



Tetracycline Resistance Associated with *tet* Genes or Integrons among *Enterobacter cloacae* Strains Isolated from Patients with Urinary Tract Infections

ARTICLE INFO

Article Type Original Research

Authors

Mona Banihashemi, $MSc^{1,2}$ Ali Majidpour, $PhD^{1,3}$ Mina Boustanshenas, MSc^{1*} Samaneh Mazar-Atabaki, MSc^{1}

How to cite this article

Banihashem M., Majidpour A., Boustanshenas M., Mazar-Atabaki S., Tetracycline Resistance Associated with tet Genes or Integrons among Enterobacter cloacae Strains Isolated from Patients with Urinary Tract Infections. Infection Epidemiology and Microbiology. 2020;6(1):1-9

¹ Antimicrobial Resistance Research Center, Institute of Immunology and Infectious Diseases, Iran University of Medical Sciences, Tehran, Iran ²Department of Microbiology, Faculty of Basic Sciences, Islamic Azad University, Science and Research Branch, Tehran, Iran ³ Department of Infectious Diseases, School of Medicine, Iran University of Medical Sciences, Tehran, Iran

* Correspondence

Address: Antimicrobial Resistance Research Center; Institute of Immunology and Infectious Diseases, Iran University of Medical Sciences, Tehran, Iran. bustanshenas.m@tak.iums.ac.ir

Article History

Received: January 15,2020 Accepted: March 15,2020 Published: March 22,2020

ABSTRACT

Background: This study aimed to evaluate the prevalence of *tet* genes and Class I and 2 integrons in *Enterobacter cloacae* strains isolated from patients with urinary tract infections (UTIs).

Materials & Method: A total of 50 *E. cloacae* isolates were collected. Antimicrobial susceptibility pattern and tetracycline MIC were determined. The presence of *tet* genes (tetA, tetB, tetC, tetD) and Class 1 and 2 integrons and the content of Class 1 integron were determined.

Findings: Tetracycline MIC pattern classified 36 % of the *E. cloacae* isolates as resistant. The most common *tet* gene was *tet*C (22%), followed by *tet*D, *tet*A, and *tet*B. Class 1 integron was detected in 64% of the isolates. Class 1 integron content analysis showed two variable gene cassettes (*aadA*1 and *aadA*5/*dfrA*17 genes). The frequency of *aadA*5/*dfrA*17 was 18.75%, which was more common than aadA1 gene (6.25%).

Conclusion: The most important genetic markers for tetracycline resistance in *E. cloacae* isolates were *tet*C and Class 1 integron. Harboring Class 1 integron and resistance to streptomycin and ciprofloxacin were significantly correlated.

Keywords: Integron, *Enterobacter cloacae*, Tetracycline resistance genes, Urine, Integron content, Class 1 and 2 integrons

CITATION LINKS

[1] Salimian Rizi K, Najar Peerayeh S, Bakhshi B, Rahbar M. Prevalence of integrons ... [2] Flynn DM, Weinstein RA, Nathan C, Gaston MA, Kabins SA. Patients' endogenous flora ... [3] Gaston M. Enterobacter: An emerging ... [4] Xu Z, Li L, Shirtliff M, Peters BM, Li B, Peng Y, et al. Resistance ... [5] You R, Gui Z, Xu Z, Shirtliff ME, Yu G, Zhao X, et al. Methicillin-resistance Staphylococcus... [6] Deng Y, Bao X, Ji L, Chen L, Liu J, Miao J, et al. Resistance... [7] Xu Z, Li L, Shi L, Shirtliff ME. Class 1 integron in... [8] Giakkoupi P, Tzouvelekis LS, Tsakris A, Loukova V, Sofianou D, Tzelepi E. IBC-1, ... [9] Peters SM, Bryan J, Cole MF. Enterobacterial repetitive intergenic consensus ... [10] Tzelepi E, Tzouvelekis L, Vatopoulos A, Mentis AF, Tsakris A, Legakis NJ. High prevalence of stably derepressed Class-I β-lactamase expression in multiresistant... [11] Tao R, Ying G-G, Su H-C, Zhou HW, Sidhu JP. Detection of... [12] Roberts MC. Update on acquired tetracycline resistance genes. FEMS Microbiol lett. 2005; 245(2):195-203. [13] Zhang X-X, Zhang T, Fang HH. Antibiotic resistance genes... [14] Sandalli C, Özgümüş OB, Sevim A. Characterization of tetracycline resistance genes in tetracycline-resistant... [15] Su H-C, Ying G-G, Tao R, Zhang RQ, Fogarty LR, Kolpin DW. Occurrence of... [16] Kaushik M, Kumar S, Kapoor RK, Virdi JS, Gulati P. Integrons... [17] Martínez-Carballo E, González-Barreiro C, Scharf S, Gans O. Environmental... [18] Pappas P. Laboratory in the diagnosis... [19] Wayner. Clinical and aboratory Standards... [20] Waters SH, Rogowsky P, Grinsted J, Altenbuchner J... [21] Ng LK, Martin I, Alfa M, Mulvey M. Multiplex PCR... [22] Adabi M, Bakhshi B, Goudarzi H, Zahraei SM, Pourshafie MR. Distribution... [23] White PA, McIver CJ, Deng YM, Rawlinson WD. Characterization.. [24] Uhlemann A-C, Annavajhala M, Gomez-Simmonds A. Multidrug-resistant complex... [25] Turner PJ. Meropenem activity against European isolates: Report.. [26] Mokracka J, Koczura R, Pawłowski K, Kaznowski A. Resistance patterns and integron cassette... [27] Bayani M, Siadati S, Rajabnia R, Taher AA. Drug resistance of... [28] Cattoir V, Poirel L, Rotimi V, Soussy CJ, Nordmann P. Multiplex PCR for... [29] Keeney D, Ruzin A, Bradford PA. RamA, a transcriptional regulator, and... [30] Mortazavi SH, Mansouri F, Azizi M, Alvandi A, Karbasfrushan A, Madadi-Goli N, et al. Prevalence of Class... [31] Peymani A, Farivar TN, Ghoraiian P, Najafipour R. Association between Class 1 integrons and multidrug resistance pattern among Enterobacter spp. isolated from Qazvin and Tehran teaching hospitals. J Qazvin ...

Introduction

Enterobacter is a genus of Gram-negative, facultative, anaerobic bacilli belonging to family Enterobacteriaceae which are widely found in nature [1]. The identification of this genus is routinely carried out using phenotypic and genotypic methods mainly based on the conventional biochemical tests and molecular based methods, respectively. These organisms are capable of causing opportunistic infections in hospitalized or debilitated patients [2-3]. Indiscriminate use of antibiotics in medical health and veterinary industries has led to an upward trend in antimicrobial resistance and the emergence of multidrug-resistant (MDR) strains among a massive variety of microorganisms [4-^{5]}. Before the extensive use of antibiotics, Enterobacter species were rarely considered as pathogens. But nowadays, there are many reports about the role of these organisms in causing major clinical diseases including bacteremia and urinary tract infections [6]. The increase in antimicrobial resistance among the clinical isolates is considered as a principle threat to the global public health, which could lead to enhanced clinical failures in the treatment of infectious diseases [7]. Different kinds of resistance have been reported in E. cloacae strains, including resistance to extended-spectrum cephalosporins, tetracyclines, extendedspectrum β-lactamas (ESBLs) and integronassociated class 1 [8-10]. Tetracycline is a broad spectrum bacteriostatic antibiotic which is widely used for the treatment and also as a promoter in animal husbandry industry [11]. The emergence of tetracycline resistant strains is a consequence of widespread usage of tetracycline both in medical and food industries. Resistance to tetracycline is usually due to the acquisition of *tet* resistance genes. Approximately 38 different *tet* genes have been recognized among the resistant different strains, applying resistance mechanisms. The mechanisms of these genes include: encoding efflux pump proteins, inactivating enzymes, and ribosomal protection proteins [12-13]. Based on the tet genes characteristics and mechanisms, they have been classified in different groups named with alphabetic letters. The efflux pump-mediating tet genes, including tetA, B, C, D, and E, have been reported as the most frequent genes among Enterobacteriaceae [14-15]. Overabundant use of antibiotics in the treatment of bacterial infections has led to the selection of resistant strains and increased risk of vertical and horizontal transfer of resistance genes among bacterial isolates. Until recently, it was thought that antimicrobial resistance genes transfer in bacteria is mainly occurred through the conjugation and transduction by plasmids, phages, and transposons carrying resistance genes, but later on, another mechanism was identified for antibiotic resistance genes transfer, which is done by elements called "integrons". Integrons are genetic components which are able to integrate into the alien DNA and transmit animated genetic elements called gene cassettes. As the integrons have a promoter for the genes in their contents, they could express the gene packages which were captured. Therefore, integrons not only act as a vector of gene expression but also as a natural cloning system. Integrons possess two essential genes located in conserved segment (CS), named int genes which are able to transmit and insert gene cassettes. Based on the differences in int genes, integrons have been classified into 4 different classes. Class 1 integron is the most common integron in enterobacterial isolates. Class 1, 2, and 3 integrons in bacterial isolates have been determined to be associated with antimicrobial resistance in a variety of bacterial species [16-17].

Objectives: This study aimed to evaluate the antibiotic resistance pattern, the prevalence

of tetracycline resistance genes, and the distribution and content of Class I and 2 integrons in the *E. cloacae* strains isolated from patients with urinary tract infections (UTIs).

Materials and Methods

Ethics approval and consent to participate: The study began after ethical approval by Medical Ethics Committee of Tarbiat Modares University (Code: IR.MODARES.REC). The participants provided their written informed consent in order to participate in this study.

Bacterial strains and samples: A total number of 21600 urine specimens were collected from patients with UTIs from three large academic hospitals in Tehran, Iran, during 2016-2018. Patients with symptomatic UTIs were characterized with bacteriuria (direct observation of bacteria in urine specimens) and pyuria (urinary leukocyte excretion rates of ≥400,000 leukocytes/h, which correlates with a hemocytomer count of ≥10 leukocytes/ mm³) [18]. Among 7200 culture-positive urine samples, a total of 50 *E. cloacae* strains were isolated. Urine samples were cultured on BHI agar medium and incubated at 37 °C for 18-24 hrs, discrete colonies were subjected to Gram staining. The Gram negative bacilli were subjected to biochemical tests for the presence of E. cloacae strains using API 20E identification kit in accordance with the manufacturer's manual (BioMérieux SA, Lyon, France). The standard strain of *E.* cloacae ATCC 13048 was used as positive control.

Resistance pattern of *E. cloacae* isolates: Disc diffusion and minimum inhibitory concentration (MIC) ^[2] methods were used to determine resistance to different antimicrobial agents. Results were evaluated and interpreted according to CLSI (Clinical and Laboratory Standards Institute)

guidelines (2017). Antibiotic discs including tetracycline (T, 30µg), minocycline (MIN, 30μg), docetaxel (TS, 25μg), streptomycin 10μg), gentamicin (GM, (ST, $10\mu g$), ampicillin (AP, 10µg), and ciprofloxacin (CIP, 5µg) were placed on cultured media. After incubation at 37°C for 18-24 hrs, inhibition zones diameter was accurately measured using a millimeter ruler. According to CLSI guidelines (2018), the antibiogram test results for each antibiotic were reported as susceptible, resistant, or intermediate.

Determining the tetracycline MIC by agar dilution method: The 0.5 McFarland concentration of the isolates was prepared and cultured on Muller Hinton agar containing different concentrations of tetracycline, ranging from 0.5 to 256 μg.ml⁻¹, and incubated at 37 °C for 18 hrs. The minimum concentration of each antibiotic inhibiting bacterial growth was considered as MIC of that antibiotic ^[19].

Prevalence of tetracycline resistance genes and Class I and 2 integrons: The genomic DNA of all the isolates was extracted using boiling method. The specific primers and PCR programs used in this study are shown in Table 1. The presence of four different genes including tetA, tetB, tetC, and tetD, as tetracycline resistance genes, was evaluated among the isolates. The *int*1 and in primers were used for determining Class 1 integron and its genetic content, respectively. The presence of intergron Class 2 was determined by *hep* primer. The PCRs were performed in a 25µL mixture consisting of 12.5 µL Amplicon master mix (superTag DNA polymerase, dNTPs and Taq-buffer), 10 pmol from each primer, 5 μL DNA template, and sterile distilled water up to 25 µL as a final volume of PCR mixture. The PCR products were electrophoresis on 1% agarose gel and visualized using Gel Documentation system. PCR program was set as follows: an initial denaturation step

Table 1) Primers sequences used in this study

Gene	Primer Sequence 5'3'	Amplification Condition	References
tet A	GTAATTCTGAGCACTGTCGC CTGCCTGGACAACATTGCTT	927 bp	(20)
tet B	TTGGTTAGGGGCAAGTTTTG GTAATGGGCCAATAACACCG	659 bp	(21)
tet C	CTTGAGAGCCTTCAACCCAG ATGGTCGTCATCTACCTGCC	418 bp	(21)
tet D	AAACCATTACGGCATTCTGC GACCGGATACACCATCCATC	717 bp	(21)
int 1	TGCGTGTAAATCATCGTCGT CAAGGTTCTGGACAGTTGC	900 bp	(22)
hep	CGGGATCCCGGACGGCATGCACGATTTGTA GATGCCATCGCAAGTACGAG	Variable	(23)

at 95°C for 5 min, 30 cycles of denaturation at 95°C for 1 min, annealing for 1 min (for *tet*A, *tet*D, and *int*1 at 53°C; for *tet*B and *tet*C at 55°C; for *in* and *hep* at 56°C), extension at 72°C for 1 min, and a final extension at 72°C for 10 min.

Nucleotide sequence accession numbers: One representative of each of the two gene cassettes contained in Class 1 integron was deposited in the GenBank database under Accession Numbers MN052647 and MN052648.

Statistical analysis: The sample size of this study was calculated using the sample size formula based on the prevalence of *E. cloacae* strains in urine samples collected from patients with UTIs.

Findings

A total of 2173 out of 7243 urine samples were positive for bacterial infection, of which 50 (2.3%) cases were associated with *E. cloacae* as the main pathogen according to the biochemical tests results. The resistance, intermediate, and susceptibility rates of *E. cloaceae* strains to different antibiotics

used in this study are shown in Table 2. The highest resistance was observed to ampicillin (92%) (46 out of 50 isolate), and the lowest resistance was observed to streptomycin and minocycline (12%). The tetracycline MIC was also evaluated by agar dilution method to validate the results of disc diffusion method. According to the MIC results, 18 strains (36%) were resistant to tetracycline, and only 1 strain showed susceptibility (2%); however, the tetracycline resistance rate was only 6% based on the disc diffusion method results, which was significantly different from the MIC results (P value < .05). Eleven different resistance patterns were detected among the E. cloacae isolates, and the most common one (56%) (28 isolates out of 50) belonged to the resistance to minocycline, tetracycline, and ampicillin (Table 3). Most of the isolates showing this resistance pattern also harbored Class 1 integron (53%); however, only 1 isolate harbored tetA gene, and 8 isolates harbored tetC gene, other tet genes were not detected in this resistance pattern.

Table 2) Antibiogram results of *E. cloacae* isolates using the disc diffusion method

Antibiotic Agents	ST (10mg)* N(%)	TS (25mg) N(%)	CIP (5mg) N(%)	AP (10mg) N(%)	MIN (30mg) N(%)	GM (10mg) N(%)	T (30mg) N(%)
Resistance	6 (12%)	18 (36%)	9 (18%)	46 (92%)	6 (12%)	8 (16%)	3 (6%)
Intermediate	12 (28%)	0	2 (4%)	3 (6%)	46 (92%)	0	0
Susceptible	30 (60%)	32 (64%)	39 (78%)	1 (2%)	0	42 (84%)	47 (94%)

^{*}ST: streptomycin; TS: co-trimoxazol; CIP: ciprofloxacin; AP: ampicillin; MIN: minocycline; GM: gentamicin; T: Tetracyclin **Table 3)** Resistance pattern among the *E. cloaceae* isolates

Antibiotics	No. of Strains (n=50)	No. of Strains Harboring Integron Class 1 (n=32)
AP, T	1	1
AP, MIN	1	1
AP, MIN, T	28	15
ST, AP, MIN, T	6	4
CIP, MIN, T	1	0
ST, AP, MIN, GM, T	1	0
ST, CIP, AP, MIN, T	2	1
ST, TS, AP, MIN, GM,T	1	1
ST, TS, CIP, AP, MIN,T	3	3
ST, CIP, AP, MIN,GM,T	3	3

Prevalence of tetracycline resistance determinants and integrons among the *E. cloaceae* **isolates**: In the present study, four different classes of *tet* genes, including *tet*A, B, C, and D, were investigated among the isolates using specific primers. The PCR products with the band sizes of 927, 659, 418, and 787bp belonged to *tet*A, *tet*B, *tet*C, and *tet*D, respectively. The highest prevalence rate was associated with *tet*C gene (22%) (11 out of 49), followed by *tet*D (6%), *tet*A, and

tetB (2%). Harboring *int*I and *hep* gene is the representative of Class 1 and 2 integrons. Although Class 2 integron was not found, Class 1 integron was common among the isolates (64%) (Table 4). Class 1 integron content analysis showed two variable gene cassettes at the sizes of 1000 and 1500 bp, belonging to *aadA1* and *aadA5/dfrA17* genes, respectively (Figure 1). The frequency of *aadA5/dfrA17* was 18.75%, which was more common than *aadA1* gene (6.25%).

Table 4) Tetracycline resistance genes and Class 1 and 2 integrons in the isolates

Gene	Strains No. (%)
tetA	1 (2%)
tet B	1 (2%)
tet C	11 (22%)
tet D	3 (6%)
hep	0
Int 1	32 (64%)

Discussion

Increasing frequency of antimicrobial resistance among clinically isolated bacteria is considered as a global health problem. The same trend has been reported for *E. cloacae* strains ^[24]. Various factors could influence the resistance pattern of *E. cloacae* strains, including antibiotic regimen, geographic conditions, and the year of sampling ^[25]. Many studies have demonstrated the upward trend of antibiotic resistance among

E. cloacae strains. In a study conducted by Mokracka and colleagues (2011), the tetracycline resistance was reported to be over 60% in Poland, while in the present study, the tetracycline resistance was lower [22]. As mentioned, one of the reasons for differences in antibiotic resistant patterns among E. cloacae strains is antibiotic regimen, which could vary in different geographical regions. For instance, the highest resistance rate in Iran has been reported against β-lactams, which is in agreement with the present study results [1], whereas another study by Bayani and colleagues (2013) demonstrated the highest resistance of E. cloacae isolates against cephalosporins; in their study, no MDR strain was detected [23]. In the present study, 5 strains were classified as MDR strains, underlining the treatment of MDR *E. cloacae* strains. In other studies, resistance to other antibiotic families, including tetracyclines and quinolones, was also reported with high resistance rate [28-29]. In the present study, the most prevalent *tet* gene among the *E. cloacae* isolates was tetC,

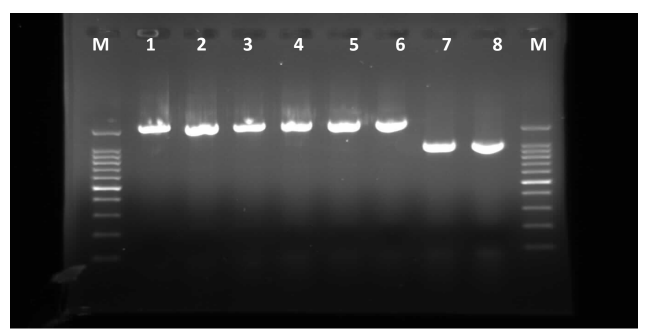


Figure 1) Class 1 integron content among integron positive strains. M:1kb DNA size marker; lane 1-6: clinical *E. cloacae* strains harboring *aadA5/dfrA17* gene cassette with 1500 bp size; lane 7 and 8: clinical *E. cloacae* strains harboring *aadA1* gene cassette with 1000 bp size.

*ST(10mg): streptomycin; TS(25mg): co-trimoxazol; CIP(5mg): ciprofloxacin; AP(10mg): ampicillin; MIN(30mg): minocycline; GM(10mg): gentamicin; T(30mg): Tetracyclin

and all the isolates harbored only one type of tet genes; no evidence of heterogeneity was observed in the tet genes. It is noteworthy that all the isolates harboring tetB and tetD showed the same resistance pattern against tetracycline and minocycline; thus, the probable relationship between *tet* genes and resistance to minocycline could be hypothesized. The isolates with tetD gene had the highest tetracycline MIC value (≥128 ug.mL⁻¹), which proved that *tet*D might have a role in intensifying the tetracycline resistance. Low prevalence of tetracycline resistance genes in phenotypically resistant isolates could be due to the presence of other resistance mechanisms, including integrons or genes which has not yet been detected in E. cloacae strains. The prevalence of MDR E. cloacae strains demonstrated an upward trend in the emergence of MDR strains in the clinical samples in Iran in such a way that the rate has increased from zero in 2012 to 75% in 2017 according to the published data [27, ^{30-31]}. Integrons are mostly responsible for the transmission of resistance determinants within species or among different bacterial species or genera. The frequency of Class 1 integron was 64% in this s1tudy, which is supported by the results of other studies conducted in Iran, while Class 2 integron was not detected. There is no evidence about the presence of Class 2 integron among *E. cloacae* isolates in published studies (1, 30). According to the statistical analysis, the correlation between harboring Class 1 integron and resistance to streptomycin and ciprofloxacin was significant (p value < .05), while resistance to other antibiotics was not correlated with harboring Class 1 integron, and other resistance mechanisms should be responsible for resistance.

Conclusion

There is a growing trend in the diagnosis of *E. cloacae* in clinical samples. The increase

in resistance to antibiotics is due to the proliferation of integrons and their resistance gene cassettes. For asmuch as MDR *E. cloacae* strains causing nosocomial infection have increased, and the mechanisms of resistance are complicated, the prevalence, resistance profile, and genotype of *E. cloacae* strains should be routinely investigated in health care units.

Acknowledgements: The author would like to extend their sincere appreciation to Antimicrobial Resistance Research Center, Iran University of Medical Sciences for supporting this study.

Ethical Permissions: not applicable

Conflict of interests: The authors declare that they have no conflict of interests.

Authors Contribution: AM: Infectious diseases consultant, MB: Main researcher, designed the study and performed molecular assays; MB: Performed microbial assays. SMA: Statistical analyst.

Fundings: This work was funded by Iran University of Medical Sciences [Grant Number 25907].

References

- 1. Salimian Rizi K, Najar Peerayeh S, Bakhshi B, Rahbar M. Prevalence of integrons and antimicrobial resistance genes among clinical isolates of Enterobacter spp. from hospitals of Tehran. Int J Enteric Pathog. 2015; 3(1):e22531.
- 2. Flynn DM, Weinstein RA, Nathan C, Gaston MA, Kabins SA. Patients' endogenous flora as the source of "nosocomial" Enterobacter in cardiac surgery. J Infect Dis. 1987; 156(2):363-8.
- 3. Gaston M. *Enterobacter*: An emerging nosocomial pathogen. J Hos Infec. 1988; 11(3):197-208.
- 4. Xu Z, Li L, Shirtliff M, Peters BM, Li B, Peng Y, et al. Resistance Class 1 integron in clinical methicillin-resistant

- Staphylococcus aureus strains in southern China, 2001–2006. Clin Microbiol Infec. 2011; 17(5):714-8.
- 5. You R, Gui Z, Xu Z, Shirtliff ME, Yu G, Zhao X, et al. Methicillin-resistance Staphylococcus aureus detection by an improved rapid polymerase chain reaction (PCR) assay. Afr J Microbiol Res. 2012; 6(43):7131-3.
- 6. Deng Y, Bao X, Ji L, Chen L, Liu J, Miao J, et al. Resistance integrons: Class 1, 2, and 3 integrons. Ann Clin Microbiol Antimicrob. 2015; 14(1):45.
- 7. Xu Z, Li L, Shi L, Shirtliff ME. Class 1 integron in Staphylococci. Mol Biology Rep. 2011; 38(8):5261-79.
- Giakkoupi P, Tzouvelekis LS, Tsakris A, Loukova V, Sofianou D, Tzelepi E. IBC-1, a novel integron-associated Class A β-lactamase with extended-spectrum properties produced by an Enterobacter cloacae clinical strain. Antimicrob Agents Chemother. 2000; 44(9):2247-53.
- 9. Peters SM, Bryan J, Cole MF. Enterobacterial repetitive intergenic consensus polymerase chain reaction typing of isolates of Enterobacter cloacae from an outbreak of infection in a neonatal intensive care unit. Am J Infec Control. 2000; 28(2):123-9.
- 10. Tzelepi E, Tzouvelekis L, Vatopoulos A, Mentis AF, Tsakris A, Legakis NJ. High prevalence of stably derepressed Class-I β-lactamase expression in multiresistant clinical isolates of Enterobacter cloacae from Greek hospitals. J Med Microbiol. 1992; 37(2):91-5.
- 11. Tao R, Ying G-G, Su H-C, Zhou HW, Sidhu JP. Detection of antibiotic resistance and tetracycline resistance genes in Enterobacteriaceae isolated from the Pearl rivers in South China. Environ Pollut. 2010; 158(6):2101-9.
- 12. Roberts MC. Update on acquired tetracycline resistance genes. FEMS

- Microbiol lett. 2005; 245(2):195-203.
- 13. Zhang X-X, Zhang T, Fang HH. Antibiotic resistance genes in water environment. Appl Microbiol Biotechnol. 2009; 82(3):397-414.
- 14. Sandalli C, Özgümüş OB, Sevim A. Characterization of tetracycline resistance genes in tetracycline-resistant Enterobacteriaceae obtained from a coliform collection. World J Microbiol Biotechnol. 2010; 26(11):2099-103.
- 15. Su H-C, Ying G-G, Tao R, Zhang RQ, Fogarty LR, Kolpin DW. Occurrence of antibiotic resistance and characterization of resistance genes and integrons in Enterobacteriaceae isolated from integrated fish farms in south China. J Environ Monit. 2011; 13(11):3229-36.
- 16. Kaushik M, Kumar S, Kapoor RK, Virdi JS, Gulati P. Integrons in Enterobacteriaceae: Diversity, distribution, and epidemiology. Int J Antimicrob Agents. 2018; 51(2):167-76.
- 17. Martínez-Carballo E, González-Barreiro C, Scharf S, Gans O. Environmental monitoring study of selected veterinary antibiotics in animal manure and soils in Austria. Environ Pollut. 2007; 148(2):570-9.
- 18. Pappas P. Laboratory in the diagnosis and management of urinary tract infections. Med Clin North Am. 1991; 75(2):313-25.
- 19. Clinical and Laboratory Standards Institute. M100: Performance standards for antimicrobial susceptibility testing. 28th Edition. Wayne P: CLSI; 2018.
- 20. Waters SH, Rogowsky P, Grinsted J, Altenbuchner J, Schmitt R. The tetracycline resistance determinants of RP1 and Tn1721: nucleotide sequence analysis. Nucleic Acids Res. 1983; 11(17):6089-105.
- 21. Ng LK, Martin I, Alfa M, Mulvey M. Multiplex PCR for the detection of tetracycline resistant genes. Mol Cell

- Probes. 2001; 15(4): 209-15.
- 22. Adabi M, Bakhshi B, Goudarzi H, Zahraei SM, Pourshafie MR. Distribution of Cclass I integron and sulfamethoxazole trimethoprim constin in vVibrio cholerae isolated from patients in Iran. Microb Drug Resist. 2009; 15(3):179-84.
- 23. White PA, McIver CJ, Deng YM, Rawlinson WD. Characterization of two new gene cassettes, aadA5 and dfrA17. FEMS Microbiol Lett. 2000; 182(2): 265–69.
- 24. Uhlemann A-C, Annavajhala M, Gomez-Simmonds A. Multidrug-resistant complexemerging as a global, diversifying threat. Front Microbiol. 2019;10:44.
- 25. Turner PJ. Meropenem activity against European isolates: Report on the MYSTIC (meropenem yearly susceptibility test information collection) 2006 results. Diagn Microbiol Infect Dis. 2008; 60(2):185-92.
- 26. Mokracka J, Koczura R, Pawłowski K, Kaznowski A. Resistance patterns and integron cassette arrays of Enterobacter cloacae complex strains of human origin. J Med Microbiol. 2011; 60(6):737-43.
- 27. Bayani M, Siadati S, Rajabnia R, Taher AA. Drug resistance of Pseudomonas aeruginosa and Enterobacter cloacae

- isolated from ICU, Babol, Northern Iran. Int J Mol Cell Med. 2013; 2(4):204.
- 28. Cattoir V, Poirel L, Rotimi V, Soussy CJ, Nordmann P. Multiplex PCR for detection of plasmid-mediated quinolone resistance *qnr* genes in ESBL-producing enterobacterial isolates. J Antimicrob Chemother. 2007; 60(2):394-7.
- 29. Keeney D, Ruzin A, Bradford PA. RamA, a transcriptional regulator, and AcrAB, an RND-type efflux pump, are associated with decreased susceptibility to tigecycline in Enterobacter cloacae. Microb Drug Resis. 2007; 13(1):1-6.
- 30. Mortazavi SH, Mansouri F, Azizi M, Alvandi A, Karbasfrushan A, Madadi-Goli N, et al. Prevalence of Class I and II integrons among MDR Enterobacter cloacae isolates obtained from clinical samples of children in Kermanshah, Iran. J Clin Diagn Res. 2018; 12(12).
- 31. Peymani A, Farivar TN, Ghoraiian P, Najafipour R. Association between Class 1 integrons and multidrug resistance pattern among Enterobacter spp. isolated from Qazvin and Tehran teaching hospitals. J Qazvin Univ Med Sci. 2014; 18(2):30-8.