

1 **Use of a Commercially Available Microarray to Characterize**
2 **Antibiotic Resistant Clinical Isolates of *Klebsiella pneumoniae***

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31 **Abstract**

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A commercially available microarray (IDENTIBAC AMR-ve) for the detection of antibiotic resistance determinants was investigated for its potential to genotype 30 clinical isolates and two control strains of *Klebsiella pneumoniae*. Resistance profiles and the production of extended spectrum β -lactamases were determined by disc diffusion and the results were compared with the microarray profiles in order to assess its scope and limitations. Genes associated with resistance to a wide-range of antibiotics, including current first line therapy options, were detected. In addition, the array also detected class 1 integrases. The array is easy to use and interpret, and is useful in providing a general description of the numbers and types of resistance determinants in *K. pneumoniae*. It also provides an indication of the potential for resistance gene acquisition. However, in most instances detected resistance to specific antibiotics could not unequivocally be assigned to hybridization with a specific array probe. We conclude that the microarray is a valuable and rapid means of investigating the presence of resistance gene-classes of therapeutic importance. It can also provide a starting point for selecting analyses of greater resolving power, such as phylogenetic subtyping by PCR-sequencing.

66 Introduction

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68 Infections caused by drug-resistant bacteria are a major problem worldwide. Resistance to antimicrobial agents
69 by members of the *Enterobacteriaceae* including *Klebsiella pneumoniae* is of particular concern. *K. pneumoniae*
70 is encountered as a saprophyte in humans and other mammals, colonizing the gastrointestinal tract, skin, and
71 nasopharynx [29]. It can cause a wide range of infections including those of the urinary and lower biliary tracts,
72 as well as of surgical wounds and the blood stream. In recent years, *K. pneumoniae* has become established as an
73 important cause of nosocomial infections, acquiring resistance to antibiotics through plasmid-encoded extended-
74 spectrum β -lactamases (ESBLs), along with a variety of genes conferring resistance to other drug classes. The
75 most prevalent ESBLs in *K. pneumoniae* are CTX-M enzymes belonging to Ambler class A [3]. In general, these
76 enzymes have hydrolytic activity against penicillins, extended-spectrum cephalosporins (e.g. cefotaxime and
77 ceftazidime) as well as monobactams (aztreonam), but are generally not effective against cephamycins (e.g.
78 cefoxitin) and carbapenems [25]. The activity against ceftazidime is variable between the different CTX-M types
79 and certain variants also have activity against 4th generation cephalosporins (e.g. cefepime) [10]. Since the initial
80 isolation of CTX-M-1 from a European patient in the late 1980s [8], >170 CTX-M allelic variants have been
81 identified [<https://www.ncbi.nlm.nih.gov/projects/pathogens/beta-lactamase-data-resources/>]. CTX-M variants
82 can be divided into 5 major phylogenetic groups, CTX-M group 1, CTX-M group 2, CTX-M group 8, CTX-M
83 group 9 and CTX-M group 25 on the basis of their amino acid sequences [32 and references therein].

84 Tzouveleki et al., [29] noted that a successive addition of genetic elements encoding resistance to
85 aminoglycosides and ESBLs, coupled with the rapid accumulation of chromosomal mutations conferring
86 resistance to fluoroquinolones, left carbapenems as of around the year 2000, as the first-choice drugs for the
87 treatment of health care-associated infections caused by *K. pneumoniae*. Subsequently there has been a rapid
88 dissemination of multidrug-resistant (MDR) *K. pneumoniae* strains producing carbapenemases encoded by
89 transmissible plasmids. The clinically most important carbapenemases include the class A enzymes of the KPC
90 group, zinc-dependent class B metallo- β -lactamases, represented mainly by the VIM, IMP, and NDM types, and
91 class D carbapenemases of the OXA-48 class [reviewed in 29]. Carbapenem-resistant *K. pneumoniae* (CRKP)
92 have emerged as an important cause of morbidity and mortality from hospital-acquired and long-term care-
93 associated infections [15, 20]. Few antimicrobial therapy options now exist for infections caused by CRKP [26].

94 The present study evaluates the potential for a commercially available DNA microarray to rapidly and
95 correctly detect resistance genes of clinical importance in *K. pneumoniae*. Furthermore, the scopes and

96 limitations of the array technique in explaining the results from antimicrobial susceptibility testing were
97 evaluated. The DNA-based assay [IDENTIBAC AMR-ve Genotyping kit Version 05; CLONDIAG/ALERE,
98 GmbH] was developed and validated for the parallel detection of resistance and integrase genes in *Escherichia*
99 *coli* and *Salmonella* [5], but has not been much used with *K. pneumoniae*. Card et al., [9] report the use of an
100 extended array, containing many of the same probes as AMR-ve, in the typing of *K. pneumoniae* (34 strains
101 distinct from those in the present study), and other *Enterobacteriaceae*. Other studies have screened respectively
102 one [27] and two [31] strains of this species.

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121 **Materials and methods**

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123 **Isolates and control strain**

124 The strain collection consisted of 30 clinical *K. pneumoniae* isolates. Strains were supplied by the Norwegian
125 National Advisory Unit on Detection of Antimicrobial Resistance (Tromsø, Norway). *K. pneumoniae* ATCC
126 700603 (SHV-18- β -lactamase producing) and *K. pneumoniae* ATCC 33495 (non-ESBL-producing) were used
127 for control purposes.

128 **Antibiotic resistance testing**

129 Disc diffusion assays were performed and interpreted according to version 6.0 of the recommendations of the
130 European Committee on Antimicrobial Susceptibility Testing (EUCAST, <http://www.eucast.org>). In brief,
131 susceptibility testing was done using Mueller–Hinton agar (Oxoid, Basingstoke UK). The inoculum was made
132 from cells grown at 37 ± 1 °C for 18-24 h on tryptone soya agar (TSA, Oxoid) and adjusted to 0.5 McFarland.
133 After applying antibiotic discs (Oxoid), plates were incubated at 35 ± 1 °C for 16-20 h prior to the reading of
134 zone diameters. Based on zone diameters, strains were recorded as susceptible (S), intermediate susceptible (I) or
135 resistant (R) to each antibiotic according to the guidelines.

136 The following antibiotics and concentrations were tested: ampicillin (AMP, 10 μ g), amoxicillin/clavulanate
137 (AMC, 30 μ g), piperacillin/tazobactam (TZP, 30/6 μ g), cefotaxime (CTX, 5 μ g), ceftazidime (CAZ, 10 μ g),
138 cefuroxime (CXM, 30 μ g), meropenem (MEM, 10 μ g), ciprofloxacin (CIP, 5 μ g),
139 gentamicin (CN, 10 μ g), trimethoprim-sulphamethoxazole (SXT, 25 μ g) and aztreonam (ATM, 30 μ g).

140 **Disc-diffusion screening for β -lactamase production**

141 Phenotypic detection of ESBLs was performed on all strains using a double disc synergy test consisting of
142 cefpodoxime 10 μ g and cefpodoxime/clavulanic acid 10/1 μ g (Combination kit DD0029; Oxoid). An increase of
143 ≥ 5 mm between the cefpodoxime and cefpodoxime/clavulanic acid discs was interpreted as a positive test
144 according to the manufacturer's instructions. Based on indications from disc diffusion susceptibility and
145 microarray testing, some isolates were also screened for the presence of metallo- β -lactamases and KPC enzymes
146 (class A) using the KPC/MBL Confirm Kit (ROSCO, Denmark) and the ESBL and AmpC Screen kit (ROSCO)
147 according to the manufacturer's instructions. All tests were performed using Mueller-Hinton agar (Oxoid) as
148 described for the antibiotic susceptibility testing.

149 **AMR-ve Microarray**

150 A DNA-based assay (IDENTIBAC AMR-ve Genotyping kit Version 05; CLONDIAG/ALERE, GmbH) with 89
151 probes for different genetic markers including resistance gene families and integrases was used. The kit is able to
152 detect ~57 antimicrobial resistance genes in Gram-negative bacteria, including several ESBLs. Basic details of
153 the method and the probes used are given in the original publication [5] and the kit protocol (available at
154 www.identibac.com). In brief, *K. pneumoniae* was grown for 18-24 h at 37 °C on TSA. Genomic DNA was
155 isolated from a 1mm loopfull of cells (about 1×10^9 cells) using the DNeasy® Blood and Tissue set (Qiagen,
156 Venlo, Netherlands) as previously described [12]. The DNA concentration and purity were determined using a
157 Nanodrop-100 spectrophotometer (NanoDrop, DE, USA), and the DNA integrity and absence of intact RNA was
158 checked by agarose electrophoresis. About 0.75 µg DNA was made up in 5 µl ultrapure molecular biology grade
159 water (Sigma Aldrich) [concentration, 0.15 µg/µl]. PCR, biotin labeling, hybridization to arrays, washing and
160 development of spots were performed as described in the AMR-ve protocol. Signal intensities were read within
161 10–15 min. of the final step (addition of buffer D1). Arrays were aligned in the reader as required by manual
162 setting of array reference marker spots as described in the ATR03 2.0 installation and user guide. Results were
163 exported, and the mean signal value for 3 replicate spots per probe was determined manually. Probes with
164 intensity value ≥ 0.4 were considered positive (P in tables) whilst those <0.3 were considered negative. Values
165 between 0.3 and <0.4 were considered ambiguous (A in tables).

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179 **Results**

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181 **Overview of the results of microarray testing and the most common resistance determinants in *K.***

182 *pneumoniae*

183 Fig. 1 provides a summary of the results from the microarray tests. Table 1 shows resistance phenotypes based
184 on disc diffusion assays. All strains were positive for one or more genetic marker in the microarray. Among the
185 30 clinical isolates tested, hybridization with probes associated with resistance to the following antibiotics was
186 obtained for one or more strains: β -lactams, sulphonamides, trimethoprim, tetracyclines, aminoglycosides,
187 quinolones and chloramphenicol. In addition, hybridization with a probe indicating the presence of a class 1
188 integrase was obtained with 18 clinical isolates (60%) and both control strains. Thus the assay was able to detect
189 simultaneously resistance markers to a wide-range of antibiotic groups of clinical importance for the treatment of
190 infections caused by *K. pneumoniae*, as well as a genetic element associated with resistance-gene acquisition.

191 **The most commonly detected resistance determinants**

192 By virtue of a relatively large number of probes representing multiple antibiotic classes, the microarray was able
193 to provide an overview of which resistance gene classes were common among the clinical isolates and control
194 strains. The 5 most commonly detected resistance markers in the 30 clinical isolates were consensus-*bla_{SHV}* (β -
195 lactams; 87%), *tetA* (tetracycline; 67%), *sulI* (sulphonamides; 60%), consensus-*bla_{TEM}* (β -lactams; 53%) and
196 *bla_{CTX-M-I}*-like (β -lactams; 47%).

197 **Resistance to β -lactams**

198 All strains were resistant to ampicillin, 90% of the clinical isolates were intermediate susceptible or resistant to
199 cefuroxime (2nd generation cephalosporin), and 90% were intermediate susceptible or resistant with respect to
200 cefotaxime (3rd generation cephalosporin). All strains gave hybridization with one or more probes representing
201 classes of resistance determinants to β -lactams. Four clinical isolates and the control strain ATCC 700603 did
202 not produce hybridization with the consensus-*bla_{SHV-I}*-probe. However, these gave hybridization with other
203 determinants of resistance to β -lactams, variously consensus-*bla_{TEM}* and/or one or more of the *bla_{CTX}* and *bla_{OXA}*-
204 probes. Hybridization to the *bla_{CTX-M-I}*-like probe was common (Fig. 1), supporting the idea that the class A
205 CTX-M β -lactamases are widespread among clinical isolates of the species. All strains possessing a *bla_{CTX-M}*
206 determinant were resistant to cefuroxime and cefotaxime.

207 **Testing for β -lactamase production**

208 Detection of ESBLs:

209 ESBLs were detected in isolates 5-9, 12-20, 22-26, and 30, but not in 21, 27-29. These results correlate well or
210 are not excluded by the disc-diffusion and microarray-based tests (Table 1; Fig. 1).

211 KPC/MBL-kit (Rosco) and resistance to meropenems:

212 Based on the finding of resistance to meropenem, isolates 1-4 (Table 1), were chosen as relevant for screening
213 for KPC (or other class A enzyme) and MBL (Ambler class B) production. Strains 1, 2 and 3 were shown to be
214 MBL producers, whereas strain 4 did not possess an MBL. Strain 4 tested positive for KPC (or other Ambler
215 class A enzyme). The finding of resistance/intermediate susceptibility to meropenem in these strains
216 is thus supported by the confirmatory tests.

217 ESBL+AmpC-screen test (Rosco):

218 Isolates 10 and 11 hybridized with respectively probes detecting DHA and FOX/CMY resistance determinants
219 (Fig. 1). These were the only strains where a plasmid-mediated class C β -lactamase was indicated by the array.
220 In a previous array-based study of 29 *K. pneumoniae* clinical isolates, hybridization to the DHA- and FOX-
221 probes was not found, and only 3 instances of hybridization to the CMY-probe was obtained. However, these
222 genes were also only sporadically found in other species of *Enterobacteriaceae* [9]. In the present work, testing
223 of strains 10 and 11 using the ESBL+AmpC screen, confirmed AmpC activity in isolate 10. The results were
224 ambiguous with strain 11 perhaps owing to the presence of multiple β -lactamases (Fig. 1).

225 **Resistance to tetracycline (*tet*)**

226 Disc diffusion tests were not performed for tetracycline resistance as EUCAST zone diameter breakpoints are
227 only validated for *Escherichia coli*. The most common determinant of resistance to tetracyclines detected was
228 *tetA* (detected in 67% of clinical isolates). In addition *tetD* was detected in 3 isolates, whereas *tetB* was only
229 detected in the control strain ATCC 33495.

230 **Resistance to sulfonamides (*sul*) and trimethoprim (*dfr*)**

231 Eighty-seven percent of the clinical isolates were resistant/intermediate susceptible to SXT. Hybridization to
232 *sul1* was obtained for 60% of the clinical isolates and both control strains, whereas the corresponding number for
233 *sul2* and *sul3* was 40% and 3%, respectively (Fig. 1). The four SXT susceptible strains lacked both *sul* and *dfr*
234 ($n=3$) or only *dfr* ($n=1$). Consequently, the results of the phenotypic susceptibility testing are in accordance with
235 the microarray findings. However, resistance to SXT was found in 5 strains producing hybridization with *sul* or
236 *dfr* (but not both) and in one which did not hybridize to *sul* or *dfr* (Fig. 1; Table 1).

237 **Resistance to the aminoglycoside gentamicin**

238 Twelve of the 30 clinical isolates (40%) were resistant to gentamicin. The array has 7 probes dedicated to the
239 detection of determinants of aminoglycoside resistance. Of these, Prob_aac3Ia_1, Prob_aac3IVa_1 (3-N-
240 aminoglycoside acetyltransferases) and probe_ant2Ia_1 (aminoglycoside 2''-adenyltransferase) are generally
241 associated with gentamicin resistance [19]. However, none of the clinical isolates produced hybridization with
242 the two first-named probes. Hybridization with probe_ant2Ia_1 was obtained only with the control strain ATCC
243 700603 which also showed intermediate susceptibility to gentamicin (Table 1). Of the 12 resistant clinical
244 isolates, 10 (83%) produced hybridization with probe Prob_aac6Ib_1 which is associated with an
245 aminoglycoside 6'-N-acetyltransferase. In addition, 3 gentamicin-susceptible strains gave hybridization to this
246 probe.

247 **Resistance to the fluoroquinolone ciprofloxacin.**

248 Sixty-three percent of the clinical isolates showed resistance or intermediate susceptibility to ciprofloxacin
249 (Table 1). Of these 19 strains, only 68% hybridized with one or more of the *qnr*-probes associated with low-level
250 resistance to fluoroquinolones.

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268 **Discussion**

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270 This study describes the first systematic and extensive application of the commercially available AMR-ve
271 microarray to the determination of resistance determinants and integrases in clinical *K. pneumoniae* isolates.
272 AMR-ve has been used to characterize one [27] and two [31] *K. pneumoniae* in other studies. Data from testing
273 of a non-overlapping set of 29 clinical isolates and 5 control strains of *K. pneumoniae* using an extended version
274 of the array containing many of the same probes has been reported [9] and comparisons are discussed below. The
275 array was able to detect quickly and generally unambiguously, resistance determinants to all the major groups of
276 clinically important antibiotics in the treatment of *K. pneumoniae*. Of the total of 265 individual probe
277 hybridizations recorded for the 32 strains, only 7 reactions (< 3%) were recorded as ambiguous (given as yellow
278 cells in Fig. 1).

279 The array requires 100 – 400 ng/μl DNA (total volume 5 μl) from a single clone for linear amplification
280 prior to hybridization. This is, of course, less sensitive than a classical PCR approach where typically a few ng of
281 template DNA is required. However, array-based approaches such as AMR-ve have the advantage of being able
282 to detect many more genes (> 50 in the present study) than for example a standard multiplex PCR-analysis.
283 Furthermore, as high as a 98.8% correlation between the array and PCR-data has been reported [5].

284 Salient details concerning the origins of the array probes and the resistance determinants they may
285 reveal are given in Table 1 in the original publication describing the development of the array [5]. Amr-ve was
286 originally developed for and validated against *Escherichia coli* and *Salmonella spp.* One indication of the
287 applicability of AMR-ve to typing of *K. pneumoniae* comes from the observation in the present and previous
288 work [9], that most of the array probes gave hybridization with one or more *K pneumoniae*. However, the
289 following 13 probes did not hybridize with any strain in either study: *tetE*, *tetG* (associated with tetracycline
290 resistance); *aadA4*, *aac(3)-Ia*, *aac(3)-IV* (associated with aminoglycoside resistance); *dfrA7*, *dfrA17* (associated
291 with resistance to trimethoprim), *bla_{MOX}*, *bla_{ACC-1}*, *bla_{OXA-7}*, *bla_{CTX-M2}*, *bla_{CTX-M8}*, *bla_{CTX-M-26}* (associated with
292 resistance to β-lactams). This and other concordances between the present and previous work on non-overlapping
293 sets of strains, probably reveals important information on the typical genetic makeup of drug-resistant *K.*
294 *pneumoniae*. However, in cases of non-hybridization probe suitability might also be an issue. To assess this we
295 compared probe sequences [5] to those in the GenBank® database using the BLAST® search engine [2]. The
296 search was restricted to generally curated *K. pneumoniae* DNA and the resistance gene in question. Of the 13
297 array-probes which did hybridize to any strain, we found in the database several instances of 100% sequence

298 identity over the whole probe length for all but *tetE* (probe length 24 bp; sequence identity only over 15 bp). The
299 *tetC*-probe which did not hybridise with any of the strains investigated in the present study gave similar results to
300 *tetE* (probe length 24 bp; sequence identity only over 14 bp). However, in the study of Card et al., [9], 17% of
301 clinical isolates produced hybridization to this probe. More data is desirable in order to evaluate the limitations
302 of these probes in the investigation of resistance to tetracycline in *K. pneumoniae*.

303 In addition to the detection of a wide-range of resistance determinants to antibiotic classes with
304 relevance to the treatment of infections caused by *K. pneumoniae*, the array also detected class 1 integrases in
305 60% of the clinical isolates and both control strains. This is similar to the value of 83% for other clinical isolates
306 of the species [9] and 74% reported for multidrug resistant *K. pneumoniae* and *Escherichia coli* in Iran [28].
307 Furthermore, in a study of 136 clinical isolates of *K. pneumoniae* in India, 52 of 63 ESBL-producing isolates
308 (83%) contained a class 1 integrase [6]. The finding of a significant distribution of class 1 integrases in the
309 present and other studies, supports the notion that a potential for resistance gene acquisition is common in
310 clinical isolates of this species. Some of the studies referenced above [6, 28] used multiplex-PCR-based
311 procedures for the detection of integrases. The present study shows that similar screening can be done effectively
312 with the AMR-ve approach.

313 The most commonly indicated resistances were *bla_{SHV}* (87% clinical isolates), *tetA* and *sulI*. The *bla_{SHV}*
314 gene is chromosomally located in *K. pneumoniae* [13] and it has been suggested that SHV enzymes approach
315 ubiquity in *K. pneumoniae* [4]. The finding that the majority of clinical isolates produced hybridization with
316 *bla_{SHV}* is thus in keeping with previous work. However, the *bla_{SHV}*-probe included on the array is a consensus
317 sequence for *bla_{SHV}*-genes and this limitation of scope is discussed further below. It has been reported that *tetA*
318 encoded efflux pumps are widespread among *Enterobacteriaceae* [32], but there is limited data for *K.*
319 *pneumoniae*. In an analysis of 30 *K. pneumoniae* causing urinary tract infections in Iran [7], all of the isolates
320 tested positive for *tetA* and *tetB*. This is in line with the present array results. However, in contrast to the Iranian
321 strains, the present study strongly suggests that *tetA* (67% Norwegian clinical isolates) is more common than *tetB*
322 (0%) in Norway. In their study of 29 isolates, Card et al., [9] found that only 21% of *K. pneumoniae* hybridized
323 with *tetA*, again suggesting significant clonal differences with respect to this gene. Alternatively the low rate of
324 detection of *tetB* might be due to mutations/mismatches in the area of the probe. This could be investigated
325 further by sequencing studies.

326 β -lactam antibiotics are the most important agents in the therapy of *K. pneumoniae* infections, and the
327 array's standard set of probes is suitably dominated by those detecting determinants of resistance to these

328 antibiotics. The assay has only a single, consensus probe for *bla_{SHV}* genes. SHV-enzymes can, however, be both
329 narrow spectrum and extended-spectrum (ESBLs), depending on the subtype in question [17, 18]. Concordances
330 between hybridization with Prob_shv1_11 and measured resistances to cephalosporins and aztreonam illustrate
331 some of the limitations of the AMR-ve system: ten of the strains tested gave hybridization with only
332 Prob_shv1_11 from among the array's set of probes for β -lactam resistance genes (Table 2). If SHV-enzymes are
333 responsible for the β -lactam resistance phenotypes shown in Table 2, the array is unable to distinguish between
334 what is probably a classical narrow-spectrum enzyme in the control strain ATCC 33495, and the broad-spectrum
335 enzymes present in for example strains 8 and 14. Such a differentiation (subtyping) which is most useful to the
336 researcher can be accomplished by the PCR-sequencing approach, but is a limitation of the array in its present
337 form. In contrast to the single determinant for SHV-enzymes, the array contains several probes hybridizing to
338 *bla_{CTX-M}*-genes. For 14 clinical isolates, positive signals were obtained with Prob_ctxM1_11. This probe detects
339 among others the phylogenetic group CTX-M-15 which is currently the most widely disseminated CTX-M type
340 in the world [21]. Antibiotic resistances assayed by the disc diffusion assay were in general agreement with the
341 expected effects of the presence of *bla_{CTX-M}* genes (Table 1; Fig. 1). CTX-M-type ESBLs can exhibit activity
342 against monobactams and extended spectrum cephalosporins, including cefotaxime and ceftriaxone, but are
343 classically less effective against ceftazidime, cephamycins (e.g. ceftazidime) or carbapenems [3]. Thus, the finding
344 that the majority of clinical isolates (87%) were resistant or intermediately susceptible to ceftazidime (Table 1) is
345 somewhat surprising. However, the classical pattern of resistance conferred by CTX-M is not universal since
346 mutations can alter the specificity of CTX-enzymes [14]. Some CTX-M sub-types (e.g. CTX-M-15) have greater
347 catalytic efficiency against ceftazidime [8]. This sub-type is for example detected along with others by the array
348 ctxM1_11 probe. Given this, a more stringent probe, if practically feasible, directed at ceftazidime resistance
349 related CTX-M sub-types would be useful. Incidences of ceftazidime resistance/ intermediate susceptibility in 11
350 clinical isolates (Table 1) not showing hybridization with the ctxM1_11 probe, could potentially also be
351 explained by mutations in common plasmid-mediate SHV (or TEM) genes producing ESBLs [8]. Over-
352 production of SHV-enzymes (all 11 isolates produced hybridization with the *bla_{SHV}*-probe) and/or changes in
353 outer-membrane protein changes have also been associated with reduced susceptibility to ceftazidime in *K.*
354 *pneumoniae* [22, 23].

355 Carbapenems are in some instances the last therapeutic options for treatment of infections by *K.*
356 *pneumoniae*. It is, therefore, pertinent to look more closely at the results for the 4 strains (strains 1-4; Table 1)
357 which showed meropenem resistance/intermediate susceptibility. Although it includes probes detecting various

358 classes of Ambler A and C enzymes, the AMR-ve system is not equipped to detect determinants of class B
359 enzymes. Thus the array is not able to provide support for the finding of MBLs (detected phenotypically) in
360 strains 1, 2 and 3. Furthermore the array does not detect *bla*_{KPC} which would be useful given that KPC activity
361 was detected for strain 4.

362 Plasmid-mediated resistance to sulphonamides and trimethoprim is normally due to the acquisition of
363 dihydropteroate synthases for sulphonamides and dihydrofolate reductases for trimethoprim. Three resistance
364 genes, *sul1*, *sul2* and *sul3* encoding dihydropteroate synthases and more than 20 dihydrofolate reductase (*dfr*)
365 genes have been described [16]. Both groups of genes are associated with class 1 integrons residing on plasmids
366 and/or the chromosome [16]. The array shows 100% correlation between susceptibility to SXT and the absence
367 of hybridization to *dfr* and/or *sul* probes. However, 5 SXT-resistant strains did not produce hybridization with
368 either one or both of *sul* and *dfr* probes, possibly indicating the presence of other resistance
369 determinants/mechanisms not detected by the microarray. In summary, the microarray was able to explain
370 sensitivity but not always resistance to SXT.

371 The present study suggests that an aminoglycoside 6'-N-acetyltransferase may be associated with
372 gentamicin resistance in the current panel of strains: thirteen clinical isolates produced hybridization with this
373 probe (Fig. 1). Of these, 77% showed resistance to gentamicin (Table 1). Similarly, in a previous study of *K.*
374 *pneumoniae* isolates [9], it was found that 76% of strains producing hybridization to the probe were also resistant
375 to gentamicin. However, although AAC(6')-Ib is the most prevalent aminoglycoside-modifying enzyme known,
376 and is present in over 70% of AAC[6']-producing Gram-negative isolates [30], group I type 6'-N acetyl
377 transferases are not typically agents of resistance to gentamicin [19, 30]. In one study of carbapenem-resistant *K.*
378 *pneumoniae*, 98% of strains tested possessed *aac(6')-Ib* [1]. However, this gene was associated with resistance to
379 tobramycin and not to gentamicin. In addition, about 94% of the strains that carried *aac(6')-Ib* as the only
380 determinant of an aminoglycoside-modifying enzyme were susceptible to gentamicin [1]. The authors thus
381 proposed that *aac(6')-Ib* in association with other aminoglycoside resistance genes could explain the high
382 gentamicin MIC values found for some strains. Furthermore, they suggest that the widespread presence of
383 *aac(6')-Ib* in *K. pneumoniae* could be a consequence of it being on the same plasmids as those carrying ESBL
384 genes [1]. In the present study, the four clinical isolates which did not produce ESBLs (phenotypic tests) also
385 failed to give hybridization to Prob_aac6Ib_1, whereas every strain producing hybridization with Prob_aac6Ib_1
386 was also an ESBL producer. Thus the array test provides some support for the proposed presence of *aac(6')-Ib*
387 and ESBL-genes on the same plasmids [1]. Notwithstanding, it should be noted that some *aac(6')-Ib* subvariants

388 are known to be associated with gentamicin resistance. Among the recent variants of this enzyme with altered
389 specificity are AAC(6′)-Ib11, which confers simultaneous resistance to gentamicin and amikacin [11] and
390 AAC(6′)-Ib-cr, which has a unique extension of its substrate specificity from aminoglycosides to structurally
391 unrelated fluoroquinolones [24]. If feasible, it would be useful to include additional probes in order to ascertain
392 if the apparent correlation between gentamicin resistance and hybridization to Prob_aac6Ib_1 - probe aac (6′)-Ib
393 in the present study, indicates the presence of gene sub-types conferring gentamicin resistance

394 Of the 30 clinical isolates tested, 19 showed resistance or intermediate resistance to ciprofloxacin. Of
395 these, 32% showed resistance to ciprofloxacin without giving hybridization with probes associated with
396 resistance to quinolones. However, *qnr* genes are known to only give low-level resistance to quinolones.
397 Quinolone resistance is mainly caused by chromosomal mutations in *gyrA* and *parC*, and the contribution of a
398 *qnr*-gene can be additive to other resistance factors. The microarray is thus useful in its ability to detect *qnr* but
399 unsurprisingly it is not predicative of resistance to ciprofloxacin.

400 Susceptibility to chloramphenicol was not investigated in the present study. AMR-ve hybridizations
401 indicated that the most common determinants of resistance to this antibiotic were chloramphenicol
402 acetyltransferases (CatB3; CatA1). Genes connected with efflux proteins (FloR; CmlA) were also detected, but
403 in fewer strains (Fig.1). The same tendency was seen in a previous study of clinical isolates [9]. In the referenced
404 work, all strains producing hybridization with the *catA1*-probe and/or the *cmlA*-probe were also resistant to
405 chloramphenicol. Thus hybridization with these probes seems to be a good indicator of the resistance phenotype.

406 Summing up the results, the present study evaluates the usefulness of a commercially available
407 microarray, AMR-ve, for the detection of clinically important resistance determinants and integrases. The assay
408 was able to detect genetic determinants of integrases and a wide-range of antibiotic classes of relevance for
409 treatment of infections caused by *K. pneumoniae*. Furthermore the test was easy to perform and the results were
410 only rarely difficult to interpret ('ambiguous'). AMR-ve functions well in the genotyping of clinical isolates of
411 *K. pneumoniae* and provides a useful start point for further testing. However, the assay was shown to be too
412 restricted in scope to explain many important measured resistances to β -lactams (e.g. to carbapenems) and
413 aminoglycosides in *K. pneumoniae*. The producer of the AMR-ve has recently launched a new CarbDetect AS-1
414 Kit for multidrug-resistant Gram-negative bacteria which runs on the same platform. This test allows DNA-
415 based detection of amongst others the most important carbapenemase genes (e.g. *bla*_{OXA-23}, *bla*_{KPC}, *bla*_{NDM},
416 *bla*_{VIM}) as well as other prevalent β -lactamases (e.g. *bla*_{CTXM-1/15}, *bla*_{OXA-1}, *bla*_{OXA-2}). It also contains species-
417 specific probes for identification of *K. pneumoniae*. Thus this new system supplies some of the information

418 needed to bridge gaps between genotyping and resistance testing reported as weaknesses of the AMR-ve in the
419 present work. Additionally, whole genome sequencing studies are becoming an option for more laboratories and
420 are offered commercially. These can reveal allelic forms of genes and point mutations which are often
421 specifically associated with resistance to a particular antibiotic. The current price for the AMR-ve Genotyping
422 kit is about 1500 EUR for 50 tests (about 30 EUR per bacterium). A single analyst in our laboratory was able to
423 test about 5-10 strains per day with the AMR-ve. There are now many commercial agents offering whole
424 genome sequencing at about 300-400 EUR for a single strain with lower prices for bulk orders (including DNA
425 purification from supplied cells, libraries and bioinformatics) with a turnaround of 3-4 weeks or less. Thus the
426 total prices for the two approaches are near parity.

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Table 1 Antibiotic resistance profiles of thirty clinical isolates and two control strains of *Klebsiella pneumoniae*

| Antibiotic | Strain designation and resistance phenotype ^a | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | ATCC 700603 | ATCC 33495 | | | | |
|------------|--|---|---|---|---|---|---|---|---|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----------------|---------------|---|---|---|---|
| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | 16 | 17 | 18 | 19 | 20 | 21 | 22 | 23 | 24 | 25 | 26 | 27 | 28 | 29 | 30 | | | | | | |
| AMP | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R | R |
| AMC | R | R | R | R | R | | R | | R | R | R | R | | R | | | | | | | R | R | | R | | | | | | | | | R | | | |
| TZP | R | R | R | R | R | | I | | R | R | R | I | | R | | | | | | | R | | | I | | | | I | | | | R | | I | | |
| CTX | R | R | R | R | R | R | R | R | R | I | R | R | R | R | I | R | R | R | R | R | | R | R | R | | R | | | I | R | R | R | I | | | |
| FOX | R | R | R | R | | | | | | | R | | | | | | R | R | | | | | R | | | | | R | R | R | | | | R | | |
| CAZ | R | R | R | R | R | | R | R | R | R | R | R | | R | I | R | | R | R | R | I | I | R | R | I | I | | I | | | I | | I | R | | |
| CXM | R | R | R | R | R | R | R | R | R | R | R | R | R | | R | R | R | R | R | | R | R | R | R | R | R | R | | R | R | R | R | R | R | R | |
| MEM | R | R | I | R | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| ATM | R | R | R | R | R | I | R | R | R | I | R | R | I | R | | R | R | R | R | R | | R | | R | | | R | | | | R | R | R | | | |
| CIP | R | R | R | R | R | | I | | R | R | I | R | I | | I | I | | | R | I | | R | | I | | | R | R | | | | | | | | |
| CN | R | R | | | R | R | R | | R | R | R | R | R | | | | | | | | | | R | R | | | | | | | | | | | I | |
| STX | R | | R | R | R | R | R | | R | R | R | R | R | R | R | R | I | R | R | R | | R | R | R | R | R | R | R | | R | R | R | I | | | |

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^a (R) indicates clinical resistance and (I) indicates intermediate susceptibility. A blank entry in the table indicates full susceptibility (S) to a particular antibiotic,

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555 **Table 2** Overview of resistance phenotypes of strains producing hybridization with either the SHV-1 or the
 556 TEM-1 probe and no other determinant of resistance to β -lactams (derived from Table 1)

| Strain | Probe hybridizations | Penicillins | Cephalosporins/ Cephamycins | Monobactams (Aztreonam) | Carbapenems (Meropenem) |
|----------------|----------------------|-------------------|--------------------------------|----------------------------|----------------------------|
| 3 ^a | Prob_shv1_11 | AMP, AM/C TZP | FOX, CXM CTX, CAZ, | ATM | MEM (I) |
| 8 | Prob_shv1_11 | AMP | CXM, CTX,CAZ | ATM | - |
| 14 | Prob_shv1_11 | AMP, AM/C, TZP | CTX, CAZ, CXM | ATM | - |
| 15 | Prob_shv1_11 | AMP | CTX(I), CAZ(I) | - | - |
| 23 | Prob_shv1_11 | AMP | FOX, CXM CTX, CAZ, | - | - |
| 25 | Prob_shv1_11 | AMP | CXM, CAZ(I) | - | - |
| 26 | Prob_shv1_11 | AMP | CTX, CAZ(I), CXM | ATM | - |
| 27 | Prob_shv1_11 | AMP, TZP(I) | FOX | - | - |
| 28 | Prob_shv1_11 | AMP | FOX, CTX(I), CAZ(I) CXM, | . | . |
| ATCC 33495 | Prob_shv1_11 | AMP | - | - | - |
| 6 | Prob_tem1_1 | AMP | CTX CXM | ATM(I) | - |

557 ^a MBL detected by phenotypic tests

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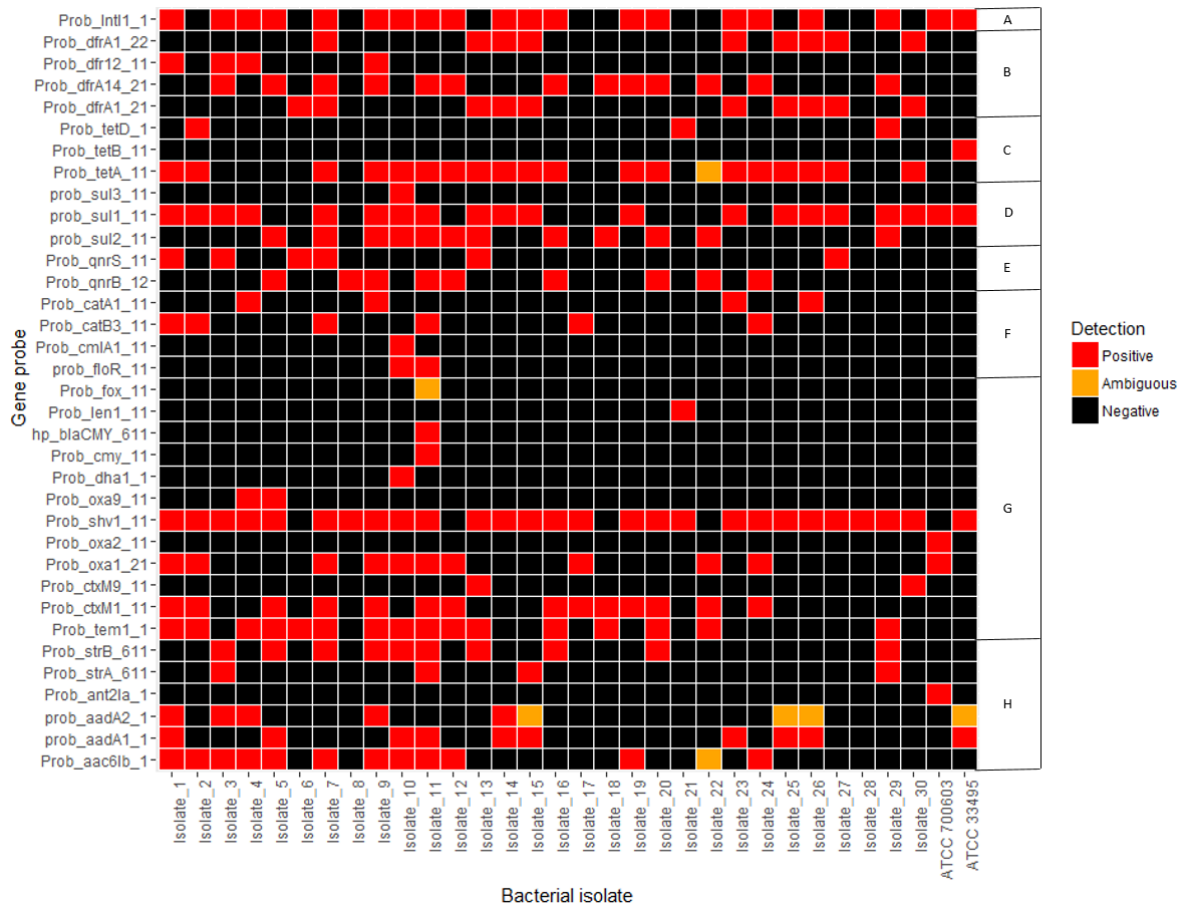
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Fig 1. Heat map showing the pattern of probe hybridizations for 32 strains of *Klebsiella pneumoniae*.

Only probes giving hybridization to one or more strain are shown.

Key: A – probe associated with class 1 integrases. B – trimethoprim. C – tetracyclines. D – sulphonamides. E – quinolones/fluoroquinolones. F – chloramphenicol. G – β -lactams. H - aminoglycosides