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VIM-2 β-LACTAMASE IN *PSEUDOMONAS AERUGINOSA* ISOLATES FROM ZAGREB, CROATIA

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- 16

18 ABSTRACT

The aim of this investigation was to characterize MBLs in *P. aeruginosa* isolates from
Zagreb, Croatia.

21 100 P. aeruginosa isolates with reduced susceptibility to either imipenem or meropenem were 22 tested for the production of MBLs by E test MBL. The susceptibility to a wide range of 23 antibiotics was determined by broth microdilution method. The presence of *bla*_{MBL} genes was detected by PCR. Hydrolysis of 0.1 mM imipenem by crude enzyme preparations of β-24 25 lactamases was monitored by UV spectrophotometer.. Outer membrane proteins were 26 prepared and analysed by SDS-PAGE. 27 Six out of 100 isolates were positive for MBLs by E test. All strains were resistant to 28 gentamicin, ceftazidime and cefotaxime and all except of one to imipenem. 29 Six strains positive for MBLs in E test were identified as VIM MBLs producers by PCR. 30 Sequencing of bla_{VIM} genes revealed the production of VIM-2 β -lactamase in all six strains. 31 This investigation proved the occurrence of VIM-2 β-lactamase among *P. aeruginosa* strains 32 from Zagreb, Croatia. VIM-2 β -lactamase with similar properties was previously described in 33 another region of Croatia, Italy, France, Spain, Greece, Taiwan and South Korea suggesting 34 that this type of enzyme is widespread in Mediteranean region of Europe and in Far East. 35 Key words: meropenem, imipenem, metallo- β -lactamases, resistance, *Pseudomonas* 36 aeruginosa

37 INTRODUCTION

38 Carbapenemases may be defined as β -lactamases that significantly hydrolyse at least

39 imipenem or/and meropenem [1]. The most clinically significant carbapenemases belong to

40 class B [2]. They are metallo- enzymes which require zinc as a cofactor and comprise four

41 families: IMP, VIM, SPM, GIM and SIM [3-6]. The VIM-type enzymes appear to the most

42 prevalent in Europe and Korea, and at least 10 different variants have been described [4]. The

worldwide spread of acquired metallo-β-lactamases (MBLs) in gram-negative bacilli has
become a great concern. MBLs possess a broad hydrolysis profile that includes carbapenems
and almost all extended-spectrum β-lactams except of aztreonam. Carbapenem resistant
strains of *P.aeruginosa* have been detected with increasing frequency in hospitals in Zagreb
and pose serious therapeutic problem. The aim of this investigation was to characterize MBLs
in *P. aeruginosa* isolates from Zagreb, Croatia.

49

50 MATERIALS AND METHODS

51 Bacteria

52 100 *P. aeruginosa* isolates with reduced susceptibility to either imipenem or meropenem were 53 tested for the production of MBLs by E test MBL (Solna, Sweden). The strains were isolated 54 during 2002 to 2004 at the Clinical Hospital Center Zagreb and University Hospital Merkur 55 in Zagreb from various clinical specimens and wards (Table 1). Isolates were identified by 56 conventional biochemical methods.

57 *E test MBL test*

58 The strip (AB Biodisk, Solna, Sweden) contains a double-sided seven-dilution range of

59 imipenem (4 to 256 mg/L) and imipenem (1 to 64 mg/L) in combination with a fixed

60 concentration of EDTA. An overnight culture of the test strains in Mueller-Hinton (MH) broth

61 was diluted to match the turbidity of 0.5 McFarland, swabbed on the plates of MH agar and

62 the plates were incubated at 35°C. A decrease of imipenem MIC by \geq 3 twofold dilutions in

63 the presence of EDTA was interpreted as being suggestive of MBL production [7].

64 Susceptibility testing

65 The susceptibility to a wide range of antibiotics was determined by twofold broth

66 microdilution method in cation supplemented MH broth in 96 well microtiter plates according

to CLSI [8]. The inoculum size was 5×10^5 CFU/ml. The plates were incubated for 18 h at 37

°C. The strains with MICs below the resistance breakpoint were tested with higher inoculum
of 10⁸ CFU/ml as well. The test was performed in triplicate. *P. aeruginosa* ATCC 27853 was
used for quality control. Antibiotic powders were obtained from the following manufacturers:
ceftazidime, gentamicin and clavulanic acid-Pliva, Zagreb; cefotaxime-Belupo, Koprivnica;
cefepime-Bristol Myers Squibb, Zagreb; imipenem-MSD, Zagreb; meropenem-AstraZeneca,
Zagreb.

74 Enzyme characterization

75 The bacterial cells were pelleted from exponential phase cultures in Lurian-Bentoni (LB) 76 broth by centrifugation at 6000 g. The pellet was washed and resuspended in cation 77 supplemented phospate buffer (0.1 mM, pH 7). Enzymes were released by sonication in ice 78 bath. Cell debris was removed by centrifugation at 10 000 g and supernatant was stored at -79 20°C as crude enzyme preparation. Hydrolysis of 0.1 mM imipenem by crude enzyme 80 preparations of β-lactamases was monitored by UV spectrophotometer at 298 nm. The change 81 of absorbance was recorded. Inhibition of enzyme activity was determined by 2 mM EDTA. 82 Enzyme activity was expressed as nmol of substrate hydrolyzed per minute expressed 83 relatively to the total protein content of the sample. Concentration of the protein in the 84 samples was determined with a commercial method (BioRad).

85 Analytical isoelectric focusing (IEF)

IEF was performed according to Matthew et al [9]. The organisms were grown overnight in
BHI broth, collected by centrifugation and β-lactamases were extracted from bacteria by
sonication. Cell debris was removed by centrifugation at 14 000 g. Supernatant was used as
crude enzyme. Isoelectric focusing was performed on polyacrylamide gels (acrylamide 7%,
bis-acrylamide 0.2%) containing ampholines with a pH range of 3.5 to 10. The β-lactamases
were detected by staining of the gel with nitrocefin, following IEF. β-lactamases of known

92 pI (isoelectric point) were used as standards: TEM-1, TEM-2, SHV-1, SHV-2, SHV-4 and
93 SHV-5.

94

95 Polymerase chain reaction and sequencing of bla_{VIM} genes

96 The presence of *bla*_{VIM} and *bla*_{IMP} genes was tested by PCR. Primers: VIM1F (5'-CAG-ATT-

97 GCC-GAT-GGT-GGT-TGG-3') and VIM1R (5'-AGG-TGG-GCC-ATT-CAG-CCA-GA-3')

98 were used to amplify bla_{VIM} genes. whereas primers IMP-A (5-GAA-GGY-GTT-TAT-GTT-

99 CAT-AC-3') and IMP-B (5'-GTA-MGT-TTC-AAG-AGT-GAT-GC-3') [1] were used for

100 detection of bla_{IMP} genes. The cycling conditions were as follows: 94°C-5min, and then 30

101 cycles-94°C-1 min, 55°C-1 min, 72°C-1-min and final extension at 72°C for 5 min. PCR

102 products were subjected to electrophoresis in agarose gel at 100 V. PCR products were

103 detected under UV light after staining with etidium bromide.

104 The amplicons were sequenced from both sides.

105 Plasmid analyis

106 Plasmids were extracted by alkaline lysis method as described previously [10]. Overnight

107 cultures of the test strains in LB broth (12 ml) were centrifuged at 14 000 rpm and the pellet

108 was resuspended in Solution I (25 mM Tris Hcl (pH 8), 50 mM Glucose, 10 mM EDTA (pH

109 8) containing 5 mg/ml lysozyme and 10 mg/ml RNA-se. The suspensions were mixed and

110 incubated on ice for 10 min before addition of the Solution II (0.2 N NaOH, 1% SDS). The

samples were then mixed genty by inversion of the tubes. After incubation on ice for 15 min,

112 Solution III was added (3 M K-Ac, pH 4.8). Solution III contained 120 ml 5 M K-Ac, 23 ml

113 gliacil acetic acid and 57 ml H₂0. Samples were again mixed by inversion of the tubes,

114 incubated on ice for 15 min and centrifuged for 15 min at 14 000 rpm at room temperature.

115 The supernatant was transferred to the clean tube avoiding any floating precipitate. Equal

116 volume of phenol/chlorophorm was added, samples were mixed and centrifuged for 5 min.

This step was repeated twice. The supernatant was transferred to the clean tube without touching the water /chlorophorm interface or the precipitate in this region and 0.6 volumes of isopropanol were added. The samples were incubated 30 min at room temperature and centrifuged at 14 000 rpm for 20 min. The supernatant was discarded and the pellet dried in the vacuum centrifuge. The pellet was washed once with 70% ethanol and resuspended in TE buffer. Samples were subjected to electrophoresis in 0.7 % agarose gel in TBE buffer. After staining with ethidium bromide, the DNA was visualised by ultraviolet light.

124

125 Genotyping of strains by pulsed-field gel electrophoresis

126 Isolation of chromosomal DNA was performed as described by Kaufman et al [11]. For each 127 isolate 1,0 ml (optical suspension density 0,6-0,7 at 540 nm) of an overnight culture grown in 128 BHI broth was pelleted by centrifugation at 10 000 rpm for 2 min. After being washed in 1 ml 129 SE buffer (75mM NaCl;25mM EDTA, Sigma), bacteria were resuspended in 500µl SE buffer 130 with 10 µl lysosime (Boehringer Mannheim GmbH). Next, 500 µl of this bacterial suspension 131 was mixed with 500 µl 2.0% low- melting-temperature agarose (InCert agarose: FMC 132 Bioproducts) and left to solidify. Solid agarose plugs were then incubated for 24h at 56^o C in 133 2ml of ESP buffer (1% N-lauril sarcosine; 0,5 M EDTANa2, pH 9,5; 500 µg/ml proteinase K, 134 Sigma). After 24h, the plugs were incubated at room temperature for 2 h in PMSF (phenylmethanesulfonyl-fluoride, *Aldrich*) and then washed three times for 30 min at 4° C 135 136 with TE buffer (10mM Tris-Hcl,pH 8, 0,1 mM EDTA, Sigma) before macrorestriction with 10U / 1 µl XbaI for 3 h at 37° C. Restriction fragments of DNA were separated by PFGE with 137 138 a CHEF-DRIII apparatus (Bio-Rad Laboratories) through 1% pulsed-field certified agarose (*Bio-Rad*) at a field strength of 6 V/cm for 20 h at 11° C; with pulses from 5 to 50 -s in 0.5 139 140 TBE buffer with thiurea (50mM, Sigma). A lambda ladder (Roche) was used as the molecular 141 size marker. After electrophoresis, gels were stained with ethidium bromide, rinsed, and

photographed under UV light. The PFGE patterns were compared following the criteria of
Tenover and coleagues for bacterial strain typing [12] and analysed by computer software
(*GelComparII*). The patterns obtained were compared by clustering methods (unweighted
pairgroup method with arithmetic averages) using the *Dice* coefficient. An optimization of
0,50% and position tolerance of 3,00% were applied during the comparison of PFGE
fingerprinting patterns.

148

149 Characterization of outer membrane proteins

150 Outer membrane proteins of six MBL positive P. aeruginosa strains were prepared as 151 described previously [13]. Cells were harvested from overnight cultures in LB broth and 152 resuspended in phosphate buffer. Proteins were released from the cells by sonication in ice 153 bath and collected by centrifugation at 10 000 g. After solubilization in 10 mM Tris HCl, 5 154 mM MgCl₂ and to 2% sodium lauroyl sarconisate for for 1 h at 37°C the insoluble OMPs 155 were recovered at 14 000 g. A second solubilization step was performed and the OMPs were 156 again pelleted as above. Prior to SDS PAGE, the OMPs were denatured by the addition of 157 3% SDS-5% β-mercaptoethanol and boiled for 10 min. Laemmli's sample buffer (62.5 mM 158 Tris/Hcl, 2% SDS, 10% glycerol, 5% mercaptoethanol) and electrode buffer (25 mM Tris (pH 159 8.3), 190 mM glycine, 0.1% SDS) were used.

Electrophoretic analysis of *P. aeruginosa* OMPs was performed in polyacrylamide gel. The gel contained 11% acrylamide, 0.54% bisacrylamide plus 0.2% SDS in 0.375 M Tris/HCl (pH 8.8). Gel was polymerised with 0.2% TEMED and 0.25% ammonium persulfate. Stacking gel contained 4% acrylamide, 0.1% bisacrylamide, 0.1% SDS in 0.125 M Tris /HCl (pH 6.8), and was polymerised as above. Staining was performed with 0.125% Coomassie brilliant blue R250 in 45% methanol, 10% acetic acid for about 30 min. Destaining was performed in 45% methanol, 10% acetic acid.

- 168 RESULTS
- 169 *E test*
- 170 Six out of 100 isolates were positive for MBLs by E test.
- 171 Prevalence
- 172 The prevalence of MBLs among carbapenem resistant isolates of *P. aeruginosa* was 6%.
- 173 Susceptibility testing
- 174 All strains were resistant to gentamicin, ceftazidime and cefotaxime. All except of strain
- 175 (132) were resistant to imipenem. Strains 6, 12, 132 and 135 were resistant to ciprofloxacin,
- strains 12, 22, 35 and 135 to aztreonam, strains 6, 12, 22 and 135 to cefoperazon and strains
- 177 22, 35 and 135 to cefepime. Resistance to piperacillin alone and combined with tazobactam
- 178 was observed in only one strain (135).
- 179 Enzyme characterization
- 180 The enzyme activity ranged from6 to 420 nmol/imipenem/min/mg of protein. Carbapenemase
- 181 activity was almost completely inhibited by 2 mM EDTA. (Table 2) .
- 182 IEF revealed a band with the pI of 5.3 in all strains.
- 183 Polymerase chain reaction and sequencing of bla_{VIM} genes
- 184 Six strains yielded an amplicon of 523 bp with primers specific for VIM β -lactamases.
- 185 Sequencing of bla_{VIM} genes revealed the production of VIM-2 β -lactamase in all six strains
- 186 (Fig. 1). No IMP MBLs producers were detected by PCR.
- 187 Plasmid analysis
- 188 No plasmid DNA was found.
- 189 Pulsed field gel electrophoresis
- 190 The strains showed distinct PFGE patterns and were not clonally related (Fig. 2).
- 191 Characterization of outer membrane proteins
- 192 Four of the tested strains lacked altered OmpD2 protein (Table 1).
- 193

194 DISCUSSION

This investigation proved the occurence of VIM-2 β-lactamase among *P. aeruginosa* strains from Zagreb, Croatia. VIM-2 β -lactamase was previously reported from another region in Croatia [14] suggesting that there is a regional spread of this resistance determinant. VIM-2 β -lactamase with similar properties was previously described in Italy [15], France [16], Spain [17], Greece [18], Taiwan [19] and South Korea [20] suggesting that this type of enzyme is widespread in Mediterreanean region of Europe and Far East. VIM-2 is the most prevalent MBL in Taiwan [21].

202 The fact that all strains displayed similar enzyme activities, but one strain (132) showed 203 markedly lower carbapenem MIC than the others suggests that there are other resistance 204 mechanisms involved in the strains with high carbapenem MIC such as loss of outer 205 membrane proteins detected in four strains (6, 12, 22, 135) or efflux. The strain with low 206 carbapenem MIC probably had higher permeability coefficient or less efficient efflux pump. 207 However, the carbapenem MICs rose above the resistance breakpoint for the strain 132 when 208 a higher inoculum which is more likely to occur in *in vivo* situation, was applied. It is possible 209 that *in vivo* in the presence of a high inoculum, high level resistance arises. Little clinical 210 information is available on the outcome of patients infected with carbapenem-susceptible 211 MBL producing *P. aeruginosa*. There is no concensus on the interpretation and reporting of 212 the antibiograms for the MBL producers. Four strains were resistant to aztreonam contrary to 213 the results obtained from other authors [16-17,22]. Aztreonam is not hydrolyzed by MBLs [1] 214 but resistance could be due to other mechanisms like overproduction of AmpC β–lactamase, 215 coexistace of other β -lactamase or efflux as reported previously [23]. The macrorestriction of 216 the Xba digested chromosomal DNA showed distinct patterns indicating that strains 217 harbouring bla_{VIM} genes are not clonally related. Thus it is more likely that the spread of MBL 218 genes is due to the genetic exchange between different clones. The occurrence of MBLs was

sporadic. The strains were obtained from various wards in two hospitals during a wide timeperiod.

221 Since all three attempts to transfer ceftazidime resistance to recipient strain and to isolate 222 plasmid DNA were unsuccessful we can conclude that gene cassettes harbouring *bla*_{VIM} genes 223 were located in the integrons [20] which are inserted in the chromosome and that their 224 dissemination in Croatia is due to the mobilization of the resistance genes. 225 Our strains harbouring VIM-2 β -lactamase were resistant to all β -lactam antibiotics, 226 aminoglycosides and fluoroquinolones and pose a serious therapeutic problem in our 227 hospitals. The fact that all strains were resistant to gentamicin suggests that *bla*_{VIM} genes are 228 located on integrons containing gene casettes with resistance determinants for 229 aminoglycosides. Since most of the strains were resistant to aminoglycosides and 230 fluoroquinolones as well only toxic compounds such as colistine are left to be administered as 231 a therapeutic agent. The prevalence of MBL positive P. aeruginosa among our carbapenem 232 resistant strains of this species is still low (6%) but there is a possibility of horizontal spread 233 of *bla*_{VIM} genes to *Enterobacteriaceae* which are more frequent pathogens. MBLs have 234 already been found in the members of the family Enterobacteriaceae [18-19]. The difficulties 235 in detecting MBLs producers in the routine laboratories together with the mobile nature of the gene cassettes carrying <u>blaving</u> genes facilitates their dissemination. Meropenem has been 236 237 extensively used in our hospitals and the presence of MBLs among carbapenem resistant 238 P. aeruginosa of different genotypes, isolated from different hospitals, although in low 239 frequency, underscores the need for restricted use of carbapenems and their systematic 240 surveillance. Constant and consistent surveillance of the MBL producing strains will be the 241 prime measure to prevent their further dissemination.

242

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	Epidemiologic chracteristics of P. aeruginosa strains							$MIC (mg/L)^{1}$									Porins	
Strain				PEGE	Date of													
No	Specimen	Hospital ²	Unit	type	isolation (D/M/Y)	IMI	MEM	CAZ	CAZ/CL	CTX	CFP	FEP	AMT	PIP	TZP	GM	CIP	OMPD ₂
6	urine	А	nephrology	1	15/01/2004	>128	64	16	>128	>128	>128	16	8	64	32	>128	16	-
12	stool	А	haematology	2	14/01/2003	64	32	>128	>128	32	>128	16	>128	64	32	>128	64	-
22	Tracheal aspirate	А	Paediatric ICU	3	15/02/2003	>128	32	32	64	>128	16	64	>128	32	16	64	0.5	-
35	Wound swab	А	oncology	4	25/02/2003	>128	32	16	32	32	4	32	>128	64	32	>128	4	+
132	Tracheal aspirate	В	Surgery ICU	4	09/08/2003	16	8	>128	>128	64	4	16	0.12	32	8	16	32	+
135	Bronchal	В	Internal ICU	5	07/01/2002	>128	64	>128	>128	>128	>128	>128	>128	>128	>128	>128	>128	-

Table1. Epidemiologic characteristics, minimum inhibitory concentrations (MIC) of various antibiotics against MBL positive P. aeruginosa strains and alteration of outer membrane proteins.

¹ Abbreviations-IMI-imipenem, MEM-meropenem, CAZ-ceftazidime, CAZ/CL-ceftazidime+clavulanic acid, CTX-cefotaxime, CFP-cefoperazone, FEP-cefepime, AMT-aztreonam, PIP-piperacillin, TZP-Tazobactam/piperacillin, GM-gentamicin, CIP-ciprofloxacin ² A- University Hospital Center-Zagreb, B-University Hospital Merkur

Strain No	Hydrolysis rate	Hydrolysis rate in the presence of 2 mM
	(nmol/min/mg protein)	EDTA
		(nmol/min/mg protein)
6	6 x 10 ⁻⁹	1 x 10 ⁻¹³
12	9 x 10 ⁻⁹	$1.8 \ge 10^{-11}$
22	4.2 x 10 ⁻⁸	$9.4 \ge 10^{-12}$
35	4 x 10 ⁻⁸	$3.8 \ge 10^{-12}$
132	3.5 x 10 ⁻⁸	2.7×10^{-10}
135	4×10^{-8}	1.2 x 10 ⁻⁹

Table 2. Hydrolysis rates against imipenem of metallo-β-lactamases produced by *P. aeruginosa* strains.