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New-Data Letter: Bacteriology	1
Rapid genotyping of Shigella sonnei using multiplex high resolution melting	2
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Running title: Multiplex-HRM typing of Shigella sonnei	4
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nucleotide polymorphism (SNP)	19

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Shigella sonnei, an emerging global cause of shigellosis, consists of four distinct lineages and the 21 current pandemic involves several geographically associated, multidrug- resistant clones that 22 belong to lineage III (1-3). A typing scheme based on high resolution melting (HRM) of six 23 chromosomal single nucleotide polymorphisms (SNPs) has been described to identify all 24 lineages/sub-lineages (4). HRM detects SNPs in small DNA fragments based on the melting 25 patterns (5, 6). HRM involves minimal labor with rapid turnaround time, and a decreased risk of 26 PCR carryover contamination (7). A sensitive intercalating fluorescent dye, EvaGreen, may yield 27 robust fluorescence intensities and strong sharp melting peaks on melting of the PCR amplicons 28 (8-10). Based on the SNPs summarized in Table 1, we have developed a multiplex-HRM, for 29 identification of lineages I, II, and III using the EvaGreen and a Rotor-Gene 6000 instrument 30 (Qiagen). 31 Ten strains (2 for each lineage/sublineage) were from a collection held at Institut Pasteur. Only 32 one lineage IV strain is reported so far (1), which is not included in the study. Primers were 33 designed to amplify fragments that contain three SNPs within kduD, deoA and emrA for typing 3 34 main lineages in a first set run, and to amplify fragments that contain SNPs within fdX and menF 35 for typing sub-lineages IIIa, IIIb, and IIIc in a second run. Primer sequences are summarized in 36 Table 1. 37 In the first set run, a 20 µl multiplex real-time PCR reaction was set up, containing 10 µl 38 SensiMixTM with EvaGreen Dye (Bioline), 20 ng of template DNA, and 2.5 μl of each primer of 39 kduD (100μM), 1 μl of each primer of deoA (100μM) and 1 μl of each primer of emrA (50μM). 40 In the second set run, the 20 μ l reaction mix was identical, except that the primers used were fdX 41 $(10 \mu M)$ and menF $(10 \mu M)$, with a quantity of 2 μ l and 1 μ l, respectively. The PCR reaction was 42 programmed as following: hold at 95°C for 10 min followed by 35 cycles of 95°C for 20s, 60°C 43 for 20s and 72°C for 22s. HRM was performed between 80 and 90°C with a stepwise increase of 44

0.05°C. The melting temperatures (Tm) were determined by the negative derivative of decrease

fluorescence over increased temperature (df/dt) for each SNP using the proprietary software	46
(version 1.7.34).	47
The first set run clearly separated lineages I, II, and III with distinctive melting curves, and the	48
Tm of each allele was at least half degree apart from that of the other allele (Fig. 1A). The second	49
set run distinguished the sub-lineages IIIa, IIIb, and IIIc with distinctive melting curves (Fig. 1B).	50
Table 2 summarizes Tm for all alleles. To assess the reproducibility, we blindly run all 10 samples	51
of various lineages randomly in same condition and revealed same results with +/- 0.05 different	52
in Tm. Sequencing of five SNPs independently verified reliability of Tm data.	53
A simple and robust genotyping report is important for rapid identification of the emerging	54
multidrug-resistant clones of lineage III (1). The multiplex-HRM approach we present here can be	55
very useful in this regard.	56

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Figure legend	93

Fig.1. Multiplex-HRM curve of two set runs. A: multiplex-HRM curve of three SNPs in kduD,
deoA and emrA respectively (from left to right), which separates lineages I, II and III (strain ID
1263, 55623 and 54184 respectively). B: multiplex-HRM curve of two SNPs in fdX (left) and
menF (right), respectively, which separates sub-lineages IIIa, IIIb, and IIIc (strain ID 54184,
2073, and 6224 respectively).

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Table 1: SNPs and primers used for multiplex-MRM-typing S. sonnei in this study:

	SNP		1	Lineage	Lineage	Sublineage	Sublineage	Sublineage	
Gene	SINE	Ancestor	Derived	Lineage	Lineage	Subilileage	Subilileage	Subilileage	
Gene	position	Ancestor	Delived	I	II	IIIa	IIIb	IIIc	
	Formor			_					
kduD	3155111	T	С	С	T	T	T	T	
deoA	4803842	G	A	G	A	G	G	G	
emrA	2973337	T	C	C	C	T	T	T	
fdx	2760031	G	A	G	G	G	A	A	
	245502								
menF	2455693	A	G	A	Α	A	A	G	
D 4	t- tD								
Primers for kduD		forward 5'-CGACGGCGAAACACTTTATC-3' reverse 5'-CGCGTATAAGAAGGCACACG-3'							
Primers f	or deoAD	forward 5'-GGAGATGCTTATCTCCGGCAAA-3'							
		reverse 5'-AGTCGGTTGGGCCTTTT-3'							
Primers for kduD		forward 5'-TGCCACCGAAGTACGTAACG-3'							
reverse 5'-CATCCACCCACATATTGGTG-3'									
Primers for kduD		forward 5'-CAAAGCCTGGGACTGGA-3' reverse 5'-CATGGTTGATAGTGTAACGC-3'							
D: C 11D									
Primers for kduD		forward 5'-TATTCTCGCGCTGGTTTTTA-3' reverse 5'-GCTTTTCTTGGCTCTTCACC-3'							

Note: These SNPs were extracted using S. sonnei strain Ss046 as the reference as described by Holt et al., 2012 (1) and Sangal et al., 2013 (4).

Journal of Clinical Microbiology

Table 2: Summary of *Tm* for each of the five SNPs used for multiplex-HMR

Main lineages	Tm (+/- 0.05)					
	kduD	deoA	emrA			
I	83.0	85.1	87.7			
II	82.5	84.8	87.7			
III	82.5	85.2	87.3			
Sub-lineages	Tm (+/- 0.05)					
	fdX		menF			
IIIa	82.5		85.7			
IIIb	82.5		86.0			
IIIc	82.9		85.7			



