Evaluation of Multilocus Variable-Number Tandem-Repeat (MLVA-8\textsubscript{Orsay}) for Typing of Carbapenem-Resistant \textit{Acinetobacter baumannii} Isolated from Patients in Tehran, Iran

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Abstract

Objectives: \textit{Acinetobacter baumannii} has emerged worldwide as a major nosocomial pathogen. The aim of this study was to evaluate the MLVA-8\textsubscript{Orsay} scheme suitability for clustering and differentiating \textit{A. baumannii} strains isolated from two hospitals in Tehran.

Methods: A panel of eight variable-number tandem-repeat (VNTR) markers was used for typing analysis (MLVA-8\textsubscript{Orsay}) of 89 carbapenem-resistant \textit{A. baumannii} strains isolated from two hospitals in Tehran.

Results: \textit{Abaum0845} (S-VNTR marker) was excluded from the MLVA scheme because of its great allelic diversity at this locus. The modified MLVA-7 scheme provided excellent discriminatory power and detected 82 MLVA types among the 83 isolates. Additionally, the great diversity associated with MLVA markers did not permit to apply the previously proposed criteria for clustering the isolates. However, six MLVA complexes were defined assuming a 71% genetic similarity cut-off, corresponding to the existence of identically sized alleles for at least five VNTR markers. The \textit{blaOXA-23}-like gene (93.2\%) was the most prevalent gene in resistant isolates collected from the two hospitals.

Conclusions: MLVA-7 discriminatory power was found to be excellent. This method was also appeared to be useful and reliable in epidemiological investigations of \textit{A. baumannii} strains. MLVA-7 analysis revealed that carbapenemase-carrying \textit{A. baumannii} isolates might be polyclonal and provided evidence for gene transfer among different strains collected from the two hospitals. Overall, the study provided evidence supporting the use of MLVA-7 scheme as an alternative or complementary method for typing of \textit{A. baumannii} isolates.

Keywords: \textit{Acinetobacter baumannii}, MLVA-8\textsubscript{Orsay}, Carbapenem Resistance

1. Background

\textit{Acinetobacter baumannii} has emerged worldwide as a major nosocomial pathogen and it is a great threat to hospitalized patients, particularly intensive care unit (ICU) patients (1, 2). Because of its ubiquitous nature, \textit{A. baumannii} is isolated from hospital environments and clinical specimens. Multidrug resistance (MDR), extensively drug resistance (XDR), involvement in hospital outbreaks, and long-term persistence in hospital settings are the hallmarks associated with \textit{A. baumannii} infections (1).

An unambiguous identification method is needed for \textit{A. baumannii} isolates in order to identify the genetic relatedness and the population structure of \textit{A. baumannii} strains, as well as to study the impact of public health measures on controlling the MDR infections. Until now, several typing methods have been developed in order to determine the genetic diversity, strains’ relatedness, and the epidemiology of \textit{A. baumannii}. Among them, multilocus sequence typing (MLST) and pulsed-field gel electrophoresis (PFGE) are common methods being used in the molecular typing of \textit{A. baumannii} (1, 3). MLST is a nucleotide sequence-based method used for unambiguous characterization of bacterial isolates. MLST provides an allelic profile or a sequence type for each isolate, which can be shared among the laboratories and throughout the world via the Internet. However, this method is expensive and inappropriate for the characterization of the isolates in the outbreaks (1, 4). In contrast, PFGE is considered the gold standard method in the molecular epidemiology of outbreaks. However, PFGE is labor-intensive, which takes several days to produce results and needs specific tools not available in all laboratories (1, 3). Therefore, in this study, a new approach of multiple-locus variable-number tandem-repeat (VNTR)
3.2. Antimicrobial Susceptibility Testing

Antimicrobial susceptibility testing was performed for all of the 89 strains of *A. baumannii* by the disk diffusion method, as recommended by the Clinical and Laboratory Standards Institute (CLSI) guidelines. The test antibiotics were as follows: imipenem (10 µg), meropenem (10 µg), gentamicin (10 µg), ciprofloxacin (5 µg), amikacin (30 µg), trimethoprim-sulfamethoxazole (23.75 + 1.25 µg), cefepime (30 µg), cefotaxime (30 µg), aztreonam (30 µg), and cef-tazidime (30 µg). *Escherichia coli* ATCC 25922 was used as a quality control strain. The minimum inhibitory concentrations (MICs) of imipenem and meropenem were determined using the agar dilution method (8). Strains resistant to at least three classes of antibiotics were considered MDR strains (9).

3.3. Molecular Detection of OXA Carbapenemase and Metallo-β-Lactamase

Four groups of OXA carbapenemase genes were detected by a multiplex PCR assay, including *bla*_{OXA-23}-like, *bla*_{OXA-24}-like, *bla*_{OXA-25}-like, and *bla*_{OXA-58}-like genes (10). Ambler class B carbapenemase genes were detected by PCR and using specific primers for *bla*_{IMP} and *bla*_{VIM} (11).

3.4. MLVA Genotyping

Genomic DNA was prepared by the boiling method (12). The MLVA-8_{Orsay} scheme was performed as described previously in a study by Pourcel et al. (6). Oligonucleotide primers targeting the flanking regions of the L-repeat VNTRs (Abaum1988, Abaum2240, Abaum3002, and Abaum3530) and S-repeat VNTRs (Abaum0826, Abaum0845, Abaum2396, and Abaum3468) were used for the amplification of *A. baumannii* genomic DNA by PCR. L (Larg)-repeat VNTR loci were the loci with a repeat unit size of above 9 bp, whereas small (S)-repeat VNTRs were the loci with a repeat unit size of up to 9 bp. The low level of polymorphism was observed in the L-repeat VNTR loci due to their slow evolving, whereas S-repeat VNTRs were very diverse. The clustering was achieved by combining L-repeat and S-repeat VNTRs (6). PCRs were performed in a 25 µL final volume containing 3 ng genomic DNA, × 1´ reaction buffer, 1.5 mM MgCl₂, 1U of Taq DNA polymerase, 200 µM of each deoxynucleoside triphosphate (dNTPs), and 20 µM of each primer. Amplification was performed with a Bio-Rad thermal cycler. PCR conditions were as follows: denaturation for 5 minutes at 94°C, followed by 36 cycles of denaturation for 30 seconds at 94°C, annealing for 30 seconds at 50°C for Abaum0826 and Abaum0845 and at 55°C for the remainder loci, elongation for 50 seconds at 72°C, and a final elongation for 7 minutes at 72°C. After PCR, 3 µL of each PCR product was loaded on 2% regular agarose gel (Invitrogen, Carlsbad, CA, USA) for L-repeats and on 2% Metaphor agarose gel (FMC Bioproduct, Rockland, ME, USA) for S-repeats. Electrophoresis was performed in 25-cm wide gels and run at 4 V/cm in 0.5 × Tris-borate-EDTA buffers for about 5 hours. A 50-bp ladder (Bio-Rad) was used as a DNA size marker for L-repeat and S-repeat VNTRs. The gels were stained with 0.5 µg/mL of ethidium bromide for 1 hour and then photographed under UV illumination. The size of the amplicon was determined using the BioDocAnalyze (BDA) software. The length of repeat, the number of repetitions, and deduced size of the flanking regions were analyzed (https://tandem.bu.edu/trf/trf.html) (13, 14). Clusters analysis of the MLVA typing data was performed by MLVA Plus online tool. The polymorphism
4. Results

4.1. Antimicrobial Susceptibility Testing

Antibiotic susceptibility testing revealed that 80.8% (72 out of 89) of the isolates were MDR. About 87.6% of the isolates were resistant to cefepime (78 out of 89), 61.7% to amikacin (55 out of 89), 80.8% to gentamicin (72 out of 89), 92.1% to trimethoprim/sulfamethoxazole (82 out of 89), and 100% to ceftazidime, aztreonam, imipenem, meropenem, cefotaxime, and ciprofloxacin (Table 1). The distribution of MICs for imipenem and meropenem in 89 A. baumannii isolates is shown in Figure 1. High-level resistance (with MICs ≥ 128 µg/mL) to imipenem, meropenem, cefotaxime, and ciprofloxacin (Table 1). The most prevalent carbapenemase was the blaOXA-24-like gene. Two isolates carried three blaOXA-23-like genes. There was no relationship between high-level MIC and co-carriage of carbapenemase genes.

4.2. Molecular Detection of Carbapenemase

The most prevalent carbapenemase was the blaOXA-23-like gene (83/89, 93.2%), followed by the blaOXA-24-like gene (5/89, 5.6%), blaIMP (5/89, 5.6%), and blaOXA-58-like (1/89, 1.1%) genes. All blaOXA-24 gene-positive strains also carried the blaOXA-23-like gene. Two isolates carried three carbapenemase determinants, one isolate carried blaOXA-23-like, blaOXA-58-like, and blaOXM, and other isolates carried a combination of blaOXA-23-like, blaOXA-24-like, and blaOXM genes. Among six blaOXA-23-like gene-negative strains, one isolate was positive for the blaOXM gene, and five isolates were positive for the blaOXA-24-like gene. There was no relationship between high-level MIC and co-carriage of carbapenemase genes.

4.3. MLVA Genotyping

The occurrence of selected VNTR loci was studied in 89 A. baumannii strains. All L-repeat and S-repeat VNTR loci yielded amplification products ranging from 154 bp to 826 bp for most of the isolates (Table 2). In one of the isolates, a half-size allelic variant was observed at locus Abaum2240. In 10 isolates, locus Abaum1998 yielded an ambiguous amplification pattern with two bands; these uncommon patterns were defined as “a”. The amplification failure was observed in the isolates at Abaum3468 (22.4%), Abaum3530 (5.6%), Abaum2240 (6.7%), Abaum1998 (6.7%), Abaum0286 (7.8%), Abaum2396 (7.8%), and Abaum3002 (4.4%). As shown in Table 2, the highest diversity was observed at S-repeat VNTRs, resulting in HGDI values ranging from 0.89 (Abaum3468) to 0.96 (Abaum0285). The locus Abaum0845 showed a wide variability and generated alleles with high diversity in repeat numbers, from one repeat up to 64 repeats; therefore, due to the high diversity associated with Abaum0845 in this study and thus no further discriminatory power by adding Abaum0845, the Abaum0845 locus was excluded from the MLVA scheme. Therefore, the isolates were analyzed by seven loci (MLVA-7).

The 83 isolates were distinguished by 25 different MLVA types, as represented by individual allelic profiles of L-repeat VNTRs (Figure 2). MLVA type 1 and 2 were the most prevalent types containing 19 and 12 members with identical allelic profiles of L-repeat VNTRs (allelic profiles of 6-7-2-3 and 7-7-3-9, respectively), followed by MLVA type 3 with seven members, MLVA type 4 with five members, MLVA type 5, 6, and 7 with four members, MLVA type 8, 9, 10, 11, and 12 with three members, and MLVA type 12 and 13 with two members. The rest of the MLVA types included only one isolate. As shown in Figure 2, with a cut-off value of 75% corresponding to the existence of identically sized alleles for at least three L-repeat VNTRs, one clonal complex and three singletons were generated.

Finally, 82 different MLVA types with individual allelic profiles for 83 isolates were defined with the MLVA-7 assay. A combined HGDI value of 0.95 was achieved, consistent with the high level of polymorphism of these VNTR loci. Figure 3 shows the minimum spanning tree (MST) clustering of A. baumannii isolates. Due to the existence of identical sized alleles for at least five VNTRs, six clonal complexes (two isolates or more) and 17 singletons were observed. As shown in Figure 3, the clonal complex 1 (CC1) with 34 isolates and CC2 with 16 isolates were the main clusters, and the rest of the CCs included two or three isolates.

Figure 1. Distribution of MIC values of IPM (imipenem) and MER (meropenem) in 89 A. baumannii isolates

Figure 2. Distribution of MIC values of IPM (imipenem) and MER (meropenem) in 89 A. baumannii isolates

Figure 3. The minimum spanning tree (MST) clustering of A. baumannii isolates.
The OXA-23 carrying isolates were dispersed in all of the six clonal complexes; four OXA-23 negative isolates were distributed to CC1, and two OXA-23 negative isolates were singletons. The other carbapenemase gene-carrying isolates were distributed to all of the CCs and/or singletons. CC5 contained only the isolates from hospital 1 and CC6 contained only the isolates from hospital 2, and the rest of the CCs contained isolates from both hospitals. MLVA-7 dendrogram and characteristics of A. baumannii strains are shown in Figure 4.

5. Discussion

A. baumannii is one of the most common pathogens associated with hospital-acquired infections, which is almost resistant to all available antibiotics (2). To some extent, the success of A. baumannii in adaptation to the hospital environment has been attributed to its ability in acquiring resistance to almost all antibiotics, including carbapenems (4). Carbapenems are used as the last resort for the treatment of MDR strains of A. baumannii; however, the acquired carbapenem-hydrolyzing OXA-type class D β-lactamases (CHDLs) and metallo β-lactamase producing strains have been reported worldwide (1, 4). The intrinsic blaOXA-51-like gene family, the acquired blaOXA-23-like, blaOXA-24/40-like, and blaOXA-58-like gene families are considered the four main families of CHDL genes that have been reported for A. baumannii (1). The emergence and rapid expansion of CHDL gene-carrying strains have been found to be associated with hospital-acquired infections and outbreaks in hospitals, particularly in intensive care units (ICUs) (4). In the present study, it was found that blaOXA-23-like gene (93.2%) was the most prevalent gene in carbapenem-resistant isolates from the two hospitals in Tehran. The emergence of the blaOXA-23-like gene in clinical isolates of A. baumannii has been reported in other studies conducted in Iran (7, 16). Clonal outbreaks of blaOXA-23-like gene-carrying A. baumannii strains have also been reported in other studies conducted in France and Brazil (17, 18). In this study, one (mainly blaOXA-23-like gene), two, or three carbapenemase genes were detected in all of the carbapenem-resistant isolates. In addition, all the carbapenemase gene-carrying isolates were resistant to imipenem and/or to meropenem. However, there was no relationship between high-level MICs and co-carriage of carbapenemase genes. While OXA carbapenemase and MBLs are considered the main resistance mechanisms in A. baumannii, the co-existence of other resistance mecha-
nisms such as increased expression of efflux pumps, loss of porins and other carbapenemase genes such as KPC carbapenemase, NDM, and SIM metallo-beta-lactamase genes can affect the levels of resistance (1, 18, 19). Thus, the presence of the aforementioned resistance mechanisms probably has affected the level of MICs in our isolates. In addition, the expression of carbapenemase genes can be affected by the presence of insertion elements in the upstream of the OXA gene (20).

MLVA has been developed for typing of various microbial species including *Escherichia coli* O157, *Salmonella enterica*, *Pseudomonas aeruginosa*, and *A. baumannii* (21-23). In the current study, MLVA-8<sub>Orsay</sub> scheme was performed in order to trace the epidemiology of the carbapenem-resistant *A. baumannii* isolates and for bacterial differentiation at the strain level. The MLVA-8<sub>Orsay</sub> scheme is based on eight VNTR markers, four of which are L-repeat VNTR markers (repeat units of 26 to 99 bp) with moderately diverse repeats, showing low levels of polymorphism, and the rest of the markers are highly diverse S-repeat VNTR markers (repeat units of 6 to 9 bp), characterized by a large number of alleles with high levels of variability (6). Among S-repeat VNTR markers, Abaum0845 was excluded from the MLVA scheme in the current study due to the high allelic diversity of this locus in the isolates tested. In the current study, the presence of Abaum0845 in the MLVA scheme caused a great diversity so that it was not possible to cluster the isolates, making it difficult to interpret the obtained results. However, when the Abaum0845 locus was excluded, the MLVA-7 scheme generated six complexes with a 71.4% genetic similarity cut-off. Abaum0845 was also absent in all *A. baumannii* isolates tested in another study conducted in Iran (16). In addition, variability associated with Abaum0845 was reported in another study conducted in Spain (24). It is possible that this locus is a rapidly evolving marker. In addition, six isolates showed no products at more than three loci including L-repeat and S-repeat VNTR markers, possibly due to the differences in the genetic structure of these isolates at the corresponding loci. These six isolates were also excluded from further analysis by MLVA-7 scheme and other VNTR markers or typing methods are needed for their genotyping. The amplification failure at various VNTR markers was shown in the registered isolates of *A. baumannii* in the MLVA bank for Acinetobacter genotyping (http://microbesgenotyping.i2bc.paris-saclay.fr/databases/view/202). The website of MLVA bank 2016.V1.3.5 is hosted by the University of Paris-Saclay, Orsay, France.

A total number of 83 *A. baumannii* isolates were typed into 25 different MLVA types using four L-repeat loci. Using a cut-off value of 75% corresponding to the existence of identically sized alleles for at least three L-repeat VNTRs, one clonal complex was achieved, showing the suitability of these markers for studying the phylogenetic relationships among the strains. Interestingly, among L-loci, Abaum1998 yielded an ambiguous amplification pattern with two bands in 10 isolates. In a study by Hu et al., two different repeat units were obtained for locus Abaum1998 in all the tested isolates; thus, this locus was excluded from their MLVA scheme (23). However, in this study, an uncommon pattern was defined by “a” for 10 isolates and analyzed as an individual pattern.

Pourcel et al. proposed the criteria for the assignment of the strains to MLVA complexes as (i) identical L-repeat VNTR profiles and (ii) a maximum of three differences of no more than two repeats at S-repeat VNTR loci (6). The intrinsic diversity of L-repeat VNTRs was higher in the isolates of the current study than in the isolates used in Pourcel et al. study; therefore, our MLVA complexes were not grouped with identical L-repeat VNTR profiles. In addition, in this study, the difference was observed in more than two repeats at S-repeat VNTR loci; thus, due to the diversity associated with MLVA-7 markers, applying the Pourcel et al.
criteria was impossible. However, the MLVA complexes in this study were defined assuming a 71% genetic similarity cut-off, corresponding to the existence of identically sized alleles for at least five VNTR markers (Figure 3). By combining seven MLVA markers, an excellent discriminatory power was achieved (HGD1 = 0.99), which is suitable for diseases surveillance and/or outbreaks investigation.

The 77 OXA-23 carrying isolates were distinguished by 76 MLVA-7 types within the six clonal complexes (Figure 4). Only did two OXA-23 gene-carrying isolates, i.e., 104T and 157T, share the same MLVA-7 allelic profile, which were isolated from hospital 1 and 2, respectively. The other families of carbapenemase gene-carrying isolates were also distributed to different clonal complexes with individual MLVA-types. No correlation was observed between MLVA-7 defined complexes and carbapenemase gene carriage and the origin of the isolates. Overall, the MLVA-7 analysis revealed that carbapenemase gene-carrying A. baumannii isolates in this study might be polyclonal (corresponding to 76 MLVA-7 types) and provided evidence for gene transfer among different MLVA-types in two hospitals in Tehran.

In conclusion, the main purpose of the study was to evaluate the MLVA-8Orsay scheme for typing of clinical A. baumannii isolates. We had some limitations with MLVA-8Orsay for typing the test isolates. First, Abaum0845 was excluded from the MLVA scheme because of its great allelic diversity observed in our isolates, making it difficult to cluster test isolates. Second, Abaum1998 yielded two bands in 10 isolates; these uncommon patterns were considered as distinct patterns. Third, nine isolates were excluded from further analysis due to amplification failure at more than three VNTR loci. Fourth, we did not apply Pourcel et al. criteria for the interpretation of the results and clustering the isolates due to the high diversity associated with MLVA-7 markers. However, the discriminatory power of MLVA-7 was found to be excellent and possibly useful and reliable in epidemiological investigations of A. baumannii. If MLVA-8Orsay was developed with new S-markers, it would be more helpful for better interpretation and clustering of the isolates.

This method is PCR-based, making it feasible in many laboratories. The results also provided evidence for the possible use of MLVA-7 scheme as an alternative or complementary method for typing of A. baumannii isolates.
### Footnotes

**Authors' Contribution:** All authors participated in the research design and contributed to different parts of the research.

**Conflict of Interests:** There is no conflict of interests.

**Ethical Approval:** The study was approved by the Faculty of Medical Sciences, Tarbiat Modares University, Tehran, Iran (ID: 91356).

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| ID  | Hospital | Specimen | Word 2010 | Word 2012 | Word 2013 | Word 2014 | Word 2015 | Word 2016 | Word 2017 | Word 2018 | Word 2019 | Word 2020 | Word 2021 | MC-1 | MC-2 | MC-3 | MC-4 | MC-5 | MC-6 | MC-7 | MC-8 | MC-9 | MC-10 | MC-11 | MC-12 |
|-----|----------|----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-------|------|------|------|------|------|------|------|------|------|------|------|
| 124T | ETT | Int | 6 | 7 | 2 | 1 | 3 | 10 | 19 | 26 | pos | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 |
| 124T | ETT | Csf | 6 | 7 | 2 | 3 | 10 | 19 | 26 | pos | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 |
| 124T | ETT | Wound | 6 | 7 | 2 | 3 | 10 | 19 | 26 | pos | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 |
| 124T | ETT | Urethra | 6 | 7 | 2 | 3 | 10 | 19 | 26 | pos | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 |
| 124T | ETT | Urine | 6 | 7 | 2 | 3 | 10 | 19 | 26 | pos | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 |
| 124T | ETT | Sputum | 6 | 7 | 2 | 3 | 10 | 19 | 26 | pos | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 |
| 124T | ETT | Sputum | 6 | 7 | 2 | 3 | 10 | 19 | 26 | pos | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 |
| 124T | ETT | Sputum | 6 | 7 | 2 | 3 | 10 | 19 | 26 | pos | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 |
| 124T | ETT | Sputum | 6 | 7 | 2 | 3 | 10 | 19 | 26 | pos | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 | 64 |

Figure 4. MLVA-7 allelic profiles and characteristics of A. baumannii strains. The categorical coefficient and UPGMA algorithm were used for the generation of the dendrogram. ETT: endotracheal tube, ICU: intensive care unit, sur: surgery, emr: emergency, int: internal medicine, infect: infectious, N: no amplification, a: two bands pattern, MIC: $\mu$g/mL.
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