Targeted next-generation sequencing of the 16S-23S rRNA region for culture-independent bacterial identification - increased discrimination of closely related species

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The aim of this study was to develop an easy-to-use culture-free diagnostic method based on next generation sequencing (NGS) of PCR amplification products encompassing whole 16S-23S rRNA region to improve the resolution of bacterial species identification. To determine the resolution of the new method 67 isolates were subjected to four identification methods: Sanger sequencing of the 16S rRNA gene; NGS of the 16S-23S rRNA region using MiSeq (Illumina) sequencer; Microflex MS (Bruker) and VITEK MS (bioMérieux). To evaluate the performance of this new method when applied directly on clinical samples, we conducted a proof of principle study with 60 urine samples from patients suspected of urinary tract infections (UTIs), 23 BacT/ALERT (bioMérieux) positive blood culture bottles and 21 clinical orthopedic samples. The resolution power of NGS of the 16S-23S rRNA region was superior to other tested identification methods. Furthermore, the new method correctly identified pathogens established as the cause of UTIs and blood stream infections with conventional culture. NGS of the 16S-23S rRNA region also showed increased detection of bacterial microorganisms in clinical samples from orthopedic patients. Therefore, we conclude that our method has the potential to increase diagnostic yield for detection of bacterial pathogenic species compared to current methods.

Timely, appropriate treatment of infection depends on rapid and specific identification of causative microorganisms. To date, identification of bacterial species highly depends on culture or molecular tests, including 16S rRNA gene Sanger sequencing1,2. Significant limitations of culture methods are that some bacteria are slow-growing or fastidious, making identification of them complicated and time and resource consuming3. Although molecular tests that target specific microorganisms are more rapid and sensitive than culture methods they need an a priori knowledge of the likely pathogenic species that could be present in a sample. If bacteriological identification by culture methods fails, Sanger sequencing of the variable 16S rRNA gene is used for identification. The gene sequence has been proven to be a reliable genetic marker as it is present in all bacteria and its function has not changed over time4. However, unequivocal identification is not always possible due to the high sequence similarities of the 16S rRNA gene in some bacterial species5. Although Sanger sequencing of the 16S rRNA gene can be applied directly on clinical materials, in polymicrobial samples it usually cannot identify more than one species simultaneously or at least this process is challenging by sorting out the ambiguous signals from mixed chromatograms using a computer program6.

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| Sample | Reported cause of UTI by conventional culture identification | Growth CFU/ml | Identification of additional colonies by culture methods | Species content by NGS of 16S-23S rRNA region (% of total reads) |
|--------|-------------------------------------------------------------|---------------|-----------------------------------------------------------|---------------------------------------------------------------|
| UR1    | *Pseudomonas aeruginosa*                                   | $10^6$        | Lactobacillus gasser (97.2%), *Pseudomonas aeruginosa* (2.4%), *Corynebacterium amycolatum* (0.4%) |                                                                |
| UR2    | *Proteus mirabilis*                                         | $10^6$        | *Proteus vulgaris* (70.3%), *Proteus mirabilis* (29.7%)     |                                                                |
| UR3    | *Escherichia coli*                                          | $10^6$        | *Escherichia coli* (99.1%), Lactobacillus delbrueckii (0.9%) |                                                                |
| UR4    | *Escherichia coli*                                          | $10^6$        | *Escherichia coli* (99.1%), *Peptoniphilus lacrimalis* (0.3%), *Bacteroides sp.* (0.6%) |                                                                |
| UR5    | *Escherichia coli*                                          | $10^6$        | *Escherichia coli* (93.2%), *Undibacterium oligoacidophilum* (6.3%), *Pseudomonas saccharophilia* (0.3%), *Phenylobacterium* sp (0.2%) |                                                                |
| UR6    | No clinical significance                                   | $10^6$        | *Bifidobacterium* sp                                       | *Actinobaculum schaalii* (100%)                                 |
| UR7    | *Escherichia coli*                                          | $10^6$        | *Escherichia coli* (80.7%), *Lactobacillus crispatus* (19.3%) |                                                                |
| UR8    | *Escherichia coli*                                          | $10^6$        | *Escherichia coli* (98.7%), *Enterococcus faecalis* (1.0%), *Aerococcus sanguinicola* (0.3%) |                                                                |
| UR9    | *Escherichia coli*                                          | $10^6$        | *Escherichia coli* (53.7%), Klebsiella oxytoca (43.6%), *Staphylococcus aureus* (2.4%), *Enterococcus faecalis* (0.3%) |                                                                |
| UR10   | No clinical significance                                   | $10^6$        | Esakella persensii (28.6%), *Fenollaria massiliensis* (4.1%), *Helcococcus sp.* (2.6%), *Peptoniphilus duodenii* (2.6%), *Mobiluncus curtisi* (2.4%), *Varrichibaculum cambriense* (2.2%), *Peptoniphilus harei* (1.3%), *Actinobaculum urinae* (0.8%), *Peptoniphilus lacrimalis* (0.8%), *Propionimicrobium* sp. (0.7%), *Facklamia sp.* (0.7%), *Finigolda magna* (0.6%), *Anaerococcus obeiensis* (0.6%), *Anaerococcus prevoti* (0.4%), *Anaerococcus degenerii* (0.4%), *Actinobaculum sp.* (0.4%), *Aerococcus urinae* (0.3%), *Fastidiosipila sanguinis* (0.2%), *Eustidiobius sanguinis* (0.2%), *Bacteroides coagulans* (0.2%), Unidentified species (50.2%) |                                                                |
| UR11   | No clinical significance                                   | $10^6$        | Staphylococcus epidermidis                                 | No PCR product                                                 |
| UR12   | No clinical significance                                   | $10^6$        | *Proteus mirabilis* (75.5%), *Proteus vulgaris* (17.5%), *Undibacterium oligoacidophilum* (4.8%), *Aerococcus urinae* (1.2%), *Corynebacterium striatum* (0.3%), *Pseudomonas saccharophilia* (0.3%), *Enterococcus faecalis* (0.2%), *Ralstonia pickettii* (0.1%) |                                                                |
| UR13   | No clinical significance                                   | $10^6$        | No PCR product                                             |                                                               |
| UR14   | No clinical significance                                   | $10^6$        | Undibacterium oligoacidophilum (36.4%), *Fenollaria massiliensis* (15.6%), *Mobiluncus curtisi* (10.6%), *Peptoniphilus lacrimalis* (5.9%), Unidentified species (5.3%), *Peptonitreroccoccus anaeobius* (4.4%), *Peptoniphilus koenosneriene* (4.1%), *Pseudomonas saccharophilia* (3.1%), *Atopobium delae* (3.7%), *Candidatus Peptoniphilus massiliensis* (2.6%), *Anaerococcus sp.* (1.8%), *Lonquetella anthrao* (1.1%), *Peptoniphilus harei* (1.1%), *Streptococcus anginosus* (0.8%), *Ralstonia pickettii* (0.7%), *Dialister propioniciaciens* (0.6%), *Methylbacterium oryzae* (0.6%), *Asimihobacurrum lactis* (0.5%), *Methylbacterium jejouli* (0.5%), *Peptoniphilus duodenii* (0.4%), *Fusobacterium nucleatum* (0.3%) |                                                                |
| UR15   | No clinical significance                                   | $10^6$        | Streptococcus anginosus                                     | Unidentified species (28.3%), *Undibacterium oligoacidophilum* (13.1%), *Peptoniphilus lacrimalis* (11.5%), *Mobiluncus curtisi* (9.5%), *Streptococcus anoginosus* (8.3%), *Fusobacterium nucleatum* (3.8%), *Propionimicrobium sp.* (2.1%), *Varrichibaculum cambriense* (2.0%), *Dialister sp.* (1.9%), *Atopobium delae* (1.2%), *Facklamia hominis* (1.1%), *Pseudomonas saccharophilia* (1.1%), *Dialister propioniciaciens* (0.9%), *Faecalbacterium prausnitzii* (0.9%), *Parvimonas micra* (0.9%), *Fenollaria massiliensis* (0.9%), *Prevotella disiens* (0.9%), *Peptoniphilus harei* (0.8%), *Actinobaculum massiliense* (0.7%), *Dialister succinatutens* (0.7%), *Gemmiger formicillus* (0.7%), *Corynebacterium pyruvicproduens* (0.6%), *Mycoplasma spermatophilus* (0.6%), *Actinobaculum sp.* (0.5%), *Asimihobacurrum lactitis* (0.5%), *Atopobium vaginiae* (0.5%), *Mobiluncus curtisi* (0.5%), *Alistipes onderdonki* (0.4%), *Anaerococcus sp.* (0.4%), *Bacteroides massiliensis* (0.4%), *Anaerococcus lactyliticus* (0.3%), *Anaerococcus obeiensis* (0.3%), *Anaerococcus prevoti* (0.3%), *Finigolda magna* (0.3%), *Methylbacterium jeouli* (0.3%), *Peptoniphilus duodenini* (0.3%), *Peptoniphilus massiliensis* (0.3%), *Ralstonia pickettii* (0.3%), *Treponema refringens* (0.3%), *Actinomyces turicensis* (0.2%), *Bacteroides sp.* (0.2%), *Dialister micraerophilus* (0.2%), *Euvacterium hallii* (0.2%), *Peptoniphilus koenosneriene* (0.2%), *Peptoniphilus obeis* (0.2%), *Phascolarctobacterium succinattens* (0.2%), *Porphyromonas bennonis* (0.2%) |                                                                |
| UR16   | No clinical significance                                   | $10^6$        | No PCR product                                             |                                                               |
| UR17   | No clinical significance                                   | $10^6$        | Atopobium delae (32.6%), *Undibacterium oligoacidophilum* (19.4%), *Lactobacillus iners* (16.4%), Unidentified species (10.5%), *Peptoniphilus coxii* (7.5%), *Anaerococcus lactyliticus* (2.7%), *Parvimonas micra* (2.2%), *Peptoniphilus lacrimalis* (1.8%), *Pseudomonas saccharophilia* (1.6%), *Peptoniphilus grossae* (1.0%), *Aerococcus christensenii* (0.9%), *Peptoniphilus duodenini* (0.9%), *Peptoniphilus harei* (0.5%), *Actinobaculum schaalii* (0.4%), *Ralstonia pickettii* (0.4%), *Solobacterium moorei* (0.4%), *Asimihobacurrum lactis* (0.3%), *Propionibacterium acnes* (0.2%), *Varrichibaculum cambriense* (0.2%) |                                                               |
| UR18   | No clinical significance                                   | $10^6$        | No PCR product                                             |                                                               |

Continued
| Sample | Reported cause of UTI by conventional culture identification | Growth CFU/ml | Identification of additional colonies by culture methods | Species content by NGS of 16S-23S rRNA region (% of total reads) |
|--------|-------------------------------------------------------------|--------------|--------------------------------------------------------|-------------------------------------------------------------|
| UR19   | No clinical significance                                   | $10^4$       | Kundibacterium oligoceanobacterium (28.1%), Sneathia sanguinegens (18.3%), Lactobacillus iners (16.5%), Atopobium vaginae (4.3%), Aerococcus chrisitensenii (4.0%), Euplasma sraealyticum (3.5%), Pelomomas saccharophila (1.8%), Ratulonia picketti (1.7%), Megaplasma indica (1.4%), Streplococcus anginosus (0.9%), Methylobacterium radiotolerans (0.6%), Prevotella amnii (0.6%), Methylbacterium aerolatum (0.5%), Dialister micraerophilus (0.4%), Ainthibacterium lactis (0.2%), Brevundimonas intermedia (0.2%), Unidentified species (16.8%) |
| UR20   | No clinical significance                                   | $10^2$       | No PCR product                                         |
| UR21   | No clinical significance                                   | $10^2$       | No PCR product                                         |
| UR22   | No clinical significance                                   | $10^2$       | Lactobacillus crispatus (78.5%), Kundibacterium oligoceanobacterium (14.4%), Pseudomonas saccharophila (2.3%), Kundibacterium ligoceanobacterium (1.5%), Ratulonia picketti (1.0%), Asinibacterium lactis (0.4%), Lactobacillus vaginalis (0.3%), Staphylococcus capitis (0.2%), Methylobacterium oryzae (0.1%), Unidentified species (1.3%) |
| UR23   | No clinical significance                                   | $10^2$       | No PCR product                                         |
| UR24   | No clinical significance                                   | $10^1$       | Staphylococcus epidermids, Streptococcus aginogus Lactobacillus jenseni (98.5%), Finegoldia magna (0.5%), Streptococcus aginogus (0.5%), Anaerococcus ohtsientis (0.5%), Propionibacterium lundihyphiformum (0.2%) |
| UR25   | No clinical significance                                   | $10^1$       | Staphylococcus hemolyticus, Proteus mirabilis, Escherichia coli Lactobacillus iners (77.8%), Acinetobacter lwofii (12.3%), Acinetobacter radioreiseters (3.8%), Escherichia coli (3.0%), Proteus mirabilis (1.1%), Enterococcus faecalis (0.9%), Peptoniphilus harei (0.4%), Strepptococcus agalactiae (0.4%), Finegoldia magna (0.2%), Proteus vulgaris (0.2%), Corynebacterium sp. (0.1%), Pseudomonas stutzeri (0.1%), Streptococcus sp. (0.1%) |
| UR26   | No clinical significance                                   | $10^1$       | Staphylococcus hominis No PCR product                  |
| UR27   | No clinical significance                                   | $10^1$       | Staphylococcus hominis Lactobacillus iners (84.1%), Lactobacillus jenseni (11.1%), Lactobacillus sp. (2.8%), Kundibacterium oligoceanobacterium (0.9%), Staphylococcus hominis (0.5%), Lactobacillus vaginalis (0.2%), Pseudomonas saccharophila (0.2%), Asinibacterium lactis (0.1%), Corynebacterium tuberculostearicum (0.1%) |
| UR28   | No clinical significance                                   | $10^1$       | Enterococcus faecalis, Streptococcus mitis/oralis No PCR product |
| UR29   | No clinical significance                                   | $10^2$       | No PCR product                                         |
| UR30   | No clinical significance                                   | $10^2$       | No PCR product                                         |
| UR31   | No clinical significance                                   | $10^2$       | No PCR product                                         |
| UR32   | No clinical significance                                   | $10^2$       | No PCR product                                         |
| UR33   | Pseudomonas aeruginosa                                     | $10^2$       | Pseudomonas aeruginosa (99.7%), Peptostreptococcaceae sp. (0.2%), Anaerococcus degeneri (0.1%) |
| UR34   | Escherichia coli                                           | $10^2$       | Escherichia coli (100%)                                |
| UR35   | Escherichia coli                                           | $10^2$       | Escherichia coli (100%)                                |
| UR36   | Proteus mirabilis                                          | $10^2$       | Unidentified species (61.4%), Peptoniphilus harei (18.0%), Proteus mirabilis (16.3%), Aerococcus urinar (4.2%), Staphylococcus aureus (0.1%) |
| UR37   | Escherichia coli                                           | $10^2$       | Escherichia coli (100%)                                |
| UR38   | Escherichia coli                                           | $10^2$       | Escherichia coli (100%)                                |
| UR39   | Klebsiella pneumonia                                       | $10^2$       | Klebsiella pneumonia (100%)                            |
| UR40   | Pseudomonas aeruginosa                                     | $10^2$       | Pseudomonas aeruginosa (100%)                          |
| UR41   | Klebsiella pneumonia                                       | $10^2$       | Klebsiella pneumonia (100%)                            |
| UR42   | Escherichia coli                                           | $10^2$       | Escherichia coli (100%)                                |
| UR43   | Klebsiella pneumonia                                       | $10^2$       | Klebsiella pneumonia (100%)                            |
| UR44   | Klebsiella pneumonia                                       | $10^2$       | Klebsiella pneumonia (100%)                            |
| UR45   | No clinical significance                                   | $10^2$       | Strepococcus mitis/oralis, Escherichia coli, Staphylococcus aureus, Staphylococcus epidermids Ureaplasma parvum (42.2%), Prevotella sp. (27.0%), Prevotella bivia (10.8%), Anaerococcus lactolyticus (7.7%), Peptostreptococcus anaerobius (6.4%), Peptoniphilus sp. (1.7%), Finegoldia magna (1.5%), Strepococcus oralis (1.5%), Escherichia coli (0.6%), Lactobacillus crispatus (0.4%), Staphylococcus aureus (0.3%) |
| UR46   | Enterococcus faecalis                                      | $10^2$       | Enterococcus faecalis (99.7%), Lactobacillus iners (0.3%) |
| UR47   | No clinical significance                                   | $10^2$       | Gordonia alkalovorans, Sphingomonas truperti, Staphylococcus schleiferi, Corynebacterium spathularum Aerococcus sanguincola (92.5%), Aerococcus sp. (4.4%), Aerococcus urinar (1.8%), Peptoniphilus harei (0.9%), Lactobacillus iners (0.3%) |
| UR48   | Staphylococcus aureus                                      | $10^2$       | Klebsiella pneumonia (99.8%), Escherichia coli (0.2%) |
| UR49   | Staphylococcus epidermids                                  | $10^2$       | Staphylococcus epidermids (97.6%), Staphylococcus sp. (1.3%), Gardnerella vaginalis (1.1%) |
| UR50   | Enterococcus faecalis                                      | $10^2$       | Enterococcus faecalis (100%)                           |
| UR51   | No clinical significance                                   | $10^2$       | Staphylococcus epidermids (100%)                        |
Next generation sequencing (NGS) allows culture free detection of a theoretically unlimited number of pathogens and thus provides insight in the full microbiome. Since the introduction of benchtop sequencers, NGS is likely to become a diagnostic tool within the next few years in microbiological laboratories. Metagenomics will be the ultimate approach in detecting all microorganisms (e.g. bacteria, viruses, fungi) in a sample. Unfortunately,
| Sample | Patient | Material                  | Culture                          | NGS of 16–23S rRNA region (% of total reads) |
|--------|---------|---------------------------|----------------------------------|---------------------------------------------|
| KM1    | Patient A | biopsy (tissue)          | Negative                         | Propionibacterium acnes (9.1%)<sup>3</sup>, Haemophilus parainfluenzae (2.3%), eukaryotic DNA (88.6%)<sup>3</sup> |
| KM2    | Patient A | punctate (fluid)         | Negative                         | eukaryotic DNA (100%)                      |
| KM3    | Patient A | punctate (fluid)         | Negative                         | Sediminibacterium salmonae (0.3%), eukaryotic DNA (99.7%)<sup>3</sup> |
| KM4    | Patient A | punctate (fluid)         | Negative                         | Gemella sanguinis (1.3%), Haemophilus parainfluenzae (1.0%), eukaryotic DNA (97.7%)<sup>3</sup> |
| KM5    | Patient A | punctate (fluid)         | Negative                         | Herminimonas sp. (10.5%), Propionibacterium acnes (9.7%)<sup>3</sup>, Moraxella catarrhalis (7.5%), eukaryotic DNA (72.3%)<sup>3</sup> |
| KM6    | Patient B | pus                      | Negative                         | Streptococcus intermedia (100%)            |
| KM7    | Patient C | biopsy (tissue)          | Negative                         | eukaryotic DNA (100%)                      |
| KM8    | Patient C | biopsy (tissue)          | Negative                         | No identification                          |
| KM9    | Patient D | joint puncture (fluid)   | Negative                         | Enhydrobacter aerosaccon (49.8%)<sup>3</sup>, Acinetobacter septicus (18.1%)<sup>3</sup>, Moraxella osloensis (14.0%), Staphylococcus sp. (5.8%), Rhenozierma soli (3.1%), Staphylococcus epidermidis (2.6%), Psychrobacter sp. (2.4%)<sup>3</sup>, Propionibacterium acnes (1.3%)<sup>3</sup>, Alkanindiges sp. (0.6%), Acinetobacter sp. (0.4%)<sup>3</sup>, Chryseobacterium sp. (0.3%)<sup>3</sup> |
| KM10   | Patient D | joint puncture (fluid)   | Negative                         | No identification                          |
| KM11   | Patient D | biopsy (tissue)          | Negative                         | Propionibacterium acnes (9.8%)<sup>3</sup>, Bacillus nealsomii (6.7%)<sup>3</sup>, Pseudomonas fluorescens (0.6%)<sup>3</sup>, eukaryotic DNA (82.9%)<sup>3</sup> |
| KM12   | Patient D | biopsy (tissue)          | Negative                         | eukaryotic DNA (100%)                      |
| KM13   | Patient D | biopsy (tissue)          | Negative                         | Undinibacterium olgocarbonphilum (3.5%)<sup>3</sup>, Propionibacterium acnes (0.7%)<sup>3</sup>, eukaryotic DNA (95.9%)<sup>3</sup> |
| KM14   | Patient D | biopsy (tissue)          | Negative                         | Propionibacterium acnes (1.1%)<sup>3</sup>, eukaryotic DNA (98.6%)<sup>3</sup> |
| KM15   | Patient D | biopsy (tissue)          | Negative                         | Veillonella parvula (0.9%), eukaryotic DNA (99.1%)<sup>3</sup> |
| KM16   | Patient D | biopsy (tissue)          | Negative                         | eukaryotic DNA (100%)                      |
| KM17   | Patient E | blood                    | n.d.                             | Bacillus ceras (0.5%)<sup>3</sup>, eukaryotic DNA (99.5%)<sup>3</sup> |
| KM18   | Obduction | formaline captured, biop (tissue) | n.d.                             | Propionibacterium acnes (64.4%)<sup>3</sup>, Staphylococcus epidermidis (25.4%), Paracoccus sanguinis (10.1%)<sup>3</sup> |
| KM19   | Obduction | formaline captured, lung biop (tissue) | n.d.                             | Staphylococcus epidermidis (30.0%), Propionibacterium acnes (34.6%)<sup>3</sup>, Pseudomonas fluorescens (29.4%)<sup>3</sup> |
| KM20   | Patient F | joint puncture (fluid)   | Negative                         | eukaryotic DNA (100%)                      |
| KM21   | Patient F | biopsy (tissue)          | Negative                         | Acinetobacter sp. (18.6%)<sup>3</sup>, Paucibacter sp. (12.8%), Herminimonas arsenicoxidans (5.2%), eukaryotic DNA (63.4%)<sup>3</sup> |

Table 3. Bacterial identification results from 21 clinical orthopedic samples based on culture and NGS of 16S-23S rRNA region. <sup>3</sup>Species present in negative control(s) and regarded as contamination introduced during sample preparation. <sup>3</sup>Genus absent in negative controls but previously reported as contamination of DNA extraction kits, PCR and other laboratory reagents<sup>3</sup>. 

The results of bacterial identification obtained by the four tested methods are shown in Supplementary Table S1. The rates of accurate identification to the species level using NGS of the 16S-23S rRNA region, Sanger sequencing of the 16S rRNA gene, Microflex MS and Vitek MS methods were 92.5%, 56.7%, 73.1% and 64.2%, respectively. The rates of accurate identification to the genus level using NGS of the 16S-23S rRNA region, Sanger sequencing of the 16S rRNA gene, Microflex MS and Vitek MS methods were 100%, 94.0%, 88.1% and 83.6%, respectively. At the species and genus level, assessment of the statistical significance of the differences in accuracy of the four methods for assigning species and genus showed statistically significant differences between NGS of the 16S-23S rRNA region versus all other tested methods for all bacterial isolates (P < 0.05). The only exception was observed between the two sequencing methods at the genus level (P = 0.2482). Moreover, the Sanger sequencing of the 16S rRNA gene was significantly more discriminative at the genus level than Vitek MS (P < 0.05).

NGS of 16S-23S rRNA region was the only method which correctly identified to the species level Shigella sonnei, Streptococcus oralis, Streptococcus mitis, Streptococcus vestibularis, Abiotrophia para-adiacens and

Results
Comparison of identification potential of the tested methods. The results of bacterial identification analysis of large datasets requires a combination of bioinformatics skills and computational resources that is nowadays mostly absent in diagnostic (medical) microbiological laboratories. Furthermore, metagenomics approaches are time consuming as the turnaround time is approximately 4–5 days. To fill the gap between the conventional methods (culture and PCR) and metagenomics, a culture free approach using targeted NGS will be an excellent approach to detect and identify bacterial species as, compared to metagenomics, it is less complicated, and cheaper and therefore more likely to get implemented in diagnostic laboratories within a short timeframe. The aim of the present study was to develop a rapid and easy-to-use culture free diagnostic method based on NGS of PCR amplification products encompassing the whole 16S-23S rRNA region to improve the resolution of bacterial species identification. Moreover, the new method was compared with three commonly used identification methods. Finally, the feasibility of the new identification method to detect and identify bacterial species in clinical specimens was evaluated.
## Sample (Growth CFU/ml) Identification Detection Frequency (n) Average (% reads) Range (% reads) SD

**UR12 (10³)**

| Identification                        | Frequency (n) | Average (% reads) | Range (% reads) | SD |
|---------------------------------------|---------------|-------------------|-----------------|----|
| Proteus mirabilis                     | 3             | 90.0              | 75.5–97.9       | 12.6 |
| Proteus vulgaris                      | 3             | 6.8               | 0.6–17.5        | 9.3  |
| Aerococcus urinae                     | 3             | 1.3               | 1.1–1.5         | 0.2  |
| Undibacterium oligocarboniphilum²    | 1             | 4.8               | n.a.            | n.a. |
| Corynebacterium striatum              | 1             | 0.3               | n.a.            | n.a. |
| Pseudomonas saccharophila             | 1             | 0.3               | n.a.            | n.a. |
| Enterococcus faecalis                | 1             | 0.2               | n.a.            | n.a. |
| *Ralstonia picketti*²                 | 1             | 0.1               | n.a.            | n.a. |
| *UR14 (10²)*                          | Undetected species | 3               | 23.6            | 5.3–42.0 | 18.4 |
| Peptoniphilus lacrimalis              | 3             | 17.9              | 5.9–33.7        | 14.3 |
| *Fenollaria massiliensis*             | 3             | 13.5              | 11.6–15.6       | 2.0  |
| Peptostreptococcus anaerobius         | 3             | 9.5               | 4.4–13.1        | 4.5  |
| Mobiluncus curtisi                    | 3             | 6.0               | 2.1–10.6        | 4.3  |
| Anaerococcus sp.                      | 3             | 3.6               | 1.8–6.6         | 2.6  |
| *Atopobium delae*                     | 3             | 2.8               | 2.3–3.7         | 0.8  |
| Dialister propionicificaciens         | 3             | 0.8               | 0.6–1.2         | 0.3  |
| *Jonquetella anthropi*                | 3             | 0.6               | 0.3–1.1         | 0.4  |
| Tissierella sp.                       | 2             | 2.3               | 2.0–2.6         | 0.4  |
| *Moryella sp.*                        | 2             | 0.8               | 0.1–1.4         | 0.9  |
| Fusobacterium nucleatum               | 2             | 0.5               | 0.3–0.7         | 0.3  |
| Ezakiella sp.                         | 2             | 0.2               | 0.1–0.2         | 0.1  |
| Undibacterium oligocarboniphilum²    | 1             | 36.4              | n.a.            | n.a. |
| *Peptoniphilus kovenoenaene*         | 1             | 4.1               | n.a.            | n.a. |
| Prevotella bivia                      | 1             | 3.4               | n.a.            | n.a. |
| *Pseudomonas saccharophila*           | 1             | 3.1               | n.a.            | n.a. |
| Candidatus Peptoniphilus massiliensis| 1             | 2.6               | n.a.            | n.a. |
| Peptoniphilus harei                   | 1             | 1.1               | n.a.            | n.a. |
| Streptococcus anginosus               | 1             | 0.8               | n.a.            | n.a. |
| *Ralstonia picketti*²                 | 1             | 0.7               | n.a.            | n.a. |
| *Methylobacterium oryzae*²           | 1             | 0.6               | n.a.            | n.a. |
| Streptococcus sp.                     | 1             | 0.6               | n.a.            | n.a. |
| *Asinibacterium lactis*              | 1             | 0.5               | n.a.            | n.a. |
| Olsenella sp.                         | 1             | 0.5               | n.a.            | n.a. |
| *Methylobacterium jeotgali*²         | 1             | 0.4               | n.a.            | n.a. |
| Peptoniphilus duerdenii               | 1             | 0.4               | n.a.            | n.a. |
| Filifactor sp.                        | 1             | 0.2               | n.a.            | n.a. |
| Proteus mirabilis                     | 1             | 0.2               | n.a.            | n.a. |
| Actinotignum sp.                      | 1             | 0.2               | n.a.            | n.a. |
| Bacteroides sp.                       | 1             | 0.2               | n.a.            | n.a. |
| *Shigella flexneri*                   | 1             | 0.1               | n.a.            | n.a. |
| *Calectella sp.*                      | 1             | 0.1               | n.a.            | n.a. |
| Howardella sp.                        | 1             | 0.1               | n.a.            | n.a. |
| Propionibacterium acnes               | 1             | 0.1               | n.a.            | n.a. |
| **UR22 (10³)**                        | Lactobacillus crispatus | 3           | 92.8            | 78.5–100 | 12.4 |
| Undibacterium oligocarboniphilum²    | 1             | 14.4              | n.a.            | n.a. |
| *Pseudomonas saccharophila*           | 1             | 2.3               | n.a.            | n.a. |
| Undibacterium ligocarboniphilum²     | 1             | 1.5               | n.a.            | n.a. |
| Undetected species                    | 1             | 1.3               | n.a.            | n.a. |
| *Ralstonia picketti*²                 | 1             | 1.0               | n.a.            | n.a. |
| *Asinibacterium lactis*              | 1             | 0.4               | n.a.            | n.a. |
| Lactobacillus vaginalis               | 1             | 0.3               | n.a.            | n.a. |
| Staphylococcus capitis                | 1             | 0.2               | n.a.            | n.a. |
| *Methylobacterium oryzae*²           | 1             | 0.1               | n.a.            | n.a. |
| **UR67 (10³)**                        | Escherichia coli | 3             | 66.1            | 62.6–70.1 | 3.9  |
| Aerococcus urinae                     | 3             | 26.8              | 22.5–29.5       | 3.8  |

Continued
**MS9.** In the remaining 4 positive samples (UR1, UR36, UR52 and UR54) bacterial species that were identified were *Staphylococcus aureus* and *Pseudomonas aeruginosa*. Escherichia coli was isolated from 2 samples (UR2, UR59). In the sample UR59 possible misidentification by conventional culture method could occur. In this sample identification by Vitek MS only was improved by NGS of the 16S-23S rRNA region. In the sample UR8 conventional culture methods could not identify *Klebsiella pneumoniae*, but culture also identified *Klebsiella oxytoca*. In the sample UR9, however the NGS-based method also revealed the presence of *Klebsiella pneumoniae*. It was shown very recently, since the year 2014, that *K. oxytoca* and *K. pneumoniae* can be difficult to distinguish due to similar morphology on agar plates. In the sample UR9 possible misidentification by conventional culture methods could occur. In this sample *K. pneumoniae* was identified by Vitek MS and Bruker MS had a problem with identification of *Staphylococcus lentus* to the species level. No method could unambiguously identify *S. pasteurii* to the species level (Table S1).

### Table 4. Reproducibility of NGS of the 16S-23S region.

| Sample (Growth CFU/ml) | Identification                  | Detection Frequency (n) | Average (% reads) | Range (% reads) | SDa                  |
|------------------------|---------------------------------|-------------------------|-------------------|----------------|---------------------|
| UR68 (10³)             | Enterococcus sp.                | 3                       | 8.5               | 7.5–9.2        | 0.9                 |
|                        | Lactobacillus gasseri           | 3                       | 55.5              | 50.9–62.4      | 6.1                 |
|                        | *Escherichia coli*              | 3                       | 36.9              | 34.6–41.8      | 4.3                 |
|                        | Streptococcus pasteuriannus     | 3                       | 5.2               | 2.3–8.7        | 3.2                 |
|                        | Enterococcus sp.                | 3                       | 1.0               | 0.3–2.0        | 0.9                 |
|                        | Veillonella sp.                 | 2                       | 0.2               | 0.2–0.2        | 0.0                 |
| Unidentified species   | 1                               | 3.6                     | n.a.              | n.a.           | n.a.                |
| Facklamia spa          |                                 | 1                       | 0.2               | n.a.           | n.a.                |
| UR69 (10³)             | *Klebsiella pneumoniae*         | 3                       | 77.5              | 73.1–85.5      | 6.9                 |
|                        | *Aerococcus urinae*             | 3                       | 12.8              | 1.8–25.1       | 11.7                |
|                        | *Actinomigium sanguinis*        | 3                       | 8.6               | 1.3–21.6       | 11.3                |
|                        | *Lactobacillus crispatus*       | 3                       | 1.0               | 0.1–2.6        | 1.4                 |

**Urine samples.**  Sixty fresh urine samples from patients suspected for urinary tract infections (UTIs) were subjected to conventional culture-based methods and culture-independent NGS of the 16S-23S rRNA region. Bacterial culture reported the growth of predominant one or two microorganisms at ≥10³ CFU/ml or growth without predominant microorganism was considered inconclusive for UTI. NGS of the 16S-23S rRNA region identified from 1 to 47 different bacterial species in each PCR-positive sample (Table 1). There was an observed association between the increasing number of bacterial genera/species identified using NGS of 16S-23S rRNA region and decreasing CFU value, median: ≥10³ CFU – 2 species/genera; 10⁴ CFU – 5 species/genera; 10⁵ CFU – 5 species/genera; 10⁶ CFU – 20 species/genera. Twenty-nine samples were reported as clinically significant. Among them NGS of 16S-23S rRNA region identified only one bacterial species in 11 samples (UR34, UR35, UR37–UR44, UR50). In the majority of these cases (n = 10) pathogenic species identified by NGS of 16S-23S rRNA region were also identified as the cause of infection by conventional culture methods. The only exception was sample UR44, for which NGS of the 16S-23S rRNA region identified solely *K. pneumoniae*, but culture also identified *Proteus mirabilis* with 10⁴ CFU/ml. In 12 samples, in which 2 or more species were identified by NGS of the 16S-23S rRNA region (UR3–UR5, UR7–UR9, UR33, UR46, UR48, UR49, UR53, UR55), contigs representing the culture identified pathogenic species consisted of the highest number of reads (53.7–99.1% of all reads). For 3 samples (UR2, UR9, UR59) NGS of the 16S-23S rRNA region showed improved identification compared to conventional culture. In sample UR2 conventional identification showed only the presence of *P. mirabilis*, however the NGS-based method also revealed the presence of the genetically related species; *Proteus vulgaris*. In the sample UR9, *Klebsiella oxytoca* was identified by NGS of 16S-23S rRNA region additionally to *E. coli*. Colonies produced by these two microorganisms can be difficult to distinguish due to similar morphology on agar plates. In the sample UR9 possible misidentification by conventional culture method could occur. In this sample *K. pneumoniae* was identified by Vitek MS and *Klebsiella variicola* by NGS of 16S-23S rRNA region. It was shown very recently, since *K. variicola* is closely related to *K. pneumoniae*, it is difficult to distinguish between these two species by commonly used methods, including Vitek MS. In the remaining 4 positive samples (UR1, UR36, UR52 and UR54) bacterial species that were identified by conventional culture methods were the cause of infection were also found by the NGS of 16S-23S rRNA method but were not predominant. Usually, only contigs containing commensal organisms consisted of higher number of reads. The urine samples classified by culture as “no clinical significance” (n = 31) contained numerous bacterial species, which probably represented the commensal flora. PCR-negative samples (n = 13) were only those samples with a growth density of 10² CFU/ml.

**Blood stream infections.**  Among the 23 positive blood culture samples, conventional culture methods and the NGS of the 16S-23S rRNA region approach produced the same identifications for 20 samples (Table 2). For 3 samples (BC11, 19 and 22) identification of bacterial organisms to the species level was improved by the
NGS-based method. In case of sample BC11 conventional culture methods were only able to identify the microorganism to the genus level of *Bacteroides*. In the other two samples, the contigs representing *Streptococcus lutetsiensis* (BC19) and *Bacteroides dorei* (BC22) showed 99.9% (4283/4284) and 100% (4433/4433) sequence homology, respectively, during BLAST analysis of the 16S-23S rRNA region. Only a 16S rRNA sequence for *Streptococcus infantis* was available in the GenBank database showing 95.4% (1413/1481) similarity and for *Bacteroides vulgatus* the whole 16S-23S rRNA region shared only 96.5% (4286/4443) similarity to the obtained contig sequence.

**Orthopedic infections.** Of the 21 clinical samples of orthopedic patients 18 were found to be culture negative and three samples were not cultured. In 5 samples (KM2, KM7, KM12, KM16 and KM20) the NGS-based method was able to detect only eukaryotic DNA and in 2 samples (KM8 and KM10) it yielded non-interpretable results most likely because of template degradation. In the remaining 14 samples the number of microorganisms detected in the orthopedic samples ranged from 1 to 3 different genera/species (Table 3). The only exception was sample KM9 in which 11 different genera/species were found. In our study, orthopedic samples had a low amount of starting material, so they were especially prone to be swamped by the contaminating DNA and result in misleading results. All negative controls contained *Pseudomonas fluorescens*. In a single negative control, *Propionibacterium acnes* was also found. Therefore, in orthopedic samples *P. fluorescens* (present in KM11 and KM19) and *P. acnes* (KM1, KM5, KM9, KM11, KM13, KM14, KM18 and KM19) were regarded as contamination introduced during sample preparation. Moreover, several bacterial genera previously reported as contamination of DNA extraction kits, PCR and other laboratory reagents were absent in all negative controls but present at orthopedic samples, including *Acinetobacter, Bacillus, Chryseobacterium, Enhydrobacter, Paracoccus, Psychrobacter* and *Undibacterium*. These bacteria can also be potentially considered as contaminations of the 16–23S rRNA NGS process introduced during sample processing.

**Reproducibility.** To assess the reproducibility of the method based on NGS of the 16S-23S rRNA region, 6 urine samples were used: 2 samples with bacterial count level at 10^4 CFU/ml, single samples with bacterial count levels at 10^4 and 10^5 CFU/ml, and 2 samples with bacterial count level at ≥10^5 CFU/ml (Table 4). The samples were analyzed in triplicate by 3 different operators, including independent PCR amplification, PCR product purification, and NGS library preparation. Subsequently, two independent MiSeq sequencing runs were conducted. The first replicate of each sample was sequenced in the first run, while the second and third replicate of the tested samples were sequenced in parallel in the second run. When an organism identified was represented by 5% or more reads in at least one replicate of a sample, this organism was always detected in its remaining two replicates (Table 4). The only exception was *Undibacterium oligocarbonophilum*, previously reported as a contaminant in DNA extraction kits, and in PCR and other laboratory reagents. In the samples with bacterial count level at ≥10^5 CFU/ml, the method achieved 100% of reproducibility with respect to the bacterial composition. Moreover, in all replicates of a sample with a bacterial count level at ≥10^5 CFU/ml always the same organism was predominant and represented by 50.9–100% of reads.

**Discussion**

As defined in the CLSI guidelines, a species identification can be assigned when the max score is 99% or higher and if the sequence similarity between best and second best species are greater than 0.5% using DNA target sequencing. However, species identification using CLSI’s criteria for the 16S rRNA gene sequences was often weak because the criteria of distance scores greater than 0.5% to the next closest species was not met for most strains. In such instances, only identification to the genus level was feasible. Therefore, Park et al. proposed a modified CLSI (mCLSI) method which was more practical and pragmatic for identification of species based on 16S rRNA sequences than the CLSI method. The mCLSI method assigns bacterial species when the similarity score is 99% or higher but irrespective of the similarity score differences. In our study, we applied the similarity score differences with the next closest species as ≥0.2%, which reflected at least 3 and 9 nucleotides difference by sequencing the 16S rRNA gene or the 16S-23S rRNA region, respectively. It allowed elimination of misidentification of very closely related species like *S. oralis* and *S. mitis* (Table S1) which in their 16S rRNA gene sequence differ only by 1 nucleotide. Using Sanger sequencing of the 16S rRNA gene, species was assigned with the similarity score 99.4% or higher (Table S1). Using NGS of the 16S-23S rRNA region, in a great majority of cases (*n* = 61), species were assigned with a similarity score above 99.2%. The exceptional 6 species with lower scores showed a similarity ranging from 96.2% to 98.8%. This could be caused by the facts that (i) a limited number of 16S-23S rRNA sequences were available in databases and (ii) the CLSI guidelines were developed for comparison of gene sequences and did not include the intergenic regions which for some genera can be endowed with higher variation. We believe that with increasing number of deposited 16S-23S rRNA sequences, it will be always possible to assign bacterial species with similarity scores of 99% or higher.

We assigned bacteria to the genus level by NGS of 16S-23S rRNA region in clinical samples when the similarity score was at least 90%. This value was determined based on the results produced during comparison of the four identification methods on pure cultures (Table S1). The 16S-23S rRNA sequence alignments produced the lowest identity for second closest match within the same genus for: *Actinobaculum* 88% (Table S1; sample 6), *Lactobacillus* 89% (sample 30), *Corynebacterium* 91% (samples 14, 17 and 18). We applied the uniform cut-off equal to 90% for all bacteria in clinical samples. However, defining 16S-23S rRNA sequences for all or at least majority species within a genus will create interpretive criteria for defining the genus and probably will vary according to the queried microorganism.

The 16S-23S rRNA sequences can differ by length even among highly genetically related species. It is caused by size variation in 16S-23S rRNA intergenic spacer region (ISR). *Staphylococcus aureus* clonal complex 75 has...
been recently renamed as *Staphylococcus argenteus* and now is a novel species of the *Staphylococcus* genus. *S. argenteus* showed identical or nearly identical 16S and 23S rRNA gene sequences to *S. aureus* but differed substantially by the ISR length. The scoring system of BLAST had been designed not to allow large gaps. The BLAST algorithm produced a set of smaller separate alignments, with the longest alignment encompassing the 23S rRNA gene (about 2.5 kb in size). The lack of an alignment for the full-length sequences allowed distinguishing *S. aureus* from *S. argenteus*.

In our study all negative controls contained *P. fluorescens*. We, however, only found this microorganism in the orthopedic samples (Table 3) and not in the urine and blood samples (Tables 1 and 2). Orthopedic samples were characterized with lower starting microbial material compared to that of the urine and blood samples. This showed that the impact of contaminating sequences is greater in low biomass samples. Moreover, *P. fluorescens* had not been associated previously with human infections and its presence can be clearly regarded as contamination. However, in a single negative control, which was introduced to monitor the impact of contaminations on identification procedure of the orthopedic samples, *P. acnes* was also identified. *P. acnes* is a common human skin-associated organism and had been previously shown as the cause of orthopedic infections. It will be highly important to limit the impact of such contaminations as *P. acnes* during samples handling as this is also a clinically significant microorganism and its contribution to an infection can be misinterpreted.

NGS of the 16S-23S rRNA region will provide enhanced information on the presence of microbial DNA within a clinically relevant time frame which is necessary for timely and accurate treatment of infections and therefore key for proper infection and antibiotic resistance control. Data of this study showed that this method correctly identified bacterial species that were identified as the cause of infection by conventional culture methods. In general, in the culture-positive samples a few additional species/genera per sample were identified by the NGS method. However, contigs representing the culture-identified pathogenic bacterial species had the highest number of sequence reads. To assess the clinical relevance of the identified species/genera in samples with a low amount of bacterial DNA, a prospective clinical validation study will be carried out including samples of complex patient groups (e.g. orthopedic patients with a clinical suspicion of a prosthetic joint infection) and samples of control groups of patients without a clinical suspicion of an infection.

Currently, NGS of the 16S-23S rRNA region suffers from two major limitations. First limitation is a lack of reference database with the 16S-23S rRNA sequences and a complementary software allowing easy and reliable species identification. Second major limitation of the method is a lack of reference sequences for many bacterial species. We currently work on the development of a reference data set for assigning clinically relevant bacterial species based on the 16S-23S rRNA sequences. The quality and amount of data accumulated in the databases is particularly important for the performance of bacterial identification using sequencing analysis of the 16S-23S rRNA region. Also, this lack of reference 16S-23S rRNA sequences in the GenBank database might have potentially introduced bias in the results when comparing 16S rRNA with 16S-23S rRNA sequences.

As a proteomic tool for microbial identification, MALDI-TOF MS is superior to NGS-based methods in cost and speed. Currently, the total costs of all reagents and consumables for NGS of 16S-23S rRNA region (not including labor) amount to ~70 € per sample with turnaround time of ≤4 days. However, NGS of the 16S-23S rRNA region was culture independent and more significantly discriminative at the species and genus level than MALDI-TOF MS approaches. Moreover, NGS of 16S-23S rRNA region, unlike all other mass spectrometry methods described above, can be extremely useful for identification of rare or unknown bacteria and bacteria with unusual phenotypic profiles.

In summary, the main objective of the present study was to develop a new diagnostic method based on NGS of 16S-23S rRNA region and assess its identification potential. Its resolution power was found to be superior to the other identification approaches commonly used in routine clinical microbiology laboratories; moreover, the method was easy in use. The method correctly identified urinary tract pathogens and blood stream pathogens previously identified as the cause of UTI and blood stream infection (BSI) with conventional culture. NGS of the 16S-23S rRNA region also showed increased sensitivity in the diagnosis of bacterial microorganisms in samples collected from patients suspected of orthopedic infections. In the analyzed samples several bacterial species had been previously reported as the cause of orthopedic infections, including *Haemophilus parainfluenzae*, *Moraxella osloensis* and *S. epidermidis*. Therefore, we conclude that our approach has the potential to increase diagnostic yield and will decrease time to result for detection of unexpected bacterial pathogens and bacterial species compared to current methods, thereby improving targeted antibiotic treatment. Furthermore, there is a huge potential of this method for detection of bacterial pathogens that can not be cultured at all, due to VNC state or due to antibiotics prior to collection of sample. Finally, with our method it will be possible to streamline processes in the laboratory and to implement it in several disciplines, like clinical, environmental and veterinary microbiology. However, this approach needs further validation and determination of its sensitivity. Furthermore, studies focused on the clinical relevance are necessary for determining the applicability of this NGS-based approach in routine diagnostics.

**Materials and Methods**

**Ethics.** All procedures were carried out according to guidelines and regulations of Certe concerning the use of patient materials for the validation of clinical methods, which are in compliance with the guidelines of the Federation of Dutch Medical Scientific Societies (FDMSS). The project was approved by the Certe medical staff under project submission 3305-0037. Every patient of the Certe is informed that samples taken may be used for research and publication purposes unless they indicate that they do not agree to it. Informed consent was obtained from all individuals or their guardians prior to study participation. All samples were used after performing and completing a conventional microbiological diagnosis and were coded to protect patients confidentiality.
**Bacterial isolates.** Pure cultures of 44 difficult to identify clinically bacterial isolates were included. Additionally, a selection of 23 isolates was used, including 15 ATCC strains (Table S1).

**Clinical samples.** Sixty fresh urine samples from patients suspected of urinary tract infections (UTI), 23 BacT/ALERT (bioMérieux) positive blood culture bottles from patients suspected of BSI and 21 clinical samples of orthopedic patients were collected. Culture was performed as part of routine diagnostics by the department of bacteriology at Certe.

**Vitek MS (bioMérieux).** Vitek MS slides were prepared and interpreted according to the manufacturer’s instructions. Strains which did not yield an identification due to unreliable results or bad spectra were repeated once.

**Bruker MS (Bruker).** Bruker Microflex slides were prepared using on-slide extraction and interpreted according to manufacturer’s instructions. Strains which did not yield an identification due to unreliable results or bad spectra were repeated once.

**Bacterial isolates DNA extraction.** Blood agar plates were used for culturing the bacterial strains. DNA isolation from pure cultures was performed using the UltraClean Microbial DNA Isolation Kit (Mobio) according to the manufacturer’s instructions.

**Urine DNA extraction.** DNA isolation of urine samples was performed using the UltraClean Microbial DNA Isolation Kit (Mobio). Briefly, 500 µl of urine was centrifuged for 30 seconds at 10,000 g. The supernatant was discarded and the pellet reconstituted in 300 µl Microbead solution. Since then the manufacturer’s instructions were followed for DNA extraction.

**Blood culture bottle DNA extraction.** DNA from 1.8 ml from positive blood culture bottles was extracted using the BioOstic Bacteraemia DNA Isolation Kit (Mobio) according to the manufactures instructions.

**DNA extraction of the clinical samples of orthopedic patients.** The Purelink Genomic DNA purification kit (Invitrogen) was used for DNA extraction of the orthopedic samples. Initial lysis was performed using lysis buffer (0.25 M NaCl, 0.5% SDS, 5x TE, 2.25 U/ml Proteinase K (Roche)). For tissue samples a small piece was digested in 200 µl lysis buffer. For fluidic samples 200 µl was added to 200 µl lysis buffer. Digestion was performed in a thermoshaker at 56 °C under light shaking for 18 hours. 200 µl Purelink Genomic lysis/binding buffer was added to 200 µl of lysed sample and vortexed to create a homogenous solution. 200 µl 96% ethanol was added and the DNA purification protocol was followed according to the manufacturer’s instructions.

**16S rRNA gene amplification and Sanger sequencing.** The DNA of the bacterial isolates and controls was amplified with 0.3 µM primers [LPWS57′-AGTTTGATCTGGCTCAG-3′ (nucleotide position 10–27) and LPWS8 5′-AGGCGCCAGGTATTCAC-3′ (nucleotide position 1370–1389)] using GoTaq® Flexi Hot Start Polymerase and Colorless GoTaq® Buffer (Promega). PCR amplification was performed on a PTC-100 thermocycler (Bio-Rad) using the following conditions: an initial incubation at 94 °C for 2 min; 25 cycles of 94 °C for 30 s, 58 °C for 30 s, and 72 °C for 1 min, followed by a final incubation at 72 °C for 5 min. PCR products were purified using DNA Clean & Concentrator (Zymo Research). DNA sequencing was performed by the GATC company (Golgen, Germany) using the PCR primers. 16S rRNA gene sequences were subjected to BLAST analysis against the NCBI nucleotide database. The sequencing data analysis allowed assigning species when the similarity score was 99% or higher and the similarity score differences with the next closest species were equal to or greater than 0.2%, which reflected 3 or more nucleotides. The genus was assigned when the similarity score was 99% or higher and the similarity score differences with the next closest species were equal to or greater than 0.2%, which reflected 3 or more nucleotides. The genus was assigned when the similarity score was 97% or higher. The BLAST analysis of the 16S rRNA gene sequences was verified against the curated leBibi-QBPP database (https://umr5558-bibiserv.univ-lyon1.fr/lebibi/lebibi.cgi) not revealing any differences in identification.

**Amplification of the 16S-23S rRNA region.** The 16S-23S rRNA region was amplified by PCR using forward primer 27F (5′-AGAGTTTGATCMTGGCTCAG-3′) and reverse primer 23S R (5′-GACATCGAGGTGCCAAAC-3′), specific for the 16S rRNA gene and adapted from the study by Sergeant et al.20, and reverse primer 2490R (5′-GACATCGAGGTGCCAAAC-3′), specific for the 23S rRNA gene and slightly modified by truncation of a single nucleotide compared to the original publication by Hunt et al.21. The amplification of the 16S-23S DNA region was carried out in a 25 µl reaction consisting of 1x Phire hotstart buffer (Thermofisher), 5 mM dNTPs (Roche), 0.5 µl Phire hotstart II DNA polymerase (Thermofisher), 600 nM of each primer and 5 µl of DNA template. PCR was performed using a Biorad PTC-200 thermocycler. An initial denaturation of 98 °C at 30 sec was followed by 25 cycles of 98 °C for 30 sec, 66 °C for 30 sec and 72 °C for 2 minutes with a final extension of 72 °C for 1 min. To increase PCR sensitivity in clinical samples the number of cycles was increased to 35. PCR products were purified using the Qiaquick PCR purification kit (Qiagen) according to the manufacturer’s instructions.

**NGS library preparation and Illumina MiSeq sequencing.** For library preparation, the Nextera XT DNA Sample Preparation Kit (Illumina) was used according to the manufacturer’s instructions. Briefly, the purified PCR amplicons quantified with a Qubit 2.0 Fluorometer (Thermofisher) were diluted to 0.2 ng/µl and a total of 1 ng of DNA was tagmented at 55 °C for 5 min. PCR amplification to introduce Illumina index sequences was performed in PCR strip tubes in a BioRad T100 thermocycler. Size distribution of fragments was estimated with a 2200 TapeStation using the Agilent DNA 1000 High Sensitivity kit according to the manufacturer’s instructions. Fragments of 200 to 1000 bp were obtained. The library DNA fragments were size selected and purified using AMPure XP beads (Beckman Coulter, Inc.). The indexed libraries were normalized, pooled and loaded onto an
Illumina MiSeq reagent cartridge using MiSeq reagent kit v3 and 600 cycles. The paired-end 2 × 300 bp sequencing was run on an Illumina MiSeq sequencer.

**Sequencing of the 16S-23S rRNA region and data analysis.** NGS generated 25,000–50,000 or 1–2 million sequencing reads for pure culture or clinical sample, respectively, to obtain a minimum coverage of 1000 per sample. The 300-nucleotide paired-end reads were de novo assembled into contigs with the SeqMan NGen software (DNASTAR) using parameters: mer size 31 nucleotides and minimum match percentage 93%. The sizes of resulting contigs produced during analysis of pure cultures ranged from 4183 bp (Bacillus cereus) up to 4856 bp (Acinetobacter parvus) with an average size of 4423 bp (Table S1). In case of clinical samples, most often an identified bacterial species/genus was represented by a single contig of expected size, around 4.5 kb. However, in some instances only smaller contigs (ranging between 1 and 3 kb in size) represented the same bacterial species/genus present in a sample. In those cases all contigs belonging to the same organism were combined and their reads were added up. Species identification was based on alignment of contig sequences with 16S-23S rRNA sequences deposited in the GenBank database using nucleotide BLAST (Basic Local Alignment Search Tool, http://www.ncbi.nlm.nih.gov/BLAST/). When a reference 16S-23S rRNA sequence was not available in the database, a reference 16S rRNA gene sequence was used for a species identification. The contig sequences were submitted via the website and bacterial species was assigned when the similarity score was 99% or higher and the similarity score differences of the first match with the next closest species was equal to or greater than 0.2%. This reflected a 9 or more nucleotides difference for a sequence of 4423 bp (the average size of the 16S-23S rRNA ampiclon). When the similarity score was between 90% and 99%, the genus could be assigned. The score below 90% was interpreted as an unidentified organism.

**Statistical analysis.** The four bacterial identification methods were compared using McNemar's test, a test of paired proportions. Only a single strain per species was taken (from Table S1) for calculation of the accuracy of bacterial identification at the species or genus level according to McNemar's test. When the P value was less than 0.05, we concluded that there was a significant difference between the methods. Statistical analysis was performed using the GraphPad software.

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**Author Contributions**
A.M.D.K.-S., E.Z., G.W., K.S. and R.F.B. designed the project, provided the strains, clinical material and epidemiological data. A.J.S., V.A., E.Z., G.W., and K.S. performed the experiments. A.J.S. and E.Z. carried out de novo assemblies. All authors interpreted the data. A.J.S. wrote the manuscript. All authors reviewed the manuscript.

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