Supplementary Data for:

"Complete assembly of *Escherichia coli* ST131 genomes using long reads demonstrates antibiotic resistance gene variation within diverse plasmid and chromosomal contexts"

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Supplementary Results

Oxford Nanopore long read quality control and long read genome assembly

Half of the reads were produced within 14 hours of sequencing, with the reminder produced over the subsequent 34 hours (Supplementary Figure 1d). A median read length of 5.5 Kb for reads Q (quality) score > 7 was achieved within one hour of sequencing (Supplementary Figure 1e), and the median Q score declined slightly as the run proceeded (Supplementary Figure 1f). An average of 30-fold theoretical coverage from 954 Mbases with Q > 7 was exceeded in this GridION run within three hours.

We compared short read-only, long read-only and hybrid assembly outputs from Unicycler v.4.6 using the long Oxford Nanopore reads and short Illumina reads to identify the most contiguous assemblies per sample across all three Unicycler modes (conservative, normal and bold).

For five samples, the long read assemblies produced 2-7 contigs (with a median of three) with nearly identical results across modes, whereas the short read assemblies resulted in 76-230 contigs (a median of 124), and the hybrid assemblies also had more contigs (6-191 with a median of 44). For VREC0739 and VREC1428, the short read libraries resulted in over-bridging of contigs making it harder to classify contigs as chromosomal or plasmid-associated, perhaps because long reads already provided sufficient genome coverage and the assembler inserted the contigs produced by short reads at short homologous repetitive regions.

VREC1013 assembly assessment and improvement

For VREC1013, the hybrid assembly improved the long read assembly such that the final optimised version had three rather than 22 contigs and a smaller length (5.36 Mb, Supplementary Table 3), after manual sequence alignment eliminated seven false-positive short contigs. Five contigs had depths of coverage <8% of the chromosomal median and may were the result of contig overbridging during assembly. Pairwise alignment of these five contigs with BLAST against the assembly showed that they had near-perfect matches (E-value < E-10) with other contigs, showing that they were effectively duplicate contigs, and thus few reads mapped to them. In contrast, the other four valid contigs acted positive controls and showed high homology to their own contigs only. As a result, duplicate contigs were removed from the VREC1013 hybrid assembly used for subsequent analyses.

Supplementary Figure 1. Overview of genome assembly using Oxford Nanopore reads to recover plasmids with antibiotic resistance genes and mobile genetic elements (MGEs). Oxford Nanopore fast5 sequences were basecalled and converted to fastq format using Albacore v.2.0 and Guppy v.0.5.1. Forward, reverse and middle adapters were removed using Porechop v.0.2.4. The genomes were assembled using Unicycler v.4.6 (optionally including Illumina short reads for comparison). The probability that the resulting contigs were chromosomal or plasmid-associated was measured using mlplasmids. Contigs were annotated using the Comprehensive Antibiotic Resistance Database (CARD) and Multiple antibiotic Resistance Annotator (MARA) to resolve precise plasmid structure, *bla*CTX-M gene alleles, copy numbers and their adjacent regions.

FAST5

Guppy/Albacore: basecalling



Raw FASTQ

Porechop: trim adapter, demultiplex



Clean FASTQ

Unicycler: assembly, bold mode



Complete assemblies

mlplasmids: extract plasmid contigs



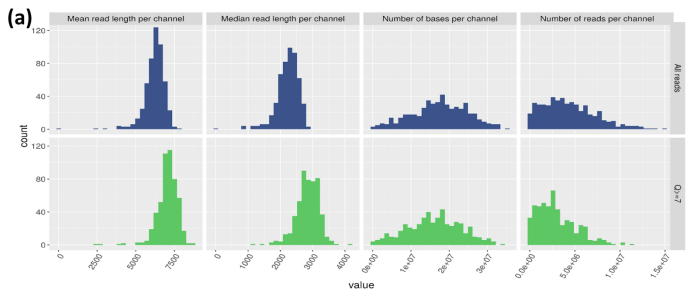
Plasmid-derived contigs

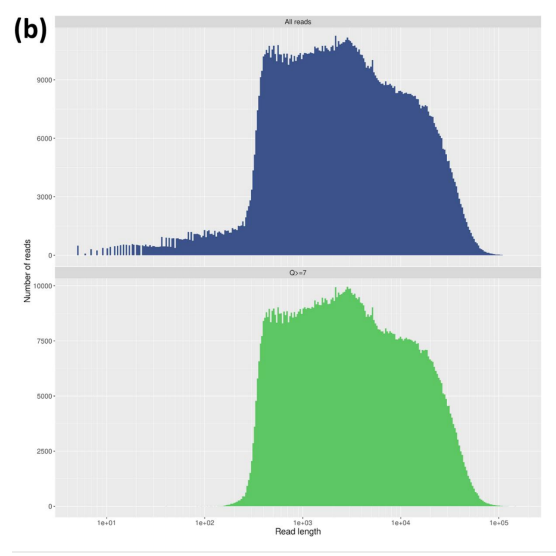
CARD, MARA: annotate plasmids

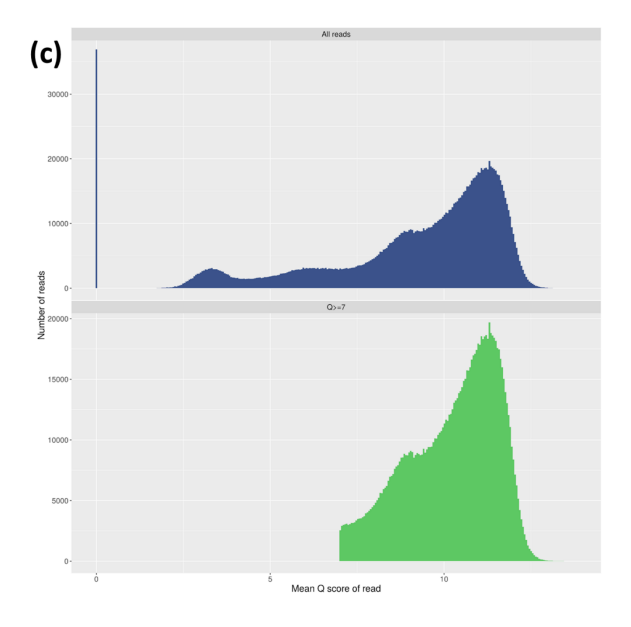


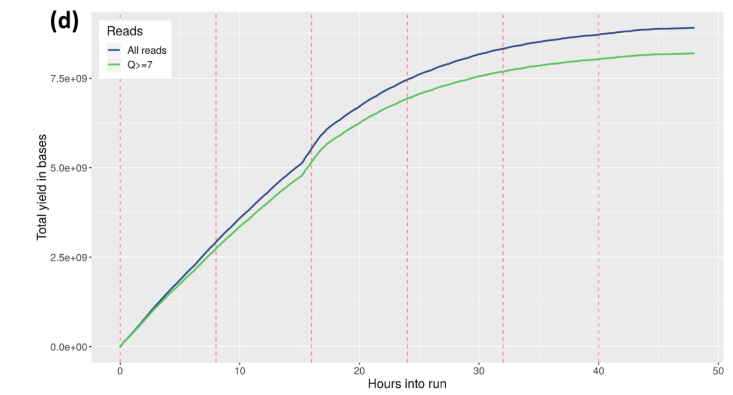
Annotated plasmids

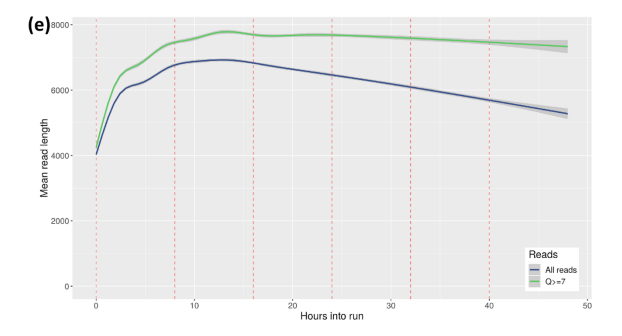
Supplementary Figure 2a-g. Summary plots of the GridION X5 sequencing run for all (blue) and filtered (green) nanopore reads generated using MinIONQC. The graphs in (a) show the read count (y-axis) with the mean and median read length and the number of bases and reads per channel (x-axis), the overall read count (y-axis) vs length (x-axis) in (b) and read count (y-axis) vs the mean Q score (x-axis) in (c). Plots were also drawn to present the total amount of bases called (x-axis; d), the mean read length (x-axis; e) and the mean Q score (x-axis; f) per hour (in their y-axes); the total amount of bases (y-axis) contained in a minimum read length (x-axis) is shown in (g).

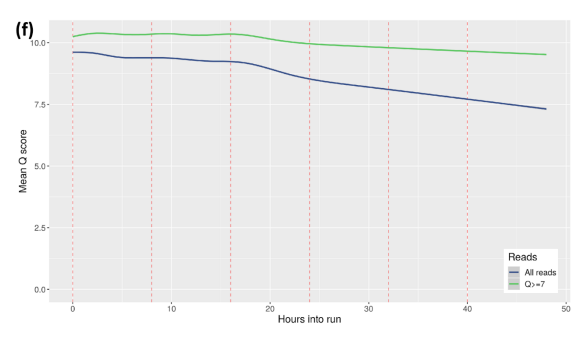


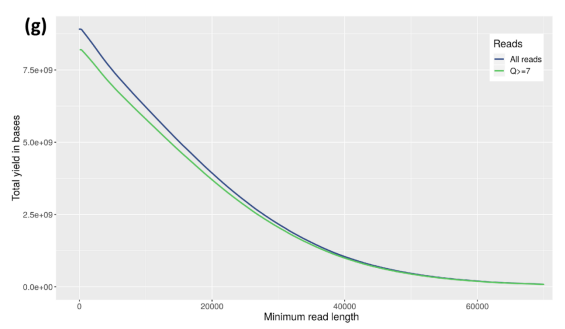




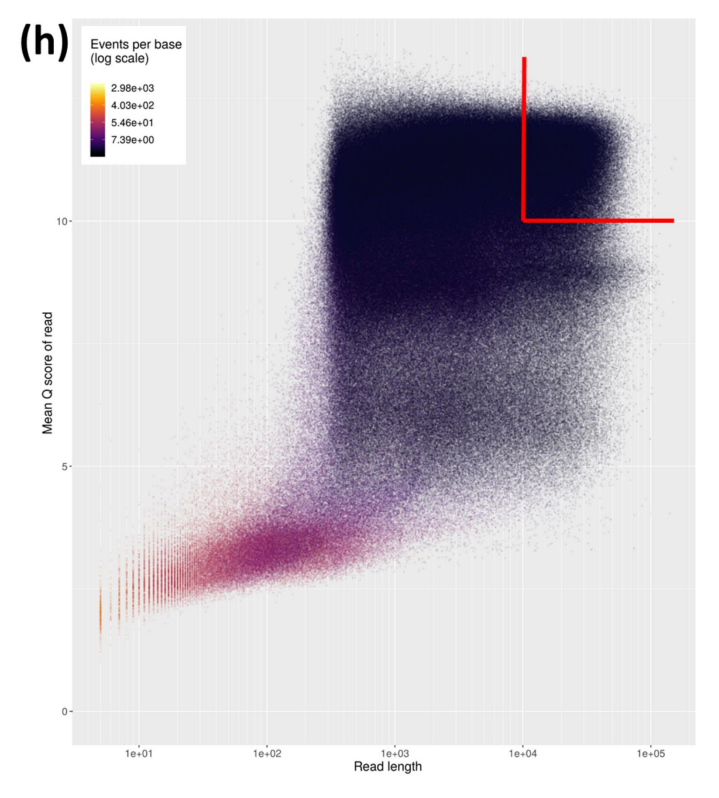








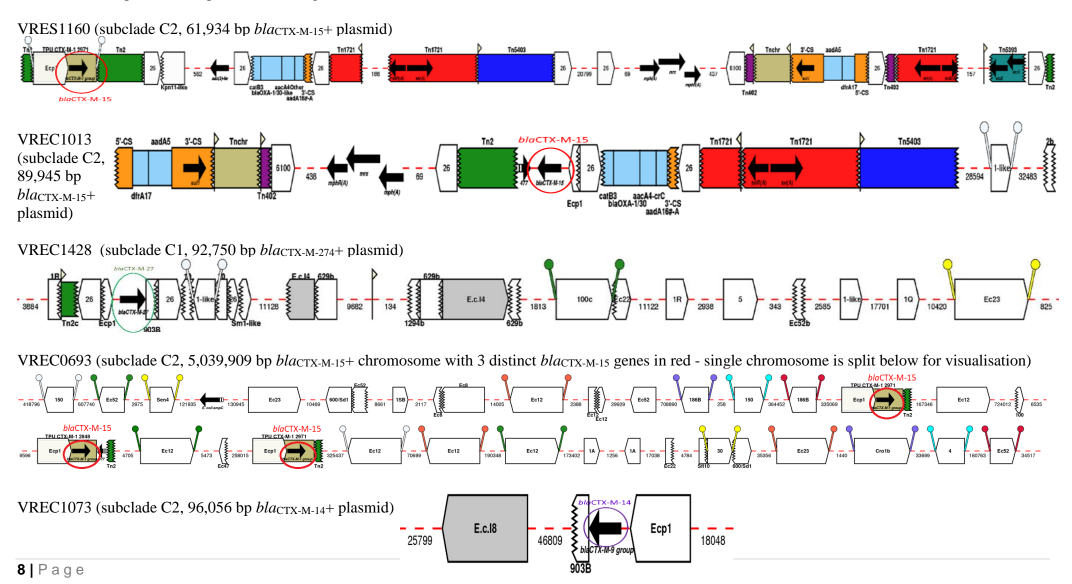
Supplementary Figure 3. Summary of the GridION X5 sequencing run output showing the read length on a log10 scale (x-axis) versus the mean Q score of each read (y-axis) where points are coloured by events per base. The horizontal red line shows reads with lengths > 10 Kb and the vertical red line read with Q scores > 10. Together, this area shows the large number of long high-quality reads generated in this study. This plot emphasises that a high proportion of the bases were accurately called: these were subsequently used for downstream analysis.

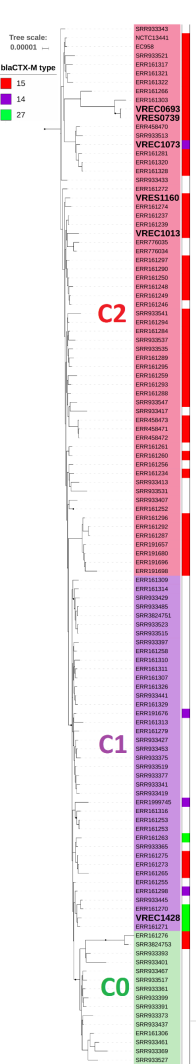


Supplementary Figure 4. The assembly graphs of six *E. coli* ST131 genomes showed many connected edges for those created from short Illumina HiSeq reads only (left) but near-complete assemblies for those made with long Oxford Nanopore read-only (centre) and the hybrid assemblies of most of the strains (right). The assemblies were generated with Unicycler v.4.6 and were visualised using Bandage. Circularized contigs indicated complete assemblies.

Sample	Short read-only	ONT long read-only	Hybrid assembly
VRES1160			
VREC0693			. + + = =
VREC0739			
VREC1013			
VREC1073			
VREC1428			

Supplementary Figure 5. The contigs from the most optimal assembly mode of Unicycler v.4.6 of five out of six *E. coli* ST131 samples were identified as chromosomal or plasmid-derived using mlplasmids. These were annotated with *bla*_{CTX-M} genes and their genetic flanking context using GalileoTM AMR based on the Multiple Antibiotic Resistance Annotator (MARA) and database [35]; all blaCTX-M variants are labelled accordingly and encircled in red (*bla*_{CTX-M-15}), purple (*bla*_{CTX-M-14}) or green (*bla*_{CTX-M-27}). The definition of the other elements are listed at https://galileoamr.arcbio.com/mara/feature/list. The long VREC0693 chromosome is split into two parts so that the gene annotation is visible.





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Supplementary Figure 6. Phylogram of the six ST131 genomes showed that all except VREC1428 were in ST131 subclade C2 (red: VRES1160, VREC1073, VRES0739, VREC0693 and VREC1013). VREC1428 clustered in subclade C1 (purple). No new isolate was in C0 (green). The phylogram was built with RAxML v.8.2.11 and iTOL v4.3 using 3,603 non-recombinant SNPs from Gubbins v.2.3.4 where branch support was performed by 100 bootstrap replicates, and the scale bar indicates the number of substitutions per site. Clade classification was based on phylogenetic analysis by [8] by including the reference NCTC13441, n=63 isolates from [8] and n=56 from [42] with associated classification and bla_{CTX-M} allele data. The right-hand part shows bla_{CTX-M-15} (red), bla_{CTX-M-14} (purple) and bla_{CTX-M-27} alleles (green). The six isolates' names are in large bold text. This mid-pointed rooted phylogeny included reference genome isolates EC958 and NCTC13441 (both in C2) and a clade B isolate as an outgroup (Figure 3). The C2 isolates were mainly bla_{CTX-M-15}-positive (48 out of 62, including VRES1160, VRES0739, VREC0693 and VREC1013), bar 13 that were bla_{CTX-M}-negative and one that was *bla*_{CTX-M-14}-positive (VREC1073). The C0 isolates were mainly bla_{CTX-M-15}-negative (13 out of 15), as were the C1 (30 out of 40) isolates except for four that were blactx-M-27-positive, three that were blactx-M-15positive and three that were *bla*_{CTX-M-14}-positive.

Supplementary Table 1. Sample collection source, sampling date and sequence read accession numbers.

Strain	Source	Sampling	Accession numbers				
Strain S	Source	date	Short reads	Long reads			
VRES1160	Faeces	26/08/2015	ERR1878359	https://ndownloader.figshare.com/files/14039495			
VREC0693	Faeces	03/06/2015	ERR2137889	https://ndownloader.figshare.com/files/14039639			
VRES0739	Faeces	05/06/2015	ERR1878196	https://ndownloader.figshare.com/files/14039354			
VREC1013	Faeces	19/08/2015	ERR2138591	https://ndownloader.figshare.com/files/14039333			
VREC1073	Blood	26/08/2015	ERR2138200	https://ndownloader.figshare.com/files/14039345			
VREC1428	Faeces	22/10/2015	ERR2138475	https://ndownloader.figshare.com/files/14039351			

Supplementary Table 2. Contigs were classified as chromosomal or plasmid-derived using the mlplasmids prediction value. Each contig were aligned against CARD to identify the presence/absence of $bla_{\text{CTX-M}}$ alleles and their copy numbers. Plasmid types were identified using PlasmidFinder.

Strain	Prediction		Contig ID	_	bla _{CTX-M} allele				Normalized
	C1	value (%)		(bp)		count	type	Depth	Depth 1.00
	Chromosome	98 - 0	1	5,126,679			-	258	1.00
	Chromosome	70	2	113,086			-	213	1.00
VRES1160	Plasmid	70	3	61,934		1	IncFIA	282	1.10
VILLETTOO	Plasmid	85	4	15,803			ColRNAI	420	1.64
	Plasmid	81	5	5,203			ColRNAI	11	0.04
	Plasmid	83	6	4,096			Col8282	473	1.85
	Chromosome	98	1	5,039,909	15	3	-	258	1.00
VREC0693	Plasmid	61	2	132,042			IncFIB	213	0.83
	Plasmid	60	3	88,790			IncB	282	1.09
	Chromosome	98	1	4,797,749			-	171	1.00
VRES0739	Plasmid	96	2	5,162			Col156	436	2.55
	Plasmid	74	3	4,001			-	303	1.77
	Chromosome	97	1	3,699,451			-	300	1.00
VREC1013	Chromosome	97	2	1,434,037			-	335	1.00
	Plasmid	84	4	89,945	15	1	IncFII	1015	3.27
	Chromosome	98	1	5,286,804			-	214	1.00
VREC1073	Plasmid	68	2	156,298			IncFIA	172	0.80
	Chromosome	60	3	96,056	14	1	-	213	1
	Chromosome	98	1	4,924,536			-	126	1.00
	Chromosome	97	2	103,034			-	57	1.00
	Chromosome	96	3	101,160			-	41	1.00
VREC1428	Plasmid	64	4	92,750	27	1	IncFIA	85	0.67
	Plasmid	92	5	5,147			ColRNAI	168	1.33
	Plasmid	99	6	5,143			Col156	207	1.64
	Plasmid	73	7	4,649			ColRNAI	239	1.90

Supplementary Table 3. Comparison of short read-only, long read-only and hybrid genome assemblies generated using the conservative, normal and bold modes of Unicycler v.04.6. Assemblies were assessed according to their total length, number of contigs produced, N50, numbers of mismatches per 100 Kb and numbers of indels per 100 Kb.

Assembly	Mode	Metric	VRES1160	VREC0693	VRES0739	VREC1013	VREC1073	VREC1428
Co		Total length (bp)	5,142,342	5,146,205	5,181,497	5,208,807	4,967,093	5,375,468
		Number of contigs	168	159	200	148	117	230
	Conservative	N50	124,175	132,865	138,725	134,439	157,528	135,303
		#mismatches / 100 Kb	1.32	1.32	65.4	1.5	285.81	0.69
		#indels / 100 Kb	0.06	0.02	1.84	0.08	261.91	0.04
		Total length (bp)	5,158,728	5,171,710	5,227,751	5,240,888	4,989,316	5,416,180
CI 4		Number of contigs	110	106	123	94	76	148
Short read-only	Normal	N50	206,138	190,908	213,071	189,184	222,158	170,443
read-only		#mismatches / 100 Kb	4.64	0.93	69.86	4.25	284.14	2.96
		#indels / 100 Kb	0.21	0.14	2.33	0.36	262.1	0.04
		Total length (bp)	5,159,662	5,163,846	5,207,686	5,226,735	4,977,746	5,411,973
		Number of contigs	124	120	146	108	86	140
	Bold	N50	206,044	190,808	212,979	190,412	222,051	184,466
		#mismatches / 100 Kb	3.07	1.78	67.11	1.96	287.37	2.03
		#indels / 100 Kb	0.16	0.06	2.03	0.13	262.26	0.11
		Total length (bp)	5,326,801	5,260,741	4,806,912	6,307,464	5,539,158	5,236,419
		Number of contigs	6	3	3	22	3	7
	Conservative	N50	5,126,679	5,039,909	4,797,749	5,073,008	5,286,804	4,924,536
		#mismatches / 100 Kb	276.23	241.39	2,772.51	344.5	0	332.79
		#indels / 100 Kb	252.29	264.7	265	306.03	0	289.71
		Total length (bp)	5,326,801	5,260,741	4,806,912	6,307,464	5,539,158	5,236,419
Long		Number of contigs	6	3	3	22	3	7
read-only	Normal	N50	5,126,679	5,039,909	4,797,749	5,073,008	5,286,804	4,924,536
reau-omy		#mismatches / 100 Kb		241.39	2772.51	344.5	0	332.79
		#indels / 100 Kb	252.29	264.7	265	306.03	0	289.71
		Total length (bp)	5,326,801	5,260,741	4,806,912	6,307,464	5,539,158	5,236,419
		Number of contigs	6	3	3	22	2	7
	Bold	N50	5,126,679	5,039,909	4,797,749	5,073,008	5,286,804	4,924,536
		#mismatches / 100 Kb		241.39	2772.51	344.5	0	332.79
		#indels / 100 Kb	252.29	264.7	265	306.03	0	289.71
		Total length (bp)	5,272,824	5,275,251	5,215,332	5,323,049	5,055,625	5,492,517
		Number of contigs	52	6	191	34	51	107
	Conservative	N50	1,444,640	5,048,264	426,378	2,673,977	1,423,856	749,550
		#mismatches / 100 Kb	1.63	242.24	2,764.2	2.04	285.57	3.7
		#indels / 100 Kb	0.32	265.38	263.44	0.09	263.18	0.02
		Total length (bp)	5,276,305	5,275,251	5,291,108	5,327,833	5,098,966	5,516,886
Hybrid	Normal	Number of contigs	42	6	110	33	44	74
		N50	1,746,191	5,048,264	72,0730	2,675,388	1,762,353	1,243,293
		#mismatches / 100 Kb	1.56	242.24	44.59	2.28	284.11	1.65
		#indels / 100 Kb	0.28	265.38	4.07	0.13	266.82	0.02
		Total length (bp)	5,293,427	5,275,251	5,267,003	5,223,433	5,115,410	5,550,270
		Number of contigs	23	6	32	3	22	47
	Bold	N50	3,801,465	5,048,264	1,222,073	3,699,451	4,958,323	1,266,683
		#mismatches / 100 Kb	271.47	242.24	2,770.38	321.64	283.97	296.99
		#indels / 100 Kb	252.55	265.38	264.11	268.47	268.27	268.29

Supplementary Table 4. Protein products encoded by the genes found in the plasmid of strains VREC1013, VRES1160 and VREC1428 (Figure 2).

Gene	Protein product
tmrB	Tunicamycin resistance protein
tetA	Tetracycline resistance protein
pinR	Serine recombinase protein
sopB	Inositol phosphate phosphatase
repE	Replication initiation protein
pifC	Transcriptional repressor protein
xerD	Tyrosine recombinase protein
agp	Glucose-1-phosphatase
chrA	Response regulator
fol P	Dihydropteroate synthase
xerC	Tyrosine recombinase protein
neo	Aminoglycoside 3'-phosphotransferase
tnpR	Transposon gamma-delta resolvase
pemK	mRNA interferase
nqrC	Na(+)-translocating NADH-quinone reductase
lolD	Lipoprotein-releasing system ATP-binding
hemR	Hemin TonB-dependent receptor
repB	Replication protein
rocC	Amino-acid permease
ccDA	Antitoxin (Plasmid maintenance)
klcA	Antirestriction protein
traD	Coupling protein
traI	Multifunctional conjugation protein
finO	Fertility inhibition protein