

Identification of Carbapenemase within Class 1 Integron Structure in Intrinsically Colistin-resistant Enterobacteriaceae

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Introduction: Identification of carbapenem resistance genes in intrinsically

colistin-resistant Enterobacteriaceae (ICRE), i.e., *Proteus* spp., *Providencia* spp., *Serratia* spp., and *Morganella* spp., is necessary for preventing such life-

threatening bacteria. Here, we characterized carbapenemase resistance genes

and their carriage within class 1 integron structures in clinical intrinsically colistin-resistant Enterobacteriaceae. **Methods:** Clinical samples were collected from six general hospitals in Tehran and Torbat-e-Hevdarieh, Iran, from 2018

to 2020. The isolates were identified by standard microbiological techniques and

16s rRNA sequencing. The resistance to imipenem and colistin was determined

by disk diffusion and minimum inhibitory concentration (MIC) assays. PCR-

sequencing was performed to detect primary carbapenemase-encoding genes,

including blaoXA-48, blaIMP, blaVIM, blaKPC, and blaNDM. Results: Three Proteus

mirabilis, three Providencia stuartii, and two Serratia marcescens isolates were

identified. The MIC_{IMP} values of all isolates were >16 μ g/ml. The *bla*_{VIM-1} gene

and bla_{KPC-2} genes were detected in five and four isolates, respectively. Cassette

arrays consisting of *blavim-1-aacA7-dhfrA1-aadA1* were detected in five isolates

and *aadB* in one isolate. The bla_{VIM-1} gene in all bla_{VIM-1} positive isolates was located on class 1 integrons, while the bla_{KPC-2} carriage was not related to these genetic elements. The *in silico* analysis of the large plasmids carrying bla_{VIM-1} *aacA7-dhfrA1-aadA1* array belonging to the IncA/C incompatibility group showed transmissibility among Enterobacteriaceae members. **Conclusions:** Transferable plasmid harboring bla_{VIM-1} in ICREs can pose a severe threat to public health. More investigations are required to depict the epidemiology of these underestimated carbapenem-resistant Enterobacteriaceae members.

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INTRODUCTION

The risk of antibiotic resistance is rapidly increasing worldwide [1]. Carbapenems are considered the most effective antibiotics for the treatment of healthcareassociated infections [2]. Carbapenem-resistant Enterobacteriaceae (CRE) are considered a major emerging threat to global public health due to the rapid emergence in clinical settings worldwide, limited treatment options, and high morbidity and mortality rates associated with infections [3]. However, the overuse of carbapenems has led to the emergence of carbapenem resistance driven mainly by the production of carbapenemases such as OXA-48, GES, KPC, IMP, VIM, and NDM [1, 2]. The CRE was listed among the first "critical priority pathogens" by the World Health Organization (WHO) in 2017 [4, 5].

The increasing number of multidrug-resistant (MDR) bacteria, particularly Gram-negative carbapenemresistant Enterobacterales, has led to using colistin as the antimicrobial of last resort [6]. However, resistance to carbapenems seems to be underestimated in less-studied Enterobacteriaceae members such as Proteus spp., Serratia spp., and Providencia spp. [2, 7]. Among the Enterobacteriaceae members, Proteus spp., Providencia spp., Edwardsiella spp., Serratia marcescens, and Morganella morganii are considered intrinsically colistin-resistant Enterobacteriaceae (ICRE). Consequently, only a few antibiotics remain available for the clinical treatment of CRE-infected patients [8, 9].

Despite the lower isolation rate among these bacteria compared to highly prevalent Enterobacteriaceae

Bolourchi et al.

e.g., *Escherichia coli* and *Klebsiella pneumoniae*, the detection and reporting of carbapenem resistance in ICREs are highly significant [10]. Because the drug of choice against these bacteria is carbapenems, they are considered almost the only therapeutic options [11]. Furthermore, the emergence of carbapenem resistance in ICREs is a critical medical situation. Therefore, early detection of ICRE isolates can significantly control their spread in healthcare systems. In the present study, we reported carbapenem-resistant intrinsically colistin-resistant *P. mirabilis, S. marcescens,* and *P. stuartii* in clinical specimens. Also, we characterized the structure of class 1 integron harboring carbapenemase genes.

MATERIAL AND METHODS

Bacterial isolates. Clinical samples were collected from six general hospitals in Tehran and Torbat-e-Heydarieh, Iran, from 2018 to 2020, and isolates were identified by standard microbiological techniques, including colony morphology, Gram stain, oxidase, and a conventional biochemical test panel. Species identification was performed by PCR-sequencing of the 16s rRNA [12].

Antimicrobial susceptibility testing. Antimicrobial susceptibility was determined using the Kirby-Bauer disk diffusion method according to Clinical and Laboratory Standards Institute (CLSI) guidelines 2020 [13]. The antimicrobials tested included amoxicillin (10 μ g), cefotaxime (30 μ g), ceftazidime (30 μ g), ciprofloxacin (5 μ g), gentamycin (10 μ g) and imipenem (10 μ g). Susceptibility to colistin was determined by the agar dilution method based on clinical breakpoints defined by CLSI [13]. The minimum inhibitory concentrations (MICs) of the CRE isolates to imipenem were determined by the microdilution method according to the recommendations and interpretation criteria of the CLSI [13]. The strain *E. coli* ATCC 25922 was used as a control in antimicrobial susceptibility testing.

Molecular detection of carbapenemases and class 1 integrons. Whole DNA was extracted from bacterial isolates by the boiling method as previously described [14]. The CRE isolates were screened for the carbapenemases including *bla*_{OXA-48}, *bla*_{IMP}, *bla*_{VIM}, $bla_{\rm KPC}$ and $bla_{\rm NDM}$ using PCR, as previously described [15], with the primers designed by others (Table 1), and PCR conditions reflected elsewhere [16]. All integronpositive strains were screened for variable regions of the class 1 integron using primers 5'-CS /3'-CS. Sequencing of the purified DNA was performed by the Bioneer Company with Sanger dideoxy chain termination method using Applied Biosystems 3730/3730X1 DNA Analyzers (Applied Biosystems). The nucleotide sequence analysis was performed using the BLAST tool at the NCBI website (https://blast.ncbi.nlm.nih.gov/Blast.cgi). The sequences were manually analyzed using CLC workbench software version 20 (CLC Bio, Aarhus, Denmark).

Plasmid analysis. A collection of plasmids carrying class 1 integron was retrieved from the GenBank database (https://www.ncbi.nlm.nih.gov/genbank/), and the plasmids incompatibility (Inc) was determined by the online web tool PLSDB (https://ccb-microbe.cs.unisaarland.de/plsdb/) [17]. The presence of relaxasome complex in plasmids was determined using the online web tool oriTfinder to determine their conjugative capability (https://bioinfo-mml.sjtu.edu.cn/oriTfinder/) [18]. This web tool was developed to facilitate the rapid detection of oriT, the origin of transfer sequence in bacterial plasmids. It can also detect the T4CP and T4SSs as components involved in plasmids horizontal transfer. The online web tool CARD identified all antimicrobial resistance genes on plasmids (https://card.mcmaster.ca/analyze/rgi) [19]. Plasmid sequence alignment was performed by Mauve version 2.4.0, an open-free software for multiple alignments of genomic sequences that exhibit the arrangement of gene cassettes within different sequences

Table 1. Primers used for am	plification of 16s rRNA	. carbapenemase-encoding	genes, and class 1 integrons

Primer	Primer sequence (5'→3')	Product size (bp)	Reference
IMP	F: GGAATAGAGTGGCTTAAYTCTC	189	[37]
	R:CCAAACYACTACGTTATCTKGAG		
VIM	F: AGTGGTGAGTATCCGACAG	261	[37]
	R:ATGAAAGTGCGTGGAGAC		
NDM	F: GGCAGCACACTTCCTATCTC	155	[37]
	R:GTTGATCTCCTGCTTGATCC		
KPC	F: ATCGCCGTCTAGTTCTGCTG	850	[37]
	R:CCCTCGAGCGCGAGTCTA		
OXA48	F: TTGGTGGCATCGATTATCGG	658	[37]
	R:CATCAAGTTCAACCCAACCG		
16S rRNA	F: AGAGTTTGATYMTGGCTC	~1500	[37]
	R:CAKAAAGGAGGTGATCC		
intI1	F: TCTCGGGTAACATCAAGG	250	[16]
	R: AGGAGATCCGAAGACCTC		
Conserved segment of class 1 integrons	5'-CS: GGCATCCAAGCAGCAAG	Variable	[16]
	3'-CS: AAAGCAGACTTGACCTGA or		
	3'-CS:GAAGCGGCGTCGGCTTGA		

RESULTS

Demographic data. Nine non-duplicated carbapenemresistant, intrinsically colistin-resistant Enterobacteriaceae isolates were identified (Table 2) in the clinical isolates originated from urine (n=5), blood (n=2), sputum (n=1), and surgery site (n=1). The patients'

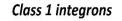
age ranged from 41 to 66 years, except for one 27-yearold male. Three species, including P. mirabilis (n=4), P. stuartii (n=3), and S. marcescens (n=2), were identified by biochemical and molecular tests. Four out of five urine samples contained P. mirabilis.

ю.	Strain	Isolation	Gender/Age	Hospital	Patient's	MIC (µg/ml) of	Antimicrobial	Carbapenemase	intI1	Variable region of
	source		source	outcome	imipenem	resistance Profile [*]	genes		class 1 integrons	
	P. stuartii	Urine	F/53	А	Discharge	>32	IPM, CTX, AMX,	bla _{VIM-1}	+	blaviM-1-aacA7-
							GEN			dhfrA1-aadA1
	P. stuartii	Tracheal	M/46	В	Death	>64	IPM, GEN, CAZ,	bla _{KPC-2}	-	-
							AMX			
	P. stuartii	Blood	M/61	В	Death	>16	IPM, CTX, GEN	bla _{VIM-1}	+	blavim-1-aacA7-
										dhfrA1-aadA1
	P. mirabilis	Urine	F/41	С	Discharge	>32	IPM, CTX, GEN,	bla _{VIM-1}	+	bla _{VIM-1} -aacA7-
							CIP, AMX			dhfrA1-aadA1
	P. mirabilis	Urine	F/55	В	Death	>64	IPM, CTX, CIP,	bla _{VIM-1}	+	blavIM-1-aacA7-
							CAZ, AMX			dhfrA1-aadA1
	P. mirabilis	Urine	M/47	С	Death	>32	IPM, GEN, CIP,	bla _{VIM-1}	+	blavIM-1-aacA7-
							AMX			dhfrA1-aadA1
	P. mirabilis	Urine	M/66	D	Discharge	>32	IPM, CTX, GEN,	bla _{KPC-2}	+	aadB
							CIP, AMX			
	S. marcescens	Surgery	F/27	Е	Discharge	>16	IPM, CTX, AMX	bla _{KPC-2}	-	-
	S. marcescens	Blood	M/58	F	Death	>32	IPM, CTX, GEN,	bla _{KPC-2}	-	-
							AMX			

*AMX: amoxicillin, CAZ: ceftazidime, CTX: ceftaxime, CIP: ciprofloxacin, GEN: gentamycin, IMP: imipenem, M: Male, F: Female.

Antimicrobial resistance test. All isolates were resistant to imipenem by disk diffusion test. The MIC of imipenem among the bacterial isolates ranged from 32 to 64 µg/ml for P. mirabilis and S. marcescens and 16 to 64 µg/ml for P. stuartii. As tested by the agar dilution method, the MIC for colistin was $>8 \mu g/ml$ for all isolates.

Molecular detection of carbapenemases. PCR screening for the carbapenem-resistant genes among the CRE isolates revealed only *bla*_{VIM-1} and *bla*_{KPC-2}. One *P*. stuartii, one P. mirabilis, and two S. marcescens isolates were positive for *bla*_{KPC-2}, and three *P. mirabilis* and two P. stuartii isolates were positive for blavIM-1. The blaIMP, $bla_{\rm NDM}$, and $bla_{\rm OXA-48}$ genes were not detected among the isolates. Table 2 shows the data on carbapenem-resistant, intrinsically colistin-resistant Enterobacteriaceae isolates.



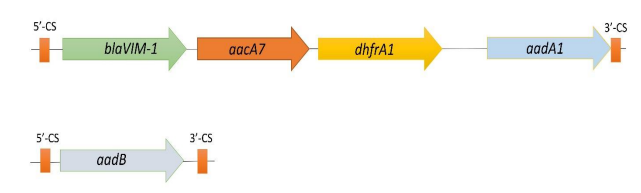


Fig. 1. The gene cassette arrays consisting of *bla*_{VIM-1}-*aacA7-dhfrA1-aadA1* and *aadB*. The *bla*_{VIM-1} gene carried on class 1 integrons in all positive VIM isolates, while the $bl_{a \text{KPC-2}}$ gene was not associated with class 1 integrons.

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Molecular detection of class 1 integron. PCR detected the presence of integrase gene intI1 in 6 out of 9 isolates. Class 1 integrons were widespread among these clinical isolates. However, there was no *intI1* in *S. marcescens*. The class 1 integrons cassette arrangements were characterized by amplification-sequencing of the internal variable regions (IVRs). Sequencing confirmed the presence of cassette arrays consisting of *bla*_{VIM-1}-*aacA7*-*dhfrA1-aadA1* (5/9) and *aadB* (1/9). The *bla*_{VIM-1} gene in all isolates was on class 1 integrons, while the *bla*_{KPC-2} gene was not related to class 1 integrons (Fig. 1).

Plasmid analysis harboring *blav*_{IM-1}*-aacA7-dhfrA1aadA1* **array.** We collected eight plasmids from the GenBank database. Two categories of plasmid harbored *blav*_{IM-1}*-aacA7-dhfrA1-aadA1* array, including plasmids (IncA/C) with >100 Kbp and ~52% CG content, isolated

from different bacteria. The second group was plasmids (IncR-1) with <100 Kbp and ~53% CG content, isolated from K. pneumoniae. Characterization of resistance genes on plasmid showed that the first category had heterogenic resistance genes compared to the second group. All plasmids (except pG06) had intact conjugal apparatus. See Table 3. Genetic similarities were observed among all eight plasmids. The alignment of plasmid sequences showed that plasmids CP031573, JO824049, CP029718, CP032168, CP031584, and CP031610 (the first group) possessed consensus genetic structures with different arrangements. The CP032168 and CP031584 (both belonging to K. pneumoniae) had identical nucleotide arrangements. KU665641 and CP023926 in the second group had consensus structures lacking large compartments of the first group. See Fig. 2.

Table 3. Plasmid analysis harboring blavIM-1-aacA7-dhfrA1-aadA1 array among Enterobacteriaceae

Plasmid	Accession	DNA	CG%	Bacterium	Inc	oriT, relaxase, T4SS	Other resistance genes
name	number	size (bp)		source	typing	and T4CP	
pIncAC2	CP031573	164095	52.45%	E. hormaechei	IncA/C	oriT (+), relaxase (+),	ant (2")-Ia, qnrA1, bla _{TEM-1} , mphA,
						and T4CP (+)	qacEdelta1 and sul1
pTC2	JQ824049	180184	52.51%	P. stuartii	IncA/C	oriT (+), relaxase (+),	mphA, bla _{SHV-5} , dfrA12, aph(3')-Ia
						and T4CP (+)	qacEdelta1 and sul1
Unnamed	CP029718	158987	52.56%	E. cloacae	IncA/C	oriT (+), relaxase (+),	mphA, QnrA1, dfrA12, qacEdelta1
						and T4CP (+)	and sull
Unnamed	CP032168	158987	52.56%	K. pneumoniae	IncA/C	oriT (+), relaxase (+),	mphA, qnrA1, dfrA12,
						and T4CP (+)	qacEdelta1and sul1
pIncAC2	CP031584	164082	52.45%	K. pneumoniae	IncA/C	oriT (+), relaxase (+),	ant (2")-Ia, qnrA1, bla _{TEM-1} , mphA
						and T4CP (+)	qacEdelta1 and sul1
pIncAC2	CP031610	160292	52.51%	E. coli	IncA/C	oriT (+), relaxase (+),	mphA, qnrA1, dfrA12, qacEdelta1
						and T4CP (+)	and sull
pG06	KU665641	53618	53.02%	K. pneumoniae	IncR	oriT (+), relaxase (-), and	ant (3")-IIa, aph(6)-Id, aph(3")-Ib,
						T4CP (-)	dfrA12, qacEdelta1 and sul1
Unnamed	CP023926	78541	53.50%	K. pneumoniae	IncR	oriT(+), relaxase (+),and	ant(3")-IIa, aph(6)-Id, aph(3")-Ib,
						T4CP (-)	dfrA12, qacEdelta1 and sul1

Nucleotide sequence accession numbers. The bla_{VIM-1} and bla_{KPC-2} gene sequences and the bla_{VIM-1} -*aacA7-dhfrA1-aadA1* and *aadB* arrays were deposited at DDBJ/ENA/GenBank under the accession MZ359738, MZ359739, MZ541989, and MZ541990, respectively.

DISCUSSION

Nosocomial infections caused by carbapenem-resistant ICRE pose a severe clinical threat and contribute to pandrug resistance. Reports suggest that the prevalence of carbapenem-resistant ICRE is increasing worldwide [20, 21].

The mortality rate of our ICRE isolates was high; five out of nine patients (55%) were expired, similar to a report from Greece [22]. Unfortunately, the patient's outcomes are rarely available in many other reports on naturally colistin-resistant Enterobacteriaceae. Therefore, determining the average mortality rate caused by carbapenem-resistant ICRE is complicated and requires further investigations.

Of the four *P. mirabilis* isolates, only one harbored $bla_{\text{KPC-2}}$, while the rest were positive for $bla_{\text{VIM-1}}$, indicating a higher prevalence of $bla_{\text{VIM-1}}$ in our *P. mirabilis* isolates. The $bla_{\text{KPC-2}}$ -harboring *P. mirabilis* was first reported in the United States (US) in 2008 [23], followed by other reports from china in 2010 and Brazil in 2015 [24, 25]. Reports of Metallo beta-lactamase $bla_{\text{VIM-1}}$ -like genes in *P. mirabilis* are available from Greece and Italy in 2006 and 2008, respectively, and

J Med Microbiol Infect Dis

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more recently from Bulgaria [25-27]. The bla_{VIM} -like carriage can be utilized to determine the levels of carbapenem resistance in *P. mirabilis*. Bontron *et al.* (2019) determined the blaVIM- $_1$ copy numbers in four susceptible P. mirabilis variants using quantitative PCR;

comparing the results with the imipenem MIC values suggested that the phenotypic resistance degree to carbapenems in bla_{VIM-1} positive *P. mirabilis* depends on the copy number. [28].

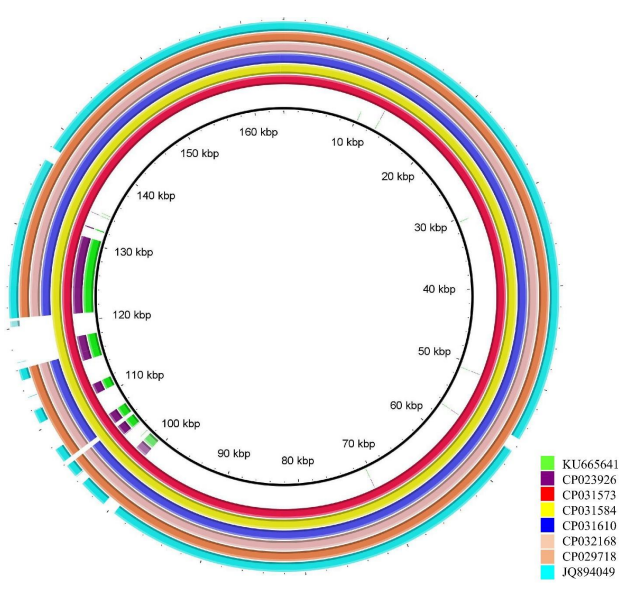


Fig. 2. Multiple sequence alignments of plasmids harboring *bla*_{VIM-1}*-aacA7-dhfrA1-aadA1* array among Enterobacterales. Sequences comparing showed that larger plasmids CP031573, JQ824049, CP029718, CP032168, CP031584, and CP031610 possessed consensus genetic structures with different arrangements (the first group), while smaller plasmids KU665641 and CP023926 were categorized in the second group.

Today, 12 carbapenem-resistant *P. stuartii* incidents comprising 39 cases and three outbreak reports are available worldwide [29]. There are reports for $bla_{\rm KPC-2}$ and $bla_{\rm VIM}$ -like for this bacterium from Saudi Arabia, Japan, Taiwan, and other countries [29-31]. We detected three non-duplicated clinical isolates with three different sources, two of which produced $bla_{\rm VIM-1}$ and one produced $bla_{\rm KPC-2}$.

In our study, both *S. marcescens* isolates encoded $bla_{\text{KPC-2}}$; this bacterium typically exhibits carbapenem resistance by encoding different bla_{KPC} genes, first reported in China [32]. Previously, *S. marcescens* was a less commonly reported bacterium in the clinical setting [33]. However, reports from Brazil and China of clinical carbapenem-resistant *S. marcescens* strains showed its clinical importance. In these studies, most isolates were

Bolourchi et al.

blaKPC-2 positive, indicating a high prevalence of this type of plasmid-mediated carbapenemase..

The sequencing results indicated that all blavIM-1 positive isolates carried this gene within the same class 1 integron structure. The blavIM-1 carriage in our intrinsically colistin-resistant CRE isolates was associated *bla*_{VIM-1}*-aacA7-dhfrA1-aadA1* cassettes. The with blaVIM₋₁ carriage by this structure was previously reported in P. stuartii isolates in 2016 [34]. Similarly, blavIM-1-aacA7-dhfrA1-aacA1 was identified in blavIM-1 positive P. mirabilis isolates in Switzerland in 2019. The integron mapping indicated that integron class 1 truncated by the IS1 element resulted in *qacED1* and *sul1* genes loss. This recombination in P. mirabilis was mediated by surrounding IS26 elements resulting in the acquisition of bla_{VIM-1} [28]. Reports of the bla_{VIM-1} carriage in P. *mirabilis* isolates by a class 1 integron containing aac(6')-I, dhfrA1, and ant(3")-Ia gene cassettes are available from Bulgaria in 2016 [35].

It seems that the IncA/C plasmids (e.g., plasmid 120kb, pPM91) carrying *blaviM-1-aacA7-dhfrA1-aadA1* gene cassettes are transferable between different Enterobacterales species [36]. Our data showed that these plasmids harbor intact conjugative elements, including the origin of transfer site oriT, T4CP, and T4SS. Thus, it can be concluded that these plasmids are effectively being transferred among bacteria. Moreover, the plasmid analysis indicated that IS elements including IS1, IS26-4, IS26-3, and IS6100 are frequently found in association of integron class 1 harboring blavIM-1-aacA7-dhfrA1-aadA1 array. The results of plasmid sequence alignment indicated that plasmids harboring blavIM-1-aacA7-dhfrA1aadA1 have high genetic similarities implying their high levels of conservation. Transferable plasmids harboring *bla*_{VIM-1} in ICREs pose a severe threat to public health. Further and continuous investigations are needed to detect CRE in clinical settings. Our epidemiological understanding requires to prevent the expansion of carbapenem-resistant Enterobacteriaceae in clinical settings.

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CONFLICT OF INTEREST

The authors declare that there are no conflicts of interest associated with this manuscript.

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J Med Microbiol Infect Dis

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