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1	Original Article
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3	Rapid detection of IMP, NDM, VIM, KPC and OXA-48-like carbapenemases from
4	Enterobacteriales and Gram negative non-fermenter bacteria by real time PCR and melt curve
5	analysis
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30	Keywords: PCR, detection, carbapenemase, Enterobacteriales, non-Fermenter

31 Abstract

32

Carbapenemase producing microorganisms are increasingly isolated and often associated with treatment failures and outbreaks. The need for reliable and timely detection and/or confirmation of carbapenemase production is paramount, therefore a real-time PCR assay targeting IMP, NDM, VIM, KPC and OXA-48-like carbapenemases was designed and validated.

37

All available allele variants of the above carbepenmases were downloaded from the Beta-Lactamase DataBase (<u>http://bldb.eu/</u>), aligned with Clustal Omega and primers designed using Primer-BLAST. Real-time PCR monoplexes were optimized for the QuantStudio 6-Flex (Applied Biosystems) using the PowerUp SYBR Green Master Mix (Life Technologies) and validated using a panel of 204 characterised strains carrying a wide range of beta-lactamases, sometimes in combination. Melt-curve analysis was used to confirm positive results.

44

The *in silico* approach allowed primers to be designed in conserved regions of the KPC and NDM alignments, while three primer sets for IMP and two for VIM were necessary to ensure amplification of the different variants. One primer set was designed for OXA-48-like, however it is unlikely to detect all variants. Expected results were obtained for all 204 tested strains, with 100% sensitivity and specificity. Melt-curve analysis showed consistent Tm results for KPC, NDM and OXA-48-like; differences were instead noted for IMP and VIM as likely consequence of higher variability in the PCR target regions. Inhibition was not observed.

52

53 The assay is rapid, easy to perform and implement. It enables unequivocal detection of IMP, NDM, VIM,

54 KPC and OXA-48-like carbapenemases even when more than one type is present simultaneously.

56 Introduction

57

Carbapenems are amongst the last line of defence to treat serious bacterial infections. However, carbapenemases (i.e. enzymes able to inactivate carbapenems) are increasingly identified in both Enterobacteriales and non-fermenter Gram-negative isolates. Carbapenemase producing organisms (CPO) are frequently associated with treatment failure and outbreaks, with the latter worsened by the fact that carbapenemases are often present on mobile genetic elements and can therefore easily spread between different bacteria [1].

64 In Wales, IMP, NDM, VIM, KPC and OXA-48-like (known as the "big five") are the carbapenemases predominantly identified and consistent with those found in the rest of the UK: in particular, IMP, NDM, 65 66 KPC and OXA-48-like in Enterobacteriales [2] and VIM in Pseudomonas species [3]. Phenotypic assays 67 employing inhibitors for the different enzymes are available for the detection of some carbapenemases. 68 For example IMP, NDM and VIM (Class B, metallo-ß-lactamase) enzymes are inhibited by chelating 69 agents like ethylenediaminetetraacetic acid (EDTA) and dipicolinic acid (DPA), while KPC (class A ß-70 lactamase) is inhibited by boronic acid; an inhibitor for OXA-48-like (class D) β -lactamases has not yet 71 been identified, although temocillin resistance can be used as an indicator of its presence [4]. These 72 phenotypic tests are both sensitive and specific predominantly for Enterobacteriales, while their 73 performance is poor for Pseudomonas species [5, 6, 7]. Furthermore, these tests require 24hrs 74 incubation to completion and can sometimes be difficult to interpret.

75 Carbapenemase screening in Enterobacteriales is performed in Wales according to EUCAST criteria, 76 i.e. meropenem inhibition zone < 28mm and/or meropenem MIC > 0.125 mg/mL [4] with isolates flagged 77 as potential CPOs sent to the Specialist Antimicrobial Chemotherapy Unit (SACU), Cardiff (UK) for 78 further investigation. For Acinetobacter and Pseudomonas species all carbapenem and carbapenem 79 plus piperacillin/tazobactam resistant isolates are referred accordingly. In SACU, phenotypic 80 confirmation of carbapenem resistance is performed and carbapenemase production investigated using 81 Neo-SensitabsTM combination disks (ROSCO, Denmark) and MBL IP/IPI (*Pseudomonas* only) and MBL 82 MP/MPI (Enterobacterales only) double ended E-Tests (bioMerieux, France), followed by in-house 83 block-based PCR to confirm positive results.

The number of isolates referred to SACU for carbapenemase confirmation doubled from approximately 250 to 500 per year between 2015 and 2018, with an accompanying 100% rise in the number of CPO

86 confirmed per year (from 19 to 38). Nevertheless, the incidence in Wales (population approximately 87 3,125,000 inhabitants estimated in June 2017 [8]) remains low when compared to data reported from 88 other Countries [9]. Referral numbers are predicted to rise further as hospitals undertake more rigorous 89 screening programmes to detect CPO in admission patients as per recommendations [10]. The increase 90 in requests for carbapenemase confirmation plus the importance of rapidly detecting CPO for effective 91 infection control management, demanded a development of the above described confirmatory approach 92 adopted by SACU. These methods are time consuming, labour intensive and involve the use of the 93 toxic and carcinogenic compound, ethidium bromide, to visualise PCR results by gel electrophoresis 94 (although safer intercalating agents are available nowadays). The need for a more rapid and reliable 95 approach has been identified with the aim to reduce the time to issue confirmatory reports and better 96 assist clinicians and infection control teams. Real-time PCR is a well-established methodology for 97 detecting genetic markers in bacteria. It can provide results much faster than block-based PCR and 98 does not normally require the use of high risk reagents. Therefore, a real-time PCR assay based on 99 SYBR Green chemistry has been designed to detect the "big five" carbapenemases and thoroughly 100 validated using a large panel of previously characterised Gram-negative isolates.

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102

103 Materials and Methods

104

105 Assay design

106 IMP, NDM, VIM, KPC and OXA-48-like allele sequences were downloaded (as of December 2018) from 107 the Beta-Lactamase DataBase [11] and aligned using Clustal Omega [12]. Alleles with less than 90% 108 sequence identity to the rest of the alignment were excluded and separate primers were designed where 109 a strain carrying them was available to the authors for *in vitro* testing. Ten different primer sets per 110 target were designed using Primer-BLAST [13] and then compared to the relevant alignment to identify 111 the pair that at least in silico was able to amplify the highest number of alleles. Where necessary, degenerate bases (a maximum of 2 per primer) were added to cover non-conserved positions. Primer 112 113 sequences and amplicon sizes are detailed in Table 1.

114

115 Bacterial strains

116 A total of 204 previously characterised Gram-negative isolates from 19 different species isolated world-117 wide were selected from the Authors', Public Health England, NCTC and ATCC collections (Supplementary data). One hundred and thirteen carried one of the "big five" carbapenemases while 118 119 22 carried two different markers simultaneously. The remaining 69 were negative for "big five" 120 carbapenemases however some carried genes for other β -lactamases (such as AmpC, CTX-M, GES, IMI, L1 & L2, OXA-10, OXA-23, OXA-51, OXA-58, TEM, SHV, SME, SPM and VEB). The 135 "big five" 121 122 CPOs carried a total of 40 IMPs (including IMP-1, IMP-4, , IMP-6, IMP-7, IMP-8, IMP-10, IMP-11, IMP-13, IMP-14, IMP-29 and IMP-62), 27 VIMs (including VIM-1, VIM-2, VIM-4, VIM-7, VIM-10, VIM-19 and 123 124 VIM-29), 31 NDMs (NDM-1, NDM-3, NDM-4, NDM-5, NDM-7), 26 KPCs (including KPC-2, KPC-3, KPC-125 4 and KPC-23) and 33 OXA-48-like markers (including OXA-48, OXA-181, OXA-204, OXA-232, OXA-126 244 and OXA-245).

127

After overnight aerobic growth on blood agar at 35±1°C, bacterial growth approximately equivalent to a 128 129 third of a loopful (10µL loop) was re-suspended in 250 µL of nuclease-free water and heat killed at 130 100°C for 10 min. Supernatant was separated by centrifugation at 12,000 g for 2 min and diluted 1:20 131 in nuclease-free water prior to testing. An extraction control (i.e. 250 µL of nuclease-free water) was 132 always included to discount cross contamination. A panel of seven strains was used as positive control in every experiment, namely Escherichia coli NCTC 13476 (IMPa), Serratia marcescens SACU 1212 133 134 (IMPb), Pseudomonas aeruginosa SACU 1205 (IMPc), Klebsiella pneumoniae NCTC 13440 (VIM), 135 Klebsiella pneumoniae NCTC 13443 (NDM), Klebsiella pneumoniae NCTC 13438 (KPC) and Klebsiella 136 pneumoniae NCTC 13442 (OXA-48).

137

138 Internal process control

An assay detecting a 76 bp fragment of the *green fluorescent protein, gfp* from *Aequorea victoria* [14]
was included to discount PCR inhibition. A custom-made plasmid (Eurofins, Germany) containing the
entire *gfp* sequence (Genbank Accession: M62653) was diluted to a working concentration of 1 pmol/μL
before being added to the GFP reaction mix.

- 143
- 144 Real Time PCR

145 Monoplex PCR assays were optimized for the QuantStudio 6-Flex (Applied Biosystems) using the 146 PowerUp SYBR Green MasterMix (Life Technologies) and Microamp Fast Optical 96-Well Reaction 147 Plate 0.1 mL (Thermo-Fisher Scientific). Briefly, reactions were performed in a final volume of 10 µL 148 containing 5 µL of MasterMix, 2.5 µL of template, 2 µL of PCR grade water and 0.5 µL of relevant primer 149 mix. The final concentration of each primer is listed in Table 2. The GFP reaction mix also contained 150 0.5 μ L per reaction of pGFP [1 pmol/ μ L], with 1.5 μ L of PCR grade water to make 10 μ L final volume. 151 After an initial uracil-DNA-glycosylase step at 50°C for 2 min, the Dual-Lock™ DNA polymerase was 152 activated at 95°C for 2 min followed by 35 cycles of denaturation at 95 °C for 1 sec and 153 annealing/extension at 60°C for 20 sec. Amplification results were analysed with the threshold set at 154 0.5 ΔRn for all targets and the baseline set between 5 and 15 cycles only for the GFP target. Melting 155 curve analysis was performed as follows: 95°C for 15 sec (ramp rate = 1.6°C/sec), 60°C for 1 min (ramp 156 rate = 1.6°C/sec) and 95°C for 15 sec (ramp rate = 0.15°C/sec) with fluorescence fluctuation analysed 157 during the latter.

- 158
- 159
- 160 Results

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162 In silico analysis

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The *in silico* approach allowed primers to be designed within 100% conserved regions of the KPC and NDM alignments. For the remaining carbapenemases, a single primer set able to amplify all alleles, was not identified. Three primer sets (tested separately) were necessary to detect IMP variants and two sets (tested in the same tube) for VIM variants. One primer set was designed for the OXA-48-like group, however despite detecting the most common variants it is unlikely to detect OXA-54, OXA-436 and OXA-535 due the suboptimal sequence identity (i.e. < 90%) to the rest of the group. . Clustal alignments of PCR fragments with primers binding sites highlighted are available in the Supplementary data.

171

Thirty-seven KPC variants were listed on the Beta-Lactamase DataBase, however two sets (i.e. KPC1 & KPC-2 and KPC-9 & KPC-23) were identical while KPC-20 was "assigned" but no sequence was
available. Specific primers were designed in a conserved region of the remaining 34 KPC alleles. Primer

binding sites were 100% conserved and no single nucleotide polymorphisms (SNPs) were identified inthe region between the primers.

177

A specific primer set was designed in a conserved region of the alignment of all available 24 NDM
variants, with no exclusion. Primer binding sites were 100% conserved and only one SNP (found in
NDM-10) was identified in the region between the primers.

181

Among the 79 described IMP variants, six (i.e. IMP-36, IMP-39, IMP-46, IMP-50, IMP-57 and IMP-65) appeared as "assigned" but no sequence was available. Moreover, IMP-8 and IMP-47 share identical sequence, consequently, the actual number of analysed variants was 72. Clustal analysis showed considerable sequence variability, therefore it was not possible to identify a single primer set able to detect all alleles. The Clustal generated tree was arbitrarily split in three sections (Figure 1) and a set of primers designed for each section (i.e. IMPa, IMPb and IMPc). Degenerate bases had to be inserted in all six primer sequences to ensure amplification of different variants.

189

Among the 60 described VIM variants, VIM-21, VIM-22 and VIM-55 were "assigned" but no sequence was available. VIM-7 showed low sequence identity with the rest of the alleles (e.g. 81% identity with VIM-1), consequently it was excluded from the alignment. Specific primers were designed in a conserved region of the remaining 56 VIM alleles and one degenerate base was introduced in the forward primer. As previously mentioned, a separate set of primers was also designed specifically for VIM-7 as this variant was available to the authors for *in vitro* testing.

196

197 Only OXA-48-like variants with a definitive assignment by NCBI (30 in total) were downloaded from the Beta-Lactamase DataBase (i.e. OXA-48, OXA-54, OXA-162, OXA-163, OXA-181, OXA-199, OXA-204, 198 199 OXA-232, OXA-244, OXA-245, OXA-247, OXA-252, OXA-370, OXA-405, OXA-416, OXA-436, OXA-200 438, OXA-439, OXA-484, OXA-505, OXA-514, OXA-515, OXA-517, OXA-519, OXA-535, OXA-538, 201 OXA-546, OXA-547, OXA-566 and OXA-657). After Clustal analysis, OXA-54, OXA-436 and OXA-535 202 were excluded from the alignment due to low sequence identity (84%, 84% and 85% respectively with OXA-48). In this instance specific primers were not designed as they were not available to the authors 203 204 for in vitro testing. Primers for the remaining 27 OXA-48-like variants were designed in a conserved 205 region with one ambiguity introduced within the forward primer while the binding site of the reverse206 primer was 100% conserved.

207

208 In vitro analysis

209

210 The eight monoplex real-time PCR assays (i.e. IMPa, IMPb, IMPc, NDM, VIM, KPC, OXA-48 and GFP) 211 were performed in 96-well plates with PCR mixes added in rows; DNA extracts were tested in column 212 1 to 10, while column 11 and 12 were used to test extraction and positive controls respectively. The 213 time necessary to complete PCR amplification and melt curve analysis was approx. 40min. Testing of 214 undiluted DNA extracts resulted in strong PCR inhibition (as revealed by absence of- or delayed GFP 215 amplification), consequently 1:20 dilutions were prepared prior to testing. Formation of primer dimers 216 was observed with the NDM assay, therefore oligonucleotide sequences were reviewed in silico and 217 concentrations decreased in vitro to improve results (Table 2).

218

DNA extracts of *Escherichia coli* NCTC 13476 (IMP), *Klebsiella pneumoniae* NCTC 13440 (VIM-1), *Klebsiella pneumoniae* NCTC 13443 (NDM-1), *Klebsiella pneumoniae* NCTC 13438 (KPC-3) and *Klebsiella pneumoniae* NCTC 13442 (OXA-48) were mixed to generate all possible target combinations (for simplicity only IMPa among the three IMP primer sets was included in this analysis). In all cases, the assay successfully detected all the carbapenemases present in the reaction mix, regardless of if only one or all five were present simultaneously (Supplementary data).

225

226 Melt-curve analysis was used to confirm positive results by comparing dissociation curves to that of the 227 relevant positive control. Minor Tm variations were observed for NDM, VIM, KPC, OXA-48-like and GFP 228 (0.15 < St Dev < 0.30), while greater differences were noted with the three IMP monoplexes (0.49 < St 229 Dev < 0.62).

230

The expected result was initially obtained for 203 of the 204 previously characterised isolates. *Klebsiella pneumoniae* SACU 27698 was meant to be negative for the "big five" carbapenemases, instead it produced unequivocal amplification with IMPa (Ct = 17.49) and IMPb (Ct = 30.49) primer sets and melting curves consistent with those of *E. coli* NCTC 13476 and *S. marcescens* SACU 1212 (i.e. IMPa

and IMPb positive controls). Further analysis with in-house block-based PCR confirmed presence of
IMP metallo-β-lactamase, while Whole Genome Sequencing (WGS) by MiSeq (Illumina) characterised
it as IMP-4 (data not shown). Indeed, the newly designed assay provided the expected result for all 204
tested isolates (100% sensitivity and specificity) including those carrying two "big five" carbapenemases
simultaneously. Cross-reaction with the other resistance markers present in the above mentioned
isolates was not observed.

241

Overall, amplification results produced mean Ct values of ca. 16 and Standard Deviation (SD) values between 0.88 and 1.45 (Table 3). Among the 40 isolates carrying IMP markers, 36 gave positive results with two of the three IMP primer sets used (i.e. a strong result in the expected Ct range and a weak result with Ct > 25 respectively). IMP alleles from part A of the Clustal generated tree gave strong IMPa results and weak IMPb results, alleles in part B gave strong IMPb results and weak IMPc results, while alleles in part C of the tree gave IMPc strong results and weaker IMPb results (Supplementary data). Four IMPs of which three IMP-13, were only amplified by one set of primers.

249

250

251 Discussion

252

253 We describe here the design and validation of an array of monoplex real-time PCRs optimised for the 254 QuantStudio 6-flex platform (ThermoFischer Scientific) to rapidly detect and differentiate the "big five" 255 carbapenemases, (i.e. IMP, NDM, VIM, KPC and OXA-48) from Enterobacteriales and non-fermenter 256 Gram-negative isolates. A rational approach that took into account all the relevant allele sequences 257 available from the Beta-Lactamase Database (as of December 2018) was applied in silico to design 258 specific primers and to then predict whether they would also detect those allele variants that were not 259 available to the authors for in vitro testing. A wide collection of previously characterised Gram-negative 260 isolates was then tested in vitro to fully validate the newly designed monoplexes. An assay targeting 261 the gfp [14] was simultaneously tested as an internal control to discount inhibition.

262

As resistance markers (especially IMP in this case) evolve rapidly causing emergence of new allele variants, SYBR Green chemistry combined with melting curve analysis in monoplex PCR reactions was 265 chosen over multiplexing with primers and probes. Newly described SNPs in primers and/or probe 266 binding sites would potentially result in assays having to be redesigned and then revalidated. This is 267 time consuming and costly. The absence of probes in this assay reduces the number of binding sites from three to two and consequently the likelihood that a newly described SNP falls into them. If indeed 268 269 a new SNP is identified in either of the two primer binding sites and it is decided that they need to be 270 redesigned, the use of separate monoplexes allows to limit the revalidation to just the affected monoplex 271 instead of having to perform a major revalidation like when a multiplex approach is used. Furthermore, 272 the use of separate monoplexes provides greater flexibility to combine testing of different panels of 273 targets if the need arises: one or more of the monoplexes described here could be quickly replaced with 274 others validated separately to detect different resistance markers (e.g. GIM, SIM, SPM, IMI, etc) without 275 the need to perform any further validation on the entire assay panel.

276

The *in silico* analysis showed that this assay should be able to detect all the IMP, NDM, VIM and KPC variants described as of December 2018. Among the OXA-48-like group, OXA-54 and OXA-535, described in the chromosome of *Shewanella species* [15, 16] and OXA-436 identified on a plasmid in *Enterobacter species*, *Citrobacter species* and *K. pneumoniae* in Denmark [17] are unlikely to be detected.

282

The in vitro experiments proved the assay to be 100% sensitive and specific on a large panel of 283 284 previously characterised isolates available to the authors. As stated by Lund et al. (2018), assays are 285 often validated using a limited selection of target variants that are available locally; furthermore, testing 286 high numbers of unsequenced isolates might simply result in redundant analysis of a small number of 287 variants which then provides little indication about the actual sensitivity and specificity of a particular 288 assay. These issues were addressed in this study by applying a robust in silico approach to confidently 289 predict amplification of those variants that were not available for *in vitro* testing and by then including 290 (where possible) isolates where the "big five" allele variant had previously been characterised (i.e. by 291 Sanger sequencing and/or by WGS).

292

293 Interpretation of melting profiles can be problematic when dissociation curves do not overlap with that 294 of the positive control. This was often observed with isolates carrying IMP genes as a likely

295 consequence of high sequence variability of the IMP PCR products. Nevertheless, when an IMP allele 296 was present, more than one primer set was often able to detect it (usually two primers sets, one producing a strong result with Ct between 14 and 18, and another producing Ct > 18) allowing confident 297 298 identification of true positives. It is noteworthy that variations in Tm results were observed also for those 299 PCR products that were not meant to contain any SNP (e.g. KPC and GFP) therefore it seems that the 300 actual Tm value cannot be reproducibly obtained by the PCR platform between different PCR runs. A 301 Tm standard deviation of 0.20 was calculated for the GFP amplified fragment (which is identical in every 302 PCR run) and this value could be considered as an indication of the Tm uncertainty of measurement 303 intrinsic to this procedure. Different Tm values were also noted for VIM alleles, however in this case, it 304 was due to VIM-7 being quite different (only 81% nucleotide identity with VIM-1), with a Tm of approx. 305 84°C compared to Tm of approx. 81°C for the remaining VIM alleles. This was consistently observed 306 and was actually helpful to differentiate between VIM-7 and the other VIM alleles.

307

308 In at least one instance, the assay provided more sensitive and reliable results than standard phenotypic 309 analysis: K. pneumoniae SACU 27698 previously characterised as negative for the "big five" 310 carbapenemases, was instead found to be IMP positive and later characterised by WGS to carry IMP-311 4. This result triggered an investigation that identified the above isolate as part of a cluster involving five 312 patients (data not shown) which clearly proves the added value of the here described real time PCR 313 assay. Retrospective analysis on the isolate showed that combination disks did not to show metallo-β-314 lactamase activity (i.e. only 2 mm difference between the meropenem and the meropenem/DPA disks) 315 while the MP/MPI double-ended strip provided a weakly positive result (MIC ratio = 12). With this in 316 mind, it was decided to modify the SACU approach for investigation of carbapenemases. The real time 317 PCR assay described here became the first line method to investigate presence of the "big five" 318 carbapenemases. Positive PCR results are reported immediately without performing further analysis, 319 however as negative PCR results cannot completely exclude presence of a carbapenemase, further 320 investigations need to be performed. As already mentioned, new "big five" allele variants might not be 321 detected by this assay or a carbapenemase not included in this assay (e.g. GIM, SIM, SPM, etc) could 322 be present. Finally, novel carbapenemases, not known as yet, would equally not be detected. 323 Therefore, all PCR negative isolates are tested with combination disks and double ended strips to 324 phenotypically exclude both carbapenemase and Extended Spectrum ß-lactamase (ESBL) activity before negative results are reported. In those cases where carbapenemase activity is observed despite a negative PCR results, alternative methods (such as sequence analysis) would be applied to investigate the cause of the resistant phenotype.

328

329 In certain cases, "big five" variants detected by this assay might actually confer either weak or 330 undetectable level of carbapenemase resistance. Indeed, KPC-14 and KPC-28 have been 331 demonstrated to have reduced carbapenemase activity [19] while OXA-163, OXA-247 and OXA-405 have been shown to have no carbapenemase activity [20, 21] due to amino acid deletions in the active 332 333 site. Isolates carrying these variants are likely to generate discrepant PCR and phenotypic results thus 334 further testing (e.g. sequence analysis of the "big 5" variant) should be performed. In particular, KPC-335 28 has also been characterised to have an amino acid substitution (i.e. H274Y) causing a 50-fold 336 increase in ceftazidime hydrolysis and consequently high MIC to ceftazidime/avibactam, an effective 337 antimicrobial option for KPC producing bacteria [19].

338

339 The assay described herein allows testing of 10 isolates at a time (plus extraction control and positive 340 control) using 96-well plates. Given the short PCR running time (ca. 40 min), DNA could be extracted 341 simultaneously from a greater number of isolates to allow a quick second PCR experiment to be 342 prepared while the first is running; up to 20 isolates could be then tested in less than 90 min. 343 Alternatively, for high through put 384-well plates may be used, allowing up to 46 strains (plus controls) 344 to be tested simultaneously. The assay was deemed easy to perform by members of staff with little or 345 no experience with real-time PCR and melt-curve analysis. Importantly, the real time PCR assay allowed unequivocal result interpretation even when more than one "big five" carbapenemase 346 347 resistance marker was present. Finally, the new algorithm improved the reference service by reducing 348 the time necessary to report positive results and at the same time reducing the risk of reporting false-349 negative results.

350

In summary the assay was easily implemented, allowed faster and more reliable positive results than previously obtained by SACU to be available for clinicians and infection control specialists. The potential wider benefits of the assay include reducing costs to the hospital from keeping potential positive patients

in side rooms, inappropriate therapy of positive patients and increased screening costs because ofdelays in positive results.

356

357

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365 Compliance with Ethical Standards

- 366
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- 368 **Conflict of interest.** The authors declare that they have no conflict of interest.
- 369 **Ethical approval.** Not required for this study.
- 370 **Informed consent.** Not required for this study.

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Table 1. Summary of primer sets used in this study (FP = Forward Primer; RP = Reverse Primer;

457 ambiguities are <u>underlined;</u> * VIM and VIM-7 tested in the same tube).

Primer set	Target	Name	Sequence (5′→3′)	Amplicon (bp)	Reference	
1	IMP(a)	IMPa_FP TGACGCAAATDTAGAAGCTTGGC		101	This study	
1		IMPa_RP	C <u>R</u> TCTCCA <u>R</u> CTTCACTGTGAC	101	This study	
	IMP(b)	IMPb_FP	GA <u>Y</u> GCCTATCT <u>R</u> ATTGACACTCC	70	This study	
2		IMPb_RP	TATA <u>R</u> CCGCGCTC <u>M</u> ACAAACCA	79	This study	
2		IMPc_FP	<u>K</u> GA <u>Y</u> GCAAATTTAGAAGCTTGGCCA	140	This study	
3	IMP(c)	IMPc_RP	CCCTTTAACAGCCTG <u>Y</u> TCCCA	142	i nis study	
	NDM	NDM_FP	CAACTTTGGCCCGCTCAAGG	100	This study	
4		NDM_RP	GCAGCCACCAAAAGCGATGT	100		
	VIM	VIM_FP	CGAAAAACACAGCGGC <u>M</u> CTTCT	70	This study	
5 *		VIM_RP	GTGGAGACTGCACGCGTTAC	73	This study	
5	VIM-7	VIM-7_FP	GGTGGCTGTGCAGTTCATGAG	100	This study	
		VIM-7_RP	ACGACCTCTGCTTCCGGATAC	122	i nis study	
6	KPC	KPC_FP	CCACTGGGCGCGCACCTATT	51	This study	
0		KPC_RP	TGTTAGGCGCCCGGGTGTAG	51	This study	
7	OXA-48-like	OXA-48_FP	CTGGTGGGT <u>Y</u> GGTTGGGTTGA	80	This study	
		OXA-48_RP	GCAGCCCTAAACCATCCGATGT	09	THIS SLUUY	
9	GFP	GFP_FP	CCTGTCCTTTTACCAGACAACCA	76	[1/]	
ð		GFP_RP	GGTCTCTCTTTCGTTGGGATCT	70	[14]	

Primer set Target		Primer Name	Final Conc. [nM]		
4	IMDo	IMP(a)_FP	500		
I	IIVIFa	IMP(a)_RP	500		
2	IMDb	IMP(b)_FP	500		
2	INFD	IMP(b)_RP	500		
3	IMPo	IMP(c)_FP	500		
3	INIEC	IMP(c)_RP	500		
4		NDM_FP	300		
4		NDM_RP	300		
	V/IM	VIM_FP	400		
5*	VIIVI	VIM_RP	400		
5	VIM-7	VIM7_FP	100		
	V 11VI-7	VIM7_RP	100		
6	KPC	KPC_FP	500		
U	KF C	KPC_RP	500		
7		OXA-48_FP	400		
· ·		OXA-48_RP	400		
8	GED	GFP_FP	300		
0	GFF	GFP_RP	300		

Table 2. Final concentration of primers used in this study (* VIM and VIM-7 tested in the same tube).

Table 3. Summary of Cycle threshold (Ct) and melting temperature (Tm) results for the primer sets
463 used in this study (* VIM and VIM-7 tested in the same tube; ** only one strain containing VIM-7 was
464 available for testing, consequently Min, Max and SD were not calculated).

Primer	Target	Ct				Tm (°C)			
set		Min	Mean	Мах	SD	Min	Mean	Мах	SD
1	IMPa	14.67	16.73	20.35	1.43	76.92	77.74	78.47	0.49
2	IMPb	14.43	16.66	18.94	1.45	75.13	76.31	77.21	0.77
3	IMPc	14.88	16.06	16.98	0.88	76.92	77.51	78.06	0.62
4	NDM	13.73	16.41	18.83	0.99	84.61	84.86	85.58	0.21
5*	VIM	14.11	16.09	18.23	0.95	80.43	80.73	81.07	0.15
5	VIM-7**	-	15.49	-	-	-	84.12	-	-
6	KPC	13.52	16.81	18.50	1.13	82.90	83.40	83.83	0.30
7	OXA-48-like	13.63	16.16	18.60	1.21	78.05	78.26	78.78	0.18
8	GFP	21.36	23.36	24.28	0.44	77.51	77.77	78.83	0.20

Figure 1. Clustal Omega generated tree of the IMP alleles available from the Beta-Lactamase 468 DataBase.

