

### **Partition locus-based classification of selected plasmids in Klebsiella pneumoniae, Escherichia coli and Salmonella enterica spp.: An additional tool**

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#### **REVISED**

**Partition locus-based classification of selected plasmids in** *Klebsiella pneumoniae***,**  *Escherichia coli* **and** *Salmonella enterica spp***: an additional tool.**

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#### **1. Introduction**

Plasmids are extrachromosomal DNA molecules capable of autonomous replication and are the main vectors of resistance and virulence genes, especially in Enterobacteriaceae. Tracing plasmids conferring drug resistance is important for analysis of evolution, epidemiology and spread of antibacterial resistance. The epidemic plasmids belong to the most frequently occurring plasmid families, however antibiotic resistance gene are not always associated with one particular replicon, transfer or partition system. Therefore is difficult to construct phylogenies of plasmids or to track the spread of particular markers.

Incompatibility (Inc) group identification has been frequently used to classify plasmids. Identification methods include the initial technique based on conjugation (Novick, 1987), hybridization with cloned replication regions (Couturier et al., 1988), PCR-based replicon typing (PBRT) (Carattoli et al., 2005) and relaxase typing (Compain et al., 2014). Moreover MLST schemes for plasmids were developed to assign plasmids to STs, in analogy to the typing developed for bacterial genomes (García-Fernández et al., 2008).

extrachromosomal DNA molecules capable of autonomous reptators of resistance and virulence genes, especially in Enterobacte fierring drug resistance is important for analysis of evolution, eprobacte fierring drug resistan Mitotic segregation of plasmids, termed partition in bacteria, is a fundamental step of the cell cycle that ensures the transmission of the whole genome to daughter cells. It is governed by specific genetic loci named *par,* first identified in low-copy-number plasmids and later found to be present as homologues in most bacterial chromosomes. *par* loci are organized into operons encoding two proteins, an ATPase and a DNA-binding protein, and including a centromeric site. These components interact with each other to direct the subcellular localization that ensures stability of their replicons. Three types of partitioning ATPases are known (Gerdes et al., 2010): the Walker-type ATPases encoded by the *par/sop* gene family (type I partitioning loci which are the most common of the *par* systems), the actin-like ATPase encoded by the *par* locus of plasmid R1 (type II partitioning locus) and the tubulinlike GTPases encoded by plasmids from *Bacillus* sp. (type III partitioning loci). Despite their similarities in genetic organization, these three *par* types use entirely different molecular mechanisms (Guynet and de la Cruz, 2011).

While the acquisition of plasmids often enables bacteria to survive in the presence of antibiotics, it is possible that plasmids also confer vulnerabilities that may be exploited in tailored antibacterial therapy (Williams and Hergenrother, 2008). In order to control the spread of multiresistance plasmids, we need to determine many more variables that affect their replication, maintenance and movement.

Recently we developed a multiplex PCR method called "plasmid relaxase gene typing" (PRaseT). This classification scheme is based on the relaxase, a key protein which is part of the mobilization region of transmissible plasmids. The aim of the present study was to identify different partition systems located on multiresistance plasmids and to design a multiplex PCR method here called "plasmid partition gene typing" (PAR-T). This method could further the classification of plasmids in *Klebsiella pneumoniae*, *Escherichia coli* and *Salmonella enterica spp.* and will constitute another option for characterising plasmids.

#### **2. Materials and methods**

#### **2.1. Database search, primer design and** *in silico* **primer assay**

Frent partition systems located on multiresistance plasmids a<br>CR method here called "plasmid partition gene typing" (PAR-1<br>r the classification of plasmids in *Klebsiella pneumoniae*, *Esch-<br>enterica spp*, and will constit An *in silico* analysis was carried out using GenBank BLAST (http://blast.ncbi.nlm.nih.gov/) on plasmids >40 kb from *K. pneumoniae* and *E. coli* conferring multidrug resistance. Due to a low number of plasmids IncHI, IncI1 from *K. pneumoniae*, the *in silico* analysis was extended to some plasmids from *Enterobacter cloacae* and *Citrobacter freundii*. For the eight Inc groups studied (IncF, IncA/C, IncL/M, IncN, IncHI, IncR, IncI1, IncX), the *par* operon was used as template; the presence of partition-specific multidomains was searched for using CD-Search (http://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi/). Multiple alignments were performed with ClustalW2 software (http://www.ebi.ac.uk/Tools/msa/clustalw2/). Primer pairs covering most sequences in each family were designed using FastPCR (http://primerdigital.com/fastpcr.html) and Primerblast (http://www.ncbi.nlm.nih.gov/tools/primer-blast) software, while minimizing codon degeneracy (Table 1). Oligonucleotide primers were tested *in silico* for hybridization with plasmids from Enterobacteriaceae referenced in GenBank.

#### **2.2. Bacterial strains and plasmids**

that (Tc) or transformants (Tf) of Enterobacteriaceae (Table 2). A<br>bonging to diverse Inc groups that encoded β-lactamases conferrition<br>cephalosporins and/or carbapenems (ESBLs, acquired ce<br>ases). They were part of four c For validation of the PCR assays, experiments were conducted with 136 *Escherichia coli* transconjugants (Tc) or transformants (Tf) of Enterobacteriaceae (Table 2). All strains carried plasmids belonging to diverse Inc groups that encoded β-lactamases conferring resistance to third-generation cephalosporins and/or carbapenems (ESBLs, acquired cephalosporinases, carbapenemases). They were part of four collections of, respectively, (i) *E. coli* strains isolated between 1997 and 2002 in various French university hospitals (Marcadé et al., 2009) (Branger et al., 2005)), (ii) *K. pneumoniae* strains from various geographical regions collected since the 1980s (D. Decré and G.Arlet, personal collection), (iii) *Salmonella enterica* subsp. *enterica* strains representing various serovars (collection of the French National Reference Center for *E. coli*, *Shigella sp.*, and *Salmonella* sp., Institut Pasteur), (iv) *E. coli*, *S. enterica*, *K. pneumoniae* strains isolated during 2013 in various Argentina University Hospitals (M. Jure, personal collection). All Tc, Tf and clinical strains used in this study were analyzed in parallel with the PAR-T and PRaseT methods. The IncR plasmids which do not encode relaxases were analyzed only by using PBRT (Carattoli et al., 2005). The Tc used as positive controls in PAR-T reactions are given in Table 2. After optimization using Tc or Tf carrying replicons of various types according to PRaseT, we applied the PAR-T method to a panel of 30 clinical strains (17 *E. coli*, 11 *K. pneumoniae* and 2 *S. enterica*) carrying replicons of one to four different types (Table 3).

#### **2.3. DNA extraction and PCR conditions**

Lysis by boiling was used for total DNA extraction as previously described (Dallenne et al., 2010). Multiplex PCR was carried out using the Qiagen Multiplex PCR kit (Qiagen, Courtaboeuf, France). The master mix contained pre-optimized concentrations of HotStarTaq DNA polymerase and  $MgCl<sub>2</sub>$ , deoxynucleotide triphosphate and buffer. To all multiplex PCRs, solution Q (Qiagen) that facilitates the reaction with difficult-to-amplify templates by modifying DNA melting was added. Total DNA in 2 μL of bacterial lysate was subjected to multiplex PCR in a 50 μL volume. The conditions for multiplex PCR were optimized to ensure that all targets were sufficiently amplified for amplicons to be easily visible on 1.5% agarose gels. The optimal primer concentrations are reported in Table 1. PCR conditions consisted of an initial activation at 95°C for 5 min, followed by 35 cycles of 95°C for 30 s, 57°C for 90 s and 72°C for 90 s with a final extension at 68°C for 10 min; for Multiplex I and Multiplex II annealing temperature was elevated to 60°C for 90 s. Simplex PCRs were

ter and 2 µL of total DNA extract. PCR conditions consisted in<br>S<sup>o</sup>C for 40 s, 72<sup>o</sup>C for 1 min], preceded by 1 cycle at 94<sup>o</sup>C for 5 nr<br>7<sup>o</sup>C for 5 min. PCR products were separated at 100 V for 90 n<br>8 v) on 1.5% agarose performed in a 50 µL mix with 2U of Taq DNA polymerase (Roche Diagnostics), 10×PCR buffer/MgCl<sub>2</sub> (Roche Diagnostics, Meylan, France), 200  $\mu$ M of each deoxynucleotide triphosphate (dNTP Mix, Eurobio, Courtaboeuf, France), 0.2 pmol/µL of each primer, 40 µL of sterile water and 2 µL of total DNA extract. PCR conditions consisted in 30 cycles [94°C for 1 min, 55°C for 40 s, 72°C for 1 min], preceded by 1 cycle at 94°C for 5 min and followed by 1 cycle at 72°C for 5 min. PCR products were separated at 100 V for 90 min (180 min for Multiplex IV) on 1.5% agarose gel electrophoresis containing ethidium bromide and visualized using GelDoc (Biorad, Marnes-La-Coquette, France). PCR products were purified using the Exosap purification kit (Illustra Exostar-1 Step, Dutscher, Brumath, France) and subjected to bidirectional DNA sequencing using the BigDye terminator v3.1 cycle sequencing kit (Applied Biosystems, Foster City, CA, USA) and an Applied Biosystems 3730 XL capillary sequencer. Sequence analysis was carried out using Sequence scanner (Applied Biosystems), GeneDoc (www.psc.edu/biomed/genedoc) and GenBank BLAST software.

#### **3. Results and discussion**

#### **3.1.** *In silico* **analysis**

We mainly focused on plasmid families previously found to be involved in the spread of resistance genes in Enterobacteriaceae (Carattoli, 2013). The majority of plasmids contain a single *par* locus, with the exception of some plasmids such as pR55 plasmid which include two *par* loci (type I *parAB* and type II *parMR*). The type I partitioning loci are the most common of the *par* systems (74%). The partition loci are often close to *rep* gene. In contrast, some of them are located at some distance, e.g. the *parAB* locus of pCTX-M-360 which is 30000 bp by away from RepA. Different designations are used in the annotated sequences for *par* loci: *par*, *sop*, *stb*. *In silico* analysis showed that some Inc groups are relatively homogeneous with respect their partition systems, such as IncL/M, N, A/C, I1 or R. The high sequence similarity within these groups (99-100%) allowed designation of a pair of primers for each group. The study of the genetic environment (500 nucleotides on either side of the *par* genes or their equivalent) of the same groups showed high similarity (75 to 100%). In contrast, other groups such as IncF showed more divergent *par* sequences. In addition, IncX plasmids from *K. pneumoniae* (e.g. pIncX-SHV, accession number JN247852 or pKPC-NY79, accession number JX104759) carried a *parA* gene encoding a Walker-type ATPase near the *rep* region and an annotated *parB* gene 25 kb apart from *parA* without a partitionspecific multidomain. *In silico* analysis of IncX plasmids in *E. coli* showed a different

organization from that found in *K. pneumoniae,* with *parA* and *parB* organized in an operon. *In silico* analysis of IncHI plasmids did not show any correlation between *par* loci and IncHI1 and HI2. Furthermore, the phylogram of *K. pneumoniae* in Fig. 1 reveals partition proteins form distinct subgroups and are connected to Inc group with the exceptions of plasmids IncF. *In silico* analysis led to the design of 18 pairs of primers targeting the *par* operons of plasmids from *K. pneumoniae*, *E. coli* and *Salmonella enterica spp.* belonging to Inc groups A/C, F, HI, I1, L/M, N, R. For plasmids belonging to the IncX group we decided to target only the *parA* gene (Table 1). The primer names for IncF (F1 to F9) and IncHI (HIa and HIb) groups were arbitrarily chosen.

#### **3.2. Primer evaluation using transconjugants and transformants**

subgroups and are connected to Inc group with the exceptions of<br>ysis led to the design of 18 pairs of primers targeting the *par* oper<br>*umoniae, E. coli* and *Salmonella enterica spp.* belonging to Inc gre<br>R. For plasmids In order to assess the sensitivity and specificity of each PCR, primers were tested using a collection of 136 recipient cells, with PRaseT as the reference method (Table 2). Each primer pair was validated using all recipient cells, first in a simplex and then a multiplex PCR and target DNA of either single cells or cell mixtures was used. PCR conditions were optimized and all amplicons were sequenced. *E. coli* strain J53 was used as negative control in PCR experiments to test for possible cross-hybridization with chromosomal DNA. An example of the results is shown in Fig S1. No non-specific amplification was observed. PAR-T results were largely consistent with the PRaseT results (98.6%) except for two strains which carried an IncX replicon (*E. coli* strains 64 and 110). These results were surprising. If the study of IncX plasmids indicates a diversity in their backbones, no data allow an explanation as to why the *par* loci are organized differently in *K. pneumoniae* and *E. coli* and why there are differences in the *parA* genes between these two species. A complete analysis of all IncX replicon present in the databases will be required (all Enterobacteriaceae).

#### **3.3. Evaluation of PAR-T using clinical strains**

on time the specificity of the designed primers (Table 3). An example Fig S1. For 24 strains (80%) there was a perfect correlation beth PAR-T and PRaseT. Six strains were positive with PRaseT bong these, five carried IncX Thirty clinical strains, each carrying replicons of one to four different types, were tested by PAR-T to confirm the specificity of the designed primers (Table 3). An example of the results is shown in Fig S1. For 24 strains (80%) there was a perfect correlation between the results obtained with PAR-T and PRaseT. Six strains were positive with PRaseT but negative with PAR-T. Among these, five carried IncX replicons (four *E. coli* strains, i.e. 17, 19, 28 and 34, and one *K. pneumoniae* strain, KpS20) and one (*E. coli* strain 105) carried an IncI1 replicon. As shown by *in silico* analysis, *par* loci in IncX plasmids are organized differently between *K. pneumonia*e and *E. coli*. It is likely that our analysis targeting plasmids >40 kb of *K. pneumoniae* and some *E. coli* did not allow the study of all the *par* loci of IncX plasmids. A complete analysis of all IncX replicon present in the databases will be required (all the Enterobacteriaceae). The strain (*E. coli* 105) that carried an IncI1 plasmid was negative with PAR-T. We considered three possibilities: (i) a divergent IncI1 *par* locus that could not hybridize with our primers was present, (ii) the *par* locus was truncated or (iii) the gene locus was absent. *In silico* analysis has shown that the *par* sequences of IncI1 replicons are well conserved and form a homogen group, thus making the first possibility (i) unlikely.

Overall the specificity were very high (>90%) for all Inc groups studied except IncX.

Finally, four strains (*E. coli* 19 and 34, and *K. pneumoniae* KpS5 and KpS26) which were negative with PRaseT were found to contain IncF and IncR plasmids, respectively, when PAR-T was used. For the *K. pneumoniae* strains, the results were not unexpected as IncR plasmids do not encode relaxases. The two *E. coli* were positive with PBRT with IncFIA, IncFIB and IncFII replicons found in both. Moreover, PCR targeting the genes of the type IV secretion system was negative and the plasmids from neither strain could be transferred to a recipient cell by conjugation (Compain et al., 2014). We considered the possibility of the loss of genes for conjugative transfer.

#### **4. Conclusions**

Considering the complexity of constant plasmid evolution and the unavailability of full-length plasmid sequencing in most laboratories, the combined use of several complementary classification methods should be a practical value. Our set of seven multiplex PCRs allowed classification of the most frequently encountered transmissible plasmids in *Klebsiella pneumoniae*, *Escherichia coli* and *Salmonella enterica spp*. by targeting their *par* loci, with the exception of IncX replicons. For this group, *in silico* analysis of all plasmids present in the databases must be carried out to design new primers and to improve the value of the method described here.

#### Conflicting interest

The authors declare that they have no conflicting interests.

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Figure 1. Phylogram of the partition proteins. Unrooted evolutionary tree showing 24 ParA and ParB proteins (or equivalent) from plasmid of *K. pneumoniae, E. coli, Salmonella spp., C. freundii, E. cloacae*. pKpQIL-IT include two *par* loci. Types I and II refer to the two types of plasmid-encoded partitioning loci shown in Fig. 1 and described in the text. The phylogram was constructed using Mega5 (Neighbor-Joining algorithm, bootstrap: 1000 replication).

Table 1. Primers used in this study.

Table 1. Primers used in this study.								
PCR name and plasmid type detected	Primer name	Sequence (5' - 3')	Le ng th (b) p)	Plasmi $\mathbf{d}$ prototy pe	<b>EMBL</b> accession number	Targe t site	Amplic on size (bp)	Primer concen tration $(\mu M)$
Multiplex I, II, III: IncFIA, IncFIB,								
IncFIC, IncFII, <b>IncFIIk</b>	-F1-for GGTC $-F1-$	<b>CAACCAGTG</b> SopAB CATAAGTGCT SopAB ACCATACGC <b>GTGAGGCGC</b>	23	pKPN 3	CP00064 8.1	sopA sopB	1370	0,4
	rev -F2-for GTC	<b>TCTCAA</b> <b>ACCAGTGCA</b> SopAB TAAACGCTG SopAB GCGTGGTTTA	24	pKF3- 21 140	FJ876827	sopA sopB	1130	0,4
	$-F2-$ rev SopAB	<b>ATCAGACGA</b> <b>TCGAA</b> <b>GATGACGAA</b> <b>GGCTATTGCC</b> -F3-for ATTGC	24 24	$\tau$	pKP18 CP00096 5.1	sopA sopB	720	0,4
Multiplex I : sop systems E. coli, K. pneumoniae, S. enterica	$-F3-$ rev -F4-for TGC	SopAB GCGCATACG <b>ACATTGATCG</b> <b>TGCCA</b> <b>TGACGAAGG</b> SopAB CTATCGCCAT	24	22 CZ	pKPN- JX424424 sopA $\cdot$ 1	sopB	928	0,4
	$-F4-$ rev	SopAB ATGCTTCCCC <b>AGGCATCCC</b> <b>GAA</b> <b>ACCAGTGCA</b> SopAB TAAACGCTG	22		pH233 KJ484626 sopA			0,4
	-F5-for GTC	SopAB CTGTAAGTGC AGCAGCTTTA 24		21 2-166	$\cdot$ 1	sopB	1533	





Table 2. Recipient cells used in validation experiments. Tc : transconjugants, Tf : transformant, PRaseT : plasmid relaxase gene typing, PBRT : PCR-based replicon typing, PAR-T : plasmid partitioning genes typing. † : Tc used as a positive control in partitioning genes typing.

genes typing.							
$N^{\circ}$ Parental strain		Tc/Tf	$\beta$ -Lactamase	Plasmid classification			
				PRaseT			
				(and/or PBRT)	PAR-T		
48	E. coli	Tc	<b>TEM-52</b>	X1	X		
51	E. coli	Tc	TEM-3	L/M	X		
52	E. coli	<b>Tf</b>	<b>TEM-52</b>	X1	X		
57	E. coli	Tc	<b>SHV-12</b>	$\boldsymbol{\mathrm{F}}$	${\bf F}$		
62	E. coli	Tc	$SHV-2$	<b>FIB</b>	${\bf F}$		
64	E. coli	Tc	SHV-5	X4			
73	E. coli	Tc	$SHV-4$	<b>FIIK</b>	<b>FIIK</b>		
85	E. coli	<b>Tf</b>	CTX-M-1	<b>FIA</b>	${\bf F}$		
91	E. coli	Tc	CTX-M-3	${\bf N}$	${\bf N}$		
94†	E. coli	Tc	CTX-M-9	<b>FII</b>	$\mathbf F$		
98	E. coli	Tc	CTX-M-1	X1	X		
100	E. coli	Tc	CTX-M-1	FIA, FII	${\bf F}$		
102	E. coli	Tc	CTX-M-1	HI <sub>2</sub>	HIa		
104	E. coli	Tf	CTX-M-1	FIA, FIB, FII	$\mathbf F$		
105	E. coli	Tc	CTX-M-1	L/M	L/M		
108	E. coli	<b>Tf</b>	$CTX-M-1$	<b>FIA</b>	${\bf F}$		
110	E. coli	Tc	CTX-M-1	X4			
111 <sup>†</sup>	E. coli	Tc	CTX-M-1	<b>FIA</b>	$\mathbf F$		
114	E. coli	Tc	CTX-M-1	L/M	L/M		
115	E. coli	Tc	CTX-M-9	FII	${\bf F}$		
118	E. coli	Tc	CTX-M-1	FIA, FIB	${\bf F}$		
120	E. coli	Tc	CTX-M-3	<b>FIB</b>	${\bf F}$		
122	E. coli	Tc	CTX-M-9	<b>FII</b>	${\bf F}$		
125	E. coli	Tc	CTX-M-1	FII	${\bf F}$		
126	E. coli	Tc	CTX-M-1	${\bf N}$	${\bf N}$		
127	E. coli	Tc	CTX-M-3	<b>FIB</b>	${\bf F}$		
Levy	E. coli	Tc	OXA-48	L/M	L/M		
$AD-48$	E. coli	$\operatorname{Tc}$	OXA-48	$L/M$	${\rm L/M}$		
$AD-50$	E. coli	Tc	<b>OXA-48</b>	L/M	L/M		
$AD-17$	E. coli	Tc	<b>OXA-48</b>	L/M	L/M		
$AD-44-2$	E. coli	Tc	NDM-1	${\bf N}$	$\mathbf N$		
$AD-2$	E. coli	Tc	<b>OXA-48</b>	L/M	L/M		
Goe-137	E. coli	Tc	<b>VIM</b>	HI <sub>2</sub>	HIa		
Goe-132	E. coli	Tc	<b>VIM</b>	A/C	A/C		
<b>KATS</b>	E. coli	Tc	NDM-1, CTX-M-1	$\mathbf F$	${\bf F}$		
$M-2$	E. coli	Tc	$KPC-2$	L/M	L/M		
$M-3$	E. coli	Tc	$KPC-2$	L/M	L/M		
$M-5$	E. coli	Tc	$KPC-2$	L/M	L/M		
Ec <sub>50</sub>	E. coli	Tc	<b>TEM-52</b>	I1	$\mathbf{I}$		
S1	K. pneumoniae	$\operatorname{Tc}$	$CMY-4$	A/C	A/C		









Table 3. Plasmid partition gene typing of 30 clinical strains of Enterobacteriaceae.



#### **Highlights**

-Low-copy-number plasmids utilize partition systems for plasmid maintenance.

-*par* loci are organized into operons encoding two proteins, an ATPase and a DNA-binding protein, and including a centromeric site.

-The method called "plasmid partition gene typing" showed high specificity for the classification of resistance plasmids (IncA/C, FIA, FIB, FIC, FIIk, FII, HI1, HI2, I1, L/M, N) except for IncX replicons.

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