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1 **Co-culture of *Methanobrevibacter smithii* with enterobacteria during urinary infection.**

2

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24 **Summary**

25 **Background.** Urinary tract infections are known to be caused by bacteria, but the potential  
26 implications of archaea have never been studied in this context.

27 **Methods.** In two different university hospital centres we used specific laboratory methods for  
28 the detection and culture of archaeal methanogens in 383 urine specimens prospectively  
29 collected for diagnosing urinary tract infection (UTI).

30 **Findings.** *Methanobrevibacter smithii* was detected by quantitative PCR and sequencing in  
31 34 (9%) of the specimens collected from 34 patients. *Escherichia coli*, *Klebsiella*  
32 *pneumoniae*, *Enterobacter* sp., *Enterococcus faecium* and mixed cultures were detected along  
33 with *M. smithii* in eighteen, six, three, one and six urine samples, respectively. Interestingly,  
34 using our specific culture method for methanogens, we also isolated *M. smithii* in 31 (91%) of  
35 the 34 PCR positive urine samples. Genotyping the 31 isolates using multispacer sequence  
36 typing revealed three different genotypes which have been previously reported in intestinal  
37 microbiota. Antibiotic susceptibility testing found the 31 isolates to be *in vitro* susceptible to  
38 metronidazole (MIC: 1 mg/L) but resistant to fosfomycin, sulfamethoxazole-trimethoprim,  
39 amoxicillin-clavulanate and ofloxacin, commonly used to treat bacterial UTI. Finally, 19  
40 (54%) of the 34 patients in whose urine samples *M. smithii* was detected were diagnosed with  
41 UTIs, including cystitis, pyelonephritis and prostatitis.

42 **Interpretation.** Our results show that *M. smithii* is part of the urinary microbiota of some  
43 individuals and could play a role in community-acquired UTI in association with enteric  
44 bacteria.

45 **Funding** This study was supported by IHU Méditerranée Infection, Marseille, France.

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49 **Introduction**

50 Forty-eight-hour culturing of a large volume of urine samples collected from women by  
51 transurethral catheter in parallel with suprapubic aspiration has shown that urine hosts a  
52 resident microbiota, which includes bacteria missed by routine procedures<sup>1</sup>. This resident  
53 urinary microbiota has been further explored using an expanded quantitative urine culture  
54 (EQUC) method<sup>2,3</sup>. Metagenomic analyses have confirmed that urine is not sterile in healthy  
55 individuals<sup>4-8</sup>.

56 An interesting concept currently emerging is that cases of urinary tract infection (UTI)  
57 may result from disequilibrium in the repertoire of the urinary microbiota rather than from the  
58 introduction of an exogenous pathogenic organism<sup>8,9</sup>. Therefore, efforts to characterise the  
59 repertoire of organisms residing in the urinary microbiota are important for clinical  
60 microbiology and medicine.

61 Current culture-dependant and culture-independent studies of the urinary microbiota  
62 rely on methods that are unable to detect archaea. These micro-organisms form a living  
63 domain distinct from that of bacteria and have been detected and cultured from the oral and  
64 gut microbiota<sup>10-12</sup>. Moreover, the specific archaea group of methanogens, characterised by  
65 the ability to produce methane from hydrogen, are recognised as emerging pathogens in a few  
66 clinical situations, including life-threatening brain abscess<sup>12-15</sup>.

67 In this study, we aimed to characterise the repertoire of human urine microbiota using  
68 laboratory protocols specifically designed for the detection and culture of methanogens in  
69 urines. Surprisingly, we found that among the six previously-described methanogens in non-  
70 urine human microbiota, only *Methanobrevibacter smithii* can be detected and cultured from  
71 urines.

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74 **Patients and methods**

75 **Sample collection.** The retrospective study was approved by the Ethics Committee of the  
76 IHU Méditerranée Infection under n° 2016-01. A total of 383 urine specimens taken for  
77 microbiological diagnosis of UTI in 383 patients without urinary catheter were prospectively  
78 collected at the IHU Méditerranée Infection Laboratory, Marseille, France (henceforth  
79 referred to as laboratory 1) (183 urines) and at the Laboratory of Bacteriology at the  
80 University Centre of Nice (henceforth referred to as laboratory 2) (200 urines) using a BD  
81 Vacutainer (Becton Dickinson, Le Pont de Claix, France). In this study, clean catch  
82 midstream urine specimen were collected by from September 2017 to November 2017 at  
83 laboratory 1 and from January 2018 to March 2018 at laboratory 2. A total of 50 BD  
84 vacutainer tubes filled with sterile phosphate buffered saline (PBS) (Thermo Fisher Scientific,  
85 Villebon-sur-Yvette, France) were intercalated in every 8 urine specimens to serve as negative  
86 controls in all further laboratory steps. The following data on each patient included in the  
87 study were anonymously collected: age, sex, underlying disease of the urinary tract, immune  
88 status, symptoms of urinary tract infection, antibiotic treatment (molecule and duration) and  
89 clinical outcome.

90 **Routine investigations.** Urine leukocyte and erythrocyte counts were obtained with a UF-  
91 1000i<sup>®</sup> analyser (Sysmex, bioMérieux, Craonne, France) in laboratory 1 and with an IQ 200  
92 IRIS analyser (Beckman Coulter, Villepinte, France) in laboratory 2. Gram staining was  
93 carried out with the automated PREVI Color Gram (bioMérieux) at both laboratories in order  
94 to stain bacteria. Indeed, methanogens are not usually coloured by using Gram staining. A  
95 200- $\mu$ L urine volume was cultured on a COLUMBIA COS medium (bioMérieux) (laboratory  
96 1) or on Uriselect medium (Bio-Rad, Marnes-la-Coquette, France) (laboratory 2) and  
97 incubated at 37 °C for 24 h (both laboratories), while visible colonies were identified by

98 matrix-assisted laser desorption ionisation–time of flight mass spectrometry (Brucker,  
99 Wissembourg, France), as previously described<sup>16</sup>.

100 **Methanogen DNA extraction, PCR and sequencing.** The manual protocol was carried out  
101 using the NucleoSpin® Tissue Mini Kit (Macherey Nagel, Hoerdt, France) as previously  
102 described<sup>11,17</sup>. Extraction of 250 µL of sterile PBS was included in each DNA extraction  
103 series as a negative control.

104 Methanogen PCR and sequencing was performed as described in previous articles<sup>11,17</sup> using  
105 the 16S rRNA broad range archaeal forward primer 5'-CCGGGTATCTAATCCGGTTC- 3'  
106 and reverse primer 5'-CTCCCAGGGTAGAGGTGAAA-3'<sup>18</sup>, and the *mcrA* gene forward  
107 primer 5'-GCTCTACGACCAGATMTGGCTTGG-3' and reverse primer 5'-  
108 CCGTAGTACGTGAAGTCATCCAGCA-3'<sup>19</sup>. All PCR products were sequenced and the  
109 various fragments were assembled using the ChromasPro1.34 software (Technelysium Pty.  
110 Ltd., Tewantin, Australia) and compared with sequences available in the GenBank database  
111 using the online NCBI BLAST program (<http://blast.ncbi.nlm.nih.gov/gate1.inist.fr/Blast.cgi>).

112 Extracted DNA was also analysed by real-time PCR targeting *M. smithii* using the following  
113 primer pair: *M. smithii*-F (5'-ACCATAACyATCAGCAGCATTAT-3') and *M. smithii*-R (5'-  
114 AGTATTGGTGAAGGATTTaCTGT-3') (Eurogentec, Seraing, Belgium) and the *M. smithii*  
115 probe (6-carboxyfluorescein [FAM]-5'ACCyTTATCAGCTTTACCA TTAATyAAAG-3')  
116 (Applied Biosystems, Courtaboeuf, France) as described in a previous study<sup>13</sup>.

117 **Methanogen isolation and culture.** A 250 µL volume of urine sample was seeded in ambient  
118 air in a sterile Hungate tube<sup>20</sup> (Dominique Dutscher, Brumath, France). The Hungate culture  
119 tube was developed specifically for growing and storing strictly anaerobic bacteria and  
120 archaea. It features an autoclavable screw cap with a 9-mm opening, a non-toxic, gas-  
121 impermeable butyl rubber stopper and a disposable screw cap. We placed 5 mL of SAB  
122 broth<sup>21</sup> into each Hungate tube, seeded it with urine then inoculated it with *Bacteroides*

123 *thetaitotaomicron* ( $10^5$  cells/mL) for hydrogen production<sup>22</sup>. The mixture in the Hungate tube  
124 was then incubated at 37 °C with agitation for seven days. Methanogen growth was inferred  
125 from the production of CH<sub>4</sub> detected by gas chromatography, as previously described<sup>11</sup>. The  
126 subculture was seeded on a Petri dish containing SAB medium supplemented with 15 g/L  
127 agar and placed in the upper chamber of a double-chamber box. Tubes inoculated with sterile  
128 PBS were used as negative controls in the archaea isolation experiments (one negative control  
129 every five tubes); culture dishes containing the SAB medium inoculated with sterile PBS were  
130 used as negative controls in the sub-culture experiments (one negative control every five  
131 dishes).

132 **Characterisation of isolates.** Isolates were further genotyped using the Multi Spacer Typing  
133 method, as previously described<sup>23</sup>. In addition, we determined the antibiotic susceptibility  
134 profile of the isolates, as previously described, by incorporating the following antibiotics:  
135 fosfomycin (100 mg/L), sulfamethoxazole-trimethoprim (50 mg/L), amoxicillin-clavulanate  
136 (100 mg/L) and ofloxacin (1 mg/L) (BIOGARAN, Colombes, France)<sup>21</sup>. *M. smithii* culture  
137 with metronidazole (1 mg/L) was used as the positive control and *M. smithii* culture without  
138 metronidazole as the negative control.

139 **Statistical analyses.** Data were analysed with Prism 7.0 (GraphPad Software) by unpaired  
140 Student's t-test and chi-square test (\*\* p < 0.01, \* p < 0.05, ns: non-significant). We used the  
141 former to compare the quantitative variables from the two centres (pH, urine salinity, white  
142 and red blood cells), and the latter to compare the categorical variables (sex ratio, % of  
143 standard cultures positive, % of urine samples positive for methanogens by molecular or  
144 culture methods).

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148 **Results**

149 **Population and routine investigations.** A total of 383 urine specimens from 383 patients  
150 were collected and analysed prospectively, 183 specimens in laboratory 1 and 200 specimens  
151 in laboratory 2. Mean patient age was 56 years (0-95 years) and 61% of the patients were  
152 women. Leukocyturia was significant in 205 of the 383 specimens ( $> 10^4$  leukocytes/mL) and  
153 142 specimens yielded bacterial growth. Comparison of the data collected from laboratory 1  
154 and from laboratory 2 is shown in Table 1.

155 **Methanogen quantitative PCR and sequencing.** Of the 383 urine samples analysed by  
156 standard PCR sequencing, 18/183 (10%) from laboratory 1 and 16/200 (8%) from laboratory  
157 2 were positive for both archaeal 16S rRNA PCR and the *mcrA* PCR, while 165/183 (90%)  
158 and 184/200 (92%), respectively, were negative in both PCR assays in the presence of 50  
159 negative controls, all of which remained negative. Sequencing the 16S rRNA PCR products  
160 revealed 99% sequence similarity with the homologous fragment of the reference 16S rRNA  
161 gene of *M. smithii* strain NVD (accession NCBI: LT223565). Further sequencing of the PCR-  
162 amplified *mcrA* gene revealed 99% sequence similarity with the homologous fragment of the  
163 reference *mcrA* gene of *M. smithii* strain NVD (accession NCBI: LT223565) and *M. smithii*  
164 ATCC 35061 (accession NCBI: NR\_074235). Quantitative PCR analyses targeting the *M.*  
165 *smithii* 16S rRNA gene yielded a median Ct of  $33.7 \pm 2.47$ , indicative of an *M. smithii* load of  
166  $1.24 \times 10^3 \pm 1.56 \times 10^3$  / mL. Detection of *M. smithii* in urine samples was significantly  
167 associated with leukocyte counts greater than  $10^4$  /mL, P-value = 0.0015 (Chi-square test).

168 **Isolation and culture of *M. smithii*.** To test the viability of the *M. smithii* organisms detected  
169 by the PCR-based methods, methanogen was cultured on 18 samples in laboratory 1 and 16  
170 samples in laboratory 2. Fifteen of the 18 cultured samples in laboratory 1 and all 16 cultured  
171 samples in laboratory 2 yielded colonies, which were obtained after 20 days of incubation. All  
172 the colonies were identified as *M. smithii* on the basis of PCR sequencing of the 16S rRNA



173 and *mcrA* genes, which revealed 100% sequence similarity with the reference homologous  
174 genes in *M. smithii* strain NVD (accession NCBI: LT223565) (Table 1). MST genotyping  
175 indicated that the *M. smithii* isolates belonged to genotype 2 (n=13), genotype 3 (n=10) and  
176 genotype 1 (n=8) (Supplementary Table 1). Antibiotic susceptibility testing indicated that  
177 fosfomycin (100 mg/L), sulfamethoxazole-trimethoprim (50 mg/L), amoxicillin-clavulanate  
178 (100 mg/L) and ofloxacin (1 mg/L) were inactive against *M. smithii*, whereas metronidazole -  
179 used as a positive control - inhibited the growth of *M. smithii*.

180 **Microbiological and clinical analyses.** *M. smithii* was detected in 34 (9%) of the 383 urine  
181 samples analysed in the study. *Escherichia coli*, *Klebsiella pneumoniae*, *Enterobacter* sp. and  
182 *Enterococcus faecium* were detected along with *M. smithii* in 18, 6, 3 and 1 urine samples,  
183 respectively. The other six samples in which *M. smithii* was detected yielded mixed cultures  
184 including *Enterobacteriaceae*. Of the 34 patients with *M. smithii* in their urine samples,  
185 63.4% were women, and mean patient age was 63 years (20-95). Ten patients (30%) had an  
186 underlying urinary tract disease, 14 (40%) were immunosuppressed and 19 (44%) had chronic  
187 renal failure (GFR < 60L/min). It is worth noting that *M. smithii* was associated with UTI in  
188 19 (56%) patients and with colonisation in 15 (44%). Of the 19 patients with UTIs, 17 (88%)  
189 were community acquired, 13 presented with cystitis, while four were diagnosed with  
190 pyelonephritis and two with prostatitis. All 19 patients received antibiotic treatments targeting  
191 the isolated bacteria and commonly used to treat UTIs, including beta-lactams,  
192 fluoroquinolones and fosfomycin-trometamol, with favourable outcomes (Table 2).

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198 **Discussion**

199 *M. smithii* was the only methanogen prospectively detected in the urines using a  
200 polyphasic approach, including molecular detection by PCR sequencing of two unrelated  
201 genes, and isolation and culture methods specifically developed for methanogenic archaea.  
202 Given that the negative controls included in every experimental procedure remained negative,  
203 that *M. smithii* was detected by different unrelated technical approaches, and that similar data  
204 were obtained in two unrelated laboratories, we can exclude the possibility of in-laboratory  
205 contamination and centre bias in the study. These pieces of experimental evidence, therefore,  
206 validate the data reported here.

207 *M. smithii* was first detected as a normal organism in the gut microbiota of the vast  
208 majority of individuals<sup>11,24</sup>, although it is depleted in children diagnosed with severe  
209 malnutrition and kwashiorkor<sup>25</sup>. It was then shown to be a component of the oral cavity  
210 microbiota in the saliva<sup>17</sup> and the dental plaque, including periodontitis<sup>26,27</sup> and peri-  
211 implantitis lesions<sup>10,28</sup>. More recently, the pathogenic potential of *M. smithii* was illustrated by  
212 its isolation and culture from a muscular abscess<sup>14</sup> and by its detection in one case of brain  
213 abscess<sup>13</sup>. Our study is of interest because it reports for the first time *M. smithii*, methanogens  
214 and archaea at large in urines, as all previous culture-based and culture-independent studies  
215 have failed to detect archaea in this type of fluid. Our success in detecting *M. smithii* was  
216 down to the use of laboratory tools that we developed specifically for detecting  
217 methanogens<sup>22, 29-31</sup>. These tools, including the technique we set up for easy isolation and  
218 culture of methanogens, could be easily implemented in other clinical microbiology  
219 laboratories enabling them to investigate the presence of methanogens in urines<sup>30</sup>.

220 We detected *M. smithii* in 9% of urine samples, and, furthermore, in the same  
221 proportions of patients in two unrelated laboratories (10% in laboratory1 and 8% in laboratory  
222 2). This supports the idea that *M. smithii* may be part of “the urinary microbiota”, a new

223 concept that has emerged since the expansion of urine metagenomic analyses and EQUIC  
224 approaches. It is worth noting that all the urine samples which cultured *M. smithii* also  
225 cultured enterobacteria, such as *E. coli*, *K. pneumoniae*, *Enterobacter sp.*, which are known to  
226 produce hydrogen as a substrate for methanogenesis<sup>32</sup>. The fact that *M. smithii* was isolated  
227 along with enterobacteria and never detected in a urine sample without bacterial growth  
228 suggests that it could play a role in or even induce dysbiosis, which facilitates the growth of  
229 enterobacteria, acknowledged agents of UTI. In our study, 19 (54%) of the 34 patients with  
230 *M. smithii* in their urine samples presented with UTIs, were most of them community  
231 acquired. These preliminary results need to be confirmed by further studies, including  
232 epidemiological, clinical and fundamental approaches, in order to determine the role of *M.*  
233 *smithii* in UTIs. Finally, our study indicates that the antibiotics commonly used to treat  
234 bacterial urinary tract infections are inactive against *M. smithii*, consistent with a previous  
235 report of its narrow spectrum of *in vitro* susceptibility<sup>33</sup>. Interestingly, the 19 patients in  
236 whose urine samples we detected *M. smithii* and who presented with UTI symptoms showed  
237 clinical improvement after receiving antimicrobial treatment active against enterobacteria but  
238 inactive against methanogens. We hypothesise that the inhibition of enterobacterial growth by  
239 appropriate antibiotic treatment induces a decrease <sup>2</sup>in hydrogen production, a major substrate  
240 for methanogenesis, and therefore destroys the metabolic cooperation between these two  
241 microorganisms.

242 In conclusion, our data show that *M. smithii* is part of the urinary microbiota in some  
243 patients and suggest that it plays a role in community-acquired UTIs in association with  
244 enterobacteria. Therefore, we encourage colleagues to use the above-described methods to  
245 investigate methanogens in urines, including *M. smithii*, in order to determine its potential  
246 contribution to the pathogenesis of UTIs.

247

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344 **Author contributions**

345 GG and RL wrote the manuscript. GG, RL cultivated the methanogens and carried out the

346 PCR. DC, RL and AC performed the clinical analyses. MD, DR and RR supervised the study.

347 **Conflicts of interests**

348 GG, RL, DC, AC, DR, MD, RR declare no potential conflict of interest.

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**Table 1. Comparison of the data collected in laboratory 1 and in laboratory 2.**

	Laboratory 1	Laboratory 2	Statistical analysis (+)
Number of patients	183	200	
Sex ratio (male/female)	56/127	93/107	p=0.0014 (Chi-square test)
Age	55 +/- 25	57 +/- 21	p=0.44 (t-test)
Number of urine samples	183	200	
Routine urine analysis			
pH*	6.36 +/- 0.97	6.31 +/- 0.97	p=0.62 (t-test)
Urine salinity ‰*	43.09 +/-18.06	42.18 +/- 16.86	p=0.61 (t-test)
White blood cells**	16.1 (0.1-3505)	8 (0-1000)	p=0.012 (t-test)
Red blood cells**	14.4 (0.1-7554)	7 (0-1000)	p=0.06 (t-test)
Standard culture +	53/183 (29%)	89/200 (45%)	p=0.002(Chi-square test)
Methanogen approach			
Archaea <i>I6S rDNA</i> PCR +	18/183 (10%)	16/200 (8%)	p=0.44(Chi-square test)
<i>mcrA</i> PCR +	18/183 (10%)	16/200 (8%)	p=0.44(Chi-square test)
Sequencing	<i>M. smithii</i> (n=18)	<i>M. smithii</i> (n=16)	
Methanogen culture +	15/183 (8.2%)	16/200 (8%)	p=0,83(Chi-square test)
Strain identification	<i>M. smithii</i> (n=15)	<i>M. smithii</i> (n=16)	

\* pH and Urine salinity values are expressed as mean +/- SD.\*\*White and red blood cells are expressed as median (range).

**Table 2. Clinical and microbiological features of 19 patients presenting with UTIs and in whose urines *M. smithii* was detected**

Type of UTI	Age (sex)	Underlying diseases of the genitourinary tract	Nosocomial	Leukocytes in urine (/μL) **	Associated bacteria (CFU/mL)	Treatment (Duration)	Course of infection
Cystitis	87 (F)	No	Yes	>1000	<i>Escherichia coli</i> (10 <sup>6</sup> )	Amoxicillin-clavulanic acid (7 days)	Favourable
	65 (F)	No	No	64	<i>Escherichia coli</i> (10 <sup>6</sup> ) and <i>Proteus mirabilis</i> (10 <sup>6</sup> )	Fosfomycin-trometamol (1 day)	Reinfection at day 45 ( <i>E. coli</i> and <i>P. mirabilis</i> )
	74 (F)	No	No	5	<i>Klebsiella pneumoniae</i> (10 <sup>4</sup> )	Pivmecillinam (7 days)	Favourable
	55 (F)	No	No	592	<i>Escherichia coli</i> (10 <sup>3</sup> )	Ofloxacin (NA)	Favourable
	25 (F)	No	No	140	<i>Escherichia coli</i> (10 <sup>6</sup> )	Pivmecillinam (5days)	Favourable
	52 (F)	Renal transplant	Yes	10	<i>Klebsiella pneumoniae</i> (10 <sup>6</sup> )	Ceftriaxone (NA)	Favourable
	50 (F)	No	No	464	<i>Escherichia coli</i> (10 <sup>6</sup> )	NA	NA
	83* (F)	No	No	23	<i>Escherichia coli</i> (10 <sup>6</sup> )	Fosfomycin and Colistin (42 days)	NA
	36 (F)	No	No	11	<i>Escherichia coli</i> (10 <sup>6</sup> )	NA	NA
	20 (F)	No	No	78	<i>Escherichia coli</i> (10 <sup>6</sup> )	Pivmecillinam (5days)	Favourable
	56* (M)	No	No	25	<i>Escherichia coli</i> (10 <sup>6</sup> )	Ciprofloxacin (3 days)	Favourable

	59 (F)	Renal transplant	No	4	<i>Klebsiella pneumoniae</i> (10 <sup>6</sup> )	Ciprofloxacin (7 days)	Favourable
	79* (F)	Chronic renal failure	No	85	<i>Enterobacter cloacae</i> (10 <sup>6</sup> )	Amoxicillin-clavulanic acid (2 days)	Favourable
Pyelonephritis	49 (F)	Renal lithiasis	No	150	<i>Enterobacter cloacae</i> (10 <sup>3</sup> )	Ceftriaxone IV and gentamicin (NA)	Favourable
	69 (M)	Renal lithiasis	No	20	<i>Escherichia coli</i> (10 <sup>6</sup> )	Ceftriaxone IV (5days)	Favourable
	75 (F)	Vesico-vaginal fistula	No	>1000	<i>Klebsiella pneumoniae</i> (10 <sup>6</sup> )	Imipenem IV (10 days)	Favourable
	83 (F)	No	No	53	<i>Escherichia coli</i> (10 <sup>6</sup> )	Ofloxacin (1day) and amoxicillin-clavulanic acid (7days)	Favourable
Prostatitis	69 (M)	No	No	221	<i>Escherichia coli</i> (10 <sup>5</sup> )	Ceftriaxone and gentamicin (NA)	Favourable
	50 (M)	No	No	>1000	<i>Escherichia coli</i> (10 <sup>4</sup> )	Pivmecillinam (15 days)	Favourable

(\*) *M. smithii* was detected in the urine of this patient using molecular methods, but not isolated in culture using the approach developed specifically for archaea.

(\*\*) Significant threshold was  $\geq 10/\mu\text{L}$  for leukocyturia. NA: not available. Favourable: patient with favourable outcome based on clinical evidence.

**Supplementary Table. Multispacer sequence typing (MST) genotyping of the 31 strains of *M. smithii* isolated during the study.**

Sample	spacer 1	spacer 2	spacer 3	spacer 4	spacer Type
13	X	X	X		1
14	X	X	X	X	2
41		X	X		3
42		X	X		3
52	X	X	X		1
58		X	X		3
63	X	X	X	X	2
75	X	X	X	X	2
99		X	X		3
106	X	X	X		1
112		X	X		3
126	X	X	X	X	2
128	X	X	X	X	2
133	X	X	X		1
149	X	X	X	X	2
167		X	X		3
4	X	X	X	X	2
6	X	X	X		1
11	X	X	X	X	2
13		X	X		3
49	X	X	X		1
67	X	X	X	X	2
81	X	X	X	X	2
87		X	X		3
108	X	X	X	X	2
114		X	X		3
115	X	X	X		1
125		X	X		3
173	X	X	X	X	2
177	X	X	X		1
188	X	X	X	X	2