Environmental dissemination of carbapenemase-producing enterobacteriaceae in rivers in Switzerland

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22 Abstract

23	The aquatic environment takes on a key role in the dissemination of antimicrobial-resistant
24	Enterobacteriaceae. This study assesses the occurrence of carbapenemase-producing
25	Enterobacteriaceae (CPE) in freshwater samples from rivers, inland canals, and streams
26	throughout Switzerland, and characterizes the isolated strains using phenotypic and NGS-
27	based genotypic methods. CPE producing KPC-2 (n=2), KPC-3 (n=1), NDM-5 (n=3), OXA-
28	48 (n=3), OXA-181 (n=6), and VIM-1 (n=2) were detected in 17/164 of the water samples.
29	Seven Escherichia coli had sequence types (STs) that belonged to extra-intestinal pathogenic
30	clonal lineages ST38, ST73, ST167, ST410, and ST648. The majority (16/17) of the
31	carbapenemase genes were located on plasmids, including the widespread IncC (n=1),
32	IncFIIA (n=1), and IncFIIB plasmids (n=4), the epidemic IncL (n=1) and IncX3 (n=5)
33	plasmids, a rare Col156 plasmid (n=1), and the mosaic IncFIB, IncR, and IncQ plasmids
34	(n=3). Plasmids were composed of elements that were identical to those of resistance
35	plasmids retrieved from clinical and veterinary isolates locally and worldwide. Our data show
36	environmental dissemination of high-risk CPE clones in Switzerland. Epidemic and mosaic-
37	like plasmids carrying clinically relevant carbapenemase genes are replicating and evolving
38	pollutants of river ecosystems, representing a threat to public health and environmental
39	integrity.
40	
41	Main finding

42 Carbapenem-resistant Enterobacteriaceae and their genetic mechanisms of resistance are
43 replicating and evolving pollutants of river ecosystems, representing a threat to public health.
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46 Keywords: carbapenems; antibiotic resistance; plasmids; aquatic environment; pollution
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48 Introduction

49 Beta-lactam antibiotics including penicillins, cephalosporins, monobactams, and carbapenems are the most frequently consumed antibiotics worldwide (WHO, 2018). 50 51 Carbapenem antibiotics, for example ertapenem, imipenem, and meropenem are classified by the World Health Organization WHO as critically important for human health and are 52 53 currently considered last resort antimicrobials to treat severe infections by multidrug resistant (MDR) Gram-negative nosocomial pathogens (WHO, 2017, van Duin and Doi, 2017). 54 55 Carbapenem resistance therefore represents a significant public health concern of global 56 dimensions. One of the most significant mechanisms of carbapenem resistance among 57 Enterobacteriaceae involves the synthesis of carbapenemases, enzymes that inactivate 58 carbapenems and other β -lactam antibiotics (Queenan and Bush, 2007). Since the first 59 isolation of carbapenemase producing Enterobacteriaceae (CPE) harboring the bla_{KPC} gene in 1996, (Yigit et al., 2001) clinical CPE carrying chromosomal or plasmid-mediated 60 carbapenemase-genes such as *bla*_{NDM}, *bla*_{OXA-48}, *bla*_{VIM} and *bla*_{IMP} have been found 61 62 worldwide (Kopotsa et al., 2019, Nordmann et al., 2011). In recent years, clinically relevant CPE have been detected in non-human sources including companion and food-producing 63 animals, the food chain, wildlife, and the environment, giving rise to health and ecological 64 65 issues at the human-animal-environmental interface (Mills and Lee, 2019). Addressing these 66 issues necessitates a holistic and multidisciplinary approach known as the One Health 67 Concept (Hernando-Amado et al., 2019). Of the One Health antibiotic resistance triad, the environment is the most dynamic, but also the least understood sector (Essack, 2018). Within 68 this sector, the aquatic environment is of particular importance because it represents a most 69 70 basic resource, and the role that it plays in the spread of antimicrobial resistance (AMR) is critical, with the genetic context of the AMR genes remaining largely unexplored (Kraemer 71

et al., 2019, Mills and Lee, 2019, Furness et al., 2017, Williams et al., 2016, Alonso et al.,
2001).

CPE producing clinically relevant carbapenemases including KPC, NDM, IMP, OXA-48-74 75 like, and VIM have been reported in European rivers since 2010 (Poirel et al., 2012), mostly rivers associated with effluent such as hospital or urban wastewater (Falgenhauer et al., 2019, 76 77 Jelić et al., 2019, Lepuschitz et al., 2019, Khan et al., 2018, Mahon et al., 2017, Zurfluh et al., 2017). Furthermore, CPE in wastewater and in surface water may include intestinal 78 pathogenic *E. coli* and extra-intestinal *E. coli* (ExPEC), which give rise to diseases in humans 79 and animals by virtue of specific virulence factors (Mahon et al., 2017; Zurfluh et al., 2017; 80 81 Kaper et al., 2004). Virulence traits include adhesins, capsular antigens, siderophores, and 82 toxins that enable pathogenic E. coli to avoid host defense systems, colonize host surfaces 83 and invade host tissues (Kaper et al., 2004). While anthropogenic influences are well recognized as major contributors of CPE to waterways, the possible pathways of transmission 84 of CPE between humans, animals including wildlife, and the freshwater ecosystem are not 85 86 well documented and potential human and animal health impacts caused by exposure to 87 environmental CPE remain unclear (ECDC, 2019, Mills and Lee, 2019). The aquatic environment provides ideal settings for carbapenemase harboring mobile genetic 88 89 elements (MGEs) including plasmids, insertion sequences, and transposons, to be retained and to disseminate via horizontal gene transfer (Gillings et al., 2018 Marti et al., 2014, 90 91 Pruden, 2014). Such MGEs contribute to what is becoming increasingly recognized as 92 xenogenetic pollution of the aquatic ecosystem, with potentially adverse impact on human 93 welfare and environmental integrity (Gillings et al., 2018). This study was designed to evaluate the occurrence of CPE in different water bodies 94 95 throughout Switzerland and to characterize the isolated strains using phenotypic and genotypic methods, including whole genome analyses. We also aimed to identify any genetic 96

97 relatedness of CPE present in the aquatic environment to CPE associated with documented
98 human and animal infections in order to assess their relevance to public and environmental
99 health. Particular emphasis was placed on identifying antimicrobial resistance genes (ARGs)
100 and MGEs.

101

102 Material and Methods

103

Sampling. Between May and August 2019, a total of 164 surface water samples were taken 104 from different water bodies including rivers (n=113), streams (n=42) and inland canals (n=9)105 106 located between 300 and 3000 m above sea level (Table S1). Water was collected from each 107 site in sterile 500 mL containers and transferred to the laboratory in a cool box. Microbiological analysis. CPE were isolated from the water samples using selective media 108 as previously described (Zurfluh et al., 2013). For more details see Supplementary Material. 109 Isolates were subjected to antimicrobial susceptibility testing (AST) according to the 110 guidelines of the Clinical and Laboratory Standards Institute (CLSI) (CLSI., 2018), as 111 detailed in Supplementary Material. For each isolate the minimal inhibitory concentration 112 (MIC) of the carbapenem antibiotics ertapenem, imipenem, and meropenem were 113 114 determined, and each isolate was tested against a panel of further 16 antimicrobials using the disk diffusion method as described in Supplementary Material. 115 116 Isolates displaying resistance to three or more classes of antimicrobials (counting β -lactams 117 as one class) were defined as multidrug-resistant (MDR) (Magiorakos et al., 2012). Whole genome sequencing (WGS) and analysis of genomic content. Prior to WGS, each 118 isolate was tested for carbapenemase production using the β -CARBATM colorimetric test 119 (Bio-Rad, Cressier, Switzerland). Isolates were screened by PCR for the presence of bla_{KPC} , 120

121 *bla*_{NDM}, *bla*_{OXA-48-like}, or *bla*_{VIM} genes, as described previously (Poirel et al., 2011, Ellington

the et al., 2007), and as outlined in Supplementary Material.

- 123 Genomic sequences were obtained using both Illumina MiniSeq (Illumina, San Diego, CA,
- 124 USA) and MinION sequencer on a R9.4 Spot On flow cell (Oxford Nanopore Technologies,
- 125 Oxford, United Kingdom). For specifics see Supplementary Material.
- 126 Reads were assembled as described previously (Stevens et al., 2019), and as further described
- 127 in Supplementary Material. In silico analyses were carried out as detailed in Supplementary
- 128 Material, to determine *Escherichia coli* core genome multilocus sequence types (cgMLST)
- and serotypes, as described by (Wirth et al., 2006) and by (Joensen et al., 2015), respectively.
- 130 Core genome alignments were performed as described earlier (Treangen et al., 2014) to
- 131 detect related strains available in public genome databases. All genomes were further
- 132 screened in silico in order to identify virulence markers, antimicrobial resistance genes and
- 133 plasmids as described previously (Xie et al., 2018, Jia et al., 2017, Carattoli et al., 2014,
- 134 McArthur et al., 2013), using databases detailed in Supplementary Material.
- 135 Plasmid sequences were compared to reference sequences using the bacterial plasmid
- 136 database PLSDB, available at <u>https://ccb-microbe.cs.uni-saarland.de/plsdb/</u> (Galata et al.,
- 137 2019), as outlined in Supplementary Material.
- 138 Geographical map. Geospatial visualization was performed by plotting GPS coordinates of
- the sampling sites onto a geographical map using the open source geographic information
- 140 system (GIS) software QGIS (<u>https://qgis.org</u>).
- 141 Accession numbers. Genome assemblies and sequence reads are deposited at Sequence Read
- 142 Archive (SRA) and GenBank hosted by the NCBI database under the BioProject ID
- 143 PRJNA604100.
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146 Res	ults
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147	Occurrence of CPE in Swiss water bodies. CPE were detected in 17 (10%) of the 164 water
148	samples, including 13 (12%) of the 113 river water samples, three (33%) of the 9 samples
149	taken from inland canals, and one (2%) of the water samples from a stream (Figure 1). CPE
150	were not detected in water bodies more than 1000 m above sea level (Figure S1/Table S1).
151	Overall, the 17 CPE included 12 E. coli, one Citrobacter freundii, one Enterobacter kobei,
152	one Klebsiella aerogenes, one Klebsiella variicola and one Raoultella ornithinolytica strain,
153	respectively (Figure 2).
154	The results of the initial PCR screening, combined with the results of WGS, identified the
155	carbapenemase genes bla_{KPC-2} , bla_{KPC-3} , bla_{OXA-48} , bla_{OXA-48} , $bla_{OXA-481}$, and bla_{VIM-1} (Figure 2).
156	
157	Antimicrobial resistance phenotypes and genotypes of the CPE. Application of the β -
158	CARBA TM test indicated the presence of carbapenem-hydrolyzing enzymes in all 17 isolates.
159	Phenotypic AST revealed that eight strains exhibited MIC values above the carbapenem
160	susceptibility breakpoints, with ertapenem non-susceptibility defined as an MIC of ≥ 1 mg/L,
161	and meropenem and imipenem non-susceptibility defined as an MIC of ≥ 2 mg/L (CLSI.,
162	2018) (Figure 2/Table S1). Resistance to other β -lactam antibiotics was common, with 14
163	(82%) and 10 (59%) of the strains exhibiting resistance to the 3^{rd} and 4^{th} generation
164	cephalosporins cefotaxime and cefepime, respectively. Moreover, 15 (88%) of the strains
165	were MDR (Figure 2).
166	
167	Phylogenetic analysis and virulence gene profiles of the carbapenemase producing E.
168	coli. To determine a possible clinical relevance of CPE cultured from the aquatic
169	environment, the 12 E. coli strains were subjected to detailed analysis. cgMLST classified the
170	strains according to nine different E. coli STs. ST167, ST410 and ST940 accounted for two

strains each (2/12), respectively, while ST38, ST73, ST205, ST648, ST656, and ST1284

172 occurred in one strain each (Figure 2).

173 *E. coli* ST38 (strain CF065) is an international AMR high risk clone responsible for the

spread of OXA-48 producing *E. coli* (Pitout et al., 2019). Notably, in contrast to the other

strains described in this study, CF065 lacked plasmid elements but contained a

176 chromosomally located bla_{OXA-48} in a genetic environment consisting of a Δ Tn1999.2-like

177 structure as described in *E. coli* ST38 clones from the UK (data not shown) (Turton et al.,

178 2016, Pitout et al., 2019).

179 E. coli ST410 strains (strain CF038 and CF124, respectively) were assigned in silico to

180 serotype O8:H9, suggesting that these strains belong to clade C which is an *E. coli* ST410

181 clade associated with humans, companion animals, and farm environments (Falgenhauer et

al., 2016). Comparison showed that NDM-5 producing CF038 was very closely related with

183 only 51 chromosomal SNPs to NDM-5 producing *E. coli* ST410 (strain ECS9) isolated from

a patient with bloodstream infection in China in 2017 (Huang et al., 2019) (GenBank

accession no. VBQE00000000) (Figure S2).

186 Moreover, there was genetic similarity (<100 different alleles in cgMLST) of NDM-5

producing *E. coli* ST410 (strain CF038) and ST167 (strains CF163 and CF164, respectively),

and of OXA-181 producing *E. coli* ST1294 (strain CF032), to isolates associated with

documented cases of human infection in Canada and India (Mataseje et al., 2018) (BioProject

190 ID PRJNA390933) (Figure S3). Further, cgMLST comparison showed that the two OXA-181

191 producing *E. coli* ST940 (strains CF061 and CF064, respectively), although retrieved from

192 geographically distinct sites, were clonal with identical cgMLST patterns (Figure S2 and

193 Figure S3). By contrast, strains CF163 and CF164 were not genetically related to an NDM-5

194 producing *E. coli* ST167 clone that infected dogs and colonized veterinary employees at a

195 Swiss veterinary clinic in 2018 (data not shown) (Endimiani et al., 2020). Further, OXA-181

producing E. coli ST410 (strain CF124) had no phylogenetic link to a clone that caused an 196 outbreak involving companion animals at one Swiss veterinary hospital in 2018 (data not 197 shown) (Nigg et al., 2019). An overview of the phylogenetic relatedness of E. coli strains 198 199 from this study is shown in Figure S2. At least one virulence gene associated with pathogenicity in intestinal and extra-intestinal E. 200 201 coli (ExPEC) diseases was detected in 11 of the 12 carbapenemase-producing E. coli (Table 1 and Figure 2). Seventeen different virulence factors were identified, whereby the most 202 frequent ones were gad (glutamate decarboxylase gene involved in gastric acid resistance), 203 *lpfA*, (long polar fimbriae gene associated with the colonization of the intestine), and *capU*, 204 205 (hexosyltransferase homolog gene associated with adhesion), which were identified in seven, 206 six, and five isolates, respectively (Table 1).

207

Plasmid analysis. To investigate the host range, epidemiology, and possible relatedness, the 208 carbapenemase-encoding plasmids were fully sequenced and compared to already published 209 clinically relevant plasmids. Overall, 16 plasmids ranging in size from 7.8 kb to 244 kb were 210 211 analyzed (Table 2). With the exception of two plasmids carrying bla_{OXA-48} genes, all plasmids harbored at least one additional ARG (Table 2). Seven plasmids contained genes for type II 212 213 toxin/antitoxins (T/As), which are genetic systems that play a role in plasmid maintenance and the dissemination of multidrug resistance in Gram-negative bacteria (Yang and Walsh, 214 215 2017) (Table 2). For further analysis, plasmids were categorized as KPC, NDM, OXA, or 216 VIM-encoding plasmids, respectively (Table 2). **KPC-encoding plasmids.** The three *bla*_{KPC} genes identified in *C. freundii*, *E. kobei*, and *K.* 217 variicola were located on plasmids p062_B-KPC-2 determined to be IncQ1, p070_A-KPC-2 218

which was typed $IncFIB_K$, and $p118_A-KPC-3$ which belonged to IncFIIB, respectively

220 (Table 2). On all three plasmids, the bla_{KPC} genes were located within the Tn3-like

transposon Tn4401*a*, which is the most common isoform of Tn4401, a genetic structure that typically surrounds bla_{KPC} genes (Naas et al., 2012).

A sequence analysis of p062_B-KPC-2 presented a hybrid structure consisting of a 170 kb

backbone which had a high degree of similarity with plasmid p1643_10 (GenBank accession

- no. KF056330) from the epidemic *Salmonella enterica* serovar Kentucky ST198 strain
- 226 1643/2010 isolated from a turkey in Poland in 2010 (Wasyl et al., 2015). Plasmid p062_B-
- **227** KPC-2 further contained a 20 kb region carrying bla_{KPC-2} identical to plasmid pKP1504-kpc

228 (GenBank accession no. KF874496), which was purified from *K. pneumoniae* ST258 strain

229 GR-1504 during the early phases of a hospital epidemic in Greece in 2008 (Papagiannitsis et

al., 2016a, Giakkoupi et al., 2009) (Figure 3). The same bla_{KPC-2} carrying structure was

identified in p079_A-KPC-2, however, in p070_A-KPC-2, the region identical to plasmid

232 pKP1504-kpc covered an ~35 kb region (data not shown). Plasmid p070_A-KPC-2 showed

233 no further sequence homology to plasmids available in PLSDB.

234 Plasmid p118_A -KPC-3 was a mosaic plasmid that shared a common region (99% identity

over a length of 120834 bp) with an unnamed 244 kb IncFIB_K plasmid from K. pneumoniae

236 ST323 strain KSB1_4E isolated from a rectal swab of a hospitalized patient in Australia in

237 2013 (Gorrie et al., 2018) (GenBank accession no. CP024500.1). Plasmid p118_A -KPC-3

further shared a 12 kb region carrying the bla_{KPC-3} gene which was identical to an unnamed

239 plasmid from *K. pneumoniae* strain AR438 registered in the culture collection of the Food

and Drug Administration/ Centers for Disease Control and Prevention (FDA/CDC)

241 Antimicrobial Resistant Isolate Bank, Atlanta, USA (GenBank accession no.

242 NZ_CP029102.1) (Figure 3).

243 NDM-encoding plasmids. The three *bla*_{NDM-5} genes from *E. coli* were located on 87 kb

IncFIA, on a 132 kb IncFIB, and on a 10 kb pKPC-CAV1193-like plasmid, which was

nontypeable by incompatibility group (Sheppard et al., 2016, Mathers et al., 2015).

246	Plasmid p038_A-NDM-5 shared 99.9% identity with pAMA1167-NDM-5, a multidrug
247	resistance plasmid from a human clinical E. coli ST410 isolate from Denmark (Overballe-
248	Petersen et al., 2018) (GenBank accession no. CP024805.1) (Figure 4). Furthermore,
249	p164_A-NDM-5 and p163_C-NDM-5, both identified in <i>E.coli</i> ST167 in this study, were
250	determined to be highly similar at the nucleotide level (99-100%), to plasmids pM309-NDM5
251	(Figure 4), and pM217_FII (data not shown), respectively. Both plasmids were detected in
252	nosocomial E. coli ST167 strains from a hematology ward in Myanmar during 2015-2016
253	(Sugawara et al., 2019) (GenBank accession nos. AP018833.1 and AP018147.1,
254	respectively). By contrast, the three NDM-5 plasmids from this study were not similar to
255	previously reported NDM-5 plasmids from dogs and veterinary employees of a Swiss
256	veterinary hospital (data not shown) (Endimiani et al., 2020, Peterhans et al., 2018).
257	OXA-48-encoding plasmids . Of the two plasmid-mediated bla_{OXA-48} genes detected in <i>R</i> .
258	ornithinolytica and E. coli ST205, the former was identified on a 63 kb IncL plasmid
259	(p023_D-OXA-48) that shared >99% identity with an IncL plasmid p704SK10_2 identified
260	in an <i>E. cloacae</i> isolated from wastewater in 2015 in Switzerland (Marti et al., 2017)
261	(GenBank accession no. CP022150). Plasmid p023_D-OXA-48 was also highly identical to
262	pEC745 identified in E. coli ST131 from Morocco (Stoesser et al., 2016) (GenBank
263	accession no. CP015075.1), and to plasmid pOXA-48_4963 which was associated with a
264	nosocomial outbreak of K. pneumoniae in 2015 in the Czech Republic (Skalova et al., 2017)
265	(GenBank accession no. KX523900) (Figure 5). As is typical for IncL plasmids harboring
266	bla_{OXA-48} , the bla_{OXA-48} gene in p023_D-OXA-48 was located within the composite
267	transposon Tn1999.2 which is a Tn1999 variant with an IS1R insertion upstream of bla_{OXA-48}
268	(Pitout et al., 2019).
269	In the second plasmid, the <i>bla</i> _{OXA-48} gene was located on a 7.8kb Col156 plasmid (p053_E-

OXA-48), that shared 99.8% nucleotide identity with pMTY17816_OXA48 identified in a

271 human K. pneumoniae isolate from a patient from Vietnam in 2017 (Honda et al., 2019) 272 (GenBank accession no. AP019554.1) (Figure 5). The *bla*_{OXA-48} gene was flanked by two 273 copies of inverted insertion sequence IS1R, corresponding to the transposon variant 274 Tn1999.3, which was described for the first time in an pOXA-48-like IncL plasmid in a clinical E. coli strain from Italy (Giani et al., 2012a). In p053_E-OXA-48 however, the two 275 copies of IS1999 and the lysR gene which are present in Tn1999.3, were missing (Figure S4). 276 **OXA-181-encoding plasmids.** The most prevalent carbapenemase gene was $bla_{OXA-181}$ which 277 was located in five of six instances on 51 kb IncX3 plasmids (Table 2). All five were >99.9% 278 279 identical to IncX3 plasmids from a human K. pneumoniae isolate from the Czech Republic 280 (pOXA-181_29144) (Skalova et al., 2017), canine and human E. coli ST410 strains from a 281 Swiss veterinary clinic (pAN-OXA-181) (Endimiani et al., 2020, Nigg et al., 2019), and K. variicola isolated from fresh vegetables imported from Asia to Switzerland (pKS22-OXA-282 181) (Zurfluh et al., 2015b) (Figure 6). 283 The remaining 155 kb plasmid, p142_A-OXA-181, was typed Inc FIB and had regions in 284 285 common to an unnamed 136.4 kb plasmid from a human E. coli isolate from Australia (GenBank accession no. LR130556.1). Plasmid p142_A-OXA-181 also shared an ~15 kb 286 region that contained the $bla_{OXA-188}$ gene with plasmid pABC260-OXA-181 from K. 287 288 pneumoniae strain ABC260 isolated from a rectal swab in the United Arab Emirates (UAE) in 2014 (Mouftah et al., 2019) (GenBank accession no. MK412915.1) (Figure 6). 289 290 **VIM-encoding plasmids**. The bla_{VIM-1} genes in this study were located on a 160 kb IncC 291 plasmid (p009_A-VIM-1), and on an 89 kb IncR/IncY plasmid, respectively (Table 2). Plasmid p009_A-VIM-1 showed 99.9% nucleotide identity to pKP-Gr642, a bla_{VIM-19}-292 containing plasmid from a K. pneumoniae isolate recovered in 2011 from a patient 293 294 hospitalized in Greece (GenBank accession no. KR559888.1) (Papagiannitsis et al., 2016b). The *bla*_{VIM-1} gene was present on the In416-like integron In4863, comprising a *bla*_{VIM}-295

- 296 $aacA7-dfrA1-\Delta aadA1-smr2$ cassette, as in pKP-Gr642 (Papagiannitsis et al., 2016b).
- Further, the presence of a bla_{CMY-4} carrying region consisting of bla_{CMY-4} -blc-sugE- $\Delta ecnR$

indicated that p009_A-VIM-1 belongs to a unique phylogenetic lineage of IncC plasmids that

- evolved from an ancestral pUMNK88_161-like plasmid that has spread among food-
- 300 producing animals worldwide (Fernández-Alarcón et al., 2011).
- 301 Plasmid p035_A-VIM-1 was a mosaic plasmid that shared a common region over a length of
- 302 ~22 kb with pENT-576 (GenBank accession no. NZ_CP008898) from a clinical *Enterobacter*
- 303 *hormaechei* subsp. *hoffmannii* ECNIH3 isolated in 2011 in the USA (Conlan et al., 2014). A
- 304 15 kb region carrying bla_{VIM-1} was identical to a resistance region located on plasmid
- 305 pMOS94 (GenBank accession no. MK671725.1) identified in clinical Pseudomonas mosseli
- isolate AM/94 in Italy in 1994 (Di Pilato et al., 2019) (Figure 7). As described for pMOS94,
- 307 the bla_{VIM-1} gene was present on a bla_{VIM} -*aacA4*-*aphA15*-*aadA15* cassette as part of an In70
- 308 integron (Di Pilato et al., 2019). Interestingly, *P. mosseli* AM/94 represents the earliest
- 309 known VIM-1-producing strain and, as an opportunistic pathogen, is thought to have
- 310 introduced *bla*_{VIM-1} from its natural soil reservoir into the clinical setting (Giani et al., 2012b).
- Finally, ten of the CPE isolates from this study contained one or more additional plasmids
- 312 harboring genes conferring resistance to aminoglycosides, extended-spectrum beta-lactams,
- 313 fluoroquinolones and macrolides (Table S2).
- 314

315 Discussion

Currently listed by the WHO as critical-priority bacteria (Tacconelli et al., 2018), CPE have

- 317 spread globally within hospital and community settings, sewage environments and other
- environmental matrices (Mills and Lee, 2019). In this nationwide study, we detected CPE in
- 319 surface water bodies in Switzerland, including rivers, inland canals and streams
- 320 predominantly localized within urbanized areas, and none at high altitudes.

321 Among the isolates, several internationally disseminated clonal lineages harboring clinically relevant carbapenemase genes were identified. For example, E. coli ST38 is an international 322 AMR ExPEC clone responsible for the spread of OXA-48 (Pitout et al., 2019). This 323 324 particular clone has previously been identified among healthy carriers in Switzerland (Zurfluh et al., 2015a). Typically for E. coli ST38, strain CF065 chromosomally carried 325 326 bla_{OXA-48} , one notable feature distinguishing it from the other strains in this study. Further, E. coli ST410, detected in two water samples in this study, is an international high-327 328 risk ExPEC clone associated with MDR human and companion animal infections (Brilhante et al., 2020, Endimiani et al., 2020, Nigg et al., 2019, Roer et al., 2018, Timofte et al., 2016). 329 330 Comparative genome analyses allowed us to disclose a epidemiologic link between a clinical 331 NDM-5 producing E. coli ST410 strain from China (Huang et al., 2019), and a non-clinical 332 strain isolated from surface water in Switzerland. Likewise, E. coli ST167 is increasingly recognized as an MDR epidemic clone of significant 333 public-health concern, predominantly in China (Zhu et al., 2016). E. coli ST167 harboring 334 335 NDM-5 has been found previously among canine *E. coli* isolates, in fecal swabs of healthy humans employed at a veterinary clinic, and in wastewater in Switzerland (Endimiani et al., 336 2020, Peterhans et al., 2018, Zurfluh et al., 2017). Comparison of WGS data revealed genetic 337 338 similarity of clinical NDM-5 producing E. coli ST167 from Canada and India with the isolates described in this study, providing further evidence for international dissemination of 339 340 this particular NDM-5 producing ExPEC clone.

341 Other potentially pathogenic STs included *E. coli* ST73 which is a uropathogenic *E. coli*

342 (UPEC) lineage associated with community acquired urinary tract infections (UTIs) (Gibreel

et al., 2012), and *E. coli* ST648 which belongs to an emerging MDR, high-risk clonal lineage

344 occurring frequently in various sources including wild bird populations, water fowl,

345 companion animals, and humans (Schaufler et al., 2019, Hornsey et al., 2011). The

occurrence in surface water highlights the potential of these pathogenic lineages to be further
disseminated into nature via watering systems affecting agriculture and food-producing
animals, as well as to spread carbapenem resistance.

349 Virulence gene profiling revealed that the majority of the strains harbored genes associated
350 with colonization of the host gut and pathogenicity in intestinal and extra-intestinal diseases,
351 further underlining the virulence potential of the environmental CPE strains from this study.
352 Taken together, these findings indicate a possible contribution of the aquatic environment to
antibiotic-resistant infectious diseases in humans.

354 Plasmids are crucial for the horizontal spread of antimicrobial resistance genes (Carattoli,

2013). Tracking MGEs, especially plasmids, is an integral component required for a better

understanding of the dissemination of clinically relevant carbapenemases. Comparative

357 sequence analysis identified several plasmids that are considered epidemic plasmids, having

been detected in other bacterial organisms, from locations worldwide, and from human and

animal sources (Pitout et al., 2019, Carattoli, 2009). The IncL plasmid p023_D-OXA-48, and

360 the five IncX3 plasmids carrying $bla_{OXA-181}$ described in this study confirm that these types of

361 plasmids are major vehicles for dissemination of OXA-48-like carbapenemases and have

become widespread in the ecosystem. The combination of the IncX3 plasmid and the *E. coli*

363 ST410 clone, both acknowledged to possess epidemic potential (Endimiani et al., 2020,

Pitout et al., 2019, Roer et al., 2018), is an especially worrisome finding in the aquaticenvironment.

366 Likewise, IncF plasmids spread *bla*_{KPC} and *bla*_{NDM} among Enterobacteriaceae (Kopotsa et al.,

2019), and IncC plasmids like plasmid p009_A-VIM-1 from this study, have been described

368 as vehicles for bla_{VIM-1} with broad host range and interspecies, interclonal and international

369 distribution (Matsumura et al., 2018). The association of an IncC plasmid harboring VIM-1

and UPEC ST73 in surface waters is of concern, since it may pose a direct risk to publichealth.

By contrast, Col156-type plasmids harboring *bla*_{OXA-48} have only been reported in clinical 372 373 isolates from Vietnam (Honda et al., 2019). Plasmids such as the Col156 plasmid p053 E-OXA-48 detected in this study therefore provide interesting epidemiological links to 374 375 temporally and geographically segregated areas. To our knowledge, environmental E. coli harboring *bla*_{OXA-48} on a Col156-type small plasmid has not been reported so far. Although 376 rare, such plasmids may the source of resistance determinants for other epidemic plasmids. 377 In the set of plasmids analyzed in this study, three plasmids appeared to be composed of 378 379 elements from various and distinct sources. Mosaic plasmids like p062_B-KPC-2, p118_A-380 KPC-3, and p035_A-VIM-1 may provide evidence for the possible rearrangement and evolution of plasmids in the aquatic environment. Although not uncommon, the impact of 381 mosaic plasmids on public health is difficult to estimate (Pesesky et al., 2019). In these, as in 382 many of the plasmids described in this study, the presence of toxin-antitoxin modules is 383 384 likely to contribute to the maintenance of the plasmid within the strains and to the spread of carbapenem resistance genes in the environment. 385 This study has several limitations. First, due to low *in vitro* hydrolytic activity of many 386 387 carbapenemases, the detection of CPE remains difficult (Bernabeu et al., 2017), thus, an underestimation of CPE cannot be excluded. Second, in this study, we did not perform 388 389 conjugation experiments to establish transmissibility of the plasmids. Although the majority of the plasmids we analyzed shared >99% identity with known transmissible plasmids, 390

further studies to assess the conjugal dynamics of all the plasmids described in this study are

392 warranted. Third, the study was conceptualized as an observational study; periodic sampling

393 at the same sites, and at additional locations would provide further information on the

394 dynamics of dissemination of CPE and their resistomes. Given the severity of the risk of

failing antimicrobial efficacy worldwide, future studies providing such data are urgentlyneeded.

397

398 Conclusions

399 Our data point to the fact that many environmental CPE may represent anthropogenic

400 contaminants of surface waters in Switzerland. The similarity of environmental and clinical

401 isolates demonstrates their geospatial and temporal persistence locally and worldwide. This

402 study demonstrates that clinically relevant carbapenemase genes are pollutants of river

403 ecosystems and represent a significant challenge to public health and to technologies to

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404 minimize the entry into the water environment.

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427	The authors declare no conflict of interest.
428	

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430 **References**

- 431 Alonso, A., Sanchez, P., Martinez, J. L. 2001. Environmental selection of antibiotic
- 432 resistance genes. Minireview. Environ Microbiol, 3(1), 1-9.
- 433 <u>https://doi.org/10.1046/j.1462-2920.2001.00161.x.</u>
- 434 Bernabeu, S., Dortet, L., Naas, T. 2017. Evaluation of the β-CARBATM test, a colorimetric
- 435 test for the rapid detection of carbapenemase activity in Gram-negative bacilli. J
- 436 Antimicrob Chemother, 72(6), 1646-1658. <u>https://doi.org/10.1093/jac/dkx061.</u>
- 437 Brilhante, M., Menezes, J., Belas, A., Feudi, C., Schwarz, S., Pomba, C., et al. 2020. OXA-
- 438 181-producing extraintestinal pathogenic *Escherichia coli* Sequence Type 410 isolated
- from a dog in Portugal. Antimicrob Agents Chemother, 64(4), pii: e02298-19.
- 440 <u>https://doi.org/10.1128/aac.02298-19.</u>
- 441 Carattoli, A. 2009. Resistance plasmid families in Enterobacteriaceae. Antimicrob Agents
 442 Chemother, 53(6), 2227-2238. https://doi.org/10.1128/AAC.01707-08.
- 443 Carattoli, A. 2013. Plasmids and the spread of resistance. I J Med Microbiol, 303(6-7), 298-
- 444 304. <u>https://www.sciencedirect.com/science/article/pii/S1438422113000167.</u>
- 445 Carattoli, A., Zankari, E., Garcìa-Fernandez, A., Larsen, M. V., Lund, O., Villa, L., et al.
- 446 2014. PlasmidFinder and pMLST: in silico detection and typing of plasmids.
- 447 Antimicrob Agents Chemother, 58, 3895-3903.
- 448 <u>https://aac.asm.org/content/aac/early/2014/04/22/AAC.02412-14.full.pdf.</u>
- 449 CLSI. 2018. Performance Standards for Antimicrobial Susceptibility Testing. 28th ed. CLSI
- 450 supplement M100. Clinical and Laboratory Standards Institute; Wayne, PA.
- 451 Conlan, S., Thomas, P. J., Deming, C., Park, M., Lau, A. F., Dekker, J. P., et al. 2014. Single-
- 452 molecule sequencing to track plasmid diversity of hospital-associated carbapenemase-
- 453 producing Enterobacteriaceae. Sci Transl Med, 6(254), 254ra126.
- 454 <u>https://doi.org/10.1126/scitranslmed.3009845.</u>

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- 455 Di Pilato, V., Antonelli, A., Giani, T., Henrici De Angelis, L., Rossolini, G. M., Pollini, S.
- 456 2019. Identification of a novel plasmid lineage associated with the dissemination of
- 457 metallo- β -lactamase genes among pseudomonads. Front Microbiol, 10, 1504.
- 458 <u>https://doi.org/10.3389/fmicb.2019.01504.</u>
- 459 Ellington, M. J., Kistler, J., Livermore, D. M., Woodford, N. 2007. Multiplex PCR for rapid
- detection of genes encoding acquired metallo- β -lactamases. J Antimicrob Chemother,

461 59(2), 321-322. <u>https://doi.org/10.1093/jac/dkl481.</u>

- 462 Endimiani, A., Brilhante, M., Bernasconi, O. J., Perreten, V., Schmidt, J. S., Dazio, V., et al.
- 463 2020. Employees of Swiss veterinary clinics colonized with epidemic clones of
- 464 carbapenemase-producing *Escherichia coli*. J Antimicrob Chemother, 75(3), 766-768.
- 465 <u>https://doi.org/10.1093/jac/dkz470.</u>
- 466 Essack, S. Y. 2018. Environment: the neglected component of the One Health triad. Lancet
- 467 Planet Health, 2(6), e238-e239. <u>https://doi.org/10.1016/S2542-5196(18)30124-4.</u>
- 468 ECDC. 2019. Carbapenem-resistant Enterobacteriaceae, second update 26 September.
- 469 European Centre for Disease Prevention and Control.
- 470 <u>https://www.ecdc.europa.eu/sites/default/files/documents/carbapenem-resistant-</u>
- 471 <u>enterobacteriaceae-risk-assessment-rev-2.pdf.</u>
- 472 Falgenhauer, L., Imirzalioglu, C., Ghosh, H., Gwozdzinski, K., Schmiedel, J., Gentil, K., et
- al. 2016. Circulation of clonal populations of fluoroquinolone-resistant CTX-M-15-
- 474 producing *Escherichia coli* ST410 in humans and animals in Germany. Int J
- 475 Antimicrob Agents, 47(6), 457-465. <u>https://doi.org/10.1016/j.ijantimicag.2016.03.019.</u>
- 476 Falgenhauer, L., Schwengers, O., Schmiedel, J., Baars, C., Lambrecht, O., Heß, S., et al.
- 477 2019. Multidrug-resistant and clinically relevant gram-negative bacteria are present in
- 478 German surface waters. Front Microbiol, 10, 2779.
- 479 <u>https://doi.org/10.3389/fmicb.2019.02779.</u>

- 480 Fernández-Alarcón, C., Singer, R. S., Johnson, T. J. 2011. Comparative genomics of
- 481 multidrug resistance-encoding IncA/C plasmids from commensal and pathogenic
- 482 *Escherichia coli* from multiple animal sources. PLoS One, 6(8), e23415.
- 483 <u>https://doi.org/10.1371/journal.pone.0023415.</u>
- 484 Furness, L. E., Campbell, A., Zhang, L., Gaze, W. H., McDonald, R. A. 2017. Wild small
- 485 mammals as sentinels for the environmental transmission of antimicrobial resistance.

486 Environ Res, 154, 28-34. <u>https://doi.org/10.1016/j.envres.2016.12.014.</u>

- 487 Galata, V., Fehlmann, T., Backes, C., Keller, A. 2019. PLSDB: a resource of complete
- 488 bacterial plasmids. Nucleic Acids Res, 47(D1), D195-D202.
- 489 <u>https://doi.org/10.1093/nar/gky1050</u>
- 490 Giakkoupi, P., Maltezou, H., Polemis, M., Pappa, O., Saroglou, G., Vatopoulos, A. 2009.
- 491 KPC-2-producing *Klebsiella pneumoniae* infections in Greek hospitals are mainly due
- to a hyperepidemic clone. Euro Surveill., 14(21), 19218.
- 493 https://www.eurosurveillance.org/content/10.2807/ese.14.21.19218-
- 494 <u>en?crawler=true&mimetype=application/pdf</u>
- 495 Giani, T., Conte, V., Di Pilato, V., Aschbacher, R., Weber, C., Larcher, C., et al. 2012a.
- 496 *Escherichia coli* from Italy producing OXA-48 carbapenemase encoded by a novel
- 497 Tn*1999* transposon derivative. Antimicrob Agents Chemother, 56(4), 2211-2213.
- 498 <u>https://doi.org/10.1128/AAC.00035-12.</u>
- 499 Giani, T., Marchese, A., Coppo, E., Kroumova, V., Rossolini, G. M. 2012b. VIM-1-
- 500 producing *Pseudomonas mosselii* isolates in Italy, predating known VIM-producing
- 501 index strains. Antimicrob Agents Chemother, 56(4), 2216-2217.
- 502 <u>https://doi.org/10.1128/AAC.06005-11.</u>
- 503 Gibreel, T. M., Dodgson, A. R., Cheesbrough, J., Fox, A. J., Bolton, F. J., Upton, M. 2012.
- 504 Population structure, virulence potential and antibiotic susceptibility of uropathogenic

- 505 *Escherichia coli* from Northwest England. J Antimicrob Chemother, 67(2), 346-356.
- 506 <u>https://doi.org/10.1093/jac/dkr451.</u>
- 507 Gillings, M. R., Westoby, M., Ghaly, T. M. 2018. Pollutants that replicate: xenogenetic
- 508 DNAs. Trends Microbiol, 26(12), 975-977.
- 509 https://www.sciencedirect.com/science/article/pii/S0966842X18301756.
- 510 Gorrie, C. L., Mirceta, M., Wick, R. R., Judd, L. M., Wyres, K. L., Thomson, N. R., et al.
- 511 2018. Antimicrobial-resistant *Klebsiella pneumoniae* carriage and infection in
- 512 specialized geriatric care wards linked to acquisition in the referring hospital. Clin
- 513 Infect Dis, 67(2), 161-170. <u>https://doi.org/10.1093/cid/ciy027.</u>
- 514 Hernando-Amado, S., Coque, T. M., Baquero, F., Martínez, J. L. 2019. Defining and
- 515 combating antibiotic resistance from One Health and Global Health perspectives. Nat
- 516 Microbiol, 4(9), 1432-1442. <u>https://doi.org/10.1038/s41564-019-0503-9.</u>
- 517 Honda, N. H., Aoki, K., Kamisasanuki, T., Matsuda, N., To, M., Matsushima, H., et al. 2019.
- 518 Isolation of three distinct carbapenemase-producing Gram-negative bacteria from a
- 519 Vietnamese medical tourist. J Infect Chemother, 25(10), 811-815.
- 520 <u>https://doi.org/10.1016/j.jiac.2019.03.020.</u>
- 521 Hornsey, M., Phee, L., Wareham, D. W. 2011. A novel variant, NDM-5, of the New Delhi
- 522 metallo-β-lactamase in a multidrug-resistant *Escherichia coli* ST648 isolate recovered
- from a patient in the United Kingdom. Antimicrob Agents Chemother, 55(12), 5952-
- 524 5954. <u>https://doi.org/10.1128/AAC.05108-11.</u>
- 525 Huang, J., Ma, S., Yu, Q., Fu, M., Shao, L., Shan, X., et al. 2019. Whole genome sequence of
- 526 an *Escherichia coli* ST410 isolate co-harbouring *bla*_{NDM-5}, *bla*_{OXA-1}, *bla*_{CTX-M-15}, *bla*_{CMY-}
- 527 $_2$, aac(3)-IIa and aac(6')-Ib-cr genes isolated from a patient with bloodstream infection
- 528 in China. J Glob Antimicrob Resist, 19, 354-355.
- 529 <u>https://doi.org/10.1016/j.jgar.2019.10.027.</u>

- 530 Jelić, M., Hrenović, J., Dekić, S., Goić-Barišić, I., Tambić Andrašević, A. 2019. First
- 531 evidence of KPC-producing ST258 *Klebsiella pneumoniae* in river water. J Hosp

532 Infect, 103(2), 147-150. <u>https://doi.org/10.1016/j.jhin.2019.04.001.</u>

- Jia, B., Raphenya, A. R., Alcock, B., Waglechner, N., Guo, P., Tsang, K. K., et al. 2017.
- 534 CARD 2017: expansion and model-centric curation of the comprehensive antibiotic
- resistance database. Nucleic Acids Res, 45(D1), D566-D573.
- 536 https://doi.org/10.1093/nar/gkw1004.
- 537 Joensen, K. G., Tetzschner, A. M. M., Iguchi, A., Aarestrup, F. M., Scheutz, F. 2015. Rapid
- and easy in silico serotyping of *Escherichia coli* isolates by use of whole-genome
- sequencing data. J Clin Microbiol, 53(8), 2410-2426.
- 540 <u>https://jcm.asm.org/content/jcm/53/8/2410.full.pdf.</u>
- 541 Kaper, J. B., Nataro, J. P., Mobley, H. L. 2004. Pathogenic *Escherichia coli*. Nature Rev
- 542 Microbiol, 2(2), 123-140. https://doi:10.1038/nrmicro818.
- 543 Khan, F. A., Hellmark, B., Ehricht, R., Söderquist, B., Jass, J. 2018. Related carbapenemase-
- 544 producing *Klebsiella* isolates detected in both a hospital and associated aquatic
- environment in Sweden. Eur J Clin Microbiol Infect Dis, 37(12), 2241-2251.
- 546 <u>https://doi.org/10.1007/s10096-018-3365-9.</u>
- 547 Kraemer, S. A., Ramachandran, A., Perron, G. G. 2019. Antibiotic pollution in the
- 548 environment: From microbial ecology to public policy. Microorganisms, 7(6), 180.
- 549 <u>https://doi.org/10.3390/microorganisms7060180.</u>
- 550 Kopotsa, K., Osei Sekyere, J., Mbelle, N. M. 2019. Plasmid evolution in carbapenemase-
- producing Enterobacteriaceae: a review. Ann N Y Acad Sci, 1457(1), 61-91.
- 552 <u>https://doi.org/10.1111/nyas.14223.</u>
- 553 Lepuschitz, S., Schill, S., Stoeger, A., Pekard-Amenitsch, S., Huhulescu, S., Inreiter, N., et al.
- 554 2019. Whole genome sequencing reveals resemblance between ESBL-producing and

555	carbapenem resistant Klebsiella pneumoniae isolates from Austrian rivers and clinical
556	isolates from hospitals. Sci Total Environ, 662, 227-235.
557	https://doi.org/10.1016/j.scitotenv.2019.01.179.
558	Magiorakos, AP., Srinivasan, A., Carey, R. B., Carmeli, Y., Falagas, M. E., Giske, C. G., et
559	al. 2012. Multidrug-resistant, extensively drug-resistant and pandrug-resistant bacteria:
560	an international expert proposal for interim standard definitions for acquired resistance.
561	Clin Microbiol Infect, 18(3), 268-281. <u>https://doi.org/10.1111/j.1469-</u>
562	<u>0691.2011.03570.x.</u>
563	Mahon, B. M., Brehony, C., McGrath, E., Killeen, J., Cormican, M., Hickey, P., et al. 2017.
564	Indistinguishable NDM-producing Escherichia coli isolated from recreational waters,
565	sewage, and a clinical specimen in Ireland, 2016 to 2017. Euro Surveill, 22(15).
566	https://doi.org/10.2807/1560-7917.ES.2017.22.15.30513.
567	Marti, E., Variatza, E., Balcazar, J. L. 2014. The role of aquatic ecosystems as reservoirs of
568	antibiotic resistance. Trends Microbiol, 22(1), 36-41.
569	https://doi.org/10.1016/j.tim.2013.11.001.
570	Marti, R., Stephan, R., Klumpp, J., Nüesch-Inderbinen, M., Hummerjohann, J., Bagutti, C., et

al. 2017. Complete genome sequence of *Enterobacter cloacae* 704SK10, an OXA-48-

572 encoding wastewater isolate. Genome Announc, 5(33), e00830-17.

- 573 <u>https://doi.org/10.1128/genomeA.00830-17.</u>
- 574 Mataseje, L. F., Boyd, D. A., Fuller, J., Haldane, D., Hoang, L., Lefebvre, B., et al. 2018.
- 575 Characterization of OXA-48-like carbapenemase producers in Canada, 2011-14. J
- 576 Antimicrob Chemother, 73(3), 626-633. <u>https://doi.org/10.1093/jac/dkx462.</u>
- 577 Mathers, A. J., Stoesser, N., Sheppard, A. E., Pankhurst, L., Giess, A., Yeh, A. J., et al. 2015.
- 578 *Klebsiella pneumoniae* carbapenemase (KPC)-producing *K. pneumoniae* at a single

- 579 institution: insights into endemicity from whole-genome sequencing. Antimicrob
- 580 Agents Chemother, 59(3), 1656-1663. <u>https://doi.org/10.1128/AAC.04292-14.</u>
- 581 Matsumura, Y., Peirano, G., Bradford, P. A., Motyl, M. R., DeVinney, R., Pitout, J. D. D.
- 582 2018. Genomic characterization of IMP and VIM carbapenemase-encoding transferable
- plasmids of Enterobacteriaceae. J Antimicrob Chemother, 73(11), 3034-3038.
- 584 <u>https://doi.org/10.1093/jac/dky303.</u>
- 585 McArthur, A. G., Waglechner, N., Nizam, F., Yan, A., Azad, M. A., Baylay, A. J., et al.
- 586 2013. The comprehensive antibiotic resistance database. Antimicrob Agents
- 587 Chemother, 57(7), 3348-3357. <u>https://doi.org/10.1128/AAC.00419-13.</u>
- 588 Mills, M. C., Lee, J. 2019. The threat of carbapenem-resistant bacteria in the environment:
- 589 Evidence of widespread contamination of reservoirs at a global scale. Environ Pollut,

590 255(Pt 1), 113143. <u>https://doi.org/10.1016/j.envpol.2019.113143.</u>

- 591 Mouftah, S. F., Pál, T., Darwish, D., Ghazawi, A., Villa, L., Carattoli, A., et al. 2019.
- 592 Epidemic IncX3 plasmids spreading carbapenemase genes in the United Arab Emirates
- and worldwide. Infect Drug Resist, 12, 1729-1742.
- 594 <u>https://doi.org/10.2147/IDR.S210554.</u>
- 595 Naas, T., Cuzon, G., Truong, H. V., Nordmann, P. 2012. Role of ISKpn7 and deletions in
- 596 bla_{KPC} gene expression. Antimicrob Agents Chemother, 56(9), 4753-4759.
- 597 <u>https://doi.org/10.1128/AAC.00334-12.</u>
- 598 Nigg, A., Brilhante, M., Dazio, V., Clément, M., Collaud, A., Gobeli Brawand, S., et al.
- 599 2019. Shedding of OXA-181 carbapenemase-producing *Escherichia coli* from
- 600 companion animals after hospitalisation in Switzerland: an outbreak in 2018. Euro
- 601 Surveill, 24(39). <u>https://doi.org/10.2807/1560-7917.ES.2019.24.39.1900071.</u>

- 602 Nordmann, P., Naas, T., Poirel, L. 2011. Global spread of Carbapenemase-producing
- Enterobacteriaceae. Emerg Infect Dis, 17(10), 1791-1798.
- 604 <u>https://doi.org/10.3201/eid1710.110655.</u>
- 605 Overballe-Petersen, S., Roer, L., Ng, K., Hansen, F., Justesen, U. S., Andersen, L. P., et al.
- 606 2018. Complete nucleotide sequence of an *Escherichia coli* Sequence Type 410 strain
- 607 carrying bla_{NDM-5} on an IncF multidrug resistance plasmid and blaOXA-181 on an
- IncX3 plasmid. Genome Announc, 6(5). <u>https://doi.org/10.1128/genomeA.01542-17.</u>
- 609 Papagiannitsis, C. C., Di Pilato, V., Giani, T., Giakkoupi, P., Riccobono, E., Landini, G., et
- al. 2016a. Characterization of KPC-encoding plasmids from two endemic settings,
- 611 Greece and Italy. J Antimicrob Chemother, 71(10), 2824-2830.
- 612 <u>https://doi.org/10.1093/jac/dkw227.</u>
- 613 Papagiannitsis, C. C., Dolejska, M., Izdebski, R., Giakkoupi, P., Skálová, A., Chudějová,
- 614 K., et al. 2016b. Characterisation of IncA/C2 plasmids carrying an In416-like integron
- 615 with the *bla*_{VIM-19} gene from *Klebsiella pneumoniae* ST383 of Greek origin. Int J
- 616 Antimicrob Agents, 47(2), 158-162. <u>https://doi.org/10.1016/j.ijantimicag.2015.12.001.</u>
- 617 Pesesky, M. W., Tilley, R., Beck, D. A. C. 2019. Mosaic plasmids are abundant and unevenly
- 618 distributed across prokaryotic taxa. Plasmid, 102, 10-18.
- 619 <u>https://doi.org/10.1016/j.plasmid.2019.02.003.</u>
- 620 Peterhans, S., Stevens, M. J. A., Nüesch-Inderbinen, M., Schmitt, S., Stephan, R., Zurfluh, K.
- 621 2018. First report of a *bla*_{NDM-5}-harbouring *Escherichia coli* ST167 isolated from a
- 622 wound infection in a dog in Switzerland. J Glob Antimicrob Resist, 15, 226-227.
- 623 <u>https://doi.org/10.1016/j.jgar.2018.10.013.</u>
- 624 Pitout, J. D. D., Peirano, G., Kock, M. M., Strydom, K. A., Matsumura, Y. 2019. The global
- 625 ascendency of OXA-48-type carbapenemases. Clin Microbiol Rev, 33(1).
- 626 <u>https://doi.org/10.1128/CMR.00102-19.</u>

- 627 Poirel, L., Barbosa-Vasconcelos, A., Simões, R. R., Da Costa, P. M., Liu, W., Nordmann, P.
- 628 2012. Environmental KPC-producing *Escherichia coli isolates in Portugal*. Antimicrob
- 629 Agents Chemother, 56(3), 1662-1663. <u>https://doi.org/10.1128/AAC.05850-11.</u>
- 630 Poirel, L., Walsh, T. R., Cuvillier, V., Nordmann, P. 2011. Multiplex PCR for detection of
- 631 acquired carbapenemase genes. Diagn Microbiol Infect Dis, 70(1), 119-123.
- 632 <u>https://doi.org/10.1016/j.diagmicrobio.2010.12.002.</u>
- 633 Pruden, A. 2014. Balancing water sustainability and public health goals in the face of
- 634 growing concerns about antibiotic resistance. Environ Sci Technol, 48(1), 5-14.
- 635 <u>https://doi.org/10.1021/es403883p.</u>
- 636 Queenan, A. M., Bush, K. 2007. Carbapenemases: the versatile beta-lactamases. Clin
- 637 Microbiol Rev, 20(3), 440-58, table of contents. <u>https://doi.org/10.1128/CMR.00001-</u>
 638 <u>07.</u>
- 639 Roer, L., Overballe-Petersen, S., Hansen, F., Schønning, K., Wang, M., Røder, B. L., et al.
- 640 2018. *Escherichia coli* Sequence Type 410 is causing new international high-risk

641 clones. mSphere, 3(4). https://doi.org/10.1128/mSphere.00337-18.

- 642 Schaufler, K., Semmler, T., Wieler, L. H., Trott, D. J., Pitout, J., Peirano, G., et al. 2019.
- 643 Genomic and functional analysis of emerging virulent and multidrug-resistant

644 *Escherichia coli* lineage. Antimicrob Agents Chemother, 63(6).

- 645 <u>https://doi.org/10.1128/AAC.00243-19.</u>
- 646 Sheppard, A. E., Stoesser, N., Sebra, R., Kasarskis, A., Deikus, G., Anson, L., et al. 2016.
- 647 Complete genome sequence of KPC-producing *Klebsiella pneumoniae* strain
- 648 CAV1193. Genome Announc, 4(1). <u>https://doi.org/10.1128/genomeA.01649-15.</u>
- 649 Skalova, A., Chudejova, K., Rotova, V., Medvecky, M., Studentova, V., Chudackova, E., et
- al. 2017. Molecular characterization of OXA-48-like-producing Enterobacteriaceae in

- the Czech Republic and evidence for horizontal transfer of pOXA-48-like plasmids.
- Antimicrob Agents Chemother, 61(2). <u>https://doi.org/10.1128/AAC.01889-16.</u>
- 653 Stevens, M. J. A., Tasara, T., Klumpp, J., Stephan, R., Ehling-Schulz, M., Johler, S. 2019.
- 654 Whole-genome-based phylogeny of *Bacillus cytotoxicus* reveals different clades within
- the species and provides clues on ecology and evolution. Sci Rep, 9(1), 1984.
- 656 https://doi.org/10.1038/s41598-018-36254-x.
- 657 Stoesser, N., Sheppard, A. E., Peirano, G., Sebra, R., Lynch, T., Anson, L., et al. 2016.
- 658 Complete sequencing of plasmids containing *bla*_{OXA-163} and *bla*_{OXA-48} in *Escherichia*
- *coli* Sequence Type 131. Antimicrob Agents Chemother, 60(11), 6948-6951.
- 660 <u>https://doi.org/10.1128/AAC.01130-16.</u>
- 661 Sugawara, Y., Akeda, Y., Hagiya, H., Sakamoto, N., Takeuchi, D., Shanmugakani, R. K., et
- al. 2019. Spreading patterns of NDM-producing Enterobacteriaceae in clinical and
- 663 environmental settings in Yangon, Myanmar. Antimicrob Agents Chemother, 63(3),
- 664 e01924-18. <u>https://doi.org/10.1128/AAC.01924-18.</u>
- 665 Tacconelli, E., Carrara, E., Savoldi, A., Harbarth, S., Mendelson, M., Monnet, D. L., et al.
- 666 2018. Discovery, research, and development of new antibiotics: the WHO priority list
- of antibiotic-resistant bacteria and tuberculosis. Lancet Infect Dis, 18(3), 318-327.
- 668 https://www.sciencedirect.com/science/article/pii/S1473309917307533.
- 669 Timofte, D., Maciuca, I. E., Williams, N. J., Wattret, A., Schmidt, V. 2016. Veterinary
- 670 hospital dissemination of CTX-M-15 extended-spectrum beta-lactamase-producing
- *Escherichia coli* ST410 in the United Kingdom. Microb Drug Resist, 22(7), 609-615.
- 672 <u>https://doi.org/10.1089/mdr.2016.0036.</u>
- Treangen, T. J., Ondov, B. D., Koren, S., Phillippy, A. M. 2014. The Harvest suite for rapid
- 674 core-genome alignment and visualization of thousands of intraspecific microbial

- 675 genomes. Genome biology, 15(11), 524.
- 676 <u>https://link.springer.com/article/10.1186/s13059-014-0524-x.</u>
- Turton, J. F., Doumith, M., Hopkins, K. L., Perry, C., Meunier, D., Woodford, N. 2016.
- 678 Clonal expansion of *Escherichia coli* ST38 carrying a chromosomally integrated OXA-
- 48 carbapenemase gene. J Med Microbiol, 65(6), 538-546.
- 680 <u>https://doi.org/10.1099/jmm.0.000248.</u>
- van Duin, D., Doi, Y. 2017. The global epidemiology of carbapenemase-producing
- Enterobacteriaceae. Virulence, 8(4), 460-469.
- 683 <u>https://doi.org/10.1080/21505594.2016.1222343.</u>
- 684 Wasyl, D., Kern-Zdanowicz, I., Domańska-Blicharz, K., Zając, M., Hoszowski, A. 2015.
- 685 High-level fluoroquinolone resistant *Salmonella enterica* serovar Kentucky ST198
- 686 epidemic clone with IncA/C conjugative plasmid carrying $bla_{CTX-M-25}$ gene. Vet
- 687 Microbiol, 175(1), 85-91. <u>https://doi.org/10.1016/j.vetmic.2014.10.014.</u>
- 688 Williams, M. R., Stedtfeld, R. D., Guo, X., Hashsham, S. A. 2016. Antimicrobial resistance
- in the environment. Water Environ Res, 88(10), 1951-1967.
- 690 <u>https://doi.org/10.2175/106143016x14696400495974.</u>
- 691 Wirth, T., Falush, D., Lan, R., Colles, F., Mensa, P., Wieler, L. H., et al. 2006. Sex and
- 692 virulence in *Escherichia coli*: an evolutionary perspective. Mol Microbiol, 60(5), 1136-
- 693 1151. <u>https://doi.org/10.1111/j.1365-2958.2006.05172.x.</u>
- 694 WHO. 2018. WHO report on surveillance of antibiotic consumption: 2016-2018 early
- 695 implementation. Geneva, Switzerland.
- 696 WHO. 2017. Critically important antimicrobials for human medicine 5th rev. Geneva,
- 697 Switzerland.

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- 698 Xie, Y., Wei, Y., Shen, Y., Li, X., Zhou, H., Tai, C., et al. 2018. TADB 2.0: an updated
- database of bacterial type II toxin–antitoxin loci. Nucleic Acids Res, 46(D1), D749-

700 D753. <u>https://academic.oup.com/nar/article/46/D1/D749/4584634.</u>

- Yang, Q. E., Walsh, T. R. 2017. Toxin-antitoxin systems and their role in disseminating and
- maintaining antimicrobial resistance. FEMS Microbiol Rev, 41(3), 343-353.
- 703 <u>https://doi.org/10.1093/femsre/fux006.</u>
- Yigit, H., Queenan, A. M., Anderson, G. J., Domenech-Sanchez, A., Biddle, J. W., Steward,
- 705 C. D., et al. 2001. Novel carbapenem-hydrolyzing beta-lactamase, KPC-1, from a
- carbapenem-resistant strain of *Klebsiella pneumoniae*. Antimicrob Agents Chemother,
- 707 45(4), 1151-1161. <u>https://doi.org/10.1128/AAC.45.4.1151-1161.2001.</u>
- 708 Zhu, Y. Q., Zhao, J. Y., Xu, C., Zhao, H., Jia, N., Li, Y. N. 2016. Identification of an NDM-
- 5-producing *Escherichia coli* Sequence Type 167 in a neonatal patient in China. Sci
- 710 Rep, 6, 29934. <u>https://doi.org/10.1038/srep29934.</u>
- 711 Zurfluh, K., Bagutti, C., Brodmann, P., Alt, M., Schulze, J., Fanning, S., et al. 2017.
- 712 Wastewater is a reservoir for clinically relevant carbapenemase- and 16s rRNA
- 713 methylase-producing Enterobacteriaceae. Int J Antimicrob Agents, 50(3), 436-440.
- 714 <u>https://doi.org/10.1016/j.ijantimicag.2017.04.017.</u>
- 715 Zurfluh, K., Nüesch-Inderbinen, M. T., Poirel, L., Nordmann, P., Hächler, H., Stephan, R.
- 716 2015a. Emergence *of Escherichia coli* producing OXA-48 β-lactamase in the
- 717 community in Switzerland. Antimicrob Resist Infect Control, 4, 9.
- 718 <u>https://doi.org/10.1186/s13756-015-0051-x.</u>
- 719 Zurfluh, K., Poirel, L., Nordmann, P., Klumpp, J., Stephan, R. 2015b. First detection of
- 720 *Klebsiella variicola* producing OXA-181 carbapenemase in fresh vegetable imported
- from Asia to Switzerland. Antimicrob Resist Infect Control, 4, 38.
- 722 <u>https://doi.org/10.1186/s13756-015-0080-5.</u>

- 723 Zurfluh, K., Hächler, H., Nüesch-Inderbinen, M., Stephan, R. 2013. Characteristics of
- β extended-spectrum β -lactamase- and carbapenemase-producing Enterobacteriaceae
- isolates from rivers and lakes in Switzerland. Appl Environ Microbiol, 79(9), 3021-
- 726 3026. <u>https://doi.org/10.1128/AEM.00054-13</u>
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- **Figure 1.** Map of Switzerland showing bodies of water, sample locations, and carbapenemase
- 736 gene status.
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742 carbapenemase producing Enterobacteriaceae isolated from surface water bodies in

- 743 Switzerland. Abbreviations: AM, ampicillin; AMC, amoxicillin-clavulanic acid; AZM,
- aztreonam; C, chloramphenicol; CIP, ciprofloxacin; CTX, cefotaxime; CZ, cefazolin; ETP,
- rtapenem; FEP, cefepime; F/M, nitrofurantoin; GM, gentamycin; IP, imipenem; K,
- 746 kanamycin; MDR, multidrug resistance; MP, meropenem; NA, nalidixic acid; S,
- streptomycin; ST, sequence type; SXT, sulfamethoxazole//trimethoprim; TE, tetracycline; -,
- not applicable or not performed; *, *E. coli* with intestinal or extraintestnal pathogenic
- virulence genes. Colors of squares categorizing antibiotic resistance profiles: Pink, resistant;
- 750 yellow, intermediate; green, susceptible, purple, multidrug resistant.







Figure 3. Comparative circular maps of bla_{KPC} -carrying plasmids generated using BRIG. The positions of the bla_{KPC} genes are indicated.

- Left panel: p062_B-KPC-2 (GenBank acc. no. CP048384.1). The rings from the inner to the
- outer represent plasmids p062_B-KPC-2 from *C. freundii* from this study (yellow), p1643_10
- 757 (GenBank acc. no. KF056330) from poultry Salmonella Kentucky isolate 1643/2010
- (orange), and pKP1504-kpc (GenBank acc. no. KF874496) from clinical K. pneumoniae
- isolate GR-1504 (yellow).



- from the inner to the outer represent plasmids p118_A-KPC-3 from K. variicola from this
- study (orange), unnamed plasmid (GenBank acc. no. NZ_CP024500.1) from K. pneumoniae
- 763 RJY9645 (green), unnamed plasmid (GenBank acc. no. NZ_CP029102.1) from *K*.
- *pneumoniae* strain AR438 (red), and pY9645-166 (GenBank acc. no. CP044029.1) from
- 765 clinical *K. pneumoniae* isolate RJY9645 (blue).
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- **Figure 4.** Comparative circular maps of *bla*_{NDM-5}-carrying plasmids generated using BRIG.
- The positions of the bla_{NDM-5} genes are indicated.
- Left panel: p038_A-NDM-5 (GenBank acc. no. CP048377.1). The rings from the inner to the
- outer represent plasmids pAMA1167-NDM-5 from *E. coli* ST410 (GenBank acc. no.
- 773 NZ_CP024805.1), and p038_A-NDM-5 from *E. coli* ST410 from this study.
- Right panel: p164_A-NDM-5 (GenBank acc. no. CP048368.1). The rings from the inner to
- the outer represent plasmids pM309-NDM5DNA from *E. coli* ST167 (GenBank acc. no.
- AP018833.1), and p164_A-NDM-5 from *E. coli* ST167 from this study.
- 777





- 781 Left panel: p023_D-OXA-48 (GenBank acc. no. CP048353.1). The rings from the inner to
- the outer represent plasmids p704SK10_2 from *E. cloacae* (GenBank acc. no. CP022150),
- pOXA-48_4963 from *K. pneumoniae* (GenBank acc. no. KX523900), p023_D-OXA-48 from
- 784 *R. ornithinolytica* from this study, and pEC745 from *E. coli* ST131 (GenBank acc. no.
- 785 CP015075.1).
- 786 Right panel: p053_E-OXA-48 (GenBank acc. no. CP048364.1). The rings from the inner to
- the outer represent plasmids p053_E-OXA-48 from *E. coli* ST205 from this study, and
- pMTY17816_OXA48 from *K. pneumoniae* isolate (GenBank acc. no. AP019554.1.
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represent plasmids p032_K-OXA-181 from *E. coli* ST1284 (GenBank acc. no. CP048321.1,

this study), pOXA-181_29144 from K. pneumoniae (GenBank acc. no.KX523903.1),

p061_A-OXA-181 from E. coli ST940 (GenBank acc. no. CP048327.1, this study), pAN-

798 OXA-181 from *E. coli* ST410 (GenBank acc. no. MK416154), p124_B-OXA-181 from *E.*

- 799 *coli* ST410 (GenBank acc. no. CP048346.1, this study), p010_B-OXA-181 *E. coli* ST656
- (GenBank acc. no.CP048332.1, this study), p064_C-OXA-181 E. coli ST940 (GenBank acc.
- no. CP048325.1, this study), and pKS22 from *K. variicola* (GenBank acc. no. KT005457).
- Right panel: Mosaic structure of p142_A-OXA181 (GenBank acc. no. CP048338.1). The
- rings from the inner to the outer represent plasmids p142_A-OXA181 from this study (pink),
- unnamed plasmid (GenBank acc. no. NZ_LR130556.1) from E. coli (green), and pABC260-
- 805 OXA-181(GenBank acc.no. MK412915.1) from *K. pneumoniae* (turquoise).
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- **Figure 7.** Comparative circular maps of *bla*_{VIM-1}-carrying plasmids generated using BRIG.
- 810 The positions of the bla_{VIM-1} genes are indicated.
- 811 Left panel: p009_A-VIM-1 (CP048305.1). The rings from the inner to the outer represent
- 812 plasmids p009_A-VIM-1 from *K. aerogenes* from this study, and pKP-Gr642 from a *K*.
- 813 *pneumoniae* isolate (GenBank acc. no. KR559888.1).
- 814 Right panel: Mosaic structure of p035_A-VIM-1 (GenBank acc. no. CP050069.1). The rings
- from the inner to the outer represent plasmids p035_A-VIM-1 from this study (green),
- 816 plasmid pENT-576 (GenBank acc. no. NZ_CP008898) from *E. cloacae* (turquoise), and
- pMOS94 (GenBank acc. no. MK671725.1) identified in clinical *P. mosseli* (green).
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821 Tables 1-2

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Table 1. Virulence factor profiles of 11 carbapenemase-producing *E. coli* strains cultured from water bodies in Switzerland

Strain	Carbapenemase	ST	Virulence factor (s)
ID			
CF038	NDM-5	410	lpfA
CF164	NDM-5	167	capU, gad, iss
CF053	OXA-48	205	astA, gad, lpfA
CF065	OXA-48	38	air, eilA, iss
CF124	OXA-181	410	lpfA
CF142	OXA-181	648	air, eilA, gad, iha, lpfA, nfaE, sat
CF010	OXA-181	656	gad, iss
CF061	OXA-181	940	capU, gad, lpfA
CF064	OXA-181	940	capU, gad, lpfA
CF032	OXA-181	1284	astA, capU, gad, iss
CF009	VIM-1	73	capU, iha, iroN, iss, mchB, mchC, mchF, mcmA, pic, sat, vat

825 *air*, enteroaggregative immunoglobulin repeat protein gene; *astA*, heat-stable toxin gene; *capU*,

826 hexosyltransferase homolog gene; eilA, Salmonella invasion gene activator hilA homolog gene; gad, glutamate

827 decarboxylase; *iha*, iron-regulated adhesin gene; *iroN*, enterobactin siderophore receptor gene; *lpfA*, long polar

fimbriae gene; mchB, gene for microcin H47 part of colicin H; mchC, MchC protein gene; mchF, ABC
transporter protein MchF gene; mcmA, gene for microcin M part of colicin H; nfaE, diffuse adherence fibri

transporter protein MchF gene; mcmA, gene for microcin M part of colicin H; nfaE, diffuse adherence fibrillar
 adhesin gene; iss, increased serum survival; pic, serine protease autotransporter gene of Enterobacteriaceae

831 (SPATE); *sat*, secreted autotransporter toxin gene; *vat*, vacuolating autotransporter toxin gene.

OUTRIC

832	Table 2. Summar	y of the features	associated with	16 carba	penemase-ei	ncoding r	plasmids from	n Entero	bacteriaceae	strains cul	tured fron	n surface wa	ter
						()							

833 bodies in Switzerland

Strain	Host species (ST)	Carbapenemase	Plasmid	Plasmid	Inc group	Other AMR genes	T/A family	Accession no.
ID				size				
CF062	C. freundii (–)	KPC-2	p062_B-KPC-2	190 kb	IncQ	aph(3')-Ia, $aph(6)$ -Id, $aph(3'')$ -Ib, $sul2$, $ant(2'')$ -Ia, $dfrA12$, $aadA$, $sul1$, bla_{OXA-9} , bla_{TYMA} , bla_{CYMA} , $again and a subscripts of the second seco$	yafQ/dinJ	CP048384.1
CF070	E. kobei (–)	KPC-2	p070 A-KPC-2	110 kb	IncFIB _v	bla _{TEM 1}	_	CP050075.1
CF118	K. variicola (–)	KPC-3	p118_A-KPC-3	244 kb	IncFIB	mphA, sul2, dfrA12, aadA, sul1, catII, tet(D), aac(3)-IId	relE/parE vapB/C	CP048380.1
CF038	E. coli (410)	NDM-5	p038_A-NDM-5	87 kb	IncFIA	<i>tet</i> (<i>D</i>), <i>sul1</i> , <i>aad</i> A5, <i>dfr</i> A32, <i>aad</i> A15, <i>dfr</i> A12, <i>bla</i> _{TEM-192} , <i>bla</i> _{TEM-118} , <i>aac</i> (6')- <i>Ib</i> - <i>cr</i> , <i>bla</i> _{OXA-140} , <i>cat</i> B3, <i>bla</i> _{CTX-M-15}	pemI/K, ccdA/B, vapC, phd/yefM hok/sok	CP048377.1
CF164	E. coli (167)	NDM-5	p164_A-NDM-5	132 kb	IncFIB	aac(6')-Ib-cr, bla _{OXA-140} , catB3, bla _{CTX-M-15} , dfrA12, aadA, sul1, brp(mbl), rmtB, bla _{TEM-1} , mphA	pemI/K, vapB/C, ccdA/B, hok,	CP048368.1
CF163	E. coli (167)	NDM-5	p163_C-NDM-5	10 kb	-	dfrA12, aadA, sul1, brp(MBL)	_	CP048374.1
CF023	R. ornithinolytica (–)	OXA-48	p023_D-OXA-48	63 kb	IncL	_	pemI/K	CP048353.1
CF053	E. coli (205)	OXA-48	p053_E-OXA-48	7.8 kb	Col156	_	_	CP048364.1
CF124	E. coli (410)	OXA-181	p124_B-OXA-181	51 kb	IncX3	qnrS1	_	CP048346.1
CF142	<i>E. coli</i> (648)	OXA-181	p142_A-OXA-181	155 kb	IncFIB	qnrS1, aadA5, dfrA32, qnrB4, dha-1, sul1, mphA, catI, tet(D)	ccdA/B, parD, pemI/K, relE/parE hok/soc	CP048338.1
CF010	E. coli (656)	OXA-181	p010_B-OXA-181	51 kb	IncX3	qnrS1	_	CP048332.1
CF061	E. coli (940)	OXA-181	p061_A-OXA-181	51 kb	IncX3	qnrS1	_	CP048327.1
CF064	E. coli (940)	OXA-181	p064_C-OXA-181	51 kb	IncX3	qnrS1	_	CP048325.1
CF032	E. coli (1284)	OXA-181	p032_K-OXA-181	51 kb	IncX3	qnrS1	_	CP048321.1
CF009	E. coli (73)	VIM-1	p009_A-VIM-1	160 kb	IncC	bla _{CMY-4} , aac(6')-1l, dfrA15, aadA12, sul1, qnrA1,	-	CP048305.1
CF035	K. aerogenes (–)	VIM-1	p035_A-VIM-1	89 kb	IncR/IncY	catB, sul1, qnrS1	relB/dinJ, vapC	CP050069.1

834 AMR, antimicrobial resistance; Inc, plasmid incompatibility; ST, sequence type determined for *E.coli*; T/A, toxin/antitoxin system; – feature not iden

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Journal Pression

Highlights

- Carbapenemase producing Enterobacteriaceae (CPE) are present in surface waters •
- Many environmental CPE are similar to clinical strains found worldwide ٠
- Clinically relevant carbapenemase genes were identified on epidemic plasmids •
- Carbapenemase genes are replicating and evolving pollutants of river ecosystems •

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Declaration of interests

 \boxtimes The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

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