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Yersiniabactin, Colibactin and Wider Resistome Contribute to Enhanced Virulence and Persistence of KPC-2-Producing Klebsiella pneumoniae CG258 in South America — Source link

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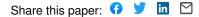
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1 Yersiniabactin, Colibactin and Wider Resistome Contribute to Enhanced Virulence and

2 Persistence of KPC-2-Producing Klebsiella pneumoniae CG258 in South America

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23

Keywords: *K. pneumoniae*; ST11; ST340; CG258; hvKp; virulence; carbapenemase, KPC-2, KPC-3,
 CTX-M; multidrug-resistance; *Galleria mellonella*; Latin America; One Health.

26

27 Abbreviations: CR-hvKp, carbapenem-resistant hypervirulent Klebsiella pneumoniae; KPC, Klebsiella 28 pneumoniae carbapenemase; CTX, cefotaximase; CG, clonal group; ST, sequence type; QACs, quaternary 29 ammonium compounds; KL, K-locus; ybt, yersiniabactin; clb, colibactin; ICEKp, integrative conjugative 30 element K. pneumoniae; pLVPK, large virulence plasmid of K. pneumoniae; CPS, capsular 31 polysaccharides; MLST, multilocus sequence typing; YbSTs, yersiniabactin sequence types; CbSTs, 32 colibactin sequence types; CR-Kp, carbapenem-resistant K. pneumoniae; MIC, minimum inhibitory 33 concentration; ESBL, extended-spectrum beta-lactamase; HM, heavy metal; ML, maximum likelihood; 34 MDR, multidrug resistance; PDR, pandrug resistance; Inc, incompatibility; IS, insertion sequence; KPZM, 35 Zn2+/Mn2+metabolism module; QRDR, quinolone-resistance determining region; PMQR, plasmid-36 mediated quinolone resistance.

37 Abstract

38 The emergence and dissemination of carbapenem-resistant hypervirulent Klebsiella 39 pneumoniae (CR-hvKp) is a worrisome public health issue compromising the treatment and 40 outcome of infections caused by this pathogen. We performed a detailed virulome and 41 resistome analysis of representative KPC- and/or CTX-M-producing K. pneumoniae 42 belonging to clonal group (CG) 258 (sequence types ST11, ST258, ST340, ST437), 43 circulating in Argentina, Brazil, Chile, Colombia and Peru; with further evaluation of the 44 virulence behavior using the Galleria mellonella infection model. Genomic analysis of K. 45 pneumoniae strains recovered from the human-animal-environment interface revealed a wide 46 resistome characterized by the presence of genes and mutations conferring resistance to 47 human and veterinary antibiotics, quaternary ammonium compounds (QACs) and heavy 48 metals. Plasmid Inc typing revealed the presence of a wide diversity of replicon types with 49 IncF, IncN, IncR and Col-like being frequently detected. Moreover, KPC-2-producing K. 50 pneumoniae belonging to ST11 (KL-64 and KL-105) and ST340 (KL-15) carried multiple 51 variants of distinct versiniabactin siderophore (vbt) and/or genotoxic colibactin (clb) genes. In 52 this regard, ICEKp3, ICEKp4 and ICEKp12 were identified in strains belonging to ST11 and 53 ST340, recovered from Argentina, Brazil, Chile and Colombia; whereas ybt 17 and a novel 54 *ybt* sequence type (YbST346) were identified together with *clb* in ICE*Kp10* structures from 55 ST11 and ST258, from Brazil and Colombia, respectively. K. pneumoniae ST11 56 (ICEKp10/YbST346 and ICEKp4/ybt 10) strains killed 100% of wax moth larvae, in a similar 57 way to hypervirulent K1/ST23 strain (vbt- and clb-negative) carrying the pLVPK-like 58 plasmid, indicating enhanced virulence. In summary, our results indicate that yersiniabactin, 59 colibactin and an expanded resistome have contributed to enhanced virulence and persistence 60 of KPC-2-producing K. pneumoniae CG258 in South America. Therefore, active surveillance 61 of hospital-associated lineages of K. pneumoniae should not only focus on clonal origin and 62 antimicrobial resistance, but also on the virulence factors ybt and clb.

63 INTRODUCTION

| 64 | Carbapenem resistance is a major public health concern worldwide, and currently Klebsiella |
|-----|---|
| 65 | pneumoniae belonging to the clonal group CG258 (which include the sequence types ST11, |
| 66 | ST258, ST340, ST437, and ST512) seem to be the main culprits for the spread of $bla_{\rm KPC}$ |
| 67 | genes (Bowers et al., 2015; Chen et al., 2014; Holt et al. 2015; Mathers et al., 2015; Paczosa |
| 68 | and Mecsas 2016; Wyres and Holt, 2016). This problem has been further exacerbated by the |
| 69 | convergence of KPC-2 production and hypervirulence, resulting in the emergence of |
| 70 | carbapenem-resistant hypervirulent K. pneumoniae (CR-hvKp) lineages, particularly in Asian |
| 71 | countries (Chen et al., 2017; Lee et al., 2017; Dong et al., 2018a; 2018b; Du et al., 2018; Gu |
| 72 | et al., 2018; Wang et al., 2018). In these countries hypervirulence has been associated with |
| 73 | the appearance and dissemination of a pLVPK-like plasmid harbouring two capsular |
| 74 | polysaccharides (CPS) upregulator genes (<i>rmpA</i> and <i>rmpA2</i>) and several siderophore gene |
| 75 | clusters (iroBCDN, iucABCD and iutA) (Struve et al., 2015; Chen et al., 2017; Du et al., |
| 76 | 2018; Gu et al., 2018). However, the acquisition of integrative conjugative elements (ICEKp) |
| 77 | harbouring yersiniabactin siderophore (ybt) is also associated with enhanced virulence, |
| 78 | whereas carriage of the genotoxic colibactin (clb) genes (in ICEKp10structures) has been |
| 79 | associated with invasive disease and colorectal cancer (Holt et al., 2015; Lam et al., 2018a). |
| 0.0 | |

80 Based on gene content variation, genomic investigation has allowed the identification 81 of 14 different structural ICEKp variants, constituting a novel target that deserves further 82 analysis for evolutionary and genomic surveillance studies (Wu et al., 2009; Lam et al., 83 2018a; Lin et al, 2008). A MLST-style approach based on diversity in eleven ybt locus genes 84 has defined yersiniabactin sequence types (YbSTs) by unique combinations of ybt gene 85 alleles and showed that YbST sequences were clustered into 17 distinct ybt lineages (Lam et 86 al., 2018a). In a similar way, variations in the *clb* locus genes have allowed definition of 87 colibactin sequence types (CbSTs), whereas phylogenetic analysis of the *clb* locus has

88 revealed three lineages that have each associated with a different ybt lineage [i.e., clb 1 (ybt

89 12), *clb* 2A (*ybt* 1) and *clb* 2B (*ybt* 17)] within the same overall structure (ICE*Kp10*).

90 Virulence in CR-Kp strains has also been associated with the type of capsular 91 polysaccharide (Cortés et al., 2002; Diago-Navarro et al., 2014; Gomez-Simmonds and 92 Uhlemann, 2017; Liu et al., 2017). In this regard, over 79 capsule (K) serotypes have been 93 described in the international K serotyping scheme (Brisse et al., 2004; Pan et al., 2015; 94 Struve et al., 2015). More recently, diversity of the capsule synthesis locus (K-locus), which 95 is 10–30 kbp in size, has been used as a novel typing method for genomic surveillance and 96 epidemiological investigations of this pathogen, and identified 134 distinct K-loci, which are 97 predictive of K serotype (Wyres et al., 2016).

98 In South American countries, KPC-2-producing K. pneumoniae has been circulating 99 in Colombia, Brazil and Argentina since at least 2005 (Villegas et al., 2006; Pavez et al., 100 2009; Gomez et al., 2011), and has more recently been reported in Ecuador, Chile, 101 Venezuela, Paraguay, Uruguay and Peru (Cifuentes et al., 2012; Zurita et al., 2013; Marquez 102 et al., 2014; Falco et al., 2016; Gomez et al., 2016; Horna et al., 2017). Although, molecular 103 epidemiology studies have confirmed predominance of the CG258 among KPC-2-producing 104 K. pneumoniae isolates collected in this region (Andrade et al., 2011; Cejas et al., 2012; 105 Pereira et al., 2013; Gomez et al., 2016; Barría-Loaiza et al., 2017; Horna et al., 2017), few 106 studies have focused on virulence determinants in these strains (Andrade et al., 2018; Araújo 107 et al., 2018). In fact, studies of biofilm formation and identification of a common set of 108 virulence genes have been restricted to KPC-2-producing K. pneumoniae from Brazil, 109 whereas sporadic identification of hypervirulent K. pneumoniae (hvKp) isolates belonging to 110 K1/ST23 and K19/ST29 have been reported in infected patients from Argentina and Brazil 111 (Cejas et al., 2014; Coutinho et al., 2014; Moura et al., 2017), and also in non-human 112 primates from Brazil (Anzai et al., 2017). In this study, using a genomic approach, we have 113 performed a detailed virulome and resistome analysis of KPC- and CTX-M-producing K.

114 pneumoniae strains belonging to CG258, recovered from the human-animal-environment 115 interface in Latin America, with further *in vivo* virulence evaluation using a *Galleria* 116 mellonella infection model.

117

118 METHODS

119 K. pneumoniae strains and genome collection

120 Laboratory studies included 19 KPC-2- and/or CTX-M-producing K. pneumoniae isolates 121 belonging to CG258 (ST11, ST258, ST340, ST437), representative of local surveillance 122 studies performed in Brazil, Peru, Chile and Argentina, between 2010 to 2016; recovered 123 from human, food-producing animals (chicken and swine) and environmental (urban rivers 124 and urban lake) samples (Oliveira et al., 2014; Martins et al., 2015; Cerdeira et al., 125 2016a;2016b; Cerdeira et al., 2017; Horna et al., 2017; Nascimento et al., 2017). K. 126 pneumoniae ATCC 13883 and hvKp K1/ST23 A58300 (Coutinho et al., 2014) were used as 127 control strains. For genome analysis, all publicly available genomes from 36 K. pneumoniae 128 CG258 strains isolated in South America were included, of which 23 were previously 129 published (Bowers et al., 2015; Araújo et al., 2018; Casella et al., 2018; Dalmolin et al., 130 2018; Pitt et al., 2018). For all 55 genomes included in this work, accession numbers are 131 listed in Table S1.

132

Antibiotic susceptibility patterns and hypermucoviscosity phenotypical identification Resistance phenotypes were determined by Kirby-Bauer method, against 30 different human and veterinary antibiotics, and the results were interpreted using the Clinical and Laboratory Standards Institute guidelines (CLSI, 2015; 2017) and The European Committee on Antimicrobial Susceptibility Testing (EUCAST, 2017). Additionally, minimum inhibitory concentrations (MICs) for ertapenem, imipenem, meropenem, enrofloxacin, ciprofloxacin,

139 levofloxacin and polymyxin B, were determined by microdilution or Etest methods (CLSI,

140 2017; EUCAST, 2017). Production of ESBL and carbapenemase enzymes was confirmed by

141 growth on CHROMagar ESBL and CHROMagar KPC, respectively.

Hypermucoviscosity phenotypes of *K. pneumoniae* isolates were determined by the string test as previously described (Moura et al., 2017). Briefly, a positive string test was defined by the formation of a viscous string > 5 mm in length when a colony was grown on a blood agar plate at 37 °C overnight and stretched by an inoculation loop.

146

147 Sequencing of *K. pneumoniae* strains

148 For 12 K. pneumoniae strains, total genomic DNA was extracted using PureLinkTM Genomic 149 DNA Mini Kit according to the Manufacturer's instructions. Library preparation was 150 performed using the Nextera XT DNA sample preparation kit (Illumina, San Diego, CA). 151 Sequencing was performed using the Illumina NextSeq platform with paired-end reads 152 (150bp). Additionally, 10 K. pneumoniae isolates selected from the 19 KPC-2 and/or CTX-M 153 producers (Table S1) were subjected to further sequencing using a PromethION R9.4.1 flow 154 cell (Oxford Nanopore Technologies). A 2D MinION library was generated from 1.5 µg 155 purified genomic DNA using the Nanopore Sequencing Kit (SQK-NSK007). DNA was 156 repaired (NEBNext FFPE RepairMix), prepared for ligation (NEBNextUltra II End-157 Repair/dA-tailing Module) and ligated with adapters (NEB Blunt/TA Ligase Master Mix).

158

159 **Bioinformatic analysis**

Forty-one *K. pneumoniae* CG258 genomes with available short-read sequence data (including genomes obtained in this study and publicly available in the GenBank) were subjected for *de novo* assemblies using Unicycler (v0.4.0) (Wick et al., 2017). For ten genomes obtained with Nanopore reads, scaffold bridging was performed, building a high-quality finished genome sequence. The contigs were annotated by Prokka v1.12 (https://github.com/tseemann/prokka).
MLSTs, YbSTs, CbSTs (Lam et al., 2018a; Diancourt et al., 2005), virulome, resistome and

166 plasmid replicon genes were screened by SRST2 (Inouye et al., 2014), using BIGSdb 167 (Bialek-Davenet et al., 2014), ARG-Annot (Gupta et al., 2014), and PlasmidFinder (Carattoli 168 et al., 2014) databases. On the other hand, since 14 of the publicly available genomes (used in 169 this study) were only available as pre-assembled sequences, Kleborate 170 (https://github.com/katholt/Kleborate) and PlasmidFinder were used to identify MLST, 171 resistome, virulome and plasmid replicon genes. Kleborate was further used to predict 172 versiniabactin ICEKp structures in all 55 genomes.

173 ResFinder 3.0 database was used to confirm resistomes (Zankari et al., 2012), and
174 while capsule and O-antigen biosynthesis loci were identified using Kaptive (Wick et al.,
175 2018), heavy metal (HM) and QAC genes were screened using BLASTN against local
176 HM/QAC and BIGSdb databases.

177 Single nucleotide variants identified using RedDog v1beta.10.3 were 178 (https://github.com/katholt/RedDog), with the reference of Κ. genome 179 pneumoniae30660/NJST258 1 (CP006923) (Bowers et al., 2015). Gubbins v.2.1.0 (Croucher 180 et al., 2014) was used to identify and exclude recombination imports. Maximum likelihood 181 (ML) trees were inferred from the recombination-masked alignment by running RaxML 182 v8.2.9 (Stamatakis et al., 2006) five times, selecting the final tree with the highest likelihood. 183 To assess branch support, we conducted 100 non-parametric bootstrap replicates using 184 RAxML.

185

186 Galleria mellonella killing assays

In order to evaluate the virulence behavior of KPC-2- and/or CTX-M-15-producing *K*. *pneumoniae* strains, *in vivo* experiments were carried out using a *Galleria mellonella* infection model (Junqueira 2012; Insua et al. 2013), with the non-virulent *K. pneumoniae* strain ATCC 13883 and the clinical hvKp K1/ST23 strain A58300 (Coutinho et al. 2014), as comparative strains. Fourteen *K. pneumoniae* strains of different lineages of CG258 and 192 origins (i.e., human, animal or environmental), circulating in Latin America, were evaluated. 193 For each experiment, a control group containing five larvae was inoculated with sterile PBS 194 in order to discard death due to physical trauma. In all experiments, groups of 250 to 350 mg G. mellonella larvae were inoculated with 10^6 CFU, and survival analysis was evaluated 195 196 during 96h (Moura et al. 2017). Survival curves were plotted using the Kaplan-Meier method, 197 and data were analyzed by the Fisher's exact test, with P < 0.001 indicating statistical 198 significance. The statistical software used was Prism7 (Graph Pad Software, San Diego, CA, 199 USA). G. mellonella larvae that did not demonstrate a response to physical stimulation and 200 had body melanization were considered dead. All experiments were performed in 201 independent triplicate assays.

202

203 RESULTS

204 Antimicrobial resistance profiles, resistome and plasmid populations

205 All 19 K. pneumoniae evaluated, in vitro, exhibited resistance to multiple antibiotics and 206 were classified as MDR or PDR phenotypes (Magiorakos et al., 2012) (Table 1). In fact, 207 resistome analysis revealed the presence of genes conferring resistance to aminoglycosides, 208 quinolones, sulphonamides, tetracycline, phenicols, fosfomycin and beta-lactam antibiotics 209 (blaoxa-1, blaoxa-2, blaoxa-9, blaoxa-10, blactx-m-15, blashy-11, blashy-12, bla_LAP-2, blatem-1A, 210 bla_{TEM-1B}, bla_{TEM-55}) (Figure 1; Table S2). Moreover, point mutations in GyrA (Ser-83-Ile), 211 GyrB (Asp-466-Glu) and ParC (Ser-80-Ile) were associated with quinolone resistance. 212 Polymyxin resistance in 5 of the 19 (26%) human and environmental isolates was associated 213 to mgrB mutations (i.e., Gly-28-Cys or Tn3 insertion at position 134) (Table S2). 214 Additionally, the presence of genes conferring resistance to silver (*sil*), copper (*pco*), arsenic 215 (ars), mercury (mer), tellurite (ter), and quaternary ammonium compounds (gacA, gacE, 216 $qacE\Delta 1$, qacL and sugE) supported a wider resistome (Figure 1, Table S2), which could 217 contribute to the apparent high versatility, persistence and adaptation of CG258 to various

218 ecosystems and hosts (Navon-Venezia et al., 2017; Dong et al., 2018). Notably, the presence

219 of tellurite resistance has also been associated with hypervirulent clonal groups of K.

220 pneumoniae (Passet and Brisse, 2015; Martin et al., 2018).

221 Additional in silico analysis of 36 genome sequences obtained from GenBank 222 confirmed the wider resistome of K. pneumoniae strains circulating in Latin America. 223 However, others CTX-M gene variants, such as bla_{CTX-M-2}, bla_{CTX-M-8}, bla_{CTX-M-9}, bla_{CTX-M-14} 224 and $bla_{CTX-M-59}$ could be identified, as well as the bla_{KPC-3} carbapenemase gene identified in 225 two Colombian K. pneumoniae strains (Colombia-2009a, Colombia-2009b) alone (Table S2). 226 Moreover, presence of *armA*, *rmtB*, *rmtD* and *rmtG*16S rRNA methyltransferases encoding 227 genes, conferring resistance to most aminoglycosides, was confirmed in 6 of the 55 (11%) 228 analyzed genomes. Worryingly, a third of isolates were predicted to be polymyxin resistant 229 based on deletions in *mgrB* and/or *pmrB* genes in 20 of the 55 analyzed genomes; whereas 230 the *mcr-1* gene was identified in a human strain isolated in Brazil (Table S2).

231 Plasmid incompatibility (Inc) typing revealed the presence of a wide diversity of 232 plasmid replicon types harbored by KPC-2- and/or CTX-M-type-producing K. pneumoniae 233 (Figure 1, Table S2). Among 55 genomes analyzed, IncF, IncN, IncR and Col-like replicons 234 were over-represented (96.4%, 54.5%, 27.3% and 27.3% respectively), whereas other Inc 235 types identified were: X3 (n=8), HI1B (n=8), L/M (n=6), U (n=5), Q1 (n=4), P6 (n=3), 236 A/C2 (n=2), I1 (n=1), I2 (n=1), HI1A (n=1) and X4 (n=1). However, bla_{KPC} plasmids 237 belonging to the IncN incompatibility group were identified in 30 of the 55 (54%) 238 carbapenem-resistant K. pneumoniae strains.

239

240 Virulome, yersiniabactin, colibactin and integrative conjugative elements (ICEs)

Among genomes analysed, lineages belonging to ST11 and ST340 carried multiple variants of distinct yersiniabactin siderophore (*ybt*) and/or genotoxic colibactin (*clb*) genes from distinct *ybt/clb* lineages and ICE*Kp* variants. In this regard, we detected *ybt* lineage 9

| 244 | (ICEKp3), ybt 10 (ICEKp4) and ybt 16 (ICEKp12) in nine ybt+ K. pneumoniae strains |
|-----|---|
| 245 | belonging to ST11 and ST340, isolated from human samples collected in Argentina, Brazil |
| 246 | and Chile (Figure 2A, Table S3); whereas ybt 17 and a novel ybt sequence type YbST346, |
| 247 | that does not belong to any of the 17 previously described ybt lineages (Lam et al, 2018a), |
| 248 | were identified together with clb lineage 2B in ICEKp10 structures, in eight Brazilian K. |
| 249 | pneumoniae ST11 strains recovered from human and environmental samples (Figure 2B). |
| 250 | ST258 lineages from Colombia harbored the classical ICEKp10 with the ybt 17 and clb 2B. |
| 251 | The alignment between the classical ICEKp10/ybt 17 and that of ICEKp10 with the |
| 252 | novel YbST346 shows that the main difference is the insertion sequence ISEc21 (IS110 |
| 253 | family) located within the Zn2 ⁺ /Mn2 ⁺ metabolism module (KPZM) (Figure 2B, Table S3). |
| 254 | Regarding other virulence determinants, the presence of the aerobactin locus (iuc) was |
| 255 | only identified in a human KPC-2-positive K. pneumoniae strain ST11 from Brazil (Figure |
| 256 | 1). |
| 257 | |
| 258 | In silico serotyping, capsule locus (KL) analysis and string test |
| 250 | Le siling construing of 55 company analyzed showed a medaminenes of 04 [K26 K15 K |

259 In silico serotyping of 55 genomes analyzed showed a predominance of O4 [K36, K15, K-260 non-typeable (NT)], O2v2 (K8, K27, K-NT) and O2v1 (K64) serotypes, which were 261 associated with ST340 (O4/K15, O4/K-NT), ST437 (O4/K36), ST11 (O2v2/K8, O2v2/27, 262 O2v2/K-NT, O2v1/K64), and ST258 (O2v2/K-NT) (Figure 1, Table S2). On the other hand, 263 we investigated the diversity of capsule synthesis loci using full locus information extracted 264 from whole genome sequences. These results show that K-loci were diverse in human and 265 environmental K. pneumoniae ST11 (i.e., KL-8, KL-27, KL-64, KL-105, KL-107, KL-127), 266 in this region (Figure 1, Table S2). Interestingly, in K. pneumoniae belonging to ST340, KL-267 15 was assigned to human and environmental strains collected in Argentina, Peru and Brazil, 268 respectively; whereas KL-151 was only identified in animal strains from Brazil. On the other 269 hand, human and environmental K. pneumoniae ST437 (from Brazil) were typed as KL-36; 270 whereas KL-106 and KL-107 accounted for strains of ST258, in Brazil and Colombia,

respectively.

KL-64/ST11 and KL-105/ST11showed a high virulence behavior in the *G. mellonella*model, the latter being identified in clinical samples from Chile and Brazil. In this regard,
KL-64 has been previously associated with strains from invasive *K. pneumoniae* infections
(Follador et al., 2016).

276 The genetic structure of the cps synthesis loci across the virulent ST11 (KL-105 and 277 KL-64) and ST340 (KL-15) was distinct from the K-loci from hvKpK1 (KL-1) and K2 (KL-278 2) (Figure 3). In this concern, for these K-loci, a conserved genetic organization at the 5' end 279 of the cps locus was observed from galF to wzc genes, whereas wzc-gnd and gnd-ugd regions 280 were variable. Moreover, while in KL-64 the gnd-ugd region is composed of genes involved 281 in GDP-D-mannose synthesis (manB and manC) and deoxythymidine diphosphate (dTDP)-L-282 rhamnose synthesis (*rmlA*, *rmlB*, *rmlC* and *rmlD*), in KL-105 the *gnd-ugd* region is only 283 composed of operon manCB (Figure 3) (Pan et al., 2015; Wyres et al., 2015; Wyres et al., 284 2016). Interestingly, dTDP-L-rhamnose is the precursor of L-rhamnose, a saccharide required 285 for the virulence of some pathogenic bacteria, being essential for resistance to serum killing 286 and for colonization (Giraud et al., 2000).

To investigate the hypermucoviscosity phenotype, all the isolates were subjected to the string test. Among the 19 KPC-2 and/or CTX-M-15 producers, only one CTX-M-15producing *K. pneumoniae* ST340/KL-151 strain (FA64), isolated from a healthy chicken sample, in Brazil, showed hypermucoviscosity. However, neither of the known hypermucoviscosity encoding genes (*rmpA* or *rmpA2*) were detected in its genome.

292

293 In vivo virulence behavior of K. pneumoniae CG258

Using the G. mellonella virulence model, ST11 CR-KP strains (n=2, KL-64/ybt+/clb+; n=2,

295 KL-105/ybt+) killed 100% of wax moth larvae inoculated with 1×10^{6} colony-forming units of

| 296 | the bacterial specimens, within 96 h, in a similar way to the known hypermucoviscous hvKp |
|-----|---|
| 297 | K1/ST23 strain which is ybt- and clb-negative and carries the pLVPK-like plasmid (P> |
| 298 | 0.9999) (Figure 4A). KPC-2- and/or CTX-M-15-producing K. pneumoniae strains belonging |
| 299 | to ST340 killed >60% of wax moth larvae (n=2, KL-15/ybt+; n=3, KL-15/ybt-; n=1, KL- |
| 300 | 151/ybt-). One ST340 KL-15/ybt+ strain isolated from a human infection killed 100% of G. |
| 301 | <i>mellonella</i> (Figure 4B). <i>K. pneumoniae</i> belonging to ST437 (<i>n</i> = 3 strains, all KL-36/ <i>ybt</i> -/ <i>clb</i> -) |
| 302 | killed ~50% of G. mellonella, compared to K. pneumoniae ATCC 13883 control (Figure 4D). |
| 303 | Overall, among 14 K. pneumoniae strains evaluated, 4/6 (83%) ybt+ isolates (4/4 |
| 304 | ST11, 1/2 ST340) killed all wax moth larvae within 96 h, compared to only 1/8 (13%) ybt- |
| 305 | strains (1/1 ST258, 0/4 ST340, 0/3 ST437) ($P = 0.03$, Fisher's exact test), suggesting that in |
| 306 | the absence of pLVPK-like plasmids, the presence of <i>ybt</i> could be enough to confer enhanced |
| 307 | virulence. However, the single ST258 strain (ybt- and clb-negative), isolated from a human |
| 308 | clinical sample also killed 100% of G. mellonella within 96 h (Figure 4C), suggesting that |
| 309 | other factors, not elucidated in this study, may also be contributing to the virulence phenotype |
| 310 | of CG258 (Araújo et al., 2018; Hennequin and Robin, 2016; Shah et al., 2017; Fu et al, 2018; |
| 311 | Marcoleta et al., 2018; Zheng et al., 2018). |

312

313 **DISCUSSION**

314 In South American countries, antimicrobial resistance has long been documented to be more 315 challenging than in developed ones (Gales et al., 2012; Sampaio and Gales, 2016). In this 316 regard, the high prevalence of carbapenem resistance in this region has occurred primarily by 317 the dissemination of KPC-producing K. pneumoniae isolates belonging to CG258, which 318 have been identified beyond the hospital setting, constituting a One Health problem (Andrade 319 et al., 2011; Gomez et al., 2011; Cejas et al., 2012; Oliveira et al., 2014; Barría-Loaiza et al., 320 2016; Rojas et al., 2017; Horna et al., 2017; Nascimento et al., 2017). In this study, we 321 performed a resistome and virulome analysis of KPC-and/or CTX-M-producing K.

pneumoniae lineages belonging to CG258, circulating in hospital settings. Additionally, environmental and animal *K. pneumoniae* isolates, recovered in Brazil, were also investigated.

325 Genome analysis revealed a wider resistome, which includes genetic determinants 326 conferring resistance to human and animal antibiotics, QACs and HMs, supporting 327 persistence and adaptation of CG258 to different hosts and anthropogenically affected 328 environments. Among MDR and PDR lineages, the presence of mutations in mgrB/pmrB 329 genes and in the quinolone resistance-determining region; as well as acquisition of 16s rRNA 330 methylases- and β -lactamases-encoding genes (includying bla_{ESBL} and bla_{KPC-2}) have 331 contributed with resistance to polymyxins, fluoroquinolones, aminoglycosides and broad-332 spectrum β -lactam antibiotics. Moreover, we have identified, for the first time, the presence 333 of the narrow-spectrum β -lactamase encoding gene bla_{LAP-2} (GenBank accession number 334 EU159120) and $bla_{\text{TEM-55}}$ ESBL gene(GenBank accession number DQ286729) in K. 335 pneumoniae strains ST340 recovered from swine and human hosts, respectively, in Brazil, 336 confirming versatility of this lineage to acquire novel genetic determinants of resistance.

337 We have identified regional $bla_{\rm KPC}$ spread consistent with high prevalence of IncN 338 plasmids, previously associated with the global spread of these genes (Stoesser et al., 2017). 339 On the other hand, the wide diversity of Inc-type plasmids, found in this study, including 340 small mobilizable Col-like replicons could be associated with the acquisition of multiple 341 resistance mechanisms, contributing to the wider resistome. Therefore, the presence of K. 342 pneumoniae in a wide range of environmental reservoirs and hosts, with plasmids that have 343 been shown to facilitate the dissemination of successful resistance genes, even in the absence 344 of selection pressures, may represent a difficult situation to control (Stoesser et al., 2017). 345 Another important issue is the identification of IncHI1-type plasmids, which have been 346 associated with the dissemination of mcr-1 and bla_{CTX-M} -type genes in Colombia and 347 Uruguay, respectively (Saavedra et al., 2017; Garcia-Fulgueiras et al., 2017).

348 Hypervirulent K. pneumoniae strains have been sporadically reported in Argentina 349 and Brazil, being associated with remarkable mortality and the production of a 350 hypermucoviscous phenotype in lineages belonging to ST23 with capsular serotype K1, and 351 ST29/K19 (Cejas et al., 2014; Coutinho et al., 2014, Moura et al., 2017). In this study, 352 virulome analysis revealed that *ybt* and *clb* genes have been acquired by strains of CG258 in 353 South America, highlighting the need to also consider these additional virulence factors 354 rather than the presence of pLVPK-like plasmids and hypermucoviscous phenotypes alone, in 355 the establishment of hypervirulence in carbapenem-resistant lineages of K. pneumoniae.

Notably, there have been increasing reports of highly virulent *K. pneumoniae* strains carrying *ybt* belonging to the international clone ST11. The emergence of CR-hvKp strains carrying *ybt* plus a deletion variant of the pLVPK-like plasmid belonging to ST11 has been associated with outbreak of fatal nosocomial infection in China (Gu et al., 2018), raising an epidemiological alert in response to the increased number of cases reported, in the last year (Lee et al., 2017; Zhan et al., 2017; Chen and Kreiswirth, 2018; Du et al., 2018; Wong et al.,

362 2018; Yao et al., 2018).

363 In summary, these results (available for interactive exploration in Microreact at 364 https://microreact.org/project/H1LSZsRz7) confirm the enhanced virulence of KPC-2- and/or 365 CTX-M-producing K. pneumoniae belonging to the international high-risk clone CG258 in 366 South America, where acquisition of ICEKp encoding versiniabactin and colibactin, and 367 wider resistome have likely contributed to enhanced virulence and persistence of ST11 (KL-368 64 and KL-105) and ST340 (KL-15) lineages, in the human-environment interface. While 369 capsule composition deserves further investigation, active surveillance should not only focus 370 on clonal origin, antimicrobial resistance and presence of pLVPK-like plasmids, but also the 371 virulence associated with versiniabactin and colibactin, as well as other biomarkers for 372 differentiation of hvKp from classical K. pneumoniae (Russo et al., 2018); and control 373 measures should be conducted to prevent the global dissemination of these lineages.

374 CONCLUSION

375 Our study points out several important issues. Firstly, interplay of yersiniabactin and/or 376 colibactin and KPC-2 production has become to be identified among K. pneumoniae 377 belonging to CG258, in South America, contributing to the emergence of highly virulent 378 lineages that pose great risk to human health (Lam et al. 2018a). Second, in South America 379 ICEKp3, ICEKp4 and ICEKp10carrying vbt and/or clb circulate among KPC-2-producingK. 380 Pneumoniae belonging to ST11 (KL-64 andKL-105), where multiple distinct K-loci often 381 indicates distinct sublineages that may correlate with independent ICEKp acquisitions 382 (supported by our phylogenetic analysis shown in Figure 1); being associated with enhanced 383 virulence, these should be considered a target for genomic surveillance along with 384 antimicrobial resistance determinants. Third, the wide resistome could be contributing to 385 adaptation of KPC-2- and/or CTX-M-producing K. pneumoniae CG258 in the human-animal-386 environment interface, highlighting the urgent need for enhanced control efforts. Finally, 387 these findings could contribute to the development of strategies for prevention, diagnosis and 388 treatment of K. pneumoniae infections.

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| | MICs (J | ug/ml) | | | | | | |
|--------|---------|--------|-----|------|-----|-----|-----|--|
| Strain | POL | ETP | IMP | MER | ENO | CIP | LVX | Kirby-Bauer |
| KP171 | 2 | 8 | 16 | 8 | 8 | 8 | 32 | CTX, CAZ, CPM, CRO, NAL, GEN, SUT, ATM, TET, CLO, FOS |
| 1194 | 2 | 4 | 8 | 8 | 8 | 8 | 32 | CTX, CAZ, CPM, CRO, NAL, AMI, GEN, SUT, ATM, TET, CLO, FOS |
| 606B | 2 | 1 | 1 | 0.25 | 4 | 4 | 16 | CTX, CAZ, CPM, CRO, NAL, GEN, SUT, ATM, TET, CLO, FOS |
| KPN535 | 16 | 4 | 8 | 4 | 8 | 8 | 32 | CTX, CAZ, CPM, CRO, NAL, GEN, SUT, ATM, CLO, FOS |
| KPC45 | 32 | 4 | 8 | 4 | 4 | 4 | 16 | CTX, CAZ, CPM, CRO, NAL, SUT, ATM |
| IBL2.4 | 8 | 4 | 8 | 8 | 4 | 8 | 32 | CTX, CAZ, CPM, CRO, CFO, NAL, ATM, TET |
| KP488 | >32 | 4 | 8 | 8 | 4 | 4 | 32 | CTX, CAZ, CPM, CRO, CFO, NAL, GEN, SUT, ATM, CLO, FOS |
| 196 | 0.25 | >32 | 8 | >32 | 8 | >32 | >32 | CTX, CAZ, CPM, CRO, CFO, NAL, SUT, ATM |
| 148 | 32 | >32 | >32 | >32 | 4 | >32 | >32 | CTX, CAZ, CPM, CRO, CFO, NAL, SUT, ATM, TET, CLO, FOS |
| 314 | 2 | >32 | >32 | >32 | 4 | >32 | >32 | CTX, CAZ, CPM, CRO, CFO, NAL, SUT, ATM, TET |
| KP411 | 0,5 | >32 | >32 | >32 | 8 | 8 | 8 | CTX, CAZ, CPM, CRO, CFO, NAL, SUT, ATM, TET, CLO |
| KP337 | 0,5 | >32 | >32 | >32 | 4 | 4 | 8 | CTX, CAZ, CPM, CRO, CFO, NAL, ATM, TET, CLO |
| KP326 | 0,5 | >32 | 8 | 8 | 8 | 8 | 16 | CTX, CAZ, CPM, CRO, NAL, SUT, ATM, TET, CLO, GEN |
| KP515 | 0,5 | >32 | >32 | >32 | 8 | 8 | 16 | CTX, CAZ, CPM, CRO, CFO, NAL, SUT, ATM, TET, CLO, FOS |
| KP870 | 0,5 | >32 | >32 | >32 | 16 | 8 | 16 | CTX, CAZ, CPM, CRO, CFO, NAL, SUT, ATM, TET, CLO, FOS |
| FA64 | 1 | 1 | 0.5 | 0.5 | 4 | 8 | 4 | CTX, CAZ, CPM, CRO, NAL, SUT, ATM |
| 2KP | 1 | 8 | 8 | 8 | 8 | 8 | 8 | CTX, CAZ, CPM, CRO, CFO, NAL, SUT, ATM, TET |
| 1ECKPC | 1 | 4 | 8 | 8 | 8 | 8 | 8 | CTX, CAZ, CPM, CRO, CFO, NAL, SUT, ATM, TET, CLO, GEN |
| N7 | 1 | 8 | 8 | 8 | 4 | 8 | 8 | CTX, CAZ, CPM, CRO, NAL, SUT, ATM, TET, CLO, GEN, AMI |

Table 1. Resistance profile of KPC-2- and/or CTX-M-producing K. pneumoniae strains belonging to CG258 circulating in South America*

*POL – polymyxin B, ETP – ertapenem, IMP – imipenem, MER- meropenem, ENO – enrofloxacin, CIP – ciprofloxacin, LVX – levofloxacin, CTX – cefotaxime, CAZ – ceftazidime, CPM – cefepime, CRO – ceftriaxone, CFO – cefoxitin, NAL – nalidixic acid, AMI - amikacin, GEN – gentamicin, SUT – sulfamethoxazole/trimethoprim, ATM - aztreonam, TET – Tetracycline, CLO – chloramphenicol, FOS – fosfomycin.

Figure legends

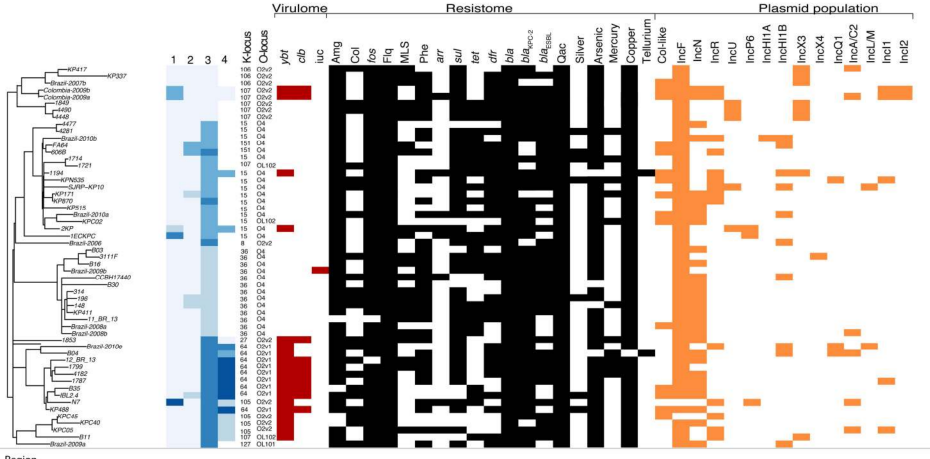
Fig. 1. Virulome, resistome and plasmid population of KPC- and/or CTX-M-producing *K*. *pneumoniae* belonging to CG258 in Latin America. Tracks indicate: (1) country, (2) strain source, (3) sequence type (ST), (4) ICE*Kp* structure. The red/black/orange regions represent the presence of the gene, and blank regions represent their absence. Amg, aminoglycosides resistance genes (i.e., transferases and 16S rRNA methylases); Col, polymyxin resistance genes (including *mgrB/pmrB* mutations and *mcr-1*); Flq, fluoroquinolone resistance genes (i.e., QRDR mutations and PMQR); MLS, macrolides resistance genes (*mphA*, *erm*); Phe, phenicols resistance genes (*cat*, *cml*, *flor*) (Table S2).

Fig. 2. Comparative analysis of integrative and conjugative elements ICEKp3, ICEKp4, ICEKp10, and ICEKp12. In A, ICEKp mobilizing versiniabactin identified in K. pneumoniae CG258 in South America. In B, alignment of ICEKp10/vbt 17 against ICEKp10 carrying a novel yersiniabactin sequence YbST346, identified in lineages belonging to ST11, isolated in Brazil. Blue blocks represent versiniabactin synthesis locus ybt, labelled with the associated *ybt* lineage. Orange represents the mobilization module. Light orange represents KPZM (Zn²⁺/Mn²⁺) metabolism module. Light blue shading denotes shared regions of homology (>95%), where the main difference between the two ICEKp10 is the presence of a novel ybt lineage, and the insertion sequence ISEc21 (IS110 family) located inside the Zn^{2+}/Mn^{2+} metabolism module (KPZM). The key differences between ICEKp3, ICEKp4 and ICEKp12 have been previously defined and are restricted to a single variable region, where ICEKp3 was constituted by restriction endonuclease, DUF4917 domain containing protein, ATP/GTP phosphatase, reverse transcriptase, DDE endonuclease, and five hypothetical proteins; whereas ICEKp4was formed by transposase, ABC transporter, type I restriction endonuclease, DNA methyltransferase and hypothetical protein; and ICEKp12 contained an additional Zn²⁺/Mn²⁺metabolism module (KPZM) (Marcoleta et al. 2016; Lam et al., 2018a).

Fig. 3. K-loci (KL-1, K-L2, KL-105, KL-64 and KL-15) structures of CR-KP lineages belonging to CG258. In *K. pneumoniae*, K-locus includes a set of genes in the terminal regions encoding for the core capsule biosynthesis machinery (i.e., *galF*, *wzi*, *wza*, *wzb*, *wzc*, *gnd* and *ugd*). The central region is highly variable, encoding for specific sugar synthesis of the capsule, processing and export proteins, plus the core assembly components Wzx (flippase) and Wzy (capsule repeat unit polymerase) (Pan et al., 2015; Wyres et al., 2016). Protein coding sequences are represented as arrows colored by predicted function of the protein product and labelled with gene names where known.

Fig. 4. *In vivo* virulence behavior of CR-KP belonging to ST11, ST340, ST437 and ST258 in a *Galleria mellonella* infection model. The virulence behavior of 1×10^6 colony-forming units of representative *K. pneumoniae* strains on *G. mellonella* survival was assessed using both, non-virulent (ATCC 13883, ST not determined in this study) and hypervirulent (A58300 K1/ST23) *K. pneumoniae* control strains. In A, *K. pneumoniae* KPC45, KP488, IBL2.4 and N7 belonging to ST11, recovered from human and environmental samples. In B, *K. pneumoniae* 1194, KP870, 1ECKPC, 2KP, FA64, KP171 strains belonging to ST340, recovered from human, animal and environmental samples. In C, *K. pneumoniae* KP337 strain belonging to ST258, recovered from a clinical sample. In D, *K. pneumoniae* KP196, KP411 and 314 strains belonging to ST437, recovered from human and environmental samples. Clinical and epidemiological characteristics of *K. pneumoniae* strains are quoted in Table S1. **P*>0.9999, indicates no statistically significant difference with respect to the hypervirulent A58300 K1/ST23 *K. pneumoniae* control strain.





Region

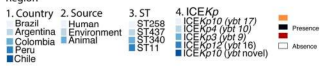
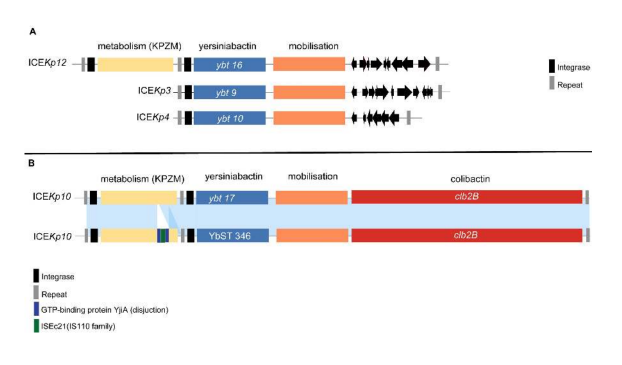
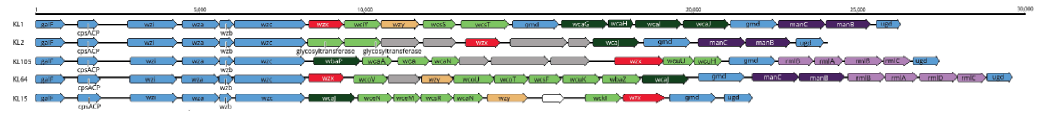


Figure 2







Common proteins including core assembly machinery Other sugar synthesis and processing WbaP/WcaJ initiating glycosyltransferase

Wzy capsule repeat unit polymerase

📕 Wzx flippase

Mannose synthesis/processing proteins

Rhamnose synthesis/processing proteins

Hypothetical protein

. providely a

Figure 4

