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Global Landscape of *Clostridioides Difficile* Phylogeography, Antibiotic Susceptibility, and Toxin Polymorphisms by Post-Hoc Whole-Genome Sequencing from the MODIFY I/II Studies

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ABSTRACT

**Introduction:** *Clostridioides* (*Clostridium*) *difficile* infection, the leading cause of healthcare-associated diarrhea, represents a significant burden on global healthcare systems. Despite being a global issue, information on *C. difficile* from a global perspective is lacking. The aim of this study is to model the global phylogeography of clinical *C. difficile*.

**Methods:** Using samples collected from the MODIFY I and II studies (NCT01241552, NCT01513239), we performed whole-genome sequencing of 1501 clinical isolates including 37 novel sequence types (STs), representing the largest worldwide collection to date.

**Results:** Our data showed ribotypes, multi-locus sequence typing clades, and whole-genome phylogeny were in good accordance. The clinical *C. difficile* genome was found to be more conserved than previously reported (61% core genes), and modest recombination rates of 1.4–5.0 were observed across clades. We observed a significant continent distribution preference among five *C. difficile* clades (Benjamini-Hochberg corrected Fisher’s exact test).

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1007/s40121-021-00426-6.
test $P < 0.01$); moreover, weak association between geographic and genetic distance among ribotypes suggested sources beyond healthcare-related transmission. Markedly different trends of antibiotic susceptibility among lineages and regions were identified, and three novel mutations (in pyridoxamine $5\’$-phosphate oxidase family protein: Tyr130Ser, Tyr130Cys, and a promoter SNP) associated with metronidazole-reduced susceptibility were discovered on a nim-related gene and its promoter by genome-wide association study. Toxin gene polymorphisms were shown to vary within and between prevalent ribotypes, and novel severe mutations were found on the tcdC toxin regulator protein.

**Conclusion:** Our systematic characterization of a global set of clinical trial C. difficile isolates from infected individuals demonstrated the complexity of the genetic makeup of this pathogenic organism. The geographic variability of clades, variability in toxin genes, and mutations associated with antibiotic susceptibility indicate a highly complex interaction of C. difficile between host and environment. This dataset will provide a useful resource for validation of findings and future research of C. difficile.

**Keywords:** Antibiotic susceptibility; CDI; Clostridioides (Clostridium) difficile; Evolution; Metronidazole; Moxifloxacin; Phylogeography; Rifaximin; Toxin; Vancomycin

**Key Summary Points**

**Why carry out this study?**

*Clostridioides (Clostridium) difficile* infection poses a significant burden on global healthcare systems; however, a global perspective on the characteristics and distribution of C. difficile lineages is lacking

Using samples from the MODIFY I and II clinical trials, this study aimed to analyze the phylogeography, antibiotic susceptibility, and toxin polymorphisms of the largest whole-genome sequenced global collection of clinical trial CDI isolates documented to date

**What was learned from the study?**

This study found ribotypes, multi-locus sequence typing clades, and whole-genome phylogeny were in good accordance, and the clinical C. difficile genome was more conserved than previously reported (> 60% core genes)

Significant continent distribution preference was observed among five C. difficile clades, with markedly different trends of antibiotic susceptibility among lineages and regions, as well as three novel mutations discovered by GWAS, providing useful insight into this highly clinically relevant population

These results provide valuable insight into the spread and characteristics of epidemic C. difficile and allow better understanding of the distribution patterns of hypervirulent strains and antibiotic resistance.

**DIGITAL FEATURES**

This article is published with digital features, including a summary slide, to facilitate understanding of the article. To view digital features for this article go to https://doi.org/10.6084/m9.figshare.14113565.

**INTRODUCTION**

*Clostridioides (Clostridium) difficile* infection (CDI) is the leading cause of healthcare-associated diarrhea, representing a significant burden upon global healthcare systems. Ribotyping and multi-locus sequence typing (MLST) have been the gold standard for the investigation of lineage prevalence and disease association. Determining phylogenetic relationships across geographic regions allows for the investigation of disease dissemination across landscapes. However, the correlation among ribotyping, MLST, and whole-genome single-nucleotide polymorphism (SNP) phylogeny has
not been explored with an extensive dataset of cases collected prospectively across multiple continents.

Clostridioides (Clostridium) difficile (C. difficile) shares a paradoxical relationship with antibiotics: CDI is recognized as an antibiotic-associated infection caused by an imbalance in the gut microbiome; conversely, antibiotics are also the first-line treatment for CDI. A detailed review of the prevalence and known mechanisms of antibiotic susceptibility/resistance in C. difficile has been published [1]. Rates of resistance to the two first-line antibiotics, metronidazole (MET) and vancomycin (VAN), are reported to be low in the published literature [2]. Nonetheless, failure of treatment with MET has been increasingly reported in recent years [3]. The resistance mechanisms employed by C. difficile against both first-line antibiotics remain poorly understood [1].

Clinical manifestations of CDI are mainly caused by virulence factors Toxin A (TcdA) and Toxin B (TcdB), encoded and regulated by genes on the pathogenicity locus including the negative regulator Toxin C (TcdC), as well as the binary toxin cytolethal distending toxin (CDT), which may potentiate the toxicity of TcdA and TcdB and lead to more severe disease [4]. Many studies have attributed the hypervirulence of the C. difficile ribotype RT027 to its tcdB sequence heterogeneity, inactivity of tcdC alleles, and presence of CDT [5].

Information on the distribution of C. difficile lineages, ribotypes, geographic associations, resistance prevalence, and toxin polymorphisms from a global perspective is lacking. Using samples collected from the MODIFY I and II studies [6], the largest whole-genome sequenced global collection of clinical trial CDI isolates documented to date, our study aims to describe the characteristics of a representative clinical C. difficile population.

METHODS

Sample Collection

The collection of the clinical C. difficile strains used in this study was previously described in detail by Wilcox et al. [6]. Briefly, 1501 CDI isolates were collected from two phase III, randomized, double-blind, placebo-controlled trials (MODIFY I [NCT01241552] and MODIFY II [NCT01513239]). No selection criteria were applied for the strains from the two trials that were analyzed in this study. This cohort represents all available strains cultured from baseline stool samples obtained from enrolled subjects [7].

DNA Extraction and Ribotyping

DNA was extracted from freshly isolated cultures generated from individual colonies, using standard large-scale DNA extraction with Qiagen Kits (QIAGEN, Venlo, The Netherlands). Ribotype analyses were conducted as previously described [8].

Whole-Genome Sequencing

Polymerase chain reaction-free index libraries, with an average insert size of 350 bp, were prepared for each isolate. Genome sequencing was performed using the Illumina® HiSeq™ 2000 platform to generate 100-bp paired-end (PE) or 90-bp PE reads (Illumina, Inc., San Diego, CA). Raw reads were filtered by SOAPnuke (https://github.com/BGI-flexlab/SOAPnuke) to remove sequencing adapters and low-quality reads, including those containing more than three unknown (denoted ‘N’) bases, those in which >50% of the bases scored <Q20, and those with read length <30 bp. After the quality-control process, high-quality data with an average of 686 Mb were generated, which has a coverage of 170-fold across the whole genome (~4 Mb).

Genome Assembly, Coding Sequence Prediction, and Core/Pangenome Identification

De novo genome assembly was performed using SPAdes v3.11.1 [9] (or SOAPdenovo2 [10] in toxin analysis for several isolates). Non-redundant coding sequence (CDS) prediction and annotation were performed using Prokka v1.13 [11]. Identification of core/pangenome was
performed with Roary v3.12.0 [12] (default setting; minimum percentage identity for blastp is 95%, percentage of isolates a gene must be in to be core is 99%).

SNP Analysis

1. Whole-genome SNP calling: high-quality reads were mapped against reference genome CD196 (NC_013315) using a Burrows-Wheeler Aligner (BWA v0.6.2) [13]. SNPs were identified using SAMtools v0.1.19 [14]. An allele was only considered to be valid if align qual \( \geq 100 \), depth \( \geq 10 \), and alt rate \( \geq 0.8 \) (found in \( \geq 80\% \) of reads); otherwise, it was treated as missing.

2. Repetitive region identification: repetitive regions in the reference genome sequence were identified using an in-house pipeline. Briefly, the reference genome was fragmented in silico into 20-bp reads using a sliding window approach; the reads generated were then mapped back to the reference, and reads mapped to multiple locations of the genome were identified. SNPs falling within these repetitive regions were excluded from all subsequent analyses \( (3,851,695/4,110,554 = 93.7\%) \). The remaining SNPs were named whole-genome SNPs in our article.

3. Validation of the SNP pipeline: artificial variants were introduced to generate a pseudo-genome based on CD630 by INDELible v1.03 [15]. The pseudo-genome and CD630 were compared to extract the SNP and insertion/deletion (indel) sites \( (31,831 \text{ SNPs and 8524 indels}) \). Repetitive regions were identified \( (3,923,283/4,290,244 = 91.4\%) \) in the pseudo-genome using the pipeline described above. Realistic sequencing reads and reads with sequencing errors (based on the sequencing quality of a real sample) were generated by iMESSi [16], accounting for a depth from 20 to 200X of the genome separately. SNPs were called using the above pipeline. Valid accuracy rate was assessed by comparing these two SNP sets (the error rate of the average data size was \( \sim 0.06\%) \).

Phylogenetic Analyses and Recombination Detection

Pseudo-whole-genome sequences were constructed for each isolate by replacing SNP positions in the CD196 reference genome with the consensus alleles identified by our SNP calling pipeline, with unknown sites being replaced by a gap character. These alignment sequences were adopted to construct the recombination-free maximum-likelihood (ML) phylogeny and detect the recombination rate (r/m ratio) within each clade by Gubbins [17] program. iTOL (https://itol.embl.de/) was used for tree visualization.

Genetic Diversity Analysis

SNP-level difference among isolates within a ribotype was calculated. Pairwise SNPs were counted based on defined bases only; missing data in either of two compared sites were ignored (for example, ‘-’ vs. ‘A’). The average number of nucleotide differences per site (nucleotide diversity, \( \pi \)) among ribotypes was calculated using DnaSP v5 [18].

MLST Analysis

Assembled contigs were aligned to a C. difficile MLST database (http://pubmlst.org/cdifficile), which organizes all allele sequences of seven C. difficile housekeeping genes to identify the sequence type and corresponding clade.

Geographical Epidemiology Analysis

The distribution pattern of the five clades across the six continents was tested by Fisher’s exact test with Monte Carlo simulation. The specific distribution preference of each clade was tested by Fisher’s exact test with the \( P \) value corrected using the Benjamini-Hochberg (BH) method. The prevalence patterns of clade 1 and clade 2 in the USA were fitted by polynomial curve (four states with only clade 1 were excluded). The genetic distance between any two isolates was calculated based on the whole-genome
SNPs, using the function of dist.dna in the “ape” R package [19] with the substitute model TN93 adopted. The geographic distance between collection sites of any two isolates was calculated by the distCosine command in R, based on the longitude and latitude, which were generated by the geocode command ‘ggmap’ using region information for each isolate. The Mantel test from the R package “ade4” was used to determine correlations between the genetic and geographic distance between isolates, with a null distribution generated by 10^5 permutations, and the Kendall rank correlation coefficient (τ) was adopted according to the distribution of the two matrices and the sample size of each ribotype. P value was corrected by BH. Restriction of gene flow between geographic areas in each ribotype was tested based on the maximum-likelihood phylogenetic tree of each ribotype using the Slatkin-Maddison test [20] under 1000 permutations in the HyPhy software [21].

Antibiotic Susceptibility/Resistance Analysis
Antibiotic susceptibility testing was performed by R.M. Alden (Culver City, CA) using the agar dilution method approved in the CLSI M11-A8 standard [22]. The resistance or reduced susceptibility of an isolate to MOX, MET, and VAN was categorized according to EUCAST ECOFF (version 7.1) breakpoints (VAN 2 mg/l; MET 2 mg/l; MOX 4 mg/l; RIF 0.004 mg/l) [23]. Isolates with MIC ≥ breakpoint were considered as ‘resistant’ (for MOX and RIF) or as reduced susceptibility (for VAN and MET). Antibiotic resistance determinants were identified using Resistance Gene Identifier 4.0.3 of the Comprehensive Antibiotic Resistance Database (CARD 2.0.0) [24], supplemented with SNPs previously published in the literature [25]. Antibiotic preference of each clade or continent was tested by Fisher’s exact test with the P value corrected by BH.

Genome-Wide Association Study (GWAS)
The antibiotic-resistant GWAS was performed with treeWAS [26] based on a SNP matrix of 1474 isolates with MIC value (n.snps.sim = 5*ncol(snps), P = 1e–05) using a ML phylogenetic tree constructed by whole-genome SNP to test the association.

Toxin Gene Analysis
De novo assembled contigs were compared with reference genes (tcdB and tcdC of CD630 [RT012], which lacks binary toxin genes; cdtA and cdtB of CD196 [RT027], which has an incomplete tcdC gene), using BLAST, and CDS were extracted according to the alignment position for each sample. The presence of indels may prevent sequence alignment, which was addressed by connecting the truncated regions to identify start and end positions and deletion region. A single amino acid sequence (or nucleotide sequence for isolates with incomplete tcdC) was selected randomly from identical sequences as a variant type (VT) within each ribotype (i.e., VT was defined by any sequence variation). VT sequences were multiple-aligned by ClustalW [27] and then used to construct the corresponding maximum-likelihood gene tree. A specific non-synonymous variant of tcdB was displayed by MEGA7 [28]. Additionally, Geneious [29] was adopted to visualize the variants for tcdC. For cdtA and cdtB, the majority base in each site within a ribotype was extracted to construct one typical sequence for that ribotype based on the whole-genome SNPs, generating the gene consensus sequence for each ribotype.

Compliance with Ethics Guidelines
MODIFY I (NCT01241552) and MODIFY II (NCT01513239) were conducted in accordance with Good Clinical Practice guidelines and the provisions of the Declaration of Helsinki. The protocols and amendments were approved by the institutional review board or independent ethics committee at each study site (BGI IRB; IRB #13029). Written informed consent for specimen collection and use was provided by all participants before the trial began.
RESULTS

Phylogeny of Global *C. difficile* Collection

We performed whole-genome sequencing (WGS) on 1501 *C. difficile* isolates from stools collected from participants enrolled in two phase III clinical trials (MODIFY I and MODIFY II) [6], which included both hospitalized and out-patients who had been diagnosed with CDI. Participants were enrolled between 2011 and 2015, from 322 sites in 30 countries/regions across six continents (Table S1 and Fig. S1).

We compared consistency between the WGS-based, recombination-free, maximum-likelihood phylogeny and the two most commonly used molecular typing methods (MLST and ribotyping). Ribotyping analysis identified 164 distinct ribotypes, of which 32 ribotypes contained 10 or more isolates (termed major ribotypes hereafter). MLST analysis identified 129 sequence types (STs), including 29 novel gene alleles and 37 novel STs, which were submitted to the *C. difficile* MLST database (http://pubmlst.org/cdifficile). Eight isolates could not be identified using the MLST database (Table S2), such that a total of 1493 isolates (99.5%), belonging to 159 ribotypes, could be assigned to five predefined MLST clades. Among these ribotypes, 151 (containing 1467 isolates) were solely found in one of the MLST clades (Table S2). Generally, we observed high concordance among the WGS-based phylogenetic tree, ribotype, and MLST clade (Fig. 1). However, our results demonstrated a relatively poor correlation between ribotype and ST, with only 78 (49%) ribotypes corresponding to unique STs. The correspondence between ribotype and MLST is summarized in Table S2.

We next investigated the prevalence of ribotypes among the five MLST clades. Clade 1 possessed the greatest number of isolates (63.8%; n = 958) and ribotypes (65.9%; n = 108). The well-known non-toxigenic ribotypes RT009, RT010, and RT039 with toxigenic profile TcdAB− CDT− [30] belong to clade 1. Clade 2 was predominantly comprised of the hypervirulent ribotype RT027 (72.1%; n = 279), along with RT176 (5.4%; n = 21) and RT198 (3.6%; n = 14), which displayed an inseparable phylogenetic relationship with RT027. Clades 3 and 4 were smaller clades, dominated by toxigenic RT023 (65.0%; n = 13) and RT017 (57.9%; n = 33), respectively. Clade 5 was another hypervirulent clade, predominantly comprised of RT078 (62.0%; n = 44) and RT126 (26.8%; n = 19).

Genetic Diversity of the *C. difficile* Genome

To elucidate the genetic diversity of *C. difficile*, we carried out pan-genome analysis on our collection. We identified a total of 29,921 CDSs, out of which 2317 CDSs were identified as core genes (defined as those present in ≥ 99% of all of the 1501 genomes), representing 61.1% of the average number of CDS per genome (3792 CDS), suggesting that clinical *C. difficile* isolates shared a conserved core genome.

To estimate the role played by homologous recombination, we calculated the rate of recombination (r) with respect to random point mutation (m) for each clade (r/m ratio). Clades 2 and 5, which include strains that have been shown to be hypervirulent, exhibited the lowest recombination rates at 1.8 and 1.4, respectively. Clades 3 and 4 showed similar recombination rates of 4.8 and 5.0, respectively, while the heterogeneous clade 1 had a slightly lower rate of 3.9. Overall, the *C. difficile* recombination rates reported here are modest.

Next, we investigated the nucleotide diversity (π) among the 32 major ribotypes. The π for all major ribotypes correlated well with results from the median pairwise SNP (PWSNP) analysis (Pearson’s correlation coefficient 0.94) (Table S3). The number of PWSNPs within each ribotype varied markedly (Fig. 2), with the greatest genetic variation observed in RT153 (π = 0.037, median PWSNP = 11,792; Table S3).

Geographical Epidemiology of *C. difficile*

The isolates were collected globally to study *C. difficile* in the most affected and representative populations. We examined the distribution pattern of the five clades across the
six continents and found a significant overall trend for clade distribution (Fisher’s exact test with Monte Carlo simulation, \( P = 1e^{-06} \)). Furthermore, we explored the specific distribution preference of each clade by Fisher’s exact test with BH corrected \( P < 0.01 \) (Fig. 3a). While the most heterogeneous clade, clade 1, was identified in all sampled countries/regions, it was the most predominant clade found in Oceania (Oceania includes only Australia and New Zealand in our data) (odds ratio [OR] 9.35, 95% confidence interval [CI] 2.99–47.08),...
followed by Asia (OR 2.14, 95% CI 1.46–3.22). Hypervirulent clades 2 and 5, primarily represented by ribotypes RT027 and RT078, were more likely to be found in North America (OR 2.68, 95% CI 2.09–3.45) and Europe (OR 3.29, 95% CI 1.97–5.56), respectively. Interestingly, the prevalence of clade 1 and clade 2 in the USA (the region with the greatest number of isolates) showed regional divergence; for clade 2, prevalence was higher on the East Coast and West Coast compared with inland, and this observation was reversed for clade 1, which was more prevalent inland (Fig. 3b). Clade 3 was most likely to be found in Europe (OR 5.1, 95% CI 1.83–16.30) (Fig. 3c), and clade 4 was predominantly observed in Asia (OR 2.89, 95% CI 1.45–5.46).

To further understand the geographic distribution of isolates from a more refined perspective, we investigated the relationship between the genetic distance between isolates and the geographic distance between collection sites for each ribotype. Of the 32 major ribotypes analyzed, 9 showed a significant correlation between genetic and geographic distances (BH-corrected Mantel test, \( P < 0.01 \)) (Table S4). The strongest significant correlation, calculated by Kendall rank correlation coefficient (\( \tau \)), was detected with RT198 (\( \tau = 0.63, P = 0.001 \)), followed by RT176 (\( \tau = 0.59, P = 0.001 \)) and RT137 (\( \tau = 0.40, P = 0.003 \)). Analysis of gene flow restriction provided support for the above findings from a different angle: significantly fewer migration events were detected for these ribotypes (BH-corrected Slatkin-Maddison test \( P < 0.01 \)) (Table S4). Although we did not see a correlation between genetic distance and geographic distance, we saw a clear restriction of gene flow across the globe. This suggests that a restriction of gene flow is driven by factors other than simple global distance.

Global Profile of Antibiotic Resistance in \textit{C. difficile}

To elucidate the distribution of antibiotic susceptibility/resistance in \textit{C. difficile} at a global level, we determined the in vitro minimum inhibitory concentrations (MICs) of four antibiotics (VAN, MET, moxifloxacin [MOX],
rifaximin (RIF) for the 1474 C. difficile isolates. Antibiotic susceptibility was determined according to European Committee on Antimicrobial Susceptibility Testing (EUCAST; version 7.1) epidemiological cut-off (ECOFF) breakpoints (VAN 2 mg/l; MET 2 mg/l; MOX 4 mg/l; RIF 0.004 mg/l). The overall reduced susceptibility/resistance rates were 32.8% for VAN, 14.5% for MET, 37.9% for MOX, and 99.5% for RIF. In total, 5.6% (83/1474) of isolates were determined to have reduced susceptibility/resistance to all four classes of antibiotics, out of which 77.1% (64/83) belonged to clade 2, having the highest mean MIC value among the population (Fig. 1). In total, 67% of RT027 strains showed reduced susceptibility or resistance to all four classes of antibiotic tested.

We studied the prevalence of reduced susceptibility/resistance to each of the four antibiotics with EUCAST ECOFF breakpoints among MLST clades and continents. Reduced susceptibility to VAN was significantly less likely among the samples from clade 4 (OR 0.23, 95% CI 0.08–0.55) and clade 5 (OR 0.15, 95% CI 0.05–0.37) as well as the samples from Europe (OR 0.61, 95% CI 0.48–0.79). The three ribotypes with the highest antibiotic susceptibility/resistance rates are RT137, 012, and 356 (0.76, 0.71, and 0.67, respectively). The reduced susceptibility/resistance to both MET and MOX was significantly more likely in clade 2 (OR 9.07, 95% CI 6.52–12.72 and OR 8.15, 95% CI 6.21–10.75 respectively); MOX resistance was significantly more likely to be observed among the samples from Europe (OR 1.58, 95% CI 1.25–1.98) and Asia (OR 2.67, 95% CI 1.91–3.75) than among those from other continents, but was not observed in the samples collected in Oceania. For MET, only RT176 has a high resistant rate (defined as > 0.5), and the high resistant regions are Mexico, Russia, and Poland. For MOX, RT052, 356, and 176 are 100% resistant, and the three regions with the highest resistance rates are Chile, Mexico, and Russia. Totally, the ribotypes which have a high resistance rate in > 3 drugs are 053, 137, 356, 176, and 198, and the regions with high resistance rates are Poland, Russia, and Mexico (Fig. 4a). Given the high prevalence of RIF resistance in our collection, no association with specific clades or locations was found.

We next sought to understand to what level the antibiotic susceptibility phenotype can be explained by known genetic determinants and predict the resistance to other drugs. We first identified the reduced susceptibility/resistance determinants in our genome collection using the Comprehensive Antibiotic Resistance Database (CARD 2.0.0). However, the predictive power with CARD annotation for C. difficile is poor for the MOX and RIF resistance: no isolate has resistance based on the CARD reports (0% sensitivity and 100% specificity). The van operons have been recognized as the genetic basis of acquired VAN resistance in Enterococci [31], but the dominant prevalence of the vanXY gene in our collection indicates that such a mechanism cannot explain the observed VAN reduced-susceptibility phenotypes. Due to the poor performance of CARD in predicting MOX and RIF resistance, we manually annotated resistance determinants reported in the literature [25] (rpoB and gyrAB mutations). This manual annotation largely improved the predictive power for MOX (sensitivity = 96.95%, specificity = 96.95%, specificity = 97.54%, specificity = 98.58%) under the previous study breakpoint of 32 mg/l [32] (Table S5). Based on the combined results of CARD and our curated annotation, clade 2 was noticeably enriched with reported determinants of fluoroquinolone (MOX; gyrA), RIF (rpoB), and macrolide (CLI; ermB) resistance (Fig. 4b). Strains from clade 5, particularly RT126, carried a broader variety of aminoglycoside and tetracycline resistance genes and a higher prevalence of the nucleoside resistance compared with the rest of the population (Fig. S2).

Because there are no known genetic determinants for C. difficile resistance to vancomycin and metronidazole, we performed a GWAS to identify potential novel markers of resistance. To test the performance of the GWAS pipeline used, we first conducted the analysis on MOX, the mechanism of resistance to which is well defined, and were able to identify significant signals for the SNPs Thr82Ile on gyrA and Asp426Asn on gyrB, which are both well-known markers of MOX resistance. Among potential
markers, three novel SNPs demonstrated strong association with MET reduced susceptibility: two mutations identified in CD196–1331, Tyr130Ser, and Tyr130Cys, and a SNP located in the promoter region of CD196–1331. CD196–1331 encodes a pyridoxamine 5'-phosphate oxidase family protein (POFP), an oxidoreductase putatively involved in the transfer of electrons during the synthesis of pyridoxal 5'-phosphate (vitamin B6).

Polymorphism of Toxin Genes

To elucidate the genetic basis of toxin-mediated hypervirulence in C. difficile, we analyzed the polymorphism of four toxin-associated genes, tcdB, tcdC, cdtA, and cdtB, across major ribotypes.

In total, 62 VTs of the tcdB gene were identified. VT diversity varied across different ribotypes, the ribotypes in clades 3, 4, and 5 having only one tcdB VT, whereas some ribotypes in clade 2 (RT027, 075, and 153) and clade 1 (50% of ribotypes) had 2–7 VTs. There was an evident clade-associated pattern for tcdB variation (Fig. S3A); clade 5 is distant from the other clades based on WGS or MLST data, but the tcdB gene was not divergent with other clades which suggests relative conservation of tcdB. More interestingly, though whole-genome phylogeny is clade-associated, the diversity of receptor-binding domain sequences (CROP domain) showed a different pattern: RT017 (clade 4) has similar variation to clade 1 (Fig. S3B). Besides, the VT019.1–0042 (12/12) clade 2 is worthy of attention, as the variant pattern could obviously be divided into two regions; the first half (about 1000 residues) was similar to VT017.1–0067 (33/33) (clade 4) with the same single amino acid insertion (at p310), while the latter half was consistent with other VTs in clade 2 isolates.

We identified 48 VTs for tcdC in the 32 major ribotypes with 14 VTs containing severe variants (stop-gain mutations or deletions), which generate a truncated TcdC protein (Fig. 5). Among these severe variants, we found the well-documented VT for RT027 [33, 34], RT078 [33, 35], and RT023 [36] and also identified novel variants. The nonsense mutation (C191A) and 36-base pair (bp) deletion have been reported without ribotype information [37]; here, we identified that 50% of RT153 isolates possess this tcdC VT. In addition, a 39-bp deletion has been reported in RT126 [35]; however, we also found a nonsense mutation at position 184 (C184T) for this ribotype, which is the same as RT078. Furthermore, Persson et al. reported that the Δ117 deletion was 100% specific for RT027 [33], whereas our data indicated that this single bp deletion was also present in one isolate of RT153 and in all isolates of RT176 and RT198. Also, an 18-bp deletion was observed in some isolates in clade 1 and clade 2, which may also produce non-functional proteins (Fig. 5).

Our data also demonstrated three types of structure for the binary genes: RT010 (clade 1) and RT017 (clade 4) lack cdtA and cdtB. Other ribotypes in clade 1 have incomplete cdtA and cdtB genes, lacking about 850 bp from the 3’ end of cdtA and about 1000 bp from the 5’ end of cdtB. All isolates in clades 2, 3, and 5 possessed complete cdtA and cdtB genes. The maximum-likelihood trees of cdtA and cdtB were congruent within the clade (Fig. S4). All VTs for tcdB, tcdC, cdtA and cdtB were summarized in Table S6.

DISCUSSION

In this study, we performed WGS of 1501 CDI isolates collected in two global phase III clinical trials. Our results demonstrated a concordant relationship among ribotypes, MLST clades, and the high-resolution phylogeny inferred by whole-genome SNP analysis. This finding was
Fig. 4 Prevalence of resistance based on in-vitro data (MIC) and in silico prediction (CARD). a Reduced antibiotic susceptibility/resistance rate across ribotypes, clades, and geographic region. Heatmap of proportions of antibiotic-reduced susceptibility/resistant samples for VAN, MET, RIF, and MOX are plotted from top to bottom against ribotypes, color-coded according to clade or continent of sampling; values are shown for proportions > 0.5. b The presence of resistance determinants, identified using the Comprehensive Antibiotic Resistance Database and literature reports, is indicated with a red line. The resistance determinants are indicated by gene name (top) and Antibiotic Resistance Ontology accession number (bottom), and the corresponding antibiotic class and clade are indicated by the color-coded X and Y axes, respectively. The heatmaps represent the proportions of antibiotic-reduced susceptibility/resistant samples; values are shown for proportions > 0.5. EUCAST European Committee on Antimicrobial Susceptibility Testing, MET metronidazole, MIC minimum inhibitory concentration, MOX moxifloxacin, RIF rifaximin, VAN vancomycin.
consistent with a previous report that ribotyping has high discriminative power to distinguish clonal relationships [38], and the relatively poor correlation between ribotype and ST is a comprehensive supplement to a previous study on a smaller dataset [39].

Our study generated the largest clinical-based collection of C. difficile genomes documented to date; the number of pangenome CDS identified was much greater than the previously reported estimate of 9640 CDSs [40]. Although conservation of the core genome of C. difficile was previously reported to be as low as 16% in studies using strains isolated from humans and animals [41, 42], our results showed a much more conserved genome content, with an average of 61% of core genome shared across our collection. Similarly, the number of core genes (2317) identified in clinical samples collected over 3 years is more than that of a recent study (1322) that analyzed samples across a longer time span and diverse host sources [43]. Therefore, it appears that C. difficile strains collected from clinical CDI cases share a much closer genetic relationship than those collected from different hosts, although our samples come from a large geographic span.

Given that recombination has been reported to play an important role in determining C. difficile genomic diversity [44], we further investigated the rate of recombination in our global collection. Our average recombination rates (r/m ratio), ranging from 1.4–5.0 across the five clades, were higher than previously reported. One previous study reported a C. difficile r/m ratio estimate of 0.2 using MLST genes [45]; given the use of only housekeeping genes, this value was perhaps understandably lower than that reported in the current study. Another study, using WGS data, reported a C. difficile r/m ratio of 0.63–1.13 under 95% CI [44]. Indeed, in our study, recombination rates appeared to vary by clade, with the hypervirulent clades 2 and 5 having lower r/m ratios than other clades. Nonetheless, the recombination rates of C. difficile reported here, and by others, are modest when compared with those of other bacterial species [45].

A significantly uneven distribution of clades was found across the globe. Clade 1, a heterogeneous group with a notable absence of hypervirulent ribotypes, was found in every sampling site and was the predominant clade in Oceania. Clades 2 and 5, predominantly represented by the hypervirulent ribotypes RT027 and RT078, had a stronger association with North America and Europe, respectively, which is in line with where these hypervirulent ribotypes were first reported [46, 47]. A previous study that focused on the isolates in the USA showed that the prevalence of RT027 (clade 2) was highest in the Northeast and the South, while RT014/020 (clade 1) was more common in the West, which is partially consistent with

Fig. 5 Variant types of tcdC. VTs with a stop-gain mutation or deletion in the tcdC gene. Each VT has two sequences; the upper sequence is the nucleotide sequence, and the lower sequence is the corresponding amino acid sequence. "-" indicates a deletion; "*" indicates presence of the same base or residue as the reference gene; color indicates bases or residues that differ from the reference gene; "*" with black background indicates a stop codon. VT variant type.
our data [48]. Clade 5 was the most genetically
distanced from the other clades, possibly
because of the well-established livestock origin
of its main ribotypes [47]; hence, it is possible
that the C. difficile ribotypes comprising this
clade may be subject to different selective pres-
sures than those in other clades. Clade 3 was
mostly comprised of the toxigenic RT023 ribo-
type and was predominantly collected from
Europe, both in our study and as reported by
others [49]. Clade 4 included the hypervirulent
RT017 ribotype and was significantly more
likely to be found in Asia in our dataset. RT017
has previously been reported to have caused
widespread CDI outbreaks in Europe, North
America, and particularly Asia [42]. Although
the distribution of clades was significantly cor-
related with continents, the correlation
between genetic distance and geographic dis-
tance between strains was not strong. In our
correlation analysis, stratified by ribotypes, a
significant correlation between genetic and
geographic distance was demonstrated in only
nine ribotypes (28%), of which one was RT027,
a well-reported hospital-associated ribotype.
Notably, in a pan-European surveillance study,
reduced ribotype diversity in individual coun-
tries was associated with increased antimicro-
bial resistance. Resistance to multiple
antibiotics was most evident in certain epi-
demic ribotypes, such as RT027 and RT001, but
was also notable in RT017, RT012, emerging
RT198 (exclusive to Hungary), and RT356
(exclusive to Italy) [50]. Our data suggest the
dissemination of closely related strains across
global regions, over long distances; possible
explanations for this include the formation of
spores to facilitate the remote transmission of
C. difficile without genetic changes, foodborne
transmission, or that gene flow is strong
between C. difficile strains.

Clade 5 exhibited a considerably different
resistance-gene pattern to other clades based on
in silico annotation, with a broad variety of
resistance-conferring alleles, likely because of
selective pressure from a range of antibiotics,
given that the predominant ribotype in this
clade, RT078, is also the predominant strain in
livestock entering the human food chain [47].
The accessory genome of RT078 is
indistinguishable between animal and human
isolates [51], and given it has a broad geo-
graphic distribution, high prevalence of resis-
tance determinants, and bidirectional exchange
between livestock and humans, the currently
rising prevalence of RT078 is a potential threat,
and continued global surveillance is important.
Fluoroquinolone resistance in RT027, the most
predominant ribotype in clade 2, is well docu-
mented, and elevated MICs of other antibiotics
have also been reported for RT027 isolates [1].
Our results showed that a significant proportion
(67%) of RT027 of C. difficile strains demon-
strated resistance or reduced susceptibility to all
four classes of antibiotics tested, including the
first-line antibiotics for CDI treatment, VAN
and MET. The high rates of reduced suscepti-
bility/resistance of RT027 may have conferred
an advantage in terms of its selective success
and expansion. Three novel SNPs in
CD196–1331 were significantly associated with
MET resistance: one at the promoter region and
two at amino acid position 130. CD196–1331
encodes a POFP, which has recently been shown
to be distantly related to nim genes (nimA–nimK)
encoding 5-nitroimidazole reductase enzymes
responsible for reduced MET susceptibility in
Bacteroides spp. [52]. CD196–1331 shared 47.5%
similarity with the nimE gene, the expression of
which has been identified in uncultured bacte-
ria from human stool samples [52], placing the
presence of the nimE gene within the habitat of
C. difficile. Functional and expressional analyses
are needed to confirm the potential role of
CD196–1331 in MET resistance in C. difficile.

Different ribotypes could share identical
toxin gene VTs (RT078 and RT126), and, con-
versely, multiple toxin gene VTs could occur
within a ribotype. The specific gene character of
ribotype 19 raised the possibility that some
genetic exchange events may have occurred
between ribotype 017 and 109. The diversity in
the CROP domain of tcdB is partially associated
with clades; ribotype 017 (clade 4) is indiscrete
with clade 1. Among all ribotypes, 153 was of
special interest because the divergence of its
toxin gene VTs was higher than that of other
ribotypes. Totally, our analysis of toxin VTs
(tcdB, tcdC, cdtAB) demonstrated the diversity
of the genetic reservoir for these genes.
Despite the aforementioned insight into the global population characteristics of C. difficile, our study has some limitations. Although the isolates were collected globally, the trials were not designed for the purpose of a balanced global sampling of C. difficile, but to investigate the safety and efficacy of monoclonal antibodies on preventing CDI recurrence in countries/regions where the disease is prevalent, so the isolates were not perfectly randomly sampled to present a representation of global epidemiology. We also expect further studies to establish WGS-based antibiotic susceptibility/resistance prediction models for C. difficile and corresponding experiment to validate the potential resistant determinates. In addition, the role of well-known toxin profiles (TcdB+, CDT+, incomplete TcdC) are partially associated with hypervirulent ribotypes, so other virulence targets should be investigated based on clinical severity phenotypes.

**CONCLUSIONS**

Using samples from participants with clinically diagnosed CDI from across the globe, this analysis improved understanding of genomic, geographic, and clinical features of CDI infection in a clinically relevant epidemic C. difficile population and will serve as a valuable resource for future studies. These findings indicated that within these epidemic populations, the core genome was more highly conserved than previously reported, with modest recombination rates. Genetic variation within the entire genome was most recapitulated by the MLST data. Moreover, material gene transfer (i.e., recombination) was a negligible signal in the genome. Highly distinct geographic prevalence of clade may be related to local selection pressures such as use of certain antibiotics. Antibiotic susceptibility rates are variable among the ribotype/clade and geographic location, which may result from localized selection pressures as suggested by the inverse correlation observed between ribotype diversity and antibiotic resistance in European surveillance samples [50]. The genetics of antibiotic resistance clearly resides on the genomic DNA, as resistance has been shown to be a co-variate with clade designation. The novel potential resistant determinants of MET susceptibility and the novel severe mutations on tcdC need further validation to understand any implications. All these results provide valuable insight into the spread and characteristics of epidemic C. difficile to potentially support future investigation. Better understanding of the spread of hypervirulent strains and antibiotic resistance may aid development of more effective treatment plans, drug development, and ultimately a better outcome for patients.

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**Compliance with Ethics Guidelines.** MODIFY I (NCT01241552) and MODIFY II (NCT01513239) were conducted in accordance with Good Clinical Practice guidelines and the provisions of the Declaration of Helsinki. The protocols and amendments were approved by the institutional review board or independent ethics committee at each study site (BGI IRB; IRB #13029). Written informed consent for specimen collection and use was provided by all participants before the trial began.

**Data Availability.** The datasets generated during and/or analysed during the current study are available in the CNGB Nucleotide Sequence Archive repository, [https://db.cngb.org/cnsa](https://db.cngb.org/cnsa) (accession number CNP0000099). To make our data accessible, user-friendly, and interactive, we developed the C. difficile Pathogen Database ([http://dev.dc.org/pathodb/](http://dev.dc.org/pathodb/)), which not only allows deposition of the genome sequence data and corresponding phenotype data for downloading but also automatically performs phylogenetic and toxin analysis for a given user’s data.

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