The emergence of a novel sequence type of MDR Acinetobacter baumannii from the intensive care unit of an Egyptian tertiary care hospital

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Abstract

Background and aim of work: Acinetobacter baumannii is known for nosocomial outbreaks worldwide. In this study, we aimed to investigate the antibiotic susceptibility patterns and the clonal relationship of A. baumannii isolates from the intensive care unit (ICU) of an Egyptian hospital.

Methods: In the present study, 50 clinical isolates of multidrug resistant (MDR)-A. baumannii were obtained from patients admitted into the ICU from June to December 2015. All isolates were analyzed for antimicrobial susceptibilities. Multiplex PCR was performed to detect genes encoding oxacillinase genes (blaOXA-51-like, blaOXA-23-like, blaOXA-24-like, and blaOXA-58-like). Multilocus sequence typing (MLST) based on the seven-gene scheme (gltA, gyrB, gdhB, recA, cpn60, gpi, rpoD) was used to examine these isolates.

Results: All A. baumannii clinical isolates showed the same resistance pattern, characterized by resistance to most common antibiotics including imipenem (MIC ≥ 8μg/mL), with the only exception being colistin. Most isolates were positive for blaOXA-51-like and blaOXA-23-like (100 and 96%, respectively); however, blaOXA-24-like and blaOXA-58-like were not detected. MLST analysis identified different sequence types (ST195, ST208, ST231, ST441, ST499, and ST723) and a new sequence type (ST13929) with other sporadic strains.

Conclusions: MDR A. baumannii strains harboring blaOXA-23-like genes were widely circulating in this ICU. MLST was a powerful tool for identifying and epidemiologically typing our strains. Strict infection control measures must be implemented to contain the worldwide spread of MDR A. baumannii in ICUs.

Keywords: MDR-A. baumannii, blaOXA-23-like, MLST

Background

The clinical care of intensive care unit (ICU) patients with infections has been complicated by the emergence and spread of extremely drug-resistant (XDR) Acinetobacter baumannii strains [1]. Due to scarce current therapeutic options; higher infection rates; poor patient outcomes because of life-threatening infections, including ventilator-related pneumonia, sepsis, urinary tract infections, and skin and soft tissue disorders may occur [1, 2].

Extensively drug-resistant (XDR) A. baumannii strains exhibiting resistance to three or more antibiotic classes, except for polymyxins, have been recently described in nosocomial outbreaks [2, 3]. The essential role of A. baumannii resistance to carbapenems, is mediated by oxacillinas (OXA-class D) and, less frequently, by metallo-β-lactamases (MBL-class B) [4, 5]. The class D carbapenemases are the most predominant carbapenemases in A. baumannii. They are categorized into six subclasses:
intrinsic chromosomal OXA-51-like, the acquired OXA-23-like, OXA-24/40-like, OXA-58-like, OXA-143-like, and OXA-235-like β-lactamases [6]. In this study, we aimed to investigate the antimicrobial susceptibility, class D carbapenemases and clonal relationship of A. baumannii strains isolated from a tertiary care hospital ICU in Egypt.

Methods
The study was carried out in EL Sheikh Zayed hospital which provides tertiary care from specialists and consultants after referral (in orthopedic, trauma, neuro/spine surgeries) from primary care and secondary care hospitals in Egypt. A lab-based surveillance was performed over a period of 6 months (June–December 2015) after the isolation of five MDR A. baumannii strains in a period of 1 week showing the same phenotypic characteristics.

Bacterial strains
All clinical samples of the patients admitted during the above-mentioned period were processed at the microbiology unit. All samples were cultured on blood agar and MacConkey agar (Oxoid Co. England). All culture plates were incubated aerobically at 35 °C for 24–48 h. Identification of isolated organisms was performed by conventional biochemical reactions. During the experimental period, 50 A. baumannii non-duplicate strains were isolated.

Antimicrobial susceptibility testing
Susceptibility testing was performed by the disc diffusion method (Modified Kirby-Bauer technique) using Mueller–Hinton agar and aerobic incubation at 35 °C for 16–18 h. Antimicrobial discs containing imipenem (10 μg), meropenem (10 μg), gentamicin (10 μg), ciprofloxacin (5 μg), amikacin (30 μg), cefotaxime (30 μg), amoxicillin/clavulanic acid (30/10 μg), aztreonam (30 μg), cefazidime (30 μg), ceftazidime/clavulanic acid (30/10 μg), amoxicillin/clavulanic acid (20/10 μg), and cefoxitin (30 μg) were obtained from Oxoid Co. (Oxoid Limited, Basingstoke, Hampshire, England) [7].

Multidrug resistance was defined in this analysis as resistance to three or more representatives of the following classes of antibiotics: fluoroquinolones, extended-spectrum cephalosporins, aminoglycosides, and carbapenemases [8].

Escherichia coli ATCC 25922, Pseudomonas aeruginosa ATCC 27853, and Staphylococcus aureus ATCC 29213 were used as reference strains for susceptibility testing per Clinical and Laboratory Standards Institute (CLSI, 2015) guidelines and interpretations [7].

Minimum inhibitory concentrations (MICs) were determined by broth microdilution and interpreted using CLSI, 2015 guidelines [7].

The presence of A. baumannii genes encoding oxacillinases (blaOXA-23-like, blaOXA-24-like, blaOXA-51-like, and blaOXA-58-like) was assessed in all 50 isolates using multiplex PCR.

Multiplex PCR assay
The sequences of blaOXA alleles encoding carbapenemases were aligned and group-specific regions were identified using BioEdit software (http://www.mbio.ncsu.edu/BioEdit/bioedit.html). The primers: 5′-TAA TGC TTT GATCGG CCT TG and 5′-TGG ATT GCA CTT CAT CTT GG were used to amplify a 353 bp fragment of genes encoding the intrinsic OXA-51-like enzymes of A. baumannii [9].

A set of primers were designed to amplify OXA-23-like genes (501 bp: 5′-GAT CGG ATT GGA ACA CCAGA and 5′-ATT TCT GAC CGC ATT TCC AT), OXA-24-like genes (246 bp: 5′-GGT TAG TTG GCC CCC TTA AA and 5′-AGT TGA GCC AAA AGG GGA TT), and OXA-58-like genes (599 bp: 5′-AAG TAT TGG GGC TTG TGC TG and 5′-CCCCTCTGGCCTACATAC) [9]. The primers were evaluated separately against control strains and then in a multiplex format. The amplification conditions were: initial denaturation at 94 °C for 5 min, 30 cycles of 94 °C for 25 s, 52 °C for 40 s, and 72 °C for 50 s, and a final elongation at 72 °C for 6 min [9].

Multilocus sequence typing
MLST analysis was performed per the protocol of the Pasteur Institute. Fragments of seven internal housekeeping genes (gltA, gyrB, gdhB, recA, cpn60, gpi, and rpoD) were amplified and sequenced as previously described [10]. Briefly, PCR amplifications were performed with a MasterCycler Nexus (Eppendorf, Hamburg, Germany) with an initial denaturation at 94 °C for 5 min, followed by 35 cycles of denaturation at 94 °C for 1 min, annealing at 55 °C for 1 min, and extension at 72 °C for 2 min, and a 4-min final extension at 72 °C. The amplicons were verified by agarose gel electrophoresis and were subsequently purified for bidirectional Sanger sequencing reactions. Multiple allele sequences were assigned for each locus with an arbitrary allele number to obtain characterization of sequence types (STs) for each A. baumannii isolate. Each sequence was compared with sequences deposited in the Institute of Pasteur MLST schema (http://pubmlst.org/perl/bigsdb/bigsdb.pl?db=pubmlst_abbaumannii_pasteur_seqdef).

Results
Out of 358 patients admitted to ICU from June to December 2015, 56 (15.6%) patients were diagnosed with various types of hospital-acquired infections (HAI) as shown in Table 1. A total of 50 non-duplicate A. baumannii strains...
were isolated from different patient samples. Phenotypic antibiotic susceptibility testing for all A. bauman-
nii isolates showed the same drug resistance pattern, characterized by resistance to all antibiotics used including imipenem, except for colistin. Genotypic analysis of \( bla_{OXA-51}\)-like, \( bla_{OXA-23}\)-like, \( bla_{OXA-24}\)-like, and \( bla_{OXA-58}\)-like genes by multiplex PCR (Fig. 1) showed that \( bla_{OXA-51}\)-like and \( bla_{OXA-23}\)-like were the most prevalent genes with 100 and 96% prevalence, respectively. However, \( bla_{OXA-24}\)-like and \( bla_{OXA-58}\)-like were not detected in the current study. Multilocus sequence typing (MLST) of the 50 clinical isolates of A. bauman-
nii has yielded different sequence types; ST195, ST208, ST231, ST441, ST499, and ST723. Interestingly, a new sequence type ST13929 was identified among A. bau-
mannii clinical isolates as shown in Table 2 and Fig. 2. A. baumannii ST13929 has been isolated from a young male (21 years old) patient who had no history of overseas travel. He was admitted to ICU at El Sheikh Zayed Specialized Hospital, Giza, Egypt on 5th of December 2014. The patient had multiple traumas due to motor car accident. After 7 days of ventilation the patient diagnosed to have ventilator associated pneumonia (VAP). Empirical antibiotic therapy of intravenous ceftriaxone/cefotax-
time had been initiated. Endotracheal aspiration has been cultured on blood, chocolate and MacConkey agars. The recovered colonies had been identified as A. baumannii. Further genotypic identification was done by restriction analysis of 16S–23S rRNA spacer sequences using \( Alu I \) and \( Nde II \). The isolate exhibited XDR towards imipenem (MIC > 32 mg/L), meropenem (MIC > 32 mg/L), ceftazi-
dime (>256 mg/L), cefepime (MICs > 256 mg/L), gentamicin (MICs > 256 mg/L), amikacin (MICs > 256 mg/L), and ciprofloxacin (MICs > 32 mg/L). Tigecycline suscept-
ibility was observed at MIC of 1 mg/L. The antimicrobial therapy was changed to tigecycline on day 11. The patient had spent 107 days in the hospital. The patient was alive after the hospitalization period. There was an ongoing XDR- A. baumannii outbreak in the institution in the same period and multiple isolates had been investigated.

**Discussion**

MDR A. baumannii is a problematic, multidrug-resistant pathogen identified in healthcare settings worldwide, especially in ICUs [12]. A. baumannii has a notable abil-
ity to capture and express resistance genes. All resist-
ance mechanisms including target modification, efflux pump expression, and enzymatic inactivation have been described in A. baumannii [13].

In the current study, five MDR A. baumannii strains were isolated over 1 week from the same ICU. All isol-
ates showed the same phenotypic characteristics which prompted us to start a survey study of the antimicrobial susceptibility and clonal relationship of A. baumannii strains isolated from this ICU.

All our isolates were resistant to imipenem. The main role of the A. baumannii resistance to carbapenems is mediated by oxacillinases and, less frequently, by metalloc-
\( \beta \)-lactamases [4, 5].

\( bla_{OXA-23}\)-like, \( bla_{OXA-24/40}\)-like, and \( bla_{OXA-58}\)-like genes have been repetitively reported in A. baumann-
nii outbreaks from diverse parts of the world. The

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**Table 1 Patient’s data**

| Age (mean ± SD) | Sex % | APACHE % | Length of stay (mean ± SD) | More than one device inserted % |
|-----------------|-------|----------|-----------------------------|---------------------------------|
|                 | Male  | Female   | <15% | >15% |                           |                                 |
| VAP             | 42.62 ± 17.63 | 72%  | 28% | 40% | 60% | (48.96 ± 77.3) | (100%) |
| CLABSI          | 41 ± 23.4 | 88.8% | 11.1% | 33.3% | 66.6% | (22.33 ± 11.4) | (55.55%) |
| CAUTI           | 44.53 ± 22.1 | 76.9% | 23.0% | 38.4% | 61.5% | (82.307 ± 67.8) | (100%) |
| P value         | 0.900 | 0.535 | 0.880 | 0.131 | 0.00 |                                  |                                 |

VAP: ventilator associated pneumonia, CLABSI: central line associated blood stream infection, CAUTI: catheter associated urinary tract infection, APACHE: acute physiology and chronic health evaluation.

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**Fig. 1** Results of multiplex PCR for detection of \( bla_{OXA-51}\)-like, \( bla_{OXA-23}\)-like, \( bla_{OXA-24}\)-like, and \( bla_{OXA-58}\)-like genes. Lane 1 100 bp DNA Ladder, Lane 2 positive control, Lanes 3–6 A. baumannii clinical isolates showing \( bla_{OXA-51}\)-like and \( bla_{OXA-23}\)-like positivity (353 and 501 bp, respectively). \( bla_{OXA-24}\)-like and \( bla_{OXA-58}\)-like were not detected at (246 and 599 bp) respectively. Lane 7 negative control.
Table 2  Sequence types (STs), allele profiles of 50 carbapenem-resistant *A. baumannii* isolates, carbapenem-hydrolyzing class D β-lactamase genes, minimum inhibitory concentration (MIC), site of isolation and patient outcome

| Sample ID | Site of isolation | Allele profile | ST | Carbapenem-hydrolyzing class D β-lactamase genes | MIC R ≥ 8 (mg/L) | Patient outcome |
|-----------|-------------------|----------------|----|--------------------------------------------------|------------------|----------------|
| A. baumannii | ETA 21 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  96  gpi  3  rpoD  195 | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 22 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  96  gpi  3  rpoD  195 | + | + | − | − | 20 | Deceased |
| A. baumannii | ETA 30 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  208 | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 33 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  208 | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 34 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  208 | + | + | − | − | 35 | Deceased |
| A. baumannii | ETA 36 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  208 | + | + | − | − | 20 | Deceased |
| A. baumannii | ETA 37 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  208 | + | + | − | − | 30 | Deceased |
| A. baumannii | Blood 44 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  208 | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 46 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  208 | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 47 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  208 | + | + | − | − | 30 | Discharged |
| A. baumannii | ETA 49 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  208 | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 5 | ghtA  12  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  1114 | + | + | − | − | 20 | Deceased |
| A. baumannii | ETA 29 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  13929a | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 24 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 25 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 38 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 30 | Deceased |
| A. baumannii | ETA 48 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 25 | Deceased |
| A. baumannii | ETA 4 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 35 | Discharged |
| A. baumannii | ETA 8 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 35 | Deceased |
| A. baumannii | ETA 11 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 20 | Discharged |
| A. baumannii | ETA 14 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 20 | Discharged |
| A. baumannii | ETA 16 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 25 | Discharged |
| A. baumannii | ETA 18 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 20 | Discharged |
| A. baumannii | ETA 31 | ghtA  3  gyrB  3  gdhB  2  recA  2  cpn60  97  gpi  3  rpoD  195 | + | + | − | − | 25 | Discharged |
Table 2 continued

| Sample ID | Site of isolation | Allele profile | ST | Carbapenem-hydrolyzing class D \( β \)-lactamase genes | MIC R ≥ 8 (mg/L) | Patient outcome |
|-----------|------------------|----------------|----|---------------------------------|-----------------|----------------|
|           |                  |                |     | \( \text{bla}_{OXA-51} \) | \( \text{bla}_{OXA-23} \) | \( \text{bla}_{OXA-24} \) | \( \text{bla}_{OXA-58} \) | IMI |                      |
| A. baumannii | ETA 32         | 10 12 4 11 4 98 5 231 |     | +      | +      | 20 Deceased |
| A. baumannii | ETA 35         | 10 12 4 11 4 98 5 231 |     | +      | +      | 25 Deceased |
| A. baumannii | ETA 40         | 10 12 4 11 4 100 5 441 |     | +      | +      | 25 Deceased |
| A. baumannii | Blood 41       | 10 12 4 11 4 100 5 441 |     | +      | +      | 25 Deceased |
| A. baumannii | ETA 45         | 10 12 4 11 4 98 5 231 |     | +      | +      | 20 Deceased |
| A. baumannii | ETA 12         | 12 17 12 1 29 102 39 236 |     | +      | +      | 35 Deceased |
| A. baumannii | ETA 17         | 12 17 12 1 29 102 39 236 |     | +      | +      | 35 Deceased |
| A. baumannii | ETA 2          | 24 92 96 11 49 162 26 499 |     | +      | +      | 20 Discharged |
| A. baumannii | ETA 4          | 24 92 96 11 49 162 26 499 |     | +      | +      | 25 Deceased |
| A. baumannii | ETA 7          | 24 92 96 11 49 162 26 499 |     | +      | +      | 20 Deceased |
| A. baumannii | ETA 13         | 24 92 96 11 49 162 26 499 |     | +      | +      | 15 Deceased |
| A. baumannii | ETA 15         | 24 92 96 11 49 162 26 499 |     | +      | +      | 15 Deceased |
| A. baumannii | ETA 39         | 24 92 96 11 49 162 26 499 |     | +      | +      | 35 Deceased |
| A. baumannii | Blood 42       | 24 92 96 11 49 162 26 499 |     | +      | +      | 25 Deceased |
| A. baumannii | Swab 50        | 24 92 96 11 49 162 26 499 |     | +      | +      | 20 Discharged |
| A. baumannii | Blood 43       | 28 38 45 1 16 66 2 1089 |     | +      | +      | 25 Deceased |
| A. baumannii | ETA 9          | 44 73 No gene 11 44 + 4 NA |     | +      | +      | 40 Deceased |
| A. baumannii | ETA 10         | 44 73 No gene 11 44 + 4 NA |     | +      | +      | 20 Discharged |

\( \text{ETA} \) endotracheal aspirate, + positive, − negative

\( ^a \) New sequence type (ST13929)

\( ^b \) gpi 173, 1 difference found. 33T → 33C
localization of numerous β-lactamase genes on plasmids facilitates their horizontal mobilization from one bacterium to another [13, 14].

All our isolates harbored the bla_{OXA-51}-like gene, which is ubiquitous in A. baumannii [15]. bla_{OXA-23} was the most universally identified gene, while bla_{OXA-24}-like and bla_{OXA-58}-like genes were not detected in any strain. bla_{OXA-23} is the most prevalent carbapenemase-encoding gene in the Mediterranean region. This might be explained by the higher carbapenemase activity of bla_{OXA-23} and/or acquisition of carbapenem resistance through horizontal gene transfer [16–18]. The bla_{OXA-23} gene was either encoded on the chromosome or on plasmids and was associated with four dissimilar genetic structures, with the most common being transposons Tn2006. bla_{OXA-23} has been reported in different regions of the Middle East, The United Arab Emirates, Algeria, Libya, Bahrain, and recently, Qatar [16, 19].

Mugnier et al. found an isolate from Egypt harboring plasmid containing bla_{OXA-23}. This finding might indicate the prevalence of the genetic environment of bla_{OXA-23} in Egyptian isolates [16]. Moreover, a recent study including three Egyptian hospitals revealed the emergence and spread of bla_{NDM-1} and bla_{OXA-23} in addition to the co-occurrence of 16S rRNA methylase armA with bla_{NDM-1} and bla_{OXA-23} in 27 distinct

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**Fig. 2** Phylogenetic tree of 50 A. baumannii strains built based on the maximum likelihood algorithm in the MEGA6 software [11] and the concatenated alleles of seven housekeeping genes with bootstrap values. The phylogenetic analysis identified ST195, ST208, ST231, ST441, ST499, and ST723 and the new sequence type ST13929. The numbers in the branches depict the sample ID of A. baumannii.
sequence types, 11 of which were novel among *A. baumannii* clinical isolates [20].

Per MLST results, ST195, ST208, ST231, ST441, ST499, and ST723 were the most prevalent isolates. Another Egyptian study illustrated the large diversity found within the strains where ten distinct sequence types (STs) were identified, ST408–ST414, ST331, ST108, and ST208 [21]. However, a study showed that the most prevalent sequence types in the gulf area were ST195, ST208, ST229, ST436, ST450, ST452, and ST499 [22].

Taking into consideration that ST208 is the ancestor strain of several STs including ST89, ST88, ST190, ST225, and ST75, it has been identified in different parts of the world such as Japan, China, Thailand, Korea, Italy, Australia, Portugal, and the Czech Republic [23].

In conclusion, MDR *A. baumannii* strains harboring the *bla*OXA-23*-like* gene were widely circulating in our ICU. MLST provided us with a powerful tool for identifying and epidemiologically typing our strains. Studying the epidemiology of HAIs is urgent to prevent the clonal dissemination of antibiotic-resistant pathogens, not only in hospital settings, but in the community, as well. Strict infection control measures and antimicrobial stewardship programs are necessary to contain the worldwide spread of MDR *A. baumannii*. Proving the clonal relation between clinical isolates emphasizes the importance of surveillance programs and strict IC measures that would influence decision-making and health policy.

**Authors’ contributions**

DG, and MA conceived and designed the study, carried out the collection of the bacterial strains, participated in antibiotic sensitivity and molecular genetic studies. DG drafted the manuscript. MZ participated in antibiotic sensitivity, and MHA. EA and RB carried out the MLST and participated in the design of the study. OA participated in the antimicrobial sensitivity testing and multiplex PCR. All authors read and approved the final manuscript.

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**Competing interests**

The authors declare that they have no competing interests.

**Availability of data and materials**

The raw data of this research can be made available to the interested researchers if requested.

**Ethics approval**

Ethical Committee of faculty of medicine, Cairo University approved the study and a written informed consent was not obtained from patients because the bacterial isolates studied were collected from the routine work of microbiology laboratory for patient care and no additional clinical specimens were collected for the study. It is a standard practice not to get written informed consent for use of bacterial isolates unrelated to patient identity from the routine clinical laboratory.

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