK.N
<input checked="" type="checkbox"/> 248	930.6050	2788.7932	2789.2245	-0.4313	0	57	0.03	1	K.FPENQYFFANLVDDYSSNQNDK.A

Proteins matching the same set of peptides:
[gi|60681974](#) Mass: 45773 Score: 366 Queries matched: 8
 hypothetical protein BF2494 [Bacteroides fragilis NCTC 9343]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 221	737.9210	2210.7412	2210.1115	-0.6297	0	34	6.3	1	MYILGINSVYHESAVCLLK
<input checked="" type="checkbox"/> 131	804.8580	1607.7014	1607.8125	-0.1110	0	33	14	1	MPIYNELMLPLK + Oxidation (M)
<input checked="" type="checkbox"/> 234	766.9800	2297.9182	2299.0790	-1.1609	2	32	11	1	LNGYWEIDKMRSLTDDAEK + Oxidation (M)
<input checked="" type="checkbox"/> 80	629.4150	1885.2232	1885.9707	-0.7475	0	32	32	1	SIMDNLGLPVIDDEIVK + Oxidation (M)
<input checked="" type="checkbox"/> 121	752.3550	2254.0432	2255.0205	-0.9773	1	30	51	1	LNEPDPDGDWKIEMPLR + Oxidation (M)
<input checked="" type="checkbox"/> 225	1113.0610	2224.1074	2224.0437	0.0638	0	30	25	1	DISWGSTWGAGDNVDIYQLK
<input checked="" type="checkbox"/> 44	391.1050	1170.2932	1169.6227	0.6705	0	28	59	1	LLAMEGVNPAR
<input checked="" type="checkbox"/> 88	650.7300	1299.4454	1298.7921	0.6533	2	27	44	1	KNESLKLVLQK
<input checked="" type="checkbox"/> 157	585.0580	1752.1522	1752.9312	-0.7790	2	27	40	1	VIGGNPKRFQPTPK
<input checked="" type="checkbox"/> 108	707.5460	1413.0774	1412.7848	0.2927	2	27	77	1	VAVATEGRRVAR
<input checked="" type="checkbox"/> 79	623.1710	1866.4912	1865.8263	0.6649	1	26	91	1	LWGGYAKGMTAMDYMR + Oxidation (M)
<input checked="" type="checkbox"/> 229	747.8320	2240.4742	2240.2277	0.2465	1	26	40	1	YQVDLLTLHATAGRALTAAK
<input checked="" type="checkbox"/> 126	779.9360	2336.7862	2336.1722	0.6140	0	26	1.1e+02	1	GVEVFLINALSGEGLPELMEK
<input checked="" type="checkbox"/> 209	700.9180	2099.7322	2099.9941	-0.2620	0	25	49	1	TINDMQMYIECVQLVVK + Oxidation (M)
<input checked="" type="checkbox"/> 196	681.0290	2040.0652	2040.9350	-0.8698	2	25	74	1	VERIPDMYGVKKETAET + Oxidation (M)
<input checked="" type="checkbox"/> 145	563.1340	1686.3802	1685.9365	0.4436	2	25	56	1	AGLIHAFNDRKVAK
<input checked="" type="checkbox"/> 112	721.6110	2161.8112	2162.0578	-0.2466	2	25	1.1e+02	1	ATYSDEFYMARALELARR
<input checked="" type="checkbox"/> 204	694.9280	2081.7622	2081.9795	-0.2174	1	25	55	1	MAEMVIGEKVTLSEMGGAR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 142	838.0410	2511.1012	2511.2329	-0.1317	1	25	1.4e+02	1	SVIYEMHVGGFTRHPNSGVAPEK
<input checked="" type="checkbox"/> 213	711.0680	2130.1822	2129.9721	0.2101	2	25	82	1	CDTEIRKYAICLSLQR

Mascot Search Results

Protein View

Match to: **gi|60681974** Score: **366**
hypothetical protein BF2494 [Bacteroides fragilis NCTC 9343]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (1104) 10_RJ17_01_1396.d\SSP (1104) 10_RJ17_01_1396.mgf

Nominal mass (M_r): **45773**; Calculated pI value: **5.24**
 NCBI BLAST search of **gi|60681974** against nr
 Unformatted **sequence string** for pasting into other applications

Taxonomy: **Bacteroides fragilis NCTC 9343**
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60493408](#) from **Bacteroides fragilis NCTC 9343**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **27%**

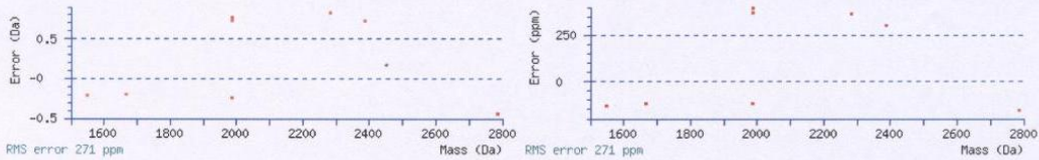
Matched peptides shown in **Bold Red**

1 MVL~~L~~MAVSFA FAQEK~~N~~VKEA KSTAGEV~~K~~PD FAKA**EQ**LINE ALTN**PE**TKDN
 51 AAT**W**DVAGYI QKRINEKEME NAYLRKPYDT LKVVNSVLNM YNYYVKDEL
 101 AQIPNEK**G**KI K~~N~~KYRSANSK **TIL**AERPNLI NGG**IQ**YFNLN KNEDALYKFA
 151 AYVDAATL**E**M MEKENLLEKD TILPQVAYYA TLAADRVGDK DAVMKYAQYA
 201 LK**D**KENGQFA MQL**L**TDAYKA KGDTAKWVEK LQEGIV**K**FPE NQY**F**FANLVD
 251 Y**Y**SSSNQNDK AMQFAD**D**MLA KDPNNKLYLY VKAYLYHNMK DYEKAIEFYK
 301 KTL**D**IDPAYA EACSNLGLVY LLQAQ**E**YADK APADINDPNY ATAQA**E**IKKK
 351 YEA**A**KPYEK ARELKPQDK LWLQGLYRVY YNL**M**GF**E**PE **E**IE**K**MM

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
34 - 48	835.8360	1669.6574	1669.8522	-0.1948	0 K.AEQLINEALTN PE TK.D (Ions score 74)
49 - 62	776.2740	1550.5334	1550.7365	-0.2031	0 K.DNAAT W DVAGYIQK.R (Ions score 64)
121 - 141	797.0170	2388.0292	2387.2961	0.7331	0 K.TIL A ERP N LINGG IQ YFNLN.K (Ions score 32)
203 - 219	663.5640	1987.6702	1986.9357	0.7345	1 K.DKENGQFAMQL L TDAYK.A Oxidation (M) (Ions score 51)
203 - 219	994.8670	1987.7194	1986.9357	0.7838	1 K.DKENGQFAMQL L TDAYK.A Oxidation (M) (Ions score 26)
238 - 260	930.6050	2788.7932	2789.2245	-0.4313	0 K.FPEN Q YF F ANLVDY Y SSSNQNDK.A (Ions score 57)
379 - 394	995.8420	1989.6694	1989.9030	-0.2335	0 R.VVYN L MG PE FE E IEK.M Oxidation (M) (Ions score 68)
379 - 396	762.6100	2284.8082	2283.9738	0.8344	1 R.VVYN L MG PE FE E IEKMM.- 3 Oxidation (M) (Ions score 19)



LOCUS YP_212118 396 aa linear BCT 01-MAY-2009
 DEFINITION hypothetical protein BF2494 [Bacteroides fragilis NCTC 9343].
 ACCESSION YP_212118
 VERSION YP_212118.1 GI:60681974
 DBLINK Project:46
 DBSOURCE REFSEQ: accession [NC_003228.3](#)
 KEYWORDS
 SOURCE Bacteroides fragilis NCTC 9343
 ORGANISM **Bacteroides fragilis NCTC 9343**
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 396)
 AUTHORS Cerdeno-Tarraga,A.M., Patrick,S., Crossman,L.C., Blakely,G.,
 Abratt,V., Lennard,N., Poxton,I., Duerden,B., Harris,B.,
 Quail,M.A., Barron,A., Clark,L., Corton,C., Doggett,J.,
 Holden,M.T., Larke,N., Line,A., Lord,A., Norbertczak,H., Ormond,D.,
 Price,C., Rabinowitsch,E., Woodward,J., Barrell,B. and Parkhill,J.
 TITLE Extensive DNA inversions in the B. fragilis genome control variable
 gene expression
 JOURNAL Science 307 (5714), 1463-1465 (2005)
 PUBMED [15746427](#)
 REFERENCE 2 (residues 1 to 396)
 AUTHORS Cerdeno-Tarraga,A.M.
 TITLE Direct Submission
 JOURNAL Submitted (29-JUL-2004) Pathogen Sequencing Unit, Sanger Institute,
 Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SA, United
 Kingdom
 REFERENCE 3 (residues 1 to 396)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (08-APR-2002) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA

COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
NCBI review. The reference sequence was derived from CAH08194.
Method: conceptual translation.

FEATURES Location/Qualifiers

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/organism="Bacteroides fragilis NCTC 9343"
/strain="ATCC 25285; NCTC 9343"
/db_xref="ATCC:25285"
/db_xref="taxon:272559"

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/note="putative PEP-CTERM system TPR-repeat lipoprotein;
TIGR02917"
/db_xref="CDD:131963"

Region 247..323
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/note="Tetratricopeptide repeat domain; typically contains
34 amino acids
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X(2)-[ASL]-X(4)-[PKE] is the consensus sequence; found in
a variety of organisms including bacteria, cyanobacteria,
yeast, fungi...; cd00189"
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Site order(249,261,265,268,283,295,299,302,317)
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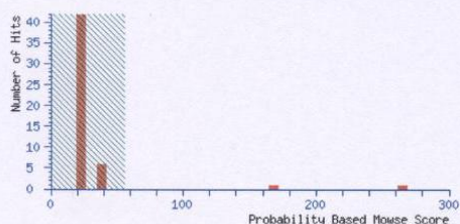
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 2101
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2101) 10_RI4_01_1304.d\SSP (2101) 10_RI4_01_1304.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:14:56 GMT
 Protein hits : [gi|53715478](#) 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]
 [gi|189460676](#) hypothetical protein BACOP_01323 [Bacteroides coprocola DSM 17136]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 56 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53715478](#) Mass: 12688 Score: 265 Queries matched: 4 emPAI: 0.61
 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 60	565.2530	1128.4914	1128.6390	-0.1476	0	33	13	1	K.EVNELATILK.E
<input checked="" type="checkbox"/> 71	644.2390	1286.4634	1287.6558	-1.1923	0	52	0.14	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/> 72	666.8020	1331.5894	1331.7449	-0.1554	0	88	3.6e-05	1	K.AFAEQLVNLTVK.E
<input checked="" type="checkbox"/> 234	852.4230	2554.2472	2554.2550	-0.0079	0	91	1.1e-05	1	K.EEYGIPEAAAVAVAGPAAGAAAEEK.S

Proteins matching the same set of peptides:
[gi|150003396](#) Mass: 12704 Score: 265 Queries matched: 4
 50S ribosomal protein L7/L12 [Bacteroides vulgatus ATCC 8482]
[gi|153805949](#) Mass: 12686 Score: 265 Queries matched: 4
 hypothetical protein BACCAC_00194 [Bacteroides caccae ATCC 43185]

2. [gi|189460676](#) Mass: 12771 Score: 165 Queries matched: 3 emPAI: 0.27
 hypothetical protein BACOP_01323 [Bacteroides coprocola DSM 17136]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
60	565.2530	1128.4914	1128.6390	-0.1476	0	33	13	1	K.EVNELATILK.E
71	644.2390	1286.4634	1285.6765	0.7869	0	43	1	2	K.VLEEAGAEVELK.-
72	666.8020	1331.5894	1331.7449	-0.1554	0	88	3.6e-05	1	K.AFAEQLVNLTVK.E

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 180	725.9040	2174.6902	2175.0960	-0.4059	0	33	6.2	1	GPLTQGYLGTFOYSAPFQLR
<input checked="" type="checkbox"/> 160	1041.9470	2081.8794	2083.0334	-1.1540	2	32	9.8	1	SGYERVDIVENKGFVSR
<input checked="" type="checkbox"/> 213	763.0340	2286.0802	2287.0361	-0.9560	1	29	20	1	SGGIGLTQAFYRAMMEAQNGGE
<input checked="" type="checkbox"/> 146	673.3010	2016.8812	2017.0844	-0.2032	0	29	21	1	ITQLYEGTNGIQALDLIR
<input checked="" type="checkbox"/> 250	970.7500	2909.2282	2908.6320	0.5962	0	29	13	1	QRPILAEGPAILLVNLPENIGMVAR
65	584.5490	1750.6252	1749.9050	0.7202	0	27	51	1	HLVTALFELEDGGPPR

MASCOT Mascot Search Results

Protein View

Match to: **gi|53715478** Score: **265**
50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2101) 10_RI4_01_1304.d\SSP (2101) 10_RI4_01_1304.mgf

Nominal mass (M_r): **12688**; Calculated pI value: **4.71**
 NCBI BLAST search of [gi|53715478](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60683451](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253566657](#) from [Bacteroides sp. 3_2_5](#)
[gi|255011634](#) from [Bacteroides fragilis 3_1_12](#)
[gi|265767535](#) from [Bacteroides sp. 2_1_16](#)
[gi|81313501](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|81608231](#) from [Bacteroides fragilis](#)
[gi|52218343](#) from [Bacteroides fragilis YCH46](#)
[gi|60494885](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251944829](#) from [Bacteroides sp. 3_2_5](#)
[gi|263252840](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **50%**

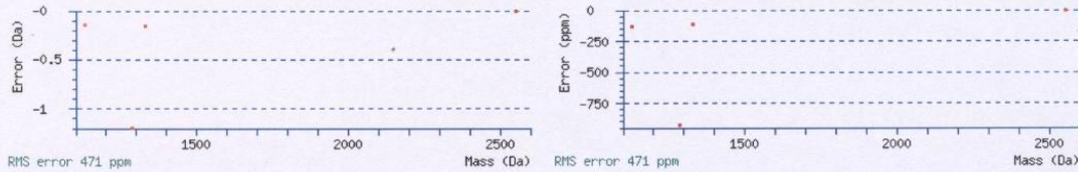
Matched peptides shown in **Bold Red**

1 MADLKAFAEQ LVNLTVKEVN ELATILKEEY GIEPAAAAVA VAAGPAAGAA
51 AAEKSSPDV VLKSAGAAKL QVVKAVKEAC GLGLKEAKDM VDGAPSVVKE
101 GLAKDEAESL KKTLEEGAE VELK

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss Sequence
6 - 17	666.8020	1331.5894	1331.7449	-0.1554	0 K.AFAEQLVNLTVK.E (Ions score 88)
18 - 27	565.2530	1128.4914	1128.6390	-0.1476	0 K.EVNELATILK.E (Ions score 33)
28 - 55	852.4230	2554.2472	2554.2550	-0.0079	0 K.EEYGIEPAAAAVVAAGPAAGAAAAEEK.S (Ions score 91)
113 - 124	644.2390	1286.4634	1287.6558	-1.1923	0 K.TLEEAGAEVELK.- (Ions score 52)



LOCUS YP_101470 124 aa linear BCT 26-APR-2009
 DEFINITION 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46].
 ACCESSION YP_101470
 VERSION YP_101470.1 GI:53715478
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS
 SOURCE Bacteroides fragilis YCH46
 ORGANISM Bacteroides fragilis YCH46
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 124)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 124)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 124)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT REVIEWED REFSEQ: This record has been curated by NCBI staff. The
 reference sequence was derived from BAD50936.

Method: conceptual translation.

FEATURES

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 /strain="YCH46"
 /db_xref="taxon:295405"

Protein 1..124
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 /calculated_mol_wt=12506

Region 2..123
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 /db_xref="CDD:100102"

Site order(5,8,12,25,30)
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 /note="peripheral dimer interface"
 /db_xref="CDD:100102"

Site order(19,22..23,27,30)
 /site_type="other"
 /note="L10 interface"
 /db_xref="CDD:100102"

Site order(69..70,73..74,77,84..85,88)
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 /note="L11 interface"
 /db_xref="CDD:100102"

Site order(70,74,77,84..85,88)
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 /note="putative EF-Tu interaction site"
 /db_xref="CDD:100102"

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 /db_xref="CDD:100102"

CDS 1..124
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 /note="present in two forms; L12 is normal, while L7 is aminoacylated at the N-terminal serine; the only multicopy ribosomal protein; 4:1 ratio of L7/L12 per ribosome; two L12 dimers bind L10; critically important for translation efficiency and fidelity; stimulates GTPase activity of translation factors"
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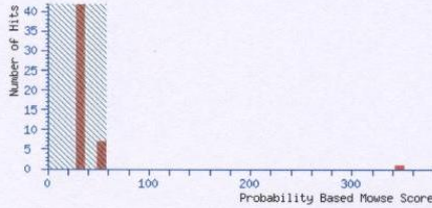
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 2102
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2102)_RE11_01_1007.d\SSP (2102)_RE11_01_1007.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:16:07 GMT
 Protein hits : [gi|53715478](#) 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: **Peptide Summary** [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53715478](#) Mass: 12688 Score: 347 Queries matched: 30 emPAI: 1.05
 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
33	447.6820	893.3494	893.4858	-0.1364	0	(35)	8.2	2	K.SSFDVVLK.S
<input checked="" type="checkbox"/>	34	447.6950	893.3754	893.4858	-0.1104	0 (34)	11	1	K.SSFDVVLK.S
<input checked="" type="checkbox"/>	35	447.7160	893.4174	893.4858	-0.0684	0 47	0.54	1	K.SSFDVVLK.S
<input checked="" type="checkbox"/>	36	447.7320	893.4494	893.4858	-0.0364	0 (41)	2.2	2	K.SSFDVVLK.S
<input checked="" type="checkbox"/>	37	447.7340	893.4534	893.4858	-0.0324	0 (36)	6.3	1	K.SSFDVVLK.S
<input checked="" type="checkbox"/>	38	447.7350	893.4554	893.4858	-0.0304	0 (42)	1.9	1	K.SSFDVVLK.S
<input checked="" type="checkbox"/>	39	447.7600	893.5054	893.4858	0.0196	0 (32)	1.6	1	K.SSFDVVLK.S
<input checked="" type="checkbox"/>	65	565.2660	1128.5174	1128.6390	-0.1216	0 34	12	1	K.EVVELATILK.E
<input checked="" type="checkbox"/>	66	567.2240	1132.4334	1132.5434	-0.1100	0 41	2.6	1	K.DMVDGAPSVVK.E + Oxidation (M)
<input checked="" type="checkbox"/>	69	644.7600	1287.5054	1287.6558	-0.1503	0 (48)	0.42	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/>	70	644.7780	1287.5414	1287.6558	-0.1143	0 (63)	0.016	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/>	71	644.7810	1287.5474	1287.6558	-0.1083	0 (51)	0.24	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/>	72	644.7840	1287.5534	1287.6558	-0.1023	0 (69)	0.0041	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/>	73	644.7860	1287.5574	1287.6558	-0.0983	0 82	0.00022	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/>	74	644.8000	1287.5854	1287.6558	-0.0703	0 (68)	0.0054	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/>	75	644.8140	1287.6134	1287.6558	-0.0423	0 (44)	2.2	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/>	76	644.8420	1287.6694	1287.6558	0.0137	0 (66)	0.0086	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/>	82	444.8610	1331.5612	1331.7449	-0.1837	0 (42)	1.7	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	83	666.8020	1331.5894	1331.7449	-0.1554	0 (84)	0.00011	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	84	666.8030	1331.5914	1331.7449	-0.1534	0 97	5.7e-06	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	85	666.8080	1331.6014	1331.7449	-0.1434	0 (75)	0.00091	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	86	666.8130	1331.6114	1331.7449	-0.1334	0 (77)	0.00054	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	87	666.8320	1331.6494	1331.7449	-0.0954	0 (89)	3.8e-05	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	88	666.8340	1331.6534	1331.7449	-0.0914	0 (88)	5.1e-05	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	89	666.8400	1331.6654	1331.7449	-0.0794	0 (80)	0.00033	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	90	666.8420	1331.6694	1331.7449	-0.0754	0 (88)	4.6e-05	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	91	666.8660	1331.7174	1331.7449	-0.0274	0 (94)	1.2e-05	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	92	666.8780	1331.7414	1331.7449	-0.0034	0 (80)	0.00029	1	K.AFAEQLVNLTK.E
<input checked="" type="checkbox"/>	107	730.8450	1459.6754	1460.7181	-1.0426	1 36	7.8	1	K.EAKMVDGAPSVVK.E + Oxidation (M)
<input checked="" type="checkbox"/>	136	880.4590	1758.9034	1758.9880	-0.0845	1 12	1.6e+03	1	M.ADLKAFAEQLVNLTK.E

Peptide matches not assigned to protein hits: (no details means no match)

Query Observed Mr (expt) Mr (calc) Delta Miss Score Expect Rank Peptide

Mascot Search Results

Protein View

Match to: **gi|53715478** Score: **347**
50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2102)_RE11_01_1007.d\SSP (2102)_RE11_01_1007.mgf

Nominal mass (M_n): **12688**; Calculated pI value: **4.71**
 NCBI BLAST search of **gi|53715478** against nr
 Unformatted **sequence string** for pasting into other applications

Taxonomy: **Bacteroides fragilis YCH46**
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60693451](#) from **Bacteroides fragilis NCTC 9343**
[gi|253566657](#) from **Bacteroides sp. 3_2_5**
[gi|255011634](#) from **Bacteroides fragilis 3_1_12**
[gi|265767535](#) from **Bacteroides sp. 2_1_16**
[gi|81313501](#) from **Bacteroides fragilis NCTC 9343**
[gi|81608231](#) from **Bacteroides fragilis**
[gi|52218343](#) from **Bacteroides fragilis YCH46**
[gi|60494885](#) from **Bacteroides fragilis NCTC 9343**
[gi|251944829](#) from **Bacteroides sp. 3_2_5**
[gi|263252840](#) from **Bacteroides sp. 2_1_16**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **48%**

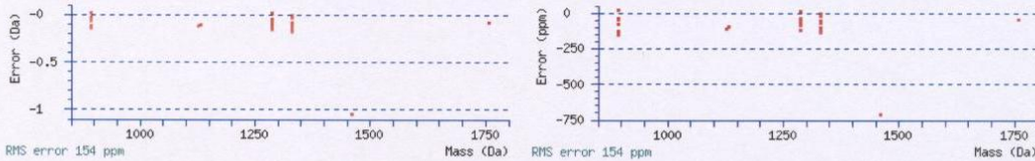
Matched peptides shown in **Bold Red**

- 1** MADLKAFAEQ LVNLTVKEVN ELATILKEEY GIEPAAAAVA VAAGPAAGAA
- 51** AAEEKSSFVDV VLK SAGA AKL QVVKAVKEAC GLGLKEAKDM VDGAPSVVKE
- 101** GLAKDEAESL KKTLEEGAE VELK

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
2 - 17	880.4590	1758.9034	1758.9880	-0.0845	1 M.ADLKAFAEQLVNLTVK.E (Ions score 12)
6 - 17	444.8610	1331.5612	1331.7449	-0.1837	0 K.AFAEQLVNLTVK.E (Ions score 42)
6 - 17	666.8020	1331.5894	1331.7449	-0.1554	0 K.AFAEQLVNLTVK.E (Ions score 84)
6 - 17	666.8030	1331.5914	1331.7449	-0.1534	0 K.AFAEQLVNLTVK.E (Ions score 97)
6 - 17	666.8080	1331.6014	1331.7449	-0.1434	0 K.AFAEQLVNLTVK.E (Ions score 75)
6 - 17	666.8130	1331.6114	1331.7449	-0.1334	0 K.AFAEQLVNLTVK.E (Ions score 77)
6 - 17	666.8320	1331.6494	1331.7449	-0.0954	0 K.AFAEQLVNLTVK.E (Ions score 89)
6 - 17	666.8340	1331.6534	1331.7449	-0.0914	0 K.AFAEQLVNLTVK.E (Ions score 88)
6 - 17	666.8400	1331.6654	1331.7449	-0.0794	0 K.AFAEQLVNLTVK.E (Ions score 80)
6 - 17	666.8420	1331.6694	1331.7449	-0.0754	0 K.AFAEQLVNLTVK.E (Ions score 88)
6 - 17	666.8660	1331.7174	1331.7449	-0.0274	0 K.AFAEQLVNLTVK.E (Ions score 94)
6 - 17	666.8780	1331.7414	1331.7449	-0.0034	0 K.AFAEQLVNLTVK.E (Ions score 80)
18 - 27	565.2660	1128.5174	1128.6390	-0.1216	0 K.EVNELATILK.E (Ions score 34)
56 - 63	447.8620	893.3494	893.4858	-0.1364	0 K.SSFDVVLK.S (Ions score 35)
56 - 63	447.6950	893.3754	893.4858	-0.1104	0 K.SSFDVVLK.S (Ions score 34)
56 - 63	447.7160	893.4174	893.4858	-0.0684	0 K.SSFDVVLK.S (Ions score 47)
56 - 63	447.7320	893.4494	893.4858	-0.0364	0 K.SSFDVVLK.S (Ions score 41)
56 - 63	447.7340	893.4534	893.4858	-0.0324	0 K.SSFDVVLK.S (Ions score 36)
56 - 63	447.7350	893.4554	893.4858	-0.0304	0 K.SSFDVVLK.S (Ions score 42)
56 - 63	447.7600	893.5054	893.4858	0.0196	0 K.SSFDVVLK.S (Ions score 32)
86 - 99	730.8450	1459.6754	1460.7181	-1.0426	1 K.EAKDMVDGAPSVVK.E Oxidation (M) (Ions score 36)
89 - 99	567.2240	1132.4334	1132.5434	-0.1100	0 K.DMVDGAPSVVK.E Oxidation (M) (Ions score 41)
113 - 124	644.7600	1287.5054	1287.6558	-0.1503	0 K.TLEEAGAEVELK.- (Ions score 48)
113 - 124	644.7780	1287.5414	1287.6558	-0.1143	0 K.TLEEAGAEVELK.- (Ions score 63)
113 - 124	644.7810	1287.5474	1287.6558	-0.1083	0 K.TLEEAGAEVELK.- (Ions score 51)
113 - 124	644.7840	1287.5534	1287.6558	-0.1023	0 K.TLEEAGAEVELK.- (Ions score 69)
113 - 124	644.7860	1287.5574	1287.6558	-0.0983	0 K.TLEEAGAEVELK.- (Ions score 82)
113 - 124	644.8000	1287.5854	1287.6558	-0.0703	0 K.TLEEAGAEVELK.- (Ions score 68)
113 - 124	644.8140	1287.6134	1287.6558	-0.0423	0 K.TLEEAGAEVELK.- (Ions score 44)
113 - 124	644.8420	1287.6694	1287.6558	0.0137	0 K.TLEEAGAEVELK.- (Ions score 66)



LOCUS YP_101470 124 aa linear BCT 26-APR-2009
 DEFINITION 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46].
 ACCESSION YP_101470

VERSION YP_101470.1 GI:53715478
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM Bacteroides fragilis YCH46
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 124)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 124)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 124)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
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 Method: conceptual translation.
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 /coded_by="complement(NC_006347.1:4790804..4791178)"
 /note="present in two forms; L12 is normal, while L7 is
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 efficiency and fidelity; stimulates GTPase activity of
 translation factors"
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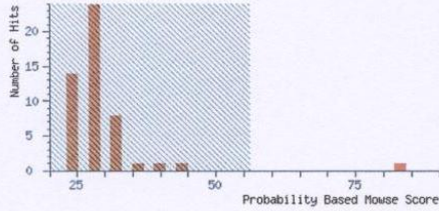
Mascot: <http://www.matrixscience.com/>

{MATRIX} Mascot Search Results
{SCIENCE}

User : LAKSHMY MANICKAN
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 2401
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2401) 10_RK3_01_1421.d\SSP (2401) 10_RK3_01_1421.mgf
 Database : NCBIInr 20090522 (8876587 sequences; 3036162093 residues)
 Taxonomy : Bacteria (Eubacteria) (4773688 sequences)
 Timestamp : 30 May 2009 at 10:35:03 GMT
 Protein hits : [gi|53713032](#) GrpE protein [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 56 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53713032](#) Mass: 22007 Score: 83 Queries matched: 2
 GrpE protein [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 119	652.2180	1302.4214	1302.6489	-0.2275	0	48	0.3	1	K.SILPVIDDMER.A + Oxidation (M)
<input checked="" type="checkbox"/> 240	860.6480	2578.9222	2579.2391	-0.3169	0	35	3.7	1	K.DQPLDLDYHEAIAVIPAPTEEQK.G

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 205	694.9240	2081.7502	2080.9338	0.8164	0	32	8.8	1	TGTGESEAEAVGFVQVR
<input checked="" type="checkbox"/> 228	1112.9430	2223.8714	2223.0696	0.8019	1	31	11	1	DYLRDDVDQVLIDSPDAFK
<input checked="" type="checkbox"/> 224	737.6170	2209.8292	2209.1062	0.7230	2	30	13	1	NSSNIFQIKKPHNTQTMR + Oxidation (M)
<input checked="" type="checkbox"/> 192	989.4900	1976.9654	1975.9455	1.0199	2	30	40	1	DMPEGVDTDKVIQRAMR + Oxidation (M)
<input checked="" type="checkbox"/> 227	742.2680	2223.7822	2223.0049	0.7773	0	29	17	1	QMPAGYETQIGEGGMLSGGQR
<input checked="" type="checkbox"/> 124	660.2840	1318.5534	1317.6421	0.9114	0	29	34	1	ANGLPVICMETR
<input checked="" type="checkbox"/> 54	362.3120	1083.9142	1084.6142	-0.7000	1	28	45	1	AVPIPGYGRR
<input checked="" type="checkbox"/> 162	562.7890	1685.3452	1684.7800	0.5651	0	27	30	1	MFTLADDATCIIGTR
<input checked="" type="checkbox"/> 129	671.2260	1340.4374	1339.6884	0.7490	0	27	35	1	EALTFPLPEHR
<input checked="" type="checkbox"/> 212	700.9180	2099.7322	2099.0932	0.6389	1	26	39	1	QSLDLNIIIVTHDKESMK + Oxidation (M)
<input checked="" type="checkbox"/> 106	555.3090	1662.9052	1661.8195	1.0857	2	25	1.3e+02	1	DDWIGQAKALMRDK + Oxidation (M)
<input checked="" type="checkbox"/> 144	750.8970	2249.6692	2250.0773	-0.4081	0	25	1.2e+02	1	SSGAAVAMVFPVQNLTPQDMR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 109	579.1220	1734.3442	1733.8836	0.4606	0	24	1.4e+02	1	LIELVSENFTDPTR
<input checked="" type="checkbox"/> 78	438.3790	1312.1152	1311.6129	0.5023	2	24	1.6e+02	1	YGESPSQSMKKA
<input checked="" type="checkbox"/> 173	910.6630	2728.9672	2728.3160	0.6512	2	24	1.2e+02	1	QYSAQSAAMRETGVANSTLMDPKLE + 2 Oxidation (M)
<input checked="" type="checkbox"/> 142	705.5310	2113.5712	2113.0209	0.5503	1	23	1.3e+02	1	MFTTDRITDTATAVADVLE + Oxidation (M)
<input checked="" type="checkbox"/> 189	650.5210	1948.5412	1947.9915	0.5497	2	23	66	1	SSEAAPGRGSPAGVAPPVW
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<input checked="" type="checkbox"/> 191	656.3610	1966.0612	1967.1316	-1.0704	1	23	1.1e+02	1	LQAAGIGAVRGGALYFVLNK
<input checked="" type="checkbox"/> 188	970.8420	1939.6694	1940.1129	-0.4434	1	23	76	1	QKVIKPLVSEVQVQSMK
<input checked="" type="checkbox"/> 195	670.0460	2007.1162	2008.0710	-0.9549	2	23	1.1e+02	1	IVRVGGMMVAPGTGRNVTK + Oxidation (M)
<input checked="" type="checkbox"/> 241	866.3330	2595.9772	2596.2519	-0.2748	1	23	68	1	APFTAELDASDIEKGYPHFMLK + Oxidation (M)
<input checked="" type="checkbox"/> 219	1069.6480	3205.9222	3205.5251	0.3971	2	22	1.6e+02	1	EAVHETADGRQFLVMHGDVSVVRYAK
<input checked="" type="checkbox"/> 217	709.5750	2125.7032	2126.1372	-0.4340	0	22	88	1	VAVVGFAPVANQDIGNVNLTK
<input checked="" type="checkbox"/> 169	904.6440	2710.9102	2711.3914	-0.4812	0	22	1.7e+02	1	TELSQYAIISVLIDMGLSLMGLLWQK + Oxidation (M)
<input checked="" type="checkbox"/> 235	766.9700	2297.8882	2298.0223	-0.1341	0	22	93	1	YDADGQFAGYATDADGKPVNTR

Mascot Search Results

Protein View

Match to: gi|53713032 Score: 83
 GrpE protein [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2401) 10_RK3_01_1421.d\SSP (2401) 10_RK3_01_1421.mgf

Nominal mass (M_r): 22007; Calculated pI value: 4.71
 NCBI BLAST search of gi|53713032 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60681311](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|81315621](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|81690697](#) from [Bacteroides fragilis](#)
[gi|52215897](#) from [Bacteroides fragilis YCH46](#)
[gi|60492745](#) from [Bacteroides fragilis NCTC 9343](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 17%

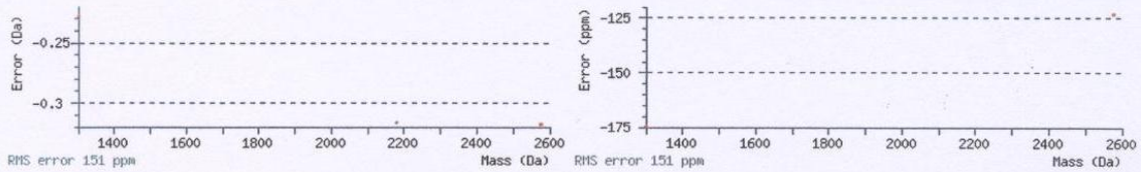
Matched peptides shown in **Bold Red**

1 MDPKEKKTQ EEELKVDDIQ DTVEGQSQNE EATEATEPLT AEEKLEKELK
 51 EAQAQIEDQK DKYLRLSAEF DNYRKRITVKE KAEILINGGE KSIKSI~~LPVI~~
 101 **DDMERALTTM** ETATDVNAVK EGVELIYNKF LSILSQDGVK VIETKQ**QPLD**
 151 **TDYHEALAVI** **PAPTEEQK**GK ILDCVQTGYT LNGKVI~~RHAK~~ VVVGE

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
95 - 105	652.2180	1302.4214	1302.6489	-0.2275	0	K.SILPVIDDMER.A Oxidation (M) (Ions score 48)
146 - 168	860.6480	2578.9222	2579.2391	-0.3169	0	K.DQPLD TDYHEALAVI PAPTEEQK.G (Ions score 35)



LOCUS YP_099024 195 aa linear BCT 26-APR-2009
 DEFINITION GrpE protein [Bacteroides fragilis YCH46].
 ACCESSION YP_099024
 VERSION YP_099024.1 GI:53713032
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 195)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 195)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 195)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD48490](#).
 Method: conceptual translation.
 FEATURES Location/Qualifiers

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            /strain="YCH46"
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Region     56..193
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            /note="GrpE is the adenine nucleotide exchange factor of
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            which enables rebinding of ATP, one step in the Hsp70
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            /note="hsp70 (ATPase domain) interactions"
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CDS        1..195
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            /note="similar to gp:AE016931_34 [Bacteroides
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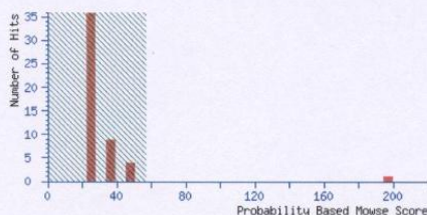
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 2701
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2701) 100_RH7_01_1762.d\SSP (2701) 100_RH7_01_1762.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Bubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:17:22 GMT
 Protein hits : [gi|53714026](#) ATP synthase subunit E [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|53714026](#) Mass: 21789 Score: 197 Queries matched: 4 empAI: 0.54
 ATP synthase subunit E [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 120	566.2910	1130.5674	1130.6448	-0.0773	0	58	0.054	1	K.LFAGQAVNALK.S
<input checked="" type="checkbox"/> 160	671.2750	1340.5354	1340.6612	*-0.1258	0	40	2.5	1	K.ALFSVSPADGSYK.V
<input checked="" type="checkbox"/> 171	740.3800	1478.7454	1478.7955	-0.0501	0	34	12	1	K.AFLRPQLVEMLP.- + Oxidation (M)
<input checked="" type="checkbox"/> 192	835.2550	1668.4954	1668.7130	-0.2175	0	65	0.0053	1	K.VNFGEEEPNVYK.A + Oxidation (M)

Proteins matching the same set of peptides:
[gi|255009692](#) Mass: 21804 Score: 197 Queries matched: 4
 putative V-type ATP synthase subunit E [Bacteroides fragilis 3_1_12]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 154	651.8680	1301.7214	1301.6728	0.0487	1	41	2.4	1	VTDLDKAGAWAR
<input checked="" type="checkbox"/> 170	738.3230	1474.6314	1475.8209	-1.1894	1	36	6.3	1	TRTFVQQLINR
<input checked="" type="checkbox"/> 132	613.0830	1836.2272	1835.9377	0.2894	0	31	28	1	QSPQPALATIPVSEQDR
<input checked="" type="checkbox"/> 187	535.3750	1603.1032	1602.8035	0.2996	1	31	17	1	AEMDAGAVRALDQLK + Oxidation (M)
<input checked="" type="checkbox"/> 108	350.2350	1047.6832	1046.5032	1.1799	0	29	37	1	YQSSLLEHA
<input checked="" type="checkbox"/> 90	420.6080	1258.8022	1259.6292	-0.8270	2	29	59	1	KEHNKTMQTK + Oxidation (M)
<input checked="" type="checkbox"/> 117	551.2270	1650.6592	1650.7777	-0.1185	0	29	73	1	YYASLETFPNELSSK
<input checked="" type="checkbox"/> 72	378.4320	1132.2742	1132.5989	-0.3247	0	28	53	1	AAGITQAFAR
<input checked="" type="checkbox"/> 246	837.6220	2509.8442	2510.9844	-1.1403	1	28	22	1	ERNFCCGGGSGILMDEMMEIR + 3 Oxidation (M)
<input checked="" type="checkbox"/> 163	696.3180	1390.6214	1391.7595	-1.1380	1	28	47	1	FRLNNDLLMLK + Oxidation (M)
<input checked="" type="checkbox"/> 212	974.9210	1947.8274	1947.0360	0.7915	1	28	39	1	LLAFRDGSRPATIMDR + Oxidation (M)
<input checked="" type="checkbox"/> 98	468.2630	934.5114	934.4066	0.1049	0	27	69	1	RANCDVVK
<input checked="" type="checkbox"/> 131	612.9790	1835.9152	1835.8148	0.1003	1	26	1e+02	1	GFFEAAKDFMGSDATSR
<input checked="" type="checkbox"/> 127	599.9810	1796.9212	1796.7934	0.1278	1	26	1.1e+02	1	MHSYSPTGMSDRIGSR + Oxidation (M)
<input checked="" type="checkbox"/> 193	843.8680	2528.5822	2529.3663	-0.7841	0	26	1e+02	1	LGQLNLLAAGQAPPAAGASAPGTAVR
<input checked="" type="checkbox"/> 118	559.4540	1675.3402	1675.8601	-0.5200	2	26	1e+02	1	RASDRIASDTASRLR
<input checked="" type="checkbox"/> 58	351.0500	1050.1282	1049.5036	0.6246	1	26	1.3e+02	1	ARVGGNSMR + Oxidation (M)
<input checked="" type="checkbox"/> 174	752.4730	2254.3972	2254.0504	0.3467	2	25	1.3e+02	1	MSTRDMVVIGMDGRVVEGDR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 182	779.3900	2335.1482	2335.1334	0.0148	1	25	1.5e+02	1	MKITELLGAMVQAGMAPSSNDR + Oxidation (M)
<input checked="" type="checkbox"/> 56	350.1610	1047.4612	1048.4430	-0.9818	0	25	1.7e+02	1	SDAHLMLGR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 215	659.1260	1974.3562	1975.0487	-0.6925	1	25	55	1	DEVVDAAKGLIASGIAPGHR
<input checked="" type="checkbox"/> 116	550.4870	1648.4392	1648.8315	-0.3923	2	25	1.3e+02	1	MTNATNARAVARETK + Oxidation (M)
<input checked="" type="checkbox"/> 86	414.2450	1239.7132	1238.5457	1.1674	0	25	1.9e+02	1	ISQNMPGMAMK + 2 Oxidation (M)

MASCOT **SCIENCE** Mascot Search Results

Protein View

Match to: [gi|53714026](#) Score: 197
ATP synthase subunit E [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2701) 100_RH7_01_1762.d\SSP (2701) 100_RH7_01_1762.mgf

Nominal mass (M_n): 21789; Calculated pI value: 5.08
 NCBI BLAST search of [gi|53714026](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60682222](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253566982](#) from [Bacteroides sp. 3_2_5](#)
[gi|265764376](#) from [Bacteroides sp. 2_1_16](#)
[gi|52216891](#) from [Bacteroides fragilis YCH46](#)
[gi|60493656](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251944106](#) from [Bacteroides sp. 3_2_5](#)
[gi|263256984](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 25%

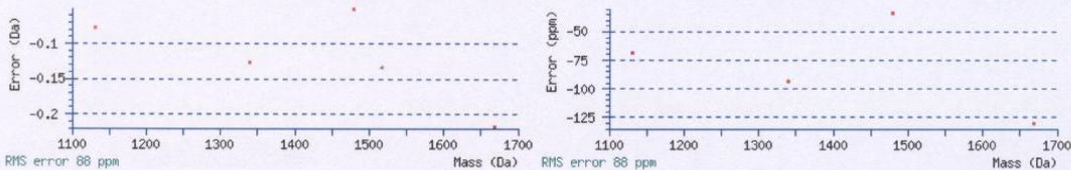
Matched peptides shown in **Bold Red**

- 1 MENKIQELTD KIYREGVEKG NEEARRLIAN AQEAKKIVE DAHKEAESII
- 51 ASSRKSADDEL TENTKSELKL **FAGQAVNALK** SEIATMVTDK IVTAPVKEFA
- 101 QNKDFLNAPI VALASKWVD EPIIISTSDA ESLKKYFAAN AKALLDKGV
- 151 IEQVNGIKAL **FSVSPADGSY KVNFGEEEFM NYFKAPLRQ LVEMLF**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
70 - 80	566.2910	1130.5674	1130.6448	-0.0773	0	K.LFAGQAVNALK.S (Ions score 58)
159 - 171	671.2750	1340.5354	1340.6612	-0.1258	0	K.ALFSVSPADGSYK.V (Ions score 40)
172 - 184	835.2550	1668.4954	1668.7130	-0.2175	0	K.VNFGEEEFMNYFK.A Oxidation (M) (Ions score 65)
185 - 196	740.3800	1478.7454	1478.7955	-0.0501	0	K.AFLRPQLVEMLF.- Oxidation (M) (Ions score 34)



LOCUS YP_100018 196 aa linear BCT 26-APR-2009
 DEFINITION ATP synthase subunit E [Bacteroides fragilis YCH46].
 ACCESSION YP_100018
 VERSION YP_100018.1 GI:53714026
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 196)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 196)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 196)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD49484](#).
 Method: conceptual translation.

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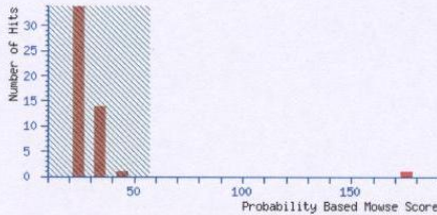
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 2704
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2704) 1000_RG17_01_1193.d\SSP (2704) 1000_RG17_01_1193.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:18:32 GMT
 Protein hits : [gi|53714026](#) ATP synthase subunit E [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ion score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ion scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ion scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold p<: Max. number of hits:

Standard scoring: MudPIT scoring Ions score or expect cut-off: Show sub-sets:

Show pop-ups: Suppress pop-ups Sort unassigned: Require bold red:

Select All Select None Search Selected Error tolerant

1. [gi|53714026](#) Mass: 21789 Score: 175 Queries matched: 5 emPAI: 0.33
 ATP synthase subunit E [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 74	566.3400	1130.6654	1130.6448	0.0207	0	54	0.12	1	K.LFAGQAVNALK.S
<input checked="" type="checkbox"/> 75	566.7860	1131.5574	1130.6448	0.9127	0	(26)	82	1	K.LFAGQAVNALK.S
<input checked="" type="checkbox"/> 87	671.3080	1340.6014	1340.6612	-0.0598	0	30	26	1	K.ALFSVSPADGSYK.V
<input checked="" type="checkbox"/> 101	740.3580	1478.7014	1478.7955	-0.0941	0	33	12	1	K.AFLRPQLVEMLF.- + Oxidation (M)
<input checked="" type="checkbox"/> 155	1059.9960	2117.9774	2117.0892	0.8883	1	57	0.045	1	K.WSVDEPIIISTDAESLKK.Y

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 100	738.3090	1474.6034	1474.8395	-0.2360	1	49	0.38	1	ELIEISKFLDDR
<input checked="" type="checkbox"/> 167	767.3490	2299.0252	2298.2947	0.7305	1	34	7.7	1	QRPNQAQALETLDTIKFLPK
<input checked="" type="checkbox"/> 80	651.8460	1301.6774	1301.6463	0.0311	0	31	23	1	VTDIDVNVNECK
<input checked="" type="checkbox"/> 107	786.0440	2355.1102	2354.2376	0.8725	2	31	34	1	TGDLAAGPVLGKVRAMVDDSGAR
<input checked="" type="checkbox"/> 83	657.1320	1312.2494	1312.6987	-0.4492	0	29	23	1	TLFAGTSAQAPATAK
<input checked="" type="checkbox"/> 161	1106.0070	2209.9994	2210.1306	-0.1312	1	28	36	1	CQIVYNVLRKHIQFGAYR
<input checked="" type="checkbox"/> 35	370.1030	738.1914	737.3014	0.8900	0	28	59	1	SMGGEGGK + Oxidation (M)
<input checked="" type="checkbox"/> 163	742.6100	2224.8082	2224.0318	0.7764	2	27	31	1	DKDNDFGEIVSEAVEVCKR
<input checked="" type="checkbox"/> 39	379.1450	1134.4132	1135.5696	-1.1564	0	26	1.4e+02	1	HTIYTLMNK + Oxidation (M)
<input checked="" type="checkbox"/> 174	1209.6530	3625.9372	3624.9913	0.9459	1	26	89	1	QSNILLTBLPLMTPTGTFVINGTERVVVSQIVR
<input checked="" type="checkbox"/> 116	836.3120	1670.6094	1671.7351	-1.1257	1	26	52	1	ESFADEHPFYRMK + Oxidation (M)
<input checked="" type="checkbox"/> 102	740.9110	2219.7112	2220.1653	-0.4541	0	26	1.1e+02	1	AYVLIPLLGSLSLGMGNFK
<input checked="" type="checkbox"/> 153	694.3570	2080.0492	2079.9248	0.1244	2	25	72	1	MREMMEQAEARNAQNK + Oxidation (M)
<input checked="" type="checkbox"/> 112	808.6520	2422.9642	2423.3830	-0.4188	0	25	1.1e+02	1	LQQALLGPAGMLSLGLVLMMLR
<input checked="" type="checkbox"/> 108	795.2000	2382.5782	2383.1655	-0.5874	2	24	1.3e+02	1	EDKSIIGYLESKEGNDFIGR
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<input checked="" type="checkbox"/> 71	562.3360	1122.6574	1123.7077	-1.0503	2	24	2.1e+02	1	VVASPAKRALL
<input checked="" type="checkbox"/> 136	641.2960	1920.8662	1920.0316	0.8346	0	23	1.1e+02	1	LTLNLNSIDSLNAIYTR
<input checked="" type="checkbox"/> 120	842.6510	2524.9612	2524.3618	0.5994	1	23	1.5e+02	1	MPIAPGSTLGLGGQLGRMTALAAR + Oxidation (M)
<input checked="" type="checkbox"/> 124	870.3300	2607.9682	2607.3101	0.6580	0	23	1.6e+02	1	SVLEVSEIINEGLSAPLVNNDMK + Oxidation (M)
<input checked="" type="checkbox"/> 88	672.3260	2013.9562	2013.9322	0.0240	0	23	2.7e+02	1	MPAGMSTASPSPLPAPMR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 144	664.5690	1990.6852	1991.0146	-0.3294	1	23	86	1	KNVFMNTFDKPELSSR + Oxidation (M)
<input checked="" type="checkbox"/> 31	360.2380	1077.6922	1077.6546	0.0375	0	23	3.3e+02	1	LALPPVDVVR
<input checked="" type="checkbox"/> 115	827.1690	2478.4852	2478.2074	0.2778	1	23	1.8e+02	1	IRLDGQDLGAQMVQAGQAWSYR + Oxidation (M)

Mascot Search Results

Protein View

Match to: [gi|53714026](#) Score: 175
ATP synthase subunit E [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2704) 1000_RG17_01_1193.d\SSP (2704) 1000_RG17_01_1193.mgf

Nominal mass (M_n): **21789**; Calculated pI value: **5.08**
 NCBI BLAST search of [gi|53714026](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60682222](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253566982](#) from [Bacteroides sp. 3_2_5](#)
[gi|265764376](#) from [Bacteroides sp. 2_1_16](#)
[gi|52216891](#) from [Bacteroides fragilis YCH46](#)
[gi|60493656](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251944106](#) from [Bacteroides sp. 3_2_5](#)
[gi|263256984](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **28%**

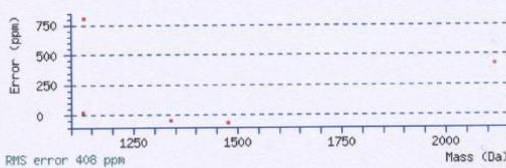
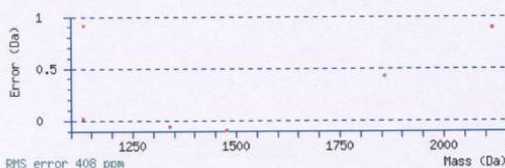
Matched peptides shown in **Bold Red**

- 1 MENKIQLTD K IYREGVEKG NEEARRLIAN AQEEAKKIVE DAKKEAESII
- 51 ASSRKSADLEL TENTKSELKL **FAGQAVNALK** SBIATMVTDK IVTAPVKEFA**
- 101 QNKDFLNAFI VALASKWSVD **EPIIISTSDA** ESLKYYFAAN AKALLDKGVT**
- 151 IEQVNGIKAL **FSVSPADGSY** KVNFGEEEFM NYFKALRPQ LVEMLF**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
70 - 80	566.3400	1130.6654	1130.6448	0.0207	0	K.LFAGQAVNALK.S (Ions score 54)
70 - 80	566.7860	1131.5574	1130.6448	0.9127	0	K.LFAGQAVNALK.S (Ions score 26)
117 - 135	1059.9960	2117.9774	2117.0892	0.8883	1	K.WSVDEPIIISTSDAESLKK.Y (Ions score 57)
159 - 171	671.3080	1340.6014	1340.6612	-0.0598	0	K.ALFSVSPADGSYK.V (Ions score 30)
185 - 196	740.3580	1478.7014	1478.7955	-0.0941	0	K.AFLRPQLVEMLF.- Oxidation (M) (Ions score 33)



LOCUS YP_100018 196 aa linear BCT 26-APR-2009
 DEFINITION ATP synthase subunit E [Bacteroides fragilis YCH46].
 ACCESSION YP_100018
 VERSION YP_100018.1 GI:53714026
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 196)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 196)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 196)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD49484](#).
 Method: conceptual translation.
 FEATURES Location/Qualifiers

source 1..196
/organism="Bacteroides fragilis YCH46"
/strain="YCH46"
/db_xref="taxon:295405"
Protein 1..196
/product="ATP synthase subunit E"
/calculated_mol_wt=21672
Region 1..195
/region_name="ATP-synt_B"
/note="ATP synthase B/B' CF(0); cl07975"
/db_xref="CDD:143711"
Region 18..196
/region_name="vATP-synt_E"
/note="ATP synthase (E/31 kDa) subunit; pfam01991"
/db_xref="CDD:110943"
CDS 1..196
/locus_tag="BF2734"
/coded_by="complement(NC_006347.1:3139699..3140289)"
/note="similar to gp:AE016931_92 [Bacteroides
thetaiotaomicron VPI-5482], percent identity 89 in 196 aa,
BLASTP E(): 3e-95"
/transl_table=11
/db_xref="GeneID:3083533"

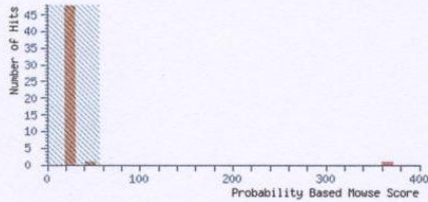
Mascot: <http://www.matrixscience.com/>

MASCOT (SCIENCE) **Mascot Search Results**

User : LAKSHMY MANICKAN
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 2804
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2804) 10_RJ17_01_1396.d\SSP (2804) 10_RJ17_01_1396.mgf
 Database : NCBInr 20090522 (8876587 sequences; 3036162093 residues)
 Taxonomy : Bacteria (Eubacteria) (4773688 sequences)
 Timestamp : 30 May 2009 at 10:39:02 GMT
 Protein hits : [gi|53713703](#) TPR repeat-containing protein [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 56 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p < 0.05$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53713703](#) Mass: 46634 Score: 366 Queries matched: 8 emPAI: 0.51
 TPR repeat-containing protein [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 125	776.2740	1550.5334	1550.7365	-0.2031	0	64	0.0061	1	K.DNAATWDVAGYIQK.R
<input checked="" type="checkbox"/> 140	835.8360	1669.6574	1669.8522	-0.1948	0	74	0.00071	1	K.ABQLINEALTNPETK.D
<input checked="" type="checkbox"/> 185	663.5640	1987.6702	1986.9357	0.7345	1	51	0.11	1	K.DKENGQFAMQLLTDAYK.A + Oxidation (M)
<input checked="" type="checkbox"/> 186	994.8670	1987.7194	1986.9357	0.7838	1	(26)	81	1	K.DKENGQFAMQLLTDAYK.A + Oxidation (M)
<input checked="" type="checkbox"/> 187	995.8420	1989.6694	1989.9030	-0.2335	0	68	0.0022	1	R.VVYNLNMGPFEIEIK.M + Oxidation (M)
<input checked="" type="checkbox"/> 233	762.6100	2284.8082	2283.9738	0.8344	1	19	1.7e+02	2	R.VVYNLNMGPFEIEIK.M - + 3 Oxidation (M)
<input checked="" type="checkbox"/> 238	797.0170	2388.0292	2387.2961	0.7331	0	32	9.9	1	K.TYLAERPNLNGIQYFNLK.N
<input checked="" type="checkbox"/> 248	930.6050	2788.7932	2789.2245	-0.4313	0	57	0.025	1	K.FPENQYFFANLVDDYSSMQNDK.A

Proteins matching the same set of peptides:
[gi|60681974](#) Mass: 45773 Score: 366 Queries matched: 8
 hypothetical protein BP2494 [Bacteroides fragilis NCTC 9343]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 221	737.9210	2210.7412	2210.1115	0.6297	0	34	5.2	1	MYILGINSVYHESAVCLLK
<input checked="" type="checkbox"/> 131	804.8580	1607.7014	1607.8125	-0.1110	0	33	12	1	MPIYNELMLPLK + Oxidation (M)
<input checked="" type="checkbox"/> 234	766.9000	2297.9182	2299.0790	-1.1609	2	32	9	1	LNQYWEIDKMRSLTDAEK + Oxidation (M)
<input checked="" type="checkbox"/> 80	629.4150	1885.2232	1885.9707	-0.7475	0	32	27	1	SIMDNLGLPVIDEIVK + Oxidation (M)
<input checked="" type="checkbox"/> 121	752.3550	2254.0432	2255.0205	-0.9773	1	30	43	1	LNKDFDGGDDKIEFLR + Oxidation (M)
<input checked="" type="checkbox"/> 225	1113.0610	2224.1074	2224.0437	0.0638	0	30	21	1	DISWGSTWAGDNDVIYQLK
<input checked="" type="checkbox"/> 44	391.1050	1170.2932	1169.6227	0.6705	0	28	49	1	LLAMEGVNPAR
<input checked="" type="checkbox"/> 88	650.7300	1299.4454	1298.7921	0.6533	2	27	37	1	ENESLKLVLQK
<input checked="" type="checkbox"/> 157	585.0580	1752.1522	1752.9312	-0.7790	2	27	33	1	VIGGNPKRFQTPDK
<input checked="" type="checkbox"/> 229	747.8320	2240.4742	2240.2277	0.2465	1	26	33	1	YQVDLLTLHATAGRALTAAK
<input checked="" type="checkbox"/> 196	681.0290	2040.0652	2040.9350	-0.8698	2	25	61	1	VREIPDMYGNKETAET + Oxidation (M)
<input checked="" type="checkbox"/> 145	563.1340	1686.3802	1685.9365	0.4436	2	25	46	1	AGLIHAFNDRKVAK
<input checked="" type="checkbox"/> 204	694.9280	2081.7622	2081.9795	-0.2174	1	25	46	1	MAEIVIGKVTLEEMGGAR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 209	700.9180	2099.7322	2099.0106	0.7216	1	25	45	1	YNIAGMAGLIGNDKFNNAK + Oxidation (M)
<input checked="" type="checkbox"/> 142	838.0410	2511.1012	2511.2329	-0.1317	1	25	1.2e+02	1	SVIYEMHVGGRTRHPNSGVAPEK
<input checked="" type="checkbox"/> 213	711.0680	2130.1822	2129.9721	0.2101	2	25	68	1	CDTEIRRYAICLSQR
<input checked="" type="checkbox"/> 82	632.9100	1895.7082	1896.0693	-0.3612	2	24	1.5e+02	1	FAVPDERAARNIGLIVR
<input checked="" type="checkbox"/> 108	707.5460	2119.6162	2119.1021	0.5140	2	24	1.2e+02	1	SGEISEQRINEAYQRIK
<input checked="" type="checkbox"/> 129	535.0980	1602.2722	1601.7005	0.5717	1	24	68	1	FDRHAMAGHGESGSK + Oxidation (M)
<input checked="" type="checkbox"/> 27	350.2230	1047.6472	1048.5414	-0.8942	0	24	2.1e+02	1	DLNPLANHR
<input checked="" type="checkbox"/> 105	703.1500	2106.4282	2105.9299	0.4982	0	24	1.4e+02	1	NDPGAGNGVPGAVMDMDVFR + Oxidation (M)
<input checked="" type="checkbox"/> 86	646.1180	1935.3322	1935.0690	0.2632	1	23	1.5e+02	1	APSIESLRQAQPVVVWR

Mascot Search Results

Protein View

Match to: gi|53713703 Score: 366
 TPR repeat-containing protein [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2804) 10_RJ17_01_1396.d\SSP (2804) 10_RJ17_01_1396.mgf

Nominal mass (M_r): 46634; Calculated pI value: 5.42
 NCBI BLAST search of gi|53713703 against nr
 Unformatted sequence string for pasting into other applications

Taxonomy: Bacteroides fragilis YCH46
 Links to retrieve other entries containing this sequence from NCBI Entrez:
 gi|253565651 from Bacteroides sp. 3_2_5
 gi|265764027 from Bacteroides sp. 2_1_16
 gi|52216568 from Bacteroides fragilis YCH46
 gi|251945930 from Bacteroides sp. 3_2_5
 gi|263256635 from Bacteroides sp. 2_1_16

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 26%

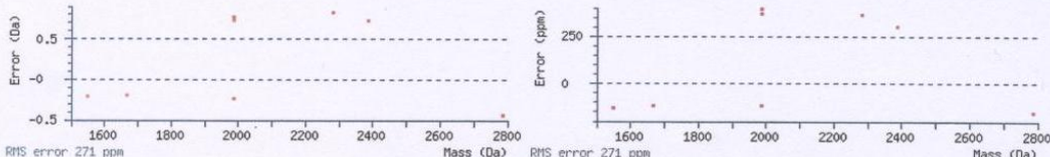
Matched peptides shown in **Bold Red**

1 MKRVLFSMVL LMAVSFAFAQ EKNVKEAKSI AGEVKPDKFAK **AEQLINEALT**
 51 **NPETKDNAAT** WDVAGYIQKR INEKEMENAY LRPYDTLKV YNSVLNMYNY
 101 YVKCDELAQI PNEKGIKIKN YRSANSKTIL **AERPFLINGG IQYFNLNKN**
 151 DALKYFAAYV DAATLPMMEK ENLLEKDTIL PQVAYATLA ADRVGDKDAV
 201 MKYAQYALKD **KENGQFAMQL** LTDAYRAKGD TAKWVEKLQE GIVK**FPENQY**
 251 **FPANLVDYYS** SSSNQDRAMQ FADDMLEKDP NNNKLYLVKA YLYHNMKDYE
 301 KAIEFYKKTLL DIDPAYAEAC SNLGLVYLLQ AQEYADKAPA DINDPNYATA
 351 QAEIKKFFVEA AKPYEKARE LKPDQKDLWL QGLYRVYYNL **NMGPEFEBIE**
 401 **KMM**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
41 - 55	835.8360	1669.6574	1669.8522	-0.1948	0	K.AEQLINEALTNPETK.D (Ions score 74)
56 - 69	776.2740	1550.5334	1550.7365	-0.2031	0	K.DNAATWDVAGYIQK.R (Ions score 64)
128 - 148	797.0170	2388.0292	2387.2961	0.7331	0	K.TILAERPFLINGGIQYFNLKN.N (Ions score 32)
210 - 226	663.5640	1987.6702	1986.9357	0.7345	1	K.DKENGQFAMQLLTDAYK.A Oxidation (M) (Ions score 51)
210 - 226	994.8670	1987.7194	1986.9357	0.7838	1	K.DKENGQFAMQLLTDAYK.A Oxidation (M) (Ions score 26)
245 - 267	930.6050	2788.7932	2789.2245	-0.4313	0	K.FPENQYFPANLVDYYSNSSNQDK.A (Ions score 57)
386 - 401	995.8420	1989.6694	1989.9030	-0.2335	0	R.VYYNLNMGPEFEBIEK.M Oxidation (M) (Ions score 68)
386 - 403	762.6100	2284.8082	2283.9738	0.8344	1	R.VYYNLNMGPEFEBIEKMM.- 3 Oxidation (M) (Ions score 19)



LOCUS YP_099695 403 aa linear BCT 26-APR-2009
 DEFINITION TPR repeat-containing protein [Bacteroides fragilis YCH46].
 ACCESSION YP_099695
 VERSION YP_099695.1 GI:53713703
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS
 SOURCE Bacteroides fragilis YCH46
 ORGANISM Bacteroides fragilis YCH46
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 403)
 AUTHORS Kuwahara, T., Yamashita, A., Hirakawa, H., Nakayama, H., Toh, H.,
 Okada, N., Kuhara, S., Hattori, M., Hayashi, T. and Ohnishi, Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 403)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 403)
 AUTHORS Hattori, M., Yamashita, A., Toh, H., Oshima, K. and Shiba, T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,

1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD49161](#).
 Method: conceptual translation.

FEATURES

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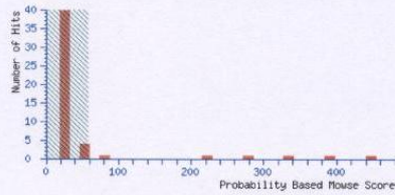
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unm.ac.uk
 Search title : SSP 2804
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2804) 1000_RE7_01_1712.d\SSP (2804) 1000_RE7_01_1712.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:12:18 GMT
 Protein hits : [gi|239636400](#) dihydrolipoyl dehydrogenase [Staphylococcus warneri L37603]
[gi|53713703](#) TPR repeat-containing protein [Bacteroides fragilis YCH46]
[gi|223043880](#) dihydrolipoyl dehydrogenase [Staphylococcus capitis SK14]
[gi|70726857](#) dihydrolipoamide dehydrogenase [Staphylococcus haemolyticus JCSC1435]
[gi|150011035](#) dihydrolipoamide dehydrogenase [Staphylococcus xylosum]
[gi|52080062](#) dihydrolipoamide dehydrogenase [Bacillus licheniformis ATCC 14580]
[gi|50084222](#) dihydrolipoamide dehydrogenase (E3 component of pyruvate and 2-oxoglutarate dehydrogenase complexes) [Acinetobacte

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As **Peptide Summary** Help

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|239636400](#) Mass: 49727 Score: 447 Queries matched: 11 emPAI: 0.47
 dihydrolipoyl dehydrogenase [Staphylococcus warneri L37603]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
113	493.7310	985.4474	985.5808	-0.1333	0	21	2.9e+02	6	K.LTGGVEGLLK.G
141	640.7770	1279.5394	1278.6456	0.8939	0	18	4.2e+02	2	R.ALSLDDTNGFVK.L
<input checked="" type="checkbox"/> 123	547.5700	1639.6882	1639.8781	-0.1899	0	(22)	2.5e+02	1	R.VIDSTGALNLQEVPGK.L
<input checked="" type="checkbox"/> 182	820.8600	1639.7054	1639.8781	-0.1726	0	79	0.00029	1	R.VIDSTGALNLQEVPGK.L
<input checked="" type="checkbox"/> 127	561.9150	1682.7232	1682.8839	-0.1607	0	(52)	0.27	1	R.RPNTDELGLBELGLK.F
<input checked="" type="checkbox"/> 187	842.3690	1682.7234	1682.8839	-0.1604	0	60	0.026	1	R.RPNTDELGLBELGLK.F
<input checked="" type="checkbox"/> 230	1103.5810	2205.1474	2205.2157	-0.0683	0	37	5.3	1	R.TSINNIYAIGDIVPGLPLANK.A
<input checked="" type="checkbox"/> 169	751.3460	2251.0162	2251.1597	-0.1435	2	22	3.5e+02	4	K.GHKVEIVKGAIFYFVNNSLR.V
<input checked="" type="checkbox"/> 238	772.6800	2315.0182	2315.2022	-0.1840	0	104	9.4e-07	1	K.NAIITGSRPIEIPNPFEGNR.V
<input checked="" type="checkbox"/> 246	854.9440	2561.8102	2562.2602	-0.4500	0	43	0.73	1	R.FVRAQHSNGLVIAESVSLNYEK.V
<input checked="" type="checkbox"/> 247	863.3940	2587.1602	2586.3694	0.7908	0	68	0.0028	1	M.VVGDFFPIETDITVIGAGPGGYVAAIR.A

2. [gi|53713703](#) Mass: 46634 Score: 393 Queries matched: 7 emPAI: 0.41
 TPR repeat-containing protein [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 129	582.2570	1162.4994	1162.6135	-0.1140	0	44	1.3	1	K.DLMLQGLYR.V
<input checked="" type="checkbox"/> 174	776.2780	1550.5414	1550.7365	-0.1951	0	64	0.0087	1	K.DNAATWDVAGYIQK.E
<input checked="" type="checkbox"/> 185	835.8410	1669.6674	1669.8522	-0.1848	0	70	0.0026	1	K.AEQLINEALTNFETK.D
<input checked="" type="checkbox"/> 198	893.3540	1784.6934	1784.8443	-0.1509	0	77	0.00044	1	K.VYNSVLNMYNYVYK.C + Oxidation (M)
<input checked="" type="checkbox"/> 211	994.4230	1986.8314	1986.9357	-0.1042	1	72	0.0014	1	K.DKENGQFAMQLLTDAYK.A + Oxidation (M)
<input checked="" type="checkbox"/> 236	1142.9710	2283.9274	2283.9738	-0.0463	1	46	0.52	1	R.VYVNLNMGPFEEIEKQM.- + 3 Oxidation (M)
<input checked="" type="checkbox"/> 242	797.0000	2387.9782	2387.2961	0.6821	0	21	1.3e+02	1	K.TTLAERPNLNGGIQVPLNK.N

Proteins matching the same set of peptides:

[gi|60681974](#) Mass: 45773 Score: 393 Queries matched: 7
 hypothetical protein BP2494 [Bacteroides fragilis NCTC 9343]

3. [gi|223043880](#) Mass: 49736 Score: 337 Queries matched: 10 emPAI: 0.38
 dihydrolipoyl dehydrogenase [Staphylococcus capitis SK14]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
113	493.7310	985.4474	985.5808	-0.1333	0	21	2.9e+02	6	K.LTGGVEGLLK.G
141	640.7770	1279.5394	1278.6456	0.8939	0	18	4.2e+02	2	R.ALSLDDTNGFVK.L
123	547.5700	1639.6882	1639.8781	-0.1899	0	(22)	2.5e+02	1	R.VIDSTGALNLQEVPGK.L

{MATRIX}
{SCIENCE} Mascot Search Results

Protein View

Match to: [gi|60681974](#) Score: 393
hypothetical protein BF2494 [Bacteroides fragilis NCTC 9343]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2804) 1000_RE7_01_1712.d\SSP (2804) 1000_RE7_01_1712.mgf

Nominal mass (M_r): 45773; Calculated pI value: 5.24
 NCBI BLAST search of [gi|60681974](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis NCTC 9343](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60493408](#) from [Bacteroides fragilis NCTC 9343](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 27%

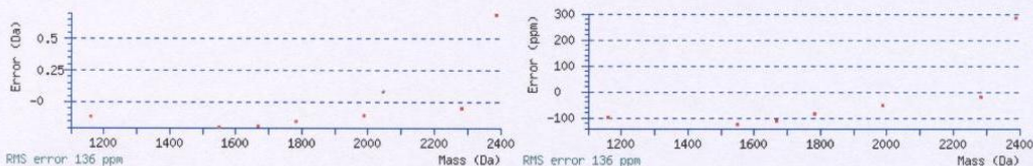
Matched peptides shown in **Bold Red**

1 MVLMLAVSFA FAQEKNVKEA KSIAGEVKPD FAKA**BQLINE** ALTN**PETKDN**
 51 AATWDVAGYI **QKR**INEKEME NAYLRKPYDT LKVYNSVLNM YNYVVK**CDEL**
 101 AQIPNEKGI KNKYRSANSK **TILAERP**NLI **NGGIQYF**NLN KNEDAL**KYFA**
 151 AYVDAATLPM MEKENLLEKD TILFQVAYYA TLAADRVGDK DAVMFKYAQYA
 201 LK**DKENGQ**FA **MQLLT**DAYKA KGD**TAKVVEK** LQEGIVK**FPE** NQYFFANLVD
 251 YSSSNQNDK AMQFAD**DMLA** KDPNNKLYLY VKAYLHNMK DYEKAI**EFYK**
 301 KTLDIDPAYA EACSNLGLVY LLQAQ**EYADK** APADINDPNY ATAQAE**IKFK**
 351 YEAAKPYEK ARELKP**DQKD** **LWLQGLYRVY** YNLNMG**PEFE** **RIEKMM**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
34 - 48	835.8410	1669.6674	1669.8522	-0.1848	0	K.ABQLINEALTN PETK .D (Ions score 70)
49 - 62	776.2780	1550.5414	1550.7365	-0.1951	0	K.DNAATWDVAGYI KR .R (Ions score 64)
83 - 96	893.3540	1784.6934	1784.8443	-0.1509	0	K.VYNSVLNMYNYV K .C Oxidation (M) (Ions score 77)
121 - 141	797.0000	2387.9782	2387.2961	0.6821	0	K.TILAERP NLI NGGIQYF NLN .N (Ions score 21)
203 - 219	994.4230	1986.8314	1986.9357	-0.1042	1	K.DKENGQ FA MQLLT DAYK .A Oxidation (M) (Ions score 72)
370 - 378	582.2570	1162.4994	1162.6135	-0.1140	0	K.DLWLQGLY R .V (Ions score 44)
379 - 396	1142.9710	2283.9274	2283.9738	-0.0463	1	R.VYVYLN MGPEFE RIEK MM .- 3 Oxidation (M) (Ions score 46)



LOCUS YP_212118 396 aa linear ECT 01-MAY-2009
 DEFINITION hypothetical protein BF2494 [Bacteroides fragilis NCTC 9343].
 ACCESSION YP_212118
 VERSION YP_212118.1 GI:60681974
 DBLINK Project:46
 DBSOURCE REFSEQ: accession [NC_003228.3](#)
 KEYWORDS
 SOURCE Bacteroides fragilis NCTC 9343
 ORGANISM [Bacteroides fragilis NCTC 9343](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 396)
 AUTHORS Cerdeno-Tarraga,A.M., Patrick,S., Crossman,L.C., Blakely,G.,
 Abratt,V., Lennard,N., Poxton,I., Duerden,B., Harris,B.,
 Quail,M.A., Barron,A., Clark,L., Corton,C., Doggett,J.,
 Holden,M.T., Larke,N., Line,A., Lord,J., Norbertczak,H., Ormond,D.,
 Price,C., Rabinowitsch,E., Woodward,J., Barrell,B. and Parkhill,J.
 TITLE Extensive DNA inversions in the B. fragilis genome control variable
 gene expression
 JOURNAL Science 307 (5714), 1463-1465 (2005)
 PUBMED [15746427](#)
 REFERENCE 2 (residues 1 to 396)
 AUTHORS Cerdeno-Tarraga,A.M.
 TITLE Direct Submission
 JOURNAL Submitted (29-JUL-2004) Pathogen Sequencing Unit, Sanger Institute,
 Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SA, United
 Kingdom
 REFERENCE 3 (residues 1 to 396)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (08-APR-2002) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 COMMENT PROVISIONAL [REFSEQ](#): This record has not yet been subject to final

NCBI review. The reference sequence was derived from CAH08194.
Method: conceptual translation.

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FEATURES             Location/Qualifiers
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                        /note="putative PEP-CTERM system TPR-repeat lipoprotein;
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                        /db_xref="CDD:131963"
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                        34 amino acids
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                        X(2)-[ASL]-X(4)-[PKE] is the consensus sequence; found in
                        a variety of organisms including bacteria, cyanobacteria,
                        yeast, fungi...; cd00189"
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                        318..319,322..323)
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                        /note="binding surface"
                        /db_xref="CDD:29151"
    Site            order(249,261,265,268,283,295,299,302,317)
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                        /coded_by="NC_003228.3:2901913..2903103"
                        /note="Similar to Bacteroides thetaiotaomicron putative
                        tetratricopeptide repeat family protein BT0900
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                        4.1e-110, 72.79% id in 397 aa"
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                        /db_xref="GeneID:3287481"

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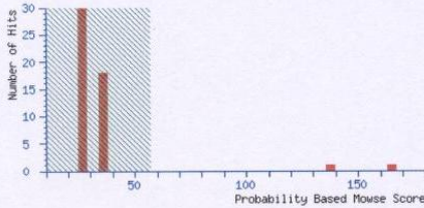
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 2806
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2806) 10_RH14_01_1275.d\SSP (2806) 10_RH14_01_1275.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:19:44 GMT
 Protein hits : [gi|29348144](#) 50S ribosomal protein L7/L12 [Bacteroides thetaiotaomicron VPI-5482]
 [gi|189460676](#) hypothetical protein BACCOP_01323 [Bacteroides coprocola DSM 17136]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold $p < 0.05$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

- [gi|29348144](#) Mass: 12730 Score: 165 Queries matched: 6 emPAI: 0.61
 50S ribosomal protein L7/L12 [Bacteroides thetaiotaomicron VPI-5482]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
89	565.2830	1128.5514	1128.6390	-0.0876	0	36	7.8	2	K.EVNELATILK.E
106	644.2710	1286.5274	1287.6558	-1.1283	0	(14)	1.2e+03	5	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/> 107	644.7870	1287.5594	1287.6558	-0.0963	0	46	0.7	1	K.TLEEAGAEVELK.-
<input checked="" type="checkbox"/> 113	666.8140	1331.6134	1331.7449	-0.1314	0	85	0.0001	1	K.AFAEQLVNLTVK.E
<input checked="" type="checkbox"/> 114	444.8790	1331.6152	1331.7449	-0.1297	0	(41)	2.5	1	K.AFAEQLVNLTVK.E
<input checked="" type="checkbox"/> 115	666.8760	1331.7374	1331.7449	-0.0074	0	(61)	0.024	1	K.AFAEQLVNLTVK.E

- Proteins matching the same set of peptides:
- [gi|53715478](#) Mass: 12688 Score: 165 Queries matched: 6
50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]
 - [gi|150003396](#) Mass: 12704 Score: 165 Queries matched: 6
50S ribosomal protein L7/L12 [Bacteroides vulgatus ATCC 8482]
 - [gi|153805949](#) Mass: 12686 Score: 165 Queries matched: 6
hypothetical protein BACCAC_00194 [Bacteroides caccae ATCC 43185]
 - [gi|160888407](#) Mass: 12775 Score: 165 Queries matched: 6
hypothetical protein BACUNI_00820 [Bacteroides uniformis ATCC 8492]
 - [gi|189465410](#) Mass: 12759 Score: 165 Queries matched: 6
hypothetical protein BACINT_01763 [Bacteroides intestinalis DSM 17393]
 - [gi|198277435](#) Mass: 12757 Score: 165 Queries matched: 6
hypothetical protein BACPLE_03654 [Bacteroides plebeius DSM 17135]
 - [gi|224023599](#) Mass: 12771 Score: 165 Queries matched: 6
hypothetical protein BACCOPRO_00306 [Bacteroides coprophilus DSM 18228]
 - [gi|224537808](#) Mass: 12757 Score: 165 Queries matched: 6
hypothetical protein BACCELL_02695 [Bacteroides cellulosilyticus DSM 14838]

- [gi|189460676](#) Mass: 12771 Score: 135 Queries matched: 5 emPAI: 0.27
 hypothetical protein BACCOP_01323 [Bacteroides coprocola DSM 17136]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
89	565.2830	1128.5514	1128.6390	-0.0876	0	36	7.8	2	K.EVNELATILK.E
106	644.2710	1286.5274	1285.6765	0.8509	0	14	1.2e+03	5	K.VLEEAGAEVELK.-
113	666.8140	1331.6134	1331.7449	-0.1314	0	85	0.0001	1	K.AFAEQLVNLTVK.E
114	444.8790	1331.6152	1331.7449	-0.1297	0	(41)	2.5	1	K.AFAEQLVNLTVK.E
115	666.8760	1331.7374	1331.7449	-0.0074	0	(61)	0.024	1	K.AFAEQLVNLTVK.E

MATRIX
SCIENCE Mascot Search Results

Protein View

Match to: [gi|53715478](#) Score: 165
50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (2806) 10_RH14_01_1275.d\SSP (2806) 10_RH14_01_1275.mgf

Nominal mass (M_r): **12688**; Calculated pI value: **4.71**
 NCBI BLAST search of [gi|53715478](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60683451](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253566657](#) from [Bacteroides sp. 3_2_5](#)
[gi|255011634](#) from [Bacteroides fragilis 3_1_12](#)
[gi|265767535](#) from [Bacteroides sp. 2_1_16](#)
[gi|81313501](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|81608231](#) from [Bacteroides fragilis](#)
[gi|52218343](#) from [Bacteroides fragilis YCH46](#)
[gi|60494885](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251944829](#) from [Bacteroides sp. 3_2_5](#)
[gi|263252840](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **27%**

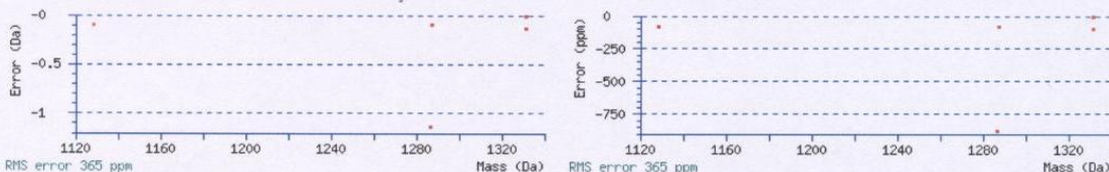
Matched peptides shown in **Red**

- 1 **MADLKAFAEQ LVNLTVKEVN ELATILK**EEY GIEPAAAAVA VAAGPAAGAA
- 51 **AABEKSSPDV VLKSAGAACL QVVKAVKEAC** GLGLKEAKDM VDGAPSVVKE
- 101 **GLAKDEAESL KKTLEEAGAE VELK**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
6 - 17	666.8140	1331.6134	1331.7449	-0.1314	0	K.AFAEQLVNLTVK.E (Ions score 85)
6 - 17	444.8790	1331.6152	1331.7449	-0.1297	0	K.AFAEQLVNLTVK.E (Ions score 41)
6 - 17	666.8760	1331.7374	1331.7449	-0.0074	0	K.AFAEQLVNLTVK.E (Ions score 61)
18 - 27	565.2830	1128.5514	1128.6390	-0.0876	0	K.EVNELATILK.E (Ions score 36)
113 - 124	644.2710	1286.5274	1287.6558	-1.1283	0	K.TLEEAGAEVELK.- (Ions score 14)
113 - 124	644.7870	1287.5594	1287.6558	-0.0963	0	K.TLEEAGAEVELK.- (Ions score 46)



LOCUS YP_101470 124 aa linear BCT 26-APR-2009
 DEFINITION 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46].
 ACCESSION YP_101470
 VERSION YP_101470.1 GI:53715478
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 124)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 124)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 124)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,

1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 REVIEWED REFSEQ: This record has been curated by NCBI staff. The
 reference sequence was derived from [BAD50936](#).
 Method: conceptual translation.

FEATURES
 Location/Qualifiers

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 /strain="YCH46"
 /db_xref="taxon:295405"

Protein 1..124
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 /calculated_mol_wt=12506

Region 2..123
 /region_name="Ribosomal L7_L12"
 /note="Ribosomal protein L7/L12. Ribosomal protein L7/L12
 refers to the large ribosomal subunit proteins L7 and L12,
 which are identical except that L7 is acetylated at the N
 terminus. It is a component of the L7/L12 stalk, which is
 located at the surface of...; cd00387"

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 80..81,83,85..86,100..101,103,106)
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 /db_xref="CDD:100102"

Site order(5,8,12,25,30)
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 /note="peripheral dimer interface"
 /db_xref="CDD:100102"

Site order(19,22..23,27,30)
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 /db_xref="CDD:100102"

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 L12 dimers bind L10; critically important for translation
 efficiency and fidelity; stimulates GTPase activity of
 translation factors"
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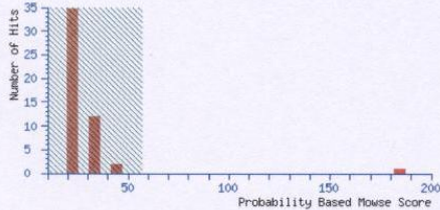
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 3303
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (3303) 10_RJ5_01_1367.d\SSP (3303) 10_RJ5_01_1367.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:24:24 GMT
 Protein hits : [gi|53711794](#) cationic outer membrane protein precursor [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

- [gi|53711794](#) Mass: 19149 Score: 184 Queries matched: 8 emPAI: 1.24
 cationic outer membrane protein precursor [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
110	593.7370	1185.4594	1185.5706	-0.1112	0	20	3.3e+02	7	K.YFGPEGELFK.K
<input checked="" type="checkbox"/> 126	657.7750	1313.5354	1313.6656	-0.1301	1	43	1.3	1	R.KYFGPEGELFK.K
127	438.8580	1313.5522	1313.6656	-0.1134	1	(42)	2.1	2	R.KYFGPEGELFK.K
<input checked="" type="checkbox"/> 138	686.3240	1370.6334	1370.7155	-0.0821	0	53	0.14	1	K.FALIDMEYILK.N + Oxidation (M)
<input checked="" type="checkbox"/> 139	686.8900	1371.7654	1370.7155	1.0499	0	(19)	4.1e+02	3	K.FALIDMEYILK.N + Oxidation (M)
<input checked="" type="checkbox"/> 197	968.3630	1934.7114	1934.9659	-0.2544	0	35	6.1	1	R.EELMKPIQDEIYNAVK.A + Oxidation (M)
<input checked="" type="checkbox"/> 198	645.9190	1934.7352	1934.9659	-0.2307	0	(33)	9.1	1	R.EELMKPIQDEIYNAVK.A + Oxidation (M)
<input checked="" type="checkbox"/> 142	697.9610	2090.8612	2091.0670	-0.2058	1	36	12	1	K.REELMKPIQDEIYNAVK.A + Oxidation (M)

Proteins matching the same set of peptides:
[gi|255009916](#) Mass: 19218 Score: 184 Queries matched: 8
 putative cationic 19 kDa outer membrane protein precursor [Bacteroides fragilis 3_1_12]
[gi|265765175](#) Mass: 19077 Score: 184 Queries matched: 8
 cationic outer membrane protein [Bacteroides sp. 2_1_16]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 127	438.8580	1313.5522	1313.5962	-0.0440	0	43	1.5	1	QYFGPEGELMK + Oxidation (M)
<input checked="" type="checkbox"/> 221	700.9150	2099.7232	2100.1262	-0.4031	2	39	2.2	1	YLMVGGDVVDPVAAQVRR
<input checked="" type="checkbox"/> 233	741.9410	2222.8012	2223.1310	-0.3299	2	37	3.2	1	ELLIGKTDYDFPFKEADK
<input checked="" type="checkbox"/> 82	431.2590	1290.7552	1289.6364	1.1188	1	32	40	1	QKPDAGPFDKR
<input checked="" type="checkbox"/> 217	694.9340	2081.7802	2080.9371	0.8430	0	31	15	1	SGETIFENNQANDLTGPMK + Oxidation (M)
<input checked="" type="checkbox"/> 108	584.9800	1751.9182	1751.6879	0.2302	0	31	37	1	ATCSDACDHLFPFSGGR
<input checked="" type="checkbox"/> 111	607.0930	1818.2572	1817.9035	0.3537	0	29	47	1	WAATHAPEHWGLMLAK
<input checked="" type="checkbox"/> 215	687.9070	2060.6992	2060.0659	0.6333	2	29	22	1	MSRPKKVPMPELPPHER + 2 Oxidation (M)
<input checked="" type="checkbox"/> 128	658.3940	1972.1602	1972.9784	-0.8182	0	29	74	1	MVAIGMNSMLLSYILR + 3 Oxidation (M)
<input checked="" type="checkbox"/> 231	1105.9580	2209.9014	2210.2232	-0.3218	0	28	28	1	GVLPLALLVLSGSLVLAGCDDK
<input checked="" type="checkbox"/> 235	748.5640	2242.6702	2242.1600	0.5101	2	27	29	1	TIEGRPALPDTRMTTQRR + Oxidation (M)
<input checked="" type="checkbox"/> 109	593.4080	1777.2022	1777.9323	-0.7301	0	27	87	1	TEQVVGTDHPNTLVLR
<input checked="" type="checkbox"/> 203	979.4010	2935.1812	2934.4164	0.7648	2	27	66	1	HMGVSLRVINDMVMGKGFQTHGR + Oxidation (M)
<input checked="" type="checkbox"/> 134	666.8440	1331.6734	1331.7231	-0.0497	0	27	58	1	AMLQSGSGSLLLR
<input checked="" type="checkbox"/> 178	565.9430	1694.8072	1695.8767	-1.0695	0	26	63	1	AKPVAVSYGGMAGGLR + Oxidation (M)
<input checked="" type="checkbox"/> 213	685.9740	2054.9002	2056.0126	-1.1125	1	25	68	1	SGQPHTKAPIYSHVYDR
<input checked="" type="checkbox"/> 230	737.6170	2209.8292	2210.0170	-0.1878	1	25	55	1	MNTKVAIIMGSHSDWETMK + 2 Oxidation (M)

Mascot Search Results

Protein View

Match to: **gi|53711794** Score: **184**
cationic outer membrane protein precursor [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (3303) 10_RJ5_01_1367.d\SSP (3303) 10_RJ5_01_1367.mgf

Nominal mass (M₀): **19149**; Calculated pI value: **6.43**
 NCBI BLAST search of **gi|53711794** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: **Bacteroides fragilis YCH46**
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60680025](#) from **Bacteroides fragilis NCTC 9343**
[gi|253564138](#) from **Bacteroides sp. 3_2_5**
[gi|52214659](#) from **Bacteroides fragilis YCH46**
[gi|60491459](#) from **Bacteroides fragilis NCTC 9343**
[gi|251947914](#) from **Bacteroides sp. 3_2_5**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **22%**

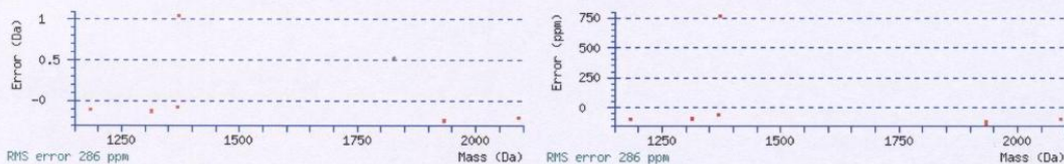
Matched peptides shown in **Bold Red**

1 MKKSVLFIIIL LFAVGMTAQA QK**FALIDMEY ILKNIPAYER** ANEQLSQATK
51 QWQGEVEVLA KEAQTFPKDY QAASAKLTAA QKTQKEDAIV EKEKAASELK
101 RKYFGPEGEL **FKKREELMKP IQDEIYNAVK** AVAEENGYAV VVDRASASSI
151 IPATPRIDVS NEVLAKLGYS N

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
23 - 33	686.3240	1370.6334	1370.7155	-0.0821	0	K.FALIDMEYILK.N Oxidation (M) (Ions score 53)
23 - 33	686.8900	1371.7654	1370.7155	1.0499	0	K.FALIDMEYILK.N Oxidation (M) (Ions score 19)
102 - 112	657.7750	1313.5354	1313.6656	-0.1301	1	R.RYFGPEGELFK.K (Ions score 43)
102 - 112	438.8580	1313.5522	1313.6656	-0.1134	1	R.RYFGPEGELFK.K (Ions score 42)
103 - 112	593.7370	1185.4594	1185.5706	-0.1112	0	K.YFGPEGELFK.K (Ions score 20)
114 - 130	697.9610	2090.8612	2091.0670	-0.2058	1	K.REELMKPIQDEIYNAVK.A Oxidation (M) (Ions score 36)
115 - 130	968.3630	1934.7114	1934.9659	-0.2544	0	R.EELMKPIQDEIYNAVK.A Oxidation (M) (Ions score 35)
115 - 130	645.9190	1934.7352	1934.9659	-0.2307	0	R.EELMKPIQDEIYNAVK.A Oxidation (M) (Ions score 33)



LOCUS YP_097786 171 aa linear BCT 26-APR-2009
 DEFINITION cationic outer membrane protein precursor [Bacteroides fragilis YCH46].
 ACCESSION YP_097786
 VERSION YP_097786.1 GI:53711794
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM **Bacteroides fragilis YCH46**
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 171)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H., Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 171)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 171)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences, 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final NCBI review. The reference sequence was derived from [BAD47252](#).

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Method: conceptual translation.
FEATURES             Location/Qualifiers
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                     /organism="Bacteroides fragilis YCH46"
                     /strain="YCH46"
                     /db_xref="taxon:295405"
  Protein            1..171
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                     /calculated_mol_wt=19030
  Region             4..167
                     /region_name="HlpA"
                     /note="Outer membrane protein [Cell envelope biogenesis,
                     outer membrane]; cl08146"
                     /db_xref="CDD:142087"
  Region             7..168
                     /region_name="OmpH"
                     /note="Outer membrane protein (OmpH-like); pfam03938"
                     /db_xref="CDD:112737"
  CDS                1..171
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                     /coded_by="complement(NC_006347.1:578464..578979)"
                     /note="similar to gp:AE016941_210 [Bacteroides
                     thetaiotaomicron VPI-5482], percent identity 71 in 171 aa,
                     BLASTP E(): 2e-63"
                     /transl_table=11
                     /db_xref="GeneID:3082496"

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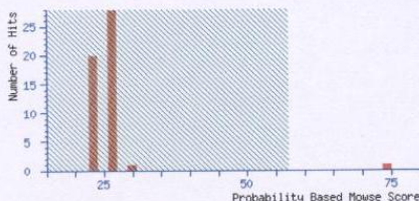
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 3404
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (3404) 10_RJ16_01_1392.d\SSP (3404) 10_RJ16_01_1392.mgf
 Database : NCBIInr 20100108 (10291680 sequences; 3511877860 residues)
 Taxonomy : Bacteria (Eubacteria) (5718488 sequences)
 Timestamp : 13 Jan 2010 at 10:33:08 GMT
 Protein hits : [gi|143945](#) Fe-superoxide dismutase [Bacteroides fragilis]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As **Peptide Summary** [Help](#)

Significance threshold $p < 0.05$ Max. number of hits **AUTO**

Standard scoring MudPIT scoring Ions score or expect cut-off **0** Show sub-sets **0**

Show pop-ups Suppress pop-ups Sort unassigned **Decreasing Score** Require bold red

Select All Select None Search Selected Error tolerant

- [gi|143945](#) Mass: 21736 Score: 74 Queries matched: 2 emPAI: 0.15
 Fe-superoxide dismutase [Bacteroides fragilis]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 138	545.5980	1089.1814	1089.4767	-0.2953	0	14	8.6e+02	7	R.DFGSFENFK.K
<input checked="" type="checkbox"/> 174	773.3660	1544.7174	1543.7922	0.9252	0	60	0.048	1	K.LWEIWDVVEK.R

Proteins matching the same set of peptides:

- [gi|533307](#) Mass: 21715 Score: 74 Queries matched: 2
 superoxide dismutase [Bacteroides fragilis]
- [gi|53713818](#) Mass: 23124 Score: 74 Queries matched: 2
 superoxide dismutase [Bacteroides fragilis YCH46]
- [gi|60682036](#) Mass: 21758 Score: 74 Queries matched: 2
 superoxide dismutase [Fe] [Bacteroides fragilis NCTC 9343]
- [gi|150006035](#) Mass: 23299 Score: 74 Queries matched: 2
 superoxide dismutase [Bacteroides vulgatus ATCC 8482]
- [gi|212692986](#) Mass: 21962 Score: 74 Queries matched: 2
 hypothetical protein BACDOR_02487 [Bacteroides dorei DSM 17855]
- [gi|237709817](#) Mass: 23300 Score: 74 Queries matched: 2
 superoxide dismutase [Bacteroides sp. 9_1_42FAA]
- [gi|237725616](#) Mass: 23274 Score: 74 Queries matched: 2
 superoxide dismutase [Bacteroides sp. D4]
- [gi|255009481](#) Mass: 21757 Score: 74 Queries matched: 2
 superoxide dismutase [Bacteroides fragilis 3_1_12]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 171	509.2150	1524.6232	1523.6596	0.9636	0	28	41	1	MMTQSGNPEIQEK + 2 Oxidation (M)
<input checked="" type="checkbox"/> 223	663.3920	1987.1542	1987.9851	-0.8309	1	27	44	1	DAEKAGQNPVDVYVENLK
<input checked="" type="checkbox"/> 81	389.8350	1166.4832	1167.5455	-1.0623	0	27	97	1	RPAGTYSGGMR + Oxidation (M)
<input checked="" type="checkbox"/> 116	473.3980	944.7814	944.5443	0.2371	1	26	56	1	HKLLGYSK
<input checked="" type="checkbox"/> 190	563.4840	1687.4302	1687.9522	-0.5220	1	26	52	1	AIGHIAPIQALDWR
<input checked="" type="checkbox"/> 180	803.5730	2407.6972	2407.3084	0.3888	1	25	1e+02	1	AVAAHVASTAGIAGLNVEGAAPVRR
<input checked="" type="checkbox"/> 225	1006.5690	3016.6852	3017.7391	-1.0540	0	25	1.2e+02	1	LLLGIFALFGLATLACAVAPSPISLLVAR
<input checked="" type="checkbox"/> 192	847.9400	2540.7982	2540.1826	0.6155	1	25	1.4e+02	1	QGCIESWVSGPGVAADHARTTQGR
<input checked="" type="checkbox"/> 204	887.8930	1773.7714	1773.9593	-0.1879	2	24	1.5e+02	1	VRKSQVAMAEVAQVK
<input checked="" type="checkbox"/> 145	590.9250	1769.7532	1769.0352	0.7180	0	24	1.8e+02	1	VIAAVHPGLWQPAVLAK
<input checked="" type="checkbox"/> 247	1313.3280	3936.9622	3935.8128	1.1494	2	24	1.1e+02	1	QDHPGQCPCGNTLEPILPSVDEGENPELIDFRR
<input checked="" type="checkbox"/> 240	777.6720	2329.9942	2330.0916	-0.0974	2	24	82	1	SSTQIVEMSTORMMVKDTEK + Oxidation (M)
<input checked="" type="checkbox"/> 183	811.4540	2431.3402	2431.0063	0.3339	1	24	2.1e+02	1	DEYSQYKDYDLQDFYHR + Oxidation (M)
<input checked="" type="checkbox"/> 118	473.9190	1418.7352	1417.7599	0.9753	1	24	2.3e+02	1	AMTIGSKSLIER + Oxidation (M)

MASCOT Mascot Search Results

Protein View

Match to: **gi|60682036** Score: **74**
superoxide dismutase [Fe] [Bacteroides fragilis NCTC 9343]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (3404) 10_RJ16_01_1392.d\SSP (3404) 10_RJ16_01_1392.mgf

Nominal mass (M_r): **21758**; Calculated pI value: **6.07**
 NCBI BLAST search of [gi|60682036](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis NCTC 9343](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|253565789](#) from [Bacteroides sp. 3_2_5](#)
[gi|265764163](#) from [Bacteroides sp. 2_1_16](#)
[gi|55977805](#) from [Bacteroides fragilis](#)
[gi|60493470](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251946068](#) from [Bacteroides sp. 3_2_5](#)
[gi|263256771](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **10%**

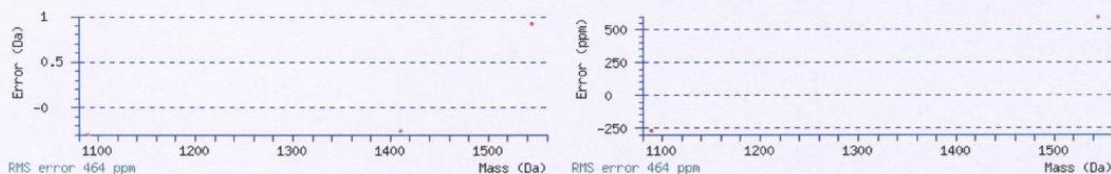
Matched peptides shown in **Bold Red**

1 MTYEMPKLPY ANNALEPVIS QOTIDYHYGK HLQTYVNNLN SLVPGTEYEG
 51 KTVEAIVASA PDGAIFNNAG QVLNHTLYPL QFAPKPAKNE PAGKLGKAIK
101 RDFGSFENFK**** KEFNAASVGL FSGGWANLSV DKDGKLHITK EPNGSNPVRT
 151 GLKPLLGFDV WEHAYYLDYQ NRRADHVNKL **WEIIDWDVVE** KRL

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
102 - 110	545.5980	1089.1814	1089.4767	-0.2953	0	R.DFGSFENFK.K (Ions score 14)
180 - 191	773.3660	1544.7174	1543.7922	0.9252	0	K.LWEIIDWDVVEK.R (Ions score 60)



LOCUS YP_212180 193 aa linear BCT 01-MAY-2009
 DEFINITION superoxide dismutase [Fe] [Bacteroides fragilis NCTC 9343].
 ACCESSION YP_212180
 VERSION YP_212180.1 GI:60682036
 DBLINK Project:46
 DBSOURCE REFSEQ: accession [NC_003228.3](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis NCTC 9343
 ORGANISM [Bacteroides fragilis NCTC 9343](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 193)
 AUTHORS Cerdeno-Tarraga,A.M., Patrick,S., Crossman,L.C., Blakely,G.,
 Abratt,V., Lennard,N., Poxton,I., Duerden,B., Harris,B.,
 Quail,M.A., Barron,A., Clark,L., Corton,C., Doggett,J.,
 Holden,M.T., Larke,N., Line,A., Lord,A., Norbertczak,H., Ormond,D.,
 Price,C., Rabinowitsch,E., Woodward,J., Barrell,B. and Parkhill,J.
 TITLE Extensive DNA inversions in the B. fragilis genome control variable
 gene expression
 JOURNAL Science 307 (5714), 1463-1465 (2005)
 PUBMED [15746427](#)
 REFERENCE 2 (residues 1 to 193)
 AUTHORS Cerdeno-Tarraga,A.M.
 TITLE Direct Submission
 JOURNAL Submitted (29-JUL-2004) Pathogen Sequencing Unit, Sanger Institute,
 Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SA, United
 Kingdom
 REFERENCE 3 (residues 1 to 193)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (08-APR-2002) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 COMMENT PROVISIONAL [REFSEQ](#): This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [CAH08256](#).

Method: conceptual translation.

FEATURES Location/Qualifiers

source 1..193
 /organism="Bacteroides fragilis NCTC 9343"
 /strain="ATCC 25285; NCTC 9343"
 /db_xref="ATCC:25285"
 /db_xref="taxon:272559"

Protein 1..193
 /product="superoxide dismutase [Fe]"
 /EC_number="1.15.1.1"
 /calculated_mol_wt=21640

Region 1..193
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 /note="Superoxide dismutase [Inorganic ion transport and metabolism]; COG0605"
 /db_xref="CDD:30950"

Region 2..83
 /region_name="Sod_Fe_N"
 /note="Iron/manganese superoxide dismutases, alpha-hairpin domain; pfam00081"
 /db_xref="CDD:109149"

Region 90..192
 /region_name="Sod_Fe_C"
 /note="Iron/manganese superoxide dismutases, C-terminal domain; pfam02777"
 /db_xref="CDD:111650"

CDS 1..193
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 /locus_tag="BF2556"
 /gene_synonym="sod"
 /coded_by="NC_003228.3:2982352..2982933"
 /note="Similar to Bacteroides fragilis superoxide dismutase [Fe] SodB or Sod SWALL:SODF_BACFR (SWALL:P53638) (193 aa) fasta scores: E(): 1.2e-77, 99.48% id in 193 aa, and to Escherichia coli, Escherichia coli O6, Escherichia coli O157:H7, and Shigella flexneri superoxide dismutase [Fe] SodB or B1656 or C2050 or Z2678 or ECS2365 or SF1684 or S1816 SWALL:SODF_ECOLI (SWALL:P09157) (192 aa) fasta scores: E(): 1.6e-37, 51.04% id in 192 aa"
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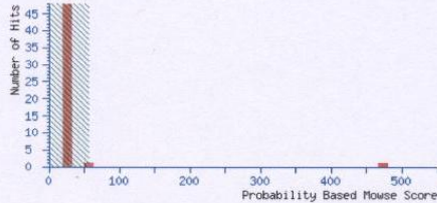
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4202
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4202) 10_RN10_01_1584.d\SSP (4202) 10_RN10_01_1584.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Subacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:25:46 GMT
 Protein hits : [gi|53711794](#) cationic outer membrane protein precursor [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|53711794](#) Mass: 19149 Score: 473 Queries matched: 13 emPAI: 3.28
 cationic outer membrane protein precursor [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 89	593.7220	1185.4294	1185.5706	-0.1412	0	48	0.38	1	K.YFGPEGELFK.K
<input checked="" type="checkbox"/> 90	610.7910	1219.5674	1219.6561	-0.0886	0	56	0.071	1	R.ASASSIIFATPR.I
<input checked="" type="checkbox"/> 91	610.8180	1219.6214	1219.6561	-0.0346	0	(18)	7.2e+02	4	R.ASASSIIFATPR.I
<input checked="" type="checkbox"/> 92	611.2710	1220.5274	1219.6561	0.8714	0	(45)	0.86	1	R.ASASSIIFATPR.I
<input checked="" type="checkbox"/> 105	438.8460	1313.5162	1313.6656	-0.1494	1	49	0.34	1	R.KYFGPEGELFK.K
<input checked="" type="checkbox"/> 106	657.7710	1313.5274	1313.6656	-0.1381	1	(42)	1.6	1	R.KYFGPEGELFK.K
<input checked="" type="checkbox"/> 113	686.3220	1370.6294	1370.7155	-0.0861	0	73	0.0016	1	K.FALIDMEYILK.N + Oxidation (M)
<input checked="" type="checkbox"/> 126	746.7550	1491.4954	1490.7365	0.7589	0	50	0.19	1	K.AVAEENGYAVVVD.R.A
<input checked="" type="checkbox"/> 167	968.3790	1934.7434	1934.9659	-0.2224	0	37	3.5	1	R.BELMKPIQDEIYNAVK.A + Oxidation (M)
<input checked="" type="checkbox"/> 203	697.9660	2090.8762	2091.0670	-0.1908	1	45	0.67	1	K.BEELMKPIQDEIYNAVK.A + Oxidation (M)
<input checked="" type="checkbox"/> 233	786.3270	2355.9592	2356.2023	-0.2431	1	46	0.44	1	R.ANQQLSQATKQWQGEVVLAK.E
<input checked="" type="checkbox"/> 241	898.4020	2692.1842	2692.3820	-0.1978	1	47	0.36	1	K.AVAEENGYAVVVD.RASASSIIFATPR.I
<input checked="" type="checkbox"/> 251	991.4350	2971.2832	2971.4385	-0.1554	2	23	85	1	K.QWQGEVVLAKAEQTMFKDYQAASAK.L + Oxidation (M)

Proteins matching the same set of peptides:
[gi|265765175](#) Mass: 19077 Score: 473 Queries matched: 13
 cationic outer membrane protein [Bacteroides sp. 2_1_16]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 100	635.2560	1268.4974	1268.5997	-0.1022	0	40	2.6	1	SVAGDEVADAHAK
<input checked="" type="checkbox"/> 182	666.0470	1995.1192	1994.0666	1.0526	0	35	8.3	1	YAYVPYSHFFIGAVLVAK
<input checked="" type="checkbox"/> 132	805.8850	1609.7554	1609.9001	-0.1446	0	33	12	1	ISLDTYMSLLVVLK + Oxidation (M)
<input checked="" type="checkbox"/> 194	1028.3360	3081.9862	3081.5739	0.4123	1	33	14	1	VACPPRAEIVAGLQAMPFGAAILVAEHR
<input checked="" type="checkbox"/> 234	1181.9720	3542.8942	3542.9397	-0.0456	1	32	20	1	IGLCGLVSPVGVISIVKLFHAPETAIPALAMFR + Oxidation (M)
<input checked="" type="checkbox"/> 168	646.8500	1937.5282	1936.8917	0.6364	2	32	11	1	SGADDLLICGPRMGR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 160	947.3360	2838.9862	2838.3818	0.6044	0	31	26	1	DLANAGGNASDILMNLPSIAVDPEGNVR + Oxidation (M)
<input checked="" type="checkbox"/> 175	979.3210	2934.9412	2935.4493	-0.5081	1	29	41	1	QMILMARMLTSEQDINDLLTHINTF + Oxidation (M)
<input checked="" type="checkbox"/> 180	663.0380	1986.0922	1986.0945	-0.0023	1	28	37	1	QARETSLIPVHPMVRPR
<input checked="" type="checkbox"/> 173	650.3710	1948.0912	1946.9230	1.1682	0	28	38	1	MIDIHCHIFPGIDGPK + Oxidation (M)
<input checked="" type="checkbox"/> 190	674.2500	2019.7282	2019.8713	-0.1431	1	28	27	1	CPAFAAAEAGMARAMADR + Oxidation (M)
<input checked="" type="checkbox"/> 110	673.9490	1345.8834	1345.7388	0.1447	0	28	81	1	GLTLAQTSMVGIR
<input checked="" type="checkbox"/> 108	662.2560	1322.4974	1322.6983	-0.2008	1	28	42	1	VPDGLNKDLFR
<input checked="" type="checkbox"/> 246	957.2550	2868.7432	2868.3283	0.4149	1	26	33	1	MADLSLHIAQSARGPYEGDLAQAMAR + Oxidation (M)

Mascot Search Results

Protein View

Match to: gi|53711794 Score: 473
cationic outer membrane protein precursor [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4202) 10_RN10_01_1584.d\SSP (4202) 10_RN10_01_1584.mgf

Nominal mass (M_r): 19149; Calculated pI value: 6.43
 NCBI BLAST search of gi|53711794 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60680025](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253564138](#) from [Bacteroides sp. 3_2_5](#)
[gi|52214659](#) from [Bacteroides fragilis YCH46](#)
[gi|60491459](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251947914](#) from [Bacteroides sp. 3_2_5](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 59%

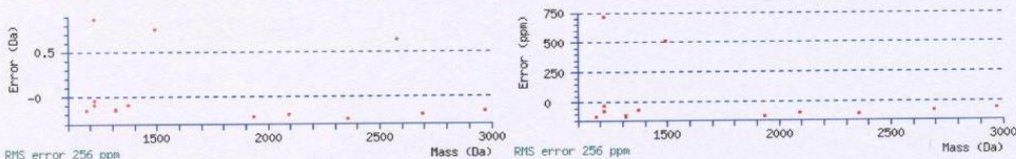
Matched peptides shown in **Bold Red**

1 MKKSVLPFIL LFAVGMTAQA QK**FALIDMEY** ILKNIPAYER ANEQLSQATK
51 QWQGEVEVLA KEAQT**MFKDY** QAASAKLTAA QKTQKEDAI**V** EKEKAASELK
101 RKYFGPEGEL F**KRELMKP** IQDEIYNA**V** AVAENGYAV VVDRASASSI
151 IFATPRIDVS NEVLAKLGYS N

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
23 - 33	686.3220	1370.6294	1370.7155	-0.0861	0 K.FALIDMEYILK.N Oxidation (M) (Ions score 73)
41 - 61	786.3270	2355.9592	2356.2023	-0.2431	1 R.ANEQLSQATKQWQGEVEVLAK.E (Ions score 46)
51 - 76	991.4350	2971.2832	2971.4385	-0.1554	2 K.QWQGEVEVLAKEAQT MFKDY QAASAK.L Oxidation (M) (Ions score 23)
102 - 112	438.8460	1313.5162	1313.6656	-0.1494	1 R.KYFGPEGELFK.K (Ions score 49)
102 - 112	657.7710	1313.5274	1313.6656	-0.1381	1 R.KYFGPEGELFK.K (Ions score 42)
103 - 112	593.7220	1185.4294	1185.5706	-0.1412	0 K.YFGPEGELFK.K (Ions score 48)
114 - 130	697.9660	2090.8762	2091.0670	-0.1908	1 K.RELMKPIQDEIYNA V .A Oxidation (M) (Ions score 45)
115 - 130	968.3790	1934.7434	1934.9659	-0.2224	0 R.EELMKPIQDEIYNA V .A Oxidation (M) (Ions score 37)
131 - 144	746.7550	1491.4954	1490.7365	0.7589	0 K.AVAENGYAVVDRASASSI FATPR .I (Ions score 50)
131 - 156	898.4020	2692.1842	2692.3820	-0.1978	1 K.AVAENGYAVVDRASASSI FATPR .I (Ions score 47)
145 - 156	610.7910	1219.5674	1219.6561	-0.0886	0 R.ASASSI FATPR .I (Ions score 56)
145 - 156	610.8180	1219.6214	1219.6561	-0.0346	0 R.ASASSI FATPR .I (Ions score 18)
145 - 156	611.2710	1220.5274	1219.6561	0.8714	0 R.ASASSI FATPR .I (Ions score 45)



LOCUS YP_097786 171 aa linear BCT 26-APR-2009
 DEFINITION cationic outer membrane protein precursor [Bacteroides fragilis YCH46].
 ACCESSION YP_097786
 VERSION YP_097786.1 GI:53711794
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 171)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H., Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 171)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 171)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences, 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final NCBI review. The reference sequence was derived from [BAD47252](#).
 Method: conceptual translation.
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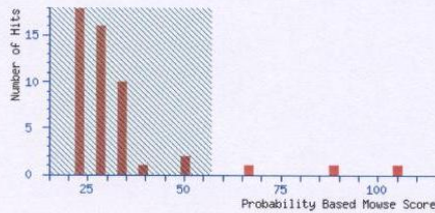
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4302
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4302) 10_RJ2_01_1361.d\SSP (4302) 10_RJ2_01_1361.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:32:16 GMT
 Protein hits : [gi|29349795](#) adenylate kinase [Bacteroides thetaiotaomicron VPI-5482]
 [gi|260910267](#) adenylate kinase [Prevotella sp. oral taxon 472 str. F0295]
 [gi|260593579](#) adenylate kinase [Prevotella veroralis F0319]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: Peptide Summary Help

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

- 1. [gi|29349795](#) Mass: 21073 Score: 105 Queries matched: 2
 adenylate kinase [Bacteroides thetaiotaomicron VPI-5482]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 117	504.2000	1006.3854	1006.5236	-0.1382	0	51	0.23	1	K.GVIFDGFPR.T
<input checked="" type="checkbox"/> 171	710.2580	1418.5014	1418.7592	-0.2577	0	54	0.088	1	-.MLNIVIFGAPGSGK.G + Oxidation (M)

Proteins matching the same set of peptides:

- [gi|53712412](#) Mass: 21018 Score: 105 Queries matched: 2
adenylate kinase [Bacteroides fragilis YCH46]
- [gi|150004241](#) Mass: 20871 Score: 105 Queries matched: 2
adenylate kinase [Bacteroides vulgatus ATCC 8482]
- [gi|153809272](#) Mass: 21085 Score: 105 Queries matched: 2
hypothetical protein BACCAC_03584 [Bacteroides caccae ATCC 43185]
- [gi|154495021](#) Mass: 20930 Score: 105 Queries matched: 2
hypothetical protein PARMER_04067 [Parabacteroides merdae ATCC 43184]
- [gi|160887206](#) Mass: 21071 Score: 105 Queries matched: 2
hypothetical protein BACOVA_05222 [Bacteroides ovatus ATCC 8483]
- [gi|160891012](#) Mass: 21165 Score: 105 Queries matched: 2
hypothetical protein BACUNI_03459 [Bacteroides uniformis ATCC 8492]
- [gi|167765207](#) Mass: 20977 Score: 105 Queries matched: 2
hypothetical protein BACSTE_03595 [Bacteroides stercoris ATCC 43183]
- [gi|189462905](#) Mass: 20867 Score: 105 Queries matched: 2
hypothetical protein BACOP_03606 [Bacteroides coprocola DSM 17136]
- [gi|189467969](#) Mass: 21048 Score: 105 Queries matched: 2
hypothetical protein BACINT_04363 [Bacteroides intestinalis DSM 17393]
- [gi|198274218](#) Mass: 20879 Score: 105 Queries matched: 2
hypothetical protein BACPLE_00358 [Bacteroides plebeius DSM 17135]
- [gi|212690825](#) Mass: 20901 Score: 105 Queries matched: 2
hypothetical protein BACDOR_00312 [Bacteroides dorei DSM 17855]
- [gi|218129349](#) Mass: 20933 Score: 105 Queries matched: 2
hypothetical protein BACEGG_00926 [Bacteroides eggerthii DSM 20697]
- [gi|218262912](#) Mass: 20944 Score: 105 Queries matched: 2
hypothetical protein PRABACTJOHN_02899 [Parabacteroides johnsonii DSM 18315]
- [gi|224025342](#) Mass: 20870 Score: 105 Queries matched: 2
hypothetical protein BACOPRO_02081 [Bacteroides coprophilus DSM 18228]
- [gi|224538342](#) Mass: 21028 Score: 105 Queries matched: 2

hypothetical protein BACCELL_03233 [Bacteroides cellulosilyticus DSM 14838]
 gi|237708208 Mass: 21143 Score: 105 Queries matched: 2
 LOW QUALITY PROTEIN: adenylate kinase [Bacteroides sp. 9_1_42FAA]
 gi|237715693 Mass: 21657 Score: 105 Queries matched: 2
 adenylate kinase [Bacteroides sp. D1]
 gi|254884741 Mass: 20813 Score: 105 Queries matched: 2
 adenylate kinase [Bacteroides sp. 4_3_47FAA]
 gi|255007910 Mass: 21018 Score: 105 Queries matched: 2
 adenylate kinase [Bacteroides fragilis 3_1_12]
 gi|255693744 Mass: 20986 Score: 105 Queries matched: 2
 adenylate kinase [Bacteroides finegoldii DSM 17565]

2. gi|260910267 Mass: 21197 Score: 89 Queries matched: 2
 adenylate kinase [Prevotella sp. oral taxon 472 str. F0295]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
117	504.2000	1006.3854	1006.5236	-0.1382	0	51	0.23	1	K.GVIFDGFPR.T
171	710.2580	1418.5014	1417.7751	0.7263	1	38	3.7	3	-.MKNIVIFGAPGSGK.G

Proteins matching the same set of peptides:
 gi|261880713 Mass: 21228 Score: 89 Queries matched: 2
 adenylate kinase [Prevotella bergensis DSM 17361]
 gi|281298521 Mass: 21183 Score: 89 Queries matched: 2
 adenylate kinase [Prevotella buccalis ATCC 35310]

3. gi|260593579 Mass: 21280 Score: 69 Queries matched: 2
 adenylate kinase [Prevotella veroralis F0319]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
117	504.2000	1006.3854	1006.5236	-0.1382	0	51	0.23	1	K.GVIFDGFPR.T
105	467.9130	1400.7172	1401.7802	-1.0631	1	18	7.9e+02	5	-.MKNIVIFGAPGAGK.G

Proteins matching the same set of peptides:
 gi|252119831 Mass: 21127 Score: 67 Queries matched: 2
 adenylate kinase [Prevotella melaninogenica ATCC 25845]
 gi|281423652 Mass: 20936 Score: 67 Queries matched: 2
 adenylate kinase [Prevotella oris F0302]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
182	496.5760	1486.7062	1485.8151	0.8911	1	35	10	1	QSTTNKQITLPAGK
72	390.1580	778.3014	779.4654	-1.1639	0	33	15	1	IAPAGFVR
222	659.5860	1975.7362	1976.1167	-0.3805	2	31	16	1	LLKIGEEHGAIAKLTGGGR
224	1002.7190	2003.4234	2004.0316	-0.6082	1	29	33	1	SFIITLGEALHEEPRDK
130	570.2700	1707.7882	1706.8913	0.8969	0	29	72	1	EIMVYSSLIGVDPR + Oxidation (M)
202	563.1430	1686.4072	1685.7971	0.6101	0	27	39	1	LNEMIFDFNTDVK
213	949.7190	1897.4234	1896.9074	0.5161	0	26	74	1	MVSIENFGMLSNQQQVK + Oxidation (M)
139	415.1260	1242.3562	1241.6616	0.6946	0	26	57	1	VSIPLSLAGQGEK
189	774.6890	1547.3634	1546.8103	0.5531	1	26	51	1	AGEALSAVGLDGFGR
220	654.1620	1959.4642	1958.9957	0.4684	1	25	50	1	LMTAYMAFAALESQKIR + Oxidation (M)
128	562.7700	1123.5254	1124.6441	-1.1187	0	25	75	1	EPGVAEIIIGK
113	493.8410	1478.5012	1477.7446	0.7566	2	25	1.7e+02	1	DIMDKETELTRK
211	918.5100	2752.5082	2751.3254	1.1827	2	25	1.5e+02	1	MDAEDAHHLTLRMLGAAGRTGLACAL
216	643.2610	1926.7612	1927.0091	-0.2479	1	24	75	1	TATAAYLKIPDQYTPFK
157	661.2900	1980.8482	1980.1269	0.7213	1	24	2.2e+02	1	SIINKAAIIGGGVIGAGWVAR
124	362.0050	1082.9932	1082.6448	0.3484	1	23	84	1	KVLTEIAGPR
61	358.1450	1071.4132	1072.6063	-1.1931	0	23	3.4e+02	1	QLLLTMAQR
190	776.2270	2325.6592	2325.2403	0.4189	0	23	1.7e+02	1	GSTLFPVIFLTGHGDVPMVAVALK + Oxidation (M)
121	354.9720	1061.8942	1062.5128	-0.6186	1	23	1.1e+02	1	ADVQEAERK + Oxidation (M)
163	689.3310	2064.9712	2065.0184	-0.0472	1	22	3e+02	1	EDKVAQMVSAIQGGALGMK + 2 Oxidation (M)
136	617.4450	1232.8754	1232.6976	0.1779	1	22	2.4e+02	1	EIKLALASSSSK
125	549.2190	1644.6352	1644.9093	-0.2742	2	22	2.7e+02	1	NLAGSVLVMGKGRATR + Oxidation (M)
215	640.2460	1917.7162	1918.8956	-1.1794	1	22	1.1e+02	1	MTDSPPAANGGREAFVQR + Oxidation (M)
235	1097.0470	2192.0794	2192.1664	-0.0869	1	22	2.2e+02	1	ATNILLAGKTIIVVAGYGCWK
86	422.3290	1263.9652	1263.6360	0.3292	0	22	3.4e+02	1	QLLASAYAAR
244	820.3240	2457.9502	2458.1984	-0.2482	2	22	1e+02	1	MMYAVGNSNERAAELAGAPVKAYK + 2 Oxidation (M)
105	467.9130	1400.7172	1401.7576	-1.0404	1	22	3.5e+02	1	RGETIGIIGTSGSK
181	743.4140	2227.2202	2227.2259	-0.0057	2	22	3.6e+02	1	AGKNVKAHLHQGAHITPTMPK + Oxidation (M)
93	867.4750	866.4677	865.4215	1.0462	0	22	1.7e+02	1	TLMEQTK + Oxidation (M)
219	974.3510	1946.6874	1945.9489	0.7386	0	21	1.3e+02	1	LQGDLDLMPATAVEEMIR + Oxidation (M)
116	497.2930	1488.8572	1489.7810	-0.9239	1	21	4.3e+02	1	TDGVQLVIKMSGK
184	751.3550	1500.6954	1499.7256	0.9698	0	21	3.8e+02	1	FTSTSVTNFSANPK
241	1162.2730	3483.7972	3482.7472	1.0499	1	21	2.3e+02	1	MNTLPTDNDPVQAWMALGVQSAILARTMPAVR + Oxidation (M)
70	387.1380	772.2614	772.4416	-0.1801	2	21	3.3e+02	1	GASARRR
242	1168.2350	3501.6832	3501.8752	-0.1921	2	21	2.7e+02	1	VHMASRGHALLADALYGRPGLGLTRQALHAHR

MASCOT Mascot Search Results

Protein View

Match to: **gi|53712412** Score: **105**
adenylate kinase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4302) 10_RJ2_01_1361.d\SSP (4302) 10_RJ2_01_1361.mgf

Nominal mass (M_r): **21018**; Calculated pI value: **5.10**
 NCBI BLAST search of [gi|53712412](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60680574](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253563556](#) from [Bacteroides sp. 3_2_5](#)
[gi|265762597](#) from [Bacteroides sp. 2_1_16](#)
[gi|68568759](#) from [Bacteroides fragilis](#)
[gi|81316357](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|52215277](#) from [Bacteroides fragilis YCH46](#)
[gi|60492008](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251947332](#) from [Bacteroides sp. 3_2_5](#)
[gi|263255205](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **12%**

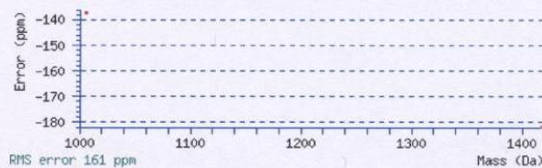
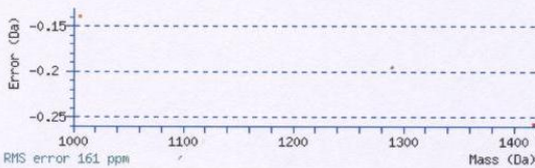
Matched peptides shown in **Bold Red**

- 1** MLNIVIFGAP GSGKGTQSER IVEKYGINHI STGDVLR AEI KNGTELGKTA
- 51** KGYIDGQQLI PDELMVDLLA SVFDSFKDSK **GVIFDGFPR** IPQAEALKVM
- 101** LKERGQDISV MLDLDPPEE LMTRLIKRGK ESGRADDNEE TIKKRLVVYV
- 151** TQTSPLKBYI KEGRYQHIN GLGTMGIFP DICKAVDTL

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
1 - 14	710.2580	1418.5014	1418.7592	-0.2577	0 -.MLNIVIFGAPGSGK.G Oxidation (M) (Ions score 54)
81 - 89	504.2000	1006.3854	1006.5236	-0.1382	0 K.GVIFDGFPR.T (Ions score 51)



LOCUS YP_098404 189 aa linear BCT 26-APR-2009
 DEFINITION adenylate kinase [Bacteroides fragilis YCH46].
 ACCESSION YP_098404
 VERSION YP_098404.1 GI:53712412
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 189)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 189)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 189)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT REVIEWED REFSEQ: This record has been curated by NCBI staff. The
 reference sequence was derived from [BAD47870](#).
 Method: conceptual translation.

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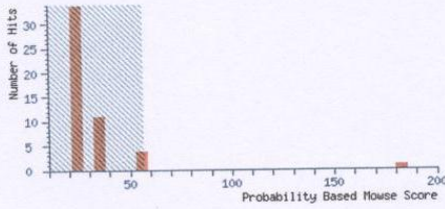
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : LAKSHMY MANICKAN
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4502
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4502) 10_RN2_01_1570.d\SSP (4502) 10_RN2_01_1570.mgf
 Database : NCBI nr 20090522 (8876587 sequences; 3036162093 residues)
 Taxonomy : Bacteria (Eubacteria) (4773688 sequences)
 Timestamp : 30 May 2009 at 10:44:47 GMT
 Protein hits : [gi|53715240](#) triosephosphate isomerase [Bacteroides fragilis YCH46]
 [gi|21674265](#) triosephosphate isomerase [Chlorobium tepidum TLS]
 [gi|34763378](#) Triosephosphate isomerase [Fusobacterium nucleatum subsp. vincentii ATCC 49256]
 [gi|78223155](#) triosephosphate isomerase [Geobacter metallireducens GS-15]
 [gi|189461507](#) hypothetical protein BACCOP_02166 [Bacteroides coprocola DSM 17136]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 56 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)
 Significance threshold p< Max. number of hits
 Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets
 Show pop-ups Suppress pop-ups Sort unassigned Require bold red
 Error tolerant

1. [gi|53715240](#) Mass: 26867 Score: 183 Queries matched: 4 emPAI: 0.42
 triosephosphate isomerase [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
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<input checked="" type="checkbox"/> 88	655.2560	1308.4974	1308.6714	-0.1740	1	54	0.074	1	K.VADFKGIIDAPN.-
<input checked="" type="checkbox"/> 118	808.8830	1615.7514	1615.8974	-0.1459	0	59	0.027	1	K.IVLAYEPVWAIPTGK.T
<input checked="" type="checkbox"/> 123	547.2460	1638.7162	1638.8478	-0.1316	0	39	2.7	1	K.TASPAQAEIHAFIR.S

2. [gi|21674265](#) Score: 59 Queries matched: 1
 triosephosphate isomerase [Chlorobium tepidum TLS]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
118	808.8830	1615.7514	1615.8974	-0.1459	0	59	0.027	1	K.LVIAYEPVWAIPTGK.T

Proteins matching the same set of peptides:
[gi|218295498](#) Score: 59 Queries matched: 1
[gi|227987582](#) Score: 59 Queries matched: 1
[gi|229231135](#) Score: 59 Queries matched: 1

3. [gi|34763378](#) Score: 59 Queries matched: 1
 Triosephosphate isomerase [Fusobacterium nucleatum subsp. vincentii ATCC 49256]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
118	808.8830	1615.7514	1615.8974	-0.1459	0	59	0.027	1	K.VIIAYEPVWAIPTGK.T

Proteins matching the same set of peptides:
[gi|1350021144](#) Score: 59 Queries matched: 1
[gi|197737446](#) Score: 59 Queries matched: 1
[gi|222055981](#) Score: 59 Queries matched: 1
[gi|237740886](#) Score: 59 Queries matched: 1
[gi|237745353](#) Score: 59 Queries matched: 1

{MATRIX}
{SCIENCE} Mascot Search Results

Protein View

Match to: gi|53715240 Score: 183
triosephosphate isomerase [Bacteroides fragilis YCH46]
Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4502) 10_RN2_01_1570.d\SSP (4502) 10_RN2_01_1570.mgf

Nominal mass (M_r): 26867; Calculated pI value: 5.14
NCBI BLAST search of gi|53715240 against nr
Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
Links to retrieve other entries containing this sequence from NCBI Entrez:
gi|60683174 from [Bacteroides fragilis NCTC 9343](#)
gi|81313773 from [Bacteroides fragilis NCTC 9343](#)
gi|81824915 from [Bacteroides fragilis](#)
gi|52218105 from [Bacteroides fragilis YCH46](#)
gi|60494608 from [Bacteroides fragilis NCTC 9343](#)

Fixed modifications: Carboxymethyl (C)
Variable modifications: Oxidation (M)
Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
Sequence Coverage: 21%

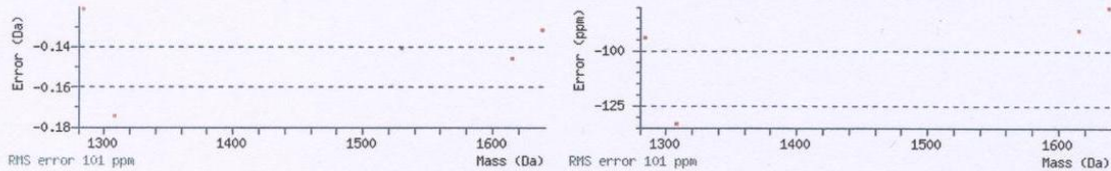
Matched peptides shown in **Bold Red**

1 MRKNIVAGNW KMNKTLQEGI ALAKELNEAL ANEKPNCVDI ICTPFIHLAS
51 VTPPLVDAKI GVGAENCADK ESGAYTGEVS AAMVASTGAK YVILGHSERR
101 **AYYGETVEIL** RDKVKLALAN GLTPIFCIGE VLEERRANKQ NEVVAAQLAS
151 VFDLSAEDFS KIVLAYEVPVW AIGTGKTASF AQAQEIHAPI RSAVAEKYKQ
201 ETADNTSILY GGSCKPSNAK ELFANPDVDG GLIGGAALKV ADFPKGIIDAF
251 N

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
101 - 111	643.2770	1284.5394	1284.6601	-0.1207	0	R.AYYGETVEILK.D (Ions score 30)
162 - 176	808.8830	1615.7514	1615.8974	-0.1459	0	K.IVLYEYVVAIGTK.T (Ions score 59)
177 - 191	547.2460	1638.7162	1638.8478	-0.1316	0	K.TASPAQAEIHAFIR.S (Ions score 39)
240 - 251	655.2560	1308.4974	1308.6714	-0.1740	1	K.VADFKGIIDAFN.- (Ions score 54)



LOCUS YP_101232 251 aa linear BCT 26-APR-2009
DEFINITION triosephosphate isomerase [Bacteroides fragilis YCH46].
ACCESSION YP_101232
VERSION YP_101232.1 GI:53715240
DBLINK Project:13067
DBSOURCE REFSEQ: accession NC_006347.1
KEYWORDS .
SOURCE Bacteroides fragilis YCH46
ORGANISM [Bacteroides fragilis YCH46](#)
Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
Bacteroidaceae; Bacteroides.
REFERENCE 1 (residues 1 to 251)
AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
inversions regulating cell surface adaptation
JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
PUBMED 15466707
REFERENCE 2 (residues 1 to 251)
CONSTRM NCBI Genome Project
TITLE Direct Submission
JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
Information, NIH, Bethesda, MD 20894, USA
REFERENCE 3 (residues 1 to 251)
AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
TITLE Direct Submission
JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan

COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from BAD50698.
 Method: conceptual translation.

FEATURES Location/Qualifiers

source 1..251
 /organism="Bacteroides fragilis YCH46"
 /strain="YCH46"
 /db_xref="taxon:295405"

Protein 1..251
 /product="triosephosphate isomerase"
 /EC_number="5.3.1.1"
 /calculated_mol_wt=26462

Region 5..248
 /region_name="TIM"
 /note="Triosephosphate isomerase (TIM) is a glycolytic
 enzyme that catalyzes the interconversion of
 dihydroxyacetone phosphate and
 D-glyceraldehyde-3-phosphate. The reaction is very
 efficient and requires neither cofactors nor metal ions.
 TIM, usually...; cd00311"
 /db_xref="CDD:73362"

Site order(9,11,96,167,173,213,232,234..235)
 /site_type="other"
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 /db_xref="CDD:73362"

Site order(9,12,45..47,49,52,65,83,86..87,98..99)
 /site_type="other"
 /note="dimer interface"
 /db_xref="CDD:73362"

Site order(11,96,167)
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 /note="catalytic triad"
 /db_xref="CDD:73362"

CDS 1..251
 /gene="tpiA"
 /locus_tag="BF3956"
 /coded_by="complement(NC_006347.1:4511582..4512337)"
 /note="Reversibly isomerizes the ketone sugar
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 glyceraldehyde-3-phosphate"
 /transl_table=11
 /db_xref="GeneID:3085037"

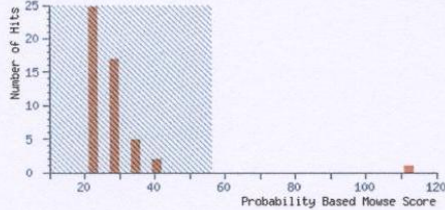
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : LAKSHMY MANICKAN
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4601
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4601) 10_RJ4_01_1365.d\SSP (4601) 10_RJ4_01_1365.mgf
 Database : NCBI nr 20090522 (8876587 sequences; 3036162093 residues)
 Taxonomy : Bacteria (Eubacteria) (4773688 sequences)
 Timestamp : 30 May 2009 at 10:46:21 GMT
 Protein hits : [gi|29346739](#) putative thiol peroxidase [Bacteroides thetaiotaomicron VPI-5482]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 56 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|29346739](#) Mass: 18208 Score: 112 Queries matched: 2 emPAI: 0.19
 putative thiol peroxidase [Bacteroides thetaiotaomicron VPI-5482]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 104	552.2650	1102.5154	1102.6386	-0.1232	0	46	0.72	1	K.LIGEFIQVGK.V
<input checked="" type="checkbox"/> 114	600.7170	1199.4194	1199.6332	-0.2138	0	67	0.0046	1	R.MADGFLAGLLAR.A + Oxidation (M)

Proteins matching the same set of peptides:
[gi|53714055](#) Mass: 18042 Score: 112 Queries matched: 2
 putative thiol peroxidase [Bacteroides fragilis YCH46]
[gi|153808277](#) Mass: 18064 Score: 112 Queries matched: 2
 hypothetical protein BACCAC_02565 [Bacteroides caccae ATCC 43185]
[gi|160882643](#) Mass: 18079 Score: 112 Queries matched: 2
 hypothetical protein BACOVA_00596 [Bacteroides ovatus ATCC 8483]
[gi|212700035](#) Mass: 17981 Score: 112 Queries matched: 2
 hypothetical protein BACFIN_02519 [Bacteroides finegoldii DSM 17565]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 237	766.9440	2297.8102	2298.1380	-0.3278	1	38	2	1	FDFRGVDASIELSGDDTITLK
<input checked="" type="checkbox"/> 108	568.4830	1702.4272	1702.8176	-0.3904	1	37	5.8	1	HFGVYEPAQESARGR
<input checked="" type="checkbox"/> 225	742.2740	2223.8002	2224.2467	-0.4465	2	34	5.8	1	AKKYLTVQVSEGGSLVLPVGTK
<input checked="" type="checkbox"/> 195	647.5700	1939.6882	1939.0093	0.6789	1	31	11	1	MLFSMAVGLNAVSMAAKAK
<input checked="" type="checkbox"/> 231	748.5940	2242.7602	2243.0892	-0.3290	0	31	10	1	TLDNMGAPEPNITLLWSER + Oxidation (M)
<input checked="" type="checkbox"/> 226	1113.5220	2225.0294	2224.0035	-1.0260	2	30	19	1	MCAEVARRGGELVDGGLMR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 241	1171.8740	3512.6002	3511.6639	0.9363	2	27	50	1	NESIRKSFIDLYENNTPIYAECCGLMYLQK + Oxidation (M)
<input checked="" type="checkbox"/> 223	1105.9620	2209.9094	2210.2232	-0.3138	0	27	34	1	GVLPLALLVLSGSLVLAGCDDK
<input checked="" type="checkbox"/> 230	746.9420	2237.8042	2238.1408	-0.3366	1	27	29	1	MFIGHFAPAFIAAAAYSRGPK + Oxidation (M)
<input checked="" type="checkbox"/> 134	679.3990	2035.1752	2035.0347	0.1405	2	25	1.3e+02	1	RYEHRALAEVQIEHER
<input checked="" type="checkbox"/> 211	700.9210	2099.7412	2098.9807	0.7605	2	25	47	1	ETNIKETFWESRTDSEK
<input checked="" type="checkbox"/> 146	719.7530	2156.2372	2156.1590	0.0782	0	25	1.2e+02	1	VYDLVVGGINGVAVADAAGR
<input checked="" type="checkbox"/> 222	737.6270	2209.8592	2211.0221	-1.1629	1	25	51	1	QILCGGCSITEAQADALMKK + Oxidation (M)
<input checked="" type="checkbox"/> 89	479.3080	1434.9022	1434.6959	0.2063	0	24	2e+02	1	IQQMEAGHLVQR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 111	592.2770	1182.5394	1182.5591	-0.0196	0	24	1e+02	1	MITSVFEDNK
<input checked="" type="checkbox"/> 243	1222.4520	2442.8894	2443.3798	-0.4904	2	23	95	1	LGTDAINKIADPHVLDELKLR
<input checked="" type="checkbox"/> 56	366.2100	1095.6082	1096.5149	-0.9067	0	23	2.8e+02	1	IGAGSGSYTER
<input checked="" type="checkbox"/> 116	612.3190	1222.6234	1222.6167	0.0068	1	23	1.1e+02	1	LHSTGPERNGR

Mascot Search Results

Protein View

Match to: gi|53714055 Score: 112
 putative thiol peroxidase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4601) 10_RJ4_01_1365.d\SSP (4601) 10_RJ4_01_1365.mgf

Nominal mass (M_r): 18042; Calculated pI value: 5.55
 NCBI BLAST search of gi|53714055 against nr
 Unformatted sequence string for pasting into other applications

Taxonomy: Bacteroides fragilis YCH46
 Links to retrieve other entries containing this sequence from NCBI Entrez:
 gi|60682249 from Bacteroides fragilis NCTC 9343
 gi|253564697 from Bacteroides sp. 3 2 5
 gi|265766326 from Bacteroides sp. 2 1 16
 gi|52216920 from Bacteroides fragilis YCH46
 gi|60493683 from Bacteroides fragilis NCTC 9343
 gi|251946162 from Bacteroides sp. 3 2 5
 gi|263253994 from Bacteroides sp. 2 1 16

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 13%

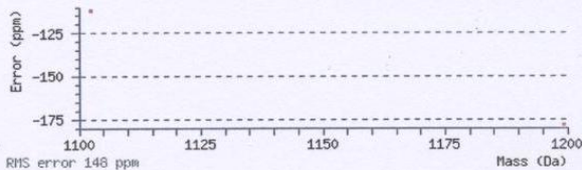
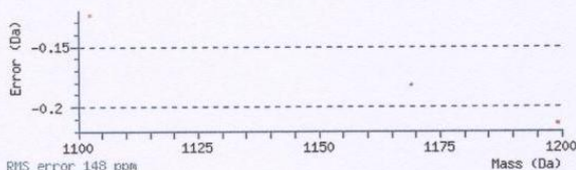
Matched peptides shown in **Bold Red**

1 MATTNFKGQP **VKLIGEFIQV** **GKVAPDFELV** KSDLSSFALK DLKGNIVLN
 51 IFPSLDTGVC ATSVRKFNKM AAGMKDVTVL AISKDLPFAQ GRFCTTEGIE
 101 NVIPLSDFRF SDFDESYGVR **MADGPLAGLL** ARAVVVIGKD GKVAYTELVP
 151 EITQEPDYEK ALAAVK

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
13 - 22	552.2650	1102.5154	1102.6386	-0.1232	0	K.LIGEFIQVK.V (Ions score 46)
121 - 132	600.7170	1199.4194	1199.6332	-0.2138	0	R.MADGPLAGLLAR.A Oxidation (M) (Ions score 67)



LOCUS YP_100047 166 aa linear BCT 26-APR-2009
 DEFINITION putative thiol peroxidase [Bacteroides fragilis YCH46].
 ACCESSION YP_100047
 VERSION YP_100047.1 GI:53714055
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS
 SOURCE Bacteroides fragilis YCH46
 ORGANISM Bacteroides fragilis YCH46
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 166)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 166)
 CONSRIM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 166)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from BAD49513.

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Method: conceptual translation.
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                    /strain="YCH46"
                    /db_xref="taxon:295405"
  Protein            1..166
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                    subfamily; composed of PRXs containing peroxidatic and
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                    specific antioxidant (TSA) protein also known as
                    TRX-dependent thiol peroxidase (Tpx). Tpx is a...;
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  Site                order(54,56,85..86,92,108,127..128)
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                    /note="dimer interface"
                    /db_xref="CDD:48563"
  Site                order(57,60,132)
                    /site_type="other"
                    /note="catalytic triad"
                    /db_xref="CDD:48563"
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                    /site_type="other"
                    /note="peroxidatic and resolving cysteines"
                    /db_xref="CDD:48563"
  CDS                1..166
                    /locus_tag="BF2763"
                    /coded_by="complement(NC_006347.1:3175264..3175764)"
                    /note="similar to gp:AE016931_120 [Bacteroides
                    thetaiotaomicron VPI-5482], percent identity 92 in 166 aa,
                    BLASTP E(): 9e-84"
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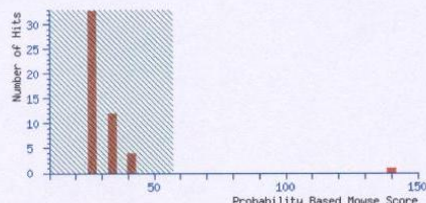
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4605
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4605) 10_R17_01_1311.d\SSP (4605) 10_R17_01_1311.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:36:18 GMT
 Protein hits : [gi|53712002](#) thioredoxin [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|53712002](#) Mass: 11637 Score: 140 Queries matched: 5 emPAI: 0.30
 thioredoxin [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 70	539.7250	1077.4354	1077.6223	-0.1868	0	34	11	1	R.NIPTVLFPPK.N
<input checked="" type="checkbox"/> 80	601.2670	1200.5194	1200.6424	-0.1229	0	48	0.47	1	K.MVGPIDELAK.E + Oxidation (M)
<input checked="" type="checkbox"/> 81	601.2980	1200.5814	1200.6424	-0.0609	0	(41)	2.6	1	K.MVGPIDELAK.E + Oxidation (M)
<input checked="" type="checkbox"/> 152	918.8010	1835.5874	1836.7836	-1.1962	0	58	0.031	1	K.CDVENSDDLPAEFGIR.N
<input checked="" type="checkbox"/> 153	612.8830	1835.6272	1836.7836	-1.1564	0	(24)	66	3	K.CDVENSDDLPAEFGIR.N

Proteins matching the same set of peptides:
[gi|255007522](#) Mass: 11635 Score: 138 Queries matched: 5
 putative thioredoxin [Bacteroides fragilis 3_1_12]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 90	665.2880	1328.5614	1328.7187	-0.1573	0	38	4.2	1	VINSDTVIADLK
<input checked="" type="checkbox"/> 126	819.8490	1637.6834	1637.7897	-0.1062	0	38	3.5	1	SESASFTPNTDIITR
<input checked="" type="checkbox"/> 96	692.2880	1382.5614	1382.8245	-0.2631	2	33	13	1	LSIALGKSGKPNPAK
<input checked="" type="checkbox"/> 83	609.2310	1216.4474	1215.6611	0.7863	1	32	18	1	KAYDPLIGPSR
<input checked="" type="checkbox"/> 89	651.7980	1301.5814	1302.7408	-1.1594	1	31	23	1	FIRGIVTDLGGK
<input checked="" type="checkbox"/> 227	761.9600	2282.8582	2282.0558	0.8023	2	31	13	1	DMKEASKSIWLDMENQVDK + Oxidation (M)
<input checked="" type="checkbox"/> 106	738.2980	1474.5814	1474.8143	-0.2329	1	30	27	1	DABELKQAPTALR
<input checked="" type="checkbox"/> 173	997.4120	1992.8094	1992.8999	-0.0905	2	29	25	1	AGRYKEASEYMPYEQK + Oxidation (M)
<input checked="" type="checkbox"/> 153	612.8830	1835.6272	1835.8975	-0.2703	0	28	30	1	CNLSSEPIENAILFTK
<input checked="" type="checkbox"/> 71	548.7450	1095.4754	1095.5448	-0.0693	1	28	43	1	DAETLEKYK
<input checked="" type="checkbox"/> 87	639.2820	1276.5494	1277.6867	-1.1372	1	27	57	1	GKLIIDELFSEK
<input checked="" type="checkbox"/> 186	1032.4920	2062.9694	2062.9636	0.0058	2	27	48	1	QRDVQRELAAMSGDNAR + Oxidation (M)
<input checked="" type="checkbox"/> 156	926.7690	1851.5234	1851.8859	-0.3624	1	27	38	1	MKIMTHSNDGFVISEK + Oxidation (M)
<input checked="" type="checkbox"/> 166	646.0030	1934.9872	1935.0178	-0.0306	2	26	62	1	KMGGMGTILGLMPGMGKIK + Oxidation (M)
<input checked="" type="checkbox"/> 174	665.2790	1992.8152	1991.9478	0.8673	0	26	51	1	MNDLTQTPLEMIATMGAR
<input checked="" type="checkbox"/> 240	837.2560	2508.7462	2508.1382	0.6080	2	26	41	1	MAGGAQMPSPSPKSTDMRIGAR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 112	782.0590	2343.1552	2342.1801	0.9751	2	25	1.3e+02	1	REVLAMTAIVIARYSSDNQR
<input checked="" type="checkbox"/> 190	1045.0920	3132.2542	3132.4803	-0.2262	0	25	1.2e+02	1	IFEHSGHAPHIEPEAFMNYLNLFLK
<input checked="" type="checkbox"/> 73	551.6520	1651.9342	1651.8286	0.1055	2	25	1.6e+02	1	MNKVIFACVRNAGR + Oxidation (M)
<input checked="" type="checkbox"/> 78	572.7520	1143.4894	1143.5448	-0.0554	0	25	1.7e+02	1	VTDDYAVFSK
<input checked="" type="checkbox"/> 216	742.2870	2223.8392	2224.2467	-0.4075	2	25	60	1	AKKYLTVQSEGGSLVLFVGTK
<input checked="" type="checkbox"/> 213	737.9170	2210.7292	2210.0718	0.6574	0	24	62	1	SEVEFAMYPFVHLLNAAK + Oxidation (M)
<input checked="" type="checkbox"/> 188	696.6030	2086.7872	2087.9768	-1.1897	1	24	73	1	GTDTNMPQSRSMNLGLPK + Oxidation (M)

MASCOT Mascot Search Results

Protein View

Match to: gi|53712002 Score: 140
 thioredoxin [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4605) 10_RI7_01_1311.d\SSP (4605) 10_RI7_01_1311.mgf

Nominal mass (M_n): 11637; Calculated pI value: 4.51
 NCBI BLAST search of gi|53712002 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: Bacteroides fragilis YCH46
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60680203](#) from Bacteroides fragilis NCTC 9343
[gi|253563963](#) from Bacteroides sp. 3_2_5
[gi|265765340](#) from Bacteroides sp. 2_1_16
[gi|52214867](#) from Bacteroides fragilis YCH46
[gi|60491637](#) from Bacteroides fragilis NCTC 9343
[gi|251947739](#) from Bacteroides sp. 3_2_5
[gi|263254724](#) from Bacteroides sp. 2_1_16

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 34%

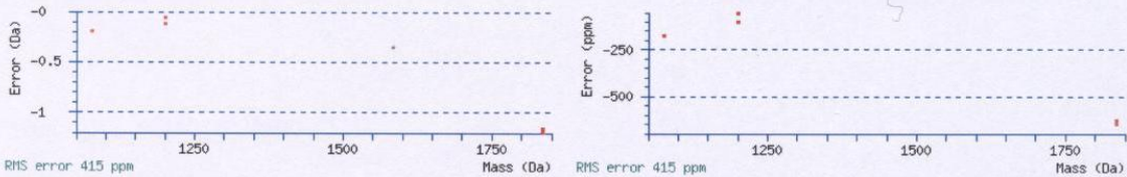
Matched peptides shown in **Bold Red**

1 MALEITDNNF KEILAEGSPV VIDFWAPWCG PCKMVGPIID ELAKEYEGKV
 51 IMGK**CDVDEN SDLPAEFGIR NIPTVLPFKN** GELVDKQVGA VGKPAFVEKV
 101 EKLL

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
34 - 44	601.2670	1200.5194	1200.6424	-0.1229	0	K.MVGPIIDELAK.E Oxidation (M) (Ions score 48)
34 - 44	601.2980	1200.5814	1200.6424	-0.0609	0	K.MVGPIIDELAK.E Oxidation (M) (Ions score 41)
55 - 70	918.8010	1835.5874	1836.7836	-1.1962	0	K.CDVDENSDLPAEFGIR.N (Ions score 58)
55 - 70	612.8830	1835.6272	1836.7836	-1.1564	0	K.CDVDENSDLPAEFGIR.N (Ions score 24)
71 - 79	539.7250	1077.4354	1077.6223	-0.1868	0	R.NIPTVLPFK.N (Ions score 34)



LOCUS YP_097994 104 aa linear BCT 26-APR-2009
 DEFINITION thioredoxin [Bacteroides fragilis YCH46].
 ACCESSION YP_097994
 VERSION YP_097994.1 GI:53712002
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 104)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 104)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 104)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan

COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD47460](#).
 Method: conceptual translation.

FEATURES Location/Qualifiers

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 /strain="YCH46"
 /db_xref="taxon:295405"

Protein 1..104
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 /calculated_mol_wt=11339

Region 8..101
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 /note="TRX family; composed of two groups: Group I, which
 includes proteins that exclusively encode a TRX domain;
 and Group II, which are composed of fusion proteins of TRX
 and additional domains. Group I TRX is a small ancient
 protein that alter the redox...; cd02947"
 /db_xref="CDD:48496"

Site order(29,32)
 /site_type="other"
 /note="catalytic residues"
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CDS 1..104
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 /coded_by="NC_006347.1:838887..839201"
 /note="similar to gp:AE016935_76 [Bacteroides
 thetaiotaomicron VPI-5482], percent identity 81 in 104 aa,
 BLASTP E(): 6e-47"
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 /db_xref="GeneID:3082512"

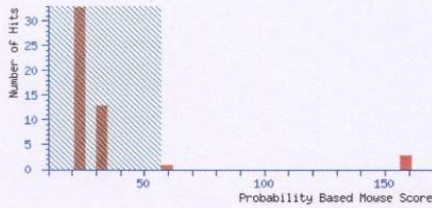
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4607
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4607) 10_RJ18_01_1401.d\SSP (4607) 10_RJ18_01_1401.mgf
 Database : NCBIInr 20100108 (10291680 sequences; 3511877860 residues)
 Taxonomy : Bacteria (Eubacteria) (5718488 sequences)
 Timestamp : 11 Jan 2010 at 10:40:03 GMT
 Protein hits : [gi|53715262](#) malate dehydrogenase [Bacteroides fragilis YCH46]
[gi|224536272](#) hypothetical protein BACCELL_01139 [Bacteroides cellulosilyticus DSM 14838]
[gi|160890687](#) hypothetical protein BACUNI_03132 [Bacteroides uniformis ATCC 8492]
[gi|150002856](#) malate dehydrogenase [Bacteroides vulgatus ATCC 8482]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|53715262](#) Mass: 32945 Score: 159 Queries matched: 4 emPAI: 0.21
 malate dehydrogenase [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 121	621.2850	1240.5554	1240.7027	-0.1472	0	41	2.3	1	R.EELIGVNAGIVK.S
<input checked="" type="checkbox"/> 175	853.8230	1705.6314	1705.8920	-0.2606	0	56	0.055	1	K.LNEVVASTMVGATLTK.L + Oxidation (M)
<input checked="" type="checkbox"/> 178	862.6630	2584.9672	2585.3121	-0.3449	0	21	2.9e+02	2	K.YSPNAILVVISNPMDTMTYLALK.S + 2 Oxidation (M)
<input checked="" type="checkbox"/> 248	906.0490	2715.1252	2715.4020	-0.2769	0	41	1.2	1	K.LLGTSAWYAPGAAGYVVESIIHNQK.K

Proteins matching the same set of peptides:
[gi|255011805](#) Mass: 32932 Score: 159 Queries matched: 4
 malate dehydrogenase [Bacteroides fragilis 3_1_12]

2. [gi|224536272](#) Mass: 33016 Score: 159 Queries matched: 4 emPAI: 0.21
 hypothetical protein BACCELL_01139 [Bacteroides cellulosilyticus DSM 14838]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 121	621.2850	1240.5554	1240.7027	-0.1472	0	41	2.3	1	R.EELIGVNAGIVK.T
<input type="checkbox"/> 175	853.8230	1705.6314	1705.8920	-0.2606	0	56	0.055	1	K.LNEVVASTMVGATLTK.L + Oxidation (M)
<input type="checkbox"/> 178	862.6630	2584.9672	2585.3121	-0.3449	0	21	2.9e+02	2	K.YSPNAILVVISNPMDTMTYLALK.S + 2 Oxidation (M)
<input type="checkbox"/> 248	906.0490	2715.1252	2715.4020	-0.2769	0	41	1.2	1	K.LLGTSAWYAPGAAGYVVESIIHNQK.K

3. [gi|160890687](#) Mass: 33051 Score: 159 Queries matched: 4 emPAI: 0.21
 hypothetical protein BACUNI_03132 [Bacteroides uniformis ATCC 8492]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 121	621.2850	1240.5554	1240.7027	-0.1472	0	41	2.3	1	R.EELIGVNAGIVK.S
<input type="checkbox"/> 175	853.8230	1705.6314	1705.8920	-0.2606	0	56	0.055	1	K.LNEVVASTMVGATLTK.L + Oxidation (M)
<input type="checkbox"/> 178	862.6630	2584.9672	2585.3121	-0.3449	0	20	3e+02	4	K.YSPNAILVVISNPMDTMTYLSLK.S + Oxidation (M)
<input type="checkbox"/> 248	906.0490	2715.1252	2715.4020	-0.2769	0	41	1.2	1	K.LLGTSAWYAPGAAGYVVESIIHNQK.K

4. [gi|150002856](#) Mass: 33174 Score: 62 Queries matched: 2
 malate dehydrogenase [Bacteroides vulgatus ATCC 8482]
 Check to include this hit in error tolerant search

{MATRIX} Mascot Search Results
{SCIENCE}

Protein View

Match to: gi|53715262 Score: 159
malate dehydrogenase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4607) 10_RJ18_01_1401.d\SSP (4607) 10_RJ18_01_1401.mgf

Nominal mass (M_n): 32945; Calculated pI value: 5.14
 NCBI BLAST search of gi|53715262 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: **Bacteroides fragilis YCH46**
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60683199](#) from **Bacteroides fragilis NCTC 9343**
[gi|253567153](#) from **Bacteroides sp. 3_2_5**
[gi|265767749](#) from **Bacteroides sp. 2_1_16**
[gi|73920985](#) from **Bacteroides fragilis NCTC 9343**
[gi|73920986](#) from **Bacteroides fragilis**
[gi|52218127](#) from **Bacteroides fragilis YCH46**
[gi|60494633](#) from **Bacteroides fragilis NCTC 9343**
[gi|251943984](#) from **Bacteroides sp. 3_2_5**
[gi|263252421](#) from **Bacteroides sp. 2_1_16**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **24%**

Matched peptides shown in **Bold Red**

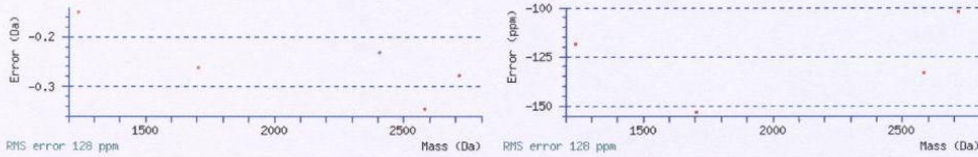
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1 MSKVTVVVAG NVGATCANVL APNEVADEVV MLDVKEGVSE GKAMDDMQTA
51 QLLGFDTTIV GCTNDYAQTA NSDVVVITSG IPRKPGMTR ELIGVNAGIV
101 ESVAENLLKY SPNALIVVIS NPMDTMTYLA LKSLGLPKNR VIGMGGALDS
151 SRFKYPLSQ LGCNANEVEG MVIGGHGDDT MIPLARLATY KGQFVSTLLS
201 EKLNVEVVA TMVGGATLTK LLGTSAWYAP GAAGAYVVES IIHNQKMMVP
251 CSVMLEGEYG ESDLCIGVFP ILGKNGIEKI VELELNADK AKFAASAAAV
301 HKTNAALKEV GAL
    
```

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
90 - 101	621.2850	1240.5554	1240.7027	-0.1472	0 R.ELIGVNAGIVK.S (Ions score 41)
110 - 132	862.6630	2584.9672	2585.3121	-0.3449	0 K.YSPNALIVVISNPMDTMTYLALK.S 2 Oxidation (M) (Ions score 21)
204 - 220	853.8230	1705.6314	1705.8920	-0.2606	0 K.LNEVVASTMVGGATLTK.L Oxidation (M) (Ions score 56)
221 - 246	906.0490	2715.1252	2715.4020	-0.2769	0 K.LLGTSAWYAPGAAGAYVVESIIHNQK.K (Ions score 41)



LOCUS YP_101254 313 aa linear BCT 26-APR-2009
 DEFINITION malate dehydrogenase [Bacteroides fragilis YCH46].
 ACCESSION YP_101254
 VERSION YP_101254.1 GI:53715262
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS
 SOURCE Bacteroides fragilis YCH46
 ORGANISM **Bacteroides fragilis YCH46**
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 313)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H., Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 313)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 313)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences, 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final NCBI review. The reference sequence was derived from [BAD50720](#). Method: conceptual translation.
 FEATURES
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/strain="YCH46"
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Region 20..286
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/note="L-lactate dehydrogenase-like malate dehydrogenase
proteins: cd01339"
Site /db_xref="CDD:133424"
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156,221,227..229,232..233,236)
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Site /db_xref="CDD:133424"
order(33..34,78..81,99,119,121,144,148,176)
/site_type="other"
/note="NAD(P) binding site"
Site /db_xref="CDD:133424"
order(54,56,162,167,169,171,189,191..193,243,245..248,274,
283)
/site_type="other"
/note="tetramer (dimer of dimers) interface"
Site /db_xref="CDD:133424"
order(89,121,152,176,215,226)
/site_type="other"
/note="substrate binding site"
/db_xref="CDD:133424"
CDS 1..313
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/coded_by="complement (NC_006347.1:4546190..4547131)"
/note="similar to gp:AB016942_184 [Bacteroides
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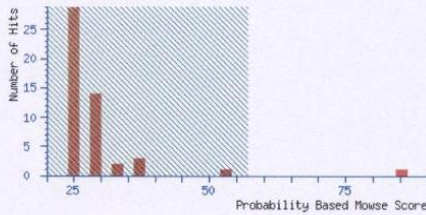
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4608
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4608) 10_RJ14_01_1388.d\SSP (4608) 10_RJ14_01_1388.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 16:28:30 GMT
 Protein hits : [gi|53715262](#) malate dehydrogenase [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p < 0.05$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

- 1. [gi|53715262](#) Mass: 32945 Score: 85 Queries matched: 2 emPAI: 0.10
 malate dehydrogenase [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
<input checked="" type="checkbox"/> 99	621.2790	1240.5434	1240.7027	-0.1592	0 34 12 1	R.EELIGVNAVIGV.S
<input checked="" type="checkbox"/> 246	906.0440	2715.1102	2715.4020	-0.2919	0 51 0.12 1	K.LLGTSAWYAPGAGAYVVESIIHNQK.K

Proteins matching the same set of peptides:

- [gi|160890687](#) Mass: 33051 Score: 85 Queries matched: 2
 hypothetical protein BACUNI_03132 [Bacteroides uniformis ATCC 8492]
- [gi|167765305](#) Mass: 33007 Score: 85 Queries matched: 2
 hypothetical protein BACSTE_03693 [Bacteroides stercoris ATCC 43183]
- [gi|189463781](#) Mass: 32987 Score: 85 Queries matched: 2
 hypothetical protein BACINT_00114 [Bacteroides intestinalis DSM 17393]
- [gi|224536272](#) Mass: 33016 Score: 85 Queries matched: 2
 hypothetical protein BACCELL_01139 [Bacteroides cellulosilyticus DSM 14838]
- [gi|255011805](#) Mass: 32932 Score: 85 Queries matched: 2
 malate dehydrogenase [Bacteroides fragilis 3_1_12]
- [gi|154494237](#) Mass: 33179 Score: 83 Queries matched: 2
 hypothetical protein PARMER_03587 [Parabacteroides merdae ATCC 43184]
- [gi|218263133](#) Mass: 33190 Score: 83 Queries matched: 2
 hypothetical protein PRABACTJOHN_03033 [Parabacteroides johnsonii DSM 18315]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect Rank	Peptide
<input checked="" type="checkbox"/> 245	878.6520	2632.9342	2633.3125	-0.3784	0 53 0.079 1	LPYANNALEPVISQQTIDYHYGK
<input checked="" type="checkbox"/> 192	1011.0100	2020.0054	2019.9949	0.0106	1 39 3 1	MGRIFVSPHAAEGGALADR + Oxidation (M)
<input checked="" type="checkbox"/> 211	1062.9500	2123.8854	2124.0535	-0.1680	1 38 3.6 1	VPEGLLSPIYGGGGGHDMRR
<input checked="" type="checkbox"/> 136	738.7680	1475.5214	1475.7330	-0.2116	0 35 6.2 1	GGEVFLDMGPEVK
<input checked="" type="checkbox"/> 112	645.4580	1933.3522	1932.1600	1.1921	0 34 16 1	NILFIPILAALIGYFVR
<input checked="" type="checkbox"/> 202	694.9150	2081.7232	2080.9405	-0.7827	1 33 8.5 1	TGVMQAEDDLNLESMEKR + Oxidation (M)
<input checked="" type="checkbox"/> 157	854.1340	2559.3802	2560.2445	-0.8644	1 31 32 1	KVGYESDFPESGGTPSLASSLLNR
<input checked="" type="checkbox"/> 108	427.1590	1278.4552	1278.7085	-0.2533	2 30 23 1	RKTGFEGLPFK
<input checked="" type="checkbox"/> 121	678.7620	1355.5094	1354.7681	0.7414	2 30 27 1	DRKNVSLTGIPR
<input checked="" type="checkbox"/> 137	740.6230	2218.8472	2219.1474	-0.3002	0 28 50 1	DYTLQVVLVDASGTPLYTHK
<input checked="" type="checkbox"/> 199	1023.3230	3066.9472	3067.6376	-0.6904	1 28 44 1	LNDIDTLFGLPGIPITDLLRMAQAEGLR + Oxidation (M)
<input checked="" type="checkbox"/> 68	468.2760	1401.8062	1400.8286	0.9776	2 28 1.1e+02 1	VRALGLRMLSSK

MASCOT Mascot Search Results

Protein View

Match to: [gi|53715262](#) Score: 85
malate dehydrogenase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4608) 10_RJ14_01_1388.d\SSP (4608) 10_RJ14_01_1388.mgf

Nominal mass (M_n): 32945; Calculated pI value: 5.14
 NCBI BLAST search of [gi|53715262](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60683199](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253567153](#) from [Bacteroides sp. 3_2_5](#)
[gi|265767749](#) from [Bacteroides sp. 2_1_16](#)
[gi|73920985](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|73920986](#) from [Bacteroides fragilis](#)
[gi|52218127](#) from [Bacteroides fragilis YCH46](#)
[gi|60494633](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251943984](#) from [Bacteroides sp. 3_2_5](#)
[gi|263252421](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 12%

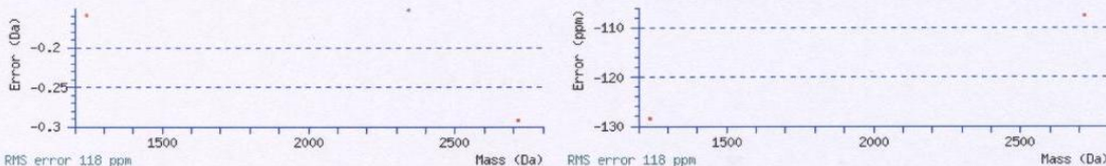
Matched peptides shown in **Bold Red**

- 1 MSKVTVVGAG NVGATCANVL AFNEVADEVV MLDVKEGVSE GKAMDMMQTA
- 51 QLLGPDTTIV GCTNDYAQTA NSDVVVITSG IPRKFGMTR**E ELIGVNAGIV**
- 101 **K**SVAENLLKY SPNAIIVVIS NPMDTMTYLA LKSLGLPKNR VIGMGGALDS
- 151 SRPKYFLSQA LGCNANEVEG MVIGGHGDTT MIPLARLATY KGQPVSTLLS
- 201 EEKLN**B**VVAS TMVGGATLTK **LLGTSAWYAP GAAGAYV**V**ES I**I**HNQ**K**MMVP**
- 251 CSVMLEGEY**G** ESDLCIGVPV ILGKNGIEKI VELELNAD**E**K AKFAASAA**V**
- 301 HKTNAALKEV GAL

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
90 - 101	621.2790	1240.5434	1240.7027	-0.1592	0	R .EELIGVNAGIVK.S (Ions score 34)
221 - 246	906.0440	2715.1102	2715.4020	-0.2919	0	K .LLGTSAWYAPGAAGAYV V ES I IHNQ K .K (Ions score 51)



LOCUS YP_101254 313 aa linear BCT 26-APR-2009
 DEFINITION malate dehydrogenase [Bacteroides fragilis YCH46].
 ACCESSION YP_101254
 VERSION YP_101254.1 GI:53715262
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 313)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 313)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 313)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan

COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final NCBI review. The reference sequence was derived from [BAD50720](#).
Method: conceptual translation.

FEATURES Location/Qualifiers

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/organism="Bacteroides fragilis YCH46"
/strain="YCH46"
/db_xref="taxon:295405"

Protein 1..313
/product="malate dehydrogenase"
/calculated_mol_wt=32545

Region 20..286
/region_name="LDH-like MDH"
/note="L-lactate dehydrogenase-like malate dehydrogenase proteins; cd01339"
/db_xref="CDD:133424"

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/site_type="other"
/note="dimer interface"
/db_xref="CDD:133424"

Site order(33..34,78..81,99,119,121,144,148,176)
/site_type="other"
/note="NAD(P) binding site"
/db_xref="CDD:133424"

Site order(54,56,162,167,169,171,189,191..193,243,245..248,274,283)
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/note="tetramer (dimer of dimers) interface"
/db_xref="CDD:133424"

Site order(89,121,152,176,215,226)
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/note="substrate binding site"
/db_xref="CDD:133424"

CDS 1..313
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/coded_by="complement(NC_006347.1:4546190..4547131)"
/note="similar to gp:AE016942_184 [Bacteroides thetaiotaomicron VPI-5482], percent identity 92 in 313 aa, BLASTP E(): e-163"
/transl_table=11
/db_xref="GeneID:3085059"

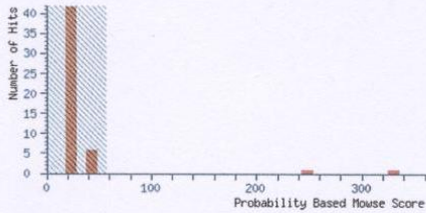
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4702
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4702) 10_RJ11_01_1379.d\SSP (4702) 10_RJ11_01_1379.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:37:28 GMT
 Protein hits : [gi|53715294](#) elongation factor Ts [Bacteroides fragilis YCH46]
 [gi|255011774](#) elongation factor Ts [Bacteroides fragilis 3_1_12]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|53715294](#) Mass: 36339 Score: 331 Queries matched: 7 emPAI: 0.69
 elongation factor Ts [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 122	623.7650	1245.5154	1245.6969	-0.1814	0	68	0.0044	1	K.VEEGFGAIIALK.C
<input checked="" type="checkbox"/> 139	686.3340	1370.6534	1370.7405	-0.0871	0	61	0.023	1	K.LTQDILDAAVANK.C
<input checked="" type="checkbox"/> 182	847.2930	1692.5714	1693.7539	-1.1824	0	39	2.5	1	K.EVCLLNQEDIMDAK.K + Oxidation (M)
<input checked="" type="checkbox"/> 244	772.6100	2314.8082	2315.1679	-0.3597	0	41	1.2	1	K.TLEEVLALPMGDATVAQAVTDR.T + Oxidation (M)
<input checked="" type="checkbox"/> 184	849.5650	2545.6732	2546.1239	-0.4508	0	61	0.026	1	K.MELDGYMVLLEGATIAAYNHMNR.N + 3 Oxidation (M)
<input checked="" type="checkbox"/> 248	849.6500	2545.9282	2546.1239	-0.1958	0	(52)	0.089	2	K.MELDGYMVLLEGATIAAYNHMNR.N + 3 Oxidation (M)
<input checked="" type="checkbox"/> 249	854.6580	2560.9522	2561.1989	-0.2467	0	64	0.0067	1	K.QVAMQVAAMNPIAVDEGVSEEVK.Q + 2 Oxidation (M)

2. [gi|255011774](#) Mass: 36340 Score: 257 Queries matched: 6 emPAI: 0.55
 elongation factor Ts [Bacteroides fragilis 3_1_12]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 139	686.3340	1370.6534	1370.7405	-0.0871	0	61	0.023	1	K.LTQDILDAAVANK.C
<input type="checkbox"/> 182	847.2930	1692.5714	1693.7539	-1.1824	0	39	2.5	1	K.EVCLLNQEDIMDAK.K + Oxidation (M)
<input type="checkbox"/> 244	772.6100	2314.8082	2315.1679	-0.3597	0	41	1.2	1	K.TLEEVLALPMGDATVAQAVTDR.S + Oxidation (M)
<input type="checkbox"/> 184	849.5650	2545.6732	2546.1239	-0.4508	0	(52)	0.24	2	K.MELDGYMVLLEGATISAYNHMNR.N + 2 Oxidation (M)
<input checked="" type="checkbox"/> 248	849.6500	2545.9282	2546.1239	-0.1958	0	56	0.04	1	K.MELDGYMVLLEGATISAYNHMNR.N + 2 Oxidation (M)
<input checked="" type="checkbox"/> 249	854.6580	2560.9522	2561.1989	-0.2467	0	64	0.0067	1	K.QVAMQVAAMNPIAVDEGVSEEVK.Q + 2 Oxidation (M)

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 152	742.8350	2225.4832	2225.2168	0.2663	1	34	19	1	RLDVRPLASGVVILTYGPDG
<input checked="" type="checkbox"/> 234	742.3310	2223.9712	2225.0825	-1.1113	2	30	22	1	SSPFGNDTKVGSAPQANKR
<input checked="" type="checkbox"/> 101	537.8890	1610.6452	1610.8338	-0.1886	1	30	56	1	DKVSIITYGSMAAK
<input checked="" type="checkbox"/> 168	822.2630	1642.5114	1641.7742	0.7372	0	29	25	1	YVELMTPFVIMDGR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 64	371.0920	1110.2542	1109.5023	0.7519	0	29	43	1	SMTGDVANATK + Oxidation (M)
<input checked="" type="checkbox"/> 132	665.2920	1328.5694	1328.7122	-0.1428	0	27	62	1	QVDIPAMLVATR + Oxidation (M)
<input checked="" type="checkbox"/> 233	742.2680	2223.7822	2223.2725	0.5096	2	26	37	1	KNILPQEKLTLLIEVSEK
<input checked="" type="checkbox"/> 138	679.3950	2035.1632	2033.9952	1.1679	1	26	1.2e+02	1	ARNVTIAEFPMSADAAGAAR + Oxidation (M)
<input checked="" type="checkbox"/> 208	970.7680	1939.5214	1939.9053	-0.3839	1	25	49	1	MDKEMVIDLVTQMAR + 2 Oxidation (M)

Mascot Search Results

Protein View

Match to: gi|53715294 Score: 331
 elongation factor Ts [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4702) 10_RJ11_01_1379.d\SSP (4702) 10_RJ11_01_1379.mgf

Nominal mass (M_n): 36339; Calculated pI value: 5.09
 NCBI BLAST search of gi|53715294 against nr
 Unformatted sequence string for pasting into other applications

Taxonomy: *Bacteroides fragilis* YCH46
 Links to retrieve other entries containing this sequence from NCBI Entrez:
 gi|60683229 from *Bacteroides fragilis* NCTC 9343
 gi|253567185 from *Bacteroides* sp. 3_2_5
 gi|265767781 from *Bacteroides* sp. 2_1_16
 gi|60389524 from *Bacteroides fragilis*
 gi|81313719 from *Bacteroides fragilis* NCTC 9343
 gi|52218159 from *Bacteroides fragilis* YCH46
 gi|60494663 from *Bacteroides fragilis* NCTC 9343
 gi|251944016 from *Bacteroides* sp. 3_2_5
 gi|263252453 from *Bacteroides* sp. 2_1_16

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 32%

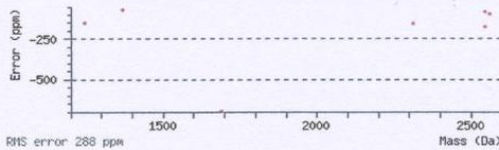
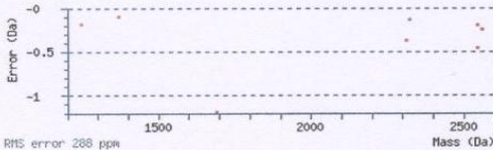
Matched peptides shown in Bold Red

1 MAVTMADITK LRKMTGAGMM DCKNALTDAE GDFDKAMKII REKQQA~~VA~~AK
 51 RSDREASEGC VLVKVEEGFG AIIALKCETD FVAQNAD~~FK~~ LTQDILDAAV
 101 ANKCKTLEEV LALPMGDATV AQAVTDRGTI TGEKMELDGY MVLEGATIAA
 151 YNHMNRNGLC TMVAFNKVD EQLAKQVAMQ VAAMNPIAVD EDGVSEEVKQ
 201 KEIEVAVEKT KVEQVQKAVE AALKKANINP AHVDS~~ED~~HME SNMAKGWITA
 251 EDVAKAKEII ATVSAEKAAN MPEQMIONIA KGRLAK~~PL~~KE VCLLNQEDIM
 301 DAKKT~~VR~~EVL KEADPELKVV DPKRPTLRAE

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss Sequence
65 - 76	623.7650	1245.5154	1245.6969	-0.1814	0 K.VEEFGGAIILK.C (Ions_score 68)
91 - 103	686.3340	1370.6534	1370.7405	-0.0871	0 K.LTQDILDAAVANK.C (Ions_score 61)
106 - 127	772.6100	2314.8082	2315.1679	-0.3597	0 K.TLEVLALPMGDATVAQAVTDR.T Oxidation (M) (Ions_score 41)
135 - 156	849.5650	2545.6732	2546.1239	-0.4508	0 K.MELDGYMVLEGATIAAYNHMNR.N 3 Oxidation (M) (Ions_score 61)
135 - 156	849.6500	2545.9282	2546.1239	-0.1958	0 K.MELDGYMVLEGATIAAYNHMNR.N 3 Oxidation (M) (Ions_score 52)
176 - 199	854.6580	2560.9522	2561.1989	-0.2467	0 K.QVAMQVAAMNPIAVDE GV SEEVK.Q 2 Oxidation (M) (Ions_score 64)
290 - 303	847.2930	1692.5714	1693.7539	-1.1824	0 K.EVCLLNQEDIMDAK.K Oxidation (M) (Ions_score 39)



LOCUS YP_101286 330 aa linear BCT 26-APR-2009
 DEFINITION elongation factor Ts [Bacteroides fragilis YCH46].
 ACCESSION YP_101286
 VERSION YP_101286.1 GI:53715294
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM Bacteroides fragilis YCH46
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 330)
 AUTHORS Kuwahara,T., Yamashita,A., Hiraoka,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 330)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 330)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from BAD50752.
 Method: conceptual translation.
 FEATURES Location/Qualifiers

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/strain="YCH46"
/db_xref="taxon:295405"
Protein 1..330
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/note="elongation factor Ts; Provisional; PRK09377"
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Region 56..225
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/note="EF-Ts; functions during elongation stage of protein
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complex and promotes exchange of GDP to GTP resulting in
regeneration of the active form of EF-Tu"
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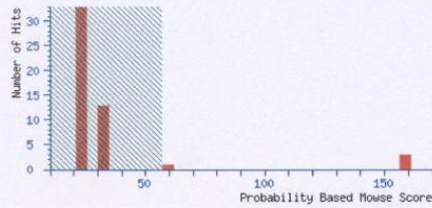
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4705
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4705) 10_RJ18_01_1401.d\SSP (4705) 10_RJ18_01_1401.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:38:38 GMT
 Protein hits : [gi|53715262](#) malate dehydrogenase [Bacteroides fragilis YCH46]
[gi|224536272](#) hypothetical protein BACCELL_01139 [Bacteroides cellulosilyticus DSM 14838]
[gi|160890687](#) hypothetical protein BACUNI_03132 [Bacteroides uniformis ATCC 8492]
[gi|150002856](#) malate dehydrogenase [Bacteroides vulgatus ATCC 8482]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event.
 Individual ions scores > 57 indicate identity or extensive homology (p<0.05).
 Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As Peptide Summary Help

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

- [gi|53715262](#) Mass: 32945 Score: 159 Queries matched: 4 empAI: 0.21
 malate dehydrogenase [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 121	621.2850	1240.5554	1240.7027	-0.1472	0	41	2.2	1	R.EELIGVNAGIVK.S
<input checked="" type="checkbox"/> 175	853.8230	1705.6314	1705.8920	-0.2606	0	56	0.054	1	K.LNEVVASTMVGATLTK.L + Oxidation (M)
<input checked="" type="checkbox"/> 178	862.6630	2584.9672	2585.3121	-0.3449	0	21	2.9e+02	2	K.YSPNAILVVISNPMDTMTYLALK.S + 2 Oxidation (M)
<input checked="" type="checkbox"/> 248	906.0490	2715.1252	2715.4020	-0.2769	0	41	1.2	1	K.LLGTSAWYAPGAAGAYVVESIIHNQK.K

Proteins matching the same set of peptides:
[gi|255011805](#) Mass: 32932 Score: 159 Queries matched: 4
 malate dehydrogenase [Bacteroides fragilis 3_1_12]
- [gi|224536272](#) Mass: 33016 Score: 159 Queries matched: 4 empAI: 0.21
 hypothetical protein BACCELL_01139 [Bacteroides cellulosilyticus DSM 14838]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
121	621.2850	1240.5554	1240.7027	-0.1472	0	41	2.2	1	R.EELIGVNAGIVK.T
175	853.8230	1705.6314	1705.8920	-0.2606	0	56	0.054	1	K.LNEVVASTMVGATLTK.L + Oxidation (M)
178	862.6630	2584.9672	2585.3121	-0.3449	0	21	2.9e+02	2	K.YSPNAILVVISNPMDTMTYLALK.S + 2 Oxidation (M)
248	906.0490	2715.1252	2715.4020	-0.2769	0	41	1.2	1	K.LLGTSAWYAPGAAGAYVVESIIHNQK.K
- [gi|160890687](#) Mass: 33051 Score: 159 Queries matched: 4 empAI: 0.21
 hypothetical protein BACUNI_03132 [Bacteroides uniformis ATCC 8492]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
121	621.2850	1240.5554	1240.7027	-0.1472	0	41	2.2	1	R.EELIGVNAGIVK.S
175	853.8230	1705.6314	1705.8920	-0.2606	0	56	0.054	1	K.LNEVVASTMVGATLTK.L + Oxidation (M)
178	862.6630	2584.9672	2585.3121	-0.3449	0	20	3e+02	4	K.YSPNAILVVISNPMDTMTYLSLK.S + Oxidation (M)
248	906.0490	2715.1252	2715.4020	-0.2769	0	41	1.2	1	K.LLGTSAWYAPGAAGAYVVESIIHNQK.K
- [gi|150002856](#) Mass: 33174 Score: 62 Queries matched: 2
 malate dehydrogenase [Bacteroides vulgatus ATCC 8482]
 Check to include this hit in error tolerant search

Mascot Search Results

Protein View

Match to: [gi|53715262](#) Score: 159
malate dehydrogenase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTBOMICS 3- 151208\SSP (4705) 10_RJ18_01_1401.d\SSP (4705) 10_RJ18_01_1401.mgf

Nominal mass (M_r): 32945; Calculated pI value: 5.14
 NCBI BLAST search of [gi|53715262](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60683199](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253567153](#) from [Bacteroides sp. 3_2_5](#)
[gi|265767749](#) from [Bacteroides sp. 2_1_16](#)
[gi|73920985](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|73920986](#) from [Bacteroides fragilis](#)
[gi|52218127](#) from [Bacteroides fragilis YCH46](#)
[gi|60494633](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251943984](#) from [Bacteroides sp. 3_2_5](#)
[gi|263252421](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 24%

Matched peptides shown in **Bold Red**

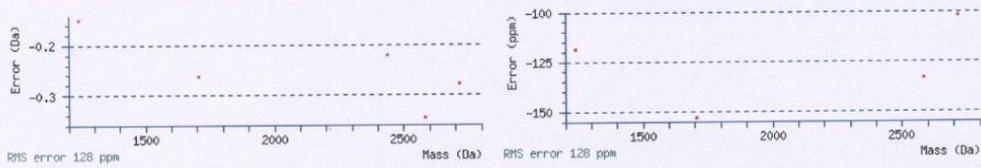
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1 MSKVTVVGAG NVGATCANVL AFNEVADEVV MLDVKEGVSE GKAMDMMQTA
51 QLLGFDTTIV GCTNDYAQTA NSDVVVITSG IPRKPGMTRELIGVNAGIV
101 KSVANLLKY SFNAIIVVIS NFMDTMTYLA LKSLGLPKNR VIGMGGALDS
151 SRFYFLSQA LGCNANEVVG MVVGGHDTT MIPLARLATY KGQPVSTLLS
201 BEKLINEVVAS TMVGGATLTK LLGTSAWYAP GAAGAYVVES IIHNQKKMVP
251 CSVMLEGBYQ ESDLICIGVPV ILKNGIEKI VELELNADEK AKFAASAAAV
301 HKTNAALKEV GAL
  
```

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
90 - 101	621.2850	1240.5554	1240.7027	-0.1472	0 R.BELIGVNAGIVK.S (Ions score 41)
110 - 132	862.6630	2584.9672	2585.3121	-0.3449	0 K.YSPNAIIVVISNFMDDTMTYLALK.S 2 Oxidation (M) (Ions score 21)
204 - 220	853.8230	1705.6314	1705.8920	-0.2606	0 K.LNEVVASTMVGGATLTK.L Oxidation (M) (Ions score 56)
221 - 246	906.0490	2715.1252	2715.4020	-0.2769	0 K.LLGTSAWYAPGAAGAYV V ES I IHNQK K (Ions score 41)



LOCUS YP_101254 313 aa linear BCT 26-APR-2009
 DEFINITION malate dehydrogenase [Bacteroides fragilis YCH46].
 ACCESSION YP_101254
 VERSION YP_101254.1 GI:53715262
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 313)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H., Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 313)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 313)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences, 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final NCBI review. The reference sequence was derived from [BAD50720](#). Method: conceptual translation.
 FEATURES Location/Qualifiers
 source 1..313 /organism="Bacteroides fragilis YCH46"

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/strain="YCH46"
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Region 20..286
/region_name="LDH-like MDH"
/note="L-lactate dehydrogenase-like malate dehydrogenase
proteins; cd01339"
/db_xref="CDD:133424"
Site order(22,37..38,41,44..45,47..48,52,54..56,151..152,154,
156,221,227..229,232..233,236)
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Site order(33..34,78..81,99,119,121,144,148,176)
/site_type="other"
/note="NAD(P) binding site"
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Site order(54,56,162,167,169,171,189,191..193,243,245..248,274,
283)
/site_type="other"
/note="tetramer (dimer of dimers) interface"
/db_xref="CDD:133424"
Site order(89,121,152,176,215,226)
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CDS 1..313
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/note="similar to gp:AB016942_184 [Bacteroides
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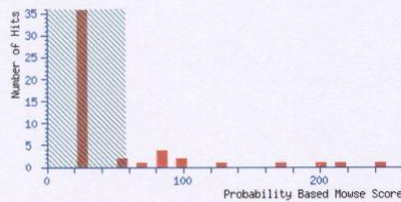
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 4804
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4804) 100_RE5_01_1707.d\SSP (4804) 100_RE5_01_1707.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:39:46 GMT
 Protein hits : [gi|223043068](#) pyridoxine biosynthesis protein [Staphylococcus capitis SK14]
[gi|228475242](#) pyridoxine biosynthesis protein [Staphylococcus hominis SK119]
[gi|15923509](#) pyridoxal biosynthesis lyase PdxS [Staphylococcus aureus subsp. aureus Mu50]
[gi|53715262](#) malate dehydrogenase [Bacteroides fragilis YCH46]
[gi|167765305](#) hypothetical protein BACSTE_03693 [Bacteroides stercoris ATCC 43183]
[gi|224536272](#) hypothetical protein BACCELL_01139 [Bacteroides cellulosilyticus DSM 14838]
[gi|160890687](#) hypothetical protein BACUNI_03132 [Bacteroides uniformis ATCC 8492]
[gi|229542250](#) pyridoxine biosynthesis protein [Bacillus coagulans 36D1]
[gi|23100142](#) pyridoxine biosynthesis protein [Oceanobacillus ihyensensis HTE831]
[gi|160947252](#) hypothetical protein PEPMIC_01185 [Parvimonas micra ATCC 33270]
[gi|239617197](#) pyridoxine biosynthesis protein [Kosmotoga olearia TBF 19.5.1]
[gi|83592546](#) malate dehydrogenase [Rhodospirillum rubrum ATCC 11170]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: **Peptide Summary** Help

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|223043068](#) Mass: 32030 Score: 244 Queries matched: 5 emPAI: 0.48
 pyridoxine biosynthesis protein [Staphylococcus capitis SK14]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 155	693.7620	1385.5094	1385.7150	-0.2056	0	55	0.083	1	K.GLDINQISLEER.M
<input checked="" type="checkbox"/> 169	773.3010	1544.5874	1544.7868	-0.1994	0	83	0.00013	1	K.IAEAGAVAVMALER.V + Oxidation (M)
<input checked="" type="checkbox"/> 179	830.2580	1658.5014	1658.7532	-0.2517	0	64	0.0068	1	R.LTVMNDDEIMTPAK.D + 2 Oxidation (M)
<input checked="" type="checkbox"/> 202	650.5870	1948.7392	1948.9894	-0.2503	0	(21)	1.4e+02	2	K.AIVQATTHYQDYELIGK.L
<input checked="" type="checkbox"/> 203	975.4180	1948.8214	1948.9894	-0.1680	0	42	1.4	1	K.AIVQATTHYQDYELIGK.L

Proteins matching the same set of peptides:
[gi|239636916](#) Mass: 31970 Score: 244 Queries matched: 5
 pyridoxine biosynthesis protein [Staphylococcus warneri L37603]
[gi|242372735](#) Mass: 32072 Score: 244 Queries matched: 5
 pyridoxine (vitamin B6) biosynthesis protein [Staphylococcus epidermidis M23864:W1]
[gi|27469180](#) Mass: 32048 Score: 242 Queries matched: 5
 pyridoxine biosynthesis protein [Staphylococcus epidermidis ATCC 12228]
[gi|251809935](#) Mass: 32038 Score: 242 Queries matched: 5
 pyridoxine (vitamin B6) biosynthesis protein [Staphylococcus epidermidis BCM-HMP0060]

2. [gi|228475242](#) Mass: 32014 Score: 215 Queries matched: 5 emPAI: 0.35
 pyridoxine biosynthesis protein [Staphylococcus hominis SK119]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 155	693.7620	1385.5094	1385.7514	-0.2420	1	55	0.083	1	K.GLDINKISLEER.M
<input checked="" type="checkbox"/> 169	773.3010	1544.5874	1544.7868	-0.1994	0	83	0.00013	1	R.IAEAGAVAVMALER.V + Oxidation (M)
<input checked="" type="checkbox"/> 179	830.2580	1658.5014	1658.7531	-0.2517	0	35	5.4	2	R.LTVMNDDEIMTYAK.E + Oxidation (M)
<input checked="" type="checkbox"/> 202	650.5870	1948.7392	1948.9894	-0.2503	0	(21)	1.4e+02	2	K.AIVQATTHYQDYELIGK.L
<input checked="" type="checkbox"/> 203	975.4180	1948.8214	1948.9894	-0.1680	0	42	1.4	1	K.AIVQATTHYQDYELIGK.L

3. [gi|15923509](#) Mass: 32088 Score: 202 Queries matched: 3 emPAI: 0.34
 pyridoxal biosynthesis lyase PdxS [Staphylococcus aureus subsp. aureus Mu50]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
155	693.7620	1385.5094	1385.7150	-0.2056	0	55	0.083	1	K.GLDINQLSLEER.M
169	773.3010	1544.5874	1544.7868	-0.1994	0	83	0.00013	1	R.IAEAGAVAVMALER.V + Oxidation (M)
179	830.2580	1658.5014	1658.7532	-0.2517	0	64	0.0068	1	R.LIVMDEIHTFAK.D + 2 Oxidation (M)

Proteins matching the same set of peptides:

[gi|82750226](#) Mass: 31832 Score: 202 Queries matched: 3
 pyridoxine biosynthesis protein [Staphylococcus aureus RF122]

4. [gi|53715262](#) Mass: 32945 Score: 171 Queries matched: 5 emPAI: 0.21
 malate dehydrogenase [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 136	621.2730	1240.5314	1240.7027	-0.1712	0	33	15	1	R.BELIGVNAGIVK.S
<input checked="" type="checkbox"/> 141	636.7330	1271.4514	1271.6609	-0.2094	0	78	0.0004	1	K.IVLELNNADEK.A
<input checked="" type="checkbox"/> 247	862.6770	2585.0092	2585.3121	-0.3029	0	21	1.2e+02	1	K.YSPNAILVVISNPMDTMTYLALK.S + 2 Oxidation (M)
<input checked="" type="checkbox"/> 250	992.4190	2974.2352	2974.5533	-0.3181	1	40	1.6	1	K.GQPVSTLLSEKLENEVVASTMVGATLTK.L + Oxidation (M)
251	992.4990	2974.4752	2974.5533	-0.0781	1	(17)	3.6e+02	2	K.GQPVSTLLSEKLENEVVASTMVGATLTK.L + Oxidation (M)

5. [gi|167765305](#) Mass: 33007 Score: 130 Queries matched: 3 emPAI: 0.10
 hypothetical protein BACSTE_03693 [Bacteroides stercoris ATCC 43183]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
136	621.2730	1240.5314	1240.7027	-0.1712	0	33	15	1	R.BELIGVNAGIVK.S
141	636.7330	1271.4514	1271.6609	-0.2094	0	78	0.0004	1	K.IVLELNNADEK.A
247	862.6770	2585.0092	2585.3121	-0.3029	0	20	1.6e+02	3	K.YSPNAILVVISNPMDTMTYLSLK.S + Oxidation (M)

Proteins matching the same set of peptides:

[gi|218128506](#) Mass: 32901 Score: 130 Queries matched: 3
 hypothetical protein BACEGG_00076 [Bacteroides eggerthii DSM 20697]

6. [gi|224536272](#) Mass: 33016 Score: 93 Queries matched: 4 emPAI: 0.10
 hypothetical protein BACCELL_01139 [Bacteroides cellulosilyticus DSM 14838]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
136	621.2730	1240.5314	1240.7027	-0.1712	0	33	15	1	R.BELIGVNAGIVK.T
247	862.6770	2585.0092	2585.3121	-0.3029	0	21	1.2e+02	1	K.YSPNAILVVISNPMDTMTYLALK.S + 2 Oxidation (M)
250	992.4190	2974.2352	2974.5533	-0.3181	1	40	1.6	1	K.GQPVSTLLSEKLENEVVASTMVGATLTK.L + Oxidation (M)
251	992.4990	2974.4752	2974.5533	-0.0781	1	(17)	3.6e+02	2	K.GQPVSTLLSEKLENEVVASTMVGATLTK.L + Oxidation (M)

7. [gi|160890687](#) Mass: 33051 Score: 92 Queries matched: 4 emPAI: 0.10
 hypothetical protein BACUNI_03132 [Bacteroides uniformis ATCC 8492]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
136	621.2730	1240.5314	1240.7027	-0.1712	0	33	15	1	R.BELIGVNAGIVK.S
247	862.6770	2585.0092	2585.3121	-0.3029	0	20	1.6e+02	3	K.YSPNAILVVISNPMDTMTYLSLK.S + Oxidation (M)
250	992.4190	2974.2352	2974.5533	-0.3181	1	40	1.6	1	K.GQPVSTLLSEKLENEVVASTMVGATLTK.L + Oxidation (M)
251	992.4990	2974.4752	2974.5533	-0.0781	1	(17)	3.6e+02	2	K.GQPVSTLLSEKLENEVVASTMVGATLTK.L + Oxidation (M)

8. [gi|229542250](#) Score: 83 Queries matched: 1
 pyridoxine biosynthesis protein [Bacillus coagulans 36D1]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
169	773.3010	1544.5874	1544.7868	-0.1994	0	83	0.00013	1	K.LAEAGAVAVMALER.V + Oxidation (M)

9. [gi|23100142](#) Score: 82 Queries matched: 1
 pyridoxine biosynthesis protein [Oceanobacillus iheyensis HTE831]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
169	773.3010	1544.5874	1543.8028	0.7847	0	82	0.00013	3	K.IAEQAGAVAVMALER.V + Oxidation (M)

10. [gi|160947252](#) Score: 82 Queries matched: 1
 hypothetical protein PEPMIC_01185 [Parvimonas micra ATCC 33270]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
169	773.3010	1544.5874	1543.8392	0.7483	1	82	0.00013	3	K.IAEKAGAVAVMALER.I + Oxidation (M)

11. [gi|239617197](#) Score: 80 Queries matched: 1

{MATRIX} SCIENCE Mascot Search Results

Protein View

Match to: gi|53715262 Score: 171
malate dehydrogenase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (4804) 100_RE5_01_1707.d\SSP (4804) 100_RE5_01_1707.mgf

Nominal mass (M_r): 32945; Calculated pI value: 5.14
 NCBI BLAST search of gi|53715262 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: **Bacteroides fragilis YCH46**
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60683199](#) from **Bacteroides fragilis NCTC 9343**
[gi|253567153](#) from **Bacteroides sp. 3_2_5**
[gi|265767749](#) from **Bacteroides sp. 2_1_16**
[gi|73920985](#) from **Bacteroides fragilis NCTC 9343**
[gi|73920986](#) from **Bacteroides fragilis**
[gi|52218127](#) from **Bacteroides fragilis YCH46**
[gi|60494633](#) from **Bacteroides fragilis NCTC 9343**
[gi|251943984](#) from **Bacteroides sp. 3_2_5**
[gi|263252421](#) from **Bacteroides sp. 2_1_16**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 23%

Matched peptides shown in **Bold Red**

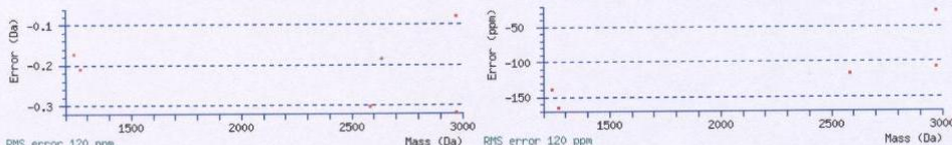
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51 QLLGFDTTIV GCTNDYQAQTA NSDVVVITSG IPRKPGMIRE ELIGVNAGIV
101 KVAENLLKY SPNALIVVIS NPMDTMTYLA LKSLGLPKNR VIGMGGLDLS
151 SRFYFLSQA LGCNANEVEG MVIGGHGDTT MIPLARLATY KQPFVSTLLS
201 BEKLEVVVAS TMVGGATLTK LLGTSAWYAP GAAGAYVVES IHNKQKMVP
251 CSVMLEGEYG ESDLGIGVEV ILGRNGIEKI VELELNADEK AKFAASAAAV
301 HKTNAALKEV GAL
    
```

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
90 - 101	621.2730	1240.5314	1240.7027	-0.1712	0 R.EELIGVNAGIVK.S (Ions score 33)
110 - 132	862.6770	2585.0092	2585.3121	-0.3029	0 K.YSPNALIVVISNPMDTMTYLALK.S 2 Oxidation (M) (Ions score 21)
192 - 220	992.4190	2974.2352	2974.5533	-0.3181	1 K.GQPVSTLLSEKLINEVVASTMVGGATLTK.L Oxidation (M) (Ions score 40)
192 - 220	992.4990	2974.4752	2974.5533	-0.0781	1 K.GQPVSTLLSEKLINEVVASTMVGGATLTK.L Oxidation (M) (Ions score 17)
280 - 290	636.7330	1271.4514	1271.6609	-0.2094	0 K.IVELELNADEK.A (Ions score 78)



LOCUS YP_101254 313 aa linear BCT 26-APR-2009
 DEFINITION malate dehydrogenase [Bacteroides fragilis YCH46].
 ACCESSION YP_101254
 VERSION YP_101254.1 GI:53715262
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM **Bacteroides fragilis YCH46**
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 313)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 313)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 313)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
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 /site_type="other"
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 /db_xref="CDD:133424"
Site order(54,56,162,167,169,171,189,191..193,243,245..248,274,283)
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 /note="tetramer (dimer of dimers) interface"
 /db_xref="CDD:133424"
Site order(89,121,152,176,215,226)
 /site_type="other"
 /note="substrate binding site"
 /db_xref="CDD:133424"
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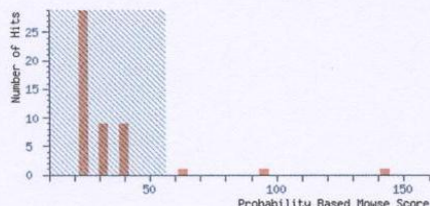
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : LAKSHMY MANICKAN
 Email : lakshmy.manickan@unn.ac.uk
 Search title : 5106 bf
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (5106) 10_R15_01_1306.d\SSP (5106) 10_R15_01_1306.mgf
 Database : NCBI nr 20090528 (8915381 sequences; 3049521622 residues)
 Taxonomy : Bacteria (Eubacteria) (4801653 sequences)
 Timestamp : 6 Jun 2009 at 11:30:52 GMT
 Protein hits : [gi|126496](#) RecName: Full=Lysostaphin; AltName: Full=Glycyl-glycine endopeptidase; Flags: Precursor
 [gi|29348144](#) 50S ribosomal protein L7/L12 [Bacteroides thetaiotaomicron VPI-5482]
 [gi|53713908](#) hypothetical protein BF2616 [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ion scores > 56 indicate identity or extensive homology (p<0.05). Protein scores are derived from ion scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|126496](#) Mass: 51656 Score: 142 Queries matched: 3 emPAI: 0.13
 RecName: Full=Lysostaphin; AltName: Full=Glycyl-glycine endopeptidase; Flags: Precursor
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 104	819.8400	1637.6654	1637.7897	-0.1242	0	57	0.044	1	K.SESASFTPTNDIITR.T
<input checked="" type="checkbox"/> 166	1023.3990	2044.7934	2044.9234	-0.1400	0	86	4.2e-05	1	R.MVNSPNSNTAQDPMPLK.S + 2 Oxidation (M)
<input checked="" type="checkbox"/> 167	682.6050	2044.7932	2044.9234	-0.1302	0	(30)	16	1	R.MVNSPNSNTAQDPMPLK.S + 2 Oxidation (M)

Proteins matching the same set of peptides:
[gi|153047](#) Mass: 42213 Score: 142 Queries matched: 3
 lysostaphin (ttg start codon)
[gi|3287967](#) Mass: 53058 Score: 142 Queries matched: 3
 RecName: Full=Lysostaphin; AltName: Full=Glycyl-glycine endopeptidase; Flags: Precursor

2. [gi|29348144](#) Mass: 12730 Score: 93 Queries matched: 1 emPAI: 0.27
 50S ribosomal protein L7/L12 [Bacteroides thetaiotaomicron VPI-5482]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 83	666.8460	1331.6774	1331.7449	-0.0674	0	93	1.2e-05	1	K.AFAEQLVNLTVK.E

Proteins matching the same set of peptides:
[gi|53715478](#) Mass: 12688 Score: 93 Queries matched: 1
 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]
[gi|110639543](#) Mass: 12652 Score: 93 Queries matched: 1
 50S ribosomal protein L7/L12 [Cytophaga hutchinsonii ATCC 33406]
[gi|149278891](#) Mass: 12813 Score: 93 Queries matched: 1
 50S ribosomal protein L7/L12 [Pedobacter sp. BAL39]
[gi|150003396](#) Mass: 12704 Score: 93 Queries matched: 1
 50S ribosomal protein L7/L12 [Bacteroides vulgatus ATCC 8482]
[gi|150008875](#) Mass: 12727 Score: 93 Queries matched: 1
 50S ribosomal protein L7/L12 [Parabacteroides distasonis ATCC 8503]
[gi|153805949](#) Mass: 12686 Score: 93 Queries matched: 1
 hypothetical protein BACCAC_00194 [Bacteroides caccae ATCC 43185]
[gi|154491722](#) Mass: 12679 Score: 93 Queries matched: 1
 hypothetical protein PARMER_01333 [Parabacteroides merdae ATCC 43184]
[gi|160888407](#) Mass: 12775 Score: 93 Queries matched: 1

hypothetical protein BACUNI_00820 [Bacteroides uniformis ATCC 8492]
 gi|189460676 Mass: 12771 Score: 93 Queries matched: 1
 hypothetical protein BACCOPI_01323 [Bacteroides coprocola DSM 17136]
 gi|189465410 Mass: 12759 Score: 93 Queries matched: 1
 hypothetical protein BACINT_01763 [Bacteroides intestinalis DSM 17393]
 gi|198277435 Mass: 12757 Score: 93 Queries matched: 1
 hypothetical protein BACPLE_03654 [Bacteroides plebeius DSM 17135]
 gi|224023599 Mass: 12771 Score: 93 Queries matched: 1
 hypothetical protein BACCOPRO_00306 [Bacteroides coprophilus DSM 18228]
 gi|224537808 Mass: 12757 Score: 93 Queries matched: 1
 hypothetical protein BACCELL_02695 [Bacteroides cellulosilyticus DSM 14838]
 gi|227413422 Mass: 13015 Score: 93 Queries matched: 1
 LSU ribosomal protein L12P [Dyadobacter fermentans DSM 18053]

3. gi|53713908 Mass: 12967 Score: 60 Queries matched: 3
 hypothetical protein BF2616 [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
✓ 120	590.5810	1768.7212	1768.9723	-0.2511	0	33	8.8	1	K.LPKFNLIAAQYVEGK.I
✓ 121	885.4150	1768.8154	1768.9723	-0.1568	0	(17)	4.9e+02	1	K.LPKFNLIAAQYVEGK.I
✓ 219	763.3310	2286.9712	2287.0756	-0.1045	1	28	25	1	K.KLAEFASYNLNLEVPENR.E

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
✓ 88	681.0850	2040.2332	2040.0528	0.1804	1	39	4	1	EVFVSGLEREYLGDTAGSLK
✓ 108	559.9450	1676.8132	1676.8958	-0.0827	1	37	4.7	1	DGVVAVVRSHSPDLAR
✓ 179	1041.9630	2081.9114	2080.9955	0.9159	1	36	5	1	TMREIDAMEVMGVSVIR + Oxidation (M)
✓ 109	843.7830	2528.3272	2528.2693	0.0579	1	30	33	1	IWSAGVAASPLGKMQVABQAGVRAER + Oxidation (M)
✓ 204	737.6530	2209.9372	2211.0598	-1.1227	0	29	23	1	EHRPPVMMSTASQMLVER + Oxidation (M)
✓ 209	742.3440	2224.0102	2225.2063	-1.1961	2	28	32	1	MLLSLPVQQGGVNLSSRRR + Oxidation (M)
✓ 205	1105.9790	2209.9434	2210.1306	-0.1872	1	28	30	1	COIVVNLKTHIQFGAYR
✓ 243	929.9330	2786.7772	2787.4627	-0.6855	1	26	30	1	HSIPLTAALALGAGAAQAAPTSTDEAR
✓ 236	878.4340	2632.2802	2633.3517	-1.0715	1	26	48	1	MDLNLQDKVIVTGGASGIGGATSMR + 2 Oxidation (M)
✓ 37	819.1320	818.1247	817.4480	0.6767	1	26	55	1	MGIDKVR
✓ 202	731.8270	2192.4592	2192.1114	0.3478	0	26	39	1	QLDPSGAVGWVVAALADLGLGAE
✓ 70	572.2670	1713.7792	1713.8210	-0.0418	0	25	1.6e+02	1	DIIEHPBEGSGGYIK
✓ 231	834.3320	2499.9742	2499.3261	0.6481	1	25	43	1	LTAEQVLDKTIPLFNIVYENK
✓ 247	968.7290	2903.1652	2903.5604	-0.3952	2	24	42	1	RARLFQDEASVEDLATILTIILSSDK
✓ 184	700.9170	2099.7292	2100.1263	-0.3971	1	24	52	1	GGFRHLIVTSGGGVAGILSMR + Oxidation (M)
✓ 126	608.5840	1822.7302	1821.8508	0.8793	1	24	68	1	FARGMWARVGYGTAYK + Oxidation (M)
✓ 163	675.9900	2024.9482	2026.0769	-1.1287	0	24	86	1	GAVINMTGPSVDVNLGALTVV
✓ 172	688.2020	2061.5842	2062.0272	-0.4431	1	24	61	1	WPGSDYDGLVWVVTIRNGK
✓ 178	594.9630	2081.8672	2081.0800	0.7872	2	24	78	1	GHDLSGRRITVSTCGIVPR
✓ 235	866.7260	2597.1562	2597.2386	-0.0825	2	23	79	1	MIADGEGATKLIECTVMNVSGEK + Oxidation (M)
✓ 73	592.7900	1183.5654	1184.6765	-1.1110	1	23	1.2e+02	1	LKPTGNKVTLD
✓ 171	688.1910	2061.5512	2062.0259	-0.4747	0	23	70	1	IDVLNNSLIGSTFYSYBYK
✓ 240	914.1200	2739.3382	2739.3717	-0.0335	0	23	88	1	YFIVPEYHILTLTTPDISIMEDLK + Oxidation (M)
✓ 146	647.6210	1939.8412	1940.0699	-0.2287	1	23	1e+02	1	LRLLANVLPAMDQMAR + Oxidation (M)
✓ 139	945.0180	2832.0322	2831.4424	0.5898	1	23	1.7e+02	1	MBIAALVIVMGHPAVYAPIYKADR + 2 Oxidation (M)
✓ 246	937.0270	2808.0592	2807.4449	0.6143	1	23	62	1	TSAITGWKIVGVSPAEAMTSQVLMLSK + Oxidation (M)
✓ 203	1103.5350	3307.5832	3307.7346	-0.1514	2	22	1.9e+02	1	VLNAKYHEKEARIIAQAGAPGALTIATMMAGR
✓ 220	767.3070	2298.8992	2299.1193	-0.2201	1	22	83	1	KVSNWGHNLTVSGNVTSNK
✓ 248	978.0340	2931.0802	2930.4670	0.6132	2	22	67	1	LKADCDSPKLVYAHISPYMGPEIIR
✓ 218	1130.4240	3388.2502	3387.5509	0.6993	2	22	1.4e+02	1	EQTEVPMVMWSPGFAASFGVNDKCLQRR + Oxidation (M)
✓ 227	813.3450	2437.0132	2438.1060	-1.0928	0	22	1e+02	1	PGFVEMPNDEBEGNAIAALNEK + Oxidation (M)
✓ 148	976.3900	2926.1482	2926.4519	-0.3038	1	22	1.9e+02	1	SRDIDREGLALALAAAADLTGDDLEAGLDR
✓ 113	864.0980	2589.2722	2588.3996	0.8726	1	21	2.5e+02	1	LGMGKPLGLGSAVAINYELQLTDRK + Oxidation (M)
✓ 140	632.7910	1895.3512	1894.8447	0.5064	2	21	1.1e+02	1	SADRANMNMGMRRDIK
✓ 180	695.6620	2083.9642	2083.9844	-0.0203	1	21	1.5e+02	1	DNIGKMLSPFTTENGQGTK + Oxidation (M)
✓ 101	804.1330	2409.3772	2409.3168	0.0603	2	21	2.6e+02	1	LVELVEALPFGARDGWLYPGRK
✓ 154	663.5160	1987.5262	1988.0626	-0.5364	2	21	1.2e+02	1	VRVDYANGVLQVMGPKAR + Oxidation (M)
✓ 102	809.7900	2426.3482	2425.3428	1.0054	2	21	2.8e+02	1	ANIIAKSGKAVEVAGDITVVLDK
✓ 250	989.7350	2966.1832	2966.5059	-0.3227	2	21	93	1	NSDYVIFVQNKIYVSLNKLMSSEK + Oxidation (M)
✓ 133	618.7790	1853.3152	1853.8465	-0.5313	0	21	1.2e+02	1	LESVLDEAEMANSFGR + Oxidation (M)
✓ 196	1081.5960	3241.7662	3240.8632	0.9030	2	21	2.8e+02	1	MTLDKLGSAFLGLGIALASLGVLPALARGISTLR + Oxidation (M)
✓ 74	623.6370	1867.8892	1866.9112	0.9780	0	21	3.3e+02	1	YLNDFSYAGSIIGTSGR
✓ 115	874.7990	2621.3752	2621.3418	0.0334	1	21	2.9e+02	1	GHEVAAAGHNLMTGPPGAGKTLAR + 2 Oxidation (M)
✓ 221	771.7050	2312.0932	2312.0590	0.0341	1	21	1.7e+02	1	TADLGCSRQDIVNAYDNLGL
✓ 190	704.7750	2111.3032	2111.0371	0.2661	1	21	1.4e+02	1	PSHPMGAVVLEARGGFSAGR + Oxidation (M)
✓ 149	656.6840	1967.0302	1966.9935	0.0367	1	21	1.9e+02	1	MPDGKHWVIGTVDAQK + Oxidation (M)
✓ 194	1075.9480	3224.8222	3224.6032	0.2190	0	20	2.7e+02	1	MNGLAVSGAVLATAVGGFDPAPMIQMAALGAR + 3 Oxidation (M)
✓ 75	624.5210	1870.5412	1870.9398	-0.3986	2	20	3.3e+02	1	DFGATHEELAGRLGRSR
✓ 206	1106.2650	3315.7732	3315.5621	0.2111	1	20	2.8e+02	1	GGGIANIVNGFFGGMAGCAMIQSINVKSSGR + Oxidation (M)
✓ 111	854.3570	1706.6994	1706.8125	-0.1130	1	20	2.1e+02	1	NARYASSFDVGEHVR
✓ 71	581.2250	1740.6532	1741.7968	-1.1436	0	20	4.4e+02	1	ILDDYIEMBEELSK + Oxidation (M)
✓ 197	1082.6650	3244.9732	3245.5746	-0.6014	1	20	2.8e+02	1	VAGVGDAMVNGVDYSMRILWLPDVMQAQYK + Oxidation (M)

Mascot Search Results

Protein View

Match to: gi|53715478 Score: 93
 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (5106) 10_RI5_01_1306.d\SSP (5106) 10_RI5_01_1306.mgf

Nominal mass (M_r): 12688; Calculated pI value: 4.71
 NCBI BLAST search of gi|53715478 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60683451](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253566657](#) from [Bacteroides sp. 3_2_5](#)
[gi|255011634](#) from [Bacteroides fragilis 3_1_12](#)
[gi|265767535](#) from [Bacteroides sp. 2_1_16](#)
[gi|81313501](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|81609231](#) from [Bacteroides fragilis](#)
[gi|52218343](#) from [Bacteroides fragilis YCH46](#)
[gi|60494885](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251944829](#) from [Bacteroides sp. 3_2_5](#)
[gi|263252840](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 9%

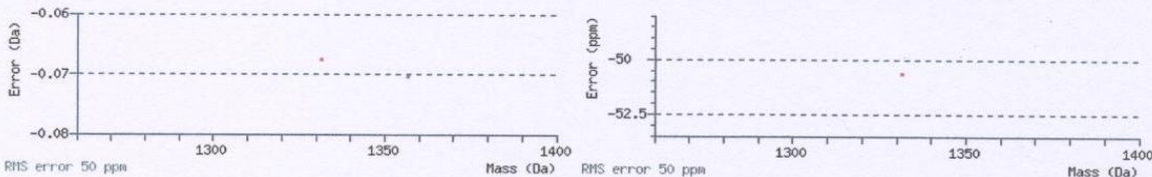
Matched peptides shown in **Bold Red**

1 MADLK**APAEQ** LVN**LTVK**EVN ELATILKEEY GIEPAAAAVA VAAGPAAGAA
 51 AAEKSSFDV VLKSAGAAKL QVVKAVKEAC GLGLKEAKDM VDGAPSVVKE
 101 GLAKDEAESL KKTLEEAGAE VELK

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
6 - 17	666.8460	1331.6774	1331.7449	-0.0674	0	K.APAEQLVNLTVK.E (Ions score 93)



LOCUS YP_101470 124 aa linear BCT 26-APR-2009
 DEFINITION 50S ribosomal protein L7/L12 [Bacteroides fragilis YCH46].
 ACCESSION YP_101470
 VERSION YP_101470.1 GI:53715478
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM Bacteroides fragilis YCH46
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 124)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 124)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 124)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT REVIEWED REFSEQ: This record has been curated by NCBI staff. The

reference sequence was derived from [BAD50936](#).
Method: conceptual translation.

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                  /strain="YCH46"
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                  L12 dimers bind L10; critically important for translation
                  efficiency and fidelity; stimulates GTPase activity of
                  translation factors"
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Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

Protein View

Match to: gi|53713908 Score: 60
 hypothetical protein BF2616 [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (5106) 10_RI5_01_1306.d\SSP (5106) 10_RI5_01_1306.mgf

Nominal mass (M_r): 12967; Calculated pI value: 5.28
 NCBI BLAST search of gi|53713908 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60682117](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253567099](#) from [Bacteroides sp. 3 2 5](#)
[gi|265764254](#) from [Bacteroides sp. 2 1 16](#)
[gi|52216773](#) from [Bacteroides fragilis YCH46](#)
[gi|60493551](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251944223](#) from [Bacteroides sp. 3 2 5](#)
[gi|263256862](#) from [Bacteroides sp. 2 1 16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 31%

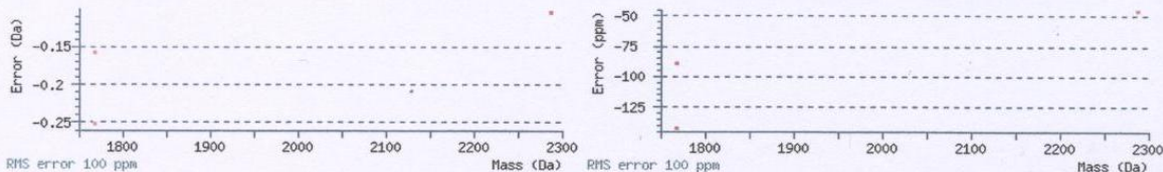
Matched peptides shown in **Bold Red**

1 MDYKKTNAPT NTITRDMDL CADTGNVYET VAIIGKRANQ ISVEIKNDLS
51 KKLAEFASYN DNLEEVFENR EQIEISRYE KLPKPNLIAA QEYVEGKIY
 101 RNPAAKEKEKL Q

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss Sequence
52 - 70	763.3310	2286.9712	2287.0756	-0.1045	1 K.KLAEFASYN DNLEEVFENR.E (Ions score 28)
82 - 97	590.5810	1768.7212	1768.9723	-0.2511	0 K.LPKPNLIAAQEYVEGK.I (Ions score 33)
82 - 97	885.4150	1768.8154	1768.9723	-0.1568	0 K.LPKPNLIAAQEYVEGK.I (Ions score 17)



LOCUS YP_099900 111 aa linear BCT 26-APR-2009
 DEFINITION hypothetical protein BF2616 [Bacteroides fragilis YCH46].
 ACCESSION YP_099900
 VERSION YP_099900.1 GI:53713908
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM Bacteroides fragilis YCH46
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 111)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 111)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 111)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD49366](#).

Method: conceptual translation.
FEATURES Location/Qualifiers
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/strain="YCH46"
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Protein 1..111
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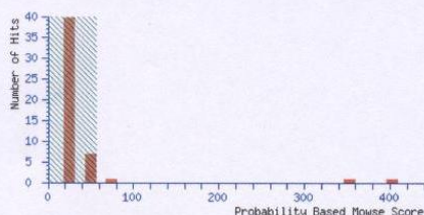
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 5205
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (5205) 10_RJ7_01_1371.d\SSP (5205) 10_RJ7_01_1371.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:41:02 GMT
 Protein hits : [gi|53712495](#) hypothetical protein BF1203 [Bacteroides fragilis YCH46]
 [gi|255007972](#) hypothetical protein Bfra3_02458 [Bacteroides fragilis 3_1_12]
 [gi|153807766](#) hypothetical protein BACCAC_02049 [Bacteroides caccae ATCC 43185]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As **Peptide Summary** [Help](#)
 Significance threshold p< Max. number of hits
 Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets
 Show pop-ups Suppress pop-ups Sort unassigned Require bold red
 Error tolerant

1. [gi|53712495](#) Mass: 15517 Score: 403 Queries matched: 7 emPAI: 1.69
 hypothetical protein BF1203 [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 132	667.2930	1332.5714	1332.6959	-0.1244	0	49	0.35	1	K.NMLIVETIDAAK.K + Oxidation (M)
<input checked="" type="checkbox"/> 146	688.8010	1375.5874	1375.8075	-0.2200	0	40	4.6	1	K.TILSISGKPGLYK.L
<input checked="" type="checkbox"/> 159	733.7710	1465.5274	1465.7089	-0.1814	0	56	0.054	1	R.EYLAEVLPDFDR.D
<input checked="" type="checkbox"/> 186	579.8860	1736.6362	1736.8369	-0.2008	1	37	3.8	2	R.EYLAEVLPDFDRDR.V
<input checked="" type="checkbox"/> 224	712.9460	2135.8162	2136.0772	-0.2611	0	(60)	0.017	1	K.IISLADIAMTYNDSEVPLR.D + Oxidation (M)
<input checked="" type="checkbox"/> 225	1068.9160	2135.8174	2136.0772	-0.2598	0	102	1.1e-06	1	K.IISLADIAMTYNDSEVPLR.D + Oxidation (M)
<input checked="" type="checkbox"/> 248	874.0220	2619.0442	2619.3578	-0.3136	1	119	2e-08	1	K.IISLADIAMTYNDSEVPLRDVLR.S + Oxidation (M)

2. [gi|255007972](#) Mass: 15498 Score: 352 Queries matched: 6 emPAI: 1.69
 hypothetical protein Bfra3_02458 [Bacteroides fragilis 3_1_12]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 146	688.8010	1375.5874	1375.8075	-0.2200	0	40	4.6	1	K.TILSISGKPGLYK.L
<input type="checkbox"/> 159	733.7710	1465.5274	1464.7249	0.8026	0	56	0.055	2	R.QYLAEVLPDFDR.D
<input checked="" type="checkbox"/> 186	579.8860	1736.6362	1735.8529	0.7832	1	37	3.7	1	R.QYLAEVLPDFDRDR.V
<input type="checkbox"/> 224	712.9460	2135.8162	2136.0772	-0.2611	0	(60)	0.017	1	K.IISLADIAMTYNDSEVPLR.D + Oxidation (M)
<input type="checkbox"/> 225	1068.9160	2135.8174	2136.0772	-0.2598	0	102	1.1e-06	1	K.IISLADIAMTYNDSEVPLR.D + Oxidation (M)
<input type="checkbox"/> 248	874.0220	2619.0442	2619.3578	-0.3136	1	119	2e-08	1	K.IISLADIAMTYNDSEVPLRDVLR.S + Oxidation (M)

3. [gi|153807766](#) Mass: 16006 Score: 77 Queries matched: 2 emPAI: 0.21
 hypothetical protein BACCAC_02049 [Bacteroides caccae ATCC 43185]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 146	688.8010	1375.5874	1375.8075	-0.2200	0	40	4.6	1	K.TILSISGKPGLYK.L
<input type="checkbox"/> 186	579.8860	1736.6362	1735.8529	0.7833	1	37	4.2	3	R.EYLAEVLPDFDRDR.V

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 150	695.1280	1388.2414	1388.6969	-0.4555	1	35	6.5	1	MKANLTLGDGPAK + Oxidation (M)
<input type="checkbox"/> 227	713.9140	2138.7202	2139.0194	-0.2992	0	34	6.4	1	MYQELSDPPVIFLNNSK + Oxidation (M)

MASCOT Mascot Search Results

Protein View

Match to: [gi|53712495](#) Score: 403
hypothetical protein BP1203 [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (5205) 10_RJ7_01_1371.d\SSP (5205) 10_RJ7_01_1371.mgf

Nominal mass (M_n): 15517; Calculated pI value: 5.15
 NCBI BLAST search of [gi|53712495](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60680695](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253563474](#) from [Bacteroides sp. 3_2_5](#)
[gi|265762658](#) from [Bacteroides sp. 2_1_16](#)
[gi|52215360](#) from [Bacteroides fragilis YCH46](#)
[gi|60492129](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251947250](#) from [Bacteroides sp. 3_2_5](#)
[gi|263255266](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 44%

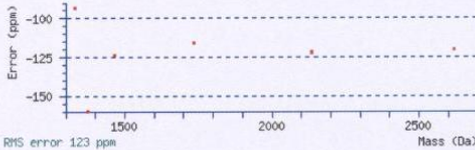
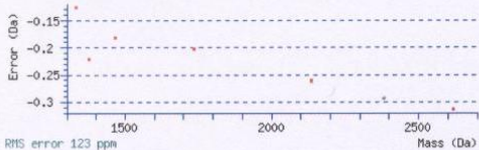
Matched peptides shown in **Bold Red**

1 MLK**TILSISG** K**PGLYKLISQ** GKNMLIV**ETI** DAAK**RFPAY** GNEKI**ISLAD**
51 IAMY**TNDSEV** PL**RDVLR**SIK EK**ENAAIASI** DV**KKATSEQL** REY**LAEVLDP**
101 **PDRDR**VY**TND** IK**KLILWYNI** LV**SNGITDFG** E**TAVEAE**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
4 - 16	688.8010	1375.5874	1375.8075	-0.2200	0	K.TILSISGK PGLYK .L (Ions score 40)
23 - 34	667.2930	1332.5714	1332.6959	-0.1244	0	K.NMLIVETIDAAK.K Oxidation (M) (Ions score 49)
45 - 63	712.9460	2135.8162	2136.0772	-0.2611	0	K.IISLADIAMY TNDSEVPLR .D Oxidation (M) (Ions score 60)
45 - 63	1068.9160	2135.8174	2136.0772	-0.2598	0	K.IISLADIAMY TNDSEVPLR .D Oxidation (M) (Ions score 102)
45 - 67	874.0220	2619.0442	2619.3578	-0.3136	1	K.IISLADIAMY TNDSEVPLRDVLR .S Oxidation (M) (Ions score 119)
92 - 103	733.7710	1465.5274	1465.7089	-0.1814	0	R.EYLA EVLPDFDR .D (Ions score 56)
92 - 105	579.8860	1736.6362	1736.8369	-0.2008	1	R.EYLA EVLPDFDRDR .V (Ions score 37)



LOCUS YP_098487 138 aa linear BCT 26-APR-2009
 DEFINITION hypothetical protein BP1203 [Bacteroides fragilis YCH46].
 ACCESSION YP_098487
 VERSION YP_098487.1 GI:53712495
 DBLINK Project:11067
 DBSOURCE REFSQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 138)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 138)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 138)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD47953](#).
 Method: conceptual translation.
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 Protein 1..138

CDS

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BLASTP E(): 8e-56"  
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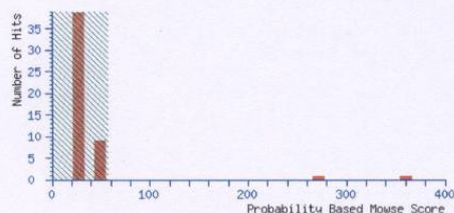
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 5402
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (5402) 10_RF5_01_1076.d\SSP (5402) 10_RF5_01_1076.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:42:08 GMT
 Protein hits : [gi|53712495](#) hypothetical protein BF1203 [Bacteroides fragilis YCH46]
 [gi|255007972](#) hypothetical protein Bfra3_02458 [Bacteroides fragilis 3_1_12]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: Peptide Summary Help

Significance threshold $p < 0.05$ Max. number of hits AUTO

Standard scoring MudPIT scoring Ions score or expect cut-off 0 Show sub-sets 0

Show pop-ups Suppress pop-ups Sort unassigned Decreasing Score Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53712495](#) Mass: 15517 Score: 361 Queries matched: 9 emPAI: 1.20
 hypothetical protein BF1203 [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect	Rank	Peptide
103	565.3110	1128.6074	1129.5979	-0.9904	0	30	K.ENAAIASIDVK.K
<input checked="" type="checkbox"/> 120	659.3070	1316.5994	1316.7010	-0.1015	0	(49)	K.NMLIVETIDAAK.K
<input checked="" type="checkbox"/> 122	667.2980	1332.5814	1332.6959	-0.1144	0	54	K.NMLIVETIDAAK.K + Oxidation (M)
<input checked="" type="checkbox"/> 142	733.8000	1465.5854	1465.7089	-0.1234	0	56	R.EYLAEVLPDFDR.D
<input checked="" type="checkbox"/> 173	579.8940	1736.6602	1736.8369	-0.1768	1	43	R.EYLAEVLPDFDRR.V
<input checked="" type="checkbox"/> 220	712.9900	2135.9482	2136.0772	-0.1291	0	(42)	K.IISLADIAMYTNDSEVPLR.D + Oxidation (M)
<input checked="" type="checkbox"/> 221	1068.9820	2135.9494	2136.0772	-0.1278	0	86	K.IISLADIAMYTNDSEVPLR.D + Oxidation (M)
<input checked="" type="checkbox"/> 222	713.0160	2136.0262	2136.0772	-0.0511	0	(33)	K.IISLADIAMYTNDSEVPLR.D + Oxidation (M)
<input checked="" type="checkbox"/> 247	873.9560	2618.8462	2619.3578	-0.5116	1	93	K.IISLADIAMYTNDSEVPLRDVLR.S + Oxidation (M)

2. [gi|255007972](#) Mass: 15498 Score: 271 Queries matched: 6 emPAI: 0.81
 hypothetical protein Bfra3_02458 [Bacteroides fragilis 3_1_12]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect	Rank	Peptide
142	733.8000	1465.5854	1464.7249	0.8606	0	49	R.QYLAEVLPDFDR.D
173	579.8940	1736.6602	1735.8529	0.8072	1	43	R.QYLAEVLPDFDRR.V
220	712.9900	2135.9482	2136.0772	-0.1291	0	(42)	K.IISLADIAMYTNDSEVPLR.D + Oxidation (M)
221	1068.9820	2135.9494	2136.0772	-0.1278	0	86	K.IISLADIAMYTNDSEVPLR.D + Oxidation (M)
222	713.0160	2136.0262	2136.0772	-0.0511	0	(33)	K.IISLADIAMYTNDSEVPLR.D + Oxidation (M)
247	873.9560	2618.8462	2619.3578	-0.5116	1	93	K.IISLADIAMYTNDSEVPLRDVLR.S + Oxidation (M)

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta Miss Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 59	409.7360	817.4574	817.4545	0.0029	0	34	DVITVSGK
<input checked="" type="checkbox"/> 103	565.3110	1128.6074	1128.6437	-0.0363	1	32	IEAALVRMAR
<input checked="" type="checkbox"/> 176	592.5390	1774.5952	1774.9478	-0.3527	1	32	LYGINASGWTPARLTR
<input checked="" type="checkbox"/> 151	521.1690	1560.4852	1559.9623	0.5228	2	31	VNIIQLAQKHKR
<input checked="" type="checkbox"/> 125	677.4280	2029.2622	2029.2339	0.0282	2	30	KGLTIPLSIYLRKGSWIK
<input checked="" type="checkbox"/> 188	944.1040	2829.2902	2830.2758	-0.9856	1	30	MNTGQWMTNLTESGAGSDLGVMKAK + Oxidation (M)
<input checked="" type="checkbox"/> 126	678.3130	1354.6114	1355.7159	-1.1044	0	29	MLFSLFNILDK + Oxidation (M)
148	763.1340	2286.3802	2286.0872	0.2930	0	28	MQYDALMDESTTLLSLAIGAR

MASCOT Mascot Search Results

Protein View

Match to: gi|53712495 Score: 361
 hypothetical protein BF1203 [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (5402) 10_RF5_01_1076.d\SSP (5402) 10_RF5_01_1076.mgf

Nominal mass (M_r): 15517; Calculated pI value: 5.15
 NCBI BLAST search of gi|53712495 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60680695](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253563474](#) from [Bacteroides sp. 3_2_5](#)
[gi|265762658](#) from [Bacteroides sp. 2_1_16](#)
[gi|52215360](#) from [Bacteroides fragilis YCH46](#)
[gi|60492129](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251947250](#) from [Bacteroides sp. 3_2_5](#)
[gi|263255266](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 43%

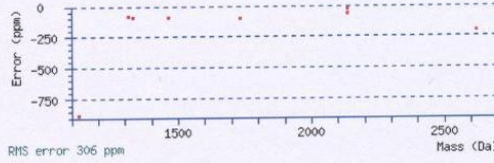
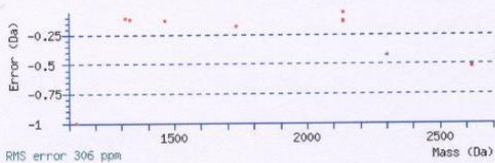
Matched peptides shown in **Bold Red**

1 MLKTIISISG KPGLYKLISQ GKNMLIVETI DAAKRFPAY GNEKIISLAD
51 IAMYTNDSEV PLRDVLRSLK EKENAAIASI DVKKATSEQL REYLAEVLDP
101 FDRDRVYVND IKKLLWYNI LVSNGITDFG EETAVEAE

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss Sequence
23 - 34	659.3070	1316.5994	1316.7010	-0.1015	0 K.NMLIVETIDAAK.K (Ions score 42)
23 - 34	667.2980	1332.5814	1332.6959	-0.1144	0 K.NMLIVETIDAAK.K Oxidation (M) (Ions score 54)
45 - 63	712.9900	2135.9482	2136.0772	-0.1291	0 K.IISLADIAMYTNDSEVPLR.D Oxidation (M) (Ions score 42)
45 - 63	1068.9820	2135.9494	2136.0772	-0.1278	0 K.IISLADIAMYTNDSEVPLR.D Oxidation (M) (Ions score 86)
45 - 63	713.0160	2136.0262	2136.0772	-0.0511	0 K.IISLADIAMYTNDSEVPLR.D Oxidation (M) (Ions score 33)
45 - 67	873.9560	2618.8462	2619.3578	-0.5116	1 K.IISLADIAMYTNDSEVPLRDVLR.S Oxidation (M) (Ions score 93)
73 - 83	565.3110	1128.6074	1129.5979	-0.9904	0 K.ENAAIASIDVK.K (Ions score 30)
92 - 103	733.8000	1465.5854	1465.7089	-0.1234	0 R.EYLAEVLDPFDR.D (Ions score 56)
92 - 105	579.8940	1736.6602	1736.8369	-0.1768	1 R.EYLAEVLDPFDRD.V (Ions score 43)



LOCUS YP_098487 138 aa linear BCT 26-APR-2009
 DEFINITION hypothetical protein BF1203 [Bacteroides fragilis YCH46].
 ACCESSION YP_098487
 VERSION YP_098487.1 GI:53712495
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 138)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 138)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 138)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL [REFSEQ](#): This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD47953](#).
 Method: conceptual translation.
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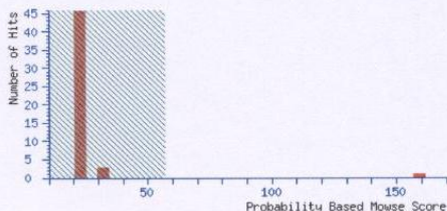
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6102
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6102) 10_RF4_01_1074.d\SSP (6102) 10_RF4_01_1074.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:44:14 GMT
 Protein hits : [gi|53714011](#) putative RNA-binding protein [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As **Peptide Summary** [Help](#)

Significance threshold $p < 0.05$ Max. number of hits **AUTO**

Standard scoring MudPIT scoring Ions score or expect cut-off **0** Show sub-sets **0**

Show pop-ups Suppress pop-ups Sort unassigned **Decreasing Score** Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53714011](#) Mass: 12893 Score: 159 Queries matched: 4 emPAI: 1.03
 putative RNA-binding protein [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 231	813.6560	2437.9462	2438.1060	-0.1598	0	50	0.15	1	K.GFGFVEMPNDDEGNAAIAALNEK.E + Oxidation (M)
<input checked="" type="checkbox"/> 244	885.2950	2652.8632	2653.2330	-0.3698	1	70	0.0013	1	R.SKGGFVEMPNDDEGNAAIAALNEK.E + Oxidation (M)
<input checked="" type="checkbox"/> 250	994.3830	2980.1272	2980.3760	-0.2488	1	39	1.8	1	K.GFGFVEMPNDDEGNAAIAALNEKEIDGK.T + Oxidation (M)
<input checked="" type="checkbox"/> 251	994.8610	2981.5612	2980.3760	1.1852	1	(14)	7.5e+02	7	K.GFGFVEMPNDDEGNAAIAALNEKEIDGK.T + Oxidation (M)

Proteins matching the same set of peptides:
[gi|255009677](#) Mass: 12909 Score: 159 Queries matched: 4
 putative RNA-binding protein [Bacteroides fragilis 3_1_12]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 131	665.0350	1992.0832	1991.9775	0.1057	1	30	43	1	IYKSSFTAMYTAVTTHR + Oxidation (M)
<input checked="" type="checkbox"/> 146	723.3000	2166.8782	2165.9794	0.8988	2	30	52	1	VREVRGGDGGATMDSMDLER + Oxidation (M)
<input checked="" type="checkbox"/> 28	370.9340	1109.7802	1110.6761	-0.8959	0	27	46	1	GILAGLGLLER
<input checked="" type="checkbox"/> 110	609.5880	1825.7422	1824.9734	0.7688	1	26	1e+02	1	FGAGNDSGPPPPQVTLKIK
<input checked="" type="checkbox"/> 201	655.2180	1962.6322	1961.9806	0.6516	1	26	43	1	EATGALNDEVLRNLRYER
<input checked="" type="checkbox"/> 63	385.3540	1153.0402	1153.5074	-0.4672	0	25	98	1	DPSMDALYAR + Oxidation (M)
<input checked="" type="checkbox"/> 62	383.3540	1147.0402	1147.6237	-0.5836	0	25	1.2e+02	1	LVLVGDYVDR
<input checked="" type="checkbox"/> 180	909.1040	2724.2902	2725.4109	-1.1207	1	24	1.4e+02	1	LDPVAKVAMITNLLTEDNLPYSYHR + Oxidation (M)
<input checked="" type="checkbox"/> 130	658.3130	1314.6114	1314.6568	-0.0454	0	24	1.1e+02	1	LISATEPDWQR
<input checked="" type="checkbox"/> 99	563.1690	1686.4852	1686.7631	-0.2780	1	24	1.4e+02	1	TSGSPSSLFSSSARCR
<input checked="" type="checkbox"/> 172	878.1610	2631.4612	2630.3486	1.1126	1	24	1.4e+02	1	IAPDENAGSLHDLRLMHLGSDLILK + Oxidation (M)
<input checked="" type="checkbox"/> 145	715.1790	1428.3434	1427.7732	0.5702	2	24	1.5e+02	1	ERLNVGKAEDAVK
<input checked="" type="checkbox"/> 139	678.9820	2033.9242	2034.0343	-0.1101	1	24	2e+02	1	DIMQELNYELPDLKAVK + Oxidation (M)
<input checked="" type="checkbox"/> 165	837.5110	2509.5112	2508.3635	1.1477	2	24	1.8e+02	1	WMQNSITVPLKLAQVAHSVSR + Oxidation (M)
<input checked="" type="checkbox"/> 113	622.6150	1243.2154	1243.6772	-0.4617	0	24	2e+02	1	TDLAALLAATER
<input checked="" type="checkbox"/> 25	521.4730	1561.3972	1561.8100	-0.4128	0	23	1.8e+02	1	LAGIAGLGSAAFSGSENK
<input checked="" type="checkbox"/> 133	666.6390	1996.8952	1996.8943	0.0009	2	23	2e+02	1	KGSDGMGGSGGSGGKSDKPK + Oxidation (M)
<input checked="" type="checkbox"/> 226	763.6760	2288.0062	2287.0903	0.9158	1	23	97	1	MTSPDRFGGTGVNVIQSGFSVK + Oxidation (M)
<input checked="" type="checkbox"/> 108	603.4090	1807.2052	1806.9224	0.2828	1	23	2.2e+02	1	LDGEAIHGKSPNELATR
<input checked="" type="checkbox"/> 61	377.1540	752.2934	752.3276	-0.0341	0	23	1e+02	1	AMFGGDR
<input checked="" type="checkbox"/> 232	819.3970	2455.1692	2455.1914	-0.0222	1	23	1.1e+02	1	SAPPGFARELAGHDLAAGLSAMSR + Oxidation (M)
<input checked="" type="checkbox"/> 213	1072.7390	3215.1952	3215.7774	-0.5822	1	23	1.5e+02	1	VTLACVGLLIGLVYVLAASPASRIGASR + Oxidation (M)

MASCOT Mascot Search Results

Protein View

Match to: [gi|53714011](#) Score: 159
putative RNA-binding protein [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6102) 10_RF4_01_1074.d\SSP (6102) 10_RF4_01_1074.mgf

Nominal mass (M_r): 12893; Calculated pI value: 5.24
 NCBI BLAST search of [gi|53714011](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60682207](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|265764361](#) from [Bacteroides sp. 2_1_16](#)
[gi|52216876](#) from [Bacteroides fragilis YCH46](#)
[gi|60493641](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|263256969](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 24%

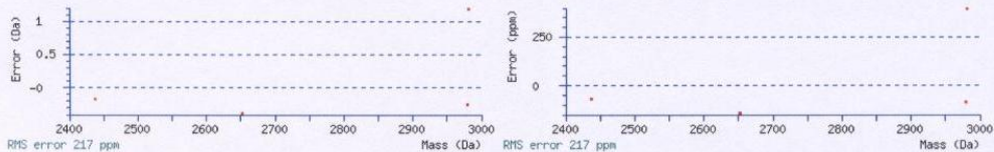
Matched peptides shown in **Bold Red**

1 MNIFIAGISY NLSNADLDEL FEEFGEVISA KIVMDRETGR **SKGPGFVEMP**
51 NDEEGNAALA ALNEKEIDGR TLAVSVARPR EEGPRNSNY GGGNRGGYGN
101 NRGGGYGGGN RGGYGGGRD RY

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
41 - 65	885.2950	2652.8632	2653.2330	-0.3698	1	R.SKGPGFVEMPNDDEEGNAIAALNEK.E Oxidation (M) (Ions score 70)
43 - 65	813.6560	2437.9462	2438.1060	-0.1598	0	K.GPGFVEMPNDDEEGNAIAALNEK.E Oxidation (M) (Ions score 50)
43 - 70	994.3830	2980.1272	2980.3760	-0.2488	1	K.GPGFVEMPNDDEEGNAIAALNEKEIDGK.T Oxidation (M) (Ions score 39)
43 - 70	994.8610	2981.5612	2980.3760	1.1852	1	K.GPGFVEMPNDDEEGNAIAALNEKEIDGK.T Oxidation (M) (Ions score 14)



LOCUS YP_100003 122 aa linear BCT 26-APR-2009
 DEFINITION putative RNA-binding protein [Bacteroides fragilis YCH46].
 ACCESSION YP_100003
 VERSION YP_100003.1 GI:53714011
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 122)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H., Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 122)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 122)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences, 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final NCBI review. The reference sequence was derived from [BAD49469](#).
 Method: conceptual translation.
 FEATURES Location/Qualifiers
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 /strain="YCH46"
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 Protein 1..122
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 /calculated_mol_wt=12770
 Region 2..75
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 /note="RRM (RNA recognition motif), also known as RBD (RNA binding domain) or RNP (ribonucleoprotein domain), is a highly abundant domain in eukaryotes found in proteins involved in post-transcriptional gene expression processes"

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including mRNA and rRNA...; cd00590"
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order(4,44,46)
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Site 74..75
      /site_type="other"
      /note="RRM dimerization site"
      /db_xref="CDD:100104"
CDS 1..122
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     /coded_by="complement(NC_006347.1:3121075..3121443)"
     /note="similar to gpu:BX842649_260 [Bdellovibrio
bacteriovorus], percent identity 54 in 114 aa, BLASTP E():
1e-27"
     /transl_table=11
     /db_xref="GeneID:3083599"
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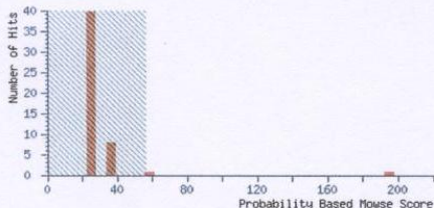
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : LAKSHMY MANICKAN
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6103
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6103) 10_RH24_01_1296.d\SSP (6103) 10_RH24_01_1296.mgf
 Database : MCBInr 20090522 (8876587 sequences; 3036162093 residues)
 Taxonomy : Bacteria (Subbacteria) (4773688 sequences)
 Timestamp : 30 May 2009 at 10:57:36 GMT
 Protein hits : [gi|53714574](#) lactoylglutathione lyase and related protein [Bacteroides fragilis YCH46]
[gi|126496](#) RecName: Full=Lysostaphin; AltName: Full=Glycyl-glycine endopeptidase; Flags: Precursor

Probability Based Mowse Score

Ions score is $-10 * \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 56 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p < 0.05$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53714574](#) Mass: 14857 Score: 195 Queries matched: 3 emPAI: 0.85
 lactoylglutathione lyase and related protein [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 154	498.9330	1493.7772	1493.8354	-0.0582	1	44	1	1	R.KGAEGLNIAFLHPK.S
<input checked="" type="checkbox"/> 172	919.4290	1836.8434	1836.9509	-0.1074	0	62	0.015	1	K.SIEALPYEIVLGLK.C
<input checked="" type="checkbox"/> 241	821.6930	2462.0572	2462.2190	-0.1618	0	90	1.7e-05	1	K.GAGVHHVAFIADGVANALAEAEK.E

2. [gi|126496](#) Mass: 51656 Score: 64 Queries matched: 2 emPAI: 0.06
 RecName: Full=Lysostaphin; AltName: Full=Glycyl-glycine endopeptidase; Flags: Precursor
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 161	819.8350	1637.6554	1637.7897	-0.1342	0	37	3.6	1	K.SESASFTPTDIITR.T
<input checked="" type="checkbox"/> 209	1023.0460	2044.0774	2044.9234	-0.8460	0	27	65	1	R.MVNSFNSSTAQDPMFPLK.S + 2 Oxidation (M)

Proteins matching the same set of peptides:
[gi|153047](#) Mass: 42213 Score: 64 Queries matched: 2
 lysostaphin (ttg start codon)
[gi|3287967](#) Mass: 53058 Score: 64 Queries matched: 2
 RecName: Full=Lysostaphin; AltName: Full=Glycyl-glycine endopeptidase; Flags: Precursor

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 164	850.9000	1699.7854	1699.8814	-0.0960	0	42	1.5	1	ANVAEIMNIIGIEGK
<input checked="" type="checkbox"/> 167	858.8630	1715.7114	1715.8777	-0.1662	2	41	1.8	1	ELLNKNHYGMDRVK
<input checked="" type="checkbox"/> 175	616.5650	1846.6732	1846.8479	-0.1747	2	30	16	1	MKQEAENAEADKQER
<input checked="" type="checkbox"/> 108	374.4570	1120.3492	1121.5135	-1.1643	0	29	22	1	ASVGMHAAETR + Oxidation (M)
<input checked="" type="checkbox"/> 205	679.2120	2034.6142	2034.0204	0.5938	1	29	17	1	LIAEEDAQRVIVPCNTR
<input checked="" type="checkbox"/> 184	645.0580	1932.1522	1930.9611	1.1911	0	28	29	1	MWITNGPHADTLVVYAK + Oxidation (M)
<input checked="" type="checkbox"/> 93	465.3710	1393.0912	1392.6496	0.4416	1	28	72	1	NEEKYHWIHK + Oxidation (M)
<input checked="" type="checkbox"/> 84	394.2260	1179.6562	1179.4835	0.1727	0	28	1e+02	1	LCMMSGHVR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 197	667.5230	1999.5472	1999.9018	-0.3546	1	27	26	1	MAVDGTDHDSPTLDQARR + Oxidation (M)
<input checked="" type="checkbox"/> 121	636.6970	1907.0692	1907.9476	-0.8785	0	27	81	1	EVTLSFSEAIAGVDDISIR
<input checked="" type="checkbox"/> 213	694.0640	2079.1702	2079.9578	-0.7876	1	26	47	1	NOGMEQRYANEIVGANMR
<input checked="" type="checkbox"/> 234	767.3330	2298.9772	2300.1185	-1.1413	1	26	42	1	RYEHYNTIEHIDLENK
<input checked="" type="checkbox"/> 248	930.1100	2787.3082	2787.2996	0.0085	1	26	43	1	NMNLAYADVDRMAGPEVPSGSGFK + Oxidation (M)

Mascot Search Results

Protein View

Match to: [gi|53714574](#) Score: 195
lactoylglutathione lyase and related protein [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6103) 10_RH24_01_1296.d\SSP (6103) 10_RH24_01_1296.mgf

Nominal mass (M_r): 14857; Calculated pI value: 5.50
 NCBI BLAST search of [gi|53714574](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60682596](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253565078](#) from [Bacteroides sp. 3_2_5](#)
[gi|265765962](#) from [Bacteroides sp. 2_1_16](#)
[gi|52217439](#) from [Bacteroides fragilis YCH46](#)
[gi|60494030](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251946543](#) from [Bacteroides sp. 3_2_5](#)
[gi|263253630](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 41%

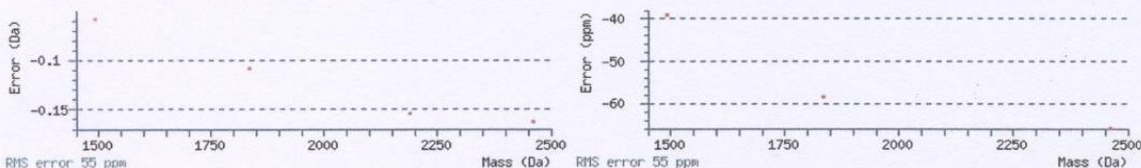
Matched peptides shown in **Bold Red**

1 MKISHIEHLG IAVK**SIEEAL** **PYYENVLGLK** CYNIVETVEDQ KVRTAPLKVQ
 51 DTKIELLEPT CPESTIAKPI ENKGAGVHHV **AF**AIEDGVAN ALAAEAEKSI
 101 RLIDKAPRKG **AEGLNIAFLH** PKSTLGLVLTE LCEH

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Sequence
15 - 30	919.4290	1836.8434	1836.9509	-0.1074	0	K.SIEEALPYYENVLGLK.C (Ions score 62)
74 - 98	821.6930	2462.0572	2462.2190	-0.1618	0	K.GAGVHHVAF AIEDGVANALAAEAEK.E (Ions score 90)
109 - 122	498.9330	1493.7772	1493.8354	-0.0582	1	R.KGAEGLNIAFLHPK.S (Ions score 44)



LOCUS YP_100566 134 aa linear BCT 26-APR-2009
 DEFINITION lactoylglutathione lyase and related protein [Bacteroides fragilis YCH46].
 ACCESSION YP_100566
 VERSION YP_100566.1 GI:53714574
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 134)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H., Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 134)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 134)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences, 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final NCBI review. The reference sequence was derived from [BAD50032](#). Method: conceptual translation.
 FEATURES Location/Qualifiers

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superfamily; cl00411"
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CDS 1..134
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/note="similar to gp:AE016932_217 [Bacteroides
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BLASTP E(): 5e-71"
/transl_table=11
/db_xref="GeneID:3082133"

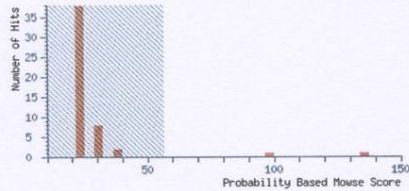
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : LAKSHMY MANICKAN
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6206
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6206) 10_RJ13_01_1386.d\SSP (6206) 10_RJ13_01_1386.mgf
 Database : NCBI nr 20090522 (8876587 sequences; 3036162093 residues)
 Taxonomy : Bacteria (Eubacteria) (4773688 sequences)
 Timestamp : 30 May 2009 at 10:59:43 GMT
 Protein hits : [gi|53711592](#) hypothetical protein BF0301 [Bacteroides fragilis YCH46]
 [gi|143945](#) Fe-superoxide dismutase [Bacteroides fragilis]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 56 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As Help

Significance threshold $p < 0.05$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53711592](#) Mass: 16013 Score: 136 Queries matched: 2 emPAI: 0.47
 hypothetical protein BF0301 [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 202	1021.4130	2040.8114	2041.0691	-0.2577	1	26	43	1	K.TAPGANDLTDADALKIVQK.L
<input checked="" type="checkbox"/> 234	1139.9630	2277.9114	2277.1198	0.7916	0	112	1.8e-07	1	R.QDLADAEALQVQVMETYLEFK.Q + Oxidation (M)

2. [gi|143945](#) Mass: 21736 Score: 100 Queries matched: 3 emPAI: 0.15
 Fe-superoxide dismutase [Bacteroides fragilis]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 239	792.5960	2374.7662	2375.1757	-0.4095	0	27	25	1	K.HLQTIYVNNLNSLVPGETEYEGK.T
<input checked="" type="checkbox"/> 245	878.6220	2632.8442	2633.3125	-0.4684	0	(25)	35	1	K.LPYANNALEPVISQQTIDYHYGK.H
<input checked="" type="checkbox"/> 246	878.6530	2632.9372	2633.3125	-0.3754	0	73	0.00055	1	K.LPYANNALEPVISQQTIDYHYGK.H

Proteins matching the same set of peptides:
[gi|533307](#) Mass: 21715 Score: 100 Queries matched: 3
 superoxide dismutase [Bacteroides fragilis]
[gi|53713818](#) Mass: 23124 Score: 100 Queries matched: 3
 superoxide dismutase [Bacteroides fragilis YCH46]
[gi|60682036](#) Mass: 21758 Score: 100 Queries matched: 3
 superoxide dismutase [Fe] [Bacteroides fragilis NCTC 9343]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
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<input checked="" type="checkbox"/> 143	778.5110	2332.5112	2333.1798	-0.6686	1	35	11	1	MIDNIQNVTDQLRTPFER
<input checked="" type="checkbox"/> 235	767.2700	2298.7882	2299.2859	-0.4978	2	31	9.5	1	SGTIKVIQGNLIRNVTITQEAQ
<input checked="" type="checkbox"/> 153	549.2230	1644.6472	1643.8519	0.7953	0	27	39	1	AFVNFDTVAEVQVAK
<input checked="" type="checkbox"/> 209	694.9150	2081.7232	2080.9338	-0.7894	0	27	29	1	TGTGESEEAEEAVGPMQVR
<input checked="" type="checkbox"/> 135	742.1990	1482.3834	1481.6425	-0.7410	0	26	36	1	THMGLQNCINQR + Oxidation (M)
<input checked="" type="checkbox"/> 60	396.1590	1185.4552	1184.5509	-0.9043	1	26	1.2e+02	1	ANWEGHGKMR
<input checked="" type="checkbox"/> 138	757.2680	2268.7822	2268.2114	-0.5708	0	26	83	1	KPHLIAILVGNDDGGSETTVASK
<input checked="" type="checkbox"/> 118	666.3070	1995.8992	1994.9863	-0.9128	2	25	1.3e+02	1	AGPGEAAARRPEMPAPR
<input checked="" type="checkbox"/> 184	940.8210	2819.4412	2820.4004	-0.9592	0	25	93	1	DVIALAASVALSHNTYDAAVFLGVCDK

Mascot Search Results

Protein View

Match to: gi|53711592 Score: 136
 hypothetical protein BF0301 [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6206) 10_RJ13_01_1386.d\SSP (6206) 10_RJ13_01_1386.mgf

Nominal mass (M_r): 16013; Calculated pI value: 5.67
 NCBI BLAST search of gi|53711592 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60679842](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253564357](#) from [Bacteroides sp. 3_2_5](#)
[gi|265764976](#) from [Bacteroides sp. 2_1_16](#)
[gi|52214457](#) from [Bacteroides fragilis YCH46](#)
[gi|60491276](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251948133](#) from [Bacteroides sp. 3_2_5](#)
[gi|263254360](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 26%

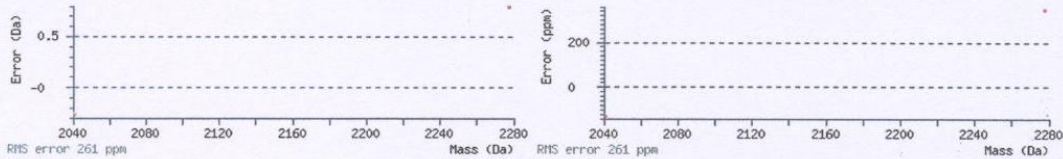
Matched peptides shown in **Bold Red**

1 MDLPERVSED IKNAMKAKDK VALETLRNVK KFFLEAKTAP GANDTLTDAD
 51 **ALKIVQK**LVK QKDAAEIYI GQGRQDLADA ELAQVQVMET YLPKQMSAEE
 101 LEAALKEIIA EVGATSGKDM GKVMGVASK LAGLAEGRAI SAKVKELLG

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss Sequence
38 - 57	1021.4130	2040.8114	2041.0691	-0.2577	1 K.TAPGANDTLTDADALKIVQK.L (Ions score 26)
75 - 94	1139.9630	2277.9114	2277.1198	0.7916	0 R.ODLADAEALAQVQVMETYLPK.Q Oxidation (M) (Ions score 112)



LOCUS YP_097584 149 aa linear BCT 26-APR-2009
 DEFINITION hypothetical protein BF0301 [Bacteroides fragilis YCH46].
 ACCESSION YP_097584
 VERSION YP_097584.1 GI:53711592
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 149)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 149)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 149)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagamiara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD47050](#).
 Method: conceptual translation.
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 /strain="YCH46"
 /db_xref="taxon:295405"
 Protein 1..149
 /product="hypothetical protein"
 /calculated_mol_wt=15891

Region 5..148
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/note="GatB domain; cl11497"
/db_xref="CDD:142702"
CDS 1..149
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BLASTP E(): 1e-62"
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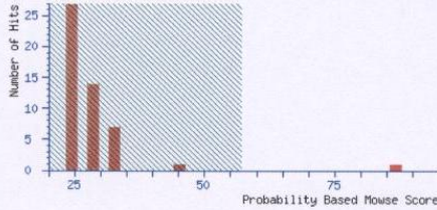
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6301
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP 6301 (1000) RN13_01_1592.d\SSP 6301 (1000) RN13_01_1592.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:46:45 GMT
 Protein hits : [gi|53712002](#) thioredoxin [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: Peptide Summary Help

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53712002](#) Mass: 11637 Score: 87 Queries matched: 2 emPAI: 0.30
 thioredoxin [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 125	539.7950	1077.5754	1077.6223	-0.0468	0	16	7.2e+02	3	R.NIPTVLFFK.N
<input checked="" type="checkbox"/> 184	918.8160	1835.6174	1836.7836	-1.1662	0	70	0.0016	1	K.CDVENSDDLPAEFGIR.N

Proteins matching the same set of peptides:
[gi|255007522](#) Mass: 11635 Score: 85 Queries matched: 2
 putative thioredoxin [Bacteroides fragilis 3_1_12]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 129	582.2650	1162.5154	1162.6598	-0.1443	0	47	0.6	1	LVNEITFSK
<input checked="" type="checkbox"/> 205	652.9580	1955.8522	1956.9323	-1.0802	0	35	8.2	1	MTDSAGIPGGAGAAGGSSKPPR + Oxidation (M)
<input checked="" type="checkbox"/> 126	552.0970	1653.2692	1652.8780	0.3911	2	34	15	1	TSRMLGVSRTIYR
<input checked="" type="checkbox"/> 83	387.2600	1158.7582	1158.5265	0.2317	1	31	52	1	NSDEKSTHMK
<input checked="" type="checkbox"/> 114	489.3880	1465.1422	1465.7096	-0.5674	1	29	56	1	GMAARASGQTFDVR
<input checked="" type="checkbox"/> 189	934.2050	2799.5932	2799.3940	0.1992	2	28	60	1	DAEALGHRLLQQWGAQKADIYSSDLK
<input checked="" type="checkbox"/> 134	626.2680	1875.7822	1876.0128	-0.2306	1	27	89	1	VKINPIQTMTFEQAK + Oxidation (M)
<input checked="" type="checkbox"/> 73	366.2200	1095.6382	1094.6448	0.9934	0	27	1.2e+02	1	IILASHGGLSK
<input checked="" type="checkbox"/> 137	638.7210	1913.1412	1913.0694	0.0718	1	27	92	1	ALTTQPSLINALRSTTAR
<input checked="" type="checkbox"/> 142	652.9630	1303.9114	1303.7347	0.1767	1	26	57	1	TIKQSLSIDTAK
<input checked="" type="checkbox"/> 241	1133.4750	3397.4032	3397.6937	-0.2906	1	26	76	1	IEYIEEPYVRASIMVNDYVGVAVMEIAQAK
<input checked="" type="checkbox"/> 124	539.2530	1614.7372	1614.9457	-0.2085	2	26	1.5e+02	1	GLYNALKIPKITER
<input checked="" type="checkbox"/> 69	354.1940	1059.5602	1060.6063	-1.0461	2	26	1.8e+02	1	AKKNGMIVGK + Oxidation (M)
<input checked="" type="checkbox"/> 160	728.9160	2183.7262	2184.2379	-0.5117	2	25	1.3e+02	1	NDGTLRNSALVTKFLLPVAR
<input checked="" type="checkbox"/> 164	747.5930	2239.7572	2239.8928	-0.1356	0	25	1.2e+02	1	MAQMFGMTISELCMDTDSR + Oxidation (M)
<input checked="" type="checkbox"/> 203	651.2730	1950.7972	1950.0898	0.7074	0	25	73	1	DLARPLGLSVEEAAAIVR
<input checked="" type="checkbox"/> 140	645.2910	1932.8512	1933.1104	-0.2592	2	24	2e+02	1	MKLVIVTGMMSGAGKTIALK + Oxidation (M)
<input checked="" type="checkbox"/> 198	644.3390	1929.9952	1928.9774	1.0178	1	24	95	1	LPTIRLCMMSVVPVEIG + Oxidation (M)
<input checked="" type="checkbox"/> 202	648.5640	1942.6702	1941.9156	0.7546	2	24	72	1	FGPDMGHLGKRVVAHPE + Oxidation (M)
<input checked="" type="checkbox"/> 230	1046.2260	3135.6562	3136.5995	-0.9433	2	24	1.5e+02	1	IHPKNFDLVVYAAFHAFPGPNEERGVPFR
<input checked="" type="checkbox"/> 210	661.0380	1980.0922	1981.0228	-0.9307	1	24	1.1e+02	1	HGASTEDLRAAVTAALAGIAS
<input checked="" type="checkbox"/> 170	510.5390	1528.5952	1527.8919	0.7033	2	24	1.1e+02	1	NAKVAAAVLRGMAK + Oxidation (M)
<input checked="" type="checkbox"/> 246	834.3670	2500.0792	2499.2614	0.8178	2	23	86	1	REEMPDMERLSGAGIPVAVPAFK
<input type="checkbox"/> 148	676.5610	2026.6612	2027.0067	-0.3456	1	23	1.6e+02	1	THDMPIIMVTAKDAEIDK

MASCOT Mascot Search Results

Protein View

Match to: **gi|53712002** Score: **87**
thioredoxin [Bacteroides fragilis YCH46]
Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP 6301 (1000)_RN13_01_1592.d\SSP 6301 (1000)_RN13_01_1592.mgf

Nominal mass (M_r): **11637**; Calculated pI value: **4.51**
NCBI BLAST search of **gi|53712002** against nr
Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60680203](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|253563963](#) from [Bacteroides sp. 3_2_5](#)
[gi|265765340](#) from [Bacteroides sp. 2_1_16](#)
[gi|52214867](#) from [Bacteroides fragilis YCH46](#)
[gi|60491637](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|251947739](#) from [Bacteroides sp. 3_2_5](#)
[gi|263254724](#) from [Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
Variable modifications: Oxidation (M)
Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
Sequence Coverage: **24%**

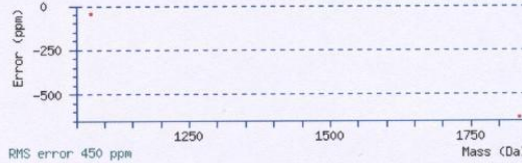
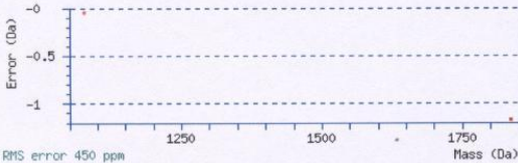
Matched peptides shown in **Bold Red**

1 MALEITDNNF KEILAEQSPV VIDFWAPWCG PCKMVGPIID ELAKEYEGKV
51 IMGK**CDVDEN** **SDLPAEFGIR** NIPTVLFFKN GELVDKQVGA VGKPAFVEKV
101 EKLL

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss Sequence
55 - 70	918.8160	1835.6174	1836.7836	-1.1662	0 K.CDVDENS DLPAEFGIR.N (Ions score 70)
71 - 79	539.7950	1077.5754	1077.6223	-0.0468	0 R.NIPTVL FFK.N (Ions score 16)



LOCUS YP_097994 104 aa linear BCT 26-APR-2009
 DEFINITION thioredoxin [Bacteroides fragilis YCH46].
 ACCESSION YP_097994
 VERSION YP_097994.1 GI:53712002
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 104)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 104)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 104)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD47460](#).
 Method: conceptual translation.
 FEATURES Location/Qualifiers
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 /strain="YCH46"
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Protein 1..104
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/calculated_mol_wt=11339

Region 8..101
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CDS 1..104
/locus_tag="BF0713"
/coded_by="NC_006347.1:838887..839201"
/note="similar to gp:AE016935_76 [Bacteroides thetaiotaomicron VPI-5482], percent identity 81 in 104 aa, BLASTP E(): 6e-47"
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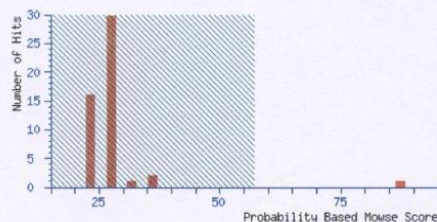
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6302
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6302) 100_RE17_01_1733.d\SSP (6302) 100_RE17_01_1733.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 16:30:06 GMT
 Protein hits : [gi|53715240](#) triosephosphate isomerase [Bacteroides fragilis YCH46]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|53715240](#) Mass: 26867 Score: 87 Queries matched: 2 emPAI: 0.12
 triosephosphate isomerase [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
197	547.2600	1638.7582	1638.8478	-0.0896	0	21	2.2e+02	4	K.TASPAQAQEIHAPIR.S
<input checked="" type="checkbox"/> 209	928.9290	1855.8434	1855.9680	-0.1245	0	67	0.0056	1	K.ELFANPDVGGILGGAAALK.V

Proteins matching the same set of peptides:
[gi|253566372](#) Mass: 28775 Score: 87 Queries matched: 2
 triosephosphate isomerase [Bacteroides sp. 3_2_5]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 211	623.0900	1866.2482	1865.9781	0.2700	1	35	5.4	1	HNKGIIALSACLAGDVAR
<input checked="" type="checkbox"/> 229	687.7600	2060.2582	2060.9875	-0.7293	2	29		1	ENNPNSSEGFITLKDGR
<input checked="" type="checkbox"/> 160	654.3460	1960.0162	1960.9524	-0.9362	0	27	1.1e+02	1	AIMTSDLGINPTNDGSAIR + Oxidation (M)
<input checked="" type="checkbox"/> 112	438.9630	1313.8672	1312.6888	1.1784	2	27	1.1e+02	1	NRVVFETKTR
<input checked="" type="checkbox"/> 182	741.7640	2222.2702	2223.2488	-0.9786	2	26		95	LGQKQGHVLTATGRALFDAK
<input checked="" type="checkbox"/> 251	920.2320	2757.6742	2756.5420	1.1322	2	26		34	LPLVISLSGIMLLSIGMWMWRRK
<input checked="" type="checkbox"/> 197	547.2600	1638.7582	1639.8603	-1.1021	1	26		63	DQKMAPPVVSAEVLK
<input checked="" type="checkbox"/> 176	713.3310	2136.9712	2137.0296	-0.0584	1	26	1.3e+02	1	MRYELVDEAGRPMLDQAK + Oxidation (M)
<input checked="" type="checkbox"/> 242	763.6450	2287.9132	2288.1908	-0.2777	1	26		46	MFLNPMPINSLPTLLGEGGK + 2 Oxidation (M)
<input checked="" type="checkbox"/> 95	365.9130	1094.7172	1095.5084	-0.7912	0	26	1.5e+02	1	AEDLAYDATK
<input checked="" type="checkbox"/> 237	718.2730	2151.7972	2151.1647	0.6324	1	26		47	AGADIAIERALAEGTSPAVAR
<input checked="" type="checkbox"/> 218	644.7310	1931.1712	1931.9741	-0.8029	2	25		67	FKSGYTLGFNRALDSEK
<input checked="" type="checkbox"/> 249	878.6190	2632.8352	2633.4615	-0.6263	2	25		47	LLFVGDVVVSLGEMVAQVVPKLG + Oxidation (M)
<input checked="" type="checkbox"/> 148	618.6470	1852.9192	1853.8982	-0.9790	1	24	1.7e+02	1	EPLMSAIDAFKAPDQR + Oxidation (M)
<input checked="" type="checkbox"/> 206	607.2630	1818.7672	1818.9588	-0.1916	2	24		94	RDGGAARKGIGYGELISGK
<input checked="" type="checkbox"/> 189	759.3340	2274.9802	2274.9789	0.0013	1	24	1.9e+02	1	MAQGGMPGMPGMPGMPGGPGROK + 2 Oxidation (M)
<input checked="" type="checkbox"/> 143	561.1460	1680.4162	1681.0515	-0.6353	2	24	1.6e+02	1	HLLLKHNIIRTLPEK
<input checked="" type="checkbox"/> 139	526.3320	1575.9742	1576.9413	-0.9671	1	24	2.3e+02	1	ASVILLHGLGSRNIK
<input checked="" type="checkbox"/> 142	556.5640	1111.1134	1111.5583	-0.4449	0	24	1.8e+02	1	SYPAMTVSLK + Oxidation (M)
<input checked="" type="checkbox"/> 234	702.0430	2103.1072	2103.0895	0.0177	2	24	1.1e+02	1	LDMRLFQAAARGGPGDVK + Oxidation (M)
<input checked="" type="checkbox"/> 225	677.3390	2028.9952	2027.9595	1.0357	2	23	1.1e+02	1	APPFNSQRMTENLNR + Oxidation (M)
<input checked="" type="checkbox"/> 119	477.1930	1428.5572	1427.7633	0.7939	2	23	2.4e+02	1	RNKDGLINQGWK
<input checked="" type="checkbox"/> 220	662.8710	1985.5912	1986.0316	-0.4404	2	23		87	RIDLMLLEAGALDEARAAR + Oxidation (M)

MASCOT Mascot Search Results

Protein View

Match to: [gi|53715240](#) Score: 87
triosephosphate isomerase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6302) 100_RE17_01_1733.d\SSP (6302) 100_RE17_01_1733.mgf

Nominal mass (M_n): 26867; Calculated pI value: 5.14
 NCBI BLAST search of [gi|53715240](#) against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60683174](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|81313773](#) from [Bacteroides fragilis NCTC 9343](#)
[gi|81824915](#) from [Bacteroides fragilis](#)
[gi|52218105](#) from [Bacteroides fragilis YCH46](#)
[gi|60494608](#) from [Bacteroides fragilis NCTC 9343](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 13%

Matched peptides shown in **Bold Red**

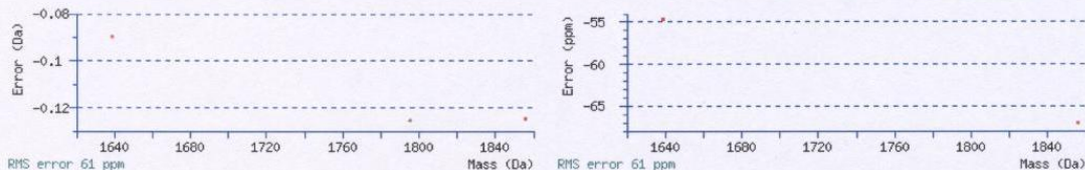
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1 MRKNIVAGNW KMNKTLQEGI ALAKELNEAL ANEKPNCDDVI ICTPFIHLAS
51 VTPLVDAAKI GVGAENCADK ESGAYTGEVS AAMVASTGAK YVILGHSERR
101 AYYGETVEIL KDKVKLALAN GLTPIFCIGE VLEEREANKQ NEVVAAQLAS
151 VFDLSAEDFS KIVLAYEPVW AIGTGKTASP AQAQEIHAFI RSAVAEKYKG
201 EIADNTSILY GGSCKPSNAK ELFANPDVDG GLIGGAALKV ADFKGIIDAF
251 N
    
```

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
177 - 191	547.2600	1638.7582	1638.8478	-0.0896	0 K.TASPAQAEIHAFI.R.S (Ions score 21)
221 - 239	928.9290	1855.8434	1855.9680	-0.1245	0 K.ELFANPDVDGGLIGGAALK.V (Ions score 67)



LOCUS YP_101232 251 aa linear BCT 26-APR-2009
 DEFINITION triosephosphate isomerase [Bacteroides fragilis YCH46].
 ACCESSION YP_101232
 VERSION YP_101232.1 GI:53715240
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE [Bacteroides fragilis YCH46](#)
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 251)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 251)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 251)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagamihara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD50698](#).
 Method: conceptual translation.
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Protein 1..251
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Region 5..248
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D-glyceraldehyde-3-phosphate. The reaction is very
efficient and requires neither cofactors nor metal ions.
TIM, usually...; cd00311"
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Site order(9,11,96,167,173,213,232,234..235)
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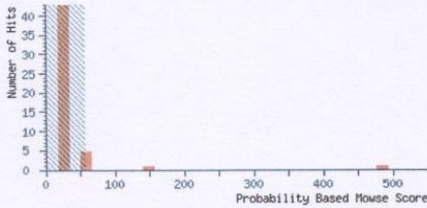
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : LAKSHMY MANICKAN
 Email : lakshmy.manickan@unn.ac.uk
 Search title : 6304 bf
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6302) 10_RJ3_01_1363.d\SSP (6302) 10_RJ3_01_1363.mgf
 Database : NCBI nr 20090528 (8915381 sequences; 3049521622 residues)
 Taxonomy : Bacteria (Eubacteria) (4801653 sequences)
 Timestamp : 6 Jun 2009 at 11:57:12 GMT
 Protein hits : [gi|53714055](#) putative thiol peroxidase [Bacteroides fragilis YCH46]
 [gi|160891883](#) hypothetical protein BACUNI_04341 [Bacteroides uniformis ATCC 8492]
 [gi|167762340](#) hypothetical protein BACSTE_00694 [Bacteroides stercoris ATCC 43183]
 [gi|218130132](#) hypothetical protein BACEGG_01719 [Bacteroides eggerthii DSM 20697]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event.
 Individual ions scores > 56 indicate identity or extensive homology ($p < 0.05$).
 Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p < 0.05$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1. [gi|53714055](#) Mass: 18042 Score: 484 Queries matched: 18 emPAI: 2.93
 putative thiol peroxidase [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 104	552.2190	1102.4234	1102.6386	-0.2152	0	76	0.00068	1	K.LIGEFIQVQK.V
<input checked="" type="checkbox"/> 117	600.7380	1199.4614	1199.6332	-0.1718	0	68	0.0038	1	R.MADGFLAGLLAR.A + Oxidation (M)
<input checked="" type="checkbox"/> 118	601.3060	1200.5974	1199.6332	0.9642	0	(11)	2.4e+03	3	R.MADGFLAGLLAR.A + Oxidation (M)
<input checked="" type="checkbox"/> 128	661.1670	1320.3194	1320.5623	-0.2428	0	(16)	7.8e+02	3	R.FSDFDESYGVR.M
<input checked="" type="checkbox"/> 129	661.1670	1320.3194	1320.5623	-0.2428	0	(67)	0.0031	1	R.FSDFDESYGVR.M
<input checked="" type="checkbox"/> 130	661.1940	1320.3734	1320.5623	-0.1888	0	69	0.0023	1	R.FSDFDESYGVR.M
<input checked="" type="checkbox"/> 131	661.2080	1320.4014	1320.5623	-0.1608	0	(23)	86	1	R.FSDFDESYGVR.M
<input checked="" type="checkbox"/> 132	661.2180	1320.4214	1320.5623	-0.1408	0	(57)	0.032	1	R.FSDFDESYGVR.M
<input checked="" type="checkbox"/> 133	661.2260	1320.4374	1320.5623	-0.1248	0	(58)	0.03	1	R.FSDFDESYGVR.M
<input checked="" type="checkbox"/> 184	610.2910	1827.8512	1829.0047	-1.1535	1	50	0.24	1	K.DTVVLAISKDLPPAQR.F
<input checked="" type="checkbox"/> 207	999.3810	1996.7474	1997.9404	-1.1930	0	(34)	5.5	1	R.FCTTEGIENVILSDFR.F
<input checked="" type="checkbox"/> 209	999.3960	1996.7654	1997.9404	-1.1750	0	(40)	1.6	1	R.FCTTEGIENVILSDFR.F
<input checked="" type="checkbox"/> 210	999.4050	1996.7954	1997.9404	-1.1450	0	42	0.96	1	R.FCTTEGIENVILSDFR.F
<input checked="" type="checkbox"/> 226	708.5960	2122.7662	2123.0310	-0.2648	0	37	2.7	1	K.VAYTELVEITQEPDYK.A
<input checked="" type="checkbox"/> 232	1088.4760	2174.9374	2176.1198	-1.1824	0	(24)	79	3	K.NIVLNIFPSLDTGVCATSVR.K
<input checked="" type="checkbox"/> 233	725.9900	2174.9482	2176.1198	-1.1716	0	48	0.29	1	K.NIVLNIFPSLDTGVCATSVR.K
<input checked="" type="checkbox"/> 244	808.6340	2422.8802	2423.1744	-0.2942	1	43	0.73	1	K.DGKVAYTELVEITQEPDYK.A
<input checked="" type="checkbox"/> 245	817.6610	2449.9612	2450.2661	-0.3050	2	54	0.055	1	K.MAAGHKDTVVLAIKDLPPAQR.F + 2 Oxidation (M)

2. [gi|160891883](#) Mass: 18030 Score: 150 Queries matched: 7 emPAI: 0.41
 hypothetical protein BACUNI_04341 [Bacteroides uniformis ATCC 8492]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 117	600.7380	1199.4614	1199.6332	-0.1718	0	68	0.0038	1	R.MADGFLAGLLAR.S + Oxidation (M)
<input checked="" type="checkbox"/> 118	601.3060	1200.5974	1199.6332	0.9642	0	(11)	2.4e+03	3	R.MADGFLAGLLAR.S + Oxidation (M)
<input checked="" type="checkbox"/> 207	999.3810	1996.7474	1997.9404	-1.1930	0	(34)	5.5	1	R.FCTTEGIENVILSDFR.F
<input checked="" type="checkbox"/> 209	999.3960	1996.7654	1997.9404	-1.1750	0	(40)	1.6	1	R.FCTTEGIENVILSDFR.F
<input checked="" type="checkbox"/> 210	999.4050	1996.7954	1997.9404	-1.1450	0	42	0.96	1	R.FCTTEGIENVILSDFR.F
<input checked="" type="checkbox"/> 232	1088.4760	2174.9374	2176.1198	-1.1824	0	(21)	1.5e+02	7	K.NVILNIFPSLDTGVCATSVR.K
<input checked="" type="checkbox"/> 233	725.9900	2174.9482	2176.1198	-1.1716	0	39	2.3	2	K.NVILNIFPSLDTGVCATSVR.K

MASCOT Mascot Search Results

Protein View

Match to: gi|53714055 Score: 484
 putative thiol peroxidase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6302) 10_RJ3_01_1363.d\SSP (6302) 10_RJ3_01_1363.mgf

Nominal mass (M_n): 18042; Calculated pI value: 5.55
 NCBI BLAST search of gi|53714055 against nr
 Unformatted [sequence_string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60682249 from Bacteroides fragilis NCTC 9343](#)
[gi|253564697 from Bacteroides sp. 3_2_5](#)
[gi|265766326 from Bacteroides sp. 2_1_16](#)
[gi|52216920 from Bacteroides fragilis YCH46](#)
[gi|60493683 from Bacteroides fragilis NCTC 9343](#)
[gi|251946162 from Bacteroides sp. 3_2_5](#)
[gi|263253994 from Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 68%

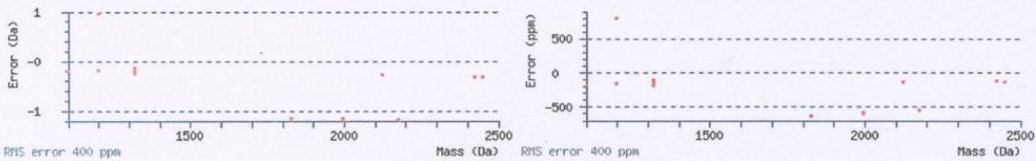
Matched peptides shown in **Red**

1 MATTNFKGQP VKLIGEFIQV GKVAPDFELV KSDLSSFALK DLKGKNIVLM
 51 IFFSLDTGVC ATSVKFNKM AAGMKDVTVL AISKDLFFAQ GRFCTTEGIE
 101 NVIPLSDFEF SDFDESYGVR MADGPLAGLL ARAVVVIGKD GKVAYTELVF
 151 EITQEPDYEK ALAAVK

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss Sequence
13 - 22	552.2190	1102.4234	1102.6386	-0.2152	0 K.LIGEFIQVGK.V (Ions_score 76)
46 - 65	1088.4760	2174.9374	2176.1198	-1.1824	0 K.NIVLNIPFSLDTGVCATSVR.K (Ions_score 24)
46 - 65	725.9900	2174.9482	2176.1198	-1.1716	0 K.NIVLNIPFSLDTGVCATSVR.K (Ions_score 48)
70 - 92	817.6610	2449.9612	2450.2661	-0.3050	2 K.MAAGMKDVTVLVLAISKDLFFAQGR.F 2 Oxidation (M) (Ions_score 54)
76 - 92	610.2910	1827.8512	1829.0047	-1.1535	1 K.DTVVLAISKDLFFAQGR.F (Ions_score 50)
93 - 109	999.3810	1996.7474	1997.9404	-1.1930	0 R.FCTTEGIENVPLSDFR.F (Ions_score 34)
93 - 109	999.3900	1996.7654	1997.9404	-1.1750	0 R.FCTTEGIENVPLSDFR.F (Ions_score 40)
93 - 109	999.4050	1996.7954	1997.9404	-1.1450	0 R.FCTTEGIENVPLSDFR.F (Ions_score 42)
110 - 120	661.1670	1320.3194	1320.5623	-0.2428	0 R.FSDFDESYGVR.M (Ions_score 16)
110 - 120	661.1670	1320.3194	1320.5623	-0.2428	0 R.FSDFDESYGVR.M (Ions_score 67)
110 - 120	661.1940	1320.3734	1320.5623	-0.1888	0 R.FSDFDESYGVR.M (Ions_score 69)
110 - 120	661.2080	1320.4014	1320.5623	-0.1608	0 R.FSDFDESYGVR.M (Ions_score 23)
110 - 120	661.2180	1320.4214	1320.5623	-0.1408	0 R.FSDFDESYGVR.M (Ions_score 57)
110 - 120	661.2260	1320.4374	1320.5623	-0.1248	0 R.FSDFDESYGVR.M (Ions_score 58)
121 - 132	600.7380	1199.4614	1199.6332	-0.1718	0 R.MADGPLAGLLAR.A Oxidation (M) (Ions_score 68)
121 - 132	601.3060	1200.5974	1199.6332	0.9642	0 R.MADGPLAGLLAR.A Oxidation (M) (Ions_score 11)
140 - 160	808.6340	2422.8802	2423.1744	-0.2942	1 K.DGKVVAYTELVPEITQEPDYEK.A (Ions_score 43)
143 - 160	708.5960	2122.7662	2123.0310	-0.2648	0 K.VAYTELVPEITQEPDYEK.A (Ions_score 37)



LOCUS YP_100047 166 aa linear BCT 26-APR-2009
 DEFINITION putative thiol peroxidase [Bacteroides fragilis YCH46].
 ACCESSION YP_100047
 VERSION YP_100047.1 GI:53714055
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession [NC_006347.1](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 166)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 166)
 CONSRTM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 166)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.

TITLE Direct Submission
JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD49513](#).
 Method: conceptual translation.

FEATURES Location/Qualifiers

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 BLASTP E(): 9e-84"
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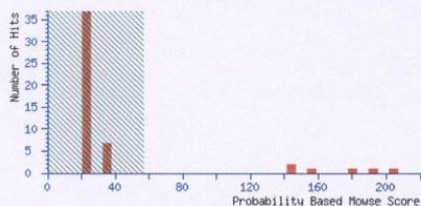
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6802
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6802) 10_RJ10_01_1377.d\SSP (6802) 10_RJ10_01_1377.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:48:01 GMT
 Protein hits : [gi|91215296](#) 3-hydroxybutyryl-CoA dehydrogenase [Psychroflexus torquis ATCC 700755]
[gi|53714585](#) fructose-bisphosphate aldolase [Bacteroides fragilis YCH46]
[gi|255010781](#) fructose-bisphosphate aldolase [Bacteroides fragilis 3_1_12]
[gi|29347101](#) fructose-bisphosphate aldolase [Bacteroides thetaiotaomicron VPI-5482]
[gi|153808476](#) hypothetical protein BACCAC_02770 [Bacteroides caccae ATCC 43185]
[gi|160886425](#) hypothetical protein BACOVA_04436 [Bacteroides ovatus ATCC 8483]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As **Peptide Summary** [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Error tolerant

1.	gi 91215296	Mass: 32980	Score: 205	Queries matched: 4	emPAI: 0.21		
	3-hydroxybutyryl-CoA dehydrogenase [Psychroflexus torquis ATCC 700755]						
	<input type="checkbox"/> Check to include this hit in error tolerant search						
	Query	Observed	Mr(expt)	Mr(calc)	Delta Miss Score	Expect Rank	Peptide
	<input checked="" type="checkbox"/> 95	393.8730	1178.5972	1178.6845	-0.0874 0 29	33 1	K.AIGMLALPLHK.E + Oxidation (M)
	<input checked="" type="checkbox"/> 166	579.9080	1736.7022	1736.8832	-0.1811 0 45	0.64 1	K.DADLLIEAVFENPDVK.T
	<input checked="" type="checkbox"/> 189	896.3390	1990.6634	1990.9312	-0.2678 0 79	0.00021 1	K.LGVSNNGEFGYTYPEPAYK.S
	<input checked="" type="checkbox"/> 229	856.0420	2565.1042	2565.3439	-0.2397 0 51	0.16 1	K.TVFTNSSTLLPSQFAEVTGRPSK.F
2.	gi 53714585	Mass: 36766	Score: 196	Queries matched: 6	emPAI: 0.30		
	fructose-bisphosphate aldolase [Bacteroides fragilis YCH46]						
	<input type="checkbox"/> Check to include this hit in error tolerant search						
	Query	Observed	Mr(expt)	Mr(calc)	Delta Miss Score	Expect Rank	Peptide
	<input checked="" type="checkbox"/> 131	693.3180	1384.6214	1384.7562	-0.1347 1 37	6 1	K.IINVLGSDNKLAE.-
	<input checked="" type="checkbox"/> 168	880.8160	1759.6174	1759.8889	-0.2714 0 (17)	3.5e+02 1	R.MVPPPLAFDVLGVMK.E + 2 Oxidation (M)
	<input checked="" type="checkbox"/> 169	587.5700	1759.6882	1759.8889	-0.2007 0 36	6.1 1	R.MVPPPLAFDVLGVMK.E + 2 Oxidation (M)
	<input checked="" type="checkbox"/> 176	919.3450	1836.6754	1837.8516	-1.1762 0 (41)	1.6 1	K.TGCDLSLAISIGTSHGAYK.F
	<input checked="" type="checkbox"/> 177	613.2370	1836.6892	1837.8516	-1.1625 0 74	0.00079 1	K.TGCDLSLAISIGTSHGAYK.F
	<input checked="" type="checkbox"/> 222	796.9540	2387.8402	2388.1242	-0.2840 0 49	0.18 1	K.GGYAIPAFNFMMEQQAIIK.A + 2 Oxidation (M)
3.	gi 255010781	Mass: 36694	Score: 179	Queries matched: 5	emPAI: 0.30		
	fructose-bisphosphate aldolase [Bacteroides fragilis 3_1_12]						
	<input type="checkbox"/> Check to include this hit in error tolerant search						
	Query	Observed	Mr(expt)	Mr(calc)	Delta Miss Score	Expect Rank	Peptide
	131	693.3180	1384.6214	1384.7562	-0.1347 1 37	6 1	K.IINVLGSDNKLAE.-
	169	587.5700	1759.6882	1758.9049	0.7833 0 19	2.9e+02 7	R.MVPPPLAFNVLGVMK.E + 2 Oxidation (M)
	176	919.3450	1836.6754	1837.8516	-1.1762 0 (41)	1.6 1	K.TGCDLSLAISIGTSHGAYK.F
	177	613.2370	1836.6892	1837.8516	-1.1625 0 74	0.00079 1	K.TGCDLSLAISIGTSHGAYK.F
	222	796.9540	2387.8402	2388.1242	-0.2840 0 49	0.18 1	K.GGYAIPAFNFMMEQQAIIK.A + 2 Oxidation (M)
4.	gi 29347101	Mass: 36791	Score: 160	Queries matched: 4	emPAI: 0.30		
	fructose-bisphosphate aldolase [Bacteroides thetaiotaomicron VPI-5482]						
	<input type="checkbox"/> Check to include this hit in error tolerant search						

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<u>131</u>	693.3180	1384.6214	1384.7562	-0.1347	1	37	6	1	K.IILNVLGSDNKLAE.-
<u>176</u>	919.3450	1836.6754	1837.8516	-1.1762	0	(41)	1.6	1	R.TGCDLSLAISIGTSHGAYK.F
<u>177</u>	613.2370	1836.6892	1837.8516	-1.1625	0	74	0.00079	1	R.TGCDLSLAISIGTSHGAYK.F
<u>222</u>	796.9540	2387.8402	2388.1242	-0.2840	0	49	0.18	1	K.GGYAIPAFNPNMMEQMQAIIK.A + 2 Oxidation (M)

5. [gi|153808476](#) Mass: 36778 Score: 148 Queries matched: 4 emPAI: 0.30
hypothetical protein BACCAC_02770 [Bacteroides caccae ATCC 43185]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<u>131</u>	693.3180	1384.6214	1383.7722	0.8493	1	25	1e+02	3	K.IILNVLGSDNKLQA.-
<u>176</u>	919.3450	1836.6754	1837.8516	-1.1762	0	(41)	1.6	1	R.TGCDLSLAISIGTSHGAYK.F
<u>177</u>	613.2370	1836.6892	1837.8516	-1.1625	0	74	0.00079	1	R.TGCDLSLAISIGTSHGAYK.F
<u>222</u>	796.9540	2387.8402	2388.1242	-0.2840	0	49	0.18	1	K.GGYAIPAFNPNMMEQMQAIIK.A + 2 Oxidation (M)

Proteins matching the same set of peptides:

[gi|255693155](#) Mass: 36831 Score: 148 Queries matched: 4
fructose-1,6-bisphosphate aldolase, class II [Bacteroides finegoldii DSM 17565]
[gi|260173446](#) Mass: 36803 Score: 148 Queries matched: 4
fructose-bisphosphate aldolase [Bacteroides sp. D2]

6. [gi|160886425](#) Mass: 36875 Score: 148 Queries matched: 4 emPAI: 0.29
hypothetical protein BACOVA_04436 [Bacteroides ovatus ATCC 8483]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<u>131</u>	693.3180	1384.6214	1383.7722	0.8493	1	25	1e+02	3	K.IILNVLGSDNKLQA.-
<u>176</u>	919.3450	1836.6754	1837.8516	-1.1762	0	(41)	1.6	1	R.TGCDLSLAISIGTSHGAYK.F
<u>177</u>	613.2370	1836.6892	1837.8516	-1.1625	0	74	0.00079	1	R.TGCDLSLAISIGTSHGAYK.F
<u>222</u>	796.9540	2387.8402	2388.1242	-0.2840	0	49	0.18	1	K.GGYAIPAFNPNMMEQMQAIIK.A + 2 Oxidation (M)

Proteins matching the same set of peptides:

[gi|237713705](#) Mass: 36861 Score: 148 Queries matched: 4
fructose-bisphosphate aldolase [Bacteroides sp. D1]
[gi|237723289](#) Mass: 36773 Score: 148 Queries matched: 4
fructose-bisphosphate aldolase [Bacteroides sp. 2_2_4]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> <u>144</u>	755.2960	2262.8662	2262.1541	0.7121	2	33	20	1	SEALFRNMTGFKYTLPIK + Oxidation (M)
<input checked="" type="checkbox"/> <u>170</u>	895.1500	2682.4282	2683.4302	-1.0020	2	29	50	1	RDGMILIPYMTLSPAIHQIVARTK + Oxidation (M)
<input checked="" type="checkbox"/> <u>197</u>	1033.8310	2065.6474	2066.0643	-0.4169	2	27	63	1	AGAEAKPEAKSEAKSAPAPEK
<input checked="" type="checkbox"/> <u>91</u>	584.8950	1751.6632	1751.8045	-0.1413	0	27	85	1	MDTGVMVYINHPTVMK + Oxidation (M)
<input checked="" type="checkbox"/> <u>147</u>	766.4820	2296.4242	2295.2773	1.1469	2	27	98	1	PLPKRALLAGFSNVLEQAMPK + Oxidation (M)
<input checked="" type="checkbox"/> <u>210</u>	735.6100	2203.8082	2202.9807	0.8275	0	26	44	1	VMSTDLNMGMTATTVOGSDVK + 2 Oxidation (M)
<input checked="" type="checkbox"/> <u>88</u>	563.1570	1686.4492	1685.7984	0.6508	0	25	1.1e+02	1	AAYGMPFHTWLTDR
<input checked="" type="checkbox"/> <u>116</u>	656.7810	1967.3212	1966.8587	0.4625	0	25	1.3e+02	1	MPADHVMVMAGFTAGNEK + 3 Oxidation (M)
<input checked="" type="checkbox"/> <u>105</u>	634.6000	1900.7782	1901.9694	-1.1913	0	25	1.4e+02	1	IDLTGSSSSNGTLSDPLLR
<input checked="" type="checkbox"/> <u>96</u>	592.6870	1775.0392	1774.0427	0.9965	0	24	1.9e+02	1	ILIVGLGVIGGSFAMALK + Oxidation (M)
<input checked="" type="checkbox"/> <u>175</u>	915.6430	2743.9072	2743.3534	0.5538	2	24	1.3e+02	1	DMPHRLQFTGPTGTNAVEAMKLAR + 2 Oxidation (M)
<input checked="" type="checkbox"/> <u>58</u>	406.0880	1215.2422	1214.6003	0.6418	1	24	1.5e+02	1	GSPQAGGSEARAK
<input checked="" type="checkbox"/> <u>64</u>	831.2410	830.2337	830.4610	-0.2273	1	24	1.4e+02	1	VRGSGDIK
<input checked="" type="checkbox"/> <u>167</u>	582.5710	1744.6912	1744.8903	-0.1991	2	24	96	1	RSNAILAMRNSYHR
<input checked="" type="checkbox"/> <u>143</u>	751.7510	2252.2312	2251.0475	1.1836	1	24	1.9e+02	1	MLFNGLCEPQASLCFRK + Oxidation (M)
<input checked="" type="checkbox"/> <u>70</u>	453.7260	905.4374	905.4463	-0.0088	0	24	1.5e+02	1	DLMLAGMR
<input checked="" type="checkbox"/> <u>109</u>	640.6990	1919.0752	1918.9232	0.1520	1	24	2e+02	1	KAVADVAGESSISDVSGDGR
<input checked="" type="checkbox"/> <u>152</u>	798.8320	2393.4742	2393.2638	0.2104	2	23	1.9e+02	1	AMLLKPTKDDVQGLGFHSHKR + Oxidation (M)
<input checked="" type="checkbox"/> <u>165</u>	865.3170	2592.9292	2592.3193	0.6099	2	23	1.6e+02	1	ANMTFHKGLYQDVVAVVCIGIK
<input checked="" type="checkbox"/> <u>223</u>	800.5750	2398.7032	2398.2566	0.4466	1	23	77	1	EYGLMRAAGGLITVATVYLYK + Oxidation (M)
<input checked="" type="checkbox"/> <u>87</u>	563.0610	1686.1612	1685.6228	0.5384	0	23	1.9e+02	1	APAAPMPDPSMGGM + 4 Oxidation (M)
<input checked="" type="checkbox"/> <u>212</u>	1126.9670	2251.9194	2251.1267	0.7928	1	23	96	1	QAFSDLTATLLDNLGMETRR
<input checked="" type="checkbox"/> <u>112</u>	647.4630	1939.3672	1939.8218	-0.4546	0	23	2e+02	1	MSEFGDSASAGTAGRPTSPD
<input checked="" type="checkbox"/> <u>137</u>	479.2240	1434.6502	1435.7558	-1.1057	0	23	1.5e+02	1	LTDADVGLASLYK
<input checked="" type="checkbox"/> <u>150</u>	780.5170	2338.5292	2339.1411	-0.6119	2	23	2.2e+02	1	YRVQMFVPTMYRMLVR + 3 Oxidation (M)
<input checked="" type="checkbox"/> <u>75</u>	498.2930	994.5714	994.4906	0.0808	0	22	1.5e+02	1	MGIFAAAGGK + Oxidation (M)
<input checked="" type="checkbox"/> <u>163</u>	858.1400	2571.3982	2571.2751	0.1230	1	22	2.3e+02	1	VRAAFEDQDAHVVVHDAIVTNK + Oxidation (M)
<input checked="" type="checkbox"/> <u>191</u>	996.9930	2987.9572	2988.4970	-0.5398	1	22	2.2e+02	1	SLHTIKSDVNLNLTDDTMSMLVNGIK + Oxidation (M)
<input checked="" type="checkbox"/> <u>173</u>	912.2700	2733.7882	2733.3771	0.4111	0	22	1.8e+02	1	VMGGLLATGIASAPFLWSMLHDYQR
<input checked="" type="checkbox"/> <u>190</u>	664.5630	1990.6672	1989.9353	0.7318	1	22	1e+02	1	EEVFPDWKMQDSILK + Oxidation (M)
<input checked="" type="checkbox"/> <u>139</u>	731.8500	2192.5282	2193.2191	-0.6909	2	22	2.8e+02	1	MPDAKPVVEKDGAKITLVNGK + Oxidation (M)
<input checked="" type="checkbox"/> <u>135</u>	475.2010	1422.5812	1423.6806	-1.0994	1	22	1.6e+02	1	YFGMKNKITSYK + Oxidation (M)
<input checked="" type="checkbox"/> <u>86</u>	562.7810	1123.5474	1122.7125	0.8350	1	22	1.7e+02	1	IVLQAQKVPK
<input checked="" type="checkbox"/> <u>181</u>	942.9120	2825.7142	2826.3705	-0.6563	2	22	2.4e+02	1	NSYDMAETSRDTAVELLKELLEQR + Oxidation (M)
<input checked="" type="checkbox"/> <u>203</u>	710.8680	2129.5822	2129.2321	0.3501	1	22	1.1e+02	1	PEPRILVLGAGPAGTATALGLR
<input checked="" type="checkbox"/> <u>208</u>	1091.6490	3271.9252	3271.6673	0.2579	1	22	2.3e+02	1	QVPGLPAPQAGGGHMSPALVAQAWNRYAGGLK + Oxidation (M)
<input checked="" type="checkbox"/> <u>184</u>	641.3350	1920.9832	1921.9720	-0.9888	1	22	1.8e+02	1	RAGYATVVMALDLDHFK + Oxidation (M)
<input checked="" type="checkbox"/> <u>76</u>	499.2150	996.4154	995.4971	0.9184	2	22	1.7e+02	1	RMKAFGDR + Oxidation (M)
<input checked="" type="checkbox"/> <u>193</u>	1009.2650	3024.7732	3024.4507	0.3224	2	21	2.2e+02	1	KLFGSQATVVTSGHAWLDMMPMSADKKG + 2 Oxidation (M)

MASCOT Mascot Search Results

Protein View

Match to: gi|53714585 Score: 196
 fructose-bisphosphate aldolase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6802) 10_RJ10_01_1377.d\SSP (6802) 10_RJ10_01_1377.mgf

Nominal mass (M_r): 36766; Calculated pI value: 5.37
 NCBI BLAST search of gi|53714585 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: [Bacteroides fragilis YCH46](#)
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60682607 from Bacteroides fragilis NCTC 9343](#)
[gi|253565089 from Bacteroides sp. 3_2_5](#)
[gi|265765951 from Bacteroides sp. 2_1_16](#)
[gi|52217450 from Bacteroides fragilis YCH46](#)
[gi|60494041 from Bacteroides fragilis NCTC 9343](#)
[gi|251946554 from Bacteroides sp. 3_2_5](#)
[gi|263253619 from Bacteroides sp. 2_1_16](#)

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 20%

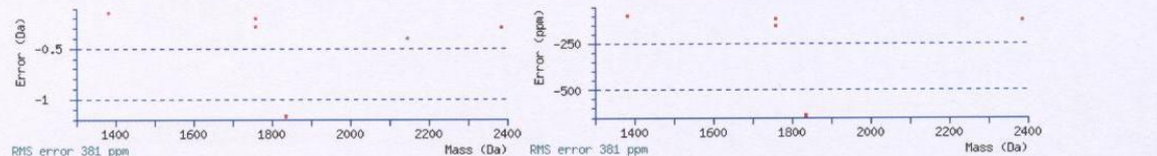
Matched peptides shown in **Bold Red**

1 MVNYKDLGLV NTRDMFAKAI KGGYAI**PAFN FNNMEQMQAI** IKA~~AVETKSP~~
 51 VILQVSKGAR QYANATLLRY MAQGA**VEYAK** ELGCKN**PRIV** LHLDHGDT**FE**
 101 TCKSCIDSGF SSMIDGSHL PYDENVALTK KVV**EYAHOFD** VT**VEGELGVL**
 151 AGVEDEVSSD HHTYTEPDEV VDFV**FKTGD** SLAIS**IGTSH** GAYK**FTPEQC**
 201 HIDPKTGRMV **FPPLAFD**VLD GVMKELPGFP IVLHGSS**VP** EEEVATIN**QF**
 251 GGALKAATGI PEEELRKA**AK** SAVCKINID**S** DSRLAMTAAI RKVFAEK**PAE**
 301 FDP**RRYL**GPA RDNMEKLYKH KI**INVLGSDN** **KLAE**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr(expt)	Mr(calc)	Delta	Miss Sequence
22 - 42	796.9540	2387.8402	2388.1242	-0.2840	0 K.GGYAI PAFN FNNMEQMQAIK.A 2 Oxidation (M) (Ions score 49)
177 - 194	919.3450	1836.6754	1837.8516	-1.1762	0 K.TGCD SLAISIGTSH GAYK.F (Ions score 41)
177 - 194	613.2370	1836.6892	1837.8516	-1.1625	0 K.TGCD SLAISIGTSH GAYK.F (Ions score 74)
209 - 224	880.8160	1759.6174	1759.8889	-0.2714	0 R.MVPP PLAFD VLDGVMK.E 2 Oxidation (M) (Ions score 17)
209 - 224	587.5700	1759.6882	1759.8889	-0.2007	0 R.MVPP PLAFD VLDGVMK.E 2 Oxidation (M) (Ions score 36)
322 - 334	693.3180	1384.6214	1384.7562	-0.1347	1 K.I INVLGSDN KLAE.- (Ions score 37)



LOCUS YP_100577 334 aa linear BCT 26-APR-2009
 DEFINITION fructose-bisphosphate aldolase [Bacteroides fragilis YCH46].
 ACCESSION YP_100577
 VERSION YP_100577.1 GI:53714585
 DBLINK Project:13067
 DBSOURCE REPSEQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM [Bacteroides fragilis YCH46](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 334)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED [15466707](#)
 REFERENCE 2 (residues 1 to 334)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 334)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT PROVISIONAL REPSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [BAD50043](#).
 Method: conceptual translation.
 FEATURES Location/Qualifiers

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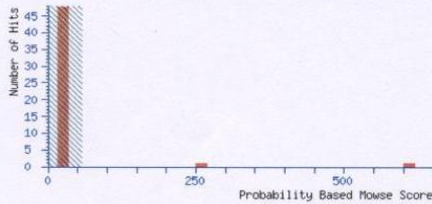
Mascot: <http://www.matrixscience.com/>

MASCOT Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6807
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6807) 10_RJ8_01_1373.d\SSP (6807) 10_RJ8_01_1373.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 13:49:18 GMT
 Protein hits : [gi|53712366](#) hypothetical protein BF1074 [Bacteroides fragilis YCH46]
 [gi|255007873](#) hypothetical protein Bfra3_01963 [Bacteroides fragilis 3_1_12]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold $p <$ Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53712366](#) Mass: 49095 Score: 610 Queries matched: 14 emPAI: 0.68
 hypothetical protein BF1074 [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 88	501.7010	1001.3874	1001.4892	-0.1017	0	38	4.8	1	K.DLFMLGYK.A + Oxidation (M)
<input checked="" type="checkbox"/> 90	547.7660	1093.5174	1093.5808	-0.0633	0	70	0.0026	1	K.AVDYIAFAPK.V
<input checked="" type="checkbox"/> 108	675.7900	1349.5654	1349.6616	-0.0961	0	55	0.082	1	K.VDVNQAYDWLK.K
<input checked="" type="checkbox"/> 121	493.5310	1477.5712	1477.7565	-0.1853	1	(20)	2.2e+02	4	K.VDVNQAYDWLKK.S
<input checked="" type="checkbox"/> 122	739.8240	1477.6334	1477.7565	-0.1231	1	21	2.4e+02	1	K.VDVNQAYDWLKK.S
<input checked="" type="checkbox"/> 123	752.3240	1502.6334	1502.8093	-0.1758	0	58	0.043	1	R.ITIGGWIGESTTIR.-
<input checked="" type="checkbox"/> 130	518.3990	1552.1752	1552.6980	-0.5228	0	12	1.1e+03	10	K.YFDELMNTHDLR.M
<input checked="" type="checkbox"/> 137	813.7950	1625.5754	1625.7573	-0.1818	0	110	2e-07	1	K.FFDEAINLETDNAK.K
<input checked="" type="checkbox"/> 185	973.3220	1944.6294	1945.8040	-1.1746	0	74	0.00066	1	K.CTDSDAYQQASFYVYK.I
<input checked="" type="checkbox"/> 102	649.2260	1944.6562	1945.8040	-1.1478	0	(24)	1.7e+02	1	K.CTDSDAYQQASFYVYK.I
<input checked="" type="checkbox"/> 205	682.6080	2044.8022	2044.9564	-0.1542	0	(30)	18	1	K.AESAAATLFYFLQMSHDK.L + Oxidation (M)
<input checked="" type="checkbox"/> 206	1023.4270	2044.8394	2044.9564	-0.1170	0	61	0.015	1	K.AESAAATLFYFLQMSHDK.L + Oxidation (M)
<input checked="" type="checkbox"/> 244	894.9800	2681.9182	2682.2847	-0.3665	0	57	0.029	1	K.DNLVALFINSGTADCESLQGIYGPK.V
<input checked="" type="checkbox"/> 251	987.7030	2960.0872	2960.3927	-0.3055	1	55	0.04	1	K.RQPIQDYLAASEYADDAIAAADKESVK.K

Proteins matching the same set of peptides:
[gi|60491969](#) Mass: 48413 Score: 610 Queries matched: 14
 conserved hypothetical protein [Bacteroides fragilis NCTC 9343]

2. [gi|255007873](#) Mass: 49151 Score: 267 Queries matched: 4 emPAI: 0.21
 hypothetical protein Bfra3_01963 [Bacteroides fragilis 3_1_12]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input type="checkbox"/> 88	501.7010	1001.3874	1001.4892	-0.1017	0	38	4.8	1	K.DLFMLGYK.A + Oxidation (M)
<input type="checkbox"/> 90	547.7660	1093.5174	1093.5808	-0.0633	0	70	0.0026	1	K.AVDYIAFAPK.M
<input type="checkbox"/> 137	813.7950	1625.5754	1625.7573	-0.1818	0	110	2e-07	1	K.FFDEAINLETDNAK.K
<input checked="" type="checkbox"/> 189	981.2920	1960.5694	1959.8196	0.7498	0	49	0.23	1	K.CTDSBAYQQASFYVYK.I

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 232	766.9760	2297.9062	2297.1321	0.7740	1	33	8.4	1	SGMIVSDNGTEPTSNILARSK
<input checked="" type="checkbox"/> 91	555.8140	1109.6134	1108.5699	1.0435	0	33	13	1	IGVGHPSMAPK + Oxidation (M)

MASCOT Mascot Search Results

Protein View

Match to: **gi|60491969** Score: **610**
conserved hypothetical protein [Bacteroides fragilis NCTC 9343]
Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6807) 10_RJ8_01_1373.d\SSP (6807) 10_RJ8_01_1373.mgf

Nominal mass (M_r): **48413**; Calculated pI value: **5.79**
NCBI BLAST search of **gi|60491969** against nr
Unformatted **sequence string** for pasting into other applications

Taxonomy: **Bacteroides fragilis NCTC 9343**

Fixed modifications: Carboxymethyl (C)
Variable modifications: Oxidation (M)
Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
Sequence Coverage: **35%**

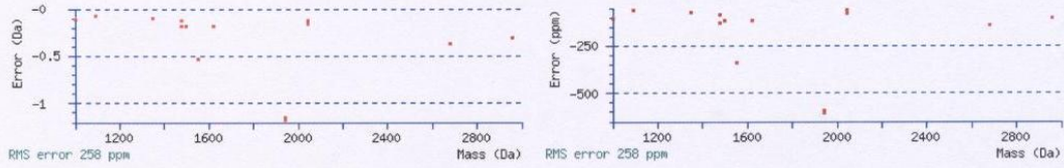
Matched peptides shown in **Bold Red**

- 1 MAVLFLSAGA TTVAQDDAN CNSNSSISHE AVKAGNFKDA YTPWKAVLEN
- 51 CPTLRPYTFT DGYKILKGLL GQIKDRNSAE YKRYFDEL MN THDLRMKYTO
- 101 EFLGKGVKVS SEDEALGIKA VDYIAFAPKV DVNQAYDWLK KSVDAAKAES
- 151 AATLFLYFLQ MSHDKLKEDE AHKEQFIQDY LAASEYADDA IAAADKESVK
- 201 KAFGGIKDNL VALFINSGTA DCESLQGIYG PKVETNQTDL NYLKKVISIM
- 251 KMMKCTDSDA YQQASFYVYK IEPSAEAAATG CAYQAYKKGIDG SVKFFDE
- 301 **AINLETDNAK** KA EKAYAAAS VLTAKKLSQ ARSYAQKAI S FNEYGAPYI
- 351 LIANLYAMSP NWSDESALNK CTYPAVIDKL QKAKSVDPVS TEEVNMKISR
- 401 **YSAYTPQAKD** **LFMLGYKAGD** **RITIGGWIGE** **STTIR**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
84 - 95	518.3990	1552.1752	1552.6980	-0.5228	0	K.YFDEL MN THDLR.M (Ions score 12)
120 - 129	547.7660	1093.5174	1093.5808	-0.0633	0	K.AVDYIAFAPK.V (Ions score 70)
130 - 140	675.7900	1349.5654	1349.6616	-0.0961	0	K.VDVNQAYDWLK.K (Ions score 55)
130 - 141	493.5310	1477.5712	1477.7565	-0.1853	1	K.VDVNQAYDWLKK.S (Ions score 20)
130 - 141	739.8240	1477.6334	1477.7565	-0.1231	1	K.VDVNQAYDWLKK.S (Ions score 21)
148 - 165	682.6080	2044.8022	2044.9564	-0.1542	0	K.AESAAATL FVFLQMSHDK.L Oxidation (M) (Ions score 30)
148 - 165	1023.4270	2044.8394	2044.9564	-0.1170	0	K.AESAAATL FVFLQMSHDK.L Oxidation (M) (Ions score 61)
174 - 200	987.7030	2960.0872	2960.3927	-0.3055	1	K.EQFIQDYLAASEYADDA IAAADKESVK.K (Ions score 55)
208 - 232	894.9800	2681.9182	2682.2847	-0.3665	0	K.DNLVALFINSGTADCE SLQGIYGPK.V (Ions score 57)
255 - 270	973.3220	1944.6294	1945.8040	-1.1746	0	K.CTSDSAYQQASFYVYK.I (Ions score 74)
255 - 270	649.2260	1944.6562	1945.8040	-1.1478	0	K.CTSDSAYQQASFYVYK.I (Ions score 24)
297 - 310	813.7950	1625.5754	1625.7573	-0.1818	0	K.FFDEAINLETDNAK.K (Ions score 110)
410 - 417	501.7010	1001.3874	1001.4892	-0.1017	0	K.DLFLMGYK.A Oxidation (M) (Ions score 38)
422 - 435	752.3240	1502.6334	1502.8093	-0.1758	0	R.ITIGGWIGESTTIR.- (Ions score 58)



LOCUS CAH06730 435 aa linear BCT 13-MAY-2009
DEFINITION conserved hypothetical protein [Bacteroides fragilis NCTC 9343].
ACCESSION CAH06730
VERSION CAH06730.1 GI:60491969
DBSOURCE embl accession CR626927.1
KEYWORDS .
SOURCE Bacteroides fragilis NCTC 9343
ORGANISM **Bacteroides fragilis NCTC 9343**
Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
Bacteroidaceae; Bacteroides.
REFERENCE 1 (residues 1 to 435)
AUTHORS Cerdeno-Tarraga,A.M., Patrick,S., Crossman,L.C., Blakely,G.,
Abratt,V., Lennard,N., Poxton,I., Duerden,B., Harris,B.,
Quail,M.A., Barron,A., Clark,L., Corton,C., Doggett,J.,
Holden,M.T., Larke,N., Line,A., Lord,A., Norbertczak,H., Ormond,D.,
Price,C., Rabinowitsch,E., Woodward,J., Barrell,B. and Parkhill,J.
TITLE Extensive DNA inversions in the B. fragilis genome control variable
gene expression
JOURNAL Science 307 (5714), 1463-1465 (2005)
PUBMED 15746427
REFERENCE 2 (residues 1 to 435)
AUTHORS Cerdeno-Tarraga,A.M.
TITLE Direct Submission
JOURNAL Submitted (29-JUL-2004) Cerdeno-Tarraga A.M., submitted on behalf
of the Pathogen Sequencing Unit, Sanger Institute, Wellcome Trust
Genome Campus, Hinxton, Cambridge CB10 1SA E-mail:

amct@sanger.ac.uk
FEATURES Location/Qualifiers
source 1..435
/organism="Bacteroides fragilis NCTC 9343"
/strain="ATCC 25285 = NCTC 9343"
/db_xref="taxon:272559"
Protein 1..435
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CDS 1..435
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/coded_by="CR626927.1:1241398..1242705"
/note="Similar to Bacteroides thetaiotaomicron
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(433 aa) fasta scores: E(): 8.7e-145, 85.21% id in 433 aa"
/transl_table=11
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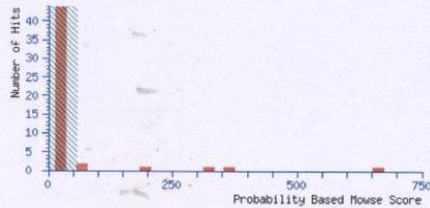
Mascot: <http://www.matrixscience.com/>

MASCOT SCIENCE Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 6901
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf
 Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 7 Jan 2010 at 16:32:37 GMT
 Protein hits :
 gi|53715725 phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]
 gi|150003549 phosphoenolpyruvate carboxykinase [Bacteroides vulgatus ATCC 8482]
 gi|150009397 phosphoenolpyruvate carboxykinase [Parabacteroides distasonis ATCC 8503]
 gi|34541315 phosphoenolpyruvate carboxykinase [Porphyromonas gingivalis W83]
 gi|123444162 phosphoenolpyruvate carboxykinase [Yersinia enterocolitica subsp. enterocolitica 8081]
 gi|281421519 phosphoenolpyruvate carboxykinase [Prevotella copri DSM 18205]

Probability Based Mowse Score

Ions score is $-10 * \log(P)$, where P is the probability that the observed match is a random event.
 Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$).
 Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: **Peptide Summary** [Help](#)

Significance threshold $p < 0.05$ Max. number of hits **AUTO**

Standard scoring MudPIT scoring Ions score or expect cut-off **0** Show sub-sets **0**

Show pop-ups Suppress pop-ups Sort unassigned **Decreasing Score** Require bold red

Error tolerant

1. [gi|53715725](#) Mass: 59339 Score: 663 Queries matched: 14 emPAI: 0.71
 phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
115	684.2770	1366.5394	1365.6677	0.8717	0	24	89	3	K.AYLVNTGWNNGSGK.R
<input checked="" type="checkbox"/> 120	708.3170	1414.6194	1414.7304	-0.1109	0	93	1.4e-05	1	R.DALLENVTVAADGK.I
<input checked="" type="checkbox"/> 141	811.3150	1620.6154	1620.7098	-0.0944	0	47	0.4	1	K.GMFSIMNMYNPLR.G + 3 Oxidation (M)
<input checked="" type="checkbox"/> 142	818.3880	1634.7614	1634.8515	-0.0901	0	110	2.5e-07	1	K.ELGLNSETATVFNLK.T
<input checked="" type="checkbox"/> 151	843.3600	1684.7054	1685.7719	-1.0665	0	32	15	1	R.LFVVDTPCGANEGTR.M
<input checked="" type="checkbox"/> 153	863.9200	1725.8254	1725.9512	-0.1258	1	60	0.029	1	R.GIIDAILDGSDKAPT.K.V
<input checked="" type="checkbox"/> 186	972.9460	1943.8774	1944.0204	-0.1429	1	53	0.14	1	K.VINLDKESRPDIFNAIK.R
<input checked="" type="checkbox"/> 190	656.6100	1966.8082	1966.9783	-0.1701	0	(46)	0.52	1	K.GQVTELGAVNVMTGVYTR.S + Oxidation (M)
<input checked="" type="checkbox"/> 191	984.4190	1966.8234	1966.9783	-0.1548	0	115	6.2e-08	1	K.GQVTELGAVNVMTGVYTR.S + Oxidation (M)
<input checked="" type="checkbox"/> 195	993.4470	1984.8794	1985.0986	-0.2191	0	(59)	0.05	1	R.VSYPYHIENIVKPVSK.G
<input checked="" type="checkbox"/> 196	662.6660	1984.9762	1985.0986	-0.1224	0	62	0.015	1	R.VSYPYHIENIVKPVSK.G
<input checked="" type="checkbox"/> 215	705.9670	2114.8792	2115.1293	-0.2501	0	(16)	5.5e+02	5	K.VIPFFDFVVPTELGVDPK.I
<input checked="" type="checkbox"/> 216	1058.5150	2115.0154	2115.1293	-0.1138	0	41	1.9	1	K.VIPFFDFVVPTELGVDPK.I
<input checked="" type="checkbox"/> 250	989.3890	2965.1452	2966.2732	-1.1281	0	33	7.2	1	R.GIASMHCASANTDMEGTSSAIFPGLSGTGK.T + 2 Oxidation (M)

2. [gi|150003549](#) Mass: 59400 Score: 366 Queries matched: 6 emPAI: 0.31
 phosphoenolpyruvate carboxykinase [Bacteroides vulgatus ATCC 8482]

Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
141	811.3150	1620.6154	1620.7098	-0.0944	0	47	0.4	1	K.GMFSIMNMYNPLR.G + 3 Oxidation (M)
142	818.3880	1634.7614	1634.8515	-0.0901	0	110	2.5e-07	1	K.ELGLNSETATVFNLK.T
153	863.9200	1725.8254	1725.9512	-0.1258	1	60	0.029	1	R.GIIDAILDGSDKAPT.K.V
190	656.6100	1966.8082	1966.9783	-0.1701	0	(46)	0.52	1	K.GQVTELGAVNVMTGVYTR.S + Oxidation (M)
191	984.4190	1966.8234	1966.9783	-0.1548	0	115	6.2e-08	1	K.GQVTELGAVNVMTGVYTR.S + Oxidation (M)
250	989.3890	2965.1452	2965.2892	-0.1440	0	33	7.2	1	R.GIASMHCASANTNMEGTSSAIFPGLSGTGK.T + 2 Oxidation (M)

Proteins matching the same set of peptides:
[gi|212692197](#) Mass: 59384 Score: 366 Queries matched: 6
 hypothetical protein BACDOR_01692 [Bacteroides dorei DSM 17855]
[gi|254881217](#) Mass: 59416 Score: 366 Queries matched: 6

3. [gi|150009397](#) Mass: 59213 Score: 334 Queries matched: 6 emPAI: 0.24
 phosphoenolpyruvate carboxykinase [Parabacteroides distasonis ATCC 8503]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
115	684.2770	1366.5394	1365.6677	0.8717	0	24	74	3	K.AYLVNTGWNGSGK.R
120	708.3170	1414.6194	1414.7304	-0.1109	0	93	1.2e-05	1	R.DALLENVTVAADGK.I
142	818.3880	1634.7614	1634.8515	-0.0901	0	110	2.1e-07	1	K.ELGLENSETATVFNLR.T
153	863.9200	1725.8254	1725.9149	-0.0894	0	44	0.91	2	R.GIIDAILDGSINEAPTK.K
195	993.4470	1984.8794	1985.0986	-0.2191	0	(59)	0.042	1	R.VSYPYIHENIVKPVSK.G
196	662.6660	1984.9762	1985.0986	-0.1224	0	62	0.013	1	R.VSYPYIHENIVKPVSK.G

4. [gi|34541315](#) Mass: 59587 Score: 178 Queries matched: 4 emPAI: 0.24
 phosphoenolpyruvate carboxykinase [Porphyromonas gingivalis W83]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
120	708.3170	1414.6194	1413.7463	0.8731	0	58	0.039	3	R.NALLENVTVAADGK.I
153	863.9200	1725.8254	1725.9512	-0.1258	1	60	0.024	1	R.GIIDAILDGSIDKAPTK.A
195	993.4470	1984.8794	1985.0986	-0.2191	0	(59)	0.042	1	R.VSYPYIHENIVKPVSK.A
196	662.6660	1984.9762	1985.0986	-0.1224	0	62	0.013	1	R.VSYPYIHENIVKPVSK.A

Proteins matching the same set of peptides:

[gi|188894298](#) Mass: 59614 Score: 178 Queries matched: 4
 phosphoenolpyruvate carboxykinase [Porphyromonas gingivalis ATCC 33277]

5. [gi|77975053](#) Mass: 59554 Score: 84 Queries matched: 3 emPAI: 0.11
 COG1866: Phosphoenolpyruvate carboxykinase (ATP) [Yersinia frederiksenii ATCC 33641]
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
151	843.3600	1684.7054	1685.7719	-1.0665	0	22	1.3e+02	2	R.LFVVDVTCGANADTR.L
195	993.4470	1984.8794	1985.0986	-0.2191	0	(59)	0.042	1	R.VSYPYIHENIVKPVSK.A
196	662.6660	1984.9762	1985.0986	-0.1224	0	62	0.013	1	R.VSYPYIHENIVKPVSK.A

Proteins matching the same set of peptides:

[gi|77976428](#) Mass: 59498 Score: 84 Queries matched: 3
 COG1866: Phosphoenolpyruvate carboxykinase (ATP) [Yersinia intermedia ATCC 29909]
[gi|123444162](#) Mass: 59550 Score: 84 Queries matched: 3
 phosphoenolpyruvate carboxykinase [Yersinia enterocolitica subsp. enterocolitica 8081]

6. [gi|223465411](#) Score: 72 Queries matched: 1
 Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
120	708.3170	1414.6194	1413.7463	0.8731	0	72	0.0014	2	R.NALLENVTVAADGK.I

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
84	502.2610	1002.5074	1002.5862	-0.0787	0	35	9.7	1	DNIFFLIR
227	737.6640	2209.9702	2211.0221	-1.0519	1	33	10	1	QILCCGCSITEAQDALMKK + Oxidation (M)
179	638.2790	1911.8152	1912.0054	-0.1902	0	32	13	1	AARHGLVPEELDALVER
210	1041.9440	2081.8734	2083.0334	-1.1600	2	31	13	1	SGYERVDIVENKGEFSVR
230	742.3140	2223.9202	2223.1834	0.7368	2	29	19	1	ELFGVMAERAKGGFIVISGR + Oxidation (M)
107	657.4030	1312.7914	1313.5935	-0.8021	0	29	32	1	AAGGGFAPVHGCCR
194	661.3090	1980.9052	1979.9193	0.9859	1	27	41	1	MPSFAAASPQIAKSCQNR + Oxidation (M)
238	1178.9000	3533.6782	3533.9259	-0.2478	2	26	56	1	ILFSLRLTPHIGPPGAFAPNVPVSRASGATVER
226	1105.9920	2209.9694	2210.0637	-0.0943	1	26	41	1	TDRGFVDSIELIAMDATAR + Oxidation (M)
58	393.3400	1176.9982	1177.5826	-0.5845	0	26	88	1	SLTEGLTSGADK
130	733.8920	2198.6542	2198.0175	0.6367	2	26	90	1	VRVYMGGGGGYDEAQRDPAR + Oxidation (M)
116	692.2780	1382.5414	1382.7994	-0.2579	1	26	51	1	LSGGNQKLIIGR
219	1068.6810	3203.0212	3202.7614	0.2597	1	26	65	1	SEAGIAVLPLSARPGIYAARIYAGIGAQVR
115	684.2770	1366.5394	1365.7140	0.8254	1	26	55	1	DFIDQSSKVVTK
43	366.2060	1095.5962	1094.4736	1.1226	0	26	1.6e+02	1	LADMRFNTR + 2 Oxidation (M)
85	506.1470	1010.2794	1010.5760	-0.2966	0	25	46	1	DLAQFLAVGK
244	905.7720	2714.2942	2715.3326	-1.0385	1	25	56	1	QTEVAIAPYDEMLFPHGLRGSAPADK
204	676.4300	2025.3682	2024.9916	0.3766	2	25	48	1	IGPGGPTDVEGDPKGGHKD
123	716.9830	2147.9272	2149.1440	-0.2168	2	25	1.3e+02	1	HDGHSIQKGLLPEYASKLR
48	381.2650	1140.7732	1140.6404	0.1328	1	25	1.8e+02	1	WVPAKLSQR
229	1112.9580	2223.9014	2224.2388	-0.3374	2	24	58	1	LPIDVLRKLVLDGIEGCK
105	651.8050	1301.5954	1300.7285	0.8669	2	24	98	1	KAGLAMAVADRAK
26	637.3450	1272.6754	1273.7176	-1.0422	1	24	1e+02	1	LLETMRLLGR + Oxidation (M)
162	897.3670	1792.7194	1793.8465	-1.1271	0	24	73	1	MTAAVAQDTLSGADAGATK + Oxidation (M)
182	965.1720	1928.3294	1928.0077	0.3217	1	24	1.3e+02	1	KQMPVYASILESIQK + Oxidation (M)
211	1042.0180	3123.0322	3123.4610	-0.4289	1	23	1.4e+02	1	WDDQAAVVASGVMTRGMVVVPCSMGTVGR
209	694.0730	2079.1972	2078.0367	1.1604	1	23	90	1	LFTRLQSAFAMLDHADR + Oxidation (M)
149	838.0810	2511.2212	2510.2660	0.9552	1	23	1.6e+02	1	MSNSPGSVFANLNGDGIKRVGVQAR + Oxidation (M)
152	853.4390	2557.2952	2558.2204	-0.9252	0	23	2.1e+02	1	TGCAVSSITMGSSSIVATALLSQR + Oxidation (M)

Mascot Search Results

Protein View

Match to: gi|53715725 Score: 663
 phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Nominal mass (M_n): 59339; Calculated pI value: 5.73
 NCBI BLAST search of gi|53715725 against nr
 Unformatted sequence string for pasting into other applications

Taxonomy: Bacteroides fragilis YCH46
 Links to retrieve other entries containing this sequence from NCBI Entrez:
 gi|60681662 from Bacteroides fragilis NCTC 9343
 gi|265767330 from Bacteroides sp. 2_1_16
 gi|60390334 from Bacteroides fragilis
 gi|81313291 from Bacteroides fragilis NCTC 9343
 gi|52218590 from Bacteroides fragilis YCH46
 gi|60495096 from Bacteroides fragilis NCTC 9343
 gi|263252635 from Bacteroides sp. 2_1_16

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 35%

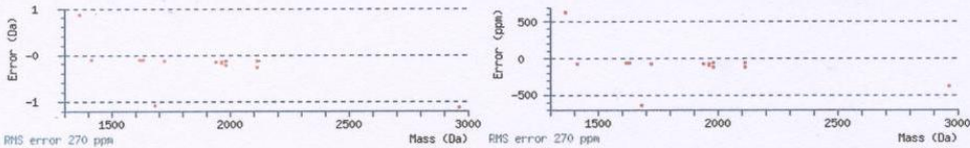
Matched peptides shown in Bold Red

1 MANLDLSKYG ITGVTEILHN PSYDVLFAEE TKPGLGFEK **QGVTELGAVN**
 51 **VMTGVYVYGR**S PKDKFFVKNE ASENSVWVTS EEYKNDNKPC SEEAWADLKA
 101 KAVKELSNKR LFFVDTFCGA NQGRMKVRF IMEVAWQAHF VTNMFIRPTA
 151 BELANYGEPD FVCFNASKAK VDNKELGLN SETATVFNLK TKEQVILNTW
 201 YGEMKKGMP SIMNYMPLR GIASHMCSAN TDMEGTSSAI FFGLSGTGKT
 251 TLSTDPKRLK IGDDEHGDWN EGVFNVEGGC YAKVINLDKE SEPDIFNAIK
 301 RDALLEENVTV AADGKINFAD KSVTENTRVS YPIYHIENIV KPVSKGPHAK
 351 QVIFLSADAF GLVPLPVSIL PEQAQYVFLS GFTAKLAGTE RGITEPTPTF
 401 SACFCGAFLS LHPTKYAEL VKKMWTGAK AYLNVTVGWSG SGRISIKDT
 451 RGIIDAILDGI SIDKAPTKVI PFFDFVVPTE LRGVDPKILD PRDLYADPAQ
 501 WNEKAKLAG RFIKNFAKFT GNEAGKKLVA AGPKL

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Sequence
41 - 59	656.6100	1966.8082	1966.9783	-0.1701	0	K.GQVTELGAVNVMTGVYVYGR.S Oxidation (M) (Ions score 46)
41 - 59	984.4190	1966.8234	1966.9783	-0.1548	0	K.GQVTELGAVNVMTGVYVYGR.S Oxidation (M) (Ions score 115)
111 - 125	843.3600	1684.7054	1685.7719	-1.0665	0	R.LFVVDVTCGANEGTR.M (Ions score 32)
176 - 190	818.3880	1634.7614	1634.8515	-0.0901	0	K.ELGLNSETATVFNLK.T (Ions score 110)
208 - 220	811.3150	1620.6154	1620.7098	-0.0944	0	K.GMFSIMNYMPLR.G 3 Oxidation (M) (Ions score 47)
221 - 249	989.3890	2965.1452	2966.2732	-1.1281	0	R.GIASHMCSANTDMEGTSSAIFFGLSGTGK.T 2 Oxidation (M) (Ions score 33)
284 - 300	972.9460	1943.8774	1944.0204	-0.1429	1	K.VINLDKESSEPDIFNAIK.R (Ions score 53)
302 - 315	708.3170	1414.6194	1414.7304	-0.1109	0	R.DALLEENVTVAADGK.I (Ions score 93)
329 - 345	993.4470	1984.8794	1985.0986	-0.2191	0	R.VSYPIYHIENIVKPVSK.G (Ions score 59)
329 - 345	662.6660	1984.9762	1985.0986	-0.1224	0	R.VSYPIYHIENIVKPVSK.G (Ions score 62)
431 - 443	684.2770	1366.5394	1365.6677	0.8717	0	K.AYLVNTGVGWSGK.R (Ions score 24)
452 - 468	863.9200	1725.8254	1725.9512	-0.1258	1	R.GIIDAILDGSIDKAPTK.V (Ions score 60)
469 - 487	705.9670	2114.8792	2115.1293	-0.2501	0	K.VIPFFDFVVPTELPVDPK.I (Ions score 16)
469 - 487	1058.5150	2115.0154	2115.1293	-0.1138	0	K.VIPFFDFVVPTELPVDPK.I (Ions score 41)



LOCUS YP_101717 535 aa linear BCT 26-APR-2009
 DEFINITION phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46].
 ACCESSION YP_101717
 VERSION YP_101717.1 GI:53715725
 DBLINK Project:13067
 DBSOURCE REFSEQ: accession NC_006347.1
 KEYWORDS .
 SOURCE Bacteroides fragilis YCH46
 ORGANISM Bacteroides fragilis YCH46
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 535)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 535)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 535)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan

COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final NCBI review. The reference sequence was derived from BAD51183. Method: conceptual translation.

FEATURES Location/Qualifiers

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 /calculated_mol_wt=58897

Region 15..532
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Site order(207,226,250,263..264,281)
 /site_type="metal-binding"
 /db_xref="CDD:29830"

Site order(245..251,292,444,450)
 /site_type="other"
 /note="ATP binding site"
 /db_xref="CDD:29830"

CDS 1..535
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 /coded_by="NC_006347.1:5088646..5090253"
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 /transl_table=11
 /db_xref="GeneID:3085522"

Mascot: <http://www.matrixscience.com/>

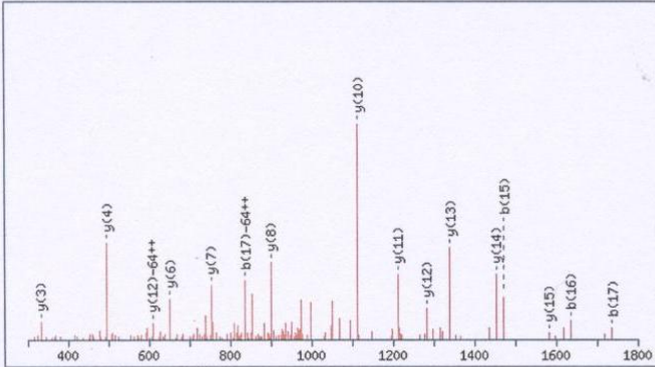
MASCOT Mascot Search Results

Peptide View

MS/MS Fragmentation of **GQVTELGAVNVMTGVYTR**
 Found in **gi|53715725**, phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

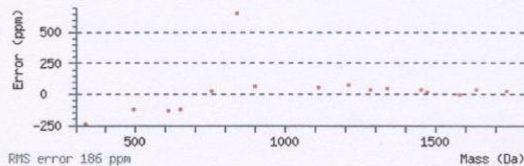
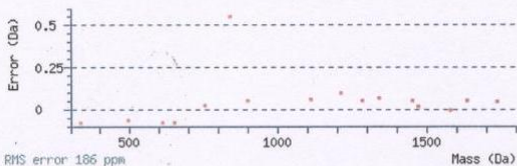
Match to Query 191: 1966.823448 from(984.419000,2+) intensity(3902720.0000)
 Title: Cmpd 36, +MSn(984.70), 6.6 min
 Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Click mouse within plot area to zoom in by factor of two about that point
 Or, Plot from to Da



Monoisotopic mass of neutral peptide $M_r(\text{calc})$: 1966.9783
 Fixed modifications: Carboxymethyl (C)
 Variable modifications:
 M12 : Oxidation (M), with neutral losses 0.0000(shown in table), 63.9983
 Ions Score: 115 Expect: 5.2e-08
 Matches (Bold Red): 16/312 fragment ions using 18 most intense peaks

#	a	a ⁺⁺	a*	a ⁺⁺⁺	b	b ⁺⁺	b*	b ⁺⁺⁺	Seq.	y	y ⁺⁺	y*	y ⁺⁺⁺	#
1	30.0338	15.5206			58.0287	29.5180			G					19
2	158.0924	79.5498	141.0659	71.0366	186.0873	93.5473	169.0608	85.0340	Q	1910.9640	955.9857	1893.9375	947.4724	18
3	257.1608	129.0840	240.1343	120.5708	285.1557	143.0815	268.1292	134.5682	V	1782.9055	891.9564	1765.8789	883.4431	17
4	358.2085	179.6079	341.1819	171.0946	386.2034	193.6053	369.1769	185.0921	T	1683.8371	842.4222	1666.8105	833.9089	16
5	487.2511	244.1292	470.2245	235.6159	515.2460	258.1266	498.2195	249.6134	E	1582.7894	791.8983	1565.7628	783.3851	15
6	600.3352	300.6712	583.3086	292.1579	628.3301	314.6687	611.3035	306.1554	L	1453.7468	727.3770	1436.7202	718.8638	14
7	657.3566	329.1819	640.3301	320.6687	685.3515	343.1794	668.3250	334.6661	G	1340.6627	670.8350	1323.6362	662.3217	13
8	728.3937	364.7005	711.3672	356.1872	756.3886	378.6980	739.3621	370.1847	A	1283.6413	642.3243	1266.6147	633.8110	12
9	827.4621	414.2347	810.4356	405.7214	855.4571	428.2322	838.4305	419.7189	V	1212.6041	606.8057	1195.5776	598.2924	11
10	941.5051	471.2562	924.4785	462.7429	969.5000	485.2536	952.4734	476.7404	N	1113.5357	557.2715	1096.5092	548.7582	10
11	1040.5735	520.7904	1023.5469	512.2771	1068.5684	534.7878	1051.5419	526.2746	V	999.4928	500.2500	982.4662	491.7368	9
12	1187.6089	594.3081	1170.5823	585.7948	1215.6038	608.3055	1198.5773	599.7923	M	900.4244	450.7158	883.3978	442.2026	8
13	1288.6566	644.8319	1271.6300	636.3186	1316.6515	658.8294	1299.6249	650.3161	T	753.3890	377.1981	736.3624	368.6849	7
14	1345.6780	673.3427	1328.6515	664.8294	1373.6729	687.3401	1356.6464	678.8268	G	652.3413	326.6743	635.3148	318.1610	6
15	1444.7464	722.8769	1427.7199	714.3636	1472.7414	736.8743	1455.7148	728.3610	V	595.3198	298.1636	578.2933	289.6503	5
16	1607.8098	804.4085	1590.7832	795.8953	1635.8047	818.4060	1618.7781	809.8927	Y	496.2514	248.6293	479.2249	240.1161	4
17	1708.8575	854.9324	1691.8309	846.4191	1736.8524	868.9298	1719.8258	860.4165	T	333.1881	167.0977	316.1615	158.5844	3
18	1765.8789	883.4431	1748.8524	874.9298	1793.8738	897.4406	1776.8473	888.9273	G	232.1404	116.5738	215.1139	108.0606	2
19									R	175.1190	88.0631	158.0924	79.5498	1



NCBI **BLAST** search of GQVTELGAVNVMTGVYTGR
(Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
115.5	1966.9783	-0.1548	GQVTELGAVNVMTGVYTGR
27.0	1966.0054	0.8180	SAEMAARAAEHVAGLPTLR
24.8	1965.9843	0.8391	AFNRSQPNMDVAGLFKR
18.9	1966.0861	0.7374	VVDAERALARFSGVALHR
18.5	1967.1350	-0.3115	AAEKERLMVLVAGLNVVR
17.8	1966.1211	0.7024	SLSPELKVTPEGRATLLR
15.6	1967.9160	-1.0925	MGQYNFDQIVDRTHTK
14.5	1965.9948	0.8286	VYTGKADWYAAGLPLER
14.4	1967.0952	-0.2718	AREVVVELVQAVAEAVGPR
13.6	1967.0298	-0.2064	MNKHPEFLLNKIENPK

Mascot: <http://www.matrixscience.com/>

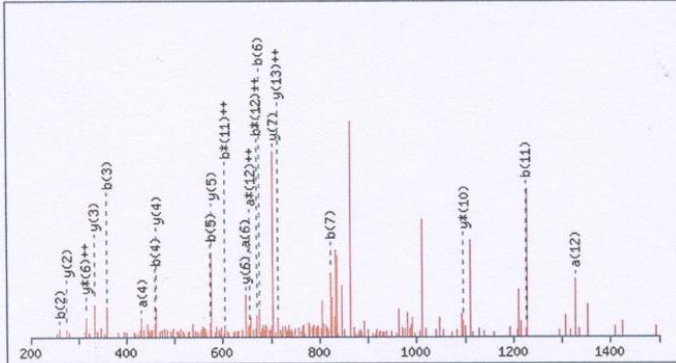
MASCOT Mascot Search Results

Peptide View

MS/MS Fragmentation of **LFVVDIFCGANEGTR**
 Found in [gi|53715725](#), phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

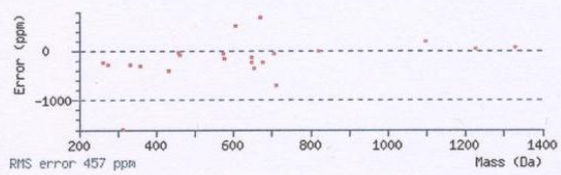
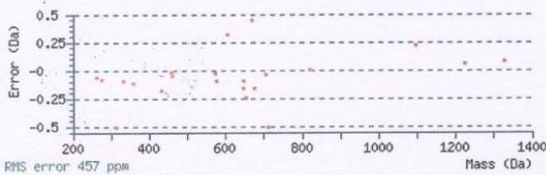
Match to Query 151: 1684.705448 from(843.360000,2+) intensity(5734853.0000)
 Title: Cmpd 46, +MSn(843.76), 6.9 min
 Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Click mouse within plot area to zoom in by factor of two about that point
 Or, Plot from to Da



Monoisotopic mass of neutral peptide $M_r(\text{calc})$: 1685.7719
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 32 Expect: 13
 Matches (Bold Red): 22/128 Fragment ions using 61 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{*++}	b	b ⁺⁺	b [*]	b ^{*++}	Seq.	y	y ⁺⁺	y [*]	y ^{*++}	#
1	86.0964	43.5519			114.0913	57.5493			L					15
2	233.1648	117.0861			261.1598	131.0835			F	1573.6951	787.3512	1556.6686	778.8379	14
3	332.2333	166.6203			360.2282	180.6177			V	1426.6267	713.8170	1409.6002	705.3037	13
4	431.3017	216.1545			459.2966	230.1519			V	1327.5583	664.2828	1310.5318	655.7695	12
5	546.3286	273.6679			574.3235	287.6654			D	1228.4899	614.7486	1211.4633	606.2353	11
6	647.3763	324.1918			675.3712	338.1892			T	1113.4630	557.2351	1096.4364	548.7218	10
7	794.4447	397.7260			822.4396	411.7234			F	1012.4153	506.7113	995.3887	498.1980	9
8	955.4594	478.2333			983.4543	492.2308			C	865.3469	433.1771	848.3203	424.6638	8
9	1012.4808	506.7441			1040.4757	520.7415			G	704.3322	352.6697	687.3056	344.1565	7
10	1083.5179	542.2626			1111.5129	556.2601			A	647.3107	324.1590	630.2842	315.6457	6
11	1197.5609	599.2841	1180.5343	590.7708	1225.5558	613.2815	1208.5292	604.7683	N	576.2736	288.6404	559.2471	280.1272	5
12	1326.6035	663.8054	1309.5769	655.2921	1354.5984	677.8028	1337.5718	669.2896	E	462.2307	231.6190	445.2041	223.1057	4
13	1383.6249	692.3161	1366.5984	683.8028	1411.6198	706.3136	1394.5933	697.8003	G	333.1881	167.0977	316.1615	158.5844	3
14	1484.6726	742.8399	1467.6461	734.3267	1512.6675	756.8374	1495.6410	748.3241	T	276.1666	138.5870	259.1401	130.0737	2
15									R	175.1190	88.0631	158.0924	79.5498	1



NCBI BLAST search of **LFVVDIFCGANEGTR**
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
32.2	1685.7719	-1.0665	LFVVDTFCGANEGTR
22.2	1685.7719	-1.0665	LFVVDTFCGANADTR
21.9	1683.9083	0.7971	FEQVSKIYSTDVVLK
21.4	1685.8083	-1.1029	NFVFDERMSEVVSK
19.4	1685.8413	-1.1359	VIFDQFNFTDIAR
18.6	1685.7865	-1.0810	ACQIMAGHLKENADK
17.4	1684.8056	-0.1002	GNSLSDTYIEVFANR
16.9	1684.7978	-0.0924	MIVEGTGAYTTVNGEK
16.4	1683.8502	0.8553	AENMKTTVNSFVLK
16.3	1685.8698	-1.1644	KALFDYEMLKDVSK

Mascot: <http://www.matrixscience.com/>

Mascot Search Results

Peptide View

MS/MS Fragmentation of **ELGLNSETATVFNLK**

Found in [gi|53715725](#), phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

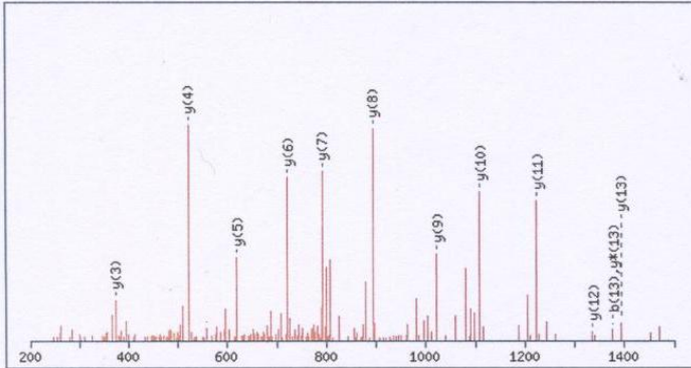
Match to Query 142: 1634.761448 from(818.388000,2+) intensity(10020720.0000)

Title: Cmpd 60, +MSn(818.83), 7.6 min

Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from to Da



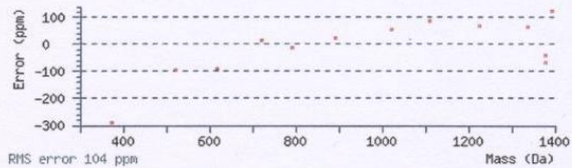
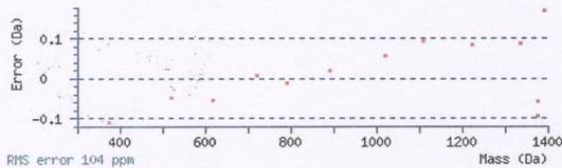
Monoisotopic mass of neutral peptide Mr(calc): 1634.8515

Fixed modifications: Carboxymethyl (C)

Ions Score: 110 Expect: 2.1e-07

Matches (Bold Red): 13/152 fragment ions using 16 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{*++}	b	b ⁺⁺	b [*]	b ^{*++}	Seq.	y	y ⁺⁺	y [*]	y ^{*++}	#
1	102.0550	51.5311			130.0499	65.5286			E					15
2	215.1390	108.0731			243.1339	122.0706			L	1506.8162	753.9118	1489.7897	745.3985	14
3	272.1605	136.5839			300.1554	150.5813			G	1393.7322	697.3697	1376.7056	688.8564	13
4	385.2445	193.1259			413.2395	207.1234			L	1336.7107	668.8590	1319.6842	660.3457	12
5	499.2875	250.1474	482.2609	241.6341	527.2824	264.1448	510.2558	255.6316	N	1223.6266	612.3170	1206.6001	603.8037	11
6	586.3195	293.6634	569.2930	285.1501	614.3144	307.6608	597.2879	299.1476	S	1109.5837	555.2955	1092.5572	546.7822	10
7	715.3621	358.1847	698.3355	349.6714	743.3570	372.1821	726.3305	363.6689	E	1022.5517	511.7795	1005.5251	503.2662	9
8	816.4098	408.7085	799.3832	400.1953	844.4047	422.7060	827.3781	414.1927	T	893.5091	447.2582	876.4825	438.7449	8
9	887.4469	444.2271	870.4203	435.7138	915.4418	458.2245	898.4153	449.7113	A	792.4614	396.7343	775.4349	388.2211	7
10	988.4946	494.7509	971.4680	486.2376	1016.4895	508.7484	999.4629	500.2351	T	721.4243	361.2158	704.3978	352.7025	6
11	1087.5630	544.2851	1070.5364	535.7719	1115.5579	558.2826	1098.5313	549.7693	V	620.3766	310.6919	603.3501	302.1787	5
12	1234.6314	617.8193	1217.6048	609.3061	1262.6263	631.8168	1245.5998	623.3035	F	521.3082	261.1577	504.2817	252.6445	4
13	1348.6743	674.8408	1331.6478	666.3275	1376.6692	688.8383	1359.6427	680.3250	N	374.2398	187.6235	357.2132	179.1103	3
14	1461.7584	731.3828	1444.7318	722.8696	1489.7533	745.3803	1472.7268	736.8670	L	260.1969	130.6021	243.1703	122.0888	2
15									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of **ELGLNSETATVFNLK**

(Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)

Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
110.3	1634.8515	-0.0901	ELGLNSETATVFNLK
22.3	1635.8688	-1.1073	IGLSEMLTALSMKAR
20.5	1635.8356	-1.0741	EYVEVTAETVEVIR
18.9	1633.8522	0.9092	TELEIRTATLSASDK
18.6	1633.6937	1.0678	AAVGARDTCNPCNAR
18.0	1634.8562	-0.0948	FNDDKTAMVRALVR
18.0	1633.8610	0.9005	NLNIALFGATGMIGSR
18.0	1634.9178	-0.1563	TLKIALFGATGMIGSR
17.8	1635.7425	-0.9811	EMVFITWDSHMPK
16.9	1634.8562	-0.0948	KNGLGGFSYRQLCAK

Mascot: <http://www.matrixscience.com/>

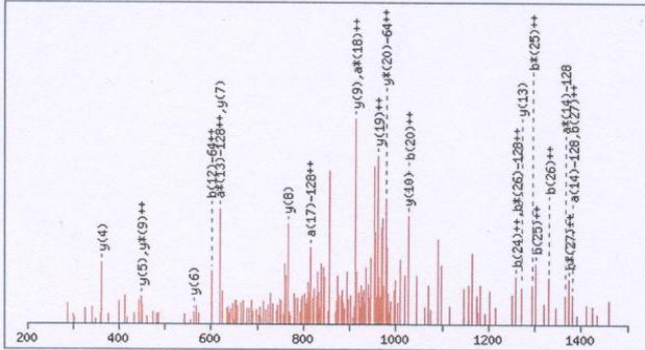
Mascot Search Results

Peptide View

MS/MS Fragmentation of **GIASMHCSANTDMEGTSSAIFFLSGTQK**
 Found in **gij53715725**, phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

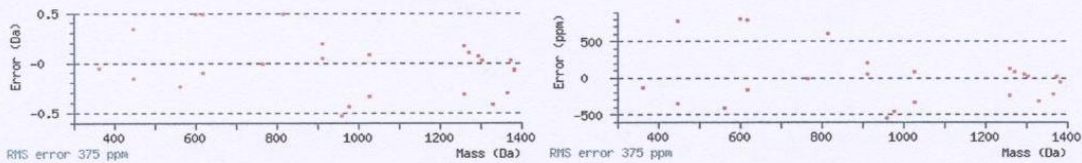
Match to Query 250: 2965.145172 from(989.389000,3+) intensity(3604278.0000)
 Title: Cmpd 55, +MSn(989.77), 7.4 min
 Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Click mouse within plot area to zoom in by factor of two about that point
 Or, to Da



Monoisotopic mass of neutral peptide **Mr(calc)**: 2966.2732
 Fixed modifications: Carboxymethyl (C)
 Variable modifications:
 M5 : Oxidation (M), with neutral losses 0.0000 (shown in table), 63.9983
 M13 : Oxidation (M), with neutral losses 0.0000 (shown in table), 63.9983
 Ions Score: 33 Expect: 6
 Matches (Bold Red): 25/520 Fragment ions using 30 most intense peaks

#	a	a ⁺⁺	a*	a ⁺⁺⁺	b	b ⁺⁺	b*	b ⁺⁺⁺	Seq.	y	y ⁺⁺	y*	y ⁺⁺⁺	#
1	30.0338	15.5206			58.0287	29.5180			G					29
2	143.1179	72.0626			171.1128	86.0600			I	2910.2590	1455.6332	2893.2325	1447.1199	28
3	214.1550	107.5811			242.1499	121.5786			A	2797.1750	1399.0911	2780.1484	1390.5779	27
4	301.1870	151.0972			329.1819	165.0946			S	2726.1379	1363.5726	2709.1113	1355.0593	26
5	448.2224	224.6149			476.2173	238.6123			M	2639.1058	1320.0566	2622.0793	1311.5433	25
6	585.2813	293.1443			613.2763	307.1418			H	2492.0704	1246.5389	2475.0439	1238.0256	24
7	746.2960	373.6516			774.2909	387.6491			C	2355.0115	1178.0094	2337.9850	1169.4961	23
8	833.3280	417.1677			861.3230	431.1651			S	2193.9969	1097.5021	2176.9703	1088.9888	22
9	904.3652	452.6862			932.3601	466.6837			A	2106.9648	1053.9861	2089.9383	1045.4728	21
10	1018.4081	509.7077	1001.3815	501.1944	1046.4030	523.7051	1029.3764	515.1919	N	2035.9277	1018.4675	2018.9012	1009.9542	20
11	1119.4558	560.2315	1102.4292	551.7182	1147.4507	574.2290	1130.4241	565.7157	T	1921.8848	961.4460	1904.8582	952.9328	19
12	1234.4827	617.7450	1217.4562	609.2317	1262.4776	631.7424	1245.4511	623.2292	D	1820.8371	910.9222	1803.8106	902.4089	18
13	1381.5181	691.2627	1364.4916	682.7494	1409.5130	705.2601	1392.4865	696.7469	M	1705.8102	853.4087	1688.7836	844.8954	17
14	1510.5607	755.7840	1493.5342	747.2707	1538.5556	769.7814	1521.5291	761.2682	E	1558.7748	779.8910	1541.7482	771.3777	16
15	1567.5822	784.2947	1550.5556	775.7814	1595.5771	798.2922	1578.5505	789.7789	G	1429.7322	715.3697	1412.7056	706.8564	15
16	1668.6298	834.8186	1651.6033	826.3053	1696.6248	848.8160	1679.5982	840.3027	T	1372.7107	686.8590	1355.6842	678.3457	14
17	1755.6619	878.3346	1738.6353	869.8213	1783.6568	892.3320	1766.6302	883.8188	S	1271.6630	636.3352	1254.6365	627.8219	13
18	1842.6939	921.8506	1825.6673	913.3373	1870.6888	935.8480	1853.6623	927.3348	S	1184.6310	592.8191	1167.6045	584.3059	12
19	1913.7310	957.3691	1896.7045	948.8559	1941.7259	971.3666	1924.6994	962.8533	A	1097.5990	549.3031	1080.5724	540.7898	11
20	2026.8151	1013.9112	2009.7885	1005.3979	2054.8100	1027.9086	2037.7834	1019.3954	I	1026.5619	513.7846	1009.5353	505.2713	10
21	2173.8835	1087.4454	2156.8569	1078.9321	2201.8784	1101.4428	2184.8519	1092.9296	F	913.4778	457.2425	896.4512	448.7293	9
22	2320.9519	1160.9796	2303.9254	1152.4663	2348.9468	1174.9770	2331.9203	1166.4638	F	766.4094	383.7083	749.3828	375.1951	8
23	2377.9734	1189.4903	2360.9468	1180.9770	2405.9683	1203.4878	2388.9417	1194.9745	G	619.3410	310.1741	602.3144	301.6608	7
24	2491.0574	1246.0324	2474.0309	1237.5191	2519.0523	1260.0298	2502.0258	1251.5165	L	562.3195	281.6634	545.2930	273.1501	6
25	2578.0895	1289.5484	2561.0629	1281.0351	2606.0844	1303.5458	2589.0578	1295.0326	S	449.2354	225.1214	432.2089	216.6081	5
26	2635.1109	1318.0591	2618.0844	1309.5458	2663.1058	1332.0566	2646.0793	1323.5433	G	362.2034	181.6053	345.1769	173.0921	4
27	2736.1586	1368.5829	2719.1321	1360.0697	2764.1535	1382.5804	2747.1270	1374.0671	T	305.1819	153.0946	288.1554	144.5813	3
28	2793.1801	1397.0937	2776.1535	1388.5804	2821.1750	1411.0911	2804.1484	1402.5779	G	204.1343	102.5708	187.1077	94.0575	2
29									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of [GIASMHC SANTDMEGTSSAIFFLSGTGK](#)
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc)	Delta	Sequence
32.7	2966.2732	-1.1281	GIASMHC SANTDMEGTSSAIFFLSGTGK
32.7	2965.2892	-0.1440	GIASMHC SANTNMEGTSSAIFFLSGTGK
32.0	2965.2416	-0.0965	DVMSMHC SANVGEDGDTTLFFFLSGTGK
23.5	2965.3586	-0.2135	ADVLPMHCAANYDDAGNTALFFFLSGTGK
21.3	2964.4916	0.6535	RGVFPMHASANVGKEGDVALFFFLSGTGK
20.8	2964.5266	0.6186	LNLTAPVTPEQKINYISIAMAGYSADK
18.6	2966.3273	-1.1821	LVEVMENYIASLDEEALAGMAGSSNSH
17.3	2964.3900	0.7551	NEFFLENLIFFCKFVKMSDMQK
17.1	2964.4949	0.6502	SEHQMAMLDOYAGHLNLLKTHVSLTK
16.9	2964.6331	0.5121	IAIHHAEGMLQIGEIPVIITASSKHRK

Mascot: <http://www.matrixscience.com/>

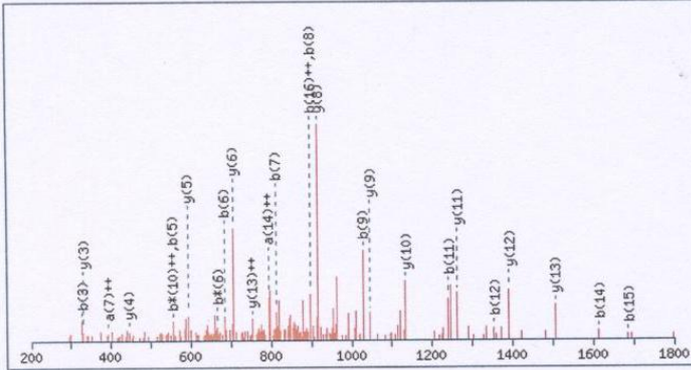
Mascot Search Results

Peptide View

MS/MS Fragmentation of **VINLDKESEPDIFNAIK**
 Found in **gi|53715725**, phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

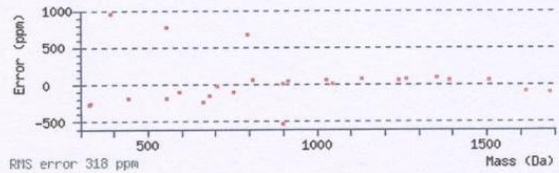
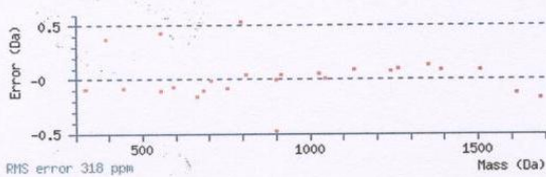
Match to Query 186: 1943.877448 from(972.946000,2+) intensity(3662039.0000)
 Title: Cmpd 62, +MSn(973.22), 7.7 min
 Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Click mouse within plot area to zoom in by factor of two about that point
 Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 1944.0204
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 53 Expect: 0.12
 Matches (bold red): 26/184 fragment ions using 52 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{*++}	b	b ⁺⁺	b [*]	b ^{*++}	Seq.	y	y ⁺⁺	y [*]	y ^{*++}	#
1	72.0808	36.5440			100.0757	50.5415			V					17
2	185.1648	93.0861			213.1598	107.0835			I	1845.9593	923.4833	1828.9327	914.9700	16
3	299.2078	150.1075	282.1812	141.5942	327.2027	164.1050	310.1761	155.5917	N	1732.8752	866.9412	1715.8487	858.4280	15
4	412.2918	206.6496	395.2653	198.1363	440.2867	220.6470	423.2602	212.1337	L	1618.8323	809.9198	1601.8057	801.4065	14
5	527.3188	264.1630	510.2922	255.6498	555.3137	278.1605	538.2871	269.6472	D	1505.7482	753.3777	1488.7217	744.8645	13
6	655.4137	328.2105	638.3872	319.6972	683.4087	342.2080	666.3821	333.6947	K	1390.7213	695.8643	1373.6947	687.3510	12
7	784.4563	392.7318	767.4298	384.2185	812.4512	406.7293	795.4247	398.2160	E	1262.6263	631.8168	1245.5998	623.3035	11
8	871.4884	436.2478	854.4618	427.7345	899.4833	450.2453	882.4567	441.7320	S	1133.5837	567.2955	1116.5572	558.7822	10
9	1000.5309	500.7691	983.5044	492.2558	1028.5259	514.7666	1011.4993	506.2533	E	1046.5517	523.7795	1029.5251	515.2662	9
10	1097.5837	549.2955	1080.5572	540.7822	1125.5786	563.2930	1108.5521	554.7797	P	917.5091	459.2582	900.4825	450.7449	8
11	1212.6107	606.8090	1195.5841	598.2957	1240.6056	620.8064	1223.5790	612.2931	D	820.4563	410.7318	803.4298	402.2185	7
12	1325.6947	663.3510	1308.6682	654.8377	1353.6896	677.3485	1336.6631	668.8352	I	705.4294	353.2183	688.4028	344.7051	6
13	1472.7631	736.8852	1455.7366	728.3719	1500.7581	750.8827	1483.7315	742.3694	F	592.3453	296.6763	575.3188	288.1630	5
14	1586.8061	793.9067	1569.7795	785.3934	1614.8010	807.9041	1597.7744	799.3909	N	445.2769	223.1421	428.2504	214.6288	4
15	1657.8432	829.4252	1640.8166	820.9120	1685.8381	843.4227	1668.8115	834.9094	A	331.2340	166.1206	314.2074	157.6074	3
16	1770.9272	885.9673	1753.9007	877.4540	1798.9222	899.9647	1781.8956	891.4514	I	260.1969	130.6021	243.1703	122.0888	2
17									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of **VINLDKESEPDIFNAIK**
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
52.5	1944.0204	-0.1429	VINLDKESEPDIFNAIK
26.7	1944.0204	-0.1429	VINLDKDSEPEIFNAIK
16.6	1944.0713	-0.1939	IRVADKETALEALQMLK
10.9	1942.9095	0.9680	EGGIDKEGIYIDENAMR
9.7	1945.0417	-1.1642	ILMSMPITPGPALLAFNK
9.7	1943.9839	-0.1065	KYAQAEEPEESPLAAALK
8.6	1943.8935	-0.0160	NATSPFEATGSLYPMAGSK
8.2	1943.9921	-0.1146	RQAAQVMESGVVAGGMIPK
7.9	1943.8275	0.0500	SDTYTMPIDEMREAMAK
7.8	1943.9921	-0.1146	KIPICGQDDVIIRNMR

Mascot: <http://www.matrixscience.com/>

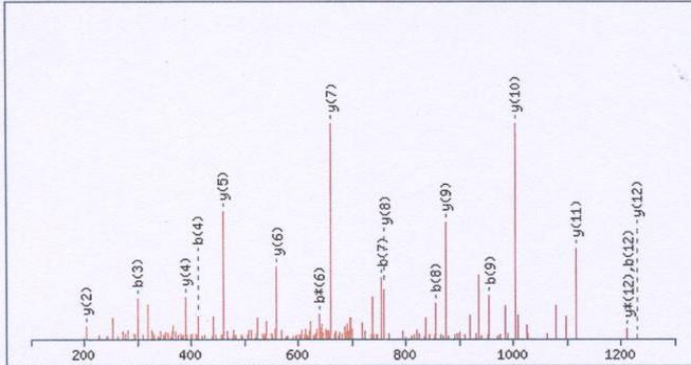
MASCOT **SCIENCE** Mascot Search Results

Peptide View

MS/MS Fragmentation of **DALLENVTVAADGK**
 Found in **gi53715725**, phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

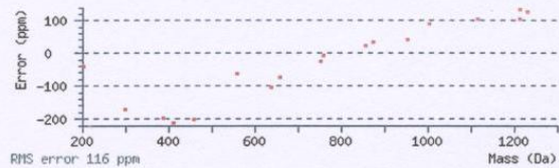
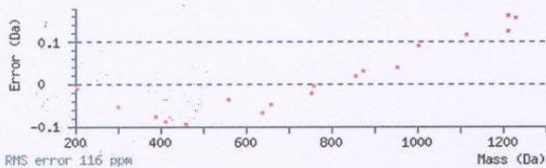
Match to Query 120: 1414.619448 from(708.317000,2+) intensity(4539975.0000)
 Title: Cmpd 11, +MSn(708.71), 5.6 min
 Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Click mouse within plot area to zoom in by factor of two about that point
 Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 1414.7304
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 93 Expect: 1.2e-05
 Matches (Bold Red): 18/136 fragment ions using 21 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{*++}	b	b ⁺⁺	b [*]	b ^{*++}	Seq.	y	y ⁺⁺	y [*]	y ^{*++}	#
1	88.0393	44.5233			116.0342	58.5207			D					14
2	159.0764	80.0418			187.0713	94.0393			A	1300.7107	650.8590	1283.6842	642.3457	13
3	272.1605	136.5839			300.1554	150.5813			L	1229.6736	615.3404	1212.6470	606.8272	12
4	385.2445	193.1259			413.2395	207.1234			L	1116.5895	558.7984	1099.5630	550.2851	11
5	514.2871	257.6472			542.2821	271.6447			E	1003.5055	502.2564	986.4789	493.7431	10
6	628.3301	314.6687	611.3035	306.1554	656.3250	328.6661	639.2984	320.1529	N	874.4629	437.7351	857.4363	429.2218	9
7	727.3985	364.2029	710.3719	355.6896	755.3934	378.2003	738.3668	369.6871	V	760.4199	380.7136	743.3934	372.2003	8
8	828.4462	414.7267	811.4196	406.2134	856.4411	428.7242	839.4145	420.2109	T	661.3515	331.1794	644.3250	322.6661	7
9	927.5146	464.2609	910.4880	455.7476	955.5095	478.2584	938.4829	469.7451	V	560.3039	280.6556	543.2773	272.1423	6
10	998.5517	499.7795	981.5251	491.2662	1026.5466	513.7769	1009.5201	505.2637	A	461.2354	231.1214	444.2089	222.6081	5
11	1069.5888	535.2980	1052.5623	526.7848	1097.5837	549.2955	1080.5572	540.7822	A	390.1983	195.6028	373.1718	187.0895	4
12	1184.6157	592.8115	1167.5892	584.2982	1212.6107	606.8090	1195.5841	598.2957	D	319.1612	160.0842	302.1347	151.5710	3
13	1241.6372	621.3222	1224.6107	612.8090	1269.6321	635.3197	1252.6056	626.8064	G	204.1343	102.5708	187.1077	94.0575	2
14									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of **DALLENVTVAADGK**
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

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Score	Mr(calc):	Delta	Sequence
93.1	1414.7304	-0.1109	DALLENVTVAADGK
72.4	1413.7463	0.8731	NALLENVTVAADGK
58.0	1413.7463	0.8731	NALLENVTVDAAGK
30.1	1414.7303	-0.1109	GIELENIDAADKK
25.9	1414.7164	-0.0970	GNQIEGRISEGQK
24.8	1414.7668	-0.1473	VTVIDGVASLAENK
24.0	1414.6875	-0.0680	DNGMGISYQELPR
22.0	1414.7667	-0.1473	EGLLVENLSVSQK
22.0	1414.6762	-0.0568	GELMQEAVPAGEGK
18.9	1414.8395	-0.2201	VSHVGVSKDLTGK

Mascot: <http://www.matrixscience.com/>

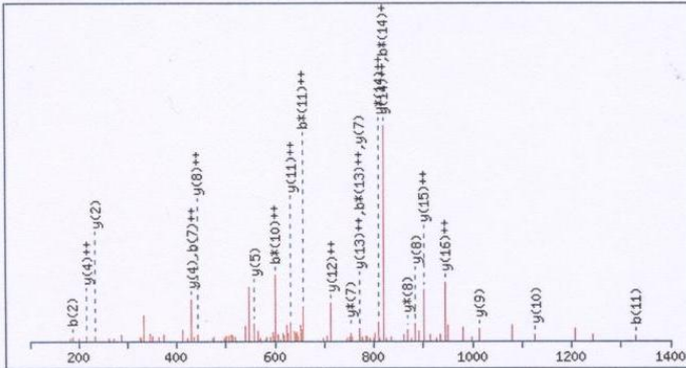
MASCOT Mascot Search Results

Peptide View

MS/MS Fragmentation of **VSYPIYHIENIVKPVSK**
 Found in **gi|53715725**, phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

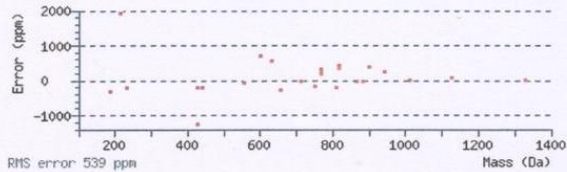
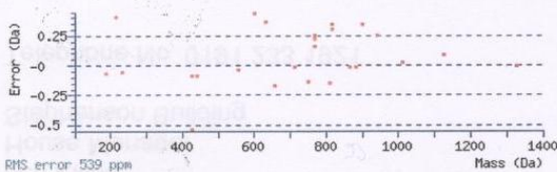
Match to Query 196: 1984.976172 from(662.666000,3+) intensity(4638027.0000)
 Title: Cmpd 53, +MSn(663.74), 7.3 min
 Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Click mouse within plot area to zoom in by factor of two about that point
 Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 1985.0986
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 62 Expect: 0.013
 Matches (Bold Red): 25/156 fragment ions using 35 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{***}	b	b ⁺⁺	b [*]	b ^{***}	Seq.	y	y ⁺⁺	y [*]	y ^{***}	#
1	72.0808	36.5440			100.0757	50.5415			V					17
2	159.1128	80.0600			187.1077	94.0575			S	1887.0375	944.0224	1870.0109	935.5091	16
3	322.1761	161.5917			350.1710	175.5892			Y	1800.0054	900.5064	1782.9789	891.9931	15
4	419.2289	210.1181			447.2238	224.1155			P	1636.9421	818.9747	1619.9156	810.4614	14
5	532.3130	266.6601			560.3079	280.6576			I	1539.8893	770.4483	1522.8628	761.9350	13
6	695.3763	348.1918			723.3712	362.1892			Y	1426.8053	713.9063	1409.7787	705.3930	12
7	832.4352	416.7212			860.4301	430.7187			H	1263.7419	632.3746	1246.7154	623.8613	11
8	945.5193	473.2633			973.5142	487.2607			I	1126.6830	563.8452	1109.6565	555.3319	10
9	1074.5619	537.7846			1102.5568	551.7820			E	1013.5990	507.3031	996.5724	498.7898	9
10	1188.6048	594.8060	1171.5782	586.2928	1216.5997	608.8035	1199.5732	600.2902	N	884.5564	442.7818	867.5298	434.2686	8
11	1301.6888	651.3481	1284.6623	642.8348	1329.6838	665.3455	1312.6572	656.8322	I	770.5135	385.7604	753.4869	377.2471	7
12	1400.7573	700.8823	1383.7307	692.3690	1428.7522	714.8797	1411.7256	706.3665	V	657.4294	329.2183	640.4028	320.7051	6
13	1528.8522	764.9298	1511.8257	756.4165	1556.8471	778.9272	1539.8206	770.4139	K	558.3610	279.6841	541.3344	271.1709	5
14	1625.9050	813.4561	1608.8784	804.9429	1653.8999	827.4536	1636.8734	818.9403	P	430.2660	215.6366	413.2395	207.1234	4
15	1724.9734	862.9903	1707.9469	854.4771	1752.9683	876.9878	1735.9418	868.4745	V	333.2132	167.1103	316.1867	158.5970	3
16	1812.0054	906.5064	1794.9789	897.9931	1840.0003	920.5038	1822.9738	911.9905	S	234.1448	117.5761	217.1183	109.0628	2
17									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of **VSYPIYHIENIVKPVSK**
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
62.3	1985.0986	-0.1224	VSYPIYHIENIVKPVSK
32.6	1985.0986	-0.1224	VSYPIYHIDNIVKPISK
24.3	1985.1018	-0.1256	VDQSRASRTGGTGLGLAIVK
24.1	1984.0993	0.8769	HDIPLLADISAALKPKAD
22.8	1984.1146	0.8616	VSYPIYHIQNIVKPVSK
22.8	1984.0782	0.8980	VSYPIYHIQNIVQPVS
22.6	1986.0157	-1.0395	VTLEDDNILINGEKVDSL
21.2	1986.0291	-1.0530	YIGNIPVSMHAVMGAVRR
20.7	1985.2481	-0.2720	WFIGVAIHLIIFINK
20.1	1985.0080	-0.0319	SMNLGPLFEPFELHNLK

Mascot: <http://www.matrixscience.com/>

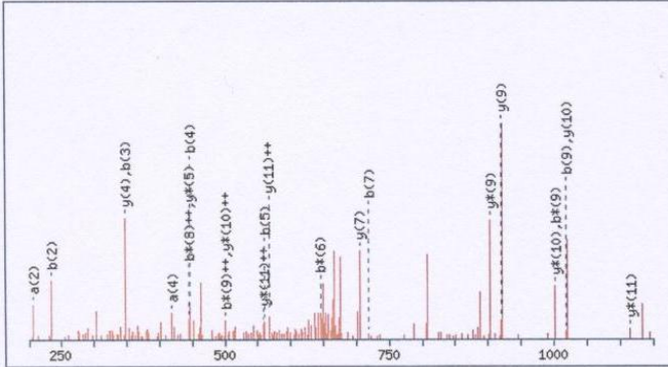
MASCOT Mascot Search Results

Peptide View

MS/MS Fragmentation of **AYLVNTGWNGSGK**
 Found in **gi53715725**, phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

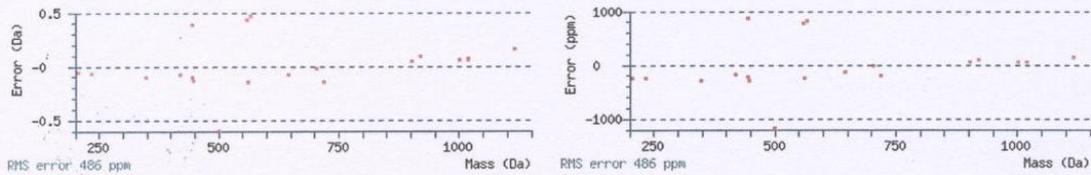
Match to Query 115: 1366.539448 from(684.277000,2+) intensity(1224761.0000)
 Title: Cmpd 4, +MSn(684.95), 5.3 min
 Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Click mouse within plot area to zoom in by factor of two about that point
 Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 1365.6677
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 24 Expect: 74
 Matches (Bold Red): 23/128 fragment ions using 42 most intense peaks

#	a	a ⁺⁺	a*	a ⁺⁺⁺	b	b ⁺⁺	b*	b ⁺⁺⁺	Seq.	y	y ⁺⁺	y*	y ⁺⁺⁺	#
1	44.0495	22.5284			72.0444	36.5258			A					13
2	207.1128	104.0600			235.1077	118.0575			Y	1295.6379	648.3226	1278.6113	639.8093	12
3	320.1969	160.6021			348.1918	174.5995			L	1132.5745	566.7909	1115.5480	558.2776	11
4	419.2653	210.1363			447.2602	224.1337			V	1019.4905	510.2489	1002.4639	501.7356	10
5	533.3082	267.1577	516.2817	258.6445	561.3031	281.1552	544.2766	272.6419	N	920.4221	460.7147	903.3955	452.2014	9
6	634.3559	317.6816	617.3293	309.1683	662.3508	331.6790	645.3243	323.1658	T	806.3791	403.6932	789.3526	395.1799	8
7	691.3774	346.1923	674.3508	337.6790	719.3723	360.1898	702.3457	351.6765	G	705.3315	353.1694	688.3049	344.6561	7
8	877.4567	439.2320	860.4301	430.7187	905.4516	453.2294	888.4250	444.7162	W	648.3100	324.6586	631.2835	316.1454	6
9	991.4996	496.2534	974.4730	487.7402	1019.4945	510.2509	1002.4680	501.7376	N	462.2307	231.6190	445.2041	223.1057	5
10	1048.5211	524.7642	1031.4945	516.2509	1076.5160	538.7616	1059.4894	530.2483	G	348.1878	174.5975	331.1612	166.0842	4
11	1135.5531	568.2802	1118.5265	559.7669	1163.5480	582.2776	1146.5215	573.7644	S	291.1663	146.0868	274.1397	137.5735	3
12	1192.5745	596.7909	1175.5480	588.2776	1220.5695	610.7884	1203.5429	602.2751	G	204.1343	102.5708	187.1077	94.0575	2
13									K	147.1128	74.0600	130.0863	65.5468	1



NCBI BLAST search of **AYLVNTGWNGSGK**
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence

25.7	1365.7140	0.8254	DFIDQSSKVVTK
25.3	1365.7504	0.7891	YVTVSSPSKALSK
24.4	1365.6677	0.8717	AYLVNTGWNGSGK
23.8	1365.6525	0.8870	FSIVDRGDENSK
19.5	1366.7391	-0.1996	MSIDSRRFVIK
18.9	1366.7642	-0.2248	YALLKLVGTGSMR
18.9	1366.6299	-0.0905	FSPDSCDRVKR
18.7	1365.7405	0.7990	HFGKPGAEPVSK
18.7	1366.7496	-0.2102	YALALFQIATEK
18.7	1366.7496	-0.2102	YALALFQLATEK

Mascot: <http://www.matrixscience.com/>

Mascot Search Results

Peptide View

MS/MS Fragmentation of **GIIDAILDGSIDKAPTK**

Found in [gi|53715725](#), phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

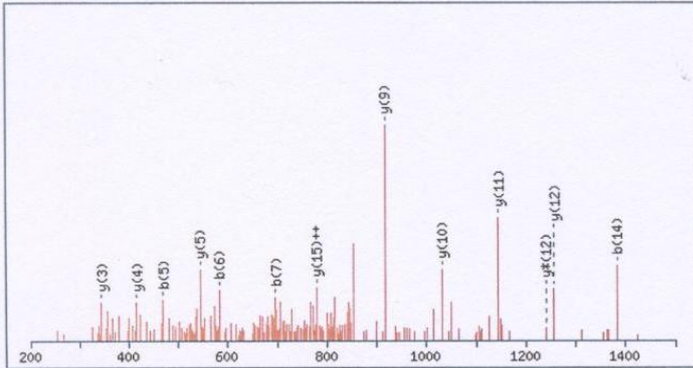
Match to Query 153: 1725.825448 from(863.920000,2+) intensity(2365091.0000)

Title: Cmpd 72, +MSn(864.71), 8.1 min

Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

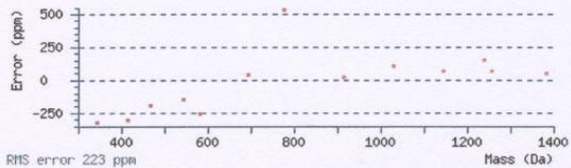
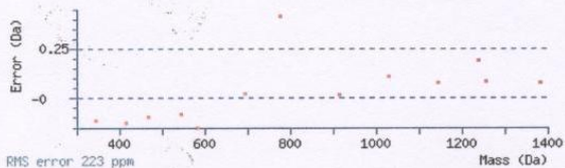
Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from to Da



Monoisotopic mass of neutral peptide Mr(calc): 1725.9512
 Fixed modifications: Carboxymethyl (C)
 Ions Score: 60 Expect: 0.024
 Matches (Bold Red): 13/144 fragment ions using 16 most intense peaks

#	a	a ⁺⁺	a [*]	a ^{*++}	b	b ⁺⁺	b [*]	b ^{*++}	Seq.	y	y ⁺⁺	y [*]	y ^{*++}	#
1	30.0338	15.5206			58.0287	29.5180			G					17
2	143.1179	72.0626			171.1128	86.0600			I	1669.9371	835.4722	1652.9105	826.9589	16
3	256.2020	128.6046			284.1969	142.6021			I	1556.8530	778.9301	1539.8265	770.4169	15
4	371.2289	186.1181			399.2238	200.1155			D	1443.7690	722.3881	1426.7424	713.8748	14
5	442.2660	221.6366			470.2609	235.6341			A	1328.7420	664.8746	1311.7155	656.3614	13
6	555.3501	278.1787			583.3450	292.1761			I	1257.7049	629.3561	1240.6783	620.8428	12
7	668.4341	334.7207			696.4291	348.7182			L	1144.6208	572.8141	1127.5943	564.3008	11
8	783.4611	392.2342			811.4560	406.2316			D	1031.5368	516.2720	1014.5102	507.7587	10
9	840.4825	420.7449			868.4775	434.7424			G	916.5098	458.7585	899.4833	450.2453	9
10	927.5146	464.2609			955.5095	478.2584			S	859.4884	430.2478	842.4618	421.7345	8
11	1040.5986	520.8030			1068.5936	534.8004			I	772.4563	386.7318	755.4298	378.2185	7
12	1155.6256	578.3164			1183.6205	592.3139			D	659.3723	330.1898	642.3457	321.6765	6
13	1283.7205	642.3639	1266.6940	633.8506	1311.7155	656.3614	1294.6889	647.8481	K	544.3453	272.6763	527.3188	264.1630	5
14	1354.7577	677.8825	1337.7311	669.3692	1382.7526	691.8799	1365.7260	683.3666	A	416.2504	208.6288	399.2238	200.1155	4
15	1451.8104	726.4088	1434.7839	717.8956	1479.8053	740.4063	1462.7788	731.8930	P	345.2132	173.1103	328.1867	164.5970	3
16	1552.8581	776.9327	1535.8316	768.4194	1580.8530	790.9301	1563.8265	782.4169	T	248.1605	124.5839	231.1339	116.0706	2
17									K	147.1128	74.0600	130.0863	65.5468	1



NCBI **BLAST** search of **GIIDAILDGSIDKAPTK**
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

All matches to this query

Score	Mr(calc):	Delta	Sequence
59.6	1725.9512	-0.1258	GIIDAILDGSIDKAPTK
43.9	1725.9149	-0.0894	GIIDAILDGSINEAPTK
21.9	1726.8923	-1.0669	NNIEIIDATCPVVL
21.8	1725.8291	-0.0036	DGPGAGKLMADLMTHGR
21.8	1726.9326	-1.1071	DLNQLLDVSSRLQR
18.7	1726.7655	-0.9400	MSDPTVMIHDPPEAR
17.7	1725.9341	-0.1087	ILNEIYWVSTYVVK
17.7	1725.9824	-0.1570	MPIQLLRHAPGAPGLR
17.7	1725.9447	-0.1193	IVEGALLAAGKPMQVAR
17.2	1725.8645	-0.0391	AVDNSSDAIASILHSR

Mascot: <http://www.matrixscience.com/>

MASCOT SCIENCE Mascot Search Results

Peptide View

MS/MS Fragmentation of **VIPFFDFVVPTELPGVDPK**

Found in **gi|53715725**, phosphoenolpyruvate carboxykinase [Bacteroides fragilis YCH46]

Match to Query 216: 2115.015448 from(1058.515000,2+) intensity(2057610.0000)

Title: Cmpd 180, +MSn(1059.16), 11.6 min

Data file D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (6901) 10_RH23_01_1294.d\SSP (6901) 10_RH23_01_1294.mgf

Click mouse within plot area to zoom in by factor of two about that point

Or, Plot from to Da

Monoisotopic mass of neutral peptide Mr(calc): 2115.1293

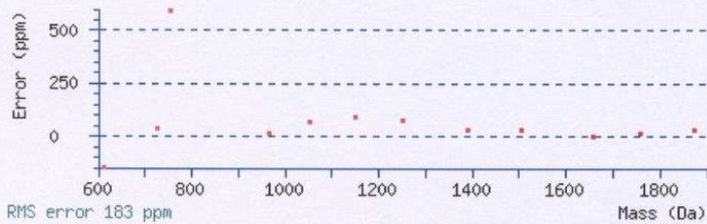
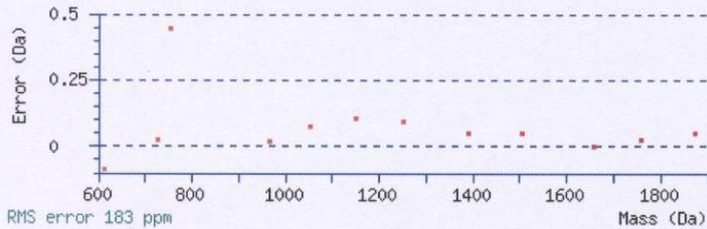
Fixed modifications: Carboxymethyl (C)

Ions Score: 41 Expect: 1.9

Matches (Bold Red): 12/144 fragment ions using 18 most intense peaks

#	a	a ⁺⁺	b	b ⁺⁺	Seq.	y	y ⁺⁺	y*	y ^{*++}	#
1	72.0808	36.5440	100.0757	50.5415	V					19
2	185.1648	93.0861	213.1598	107.0835	I	2017.0681	1009.0377	2000.0415	1000.5244	18
3	282.2176	141.6124	310.2125	155.6099	P	1903.9840	952.4957	1886.9575	943.9824	17
4	429.2860	215.1466	457.2809	229.1441	F	1806.9313	903.9693	1789.9047	895.4560	16
5	576.3544	288.6809	604.3493	302.6783	F	1659.8629	830.4351	1642.8363	821.9218	15
6	691.3814	346.1943	719.3763	360.1918	D	1512.7944	756.9009	1495.7679	748.3876	14
7	838.4498	419.7285	866.4447	433.7260	F	1397.7675	699.3874	1380.7409	690.8741	13
8	937.5182	469.2627	965.5131	483.2602	V	1250.6991	625.8532	1233.6725	617.3399	12
9	1036.5866	518.7969	1064.5815	532.7944	V	1151.6307	576.3190	1134.6041	567.8057	11
10	1133.6394	567.3233	1161.6343	581.3208	P	1052.5623	526.7848	1035.5357	518.2715	10
11	1234.6871	617.8472	1262.6820	631.8446	T	955.5095	478.2584	938.4829	469.7451	9
12	1363.7297	682.3685	1391.7246	696.3659	E	854.4618	427.7345	837.4353	419.2213	8

13	1476.8137	738.9105	1504.8086	752.9080	L	725.4192	363.2132	708.3927	354.7000	7
14	1573.8665	787.4369	1601.8614	801.4343	P	612.3352	306.6712	595.3086	298.1579	6
15	1630.8879	815.9476	1658.8829	829.9451	G	515.2824	258.1448	498.2558	249.6316	5
16	1729.9564	865.4818	1757.9513	879.4793	V	458.2609	229.6341	441.2344	221.1208	4
17	1844.9833	922.9953	1872.9782	936.9927	D	359.1925	180.0999	342.1660	171.5866	3
18	1942.0361	971.5217	1970.0310	985.5191	P	244.1656	122.5864	227.1390	114.0731	2
19					K	147.1128	74.0600	130.0863	65.5468	1



NCBI **BLAST** search of [VIPFFDFVVPTELPGVDPK](#)
 (Parameters: blastp, nr protein database, expect=20000, no filter, PAM30)
 Other BLAST [web gateways](#)

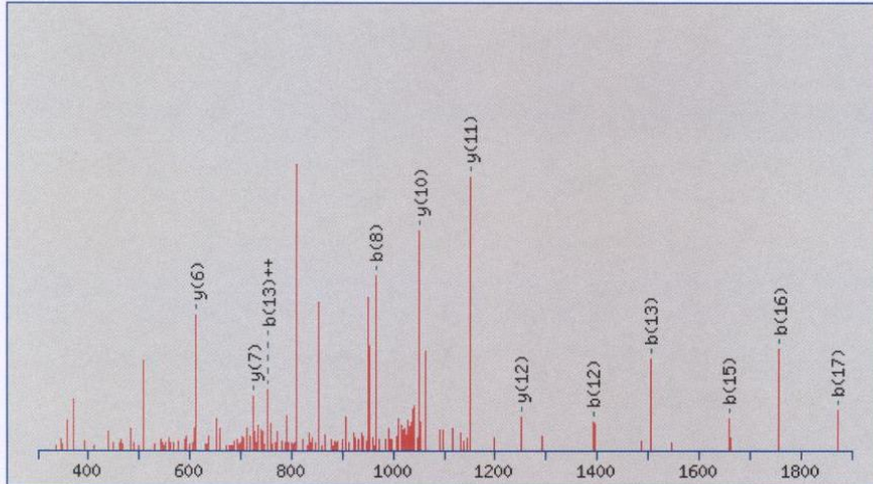
All matches to this query

Score	Mr(calc):	Delta	Sequence
41.0	2115.1293	-0.1138	VIPFFDFVVPTELPGVDPK
18.9	2115.2013	-0.1859	IKEKMPDIPVIVISAFTSK
18.0	2115.2892	-0.2737	IPISAEVAAIVRQRGLVVPK
15.8	2114.9513	0.0641	DENGVPMRDHLQPEYMK
15.4	2115.1940	-0.1785	ILDAAQIVDIVSDFVTLRK
15.4	2115.1940	-0.1785	ILDAAQIVEVVSDFVTLRK
15.0	2115.2489	-0.2335	LLPKMGTIPEVVILDPPRK
14.7	2115.1589	-0.1435	LVDVVGKIPHPGRGANFVDPK
14.1	2114.9757	0.0398	VSGNETFDDVSLSVQFESR
13.9	2114.9903	0.0252	KDEGHVTVVTVSAMAGETDR

Mascot: <http://www.matrixscience.com/>

Bacteroides fragilis SSP 6901- Peptide sequences

VIPFFDFVVPTELPGVDPK- Ion score 41

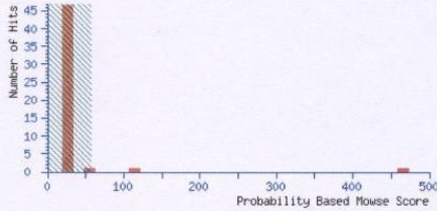


Mascot Search Results

User : Lakshmy Manickan
Email : lakshmy.manickan@unn.ac.uk
Search title : SSP 7301
MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (7301) 10_RJ12_01_1384.d\SSP (7301) 10_RJ12_01_1384.mgf
Database : NCBIInr 20100102 (10272453 sequences; 3505279183 residues)
Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
Timestamp : 6 Jan 2010 at 16:22:43 GMT
Protein hits : [gi|143945](#) Fe-superoxide dismutase [Bacteroides fragilis]
[gi|29346065](#) superoxide dismutase [Fe] [Bacteroides thetaiotaomicron VPI-5482]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event.
 Individual ions scores > 57 indicate identity or extensive homology ($p < 0.05$).
 Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As: Peptide Summary [Help](#)
 Significance threshold $p < 0.05$ Max. number of hits AUTO
 Standard scoring MudPIT scoring Ions score or expect cut-off 0 Show sub-sets 0
 Show pop-ups Suppress pop-ups Sort unassigned Decreasing Score Require bold red
 Select All Select None Search Selected Error tolerant

1. [gi|143945](#) Mass: 21736 Score: 466 Queries matched: 37 emPAI: 1.36
 Fe-superoxide dismutase [Bacteroides fragilis]

Check to include this hit in error tolerant search

Query	Observed	Mr(expt)	Mr(calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 40	545.6590	1089.3034	1089.4767	-0.1733	0	35	6.8	1	R.DFGSPENFK.K
<input checked="" type="checkbox"/> 78	772.7580	1543.5014	1543.7922	-0.2908	0	(69)	0.0026	1	K.LWEIIDWDVVEK.R
<input checked="" type="checkbox"/> 79	515.5170	1543.5292	1543.7922	-0.2631	0	(35)	6.2	1	K.LWEIIDWDVVEK.R
<input checked="" type="checkbox"/> 80	772.7800	1543.5454	1543.7922	-0.2468	0	84	7.8e-05	1	K.LWEIIDWDVVEK.R
<input checked="" type="checkbox"/> 81	772.7810	1543.5474	1543.7922	-0.2448	0	(70)	0.0019	1	K.LWEIIDWDVVEK.R
<input checked="" type="checkbox"/> 82	772.7880	1543.5614	1543.7922	-0.2308	0	(54)	0.096	1	K.LWEIIDWDVVEK.R
<input checked="" type="checkbox"/> 83	772.7930	1543.5714	1543.7922	-0.2208	0	(62)	0.025	1	K.LWEIIDWDVVEK.R
<input checked="" type="checkbox"/> 84	772.8010	1543.5874	1543.7922	-0.2048	0	(66)	0.0057	1	K.LWEIIDWDVVEK.R
<input checked="" type="checkbox"/> 85	772.8190	1543.6234	1543.7922	-0.1698	0	(54)	0.1	1	K.LWEIIDWDVVEK.R
<input checked="" type="checkbox"/> 86	772.8390	1543.6634	1543.7922	-0.1288	0	(35)	8.3	1	K.LWEIIDWDVVEK.R
<input checked="" type="checkbox"/> 181	1120.9470	2239.8794	2240.0902	-0.2108	0	68	0.0027	1	K.EFNAASVGLFGSGAWLSVDDK.L
<input checked="" type="checkbox"/> 193	792.5530	2374.6372	2375.1757	-0.5385	0	(34)	5.9	1	K.HLQTYVNNLNSLVPQTEYEGK.T
<input checked="" type="checkbox"/> 195	792.5820	2374.7242	2375.1757	-0.4515	0	(40)	1.4	1	K.HLQTYVNNLNSLVPQTEYEGK.T
<input checked="" type="checkbox"/> 196	792.5970	2374.7692	2375.1757	-0.4065	0	(27)	29	1	K.HLQTYVNNLNSLVPQTEYEGK.T
<input checked="" type="checkbox"/> 197	792.6000	2374.7782	2375.1757	-0.3975	0	(37)	3.1	1	K.HLQTYVNNLNSLVPQTEYEGK.T
<input checked="" type="checkbox"/> 198	792.6190	2374.8352	2375.1757	-0.3405	0	50	0.17	1	K.HLQTYVNNLNSLVPQTEYEGK.T
<input checked="" type="checkbox"/> 200	792.6670	2374.9792	2375.1757	-0.1965	0	(27)	37	1	K.HLQTYVNNLNSLVPQTEYEGK.T
<input checked="" type="checkbox"/> 212	847.6150	2539.8232	2540.2336	-0.4104	1	(48)	0.22	1	K.EFNAASVGLFGSGAWLSVDDKDGK.L
<input checked="" type="checkbox"/> 213	847.6320	2539.8742	2540.2336	-0.3594	1	(37)	3.2	1	K.EFNAASVGLFGSGAWLSVDDKDGK.L
<input checked="" type="checkbox"/> 214	847.6330	2539.8772	2540.2336	-0.3564	1	(49)	0.2	1	K.EFNAASVGLFGSGAWLSVDDKDGK.L
<input checked="" type="checkbox"/> 215	847.6490	2539.9252	2540.2336	-0.3084	1	(63)	0.0078	1	K.EFNAASVGLFGSGAWLSVDDKDGK.L
<input checked="" type="checkbox"/> 216	847.6530	2539.9372	2540.2336	-0.2964	1	(53)	0.082	1	K.EFNAASVGLFGSGAWLSVDDKDGK.L
<input checked="" type="checkbox"/> 217	847.7230	2540.1472	2540.2336	-0.0864	1	(66)	0.0056	1	K.EFNAASVGLFGSGAWLSVDDKDGK.L
<input checked="" type="checkbox"/> 109	847.9690	2540.8852	2540.2336	0.6516	1	83	0.00021	1	K.EFNAASVGLFGSGAWLSVDDKDGK.L
<input checked="" type="checkbox"/> 223	878.6390	2632.8952	2633.3125	-0.4174	0	(72)	0.00094	1	K.LPYANNALEPVISQQTIDYHYGK.H
<input checked="" type="checkbox"/> 224	878.6430	2632.9072	2633.3125	-0.4054	0	72	0.00092	1	K.LPYANNALEPVISQQTIDYHYGK.H
<input checked="" type="checkbox"/> 225	878.6580	2632.9522	2633.3125	-0.3604	0	(54)	0.064	1	K.LPYANNALEPVISQQTIDYHYGK.H
<input checked="" type="checkbox"/> 226	878.6660	2632.9762	2633.3125	-0.3364	0	(61)	0.011	1	K.LPYANNALEPVISQQTIDYHYGK.H
<input checked="" type="checkbox"/> 227	878.6780	2633.0122	2633.3125	-0.3004	0	(55)	0.051	1	K.LPYANNALEPVISQQTIDYHYGK.H
<input checked="" type="checkbox"/> 228	878.9610	2633.8612	2633.3125	0.5486	0	(53)	0.069	1	K.LPYANNALEPVISQQTIDYHYGK.H
<input checked="" type="checkbox"/> 229	878.9610	2633.8612	2633.3125	0.5486	0	(43)	0.74	1	K.LPYANNALEPVISQQTIDYHYGK.H
<input checked="" type="checkbox"/> 230	878.9770	2633.9092	2633.3125	0.5966	0	(57)	0.028	1	K.LPYANNALEPVISQQTIDYHYGK.H
<input checked="" type="checkbox"/> 231	878.9950	2633.9632	2633.3125	0.6506	0	(62)	0.0095	1	K.LPYANNALEPVISQQTIDYHYGK.H

✓	240	923.3140	2766.9202	2767.3758	-0.4556	0	(49)	0.17	1	R.AGLKPLLGFDVWEHAYLYDYNR.R
✓	241	923.3210	2766.9412	2767.3758	-0.4346	0	73	0.00066	1	R.AGLKPLLGFDVWEHAYLYDYNR.R
✓	242	923.3360	2766.9862	2767.3758	-0.3896	0	(30)	13	1	R.AGLKPLLGFDVWEHAYLYDYNR.R
✓	243	923.3410	2767.0012	2767.3758	-0.3746	0	(50)	0.15	1	R.AGLKPLLGFDVWEHAYLYDYNR.R

Proteins matching the same set of peptides:

[gi|53713818](#) Mass: 23124 Score: 466 Queries matched: 37
 superoxide dismutase [Bacteroides fragilis YCH46]
[gi|60682036](#) Mass: 21758 Score: 466 Queries matched: 37
 superoxide dismutase [Fe] [Bacteroides fragilis NCTC 9343]

2. [gi|29346065](#) Mass: 21762 Score: 123 Queries matched: 8 eMPIAI: 0.33
 superoxide dismutase [Fe] [Bacteroides thetaiotaomicron VPI-5482]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide	
40	545.6590	1089.3034	1089.4767	-0.1733	0	35	6.8	1	R.DFGSFENFK.K	
193	792.5530	2374.6372	2375.1757	-0.5385	0	(34)	5.9	1	K.HLQTYVNNLNSLVPGETEYEGK.T	
195	792.5820	2374.7242	2375.1757	-0.4515	0	(40)	1.4	1	K.HLQTYVNNLNSLVPGETEYEGK.T	
196	792.5970	2374.7692	2375.1757	-0.4065	0	(27)	2.9	1	K.HLQTYVNNLNSLVPGETEYEGK.T	
197	792.6000	2374.7782	2375.1757	-0.3975	0	(37)	3.1	1	K.HLQTYVNNLNSLVPGETEYEGK.T	
198	792.6190	2374.8352	2375.1757	-0.3405	0	50	0.17	1	K.HLQTYVNNLNSLVPGETEYEGK.T	
200	792.6670	2374.9792	2375.1757	-0.1965	0	(27)	3.7	1	K.HLQTYVNNLNSLVPGETEYEGK.T	
✓	245	928.6340	2782.8802	2783.3707	-0.4906	0	38	2.2	1	R.AGLKPLLGFDVWEHSYLYDYNR.R

Proteins matching the same set of peptides:

[gi|153806587](#) Mass: 23126 Score: 123 Queries matched: 8
 hypothetical protein BACCAC_00857 [Bacteroides caccae ATCC 43185]
[gi|237719542](#) Mass: 22532 Score: 123 Queries matched: 8
 superoxide dismutase [Bacteroides sp. 2_2_4]
[gi|253568517](#) Mass: 23094 Score: 123 Queries matched: 8
 conserved hypothetical protein [Bacteroides sp. 1_1_6]
[gi|160884385](#) Mass: 23105 Score: 121 Queries matched: 8
 hypothetical protein BACOVA_02363 [Bacteroides ovatus ATCC 8483]
[gi|255692463](#) Mass: 23142 Score: 121 Queries matched: 8
 superoxide dismutase [Bacteroides finegoldii DSM 17565]
[gi|260174595](#) Mass: 23119 Score: 121 Queries matched: 8
 superoxide dismutase (Fe) [Bacteroides sp. D2]

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide	
✓	89	788.7800	1575.5454	1575.8449	-0.2995	1	39	2.8	1	WLEAKDFLLWQK
✓	61	693.0460	1384.0774	1383.6228	0.4547	0	38	3.4	1	SMLPYDEIASDK + Oxidation (M)
✓	105	828.7830	1655.5514	1655.9498	-0.3984	0	32	12	1	LFTVLPBELTPAQLSK
✓	156	692.5960	2074.7662	2074.2150	0.5512	1	31	15	1	IANEPPILQIISEVKAALR
✓	194	792.5760	2374.7062	2375.2341	-0.5279	0	28	24	1	GSNAPLLDFAIAGVAAMIVMGR + 2 Oxidation (M)
✓	44	563.7460	1125.4774	1125.5198	-0.0424	0	28	41	1	EMFGAIAMPK + 2 Oxidation (M)
✓	150	1004.8860	2007.7574	2008.0068	-0.2493	1	27	39	1	WAVPGSGWGGAFGVVARR
✓	239	922.6520	2764.9342	2765.4745	-0.5404	1	26	35	1	VLINQVPGGMISNLANQLKQQALDK + Oxidation (M)
✓	60	693.0310	1384.0474	1384.5816	-0.5342	0	26	55	1	CEETGIDYEIR
✓	176	737.9140	2210.7202	2209.9698	0.7503	1	25	48	1	IADDYMPRGQWSSSDNPK
✓	37	507.1970	1012.3794	1012.4899	-0.1105	1	25	79	1	YMIKADEK + Oxidation (M)
✓	22	908.8810	907.8737	907.5603	0.3134	1	25	54	1	LRLLAESH
✓	165	696.8470	2087.5192	2087.2177	0.3015	2	25	51	1	MKVGIVGGPAGLYSAILLKK + Oxidation (M)
✓	119	884.2970	1766.5794	1766.8257	-0.2463	1	25	1e+02	1	MIRBGEAGFLRADSSR
✓	87	779.7580	2336.2522	2335.0824	1.1697	2	25	1.5e+02	1	MDQTSDDVQKIDLMKYFDR + Oxidation (M)
✓	153	1011.2920	3030.8542	3029.6814	1.1728	2	24	1.1e+02	1	YTSQFLTAAGTSPAAPLLIHKARLLR
✓	187	1142.7100	3425.1082	3425.6715	-0.5633	1	24	1.1e+02	1	GYGPLLVFSDSDGDANMLRDFADTAVGVVNR
✓	52	631.2090	1890.6052	1891.0965	-0.4913	1	24	1.5e+02	1	KTFITGAILMAAGLVTR + Oxidation (M)
✓	161	692.9330	2075.7772	2074.9676	0.8095	2	24	72	1	NYLCPFSINGANRTMKNR + Oxidation (M)
✓	219	858.9390	2573.7952	2574.3402	-0.5450	0	24	59	1	VGLAHGVSIGLQREGIDSSNFIAPTR
✓	235	903.6510	2707.9312	2708.4385	-0.5073	1	24	58	1	VEVLRDGAFTIYGSDAIAGVINPITK
✓	116	585.8940	1754.6602	1754.8873	-0.2271	1	24	89	1	EGMTTGDISKALFLTR + Oxidation (M)
✓	218	858.2970	2571.8692	2571.2421	0.6271	0	24	65	1	MQIAAPADPMVNGAQSDSASVAATIR
✓	55	673.7860	1345.5574	1346.6652	-1.1078	1	23	1.3e+02	1	KELAPMPFENR + Oxidation (M)
✓	70	734.3110	1466.6074	1466.8093	-0.2018	1	23	1.2e+02	1	NEPLKITTELPGR
✓	59	691.0200	2070.0382	2071.0256	-0.9874	0	23	2.1e+02	1	MTDSTQTPDIPLLDAVQAR
✓	88	780.7600	1559.5054	1558.8501	0.6554	1	23	97	1	LANEILREMLVLSR + Oxidation (M)
✓	159	692.6780	2075.0122	2074.0405	0.9717	1	23	1.2e+02	1	IYPSVGMKIGSGSYTLNGK + Oxidation (M)
✓	158	692.6560	2074.9462	2074.0591	0.8871	2	23	1.2e+02	1	LLTIMDELRVGCPWDEK
✓	123	905.3180	2712.9322	2713.5420	-0.6099	2	23	1.6e+02	1	MKKALAVVLAVVQLAMCVVALSGVAR + Oxidation (M)
✓	47	594.8990	1187.7834	1187.6033	0.1801	0	23	2.6e+02	1	ELQEQASLLS
✓	126	605.9000	1814.6782	1814.9738	-0.2956	2	23	1e+02	1	VADLGKVVDTLADEKSR
✓	24	910.9870	909.9797	910.4695	-0.4897	1	22	91	1	MYKQVAAR + Oxidation (M)
✓	160	692.7580	2075.2522	2076.1149	-0.8628	2	22	1.2e+02	1	RELLIALGKMAAVDYNR + Oxidation (M)
✓	96	800.5800	2398.7182	2398.3584	0.3598	0	22	2e+02	1	AATAGAVAGVSGGLAGGVAVGILYIK
✓	173	1083.2770	3246.8092	3246.4194	0.3898	1	22	1.8e+02	1	GSSLFPCAGDDQLQAGDSVYIVCARDNVDR

Mascot Search Results

Protein View

Match to: **gi|60682036** Score: **466**
superoxide dismutase [Fe] [Bacteroides fragilis NCTC 9343]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP (7301) 10_RJ12_01_1384.d\SSP (7301) 10_RJ12_01_1384.mgf

Nominal mass (M_r): **21758**; Calculated pI value: **6.07**
 NCBI BLAST search of **gi|60682036** against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: **Bacteroides fragilis NCTC 9343**
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|253565789](#) from **Bacteroides sp. 3_2_5**
[gi|265764163](#) from **Bacteroides sp. 2_1_16**
[gi|55977805](#) from **Bacteroides fragilis**
[gi|60493470](#) from **Bacteroides fragilis NCTC 9343**
[gi|251946068](#) from **Bacteroides sp. 3_2_5**
[gi|263256771](#) from **Bacteroides sp. 2_1_16**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: **58%**

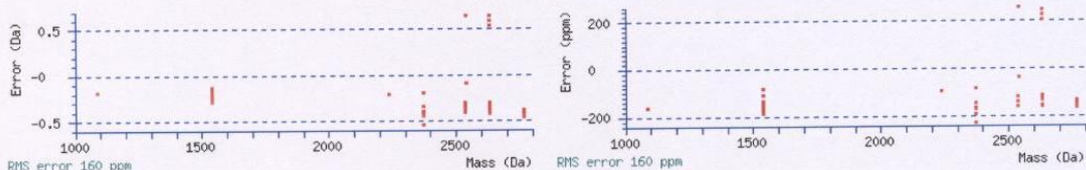
Matched peptides shown in **Bold Red**

- 1 **MTYEMPKLEPY** **ANNALEPVIS** **QQTIDYHYGK** **HLQTYVNNLN** **SLVPGTEYEG**
- 51 **KTVEAIVASA** **PDGALFNNAG** **QVLNHTLYFL** **QFAPKPAKNE** **PAGKLGKAIK**
- 101 **RDPGSFENFK** **KEFNAASVGL** **FGSGWAWLSV** **DKDGKLGK** **EPNGSNPVRA**
- 151 **GLKPLLGFDV** **WEHAYYLDYQ** **NRADHVNKL** **WEIIDWDVVE** **KRL**

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
8 - 30	878.6390	2632.8952	2633.3125	-0.4174	0 K.LPYANNALEPVISQQTIDYHYGK.H (Ions score 72)
8 - 30	878.6430	2632.9072	2633.3125	-0.4054	0 K.LPYANNALEPVISQQTIDYHYGK.H (Ions score 72)
8 - 30	878.6580	2632.9522	2633.3125	-0.3604	0 K.LPYANNALEPVISQQTIDYHYGK.H (Ions score 54)
8 - 30	878.6660	2632.9762	2633.3125	-0.3364	0 K.LPYANNALEPVISQQTIDYHYGK.H (Ions score 61)
8 - 30	878.6780	2633.0122	2633.3125	-0.3004	0 K.LPYANNALEPVISQQTIDYHYGK.H (Ions score 55)
8 - 30	878.9610	2633.8612	2633.3125	0.5486	0 K.LPYANNALEPVISQQTIDYHYGK.H (Ions score 53)
8 - 30	878.9610	2633.8612	2633.3125	0.5486	0 K.LPYANNALEPVISQQTIDYHYGK.H (Ions score 43)
8 - 30	878.9770	2633.9092	2633.3125	0.5966	0 K.LPYANNALEPVISQQTIDYHYGK.H (Ions score 57)
8 - 30	878.9950	2633.9632	2633.3125	0.6506	0 K.LPYANNALEPVISQQTIDYHYGK.H (Ions score 62)
31 - 51	792.5530	2374.6372	2375.1757	-0.5385	0 K.HLQTYVNNLNLSLVPGEYEGK.T (Ions score 34)
31 - 51	792.5820	2374.7242	2375.1757	-0.4515	0 K.HLQTYVNNLNLSLVPGEYEGK.T (Ions score 40)
31 - 51	792.5970	2374.7692	2375.1757	-0.4065	0 K.HLQTYVNNLNLSLVPGEYEGK.T (Ions score 27)
31 - 51	792.6000	2374.7782	2375.1757	-0.3975	0 K.HLQTYVNNLNLSLVPGEYEGK.T (Ions score 37)
31 - 51	792.6190	2374.8352	2375.1757	-0.3405	0 K.HLQTYVNNLNLSLVPGEYEGK.T (Ions score 50)
31 - 51	792.6670	2374.9792	2375.1757	-0.1965	0 K.HLQTYVNNLNLSLVPGEYEGK.T (Ions score 27)
102 - 110	545.6590	1089.3034	1089.4767	-0.1733	0 R.DFGSFENFK.K (Ions score 35)
112 - 132	1120.9470	2239.8794	2240.0902	-0.2108	0 K.EFNAASVGLFGSGWAWLSVDRK.L (Ions score 68)
112 - 135	847.6150	2539.8232	2540.2336	-0.4104	1 K.EFNAASVGLFGSGWAWLSVDRKDGK.L (Ions score 48)
112 - 135	847.6320	2539.8742	2540.2336	-0.3594	1 K.EFNAASVGLFGSGWAWLSVDRKDGK.L (Ions score 37)
112 - 135	847.6330	2539.8772	2540.2336	-0.3564	1 K.EFNAASVGLFGSGWAWLSVDRKDGK.L (Ions score 49)
112 - 135	847.6490	2539.9252	2540.2336	-0.3084	1 K.EFNAASVGLFGSGWAWLSVDRKDGK.L (Ions score 63)
112 - 135	847.6530	2539.9372	2540.2336	-0.2964	1 K.EFNAASVGLFGSGWAWLSVDRKDGK.L (Ions score 53)
112 - 135	847.7230	2540.1472	2540.2336	-0.0864	1 K.EFNAASVGLFGSGWAWLSVDRKDGK.L (Ions score 66)
112 - 135	847.9690	2540.8852	2540.2336	0.6516	1 K.EFNAASVGLFGSGWAWLSVDRKDGK.L (Ions score 83)
150 - 172	923.3140	2766.9202	2767.3758	-0.4556	0 R.AGLKPLLGFDVWEHAYYLDYQNR.R (Ions score 49)
150 - 172	923.3210	2766.9412	2767.3758	-0.4346	0 R.AGLKPLLGFDVWEHAYYLDYQNR.R (Ions score 73)
150 - 172	923.3360	2766.9862	2767.3758	-0.3896	0 R.AGLKPLLGFDVWEHAYYLDYQNR.R (Ions score 30)
150 - 172	923.3410	2767.0012	2767.3758	-0.3746	0 R.AGLKPLLGFDVWEHAYYLDYQNR.R (Ions score 50)
180 - 191	772.7580	1543.5014	1543.7922	-0.2908	0 K.LWEIIDWDVVEK.R (Ions score 69)
180 - 191	772.7810	1543.5474	1543.7922	-0.2448	0 K.LWEIIDWDVVEK.R (Ions score 35)
180 - 191	772.7800	1543.5454	1543.7922	-0.2468	0 K.LWEIIDWDVVEK.R (Ions score 84)
180 - 191	772.7810	1543.5474	1543.7922	-0.2448	0 K.LWEIIDWDVVEK.R (Ions score 70)
180 - 191	772.7880	1543.5614	1543.7922	-0.2308	0 K.LWEIIDWDVVEK.R (Ions score 54)
180 - 191	772.7930	1543.5714	1543.7922	-0.2208	0 K.LWEIIDWDVVEK.R (Ions score 62)
180 - 191	772.8010	1543.5874	1543.7922	-0.2048	0 K.LWEIIDWDVVEK.R (Ions score 66)
180 - 191	772.8190	1543.6234	1543.7922	-0.1688	0 K.LWEIIDWDVVEK.R (Ions score 54)
180 - 191	772.8390	1543.6634	1543.7922	-0.1288	0 K.LWEIIDWDVVEK.R (Ions score 35)



LOCUS YP_212180 193 aa linear BCT 01-MAY-2009
 DEFINITION superoxide dismutase [Fe] [Bacteroides fragilis NCTC 9343].
 ACCESSION YP_212180
 VERSION YP_212180.1 GI:60682036
 DBLINK Project:46
 DBSOURCE REFSEQ: accession [NC_003228.3](#)
 KEYWORDS .
 SOURCE Bacteroides fragilis NCTC 9343
 ORGANISM [Bacteroides fragilis NCTC 9343](#)
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 193)
 AUTHORS Cerdeno-Tarraga,A.M., Patrick,S., Crossman,L.C., Blakely,G.,
 Abratt,V., Lennard,N., Poxton,I., Duerden,B., Harris,B.,
 Quail,M.A., Barron,A., Clark,L., Corton,C., Doggett,J.,
 Holden,M.T., Larke,N., Line,A., Lord,A., Norbertczak,H., Ormond,D.,
 Price,C., Rabinowitsch,E., Woodward,J., Barrell,B. and Parkhill,J.
 TITLE Extensive DNA inversions in the B. fragilis genome control variable
 gene expression
 JOURNAL Science 307 (5714), 1463-1465 (2005)
 PUBMED [15746427](#)
 REFERENCE 2 (residues 1 to 193)
 AUTHORS Cerdeno-Tarraga,A.M.
 TITLE Direct Submission
 JOURNAL Submitted (29-JUL-2004) Pathogen Sequencing Unit, Sanger Institute,
 Wellcome Trust Genome Campus, Hinxton, Cambridge CB10 1SA, United
 Kingdom
 REFERENCE 3 (residues 1 to 193)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (08-APR-2002) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 COMMENT PROVISIONAL REFSEQ: This record has not yet been subject to final
 NCBI review. The reference sequence was derived from [CAH08256](#).
 Method: conceptual translation.
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 (193 aa) fasta scores: E(): 1.2e-77, 99.48% id in 193 aa,
 and to Escherichia coli, Escherichia coli O6, Escherichia
 coli O157:H7, and Shigella flexneri superoxide dismutase
 [Fe] SodB or B1656 or C2050 or Z2678 or ECS2365 or SF1684
 or S1816 SWALL:SODF_ECOLI (SWALL:P09157) (192 aa) fasta
 scores: E(): 1.6e-37, 51.04% id in 192 aa"
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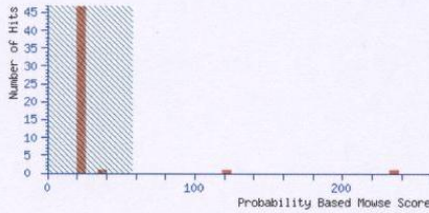
Mascot: <http://www.matrixscience.com/>

Mascot Search Results

User : Lakshmy Manickan
 Email : lakshmy.manickan@unn.ac.uk
 Search title : SSP 7401
 MS data file : D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP 7401 (10)_RN15_01_1596.d\SSP 7401 (10)_RN15_01_1596.mgf
 Database : NCBI nr 20100102 (10272453 sequences; 3505279183 residues)
 Taxonomy : Bacteria (Eubacteria) (5690016 sequences)
 Timestamp : 6 Jan 2010 at 16:24:05 GMT
 Protein hits : [gi|53711978](#) ribosome recycling factor [Bacteroides fragilis YCH46]
 [gi|29347659](#) ribosome recycling factor [Bacteroides thetaiotaomicron VPI-5482]

Probability Based Mowse Score

Ions score is $-10 \cdot \log(P)$, where P is the probability that the observed match is a random event. Individual ions scores > 57 indicate identity or extensive homology (p<0.05). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Peptide Summary Report

Format As [Help](#)

Significance threshold p< Max. number of hits

Standard scoring MudPIT scoring Ions score or expect cut-off Show sub-sets

Show pop-ups Suppress pop-ups Sort unassigned Require bold red

Select All Select None Search Selected Error tolerant

1. [gi|53711978](#) Mass: 20809 Score: 235 Queries matched: 5 emPAI: 0.82
 ribosome recycling factor [Bacteroides fragilis YCH46]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 148	542.2400	1623.6982	1623.8443	-0.1461	1	37	4.9	1	R.SITIKPWKSMFR.V + Oxidation (M)
<input checked="" type="checkbox"/> 223	695.9640	2084.8702	2085.0412	-0.1710	0	(44)	0.82	1	K.AIIDSDLGIMPENNGEIRR.I + Oxidation (M)
<input checked="" type="checkbox"/> 224	1043.9710	2085.9274	2085.0412	0.8963	0	86	6.3e-05	1	K.AIIDSDLGIMPENNGEIRR.I + Oxidation (M)
<input checked="" type="checkbox"/> 241	1133.4620	2264.9094	2265.0947	-0.1853	0	75	0.00051	1	R.VDSYSGMVPISNVAALSTPDAR.S + Oxidation (M)
<input checked="" type="checkbox"/> 250	978.4010	2932.1812	2932.4964	-0.3153	1	37	3	1	R.LLDGIRVDSYSGMVPISNVAALSTPDAR.S + Oxidation (M)

Proteins matching the same set of peptides:
[gi|153808675](#) Mass: 20841 Score: 235 Queries matched: 5
 hypothetical protein BACCAC_02974 [Bacteroides caccae ATCC 43185]
[gi|265765322](#) Mass: 20823 Score: 235 Queries matched: 5
 ribosome recycling factor [Bacteroides sp. 2_1_16]

2. [gi|29347659](#) Mass: 20827 Score: 117 Queries matched: 3 emPAI: 0.16
 ribosome recycling factor [Bacteroides thetaiotaomicron VPI-5482]
 Check to include this hit in error tolerant search

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
148	542.2400	1623.6982	1623.8443	-0.1461	1	37	4.9	1	R.SITIKPWKSMFR.A + Oxidation (M)
241	1133.4620	2264.9094	2265.0947	-0.1853	0	53	0.082	2	R.VDSYSGMVPISNVAALSTPDAR.S + Oxidation (M)
250	978.4010	2932.1812	2932.4964	-0.3152	1	26	32	3	R.LLDGIRVDSYSGMVPISNVAALSTPDAR.S + Oxidation (M)

Peptide matches not assigned to protein hits: (no details means no match)

Query	Observed	Mr (expt)	Mr (calc)	Delta	Miss	Score	Expect	Rank	Peptide
<input checked="" type="checkbox"/> 206	672.1310	2013.3712	2013.1016	0.2696	1	35	5.6	1	MPISSAPIRLPMRPIFR + 2 Oxidation (M)
<input checked="" type="checkbox"/> 150	547.6960	1640.0662	1639.9257	0.1405	1	30	22	1	TVLTGDTLRALPDLR
<input checked="" type="checkbox"/> 138	771.4460	2311.3162	2311.2172	0.0989	1	29	65	1	REGFVVVTGTEAATPPETVPVVR
<input checked="" type="checkbox"/> 115	657.2930	1312.5714	1311.6567	0.9148	0	28	44	1	LELGLTMGGFMK + Oxidation (M)
<input checked="" type="checkbox"/> 246	883.6690	2647.9852	2647.0551	0.9301	0	27	28	1	NHDGADVDELGYSCHPAEDVMR + Oxidation (M)
<input checked="" type="checkbox"/> 133	743.2970	2226.8692	2227.9435	-1.0744	0	27	93	1	QTDPAEAAVVAECNNMAFK + Oxidation (M)
<input checked="" type="checkbox"/> 193	658.3850	1972.1332	1972.0047	0.1284	1	26	57	1	LDEINLMDRLADGSAIR
<input checked="" type="checkbox"/> 162	901.3710	1800.7274	1800.9370	-0.2096	0	26	53	1	VETQTAEEVGGVIWVSR
<input checked="" type="checkbox"/> 129	721.5120	2161.5142	2162.0439	-0.5297	2	26	1e+02	1	VHMEHELRGAWERDEL
<input checked="" type="checkbox"/> 101	576.9530	1727.8372	1728.0371	-0.2000	1	26	1.2e+02	1	KTVFAALMLAIPALAK

MASCOT Mascot Search Results

Protein View

Match to: gi|53711978 Score: 235
ribosome recycling factor [Bacteroides fragilis YCH46]
 Found in search of D:\Data\Lakshmy\PROTEOMICS 3- 151208\SSP 7401 (10)_RN15_01_1596.d\SSP 7401 (10)_RN15_01_1596.mgf

Nominal mass (M_r): 20809; Calculated pI value: 5.86
 NCBI BLAST search of gi|53711978 against nr
 Unformatted [sequence string](#) for pasting into other applications

Taxonomy: **Bacteroides fragilis YCH46**
 Links to retrieve other entries containing this sequence from NCBI Entrez:
[gi|60680182](#) from **Bacteroides fragilis NCTC 9343**
[gi|253563986](#) from **Bacteroides sp. 3_2_5**
[gi|255007502](#) from **Bacteroides fragilis 3_1_12**
[gi|60390242](#) from **Bacteroides fragilis**
[gi|81316748](#) from **Bacteroides fragilis NCTC 9343**
[gi|52214843](#) from **Bacteroides fragilis YCH46**
[gi|60491616](#) from **Bacteroides fragilis NCTC 9343**
[gi|251947762](#) from **Bacteroides sp. 3_2_5**

Fixed modifications: Carboxymethyl (C)
 Variable modifications: Oxidation (M)
 Cleavage by Trypsin: cuts C-term side of KR unless next residue is P
 Sequence Coverage: 32%

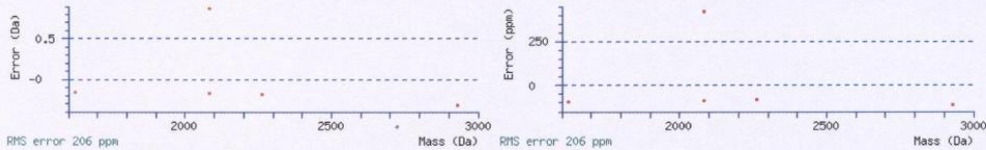
Matched peptides shown in **Bold Red**

1 MVDVKTIIIE SQEKMDMAVM YLEEALAHIR AGKASTRLLD GIRVDSYGSM
51 VPISNVAAVT TPDARSITIK PWDKSMFRVI EKAIIDSDLG IMPENNGEII
101 RIGIPLITEE RRLQAKQCK AEGETAKVSI RNARRDGIDA LKAVKDGLA
151 EDEQKNAEAK LQKVHDKYIA KIEMLAEKD KEIMTV

Show predicted peptides also

Sort Peptides By Residue Number Increasing Mass Decreasing Mass

Start - End	Observed	Mr (expt)	Mr (calc)	Delta	Miss Sequence
38 - 65	978.4010	2932.1812	2932.4964	-0.3153	1 R.LLDGIRVDSYSGMVPISNVAAVTPDAR.S Oxidation (M) (Ions score 37)
44 - 65	1133.4620	2264.9094	2265.0947	-0.1853	0 R.VDSYSGMVPISNVAAVTPDAR.S Oxidation (M) (Ions score 75)
66 - 78	542.2400	1623.6982	1623.8443	-0.1461	1 R.SITTKPDKSMFR.V Oxidation (M) (Ions score 37)
83 - 101	695.9640	2084.8702	2085.0412	-0.1710	0 K.AIIDSDLGIMPENNGEII.R Oxidation (M) (Ions score 44)
83 - 101	1043.9710	2085.9274	2085.0412	0.8863	0 K.AIIDSDLGIMPENNGEII.R Oxidation (M) (Ions score 86)



LOCUS YP_097970 186 aa linear BCT 26-APR-2009
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 ACCESSION YP_097970
 VERSION YP_097970.1 GI:53711978
 DBLINK Project:13067
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 ORGANISM Bacteroides fragilis YCH46
 Bacteria; Bacteroidetes; Bacteroidia; Bacteroidales;
 Bacteroidaceae; Bacteroides.
 REFERENCE 1 (residues 1 to 186)
 AUTHORS Kuwahara,T., Yamashita,A., Hirakawa,H., Nakayama,H., Toh,H.,
 Okada,N., Kuhara,S., Hattori,M., Hayashi,T. and Ohnishi,Y.
 TITLE Genomic analysis of Bacteroides fragilis reveals extensive DNA
 inversions regulating cell surface adaptation
 JOURNAL Proc. Natl. Acad. Sci. U.S.A. 101 (41), 14919-14924 (2004)
 PUBMED 15466707
 REFERENCE 2 (residues 1 to 186)
 CONSRM NCBI Genome Project
 TITLE Direct Submission
 JOURNAL Submitted (01-OCT-2004) National Center for Biotechnology
 Information, NIH, Bethesda, MD 20894, USA
 REFERENCE 3 (residues 1 to 186)
 AUTHORS Hattori,M., Yamashita,A., Toh,H., Oshima,K. and Shiba,T.
 TITLE Direct Submission
 JOURNAL Submitted (20-APR-2004) Kitasato Institute for Life Sciences,
 1-15-1, Kitasato, Sagami-hara, Kanagawa 228-8555, Japan
 COMMENT REVIEWED REFSEQ: This record has been curated by NCBI staff. The
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 Method: conceptual translation.
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 Region 7..184

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termination of translation. Thus ribosomes are 'recycled'
and ready for another round of...; cd00520"
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along with release factor RF-3 and elongation factor EF-G;
A GTPase-dependent process results in release of 50S from
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viability; structurally similar to tRNAs"
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Mascot: <http://www.matrixscience.com/>

G4 Blast analysis results for hypothetical proteins BF2494, BF1203 and BF0301 in *B. fragilis*

BLAST

Basic Local Alignment Search Tool

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gi|60491326|emb|CAH06074.1| putative transmembrane...

Results for:
 Your BLAST job specified more than one input sequence. This box lets you choose which input sequence to show BLAST results for.

Query ID

[|c|64607](#)
[|c|64607](#)

Description

gi|60491326|emb|CAH06074.1| putative transmembrane protein [Bacteroides fragilis NCTC 9343]

Molecule type

amino acid

Query Length

406

Database Name

nr

Description

All non-redundant GenBank CDS translations+PDB+SwissProt+PIR+PRF excluding environmental samples from WGS projects

Program

BLASTP 2.2.22+ [Citation](#)

Reference

Stephen F. Altschul, Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", *Nucleic Acids Res.* 25:3389-3402.

Reference - compositional score matrix adjustment

Stephen F. Altschul, John C. Wootton, E. Michael Gertz, Richa Agarwala, Aleksandr Morgulis, Alejandro A. Schäffer, and Yi-Kuo Yu (2005) "Protein database searches using compositionally adjusted substitution matrices", *FEBS J.* 272:5101-5109.

Other reports: [Search Summary](#) [[Taxonomy reports](#)] [[Distance tree of results](#)] [[Multiple alignment](#)] **NEW**
[Search Parameters](#)

Search parameter name Search parameter value

Program	blastp
Word size	3
Expect value	10
Hittlist size	100
Gapcosts	11,1
Matrix	BLOSUM62
Filter string	F
Genetic Code	1
Window Size	40
Threshold	11
Composition-based stats	2

Database

Database parameter name Database parameter value

Posted date	Jan 20, 2010 4:14 PM
Number of letters	3,542,056,219
Number of sequences	10,381,779
Entrez query	none

Karlin-Altschul statistics

Params Ungapped Gapped

Lambda	0.332433	0.267
K	0.144686	0.041
H	0.465429	0.14

Results Statistics

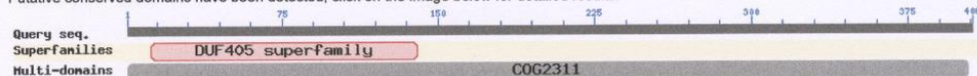
Results Statistics parameter name Results Statistics parameter value

Length adjustment	139
Effective length of query	267
Effective length of database	2098988938
Effective search space	560430046446
Effective search space used	560430046446

Graphic Summary

Show Conserved Domains

Putative conserved domains have been detected, click on the image below for detailed results.



Descriptions

Legend for links to other resources: [U](#) UniGene [E](#) GEO [G](#) Gene [S](#) Structure [M](#) Map Viewer

Sequences producing significant alignments:

(Click headers to sort columns)

YP_210036.1	putative transmembrane protein [Bacteroides fragilis NCTC 9343] >emb CAH06074.1 putative transmembrane protein [Bacteroides fragilis NCTC 9343]	824	824	100%	0.0	G
ZP_04841755.1	conserved hypothetical protein [Bacteroides sp. 3_2_5] >ref ZP_06093303.1 conserved hypothetical protein [Bacteroides sp. 2_1_16] >gb EES88356.1 conserved hypothetical protein [Bacteroides sp. 3_2_5] >gb EEZ25846.1 conserved hypothetical protein [Bacteroides sp. 2_1_16]	822	822	100%	0.0	
YP_097636.1	putative transport protein [Bacteroides fragilis YCH46] >dbj BAD47102.1 putative transport protein [Bacteroides fragilis YCH46]	822	822	100%	0.0	G
YP_001818702.1	hypothetical protein Oter_1818 [Opitutus terrae PB90-1] >gb ACE75102.1 protein of unknown function DUF418 [Opitutus terrae PB90-1]	261	261	97%	1e-67	G
ZP_05414587.1	putative membrane-associated protein [Bacteroides finegoldii DSM 17565] >gb EEX46425.1 putative membrane-associated protein [Bacteroides finegoldii DSM 17565]	241	241	95%	8e-62	
NP_813370.1	putative transport protein [Bacteroides thetaiotaomicron VPI-5482] >ref ZP_04847173.1 conserved hypothetical protein [Bacteroides sp. 1_1_6] >gb AAO79564.1 conserved hypothetical protein, putative transport protein [Bacteroides thetaiotaomicron VPI-5482] >gb EES68227.1 conserved hypothetical protein [Bacteroides sp. 1_1_6]	239	239	95%	4e-61	G
ZP_05757845.1	putative transport protein [Bacteroides sp. D2]	239	239	95%	5e-61	
ZP_02068252.1	hypothetical protein BACOVA_05266 [Bacteroides ovatus ATCC 8483] >gb EEO09405.1 hypothetical protein BACOVA_05266 [Bacteroides ovatus ATCC 8483]	238	238	95%	1e-60	
ZP_04552009.1	conserved hypothetical protein [Bacteroides sp. 2_2_4] >gb EEO55115.1 conserved hypothetical protein [Bacteroides sp. 2_2_4]	235	235	95%	7e-60	
ZP_05545092.1	conserved hypothetical protein [Parabacteroides sp. D13] >gb EEU51838.1 conserved hypothetical protein [Parabacteroides sp. D13]	233	233	95%	3e-59	
YP_001302036.1	putative membrane-associated protein [Parabacteroides distasonis ATCC 8503] >gb ABR42414.1 putative membrane-associated protein [Parabacteroides distasonis ATCC 8503]	233	233	95%	5e-59	G
ZP_05286207.1	putative membrane-associated protein [Bacteroides sp. 2_1_7]	231	231	95%	1e-58	
ZP_04846930.1	conserved hypothetical protein [Bacteroides sp. 1_1_6] >gb EES69620.1 conserved hypothetical protein [Bacteroides sp. 1_1_6]	230	230	97%	2e-58	
ZP_06075273.1	conserved hypothetical protein [Bacteroides sp. 2_1_33B] >gb EEY85242.1 conserved hypothetical protein [Bacteroides sp. 2_1_33B]	230	230	95%	2e-58	
NP_811692.1	putative membrane-associated protein [Bacteroides thetaiotaomicron VPI-5482] >gb AAO77896.1 conserved hypothetical protein, putative membrane-associated protein [Bacteroides thetaiotaomicron VPI-5482]	228	228	97%	1e-57	G
ZP_04842518.1	conserved hypothetical protein [Bacteroides sp. 3_2_5] >ref ZP_06094020.1 conserved hypothetical protein [Bacteroides sp. 2_1_16] >gb EES86904.1 conserved hypothetical protein [Bacteroides sp. 3_2_5] >gb EEZ25112.1 conserved hypothetical protein [Bacteroides sp. 2_1_16]	227	227	96%	2e-57	
YP_100550.1	putative membrane-associated protein [Bacteroides fragilis YCH46] >dbj BAD50016.1 putative membrane-associated protein [Bacteroides fragilis YCH46]	226	226	96%	4e-57	G
YP_212723.1	hypothetical protein BF3110 [Bacteroides fragilis NCTC 9343] >emb CAH08805.1 conserved hypothetical membrane protein [Bacteroides fragilis NCTC 9343]	226	226	96%	5e-57	G
ZP_03014090.1	hypothetical protein BACINT_01653 [Bacteroides intestinalis DSM 17393] >gb EDV06564.1 hypothetical protein BACINT_01653 [Bacteroides intestinalis DSM 17393]	224	224	96%	1e-56	
ZP_03678199.1	hypothetical protein BACCELL_02542 [Bacteroides cellulosilyticus DSM 14838] >gb EEF89824.1 hypothetical protein BACCELL_02542 [Bacteroides cellulosilyticus DSM 14838]	223	223	96%	3e-56	
ZP_05282878.1	hypothetical protein Bfra3_16553 [Bacteroides fragilis 3_1_12]	223	223	96%	4e-56	
ZP_03643854.1	hypothetical protein BACCOPRO_02228 [Bacteroides coprophilus DSM 18228] >gb EEF76722.1 hypothetical protein BACCOPRO_02228 [Bacteroides coprophilus DSM 18228]	218	218	95%	1e-54	
YP_001299550.1	hypothetical protein BVU_2269 [Bacteroides vulgatus ATCC 8482] >gb ABR39928.1 conserved hypothetical membrane protein [Bacteroides vulgatus ATCC 8482]	218	218	96%	1e-54	G
ZP_05255382.1	conserved hypothetical protein [Bacteroides sp. 4_3_47FAA] >gb EET15774.1 conserved hypothetical protein [Bacteroides sp. 4_3_47FAA]	217	217	96%	3e-54	
ZP_04558306.1	conserved hypothetical protein [Bacteroides sp. D4] >ref ZP_06087176.1 conserved hypothetical protein [Bacteroides sp. 3_1_33FAA] >gb EEO44758.1 conserved hypothetical protein [Bacteroides sp. D4] >gb EEZ23459.1 conserved hypothetical protein [Bacteroides sp. 3_1_33FAA]	216	216	96%	5e-54	
ZP_04540787.1	conserved hypothetical protein [Bacteroides sp. 9_1_42FAA] >gb EEO61489.1 conserved hypothetical protein [Bacteroides sp. 9_1_42FAA]	215	215	96%	1e-53	
ZP_03302157.1	hypothetical protein BACDOR_03555 [Bacteroides dorei DSM 17855] >gb EEB24135.1 hypothetical protein BACDOR_03555 [Bacteroides dorei DSM 17855]	213	213	96%	2e-53	

Alignments [Select All](#) [Get selected sequences](#) [Distance tree of results](#) [Multiple alignment](#) [NEW](#)

>ref|YP_210036.1| [G](#) putative transmembrane protein [Bacteroides fragilis NCTC 9343]
 emb|CAH06074.1| [G](#) putative transmembrane protein [Bacteroides fragilis NCTC 9343]
 Length=406

GENE ID: 3285991 BF0301 | putative transmembrane protein
 [Bacteroides fragilis NCTC 9343] (10 or fewer PubMed links)
 Score = 824 bits (2129), Expect = 0.0, Method: Compositional matrix adjust.
 Identities = 406/406 (100%), Positives = 406/406 (100%), Gaps = 0/406 (0%)

Query 1 MHTQTITPKKRINSIDALRGFALIGIMLLHCMERFDLTLAPVVEPFWQAITAVYDLSL 60
 MHTQTITPKKRINSIDALRGFALIGIMLLHCMERFDLTLAPVVEPFWQAITAVYDLSL
 Sbjct 1 MHTQTITPKKRINSIDALRGFALIGIMLLHCMERFDLTLAPVVEPFWQAITAVYDLSL 60
 Query 61 YLFLSGKSYAMFSLFLGSLFFMQMESQAAGVDFRGRFLWRLLALLFLFGYINGLYVMGEF 120
 YLFLSGKSYAMFSLFLGSLFFMQMESQAAGVDFRGRFLWRLLALLFLFGYINGLYVMGEF
 Sbjct 61 YLFLSGKSYAMFSLFLGSLFFMQMESQAAGVDFRGRFLWRLLALLFLFGYINGLYVMGEF 120
 Query 121 FMVYAVLGVFLIPLYKVPTRWLLVLCVLLFLQIPAVISFVSLSDNVANEPTAAAYMDR 180
 FMVYAVLGVFLIPLYKVPTRWLLVLCVLLFLQIPAVISFVSLSDNVANEPTAAAYMDR
 Sbjct 121 FMVYAVLGVFLIPLYKVPTRWLLVLCVLLFLQIPAVISFVSLSDNVANEPTAAAYMDR 180
 Query 181 LFERAADVFINGSLMDVLSFNTFDGQSAKCLWVFNFRYLQLLGLFIAGMLIGRQGIHKS 240
 LFERAADVFINGSLMDVLSFNTFDGQSAKCLWVFNFRYLQLLGLFIAGMLIGRQGIHKS
 Sbjct 181 LFERAADVFINGSLMDVLSFNTFDGQSAKCLWVFNFRYLQLLGLFIAGMLIGRQGIHKS 240
 Query 241 EEKMKYSRFLPYCLAFWAVFYAVAFLLPVWGVDFGALRVGQTLFKTYGNLQMMVYFC 300
 EEKMKYSRFLPYCLAFWAVFYAVAFLLPVWGVDFGALRVGQTLFKTYGNLQMMVYFC
 Sbjct 241 EEKMKYSRFLPYCLAFWAVFYAVAFLLPVWGVDFGALRVGQTLFKTYGNLQMMVYFC 300
 Query 301 GTTLLYYRYKQKVDLDRIAPVGRMSVTNYMAQSIIVGVSFLYFGGNGFAVEFNLYQSFLLG 360
 GTTLLYYRYKQKVDLDRIAPVGRMSVTNYMAQSIIVGVSFLYFGGNGFAVEFNLYQSFLLG
 Sbjct 301 GTTLLYYRYKQKVDLDRIAPVGRMSVTNYMAQSIIVGVSFLYFGGNGFAVEFNLYQSFLLG 360
 Query 361 AAFCVIQAISNWWIKRFYYPMEWLWRSITWFOVPLSRRKASLG 406
 AAFCVIQAISNWWIKRFYYPMEWLWRSITWFOVPLSRRKASLG
 Sbjct 361 AAFCVIQAISNWWIKRFYYPMEWLWRSITWFOVPLSRRKASLG 406

>ref|ZP_04841755.1| conserved hypothetical protein [Bacteroides sp. 3_2_5]
 ref|ZP_06093303.1| conserved hypothetical protein [Bacteroides sp. 2_1_16]
 gb|EES88356.1| conserved hypothetical protein [Bacteroides sp. 3_2_5]
 gb|EEZ25846.1| conserved hypothetical protein [Bacteroides sp. 2_1_16]
 Length=412

Score = 822 bits (2123), Expect = 0.0, Method: Compositional matrix adjust.
 Identities = 404/406 (99%), Positives = 405/406 (99%), Gaps = 0/406 (0%)

Query 1 MHTQTITPKKRINSIDALRGFALIGIMLLHCMERFDLTLAPVVEPFWQAITAVYDLSL 60
 MHTQTITPKKRINSIDALRGFALIGIMLLHCMERFDLTLAPVVEPFWQAITAVYDLSL
 Sbjct 7 MHTQTITPKKRINSIDALRGFALIGIMLLHCMERFDLTLAPVVEPFWQAITAVYDLSL 66
 Query 61 YLFLSGKSYAMFSLFLGSLFFMQMESQAAGVDFRGRFLWRLLALLFLFGYINGLYVMGEF 120
 YLFLSGKSYAMFSLFLGSLFFMQMESQAAGVDFRGRFLWRLLALLFLFGYINGLYVMGEF
 Sbjct 67 YLFLSGKSYAMFSLFLGSLFFMQMESQAAGVDFRGRFLWRLLALLFLFGYINGLYVMGEF 126
 Query 121 FMVYAVLGVFLIPLYKVPTRWLLVLCVLLFLQIPAVISFVSLSDNVANEPTAAAYMDR 180
 FMVYAVLGVFLIPLYKVPTRWLLVLCVLLFLQIPAVISFVSLSDNVANEPTAAAYMDR
 Sbjct 127 FMVYAVLGVFLIPLYKVPTRWLLVLCVLLFLQIPAVISFVSLSDNVANEPTAAAYMDR 186
 Query 181 LFERAADVFINGSLMDVLSFNTFDGQSAKCLWVFNFRYLQLLGLFIAGMLIGRQGIHKS 240
 LFERAADVFINGSLMDVLSFNTFDGQSAKCLWVFNFRYLQLLGLFIAGMLIGRQGIHKS
 Sbjct 187 LFERAADVFINGSLMDVLSFNTFDGQSAKCLWVFNFRYLQLLGLFIAGMLIGRQGIHKS 246
 Query 241 EEKMKYSRFLPYCLAFWAVFYAVAFLLPVWGVDFGALRVGQTLFKTYGNLQMMVYFC 300
 EEKMKYSRFLPYCLAFWAVFYAVAFLLPVWGVDFGALRVGQTLFKTYGNLQMMVYFC
 Sbjct 247 EEKMKYSRFLPYCLAFWAVFYAVAFLLPVWGVDFGALRVGQTLFKTYGNLQMMVYFC 306
 Query 301 GTTLLYYRYKQKVDLDRIAPVGRMSVTNYMAQSIIVGVSFLYFGGNGFAVEFNLYQSFLLG 360
 GTTLLYYRYKQKVDLDRIAPVGRMSVTNYMAQSIIVGVSFLYFGGNGFAVEFNLYQSFLLG
 Sbjct 307 GTTLLYYRYKQKVDLDRIAPVGRMSVTNYMAQSIIVGVSFLYFGGNGFAVEFNLYQSFLLG 366
 Query 361 AAFCVIQAISNWWIKRFYYPMEWLWRSITWFOVPLSRRKASLG 406
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 Sbjct 367 AAFCVIQAISNWWIKRFYYPMEWLWRSITWFOVPLSRRKASLG 412

>ref|YP_097636.1| [G](#) putative transport protein [Bacteroides fragilis YCH46]
 dbj|BAD47102.1| [G](#) putative transport protein [Bacteroides fragilis YCH46]
 Length=406

GENE ID: 3081573 BF0353 | putative transport protein
 [Bacteroides fragilis YCH46] (10 or fewer PubMed links)
 Score = 822 bits (2123), Expect = 0.0, Method: Compositional matrix adjust.
 Identities = 405/406 (99%), Positives = 405/406 (99%), Gaps = 0/406 (0%)

Query 1 MHTQTITPKKRINSIDALRGFALIGIMLLHCMERFDLTLAPVVEPFWQAITAVYDLSL 60
 MHTQTITPKKRINSIDALRGFALIGIMLLHCMERFDLTLAPVVEPFWQAITAVYDLSL
 Sbjct 1 MHTQTITPKKRINSIDALRGFALIGIMLLHCMERFDLTLAPVVEPFWQAITAVYDLSL 60
 Query 61 YLFLSGKSYAMFSLFLGSLFFMQMESQAAGVDFRGRFLWRLLALLFLFGYINGLYVMGEF 120
 YLFLSGKSYAMFSLFLGSLFFMQMESQAAGVDFRGRFLWRLLALLFLFGYINGLYVMGEF
 Sbjct 61 YLFLSGKSYAMFSLFLGSLFFMQMESQAAGVDFRGRFLWRLLALLFLFGYINGLYVMGEF 120
 Query 121 FMVYAVLGVFLIPLYKVPTRWLLVLCVLLFLQIPAVISFVSLSDNVANEPTAAAYMDR 180
 FMVYAVLGVFLIPLYKVPTRWLLVLCVLLFLQIPAVISFVSLSDNVANEPTAAAYMDR
 Sbjct 121 FMVYAVLGVFLIPLYKVPTRWLLVLCVLLFLQIPAVISFVSLSDNVANEPTAAAYMDR 180
 Query 181 LFERAADVFINGSLMDVLSFNTFDGQSAKCLWVFNFRYLQLLGLFIAGMLIGRQGIHKS 240
 LFERAADVFINGSLMDVLSFNTFDGQSAKCLWVFNFRYLQLLGLFIAGMLIGRQGIHKS
 Sbjct 181 LFERAADVFINGSLMDVLSFNTFDGQSAKCLWVFNFRYLQLLGLFIAGMLIGRQGIHKS 240
 Query 241 EEKMKYSRFLPYCLAFWAVFYAVAFLLPVWGVDFGALRVGQTLFKTYGNLQMMVYFC 300
 EEKMKYSRFLPYCLAFWAVFYAVAFLLPVWGVDFGALRVGQTLFKTYGNLQMMVYFC
 Sbjct 241 EEKMKYSRFLPYCLAFWAVFYAVAFLLPVWGVDFGALRVGQTLFKTYGNLQMMVYFC 300
 Query 301 GTTLLYYRYKQKVDLDRIAPVGRMSVTNYMAQSIIVGVSFLYFGGNGFAVEFNLYQSFLLG 360
 GTTLLYYRYKQKVDLDRIAPVGRMSVTNYMAQSIIVGVSFLYFGGNGFAVEFNLYQSFLLG

BLAST

Basic Local Alignment Search Tool

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gi|60492160|emb|CAH06923.1| putative anti-sigma...

Results for:

Your BLAST job specified more than one input sequence. This box lets you choose which input sequence to show BLAST results for.

Query ID

[|cl|86207](#)
[|cl|86207](#)

Description

gi|60492160|emb|CAH06923.1| putative anti-sigma factor [Bacteroides fragilis NCTC 9343]

Molecule type

amino acid

Query Length

309

Database Name

nr

Description

All non-redundant GenBank CDS translations+PDB+SwissProt+PIR+PRF excluding environmental samples from WGS projects

Program

BLASTP 2.2.22+ [Citation](#)

Reference

Stephen F. Altschul, Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", *Nucleic Acids Res.* 25:3389-3402.

Reference - compositional score matrix adjustment

Stephen F. Altschul, John C. Wootton, E. Michael Gertz, Richa Agarwala, Aleksandr Morgulis, Alejandro A. Schäffer, and Yi-Kuo Yu (2005) "Protein database searches using compositionally adjusted substitution matrices", *FEBS J.* 272:5101-5109.

Other reports: [Search Summary](#) [[Taxonomy reports](#)] [[Distance tree of results](#)] [[Multiple alignment](#)] **NEW**
[Search Parameters](#)

Search parameter name Search parameter value

Program	blastp
Word size	3
Expect value	10
Hittlist size	100
Capcosts	11,1
Matrix	BLOSUM62
Filter string	F
Genetic Code	1
Window Size	40
Threshold	11
Composition-based stats	2

Database

Database parameter name Database parameter value

Posted date	Jan 20, 2010 4:14 PM
Number of letters	3,542,056,219
Number of sequences	10,381,779
Entrez query	none

Karlin-Altschul statistics

Params Ungapped Gapped

Lambda	0.319277	0.267
K	0.135008	0.041
H	0.381721	0.14

Results Statistics

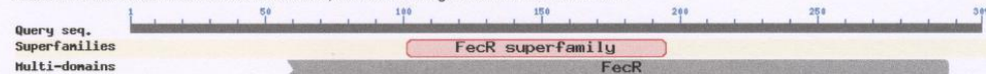
Results Statistics parameter name Results Statistics parameter value

Length adjustment	136
Effective length of query	173
Effective length of database	2130134275
Effective search space	368513229575
Effective search space used	368513229575

Graphic Summary

Show Conserved Domains

Putative conserved domains have been detected, click on the image below for detailed results.



Alignments [Select All](#) [Get selected sequences](#) [Distance tree of results](#) [Multiple alignment](#) [NEW](#)

```

>ref|YP_098536.1| G putative anti-sigma factor [Bacteroides fragilis YCH46]
ref|ZP_06091266.1| conserved hypothetical protein [Bacteroides sp. 2_1_16]
dbj|BAD48002.1| G putative anti-sigma factor [Bacteroides fragilis YCH46]
gb|EEZ26652.1| conserved hypothetical protein [Bacteroides sp. 2_1_16]
Length=321

GENE ID: 3082788 BF1252 | putative anti-sigma factor
[Bacteroides fragilis YCH46] (10 or fewer PubMed links)

Score = 635 bits (1638), Expect = 2e-180, Method: Compositional matrix adjust.
Identities = 309/309 (100%), Positives = 309/309 (100%), Gaps = 0/309 (0%)

Query 1 MTCEEKDLFDRIE SDEALKKEFLRMQNVVALTQILSRQDDSETSRKGGQHFMOQLLFRKR 60
Sbjct 13 MTCEEKDLFDRIE SDEALKKEFLRMQNVVALTQILSRQDDSETSRKGGQHFMOQLLFRKR 72

Query 61 LKRAITVSLKYAAVFAVLVVGTFYTA KLYLSEEFGKSYTIVTAPKQORVKIELPDGTIAW 120
Sbjct 73 LKRAITVSLKYAAVFAVLVVGTFYTA KLYLSEEFGKSYTIVTAPKQORVKIELPDGTIAW 132

Query 121 LSPCSRRLRFAASFNETDRKIELDGATYFDVAKNPEKPFVVSAGYRIRVLGTFKFNISAYK 180
Sbjct 133 LSPCSRRLRFAASFNETDRKIELDGATYFDVAKNPEKPFVVSAGYRIRVLGTFKFNISAYK 192

Query 181 NSKEFETDLVEGCVHIYDPADIRNEVF LQPKKAVLWGDRLMKRESDFDNEEYLNKGVVS 240
Sbjct 193 NSKEFETDLVEGCVHIYDPADIRNEVF LQPKKAVLWGDRLMKRESDFDNEEYLNKGVVS 252

Query 241 FLSEPFGRVLSVALWNDVNIKIERSVNATQ RISGKFRQSDSLESILKALQGAMPFKYKI 300
Sbjct 253 FLSEPFGRVLSVALWNDVNIKIERSVNATQ RISGKFRQSDSLESILKALQGAMPFKYKI 312

Query 301 VSEEEIIIIY 309
Sbjct 313 VSEEEIIIIY 321

>ref|YP_210870.1| G putative anti-sigma factor [Bacteroides fragilis NCTC 9343]
emb|CAH06923.1| G putative anti-sigma factor [Bacteroides fragilis NCTC 9343]
Length=309

GENE ID: 3288859 BF1203 | putative anti-sigma factor
[Bacteroides fragilis NCTC 9343] (10 or fewer PubMed links)

Score = 635 bits (1637), Expect = 3e-180, Method: Compositional matrix adjust.
Identities = 309/309 (100%), Positives = 309/309 (100%), Gaps = 0/309 (0%)

Query 1 MTCEEKDLFDRIE SDEALKKEFLRMQNVVALTQILSRQDDSETSRKGGQHFMOQLLFRKR 60
Sbjct 1 MTCEEKDLFDRIE SDEALKKEFLRMQNVVALTQILSRQDDSETSRKGGQHFMOQLLFRKR 60

Query 61 LKRAITVSLKYAAVFAVLVVGTFYTA KLYLSEEFGKSYTIVTAPKQORVKIELPDGTIAW 120
Sbjct 61 LKRAITVSLKYAAVFAVLVVGTFYTA KLYLSEEFGKSYTIVTAPKQORVKIELPDGTIAW 120

Query 121 LSPCSRRLRFAASFNETDRKIELDGATYFDVAKNPEKPFVVSAGYRIRVLGTFKFNISAYK 180
Sbjct 121 LSPCSRRLRFAASFNETDRKIELDGATYFDVAKNPEKPFVVSAGYRIRVLGTFKFNISAYK 180

Query 181 NSKEFETDLVEGCVHIYDPADIRNEVF LQPKKAVLWGDRLMKRESDFDNEEYLNKGVVS 240
Sbjct 181 NSKEFETDLVEGCVHIYDPADIRNEVF LQPKKAVLWGDRLMKRESDFDNEEYLNKGVVS 240

Query 241 FLSEPFGRVLSVALWNDVNIKIERSVNATQ RISGKFRQSDSLESILKALQGAMPFKYKI 300
Sbjct 241 FLSEPFGRVLSVALWNDVNIKIERSVNATQ RISGKFRQSDSLESILKALQGAMPFKYKI 300

Query 301 VSEEEIIIIY 309
Sbjct 301 VSEEEIIIIY 309

>ref|ZP_04840883.1| conserved hypothetical protein [Bacteroides sp. 3_2_5]
gb|EESB7484.1| conserved hypothetical protein [Bacteroides sp. 3_2_5]
Length=321

Score = 631 bits (1627), Expect = 3e-179, Method: Compositional matrix adjust.
Identities = 308/309 (99%), Positives = 308/309 (99%), Gaps = 0/309 (0%)

Query 1 MTCEEKDLFDRIE SDEALKKEFLRMQNVVALTQILSRQDDSETSRKGGQHFMOQLLFRKR 60
Sbjct 13 MTCEEKDLFDRIE SDEALKKEFLRMQNVVALTQILS QDDSETSRKGGQHFMOQLLFRKR 72

Query 61 LKRAITVSLKYAAVFAVLVVGTFYTA KLYLSEEFGKSYTIVTAPKQORVKIELPDGTIAW 120
Sbjct 73 LKRAITVSLKYAAVFAVLVVGTFYTA KLYLSEEFGKSYTIVTAPKQORVKIELPDGTIAW 132

Query 121 LSPCSRRLRFAASFNETDRKIELDGATYFDVAKNPEKPFVVSAGYRIRVLGTFKFNISAYK 180
Sbjct 133 LSPCSRRLRFAASFNETDRKIELDGATYFDVAKNPEKPFVVSAGYRIRVLGTFKFNISAYK 192

Query 181 NSKEFETDLVEGCVHIYDPADIRNEVF LQPKKAVLWGDRLMKRESDFDNEEYLNKGVVS 240
Sbjct 193 NSKEFETDLVEGCVHIYDPADIRNEVF LQPKKAVLWGDRLMKRESDFDNEEYLNKGVVS 252

Query 241 FLSEPFGRVLSVALWNDVNIKIERSVNATQ RISGKFRQSDSLESILKALQGAMPFKYKI 300
Sbjct 253 FLSEPFGRVLSVALWNDVNIKIERSVNATQ RISGKFRQSDSLESILKALQGAMPFKYKI 312

Query 301 VSEEEIIIIY 309
Sbjct 313 VSEEEIIIIY 321

>ref|ZP_05280128.1| putative anti-sigma factor [Bacteroides fragilis 3_1_12]
Length=321

Score = 586 bits (1511), Expect = 1e-165, Method: Compositional matrix adjust.
Identities = 279/309 (90%), Positives = 293/309 (94%), Gaps = 0/309 (0%)
    
```

Descriptions

Legend for links to other resources: [U](#) UniGene [E](#) GEO [G](#) Gene [S](#) Structure [M](#) Map Viewer

Sequences producing significant alignments:

(Click headers to sort columns)

YP_098536.1	putative anti-sigma factor [Bacteroides fragilis YCH46] >ref ZP_06091266.1 conserved hypothetical protein [Bacteroides sp. 2_1_16] >dbj BAD48002.1 putative anti-sigma factor [Bacteroides fragilis YCH46] >gb EEZ26652.1 conserved hypothetical protein [Bacteroides sp. 2_1_16]	635	635	100%	2e-180	G
YP_210870.1	putative anti-sigma factor [Bacteroides fragilis NCTC 9343] >emb CAH06923.1 putative anti-sigma factor [Bacteroides fragilis NCTC 9343]	635	635	100%	3e-180	G
ZP_04840883.1	conserved hypothetical protein [Bacteroides sp. 3_2_5] >gb EES87484.1 conserved hypothetical protein [Bacteroides sp. 3_2_5]	631	631	100%	3e-179	
ZP_05280128.1	putative anti-sigma factor [Bacteroides fragilis 3_1_12]	586	586	100%	1e-165	
ZP_05283939.1	hypothetical protein Bfra3_21925 [Bacteroides fragilis 3_1_12]	252	252	100%	3e-65	
YP_101246.1	putative anti-sigma factor [Bacteroides fragilis YCH46] >ref YP_213335.1 hypothetical protein BF3745 [Bacteroides fragilis NCTC 9343] >ref ZP_04844595.1 conserved hypothetical protein [Bacteroides sp. 3_2_5] >dbj BAD50712.1 putative anti-sigma factor [Bacteroides fragilis YCH46] >emb CAH09426.1 putative membrane protein [Bacteroides fragilis NCTC 9343] >gb EES84495.1 conserved hypothetical protein [Bacteroides sp. 3_2_5]	250	250	100%	1e-64	G
ZP_06095273.1	conserved hypothetical protein [Bacteroides sp. 2_1_16] >gb EEZ23941.1 conserved hypothetical protein [Bacteroides sp. 2_1_16]	250	250	100%	1e-64	
NP_811110.1	putative anti-sigma factor [Bacteroides thetaiotaomicron VPI-5482] >g AAO77304.1 putative anti-sigma factor [Bacteroides thetaiotaomicron VPI-5482]	230	230	99%	2e-58	G
ZP_03476000.1	hypothetical protein PRBACTJOHN_01664 [Parabacteroides johnsonii DSM 18315] >gb EEC96931.1 hypothetical protein PRBACTJOHN_01664 [Parabacteroides johnsonii DSM 18315]	180	180	98%	2e-43	
ZP_03677961.1	hypothetical protein BACCELL_02300 [Bacteroides cellulosilyticus DSM 14838] >gb EEF90053.1 hypothetical protein BACCELL_02300 [Bacteroides cellulosilyticus DSM 14838]	176	176	96%	4e-42	
ZP_02034096.1	hypothetical protein PARMER_04138 [Parabacteroides merdae ATCC 43184] >gb EDN84686.1 hypothetical protein PARMER_04138 [Parabacteroides merdae ATCC 43184]	168	168	99%	8e-40	
ZP_05414155.1	putative anti-sigma factor [Bacteroides finegoldii DSM 17565] >gb EEK46802.1 putative anti-sigma factor [Bacteroides finegoldii DSM 17565]	161	161	99%	7e-38	
ZP_04544887.1	conserved hypothetical protein [Bacteroides sp. D1] >ref ZP_06084784.1 conserved hypothetical protein [Bacteroides sp. 2_1_22] >gb EEO51361.1 conserved hypothetical protein [Bacteroides sp. D1] >gb EEZ02882.1 conserved hypothetical protein [Bacteroides sp. 2_1_22]	161	161	99%	8e-38	
ZP_05758326.1	putative anti-sigma factor [Bacteroides sp. D2]	159	159	99%	3e-37	
ZP_03677876.1	hypothetical protein BACCELL_02215 [Bacteroides cellulosilyticus DSM 14838] >gb EEF90135.1 hypothetical protein BACCELL_02215 [Bacteroides cellulosilyticus DSM 14838]	154	154	95%	1e-35	
ZP_03678231.1	hypothetical protein BACCELL_02574 [Bacteroides cellulosilyticus DSM 14838] >gb EEF89800.1 hypothetical protein BACCELL_02574 [Bacteroides cellulosilyticus DSM 14838]	154	154	99%	2e-35	
ZP_02435836.1	hypothetical protein BACSTE_02087 [Bacteroides stercoris ATCC 43183] >gb EDS14404.1 hypothetical protein BACSTE_02087 [Bacteroides stercoris ATCC 43183]	153	153	99%	2e-35	
ZP_02068886.1	hypothetical protein BACUNI_00286 [Bacteroides uniformis ATCC 8492] >gb EDO56000.1 hypothetical protein BACUNI_00286 [Bacteroides uniformis ATCC 8492]	152	152	99%	3e-35	
ZP_03014123.1	hypothetical protein BACINT_01686 [Bacteroides intestinalis DSM 17393] >gb EDV06597.1 hypothetical protein BACINT_01686 [Bacteroides intestinalis DSM 17393]	152	152	99%	3e-35	
ZP_03299904.1	hypothetical protein BACDOR_01271 [Bacteroides dorei DSM 17855] >ref ZP_04554581.1 conserved hypothetical protein [Bacteroides sp. D4] >gb EEB26249.1 hypothetical protein BACDOR_01271 [Bacteroides dorei DSM 17855] >gb EE047637.1 conserved hypothetical protein [Bacteroides sp. D4]	152	152	87%	5e-35	
ZP_03477618.1	hypothetical protein PRBACTJOHN_03305 [Parabacteroides johnsonii DSM 18315] >gb EEC95310.1 hypothetical protein PRBACTJOHN_03305 [Parabacteroides johnsonii DSM 18315]	152	152	96%	5e-35	
YP_001304665.1	putative anti-sigma factor [Parabacteroides distasonis ATCC 8503] >gb ABR45043.1 putative anti-sigma factor [Parabacteroides distasonis ATCC 8503]	152	152	95%	5e-35	G
YP_001297833.1	putative anti-sigma factor [Bacteroides vulgatus ATCC 8482] >gb ABR38211.1 putative anti-sigma factor [Bacteroides vulgatus ATCC 8482]	152	152	87%	6e-35	G
ZP_05255078.1	conserved hypothetical protein [Bacteroides sp. 4_3_47FAA] >gb EET15470.1 conserved hypothetical protein [Bacteroides sp. 4_3_47FAA]	151	151	87%	8e-35	
YP_101600.1	putative anti-sigma factor [Bacteroides fragilis YCH46] >ref YP_213696.1 putative anti-sigma factor [Bacteroides fragilis NCTC 9343] >ref ZP_04844033.1 conserved hypothetical protein [Bacteroides sp. 3_2_5] >ref ZP_06095115.1 conserved hypothetical protein [Bacteroides sp. 2_1_16] >dbj BAD51066.1 putative anti-sigma factor [Bacteroides fragilis YCH46] >emb CAH09804.1 putative anti-sigma factor [Bacteroides fragilis NCTC 9343] >gb EES85227.1 conserved hypothetical protein [Bacteroides sp. 3_2_5] >gb EEZ24266.1 conserved hypothetical protein [Bacteroides	150	150	81%	1e-34	G

BLAST**Basic Local Alignment Search Tool**

•

[Edit and Resubmit](#) [Save Search Strategies](#) [Formatting options](#) [Download](#)**gi|60681974|ref|YP_212118.1| hypothetical...**Results for:

Your BLAST job specified more than one input sequence. This box lets you choose which input sequence to show BLAST results for.

Query ID[lcl|73919](#)

lcl|73919

Description

gi|60681974|ref|YP_212118.1| hypothetical protein BF2494 [Bacteroides fragilis NCTC 9343]

Molecule type

amino acid

Query Length

396

Database Name

nr

Description

All non-redundant GenBank CDS translations+PDB+SwissProt+PIR+PRF excluding environmental samples from WGS projects

ProgramBLASTP 2.2.22+ [Citation](#)**Reference**Stephen F. Altschul, Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", *Nucleic Acids Res.* 25:3389-3402.[Reference - compositional score matrix adjustment](#)Stephen F. Altschul, John C. Wootton, E. Michael Gertz, Richa Agarwala, Aleksandr Morgulis, Alejandro A. Schäffer, and Yi-Kuo Yu (2005) "Protein database searches using compositionally adjusted substitution matrices", *FEBS J.* 272:5101-5109.Other reports: [Search Summary](#) [[Taxonomy reports](#)] [[Distance tree of results](#)] [[Related Structures](#)] [[Multiple alignment](#)] **NEW**[Search Parameters](#)**Search parameter name Search parameter value**

Program	blastp
Word size	3
Expect value	10
Hittist size	100
Gapcosts	11,1
Matrix	BLOSUM62
Filter string	F
Genetic Code	1
Window Size	40
Threshold	11
Composition-based stats	2

Database

Database parameter name Database parameter value

Posted date	Jan 20, 2010 4:14 PM
Number of letters	3,542,056,219
Number of sequences	10,381,779
Entrez query	none

Karlin-Altschul statistics

Params Ungapped Gapped

Lambda	0.314287	0.267
K	0.13096	0.041
H	0.371554	0.14

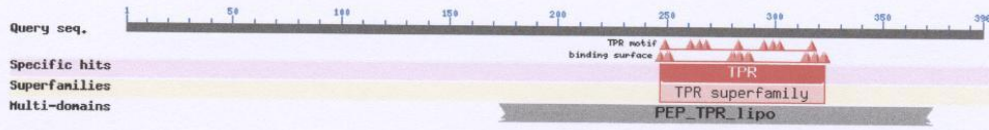
Results Statistics

Results Statistics parameter name Results Statistics parameter value

Length adjustment	139
Effective length of query	257
Effective length of database	2098988938
Effective search space	539440157066
Effective search space used	539440157066

Graphic Summary[Show Conserved Domains](#)

Putative conserved domains have been detected, click on the image below for detailed results.



Distribution of 140 Blast Hits on the Query Sequence

[?]

An overview of the database sequences aligned to the query sequence is shown. The score of each alignment is indicated by one of five different colors, which divides the range of scores into five groups. Multiple alignments on the same database sequence are connected by a striped line. Mousing over a hit sequence causes the definition and score to be shown in the window at the top, clicking on a hit sequence takes the user to the associated alignments. New: This graphic is an overview of database sequences aligned to the query sequence. Alignments are color-coded by score, within one of five score ranges. Multiple alignments on the same database sequence are connected by a dashed line. Mousing over an alignment shows the alignment definition and score in the box at the top. Clicking an alignment displays the alignment detail.



Descriptions

Sequences producing significant alignments:	Score (Bits)	E Value		
ref YP_099695.1	TPR repeat-containing protein [Bacteroides f...	795	0.0	G
ref YP_212118.1	hypothetical protein BF2494 [Bacteroides fra...	795	0.0	G
ref ZP_05281526.1	TPR repeat-containing protein [Bacteroides...	775	0.0	
ref ZP_05417548.2	tetratricopeptide repeat family protein [B...	624	9e-177	
ref ZP_01958966.1	hypothetical protein BACCAC_00554 [Bactero...	621	5e-176	
ref ZP_05756644.1	TPR repeat-containing protein [Bacteroides...	620	1e-175	
ref ZP_02063765.1	hypothetical protein BACOVA_00723 [Bactero...	620	1e-175	
ref ZP_04546993.1	TPR repeat-containing protein [Bacteroides...	617	9e-175	
ref ZP_04845680.1	TPR repeat-containing protein [Bacteroides...	598	3e-169	
ref NP_809813.1	TPR repeat-containing protein [Bacteroides t...	598	3e-169	G
ref ZP_03459917.1	hypothetical protein BACEGG_02718 [Bactero...	596	2e-168	
ref ZP_02435236.1	hypothetical protein BACSTE_01478 [Bactero...	592	2e-167	
ref ZP_06202893.1	conserved hypothetical protein [Bacteroides...	586	1e-165	
ref ZP_02070574.1	hypothetical protein BACUNI_01995 [Bactero...	586	2e-165	
ref ZP_03680531.1	hypothetical protein BACCCELL_04904 [Bacter...	585	4e-165	
ref ZP_03014415.1	hypothetical protein BACINT_01988 [Bactero...	567	9e-160	
ref ZP_03010137.1	hypothetical protein BACCOP_02007 [Bactero...	256	3e-66	
ref ZP_03644242.1	hypothetical protein BACCOPRO_02619 [Bacte...	247	2e-63	
ref YP_001300548.1	TPR repeat-containing protein [Bacteroides...	247	2e-63	G
ref ZP_03301317.1	hypothetical protein BACDOR_02699 [Bactero...	246	2e-63	
ref ZP_05254725.1	TPR domain-containing protein [Bacteroides...	246	5e-63	
ref ZP_06074897.1	TPR domain-containing protein [Bacteroides...	240	2e-61	
ref YP_001302476.1	TPR domain-containing protein [Parabacter...	238	7e-61	G
ref ZP_05545502.1	TPR domain-containing protein [Parabactero...	238	7e-61	
ref ZP_03207140.1	hypothetical protein BACPLE_00760 [Bactero...	238	1e-60	
ref ZP_06253438.1	putative TPR domain protein [Prevotella co...	230	2e-58	
ref ZP_06005642.2	conserved hypothetical protein [Prevotella...	211	2e-52	
ref ZP_06255209.1	putative TPR domain protein [Prevotella or...	206	5e-51	
ref ZP_04833032.1	putative TPR domain protein [Prevotella me...	205	9e-51	
ref ZP_06290121.1	conserved hypothetical protein [Prevotella...	198	7e-49	
ref ZP_05857835.1	putative TPR domain protein [Prevotella ve...	197	2e-48	
ref ZP_04055111.1	TPR domain protein [Porphyromonas uenonis ...	195	7e-48	
ref ZP_02031759.1	hypothetical protein PARMER_01764 [Parabac...	190	2e-46	
ref YP_001928992.1	TPR domain protein [Porphyromonas gingiva...	186	5e-45	G
ref NP_905547.1	TPR domain-containing protein [Porphyromonas...	184	2e-44	G
ref ZP_06287662.1	conserved hypothetical protein [Prevotella...	182	7e-44	
ref ZP_06269395.1	conserved hypothetical protein [Prevotella...	181	2e-43	
ref ZP_03475728.1	hypothetical protein PRABACTJOHN_01391 [Pa...	171	2e-40	
ref ZP_05918007.1	conserved hypothetical protein [Prevotella...	168	8e-40	
ref ZP_05369116.1	TPR domain protein [Propionibacterium sp. ...	147	2e-33	
ref ZP_04390828.1	TPR domain protein [Porphyromonas endodont...	147	2e-33	
ref ZP_05734739.1	putative tetratricopeptide repeat-contain...	137	3e-30	
ref ZP_06255207.1	putative TPR domain protein [Prevotella or...	132	8e-29	
ref ZP_06288289.1	tetratricopeptide repeat protein [Prevotel...	122	9e-26	
ref ZP_05918006.1	conserved hypothetical protein [Prevotella...	114	2e-23	
ref ZP_05734740.1	putative TPR domain protein [Prevotella ta...	108	8e-22	
ref ZP_05857833.1	putative TPR domain protein [Prevotella ve...	108	9e-22	
ref ZP_06287663.1	tetratricopeptide repeat protein [Prevotel...	106	4e-21	
ref ZP_04833030.1	putative TPR domain protein [Prevotella me...	104	2e-20	
ref ZP_06269393.1	tetratricopeptide repeat protein [Prevotel...	100	2e-19	
ref ZP_06255208.1	hypothetical protein HMPREF0971_01238 [Pre...	100	4e-19	
ref YP_100914.1	hypothetical protein BF3637 [Bacteroides fra...	95.5	9e-18	G
ref ZP_04843481.1	conserved hypothetical protein [Bacteroides...	95.5	1e-17	
ref ZP_03676726.1	hypothetical protein BACCCELL_01053 [Bacter...	94.0	2e-17	
ref ZP_05283200.1	hypothetical protein Bfra3_18177 [Bacteroi...	92.4	8e-17	
ref ZP_06005641.2	conserved hypothetical protein [Prevotella...	89.7	5e-16	
ref ZP_03209390.1	hypothetical protein BACPLE_03064 [Bactero...	89.4	7e-16	
ref ZP_04833031.1	TPR domain protein [Prevotella melaninogen...	87.4	3e-15	
ref ZP_06251203.1	putative TPR domain protein [Prevotella co...	84.7	2e-14	
ref ZP_05857834.1	TPR domain protein [Prevotella veroralis F...	83.6	4e-14	
ref ZP_01718101.1	hypothetical protein ALPR1_12965 [Algoriph...	67.0	4e-09	
ref ZP_02424158.1	hypothetical protein ALIPUT_00273 [Alistip...	63.2	6e-08	
ref ZP_06269394.1	tetratricopeptide repeat protein [Prevotel...	62.8	6e-08	
ref YP_003290272.1	Tetratricopeptide TPR_2 repeat protein [R...	62.4	1e-07	G
ref YP_861545.1	TPR repeat-containing protein [Gramella fors...	59.7	6e-07	G
ref ZP_02177808.1	hypothetical protein HG1285_15791 [Hydroge...	59.3	8e-07	
pdb 1NA0 A	Chain A, Design Of Stable Alpha-Helical Arrays Fro...	58.2	2e-06	S
ref YP_001933073.1	hypothetical protein TPASS_0067 [Treponem...	57.8	2e-06	G
ref NP_218507.1	hypothetical protein TP0067 [Treponema palli...	57.8	2e-06	G
ref ZP_05623358.1	TPR domain protein [Treponema vincentii AT...	56.2	6e-06	
ref NP_972674.1	TPR domain-containing protein [Treponema den...	55.8	9e-06	G
ref YP_003125075.1	hypothetical protein Cpin_5445 [Chitinoph...	55.8	9e-06	G
ref YP_001156602.1	TPR repeat-containing protein [Polynucleo...	55.1	1e-05	G
ref YP_001952243.1	Tetratricopeptide TPR_2 repeat protein [G...	54.7	2e-05	G
ref ZP_01463944.1	adventurous gliding protein T [Stigmatella...	54.7	2e-05	
ref YP_720410.1	TPR repeat-containing protein [Trichodesmium...	54.7	2e-05	G
pdb 2F07 A	Chain A, Crystal Structure Of An 8 Repeat Consensu...	54.7	2e-05	S
ref YP_002493579.1	TPR repeat-containing protein [Anaeromyxo...	54.3	2e-05	G
ref XP_001026739.1	TPR Domain containing protein [Tetrahyemen...	53.9	3e-05	G
ref YP_466187.1	TPR repeat-containing protein [Anaeromyxobac...	53.9	3e-05	G
ref YP_001996298.1	Tetratricopeptide TPR_2 repeat protein [C...	53.9	4e-05	G
ref ZP_01048923.1	TPR repeat protein [Dokdonia donghaensis M...	53.1	5e-05	

H Crystal screen compositions

1	50% (w/v) PEG 400	0.2 M Lithium Sulphate	0.1 M Sodium acetate pH 5.1	
2	20% (w/v) PEG 3000	0.1 M Sodium citrate pH 5.5		
3	20% (w/v) PEG 3350	0.2 M Diammonium hydrogen citrate pH 5.0		
4	30% (v/v) MPD	0.08 M Calcium chloride	0.1 M Sodium acetate pH 4.6	
5	20% (w/v) PEG 3350	0.2 M Magnesium formate pH 5.9		
6	20% (w/v) PEG 1000	0.2 M lithium sulphate	0.25 M Sodiumcitrate pH 4.2	0.25 M sodium dihydrogen phosphate pH 5.2
7	20% (w/v) PEG 8000	0.1M CHES pH 9.5		
8	20% (w/v) PEG 3350	0.2 M Ammonium formate pH6.6		
9	20% (w/v) PEG 3350	0.2 M Ammonium chloride pH 6.3		
10	20% (w/v) PEG 3350	0.2 M Potassium formate pH 7.3		
11	50% (v/v) MPD	0.2 M Ammonium dihydrogen phosphate	0.1M Tris pH 8.5	
12	20% (w/v) PEG 3350	0.2 M Potassium nitrate pH 6.9		
13	0.8 M Ammonium sulphate	0.1M Citric acid pH 4.0		
14	20% (w/v) PEG 3350	0.2 M Sodium		

		thiocyanate pH 6.9		
15	20% (w/v) PEG 6000	0.1M Bicine pH 9.0		
16	10% (w/v) PEG 8000	8% (v/v) Ethylene glycol	0.1 M HEPES pH 7.5	
17	40% (v/v) MPD	5% (w/v) PEG 8000	0.1M Sodium cacodylate pH 7.0	
18	40% (v/v) Ethanol	5% (w/v) PEG 1000	0.25 M Sodium citrate pH 5.2	0.25 M Sodium dihydrogen phosphate pH 5.2
19	8% (w/v) PEG 4000	0.1M Sodium acetate pH 4.6		
20	10% (w/v) PEG 8000	0.2 M Magnesium chloride	0.1 M Tris pH 7.0	
21	20% (w/v) PEG 6000	0.1M Citric acid pH 5.0		
22	50% (w/v) PEG 200	0.2 M Magnesium chloride	0.1 M Sodium cacodylate pH 6.6	
23	1.6 M Sodium citrate pH 6.5			
24	20% (w/v) PEG 3350	0.2 M Potassium citrate pH 8.3		
25	30% (v/v) MPD	0.02 M Calcium chloride	0.1M Sodium acetate pH 4.6	
26	20% (w/v) PEG 8000	0.2 M Sodium chloride	0.25 M Sodium citrate pH 4.2	0.25 M Sodium dihydrogen phosphate pH 4.2
27	20% (w/v) PEG 6000	1.0 M Lithium chloride	0.1 M Citric acid pH 4.0	
28	20% (w/v) PEG 3350	0.2 M Ammonium nitrate pH 6.3		

29	10% (w/v) PEG 6000	0.1M HEPES pH 7.0		
30	0.8 M Ammonium dihydrogen phosphate	0.8 M Potassium dihydrogen phosphate	0.1M HEPES pH 7.5	
31	40% (w/v) PEG 300	0.25 M Sodium citrate pH 5.2	0.25 M Sodium dihydrogen phosphate pH 5.2	
32	10% (w/v) PEG 3000	0.2 M Zinc acetate	0.1 M Sodium acetate pH 4.5	
33	20% (v/v) Ethanol	0.1 M Tris pH 8.5		
34	25% (v/v) 1-2-Propanediol	0.1M Sodium potassium phosphate pH 6.8	10% (v/v) glycerol	
35	10% (w/v) PEG 20000	2% (v/v) Dioxane	0.1M Bicine pH 9.0	
36	2.0 M Ammonium sulphate	0.1M Sodium acetate pH 4.6		
37	10% (w/v) PEG 1000	10% (w/v) PEG 8000		
38	24% (w/v) PEG 1000	20% (v/v) Glycerol		
39	30% (v/v) PEG 400	0.2 M Magnesium chloride	0.1M HEPES pH 7.5	
40	50% (w/v) PEG 200	0.2 M Sodium chloride	0.1 M Sodium potassium phosphate pH 7.2	
41	30% (w/v) PEG 8000	0.2 M Lithium sulphate	0.1M Sodium acetate pH 4.5	
42	70% (v/v) MPD	0.2 M Magnesium chloride	0.1 M HEPES pH 7.5	
43	20% (w/v) PEG 8000	0.1M Tris pH 8.5		

44	40% (v/v) PEG 400	0.2 M Lithium sulphate	0.1 M Tris pH 8.4	
45	40%(v/v) MPD	0.1M Tris pH 8.0		
46	25.5% (w/v) PEG 4000	0.17 M Ammonium sulphate	15% (v/v) Glycerol	
47	40% (w/v) PEG 300	0.2 M Calcium acetate	0.1M Sodium cacodylate pH 7.0	
48	14% (v/v) Isopropanol	0.14 M Calcium chloride	0.07M Sodium acetate pH 4.6	30% (v/v) glycerol
49	16% (w/v)PEG 8000	0.04 M Potassium dihydrogen phosphate	20% (v/v) Glycerol	
50	1.0 M Sodium citrate	0.1 M sodium cacodylate pH 6.5		
51	2.0 M Ammonium sulphate	0.2 M Sodium chloride	0.1 M Sodium cacodylate pH 6.5	
52	10% (v/v) Isopropanol	0.2 M Sodium chloride	0.1 M HEPES pH 7.5	
53	1.26 M Ammonium sulphate	0.2 M Lithium sulphate	0.1M Tris pH 8.5	
54	40% (v/v) MPD	0.1M CAPS pH 10.1		
55	20% (w/v)PEG 3000	0.2 M Zinc acetate	0.1M Imidazole pH 8.0	
56	10% (v/v) Isopropanol	0.2 M Zinc acetate	0.1M Sodium cacodylate pH 6.5	
57	1.0M Diammonium hydrogen phosphate	0.1 M Sodium acetate pH 4.5		
58	1.6 M Magnesium sulphate	0.1M MES pH 6.5		

59	10% (w/v) PEG 6000	0.1 M Bicine pH 9.0		
60	14.4% (w/v) PEG 8000	0.16 M Calcium acetate	0.08 M Sodium cacodylate pH 6.5	20% (v/v) glycerol
61	10% (w/v) PEG 8000	0.1 M Imidazol pH 8.0		
62	30% (v/v) Jeffamine	0.05 M Caesiumchloride	0.1 M MES pH 6.5	
63	3.2 M Ammonium sulphate	0.1 M Citric acid pH 5.0		
64	20% (v/v) MPD	0.1 M Tris pH 8.0		
65	20% (v/v) Jeffamine	0.1M HEPES pH 6.5		
66	50% (v/v) Ethylene glycol	0.2 M Magnesium chloride	0.1 M Tris pH 8.5	
67	10% (v/v) MPD	0.1 M Bicine pH 9.0		
68	0.2 M Ammonium sulphate	0.1 M Sodium acetate pH 4.6	30% (w/v) PEG MME 2000	
69	0.2 M Ammonium sulphate	0.1 M MES pH 6.5	30% (w/v) PEG MME 5000	
70	0.01 M Zinc sulphate	0.1 M MES pH 6.5	25% (w/v) PEG MME 550	
71	0.01 M Nickel chloride	0.1 M Tris pH 8.5	20% (w/v) PEG MME 2000	
72	0.1 M Sodium chloride	0.1 M Bicine pH 9.0	20% (w/v) PEG MME 550	
73	0.005 M Magnesium chloride	0.05 M HEPES pH 7.0	25% (w/v) PEG MME 550	

74	0.1 M Potassium chloride	0.015 M Magnesium chloride	0.05 M Tris pH 7.5	10% PEG MME 550
75	20% (v/v) 1-4-Butandiol	0.1 M MES pH 6.0	0.2 M Lithium sulphate	
76	1 M Sodium potassium tartrate	0.1 M Imidazole pH 8.0	0.2 M Sodium chloride	
77	20% (v/v) 1-4-Butandiol	0.1 M Sodium acetate pH 4.5		
78	1 M Sodium potassium tartrate	0.1 M CHES pH 9.5	0.2 M Lithium sulphate	
79	35% (v/v) Propanol	0.1 M Sodium cacodylate pH 6.5		
80	35% (v/v) Propanol	0.1 M Tris pH 8.5		
81	3.5 M Sodium formate			
82	0.8 M Succinic acid pH 7.0			
83	2.1 M Malic acid pH 7.0			
84	2.4 M Sodium malonate pH 7.0			
85	0.2 M Potassium chloride	0.05 M HEPES pH 7.5	35% (v/v) Pentaerythritol propoxylate	
86	0.005 M Ammonium sulphate	0.05 M Tris pH 6.5	30% (v/v) Pentaerythritol ethoxylate	
87	0.2 M Potassium	25% (w/v) PEG MME	0.1 M HEPES pH 7.5	

	bromide	2000		
88	0.2M Potassium bromide	8% (w/v) PEG 20000	8% (w/v) PEG MME 550	0.1 M Tris pH 8.5
89	1.0 M Potassium dihydrogen phosphate	0.1 M Sodium citrate pH 4.6		
90	0.5 M Potassium dihydrogen phosphate	0.1M HEPES pH 7.0		
91	20% (w/v) PEG 4000	0.005 M Cadmium chloride	0.1 M Tris pH 8.0	
92	20% (w/v) PEG 4000	0.005 M Nickel chloride	0.1 M MED pH 6.5	
93	0.8 M Sodium formate	10% (w/v) PEG 8000	10% PEG 1000	0.1 M imidazol pH 8.0
94	15% (w/v) PEG 4000	0.005 M Cadmium sulphate	0.1 M Sodium cacodylate pH 6.5	
95	20% (w/v) PEG 600	0.005 M Cobalt chloride	0.1 M HEPES pH 7.5	
96	2 M Ammonium sulphate	10% (v/v) Jeffamine	0.1 M Tris pH 8.0	

Newcastle Crystal screen formulation

PACT crystal screen formulations

PEG/Anion Screens

Well number	Salt	Buffer	PEG
1	0.2 M Sodium fluoride		20% (w/v) PEG 3350
2	0.2 M Sodium bromide		20% (w/v) PEG 3350
3	0.2 M Sodium iodide		20% (w/v) PEG 3350
4	0.2 M Potassium thiocyanate		20% (w/v) PEG 3350
5	0.2 M Sodium nitrate		20% (w/v) PEG 3350
6	0.2 M Sodium formate		20% (w/v) PEG 3350
7	0.2 M Sodium acetate		20% (w/v) PEG 3350
8	0.2 M Sodium sulphate		20% (w/v) PEG 3350
9	0.2 M Sodium potassium tartrate		20% (w/v) PEG 3350
10	0.2 M Sodium potassium phosphate		20% (w/v) PEG 3350
11	0.2 M Sodium citrate		20% (w/v) PEG 3350
12	0.2 M Sodium malonate		20% (w/v) PEG 3350
13	0.2 M Sodium fluoride	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
14	0.2 M Sodium bromide	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
15	0.2 M Sodium iodide	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
16	0.2 M Potassium thiocyanate	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
17	0.2 M Sodium nitrate	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
18	0.2 M Sodium formate	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
19	0.2 M Sodium acetate	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
20	0.2 M Sodium sulphate	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
21	0.2 M Sodium potassium tartrate	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
22	0.2 M Sodium potassium phosphate	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
23	0.2 M Sodium citrate	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
24	0.2 M Sodium malonate	0.1 M Bis-tris propane pH 6.5	20% (w/v) PEG 3350
25	0.2 M Sodium fluoride	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
26	0.2 M Sodium bromide	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
27	0.2 M Sodium iodide	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350

28	0.2 M Potassium thiocyanate	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
29	0.2 M Sodium nitrate	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
30	0.2 M Sodium formate	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
31	0.2 M Sodium acetate	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
32	0.2 M Sodium sulphate	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
33	0.2 M Sodium potassium tartrate	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
34	0.2 M Sodium potassium phosphate	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
35	0.2 M Sodium citrate	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
36	0.2 M Sodium malonate	0.1 M Bis-tris propane pH 7.5	20% (w/v) PEG 3350
37	0.2 M Sodium fluoride	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
38	0.2 M Sodium bromide	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
39	0.2 M Sodium iodide	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
40	0.2 M Potassium thiocyanate	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
41	0.2 M Sodium nitrate	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
42	0.2 M Sodium formate	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
43	0.2 M Sodium acetate	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
44	0.2 M Sodium sulphate	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
45	0.2 M Sodium potassium tartrate	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
46	0.2 M Sodium potassium phosphate	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
47	0.2 M Sodium citrate	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350
48	0.2 M Sodium malonate	0.1 M Bis-tris propane pH 8.5	20% (w/v) PEG 3350

PEG/Cation Screens

Well number		Buffer	PEG
1	0.2 M Sodium chloride	Acetate pH 5	20% (w/v) PEG 6000
2	0.2 M Ammonium chloride	Acetate pH 5	20% (w/v) PEG 6000
3	0.2 M Lithium chloride	Acetate pH 5	20% (w/v) PEG 6000
4	0.2 M Magnesium chloride	Acetate pH 5	20% (w/v) PEG 6000
5	0.2 M Calcium chloride	Acetate pH 5	20% (w/v) PEG 6000
6	0.01 M Zinc chloride	Acetate pH 5	20% (w/v) PEG 6000
7	0.2 M Sodium chloride	MES pH 6	20% (w/v) PEG 6000
8	0.2 M Ammonium chloride	MES pH 6	20% (w/v) PEG 6000
9	0.2 M Lithium chloride	MES pH 6	20% (w/v) PEG 6000
10	0.2 M Magnesium chloride	MES pH 6	20% (w/v) PEG 6000
11	0.2 M Calcium chloride	MES pH 6	20%(w/v) PEG 6000
12	0.01 M Zinc chloride	MES pH 6	20% (w/v) PEG 6000
13	0.2 M Sodium chloride	HEPES pH 7	20% (w/v) PEG 6000
14	0.2 M Ammonium chloride	HEPES pH 7	20% (w/v) PEG 6000
15	0.2 M Lithium chloride	HEPES pH 7	20% (w/v) PEG 6000
16	0.2 M Magnesium chloride	HEPES pH 7	20% (w/v) PEG 6000
17	0.2 M Calcium chloride	HEPES pH 7	20% (w/v) PEG 6000
18	0.01 M Zinc chloride	HEPES pH 7	20% (w/v) PEG 6000
19	0.2 M Sodium chloride	Tris pH 8	20% (w/v) PEG 6000
20	0.2 M Ammonium chloride	Tris pH 8	20% (w/v) PEG 6000
21	0.2 M Lithium chloride	Tris pH 8	20% (w/v) PEG 6000
22	0.2 M Magnesium chloride	Tris pH 8	20% (w/v) PEG 6000
23	0.2 M Calcium chloride	Tris pH 8	20% (w/v) PEG 6000
24	0.01 M Zinc chloride	Tris pH 8	20% (w/v) PEG 6000

PEG/pH Screen compositions

Well number	Buffer system	pH of the buffer system	PEG
1	1 M Sodium malonate, 1 M imidazole, 1 M Boric acid	4	25% (w/v) PEG 1500
2	1 M Sodium malonate, 1 M imidazole, 1 M Boric acid	5	25% (w/v) PEG 1500
3	1 M Sodium malonate, 1 M imidazole, 1 M Boric acid	6	25% (w/v) PEG 1500
4	1 M Sodium malonate, 1 M imidazole, 1 M Boric acid	7	25% (w/v) PEG 1500
5	1 M Sodium malonate, 1 M imidazole, 1 M Boric acid	8	25% (w/v) PEG 1500
6	1 M Sodium malonate, 1 M imidazole, 1 M Boric acid	9	25% (w/v) PEG 1500
7	1 M Sodium propionate, 1 M Sodium cacodylate trihydrate, 1 M Bis tris propane	4	25% (w/v) PEG 1500
8	1 M Sodium propionate, 1 M Sodium cacodylate trihydrate, 1 M Bis tris propane	5	25% (w/v) PEG 1500
9	1 M Sodium propionate, 1 M Sodium cacodylate trihydrate, 1 M Bis tris propane	6	25% (w/v) PEG 1500
10	1 M Sodium propionate, 1 M Sodium cacodylate trihydrate, 1 M Bis tris propane	7	25% (w/v) PEG 1500
11	1 M Sodium propionate, 1 M Sodium cacodylate trihydrate, 1 M Bis tris propane	8	25% (w/v) PEG 1500
12	1 M Sodium propionate, 1 M Sodium cacodylate trihydrate, 1 M Bis tris propane	9	25% (w/v) PEG 1500
13	1 M DL-Malic acid, 1 M MES, 1 M Tris	4	25% (w/v) PEG 1500
14	1 M DL-Malic acid, 1 M MES, 1 M Tris	5	25% (w/v) PEG 1500
15	1 M DL-Malic acid, 1 M MES, 1 M Tris	6	25% (w/v) PEG 1500

16	1 M DL-Malic acid, 1 M MES, 1 M Tris	7	25% (w/v) PEG 1500
17	1 M DL-Malic acid, 1 M MES, 1 M Tris	8	25% (w/v) PEG 1500
18	1 M DL-Malic acid, 1 M MES, 1 M Tris	9	25% (w/v) PEG 1500

Clear Strategy Screen 1 Conditions (CSS 1)

1 0.3 M Sodium acetate, 25% (w/v) PEG 2000 MME	2 0.2 M Lithium sulphate, 25%(w/v) PEG 2000 MME	3 0.2 M Magnesium chloride, 25% (w/v) PEG 2000 MME	4 0.2 M Potassium bromide, 25% (w/v) PEG 2000 MME	5 0.2 M Potassium thiocyanate, 25% (w/v) PEG 2000 MME	6 0.8 M Sodium formate, 25% (w/v) PEG 2000 MME
7 0.3 M Sodium acetate, 15% (w/v) PEG 4000	8 0.2 M Lithium sulphate, 15% (w/v) PEG 4000	9 0.2 M Magnesium chloride, 15% (w/v) PEG 4000	10 0.2 M Potassium bromide, 15% (w/v) PEG 4000	11 0.2 M Potassium thiocyanate, 15% (w/v) PEG 4000	12 0.8 M Sodium formate, 15% (w/v) PEG 4000
13 0.3 M Sodium acetate, 10% (w/v) PEG 8000, 10% (w/v) PEG 1000	14 0.2 M Lithium sulphate, 10% (w/v) PEG 8000, 10% (w/v) PEG 1000	15 0.2 M Magnesium chloride, 10% (w/v) PEG 8000, 10% (w/v) PEG 1000	16 0.2 M Potassium bromide, 10% (w/v) PEG 8000, 10% (w/v) PEG 1000	17 0.2 M Potassium thiocyanate, 10% (w/v) PEG 8000, 10% (w/v) PEG 1000	18 0.8 M Sodium formate, 10% (w/v) PEG 8000, 10% (w/v) PEG 1000
19 0.3 M Sodium acetate 8% (w/v) PEG 20000, 8% (w/v) PEG550 MME	20 0.2 M Lithium sulphate, 8% (w/v) PEG 20000, 8% (w/v) PEG550 MME	21 0.2 M Magnesium chloride, 8% (w/v) PEG 20000, 8% (w/v) PEG550 MME	22 0.2 M Potassium bromide, 8% (w/v) PEG 20000, 8% (w/v) PEG550 MME	23 0.2 M Potassium thiocyanate, 8% (w/v) PEG 20000, 8% (w/v) PEG550 MME	24 0.8 M Sodium formate, 8% (w/v) PEG 20000, 8% (w/v) PEG550 MME

Clear Strategy Screen 2 conditions (MDCS2)

[A1] Tube : 1 1.5 M Ammonium sulphate	[A2] Tube : 2 0.8 M Lithium sulphate	[A3] Tube : 3 2 M Sodium formate	[A4] Tube : 4 0.5 M Potassium phosphate	[A5] Tube : 5 25% (w/v) PEG 2000 MME, 0.2 M Calcium acetate	[A6] Tube : 6 15% (w/v) PEG 4000, 0.2 M Calcium acetate
[B1] Tube : 7 2.7 M Ammonium sulphate	[B2] Tube : 8 1.8 M Lithium sulphate	[B3] Tube : 9 4 M Sodium formate	[B4] Tube : 10 1.0 M Potassium dihydrogen phosphate	[B5] Tube : 11 10% (w/v) PEG 8000, 10% (w/v) PEG 1000, 0.2 M Calcium acetate	[B6] Tube : 12 8% (w/v) PEG 20000, 8% (w/v) PEG 550 MME, 0.2 M Calcium acetate
[C1] Tube : 13 40%v/v Methanepentanediol	[C2] Tube : 14 40%v/v Butanediol	[C3] Tube : 15 20% (w/v) PEG 4000, 0.005 M Cadmium chloride	[C4] Tube : 16 20% (w/v) PEG 550 MME, 0.15 M Potassium thiocyanate	[C5] Tube : 17 20% (w/v) PEG 600, 0.15 M Potassium thiocyanate	[C6] Tube : 18 20% (w/v) PEG 15000, 0.15 M Potassium thiocyanate
[D1] Tube : 19 35%v/v Isopropanol	[D2] Tube : 20 30% (v/v) Jeffamine 600M	[D3] Tube : 21 20% (w/v) PEG 4000, 0.005 M Nickel chloride	[D4] Tube : 22 18% (w/v) PEG 3350, 0.15 M Potassium thiocyanate	[D5] Tube : 23 18% (w/v) PEG 5000 MME, 0.15 M Potassium thiocyanate	[D6] Tube : 24 15% (w/v) PEG 6000, 0.15 M Potassium thiocyanate

Hampton screen 1 formulations (HS 1)

Tube	SALT	BUFFER	Precipitant
1	0.02 M Calcium Chloride	0.1 M Sodium acetate pH 4.6	30% (w/v) 2-methyl-2,4-pentanediol
2	None	None	0.4 M Potassium sodium tartrate tetrahydrate
3	None	None	0.4 M Ammonium dihydrogen phosphate
4	None	0.1 M Tris-HCl pH 8.5	2.0 M Ammonium Sulphate
5	0.2 M tri-sodium citrate	0.1 M sodium HEPES pH 7.5	30% (w/v) 2-methyl-2,4-pentanediol
6	0.2 M Magnesium chloride	0.1 M Tris-HCl pH 8.5	30% (w/v) PEG 4000
7	None	0.1 M sodium cacodylate pH 6.5	1.4 M Sodium acetate trihydrate
8	0.2 M tri-sodium citrate	0.1 M sodium cacodylate pH 6.5	30% (v/v) 2-propanol
9	0.2 M Ammonium acetate	0.1 M sodium citrate pH 5.6	30% (w/v) PEG 4000
10	0.2 M Ammonium acetate	0.1 M sodium acetate pH 4.6	30% (w/v) PEG 4000
11	None	0.1 M sodium citrate pH 5.6	1.0 M Ammonium dihydrogen phosphate
12	0.2 M Magnesium chloride	0.1 M sodium HEPES pH 7.5	30% (v/v) 2-propanol
13	0.2 M tri-sodium citrate	0.1 M Tris-HCl pH 8.5	30% (v/v) PEG 400
14	0.2 M Calcium Chloride	0.1 M sodium HEPES pH 7.5	28% (v/v) PEG 400
15	0.2 M Ammonium acetate	0.1 M sodium cacodylate pH 6.5	30% (w/v) PEG 8000
16	None	0.1 M sodium HEPES pH 7.5	1.5 M Lithium sulphate monohydrate
17	0.2 M Lithium sulphate	0.1 M Tris-HCl pH 8.5	30% (w/v) PEG 4000
18	0.2 M Magnesium acetate	0.1 M sodium cacodylate pH 6.5	20% (w/v) PEG 8000
19	0.2 M Ammonium acetate	0.1 M Tris-HCl pH 8.5	30% (v/v) 2-propanol
20	0.2 M Ammonium sulphate	0.1 M sodium acetate pH 4.6	25% (w/v) PEG 4000
21	0.2 M Magnesium acetate	0.1 M sodium cacodylate pH 6.5	30% (v/v) 2-methyl-2,4-pentanediol
22	0.2 M Sodium acetate	0.1 M Tris-HCl pH 8.5	30% (w/v) PEG 4000
23	0.2 M Magnesium chloride	0.1 M sodium HEPES pH 7.5	30% (v/v) PEG 400
24	0.2 M Calcium Chloride	0.1 M sodium acetate pH 4.6	20% (v/v) 2-propanol
25	None	0.1 M Imidazole pH 6.5	1.0 M Sodium acetate trihydrate
26	0.2 M Ammonium acetate	0.1 M sodium citrate pH 5.6	30% (v/v) 2-methyl-2,4-pentanediol
27	0.2 M tri-sodium citrate	0.1 M sodium HEPES pH 7.5	20% (v/v) 2-propanol
28	0.2 M Sodium acetate	0.1 M sodium Cacodylate pH 6.5	30% (w/v) PEG 8000
29	None	0.1 M sodium HEPES pH 7.5	0.8 M Potassium sodium tartrate
30	0.2 M Ammonium acetate	None	30% (w/v) PEG 8000
31	0.2 M Ammonium acetate	None	30% (w/v) PEG 4000
32	None	None	2.0 M Ammonium Sulphate
33	None	None	4.0 M Sodium formate
34	None	0.1 M sodium Acetate pH 4.6	2.0 M Sodium formate

35	None	0.1 M sodium HEPES pH 7.5	1.6 M Potassium sodium tartrate
36	None	0.1 M Tris-HCl pH 8.5	8% (w/v) PEG 8000
37	None	0.1 M sodium acetate pH 4.6	8% (w/v) PEG 4000
38	None	0.1 M sodium HEPES pH 7.5	1.4 M Sodium citrate dihydrate
39	None	0.1 M sodium HEPES pH 7.5	2% (v/v) PEG 400 & 2.0 M Ammonium Sulphate
40	None	0.1 M sodium citrate pH 5.6	20% (v/v) 2-propanol & 20% (w/v) PEG 4000
41	None	0.1 M sodium HEPES pH 7.5	10% (v/v) 2-propanol & 20% (w/v) PEG 4000
42	0.05 M potassium phosphate monobasic	None	20% (w/v) PEG 8000
43	None	None	30% (w/v) PEG 1500
44	None	None	0.2 M Magnesium formate
45	0.2 M Zinc acetate	0.1 M sodium cacodylate pH 6.5	18% (w/v) PEG 8000
46	0.2 M Calcium acetate	0.1 M sodium cacodylate pH 6.5	18% (w/v) PEG 8000
47	None	0.1 M sodium acetate pH 4.6	2.0 M Ammonium Sulphate
48	None	0.1 M Tris-HCl pH 8.5	2.0 M Ammonium dihydrogen phosphate

Hampton Crystal Screen II formulations (HCS2)

[A1] Tube : 1 2.0 M sodium chloride, 10% PEG 6000	[A2] Tube : 2 0.5 M sodium chloride, 0.01 M hexadecyltrimethylammonium bromide, 0.01 M magnesium chloride hexahydrate	[A3] Tube : 3 25% (v/v) ethylene glycol	[A4] Tube : 4 35% (v/v) dioxane	[A5] Tube : 5 2.0 M ammonium sulphate, 5% (v/v) iso- propanol
[B1] Tube : 7 10% (w/v) PEG 1000, 10% (w/v) PEG 8000	[B2] Tube : 8 1.5 M sodium chloride, 10% (v/v) ethanol	[B3] Tube : 9 0.1 M sodium acetate trihydrate pH 4.6, 2 M sodium chloride	[B4] Tube : 10 0.2 M sodium chloride, 0.1 M sodium acetate trihydrate pH 4.6, 30% (v/v) MPD	[B5] Tube : 11 0.01 M cobalt chloride hexahydrate, 0.1 M sodium acetate trihydrate pH 4.6, 1 M 1,6 hexanediol
[C1] Tube : 13 0.2 M ammonium sulphate, 0.1 M sodium acetate trihydrate pH 4.6, 30% (w/v) PEG monomethyl ether 2000	[C2] Tube : 14 0.2 M potassium sodium tetrahydrate, 0.1 M tri-sodium citrate dihydrate pH 5.6, 2 M ammonium sulfate	[C3] Tube : 15 0.5 M ammonium sulphate, 0.1 M tri-sodium citrate dihydrate pH 5.6, 1 M lithium sulfate monohydrate	[C4] Tube : 16 0.5 M sodium chloride, 0.1 M tri-sodium citrate dihydrate pH 5.6, 2% (v/v) ethylene imine polymer	[C5] Tube : 17 0.1 M tri-sodium citrate dihydrate pH 5.6, 35% (v/v) tert- butanol
[D1] Tube : 19 0.1 M tri-sodium citrate dehydrate, pH 5.6 2.5 M 1,6 hexanediol	[D2] Tube : 20 0.1 M MES pH 6.5, 1.6 M magnesium sulfate heptahydrate	[D3] Tube : 21 0.1 M sodium dihydrogen phosphate, 0.1 M MES pH 6.5, 2 M sodium chloride, 0.1 M potassium dihydrogen phosphate	[D4] Tube : 22 0.1 M MES pH 6.5, 12% (w/v) PEG 20,000	[D5] Tube : 23 1.6 M ammonium sulphate, 0.1 M MES pH 6.5 10% (v/v) dioxane
[E1] Tube : 25 0.01 M cobaltous chloride hexahydrate, 0.1 M MES pH 6.5, 1.8 M ammonium sulfate	[E2] Tube : 26 0.2 M ammonium sulphate 0.1 M MES pH 6.5, 30% (w/v) PEG monomethyl ether 5000	[E3] Tube : 27 0.01 M zinc sulfate heptahydrate, 0.1 M MES pH 6.5 25% (v/v) PEG monomethyl ether 550	[E4] Tube : 28 pH 6.5 1.6 M tri-sodium citrate dihydrate	[E5] Tube : 29 0.5 M ammonium sulphate, 0.1 M HEPES pH 7.5, 30% (v/v) MPD
[F1] Tube : 31 0.1 M HEPES pH 7.5, 20% (v/v) jeffamine M- 600	[F2] Tube : 32 0.1 M sodium chloride, 0.1 M HEPES pH 7.5, 1.6 M ammonium sulfate	[F3] Tube : 33 0.1 M HEPES pH 7.5, 2 M ammonium formate	[F4] Tube : 34 0.05 M cadmium sulfate hydrate, 0.1 M HEPES	[F5] Tube : 35 0.1 M HEPES pH 7.5, 70% (v/v) MPD

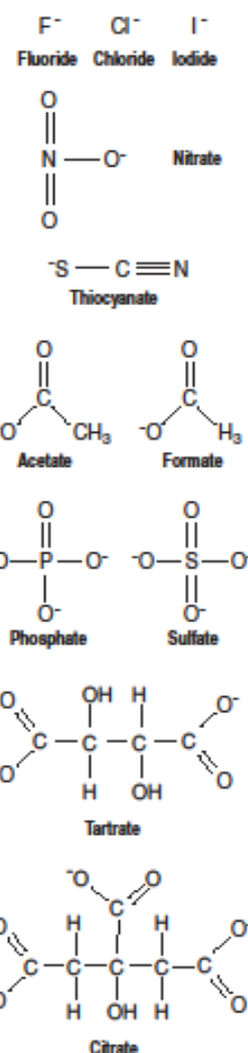
			pH 7.5, 1 M sodium acetate trihydrate	
[G1] Tube : 37 0.1 M HEPES pH 7.5, 10% (w/v) PEG 8000	[G2] Tube : 38 0.1 M HEPES pH 7.5, 20% (w/v) PEG 10000, 8% (v/v) ethylene glycol	[G3] Tube : 39 0.2 M magnesium chloride hexahydrate, 0.1 M tris pH 8.5, 3.4 M 1,6 hexanediol	[G4] Tube : 40 0.1 M tris pH 8.5, 25% (v/v) tert- butanol	[G5] Tube : 41 0.01 M nickel (II) chloride hexahydrate, 0.1 M tris pH 8.5, 1 M lithium sulfate monohydrate
[H1] Tube : 43 0.2 M ammonium dihydrogen phosphate, 0.1 M tris pH 8.5, 50% (v/v) MPD	[H2] Tube : 44 0.1 M tris pH 8.5, 20% (v/v) ethanol	[H3] Tube : 45 0.01 M nickel (II) chloride hexahydrate, 0.1 M tris pH 8.5, 20% (w/v) PEG monomethyl ether 2000	[H4] Tube : 46 0.1 M sodium chloride, 0.1 M bicine pH 9.0, 20% (v/v) PEG monomethyl ether 550	[H5] Tube : 47 0.1 M bicine, pH 9.0, 2 M magnesium chloride hexahydrate

Recipe courtesy of [Hampton Research](#)

Reference website

http://xray.bmc.uu.se/markh/php/xtalscreens.php?func=lookup&screen_name=Expand+List

Tube #	Salt	Tube #	Polymer	Tube #	pH ◊
1.	0.2 M Sodium fluoride	1.	20% w/v Polyethylene glycol 3,350	1.	7.3
2.	0.2 M Potassium fluoride	2.	20% w/v Polyethylene glycol 3,350	2.	7.3
3.	0.2 M Ammonium fluoride	3.	20% w/v Polyethylene glycol 3,350	3.	6.2
4.	0.2 M Lithium chloride	4.	20% w/v Polyethylene glycol 3,350	4.	6.8
5.	0.2 M Magnesium chloride hexahydrate	5.	20% w/v Polyethylene glycol 3,350	5.	5.9
6.	0.2 M Sodium chloride	6.	20% w/v Polyethylene glycol 3,350	6.	6.9
7.	0.2 M Calcium chloride dihydrate	7.	20% w/v Polyethylene glycol 3,350	7.	5.1
8.	0.2 M Potassium chloride	8.	20% w/v Polyethylene glycol 3,350	8.	7.0
9.	0.2 M Ammonium chloride	9.	20% w/v Polyethylene glycol 3,350	9.	6.3
10.	0.2 M Sodium iodide	10.	20% w/v Polyethylene glycol 3,350	10.	7.0
11.	0.2 M Potassium iodide	11.	20% w/v Polyethylene glycol 3,350	11.	7.0
12.	0.2 M Ammonium iodide	12.	20% w/v Polyethylene glycol 3,350	12.	6.2
13.	0.2 M Sodium thiocyanate	13.	20% w/v Polyethylene glycol 3,350	13.	6.9
14.	0.2 M Potassium thiocyanate	14.	20% w/v Polyethylene glycol 3,350	14.	7.0
15.	0.2 M Lithium nitrate	15.	20% w/v Polyethylene glycol 3,350	15.	7.1
16.	0.2 M Magnesium nitrate hexahydrate	16.	20% w/v Polyethylene glycol 3,350	16.	5.9
17.	0.2 M Sodium nitrate	17.	20% w/v Polyethylene glycol 3,350	17.	6.8
18.	0.2 M Potassium nitrate	18.	20% w/v Polyethylene glycol 3,350	18.	6.8
19.	0.2 M Ammonium nitrate	19.	20% w/v Polyethylene glycol 3,350	19.	6.2
20.	0.2 M Magnesium formate dihydrate	20.	20% w/v Polyethylene glycol 3,350	20.	7.0
21.	0.2 M Sodium formate	21.	20% w/v Polyethylene glycol 3,350	21.	7.2
22.	0.2 M Potassium formate	22.	20% w/v Polyethylene glycol 3,350	22.	7.3
23.	0.2 M Ammonium formate	23.	20% w/v Polyethylene glycol 3,350	23.	6.6
24.	0.2 M Lithium acetate dihydrate	24.	20% w/v Polyethylene glycol 3,350	24.	7.9
25.	0.2 M Magnesium acetate tetrahydrate	25.	20% w/v Polyethylene glycol 3,350	25.	7.9
26.	0.2 M Zinc acetate dihydrate	26.	20% w/v Polyethylene glycol 3,350	26.	6.4
27.	0.2 M Sodium acetate trihydrate	27.	20% w/v Polyethylene glycol 3,350	27.	8.0
28.	0.2 M Calcium acetate hydrate	28.	20% w/v Polyethylene glycol 3,350	28.	7.5
29.	0.2 M Potassium acetate	29.	20% w/v Polyethylene glycol 3,350	29.	8.1
30.	0.2 M Ammonium acetate	30.	20% w/v Polyethylene glycol 3,350	30.	7.1
31.	0.2 M Lithium sulfate monohydrate	31.	20% w/v Polyethylene glycol 3,350	31.	6.0
32.	0.2 M Magnesium sulfate heptahydrate	32.	20% w/v Polyethylene glycol 3,350	32.	6.0
33.	0.2 M Sodium sulfate decahydrate	33.	20% w/v Polyethylene glycol 3,350	33.	6.7
34.	0.2 M Potassium sulfate	34.	20% w/v Polyethylene glycol 3,350	34.	6.8
35.	0.2 M Ammonium sulfate	35.	20% w/v Polyethylene glycol 3,350	35.	6.0
36.	0.2 M Sodium tartrate dibasic dihydrate	36.	20% w/v Polyethylene glycol 3,350	36.	7.3
37.	0.2 M Potassium sodium tartrate tetrahydrate	37.	20% w/v Polyethylene glycol 3,350	37.	7.4
38.	0.2 M Ammonium tartrate dibasic	38.	20% w/v Polyethylene glycol 3,350	38.	6.6
39.	0.2 M Sodium phosphate monobasic monohydrate	39.	20% w/v Polyethylene glycol 3,350	39.	4.7
40.	0.2 M Sodium phosphate dibasic dihydrate	40.	20% w/v Polyethylene glycol 3,350	40.	9.1
41.	0.2 M Potassium phosphate monobasic	41.	20% w/v Polyethylene glycol 3,350	41.	4.8
42.	0.2 M Potassium phosphate dibasic	42.	20% w/v Polyethylene glycol 3,350	42.	9.2
43.	0.2 M Ammonium phosphate monobasic	43.	20% w/v Polyethylene glycol 3,350	43.	4.6
44.	0.2 M Ammonium phosphate dibasic	44.	20% w/v Polyethylene glycol 3,350	44.	8.0
45.	0.2 M Lithium citrate tribasic tetrahydrate	45.	20% w/v Polyethylene glycol 3,350	45.	8.4
46.	0.2 M Sodium citrate tribasic dihydrate	46.	20% w/v Polyethylene glycol 3,350	46.	8.3
47.	0.2 M Potassium citrate tribasic monohydrate	47.	20% w/v Polyethylene glycol 3,350	47.	8.3
48.	0.2 M Ammonium citrate dibasic	48.	20% w/v Polyethylene glycol 3,350	48.	5.1



◊ Measured pH at 25 ° C

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34 Journey
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 E-mail: tcob@hampton.com
 Website: www.hamptonresearch.com

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Solutions for Crystal Growth

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Tube Number	Salt	Tube Number	Buffer †
A1.	1.8 M Sodium Acetate pH 7.0	A1.	0.1 M Bis-Tris Propane pH 7.0
A2.	2.8 M Sodium Acetate pH 7.0	A2.	0.1 M Bis-Tris Propane pH 7.0
A3.	1.5 M Ammonium Chloride	A3.	0.1 M Sodium Acetate pH 4.6
A4.	1.5 M Ammonium Chloride	A4.	0.1 M Bis-Tris Propane pH 7.0
A5.	1.5 M Ammonium Chloride	A5.	0.1 M Tris pH 8.5
A6.	3.5 M Ammonium Chloride	A6.	0.1 M Sodium Acetate pH 4.6
A7.	3.5 M Ammonium Chloride	A7.	0.1 M Bis-Tris Propane pH 7.0
A8.	3.5 M Ammonium Chloride	A8.	0.1 M Tris pH 8.5
A9.	2.2 M Sodium Chloride	A9.	0.1 M Sodium Acetate pH 4.6
A10.	2.2 M Sodium Chloride	A10.	0.1 M Bis-Tris Propane pH 7.0
A11.	2.2 M Sodium Chloride	A11.	0.1 M Tris pH 8.5
A12.	3.2 M Sodium Chloride	A12.	0.1 M Sodium Acetate pH 4.6
B1.	3.2 M Sodium Chloride	B1.	0.1 M Bis-Tris Propane pH 7.0
B2.	3.2 M Sodium Chloride	B2.	0.1 M Tris pH 8.5
B3.	1.0 M di-Ammonium hydrogen Citrate	B3.	0.1 M Sodium Acetate pH 4.6
B4.	1.8 M di-Ammonium hydrogen Citrate	B4.	0.1 M Sodium Acetate pH 4.6
B5.	1.0 M tri-Ammonium Citrate pH 7.0	B5.	0.1 M Bis-Tris Propane pH 7.0
B6.	2.0 M tri-Ammonium Citrate pH 7.0	B6.	0.1 M Bis-Tris Propane pH 7.0
B7.	0.7 M tri-Sodium Citrate dihydrate	B7.	0.1 M Bis-Tris Propane pH 7.0
B8.	0.7 M tri-Sodium Citrate dihydrate	B8.	0.1 M Tris pH 8.5
B9.	1.2 M tri-Sodium Citrate dihydrate	B9.	0.1 M Bis-Tris Propane pH 7.0
B10.	1.2 M tri-Sodium Citrate dihydrate	B10.	0.1 M Tris pH 8.5
B11.	0.4 M Magnesium Formate	B11.	0.1 M Sodium Acetate pH 4.6
B12.	0.4 M Magnesium Formate	B12.	0.1 M Bis-Tris Propane pH 7.0
C1.	0.4 M Magnesium Formate	C1.	0.1 M Tris pH 8.5
C2.	0.7 M Magnesium Formate	C2.	0.1 M Bis-Tris Propane pH 7.0
C3.	2.0 M Sodium Formate	C3.	0.1 M Sodium Acetate pH 4.6
C4.	2.0 M Sodium Formate	C4.	0.1 M Bis-Tris Propane pH 7.0
C5.	2.0 M Sodium Formate	C5.	0.1 M Tris pH 8.5
C6.	3.5 M Sodium Formate	C6.	0.1 M Sodium Acetate pH 4.6
C7.	3.5 M Sodium Formate	C7.	0.1 M Bis-Tris Propane pH 7.0
C8.	3.5 M Sodium Formate	C8.	0.1 M Tris pH 8.5
C9.	1.2 M DL-Malic Acid pH 7.0	C9.	0.1 M Bis-Tris Propane pH 7.0
C10.	2.2 M DL-Malic Acid pH 7.0	C10.	0.1 M Bis-Tris Propane pH 7.0
C11.	1.4 M Sodium Malonate pH 7.0	C11.	0.1 M Bis-Tris Propane pH 7.0
C12.	2.4 M Sodium Malonate pH 7.0	C12.	0.1 M Bis-Tris Propane pH 7.0
D1.	2.5 M Ammonium Nitrate	D1.	0.1 M Sodium Acetate pH 4.6
D2.	2.5 M Ammonium Nitrate	D2.	0.1 M Bis-Tris Propane pH 7.0
D3.	2.5 M Ammonium Nitrate	D3.	0.1 M Tris pH 8.5
D4.	6.0 M Ammonium Nitrate	D4.	0.1 M Sodium Acetate pH 4.6
D5.	6.0 M Ammonium Nitrate	D5.	0.1 M Bis-Tris Propane pH 7.0
D6.	6.0 M Ammonium Nitrate	D6.	0.1 M Tris pH 8.5
D7.	1.5 M Sodium Nitrate	D7.	0.1 M Sodium Acetate pH 4.6
D8.	1.5 M Sodium Nitrate	D8.	0.1 M Bis-Tris Propane pH 7.0
D9.	1.5 M Sodium Nitrate	D9.	0.1 M Tris pH 8.5
D10.	4.0 M Sodium Nitrate	D10.	0.1 M Sodium Acetate pH 4.6
D11.	4.0 M Sodium Nitrate	D11.	0.1 M Bis-Tris Propane pH 7.0
D12.	4.0 M Sodium Nitrate	D12.	0.1 M Tris pH 8.5

† Buffer pH is that of a 1.0 M stock prior to dilution with other reagent components, pH with HCl or NaOH.

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I Bioinformatics analysis of GH95 and GH97 proteins, gel filtration graphs for CAH06598 and CAH09443 and PCR amplification conditions used in the cloning of BF3763 and BF0855 genes.

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Search for

ProtParam CAH06598

User-provided sequence:

```

10          20          30          40          50          60
MKIKLLLLLC CGLWSSCNSY DYCPVTPSES DLVFTGLARS WDEAMPLGNA TVGALVWQRD
70          80          90         100         110         120
STLRLSLDRT DLWDLRPVDS LSGDNFRFSW VKEHIRQKNY LPVQKKLDWP YDMNPAPSKI
130         140         150         160         170         180
PGAAIEFPLE QIGTPTQVRL YLNNALCEAD WADGTQMOTF VHATEPIGWF VFRNLKTPIE
190         200         210         220         230         240
PSIITPVYNK TKPDGSLDPV SGQDLHRLGY QQGKVVREGN QITYHQKGYG DFSYDVTVCW
250         260         270         280         290         300
KQEGETLYGT WSVTSSLSGE QASEKAEAAAL QRGLKHDYQA HLEYWDKYWA QSSITLPDSV
310         320         330         340         350         360
LQKQYQNEHY KFGSTTREHS YPISLQAVWT ADNGKLPPWK GDYHHDINTQ LSYWPAYTGN
370         380         390         400         410         420
HLTEGMGYLN TLWNQRDAYK RYTRRYFGTE GMNIPGVCTL TGEPMGGWIQ YSMSQTVAAW
430         440         450         460         470         480
LAQHFYLQWK YSADRTFLKE RAYPFIKDVA IYLEQISEVT PEGVRKLEFS SSPEIFDNSL
490         500         510         520         530         540
QAWFSDMTNY DLAMMHFLFK ATSELAHELN LADEAGHWAS LEAQLPDYDI DEEGCLTFBK
550         560         570         580         590         600
GYPKESHHRH FSHAMAIHPL GLIDWSDGKQ SQHIIRATLK RLDKVGPDYW TGYSYSWLAN
610         620         630         640         650         660
MKARAFDGEA AAQALKTFAE CFCLKNTFHA NGDQTQSGKS RFTYRPFMLE GNFAFAAGIQ
670         680         690         700         710         720
EMLLQSHSTG V IRIFPAIPKE WKDVSFENLR AMGAFLVSAR MEGGEINRVR IYSEKGGMLK
730         740         750
MARPGTLKPN KNYTSLGTDI LNIDTQAGEW IELNP
    
```

References and documentation are available.

new Please note the modified algorithm for extinction coefficient.

Number of amino acids: 755

Molecular weight: 86102.1

Theoretical pI: 5.81

Amino acid composition:

Ala (A)	56	7.4%
Arg (R)	32	4.2%
Asn (N)	30	4.0%
Asp (D)	43	5.7%
Cys (C)	10	1.3%
Gln (Q)	37	4.9%
Glu (E)	45	6.0%
Gly (G)	55	7.3%
His (H)	21	2.8%
Ile (I)	32	4.2%
Leu (L)	72	9.5%
Lys (K)	41	5.4%
Met (M)	19	2.5%
Phe (F)	31	4.1%

Pro (P)	37	4.9%
Ser (S)	52	6.9%
Thr (T)	48	6.4%
Trp (W)	26	3.4%
Tyr (Y)	39	5.2%
Val (V)	29	3.8%
Pyl (O)	0	0.0%
Sec (U)	0	0.0%

(B)	0	0.0%
(Z)	0	0.0%
(X)	0	0.0%

Total number of negatively charged residues (Asp + Glu): 88

Total number of positively charged residues (Arg + Lys): 73

Atomic composition:

Carbon	C	3887
Hydrogen	H	5847
Nitrogen	N	1027
Oxygen	O	1138
Sulfur	S	29

Formula: C₃₈₈₇H₅₈₄₇N₁₀₂₇O₁₁₃₈S₂₉

Total number of atoms: 11928

Extinction coefficients:

Extinction coefficients are in units of M⁻¹ cm⁻¹, at 280 nm measured in water.

Ext. coefficient 201735
 Abs 0.1% (=1 g/l) 2.343, assuming ALL Cys residues appear as half cystines

Ext. coefficient 201110
 Abs 0.1% (=1 g/l) 2.336, assuming NO Cys residues appear as half cystines

Estimated half-life:

The N-terminal of the sequence considered is M (Met).

The estimated half-life is: 30 hours (mammalian reticulocytes, in vitro).
 >20 hours (yeast, in vivo).
 >10 hours (Escherichia coli, in vivo).

Instability index:

The instability index (II) is computed to be 34.84
 This classifies the protein as stable.

Aliphatic index: 72.28

Grand average of hydropathicity (GRAVY): -0.472



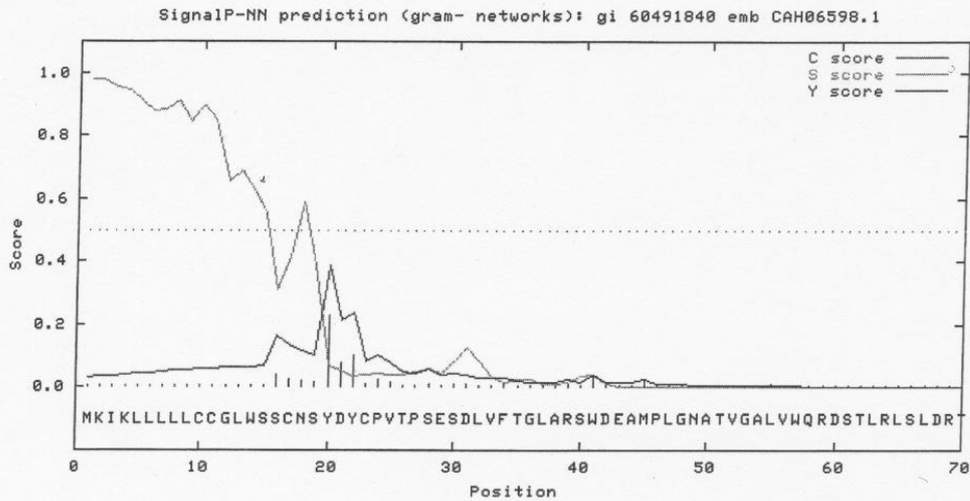
SignalP 3.0 Server - prediction results

Technical University of Denmark

Using neural networks (NN) and hidden Markov models (HMM) trained on Gram-negative bacteria

>gi_60491840_emb_CAH06598.1_conserved hypothetical protein_Bacteroides fragilis NCTC 9343_

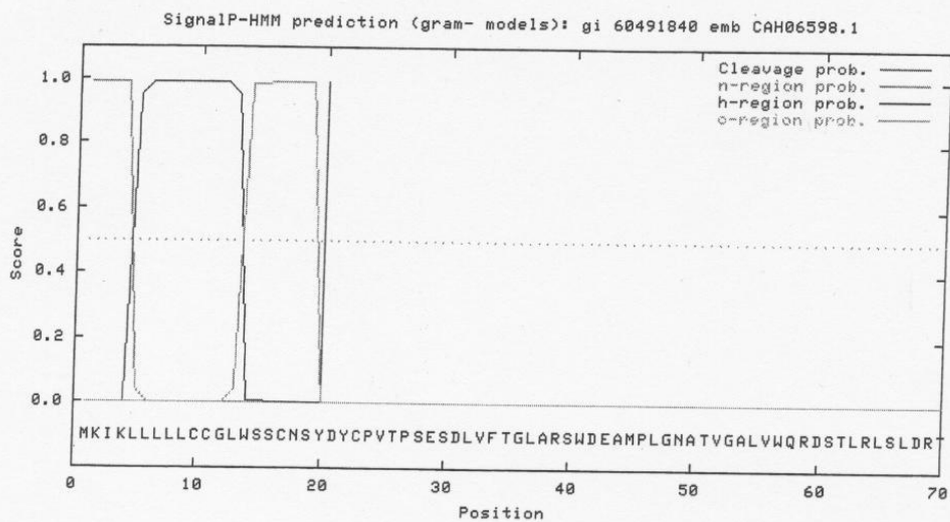
SignalP-NN result:



data

```
>gi_60491840_emb_CAH0 length = 70
# Measure Position Value Cutoff signal peptide?
max. C 20 0.227 0.52 NO
max. Y 20 0.386 0.33 YES
max. S 2 0.982 0.92 YES
mean S 1-19 0.751 0.49 YES
D 1-19 0.568 0.44 YES
# Most likely cleavage site between pos. 19 and 20: CNS-YD
```

SignalP-HMM result:



data

```
>gi_60491840_emb_CAH06598.1_
Prediction: Signal peptide
Signal peptide probability: 0.991
Max cleavage site probability: 0.988 between pos. 19 and 20
```

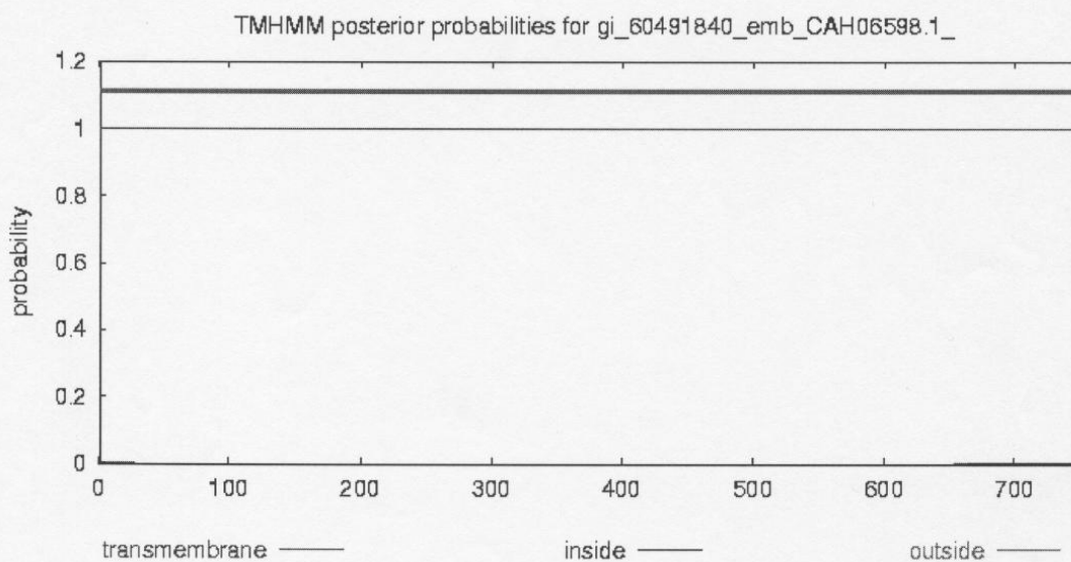
gnuplot script
for making the plot(s)

Explain the output. Go back.

TMHMM result

[HELP](#) with output formats

```
# gi_60491840_emb_CAH06598.1_ Length: 755
# gi_60491840_emb_CAH06598.1_ Number of predicted TMHs: 0
# gi_60491840_emb_CAH06598.1_ Exp number of AAs in TMHs: 0.0179
# gi_60491840_emb_CAH06598.1_ Exp number, first 60 AAs: 0.01192
# gi_60491840_emb_CAH06598.1_ Total prob of N-in: 0.00063
gi_60491840_emb_CAH06598.1_ TMHMM2.0 outside 1 755
```



```
# plot in postscript, script for making the plot in gnuplot, data for plot
```

Search for

ProtParam CAH09443

User-provided sequence:

```

10           20           30           40           50           60
MKRKMMSLLL ALAVISGSSV YAKVIDVMSP NGAIKVSVDI KDRIYYSVSY DNDQLLKDCY
70           80           90          100          110          120
LNLQLQNETL GTNPHLRSTK RGTIDESVKR EIPFKNAIVR NHCNTLRMNF SGNYAVEFRV
130          140          150          160          170          180
FDNGIAYRFV TDKKGDNIVM GEDFAINFPT NYKAHLSQPD GFKTSYECPY THVDTEKYAA
190          200          210          220          230          240
TDRMSYLPVL IETDKAYKIL ISEADLSDYP CMFLKSTGKN GMQSIFPKAP LAFGEDGDRS
250          260          270          280          290          300
LKITEEADYI AKTDGKRSFP WRMMVISKED KELIENEMVY NLSAPCVLED YSWIKPGQVS
310          320          330          340          350          360
WEWWHDARLY GVDFRSGFNM DSYKYYIDFA SKFGIPYIIM DEGWAKNTRD PFTPNPTINL
370          380          390          400          410          420
TELIKYGKDR NVKIVLWLPW LTVENHFDLF KTFADWGIAG VKIDFMDRSD QWMVNYERYV
430          440          450          460          470          480
AKEAAKHKLF VDFHGAFKPA GLERKYPNVL SYEGVLGMEQ GGNCKPENSI YLPFMRNAVQ
490          500          510          520          530          540
PMDFTPGSMI SAQPEDNRST RANAMGSGTR AFQMALFIIF ESGLQMLADN PVYYYRELPC
550          560          570          580          590          600
TEFITSVPVT WDETKVLYAK VGEAVVAKR KGEQWFIGGI TGNQPQNIIE DLGFIPAGQS
610          620          630          640
FTLTSFEDGI NADRQAMDYK KKESTVNNQT RMTLKMVRNG GWAGTIKMK

```

References and documentation are available.

new Please note the modified algorithm for extinction coefficient.

Number of amino acids: 649

Molecular weight: 74054.6

Theoretical pI: 6.33

Amino acid composition:

Ala (A)	42	6.5%
Arg (R)	29	4.5%
Asn (N)	37	5.7%
Asp (D)	44	6.8%
Cys (C)	7	1.1%
Gln (Q)	17	2.6%
Glu (E)	37	5.7%
Gly (G)	44	6.8%
His (H)	8	1.2%
Ile (I)	41	6.3%
Leu (L)	43	6.6%
Lys (K)	49	7.6%
Met (M)	27	4.2%
Phe (F)	34	5.2%
Pro (P)	30	4.6%
Ser (S)	39	6.0%
Thr (T)	35	5.4%
Trp (W)	13	2.0%

Tyr (Y)	33	5.1%
Val (V)	40	6.2%
Pyl (O)	0	0.0%
Sec (U)	0	0.0%

(B)	0	0.0%
(Z)	0	0.0%
(X)	0	0.0%

Total number of negatively charged residues (Asp + Glu): 81
Total number of positively charged residues (Arg + Lys): 78

Atomic composition:

Carbon	C	3337
Hydrogen	H	5118
Nitrogen	N	868
Oxygen	O	973
Sulfur	S	34

Formula: $C_{3337}H_{5118}N_{868}O_{973}S_{34}$
Total number of atoms: 10330

Extinction coefficients:

Extinction coefficients are in units of $M^{-1} cm^{-1}$, at 280 nm measured in water.

Ext. coefficient 121045
Abs 0.1% (=1 g/l) 1.635, assuming ALL Cys residues appear as half cystines

Ext. coefficient 120670
Abs 0.1% (=1 g/l) 1.629, assuming NO Cys residues appear as half cystines

Estimated half-life:

The N-terminal of the sequence considered is M (Met).

The estimated half-life is: 30 hours (mammalian reticulocytes, in vitro).
>20 hours (yeast, in vivo).
>10 hours (Escherichia coli, in vivo).

Instability index:

The instability index (II) is computed to be 32.76
This classifies the protein as stable.

Aliphatic index: 74.82

Grand average of hydropathicity (GRAVY): -0.370

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Search for



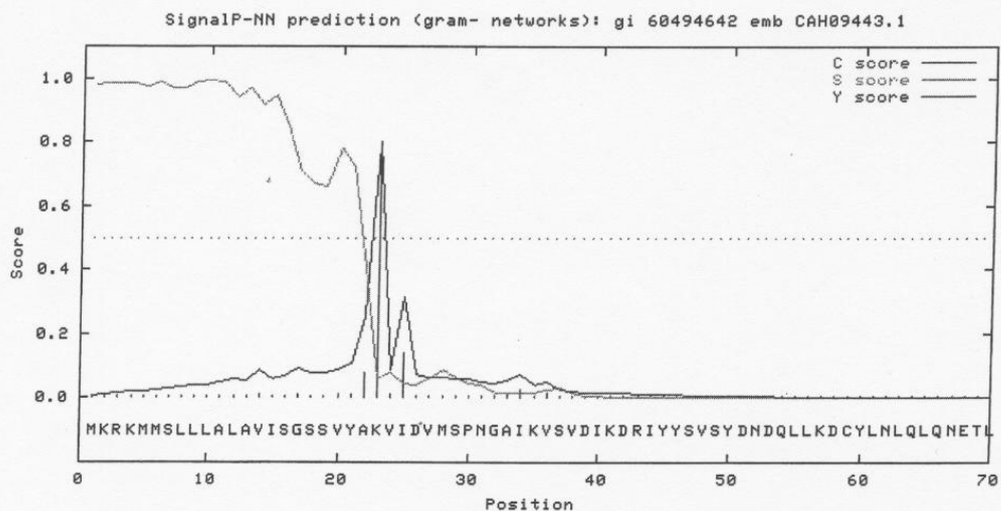
SignalP 3.0 Server - prediction results

Technical University of Denmark

Using neural networks (NN) and hidden Markov models (HMM) trained on Gram-negative bacteria

>gi_60494642_emb_CAH09443.1_putative exported protein_Bacteroides fragilis NCTC 9343_

SignalP-NN result:



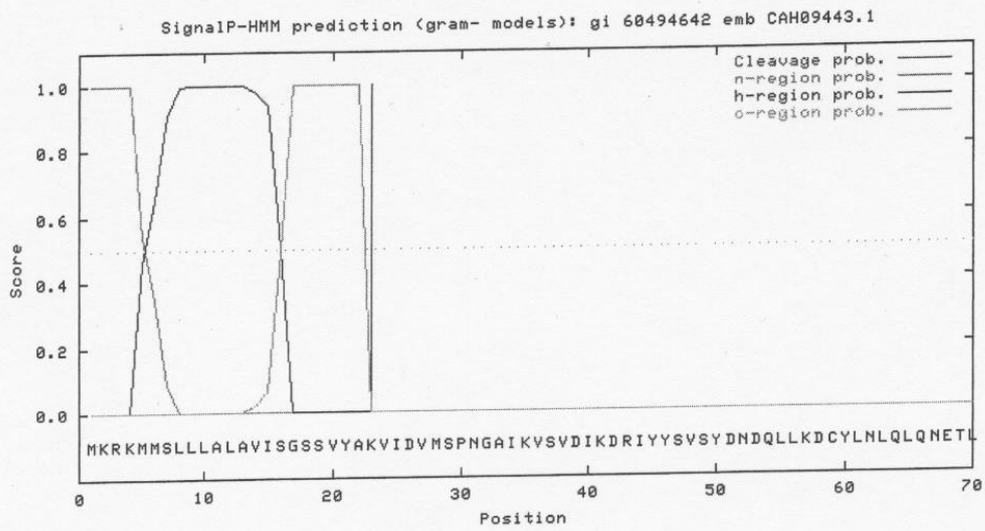
data

>gi_60494642_emb_CAH0 length = 70

# Measure	Position	Value	Cutoff	signal peptide?
max. C	23	0.797	0.52	YES
max. Y	23	0.800	0.33	YES
max. S	10	0.993	0.92	YES
mean S	1-22	0.882	0.49	YES
D	1-22	0.841	0.44	YES

Most likely cleavage site between pos. 22 and 23: VYA-KV

SignalP-HMM result:



data

```
>gi_60494642_emb_CAH09443.1_
Prediction: Signal peptide
Signal peptide probability: 1.000
Max cleavage site probability: 0.999 between pos. 22 and 23
```

gnuplot script
for making the plot(s)

Explain the output. Go back.

gel11148_MI_OV_20VMM_01

Fractions_1

Injection

gel11148_Flow_01

CAH06598 gel filtration graph

40

30

20

10

10

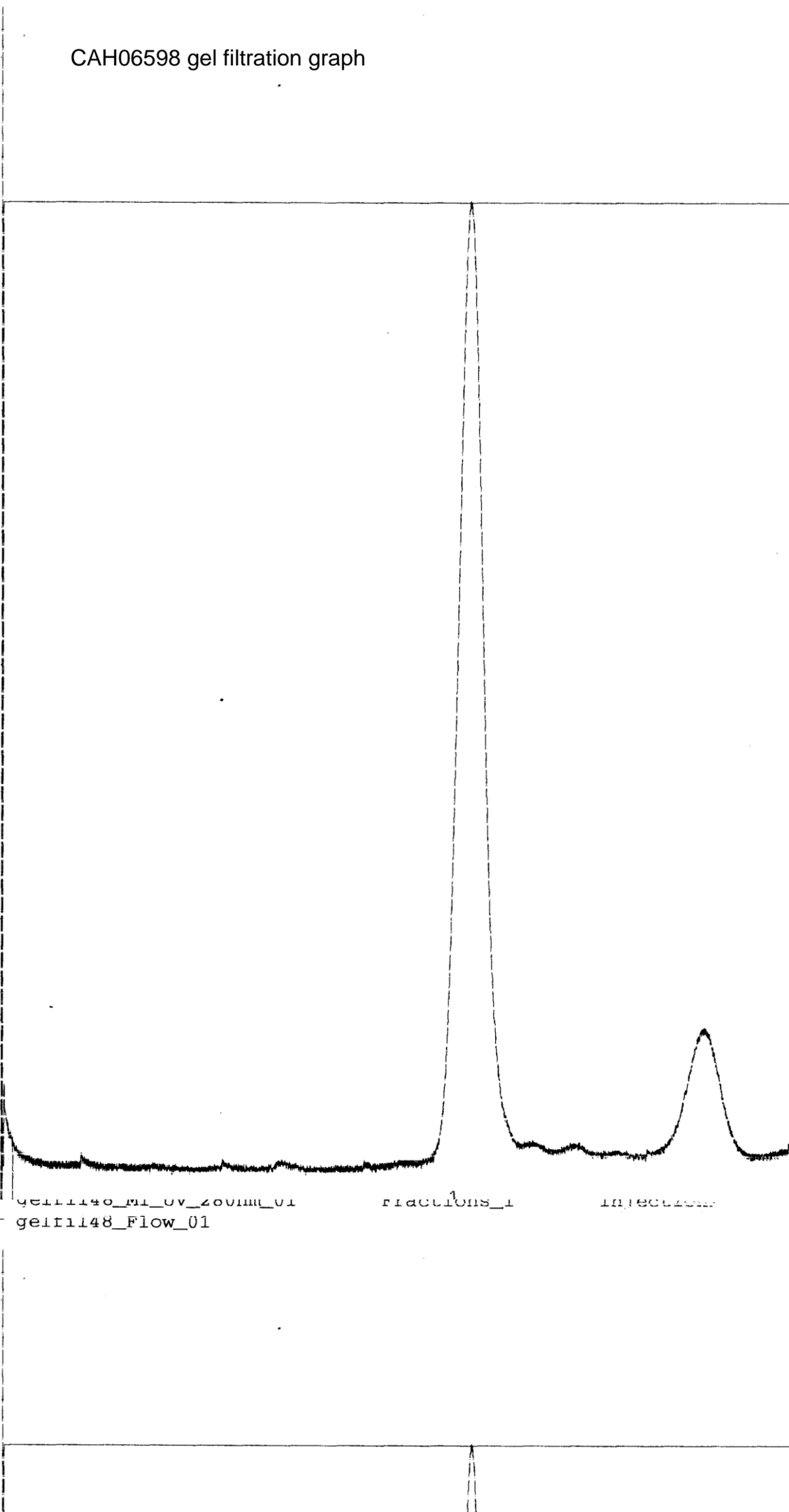
gel11148_MI_OV_20VMM_01

Fractions_1

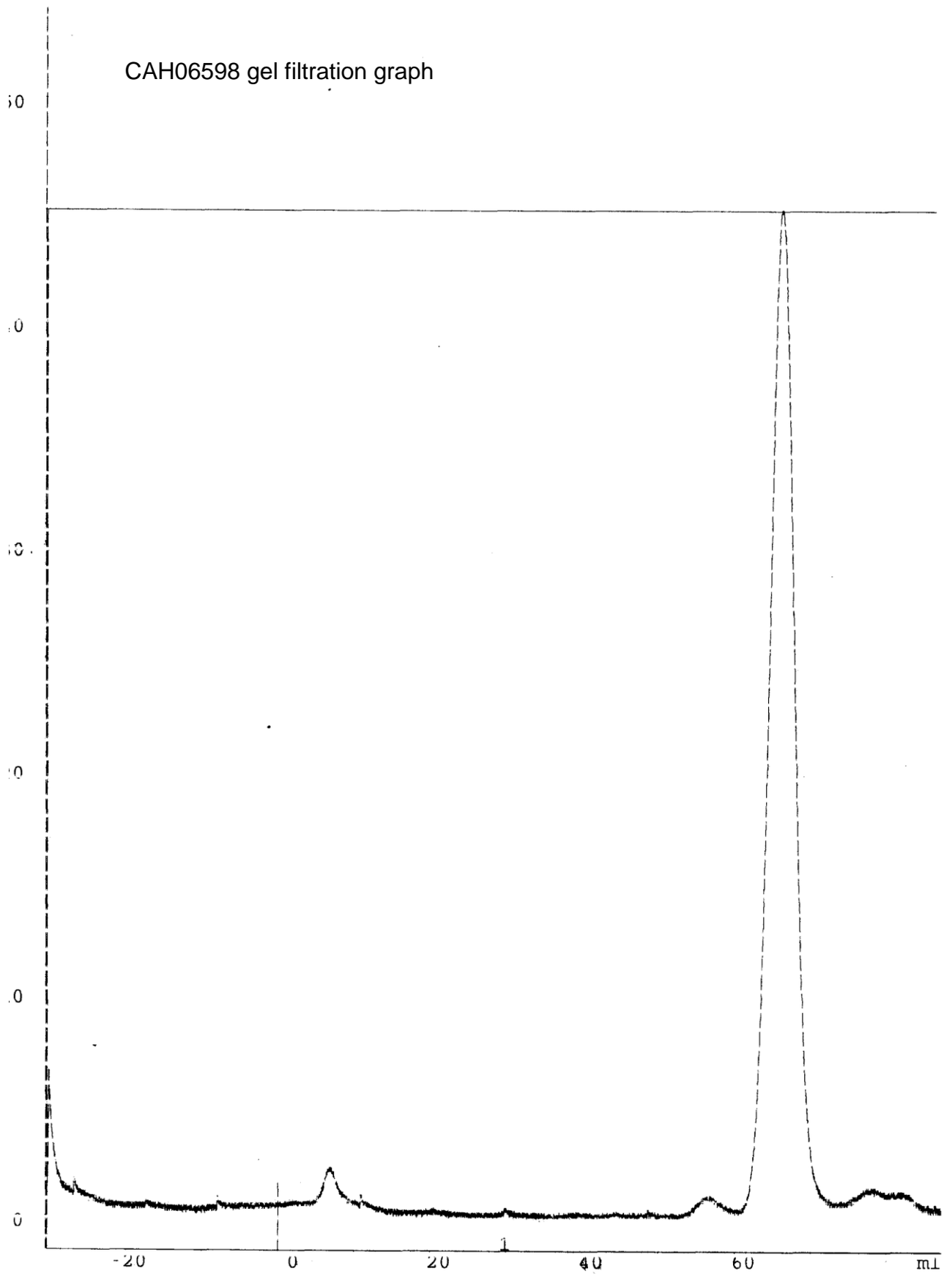
Injection

gel11148_Flow_01

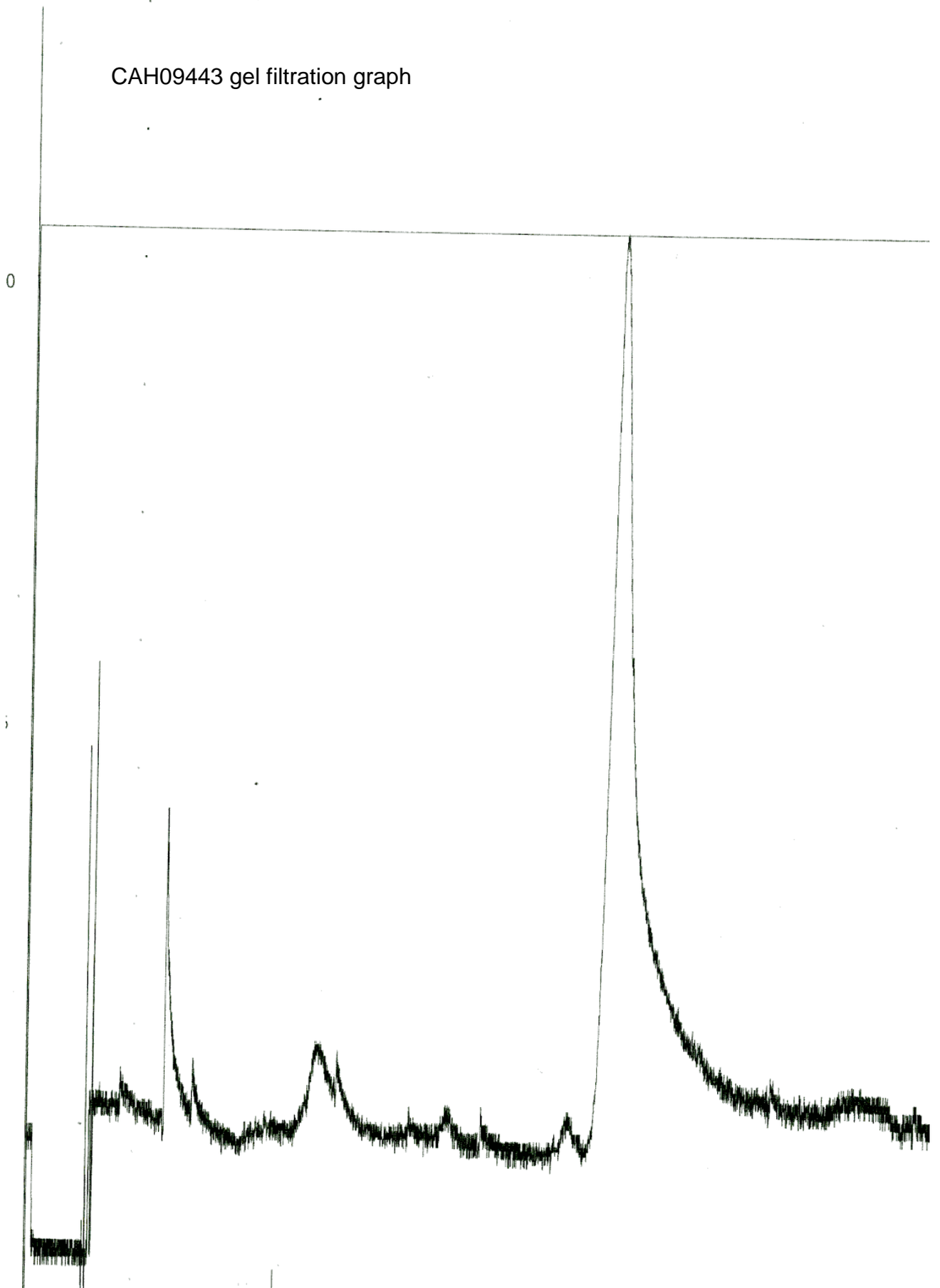
40



gelfil49_MI_OV_280nm_01 Fractions 1 Injections
gelfil49_Flow_01



CAH09443 gel filtration graph



AU

CAH09443 gel filtration graph- selenomethionine derivative

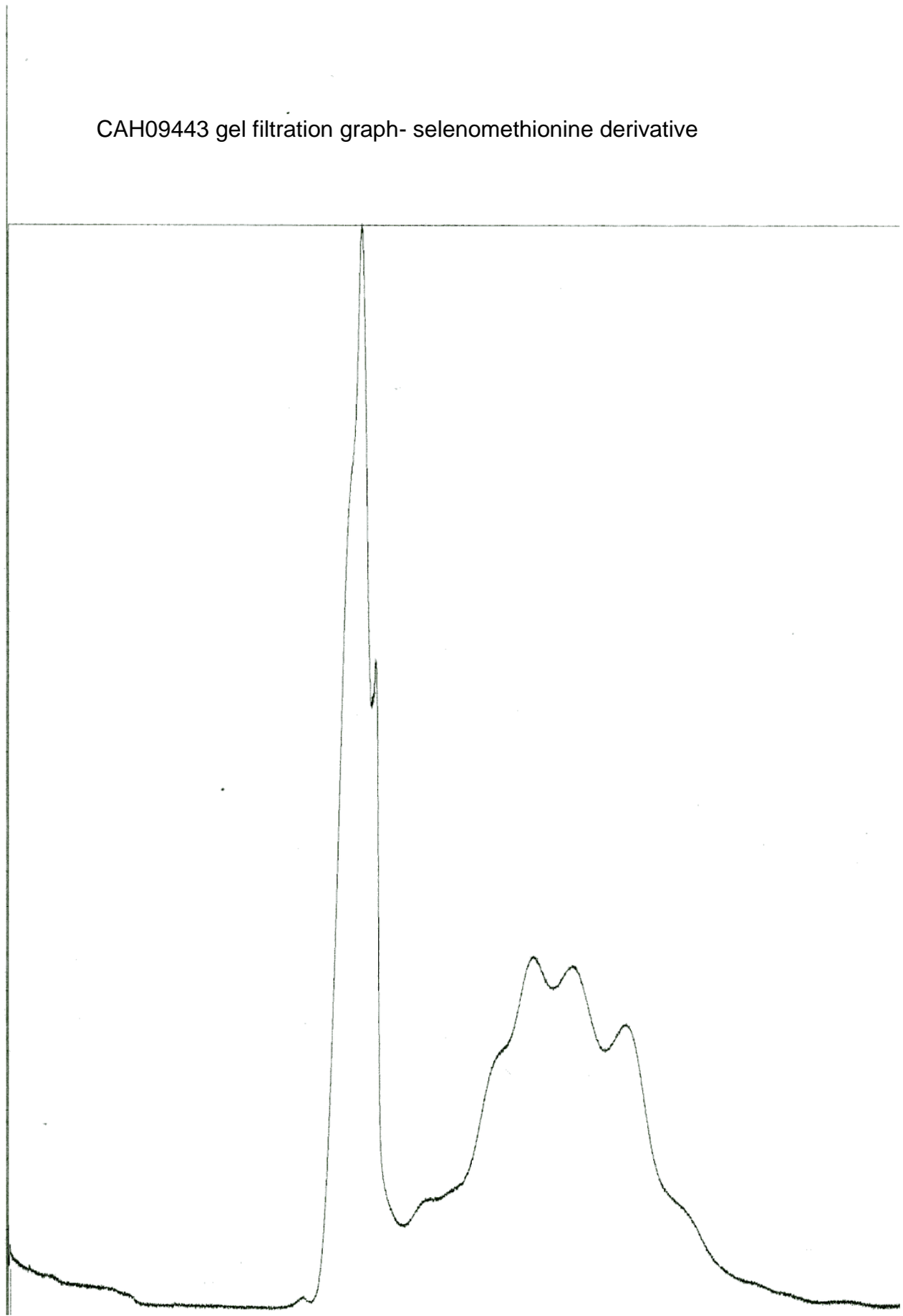
0.8

0.6

0.4

0.2

0.0



676

Step	Temperature	Time
Denature	94°C	2 min
Denature	94°C	15 s
Annealing	60.27°C	30 s
Extension	68°C	2 min 39 s
Repeated 4 more times		
Denature	94°C	15 s
Annealing	70.03°C	30 s
Extension	68°C	2 min 39 s
Repeated 24 more times		
Extension	68°C	10 min
Hold at 10°C		

PCR conditions for the amplification of the BF3763 gene from *B. fragilis*

Step	Temperature	Time
Denature	94°C	2 min
Denature	94°C	15 s
Annealing	60.27°C	30 s
Extension	68°C	2 min 39 s
Denature	94°C	15 s
Annealing	70.03°C	30 s
Extension	68°C	2 min 39 s
Extension	68°C	10 min
Hold at 10°C		

KOD polymerase based PCR amplification of BF3763 gene in *B. fragilis*

Step	Temperature	Time
Denature	94°C	2 min
Denature	94°C	15 s
	61 °C	30 s
Extension	72°C	1 min 53 s
	94°C	15 s
	80°C	30 s
Annealing	72°C	1 min 53 s
Denature	94°C	15 s
Annealing	73°C	30 s
Extension	72°C	1 min 53 s
Extension	72°C	10 min
Hold at 10°C		

PCR conditions for the amplification of the BF0855 gene from *B. fragilis*

Step	Temperature	Time
Denature	94°C	2 min
Denature	94°C	15 s
Annealing	61 °C	30 s
Extension	72°C	1 min 53 s
Denature	94°C	15 s
	80°C	30 s
Annealing	72°C	1 min 53 s
Denature	94°C	15 s
Annealing	73°C	30 s
Extension	72°C	1 min 53 s
Extension	72°C	10 min
Hold at 10°C		

Amplification of the BF3763 gene from transformed *E. coli* to screen for colonies containing successful inserts

Appendix J

Figure represents the standard curve for the Bradford assay

