Background: Multidrug resistance Pseudomonas aeruginosa (MDR-\(P.\) aeruginosa) is a worldwide threat for public health. Hyperexpression of efflux pump systems (MexAB-OprM and MexCD-OprJ), which is a well-known mechanisms for MDR emerging, is controlled by regulatory genes, \(mexR\) and \(nfxB\), respectively. The aim of this study was to evaluate point mutations in \(mexR\) and \(nfxB\) genes in MDR-\(P.\) aeruginosa isolated from wound infections.

Materials and Methods: A total of 34 \(P.\) aeruginosa cultures obtained from wound infections were analyzed. Among them eight isolates identified as MDR-\(P.\) aeruginosa and were subjected to determination of mutations in \(mexR\) and \(nfxB\) genes.

Results: We detected eight-point mutations in \(mexR\) and 12-point mutations in \(nfxB\). The most common mutations were common G327-A (eight isolates), G384-A (eight isolates), G411-A (eight isolates). Mutations in A371-C and A372-C were the predominant substitution which was seen in \(nfxB\). Amino acid substitutions were also found at position 124 and 126 for NfxB and MexR, respectively.

Conclusions: \(P.\) aeruginosa isolates with mutation in efflux pump regulatory genes such as \(mexR\) and \(nfxB\) could be a main factor contributed to antibiotic resistance and must be considered in antibiotic treatment.

Key Words: Efflux pump, \(mexR\), \(nfxB\), Pseudomonas aeruginosa

INTRODUCTION

\(P.\) aeruginosa, which is known as a nonfermentative gram negative opportunistic pathogen, is the leading cause of diverse infections including pneumonia, wound and urinary tract infection (UTI). Because of resistance to many antibiotics, treatment of infections caused by \(P.\) aeruginosa is difficult. Several mechanisms have been known for antibiotic resistance, for example, inactivation or modification of the antibiotic, alteration, or modification in the target site of the antibiotic, and decreased intracellular drug concentration by overexpression of efflux pump. Until recently, in \(P.\) aeruginosa five super family of efflux pumps including (i) the ATP-binding cassette (ABC) family, (ii) the small multidrug
resistance family, (iii) the major facilitator super family, (iv) the resistance-nodulation-division (RND) family, and (v) the multidrug and toxic compound extrusion family have been recognized.\(^4\) Analysis of P. aeruginosa genome showed that all five super families are presented but RND family is predominant.\(^5\) RND pumps consist of three subunit including membrane fusion protein (MFP), an outer membrane factor (OMF), and a cytoplasmic membrane transporter.\(^5,6\) MexAB-OprM and MexCD-OprJ, which are members of RND efflux pumps family, have the wide substrate of antibiotics such as, β-lactam class (aztreonam, ceftazidime) and fluoroquinolones.\(^7\) Active efflux pumps decrease intracellular concentration of drug by pumping it to out. Expression of MexAB-OprM and MexCD-OprJ are regulated by pumping it to out. Expression of MexAB-OprM and MexCD-OprJ is regulated by mexR and nfxB, respectively.\(^17\) MexR consisting of 147 amino acid residues, negatively regulate MexAB-OprM expression.\(^8\) Mutation in mexR (nalB strains) and nfxB compromised regulatory functions and therefore resulting in efflux pumps hyperexpression.\(^8,9\) Several independent studies have revealed that overexpression of mentioned efflux pumps contributed to multi-drug resistance (MDR) phenotype.\(^10-12\) MDR P. aeruginosa is defined as resistance to at least three antibiotics belonged to different classes especially, carbapenems, fluoroquinolones, and aminoglycosides.\(^13\) Due to limited choice of effective drug, mortality rates in patients infected with MDR P. aeruginosa is higher in comparison with multiple drugs susceptible P. aeruginosa.\(^14,15\) Therefore, in order to have appropriate therapy for patients infected with P. aeruginosa, identification of multidrug resistance isolates is essential. In this study, our aim was to detect mexR and nfxB mutations in MDR- P. aeruginosa isolated from wound infections.

**MATERIALS AND METHODS**

**Sample collection and identification**

This study was conducted between January 2013 and March 2013 at three major hospitals of Isfahan, Iran. During the study, a total of 100 wound samples that were obtained from wound infections were analyzed. P. aeruginosa identification performed based on standard test such as Gram staining, catalase, oxidase, oxidative-fermentative (OF) test, pigment production, and growth at 42°C.\(^1,16\) Subsequently, primary identification confirmed with PCR by using ITS (16s-23s rRNA internal transcribed spacer) gene specific primer.\(^17\)

**Antibiotic susceptibility testing**

Antibiotic susceptibility of the isolates was ascertained by Kirby-Bauer disk diffusion method according to CLSI (clinical laboratory standard institute) guidelines.\(^18\) In this study, following antibiotic disks (MAST, UK) were used: ceftazidime (30 μg), cefotaxime (30 μg) aztreonam (30 μg), ciprofloxacin (5 μg), imipenem (10 μg), meropenem (10 μg), and amikacin (30 μg). P. aeruginosa ATCC 27853 was used for quality control.

**Preparation of genomic DNA**

For DNA extraction, two or three colony of fresh culture of P. aeruginosa was dissolved in 300 ml of lysis buffer containing (Tris 100 mmol, NaCl 50 mmol, and EDTA 25 mmol, pH = 7.5) completely. Subsequently, suspension was boiled at 95°C for 15 min. Equal volumes of phenol and chloroform (25:24, pH = 7.5) was added, mixed thoroughly, and centrifuged at 9000 g for 5 min. Aqueous–viscous supernatant was transformed to a fresh micro tube, phenol/chloroform (25:24) was added again and centrifuged at 9000 g for 5 min. To DNA precipitation, 600 μl of cold pure ethanol (Merck, Germany) was added and centrifuged at 13000 g (4°C, 20 min). Obtained DNA after washing with 70% ethanol was stored at –20°C.

**mexR and nfxB amplifications and sequencing**

PCR for mexR (503 bp) and nfxB (731 bp) amplification were carried out in separate 25 μl reaction mixture consist of 10 pmol of each primer (Metabion, Germany), mexR-F (5’-CTGATCAAACACATTTACA-3’), mexR-R (5’-CTTCGAAAAGAATGTTCTTAAA-3’), nfxB-F (5’-ACGGAGCCAGTGGTTCT-3’) and nfxB-R (5’-ACTGATCTTCCGCAGTGTCG-3’), 2.5 μl (25 mMol Mgcl₂), 200 μM dNTP, 2.5 μl PCR buffer 10X, 1.25 u taq DNApol (Cinna gen, Iran), and 5 μl DNA template.\(^19,20\) Cycling condition performed in 35 cycles including, denaturation at 95°C for 1 min, annealing at 55°C for mexR and 60°C for nfxB, 45 s, extension at 72°C for 1 min and final extension at 72°C for 10 min. PCR products were visualized on 1.5% agarose gel stained with DNA green viewer dye (Figures 1 and 2). PCR products were sent for BIO NEER (Korea) company for purification and sequencing on both strands. We used Mega4 as multiple sequence alignment software for sequences analyzing.

**RESULTS**

Eight of 34 nonrepetitive P. aeruginosa were identified as MDR-P. aeruginosa. All isolates showed resistance to ceftazidime, cefotaxime, aztreonam, ciprofloxacin, imipenem, and meropenem. Our results showed seven silent mutations including G327-A (eight isolates), G384-A (eight isolates), G411-A (eight isolates), C223-T (five isolates), C264-T (two isolates), G60-A (one isolate), C168-T (one isolate) in mexR gene. Only one missense mutation which changed val
126 to Glu was seen [Table 1]. Four isolates had five different point mutations simultaneously [Table 1]. In nfxB, 12-point mutations were found. Mutations at position A371-C and A372-C were the predominant substitution. We also found missense mutations in codon 124 that changed Glu to Ala [Table 1]. The GenBank accession numbers of mexR and nfxB which determined in this work are as follow: KF675781-88.

**DISCUSSION**

Resistance to multiple antibiotics in *P. aeruginosa* as one of the most common etiological agents of wound infections is well-known property. Treatment of MDR-*P. aeruginosa* is difficult due to limited choice of antibiotics. Generally, antibiotic resistance mechanism in *P. aeruginosa* is divided into two categories, intrinsic and acquired resistance. Low permeability of outer membrane protein, constitutive

![Figure 1: Gel image of representative PCR of mexR gene. Line1 Ladder, Lane 2 negative control, line 3 positive, line 4 to 10 clinical specimens](image)

![Figure 2: Gel image of representative PCR of nfxB gene. Line1, 7 Ladder, Lane 2 positive control, line 3 to 6 clinical specimens, line 8 negative control](image)

### Table 1: Mutations in mexR, nfxB in MDR-*Pseudomonas aeruginosa* isolated from wound infections

| Isolates | mexR | nfxB | Antibiotic resistance |
|----------|------|------|-----------------------|
| 1        | G60-A | A61-C | R: AZ, CZ, CT, CP, MR, IM |
|          | G327-A | (Glu124-Ala) | |
|          | T377-A | A372-C | |
|          | (val126-Glu) | A373-G | |
|          | G384-A | T555-G | |
|          | G411-A | | |
| 2        | C223-T | A371-C | R: AZ, CZ, CT, CP, MR |
|          | G327-A | (Glu124-Ala) | IM, AM |
|          | T377-A | A372-C | |
|          | (val126-Glu) | G423-A | |
|          | G384-A | T480-C | |
|          | G411-A | T555-G | |
| 3        | C223-T | A371-C | R: AZ, CZ, CT, CP, MR |
|          | G327-A | (Glu124-Ala) | IM, AM |
|          | T377-A | A372-C | |
|          | (val126-Glu) | G423-A | |
|          | G384-A | T480-C | |
|          | G411-A | T555-G | |
| 4        | C223-T | G39-A | R: AZ, CZ, CT, CP, MR |
|          | G327-A | A183-G | IM, AM |
|          | T377-A | A371-C | |
|          | (val126-Glu) | (Glu124-Ala) | |
|          | G384-A | A372-C | |
|          | G411-A | G423-A | |
|          | | T480-C | |
|          | | A486-T | |
|          | | T537-C | |
|          | | C543-T | |
|          | | T555-G | |
| 5        | C223-T | G39-A | R: AZ, CZ, CT, CP, MR |
|          | G327-A | A183-G | IM, AM |
|          | T377-A | A371-C | |
|          | (val126-Glu) | (Glu124-Ala) | |
|          | G384-A | A372-C | |
|          | G411-A | G423-A | |
|          | | T480-C | |
|          | | A486-T | |
|          | | A536, T537 deletion | |
|          | | C543-T | |
|          | | T555-G | |
| 6        | C223-T | A371-C | R: AZ, CZ, CT, CP, MR |
|          | G327-A | (Glu124-Ala) | IM, AM |
|          | T377-A | A372-C | |
|          | (val126-Glu) | G423-A | |
|          | G384-A | T480-C | |
|          | G411-A | T555-G | |
| 7        | C168-T | A371-C | R: AZ, CZ, CT, CP |
|          | C264-T | (Glu124-Ala) | MR, IM, AM |
|          | G327-A | A372-C | |
expression of AmpC and efflux pumps is example of intrinsic resistance mechanism.[23-25] MexAB-OprM and MexCD-OprJ are two preponderant efflux pumps that contribute to intrinsic resistance of P. aeruginosa to different antibiotics.[3] MexAB-OprM and MexCD-OprJ are negatively regulated by MexR and NfxB, respectively; hence, mutations in these regulatory genes lead to overexpression of mentioned efflux pumps and multiple drug resistance emerging.[19,26] In our study mutations of mexR and nfxB in MDR-P. aeruginosa isolated from wound infections were investigated. In mexR, 8-point mutations were detected, seven of them were silent. Only the mutation in thymine 377 and conversion to adenine leads to substitution of Val with Glu in codon 126. This substitution had related to antibiotic resistance, according to similar studies.[19,27,28] Results of several independent studies revealed different mutations, for example, Suman and colleagues reported 24 silent and four missense mutations in 14 clinical isolates of P. aeruginosa.[29] Different amino acids substitution with relation to antibiotic resistance was seen in codon 21, 95, 30, 79, 106, 114, 78, 107, 53, and 103 of mexR gene.[10,12,29,30] Although mutation in mexR is one of the main cause of MexAB-OprM overexpression but its expression and DNA binding activity in order to prediction of antibiotic effectiveness should be evaluated. Based on our results, mutations in nfxB were relatively high. The nfxB mutations consisted of 11 silent mutations, one missense mutation and a deletion from position 536 to 537. Mutations in position 371 and 372 were the most frequently replacement in all isolates [Table 1]. Analysis of NfxB showed that DNA binding domain is located between amino acid residues 26 and 42. [12] There was not any mutation mentioned region in this study. Missense mutation was seen in Glu 124-Ala of NfxB. Similar study revealed that change in codon 124 was related to ciprofloxacin resistance. [12] Results of another study of Fluoroquinolone resistance P. aeruginosa showed that mutation in codon 82 (Arg to Leu) was the most frequently substitution in nfxB. [31] Hyperexpression of efflux pumps such as MexAB-OprM and MexCD-OprJ are complicated. Isolates of nalC and nalD had mutations in PA-3721 and PA-3574 genes showed overexpression of MexAB-OprM independent from mexR mutations. [32,33] Therefore, expression of MexAB-OprM is not only regulated by MexR but also affected by PA-3721 and PA-3574 genes. In order to know preponderant mutations, further study for evaluation of expression mexB, mexC, oprM and oprJ by realtime PCR is necessary.

Table 1: Contd...

| Isolates | mexR | nfxB | Antibiotic resistance |
|----------|------|------|-----------------------|
| T377-A   | T480-C |     | AZ, CZ, CT, CP, MR IM |
| (val 126-Glu) | T537-C |     |                       |
| G384-A | C543-T |     |                       |
| G411-A | T555-G |     |                       |
| C264-T | A371-C |     |                       |
| (Glu124-Ala) | T377-A | G411-A | C543-T | T555-G |

a: R: Resistance, b: AZ: Aztreonam, CZ: Ceftazidime, CT: Cefotaxime, CP: Ciprofloxacin, MR: Meropenem, IM: Imipenem, AM: Amikacin

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REFERENCES

1. Forbes BA, Sahm DF, Weissfeld AS. Bailey and Scott's Diagnostic Microbiology, 12th ed. USA: Elsevier; 2007. p. 340-50.
2. Strateva T, Yordanov D. Pseudomonas aeruginosa: A phenomenon of bacterial resistance. J Med Microbiol 2009;58:1133-48.
3. Lister PD, Wolter DG, Hanson ND. Antibacterial-resistant Pseudomonas aeruginosa: Clinical impact and complex regulation of chromosomally encoded resistance mechanisms. Clin Microbiol Rev 2009;22:582-610.
4. Schweizer HP. Efflux as mechanism of resistance to antimicrobials in Pseudomonas aeruginosa and related bacteria: Unanswered questions. Genet Mol Res 2003;2:48-62.
5. Alekshun MN, Levy SB. Molecular mechanisms of antibacterial multidrug resistance. Cell 2007;128:1037-150.
6. Ozer B, Duran N, Onlen Y, Savas L. Efflux pump genes and antimicrobial resistance of Pseudomonas aeruginosa strains isolated from lower respiratory tract infections acquired in an intensive care unit. J Antimicrob Chemother (2012) 2016:65:9-13.
7. Llanes L, Hocquet D, Vogne C, Baltich DB, Neuwirth C, Plesiart P. Clinical strains of Pseudomonas aeruginosa overproducing MexAB-OprM and MexXY efflux pumps simultaneously. Antimicrob Agents Chemother 2004;48:1797-802.
8. Poole K, Tetro K, Zhao Q, Neshat S, Heinrichs DE, Bianco N. Expression of the multidrug resistance operon mexA-mexB oprM Pseudomonas aeruginosa: mexR encodes a regulator of operon expression. Antimicrob Agents Chemother 1996;40:2021-8.
9. Jeannot K, Elsen S, Kohler T, Delden CV, Plesiart P. Resistance and virulence of Pseudomonas aeruginosa clinical strains producing the MexCD-OprJ efflux pump. Antimicrob Agents Chemother 2008;52:2455-62.
10. Adewoye L, Sutherland A, Srikumar R, Poole K, The MexR repressor of the mexAB-oprM multidrug efflux operon in Pseudomonas aeruginosa: Characterization of mutations compromising activity. J Bacteriol 2002;184:4308-12.
11. Dumas JL, Delden CV, Perron K, Kohler T. Analysis of antibiotic resistance gene expression in Pseudomonas aeruginosa by quantitative real-time-PCR. FEMS Microbiol Lett 2006;254:217-25.
12. Higgins PG, Fluit AC, Milatovic D, Verhoef J, Schmitz FJ, Mutations in gyrA, ParC, MexR and NfxB in clinical isolates Pseudomonas aeruginosa. Int J Antimicrob Agents 2003;21:409-13.
13. Majigoros AP, Srinivasan A, Carey RB, Carmeli Y, Falagas ME, Giske CG, et al. Multidrug-resistant, extensively drug-resistant and pandrug-resistant bacteria: An international expert proposal for interim standard definitions for acquired resistance. Clin Microbiol Infect 2012;18:268-61.
14. Pena C, Zorrilla SG, Oriol I, Tubau F, Dominguez MA, Pujol M. Impact of multidrug resistance on *Pseudomonas aeruginosa* ventilator-associated pneumonia outcome: Predictors of early and crude mortality. Eur J Clin Microbiol Infect Dis 2013;32:413-20.

15. Morales E, Cots F, Sala M, Corom M, Belvis F, Riu M. Hospital costs of nosocomial multi-drug resistant *Pseudomonas aeruginosa* acquisition. BMC Health Serv Res 2012;12:1-8.

16. Mahon C, Lehman D, Manuselis G. *Text Book of Diagnostic Microbiology*, 4th ed. USA: Elsevier; 2011. p. 480-90.

17. Tyler SD, Strathdee CA, Rozee KR, Johnson WM. Oligonucleotide primers designed to differentiate pathogenic *Pseudomonas* on the basis of the sequencing of genes coding for 16S-23S rRNA internal transcribed spacers. Clin Diagn Lab Immunol 1995;2:448-53.

18. National Committee for Clinical Laboratory Standards, Ed., “Performance standards for antimicrobial disk susceptibility tests,” Approved Standards M100-S22, National Committee for Clinical Laboratory Standards, Wayne, Pa, USA, 2012.

19. Gorgani N, Ahlbrand S, Patterson A, Pourmand N. Detection of point mutations associated with antibiotic resistance in *Pseudomonas aeruginosa*. Int J Antimicrob Agents 2009;34:414-8.

20. Jeannot K, Elsen S, Kohler T, Attree I, Delden CV, Plesiat P. Resistance and virulence of *Pseudomonas aeruginosa* clinical strains overproducing the MexCD-OprJ efflux pump. Antimicrob Agents Chemother 2008;52:2455-62.

21. Breidenstein EB, Nunez CD, Hancock RE. *Pseudomonas aeruginosa* isolates from cystic fibrosis patients. Antimicrob Agents Chemother 2007;51:4062-70.

22. Hirsch EB, Tam VH. Impact of multidrug-resistant *Pseudomonas aeruginosa* infection on patient outcomes. Expert Rev Pharmacoecon Outcomes Res 2010;10:441-51.

23. Soto SM. Role of efflux pumps in the antibiotic resistance of bacteria embedded in a biofilm. Virulence 2013;4:1-7.

24. Livermore DM. Multiple mechanisms of antimicrobial resistance in *Pseudomonas aeruginosa*: Our worst nightmare?. Clin Infect Dis 2002;34:634-40.

25. Hooper DC. Efflux pumps and nosocomial antibiotic resistance: A primer for hospital epidemiologists. Clin Infect Dis 2005;40:1811-7.

26. Anderson C, Jalal S, Ali D, Wang Y, Islam S, Jari A. Critical biophysical properties in *Pseudomonas aeruginosa* efflux gene regulator MexR are targeted by mutations conferring multidrug resistance. Protein Sci 2010;19:680-92.

27. Tomas M, Dounith M, Warner M, Turton JF, Becero A, Bou G. Efflux pumps. OprD Porin, AmpC β-Lactamase, and multiresistance in *Pseudomonas aeruginosa* isolates from cystic fibrosis patients. Antimicrob Agents Chemother 2010;54:2219-24.

28. Henrichfreise B, Wiegard I, Pfister W, Wiedemann B. Resistance mechanisms of multiresistant *Pseudomonas aeruginosa* strains from Germany and correlation with hypermutation. Antimicrob Agents Chemother 2007;51:4062-70.

29. Suman G, Khan M, Sabitha K, Jamil K. Mutation in mexR-gene leading to drug resistance in corneal keratitis in human. Indian J Exp Biol 2006;44:929-36.

30. Hocquet D, Bertrand X, Kohler T, Talon D, Plesiat P. Genetic and phenotypic variations of a resistant *Pseudomonas aeruginosa* epidemic clone. Antimicrob Agents Chemother 2003;47:1887-94.

31. Jalal S, Ciofu O, Heiby N, Gotli N, Wretlind B. Molecular mechanisms of fluoroquinolone resistance in *Pseudomonas aeruginosa* isolates from cystic fibrosis patients. Antimicrob Agents Chemother 2000;44:710-2.

32. Morita Y, Cao L, Gould VC, Avison MB, Poole K. NaID encodes a second repressor of the MexAB-OprM multidrug efflux operon of *Pseudomonas aeruginosa*. J Bacteriol 2006;188:8649-54.

33. Sobel ML, Hocquet D, Cao L, Plesiat P, Poole K. Mutations in PA3574 (nalD) lead to increased MexAB-OprM expression and multidrug resistance in laboratory and clinical isolates of *Pseudomonas aeruginosa*. Antimicrob Agents Chemother 2005;49:1782-6.

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