






Penicillin Binding Protein Substitutions Cooccur with Fluoroquinolone Resistance in Epidemic Lineages of Multidrug-Resistant *Clostridioides difficile*

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ABSTRACT *Clostridioides difficile* remains a key cause of healthcare-associated infection, with multidrug-resistant (MDR) lineages causing high-mortality ($\geq 20\%$) outbreaks. Cephalosporin treatment is a long-established risk factor, and antimicrobial stewardship is a key control. A mechanism underlying raised cephalosporin MICs has not been identified in *C. difficile*, but among other species, this is often acquired via amino acid substitutions in cell wall transpeptidases (penicillin binding proteins [PBPs]). Here, we investigated five *C. difficile* transpeptidases (PBP1 to PBP5) for recent substitutions, associated cephalosporin MICs, and co-occurrence with fluoroquinolone resistance. Previously published genome assemblies ($n = 7,096$) were obtained, representing 16 geographically widespread lineages, including healthcare-associated ST1(027). Recent amino acid substitutions were found within PBP1 ($n = 50$) and PBP3 ($n = 48$), ranging from 1 to 10 substitutions per genome. β -Lactam MICs were measured for closely related pairs of wild-type and PBP-substituted isolates separated by 20 to 273 single nucleotide polymorphisms (SNPs). Recombination-corrected phylogenies were constructed to date substitution acquisition. Key substitutions such as PBP3 V497L and PBP1 T674I/N/V emerged independently across multiple lineages. They were associated with extremely high cephalosporin MICs; 1 to 4 doubling dilutions $>$ wild-type, up to 1,506 $\mu\text{g/mL}$. Substitution patterns varied by lineage and clade, showed geographic structure, and occurred post-1990, coincident with the *gyrA* and/or *gyrB* substitutions conferring fluoroquinolone resistance. In conclusion, recent PBP1 and PBP3 substitutions are associated with raised cephalosporin MICs in *C. difficile*. Their co-occurrence with fluoroquinolone resistance hinders attempts to understand the relative importance of these drugs in the dissemination of epidemic lineages. Further controlled studies of cephalosporin and fluoroquinolone stewardship are needed to determine their relative effectiveness in outbreak control.

IMPORTANCE Fluoroquinolone and cephalosporin use in healthcare settings has triggered outbreaks of high-mortality, multidrug-resistant *C. difficile* infection. Here, we identify a mechanism associated with raised cephalosporin MICs in *C. difficile* comprising amino acid substitutions in two cell wall transpeptidase enzymes (penicillin binding proteins). The higher the number of substitutions, the greater the impact on phenotype. Dated phylogenies revealed that substitutions associated with raised cephalosporin and fluoroquinolone MICs were co-acquired immediately before clinically important outbreak strains emerged. PBP substitutions were geographically structured within genetic

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lineages, suggesting adaptation to local antimicrobial prescribing. Antimicrobial stewardship of cephalosporins and fluoroquinolones is an effective means of *C. difficile* outbreak control. Genetic changes associated with raised MIC may impart a “fitness cost” after antibiotic withdrawal. Our study therefore identifies a mechanism that may explain the contribution of cephalosporin stewardship to resolving outbreak conditions. However, due to the co-occurrence of raised cephalosporin MICs and fluoroquinolone resistance, further work is needed to determine the relative importance of each.

KEYWORDS AMR mechanism, antimicrobial resistance, *Clostridioides difficile*, PBPs, penicillin binding proteins, cell wall transpeptidase, cephalosporin MIC, cephalosporin resistance

Clostridioides difficile is among the leading causes of health care-associated infection, with symptoms ranging from diarrhea to potentially fatal pseudomembranous colitis (1). Over the last 30 years, unrestricted antimicrobial use has selected multidrug-resistant (MDR) *C. difficile* lineages, which can be identified by multilocus sequence type (ST) and/or PCR ribotype (2–9). Uncontrolled prescribing of antimicrobials such as fluoroquinolones and cephalosporins, which are associated with a high risk of *C. difficile* infection (CDI), creates conditions under which MDR lineages can cause persistent, high-mortality ($\geq 20\%$) outbreaks (9–18). Such health care-associated transmission may be geographically widespread, as in the “hypervirulent” ST1 ribotype 027 [ST1(027)] lineage FQ-R1 (19), and/or prolonged, as in ST17(018), predominating in Japanese and Italian health care settings since the 1990s (17, 20). Cases associated with the rapid transmission of MDR lineages are typically superimposed on a background of sporadic, unlinked cases caused by diverse *C. difficile* strains, which lack acquired antimicrobial resistance (AMR) (21, 22).

Antimicrobial stewardship is an extremely effective means of preventing or resolving CDI outbreaks in health care settings (23–29). This approach contributed to the marked decline in fluoroquinolone-resistant lineages in the United Kingdom a decade ago, with resistant ST1(027), ST3(001), ST42(106), and ST37(017) falling from 67% to $\sim 3\%$ of cases (30, 31). Fluoroquinolone resistance can be predicted from whole-genome sequences by characteristic single nucleotide polymorphisms (SNPs) in the chromosomal *gyrA* and/or *gyrB* genes (30, 32). Equivalent analysis for cephalosporins is lacking because the genetic mechanism(s) influencing cephalosporin susceptibility in *C. difficile* is unknown. Wild-type cephalosporin MICs, defined as the phenotype conferred by genes in their ancestral, nonmutated form, as found in the natural *C. difficile* population, are already moderate to high for this species, up to and greater than 256 $\mu\text{g/mL}$ (33–36). This has led to the widely accepted concept that *C. difficile* cephalosporin MICs are intrinsically high (10). For this reason, and the practical difficulties of determining MICs approaching the limit of drug solubility, cephalosporin MICs are rarely measured for *C. difficile*.

Studies aiming to understand the mechanism of *C. difficile* cephalosporin resistance have focused on the endogenous *C. difficile* class D β -lactamase, but findings have been inconclusive (37, 38). In many bacterial species, reduced susceptibility to cephalosporins and other β -lactams is conferred by amino acid substitutions in penicillin binding proteins (PBPs). These are enzymes catalyzing cell wall peptidoglycan biosynthesis that are classified according to molecular weight (high [HMW] or low [LMW]) and enzymatic activity (transpeptidase or carboxypeptidase) (39). β -Lactam antibiotics target PBPs by acting as inhibitory substrate analogues (39), binding covalently to the active site serine (40) in the first of three conserved motifs, SXXK, (S/Y)XN, and (K/H)(S/T)G (41). Exposure to β -lactams selects substitutions which reduce the affinity of the drug for the PBP, increasing the MIC (42, 43). However, the nature and frequency of PBP substitutions among clinically important *C. difficile* lineages, and their impact on cephalosporin MICs, have not been investigated systematically. Here, our aims were to identify and characterize PBP substitutions in *C. difficile*, determine the extent to which they

cooccur with fluoroquinolone resistance in epidemic *C. difficile* lineages, and date this phylogenetically. Additionally, we aimed to assess the phenotype of both PBP-substituted strains and closely related wild-type controls in terms of their cephalosporin MICs.

RESULTS

The study was designed as follows. A globally distributed collection of published *C. difficile* genomes was assembled ($n = 7,094$) representing 16 genetic lineages, 14 commonly associated with CDI in health care settings and 2 carried asymptomatically (non-toxicogenic). The occurrence of recent, within-lineage PBP substitutions was investigated and compared to the occurrence of fluoroquinolone resistance. Cephalosporin (and other β -lactam) MICs were measured for representative strains containing PBP substitutions and closely related “wild-type” ancestors. Finally, the timing and sequence of PBP substitution and fluoroquinolone resistance acquisition events were investigated phylogenetically.

Lineages studied. The 7,094 genomes represented lineages ST1(ribotypes 027/198/176/181) ($n = 1,918$), ST17(018) ($n = 279$), ST42(106) ($n = 563$), ST3(001) ($n = 411$), ST37(017) ($n = 424$) and its recent descendant ST81(369) ($n = 39$), ST63(053) ($n = 37$), ST54(012) ($n = 148$), ST8(002) ($n = 593$) and its recent descendant ST183 ($n = 14$), ST11(078) ($n = 628$), ST2(014/020) ($n = 790$), ST6(005) ($n = 404$), ST10(015) ($n = 263$), ST7(026) ($n = 190$), ST56(058) ($n = 16$), and two prevalent nontoxicogenic genotypes, ST26(039) ($n = 175$) and ST15(010) ($n = 202$). Four genetically distinct *C. difficile* clades, 1, 2, 4, and 5, were represented (44). Genomes, accession numbers, AMR predictions from genotype and references are listed per lineage (see Data Set S1, sheets 1 to 18, in the supplemental material).

***C. difficile* PBPs.** To date, nine PBPs have been described within the *C. difficile* genome (Table 1), five of which are transpeptidases (PBP1 to PBP5 [PBP1–5]) (45). HMW PBP1 and PBP3 are essential for growth *in vitro* (46), while LMW PBP2 and PBP4 are not essential for growth but are required for sporulation (46). Only LMW PBP5 has been described as variably present (45). PBP1 to PBP4 were present in all genomes studied, while PBP5 occurred in lineages ST3(001), ST11(078), and ST37(017)/ST81(369). No additional PBPs were identified using known *C. difficile* PBP sequences in low-stringency BLAST searches.

Recent PBP substitutions. PBP gene sequences were compared within each lineage to identify recent SNPs. These were absent or rare in LMW PBP2, PBP4, and PBP5. In contrast, multiple SNPs occurred in HMW PBP1 and PBP3, almost all of which were nonsynonymous. The resultant amino acid substitutions affected a total of 48/993 (4.8%) positions in PBP3 and 50/855 to 925 (5.8 to 5.4%) positions in PBP1. Substitution data are shown per isolate (Data Set S1). The variable PBP1 size reflected differing numbers of repeats of the 14 amino-acid sequence, TPPDNGGNNGGGST, which occurred between 1 and 6 times near the C terminus of the protein (Table S1).

The frequency of each PBP amino acid substitution was recorded per lineage (Table 2). This identified the most common substitutions within and among lineages. In PBP3, V497L was most frequent, occurring in 2,897 genomes and 10 lineages, followed by A778V in 664 genomes and 10 lineages. In PBP1, substitution T674I/N/V was most frequent, occurring in 1,379 genomes of 10 lineages, followed by A555T in 442 genomes of 7 lineages. These data were plotted to visualize the relative positions and frequencies of substitutions within PBP1 and PBP3 (Fig. 1). Virtually all substitutions occurred within the conserved transpeptidase domains, flanking the active site motifs.

Association between PBP substitutions and fluoroquinolone resistance. In addition to PBP substitutions, the presence of fluoroquinolone resistance was investigated in all lineages (Fig. 2; Data Set S1). The occurrence of PBP substitutions was significantly associated with fluoroquinolone resistance in 8 of the 14 clinically important lineages: ST1(027/198/176/181), ST17(018), ST42(106), ST3(001), ST37(017)/ST81(369), ST63(053), ST54(012), and ST8(002)/ST183 (Fig. 2, $P < 0.001$), suggesting the possibility of almost simultaneous acquisition. There was no evidence of association for the remaining six clinically important lineages, ST11(078) ($P = 1.00$), ST2(014/020) ($P = 0.281$), and ST6(005)

TABLE 1 PBPs and β -lactamases of *C. difficile* reference genomes^{a,i}

Strain	Strain CD630 locus	Alternate designation	PBP classification	Size (aa)	blastp-predicted function/family	blastp E value ^h
R20291 locus NC_013316.1	NC_009089.1 (former locus)					
0712 ^b	RS04495 (07810) ^c	PBP1 ^g	HMW class A	897	Bifunctional transglycosylase/transpeptidase pbp_1A_fam	2.14e-166
0985 ^b	RS06420 (11480) ^c	PBP3 ^g	HMW class B	992	Transpeptidase Pbp2_mrdA for cell elongation	1.53e-127
1067 ^c	RS06830 (12290)	PBP2 ^g	LMW class B	554	FtsI/Pbp2 Transpeptidase Pbp2_mrdA for cell elongation	2.85e-101 1.81e-93
2544 ^c	RS14215 (26560) ^c	PBP4 ^g spoVD	LMW class B	659	spoVD_pbp transpeptidase	0e+00
-	-	PBP5 ^g strain M68	LMW class B	696	FtsI/Pbp2 Pbp2_mrdA transpeptidase	3.23e-165 1.00e-110 2.67e-104
1131 ^c	RS07160 (12910) ^c	<i>dacF</i>	LMW class C	387	D-Alanyl-D-alanine carboxypeptidase	1.68e-116
2048 ^c	RS11615 (21410)		LMW class C	397	D-Alanyl-D-alanine carboxypeptidase	2.31e-88
0441 ^d	RS03150 (05150)		LMW class C	414	D-Alanyl-D-alanine carboxypeptidase	8.55e-94
2390 ^d	RS13415 (24980)	<i>dacF1</i>	LMW class C	429	D-Alanyl-D-alanine carboxypeptidase	7.59e-97
3056 ^c	RS17015 (31960)		PBP or β -lactamase?	340	blastp: CubicO group peptidase, β -lactamase class C (R20291 "put. PBP," CD630 "serine hydrolase")	3.06e-40
1318 ^d	RS08060 (14690)	<i>cwp20</i>	PBP or β -lactamase?	1013	blastp: β -lactamase, cell wall binding protein repeats (R20291 "put. PBP," "cell surface protein")	2.40e-55
2283	RS12870 (23930)			338	Transglycosylase domain-containing protein	1.37e-74
0399	RS02840 (04580)	blaCDD, CDD1/2	β -Lactamase	312	YbaI class D β -lactamase (37)	5.94e-51

^aGray shading indicates proteins containing transpeptidase domains. MDR reference strains contain the following PBP substitutions relative to the wild-type of the identical genotype: strain R20291, ST1(027) UK 2006 (96) contains PBP3 V497L; strain CD630, ST54(012) Switzerland 1982 (99) contains PBP1 T674I and PBP3 N537K; strain M68, ST37 (017) Ireland 2006 (99) contains PBP3 Y721C.

^bEssential for growth *in vitro* (46).

^cNot essential for growth *in vitro* but required for sporulation (46).

^dNot essential for growth *in vitro* (46).

^eExistence shown experimentally by mass spectrometry (104, 105).

^fPBP5 absent in R20291 and CD630 but present in M68 (NC_017175.1) chromosomal locus RS02615, coordinates 501965 to 504052.

^gPBP1-5 (45).

^hblastp E values for R20291 sequences: the closer the value is to zero, the more significant the match.

ⁱDashes indicate that PBP5 is absent from strain R20291 and strain CD630.

($P = 1.00$), with ST10(015), ST7(026), and ST56(058) containing no isolates with PBP substitutions ($P = 0.865$ for the group overall). Similarly, there was no evidence of association for the two nontoxigenic lineages, ST15(010) ($P = 0.426$) and ST26(039) (no isolates with fluoroquinolone resistance substitutions) ($P = 0.531$ for the group overall).

Among the 14 clinically important lineages studied, PBP substitutions occurred less often in the absence of fluoroquinolone resistance (Fig. 2; Data Set S1). Genomes in this category reached between 6 and 30% of the total examined only in lineages ST42(106), ST37(017), ST54(012), ST2(014/020), and ST6(005). Only two toxigenic lineages, ST37 (017) and ST11(078), were notable for fluoroquinolone resistance in the absence of PBP substitutions, at 20 and 12% of the total genomes examined, respectively (Fig. 2; Data Set S1). PBP substitutions occurred without fluoroquinolone resistance in over 90% of nontoxigenic lineage genomes ST26(039) and ST15(010) (Fig. 2; Data Set S1).

Association between PBP substitutions and β -lactam MICs. β -Lactam MICs were measured for isolates representing eight of the lineages studied. Four lineages, ST1 (027) (FQ-R1 and FQ-R2 [19]), ST3(001), ST42(106), and ST17(018), were chosen due to their clinical importance and high proportion of genomes containing both PBP substitutions and fluoroquinolone resistance (Fig. 2). Specific PBP substitution combinations were chosen for phenotyping on the basis of high frequency within available collections (Data Set S1). Individual isolate choices were based on the low numbers of SNP differences (Fig. 3A) between wild-type and PBP-substituted genomes. The remaining four lineages we phenotyped were ST10(015), ST6(005), ST56(058), and ST7(026). These were chosen because while their prevalence varied (Fig. 2), they all lacked PBP

[illegible][illegible]

substitutions and therefore represented additional wild-type controls. Certain PBP substitutions/combinations were identified only in genomes we obtained from publicly available sequence databases (for example, the Southeast Asian clades ST81(369) and ST183). For this reason, their phenotype could not be assessed.

Among the isolates phenotyped from lineages ST1(027) (FQ-R1 and FQ-R2 [19]), ST3(001), ST42(106), and ST17(018), 10 different individual PBP substitutions were represented, three in PBP3 and seven in PBP1 (Fig. 3A and B). These included the four substitutions identified most frequently in the study as a whole (Fig. 1A and B; Data Set S1; Fig. 3B). For ST1(027) FQ-R2, MICs were determined for 3 different PBP substitution combinations (1, 2, or 3 substitutions) versus the wild-type; for ST1(027) FQ-R1, MICs were determined for 2 PBP combinations (1 or 2 substitutions) versus the wild-type; for ST17(018), MICs were determined for 1 PBP combination (containing 5 substitutions) versus the wild-type; for ST3(001), MICs were determined for 3 PBP combinations (containing 1, 1, or 4 substitutions) versus the wild-type; and finally, for ST42(106), MICs were determined for 1 PBP substitution combination (containing 2 substitutions) versus the wild-type (Fig. 3A). For the lineages lacking PBP substitutions, the numbers of isolates phenotyped were as follows: ST10(015), $n = 11$; ST6(005), $n = 10$; ST56(058), $n = 4$; and ST7(026), $n = 5$ (Fig. 3A).

The number of PBP substitutions per isolate was significantly correlated with MIC for cephalosporins (for cephadrine, $r = 0.88$, $P < 0.001$; cefuroxime, $r = 0.76$, $P = 0.001$; cefotaxime, $r = 0.78$, $P < 0.001$) and slightly less strongly for carbapenems (for meropenem, $r = 0.65$, $P = 0.009$; imipenem, $r = 0.50$, $P = 0.059$). In comparison, there was no significant correlation for penicillins (amoxicillin: $r = 0.17$, $P = 0.55$; co-amoxiclav: $r = 0.33$, $P = 0.237$; piperacillin-tazobactam: $r = 0.39$, $P = 0.149$). Actual PBP substitutions and MICs are shown in Fig. 3A. The greatest increases in cephalosporin MICs were associated with the highest numbers of substitutions; for example, the cefuroxime MIC increased from 376 to 1,506 $\mu\text{g/mL}$ in ST3(001) (four substitutions) and ST17(018) (five substitutions). The cephadrine MIC increased from 36 to 239 $\mu\text{g/mL}$ in the latter. Intriguingly, the wild-type ancestors of these four PBP-substituted lineages had cefotaxime MICs which were still higher

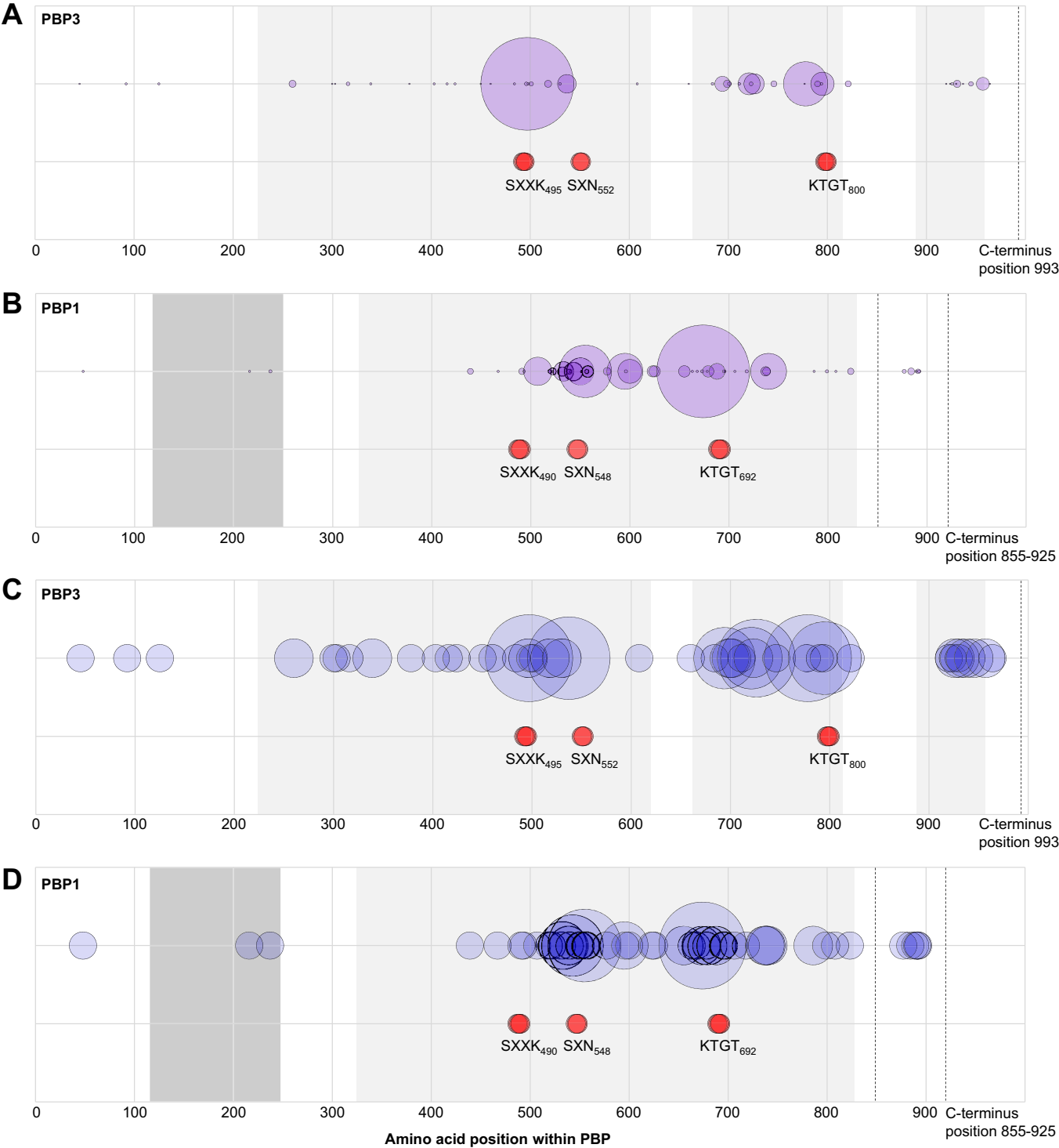


FIG 1 Positions and relative frequency of amino acid substitutions within PBP3 (993 amino acids) and PBP1 (855 to 925 amino acids). (A) Substitutions within PBP3 ($n = 48$) are represented by purple circles, plotted according to locations within the PBP (x axis). Circles are scaled according to substitution frequency within the entire data set. Light gray shading indicates the position of conserved transpeptidase domains identified by BLASTP. Red circles indicate transpeptidase catalytic motifs. (B) Same as described for panel A but for PBP1 (n substitutions = 50) and dark gray indicates the N-terminal glycosyl transferase domain. (C) Same as described for panel A except that relative sizes of blue circles indicate the number of lineages in which each substitution was identified. (D) Same as described for panel B, with blue circles again indicating the number of lineages in which each substitution was identified. Raw data, including the identity of each substitution, are shown in Table 2.

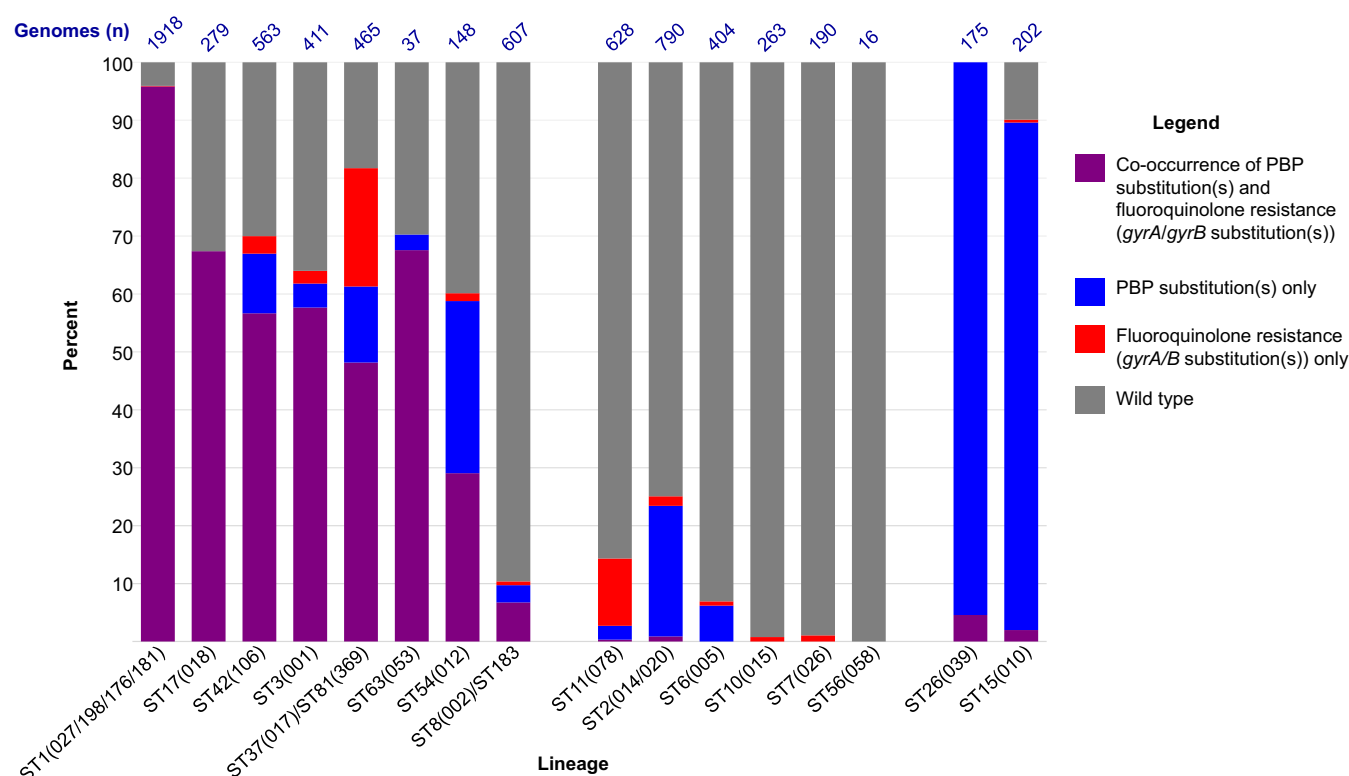


FIG 2 Occurrence of PBP substitutions and fluoroquinolone resistance in the 16 lineages studied.

than those of the four lineages which have not yielded PBP-substituted strains: cefuroxime, 128 $\mu\text{g/mL}$ versus 376 $\mu\text{g/mL}$, and cefotaxime, 128 $\mu\text{g/mL}$ versus 256 $\mu\text{g/mL}$.

Phylogenetic analyses. Recombination-corrected phylogenies were constructed to identify and date the order in which PBP substitutions and fluoroquinolone resistance were acquired by seven genetic lineages. These included the four PBP-substituted lineages which had been phenotyped [ST1(027), ST17(018), ST3(001), and ST42(106)] and a further three lineages containing notable MDR PBP-substituted fluoroquinolone-resistant strains. These were ST8(002)/ST183 and ST37(017)/ST81, both important in Southeast Asia, and ST54(012), which is notable in Costa Rica (47–51) (Fig. 4 to 7).

Irrespective of lineage, the sequence of PBP substitution acquisitions in epidemic strains typically started with PBP3 V497L and/or A778V (i.e., the most frequent PBP3 substitutions) (Table 2). Then, further PBP substitutions followed, yielding a variety of patterns. Among lineages well known for epidemic spread, the initial PBP3 V497L substitution occurred simultaneously with fluoroquinolone resistance (Fig. 4 to 7). One notable exception was the ST1(027) FQ-R1 lineage in which the PBP3 A726V substitution occurred first, while PBP3 V497L was absent (Fig. 4B).

PBP-substituted, fluoroquinolone-resistant clades evolved more than once in ST1(027), ST3(001), ST37(017)/ST81(369), ST42(106), and ST54(012) (Fig. 4A and B, 5B, 6B, and 7A and B). Their PBP substitution patterns each showed geographic structure (Fig. 4 to 8). MDR clades with the highest numbers of PBP substitutions were identified within ST17(018) in Italy and Southeast Asia (Fig. 5A), ST3(001) in United Kingdom/Germany (Fig. 5B), ST8(002)/ST183 in Japan (Fig. 6A), and ST37(017)/ST81(369) in Southeast Asia (Fig. 6B).

The occurrence of PBP substitutions in the absence of fluoroquinolone resistance was investigated phylogenetically in ST2(014/020) (Fig. 8) and to a lesser extent in ST42(106) and ST54(012) (Fig. 7). For ST2(014/020), a dated phylogeny was constructed using PBP-substituted genomes from six independent locations together with wild-type genomes from the same locations and dates (Fig. 8). The PBP-substituted genomes clustered by location, while the wild-type strains did not. It is not known whether the clusters

A

Lineage	Isolate	Location	SNPs	PBP Substitutions				Cephadrine (first)	Cefuroxime (second)	Cefotaxime (third)	Ampicillin	Co-amoxiclav	Meropenem	Imipenem	Piperacillin-tazobactam	
				PBP3	PBP1											
ST1(027)	TN145	Oxfordshire, UK		WT		WT		72	376	256	2	2	4	8	16	
ST1(027)-FQ-R2	1421	Oxfordshire, UK	30	V497L		WT		239	627	284	2	2	8	8	16	
ST1(027)-FQ-R2	1582	Oxfordshire, UK	28	V497L		T674I		239	537	284	2	2	16	8	32	
ST1(027)-FQ-R2	1503a	Oxfordshire, UK	29	V497L		T674I	A555T	239	627	284	2	2	16	8	32	
ST1(027)	OPT_1687	Calgary, Canada		WT		WT		72	376	256	2	2	4	8	16	
ST1(027) FQ-R1	OPT_1905	Columbus, USA	20	A726V		WT		143	752	365	2	2	8	8	32	
ST1(027) FQ-R1	OPT_2787	Detroit, USA	27	A726V		A555T		143	752	512	2	2	8	8	16	
ST17(018)	Oxf79	Oxfordshire, UK		WT		WT		36	376	256	1	1	4	8	8	
ST17(018)	OPT_2644	Italy	118	V497L	A778V	I550L	K595N	T740N	573	1506	>512	2	2	8	16	32
ST3(001)	2915	Oxfordshire, UK		WT		WT		36	376	256	1	1	4	8	8	
ST3(001)	Oxf746b	Oxfordshire, UK	162	A778V		WT		72	470	284	1	1	4	4	8	
ST3(001)	1172-p1	Oxfordshire, UK	273	V497L		WT		143	752	284	1	1	4	8	8	
ST3(001)	OPT_2456	Germany	134	V497L	A778V	T600I	A507T	239	1506	512	1	1	4	16	8	
ST42(106)	Oxf1499	Oxfordshire, UK		WT		WT		72	376	256	2	1	4	8	16	
ST42(106)	L.15.7921787	Leeds, UK	20	V497L		T674I		143	376	256	2	2	8	8	16	
Lineage	Isolates (n)			PBP3		PBP1										
ST10(015)	11	UK	N/A	WT		WT		NT	128	128	0.5 ¹ , 1 ¹⁰	1	8	4	8	
ST6(005)	10	UK	N/A	WT		WT		NT	128	128	1	1	8	4	8	
ST56(058)	4	UK	N/A	WT		WT		NT	128	128	1	1	8	4	8	
ST7(026)	5	UK	N/A	WT		WT		NT	128	64 ³ , 128 ²	0.5 ¹ , 1 ⁴	1	8	2 ² , 4 ³	8	

*An additional third generation cephalosporin was tested, ceftazidime, all isolates MIC 128µg/ml.

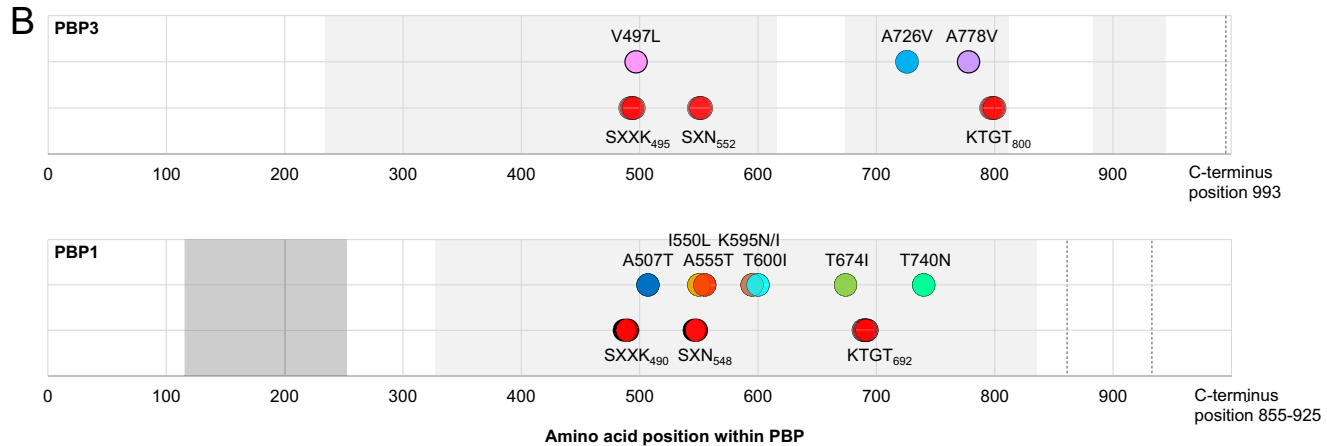
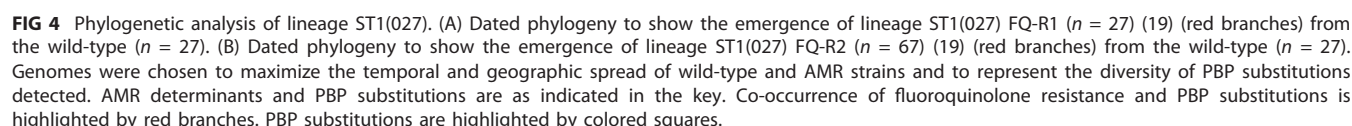


FIG 3 β -Lactam MICs of wild-type and PBP-substituted *C. difficile* isolates. (A) Upper panel, MICs for the β -lactams shown, measured for isolates belonging to the toxigenic MDR lineages ST1(027) (FQ-R2 and FQ-R1), ST17(018), ST3(001), and ST42(106), with one row per isolate tested. The column headed “SNPs” refers to the number of single nucleotide polymorphisms between each isolate and its wild-type (WT) within-lineage comparator. Identical PBP substitutions found in different isolates are highlighted by identical colors. The intensity of gray shading indicates the fold increase in MIC relative to the MIC of the wild type. Lower panel, MICs measured for the toxigenic non-MDR lineages ST10(015), ST6(005), ST56(058), and ST7(026), with one row per lineage. The number of isolates tested per lineage, if greater than 1, is indicated as a superscript. NT, not tested. (B) Positions of the PBP substitutions (colored as shown in panel A) contained in the isolates that underwent phenotyping (A), relative to the conserved transpeptidase domains (gray) and the active site motifs (red circles).

represented within-hospital nosocomial outbreaks; their small scale suggests that at the time they occurred, such outbreaks may have been difficult to distinguish from background sporadic cases. Interestingly, both ST42(106) and ST54(012) phylogenies (Fig. 7) contained clades in which PBP substitution acquisition preceded fluoroquinolone resistance. ST42(106) recently replaced ST1(027) as the most prevalent lineage in North America (8), but a single MDR clade was not apparent, with PBP3 V497L occurring on multiple independent occasions within the ST42(106) phylogeny.



DISCUSSION

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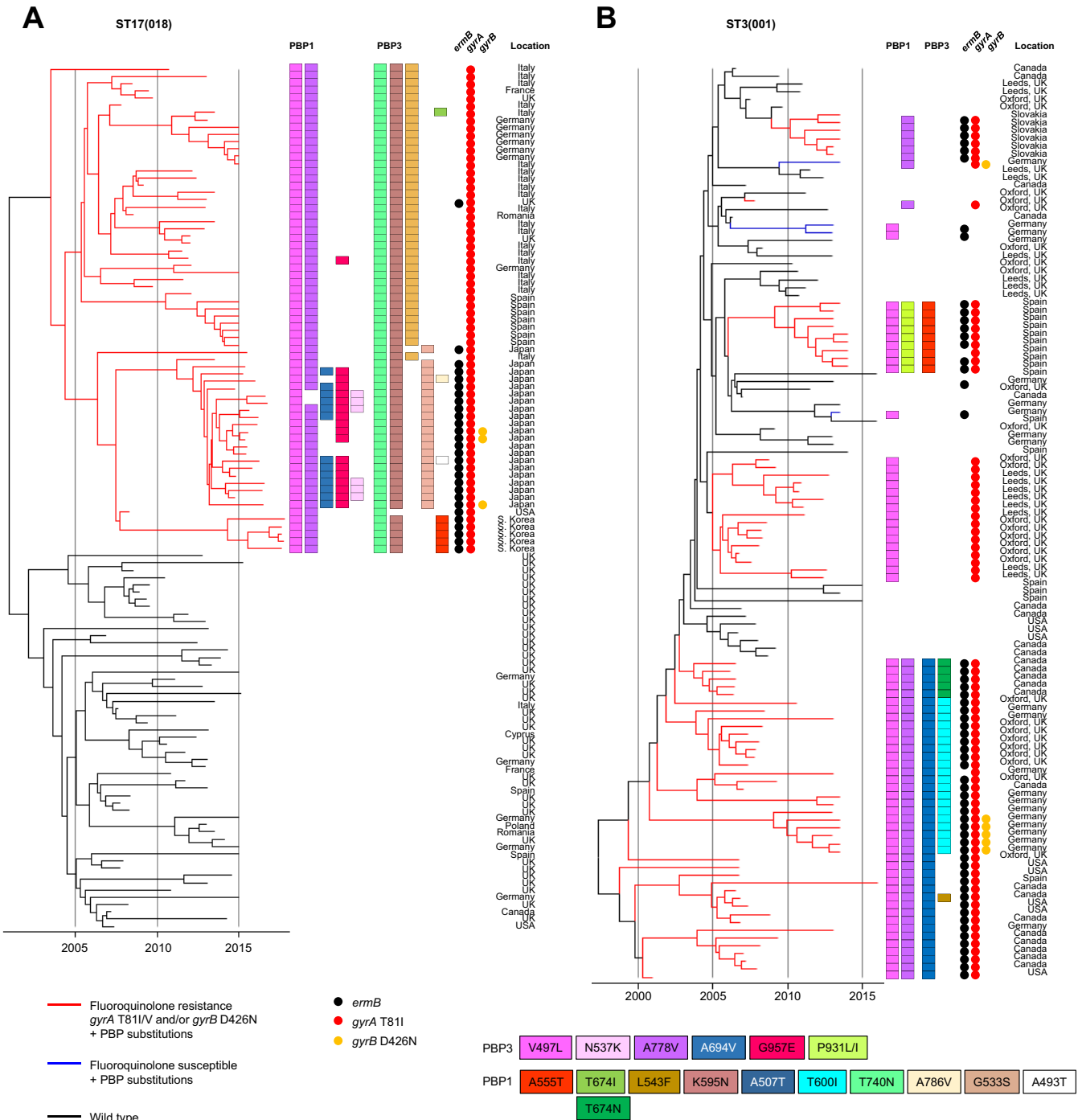
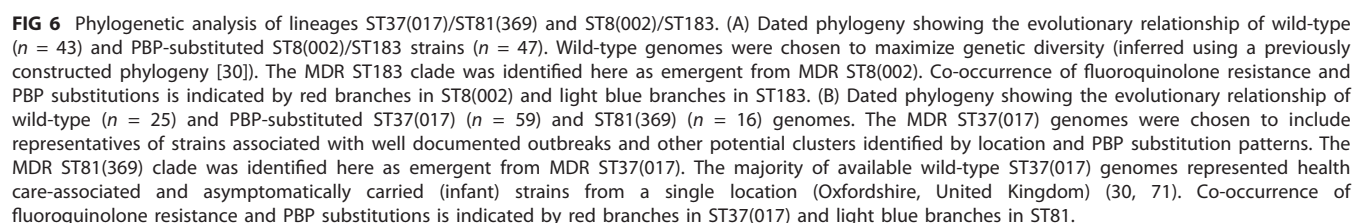
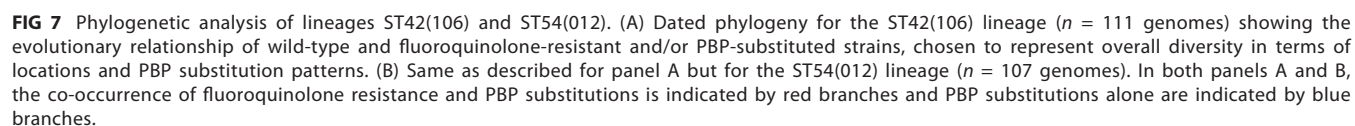


FIG 5 Phylogenetic analysis of lineages ST17(018) and ST3(001). (A) Dated phylogeny to show the emergence of MDR lineage ST17(018) ($n = 66$) (red branches) from the wild-type ($n = 53$). MDR strains from Europe, Southeast Asia, and North America were chosen to maximize the geographic spread and PBP substitutions. These and other AMR determinants and PBP substitutions are as indicated in the key. Co-occurrence of fluoroquinolone resistance and PBP substitutions is highlighted by red branches. (B) Dated phylogeny showing the evolutionary relationship of wild-type ($n = 40$) and PBP-substituted ST3 (001) genomes ($n = 77$).

have been reported only occasionally, associated with raised carbapenem MICs in a single lineage (45, 54). PBPs with reduced β -lactam affinity are clinically important in other Gram-positive pathogens, for example, *Streptococcus pneumoniae* (43) and methicillin-resistant *Staphylococcus aureus* (55). The present study was therefore performed to investigate systematically, and phenotypically, recent PBP substitutions among clinically important *C. difficile* lineages.





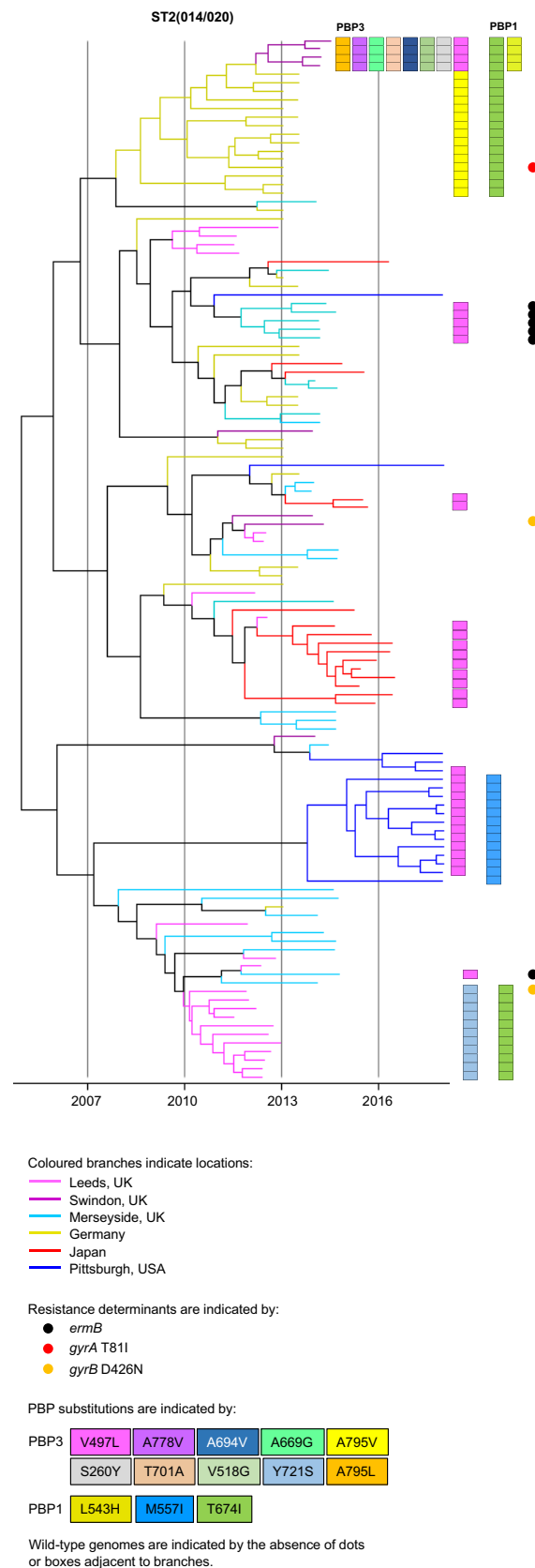


FIG 8 Phylogenetic analysis of ST2(014/020) genomes. Dated phylogeny showing the evolutionary relationship between wild-type ($n = 63$) and PBP-substituted ST2 (014/020) strains ($n = 60$) mostly lacking fluoroquinolone resistance. Branch color indicates one of six locations indicated in the key. Clustering of PBP-substituted genomes is compared with that of the wild-type for six independent geographic locations. Wild-type genomes from each location were collected concurrently with the PBP-substituted strains. These are indicated by the absence of AMR determinants (boxes or dots). Occurrence of AMR determinants is as defined in the key.

We identified multiple, recent PBP substitutions, which are focused in the conserved functional domains of the two HMW *C. difficile* transpeptidases, PBP1 and PBP3 (Table 2; Fig. 1). The locations of the substitutions suggest a possible response to β -lactam selective pressure, leading to changes which could (i) reduce β -lactam access to the transpeptidase active site and (ii) represent compensatory changes preserving the efficiency of cell wall biosynthesis. Further insights may be gained if the structures of these proteins are determined. Equivalent changes were not detected in LMW PBP2 and PBP4. This difference between the two PBP classes may reflect differences in function (46); for example, HMW PBP1 and PBP3 are essential for growth *in vitro*, while LMW PBP2 and PBP4 are not essential for growth but are required for sporulation (46). Although PBP5 transpeptidase was variably present, it was not recently acquired by MDR *C. difficile* lineages. Its constant chromosomal location suggests that gradual loss rather than recent acquisition may explain its variable presence.

PBP1 and PBP3 substitutions were significantly associated with raised cephalosporin MICs, relative to closely related wild-type strains, and the higher the number of substitutions, the higher the MIC (Fig. 3A). The mechanism underlying substitution acquisition was not recombination but rather the accumulation of *de novo* chromosomal mutations. This was indicated because virtually all SNPs were nonsynonymous, flanked the catalytic domains (Fig. 1), and arose multiple times (Table 2). Only two major recombination events were found involving ST81(369) (see Fig. S1 in the supplemental material) and ST1(181). While we have shown statistically significant associations between PBP substitutions and raised beta-lactam MICs in isolates separated from the wild-type by small numbers of SNPs (Fig. 3A), the demonstration of a causal association would require controlled laboratory allelic exchange experiments, which are beyond the scope of the present study.

The Co-occurrence of PBP substitutions and fluoroquinolone resistance in the clinically important epidemic lineages (Fig. 2) was statistically significant. Stewardship of cephalosporins may represent an additional tool for outbreak control, potentially mirroring the success of fluoroquinolone stewardship (30). In support of this, studies performed between the introduction of first-generation cephalosporins (mid-1960s) and the emergence of widespread fluoroquinolone resistance (late 1980s onwards) reported cephalosporin stewardship alone to be successful (24, 56–58). We found here that the cephalosporin MICs of wild-type strains differed according to whether they later went on to acquire PBP substitutions (Fig. 3A). For example, wild type cefuroxime and cefotaxime MICs were 128 μ g/mL when no PBP substitutions subsequently acquired, versus wild type MICs of 376 μ g/mL or 256 μ g/mL when PBP substitutions were subsequently acquired. The mechanism underlying these differences must predate PBP substitution acquisition and is currently unknown. It suggests that wild-type precursors of PBP-substituted clades are better adapted to cephalosporin exposure and therefore perhaps more likely to acquire PBP substitutions, as observed for the multiple clades detected phylogenetically in specific lineages. The co-occurrence of PBP substitutions and fluoroquinolone resistance currently hinders attempts to understand their relative importance in epidemic spread. Further controlled studies of cephalosporin and fluoroquinolone stewardship are needed.

Given the large number of PBP-substituted, clinically important MDR clades which have emerged over the last 30 years in different geographic regions (Fig. 4 to 8), it is surprising that their elevated cephalosporin MICs have not been highlighted previously. This likely reflects acceptance of the concept that high cephalosporin MICs (\sim 256 μ g/mL) are inherent to *C. difficile* (10, 33–36) and that their determination is consequently uninformative, particularly since wild-type MICs are thought to exceed clinically relevant concentrations and a mechanism linked to raised MICs has not been defined.

In addition to PBP substitutions, our phylogenies revealed the sequence and timing of MDR acquisition by clinically important lineages. The co-occurrence of PBP substitutions and fluoroquinolone resistance predated epidemic spread, which was reflected in short-branched, geographically structured clades (Fig. 4 to 7). The first PBP substitution was typically PBP3 V497L, followed by others, yielding a variety of final combinations.

The more highly PBP-substituted, fluoroquinolone-resistant clades were frequently positive for additional AMR determinants, particularly the *ermB* gene (clindamycin resistance) and *rpoB* substitutions (rifampin resistance) (Data Set S1; Fig. 5 and 6).

We dated the emergence of the two MDR ST1(027) clades (FQ-R1 and FQ-R2) to the mid- to late 1990s, as previously described (19) (Fig. 4A and B). The emergence date of the MDR ST17(018) clade is compatible with the first reports of outbreaks in 1996 to 1999 (59), the phylogeny root having a 95% credible interval dating of December 1998 to July 2002 (Fig. 5A). European and Asian ST17(018) clades then diverged, acquiring further region-specific PBP substitutions (Fig. 5A), arguing against their recent intercontinental spread.

Greater numbers of PBP substitutions were significantly associated with the highest cephalosporin MICs (Fig. 3A). Consistent with this, highly substituted clades of multiple lineages [ST17(018), ST81(369)/ST37(018), and ST183/ST8(002)] (Fig. 5A and 6A and B) predominated in Southeast Asia, where cephalosporin use is high (60–62). Adaptation to local prescribing conditions through PBP substitution acquisition offers a possible explanation for the temporal and geographic variation in prevalent *C. difficile* lineages (6, 8, 19, 47, 58, 63). For example, United Kingdom and local (Oxfordshire) cephalosporin prescribing levels from 1998 to 2013 were described previously (30) and were higher (as was fluoroquinolone prescribing) when epidemic lineages (ST1(027), ST3(001), and ST42(106)) predominated. This concept may extend to competitive exclusion of lesser PBP-substituted strains by more highly substituted ones, a scenario requiring greater numbers of PBP substitutions to carry a fitness cost. This appears possible, as in *Clostridium perfringens* *in vitro*, where PBP substitutions are associated with slower growth (64). We hypothesize that local levels of cephalosporin (and fluoroquinolone) prescribing determine the prevalent *C. difficile* MDR strain(s) in a given region. For example, the unusually low levels of ST1(027) seen in Asia (65) may reflect competitive exclusion, under local prescribing conditions, by the more highly PBP-substituted clades which predominate here. The relative geographic restriction of ST1(027) FQ-R1 (in the United States, South Korea, and Germany) in comparison to the more globally distributed FQ-R2 may also reflect the different PBP substitutions of the two clades (Fig. 4) and variations in MIC (Fig. 3A).

MDR strains exhibit high transmissibility in clinical settings when prescribing is uncontrolled (17). The PBP1 and PBP3 transpeptidases function in cell wall biosynthesis, and therefore, substitutions impacting their catalytic domain could affect transmissibility via sporulation. A high sporulation phenotype has been reported in at least two epidemic lineages: ST3(001) (United Kingdom) and ST81(369) (Asia) (66, 67). Sporulation phenotype is reportedly variable in ST1(027) (68, 69), and we have observed variation in its PBP substitutions (Fig. 4A and B; Data Set S1). However, the possibility of a link remains to be investigated. It is relevant to future experimental design that the MDR laboratory reference strains CD630 [ST54(012)] and R20291 [ST1(027)] both contain PBP substitutions (Table 1).

PBP substitutions occurred without fluoroquinolone resistance in a minority of the toxigenic lineages studied: ST42(106), ST37(017), ST54(012), ST2(014/020), and ST6(005) (Fig. 2; Data Set S1). ST42(106) was of interest since it recently exceeded the prevalence of ST1(027) in North America (8, 70). Visual inspection of phylogenies was used to assess whether PBP substitutions might enhance transmissibility in the absence of fluoroquinolone resistance. An ST2(020/014) phylogeny showed some possibility of locally enhanced transmission (Fig. 8), as did ST54(012) and ST42(106) (U.S. genomes) (Fig. 7A and B). However, these events were small scale, and this question remains to be answered. PBP substitutions (without fluoroquinolone resistance) were, however, widespread within the two nontoxigenic lineages ST15(010) and ST26(039) (Table S1), potentially explaining their high prevalence over other nontoxigenic strains. As well as being isolated coincidentally from patients with CDI caused by toxigenic strains, they are frequently carried by human infants (71) and potentially pets and farm/wild animals. The niches they colonize are likely to be more extensive than, and distinct from, the MDR toxigenic lineages, leading to potential opportunities for β -lactam exposure in the absence of exposure to fluoroquinolones, for example, in community or veterinary settings or in the environment. In contrast, the MDR toxigenic lineages containing

both PBP substitutions and fluoroquinolone resistance-associated SNPs exist in a much narrower ecological niche, persisting only in health care settings when prescribing of β -lactams and/or fluoroquinolones is uncontrolled (30). Examples of isolates containing PBP substitutions but lacking other AMR determinants were not included among those we phenotyped. This limitation is mitigated to some extent by the fact that such genomes do have specific PBP substitutions in common with the phenotyped strains, such as PBP3 V497L, but would be interesting to examine in the future.

The cephalosporin MICs for PBP-substituted strains were extremely high for certain antibiotics (for example, $>512 \mu\text{g/mL}$ for cefotaxime and up to $1,506 \mu\text{g/mL}$ for cefuroxime) (Fig. 3A). This raises questions about the *in vivo* conditions required for PBP substitution selection. Intravenous β -lactams are eliminated in active form by biliary excretion, resulting in highly variable intestinal concentrations (72). Intestinal concentrations ranging from 1.01 to $1,345 \mu\text{g/mL}$ have been reported (73), and so the potential exists for *C. difficile* to be exposed *in vivo* to cephalosporin concentrations reaching the MICs measured here. Bacteria with raised MICs can also be selected experimentally at antimicrobial concentrations up to several hundred-fold below lethal levels. However, the overall contribution made by such “sub-MIC selection” to resistance in clinically important bacteria is unknown (74–79).

To date, $<1\%$ of known *C. difficile* lineages (1,042 STs identified as at 20 January 2023; <https://pubmlst.org/organisms/clostridioides-difficile/>) have evolved a PBP-substituted clade(s). Furthermore, each lineage has tended to evolve more than one such clade (Fig. 4 to 7). This suggests that the wild-type phenotype of such lineages may favor the acquisition of chromosomal SNPs which are associated with raised cephalosporin MICs. As discussed above, the wild-type cephalosporin MICs for PBP-substituted lineages were indeed higher than for those lacking such strains (Fig. 3A). This potentially favors survival of these lineages in low cephalosporin concentrations *in vivo*, which could allow selection of PBP substitutions to occur. Since such strains also frequently contain additional SNPs associated with MDR, an alternative or additional mechanism might be a hypermutator phenotype, as in *S. pneumoniae* (74). An interesting area for further work could be a genome-wide association study to detect SNPs associated with raised cephalosporin MICs, an approach already employed to identify three novel mutations associated with reduced susceptibility to metronidazole (80).

In summary, our findings identify a potential role for cephalosporin selection in the evolution of epidemic CDI lineages. Specific regional prescribing practices may determine the locally predominant epidemic strains, potentially explaining the marked international variation in *C. difficile* molecular epidemiology. Since antimicrobial stewardship typically targets multiple drug classes (29, 81) and epidemic strains have raised MICs for fluoroquinolones, cephalosporins (Fig. 2 and 3), and more variably clindamycin (*ermB*) (Data Set S1), it is difficult to determine the relative contributions made by stewardship of each drug to CDI control. The timing of cephalosporin and fluoroquinolone resistance acquisition, immediately before the emergence of multiple epidemic strains from divergent *C. difficile* genetic backgrounds, suggests that AMR may be equally important as, or even exceed, strain-specific virulence determinants in driving epidemic CDI.

MATERIALS AND METHODS

Whole-genome sequencing (WGS) from 7,094 *C. difficile* isolates, predominantly cultured from humans with CDI, was obtained. Clinical isolates from hospital and community patients from Europe, North and South America, Southeast Asia, and Australia were included. Fourteen CDI lineages and two nontoxigenic lineages were represented. A complete list of genomes, their identifiers in public databases, and references are provided (see Data Set S1 in the supplemental material). Raw sequence reads were assembled *de novo* as required using Velvet (version 1.0.7 to 1.0.18) (82) and VelvetOptimiser with default settings (2.1.7) (83). A minority of genomes were obtained assembled, either from the NCBI database (<https://www.ncbi.nlm.nih.gov/genome/browse#!/prokaryotes/535/>) or Enterobase (<https://enterobase.warwick.ac.uk/species/index/clostridium>) (84). Assemblies were imported to a BIGSdb database (85) which was used to identify the seven loci used in multilocus sequence typing (44). Sequence types (STs) were assigned using the *C. difficile* PubMLST database (<https://pubmlst.org/organisms/clostridioides-difficile/>). STs and PCR ribotypes were used to indicate genetic lineages, identified by, e.g., the notation ST1(027) [sequence type 1 (PCR ribotype 027)].

BLAST searches performed within BIGSdb (85) were used to identify and extract chromosomal gene sequences for PBP transpeptidases (PBP1–5) (45), together with *gyrA*, *gyrB*, and *rpoB*, specific mutations in which confer AMR. Established amino acid substitutions scored as conferring resistance to fluoroquinolones were *GyrA* T811 and *GyrB* D426N, and those scored as conferring resistance to rifampin were *RpoB* R505K, H502N, and S498T. Acquisition of *ermB*, conferring clindamycin resistance, was also noted (86–89). Each unique allele sequence identified at these loci (PBP1–5, *gyrA*, *gyrB*, *rpoB*, and *ermB*) was assigned a number (Data Set S1) and can be downloaded at <https://pubmlst.org/organisms/clostridioides-difficile/> (44, 85). Newly extracted gene sequences were queried against this database, and the allele numbers were recorded for each genome together with the substitutions relevant to AMR (Data Set S1).

Identification of recent PBP substitutions. Identification of recent PBP substitutions was achieved using MEGA (<https://www.megasoftware.net/>) (90), which facilitated within-lineage comparisons of the nucleotide and amino acid sequences of PBP1–5 alleles. Comparisons were made relative to the wild-type PBP sequence for each lineage, wild-type alleles being taken from non-MDR genomes within the lineage.

Phenotyping. Isolates were chosen for phenotyping from a total of eight lineages, four containing both PBP-substituted and wild-type strains and four containing wild-type strains only. To control for any potential confounding due to population structure, the isolates containing PBP substitutions were chosen to minimize SNP distances from their equivalent wild-type (Fig. 3A). The following numbers of isolates were phenotyped per lineage: ST1(027) FQ-R1 and FQ-R2, total phenotyped = 7, comprising 2 wild-type and 5 PBP substituted; ST3(001), total phenotyped = 4, comprising 1 wild-type and 3 PBP substituted; ST17(017), total phenotyped = 2, comprising 1 wild-type and 1 PBP substituted; and ST42(106), total phenotyped = 2, comprising 1 wild-type and 1 PBP substituted. Among the wild-type-only lineages, the following were phenotyped: ST10(015) (*n* = 11), ST6(005) (*n* = 10), ST56(058) (*n* = 4), and ST7(026) (*n* = 5).

MICs of cefotaxime, cefuroxime, cephadrine, amoxicillin, amoxicillin-clavulanate, meropenem, imipenem, and piperacillin-tazobactam were determined by Wilkins-Chalgren agar dilution methods (91, 92). Briefly, *C. difficile* isolates and controls (*C. difficile* ATCC 700057, *C. difficile* E4 [PCR ribotype 010] and *Bacteroides fragilis* ATCC 25285) were cultured in prereduced Schaedler anaerobic broths at 37°C for 24 h anaerobically. Isolates and controls were diluted in prereduced saline to a McFarland standard 1 equivalence and multipoint inoculated onto prepared antibiotic-containing agar plates and controls. Agar plates were incubated at 37°C for 24 h, anaerobically, prior to MIC determination. The MIC was defined as the lowest concentration of antimicrobial that completely inhibited growth, showed a marked reduction in growth, showed only 1 or 2 colonies, or left a faint haze of growth on the plate, according to Clinical and Laboratory Standards Institute (CLSI) guidelines (93).

Antimicrobial concentrations were prepared using solvents and diluents recommended in the CLSI guidelines (93, 94). For amoxicillin-clavulanate and piperacillin-tazobactam, clavulanic acid and tazobactam were added to agar at fixed concentrations of 2 mg/L and 4 mg/L, respectively. In order to test susceptibility within normal doubling dilutions, additional antibiotic concentrations were prepared for the antibiotic plate range. All antibiotics were tested at the following dilutions: 0.125, 0.25, 0.5, 1, 2, 4, 8, 16, 32, 36, 40, 46, 53, 64, 71, 80, 91, 107, 128, 142, 160, 182, 213, 256, 284, 320, 366, 427, and 512 mg/L. Additionally, cefuroxime and cephadrine were prepared up to their limit of solubility, and the following ranges of dilutions were prepared: for cefuroxime, 102, 120, 143, 160, 205, 213, 240, 287, 319, 409, 478, and 572 mg/L; for cephadrine, 105, 118, 134, 157, 188, 209, 235, 269, 314, 376, 418, 471, 538, 627, 753, 837, 941, 1,076, 1,255, and 1,506 mg/L.

Statistical methods. The association between the presence of PBP substitutions and the presence of fluoroquinolone resistance substitutions was tested using a two-tailed Fisher's exact test. This was done for each lineage containing at least one isolate with PBP and/or fluoroquinolone resistance substitutions. Lineages were further combined into three groups: (i) clinically important, toxigenic lineages with evidence of multidrug resistance, (ii) other toxigenic lineages lacking MDR, and (iii) nontoxigenic lineages. The correlation between the number of PBP substitutions and β -lactam MICs was calculated for each antibiotic, using Spearman's rank correlation coefficient. This was done for all isolates in lineages containing PBP substitutions [ST1(027), ST17(018), ST3(001), and ST42(106)].

Construction of dated phylogenies. Dated phylogenies were constructed using genomes chosen to maximize the geographic and temporal spread of wild-type and AMR strains, as well as to represent the diversity of PBP substitutions detected in each lineage. Each set of genomes was first aligned to a reference using MuMMER version 3.1 (95) to produce a genome-wide alignment. The following genomes were used as references; ST1(027) strain R20291, accession no. [FN545816](#) (96) (Fig. 4); ST17(018) strain CBA7209, accession no. [QLOB00000000](#); (97) (Fig. 5A); ST3(001) strain B19, accession no. [FN668944.1](#) (96) (Fig. 5B); ST8(002) and descendant ST183 strain W0003a, accession no. [CP025047.1](#) (98) (Fig. 6A); ST37(017) and descendant ST81(369) strain M68, accession no. [NC_017175.1](#) (96) (Fig. 6B); ST42(106) strain W0023a, accession no. [CP025045.1](#), (98) (Fig. 7A); ST54(012) strain CD630, accession no. [AM180355.1](#) (99) (Fig. 7B); and ST2(014/020) strain W0022a, accession no. [CP025046.1](#) (98) (Fig. 8).

Initial phylogenies, built using PhyML version 3.3 (100), were then corrected for recombination using ClonalFrameML version 1.12 (101). Finally, these phylogenies were dated using BactDating version 1.1 (102) by assuming a mean evolutionary rate of 1.4 mutations per year per genome as in previous similar studies (103).

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

DATA SET S1, XLSX file, 1.6 MB.

FIG S1, PDF file, 0.6 MB.

TABLE S1, DOCX file, 0.02 MB.

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REFERENCES

- Smits WK, Lyras D, Lacy DB, Wilcox MH, Kuijper EJ. 2016. *Clostridium difficile* infection. Nat Rev Dis Primers 2:16020. <https://doi.org/10.1038/nrdp.2016.20>.
- McDonald LC, Killgore GE, Thompson A, Owens RC, Jr, Kazakova SV, Sambol SP, Johnson S, Gerding DN. 2005. An epidemic, toxin gene-variant strain of *Clostridium difficile*. N Engl J Med 353:2433–2441. <https://doi.org/10.1056/NEJMoa051590>.
- Zaiss NH, Witte W, Nübel U. 2010. Fluoroquinolone resistance and *Clostridium difficile*, Germany. Emerg Infect Dis 16:675–677. <https://doi.org/10.3201/eid1604.090859>.
- Barbanti F, Spigaglia P. 2016. Characterization of *Clostridium difficile* PCR-ribotype 018: a problematic emerging type. Anaerobe 42:123–129. <https://doi.org/10.1016/j.anaerobe.2016.10.003>.
- Imwattana K, Knight DR, Kullin B, Collins DA, Putsathit P, Kiratisin P, Riley TV. 2020. Antimicrobial resistance in *Clostridium difficile* ribotype 017. Expert Rev Anti Infect Ther 18:17–25. <https://doi.org/10.1080/14787210.2020.1701436>.
- Freeman J, Vernon J, Pilling S, Morris K, Nicolson S, Shearman S, Clark E, Palacios-Fabrega JA, Wilcox M, Pan-European Longitudinal Surveillance of Antibiotic Resistance among Prevalent *Clostridium difficile* Ribotypes' Study Group. 2020. Five-year Pan-European, longitudinal surveillance of *Clostridium difficile* ribotype prevalence and antimicrobial resistance: the extended ClosER study. Eur J Clin Microbiol Infect Dis 39:169–177. <https://doi.org/10.1007/s10096-019-03708-7>.
- Lew T, Putsathit P, Sohn KM, Wu Y, Ouchi K, Ishii Y, Tateda K, Riley TV, Collins DA. 2020. Antimicrobial susceptibilities of *Clostridium difficile* isolates from 12 Asia-Pacific countries in 2014 and 2015. Antimicrob Agents Chemother 64:e00296–20. <https://doi.org/10.1128/AAC.00296-20>.
- Carlson TJ, Blasingame D, Gonzales-Luna AJ, Alnezary F, Garey KW. 2020. *Clostridioides difficile* ribotype 106: a systematic review of the antimicrobial susceptibility, genetics, and clinical outcomes of this common worldwide strain. Anaerobe 62:102142. <https://doi.org/10.1016/j.anaerobe.2019.102142>.
- Owens RC, Jr, Donskey CJ, Gaynes RP, Loo VG, Muto CA. 2008. Antimicrobial-associated risk factors for *Clostridium difficile* infection. Clin Infect Dis 46(Suppl 1):S19–S31. <https://doi.org/10.1086/521859>.
- Gerding DN. 2004. Clindamycin, cephalosporins, fluoroquinolones, and *Clostridium difficile*-associated diarrhea: this is an antimicrobial resistance problem. Clin Infect Dis 38:646–648. <https://doi.org/10.1086/382084>.
- Muto CA, Pokrywka M, Shutt K, Mendelsohn AB, Nouri K, Posey K, Roberts T, Croyle K, Krystofiak S, Patel-Brown S, Pasculle AW, Paterson DL, Saul M, Harrison LH. 2005. A large outbreak of *Clostridium difficile*-associated disease with an unexpected proportion of deaths and colectomies at a teaching hospital following increased fluoroquinolone use. Infect Control Hosp Epidemiol 126:273–280. <https://doi.org/10.1086/502539>.
- Johnson S, Samore MH, Farrow KA, Killgore GE, Tenover FC, Lyras D, Rood JI, DeGirolami P, Baltch AL, Rafferty ME, Pear SM, Gerding DN. 1999. Epidemics of diarrhea caused by a clindamycin-resistant strain of *Clostridium difficile* in four hospitals. N Engl J Med 341:1645–1651. <https://doi.org/10.1056/NEJM199911253412203>.
- Loo VG, Poirier L, Miller MA, Oughton M, Libman MD, Michaud S, Bourgault AM, Nguyen T, Frenette C, Kelly M, Vibien A, Brassard P, Fenn S, Dewar K, Hudson TJ, Horn R, René P, Monczak Y, Dascal A. 2005. A predominantly clonal multi-institutional outbreak of *Clostridium difficile*-associated diarrhea with high morbidity and mortality. N Engl J Med 353:2442–2449. <https://doi.org/10.1056/NEJMoa051639>.
- Arvand M, Hauri AM, Zaiss NH, Witte W, Bettge-Weller G. 2009. *Clostridium difficile* ribotypes 001, 017, and 027 are associated with lethal *C. difficile* infection in Hesse, Germany. Euro Surveill 14:19403. <https://doi.org/10.2807/es.14.45.19403-en>.
- Goorhuis A, Debast SB, Dutilh JC, van Kinschot CM, Harmanus C, Cannegieter SC, Hagen EC, Kuijper EJ. 2011. Type-specific risk factors and outcome in an outbreak with two different *Clostridium difficile* types simultaneously in one Hospital. Clin Infect Dis 53:860–869. <https://doi.org/10.1093/cid/cir549>.
- Walker AS, Eyre DW, Wyllie DH, Dingle KE, Griffiths D, Shine B, Oakley S, O'Connor L, Finney J, Vaughan A, Crook DW, Wilcox MH, Peto TE, Infections in Oxfordshire Research Database. 2013. Relationship between bacterial strain type, host biomarkers, and mortality in *Clostridium difficile* infection. Clin Infect Dis 56:1589–1600. <https://doi.org/10.1093/cid/cit127>.
- Baldan R, Trovato A, Bianchini V, Biancardi A, Cichero P, Mazzotti M, Nizzero P, Moro M, Ossi C, Scarpellini P, Cirillo DM. 2015. *Clostridium difficile* PCR ribotype 018, a successful epidemic genotype. J Clin Microbiol 53:2575–2580. <https://doi.org/10.1128/JCM.00533-15>.
- Serafino S, Consonni D, Migone De Amicis M, Sisto F, Domeniconi G, Formica S, Zarantonello M, Maraschini A, Cappellini MD, Spigaglia P, Barbanti F, Castaldi S, Fabio G. 2018. Clinical outcomes of *Clostridium difficile* infection according to strain type. A prospective study in medical wards. Eur J Intern Med 54:21–26. <https://doi.org/10.1016/j.ejim.2018.03.015>.
- He M, Miyajima F, Roberts P, Ellison L, Pickard DJ, Martin MJ, Connor TR, Harris SR, Fairley D, Bamford KB, D'Arc S, Brazier J, Brown D, Coia JE, Douce G, Gerding D, Kim HJ, Koh TH, Kato H, Senoh M, Louie T, Michell S, Butt E, Peacock SJ, Brown NM, Riley T, Songer G, Wilcox M, Pirmohamed M, Kuijper E, Hawkey P, Wren BW, Dougan G, Parkhill J, Lawley TD. 2013. Emergence and global spread of epidemic healthcare-associated *Clostridium difficile*. Nat Genet 45:109–113. <https://doi.org/10.1038/ng.2478>.
- Senoh M, Haru Kato H. 2022. Molecular epidemiology of endemic *Clostridioides difficile* infection in Japan. Anaerobe 74:102510. <https://doi.org/10.1016/j.anaerobe.2021.102510>.
- Hensgens MP, Goorhuis A, van Kinschot CM, Crobach MJ, Harmanus C, Kuijper EJ. 2011. *Clostridium difficile* infection in an endemic setting in the Netherlands. Eur J Clin Microbiol Infect Dis 30:587–593. <https://doi.org/10.1007/s10096-010-1127-4>.
- Eyre DW, Davies KA, Davis G, Fawley WN, Dingle KE, De Maio N, Karas A, Crook DW, Peto TEA, Walker AS, Wilcox MH, EUCLID Study Group. 2018. Two distinct patterns of *Clostridium difficile* diversity across Europe indicating contrasting routes of spread. Clin Infect Dis 67:1035–1044. <https://doi.org/10.1093/cid/ciy252>.

23. Pear SM, Williamson TH, Bettin KM, Gerding DN, Galgiani JN. 1994. Decrease in nosocomial *Clostridium difficile*-associated diarrhea by restricting clindamycin use. *Ann Intern Med* 120:272–277. <https://doi.org/10.7326/0003-4819-120-4-199402150-00003>.
24. McNulty C, Logan M, Donald IP, Ennis D, Taylor D, Baldwin RN, Bannerjee M, Cartwright KA. 1997. Successful control of *Clostridium difficile* infection in an elderly care unit through use of a restrictive antibiotic policy. *J Antimicrob Chemother* 40:707–711. <https://doi.org/10.1093/jac/40.5.707>.
25. Valiquette L, Cossette B, Garant MP, Diab H, Pépin J. 2007. Impact of a reduction in the use of high-risk antibiotics on the course of an epidemic of *Clostridium difficile*-associated disease caused by the hypervirulent NAP1/027 strain. *Clin Infect Dis* 45:S112–S121. <https://doi.org/10.1086/519258>.
26. Debast SB, Vaessen N, Choudry A, Wieggers-Ligtvoet EAJ, van den Berg RJ, Kuijper EJ. 2009. Successful combat of an outbreak due to *Clostridium difficile* PCR ribotype 027 and recognition of specific risk factors. *Clin Microbiol Infect* 15:427–434. <https://doi.org/10.1111/j.1469-0691.2009.02713.x>.
27. Feazel LM, Malhotra A, Perencevich EN, Kaboli P, Diekema DJ, Schweizer ML. 2014. Effect of antibiotic stewardship programmes on *Clostridium difficile* incidence: a systematic review and meta-analysis. *J Antimicrob Chemother* 69:1748–1754. <https://doi.org/10.1093/jac/dku046>.
28. Sarma JB, Marshall B, Cleeve V, Tate D, Oswald T, Woolfrey S. 2015. Effects of fluoroquinolone restriction (from 2007 to 2012) on *Clostridium difficile* infections: interrupted time-series analysis. *J Hosp Infect* 91:74–80. <https://doi.org/10.1016/j.jhin.2015.05.013>.
29. Muto CA, Blank MK, Marsh JW, Vergis EN, O'Leary MM, Shutt KA, Pasculle AW, Pokrywka M, Garcia JG, Posey K, Roberts TL, Potoski BA, Blank GE, Simmons RL, Veldkamp P, Harrison LH, Paterson DL. 2007. Control of an outbreak of infection with the hypervirulent *Clostridium difficile* BI strain in a university hospital using a comprehensive “bundle” approach. *Clin Infect Dis* 45:1266–1273. <https://doi.org/10.1086/522654>.
30. Dingle KE, Didelot X, Quan TP, Eyre DW, Stoesser N, Golubchik T, Harding RM, Wilson DJ, Griffiths D, Vaughan A, Finney JM, Wyllie DH, Oakley SJ, Fawley WN, Freeman J, Morris K, Martin J, Howard P, Gorbach S, Goldstein EJC, Citron DM, Hopkins S, Hope R, Johnson AP, Wilcox MH, Peto TEA, Walker AS, Crook DW, Modernising Medical Microbiology Informatics Group. 2017. Effects of control interventions on *Clostridium difficile* infection in England: an observational study. *Lancet Infect Dis* 17:411–421. [https://doi.org/10.1016/S1473-3099\(16\)30514-X](https://doi.org/10.1016/S1473-3099(16)30514-X).
31. Lawes T, Lopez-Lozano JM, Nebot CA, Macartney G, Subbarao-Sharma R, Wares KD, Sinclair C, Gould IM. 2017. Effect of a national 4C antibiotic stewardship intervention on the clinical and molecular epidemiology of *Clostridium difficile* infections in a region of Scotland: a non-linear time-series analysis. *Lancet Infect Dis* 17:194–206. [https://doi.org/10.1016/S1473-3099\(16\)30397-8](https://doi.org/10.1016/S1473-3099(16)30397-8).
32. Huang H, Weintraub A, Fang H, Nord CE. 2009. Antimicrobial resistance in *Clostridium difficile*. *Int J Antimicrob Agents* 34:516–522. <https://doi.org/10.1016/j.ijantimicag.2009.09.012>.
33. Dzik J, Bartlett JG. 1980. In vitro susceptibility of *Clostridium difficile* isolates from patients with antibiotic-associated diarrhea or colitis. *Antimicrob Agents Chemother* 17:695–698. <https://doi.org/10.1128/AAC.17.4.695>.
34. Shuttlesworth R, Taylor M, Jones DM. 1980. Antimicrobial susceptibilities of *Clostridium difficile*. *J Clin Pathol* 33:1002–1005. <https://doi.org/10.1136/jcp.33.10.1002>.
35. Greenfield RA, Kurzynski TA, Craig WA. 1982. In vitro susceptibility of *Clostridium difficile* isolates to cefotaxime, moxalactam, and cefoperazone. *Antimicrob Agents Chemother* 21:846–847. <https://doi.org/10.1128/AAC.21.5.846>.
36. Chow AW, Cheng N, Bartlett KH. 1985. In vitro susceptibility of *Clostridium difficile* to new beta-lactam and quinolone antibiotics. *Antimicrob Agents Chemother* 28:842–844. <https://doi.org/10.1128/AAC.28.6.842>.
37. Toth M, Stewart NK, Smith C, Vakulenko SB. 2018. Intrinsic class D β -lactamases of *Clostridium difficile*. *mBio* 9:e01803-18. <https://doi.org/10.1128/mBio.01803-18>.
38. Sandhu BK, Edwards AN, Anderson SE, Woods EC, McBride SM. 2019. Regulation and anaerobic function of the *Clostridioides difficile* β -lactamase. *Antimicrob Agents Chemother* 64:e01496-19. <https://doi.org/10.1128/AAC.01496-19>.
39. Fisher JF, Mobashery S. 2016. β -Lactam resistance mechanisms: Gram-positive bacteria and *Mycobacterium tuberculosis*. *Cold Spring Harb Perspect Med* 6:a025221. <https://doi.org/10.1101/cshperspect.a025221>.
40. Sauvage E, Kerff F, Terrak M, Ayala JA, Charlier P. 2008. The penicillin-binding proteins: structure and role in peptidoglycan biosynthesis. *FEMS Microbiol Rev* 32:234–258. <https://doi.org/10.1111/j.1574-6976.2008.00105.x>.
41. Zapun A, Contreras-Martel C, Vernet T. 2008. Penicillin-binding proteins and β -lactam resistance. *FEMS Microbiol Rev* 32:361–385. <https://doi.org/10.1111/j.1574-6976.2007.00095.x>.
42. Kwon DH, Dore MP, Kim JJ, Kato M, Lee M, Wu JY, Graham DY. 2003. High-level beta-lactam resistance associated with acquired multidrug resistance in *Helicobacter pylori*. *Antimicrob Agents Chemother* 47:2169–2178. <https://doi.org/10.1128/AAC.47.7.2169-2178.2003>.
43. Dewé TCM, D'Aeth JC, Croucher NJ. 2019. Genomic epidemiology of penicillin-non-susceptible *Streptococcus pneumoniae*. *Microb Genom* 5:e000305. <https://doi.org/10.1099/mgen.0.000305>.
44. Griffiths D, Fawley W, Kachrimanidou M, Bowden R, Crook DW, Fung R, Golubchik T, Harding RM, Jeffery KJ, Jolley KA, Kirtan R, Peto TE, Rees G, Stoesser N, Vaughan A, Walker AS, Young BC, Wilcox M, Dingle KE. 2010. Multilocus sequence typing of *Clostridium difficile*. *J Clin Microbiol* 48:770–778. <https://doi.org/10.1128/JCM.01796-09>.
45. Isidro J, Santos A, Nunes A, Borges V, Silva C, Vieira L, Mendes AL, Serrano M, Henriques AO, Gomes JP, Oleastro M. 2018. Imipenem resistance in *Clostridium difficile* ribotype 017, Portugal. *Emerg Infect Dis* 24:741–745. <https://doi.org/10.3201/eid2404.170095>.
46. Dembek M, Barquist L, Boinett CJ, Cain AK, Mayho M, Lawley TD, Fairweather NF, Fagan RP. 2015. High-throughput analysis of gene essentiality and sporulation in *Clostridium difficile*. *mBio* 6:e02383-14. <https://doi.org/10.1128/mBio.02383-14>.
47. Aoki K, Takeda S, Miki T, Ishii Y, Tateda K. 2019. Antimicrobial susceptibility and molecular characterization using whole-genome sequencing of *Clostridioides difficile* collected in 82 hospitals in Japan between 2014 and 2016. *Antimicrob Agents Chemother* 63:e01259-19. <https://doi.org/10.1128/AAC.01259-19>.
48. Jia H, Du P, Yang H, Zhang Y, Wang J, Zhang W, Han G, Han N, Yao Z, Wang H, Zhang J, Wang Z, Ding Q, Qiang Y, Barbut F, Gao GF, Cao Y, Cheng Y, Chen C. 2016. Nosocomial transmission of *Clostridium difficile* ribotype 027 in a Chinese hospital, 2012–2014, traced by whole genome sequencing. *BMC Genomics* 17:405. <https://doi.org/10.1186/s12864-016-2708-0>.
49. Qin J, Dai Y, Ma X, Wang Y, Gao Q, Lu H, Li T, Meng H, Liu Q, Li M. 2017. Nosocomial transmission of *Clostridium difficile* genotype ST81 in a general teaching hospital in China traced by whole genome sequencing. *Sci Rep* 7:9627. <https://doi.org/10.1038/s41598-017-09878-8>.
50. Wu Y, Liu C, Li WG, Xu JL, Zhang WZ, Dai YF, Lu JX. 2019. Independent microevolution mediated by mobile genetic elements of individual *Clostridium difficile* isolates from clade 4 revealed by whole-genome sequencing. *mSystems* 4:e00252-18. <https://doi.org/10.1128/mSystems.00252-18>.
51. Ramírez-Vargas G, Quesada-Gómez C, Acuña-Amador L, López-Ureña D, Murillo T, Del Mar Gamboa-Coronado M, Chaves-Olarte E, Thomson N, Rodríguez-Cavallini E, Rodríguez C. 2017. A *Clostridium difficile* lineage endemic to Costa Rican hospitals is multidrug resistant by acquisition of chromosomal mutations and novel mobile genetic elements. *Antimicrob Agents Chemother* 61:e02054-16. <https://doi.org/10.1128/AAC.02054-16>.
52. Kachrimanidou M, Baktash A, Metallidis S, Tsachouridou O, Netsika F, Dimoglou D, Kassomenaki A, Mouza E, Haritonidou M, Kuijper E. 2020. An outbreak of *Clostridioides difficile* infections due to a 027-like PCR ribotype 181 in a rehabilitation centre: epidemiological and microbiological characteristics. *Anaerobe* 65:102252. <https://doi.org/10.1016/j.anaerobe.2020.102252>.
53. Boekhoud IM, Sidorov I, Nooij S, Harmanus C, Bos-Sanders IMJG, Viprey V, Spittal W, Clark E, Davies K, Freeman J, Kuijper EJ, Smits WK, COMBACTE-CDI Consortium. 2021. Haem is crucial for medium-dependent metronidazole resistance in clinical isolates of *Clostridioides difficile*. *J Antimicrob Chemother* 76:1731–1740. <https://doi.org/10.1093/jac/dkab097>.
54. Imwattana K, Putsathit P, Knight DR, Kiratisin P, Riley TV. 2021. Molecular characterization of, and antimicrobial resistance in, *Clostridioides difficile* from Thailand, 2017–2018. *Microb Drug Resist* 27:1505–1512. <https://doi.org/10.1089/mdr.2020.0603>.
55. Hiramatsu K, Cui L, Kuroda M, Ito T. 2001. The emergence and evolution of methicillin-resistant *Staphylococcus aureus*. *Trends Microbiol* 9:486–493. [https://doi.org/10.1016/S0966-842X\(01\)02175-8](https://doi.org/10.1016/S0966-842X(01)02175-8).
56. Ludlam H, Brown N, Sule O, Redpath C, Coni N, Owen G. 1999. An antibiotic policy associated with reduced risk of *Clostridium difficile*-associated diarrhoea. *Age Ageing* 28:578–580. <https://doi.org/10.1093/ageing/28.6.578>.

57. O'Connor KA, Kingston M, O'Donovan M, Cryan B, Twomey C, O'Mahony D. 2004. Antibiotic prescribing policy and *Clostridium difficile* diarrhoea. QJM 97:423–429. <https://doi.org/10.1093/qjmed/hch076>.
58. Belmares J, Johnson S, Parada JP, Olson MM, Clabots CR, Bettin KM, Peterson LR, Gerding DN. 2009. Molecular epidemiology of *Clostridium difficile* over the course of 10 years in a tertiary care hospital. Clin Infect Dis 49:1141–1147. <https://doi.org/10.1086/605638>.
59. Kato H, Kita H, Karasawa T, Maegawa T, Koino Y, Takakuwa H, Saikai T, Kobayashi K, Yamagishi T, Nakamura S. 2001. Colonisation and transmission of *Clostridium difficile* in healthy individuals examined by PCR ribotyping and pulsed-field gel electrophoresis. J Med Microbiol 50:720–727. <https://doi.org/10.1099/0022-1317-50-8-720>.
60. Muraki Y, Yagi T, Tsuji Y, Nishimura N, Tanabe M, Niwa T, Watanabe T, Fujimoto S, Takayama K, Murakami N, Okuda M. 2016. Japanese antimicrobial consumption surveillance: first report on oral and parenteral antimicrobial consumption in Japan (2009–2013). J Glob Antimicrob Resist 7:19–23. <https://doi.org/10.1016/j.jgar.2016.07.002>.
61. Kim YA, Park YS, Youk T, Lee H, Lee K. 2018. Changes in antimicrobial usage patterns in Korea: 12-year analysis based on database of the National Health Insurance Service-National Sample Cohort. Sci Rep 8:12210. <https://doi.org/10.1038/s41598-018-30673-6>.
62. Qu X, Yin C, Sun X, Huang S, Li C, Dong P, Lu X, Zhang Z, Yin A. 2018. Consumption of antibiotics in Chinese public general tertiary hospitals (2011–2014): trends, pattern changes and regional differences. PLoS One 13:e0196668. <https://doi.org/10.1371/journal.pone.0196668>.
63. Collins DA, Hawkey PM, Riley TV. 2013. Epidemiology of *Clostridium difficile* infection in Asia. Antimicrob Resist Infect Control 2:21. <https://doi.org/10.1186/2047-2994-2-21>.
64. Park M, Rafii F. 2017. Exposure to beta-lactams results in the alteration of penicillin-binding proteins in *Clostridium perfringens*. Anaerobe 45:78–85. <https://doi.org/10.1016/j.anaerobe.2017.02.004>.
65. Cheng JW, Xiao M, Kudinha T, Xu Z, Hou X, Sun L, Zhang L, Fan X, Kong F, Xu Y. 2016. The first two *Clostridium difficile* ribotype 027/ST1 isolates identified in Beijing, China—an emerging problem or a neglected threat? Sci Rep 6:18834. <https://doi.org/10.1038/srep18834>.
66. Wilcox MH, Fawley WN. 2000. Hospital disinfectants and spore formation by *Clostridium difficile*. Lancet 356:1324. [https://doi.org/10.1016/S0140-6736\(00\)02819-1](https://doi.org/10.1016/S0140-6736(00)02819-1).
67. Wang B, Peng W, Zhang P, Su J. 2018. The characteristics of *Clostridium difficile* ST81, a new PCR ribotype of toxin A⁺ B⁺ strain with high-level fluoroquinolones resistance and higher sporulation ability than ST37/PCR ribotype 017. FEMS Microbiol Lett 365. <https://doi.org/10.1093/femsle/fny168>.
68. Burns DA, Heap JT, Minton NP. 2010. The diverse sporulation characteristics of *Clostridium difficile* clinical isolates are not associated with type. Anaerobe 16:618–622. <https://doi.org/10.1016/j.anaerobe.2010.10.001>.
69. Burns DA, Heeg D, Cartman ST, Minton NP. 2011. Reconsidering the sporulation characteristics of hypervirulent *Clostridium difficile* BI/NAP1/027. PLoS One 6:e24894. <https://doi.org/10.1371/journal.pone.0024894>.
70. Karłowsky JA, Adam HJ, Baxter MR, Dutka CW, Nichol KA, Laing NM, Golding GR, Zhanel GG. 2020. Antimicrobial susceptibility of *Clostridioides difficile* isolated from diarrhoeal stool specimens of Canadian patients: summary of results from the Canadian *Clostridioides difficile* (CAN-DIFF) surveillance study from 2013 to 2017. J Antimicrob Chemother 75:1824–1832. <https://doi.org/10.1093/jac/dkaa118>.
71. Stoesser N, Eyre DW, Quan TP, Godwin H, Pill G, Mbuvi E, Vaughan A, Griffiths D, Martin J, Fawley W, Dingle KE, Oakley S, Wanelik K, Finney JM, Kachrimanidou M, Moore CE, Gorbach S, Riley TV, Crook DW, Peto TEA, Wilcox MH, Walker AS, Modernising Medical Microbiology Informatics Group (MMMIG). 2017. Epidemiology of *Clostridium difficile* in infants in Oxfordshire, UK: risk factors for colonization and carriage, and genetic overlap with regional *C. difficile* infection strains. PLoS One 12:e0182307. <https://doi.org/10.1371/journal.pone.0182307>.
72. Karachalios G, Charalabopoulos K. 2002. Biliary excretion of antimicrobial drugs. Chemotherapy 48:280–297. <https://doi.org/10.1159/000069712>.
73. Kokai-Kun JF, Roberts T, Coughlin O, Sicard E, Rufiange M, Fedorak R, Carter C, Adams MH, Longstreth J, Whalen H, Sliman J. 2017. The oral beta-lactamase SYN-004 (ribaxamase) degrades ceftriaxone excreted into the intestine in phase 2a clinical studies. Antimicrob Agents Chemother 61:e02197-16. <https://doi.org/10.1128/AAC.02197-16>.
74. Negri MC, Morosini MI, Baquero MR, del Campo R, Blázquez J, Baquero F. 2002. Very low cefotaxime concentrations select for hypermutable *Streptococcus pneumoniae* populations. Antimicrob Agents Chemother 46:528–530. <https://doi.org/10.1128/AAC.46.2.528-530.2002>.
75. Gullberg E, Cao S, Berg OG, Ilbäck C, Sandegren L, Hughes D, Andersson DI. 2011. Selection of resistant bacteria at very low antibiotic concentrations. PLoS Pathog 7:e1002158. <https://doi.org/10.1371/journal.ppat.1002158>.
76. Gullberg E, Albrecht LM, Karlsson C, Sandegren L, Andersson DI. 2014. Selection of a multidrug resistance plasmid by sublethal levels of antibiotics and heavy metals. mBio 5:e01918-14. <https://doi.org/10.1128/mBio.01918-14>.
77. Andersson DI, Hughes D. 2012. Evolution of antibiotic resistance at non-lethal drug concentrations. Drug Resist Updat 15:162–172. <https://doi.org/10.1016/j.drug.2012.03.005>.
78. Murray AK, Zhang L, Yin X, Zhang T, Buckling A, Snape J, Gaze WH. 2018. Novel insights into selection for antibiotic resistance in complex microbial communities. mBio 9:e00969-18. <https://doi.org/10.1128/mBio.00969-18>.
79. Sandegren L. 2014. Selection of antibiotic resistance at very low antibiotic concentrations. Ups J Med Sci 119:103–107. <https://doi.org/10.3109/03009734.2014.904457>.
80. Zhao H, Nickle DC, Zeng Z, Law PYT, Wilcox MH, Chen L, Peng Y, Meng J, Deng Z, Albright A, Zhong H, Xu X, Zhu S, Shen J, Blanchard RL, Dorr MB, Shaw PM, Li J. 2021. Global landscape of *Clostridioides difficile* phylogeography, antibiotic susceptibility, and toxin polymorphisms by post-hoc whole-genome sequencing from the MODIFY I/II studies. Infect Dis Ther 10:853–870. <https://doi.org/10.1007/s40121-021-00426-6>.
81. Department of Health, Health Protection Agency. 2008. *Clostridium difficile* infection: how to deal with the problem. https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/340851/Clostridium_difficile_infection_how_to_deal_with_the_problem.pdf.
82. Zerbino DR, Birney E. 2008. Velvet: algorithms for de-novo short read assembly using de Bruijn graphs. Genome Res 18:821–829. <https://doi.org/10.1101/gr.074492.107>.
83. Gladman S, Seemann T. 2008. VelvetOptimiser, 2.1.7. Monash University, Victoria, Australia.
84. Frentrop M, Zhou Z, Steglich M, Meier-Kolthoff JP, Göker M, Riedel T, Bunk B, Spröer C, Overmann J, Blaschitz M, Indra A, von Müller L, Kohl TA, Niemann S, Seyboldt C, Klawonn F, Kumar N, Lawley TD, García-Fernández S, Cantón R, Del Campo R, Zimmermann O, Groß U, Achtman M, Nübel U. 2020. A publicly accessible database for *Clostridioides difficile* genome sequences supports tracing of transmission chains and epidemics. Microb Genom 6:mgen000410. <https://doi.org/10.1099/mgen.0.000410>.
85. Jolley KA, Bray JE, Maiden MCJ. 2018. Open-access bacterial population genomics: BIGSdb software, the PubMLST.org website and their applications. Wellcome Open Res 3:124. <https://doi.org/10.12688/wellcomeopenres.14826.1>.
86. Spigaglia P, Barbanti F, Mastrantonio P, Brazier JS, Barbut F, Delmée M, Kuijper E, Poxton IR, on behalf of The European Study Group on ESGCD. 2008. Fluoroquinolone resistance in *Clostridium difficile* isolates from a prospective study of *C. difficile* infections in Europe. J Med Microbiol 57:784–789. <https://doi.org/10.1099/jmm.0.47738-0>.
87. Drudy D, Quinn T, O'Mahony R, Kyne L, O'Gaora P, Fanning S. 2006. High-level resistance to moxifloxacin and gatifloxacin associated with a novel mutation in *gyrB* in toxin-A-negative, toxin-B-positive *Clostridium difficile*. J Antimicrob Chemother 58:1264–1267. <https://doi.org/10.1093/jac/dkl398>.
88. Curry SR, Marsh JW, Shutt KA, Muto CA, O'Leary MM, Saul MI, Pasculle AW, Harrison LH. 2009. High frequency of rifampin resistance identified in an epidemic *Clostridium difficile* clone from a large teaching hospital. Clin Infect Dis 48:425–429. <https://doi.org/10.1086/596315>.
89. Spigaglia P, Barbanti F, Mastrantonio P, European Study Group on *Clostridium difficile* (ESGCD). 2011. Multidrug resistance in European *Clostridium difficile* clinical isolates. J Antimicrob Chemother 66:2227–2234. <https://doi.org/10.1093/jac/dkr292>.
90. Tamura K, Stecher G, Kumar S. 2021. MEGA11: Molecular Evolutionary Genetics Analysis version 11. Mol Biol Evol 38:3022–3027. <https://doi.org/10.1093/molbev/msab120>.
91. Baines SD, O'Connor R, Freeman J, Fawley WN, Harmanus C, Mastrantonio P, Kuijper EJ, Wilcox MH. 2008. Emergence of reduced susceptibility to metronidazole in *Clostridium difficile*. J Antimicrob Chemother 62:1046–1052. <https://doi.org/10.1093/jac/dkn313>.
92. Freeman J, Vernon J, Morris K, Nicholson S, Todhunter S, Longshaw C, Wilcox MH. 2015. Pan-European longitudinal surveillance of antibiotic resistance among prevalent *Clostridium difficile* ribotypes. Clin Microbiol Infect 21:248.e9–248.e16. <https://doi.org/10.1016/j.cmi.2014.09.017>.

93. Clinical and Laboratory Standards Institute. 2012. M11-A8. Methods for antimicrobial susceptibility testing of anaerobic bacteria; approved standard—8th ed. Clinical and Laboratory Standards Institute, Wayne, PA.
94. Clinical and Laboratory Standards Institute. 2012. M100S. Performance standards for antimicrobial susceptibility testing. Clinical and Laboratory Standards Institute, Wayne, PA.
95. Kurtz S, Phillippy A, Delcher AL, Smoot M, Shumway M, Antonescu C, Salzberg SL. 2004. Versatile and open software for comparing large genomes. *Genome Biol* 5:R12. <https://doi.org/10.1186/gb-2004-5-2-r12>.
96. He M, Sebaihia M, Lawley TD, Stabler RA, Dawson LF, Martin MJ, Holt KE, Seth-Smith HM, Quail MA, Rance R, Brooks K, Churcher C, Harris D, Bentley SD, Burrows C, Clark L, Corton C, Murray V, Rose G, Thurston S, van Tonder A, Walker D, Wren BW, Dougan G, Parkhill J. 2010. Evolutionary dynamics of *Clostridium difficile* over short and long time scales. *Proc Natl Acad Sci U S A* 107:7527–7532. <https://doi.org/10.1073/pnas.0914322107>.
97. Ahn SW, Lee SH, Kim UJ, Jang HC, Choi HJ, Choy HE, Kang SJ, Roh SW. 2021. Genomic characterization of nine *Clostridioides difficile* strains isolated from Korean patients with *Clostridioides difficile* infection. *Gut Pathog* 13:55. <https://doi.org/10.1186/s13099-021-00451-3>.
98. Yin C, Chen DS, Zhuge J, McKenna D, Sagurton J, Wang G, Huang W, Dimitrova N, Fallon JT. 2018. Complete genome sequences of four toxigenic *Clostridium difficile* clinical isolates from patients of the lower Hudson Valley, New York, USA. *Genome Announc* 6:e01537-17. <https://doi.org/10.1128/genomeA.01537-17>.
99. Sebaihia M, Wren BW, Mullany P, Fairweather NF, Minton N, Stabler R, Thomson NR, Roberts AP, Cerdeño-Tárraga AM, Wang H, Holden MT, Wright A, Churcher C, Quail MA, Baker S, Bason N, Brooks K, Chillingworth T, Cronin A, Davis P, Dowd L, Fraser A, Feltwell T, Hance Z, Holroyd S, Jagels K, Moule S, Mungall K, Price C, Rabinowitsch E, Sharp S, Simmonds M, Stevens K, Unwin L, Whithead S, Dupuy B, Dougan G, Barrell B, Parkhill J. 2006. The multidrug-resistant human pathogen *Clostridium difficile* has a highly mobile, mosaic genome. *Nat Genet* 38:779–786. <https://doi.org/10.1038/ng1830>.
100. Guindon S, Dufayard J-F, Lefort V, Anisimova M, Hordijk W, Gascuel O. 2010. New algorithms and methods to estimate maximum-likelihood phylogenies: assessing the performance of PhyML 3.0. *Syst Biol* 59: 307–321. <https://doi.org/10.1093/sysbio/syq010>.
101. Didelot X, Wilson DJ. 2015. ClonalFrameML: efficient inference of recombination in whole bacterial genomes. *PLoS Comput Biol* 11:e1004041. <https://doi.org/10.1371/journal.pcbi.1004041>.
102. Didelot X, Croucher NJ, Bentley SD, Harris SR, Wilson DJ. 2018. Bayesian inference of ancestral dates on bacterial phylogenetic trees. *Nucleic Acids Res* 46:e134. <https://doi.org/10.1093/nar/gky783>.
103. Didelot X, Eyre DW, Cule M, Ip CLC, Ansari MA, Griffiths D, Vaughan A, O'Connor L, Golubchik T, Batty EM, Piazza P, Wilson DJ, Bowden R, Donnelly PJ, Dingle KE, Wilcox M, Walker AS, Crook DW, Peto TEA, Harding RM. 2012. Microevolutionary analysis of *Clostridium difficile* genomes to investigate transmission. *Genome Biol* 13:R118. <https://doi.org/10.1186/gb-2012-13-12-r118>.
104. Lawley TD, Croucher NJ, Yu L, Clare S, Sebaihia M, Goulding D, Pickard DJ, Parkhill J, Choudhary J, Dougan G. 2009. Proteomic and genomic characterization of highly infectious *Clostridium difficile* 630 spores. *J Bacteriol* 191:5377–5386. <https://doi.org/10.1128/JB.00597-09>.
105. Pettit LJ, Browne HP, Yu L, Smits WK, Fagan RP, Barquist L, Martin MJ, Goulding D, Duncan SH, Flint HJ, Dougan G, Choudhary JS, Lawley TD. 2014. Functional genomics reveals that *Clostridium difficile* Spo0A coordinates sporulation, virulence and metabolism. *BMC Genomics* 15:160. <https://doi.org/10.1186/1471-2164-15-160>.