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| 1 | An IncP plasmid carrying the colistin resistance gene <i>mcr-1</i> |
|----|---|
| 2 | in Klebsiella pneumoniae from hospital sewage |
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| 19 | |

20 Abstract

A *Klebsiella pneumoniae* strain of ST313 recovered from hospital sewage was found to carry the plasmid-borne colistin resistance gene *mcr-1*, which was bracketed by two copies of the insertion sequence IS*Apl1* on a 57-kb self-transmissible IncP type plasmid of a new IncP-1 clade. The carriage of *mcr-1* on a self-transmissible broad-host-range plasmid highlights that *mcr-1* has the potential to be spread beyond the *Enterobacteriaceae*.

27 Colistin is the last resort antimicrobial agents to treat the infections caused by many 28 Gram-negative bacteria. Recently, a plasmid-borne colistin resistance gene, mcr-1, has 29 been found in Escherichia coli and Klebsiella pneumonia from human and animals in 30 China (1). A few follow-up studies have found mcr-1-carrying E. coli in many 31 countries in Africa (2), Europe (3-8), Asia (1-3, 9, 10) and North America (9, 11). In 32 addition, mcr-1 has also been found in Kluyvera ascorbata (12) and several species of 33 Salmonella (13-15). All of above findings suggest that mcr-1 has been widely distributed and imposes an emerging threat for clinical management and public and 34 35 animal health. During a screening study for the presence of colistin-resistant Enterobacteriaceae in hospital sewage, we found a mcr-1-carrying K. pneumonia, 36 which is reported here. 37

38

39 K. pneumoniae strain WCHKP1511 was recovered from the influx mainstream of 40 hospital sewage at West China Hospital, Chengdu, western China, in November 2015. 41 Strain WCHKP1511 grew on CHROMAgar Orientation agar plates (CHROMAgar, 42 Paris, France) containing 4 µg/ml colistin and 64 µg/ml linezolid. Species 43 identification was established by partially sequencing the gyrB gene (16). Strain WCHKP1511 was resistant to colistin (MIC, 8 µg/ml), polymyxin B (MIC, 8 µg/ml), 44 45 chloramphenicol (MIC, 128 µg/ml) and tetracycline (MIC, 64 µg/ml) but was 46 susceptible to amikacin (MIC, 0.5 μ g/ml), ceftazidime (MIC, $\leq 0.5 \mu$ g/ml), cefotaxime (MIC, 0.03 µg/ml), ciprofloxacin (MIC, 0.03 µg/ml), imipenem (MIC, 0.125 µg/ml) 47

| 48 | and tigecycline (MIC, 1 μ g/ml) as determined using the microdilution broth method |
|----|--|
| 49 | following recommendations of the Clinical Laboratory Standards Institute (CLSI) |
| 50 | (17). WCHKP1511 was susceptible to fosfomycin (MIC, 32 μ g/ml) as determined |
| 51 | using the agar dilution method following recommendations of CLSI (17). In addition, |
| 52 | strain WCHKP1511 was resistant to ampicillin and trimethoprim-sulphamethoxazole, |
| 53 | intermediate to amoxicillin-clavuanate, gentamicin, tobramycin and nitrofurantoin, |
| 54 | and susceptible to aztreonam, cefazolin, cefepime, cefoxitin, ceftriaxone, ertapenem, |
| 55 | levofloxacin and piperacillin-tazobactam as determined using the Vitek II automated |
| 56 | system (bioMerieux, Lyon, France). Breakpoints defined by FDA and by EUCAST |
| 57 | were used for tigecycline and colistin, respectively; otherwise, those defined by CLSI |
| 58 | were applied. Strain WCHKP1511 was found to carry mcr-1 by PCR and sequencing |
| 59 | (1, 12). Although $mcr-1$ has been widely found in E. coli, $mcr-1$ -carrying K. |
| 60 | pneumoniae remains uncommon and had only been found in eastern China (Jiangsu |
| 61 | and Zhejiang provinces) before (1, 18). |

Strain WCHKP1511 was subjected to the 150-bp paired-end whole genome
sequencing with a ca. 200× coverage using the HiSeq 2500 Sequencer (Illumina, San
Diego, CA, USA). A total of 3,784,972 reads and 567,745,800 clean bases were
generated, which were assembled into 275 contigs (230 contigs ≥ 1,000 bp) in length
(N50, 67,217 bp) with a 57.46% GC content using the Spades program (19).

| 69 | WCHKP1511 belonged to ST313, which was determined by using the genomic |
|----|--|
| 70 | sequence to query the multi-locus sequence typing database of K. pneumoniae |
| 71 | (http://bigsdb.web.pasteur.fr/klebsiella/klebsiella.html). In the K. pneumoniae MLST |
| 72 | database, only one ST313 strain, KML 2185, which was recovered from human blood |
| 73 | in the Netherlands in 2007, has been deposited. K. pneumoniae strains carrying mcr-1 |
| 74 | in Jiangsu province belonged to ST25 (20), while the STs of those in Zhejiang |
| 75 | province were unknown. ST313 was not closely related to ST25, as only 2 out of 7 |
| 76 | alleles were identical between the two STs. |

77

78 Antimicrobial resistance genes were predicted using ResFinder from the Center for 79 Genomic Epidemiology (http://genomicepidemiology.org/). In addition to mcr-1, 80 WCHKP1511 had multiple genes mediating resistance to aminoglycosides 81 (aac(3)-Iva, aadA2, aph(3')-Ia and aph(4)-I), β -lactams $(bla_{\text{TEM-135}} \text{ and } a \text{ new } bla_{\text{SHV}})$ 82 variant), chloramphenicol (floR), fosfomycin (fosA), quinolones (oqxA and oqxB), 83 sulphonamides (sul2), trimethoprim (dfrA12) and tetracycline (tet(A)) as predicted by 84 ResFinder. $bla_{\text{TEM-135}}$ encodes a broad-spectrum β -lactamase (21). Of note, although 85 strain WCHKP1511 carried fosA, it was susceptible to fosfomycin, which warrants 86 further investigation. *bla*_{SHV} in strain WCHKP1511 is a new variant, which encodes a 87 SHV enzyme with an amino acid difference (Thr14Asn, the position is based on the 88 ATG start condon) from SHV-111, the closest match. The new SHV has been assigned 89 SHV-195 NCBI classification by the β-lactamase system

90 (www.ncbi.nlm.nih.gov/pathogens/submit_beta_lactamase/). As strain WCHKP1511 was susceptible to third generation cephalosporins, $bla_{SHV-195}$ is unlikely to encode an 91 92 extended-spectrum β -lactamase (ESBL). The complete coding sequence of $bla_{SHV-195}$ 93 was cloned onto the pBC SK vector (Agilent, Santa Clara, CA, USA), which was electroporated into E. coli DH5a. E. coli transformants containing bla_{SHV-195} were 94 95 resistant to ampicillin (MIC, >256 µg/mL) and cephalothin (32 µg/mL) but were susceptible to aztreonam, ceftazidime, cefotaxime, cefoxitin and imipenem 96 97 determined using the broth microdilution method (17). This confirmed that $bla_{SHV-195}$ 98 encodes a broad-spectrum rather than an ESBL.

99

In addition, there was a predicted bleomycin-resistance gene, designated ORFble here, which was not identified by Resfinder but was identified by the Prokka annotation tool (22) and was confirmed by Protein-BLAST in strain WCHKP1511. The complete coding sequence of ORFble was amplified with primers 1511_ble_BamHI_F (CGC<u>GGATCC</u>TTGGTTCACCATGAAGATG)/1511_ble_EcoRI_R

105 (CC<u>GGAATT</u>CCGGCCGATTGCTGAACAGATTA), was cloned onto pBC SK and 106 was electroporated into *E. coli* DH5 α . ORFble-containing transformants were selected 107 on LB agar plates containing 25 µg/mL chloramphenicol and the presence of ORFble 108 in transformants was confirmed by PCR and sequencing. However, MIC (0.25 µg/mL) 109 of zeocin (Thermo Fisher Scientific, Waltham, MA, USA), a bleomycin, against *E.* 110 *coli* DH5 α transformant containing ORFble was the same as that against DH5 α as

- 111 determined using the broth microdilution method (17). This suggests that ORFble did
- 112 not mediate resistance to bleomycin and its function remains undetermined.
- 113

| 114 | Conjugation experiments were carried out in broth using azide-resistant E. coli strain |
|-----|--|
| 115 | J53 as the recipient and 2 $\mu\text{g/ml}$ colistin plus 150 $\mu\text{g/ml}$ sodium azide was used for |
| 116 | selecting transconjugants. The presence of mcr-1 in transconjugants was confirmed |
| 117 | using PCR. In strain WCHKP1511, mcr-1 could be transferred to E. coli J53 at a |
| 118 | frequency of 10^{-2} cells per donor cell by mating, suggesting that <i>mcr-1</i> was carried by |
| 119 | a self-transmissible plasmid, which was assigned pMCR_1511. The sequence of |
| 120 | pMCR_1511 was completely circularised with gaps between contigs closed by Sanger |
| 121 | sequencing of amplicons from PCRs using primers designed based on available contig |
| 122 | sequences. pMCR_KP1511 was 57,278 bp in size and had no known antimicrobial |
| 123 | resistance genes other than mcr-1. Unlikely the previously-described mcr-1-carrying |
| 124 | IncI2 plasmid pHNSHP45 (GenBank accession number KP347127) (1), |
| 125 | pMCR_KP1511 belonged to the IncP type, a broad-host-range incompatibility group. |
| 126 | Plasmid pKH-457-3-BE carrying mcr-1 in E. coli from Belgium was found to have an |
| 127 | IncP backbone (3). However, the sequence of pKH-457-3-BE was not available for |
| 128 | further analysis. Nonetheless, it has been suggested that pKH-457-3-BE was 99% |
| 129 | similarity and 73% coverage with the IncHI2 plasmid pHXY0908 (GenBank |
| 130 | accession number KM877269) in Salmonella enterica serotype Typhimurium (8). |
| 131 | pMCR_KP1511 had only a 6% coverage with pHXY0908, indicating that |

pMCR_1511 was very different from pKH-457-3-BE and pKH-457-3-BE may not bea true IncP plasmid but is likely of IncHI2.

134

135 pMCR_1511 has the typical IncP-1 plasmid backbone (23) containing the trfA encoding the replication initiation protein, two par modules for plasmid partitioning, 136 137 two conjugative regions tra (17.6 kb) and trb (12.7 kb), the host-lethal protein-encoding kil genes and their regulator kor (stands for kil-override) genes, and a 138 139 toxin-antitoxin *higA-B* system (Figure 1). The backbone of pMCR 1511 was highly 140 similar (99% identity) with that on plasmid pHNFP671 (GenBank accession number 141 KP324830), which was an IncP plasmid in E. coli isolate FP671 from Guangzhou, 142 China but did not carry *mcr-1*. IncP-1 plasmid has six assigned clades, i.e., α , β , γ , δ , ε 143 and ζ , among which β clade has β 1 and β 2 two subclades (24). To determine of which 144 clade pMCR_1511 was, the sequence of 30 genes belonging to the IncP-1 backbone 145 was retrieved from pMCR 1511, concatenated and then aligned to the counterparts of 146 one representative plasmid of each clade including $\beta 1$ and $\beta 2$ subclades as described 147 previously (25). Phylogenetic analysis of the IncP-1 plasmid backbone revealed that 148 pMCR_1511 and pHNFP671 belonged to a new IncP-1 clade (Figure S1 in the 149 supplementary file).

150

151 When compared the sequence of pHNFP671, pMCR_1511 has two unique regions
152 (Figure 1). One of the unique region harbored *mcr-1* and the other contained ORFble.

| 153 | The sequence comparison of pMCR_1511 and pHNFP671 allowed us to analyze the |
|-----|---|
| 154 | genetic context of $mcr-1$ in detail. Like most genetic contexts available in the |
| 155 | GenBank, mcr-1 was located downstream of the insertion sequence ISApl1 on |
| 156 | pMCR_1511. However, there was another ISApl1, which was interrupted (see below), |
| 157 | downstream and therefore mcr-1 was bracketed by two copies of ISApl1 on |
| 158 | pMCR_1511. It has been known that ISApl1 is able to generate 2-bp direct target |
| 159 | repeats (DR) upon insertion (26) |
| 160 | (https://www-is.biotoul.fr/index.html?is_special_name=ISApl1). The 2-bp flanking |
| 161 | sequences of the region bracketed by the two copies of ISApl1 were identical (AC, |
| 162 | Figure 2). When the region formed by two copies of ISApl1 and one of the 2-bp |
| 163 | flanking sequences were subtracted artificially, the joined sequence perfectly matched |
| 164 | that of an open reading frame (orf) with unknown function on plasmid pHNFP671. It |
| 165 | therefore proved that the 2-bp sequence was truly DR generated by ISApl1 rather than |
| 166 | coincidence and the two ISApl1 formed a composite transposon to mobilize mcr-1 |
| 167 | gene. A very recent analysis revealed that the ISApl1-formed composite transposon |
| 168 | carrying mcr-1 has also been seen on either the chromosome or a plasmid (IncH or |
| 169 | unknown Inc groups) of seven E. coli strains (27). The ISApl1-formed composite |
| 170 | transposon carrying mcr-1 is all located at different locations, which are also different |
| 171 | from the location on pMCR_1511, in the seven E. coli strains and is flanked by 2-bp |
| 172 | DR in five strains (27). The previous analysis (27) and the findings in the present |

study suggest that the IS*Apl1*-formed composite transposon is a common vehicle formediating the spread of *mcr-1*.

175

176 The ISApl1 downstream of mcr-1 on pMCR_1511 was interrupted by the insertion of 177 Tn3 with the characteristic 5-bp DR (Figure 2). The Tn3 was also disrupted by IS26 178 and most of the Tn3 was absent, which may be due to the action of IS26. It is well 179 known that the insertion of IS26 can lead to the deletion of the adjacent sequence of 180 the insertion site (28). Alternative explanation for the absence of most part of Tn3 is 181 that the insertion of the second IS26 and the recombination between the two IS26 182 could lead to the loss of the intervening region. Although ISApl1 downstream of mcr-1 was interrupted by Tn3, the right-end inverted repeat (IRR) of the ISApl1 183 184 remained intact (Figure 2). The transposase encoded by the ISApl1 upstream of mcr-1 185 had the potential to recognize the IRR of the ISApl1 downstream of mcr-1 and then 186 could realize the mobilization of the region bracketed by the two copies of the ISApl1. 187

In conclusion, the plasmid-borne colistin resistance gene *mcr-1* was found in a *K*. *pneumoniae* of an infrequently encountered ST from hospital sewage. *mcr-1* was carried by a self-transmissible IncP plasmid, which is a broad-host-range type of plasmids and has the potential to mediate the dissemination of *mcr-1* from the *Enterobacteriaceae* to other Gram-negative bacteria such as *Pseudomonas aeruginosa*. *mcr-1* was bracketed by two copies of IS*Apl1*, which were able to form a composite transposon and represented a common mechanism for mediating the mobilization of*mcr-1*.

196

| 197 | Nucleotide sequence accession number. Reads and the Whole Genome Shotgun |
|-----|---|
| 198 | Sequencing project of K. pneumoniae strain WCHKP1511 have been deposited into |
| 199 | DDBJ/EMBL/GenBank under accession number SRR3170679 and LSMF00000000, |
| 200 | respectively. The sequence of pMCR_1511 has been deposited into |
| 201 | DDBJ/EMBL/GenBank under accession number KX377410. |
| 202 | |

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305
306

307 Figure legends

Figure 1. Genetic structure of IncP plasmid pMCR_1511 carrying mcr-1 and the 308 comparison with IncP plasmid pHNFP671. Regions and genes that are indicated are 309 310 ORFble (a predicted [but actually not] bleomycin resistance gene), higA-B (encoding 311 a toxin/antitoxin system), mcr-1, kor-par-kli (for plasmid maintenance), tra and trb 312 (the two conjugation-encoding regions), and *trfA* (encoding the plasmid replication initiation protein). Backbones of the two IncP plasmids are almost identical. 313 314 Compared to pHNFP671, pMCR 1511 carried two additional regions that contained 315 either mcr-1 or ORFble.

316

Figure 2. Genetic context of *mcr-1* on pMCR_1511. Genetic context of *mcr-1* on the 317 318 IncI2 plasmid pHNSHP45 (GenBank accession number KP347127) and the corresponding region on the IncP plasmid pHNFP671 (GenBank accession number 319 320 KP324830) are shown for comparison. orfs that encode hypothetical proteins with 321 unknown function are indicated in white except that the one disrupted by the 322 ISApl1-formed composite transposon on pMCR 1511 is shown in black ($\Delta 1$ and $\Delta 2$). 323 Other genes shown are nikB (encoding relaxase of the plasmid), ydgA (DNA 324 topoisomerase III), ydfA (transcriptional regulator), parA (resolvase), bla_{TEM} (shown 325 as a white arrow in Tn3) and *traB* (conjugative protein). The 2-bp direct repeat (GA) 326 abutting the ISApl1-mcr-1-pho region on pHNSHP45 and the 2-bp direct repeat (AC) 327 abutting the ISApl1-formed composite transposon on pMCR 1511 are shown. On

- 328 pMCR_1511, the ISApl1 downstream of mcr-1 was interrupted by Tn3, which was
- 329 interrupted by IS26.