Northumbria Research Link

Citation: Carro, Lorena, Nouioui, Imen, Sangal, Vartul, Meier-Kolthoff, Jan P., Trujillo, Martha E., Montero-Calasanz, Maria del Carmen, Sahin, Nevzat, Smith, Darren, Kim, Kristi E., Peluso, Paul, Deshpande, Shweta, Woyke, Tanja, Shapiro, Nicole, Kyrpides, Nikos, Klenk, Hans-Peter, Göker, Markus and Goodfellow, Michael (2018) Genome-based classification of micromonosporae with a focus on their biotechnological and ecological potential. Scientific Reports, 8 (1). ISSN 2045-2322

Published by: Nature Publishing

URL: https://doi.org/10.1038/s41598-017-17392-0 https://doi.org/10.1038/s41598-017-17392-0 <a href="https://doi.org/10.1038/s41598-0-1739

This version was downloaded from Northumbria Research Link: http://nrl.northumbria.ac.uk/id/eprint/33321/

Northumbria University has developed Northumbria Research Link (NRL) to enable users to access the University's research output. Copyright © and moral rights for items on NRL are retained by the individual author(s) and/or other copyright owners. Single copies of full items can be reproduced, displayed or performed, and given to third parties in any format or medium for personal research or study, educational, or not-for-profit purposes without prior permission or charge, provided the authors, title and full bibliographic details are given, as well as a hyperlink and/or URL to the original metadata page. The content must not be changed in any way. Full items must not be sold commercially in any format or medium without formal permission of the copyright holder. The full policy is available online: http://nrl.northumbria.ac.uk/policies.html

This document may differ from the final, published version of the research and has been made available online in accordance with publisher policies. To read and/or cite from the published version of the research, please visit the publisher's website (a subscription may be required.)







Received: 14 August 2017 Accepted: 8 November 2017 Published online: 11 January 2018

OPEN Genome-based classification of micromonosporae with a focus on their biotechnological and ecological potential

Lorena Carro 101, Imen Nouioui1, Vartul Sangal2, Jan P. Meier-Kolthoff 103, Martha E. Trujillo4, Maria del Carmen Montero-Calasanz 1, Nevzat Sahin 5, Darren Lee Smith2, Kristi E. Kim6, Paul Peluso⁶, Shweta Deshpande⁷, Tanja Woyke¹, Nicole Shapiro⁷, Nikos C. Kyrpides⁷, Hans-Peter Klenk¹, Markus Göker 63 & Michael Goodfellow¹

There is a need to clarify relationships within the actinobacterial genus Micromonospora, the type genus of the family Micromonosporaceae, given its biotechnological and ecological importance. Here, draft genomes of 40 Micromonosporα type strains and two non-type strains are made available through the Genomic Encyclopedia of Bacteria and Archaea project and used to generate a phylogenomic tree which showed they could be assigned to well supported phyletic lines that were not evident in corresponding trees based on single and concatenated sequences of conserved genes. DNA G+C ratios derived from genome sequences showed that corresponding data from species descriptions were imprecise. Emended descriptions include precise base composition data and approximate genome sizes of the type strains. antiSMASH analyses of the draft genomes show that micromonosporae have a previously unrealised potential to synthesize novel specialized metabolites. Close to one thousand biosynthetic gene clusters were detected, including NRPS, PKS, terpenes and siderophores clusters that were discontinuously distributed thereby opening up the prospect of prioritising gifted strains for natural product discovery. The distribution of key stress related genes provide an insight into how micromonosporae adapt to key environmental variables. Genes associated with plant interactions highlight the potential use of micromonosporae in agriculture and biotechnology.

Prokaryotic systematics is a core scientific discipline that encompasses classification, nomenclature, identification, and evolutionary processes¹. The subject is practiced by few but its applications are relevant to most, if not all, microbiologists². The discipline began as a largely empirical science but became increasingly objective due to the introduction of new concepts and practices, especially the development of chemotaxonomic, numerical phenetic and molecular systematic methods^{3,4}. These developments led to the concept of polyphasic taxonomy, that is, the integrated use of genotypic and phenotypic data to generate classifications of prokaryotes⁵, an approach that was dependent on rapid data acquisition and improved data handling techniques^{6,7}. Genotypic data tend to be derived from analyses of nucleic acids and phenotypic characteristics from chemotaxonomic, cultural, morphological and other expressed features8. The selection of methods for polyphasic studies, while critical, is somewhat subjective though 16S rRNA gene sequencing has proved to be a powerful tool for establishing relationships between prokaryotes at generic and suprageneric ranks⁹⁻¹¹, but tends to be of limited use in distinguishing between closely related species 12-14. In contrast, DNA-DNA pairing, molecular fingerprinting, multilocus sequence typing and phenotypic studies provide valuable data for circumscribing such species^{14–18}.

¹School of Biology, Newcastle University, Newcastle upon Tyne, UK. ²Department of Biomedical Sciences, Northumbria University, Newcastle upon Tyne, UK. ³Leibniz Institute DSMZ-German Collection of Microorganisms and Cell Cultures, Inhoffenstraße 7B, Braunschweig, Germany. Departamento de Microbiologia y Genetica, Lab 214, Universidad de Salamanca, Salamanca, Spain. 5 Department of Biology, Faculty of Art and Science, Ondokuz Mayis University, Kurupelit-Samsun, Turkey. ⁶Pacific Biosciences, 1380 Willow Rd, Menlo Park, California, USA. ⁷DOE Joint Genome Institute, Walnut Creek, California, USA. Correspondence and requests for materials should be addressed to L.C. (email: lcq@usal.es) or H.-P.K. (email: hans-peter.klenk@newcastle.ac.uk)

The widespread application of polyphasic taxonomy led to marked improvements in the classification of archaea and bacteria which, in turn, provided a sound basis for a stable nomenclature and improved identification, as exemplified by the current state of actinobacterial systematics¹⁹. The need to build upon these developments has been raised by those pressing for step-changes in prokaryotic systematics through "embracing the genome"^{20–23}.

The application of low cost whole genome sequencing (WGS) technologies and associated bioinformatic tools is not only providing grist to the taxonomic mill²⁴⁻²⁶, but is furthering our knowledge of developmental and evolutionary processes²⁷⁻²⁹, as well as underpinning the ecological, physiological and biotechnological potential of prokaryotes^{25,26,30-32} thereby repositioning prokaryotic systematics as a fundamental scientific discipline. However, it is essential that taxonomies based on whole genome sequence data follow sound taxonomic practice, notably by following the nomenclatural type concept and the requirement to deposit type strains in two public culture collections in different countries^{33,34}. In this context, the analysis of whole genome sequences of type strains under the auspices of the *Genetic Encyclopaedia of Bacteria and Archaea* (GEBA) project is greatly improving our understanding of phylogenetic relationships within and between these taxa, as well as generating an invaluable framework, technology and organisation for large scale genome sequencing of prokaryotes that will lead to an unprecedented coverage of prokaryotic diversity on the planet³⁵⁻³⁹. The application of innovative phylogenetic and taxonomic methods is also providing new metrics for the recognition of generic and species boundaries^{20,23,40,41}, as well as resolving the structure of complex prokaryotic taxa, such as the actinobacterial genera *Amycolatopsis, Rhodococcus* and *Streptomyces*^{18,25,26}. Members of all of these taxa are a rich source of novel specialized metabolites, notably antibiotics^{42,43}.

This study was designed to explore the extent to which whole genome sequence data derived from type strains of the genus Micromonospora can be used to clarify relationships within this taxon and provide insights into the biological properties and biotechnological potential of micromonosporae. The genus Micromonospora⁴⁴ is the type genus of the family Micromonosporaceae⁴⁵ of the order Micromonosporales⁴⁶; the family encompasses 31 validly named genera which can be distinguished using a combination of chemotaxonomic, morphological and phylogenetic criteria^{45,47}. The genus was proposed by Ørskov in 1923⁴⁸ for strains isolated from air that had been designated as "Streptothrix chalcea" by Foulerton 49 and then reclassified as Micromonospora chalcea, the type species of the genus. At the time of writing the genus encompasses 79 species with validly published names (http://www.bacterio.net/micromonospora.html)50, the majority of which have been described using polyphasic methods^{44,51} though there is evidence that the taxon remains underspeciated^{52,53}. Initially, micromonosporae were associated with soil, freshwater and marine habitats⁴⁴ but novel strains have been isolated from animal^{54–56} and plant tissues^{57–64}, as well as from limestone⁶⁵, Antarctic sandstone⁶⁶ and from a nickel mining site⁶⁷. Micromonosporae form a tight cluster within the Micromonosporaceae 16S rRNA gene tree^{44,51} though 16S rRNA gene sequences are not sufficiently divergent to distinguish between closely related strains thereby drawing upon the need for associated DNA-DNA relatedness studies^{68,69}. It is now apparent that phylogenies showing greater resolution between Micromonospora species can be generated using gyrB sequences⁷⁰ and multilocus sequence analysis (MLSA) of housekeeping genes⁵³.

Despite the advances outlined above there is a clear need to devise an improved framework for the classification and identification of *Micromonospora* strains, partly because of their importance in biotechnology, bioprospecting and ecology^{42,44}. Amongst actinobacteria, micromonosporae are second only to streptomycetes in their ability to synthesize specialized metabolites; they are a particularly rich source of antibiotics, as exemplified by the production of the aminoglycosides: gentamicin, sagamicin, sisomicin and verdamicin from *Micromonospora purpurea*⁷¹ (reclassified as *Micromonospora echinospora*⁷⁰), *Micromonospora sagamiensis*⁷², *Micromonospora inyonensis*⁷³ and "Micromonospora grisea"⁷⁴, respectively; everninomicin, an oligosaccharide antibiotic from *Micromonospora carbonacea*⁷⁵; the ansamycin antibiotic halomicin from *Micromonospora halophytica*⁷⁶; and the new macrolide antibiotics megalomicin⁷⁷ and mycinamicin⁷⁸ from *Micromonospora nigra* and "Micromonospora griseorubida", respectively. Other specialized metabolites synthesized by micromonosporae include the antitumour compounds calicheamicin and lupinacidin C, these enediyne and anthraquinone antibiotics are produced by a *M. echinospora* NRRL 15839⁷⁹ and *Micromonospora lupini*⁸⁰, respectively; and retymicin, galtamycin B, saquayamycin Z and ribofuranosyllumichrome from *Micromonospora* strain Tü 6368⁸¹. On a broader front *Micromonospora* strains have been considered to be a potential source of biocontrol agents, biofuels, plant growth products and plant probiotics^{82–85}.

The metabolic potential of micromonosporae has been underlined in a few whole genome studies which show that a large proportion of the genetic potential of the tested strains code for the biosynthesis of natural products^{85–87}. One of the drivers of the present study was to build upon these pioneering investigations to provide an insight into the potential of micromonosporae to produce new natural products thereby paving the way for developments in applied genomics with particular reference to genome mining^{88–90} and methods for activating silent biosynthetic gene clusters^{91–93}. Investigations like these also highlight genomic features of potential ecological significance, as exemplified by the work on *M. lupini* strain Lupac 08, an endophyte able to colonise internal plant tissues⁹⁴.

Here, whole genome sequences generated from 40 *Micromonospora* type strains and two strains related to *Micromonospora aurantiaca* and *M. echinospora* were generated and used to construct a phylogenomic tree together with the available genomes of *M. aurantiaca* ATCC 27029^T and L5⁸⁷, and *M. lupini* Lupac 08⁸⁶. The resultant data were used to determine the distribution of genes considered to code for natural products and for environmental adaptation, including stress responses. Little congruence was found between the structure of the phylogenomic tree and corresponding single gene trees based on 16S rRNA and conserved housekeeping gene sequences but congruence considerably increased when the single genes were combined in an MLSA of the conserved genes. The genomes of the strains were found to be rich in biosynthetic gene clusters many of which were discontinuously distributed. This study provides further evidence that the taxogenomic-approach to prokaryotic

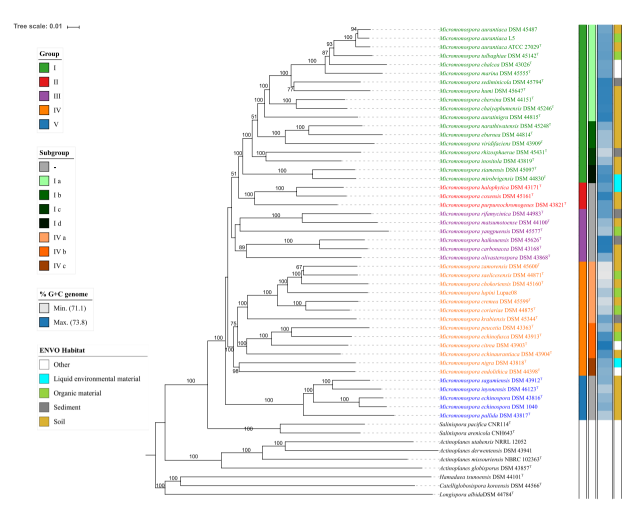


Figure 1. *Micromonospora* phylogeny inferred using the Genome BLAST Distance Phylogeny (GBDP) approach. The tree was inferred using the FastME from the GBDP intergenomic distances calculated from whole proteomes. The numbers above branches are GBDP pseudo-bootstrap support values from a 100 replicates, only values above 50% are shown. Tip colours on the right indicate the habitats from which the strains were isolated, those in the middle-right indicate genomic DNA G+C content, as embedded in the legends. Tip colours on the left indicate selected clades within the genus and those on the middle-left indicate well-supported subgroups within this clades.

systematics can clarify relationships with complex actinobacteria taxa and provide invaluable insights into the biotechnological and ecological potential of the defined groups.

Results

General genome properties. High quality draft genomes were obtained for 40 Micromonospora type strains, 17 of which were completely closed. Approximate genome sizes of the investigated strains varied from 6.1 Mbp for *Micromonospora marina* DSM $4555\overline{5^{T}}$, a strain isolated from sea sand in Thailand⁹⁵ to 7.9 Mbp for *M*. carbonacea DSM 43168^T, a strain recovered from a soil sample in the United States⁹⁶ (Fig. 1); the average genome size for all of the *Micromonospora* strains was 7 ± 0.4 Mbp (Supplementary Table 1). In the following sections, we report on gene numbers indicated by IMG annotation, but these numbers need to be interpreted cautiously as not all of the genome sequences were complete. The number of genes ranged from 5,550 in the genome of the type strain of M. marina to 7,388 in that of Micromonospora cremea DSM 45599^T, a strain isolated from the rhizosphere of *Pisum sativum*⁹⁷. RNA genes represented 1–2% of the whole genome sequences ranging from 63 genes in the type strain of M. aurantiaca to 133 genes in Micromonospora humi DSM 45647^T, a strain isolated from peat swamp forest soil⁹⁸. An average of ten genes were identified as encoding rRNA's (from 6 to 15 genes) and an average of 58 for tRNA's (from 48 to 87 genes). The number of pseudogenes varied from 0 in 14 out of the 45 genomes up to 445 in the genome of the M. cremea type strain. The number of genes with a predicted function averaged 4,600, these ranged from 3,934 in the genome of M. nigra DSM 43818^T to 5,266 in that of M. cremea DSM 45599^T. Between 4 and 10% of the genes were associated with the expression of signal peptides while the percentage of transmembrane proteins varied from 21 to 27% (Supplementary Table 1). The number of Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR) rose from nought in the genome of Micromonospora inositola DSM 43819^T to over ten, as exemplified by M. sagamiensis DSM 43912^T (11), Micromonospora yangpuensis DSM 45577^T (12), Micromonospora olivasterospora DSM 43868^T (16), Micromonospora peucetia DSM 43363^T (17) and

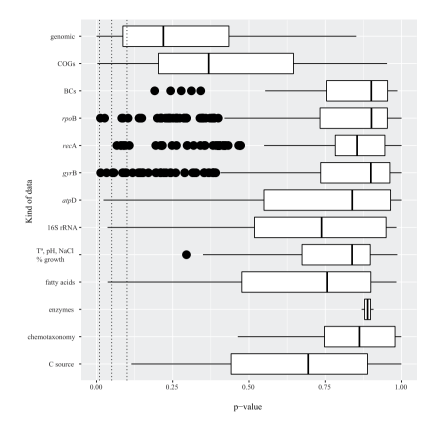


Figure 2. Tip permutation test analysis of *Micromonospora* features respect to phylogenomics. Shown are the p-values from the tests of individual characters arranged by kind of character. The dotted lines represent alpha levels (0.01, 0.05, 0.10). With exceptions for some fatty acids, no correlation was observed between phenotypic data (C source: carbon sources; chemotaxonomy: other than fatty acid data; enzyme production; fatty acids; and growth at several temperatures, pH and percentage of salinity) and genome scale phylogeny. The single genes (16S rRNA, *atp*D, *gyr*B, *rec*A, *rpo*B) presented few significant phylogenetically conserved characters. The number of BCs (biosynthetic gene clusters) classified by type of organic compounds presented no detectable phylogeny conservation, whereas some COGs categories and the GC content (as part of the group of other genomic characters) were significantly conserved.

Micromonospora viridifaciens DSM 43909^T (18) with an average of five per genome (Supplementary Table 1). None of these genomic characteristics were found to be phylogenetically conserved (α = 0.01) in the tip permutation test (Fig. 2, Supplementary Table 2).

More than 40% of the average number of 6,361 genes detected in the *Micromonospora* genomes were conserved as the core genome. The size of the core genome corresponded to around 50% of the smallest genome, as exemplified by *M. marina* DSM 45555^T . Large differences were found in the Clusters of Orthologous Groups (COGs), notably in categories G (carbohydrate transport and metabolism), K (transcription), N (cell motility), S (function unknown), W (extracellular structures), and X (mobilome: prophages, transposons) (Supplementary Figure 1; Supplementary Table 3). Particularly large variations were seen in category X, ranging from 123 genes in the *M. inyonensis* DSM 46123^T genome, which mainly coded for transposases, to four phage related proteins in the genome of *Micromonospora chersina* DSM 44151^T , none of which coded for transposases (Supplementary Figure 1; Supplementary Table 3); however, it is known that the number of transposases can increase quickly through autoreplication⁹⁹. For instance, one of the largest bacterial genomes encountered up to date, *Ktedonobacter racemifer* SOSP1–21^T, contains a huge number of transposases¹⁰⁰. Among the COG counts, categories I (lipid transport and metabolism) and M (cell wall/membrane biogenesis) were seen to be phylogenetically conserved ($\alpha = 0.01$) in the tip-permutation test (Fig. 2, Supplementary Table 2).

Genome based classification. It can be seen from the phylogenomic tree (Fig. 1) that the *Micromonospora* strains form a monophyletic group supported by a 100% bootstrap value. This taxon is clearly separated from an adjacent lineage that encompasses the type strains of *Salinispora arenicola* and *Salinispora tropica*. It is also evident from Fig. 1 that the *Micromonospora* strains fall into four well supported sublineages, groups I, II, IV, and V, and the less well supported strains that for the sake of clarity, have been classified into group III, a taxon that may prove to be heterogeneous. The largest taxon, group I, encompasses 18 strains, including *M. chalcea* DSM 43026^T, the type strain of the type species of the genus *Micromonospora*. These strains were assigned to four subgroups that were supported by 100% bootstrap values, group Ia encompasses the three *M. aurantiaca* strains and the type strains of *Micromonospora auratinigra*, *M. chalcea*, *Micromonospora chaiyaphumensis*, *M. chersina*, *M. humi*,

M. marina, Micromonospora sediminicola and Micromonospora tulbaghiae, closely related organisms isolated from sea sand⁹⁵, marine sediment¹⁰¹, plants^{59,87}, peat swamp forest^{98,102}, air⁴⁹ and soil^{44,103,104}; group Ib is composed of the type strains of Micromonospora eburnea, Micromonospora narathiwatensis and M. viridifaciens, also from soil^{68,105,106}; group Ic includes the type strains of M. inositola and Micromonospora rhizosphaerae from soil¹⁰⁷ and rhizosphere¹⁰⁸, respectively, and group Id the type strains of Micromonospora mirobrigensis and Micromonospora siamensis, two highly related organisms isolated, in turn, from a pond¹⁰⁹ and peat swamp forest soil¹¹⁰.

Group II encompasses the type strains of *Micromonospora coxensis*, *M. halophytica* and *Micromonospora purpureochromogenes*, isolates from saline habitats^{76,111} and adobe soil⁴⁴, respectively, and group III *M. carbonacea* DSM 43168^T and *Micromonospora haikouensis* DSM 45626^T, two highly related strains isolated from soil^{44,112}, *Micromonospora matsumotoense* DSM 44100^T and *Micromonospora rifamycinica* DSM 44983^T from rhizosphere soil^{107,108}, *M. olivasterospora* DSM 43868^T from soil¹¹³ and *M. yangpuensis* DSM 45577^T, an isolate from a sponge⁵⁶ that lies towards the periphery of the taxon. Group IV, the second largest taxon, contains thirteen strains which were recovered in three subgroups, the first of which, IVa, contains *M. lupini* Lupac 08 and the type strains of *Micromonospora coriariae*, *M. cremea, Micromonospora saelicesensis* and *Micromonospora zamorensis*, all of which were isolated from ecto- and endo-rhizospheres^{62,69,97}, and *Micromonospora chokoriensis* DSM 45160^T and *Micromonospora krabiensis* DSM 45344^T isolated from sandy and marine soils^{111,114}; in turn, group IVb is composed of the type strains of *Micromonospora citrea, Micromonospora echinaurantiaca, Micromonospora echinofusca* and *M. peucetia*, isolates from soil, chukar excrement and lake mud⁶⁸, respectively, while group IVc encompasses *Micromonospora. endolithica* DSM 44398^T and *M. nigra* DSM 43818^T, strains recovered from Antarctic sandstone⁴⁴ and a saline pond, respectively. Group V was composed of the two strains of *M. echinospora* and the type strains of *M. inyonensis, Micromonospora pallida* and *M. sagamiensis*, all of which were isolated from soil^{44,68}.

The *in silico* DNA G+C content of the *Micromonospora* genomes fell within the range 71.1 to 73.8 mol % though narrower ranges are apparent within some groups, as exemplified by the group Ia strains, which showed values within the limit 72.8–73.6 mol % (Fig. 1). The tip permutation test (Fig. 2, Supplementary Table 2) indicated that the G+C content is phylogenetically conserved (α =0.01) when calculated from the genome sequences. The genomes of 8 strains showed differences of more than one percent in G+C content when the *in silico* data were compared with results derived using experimental procedures, namely *M. aurantiaca* ATCC 27029^T (72.9% against 71.6%), *M. coriariae* DSM 44875^T (71.8% against 70.2%), *M. endolithica* DSM 44398^T (72.4% against 70%), *M. haikouensis* DSM 45626^T (73.7% against 71.5%), *M. matsumotoense* DSM 44100^T (72.3% against 71%), *Micromonospora mirobrigenesis* DSM 44830^T (72.4% against 70%), *M. rifamycinica* DSM 44983^T (73.3% against 68.6%) and *M. sediminicola* DSM 45794^T (73.6% against 74.8%). The *in silico* G+C contents of *M. citrea* DSM 43903^T, *M. echinaurantiaca* DSM 43904^T, *M. echinofusca* DSM 43913^T, *M. inyonensis* DSM 46123^T, *M. peucetia* DSM 43363^T, *M. sagamiensis* DSM 43912^T, *M. tulbaghiae* DSM 45142^T and *M. viridifaciens* DSM 43909^T were 73.8%, 73.2%, 73.3%, 71.9%, 72.3%, 72.5%, 73.0% and 72.1%, respectively; G+C contents had not been previously estimated for these strains.

Six pairs of Micromonospora type strains were considered to be closely related as their GBDP distances (the log-transformed ratios of the total number of non-identical amino-acids within the hits to the overall length of the hits in their genomes) were below 0.09; in each case digital DNA-DNA (dDDH) values were determined. Each pair, namely M. coriariae DSM 44875^T and M. cremea DSM 45599^T, M. carbonacea DSM 43168^T and M. haikouensis DSM45626^T, M. coxensis DSM 45161^T and M. halophytica DSM 43171^T, M. inyonensis DSM 46123^T and M. sagamiensis DSM 43912^T, M. mirobrigensis DSM 44830^T and M. siamensis DSM 45097^T, was found to share dDDH similarities values below the recommended 70% cut-off for the delineation of species¹¹⁵, namely 53.8%, 59.1%, 52.2%, 69.8% and 53.6%, respectively. Consequently, all of these strains can be considered to represent bona fide species. The group encompassing \dot{M} . aurantiaca ATCC 27029^T, \dot{M} . chalcea DSM 43026^T and \dot{M} . tulbaghiae DSM 45142^T gave the following dDDH values: 51.5% between M. aurantiaca ATCC 27029^T and M. chalcea DSM 43026^T, 51.3% between M. chalcea DSM 43026^T and M. tulbaghiae DSM 45142^T, and 60.1% between M. aurantiaca ATCC 27029^T and M. tulbaghiae DSM 45142^T indicating that all of these taxa should retain their species status. Corresponding dDDH values were obtained for the three M. aurantiaca strains: M. aurantiaca DSM 45487 and L5 shared 89.8% and 89.9% dDDH values with the type strain of M. aurantiaca and a 91.6% with one another indicating that they all belong to the same genomic species 115. Similarly, M. echinospora DSM 43816^T and DSM 1040 are members of the same genomic species as they shared a 78.4% dDDH value.

Insights from genome sequences. Classification. In general, little correlation was found between the groups circumscribed in the phylogenomic tree (Fig. 1) and those recovered in the trees based on single and concatenated gene sequences (Supplementary Figures 2–7). All of the groups were well supported in the GBDP analysis, apart from group III. Few of the groups delineated in the single and concatenated gene trees were supported by high bootstrap values though the five strains assigned to group V in the whole genome tree were supported by high bootstrap values in all of the other trees. The two largest groups recovered in the whole-genome tree, I and IV, were particularly fragmented in the individual and concatenated gene trees though the initial six strains assigned to group Ia, M. aurantiaca ATCC 27029^T, DSM 45487 and L5, M. chalcea DSM 43026^T, M. marina DSM 45555^T and M. tulbaghiae DSM 45142^T, were found intact in all of the single and concatenated gene trees. Similarly, the type strains of M. chokoriensis, M. coriariae, M. cremea, M. lupini, M. saelicesensis and M. zamorensis (group IVa) were recovered with high bootstrap support in all but the 16S rRNA gene analysis. More importantly, it can be seen from the principal coordinates plot (Fig. 3) that there is a closer correspondence between the whole genome and MLSA trees than with any of the those based on individual gene sequences, notably with respect to the 16S rRNA gene tree. These results are in line with those from the tip-permutation test which revealed comparatively few significantly phylogenetically conserved characters within these genes (Fig. 2).

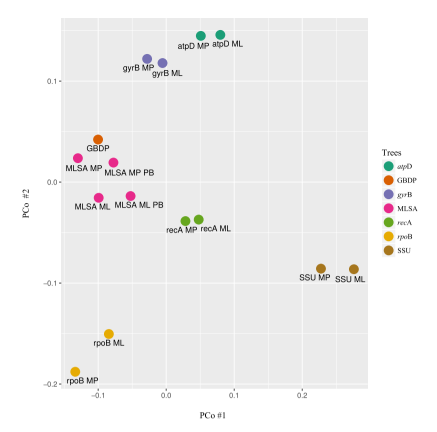


Figure 3. Principal coordinate analysis of topological distances. The analysis of the bootstrap-weighted relative Robinson-Foulds topological distances as calculated by RAxML shows that the lowest distances were between the whole genome sequence phylogeny (GBDP) and the MLSA phylogenies and the highest ones between the 16S rRNA gene phylogenies (SSU) and the GBDP tree; the distances with the other individual trees lay within these ranges. ML, maximum likelihood; MP, maximum parsimony; PB, partition bootstrap.

Concatenating such moderately informative genes apparently had the expected effect that the signal added up whereas the noise cancelled out²⁰.

Phenotypic properties. The Micromonospora strains share similar chemotaxonomic and morphological features. All of them contain meso-diaminopimelic in the cell wall peptidoglycan, xylose in whole-organism hydrolysates, complex mixtures of iso- and anteiso- branched fatty acids with predominant proportions of iso- $C_{15:0}$ and iso- $C_{16:0}$ and polar lipid patterns containing phophatidylethanolamine (diagnostic lipid), as shown in Supplementary Figure 8 and Supplementary Table 4. Most of the strains contain tetra- and hexa-hydrogenated menaquinones with ten isoprene units (MK-10 [H₄, H₆]), as shown in Supplementary Figure 8. In general, the strains grew at 20 and 37 °C, at pH 8.0 and 9.0 and in the presence of 1%, w/v sodium chloride, and produced catalase, hydrolysed aesculin and arbutin, degraded casein, starch, Tween 20 and xylan, but do not grow at 4 °C, pH 4.4 or in the presence of 5%, w/v sodium chloride (Supplementary Table 5). The tip permutation test showed that there was little evidence of correlation between the distribution of phenotypic features within the genus Micromonospora and the topology of the phylogenomic tree (Fig. 2, Supplementary Table 2). The lowest p-value recorded was for iso- $C_{17:0}$ (0.0394).

There was little sign that the distribution of phenotypic properties across the *Micromonospora* strains was influenced by the habitats from which they were isolated (Supplementary Figure 8, Supplementary Table 6) though none of the isolates from aquatic habitats contained arabinose, rhamnose or ribose in whole-organism hydrolysates or used trehalose or xylose as sole carbon sources (Supplementary Figure 8). There was some indication that strains from habitats rich in organic matter, notably from plant tissues, grew within a narrower pH range than those from soil samples, at 4°C and used alanine, propionic acid and valine as sole carbon sources (Supplementary Figure 8). The Chi-2-test shows that some of the phenotypic features were correlated, mainly due to the presence of common pathways, as exemplified by the utilization of cellobiose, melibiose, maltose and raffinose as carbon sources (Supplementary Figure 9).

Genes potentially associated with environmental adaptation. The genomes of M. citrea DSM 43903^T, M. coxensis DSM 45161^T M. echinofusca DSM 43913^T, M. endolithica DSM 44398^T, M. halophytica DSM 43171^T, M. marina DSM 45555^T, M. mirobrigensis DSM 44830^T, M. nigra DSM 43818^T and M. siamensis DSM 45097^T, isolates from diverse habitats (Supplementary Table 6), contained genes associated with photosynthesis, as described for marine bacteria ¹¹⁶; these genes belong to the proteorhodopsin family, which includes light-regulated transmembrane

proteins. The genomes of all of these strains contained genes associated with the production of β-carotene ketolase (*crt*O), lycopene β-cyclase (*crt*Y), octaprenyl diphosphate synthase (*isp*B), phytoene dehydrogenase (*crt*I), phytoene synthase (*crt*B), proteorhodopsin (*prot*) and spheroidene monooxygenase (*crt*A). Similarly, the genomes of the *M. coxensis*, *M. echinosfusca*, *M. halophytica* and *M. siamensis* strains include genes that encode for sensory rhodopsin II (SRII). The genomes of these organisms also contained 15 out of 25 genes implicated in carbon fixation in photosynthetic bacteria, 20 out of 41 genes associated with glycolysis/gluconeogenesis, 12–14 out of 58 genes associated with dicarboxylate and glyoxylate metabolism, 19 out of 31 genes implicated in phenylalanine, tyrosine and tryptophan biosynthesis and 11 out of 13 genes associated with CO₂ fixation, according to the KEGG (Kyoto Encyclopedia of Genes and Genomes) pathway database¹¹⁷ (Supplementary Table 10). It is also evident from this Table that the genomes of 23 of the strains contained a CO dehydrogenase maturation factor gene (*cox*F) associated with CO₂ fixation. In addition, the genomes of *M. aurantiaca* DSM 45487 and L5, *M. chalcea* DSM 43026^T and *M. tulbaghiae* DSM 45142^T contained a *cox*D gene, which codes for a carbon monoxide oxidation accessory protein (Supplementary Table 10).

Micromonospora strains are rich in degrading enzymes (Supplementary Table 10), as exemplified by the ability of the plant endophyte M. lupini Lupac 08 to produce amylases, cellulases, chitinases, pectinases and xylanases94. The genomes of all of the *Micromonospora* strains contained genes coding for amylases, notably α -amylases and glucoamylases. All of the micromonosporal genomes presented genes associated with cellulase production with the exception of those of M. echinaurantiaca DSM 43904^T, M. inositola DSM 43819^T and M. peucetia DSM 43363^T. The genomes of M. chalcea DSM 43026^T, M. chokoriensis DSM 45160^T, M. eburnea DSM 44814^T, M. echinospora DSM 1040 and DSM 43816^T, M. haikouensis DSM 45626^T and M. rifamycinica DSM 44983^T contained a gene encoding for a putative secreted cellulase. In turn, all of the genomes contained chiC genes, coding for chitinases, the number of these genes ranged from three in the genomes of M. pallida DSM $43817^{\rm T}$ and M. rhizosphaerae DSM 45431^T up to 13 in the genome of M. cremea DSM 45599^T. The genomes of most of the Micromonospora strains contained genes associated with the production of pectate lyases, including the hrpW gene, which codes for a harpin secreted effector that elicits the hypersensitive response in plants¹¹⁸, this gene was detected in M. coxensis DSM 45161^T, M. echinofusca DSM 43913^T and M. yangpuensis DSM 45577^T. The genomes of M. aurantiaca ATCC 27029^T, DSM 45487 and L5, M. carbonacea DSM 43168^T, M. echinospora DSM 43816^T, M. haikouensis DSM 45626^T, M. matsumotoense DSM 44100^T, M. rifamycinica DSM 44983^T and M. sagamiensis DSM 43912^T also presented genes coding for pectinesterase. Similarly, all but the type strain of M. olivasterospora have genomes associated with the production of xylanases, notably for endo-1,4-β-xylanase A precursors with an average of 21 genes per genome. All of the micromonosporal genomes contained genes that code for ß-phosphoglucomutases, enzymes associated with starch degradation, as well as those that encode for trehalose phosphorylases, enzymes associated with trehalose degradation⁹⁴. The genomes of the type strains of M. carbonacea, M. chokoriensis, M. haikouensis, M. humi, M. lupini, M. matsumotoense, M. rifamycinica, M. saelicesensis and M. zamorensis contained the trehalase gene (treA) while that of M. pallida DSM 43817T was alone in coding for trehalose 6-phosphate hydrolase (treC).

The *Micromonospora* strains have the capacity to produce plant-related hormones (Supplementary Table 10). The genomes of all of the strains contained genes predicted to code for indole-3-glycerol phosphate synthase (trpD), an intermediate in the tryptophan synthetic pathway associated with the production of indol-acetic acid (IAA) which stimulates plant growth 119. In addition, the genomes of most of the strains contained genes that coded for acetoin synthesis, which induces systemic resistance in Arabidopsis 120, exemplified by the acetolactate synthase large and small subunit genes, as well as for a gene enconding for acetoin dehydrogenase; M. aurantiaca L5, M. citrea DSM 43903^T, M. coriariae DSM 44875^T, M. cremea DSM 45599^T, M. nigra DSM 43818^T, M. olivasterospora DSM 43868^T, M. rhizosphaerae DSM 45431^T, and M. yangpuensis DSM 45577^T lack this gene. The genomes of the group V strains (M. echinospora DSM 1040 and DSM 43816^T, M. inyonensis DSM 46123^T, M. pallida DSM 43817^T and M. sagamiensis DSM 43912^T) included genes predicted to produce 2,3-butanediol dehydrogenase, an enzyme associated with the plant growth promoting hormone 2,3-butanediol and acetoin production¹²¹. However, only M. coriariae DSM 44875^T and M. krabiensis DSM 45344^T have genes predicted to encode for 1-aminocyclopropane-1-carboxylate (ACC) deaminase, a plant-growth promotor associated with the reduction of ethylene levels which lead to a reduction in plant stress¹²². Other characteristics involved in plant growth promotion include the ability to solubilize phosphates and the production of siderophores that scavenge phosphate and iron from soil making them available for plants¹²²; genes coding for the production of phosphatases and siderophores were detected in all of the Micromonospora genomes. In contrast, none of the genomes contained genes associated with the ability to fix atmospheric nitrogen.

The *Micromonospora* strains produced a well-developed substrate mycelium that carried single spores either directly or on short sporophores. None of the strains formed aerial hyphae though the genomes of almost half of them showed the presence of a predicted surface active peptide cluster (sapB) that encodes for a lantabiotic-like peptide which has been considered to trigger the formation of aerial hyphae when strains are grown on rich media¹²³. Other genes related to sporulation were found in all of the strains, namely whiB and whiD genes, which are required for the differentiation of aerial hyphae into mature spores in $Streptomyces^{124}$. Another characteristic of Micromonospora strains is their ability to produce a range of pigments at the onset of spore production. The genomes of all of the tested strains, apart from M. cremea DSM 45599^T , contained whiE-ks, whiE-clf, whiEI, whiEII, whiEVI, and whiEVII genes which are associated with spore pigment production in $Streptomyces^{125}$. All of the strains contained genes coding for the production of pigments, as well as biosynthetic gene clusters associated with the production of carotenoid, isorenieratene and sioxanthin compounds (Supplementary Table 7). Between the genes implicated in carotenoid production pathway detected in most of the Micromonospora genomes were the putative genes encoding for β -carotene ketolases, phi-carotenoid synthases, geranylgeranyl pyrophosphate synthetases, lycopene cyclases, phytoene synthases and squalene-hopene cyclases (Supplementary Table 10).

The genomes of the *Micromonospora* strains contained a range of genes associated with DNA repair systems (Supplementary Table 10). All of the genomes included at least one copy of excinuclease subunits A, B and C (*uvr*A, *uvr*B, *uvr*C genes) and three copies of ATP-dependent DNA helicase (*uvr*D), one of which has only been associated with actinobacteria (*uvr*D-actino). The *Micromonospora* genomes were also rich in *rec* genes, implicated in recombination, in the production of helicases, and for general DNA repair, but only the genome of *M. nigra* DSM 43818^T presented the *rec*B gene, which codes for an exodeoxyribonuclease. Similarly, all of the genomes contained exodeoxyribonuclease genes (*xse*A, *xse*B, and *exo*III), as well as genes associated with the production of several exo- and endonucleases, namely *sbc*C, *sbc*D, *end*1, *endo*IV, *endo*V.

Genes associated with stress responses. The genomes of Micromonospora strains annotated by RAST^{126,127} and analyzed through the SEED viewer¹²⁸ showed between 115 and 144 putative genes known to be associated with stress responses, notably those encoding for carbon starvation, heat shock responses, osmoregulation and oxidative stress (Supplementary Table 10). The genomes of all of the strains contained cspA and cspC genes, which encode for families of proteins that respond to cold shock 129 , and dnaK, grpE and hrcA genes involved in heat shock responses¹³⁰. In contrast, cspG genes, that encode for a cold shock protein associated with cellular SOS repair systems¹³¹, were restricted to the genomes of M. aurantiaca DSM 45487, M. echinospora DSM 43816^T and M. krabiensis DSM 45344^T. All of the Micromonospora genomes contained betC and proU genes which govern the uptake of betaine and choline, metabolites that contribute to responses to oxidative stress^{132,133}. Other universally distributed stress related genes include those that encode for alkyl hydroperoxidase reductases 134 (ahpC genes), redox sensitive transcriptional regulators 135,136 (rex and sox genes), iron-stress related fur genes 137, and the nitric oxide dioxygenase gene (hmpX), which is induced by the presence of NO and prevents the inhibition of growth caused by nitrosative stress¹³⁸; around half of the genomes showed the presence of superoxide dismutase genes (sod). Genes associated with ectoine biosynthesis (ect) for osmoregulation were found in the genomes of six Micromonospora strains (Supplementary Table 10); genes encoding for diaminobutyrate-pyruvate aminotransferases (ectB genes) were present in the genomes of M. chersina DSM 44151^T, M. echinospora DSM 43816^T, M. endolithica DSM 44398^T, M. matsumotoense DSM 44100^T and M. peucetia DSM 43363^T, the genome of the remaining strain, M. eburnea DSM 44814^T contained ectC genes that encode for L-ectoine synthases. The genomes of most of the Micromonospora strains contained rspA genes that code for starvation sensing protein A which may help them to survive in low carbon habitats by activating peptide uptake^{139–141}; these genes were not detected in the genomes of the type strains of M. auratinigra, M. coxensis, M. halophytica, M. inyonensis, M. marina, M. nigra, M. olivasterospora, M. sagamiensis, M. sediminicola or M. siamensis (Supplementary Table 10).

Biosynthetic gene clusters coding for specialized metabolites. All of the Micromonospora genomes were screened for candidate biosynthetic gene clusters using the specialized metabolite identification pipeline antiSMASH. The number of such putative bioclusters ranged from 7 in the genomes of M. cremea DSM 45599^T and M. rhizosphaerae DSM 45431^{T} to 48 in that of M. matsumotoense DSM 44100^{T} ; the genomes of the type strains of M. carbonacea, M. echinospora, M. haikouensis, M. marina, M. pallida and M. sagamiensis were also rich in such biosynthetic gene clusters (Fig. 4). The average numbers of biosynthetic gene clusters detected in the genomes of the Micromonospora strains was twenty, most of which seem to be related with antibiotic, siderophore and terpene production (Fig. 4, Supplementary Table 7). Just over 22% of the biosynthetic gene clusters present in the micromonosporal genomes (206 out of 915) lacked any homology with known bioclusters; these biosynthetic gene clusters belong to several cluster types, notably terpenes (82), non ribosomal peptides (27), lantipeptides (26) and polyketide synthases (16) (Fig. 4, Supplementary Table 7). The remaining bioclusters showed similarities to a greater or lesser extent for known compounds though most of them (85%) showed less than 75% similarity of their genes with known bioclusters. A total of 172 different bioclusters were detected in all the genomes, most of them related to antibiotic production though only 33 presented a similarity of over 50% with known compounds (Supplementary Table 7). Thirteen of these compounds were related to known antibiotics (actinorhodin, chloramphenicol, diazepinomicin, leucanicidin, livipeptin, lobosamide, micromonolactam, rishirilide B, salinilactam, sibiromycin, streptazone tiacumicin B, TLN-05220) with similarity values of their genes over 75% (Supplementary Table 7).

Two biosynthethic gene clusters were found in the genomes of all of the *Micromonospora* strains, one related to the production of alkyl-O-dihydrogeranyl-methoxyhydroquinone (with ~70% similarity) and the other with a bacteriocin-terpene related to the production of lymphostin (with ~40% similarity). Similarly, the sioxanthin biosynthetic gene cluster was found in all of the *Micromonospora* strains, apart from *M. inositola* DSM 43819^T and *M. pallida* DSM 43817^T. In contrast, most of the biosynthetic gene clusters had a limited distribution, 83 were found in a single genome, as exemplified by the one related to the carotenoid biosynthetic gene cluster present in the *M. pallida* genome; 34 biosynthetic gene clusters were detected in just two of the *Micromonospora* genomes, as illustrated by the chlorothricin biosynthetic gene cluster present in the genomes of the type strains of *M. eburnea* and *M. endolithica*. The sequences of five biosynthetic gene clusters, apart from the one encoding for sioxanthin, were identical to ones known to code for geosmin¹⁴² as found in the genome of *M. pallida* DSM 43817^T; leucanicidin, a potent nematocide¹⁴³, as seen in the genome of *M. carbonacea* DSM 43168^T; livipeptin, an aldehyde peptide¹⁴⁴, as detected in the genomes of *M. echinofusca* DSM 43913^T and *M. peucetia* DSM 43365^T; micromonolactam, a polyene macrolactam¹⁴⁵, as found in the genome of *M. haikouensis* DSM 45626^T while SapB, which is associated with aerial hyphae formation¹²³, was found in almost half of the *Micromonospora* genomes (Supplementary Table 7).

The genomes of several *Micromonospora* strains assigned to taxa defined in the whole genome tree (Fig. 1) included specific biosynthetic gene clusters associated with the synthesis of known bioactive compounds. This was particularly so with the group V strains, namely *M. echinospora* DSM 1040 and DSM 43816^T, *M. inyonensis* DSM 46123^T, *M. pallida* DSM 43817^T and *M. sagamiensis* DSM 43912^T. The genomes of these strains contain bioclusters which show similarities against (i) feglymycin, a peptide antibiotic produced by *Streptomyces* sp. DSM

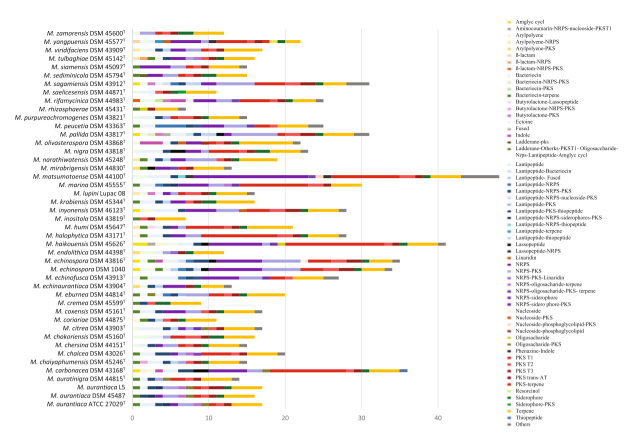


Figure 4. Biosynthetic gene clusters found in the *Micromonospora* genomes using antiSMASH 3.0. Highly variable profiles were found between the strains. The genomes of the *Micromonospora* strains were found to be especially rich in NRPS, PKS and terpene clusters; whereas there was also an abundance of bacteriocin, lantipeptide and siderophores clusters.

11171¹⁴⁶ that inhibits HIV cell to cell transfer¹⁴⁷ (this biosynthetic gene cluster was also detected in the genome of *M. rifamycinica* DSM 44983^T); (ii) gentamicin produced by *M. echinospora* NRRL 2953 and NRRL 2985^{T 71} (this biocluster was also found in the genomes of *M. carbonacea* DSM 43168^T, *M. haikouensis* DSM 45626^T, *M. matsumotoense* DSM 44100^T, *M. peucetia* DSM 43363^T and *M. yangpuensis* DSM 45577^T; (iii) herbimycin, isolated from *Streptomyces* sp. RM-7–15¹⁴⁸, which shows activity against herbs and heat shock protein 90, this biosynthetic gene cluster was present in the genome of *M. narathiwatensis* DSM 45248^T, and (iv) TLN-05220, a product of *M. echinospora* NRRL 1225, which shows activity against methicillin-resistant strains of *Staphylococcus aureus*, vancomycin-resistant enterococci and several human cell lines¹⁴⁹. Further, apart from the type strain on *M. inyonensis*, the genomes of the group V strains had a biosynthethic gene cluster related to crocacin, an electron transport inhibitor isolated from *Chondromyces crocatus* CM c3 that inhibits Gram-positive bacteria and fungi¹⁵⁰ while the genomes of *M. echinospora* DSM 1040 and *M. sagamiensis* DSM 43912^T contain a biocluster associated with the production of muraymycin, a nucleoside-lipopeptide antibiotic synthesized by *Streptomyces* sp. LL-AA896 that inhibits peptidoglycan biosynthesis¹⁵¹.

The genomes of the initial six strains of the group 1a strains, namely M. aurantiaca ATCC 27029^T, DSM 45487 and L5, M. chalcea DSM 43026^T, M. marina DSM 45555^T, and M. tulbaghiae DSM 45142^T, include biosynthetic gene clusters with the potential to synthesize known specialized metabolites. These strains, apart from M. chalcea DSM 43026^T, contain a biocluster which present some similarity with the biocluster responsible of leinamycin production, a potent antitumor antibiotic produced by Streptomyces strains 152 (this biosynthetic gene cluster is also present in the genomes of M. haikouensis DSM 45626^T, M. matsumotoense DSM 44100^T and M. purpureochromogenes DSM 43821^T). In turn, the genomes of all but the M. marina strain contain a biocluster related to nocathiacin, a thiazole peptide antibiotic produced by Nocardia sp. WW-12651153, derivatives of which have been used to treat multidrug-resistant bacterial infections¹⁵⁴ although in this instance the similarity between the two bioclusters is low (this biocluster is also present in the genomes of M. coxensis DSM 45161^T, M. humi DSM 45647^T, M. peucetia DSM 43363^T and M. sediminicola DSM 45794^T). The genomes of the three M. aurantiaca strains have a biocluster associated with the production of dynemicin, a 1,5-diyn-3-ene-containing antibiotic produced by M. chersina¹⁵⁵ with antibacterial and antitumor activities (this biocluster was also detected in the genomes of M. chersina DSM 44151^T and M. yangpuensis DSM 45577^T). However, none of the different types of biosynthetic gene clusters appeared to be phylogenetically conserved ($\alpha = 0.01$) in the tip permutation test (Fig. 2, Supplementary Table 2). The distribution of bioclusters was not related either to the origin of the strains, though there was an average of 21 bioclusters in the genomes of the strains isolated from soil, sediment and liquid environments strains and 17 in the genomes of those isolated from organic material. In contrast, only the genome of *M. marina* DSM 45555^T contained the kiamycin biosynthetic cluster which has been detected in marine-related organisms¹⁵⁶.

Discussion

The results of this study provide further evidence that data generated in whole genome sequencing studies provide an essential framework for the reclassification of taxonomically complex prokaryotic taxa previously defined from analyses of relatively few taxonomic features^{25,26,159}. It is evident from the *Micromonospora* phylogenomic tree that the tested strains not only form a monophyletic group but fall into several well supported phyletic lines, only two of which were recognised in their entirety in the corresponding trees based on single gene sequences. The six initial members of group 1a, namely *M. aurantiaca* ATCC 27029^T, DSM 45487 and L5, *M. chalcea* DSM 43026^T, *M. marina* DSM 45555^T and *M. tulbaghiae* DSM 45142^T, were defined in the *atp*D, *gyr*B, *rec*A, *rpo*B and 16S rRNA gene trees, as well as in the MLSA tree based on all of the individual gene sequences. Similarly, the group V strains, *M. echinospora* DSM 1040 and DSM 43816^T, *M. inyonensis* DSM 46123^T, *M. pallida* DSM 43817^{TT}, and *M. sagamiensis* DSM 43912^T, was recovered in all the phylogenetic trees.

Genomic DNA G+C content, that is, the proportion of cytosine and guanine moieties over the overall number of nucleotides in the genome, feature prominently in the description of prokaryotic genera and species^{8,160,161}. DNA base composition values based on the application of conventional methods are considered to be indirect values as they do not count nucleotides, but estimate genomic G+C content from physical properties drawn from analyses of extracted and/or digested DNA162. However, it is becoming increasingly apparent that estimates of G+C content taken directly from whole genome sequences are of higher quality than those derived from well known experimental methods¹⁶³. Indeed, these workers have shown that strains within a species have G+C values within a 1% range. It is, therefore, encouraging that in the present study a statistically sound relationship was found between in silico G+C values and the distribution of the Micromonospora strains within the phylogenomic tree. Moreover, in silico G+C values of the tested type strains fell within the range 71.1–73.8 mol % with narrower ranges found for strains assigned to well supported phyletic lines, as exemplified by the group 1a and IVa strains which showed values of 72.8-73.6 and 71.1-72.0 mol %, respectively. The genomes of eight of the Micromonospora type strains showed more than a 1% difference when in silico G+C values where compared with corresponding results found using conventional laboratory based methods. It is important to resolve such discrepancies between G+C values so that differences between closely related species are not obscured 159,164. Emended descriptions are given for these Micromonospora species and for an additional eight species that previously lacked estimates of DNA G+C values.

DNA-DNA hybridization (DDH) is still widely used to estimate genetic relatedness between closely related bacteria as it is seen to be the "gold standard" for species delineation between prokaryotes¹⁶⁵. Indeed, the recommendation of Wayne and his colleagues that a DDH of 70% for the prokaryotic species boundary has been widely followed by the systematic community¹¹⁵. It is now evident that dDDH methods based on comparisons of whole genome sequences provide better quality data for discriminating between closely related strains than corresponding values derived from the application of experimental methods that are well known to be expensive, labour-intensive and prone to experimental error^{166–168}. dDDH values estimated from the genomes of the six pairs of closely related type strains showed that they fell below the 70% threshold¹¹⁵ indicating that *M. coriariae*⁶² and *M. cremea*⁹⁷, *M. carbonacea*⁹⁶ and *M. haikouensis*¹¹², *M. coxensis*¹¹¹ and *M. halophytica*⁷⁶, *M. mirobrigensis*¹⁰⁹ and *M. siamensis*¹¹⁰, *M. inyonensis*⁶⁸ and *M. sagamiensis*⁶⁸ are validly named species.

Kroppenstedt and his colleagues⁶⁸ recognised that the type and only representatives of *M. inyonensis* and *M. sagamiensis* were closely related but could be distinguished based on cultural and phenotypic properties, by their fatty acid and MALDI-TOF mass spectrometric profiles and by a DDH value of 61.3%¹¹⁵. In the present study, these strains were found to share a dDDH value marginally below the recommended cut-off point, but were distinguished readily by the number and type of their biosynthetic gene clusters, by the presence of different stress genes in their genomes and by differences in the composition of nine of the COG groups, notably those belonging to the categories G, R and X. In light of all of these data it can be concluded that *M. inyonensis* and *M. sagamiensis* strains belong to different, but closely related species.

The family *Micromonosporaceae* encompasses several genera, such as *Salinispora*^{169,170}, that are difficult to distinguish from *Micromonospora* strains using conventional genotype and phenotype procedures^{45,171}. The phylogenomic classification of the representative *Micromonospora* type strains not only provides a framework for clarifying relationships with those from related genera but also allows the taxonomic provenance of *Micromonospora* strains to be established. It is interesting that the genera *Micromonospora* and *Salinispora* are quite sharply separated, albeit closely related, in the phylogenomic tree though the genomes of additional representatives of these taxa need to be examined to underscore precise relationships between them. It is encouraging that the authenticity of *M. aurantiaca* DSM 45487 and L5, and *M. echinospora* DSM 1040 were confirmed in the present study.

There were few signs of concordance between the distribution of chemotaxonomic and other phenotypic markers drawn from the original descriptions of the *Micromonospora* type strains and their assignment to taxa in the phylogenomic tree. This lack of congruence can be attributed to factors such as the use of such a small sample

of strains and tests, reliance on inappropriate and/or unreliable phenotypic tests and failure to use appropriate reference material. Sutcliffe and his colleagues have stressed the need to address such issues. A better understanding of the relationship between genotype and phenotype can be expected to provide a way forward on such matters. In sharp contrast to the issues raised above, all of the type strains produced whole organism hydrolysates rich in meso-diaminopimelic acid and xylose, major amounts of saturated and unsaturated fatty acids, notably iso- $C_{15:0}$ and iso- $C_{16:0}$, a polar lipid pattern containing phosphatidylethanolamine (diagnostic lipid) and usually diphosphatidylgycerol and phosphatidylinositol (phospholipid pattern 2 sensu Lechevalier et al.¹⁷²) and tetraand hexa- hydrogenated menaquinones with either nine or ten isoprene units as predominant isoprenologues. Such genus specific properties are of particular value in distinguishing Micromonospora from most of the other genera classified in the family $Micromonosporaceae^{45}$. Similarly, many of the Micromonospora strains share phenotypic features, as exemplified by their ability to hydrolyse aesculin and arbutin and degrade casein, starch, Tween 20 and xylan. In contrast, very few of the Micromonospora strains grew at 4 °C, pH 4.4 or in the presence of 5% w/v sodium chloride.

In general, *Micromonospora* species have been associated with aquatic and terrestrial habitats across diverse geographical regions thereby underscoring their adaptability⁴⁴. More recently, they have been recovered from the tissues of a broad range of plants^{173–175}. In the present study, little correlation was found between the source of the *Micromonospora* strains and their distribution to taxa delineated in the phylogenomic tree. However, it is interesting that all of the strains isolated from ecto- or endo-rhizospheres, namely *M. coriariae* DSM 44875^T from a root nodule of *Coriaria myrtifolia*, *M. lupini* Lupac 08 and *M. saelicesensis* DSM 44871^T from root nodules of *Lupinus angustifolius*, and *M. cremea* DSM45599^T and *M. zamorensis* DSM 45600^T from the rhizosphere of *Pisum sativum* were recovered in the well delineated subgroup IVa. Associations such as these would be much easier to establish if more details were given on the sources of strains in species description of prokaryotes. Indeed, such information is a prerequisite for data-driven prokaryotic systematics¹⁷⁶.

It is now well known that micromonosporae are associated with roots of diverse plant species^{173–175}, notably nodules of healthy leguminous plants^{53,57,69,94,177}. The discovery that *Micromonospora* strains occupy nitrogen-fixing nodules poses several intriguing questions such as whether they are in transition from a saprophytic to a facultatively endophytic lifestyle and whether they have a beneficial effect on the plant. In general, deductions drawn from the genomes of the tested strains underpin key genome features captured by Trujillo and her colleagues for *M. lupini* strain Lupac 08⁹⁴. Some, if not all of the genomes of the *Micromonospora* strains, like the *M. lupini* strain, have putative genes that encode for acetoin, 2,3-butanediol dehydrogenase and indol-3-acetic acid, auxinic phytohormones implicated in phytostimulation^{119,122,178,179}. Along similar lines, the genomes of all of these strains are rich in putative genes that code for antibiotics, chitin degradation and siderophores, compounds that may contribute to the defence of the host plant against root infecting fungi. It is also interesting that the genomes of a few *Micromonospora* strains, including *M. lupini* strain Lupac 08, contained genes encoding trehalase, an enzyme that degrades trehalose and is implicated in nodule growth regulation^{180,181}. All of these observations indicate that micromonosporae confer protection to the plant. It has also been shown that inoculation of strain Lupac 08 into legumes contributes to the welfare of the host plant⁹⁴. An important conclusion drawn from this study is that micromonosporal genomes lack *nif*H-like fragments, despite early claims to the contrary^{174,177}.

It is still too early to draw far reaching conclusions about the ecological roles of facultatively endophytic micromonosporae, as their genomes have an array of putative genes that code for degradative enzymes involved in the turnover of plant polymers, notably amylases, cellulases, chitinases, pectinases and xylanases. Indeed, *Micromonospora* strains may have the capacity, proven in the case of strain Lupac 08, to produce a range of degradative enzymes that are characteristic of saprophytic bacteria. This picture is clouded even further as the genomes of the *Micromonospora* strains isolated from diverse habitats encoded for much the same traits as the endophytic strains. It could be that micromonosporae have the capacity to colonize multiple ecological niches though additional studies are required to address this point.

There are several reports that *Micromonospora* strains can form sterile aerial hypahe^{45,182-185} and one which presented evidence that on certain nutrient media micromonosporae from marine sediments form aerial mycelia that can be used to propagate fresh colonies¹⁸⁶. Baldacci and Locci¹⁸⁷ found that strains designated as "Micromonospora melanosporea" formed aerial mycelia with short branching sporophores bearing single spores. In light of these observations it is interesting that the genomes of many of the *Micromonospora* type strains showed the presence of putative genes associated with aerial hyphae formation and spore maturation in streptomycetes^{123,125,188} though *whi*B and *whi*D like genes have been shown to have a role as transcription factors in mycobacteria¹⁸⁹. The *whi*E genes detected in the genomes of all of the *Micromonospora* strains, apart from *M. cremea* DSM 45599^T, are associated with the final stage of sporulation, when polyketide pigments are formed in the spore coat¹²⁵. It is possible that the *whi*E genes may be involved in the formation of black pigments that are produced towards the end of the micromonosporal growth cycle. It is also plausable that over evolutionary time micromonosporae have lost the capacity to form spores on aerial hyphae.

The genomes of most of the *Micromonospora* strains contained a broad range of genes associated with the synthesis of pigments, notably, carotenoids, isorenieratene and sioxanthin. Although the biosynthetic cluster associated with the production of carotenoids was only found in the genome of *M. pallida* DSM 43817^T, the biocluster for the synthesis of sioxanthin has been associated with the production of a novel glycosylated carotenoid in *Salinispora* strains¹⁹⁰. This sioxanthin biosynthetic gene cluster was found in all of the *Micromonospora* genomes, apart from those of the type strains of *M. pallida* and *M. inositola*; only the *M. inositola* strain was shown to have the capacity to produce isorenieratene, an aromatic carotenoid produced by green photosynthetic bacteria and a few actinobacteria ¹⁹¹. All of the *Micromonospora* genomes contained genes implicated in carotenoid biosynthesis, as exemplified by those coding for the production of β-carotene ketolases, phi-carotenoid synthases and lycopene β-cyclases^{192,193}. The presence of such compounds in non-photosynthetic organisms has been associated with UV protection¹⁹⁴ and in the case of photosynthetic bacteria with light harvesting complexes¹⁹⁵. Additional work is

needed to account for the presence of genes associated with photosynthesis that were detected in the genomes of strains assigned to groups Id (*M. mirobrigensis* DSM 44830^T and *M. siamensis* DSM 45097^T), II (*M. coxensis* DSM 45161^T and *M. halophytica* DSM 43171^T), IVb (*M. citrea* DSM 43903^T and *M. echinofusca* DSM 43913^T) and IVc (*M. endolithica* DSM 44398^T and *M. nigra* DSM 43818^T).

It was particularly interesting that the genome of most, if not all, of the *Micromonospora* strains were replete with genes relevant to their ability to adapt to low levels of carbon^{139–141}, temperature fluxes^{129,130}, and changes in the osmotic environment^{132,133}, a combination of key environmental variables that lend further weight to the suggestion that micromonosporae may be able to colonise multiple microhabitats⁴⁵. In addition, the micromonosporal genomes included genes associated with protection against UV-radiation and for repairing DNA damage. Indeed, all of the strains were found to have the potential to protect and repair damage caused by UV radiation as they have genes associated with the synthesis of Uvr ABCD proteins, excision proteins that have been reported in several bacteria¹⁹⁶. Further, mutations in uvr ABC genes have been associated with UV sensitivity in *Rhodobacter sphaeroides*¹⁹⁷. Genes associated with desiccation were not detected in the genomes of the *Micromonospora* strains though several genes involved in the biosynthesis and uptake of trehalose were seen, trehalose has been linked with tolerance to heat and desiccation in bacteria¹⁹⁸.

Since the discovery of gentamicin from "M. purpurea" INMI 632 in 1963⁷¹ hundreds of bioactive molecules with diverse properties and structures have been isolated from *Micromonospora* species^{83,199–202}. Major classes of clinically significant specialized metabolites synthesized by micromonosporae include aminoglycosides (gentamicins), anthracyclines (daunorubicin), ansamycins (rifampicins), macrolides (erythromicins), as well as enediyne (calichenomicins) and oligosaccharide (everninomicins) antibiotics. It is not surprising in light of these observations, those drawn from earlier whole genome studies on micromonosporae^{85–87} and from corresponding work on other filamentous actinobacteria^{39,203} that the genomes of the tested strains were rich in biosynthetic gene clusters encoding for known and predicted specialized metabolites, notably antibiotics, siderophores and terpenes. The analysis of the micromonosporal genomes also confirmed the relationship between presence of the aminoshikimate pathway and the capacity of *Micromonospora* strains to synthesize ansamycins; the genomes of all the rifamycin-like producers contained the AHBA synthase-like gene, a key enzyme of this variant of the shikimate pathway implicated in the production of aromatic antibiotics. It is particularly interesting that many of the bioclusters were found only in a few of the micromonosporal genomes, an observation that underlines the merit of selecting representatives of novel actinobacterial taxa in the search for new specialized metabolites^{204,205}, thereby providing further evidence that comparative analysis of actinobacterial genomes can be used to select gifted strains for gene mining and natural product discovery^{30,206,207}. In contrast, it was not possible to detect any relationship between the phylogeny of Micromonospora strains and their source through such an association has been found for Salinispora species^{208–211}. Indeed, strains assigned to most of the clades and subclades were isolated from diverse geographical regions. It can also be concluded from the analyses of the genomes generated in this study that micromonosporae have a very much greater potential to synthesize specialized metabolites, notably antibiotics, than previously realised. Consequently, Micromonospora and other genera classified in the family Micromonosporaceae⁴⁵ should feature much more prominently in the search for new classes of bioactive compounds that are urgently needed to control drug resistant pathogens.

Revision to descriptions of *Micromonospora* **species.** *Emended description of Micromonospora aurantiaca Sveshnikova et al.* 1969. The species description is as given by Sveshnikova *et al.*²¹² with the following changes: The approximate genome size of the type strain is 7.03 Mbp and its genome G+C content is 72.9%.

Emended description of Micromonospora auratinigra Thawai et al. 2004. The species description is as given by Thawai et al. 102 with the following changes: The approximate genome size of the type strain is 6.76 Mbp.

Emended description of Micromonospora carbonacea Luedemann and Brodsky 1965. The description is as given by Luedemann and Brodsky 96 with the following changes: The approximate genome size of the type strain is 7.94 Mbp and its genome G+C content is 73.8%.

Emended description of Micromonospora chaiyaphumensis Jongrungruangchok et al. 2008. The species description is as given by Jongrungruangchok et al. 104 with the following changes: The approximate genome size of the type strain is 6.74 Mbp.

Emended description of Micromonospora chalcea (Foulerton 1905) Ørskov 1923. The species description is as given by Genilloud⁴⁴ with the following changes: The approximate genome size of the type strain is 6.99 Mbp.

Emended description of Micromonospora chersina Tomita et al. 1992. The species description is as given by Tomita et al. 103 with the following changes: The approximate genome size of the type strain is 6.68 Mbp and its genome G+C content is 73.6%.

Emended description of Micromonospora chokoriensis Ara & Kudo 2007. The species description is as given by Ara and Kudo¹¹¹ with the following changes: The approximate genome size of the type strain is 6.89 Mbp.

Emended description of Micromonospora citrea Kroppenstedt et al. 2005. The species description is as given by Kroppenstedt et al. 68 with the following changes: The approximate genome size of the type strain is 7.21 Mbp and its genome G+C content is 73.8%.

Emended description of Micromonospora coriariae Trujillo et al. 2006. The species description is as given by Trujillo et al. 62 with the following changes: The approximate genome size of the type strain is 6.93 Mbp and its genome G+C content is 71.8%.

Emended description of Micromonospora coxensis Ara & Kudo 2007. The species description is as given by Ara and Kudo¹¹¹ with the following changes: The approximate genome size of the type strain is 6.77 Mbp.

Emended description of Micromonospora cremea Carro et al. 2012. The species description is as given by Carro *et al.*⁹⁷ with the following changes: The approximate genome size of the type strain is 7.76 Mbp.

Emended description of Micromonospora eburnea Thawai et al. 2005. The species description is as given by Thawai *et al.* 105 with the following changes: The approximate genome size of the type strain is 7.18 Mbp.

Emended description of Micromonospora echinaurantiaca Kroppenstedt et al. 2005. The species description is as given by Kroppenstedt et al. 68 with the following changes: The approximate genome size of the type strain is 7.20 Mbp and its genome G+C content is 73.2%.

Emended description of Micromonospora echinofusca Kroppenstedt et al. 2005. The species description is as given by Kroppenstedt et al. 68 with the following changes: The approximate genome size of the type strain is 7.00 Mbp and its genome G+C content is 73.3%.

Emended description of Micromonospora echinospora Luedemann and Brodsky 1964 emend. Kasai et al. 2000. The species description is as given by Kasai et al. 70 with the following changes: The approximate genome size of the type strain is 7.78 Mbp and its genome G+C content is 72.3%.

Emended description of Micromonospora endolithica Hirsch et al. 2004. The species description is as given by Hirsch et al. 66 with the following changes: The approximate genome size of the type strain is 7.03 Mbp and its genome G+C content is 72.5%.

Emended description of Micromonospora haikouensis Xie et al. 2012. The species description is as given by Xie *et al.*¹¹² with the following changes: The approximate genome size of the type strain is 7.58 Mbp and its genome G+C content is 73.7%.

Emended description of Micromonospora halophytica Weinstein et al. 1968. The species description is as given by Weinstein et al. 76 with the following changes: The approximate genome size of the type strain is 6.27 Mbp and its genome G+C content is 73.0%.

Emended description of Micromonospora inositola Kawamoto et al. 1974. The species description is as given by Kawamoto et al. 107 with the following changes: The approximate genome size of the type strain is 6.71 Mbp and its genome G+C content is 72.2%.

Emended description of Micromonospora inyonensis Kroppenstedt et al. 2005. The species description is as given by Kroppenstedt et al. 68 with the following changes: The approximate genome size of the type strain is 6.92 Mbp and its genome G+C content is 71.9%.

Emended description of Micromonospora krabiensis Jongrungruangchok et al. 2008. The species description is as given by Jongrungruangchok *et al.*¹¹⁴ with the following changes: The approximate genome size of the type strain is 7.07 Mbp.

Emended description of Micromonospora marina Tanasupawat et al. 2010. The species description is as given by Tanasupawat *et al.*⁹⁵ with the following changes: The approximate genome size of the type strain is 6.06 Mbp.

Emended description of Micromonospora matsumotoense (Asano et al. 1989) Lee et al. 1999. The species description is as given by Lee et al. 213 with the following changes: The approximate genome size of the type strain is 7.75 Mbp and its genome G+C content is 72.3%.

Emended description of Micromonospora mirobrigensis Trujillo et al. 2005. The species description is as given by Trujillo et al. 109 with the following changes: The approximate genome size of the type strain is 6.17 Mbp and its genome G+C content is 73.3%.

Emended description of Micromonospora narathiwatensis Thawai et al. 2008. The species description is as given by Thawai et al. 106 with the following changes: The approximate genome size of the type strain is 6.61 Mbp and its genome G+C content is 72.6%.

Emended description of Micromonospora nigra (Weinstein et al. 1968) Kasai et al. 2000. The species description is as given by Kasai et al. 70 with the following changes: The approximate genome size of the type strain is 6.36 Mbp and its genome G+C content is 72.6%.

Emended description of Micromonospora olivasterospora Kawamoto et al. 1983. The species description is as given by Kawamoto *et al.*¹¹³ with the following changes: The approximate genome size of the type strain is 7.07 Mbp and its genome G+C content is 72.5%.

Emended description of Micromonospora pallida (Luedemann and Brodsky 1964) Kasai et al. 2000. The species description is as given by Kasai et al. 70 with the following changes: The approximate genome size of the type strain is 7.76 Mbp and its genome G+C content is 71.9%.

Emended description of Micromonospora peucetia Kroppenstedt et al. 2005. The species description is as given by Kroppenstedt et al. 68 with the following changes: The approximate genome size of the type strain is 7.37 Mbp and its genome G+C content is 72.3%.

Emended description of Micromonospora purpure ochromogenes (Waksman and Curtis 1916) Luedemann 1971. The species description is as given by Luedemann 214 with the following changes: The approximate genome size of the type strain is 6.67 Mbp and its genome G+C content is 73.0%.

Emended description of Micromonospora rhizosphaerae Wang et al. 2011. The species description is as given by Wang *et al.* 108 with the following changes: The approximate genome size of the type strain is 7.18 Mbp.

Emended description of Micromonospora rifamycinica Huang et al. 2008. The species description is as given by Huang et al. 215 with the following changes: The approximate genome size of the type strain is 7.01 Mbp and its genome G+C content is 73.0%.

Emended description of Micromonospora saelicesensis Trujillo et al. 2007. The species description is as given by Trujillo *et al.* 69 with the following changes: The approximate genome size of the type strain is 7.10 Mbp.

Emended description of Micromonospora sagamiensis Kroppenstedt et al. 2005. The species description is as given by Kroppenstedt et al. 68 with the following changes: The approximate genome size of the type strain is 6.93 Mbp and its genome G+C content is 72.5%.

Emended description of Micromonospora sediminicola Supong et al. 2013. The species description is as given by Supong et al. 101 with the following changes: The approximate genome size of the type strain is 6.89 Mbp and its genome G+C content is 73.6%.

Emended description of Micromonospora siamensis Thawai et al. 2006. The species description is as given by Thawai *et al.*¹¹⁰ with the following changes: The approximate genome size of the type strain is 6.25 Mbp.

Emended description of Micromonospora tulbaghiae Kirby and Meyers 2010. The species description is as given by Kirby and Meyers⁵⁹ with the following changes: The approximate genome size of the type strain is 6.49 Mbp and its genome G+C content is 73.0%.

Emended description of Micromonospora viridifaciens Kroppenstedt et al. 2005. The species description is as given by Kroppenstedt et al. 68 with the following changes: The approximate genome size of the type strain is 7.07 Mbp and its genome G+C content is 72.1%.

Emended description of Micromonospora yangpuensis Zhang et al. 2012. The species description is as given by Zhang et al. 56 with the following changes: The approximate genome size of the type strain is 6.52 Mbp.

Emended description of Micromonospora zamorensis Carro et al. 2012. The species description is as given by Carro *et al.*⁹⁷ with the following changes: The approximate genome size of the type strain is 7.09 Mbp.

Conclusions

The results of this and corresponding taxonomic analyses based on a comparison of whole genome sequences of bacterial taxa^{159,216,217} are a timely reminder that classification and identification of prokaryotes are markedly data dependent and hence are in a constant state of development due to the introduction of new technologies³. To date, much of the emphasis in the GEBA project has been on the analysis of genomic sequences generated from the type strains of diverse taxa in order to expand coverage of the tree of life^{35–39} while other sequence based studies have been focused on many representatives of individual clinically significant bacterial species in order to enhance understanding of pathogenesis^{218–220}. It is evident from the present study that the analysis of genome sequences of taxonomically complex genera offers a halfway house between these contrasting approaches to phylogenomics, one which has led to substantial improvements in the classification of the genus *Micromonospora*. In addition, the associated wealth of biological information provides a unique platform for the search and discovery of novel natural products, using genome mining and genetic engineering procedures, while providing leads to unravelling the ecological roles of micromonosporae.

Methods

Strains and DNA isolation. To cover the ecologic diversity of micromonosporae, strains of forty *Micromonospora* species (40 type and 2 non-type strains) were obtained from the DSMZ collection (Supplementary Table 8). All of the strains were grown in DSM medium 65 at 28 °C for 7 days when the biomass was harvested. Genomic DNA was extracted from the biomass preparations using a MasterPure ™ Gram Positive

DNA Purification Kit (Epicentre MGP04100) following the instructions of the manufacturer, albeit with modifications, namely incubation overnight on a shaker with $10\,\text{mg}$ proteinase K, 7.5 units achromopeptidase, $7.5\,\mu\text{g}/\mu\text{l}$ lysostaphin, 1050 units lysozyme, and 7.5 units mutanolysin to improve cell lysis.

Genome sequencing and assembly. The genome sequences of all of the *Micromonospora* strains, apart from the type strain of *M. chalcea*, were generated under the auspices of two "GEBA" projects, namely KMG-II, "From individual species to whole genera" and ACTINO 1000 "Exploiting the genomes of the *Actinobacteria*: plant growth promoters and producers of natural products and energy relevant enzymes united in a taxonomically unresolved phylum"; details on these projects are given in Supplementary Table 9. The genome of *M. chalcea* DSM 43026^T was sequenced, trimmed and assembled at Northumbria University using Illumina technology and protocols described by Sangal *et al.*²⁵. General aspects related to library construction and sequencing can be found at the JGI website (https://img.jgi.doe.gov/); the number of scaffolds and assembly methods are shown in Supplementary Tables 1 and 9.

Genome annotation. All of the genomes were annotated through the pipeline developed by the Joint Genome Institute (JGI) at the Department of Energy (DOE) using the Integrated Microbial Genomes Expert Review (IMG-ER) then compared with publicly available genomes of *M. aurantiaca* strains ATCC 27029^T and L5 and M. *lupini* Lupac 08. The JGI genome annotation pipeline, which includes Prodigal²²¹, was used followed by manual curation using GenePRIMP²²² for finished genomes and draft genomes. The predicted translation of proteins were analyzed using the National Center for Biotechnology Information (NCBI) non-redundant database, UniProt, TIGRFam, Pfam, PRIAM, KEGG, COG and InterPro databases. RNA gene identification was realised using the tRNAscanSE 21.23²²³ and HMMER 3.0rcl²²⁴ programs. Prediction of non-coding genes was determine under INFERNAL 1.0.2²²⁵. The Integrated Microbial Genomes – Expert Review (IMG-ER) platform²²⁶ permitted additional gene prediction analysis and functional annotation. CRT²²⁷ and PILER-CR²²⁸ allowed CRISPR element detection.

Genome analyses. The core genome was determined through the JGI tool Phylogenetic Profiler for Single Genes using the default options. CRISPRFinder²²⁹ was used to compare the analysis of CRISPR elements. antiS-MASH 3.0²³⁰ was used to determine and compare biosynthetic gene clusters. Presence of other genes was analyzed through the SEED viewer¹²⁸ after RAST annotation^{126,127} of the genomes.

Phylogenetic analyses. Genome-scale phylogenies were inferred from whole proteomes using the Genome BLAST Distance Phylogeny (GBDP) method, as previously described¹⁵⁹. Individual gene trees and multilocus sequence analysis (MLSA) trees were inferred using the phylogenies and gene similarities platform at the GGDC web server¹⁶⁶ available at https://ggdc.dsmz.de/ phylogeny-service.php through the DSMZ phylogenomics pipeline¹⁶³ adapted to single genes. Multiple sequence alignments were generated using MUSCLE software²³¹, maximum-likelihood (ML) and maximum-parsimony (MP) trees were inferred from the alignments with RAxML²³² and TNT²³³, respectively. For ML, rapid bootstrapping in conjunction with the autoMRE bootstrapping criterion²³⁴ was followed by a search for the best tree; for MP, 1000 bootstrapping replicates were used in conjunction with tree-bisection-and-reconnection branch swapping and ten random sequence replicates. The sequences were checked for compositional bias using the X² test, as implemented in PAUP*²³⁵. For the MLSA data set, the partition bootstrap (PB) was applied in addition to ordinary bootstrap²³⁶.

Phenotypic tests. All of the type strains were examined for a broad range of phenotypic properties known to be of value in *Micromonospora* systematics⁴⁴, namely catalase and oxidase production⁶²; degradation of organic compounds¹⁰⁹; carbon substrate utilization²³⁷; growth at different temperatures (4, 10, 20, 28, 37 and 45 °C), NaCl concentrations (1, 3, 5, 7 and 9%, w/v) and pH values (4.5, 5.5, 6.5, 8.0 and 9.0) using SA1 agar¹⁰⁹ as the basal medium; pH values were determined using appropriate buffers, as previously described⁹⁷. Chemotaxonomic properties of the strains were drawn from species descriptions, as indicated in Supplementary Tables 4 and 5.

Habitat classification. The strains were classified in groups according to the substrates from which they were isolated following the ENVO B classification (https://bioportal.bioontology.org/ontologies/ENVO).

Statistical analysis. A tip-permutation test in conjunction with the calculation of maximum-parsimony scores was carried out as previously described²³⁶ to evaluate the phylogenetic conservation of phenotypic and genotypic features of the *Micromonospora* strains with respect to the GBDP tree. To this end, the tips of the tree reduced to the *Micromonospora* clade were permuted 10,000 times (including the original arrangement) and maximum-parsimony scores of the selected characters and each of the trees determined with Tree analysis using New Technology (TNT). The number of times the score of a permuted tree was as low or lower than the score of the original tree yielded the p-value. More sophisticated tests are available for phylogenetic conservation, particularly for continuous characters, but our approach allowed an easy comparison of binary and continuous characters as TNT deals with both. Proportion and count data were brought to the same scale using opm²³⁸, which generates TNT input files.

A principal coordinate analysis of bootstrap-weighted relative Robinson-Foulds distances between trees was calculated with RAxML. The distances were visualized as principal coordinates, as implemented in R²³⁹. The Chi-2 test, as implemented in R, was used to determine correlations between binary phenotypic features of the *Micromonospora* strains.

Data availability. All data generated or analyzed during this study are included in this published article (or in the Supplementary Information files), the genomes are available at the JGI website and with the following accession codes at the NCBI database: M. aurantiaca DSM 45487: FMHX01000001-FMHX01000004; M. auratinigra DSM 44815^T: LT594323-LT594323; M. carbonacea DSM 43168^T: FMCT1000001-FMCT01000051; M. chaiyaphumensis DSM 45246^T: FMCS01000001-FMCS01000023; M. chalcea DSM 43026^T: MAGP000000000; M. chersina DSM 44151^T: FMIB01000001-FMIB01000002; M. chokoriensis DSM 45160^T: LT607409-LT607409; M. citrea DSM 43903^T: FMHZ01000001-FMHZ01000002; M. coriariae DSM 44875^T: LT607412-LT607412; M. coxensis DSM 45161^T: LT607753-LT607753; M. cremea DSM 45599^T: FSQT00000000; M. eburnea DSM 44814^T: FMHY01000001-FMHY01000002; M. echinaurantiaca DSM 43904^T: LT607750-LT607750; M. echinofusca DSM 43913^T: LT607733-LT607733; M. echinospora DSM 43816^T: LT607413-LT607413; M. haikouensis DSM 45626^T: FMCW01000001-FMCW01000097; M. halophytica DSM 43171^T: FMDN01000001-FMDN01000064; M. humi DSM 45647^T: FMDM01000001-FMDM01000037; M. inositola DSM 43819^T: LT607754-LT607754; M. inyonensis DSM 46123^T: FMHU01000001-FMHU01000004; M. krabiensis DSM 45344^T: LT598496-LT598496; M. marina DSM 45555^T: FMCV01000001-FMCV01000074; M. matsumotoense DSM 44100^T: FMCU01000001-FMCU01000057; M. mirobrigensis DSM 44830^T: FMCX01000001-FMCX01000022; M. narathiwatensis DSM 45248^T: LT594324-LT594324; M. nigra DSM 43818^T: FMHT01000001-FMHT01000003; M. pallida DSM 43817^T: FMHW01000001-FMHW01000004; M. peucetia DSM 43363^T: FMIC01000001-FMIC01000002; M. purpureochromogenes DSM 43821^T: LT607410-LT607410; M. rhizosphaerae DSM 45431^T: FMHV01000001-FMHV01000003; M. rifamycinica DSM 44983^T: LT607752-LT607752; M. saelicesensis DSM 44871^T: FMCR01000001-FMCR01000011; M. sediminicola DSM 45794^T: FLRH01000001-FLRH01000005; M. siamensis DSM 45097^T: LT607751-LT607751; M. tulbaghiae DSM 45142^T: FMCQ01000001-FMCQ01000019; M. viridifaciens DSM 43909^T: LT607411-LT607411; M. yangpuensis DSM 45577^T: FMIA01000001-FMIA01000002; *M. zamorensis* DSM 45600^T: LT607755-LT607755.

References for Supplementary Tables 4 and 5²⁴⁰⁻²⁴⁵.

References

- 1. Priest, F. & Goodfellow, M. (Eds) Applied Microbial Systematics. (Kluwer Academic Publishers, 2000).
- 2. Goodfellow, M., Sutcliffe, I. C. & Chun, J. (Eds) New Approaches to Prokaryotic Systematics. (Academic Press, 2014).
- 3. Goodfellow, M. & O'Donnell, A. G. (Eds) New Bacterial Systematics. (Academic Press, 1993).
- Goodfellow, M., Freeman, R. & Sisson, P. R. Curie-point pyrolysis mass spectrometry as a tool in clinical microbiology. Zentralbl Bakteriol 285, 133–156 (1997).
- Colwell, R. R. Polyphasic taxonomy of the genus Vibrio: numerical taxonomy of Vibrio cholerae, Vibrio parahaemolyticus, and related Vibrio species. J Bacteriol 104, 410–433 (1970).
- Canhos, V. P., Manfio, G. P. & Blaine, L. D. Software tools and databases for bacterial systematics and their dissemination via global networks. Antonie Van Leeuwenhoek 64, 205–229 (1993).
- 7. Vandamme, P. et al. Polyphasic taxonomy, a consensus approach to bacterial systematics. Microbiol Rev 60, 407-438 (1996).
- 8. Tindall, B. J., Rosselló-Móra, R., Busse, H. J., Ludwig, W. & Kämpfer, P. Notes on the characterization of prokaryote strains for taxonomic purposes. *Int J Syst Evol Microbiol* **60**, 249–266, https://doi.org/10.1099/ijs.0.016949-0 (2010).
- 9. Woese, C. R. Bacterial evolution. Microbiol Rev 51, 221–271 (1987).
- Yarza, P. et al. Uniting the classification of cultured and uncultured Bacteria and Archaea using 16S rRNA gene sequences. Nat Rev Microbiol 12, 635–645, https://doi.org/10.1038/nrmicro3330 (2014).
- Labeda, D. P. et al. Phylogenetic study of the species within the family Streptomycetaceae. Antonie Van Leeuwenhoek 101, 73–104, https://doi.org/10.1007/s10482-011-9656-0 (2012).
- 12. Fox, G. E., Wisotzkey, J. D. & Jurtshuk, P. Jr. How close is close: 16S rRNA sequence identity may not be sufficient to guarantee species identity. *Int J Syst Bacteriol* 42, 166–170, https://doi.org/10.1099/00207713-42-1-166 (1992).
- 13. Kim, H. et al. Differentiation of Mycobacterium species by analysis of the heat-shock protein 65 gene (hsp65). Int J Syst Evol Microbiol 55, 1649–1656, https://doi.org/10.1099/ijs.0.63553-0 (2005).
- 14. Kämpfer, P. Continuing importance of the "phenotype" in the genomic era. *Methods Microbiol* 41, 307–320, https://doi.org/10.1016/bs.mim.2014.07.005 (2014).
- 15. Stackebrandt, E. et al. Re-evaluation of the status of the genus Oerskovia, reclassification of Promicromonospora enterophila (Jager et al. 1983) as Oerskovia enterophila comb. nov. and description of Oerskovia jenensis sp. nov. and Oerskovia paurometabola sp. nov. Int J Syst Evol Microbiol 52, 1105–1111, https://doi.org/10.1099/00207713-52-4-1105 (2002).
- 16. Mignard, S. & Flandrois, J. P. A seven-gene, multilocus, genus-wide approach to the phylogeny of mycobacteria using supertrees. *Int J Syst Evol Microbiol* **58**, 1432–1441, https://doi.org/10.1099/ijs.0.65658-0 (2008).
- 17. Nouioui, I., Ghodhbane-Gtari, F., Beauchemin, N. J., Tisa, L. S. & Gtari, M. Phylogeny of members of the *Frankia* genus based on *gyrB*, *nifH* and *glnII* sequences. *Antonie van Leeuwenhoek* **100**, 579, https://doi.org/10.1007/s10482-011-9613-y (2011).
- 18. Labeda, D. P. et al. Phylogenetic relationships in the family Streptomycetaceae using multi-locus sequence analysis. Antonie Van Leeuwenhoek 110, 563–583, https://doi.org/10.1007/s10482-016-0824-0 (2017).
- Goodfellow, M. et al. (Eds) Bergey's Manual of Systematic Bacteriology, 2nd edn., Vol 5, The Actinobacteria, Part B. (Springer, 2012).
- 20. Klenk, H. P. & Göker, M. En route to a genome-based classification of Archaea and Bacteria? Syst Appl Microbiol 33, 175–182, https://doi.org/10.1016/j.syapm.2010.03.003 (2010).
- Sutcliffe, I. C., Trujillo, M. E. & Goodfellow, M. A call to arms for systematists: revitalising the purpose and practises underpinning the description of novel microbial taxa. Antonie Van Leeuwenhoek 101, 13–20, https://doi.org/10.1007/s10482-011-9664-0 (2012).
- 22. Whitman, W. B. The need for change: embracing the genome. Methods Microbiol 41, 1-12, https://doi.org/10.1016/bs.mim.2014.08.002 (2014).
- 23. Chun, J. & Rainey, F. A. Integrating genomics into the taxonomy and systematics of the *Bacteria* and *Archaea*. *Int J Syst Evol Microbiol* 64, 316–324, https://doi.org/10.1099/ijs.0.054171-0 (2014).
- 24. Stephens, W. Z. et al. Identification of population bottlenecks and colonization factors during assembly of bacterial communities within the Zebrafish intestine. MBio 6, e01163–01115, https://doi.org/10.1128/mBio.01163-15 (2015).
- Sangal, V. et al. Next-generation systematics: An innovative approach to resolve the structure of complex prokaryotic taxa. Sci Rep 6, 38392, https://doi.org/10.1038/srep38392 (2016).
- 26. Tang, B. et al. A systematic study of the whole genome sequence of Amycolatopsis methanolicastrain 239T provides an insight into its physiological and taxonomic properties which correlate with its position in the genus. Synth Syst Biotech 1, 169–186, https://doi.org/10.1016/j.synbio.2016.05.001 (2016).

- Girard, G. et al. A novel taxonomic marker that discriminates between morphologically complex actinomycetes. Open Biol 3, 130073, https://doi.org/10.1098/rsob.130073 (2013).
- Chandra, G. & Chater, K. F. Developmental biology of Streptomyces from the perspective of 100 actinobacterial genome sequences. FEMS Microbiol Rev 38, 345–379, https://doi.org/10.1111/1574-6976.12047 (2014).
- Kamneva, O. K. & Ward, N. L. Reconciliation approaches to determining HGT, duplications, and losses in gene trees. Methods Microbiol 41, 183–199, https://doi.org/10.1016/bs.mim.2014.08.004 (2014).
- 30. Doroghazi, J. R. & Metcalf, W. W. Comparative genomics of actinomycetes with a focus on natural product biosynthetic genes. BMC Genomics 14, 611, https://doi.org/10.1186/1471-2164-14-611 (2013).
- Creason, A. L., Davis, E. W., Putnam, M. L., Vandeputte, O. M. & Chang, J. H. Use of whole genome sequences to develop a
 molecular phylogenetic framework for *Rhodococcus fascians* and the *Rhodococcus* genus. *Front Plant Sci* 5, https://doi.org/10.3389/
 fpls.2014.00406 (2014).
- 32. Lopez-Modéjar, R., Kostovčík, M., Lladó, S., Carro, L. & García-Fraile, P. In *Probiotics in Agroecosystem* (eds V. Kumar, M. Kumar, S. Sharma, & R. Prasad) (Springer Nature, 2017).
- 33. Lapage, S. P. et al. (Eds) International Code of Nomenclature of Bacteria: Bacteriological Code, 1990 Revision. (ASM Press, 1992).
- 34. Stackebrandt, E. et al. Deposit of microbial strains in public service collections as part of the publication process to underpin good practice in science. Springer PLUS 3, 208, https://doi.org/10.1186/2193-1801-3-208 (2014).
- 35. Kyrpides, N. C. et al. Genomic encyclopedia of Bacteria and Archaea: sequencing a myriad of type strains. PLoS Biol 12, e1001920, https://doi.org/10.1371/journal.pbio.1001920 (2014).
- Kyrpides, N. C. et al. Genomic Encyclopedia of Type Strains, Phase I: The one thousand microbial genomes (KMG-I) project. Stand Genomic Sci 9, 1278–1284, https://doi.org/10.4056/sigs.5068949 (2014).
- 37. Whitman, W. B. et al. Genomic Encyclopedia of Bacterial and Archaeal Type Strains, Phase III: the genomes of soil and plant-associated and newly described type strains. Stand Genomic Sci 10, 26, https://doi.org/10.1186/s40793-015-0017-x (2015).
- 38. Wu, D. et al. A phylogeny-driven genomic encyclopaedia of Bacteria and Archaea. Nature 462, 1056–1060, https://doi.org/10.1038/nature08656 (2009).
- Mukherjee, S. et al. 1,003 reference genomes of bacterial and archaeal isolates expand coverage of the tree of life. Nat Biotech, https://doi.org/10.1038/nbt.3886 (2017).
- Sangal, V., Nieminen, L., Tucker, N. P. & Hoskisson, P. A. Revolutionizing prokaryotic systematics through next-generation sequencing. Methods Microbiol 41, 75–101, https://doi.org/10.1016/bs.mim.2014.07.001 (2014).
- 41. Qin, Q.-L. et al. A proposed genus boundary for the prokaryotes based on genomic insights. J Bacteriol 196, 2210–2215, https://doi.org/10.1128/jb.01688-14 (2014).
- doi.org/10.1128/jb.01688-14 (2014).
 42. Barka, E. A. *et al.* Taxonomy, physiology, and natural products of *Actinobacteria*. *Microbiol Mol Biol Rev* **80**, 1–43, https://doi.
- org/10.1128/MMBR.00019-15 (2016).
 43. Bull, A. T., Asenjo, J. A., Goodfellow, M. & Gomez-Silva, B. The Atacama Desert: Technical resources and the growing importance
- of novel microbial diversity. *Annu Rev Microbiol* **70**, 215–234, https://doi.org/10.1146/annurev-micro-102215-095236 (2016).

 44. Genilloud, O. In *Bergey's Manual of Systematic Bacteriology, 2nd edn., Vol 5, The Actinobacteria, Part B* (eds M. Goodfellow *et al.*) 1039–1057 (Springer, 2012).
- 45. Genilloud, O. In Bergey's Manual of Systematic Bacteriology, 2nd edn., Vol 5, The Actinobacteria, Part B (eds M. Goodfellow et al.)
- 1035–1038 (Springer, 2012).
 46. Genilloud, O. In Bergey's Manual of Systematic Bacteriology, 2nd edn., Vol 5, The Actinobacteria, Part B (eds M. Goodfellow et al.)
- 1035 (Springer, 2012).
 47. Mingma, R., Tanaka, K., Ōmura, S., Takahashi, Y. & Matsumoto, A. Actinorhabdospora filicis gen. nov., sp. nov., a new member of the family Micromonosporaceae. Int J Syst Evol Microbiol 66, 3071–3077, https://doi.org/10.1099/ijsem.0.001145 (2016).
- 48. Ørskov, J. Investigations into the Morphology of the Ray Fungi. (Levin and Munksgaard, 1923).
- 49. Foulerton, A. New species of *Streptothrix* isolated from the air. *Lancet* 1199–1200 (1905).
- Parte, A. C. LPSN—list of prokaryotic names with standing in nomenclature. Nucleic Acids Research 42, D613–D616, https://doi. org/10.1093/nar/gkt1111 (2014).
- 51. Carro, L., Riesco, R., Spröer, C. & Trujillo, M. E. *Micromonospora luteifusca* sp. nov. isolated from cultivated *Pisum sativum. Syst Appl Microbiol* 39, 237–242, https://doi.org/10.1016/j.syapm.2016.04.003 (2016).
- Betancur, L. A. et al. Marine actinobacteria as a source of compounds for phytopathogen control: An integrative metabolic-profiling/bioactivity and taxonomical approach. PLOS ONE 12, e0170148, https://doi.org/10.1371/journal.pone.0170148 (2017).
- Carro, L., Spröer, C., Alonso, P. & Trujillo, M. E. Diversity of Micromonospora strains isolated from nitrogen fixing nodules and rhizosphere of Pisum sativum analyzed by multilocus sequence analysis. Syst Appl Microbiol 35, 73–80, https://doi.org/10.1016/j. syapm.2011.11.003 (2012).
- 54. Fang, B. et al. Two new species of the genus Micromonospora: Micromonospora palomenae sp. nov. and Micromonospora harpali sp. nov. isolated from the insects. Antonie Van Leeuwenhoek 108, 141–150, https://doi.org/10.1007/s10482-015-0472-9 (2015).
- 55. Xiang, W. et al. Micromonospora polyrhachis sp. nov., an actinomycete isolated from edible Chinese black ant (*Polyrhachis vicina* Roger). Int J Syst Evol Microbiol **64**, 495–500, https://doi.org/10.1099/ijs.0.055863-0 (2014).
- 56. Zhang, L., Xi, L., Ruan, J. & Huang, Y. Micromonospora yangpuensis sp. nov., isolated from a sponge. Int J Syst Evol Microbiol 62, 272–278, https://doi.org/10.1099/ijs.0.029439-0 (2012).
- 57. Carro, L., Riesco, R., Spröer, C. & Trujillo, M. E. Three new species of *Micromonospora* isolated from *Pisum sativum* nodules: *Micromonospora ureilytica* sp. nov., *Micromonospora noduli* sp. nov., and *Micromonospora vinacea* sp. nov. *Int J Syst Evol Microbiol*. https://doi.org/10.1099/ijsem.0.001231 (2016).
- Garcia, L. C., Martinez-Molina, E. & Trujillo, M. E. Micromonospora pisi sp. nov., isolated from root nodules of Pisum sativum. Int J Syst Evol Microbiol 60, 331–337, https://doi.org/10.1099/ijs.0.012708-0 (2010).
- 59. Kirby, B. M. & Meyers, P. R. Micromonospora tulbaghiae sp. nov., isolated from the leaves of wild garlic, Tulbaghia violacea. Int J Syst Evol Microbiol 60, 1328–1333, https://doi.org/10.1099/ijs.0.013243-0 (2010).
- Kittiwongwattana, C. et al. Micromonospora oryzae sp. nov., isolated from roots of upland rice. Int J Syst Evol Microbiol https://doi. org/10.1099/ijsem.0.000500 (2015).
- 61. Li, L., Mao, Y. J., Xie, Q. Y., Deng, Z. & Hong, K. Micromonospora avicenniae sp. nov., isolated from a root of Avicennia marina. Antonie Van Leeuwenhoek 103, 1089–1096, https://doi.org/10.1007/s10482-013-9888-2 (2013).
- 62. Trujillo, M. E., Kroppenstedt, R. M., Schumann, P., Carro, L. & Martinez-Molina, E. Micromonospora coriariae sp. nov., isolated from root nodules of Coriaria myrtifolia. Int J Syst Evol Microbiol 56, 2381–2385, https://doi.org/10.1099/ijs.0.64449-0 (2006).
- 63. Kaewkla, O., Thamchaipinet, A. & Franco, C. M. M. Micromonospora terminaliae sp. nov., an endophytic actinobacterium isolated from the surface-sterilized stem of the medicinal plant Terminalia mucronata. Int J Syst Evol Microbiol 67, 225–230, https://doi.org/10.1099/ijsem.0.001600 (2017).
- 64. Zhao, S. et al. Micromonospora parathelypteridis sp. nov., an endophytic actinomycete with antifungal activity isolated from the root of Parathelypteris beddomei (Bak.) Ching. Int J Syst Evol Microbiol 67, 268–274, https://doi.org/10.1099/ijsem.0.001614 (2017).
- 65. Nimaichand, S. et al. Micromonospora kangleipakensis sp. nov., isolated from a sample of limestone quarry. Int J Syst Evol Microbiol 63, 4546–4551, https://doi.org/10.1099/ijs.0.052746-0 (2013).

- 66. Hirsch, P., Mevs, U., Kroppenstedt, R. M., Schumann, P. & Stackebrandt, E. Cryptoendolithic actinomycetes from antarctic sandstone rock samples: *Micromonospora endolithica* sp. nov. and two isolates related to *Micromonospora coerulea* Jensen 1932. *Syst Appl Microbiol* 27, 166–174, https://doi.org/10.1078/072320204322881781 (2004).
- Lin, Y. B. et al. Micromonospora nickelidurans sp. nov., isolated from soil from a nickel-mining site. Int J Syst Evol Microbiol 65, 4615–4620, https://doi.org/10.1099/ijsem.0.000623 (2015).
- 68. Kroppenstedt, R. M. et al. Eight new species of the genus Micromonospora, Micromonospora citrea sp. nov., Micromonospora echinaurantiaca sp. nov., Micromonospora echinofusca sp. nov. Micromonospora fulviviridis sp. nov., Micromonospora inyonensis sp. nov., Micromonospora peucetia sp. nov., Micromonospora sagamiensis sp. nov., and Micromonospora viridifaciens sp. nov. Syst Appl Microbiol 28, 328–339, https://doi.org/10.1016/j.syapm.2004.12.011 (2005).
- 69. Trujillo, M. E., Kroppenstedt, R. M., Fernandez-Molinero, C., Schumann, P. & Martinez-Molina, E. *Micromonospora lupini* sp. nov. and *Micromonospora saelicesensis* sp. nov., isolated from root nodules of *Lupinus angustifolius*. *Int J Syst Evol Microbiol* 57, 2799–2804, https://doi.org/10.1099/ijs.0.65192-0 (2007).
- Kasai, H., Tamura, T. & Harayama, S. Intrageneric relationships among Micromonospora species deduced from gyrB-based phylogeny and DNA relatedness. Int J Syst Evol Microbiol 50 Pt 1, 127–134, https://doi.org/10.1099/00207713-50-1-127 (2000).
- 71. Weinstein, M. J. et al. Gentamicin, a new antibiotic complex from. Micromonospora. J Med Chem 6, 463-464 (1963).
- 72. Okachi, R., Kawamoto, I., Takasawa, S., Yamamoto, M. & Sato, S. A new antibiotic XK-62-2 (Sagamicin). I. Isolation, physicochemical and antibacterial properties. *J Antibiot (Tokyo)* 27, 793–800 (1974).
- 73. Weinstein, M. J. et al. Antibiotic 6640, a new Micromonospora-produced aminoglycoside antibiotic. J Antibiot (Tokyo) 23, 551–554 (1970)
- Weinstein, M. J., Wagman, G. H., Marquez, J. A., Testa, R. T. & Waitz, J. A. Verdamicin, a new broad spectrum aminoglycoside antibiotic. Antimicrob Agents Chemother 7, 246–249 (1975).
- 75. Weinstein, M. J., Luedemann, G. M., Oden, E. M. & Wagman, G. H. Everninomicin, a new antibiotic complex from *Micromonospora carbonacea*. *Antimicrob Agents Chemother (Bethesda)* 10, 24–32 (1964).
- 76. Weinstein, M. J., Luedemann, G. M., Oden, E. M. & Wagman, G. H. Halomicin, a new *Micromonospora*-produced antibiotic. *Antimicrob Agents Chemother (Bethesda)* 7, 435–441 (1967).
- 77. Weinstein, M. J. et al. Preliminary studies on megalomicin, a new Micromonospora-produced macrolide antibiotic complex. Antimicrob Agents Chemother (Bethesda) 8, 260–261 (1968).
- 78. Satoi, S., Muto, N., Hayashi, M., Fujii, T. & Otani, M. Mycinamicins, new macrolide antibiotics. I. Taxonomy, production, isolation, characterization and properties. *J Antibiot (Tokyo)* 33, 364–376 (1980).
- 79. Lee, M. D. et al. Calicheamicins, a novel family of antitumor antibiotics. 3. Isolation, purification and characterization of calicheamicins beta 1Br, gamma 1Br, alpha 2I, alpha 3I, beta 1I, gamma 1I and delta 1I. J Antibiot (Tokyo) 42, 1070–1087 (1989).
- Igarashi, Y. et al. Lupinacidin C, an inhibitor of tumor cell invasion from Micromonospora lupini. J Nat Prod 74, 862–865, https://doi.org/10.1021/np100779t (2011).
- 81. Antal, N. *et al.* Retymicin, galtamycin B, saquayamycin Z and ribofuranosyllumichrome, novel secondary metabolites from *Micromonospora* sp. Tu 6368. I. Taxonomy, fermentation, isolation and biological activities. *J Antibiot (Tokyo)* 58, 95–102, https://doi.org/10.1038/ja.2005.12 (2005).
- 82. Das, S., Ward, L. R. & Burke, C. Prospects of using marine actinobacteria as probiotics in aquaculture. *Appl Microbiol Biotechnol* 81, 419–429, https://doi.org/10.1007/s00253-008-1731-8 (2008).
- 83. Hirsch, A. M. & Valdés, M. Micromonospora: An important microbe for biomedicine and potentially for biocontrol and biofuels. Soil Biol Biochem 42, 536–542, https://doi.org/10.1016/j.soilbio.2009.11.023 (2010).
- 84. Trujillo, M. E., Riesco, R., Benito, P. & Carro, L. Endophytic actinobacteria and the interaction of *Micromonospora* and nitrogen fixing plants. *Front Microbiol* 6, 1341, https://doi.org/10.3389/fmicb.2015.01341 (2015).
- 85. Talukdar, M. et al. Complete genome sequencing and comparative analyses of broad-spectrum antimicrobial-producing Micromonospora sp. HK10. Gene 594, 97–107, https://doi.org/10.1016/j.gene.2016.09.005 (2016).
- Alonso-Vega, P. et al. Genome sequence of Micromonospora lupini Lupac 08, isolated from root nodules of Lupinus angustifolius. J Bacteriol 194, 4135, https://doi.org/10.1128/JB.00628-12 (2012).
- 87. Hirsch, A. M. et al. Complete genome sequence of *Micromonospora* strain L5, a potential plant-growth-regulating actinomycete, originally isolated from *Casuarina equisetifolia* root nodules. *Genome Announc* 1, https://doi.org/10.1128/genomeA.00759-13 (2013).
- 88. Adamek, M., Spohn, M., Stegmann, E. & Ziemert, N. Mining bacterial genomes for secondary metabolite gene clusters. *Methods Mol Biol* 1520, 23–47, https://doi.org/10.1007/978-1-4939-6634-9_2 (2017).
- 89. Medema, M. H. & Fischbach, M. A. Computational approaches to natural product discovery. *Nat Chem Biol* 11, 639–648, https://doi.org/10.1038/nchembio.1884 (2015).
- 90. Ziemert, N., Alanjary, M. & Weber, T. The evolution of genome mining in microbes a review. *Nat Prod Rep* 33, 988–1005, https://doi.org/10.1039/c6np00025h (2016).
- Baltz, R. H. Renaissance in antibacterial discovery from actinomycetes. Curr Opin Pharmacol 8, 557–563, https://doi.org/10.1016/j.coph.2008.04.008 (2008).
- 92. Zhu, H., Sandiford, S. K. & van Wezel, G. P. Triggers and cues that activate antibiotic production by actinomycetes. *J Ind Microbiol Biotechnol* 41, 371–386, https://doi.org/10.1007/s10295-013-1309-z (2014).
- 93. Rutledge, P. J. & Challis, G. L. Discovery of microbial natural products by activation of silent biosynthetic gene clusters. *Nat Rev Micro* 13, 509–523, https://doi.org/10.1038/nrmicro3496 (2015).
- 94. Trujillo, M. E. et al. Genome features of the endophytic actinobacterium Micromonospora lupini strain Lupac 08: on the process of adaptation to an endophytic life style? PLoS One 9, e108522, https://doi.org/10.1371/journal.pone.0108522 (2014).
- 95. Tanasupawat, S., Jongrungruangchok, S. & Kudo, T. Micromonospora marina sp. nov., isolated from sea sand. Int J Syst Evol Microbiol 60, 648–652, https://doi.org/10.1099/ijs.0.014068-0 (2010).
- 96. Luedemann, G. M. & Brodsky, B. *Micromonospora carbonacea* sp. n., an everninomicin-producing organism. *Antimicrob Agents Chemother (Bethesda)* 10, 47–52 (1964).
- Carro, L., Pukall, R., Spröer, C., Kroppenstedt, R. M. & Trujillo, M. E. Micromonospora cremea sp. nov. and Micromonospora zamorensis sp. nov., isolated from the rhizosphere of Pisum sativum. Int J Syst Evol Microbiol 62, 2971–2977, https://doi. org/10.1099/ijs.0.038695-0 (2012).
- 98. Songsumanus, A., Tanasupawat, S., Thawai, C., Suwanborirux, K. & Kudo, T. Micromonospora humi sp. nov., isolated from peat swamp forest soil. Int J Syst Evol Microbiol 61, 1176–1181, https://doi.org/10.1099/ijs.0.024281-0 (2011).
- 99. Aziz, R. K., Breitbart, M. & Edwards, R. A. Transposases are the most abundant, most ubiquitous genes in nature. *Nucleic Acids Research* 38, 4207–4217, https://doi.org/10.1093/nar/gkq140 (2010).
- 100. Chang, Y. J. et al. Non-contiguous finished genome sequence and contextual data of the filamentous soil bacterium *Ktedonobacter racemifer* type strain (SOSP1-21). *Stand Genomic Sci* 5, 97–111, https://doi.org/10.4056/sigs.2114901 (2011).
- 101. Supong, K. et al. Micromonospora sediminicola sp. nov., isolated from marine sediment. Int J Syst Evol Microbiol 63, 570–575, https://doi.org/10.1099/ijs.0.041103-0 (2013).
- 102. Thawai, C., Tanasupawat, S., Itoh, T., Suwanborirux, K. & Kudo, T. *Micromonospora aurantionigra* sp. nov., isolated from a peat swamp forest in Thailand. *Actinomycetologica* 18, 8–14 (2004).

- Tomita, K., Hoshino, Y., Ohkusa, N., Tsuno, T. & Miyaki, T. Micromonospora chersina sp. nov. Actinomycetologica 6, 21–28, https://doi.org/10.3209/sai.6 21 (1992).
- 104. Jongrungruangchok, S., Tanasupawat, S. & Kudo, T. *Micromonospora chaiyaphumensis* sp. nov., isolated from Thai soils. *Int J Syst Evol Microbiol* **58**, 924–928, https://doi.org/10.1099/ijs.0.65594-0 (2008).
- 105. Thawai, C. et al. Micromonospora eburnea sp. nov., isolated from a Thai peat swamp forest. Int J Syst Evol Microbiol 55, 417–422, https://doi.org/10.1099/ijs.0.63217-0 (2005).
- Thawai, C., Tanasupawat, S., Suwanborirux, K., Itoh, T. & Kudo, T. Micromonospora narathiwatensis sp. nov., from Thai peat swamp forest soils. J Gen Appl Microbiol 53, 287–293 (2007).
- 107. Kawamoto, I., Okachi, R., Kato, H., Yamamoto, S. & Takahashi, I. The antibiotic XK-41 complex. I. Production, isolation and characterization. *J Antibiot (Tokyo)* 27, 492–501 (1974).
- 108. Wang, C. et al. Micromonospora rhizosphaerae sp. nov., isolated from mangrove rhizosphere soil. Int J Syst Evol Microbiol 61, 320–324, https://doi.org/10.1099/ijs.0.020461-0 (2011).
- Trujillo, M. E. et al. Micromonospora mirobrigensis sp. nov. Int J Syst Evol Microbiol 55, 877–880, https://doi.org/10.1099/ijs.0.63361-0 (2005).
- 110. Thawai, C., Tanasupawat, S., Itoh, T., Suwanborirux, K. & Kudo, T. *Micromonospora siamensis* sp. nov., isolated from Thai peat swamp forest. *J Gen Appl Microbiol* **51**, 229–234 (2005).
- 111. Ara, I. & Kudo, T. Two new species of the genus Micromonospora: Micromonospora chokoriensis sp. nov. and Micromonospora coxensis sp. nov., isolated from sandy soil. J Gen Appl Microbiol 53, 29–37 (2007).
- 112. Xie, Q. Y., Qu, Z., Lin, H. P., Li, L. & Hong, K. Micromonospora haikouensis sp. nov., isolated from mangrove soil. Antonie Van Leeuwenhoek 101, 649–655, https://doi.org/10.1007/s10482-011-9682-y (2012).
- 113. Kawamoto, I., Yamamoto, M. & Nara, T. Micromonosproa olivasterospora sp. nov. Int J Syst Bacteriol 33, 107-112 (1983).
- 114. Jongrungruangchok, S., Tanasupawat, S. & Kudo, T. *Micromonospora krabiensis* sp. nov., isolated from marine soil in Thailand. *J Gen Appl Microbiol* 54, 127–133 (2008).
- 115. Wayne, L. G. et al. Report of the ad hoc committee on reconciliation of approaches to bacterial systematics. Int J Syst Evol Microbiol 37, 463–464, https://doi.org/10.1099/00207713-37-4-463 (1987).
- 116. Béjà, O. et al. Bacterial rhodopsin: evidence for a new type of phototrophy in the sea. Science 289, 1902–1906, https://doi.org/10.1126/science.289.5486.1902 (2000).
- 117. Kanehisa, M., Sato, Y., Kawashima, M., Furumichi, M. & Tanabe, M. KEGG as a reference resource for gene and protein annotation. *Nucleic Acids Res* 44, D457–462, https://doi.org/10.1093/nar/gkv1070 (2016).
- 118. Charkowski, A. O. *et al.* The *Pseudomonas syringae* pv. tomato HrpW protein has domains similar to harpins and pectate lyases and can elicit the plant hypersensitive response and bind to pectate. *J Bacteriol* **180**, 5211–5217 (1998).
- Ouyang, J., Shao, X. & Li, J. Indole-3-glycerol phosphate, a branchpoint of indole-3-acetic acid biosynthesis from the tryptophan biosynthetic pathway in *Arabidopsis thaliana*. *Plant J* 24, 327–333 (2000).
- 120. Rudrappa, T. et al. The rhizobacterial elicitor acetoin induces systemic resistance in Arabidopsis thaliana. Commun. Integr Biol 3, 130–138 (2010).
- 121. Ryu, C. M. et al. Bacterial volatiles promote growth in Arabidopsis. Proc Natl Acad Sci USA 100, 4927-4932, https://doi.org/10.1073/pnas.0730845100 (2003).
- Carro, L. & Nouioui, I. Taxonomy and systematics of plant probiotic bacteria in the genomic era. AIMS Microbiology 3, 383–412, https://doi.org/10.3934/microbiol.2017.3.383 (2017).
- 123. Capstick, D. S., Willey, J. M., Buttner, M. J. & Elliot, M. A. SapB and the chaplins: connections between morphogenetic proteins in Streptomyces coelicolor. Mol Microbiol 64, 602–613, https://doi.org/10.1111/j.1365-2958.2007.05674.x (2007).
- 124. Molle, V., Palframan, W. J., Findlay, K. C. & Buttner, M. J. WhiD and WhiB, homologous proteins required for different stages of sporulation in Streptomyces coelicolor A3(2). J Bacteriol 182, 1286–1295, https://doi.org/10.1128/jb.182.5.1286-1295.2000 (2000).
- 125. Świercz, J. P. & Elliot, M. A. In *Bacterial Spores: Current Research and Applications* (ed Ernesto Abel-Santos) 39–55 (Caister Academis Press, 2012).
- 126. Aziz, R. K. et al. The RAST Server: Rapid Annotations using Subsystems Technology. BMC Genomics 9, 75, https://doi.org/10.1186/1471-2164-9-75 (2008).
- 127. Brettin, T. et al. RASTtk: A modular and extensible implementation of the RAST algorithm for building custom annotation pipelines and annotating batches of genomes. Sci Rep 5, 8365, https://doi.org/10.1038/srep08365 (2015).
- 128. Overbeek, R. et al. The SEED and the rapid annotation of microbial genomes using Subsystems Technology (RAST). Nucleic Acids Research 42, D206–D214, https://doi.org/10.1093/nar/gkt1226 (2014).
- Etchegaray, J. P., Jones, P. G. & Inouye, M. Differential thermoregulation of two highly homologous cold-shock genes, cspA and cspB, of Escherichia coli. Genes Cells 1, 171–178 (1996).
- 130. Li, J. S., Bi, Y. T., Dong, C., Yang, J. F. & Liang, W. D. Transcriptome analysis of adaptive heat shock response of Streptococcus thermophilus. PLoS One 6, e25777, https://doi.org/10.1371/journal.pone.0025777 (2011).
- 131. Verbenko, V. N., Kuznetsova, L. V., Luchkina, L. A. & Klonov, N. V. Mutation in the cspH-cspG gene cluster enhances expression of heat-shock proteins and SOS repair system of *Escherichia coli. Genetika* **45**, 1194–1202 (2009).
- 132. Boncompagni, E., Osteras, M., Poggi, M. C. & le Rudulier, D. Occurrence of choline and glycine betaine uptake and metabolism in the family *Rhizobiaceae* and their roles in osmoprotection. *Appl Environ Microbiol* 65, 2072–2077 (1999).
- 133. Nau-Wagner, G. et al. Genetic control of osmoadaptive glycine betaine synthesis in *Bacillus subtilis* through the choline-sensing and glycine betaine-responsive GbsR repressor. *J Bacteriol* 194, 2703–2714, https://doi.org/10.1128/JB.06642-11 (2012).
- 134. Lee, H. N., Lee, N. O., Han, S. J., Ko, I. J. & Oh, J. I. Regulation of the *ahpC* gene encoding alkyl hydroperoxide reductase in *Mycobacterium smegmatis. PLoS One* 9, e111680, https://doi.org/10.1371/journal.pone.0111680 (2014).
- 135. Wietzke, M. & Bahl, H. The redox-sensing protein Rex, a transcriptional regulator of solventogenesis in *Clostridium acetobutylicum*. Appl Microbiol Biotechnol **96**, 749–761, https://doi.org/10.1007/s00253-012-4112-2 (2012).
- 136. Seo, S. W., Kim, D., Szubin, R. & Palsson, B. O. Genome-wide reconstruction of OxyR and SoxRS transcriptional regulatory networks under oxidative stress in *Escherichia coli* K-12 MG1655. *Cell Reports* 12, 1289–1299, https://doi.org/10.1016/j.celrep.2015.07.043 (2015).
- 137. Delany, I., Spohn, G., Rappuoli, R. & Scarlato, V. The Fur repressor controls transcription of iron-activated and -repressed genes in *Helicobacter pylori. Mol Microbiol* 42, 1297–1309 (2001).
- 138. Kim, S. O., Orii, Y., Lloyd, D., Hughes, M. N. & Poole, R. K. Anoxic function for the *Escherichia coli* flavohaemoglobin (Hmp): reversible binding of nitric oxide and reduction to nitrous oxide. *FEBS Letters* 445, 389–394, https://doi.org/10.1016/S0014-5793(99)00157-X (1999).
- 139. Schultz, J. E. & Matin, A. Molecular and functional characterization of a carbon starvation gene of *Escherichia coli. J Mol Biol* 218, 129–140, https://doi.org/10.1016/0022-2836(91)90879-B (1991).
- 140. Lucchetti-Miganeh, C., Burrowes, E., Baysse, C. & Ermel, G. The post-transcriptional regulator CsrA plays a central role in the adaptation of bacterial pathogens to different stages of infection in animal hosts. *Microbiology* **154**, 16–29, https://doi.org/10.1099/mic.0.2007/012286-0 (2008).
- 141. Rasmussen, J. J. et al. Campylobacter jejuni carbon starvation protein A (CstA) is involved in peptide utilization, motility and agglutination, and has a role in stimulation of dendritic cells. J Med Microbiol 62, 1135–1143, https://doi.org/10.1099/jmm.0.059345-0 (2013).

- Gerber, N. N. Volatile substances from actinomycetes: their role in the odor pollution of water. CRC Crit Rev Microbiol 7, 191–214 (1979).
- 143. Isogai, A. *et al.* The structure of leucanicidin, a novel insecticidal macrolide produced by *Streptomyces halstedii*. *Agric Biol Chem* **48**, 1379–1381, https://doi.org/10.1271/bbb1961.48.1379 (1984).
- 144. Cruz Morales, P., Barona Gómez, F., Ramos Aboites, J. E. & Carreño Sánchez, L. A. Genetic system for producing a proteases inhibitor of a small peptide aldehyde type. México patent (2015).
- 145. Skellam, E. J., Stewart, A. K., Strangman, W. K. & Wright, J. L. Identification of micromonolactam, a new polyene macrocyclic lactam from two marine *Micromonospora* strains using chemical and molecular methods: clarification of the biosynthetic pathway from a glutamate starter unit. *J Antibiot (Tokyo)* 66, 431–441, https://doi.org/10.1038/ja.2013.34 (2013).
- 146. Vertesy, L. et al. Feglymycin, a novel inhibitor of the replication of the human immunodeficiency virus. Fermentation, isolation and structure elucidation. J Antibiot (Tokyo) 52, 374–382 (1999).
- 147. Ferir, G. et al. Feglymycin, a unique natural bacterial antibiotic peptide, inhibits HIV entry by targeting the viral envelope protein gp120. Virology 433, 308–319, https://doi.org/10.1016/j.virol.2012.08.007 (2012).
- 148. Shaaban, K. A. et al. Herbimycins D-F, ansamycin analogues from Streptomyces sp. RM-7-15. J Nat Prod 76, 1619–1626, https://doi.org/10.1021/np400308w (2013).
- 149. Banskota, A. H. et al. TLN-05220, TLN-05223, new echinosporamicin-type antibiotics, and proposed revision of the structure of bravomicins[ast]. J Antibiot 62, 565–570 (2009).
- 150. Kunze, B., Jansen, R., Hofle, G. & Reichenbach, H. Crocacin, a new electron transport inhibitor from *Chondromyces crocatus* (myxobacteria). Production, isolation, physico-chemical and biological properties. *J Antibiot (Tokyo)* 47, 881–886 (1994).
- 151. McDonald, L. A. et al. Structures of the muraymycins, novel peptidoglycan biosynthesis inhibitors. J Am Chem Soc 124, 10260-10261, https://doi.org/10.1021/ja017748h (2002).
- 152. Huang, S. X. et al. Leinamycin E1 acting as an anticancer prodrug activated by reactive oxygen species. Proc Natl Acad Sci USA 112, 8278–8283, https://doi.org/10.1073/pnas.1506761112 (2015).
- 153. Li, W. et al. Nocathiacins, new thiazolyl peptide antibiotics from Nocardia sp. I. Taxonomy, fermentation and biological activities. J Antibiot (Tokyo) 56, 226–231 (2003).
- 154. Sharma, I., Sullivan, M. & McCutchan, T. F. *In vitro* antimalarial activity of novel semisynthetic nocathiacin I antibiotics. Antimicrob Agents Chemother 59, 3174–3179, https://doi.org/10.1128/AAC.04294-14 (2015).
- 155. Konishi, M. *et al.* Dynemicin A, a novel antibiotic with the anthraquinone and 1,5-diyn-3-ene subunit. *J Antibiot (Tokyo)* **42**, 1449–1452 (1989).
- 156. Xie, Z. et al. Kiamycin, a unique cytotoxic angucyclinone derivative from a marine Streptomyces sp. Mar Drugs 10, 551–558, https://doi.org/10.3390/md10030551 (2012).
- 157. McAlpine, J. B. et al. Biosynthesis of diazepinomicin/ECO-4601, a Micromonospora secondary metabolite with a novel ring system. J Nat Prod 71, 1585–1590, https://doi.org/10.1021/np800376n (2008).
- 158. Floss, H. G., Yu, T. W. & Arakawa, K. The biosynthesis of 3-amino-5-hydroxybenzoic acid (AHBA), the precursor of mC7N units in ansamycin and mitomycin antibiotics: a review. J Antibiot (Tokyo) 64, 35–44, https://doi.org/10.1038/ja.2010.139 (2011).
- 159. Hahnke, R. L. et al. Genome-based taxonomic classification of *Bacteroidetes*. Front Microbiol 7, 2003, https://doi.org/10.3389/fmicb.2016.02003 (2016).
- 160. Rosselló-Móra, R. & Amann, R. The species concept for prokaryotes. FEMS Microbiol Rev 25, 39-67 (2001).
- 161. Mesbah, N. M., Whitman, W. B. & Mesbah, M. In *Taxonomy of Prokaryotes* (eds F. A. Rainey & Aharon Oren) 299–324 (Academic Press, 2011).
- 162. Mesbah, N. M., Whitman, W. B. & Mesbah, M. K. Determination of the G+C content of Prokaryotes. Methods Microbiol 38, 299-324 (2011).
- 163. Meier-Kolthoff, J. P., Klenk, H. P. & Göker, M. Taxonomic use of DNA G+C content and DNA-DNA hybridization in the genomic age. *Int J Syst Evol Microbiol* **64**, 352–356, https://doi.org/10.1099/ijs.0.056994-0 (2014).
- 164. Riedel, T. et al. Genome sequence of the pink to light reddish-pigmented Rubellimicrobium mesophilum type strain (DSM 19309(T)), a representative of the Roseobacter group isolated from soil, and emended description of the species. Stand Genomic Sci 9, 902–913, https://doi.org/10.4056/sigs.5621012 (2014).
- 165. Ramasamy, D. et al. A polyphasic strategy incorporating genomic data for the taxonomic description of novel bacterial species. Int J Syst Evol Microbiol 64, 384–391, https://doi.org/10.1099/ijs.0.057091-0 (2014).
- 166. Meier-Kolthoff, J. P., Göker, M., Spröer, C. & Klenk, H. P. When should a DDH experiment be mandatory in microbial taxonomy? Arch Microbiol 195, 413–418, https://doi.org/10.1007/s00203-013-0888-4 (2013).
- 167. Rosselló-Móra, R., Urdiain, M. & Lopez-Lopez, A. DNA-DNA hybridrization. Methods Microbiol 38, 325-347 (2011).
- 168. Gevers, D. et al. Opinion: Re-evaluating prokaryotic species. Nat Rev Microbiol 3, 733–739, https://doi.org/10.1038/nrmicro1236
- 169. Maldonado, L. A. et al. Salinispora arenicola gen. nov., sp. nov. and Salinispora tropica sp. nov., obligate marine actinomycetes belonging to the family Micromonosporaceae. Int J Syst Evol Microbiol 55, 1759–1766, https://doi.org/10.1099/ijs.0.63625-0 (2005).
- 170. Ahmed, L. et al. Salinispora pacifica sp. nov., an actinomycete from marine sediments. Antonie Van Leeuwenhoek 103, 1069–1078, https://doi.org/10.1007/s10482-013-9886-4 (2013).
- 171. Ludwig, W. et al. In Bergey's Manual of Systematic Bacteriology, 2nd edn., Vol 5, The Actinobacteria, Part B (eds M. Goodfellow et al.) 1–28 (Springer, 2012).
- 172. Lechevalier, M. P., De Biévre, C. & Lechevalier, H. Chemotaxonomy of aerobic actinomycetes: phospholipid composition. *Biochem Syst Ecol* 5, 249–260, https://doi.org/10.1016/0305-1978(77)90021-7 (1977).
- 173. Coombs, J. T. & Franco, C. M. Isolation and identification of actinobacteria from surface-sterilized wheat roots. *Appl Environ Microbiol* **69**, 5603–5608 (2003).
- 174. Valdés, M. et al. Non-Frankia actinomycetes isolated from surface-sterilized roots of Casuarina equisetifolia fix nitrogen. Appl Environ Microbiol 71, 460-466, https://doi.org/10.1128/aem.71.1.460-466.2005 (2005).
- 175. Carro, L., Pujic, P., Trujillo, M. E. & Normand, P. *Micromonospora* is a normal occupant of actinorhizal nodules. *J Biosci* 38, 685–693 (2013).
- 176. Rosselló-Móra, R., Trujillo, M. E. & Sutcliffe, I. C. Introducing a Digital Protologue: A timely move towards a database-driven systematics of *Archaea* and *Bacteria*. *Syst Appl Microbiol* 40, 121–122, https://doi.org/10.1016/j.syapm.2017.02.001 (2017).
- 177. Trujillo, M. E. *et al.* The genus *Micromonospora* is widespread in legume root nodules: the example of *Lupinus angustifolius*. *ISME J* **4**, 1265–1281, https://doi.org/10.1038/ismej.2010.55 (2010).
- 178. Farag, M. A., Zhang, H. & Ryu, C.-M. Dynamic chemical communication between plants and bacteria through airborne signals: induced resistance by bacterial volatiles. *J Chem Ecol* 39, 1007–1018, https://doi.org/10.1007/s10886-013-0317-9 (2013).
- 179. Spaepen, S., Vanderleyden, J. & Remans, R. Indole-3-acetic acid in microbial and microorganism-plant signaling. FEMS Microbiol Rev 31, 425–448, https://doi.org/10.1111/j.1574-6976.2007.00072.x (2007).
- 180. Barraza, A. et al. Down-regulation of PvTRE1 enhances nodule biomass and bacteroid number in the common bean. New Phytologist 197, 194–206, https://doi.org/10.1111/nph.12002 (2013).
- 181. Aeschbacher, R. A., Müller, J., Boller, T. & Wiemken, A. Purification of the trehalase GMTRE1 from soybean nodules and cloning of its cDNA. GMTRE1 is expressed at a low level in multiple tissues. *Plant Physiol* 119, 489–496 (1999).
- 182. Waksman, S. A. On the classification of actinomycetes. J NJ Agri Exper Stat 549-558 (1939).

- Suarez, J. E. & Hardisson, C. Morphological characteristics of colony development in *Micromonospora chalcea*. J Bacteriol 162, 1342–1344 (1985).
- 184. Waksman, S. A. The Actinomycetes: a Summary of Current Knowledge. (The Ronald Press Company, 1960).
- 185. Waksman, S. A. In The Actinomycetes Classification, Identification and Description of Genera and Species 293–297 (Williams & Wilkins Co, 1961).
- 186. Maldonado, L. A. & Quintana, E. T. Unexpected properties of micromonoporae from marine origin. *Adv Microbiol* 5, 452–456 (2015)
- 187. Baldacci, E. & Locci, R. Osservazioni e ricerche su *Micromonospora melanosporea* comb. nov. e descrizione di una nuova sottospecie: *M. melanosporea* subsp. *corymbica. Ann Microbiol Enzymol* 11, 19–30 (1961).
- 188. Kodani, S. et al. The SapB morphogen is a lantibiotic-like peptide derived from the product of the developmental gene ramS in Streptomyces coelicolor. Proc Natl Acad Sci USA 101, 11448–11453, https://doi.org/10.1073/pnas.0404220101 (2004).
- 189. Rybniker, J. et al. Insights into the function of the WhiB-like protein of mycobacteriophage TM4-a transcriptional inhibitor of WhiB2. Mol Microbiol 77, 642-657, https://doi.org/10.1111/j.1365-2958.2010.07235.x (2010).
- 190. Richter, T. K., Hughes, C. C. & Moore, B. S. Sioxanthin, a novel glycosylated carotenoid, reveals an unusual subclustered biosynthetic pathway. *Environ Microbiol* 17, 2158–2171, https://doi.org/10.1111/1462-2920.12669 (2015).
- 191. Krugel, H., Krubasik, P., Weber, K., Saluz, H. P. & Sandmann, G. Functional analysis of genes from *Streptomyces griseus* involved in the synthesis of isorenieratene, a carotenoid with aromatic end groups, revealed a novel type of carotenoid desaturase. *Biochim Biophys Acta* 1439, 57–64 (1999).
- 192. Choi, S. -k. et al. Characterization of β -carotene ketolases, CrtW, from marine bacteria by complementation analysis in Escherichia coli. Marine Biotech 7, 515–522, https://doi.org/10.1007/s10126-004-5100-z (2005).
- 193. Liang, M. H., Zhu, J. & Jiang, J.-G. Carotenoids biosynthesis and cleavage related genes from bacteria to plants. *Crit Rev Food Sci Nutr*, 00–00, https://doi.org/10.1080/10408398.2017.1322552 (2017).
- 194. Mohammadi, M., Burbank, L. & Roper, M. C. Biological role of pigment production for the bacterial phytopathogen Pantoea stewartii subsp. stewartii. Appl Environ Microbiol 78, 6859–6865, https://doi.org/10.1128/AEM.01574-12 (2012).
- 195. Kuhlbrandt, W. Structure and function of bacterial light-harvesting complexes. Structure 3, 521-525 (1995).
- Goosen, N. & Moolenaar, G. F. Repair of UV damage in bacteria. DNA Repair (Amst) 7, 353–379, https://doi.org/10.1016/j. dnarep.2007.09.002 (2008).
- 197. Mackenzie, C., Chidambaram, M., Sodergren, E. J., Kaplan, S. & Weinstock, G. M. DNA repair mutants of *Rhodobacter sphaeroides*. J Bacteriol 177, 3027–3035 (1995).
- 198. Reina-Bueno, M. et al. Role of trehalose in salinity and temperature tolerance in the model halophilic bacterium *Chromohalobacter salexigens*. PLOS One 7, e33587, https://doi.org/10.1371/journal.pone.0033587 (2012).
- Wagman, G. H. & Weinstein, M. J. Antibiotic from Micromonospora. Annu Rev Microbiol 34, 537–557, https://doi.org/10.1146/annurev.mi.34.100180.002541 (1980).
- 200. Bérdy, J. Bioactive microbial metabolites. J Antibiot (Tokyo) 58, 1-26, https://doi.org/10.1038/ja.2005.1 (2005).
- Horan, A. C. In Bioprocess Technology: Fermentation, Biocatalysis and Bioseparation. (eds Flickinger & Drew) 2333–2348 (Wiley, 1999).
- 202. Tiwari, K. & Gupta, R. K. Rare actinomycetes: a potential storehouse for novel antibiotics. *Crit Rev Biotechnol* 32, 108–132, https://doi.org/10.3109/07388551.2011.562482 (2012).
- Katz, L. & Baltz, R. H. Natural product discovery: past, present, and future. J Ind Microbiol Biotechnol 43, 155–176, https://doi. org/10.1007/s10295-015-1723-5 (2016).
- 204. Goodfellow, M. & Fiedler, H. P. A guide to successful bioprospecting: informed by actinobacterial systematics. *Antonie Van Leeuwenhoek* 98, 119–142, https://doi.org/10.1007/s10482-010-9460-2 (2010).
- 205. Wichner, D. et al. Isolation and anti-HIV-1 integrase activity of lentzeosides A-F from extremotolerant Lentzea sp. H45, a strain isolated from a high-altitude Atacama Desert soil. J Antibiot 70, 448–453, https://doi.org/10.1038/ja.2016.78 (2017).
- Baltz, R. H. Gifted microbes for genome mining and natural product discovery. J Ind Microbiol Biotechnol 44, 573–588, https://doi. org/10.1007/s10295-016-1815-x (2017).
- 207. Doroghazi, J. R. *et al.* A roadmap for natural product discovery based on large-scale genomics and metabolomics. *Nat Chem Biol* **10**, 963–968, https://doi.org/10.1038/nchembio.1659 (2014).
- Jensen, P. R. Linking species concepts to natural product discovery in the post-genomic era. J Ind Microbiol Biotechnol 37, 219–224, https://doi.org/10.1007/s10295-009-0683-z (2010).
- Millan-Aguinaga, N. et al. Phylogenomic insight into Salinispora (Bacteria, Actinobacteria) species designations. Sci Rep 7, 3564, https://doi.org/10.1038/s41598-017-02845-3 (2017).
- Jensen, P. R. Natural products and the gene cluster revolution. Trends Microbiol 24, 968-977, https://doi.org/10.1016/j.tim.2016.07.006 (2016).
- 211. Jensen, P. R., Williams, P. G., Oh, D. C., Zeigler, L. & Fenical, W. Species-specific secondary metabolite production in marine actinomycetes of the genus Salinispora. Appl Environ Microbiol 73, 1146–1152, https://doi.org/10.1128/AEM.01891-06 (2007).
- 212. Sveshnikova, M. A., Maksimova, T. S. & Kudrina, E. S. The species belonging to the genus *Micromonospora Ørskov*, 1923 and their taxonomy. *Mikrobiologiya* **38** (1969).
- 213. Lee, S. D., Goodfellow, M. & Hah, Y. C. A phylogenetic analysis of the genus *Catellatospora* based on 16S ribosomal DNA sequences, including transfer of *Catellatospora matsumotoense* to the genus *Micromonospora* as *Micromonospora matsumotoense* comb. nov. *FEMS Microbiol Lett* **178**, 349–354 (1999).
- 214. Luedemann, G. M. Micromonospora purpureochromogenes (Waksman and Curtis 1916) comb. nov. (subjective synonym: Micromonospora fusca Jensen 1932). Int J Syst Bacteriol 21, 240-247 (1971).
- Huang, H. et al. Micromonospora rifamycinica sp. nov., a novel actinomycete from mangrove sediment. Int J Syst Evol Microbiol 58, 17–20, https://doi.org/10.1099/ijs.0.64484-0 (2008).
- 216. Waite, D. W. et al. Comparative genomic analysis of the class Epsilonproteobacteria and proposed reclassification to Epsilonbacteraeota (phyl. nov.). Front Microbiol 8, 682, https://doi.org/10.3389/fmicb.2017.00682 (2017).
- 217. Scheuner, C. et al. Complete genome sequence of Planctomyces brasiliensis type strain (DSM 5305(T)), phylogenomic analysis and reclassification of Planctomycetes including the descriptions of Gimesia gen. nov., Planctopirus gen. nov. and Rubinisphaera gen. nov. and emended descriptions of the order Planctomycetales and the family. Planctomycetaceae. Stand Genomic Sci 9, 10, https://doi.org/10.1186/1944-3277-9-10 (2014).
- 218. Vinatzer, B. A. *et al.* A proposal for a genome similarity-based taxonomy for plant-pathogenic bacteria that is sufficiently precise to reflect phylogeny, host range, and outbreak affiliation applied to *Pseudomonas syringae* sensu lato as a proof of concept. *Phytopathology* **107**, 18–28, https://doi.org/10.1094/PHYTO-07-16-0252-R (2017).
- Jeffrey, B., Rose, S. J., Gilbert, K., Lewis, M. & Bermudez, L. E. Comparative analysis of the genomes of clinical isolates of *Mycobacterium avium* subsp. hominissuis regarding virulence-related genes. J Med Microbiol, https://doi.org/10.1099/jmm.0.000507 (2017).
- Stover, C. K. et al. Complete genome sequence of Pseudomonas aeruginosa PAO1, an opportunistic pathogen. Nature 406, 959–964, https://doi.org/10.1038/35023079 (2000).
- 221. Hyatt, D. et al. Prodigal: prokaryotic gene recognition and translation initiation site identification. BMC Bioinformatics 11, 119, https://doi.org/10.1186/1471-2105-11-119 (2010).

- Pati, A. et al. GenePRIMP: a gene prediction improvement pipeline for prokaryotic genomes. Nat Methods 7, 455–457, https://doi. org/10.1038/nmeth.1457 (2010).
- 223. Lowe, T. M. & Eddy, S. R. tRNAscan-SE: a program for improved detection of transfer RNA genes in genomic sequence. *Nucleic Acids Res* 25, 955–964 (1997).
- 224. Finn, R. D., Clements, J. & Eddy, S. R. HMMER web server: interactive sequence similarity searching. *Nucleic Acids Res* 39, W29–37, https://doi.org/10.1093/nar/gkr367 (2011).
- Nawrocki, E. P., Kolbe, D. L. & Eddy, S. R. Infernal 1.0: inference of RNA alignments. *Bioinformatics* 25, 1335–1337, https://doi. org/10.1093/bioinformatics/btp157 (2009).
- 226. Markowitz, V. M. et al. IMG ER: a system for microbial genome annotation expert review and curation. Bioinformatics 25, 2271–2278, https://doi.org/10.1093/bioinformatics/btp393 (2009).
- 227. Bland, C. et al. CRISPR recognition tool (CRT): a tool for automatic detection of clustered regularly interspaced palindromic repeats. BMC Bioinformatics 8, 209, https://doi.org/10.1186/1471-2105-8-209 (2007).
- 228. Petersen, J. Phylogeny and compatibility: plasmid classification in the genomics era. Arch Microbiol 193, 313–321, https://doi.org/10.1007/s00203-011-0686-9 (2011).
- 229. Grissa, I., Vergnaud, G. & Pourcel, C. CRISPRFinder: a web tool to identify clustered regularly interspaced short palindromic repeats. *Nucleic Acids Res* **35**, W52–57, https://doi.org/10.1093/nar/gkm360 (2007).
- 230. Weber, T. *et al.* antiSMASH 3.0-a comprehensive resource for the genome mining of biosynthetic gene clusters. *Nucleic Acids Res* **43**, W237–243, https://doi.org/10.1093/nar/gkv437 (2015).
- 231. Edgar, R. C. MUSCLE: multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Res* **32**, 1792–1797, https://doi.org/10.1093/nar/gkh340 (2004).
- 232. Stamatakis, Å. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics* 30, 1312–1313, https://doi.org/10.1093/bioinformatics/btu033 (2014).
- Goloboff, P. A., Farris, J. S. & Nixon, K. C. TNT, a free program for phylogenetic analysis. *Cladistics* 24, 774–786, https://doi. org/10.1111/j.1096-0031.2008.00217.x (2008).
- 234. Pattengale, N. D., Alipour, M., Bininda-Emonds, O. R., Moret, B. M. & Stamatakis, A. How many bootstrap replicates are necessary? *J Comput Biol* 17, 337–354, https://doi.org/10.1089/cmb.2009.0179 (2010).
- 235. Yang, Z. Phylogenetic analysis using parsimony and likelihood methods. J Mol Evol 42, 294–307, https://doi.org/10.1007/bf02198856 (1996).
- 236. Simon, M. *et al.* Phylogenomics of Rhodobacteraceae reveals evolutionary adaptation to marine and non-marine habitats. *ISME J* 11, 1483–1499, https://doi.org/10.1038/ismej.2016.198 (2017).
- Williams, S. T. et al. Numerical classification of Streptomyces and related genera. J Gen Microbiol 129, 1743–1813, https://doi. org/10.1099/00221287-129-6-1743 (1983).
- 238. Vaas, L. A. et al. opm: an R package for analysing OmniLog(R) phenotype microarray data. Bioinformatics 29, 1823–1824, https://doi.org/10.1093/bioinformatics/btt291 (2013).
- 239. Team, R. C. R A language and environment for statistical computing. *R Foundation for statistical computing* Viena, Austria, https://www.R-project.org/ (2016).
- 240. Vela Gurovic, M. S. et al. Micromonospora schwarzwaldensis sp. nov., a producer of telomycin, isolated from soil. Int J Syst Evol Microbiol 63, 3812–3817, https://doi.org/10.1099/ijs.0.051623-0 (2013).
- Carro, L., Pukall, R., Sproer, C., Kroppenstedt, R. M. & Trujillo, M. E. Micromonospora halotolerans sp. nov., isolated from the rhizosphere of a Pisum sativum plant. Antonie Van Leeuwenhoek 103, 1245–1254, https://doi.org/10.1007/s10482-013-9903-7
- 242. Everest, G. J. & Meyers, P. R. *Micromonospora equina* sp. nov., isolated from soil from a racecourse. *Int J Syst Evol Microbiol* 63, 879–885, https://doi.org/10.1099/ijs.0.042929-0 (2013).
- 243. Phongsopitanun, W. et al. Micromonospora fluostatini sp. nov., isolated from marine sediment. Int J Syst Evol Microbiol 65, 4417–4423, https://doi.org/10.1099/ijsem.0.000589 (2015).
- 244. Ren, J. et al. Micromonospora wenchangensis sp. nov., isolated from mangrove soil. Int J Syst Evol Microbiol 63, 2389–2395, https://doi.org/10.1099/ijs.0.045476-0 (2013).
- 245. Thawai, C., Tanasupawat, S. & Kudo, T. Micromonospora pattaloongensis sp. nov., isolated from a Thai mangrove forest. Int J Syst Evol Microbiol 58, 1516–1521, https://doi.org/10.1099/ijs.0.65410-0 (2008).

Acknowledgements

This work was performed under the auspices of the US Department of Energy's Office of Science, Biological and Environmental Research Program and the University of California, Lawrence Berkeley National Laboratory under contract No. DE-AC02-05CH11231. LC and IN were supported by postdoctoral fellowships awarded by Newcastle University. MET acknowledges financial support from the Spanish *Ministerio de Economía y Competitividad* (MICINN) under grant No. CGL2014-52735-P. The authors are indebted to Marlen Jando and Gabriele Pötter for growing cells for genome DNA extraction and to Evelyne Brambilla for extracting DNA for genome determination and quality control (all at DSMZ).

Author Contributions

H.-P.K., L.C. and Ma.G. designed the project; D.S., K.S., Ni.S., N.C.K., P.P., S.D., T.W. and V.S., were responsible for genome sequencing, assembly and annotation; J.P.M.K. and Ma.G. for the comparative genomic based phylogeny and statistical analysis; L.C. for single genes and MLSA. based phylogenies and phenotypic tests. L.C., Mi.G., M.E.T. and Ne.S. were responsible for taxonomic input; I.N. and L.C. for the analysis of the genomes. L.C. and Mi.G. drafted the manuscript which was critically reviewed by H.P.K., I.N., Ma. G., M.C.M.C., M.E.T., and V.S. All of the authors approved the final manuscript.

Additional Information

Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-017-17392-0.

Competing Interests: The authors declare that they have no competing interests.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit https://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2017