This work describes the remarkable heterogeneity of plasmids found in ten *E. coli* ST131 sequenced genomes. We find members of 8 out of the 17 main MOB plasmid groups found in γ-proteobacteria [11,53]. Specifically, we find members of Inc groups F12 (IncF), P3 (IncX), P5 (ColE1), P6 (Incl2), P11 (Incp), P12 (Incl/K/BO), Q12 (Rep_pSC101-like), Qu (Rep_pMG828-2/IGWZ12-like) plus others lacking relaxases that were identifiable as phase-related/RepFIB and no-MOB small plasmids. To date, IncF [15,17,29,56,80,92,93,119, Suzuki et al., 2009, Matsumura et al., 2013], IncFilK [81], IncI1 [93, Curiao et al., 2011]; IncN [79,120, Suzuki et al., 2009, Zong et al., 2011, Novais et al., 2012] and, sporadically, IncA/C [Naseer et al., 2010, Novais et al., 2012], IncX [94], IncU [Matsumura et al., 2013], IncY [Dhanji et al., 2011], IncK [Calhau et al., 2013], Incly [Calhau et al., 2013], IncB/O [Matsumura et al., 2013] and ColE [Matsumura et al., 2012] plasmids were identified in *E. coli* ST131 by PBRT. They were associated with the spread of class A [CTXM (-1,-2, -3, -9, -14, -14, -15, -27, -32, -61), TEM (-4, -24, -116), SHV (-2, -5, -7, -12), NDM (-1), KPC (-2, -3, -4)], class B (VIM-1), class C (CMY-2) and class D (OXA-1) beta lactamases [12,13,23-29] [26,81,83,121, Rogers et al., 2011, Ma et al., 2013, Matsumura et al., 2013, Accogli et al., 2014, Cai et al., 2014] and genes coding for resistance to different antibiotic families (tetracyclines, macrolides, aminoglycosides, trimethoprim and sulfonamides), which are located in composite genomic islands, mostly derived from IncF plasmids (see main text). Despite extensive plasmid analysis of ST131 isolates in different studies, the ST131 plasmidome was not described comprehensively, as most information comes uniquely from identification of antibiotic resistant plasmid replicons by PBRT. As of April 2014, ten plasmids were sequenced from ST131 strains, as shown in Table 3. They include four IncFII, four IncN, one IncI1 and one IncX4 plasmids. Here we provide an individualized discussion of the 8 plasmid groups (39 plasmids) found in our ST131 survey, together with the ten previously sequenced plasmids. This information underscores the relevance of the ST131 plasmidome in the adaptation of this *E. coli* clonal group by providing emergent functions (e.g., antibiotic resistance, virulence, colonization enhancement, ability to outcompete other microbial species) and thus, the usefulness of PLACNET in bacterial population studies, as explained in the main body text of the manuscript.

**MOB_{12}/IncF plasmids (Suppl. Fig. S11).** Recent plasmid history suggests that IncF plasmids represent the most abundant family among *E. coli* metapopulations. They are drivers of *E. coli* evolution because of their ability to acquire multiple adaptive traits coding for antibiotic resistance and virulence
Hanni et al., 1982, Womble et al., 1988]. Figure 5 (inset) shows that IncF plasmids identified in the ST131 genomes correspond to four dendrogram groups. The two main groups were Group I (FIB), frequent among avian pathogenic \textit{E. coli} (APEC) and extraintestinal pathogenic \textit{E. coli} (ExPEC), and linked to putative virulence traits and Group II (FII/FIA), widely disseminated among ExPEC and extensively associated with antibiotic resistance (see Table 4). Besides, there were two outlier groups, Group III and Group IV.

Group I (FIB) plasmids mostly comprises virulence plasmids (such as pAPEC-like ones) and four ST131 plasmids containing numerous virulence genes. They are compared in Fig S11A, using ST131 plasmid pJE186_2 as a reference. Three FIB (\(\Delta\text{FIA:B1}\)) plasmids, pE61BA_1 and pE35BA2+3, were identified in two strains corresponding to ST9/H22/virotpe D and ST43/H30/virotpe B, respectively. To date, only one report [56] documented FIB plasmid carriage in \textit{E. coli} ST131, despite the identification of putative virulence markers associated with CoIV plasmids often appeared in surveys of ST131 \textit{E. coli}[29,36, Kudinha et al., 2013]. Fig S11A emphasizes that all ST131 FIB plasmids (and relatives) share a 80kb genetic island (coordinates 93 kb to 21 kb in the map of Fig S11A) that includes a conserved CoIV region comprising \textit{iss, iroBCD}, \textit{iucABCD}, \textit{iutA, cvaBC} and \textit{sitC} and an \textit{ompT-hlyF-mig14} cassette[55]. While some traits of the conserved region can also be located in the chromosome (linked to PAIs), the cassette \textit{ompT-hlyF-mig14} has only been detected in plasmids[2, Billard-Pomares et al., 2011].

Group II (FII/FIA/FIB) plasmids are compared in Fig S11B, using plasmid pJJ1886_5 as a reference. The seven plasmids analyzed show extensive sequence similarity over at least 100 kb, which includes not only backbone genes but also metabolic, transport and genes associated with mobile genetic elements (MGE). Interestingly, five of the seven plasmids show extensive deletions within their Tra region (see Fig S11B from coord 40 kb to 70 kb). A wide diversity of IncFII/FIA/FIB plasmids carrying \textit{bla} genes have been described among ST131 clonal variants[17,92,93, Fiett et al., 2014], some plasmid types being apparently overrepresented at a global [29] or local level [Doumith et al., 2012, Matsumura et al., 2013] as F2:A1:B- (similar to pEK399) and F2:A-:B- (similar to pC15-1a, pHK01, pEK516), or F29:B10 (identified in ST131 from South Europe). The diversity of replicons, RFLP patterns and sequence mosaicism of \textit{E. coli} ST131 IncF plasmids, even within situations that resemble plasmid outbreak scenarios [17,80,92,119, Smet et al., 2010] suggest that multiple DNA rearrangements among IncF2 plasmids of Enterobacteriaceae are extremely frequent under selection, as described earlier[Hanni et al., 1982, Womble et al., 1988]. More recently, it has been shown that homologous recombination mediated by
IS26 and Tn2 between particular ST131 IncF2 plasmids occurs often and partly explains the mentioned diversity[80, Smet et al., 2010]. Recombination of F plasmids among them or with other plasmid groups in enterobacterial populations involved in multispecies outbreaks of $b\text{l}a_{\text{CTX-M-15}}$ and $b\text{l}a_{\text{KPC-2}}$, confirms rearrangements among large composite regions containing antibiotic resistance genes, transposons and ISs and/or transfer regions of F plasmids is extraordinarily common[Sandegren et al., 2012]. The overrepresentation of F2:A1:B- (pEK499 derivatives) [93, Dhanji et al., 2011]and F2:A:-B- plasmids (e.g. pC15-1a, pHK01, pEK516), the last group only inferred in this study in an H30 strain of “viotype B”[17,29,92, Ho et al., 2012], can reflect the expansion of different fimH30 ST131 sublineages (here designated as virotypes A, B and C) in different areas.

Group III contains only plasmid pHVH177_1, without antibiotic resistance genes and low content of virulence genes. The BRIG representation (Fig 5), which compares pHVH177_1 with its closest relatives in the dendrogram of Figure 5, shows conservation of just the MOB$_{F12}$ backbone genes (about 32 kb).

Group IV is represented by plasmid pECSF1, from the commensal ST131 strain SE15. No resistance genes and low content of virulence genes were detected. The most similar plasmids (Fig 5) are the 122 kb plasmid p1ESCUM[66], the 115 kb plasmid pUM146 (GenBank acc NC_017630) and the 114 kb plasmid pEC14_114[DebRoy et al., 2010]. Besides backbone genes, these plasmids share more than 60 additional kb that includes a number of potential transposable elements. Other close relatives are the 168 kb plasmid pIP1206 [Perichon et al., 2008] and the K. pneumoniae MDR plasmid pKF3-140[Zhao et al., 2010].

All these data taken together underscore the complex evolution of MOB$_{F12}$/IncF plasmids, facilitated by the recombinogenic potential of transposable elements[92, Hanni et al., 1982, Womble et al., 1988, Partridge et al., 2004, Johnson et al., 2010].

**Phage-related/ RepFIB plasmids (Suppl Fig S12).** The ST131 E.coli plasmids pBIDMC20B_2 and pBWH24_2, found in two ST43/H30/viotype C isolates, are closely related among them and to the antibiotic resistance plasmid pECOH89 [57] and the STEC plasmid p09EL50[5]. They all share the same RepFIB replication and ParB proteins as backbone remnants, and an extensive set of phage-related proteins (Fig S12A). This group also comprises other phage-related plasmids such as plasmid pLF82 [58] from the prototype strain of adherent invasive E. coli (AIEC), the Salmonella plasmid pHCM2[59], the
Salmonella bacteriophage SSU5 [60] and the Yersinia pestis pMT1 and MT plasmids [Hu et al., 1999]. An extensive set of phage-related genes are shared by all seven plasmids. As shown in Fig S12A, only pECOH89 plasmid harbors the bla\textsubscript{CTX-M-15} antibiotic resistance gene. As shown in the phylogenetic tree of Fig S12B, the RIP protein RepFIB (Rep3\_superfamily, pfam 01051) from ST131 plasmids is identical to those of plasmids pECOH89 and p09EL50 (here named as Group C). They are only 40% identical to Group B proteins, represented by plasmids pLF82, pHCM2, pMT1 and MT. Group A in the figure is represented by the well-known IncF (FIB) plasmids discussed in the previous section. At this point it must be stressed that, although a majority of plasmids carrying RepFIB belong to the IncF group, RepFIB also appears in more than 20 other RIP groups such as IncN, IncP or IncI groups [Gibbs et al., 1993]. Further work is required to rigorously classify RepFIB proteins (all belonging to the Rep3\_superfamily) according to plasmid RIP groups. Moreover, the biology of RepFIB/phage-related plasmids remains largely unexplored, although they might be more extended than previously thought [57].

MOB\textsubscript{p12}/IncI\textsubscript{complex} (Suppl Fig S13). The single MOB\textsubscript{p12}/Incl representative in our collection is plasmid pE2022\_1, most similar to the Inc1\_ST16 plasmid pEK204, carrying bla\textsubscript{CTX-M-15}, already isolated from a ST131 strain [93]. Plasmids pEK204 and pE2022\_1 are not phylogenetically close among themselves. Rather, they are similar to widespread IncI (bla\textsubscript{CTX-M-15}) plasmids from Europe and IncK (bla\textsubscript{CTX-M-14}) plasmids from Asia, respectively [16, 93, Zang et al., 2013, Onnberg et al., 2014].

The IncI plasmid pEK204 shares its backbone with the eight IncI (also called IncI\textalpha) plasmids used as references in Fig S13A. While pEK204 carries (bla\textsubscript{CTX-M-15} and bla\textsubscript{TEM}), these antibiotic resistance genes are absent in its closest relatives (see coord. 6 kb and 14 kb in S13A). Plasmids in this cluster include the IncI\textalpha prototype, the 93.2 kb plasmid R621a from S. enterica [Takahashi et al., 2011].

The ST131 plasmid pE2022\_1 (98.3 kb) is similar to the IncK prototype 93.6 kb plasmid pCT [61]. They share more than 80 kb including Tra regions and the bla\textsubscript{CTX-M-14} gene coding for resistance to beta-lactam antibiotics. IncK pCT-like plasmids harboring bla\textsubscript{CTX-M-14} are globally spread among animals and humans, being prevalent in UK and Spain among other countries [63, 64, Dhanji et al., 2011]. Although widely spread, they have been only sporadically described among ST131 isolates, since IncF plasmids are the main vectors carrying bla\textsubscript{CTX-M-15} and bla\textsubscript{CTX-M-14} genes in ST131 to date.

MOB\textsubscript{p0}/Incl2 plasmids (Suppl Fig S14). The two IncI2 plasmids found in our ST131 genomes (pBWH24_3 and pE61BA_7) belong to different clusters of a large plasmid family that, however, contains few
sequenced plasmids. Fig S14A shows the phylogenetic tree of MOB₉ REL proteins, underscoring the distant positions of the ST131 plasmid REL proteins. Plasmid pBWH24_3 (60.3 kb) is similar to the *E.coli* IncI2 plasmid prototype R721 (75.6 kb; [63]) and the *E.coli* APEC plasmid pChi7122_3 (56.7 kb, GenBank acc FR851304) over most of their length (Fig S14B). Plasmid R721 is the representative of a number of enterobacterial plasmids commonly isolated from *K. pneumoniae* and associated with spread of antibiotic resistance, mainly *bla*CMY-2 and *bla*KPC-2 [83]. Plasmids pBWH24_3 and pChi7122_3, on the other hand, lack any antibiotic resistance gene. These two plasmids are highly similar, the last one having a role in acid resistance and biofilm formation [64]. Besides, it is of note that IncI2 plasmids have a high number of putative integration sites, which would facilitate the acquisition of different antibiotic resistance genes, resulting in large composite multidrug platforms[83].

Plasmid pE61BA_7 (37.9 kb) is most similar (Fig S14C) to the *S. enterica* plasmid SL483 (37.9 kb; [Fricke et al., 2011] and *E. coli* plasmid pO157_Sal (37.8 kb; [65]). Plasmid pE61BA_7 belongs to a different IncI2 subcluster than pBWH24_3, as shown in Fig S14A. This group is composed of cryptic plasmids, represented by SL483 from *Salmonella agona* and pO157_Sal from *E. coli* O157:H5, and seems to be widely spread among enterobacterial isolates from animals [65]. IncI2 plasmids have been identified among CTX-M-15 producers of the ST131/H30 sublineage collected in Canada, Australia and New Zealand [16] but they were not fully characterized in any of those studies.

**MOB₉/IncX plasmids (Suppl Fig S15).** IncX plasmids are prevalent in *E. coli* and belong to a large plasmid family with four recognized subgroups X1 to X4 [Johnson et al., 2012]. The two IncX plasmids identified in this survey belong to the IncX1 (pFV9873_4) and IncX4 (pE2022_3) subclusters (Fig S15A). To these, the previously reported IncX4 plasmid pJIE143 should be added. As shown in Fig S15B, the IncX1 plasmid pFV9873_4 (33.3 kb) was most similar to the 33.8 kb plasmid p2ESCUM [66] over its entire length, and no so much to other reference hits. p2ESCUM was originally isolated from the emblematic strain UMN026, representative of a widely disseminated urinary pathogenic ST69 *E. coli* (UPEC) clone [Lescat et al., 2009]. The IncX4 plasmid pE2022_3 (35.0 kb) shares extensive homology with the *S. enterica* plasmid pSH696_34 (33.8 kb, [Gokulan et al., 2013]), which is used as a reference in Fig S15C. Compared to these two plasmids, the ST131 reference plasmid pJIE143 (34.3 kb;[94]), widely disseminated among *E. coli* from foodborne animals, lacks two important backbone genes, the conjugative coupling protein gene *traG* (around coord 19 kb) and the RIP encoding region (coord 9 kb to 10 kb). Plasmid pJIE143 codes for a completely different Pir protein (13% amino acid identity) although
also belonging to the Rep3 family. Therefore, it is probable that pSH696_34 and pE2022_3 belong to a different incompatibility group than pJIE143. IncX4 plasmids can be transferred at different temperatures and are frequent vehicles of antibiotic resistance, mainly qnr. Difficulties for detecting this plasmid group using PBRT might have underestimated its prevalence.

**MOB \(_{p11}\)/IncP1 plasmids (Suppl Fig S16).** Members of the IncP1 family are frequently isolated in the environment and often carry modules containing diverse accessory genes can be transferred between unrelated bacteria [Schluter et al., 2007, Venturini et al., 2013]. IncP1 plasmids from *E. coli* remain largely unexplored although detection of IncP sequences from animal [Dotto et al., 2014] and human *E. coli* isolates, included ST131 [16], suggest that IncP1 plasmids can be easily acquired by human associated Proteobacteria. As can be observed in the phylogenetic tree of Fig S16A, plasmids pJJ1886_4 and pE61BA_4 represent different new branches of the MOB\(_{p11}\) family and thus new additions to the ST131 plasmidome. The pJJ1886_4 REL protein is most similar to that of the 40.6 kb plasmid pDS1. However, proteome analysis resulting in the dendrogram of Fig. 5, shows plasmid pH5102707 (*E. coli*, 69.5 kb, Genbank acc NC_023907) as its closest homolog. This is confirmed in Fig S16B, where the extensive homology of pJJ1886_4 and pH5102707 is depicted. The second branch of the phylogenetic tree contains the ST131 plasmid pE61BA_4 (18.3 kb). The 37.9 kb plasmid pMBUI2 [67], from an uncultured bacterium, appears as its closest homolog at the level of REL protein. Plasmid pE61BA_4 seems a crippled plasmid, lacking most of its backbone, with just four genes shared with pMBUI2 (Suppl. Table S1), suggesting that it is the first representative of a new plasmid group. Finally, the same phylogenetic branch contains IME_E35BA REL protein. To analyze this integrative and mobilizable element (IME), we compared the relevant regions of E35BA and JJ1886 genomes using EasyFig [Sullivan et al., 2011]. Fig S16C shows the IME_E35BA (14.2 kb) inserted between a hypothetical protein and a GMP synthase encoding genes in *E.coli* JJ1886. IME_E35BA is similar to the *Burkholderia glumae* IncP island (14.7 kb; [Yoshii et al., 2012]). Both share a *tra* conjugative region, *repA*, *alpA* (encoding a DNA binding protein) and *int-P4* (coding for a P4 phage integrase family protein). They differ in their *repC* (resolvase) genes. Besides, IME_E35BA does not harbor the kasugamycin-2′-N-acetyltransferase gene [aac(2′)-lia] present in the *Burkholderia glumae* strain that confers resistance to kasugamycin, an aminoglycoside antibiotic widely used in agriculture [Yoshii et al., 2012].

**MOB\(_{c12}\) plasmids (Suppl Fig S17).** The MOB\(_{c}\) family constitutes a wide plasmid group, present in \(\gamma\)-proteobacteria, Firmicutes and Tenericutes. MOB\(_{c}\) relaxases corresponding to \(\gamma\)-proteobacterial
plasmids are clustered in the MOBc1 group, subdivided as well in two branches: MOBc11 and MOBc12[11, Garcillan-Barcia et al., 2009]. Plasmid pE61BA_2 (24.5kb), located in a ST131 ST9/H234 E.coli strain, is not close to any reference from Figure 5, but clusters in the MOBc12 group according to its REL (Fig S17A). pE61BA_2 REL protein is only 70% identical to the closest hits: the 22 kb Y. pestis cryptic plasmid pCRY [Song et al., 2004] and the 42 kb K.pneumoniae multidrug resistant (MDR) plasmid pMET1 [Soler Bistue et al., 2008]. As shown in Fig S17B, pE61BA_2 shares mobilization (mobBC –virB) and replication (repA) genes with pCRY and pMET1, but lacks the micrococcal nuclease-like and the Mpr coding genes. Since PBRT does not detect MOBc plasmids, they are probably underrepresented in the plasmidome of Enterobacteriaceae.

MOBps/ColE plasmids (Suppl Fig S18). The MOBps REL phylogenetic tree shows three different branches, constituting the MOBps1, MOBps2 and MOBps3 subfamilies [11]. The four ST131 MOBps plasmids are located in the MOBps1 subfamily (Fig S18A). Plasmid pE61BA_5 (6.6 kb) is similar to the prototype MOBps plasmid ColE1 (6.6 kb) [Tomizawa et al., 1977] over their entire length (including MOB proteins and colicin-E1), as shown in Fig S18B. The ST131 plasmids pBIIDMC38_1 (11.8 kb) and pJJ1886_3 (5.6 kb) are very similar among them, including their MOB and RIP proteins (Fig S18C). They are unique in carrying a large (2 kb) gene encoding an intriguing EamA-like transporter protein (pfam00892). Finally, plasmid pE61BA_6 (6.9 kb) is similar to enterohemorrhagic E. coli (EHEC) O157 plasmid pColD-157 (6.7 kb, [Hofinger et al., 1998]) and colicin K plasmid pColK-K235 (8.3 kb, [Rijavec et al., 2007]). As Fig S18D shows, pE61BA_6 plasmid shares MbkABCD mobilization and colicin K encoding regions with the pColK-K235 reference plasmid.

Although widely used as cloning vectors for genetic engineering, ColE plasmids represent a largely unexplored family in epidemiological studies, since they not usually carry genes coding for antibiotic resistance. On the other hand, they frequently, but not always, carry colicins [68]. The MOBps/ColE1-like plasmids identified in this study exhibited a wide size range (5.6-10.5 kb) and variable presence of colicins, two of them carrying either ColE1 or ColK, both in the same strain (Suppl figures S18A-D). These two colicins have traditionally been associated with UPEC and the B2 phylogenetic group of E. coli [68, Rijavec et al., 2007]. Although the carriage of ColE-1 like plasmids, inferred by the presence of either RIP or REL sequences, seems to be frequent among non-outbreak O25b and non-O25b ST131 isolates [Matsumura et al., 2013], the lack of full plasmid sequences and functional information preclude any conclusion about a role in the pathogenesis of B2 and ST131 clones. Besides a possible involvement of
colicin production in virulence, perhaps as a competition weapon to avoid the presence of other E. coli pathogenic strains [Budic et al., 2011], CoIE1 plasmids can promote rearrangements and contribute to the mobilization of other plasmids [Rijavec et al., 2007]. As an example, the large plasmid (pBIDMC38_1) is almost identical to the ST131 reference pJJ1886_3, with an additional type II restriction-modification system (Cfr10I) (Fig. S18). In spite of the above, CoIE plasmids carrying genes encoding antibiotic resistance are increasingly being reported although by lack of phylogenetic analysis, they cannot be accurately classified within the MOB95 group [10, Dotto et al., 2014].

**MOB\textsubscript{Q12} plasmids (Suppl Fig S19).** MOB\textsubscript{Q12} and MOB\textsubscript{Q12} plasmid clusters were previously identified by our group [11,53]. They have scarcely been documented among E. coli to date, probably because they do not carry antimicrobial resistance genes. This report is the first to document their presence in the ST131 lineage. Nevertheless, they seem to be common among UPEC, as shown in this study. The four MOB\textsubscript{Q12} plasmids identified in this study were almost identical among them and with pSE11\_6 [Oshima et al., 2008] in a DNA stretch of 4 kb, as shown in Fig S19B. Apparently they only code for RIP and MOB proteins. The role of these cryptic plasmids in the adaptation of ST131 is intriguing. Although not explored at all, a role in the adaptation by providing useful mobilization tools cannot be excluded.

**MOB\textsubscript{Q12} plasmids (Suppl Fig S20).** As seen in the REL phylogenetic tree of fig S20A, the four ST131 MOB\textsubscript{Q12} plasmids have almost identical relaxases. Three of the four MOB\textsubscript{Q12} plasmids (of about 5.2 kb) (pBIDMC38_2, pFV9873_6 and pJJ1886_2) were almost identical among them and to the UPEC plasmid pCE10B [Lu et al., 2011], as shown in Fig S20B. They constitute a cluster of plasmids within the MOB\textsubscript{Q12} family previously defined by the authors [11,53]. The fourth plasmid (pE61BA\_3; 5.5 kb) has a unique 1.6 kb DNA segment that codes for a 266 amino acid protein identical to a colicin from S. enterica (NCBI protein acc. WP\_024155916). These plasmids frequently coexist with IncF plasmids and form cointegrates with them (our unpublished data).

**Small cryptic no-MOB plasmids (Suppl Fig S21).** Small non-mobilizable cryptic plasmids remain scarcely documented among Enterobacteriaceae. Four highly similar plasmids of about 1.6 to 2.2 kb were found in our study. They are also similar to the 1.5 kb E. coli plasmid pCE10D [Lu et al., 2011], as shown in Fig S21. As can be expected, they have minimal coding capacity. Their only annotated gene codes for a RIP protein of the Rep\_HTH\_36\_superfamily (pfam13730). Similar plasmids have also been detected in various ExPEC, including ST131 [90], Shiga-toxin producing E. coli (STEC) [Lu et al., 2011, Brolund et al.,
2013] and Klebsiella [Liu et al., 2012] isolates. A fifth, 5kb no-MOB plasmid (pFV9873_3) was unique and unrelated to any reference plasmid (see Figure 5). The adaptive functions of these plasmids, if any, are unknown.

Other ST131 plasmid groups: MOB_{F11}/IncN plasmids (Suppl. Fig. S22). We carried out extensive literature analysis in the search of plasmid groups present in ST131 but not detected in our survey. There were two plasmid groups, IncN and IncA/C, for which no plasmid was identified in our study although they are being increasingly detected among ST131 isolates. Both groups have been associated with bla_{CTX-M} [120, Novais et al., 2012] and more recently with bla_{KPC} [89,81,121, Matsumura et al., 2013] and bla_{NDM} genes (GenBank acc n° KJ413946). These plasmids have spread worldwide and might have been acquired by ST131 from other enterobacterial species, e.g., during nosocomial polyclonal outbreaks [85].

Four complete sequences of MOB_{F11}/IncN plasmids originating from ST131 strains are available: the IncN1 plasmids pECN580 [79], pKC394 and pKC396 [120] (GenBank acc n° HM138652 and HM138653, respectively) and the IncN2 plasmid pNDM-ECS01 (GenBank acc n° KJ413946). Suppl Fig S22A shows the REL phylogenetic tree of the MOB_{F11}/IncN plasmid branch, which displays the IncN1 and IncN2 subclusters. BRIG comparative analysis of the three ST131 IncN1 plasmids versus reference plasmids (Suppl Fig S22B) reveals a highly conserved 34kb backbone region (coordinates 0kb to 34kb). As previously reported [11], IncN1 and IncN2 plasmid backbones, including REL proteins, are homologous but their RIP proteins are different [Poirel et al., 2011, Partridge et al, 2012]. Accordingly, the IncN2 pNDM-ECS01 plasmid RIP has no similarity to the RIPS from pECN580, pKC394 or pKC396 IncN1 plasmids. Comparative analysis of the pNDM-ECS01 plasmid (Suppl Fig S22C), harboring the bla_{NDM-1} gene, versus the reference plasmids p271A [Poirel et al, 2011], pTR3 and pTR4 [Chen et al., 2012] and pJIE137 [Partridge et al., 2012] shows that all are almost identical except pJIE137, which harbors the bla_{CTX-M-62} gene. Despite of the low prevalence of IncN1 and IncN2 plasmids in ST131 E.coli strains, these plasmids seem to be good vehicles for the transmission of several antibiotic resistance genes in Enterobacteriaceae by maintaining conserved backbones but variable regions that harbor genes encoding CTX-M, KPC or NDM beta-lactamases.
Supplemental references


