Salmonella enterica Serovar Typhi in Bangladesh: Exploration of Genomic Diversity and Antimicrobial Resistance

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ABSTRACT Typhoid fever, caused by Salmonella enterica serovar Typhi, is a global public health concern due to increasing antimicrobial resistance (AMR). Characterization of S. Typhi genomes for AMR and the evolution of different lineages, especially in countries where typhoid fever is endemic such as Bangladesh, will help public health professionals to better design and implement appropriate preventive measures. We studied whole-genome sequences (WGS) of 536 S. Typhi isolates collected in Bangladesh during 1999 to 2013 and compared those sequences with data from a recent outbreak in Pakistan reported previously by E. J. Klemm, S. Shakoor, A. J. Page, F. N. Qamar, et al. (mBio 9:e00105-18, 2018, https://doi.org/10.1128/mBio.00105-18), and a laboratory surveillance in Nepal reported previously by C. D. Britto, Z. A. Dyson, S. Duchene, M. J. Carter, et al. [PLoS Negl. Trop. Dis. 12(4):e0006408, 2018, https://doi.org/10.1371/journal.pntd.0006408]. WGS had high sensitivity and specificity for prediction of ampicillin, chloramphenicol, co-trimoxazole, and ceftriaxone AMR phenotypes but needs further improvement for prediction of ciprofloxacin resistance. We detected a new local lineage of genotype 4.3.1 (named lineage Bd) which recently diverged into a sublineage (named Bdq) containing qnr genes associated with high-level ciprofloxacin resistance. We found a ceftriaxone-resistant isolate with the blaCTX-M-15 gene and a genotype distinct from the genotypes of extensively drug-resistant (XDR) isolates from Pakistan. This result suggests a different source and geographical origin of AMR. Genotype 4.3.1 was dominant in all three countries but formed country-specific clusters in the maximum likelihood phylogenetic tree. Thus, multiple independent genetic events leading to ciprofloxacin and ceftriaxone resistance took place in these neighboring regions of Pakistan, Nepal, and Bangladesh. These independent mutational events may enhance the risk of global spread of these highly resistant clones. A short-term global intervention plan is urgently needed.

IMPORTANCE Typhoid fever, caused by Salmonella enterica serovar Typhi, is responsible for an estimated burden of approximately 17 million new episodes per year worldwide. Adequate and timely antimicrobial treatment invariably cures typhoid fever. The increasing antimicrobial resistance (AMR) of S. Typhi severely limits the treatment options. We studied whole-genome sequences (WGS) of 536 S. Typhi isolates collected in Bangladesh between 1999 and 2013 and compared those sequences with data from a recent outbreak in Pakistan and a laboratory surveillance in Nepal. The analysis suggests that multiple ancestral origins of resistance against ciprofloxacin and ceftriaxone are present in three countries. Such independent genetic events and subsequent dissemination could enhance the risk of a rapid global spread of these clones.
spread of these highly resistant clones. Given the current treatment challenges, vaccination seems to be the most appropriate short-term intervention to reduce the disease burden of typhoid fever at a time of increasing AMR.

KEYWORDS Bangladesh, *Salmonella Typhi*, antibiotic resistance, genomics

Typhoid fever is a life-threatening infectious disease caused by *Salmonella enterica* serovar Typhi. *S. Typhi* colonizes only humans, is transmitted through the fecal-oral route, and is endemic in tropical countries, especially in Africa and South and Southeast Asia. Worldwide, approximately 17 million people are infected every year by this pathogen (1–4). Though the mortality rate remains low (<1%), 1 in 20 to 25 cases experiences residual disability (5).

Adequate and timely antimicrobial treatment invariably cures typhoid fever. However, the increasing antimicrobial resistance (AMR) of *S. Typhi* limits the treatment options. In spite of suggested regional decreases in the levels of antibiotic resistance (6–8), the first cases of *S. Typhi* isolates showing multidrug resistance (MDR) (defined as co-occurring resistance to ampicillin [amp], chloramphenicol [chl], and co-trimoxazole [sxt]) were reported in the early 1970s (9, 10). Ciprofloxacin (cip) resistance first emerged in the early 1990s. At present, over 90% of clinical isolates from regions of endemcity show reduced susceptibility to ciprofloxacin (6, 7, 11). These events shifted the first-line and empirical treatments to other classes of antimicrobial agents, such as ceftriaxone (cro) and azithromycin. Alarmingly, reports of resistance against these agents have now been published (6, 12–18). Moreover, a recent report from Pakistan described the first large-scale outbreak of an *S. Typhi* clone that is extensively drug resistant (XDR; defined as MDR plus resistance to ciprofloxacin and ceftriaxone) (12).

Whole-genome sequence (WGS)-based approaches using next-generation sequencing (NGS) have become effective tools for the study of genetic diversity and prediction of resistance phenotypes (12, 19–25). Several studies have correlated WGS data with various resistance phenotypes in *S. Typhi* (12, 20, 25, 26). However, most of these studies involved small numbers of isolates from multiple countries, isolates from single outbreaks, or clusters of travel-related typhoid cases, which do not accurately represent the situation in countries where typhoid fever is endemic over longer periods of time (12, 20, 25–29). These shortcomings limit our overall understanding of the dynamics of typhoid fever in regions of endemcity, especially in South Asia, where the disease burden is high.

We generated a WGS data set of 536 *S. Typhi* strains, which were mostly isolated from the blood of pediatric patients in Bangladesh over a period of 15 years (1999 to 2013). In this study, we explored the phenotypic and genotypic diversity of these isolates using whole-genome single nucleotide polymorphism (wgSNP) analysis, classical multilocus sequence typing (MLST), and core genome MLST (cgMLST). We also examined the phylogenetic relationships between these isolates and compared the results with two published data sets from two neighboring countries, representing a hospital-based surveillance study in Nepal and an outbreak during 2016 to 2017 in Pakistan (12, 29). Additionally, we investigated the utility of NGS data for the prediction of phenotypic resistance to multiple antibiotics. We focused on the genes involved in MDR and mutations in the DNA gyrase (*gyrA* and *gyrB*) and topoisomerase IV (*parC* and *parE*) enzymes that lead to ciprofloxacin resistance.

RESULTS

Genotypic diversity of *S. Typhi* in Bangladesh. Among the 539 strains presumptively identified as *S. Typhi*, 536 (99%) were confirmed to be *S. Typhi* by WGS-based serotyping and were analyzed further. A total of 61% (329/536) of them were from hospitalized patients. Genotype 4.3.1 was dominant (65%; 350/536), followed by genotype 3.3 (13%; 69/536), genotype 3.2.2 (11%; 61/536), and 12 other genotypes (Table 1). Classical MLST analysis revealed the presence of only three different sequence types (ST) among our isolates, namely, ST1 (n = 351), ST2 (n = 166), and ST2209 (n =
TABLE 1 Genotyping and haplotyping results for the 536 S. Typhi isolates from Bangladesh based on WGS

| Genotype | No. of isolates | % of total | Haplotype |
|----------|-----------------|------------|-----------|
| 4.3.1    | 350             | 65.30      | H58       |
| 3        | 69              | 12.87      | H1        |
| 3.2.2    | 61              | 11.38      | H1        |
| 2        | 18              | 3.36       | NT        |
| 2.3.3    | 18              | 3.36       | NT        |
| 2.1.7    | 4               | 0.75       | H8        |
| 2.0.1    | 3               | 0.56       | NT        |
| 2.2      | 3               | 0.56       | NT/H39    |
| 1.2.1    | 2               | 0.37       | NA<sup>a</sup> |
| 2.5      | 2               | 0.37       | H55       |
| 3        | 2               | 0.37       | H13       |
| 3.0.1    | 2               | 0.37       | H13       |
| 3.0.2    | 1               | 0.19       | H13       |
| 4.1      | 1               | 0.19       | H52/15/10/67 |
| Total    | 536             | 100        |           |

<sup>a</sup>NA, not available.

18) (Fig. 1a), all of which had similar ratios of hospitalized and outpatient cases (~60% versus ~40%). Overall, 99% of the ST1 strains had the 4.3.1 genotype (349/351), while all of the ST2209 strains had genotype 2.3.3 (18/18; 100%). Other genotypes were within ST2 (Fig. 1a). One isolate was nontypeable (NT) by MLST analysis. With respect to the haplotyping scheme, haplotype 58 (H58) was dominant (65%; 350/536), followed by H1 (130/536; 24%) (Table 1).

**Phylogenetic relationships and new lineages.** Genotypes 3.2.2 and 3.3 clustered together in the same classical MLST type (ST2) but formed two distinct subclades in the unweighted pair group method using average linkages (UPGMA) tree based on cgMLST analyses of 3,002 core loci (Fig. 1a and b). Like genotype 2.3.3, genotype 4.3.1 also generated its own subclade in the tree, with the notable presence of multiple subgroups (Fig. 1b). The same UPGMA tree presented a distinct population structure and a similar genotypic differentiation as observed in the maximum likelihood tree (MLT), which was based on 2,328 SNPs from our WGS data compared to the S. Typhi CT18 reference genome (Fig. 1c) (30).

Genotype 1.2.1 mapped closest to the root of the MLT; genotype 4.3.1 was the most remote (Fig. 1c). As the dominant genotype, 4.3.1 formed a large subclade, with genotype 4.1 in its primary clade. Another primary clade divided into two major subclades, genotypes 3.3 and 3.2.2, which comprised the second and third most prevalent genotypes in Bangladesh. Two small subclades of genotype 2.3.3 and genotype 2.0 were also present in the MLT, rooting with genotype 2.2 and 2.1.7, respectively.

The comparative MLT created using our data and the strains from neighboring countries showed country-specific clusters inside the dominant 4.3.1 genotype and also in genotypes 3.3 and 3.2.2 (Fig. 2 and 3). This suggests different points of origin for the various lineages of S. Typhi in each country. All XDR Pakistani isolates extended into a single branch of the MLT, showing H58 lineage Ia and a minimally divergent pattern (Fig. 3; see also Fig. S1 in the supplemental material), whereas the Nepali strains were dominant in lineage II. The isolates from Bangladesh included only four isolates from lineage II but showed two distinct clusters inside genotype 4.3.1 (H58): lineage Ia (n = 223) and a previously undescribed lineage (n = 108; Fig. 3). The latter lineage did not match the SNP definition of H58 lineage I or lineage II, suggesting a previously undetected H58 lineage (genotype 4.3.1). On the basis of the MLT, this undescribed lineage could have had the same point of origin as lineage I but then clearly followed a different pattern of divergence and formed its own subclade inside genotype 4.3.1 (Fig. 3). This new H58 lineage can be distinguished by the SNPs at nucleotide position 561056 (C→A) and 2849843 (A→C) of the CT18 reference genome (this previously undescribed lineage is referred to as “lineage Bd” in the remainder of the article).
Resistance phenotypes and genotypes. On the basis of analyses performed with five different antibiotics—ampicillin (amp), chloramphenicol (chl), co-trimoxazole (sxt), ciprofloxacin (cip), and ceftriaxone (cro)—the 536 S. Typhi isolates from Bangladesh were found to harbor 12 different phenotypic resistance profiles (phenotypes; Table 2). Isolates with the “MDR, cip-R” profile (n = 202) were most prevalent in our library, followed by “cip-R only” (n = 169) and “Susceptible to all” (n = 62). A comparison of MDR and ciprofloxacin-resistant isolates with different genotypes is presented in Table 3. The single ceftriaxone-resistant (cro-R) strain (MIC > 32 μg/ml) (susceptible to sxt, chl, and cip) in our library, isolated in 2000, displayed genotype 3.3 (haplotype H1) and contained the bla_hap1_CTX_M−15 gene (ceftriaxone resistance). The other resistance genes
detected are listed in Table 4, including *bla*<sub>TEM-1B</sub> (ampicillin resistance); *catA1* (chloramphenicol resistance); *dfrA7*, *sul1*, and *sul2* (co-trimoxazole resistance); and *qnrS1* (ciprofloxacin resistance).

**Comparison of phenotypic and WGS-derived resistance profiles.** On the basis of the resistance genes identified for the five antimicrobial agents, a WGS resistance (WGS-res) profile was assigned and compared with the phenotypic profile of each isolate to evaluate the ability of the WGS approach to predict the resistance phenotype (Table 5). For all antimicrobial agents except ciprofloxacin, the two profiles corresponded at a level of 99% for the resistant isolates. In contrast, some susceptible isolates (*n* = 33) harbored resistance genes, which reduced the specificity of the method (≥91%). Three of them had truncated genes (considered inactive genes; Table 5), but the other 30 isolates had the complete coding sequences without any phenotypic resistance. This might suggest impairments (e.g., transcriptomic, translational, protein modification, etc.) in downstream steps of the resistance pathway or the presence of counteracting genes.
On the other hand, for isolates with resistant phenotypes but susceptible WGS res-profiles, we screened for mutations in genes with efflux pump or membrane permeability functions (see Table S1 in the supplemental material). However, no relevant patterns were detected for AMR.

**Ciprofloxacin resistance, background mutations, and genotypes.** The resistance gene analysis identified the *qnrS1* gene in 55 isolates (Table 4) and detected a number of different mutations (*n* = 24) in the *gyrA* and *gyrB* genes encoding DNA gyrase and in the topoisomerase IV enzyme *parC* and *parE* genes (Table 6, columns 1 to 3). The most prevalent mutation was *gyrA* D538N (*n* = 352), followed by *gyrA* S83F (*n* = 299), *gyrA* S83Y (*n* = 125), and *parE* A364V (*n* = 69). On the basis of mutations in *gyrA/B* and *parC/E* genes, 34 cip-mutation profiles were generated and compared with the ciprofloxacin MIC of each isolate (Table 6, columns 4 and 5) (Fig. 4 and 5). All of the profiles, apart from *gyrA* D538N and *parE* A364V, were associated with resistance (MIC > 0.06 μg/ml). Two different profiles with triple mutations (*gyrA*<sub>D87G</sub> *gyrA*<sub>S83F</sub> *parC*<sub>E84K</sub> for

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**FIG 3** Comparison of genotype 4.3.1 (H58) isolates from Bangladesh, Pakistan, and Nepal in a wgSNP-derived MLT. No singleton was considered in the consensus SNP data. The tree is colored by country. Different data points, including lineage, sublineage (details), presence of different gyrA-83 mutations, MDR, cip resistance, presence of *qnr* genes, and cro resistance phenotypes are indicated (by colors) in different circles around the tree.
eight isolates and gyrA<sub>D87N</sub> gyrA<sub>S83F</sub> parC<sub>S80I</sub> for one isolate) had median MICs of \( \geq 8.0 \mu g/ml \) (Fig. 5) (Table 6, columns 4 and 5). Profiles with qnr genes also had MICs of \( \geq 1.0 \mu g/ml \) (Fig. 4 and 5). No mutations were present in 18 isolates, but 3 of them showed resistance to ciprofloxacin (MIC of 0.25 to 0.5 \( \mu g/ml \); Fig. 4).

Mutations in codon 83 of gyrA (S83F and S83Y) were the most prevalent among our ciprofloxacin-resistant isolates (411/467; 88%) (Table 6, columns 1 to 3). The S83Y mutation (122/411) was closely associated with genotype 4.3.1 in Bangladesh (123/125; 98%) and was present in 96% of our H58 lineage Bd isolates (104/108; Fig. 3). In contrast, 89% of our lineage Ia isolates had the S83F mutation (198/223) and exhibited lower mean (0.74 versus 1.71 \( \mu g/ml \)) and median (0.25 versus 0.5 \( \mu g/ml \)) ciprofloxacin MIC values than the lineage Bd isolates. The latter lineage also displayed more divergence than lineage I in Bangladesh (Fig. 3; mean pairwise distances, 12.8 versus 11.2). The root of lineage Bd contained isolates detected at earlier time points (1999 to 2004) and formed a noticeable subclade at the tip composed of isolates (\( n = 55 \)) collected from 2006 onward (Fig. 3; see also Fig. 52 and 53). In addition, this small subclade had the universal presence of the qnr gene and MIC values of \( \geq 1.0 \mu g/ml \) (Fig. 3). This small, qnr-specific subclade within lineage Bd can be defined by SNPs at

### Table 2

| Phenotype                        | No. of isolates | % of total |
|----------------------------------|-----------------|------------|
| MDR, cip-R                       | 202             | 37.69      |
| MDR                              | 4               | 0.75       |
| amp-R, sxt-R, cip-R              | 1               | 0.19       |
| amp-R, chi-R, cip-R              | 2               | 0.37       |
| amp-R, cip-R                     | 53              | 9.89       |
| amp-R, cro-R                     | 1               | 0.19       |
| sxt-R, chi-R, cip-R              | 25              | 4.66       |
| sxt-R, chi-R                     | 1               | 0.19       |
| chi-R, cip-R                     | 15              | 2.80       |
| chi-R only                       | 1               | 0.19       |
| cip-R only                       | 169             | 31.53      |
| Susceptible to all               | 62              | 11.57      |
| **Total**                        | **536**         | **100%**   |

*Five different antibiotics were considered: ampicillin (amp), co-trimoxazole (sxt), chloramphenicol (chi), ciprofloxacin (cip), and ceftriaxone (cro). MDR (multidrug resistance) refers to co-occurring resistance to amp, sxt, and chi. “S” and “R” refer to susceptible and resistant phenotypes, respectively (interpretations according to EUCAST-2018).*

### Table 3

| Genotype | MDR | cip-R |
|----------|-----|-------|
|          | Yes | No    | Yes | No |
| 1.2.1    | 0   | 2     | 0   | 2  |
| 2        | 1   | 17    | 6   | 12 |
| 2.0.1    | 0   | 3     | 3   | 0  |
| 2.1.7    | 0   | 4     | 2   | 2  |
| 2.2      | 0   | 3     | 2   | 1  |
| 2.3.3    | 0   | 18    | 8   | 10 |
| 2.5      | 0   | 2     | 0   | 2  |
| 3        | 0   | 2     | 1   | 1  |
| 3.0.1    | 0   | 2     | 1   | 1  |
| 3.0.2    | 0   | 1     | 0   | 1  |
| 3.2.2    | 0   | 61    | 52  | 9  |
| 3.3      | 0   | 69    | 52  | 17 |
| 4.1      | 0   | 1     | 1   | 0  |
| 4.3.1    | 205 | 145   | 339 | 11 |
| **Total**| **206** | **330** | **467** | **69** |

**Comparison between multidrug resistance (MDR) and ciprofloxacin resistance (cip-R) with genotypes among Bangladesh isolates**
nucleotide positions 1253109 (T→G), 2385340 (A→G), 2676540 (A→T), and 2688285 (C→T) of the CT18 reference genome (and is referred to as sublineage Bdq in the rest of this article). Comparison with other lineages in our cohort of isolates from Bangladesh revealed that sublineage Bdq had a very high median ciprofloxacin MIC (4 μg/ml; Table 4).

| Resistance gene | Antibiotic class | No. of isolates | % of total | Phenotype | Matched NCBI accession no. |
|-----------------|------------------|-----------------|------------|-----------|---------------------------|
| *bla*TEM-1B     | Beta-lactam      | 271             | 50.28      | amp-R     | JF910132                  |
| *bla*CTX-M-15   | Beta-lactam      | 1               | 0.19       | cro-R     | DQ302097                  |
| catA1           | Phenicol         | 256             | 47.50      | chl-R     | V00622                    |
| dfrA7           | Trimethoprim     | 257             | 47.68      | tmp-R     | JF806498                  |
| qnrS1           | Quinolone        | 55              | 10.2       | cip-R     | AB187515                  |
| sul1            | Sulfonamide      | 257             | 47.68      | sul-R     | CP002151                  |
| sul2            | Sulfonamide      | 265             | 49.17      | sul-R     | HQ840942, FJ197818, GQ421466 |
| tet(A)          | Tetracycline     | 51              | 9.46       | tet-R     | AJS17790                  |
| tet(B)          | Tetracycline     | 46              | 8.53       | tet-R     | AF326777                  |

The table columns list each gene name, the antimicrobial class that it works against, the number of isolates that contained the gene, the percentage of isolates that contained the gene, the resulting resistance phenotype, and the NCBI gene accession number.

TABLE 5 Evaluation of the ability of WGS-res profiles to predict *S. Typhi* resistance phenotypes of our isolates

| Antimicrobial resistance category | Presence of gene(s) and WGS-res profile | No. of isolates with indicated phenotype | Sensitivity (%)<sup>b</sup> | Specificity (%)<sup>c</sup> |
|----------------------------------|-----------------------------------------|----------------------------------------|---------------------------|---------------------------|
| Ampicillin resistance            | Total                                    | 263/273                                | 99.6                      | 97.4                      |
|                                  | *bla*TEM-1B                              | 262/7                                  | 99.6                      | 97.4                      |
|                                  | Truncated *bla*TEM-1B                    | 0/2                                    | 99.6                      | 97.4                      |
|                                  | No *bla*TEM-1B                           | 1/264                                  | 99.6                      | 97.4                      |
|                                  | WGS-res profile: resistant                | 262/7                                  | 99.6                      | 97.4                      |
|                                  | WGS-res profile: susceptible              | 1/266                                  | 99.6                      | 97.4                      |
| Co-trimoxazole resistance        | Total                                    | 233/303                                | 99.1                      | 91.4                      |
|                                  | *dfrA7* + *sul1* + *sul2*                | 205/4                                 | 99.1                      | 91.4                      |
|                                  | *dfrA7* + *sul1* only                    | 26/22                                 | 99.1                      | 91.4                      |
|                                  | *sul2* only                              | 1/55                                 | 99.1                      | 91.4                      |
|                                  | None of three                            | 1/222                                 | 99.1                      | 91.4                      |
|                                  | WGS-res profile: resistant                | 231/26                                | 99.1                      | 91.4                      |
|                                  | WGS-res profile: susceptible              | 2/277                                 | 99.1                      | 91.4                      |
| Chloramphenicol resistance       | Total                                    | 250/286                                | 99.2                      | 97.6                      |
|                                  | *catA1*                                  | 248/7                                 | 99.2                      | 97.6                      |
|                                  | Truncated *catA1*                        | 0/1                                  | 99.2                      | 97.6                      |
|                                  | No *catA1*                               | 2/278                                 | 99.2                      | 97.6                      |
|                                  | WGS-res profile: resistant                | 248/7                                 | 99.2                      | 97.6                      |
|                                  | WGS-res profile: susceptible              | 2/279                                 | 99.2                      | 97.6                      |
| Ceftriaxone resistance           | Total                                    | 1/535                                 | 100.0                    | NA                       |
|                                  | *bla*CTX-M15                              | 1/0                                  | 100.0                    | NA                       |
|                                  | No *bla*CTX-M15                          | 0/535                                 | 100.0                    | NA                       |
|                                  | WGS-res profile: resistant                | 1/0                                  | 100.0                    | NA                       |
|                                  | WGS-res profile: susceptible              | 0/535                                 | 100.0                    | NA                       |

<sup>a</sup>Four antimicrobials were considered (ampicillin, co-trimoxazole, chloramphenicol, and ceftriaxone); resistance to these agents is caused mainly by acquisition of resistance genes.

<sup>b</sup>Sensitivity data represent proportions of isolates identified as phenotypically resistant by the WGS-res profile.

<sup>c</sup>Specificity data represent proportions of isolates identified as phenotypically susceptible by the WGS-res profile.

<sup>d</sup>For co-trimoxazole (sxt), we considered the presence of *dfrA7*, plus *sul1* and/or *sul2* genes to exert the resistance (R) phenotype.

<sup>e</sup>A total of 206 detected *sul2* genes matched three different GenBank IDs: FJ197818 (n = 74), GQ421466 (n = 1), and HQ840942 (n = 131). Of the four *sul2* genes, two matched FJ197818 and two HQ840942. One *sul1* gene had unreliable bases (N) in its sequence; that result was considered a sequencing error, and the complete sequence was used in calculations.

<sup>f</sup>One *sul1* gene had unreliable bases (N) in its sequence; that result was considered a sequencing error, and the complete sequence was used in calculations.

<sup>g</sup>Of the four *sul2* genes, two matched FJ197818 and two HQ840942. One *sul1* gene had unreliable bases (N) in its sequence; that result was considered a sequencing error, and the complete sequence was used in calculations.

<sup>h</sup>Of 54 genes matched GQ421466 and one HQ840942.

<sup>i</sup>Only *sul2* genes that matched HQ840942 had complete sequences. Genes that matched FJ197818 and GQ421466 were either truncated or mutated.

<sup>j</sup>All *catA1* gene sequence had one silent mutation in amino acid 195 (lysine) (CTG→TTG)

TABLE 4 List of resistance genes detected in our isolates<sup>a</sup>

| Resistance gene | Antibiotic class | No. of isolates | % of total | Phenotype | Matched NCBI accession no. |
|-----------------|------------------|-----------------|------------|-----------|---------------------------|
| *bla*TEM-1B     | Beta-lactam      | 271             | 50.28      | amp-R     | JF910132                  |
| *bla*CTX-M-15   | Beta-lactam      | 1               | 0.19       | cro-R     | DQ302097                  |
| catA1           | Phenicol         | 256             | 47.50      | chl-R     | V00622                    |
| dfrA7           | Trimethoprim     | 257             | 47.68      | tmp-R     | JF806498                  |
| qnrS1           | Quinolone        | 55              | 10.2       | cip-R     | AB187515                  |
| sul1            | Sulfonamide      | 257             | 47.68      | sul-R     | CP002151                  |
| sul2            | Sulfonamide      | 265             | 49.17      | sul-R     | HQ840942, FJ197818, GQ421466 |
| tet(A)          | Tetracycline     | 51              | 9.46       | tet-R     | AJS17790                  |
| tet(B)          | Tetracycline     | 46              | 8.53       | tet-R     | AF326777                  |
Fig. S4) and low divergence (mean pairwise distance, 8.4). Many isolates from Pakistan in lineage Ia also contained *qnr* genes (Fig. 3) but showed no specific divergence pattern.

In total, 11 ciprofloxacin-resistant isolates (11/467; 2%) did not have any other mutation in DNA gyrase and topoisomerase IV genes, leading to an estimated sensi-

| Gene | Mutation | No. of mutations | Mutation combination (profile) | No. of mutation profiles |
|------|----------|------------------|--------------------------------|-------------------------|
| *gyrA* | D538N | 352 | *gyrA*-D538N, *gyrA*-S83F | 179 |
| | S83F | 299 | *gyrA*-D538N, *gyrA*-S83Y | 120 |
| | S83Y | 125 | *gyrA*-S83F | 66 |
| | D87N | 30 | *gyrA*-S83F, *parE*-A364V | 26 |
| | N529S | 17 | *parE*-A364V | 18 |
| | D87G | 11 | *gyrA*-N529S, *gyrB*-S464F | 17 |
| | D87Y | 4 | *gyrA*-D87N, *parE*-A364V | 15 |
| | A119E | 1 | *gyrA*-D538N, *gyrA*-D87N | 12 |
| | D87A | 1 | *gyrA*-D538N, *gyrA*-S83F, *parE*-T447A | 9 |
| *gyrB* | S464F | 21 | *gyrA*-D538N | 8 |
| | S464Y | 10 | *gyrA*-D538N, *gyrA*-D87G, *gyrA*-S83F, *parC*-E84K | 8 |
| *parC* | E84K | 10 | *gyrB*-S464Y | 8 |
| | S80R | 2 | *gyrB*-S464F | 3 |
| | D69A | 2 | *gyrA*-D538N, *gyrA*-D87Y | 2 |
| | T620M | 1 | *gyrB*-S464Y, *parE*-A364V | 2 |
| | E84G | 1 | *gyrA*-D87Y | 2 |
| | S80I | 1 | *gyrB*-S83Y, *parC*-D69A, *parE*-A364V | 2 |
| *parE* | A364V | 69 | *gyrA*-D538N, *gyrA*-S83F, *parE*-L416F | 2 |
| | T447A | 9 | *gyrA*-D87N, *parE*-A364V, *parE*-S339L | 2 |
| | L416F | 2 | *gyrA*-D538N, *gyrA*-S83Y, *parE*-A365S | 2 |
| | S339L | 2 | *gyrA*-D538N, *gyrA*-S83F, *parC*-E84K | 2 |
| | A365S | 2 | Other combination pattern (one isolate for each) | 13 |
| | L502F | 1 | No mutation | 18 |
| | E460K | 1 | | |

Total 536

Table 6: Mutations detected in DNA gyrase (*gyrA* and *gyrB* genes) and topoisomerase IV (*parC* and *parE* genes) individually, and combined mutation profiles based on them.

**FIG 4** Mutation profiles detected in genes associated with ciprofloxacin resistance in our isolates and correlation with ciprofloxacin MIC. The horizontal red line indicates the threshold MIC level (0.06 μg/ml) of resistance (according to EUCAST v8.0).
tivity of 98% for the WGS method in correctly predicting ciprofloxacin resistance. In contrast, 36 of 69 ciprofloxacin-susceptible isolates had at least one mutation (not linked to a specific genotype) in one of these four genes (specificity/H11005 52%).

Comparison with neighboring countries. All genotype 4.3.1 isolates from Bangladesh (this study), Nepal (surveillance in Kathmandu), and Pakistan (outbreak in Sindh) had the same \( \text{gyrA} \) D538N mutation (Fig. S5). Likewise, the \( \text{parE} \) A364V mutation was present in all genotype 3.3 isolates from Bangladesh (70/70) and Nepal (17/19) and in genotype 3.3.1 (3/3) isolates from Nepal. Genotype 2.0 isolates from Bangladesh also had the \( \text{gyrA} \) N529S mutation present (94%; 17/18). However, none of these mutations seemed to have any association with AMR (Fig. 4).

Comparisons performed with the \( \text{bla}_{\text{CTX-M-15}} \) gene sequence of our ceftriaxone-resistant isolate revealed 92% coverage and 99% identity with the XDR isolate (GenBank accession no. LT906492.1) from the Pakistani outbreak (Table 7). In contrast, the \( \text{bla}_{\text{CTX-M-15}} \) gene from Bangladesh shared complete homology with the \( \text{Klebsiella pneumoniae} \) \( \text{bla}_{\text{CTX-M-15}} \) gene (FJ815436.1). A detailed comparison of our sequence data with the sequences of the isolates from Pakistan and Nepal is presented in Table 7.

**DISCUSSION**

5. Typhi multilocus sequence types and other genotypes in Bangladesh. Genotyping and the phylogenetic inferences agreed with the genotyping framework interpretation (27) and showed genotype 4.3.1 (haplotype 58, H58) to be dominant among the isolates from Bangladesh (Table 1). This was no surprise, as this genotype possibly emerged from South Asia in the early 1990s and now dominates in regions of typhoid...
endemicity in the world (26, 31). The same genotype was also dominant among the isolates from Nepal and Pakistan (Table 7) (12, 29). On the other hand, classical MLST revealed only three sequence types (ST), with dominance of ST1 and ST2, which accords with global MLST report (32). There are 46 complete MLST types available for S. Typhi (33). Interestingly, the third most common MLST type in our data, ST2209, had a complete match with genotype 2.3.3 (100%; 18/18) (Fig. 1a). Isolates of this genotype from 2013 (n/H11005 8) had the same mutation (gyrB S464Y). Five of 8 had a cip-resistant phenotype, which could indicate the beginning of new clonal dissemination (see Fig. S2 in the supplemental material).

Moreover, 99% (349/351) of all ST1 isolates from Bangladesh belonged to genotype 4.3.1 (Fig. 1a). This association was previously described in a small study involving 32 isolates (34). A phylogeographical report of S. Typhi included an estimate that divergence for genotype 4.3.1 commenced in the very late 1980s (26). However, the presence of ST1 could be detected before the 1980s (33, 35), as is likely the case for H58.

Presence of genotype-specific mutations. Isolates with genotype 4.3.1 from all three countries shared a common but as-yet-unreported mutation, gyrA D538N (nucleotide position 2332398 of the CT18 genome; Fig. S5). This mutation is not linked to ciprofloxacin resistance (Fig. 4 and 5) but could be crucial to the structure of the DNA gyrase enzyme, considering the associated change in the isoelectronic point (pI; D−N: 2.77 → 5.41) of the amino acid due to this mutation (36). Similar associations were also observed between genotype 2.0 and gyrA N529S (hydrophobicity, 3.47 → 1.83), and

| TABLE 7 Comparison of the isolates from Bangladesh with isolates described in other studies from two neighboring countriesa |
|-----------------------------------------------|----------------|----------------|
| **Criterion**                               | Bangladesh (present study) | Nepal (29) | Pakistan (12) |
| Sample source                               | Hospital surveillance (hospitalized and outpatient services) | Laboratory surveillance of typhoidal Salmonella | Outbreak |
| Timeline                                    | 1999–2013 | 2008–2016 | November 2016–March 2017 |
| No. of S. Typhi samples analyzed            | 536 | 198 | 100 |
| Age limit                                   | <18 yrs for hospitalized cases; no age limit for outpatient cases | <14 yrs | None |
| No. of MDR or XDR isolates                  | MDR, 206 (38%); XDR, none | MDR, 6 (0.03%); XDR, none | MDR, 89 (89%); XDR, 87 (87%) |
| No. of isolates with ciprofloxacin resistance (cro-R) | 467 (87%) | 171 (86%) | 96 (96%) |
| No. of isolates with ceftriaxone resistance (cro-R) | 1 (0.2%) (caused by bla_{CTX-M15}) | None | 88 (88%) (caused by bla_{CTX-M15}) |
| Genotype of cro-R isolate(s)                | 3.3 | NA | 4.3.1 (lineage Ia) |
| Phenotype of cro-R isolate(s)               | amp-R, cro-R | NA | XDR |
| blas_{CTX-M15} identity and coverage        | 92% coverage and 99% identity with gene sequence from Pakistan | NA | NA |
| No. of isolates with indicated dominant genotypes | 4.3.1 (H58), 350 (65%); 3.3 (H1), 69 (13%); 3.2.2 (H1), 61 (11%) | 4.3.1 (H58), 154 (78%); 3.3.0, 19 (10%) | 4.3.1 (H58), 99 (99%) |
| No. of isolates with indicated dominant H58 lineage(s) | Ia, 223 (63% of H58); Bd, 108 (31% of H58) | I, 21 (10% of H58); II 133 (67% of H58) | Ia, 92 (92% of H58) |
| Local lineage(s) detected?                  | Yes; lineage Bd (108 isolates [31% of H58]) and sublineage Bdq (55 isolates [16% of H58]) | Yes; local lineage II (no. of isolates not given) | No; a clone of lineage Ia with possible local origin |
| AMR details of local lineage                | All sublineage Bdq contain qnr genes; 88% have cip-MIC ≥1 μg/ml (median, 4 μg/ml) | (a) Intermediate resistance to CIP; (b) no MDR; (c) contains gyrA-S83F mutations | Not a lineage but a clone of lineage Ia; predominantly XDR |
| Time of emergence for local lineages        | See Fig. S3 | Possibly after 2008 | November 2016–present |

aMDR, multidrug resistance, defined as co-occurring resistance to ampicillin, chloramphenicol, and co-trimoxazole; XDR, extensive drug resistance, defined as MDR plus resistance to ciprofloxacin and ceftriaxone.
genotype 3.3 and parE A364V (hydrophobicity, 0.0 → $-0.78$; Fig. S5) (37). These mutations could have potential as markers to trace genotypes, especially the more prevalent genotypes such as 4.3.1 and 3.3.

**New H58 lineages with high-level ciprofloxacin resistance.** According to the published scheme that defines the different lineages of genotype 4.3.1 (H58) (26, 38), lineage la was dominant among the isolates from the recent Pakistan XDR outbreak, while most isolates from the Nepal surveillance belong to lineage II (Table 7). An undefined cluster within lineage II was also noticed among the Nepali isolates (Fig. 3), as has been described previously (12, 29). Among our isolates from Bangladesh, we found a new lineage of genotype 4.3.1 (H58), Bd ($n =$ 108), which represented the second most dominant lineage after la ($n =$ 223). This new lineage has decreased susceptibility to ciprofloxacin compared to lineage la (mean MIC, 1.71 versus 0.74 µg/ml). Ciprofloxacin MICs of $>0.06$ µg/ml are classified as resistant following the EUCAST guidelines. However, as the resistance breakpoint specified by the Clinical and Laboratory Standards Institute (CLSI) is 1 µg/ml, some strains could be classified as susceptible in countries that use the CLSI guidelines (12, 17, 29). Moreover, lineage Bd is probably of local origin, as it was absent in both neighboring countries (Fig. 3) and does not match the published SNP definition of lineage I or II (38). This local variant also had a higher pairwise distance in the SNP matrix (mean, 12.8 versus 11.2) than lineage la, suggesting a different pattern of divergence.

Remarkably, a sublineage of lineage Bd (Bdq; $n =$ 55) showed increased resistance compared to other isolates from the same lineage, with median ciprofloxacin MICs of 4.0 µg/ml (mean MIC, 3.4 versus 0.4 µg/ml; Fig. S4). Sublineage Bdq predominantly carried qnr genes, in addition to gyrA mutations (Fig. 3 and 5), and showed more clonality than other lineages (mean pairwise distance, 8.4 versus 11.2 for la). Moreover, sublineage Bdq emerged recently, as all isolates were from 2006 onward, but became more prevalent after 2007 (Fig. S3). Therefore, antimicrobial treatment with fluoroquinolones of infections caused by sublineage Bdq may lead to failure.

A similar highly resistant lineage with triple mutations ($gyrA_{S83F}$ $gyrA_{D87G}$ $parC_{E84G}$) but with no qnr genes was previously reported to cause failure of treatment with gatifloxacin in Nepal (28, 29). Our MLT also showed a small subclade ($n =$ 8) inside lineage la for Bangladesh, with a triple mutation ($gyrA_{S83F}$ $gyrA_{D87G}$ $parC_{E84K}$) and median ciprofloxacin MICs of 8.0 µg/ml (Fig. 4 and 5).

Notably, the number of lineage II isolates ($n =$ 4) in Bangladesh was extremely low (Fig. 3), despite the dominance of this lineage in Nepal and India (26, 27, 29). The surveillance data from Nepal, which mostly describes the isolates from Kathmandu valley, showed a shifting pattern of H58 lineages (from lineage I to lineage II) over the years (29). Such a changing pattern is not observed in Bangladesh, probably because of relatively high prevalence and dominance of local lineages, such as the previously unreported lineage Bd. The Nepal surveillance also reported association of MDR with lineage I and of cip resistance with lineage II (29). However, no such association has been found for lineage I or lineage Bd in Bangladesh (see Data Set S1 in the supplemental material).

**wgSNP analysis suggests regional clonality of S. Typhi in Bangladesh.** The wgSNP analyses of our isolates generated 2,328 SNPs, revealing that the S. Typhi population in Bangladesh is highly clonal. However, addition of the isolates from Nepal ($n =$ 198) and Pakistan ($n =$ 100) increased the number of SNPs to 3,251 but decreased the number to 627 for genotype 4.3.1 isolates only ($n =$ 603). As the filtering criteria remain the same, the number of SNPs for all Bangladesh isolates is relatively low compared to the global or multicountry context (25–27) but is similar to country-specific data. For example, 1,850 SNPs were detected in isolates from Thailand ($n =$ 44) and 2,187 SNPs in isolates from Nepal ($n =$ 198) (29, 39). The wgSNP-MLT data showed distinct differentiation of all genotypes, much like the data from the cgMLST-UPGMA tree, except the latter lacked clear inferences for different H58 lineages (Fig. 1b and c and Fig. S5). Genotype 1.2.1 mapped close to the root of the MLT, suggesting that this
genotype is one of the oldest circulating types. Likewise, being the most distantly related, genotype 4.3.1 could be one of the more recent genotypes circulating in Bangladesh (Fig. 1c) and neighboring countries (Fig. 2).

**WGS predicts AMR phenotypes with high sensitivity.** The WGS-based resistance profiles showed >99% sensitivity and >91% specificity in describing the phenotypes (for amp, sxt, chl, and cro) of AMR isolates (Table 5). Remarkably, the dfrA7 genes (involved in trimethoprim resistance) were always detected in the presence of the sul1 gene (sulfonamide resistance) and never alone. Table 5). Similarly, sul1 was never detected in the absence of dfrA7. Two isolates had discordant results, as we did not detect the concordant resistance genes in WGS analyses (Table 5). Repeating the antimicrobial susceptibility tests (ASTs) reconfirmed the resistant phenotype. Other resistance mechanisms, e.g., efflux pumps or membrane permeability changes, may be involved (40).

Ciprofloxacin resistance in 11 isolates with no mutation in DNA gyrase or topoisomerase IV genes (and no qnr genes) can suggest the presence of other mechanisms. Indeed, MDR bacteria can increase the expression of efflux pump genes, including acrAB, acrEF and tolC (through overexpression of ramA or repression of acrR genes). This enables the bacteria to expel fluoroquinolone molecules, resulting in ciprofloxacin resistance (40–43), as well as ampicillin or chloramphenicol resistance, even in the absence of bla or catA genes (44–46). On the other hand, isolates carrying a bla gene without the resistance phenotype could be the result of mutations in the promoter regions of outer membrane protein genes, such as the ompC gene, which facilitates penetration of beta-lactams through the outer membrane (47, 48). This could be the scenario for several susceptible isolates (n = 30) in our library that have the full-length resistance gene. However, transcriptomic or proteomic approaches may be required to further explore these possibilities.

**Different genotypic backgrounds of ceftriaxone resistance in Bangladesh and Pakistan.** The ceftriaxone-resistant (cro-R) strain from our library was isolated in 2000. The first report of a cro-R strain was published in 1999 (16). Interestingly, this isolate harbored the same extended-spectrum-beta-lactamase (ESBL) gene, blaCTX-M-15 (17, 49, 50), that caused the ceftriaxone-resistant phenotype in an ongoing typhoid outbreak in Pakistan (12). Other ESBL genes, including blaCMY-2 and blaCTX-M-14, have also been reported in relation with ceftriaxone resistance in other *Salmonella* species (51, 52) but never in *S. Typhi*. The sequence identity of blaCTX-M-15 between our isolate and the Pakistani isolates was 99%, with 92% coverage (Table 7). The resistance phenotype and genotype were also different from those of our isolate (Table 7). The Pakistani outbreak isolates formed a distinct cluster in the H58-specific MLT and showed high-level clonality (Fig. 3 and Fig. S1). In contrast, our ceftriaxone-resistant isolate had genotype 3.3, which suggests a different source and geographical origin. Moreover, no other ceftriaxone-resistant strains of genotype 3.3 have been reported from Bangladesh. We hypothesize that acquisition of the blaCTX-M15 gene might compromise the fitness of *S. Typhi* although as of now no data have been published in support of this. Also, no association with fitness has been found for ciprofloxacin resistance mutations in DNA gyrase genes (53). Therefore, the possibility of a global dissemination of these recently emerging variants cannot be excluded given the successful multicontinent spreading of its H58 ancestor (genotype 4.3.1).

This study had some limitations. The isolates from Pakistan are from a still-ongoing outbreak in Hyderabad and Karachi that started in 2016. The Nepal isolates are from a prospective surveillance in the area of Kathmandu valley and cover a period of 9 years (2008 to 2016). The collection of strains from Bangladesh was selected from a biobank of >3,000 strains recovered over a period of 15 years (1999 to 2013) from two different hospital settings in Dhaka. The majority (97%) of the isolates from Bangladesh are from children (<18 years old). Therefore, none of the collections cover the whole population in their respective countries. Also, there is no overlap of the isolate collection periods
between Bangladesh and Pakistan. Country-to-country comparisons of the observed data may therefore be biased.

**Conclusion.** Our study demonstrated that WGS has high sensitivity and specificity for prediction of S. Typhi resistance phenotypes. However, this genomic method still lacks sensitivity and needs fine-tuning for the detection of ciprofloxacin resistance. We detected three different mutations associated with specific genotypes that could be used to develop genotype-specific tracking tools. We report a new, local variant of genotype 4.3.1, lineage Bd, which contains a recently emerged sublineage, Bdq, that exhibits a high level of ciprofloxacin resistance. A triple mutant variant (gyrA\textsubscript{SS3F} gyrA\textsubscript{DB7G} parC\textsubscript{EB4C}) of lineage la with high ciprofloxacin resistance was also detected. A similar triple mutant variant of lineage II (gyrA\textsubscript{SS3F} gyrA\textsubscript{DB7G} parC\textsubscript{EB4C}) has been reported from Nepal and possesses the same phenotype (28, 29). Our ceftriaxone-resistant isolate contains the \textit{bla}\textsubscript{CTX-M-15} gene but has a genotype and gene sequence different from those of the same gene of XDR S. Typhi strains from the Pakistan outbreak, defining a different ancestral origin. Thus, dissemination of this isolate throughout the region from a single point is therefore less likely. However, multiple independent genetic events in neighboring countries and possible subsequent dissemination enhance the risk of the global spread of these highly resistant clones.

The data presented in this study will add to the accumulating information, from Pakistan and Nepal in particular, concerning the increasing drug resistance of S. Typhi. The emergence of XDR S. Typhi is strongly compromising effective treatment of typhoid fever. The spread of these resistant lineages and their occurrence in various Asian countries emphasize the need to inform public health professionals and sensitize the global community. Measures to implement a two-pronged approach for typhoid control need to be accelerated (54, 55). Both short-term vaccine interventions for high-risk populations and long-term water and sanitation interventions will undoubtedly be the cornerstones of a global prevention plan to address control of typhoid fever.

**MATERIALS AND METHODS**

**Isolate collection and antimicrobial susceptibility profiles.** All S. Typhi isolates used in this study were collected from the Child Health Research Foundation (CHRF) at the Department of Microbiology, Dhaka Shishu (Children’s) Hospital, in Dhaka, Bangladesh. The CHRF team has been preserving invasive \textit{Salmonella} isolates since 1999 and maintained a biobank of >3,500 S. Typhi isolates, largely from children (<18 years of age). All strains were isolated from the blood of patients diagnosed with typhoid fever in two different settings: hospital inpatients (hospitalized), and out-patients attending the consultation facility (56). Clinical and epidemiological data were collected for all isolates collected from hospital inpatients.

We selected 539 S. Typhi isolates for this study; data were available for those isolates with respect to the date of isolation (1999 to 2013), hospital setting, and phenotypic resistance for five different antibiotics (ampicillin, chloramphenicol, co-trimoxazole, ciprofloxacin, and ceftriaxone). Age data were available for 85% (456/536) cases; among those cases, 97% (443/456) patients were <18 years of age, while 76% (345/456) were <5 years of age. We checked the identity of the isolates by the use of standard biochemical tests and \textit{Salmonella} agglutinating antisera (Thermo Scientific, MA, USA). Antimicrobial susceptibility for ampicillin (amp), co-trimoxazole (sxt), and chloramphenicol (chl) was determined using the disk diffusion method (Oxoid, Thermo Scientific, MA, USA). Broth microdilution was used to determine the MIC values for ciprofloxacin (cip) and ceftriaxone (cro; Sigma-Aldrich, MO, USA). All zone diameter and MIC data were interpreted according to EUCAST v8.0 clinical breakpoints (57). Fig. S7 in the supplemental material shows the complete workflow. All sequence data have been submitted to the European Nucleotide Archive (ENA). Data Set S1 in the supplemental material summarizes relevant details of our isolates.

**DNA extraction and whole-genome sequencing.** Isolates were grown on MacConkey agar (Oxoid) overnight, and the colonies were suspended in water. The QIAamp DNA minikit (Qiagen, Hilden, Germany) was used to extract DNA from the suspension on the same day. WGS was performed using an Illumina HiSeq 4000 platform (The Oxford Genomics Centre at the Wellcome Trust Centre for Human Genetics, Oxford, United Kingdom). One \textit{Salmonella} Paratyphi isolate was also sequenced so that it could be included in comparative phylogenetic analysis (as an outgroup).

**Data quality check.** Sequence data quality was checked using FastQC v0.11.15 (58). We summarized all quality indicators using MultiQC v3 (59). If the summary revealed the presence of adapter sequences, they were removed using Trimmomatic v0.36 (60). KmerFinder was used to confirm the species of the strains (61, 62). Another tool, SeqSero, was used for WGS-based serotyping, to determine the \textit{Salmonella} serovar of the isolates and confirm the wet-lab serotyping results (63).
WGS data analyses with BioNumerics. Adaptor-free fastq files were imported into BioNumerics version 7.6.2 (Applied Maths NV, Sint-Martens-Latem, Belgium) and analyzed via the use of the integrated Calculation Engine. For the comparison with isolates from neighboring countries, we used recently published WGS data on 100 S. Typhi isolates from Pakistan (12) and 198 S. Typhi isolates from Nepal (29). The Pakistan isolates were mostly from an ongoing outbreak of XDR S. Typhi, in Hyderabad and Karachi, Sindh, Pakistan, between November 2016 and March 2017 (12). In contrast, the Nepal isolates were part of a hospital-based enteric fever surveillance performed during 2008 to 2016, based on one of the large referral hospitals in Kathmandu Valley, namely, Patan Academy of Health Sciences (PAHS). (29).

Details of the quality control of the WGS data, mapping against the reference genome, filtering the SNPs, allele calling for cgMLST, detecting the presence of acquired resistance genes, and SNP-based genotyping are described in Text S1.

Classical 7-locus MLST. The complete sequences of seven loci (aroC, dnaN, hemD, hisD, purE, sucA, and thrA) were identified in extracted contigs. All sequences were matched with Enterobase (Achtman 7-gene MLST) (http://enterobase.warwick.ac.uk/species/index/senterica) to determine the classical MLST type of each isolate.

Phylogenetic analyses. We used RaxML v8.2.10 to build maximum likelihood phylogenetic trees (MLT) (65) on the basis of the alignment of 2,328 SNPs from 536 S. Typhi isolates in our study, 3,251 SNPs from 834 isolates in the comparisons with neighboring countries, and 627 SNPs from all 603 H58 isolates. Lineages for all H58 isolates were determined as previously described (38). We employed the generalized time-reversible model and a Gamma distribution to model site-specific rate variation (the GTRGAMMA in RaxML). Support for the MLT phylogeny was assessed via 100 bootstrap pseudoanalyses. The S. Paratyphi A strain from Bangladesh (Sample: 311189, 229186) was included as an outgroup for tree rooting. All MLT and UPGMA trees were displayed and annotated using the iTOL6 online version (66). To compute the genetic distances between different groups (e.g., countries, H58 lineages, etc.), a pairwise SNP distance matrix was generated between isolates by computing the number of SNP loci at which pairs of isolates had discordant alleles. Median distances within or between groups were computed from this distance matrix.

Statistical analyses were performed using R v3.5 (64); the same application was used to generate the line graphs and box plots.

Data availability. All sequence data determined in work have been submitted to the European Nucleotide Archive (ENA) (study identifier [ID]: ERP109468).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.02112-18.

FIG S1, TIF file, 3.3 MB.
FIG S2, TIF file, 4.1 MB.
FIG S3, TIF file, 3.1 MB.
FIG S4, TIF file, 2.6 MB.
FIG S5, TIF file, 3.9 MB.
FIG S6, TIF file, 4.1 MB.
FIG S7, TIF file, 2 MB.
TABLE S1, DOCX file, 0.01 MB.
DATA SET S1, XLSX file, 0.1 MB.
DATA SET S2, XLSX file, 0.1 MB.
TEXT S1, DOCX file, 0.01 MB.

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We declare no conflicts of interest.
REFERENCES

1. Crump JA, Luby SP, Mintz ED. 2004. The global burden of typhoid fever. Bull World Health Organ 82:346–353.

2. Crump JA, Mintz ED. 2010. Global trends in typhoid and paratyphoid fever. Clin Infect Dis 50:241–246. https://doi.org/10.1086/649541.

3. Parry CM, Hien TT, Dougan G, White NJ, Farrar JJ. 2002. Typhoid fever. N Engl J Med 347:1770–1782. https://doi.org/10.1056/NEJMra022001.

4. Franco-Paredes C, Khan MI, Gonzalez-Diaz E, Santos-Preciado JJ, Rodriguez-Morales AJ, Gotuzzo E. 2016. Enteric fever: a slow response to an old plague. PLoS Negl Trop Dis 10:e0004597. https://doi.org/10.1371/journal.pntd.0004597.

5. Kirk MD, Pies SM, Black RE, Caipo M, Crump JA, Devellecchauber B, Döpfer D, Fazil A, Fischer-Walker CL, Hald T, Hall AJ, Keddy KH, Lake RJ, Lanata CF, Torgerson PR, Havelaar AH, Angulo FJ. 2015. World Health Organization estimates of the global and regional disease burden of 22 foodborne bacterial, protozoal, and viral diseases, 2010: a data synthesis. PLoS Med 12:e1001921. https://doi.org/10.1371/journal.pmed.1001921.

6. Das S, Samajpati S, Ray U, Roy I, Dutta S. 2017. Antimicrobial resistance and molecular subtype of Salmonella enterica serovar Typhi isolates from Kolkata, India over a 15 years period 1998–2012. Int J Med Microbiol 307:28–36. https://doi.org/10.1016/j.ijmm.2016.11.006.

7. Zellweger RM, Basnayat B, Shrestha P, Prapajati KG, Dongol S, Sharma PK, Koirala S, Darton TC, Dolecek C, Thompson CN, Thwaites GE, Baker SG, Karkey A. 2017. A 23-year retrospective investigation of Salmonella Typhi and Salmonella Paratyphi isolated in a tertiary Kathmandu hospital. PLoS Negl Trop Dis 11:e0005601. https://doi.org/10.1371/journal.pntd.0005601.

8. Saha S, Saha S, Ruhulamin M, Hanif M, Islam M. 1997. Decreasing trend of multiresistant Salmonella typhi in Bangladesh. J Antimicrob Chemother 39:554–556. https://doi.org/10.1093/jac/39.4.554.

9. Olarte J, Galindo E. 1973. Salmonella typhi resistant to chloramphenicol, ampicillin, and other antimicrobial agents: strains isolated during an extensive typhoid fever epidemic in Mexico. Antimicrob Agents Chemother 1:404–408. https://doi.org/10.1128/AAC.1.3.404-408.73a.

10. Smith SM, Palumbo PE, Edelson PJ. 1984. Salmonella strain resistant to multiple antibiotics: therapeutic implications. Pediatr Infect Dis J 3:455–460. https://doi.org/10.1097/00006454-198409000-00005.

11. Iyer RN, Jiangam RR, Jachint R, Venkatakalakshmi A, Nahdi FB. 2017. Prevalence and trends in the antimicrobial susceptibility pattern of Salmonella enterica serovars Typhi and Paratyphi A among children in a pediatric tertiary care hospital in South India over a period of ten years: a retrospective study. Eur J Clin Microbiol Infect Dis 36:2399–2404. https://doi.org/10.1007/s10095-016-2703-x.

12. Klemm EJ, Shakoor S, Page AJ, Qamar FN, Judge K, Saeed DK, Wong VK, Day MR, Doumith M, Do Nascimento V, Nair S, Ashton PM, Jenkins C, de Pinna EM, Godbole G. 2018. Comparison of phenotypic and WGS-derived antimicrobial resistance profiles of Salmonella enterica serovars Typhi and Paratyphi. J Antimicrob Chemother 73:365–372. https://doi.org/10.1093/jac/dkx379.

13. Anes J, Hurley D, Martins M, Fanning S. 2017. Exploring the genome and phenotype of multi-drug resistant Klebsiella pneumoniae of clinical origin. Front Microbiol 8:9193. https://doi.org/10.3389/fmicb.2017.01913.

14. Tasmin R, Hasan NA, Grim CJ, Grant AQquette, Choi SY, Alam MS, Bell R, Cavanaugh C, Balan KV, Parveen S. 2017. Genotypic and phenotypic characterization of multidrug resistant Salmonella Typhi and Salmonella enterica serovars Typhi and Paratyphi. J Antimicrob Chemother 73:365–372. https://doi.org/10.1093/jac/dkx379.

15. Anes J, Hurley D, Martins M, Fanning S. 2017. Exploring the genome and phenotype of multi-drug resistant Klebsiella pneumoniae of clinical origin. Front Microbiol 8:9193. https://doi.org/10.3389/fmicb.2017.01913.

16. Saha SK, Talukder SY, Islam M, Saha S. 1999. A highly ceftriaxone-resistant drug-resistant Salmonella enterica serovar Typhi clone harboring a plasmid encoding resistance to fluoroquinolones and third-generation cephalosporins. mBio 10:e00105-18. https://doi.org/10.1128/mBio.00105-18.

17. Guil D, Potter RF, Riaz H, Ashraf SA, Wallace MA, Munir T, All A, Burnham CA, Dantas G, Andleeb S. 2017. Draft genome sequence of a Salmonella enterica serovar Typhi strain resistant to fourth-generation cephalosporin and fluoroquinolone antibiotics. Genome Announcements 5:e00850-17. https://doi.org/10.1128/genomeA.00850-17.

18. Paris Y, Thiel S, Hildreth NA, Muller J, Dougan G, Hitzke K. 2016. Ceftriaxone resistance pattern in the recent isolates of Salmonella Typhi with special reference to cephalosporins and azithromycin in the Gangetic plain. J Clin Diagn Res 11:DM01. https://doi.org/10.7860/JCDR/2017/23330.9973.

19. Godbole GS, Day MR, Murthy S, Chattaway MA, Nair S. 2018. First report of CTX-M-15 Salmonella Typhi From England. Clin Infect Dis 66:1976–1977. https://doi.org/10.1093/cid/ciy302.

20. Saha SK, Talukder SY, Islam M, Saha S. 1999. A highly ceftriaxone-resistant Salmonella typhi in Bangladesh. Pediatr Infect Dis J 18:387. https://doi.org/10.1097/00006454-199904000-00018.

21. Djeoghout B, Saha S, Sajib MS, Tanmoy AM, Islam M, Kay GL, Langridge GC, Endtz HP, Wain J, Saha SK. 2018. Ceftriaxone-resistant Salmonella Typhi carries an IncI1-ST31 plasmid encoding CTX-M-15. J Med Microbiol 67:620–627. https://doi.org/10.1099/jmm.0.007027.

22. Choudhary A, Gopalakrishnan R, Senthur NP, Radhakrishnam O, Gha- fur KA, Thirunarayam A. 2013. Antimicrobial susceptibility of Salmonella enterica serovars in a tertiary care hospital in southern India. Indian J Medical Res 137:800–802.
Whitehead S, Barrell BG. 2001. Complete genome sequence of a multiple drug resistant Salmonella enterica serovar Typh CT18. Nature 413: 848–852. https://doi.org/10.1038/3510607.

31. Chiu C-S, Lauderdale T-L, Phung DC, Watanabe H, Kuo J-C, Wang P-J, Liu Y-Y, Liang S-Y, Chen P-C. 2014. Antimicrobial resistance in Salmonella enterica serovar Typhi isolates from Bangladesh, Indonesia, Taiwan, and Vietnam. Antimicrob Agents Chemother 58:6501–6507. https://doi.org/10.1128/AAC.03608-14.

32. Yap K-P, Ho WS, Goz-HM, Chai LC, Thong KL. 2016. Global MLST of Salmonella Typhi revisited in post-genomic era: genetic conservation, population structure, and comparative genomics of rare sequence types. Front Microbiol 7:270. https://doi.org/10.3389/fmicb.2016.00270.

33. Enterobase. 2018. University of Warwick, Coventry, United Kingdom. http://enterobase.warwick.ac.uk/species/senterica/search_strains?query_str_search. Accessed 2 August 2018.

34. Hendriksen RS, Leekitcharoenphon P, Lukjancenko O, Lukwesa-Musyani AM, Goechea JA, Albertí S. 2010. Role of porin proteins in the reversal of efflux pump activity, a rational strategy to combat gram-negative bacteria. Front Microbiol 7:270. https://doi.org/10.3389/fmicb.2016.00270.

35. Xia X, Li W-H. 1998. What amino acid properties affect protein evolution? J Mol Evol 47:557–564. https://doi.org/10.1007/PL00006412.

36. Yap K-P, Ho WS, Goz-HM, Chai LC, Chong KL. 2016. Global MLST of Salmonella enterica serovar Typhi isolates related to a massive outbreak in Zambia between 2010 and 2012. J Clin Microbiol 53:262–272. https://doi.org/10.1128/JCM.00206-14.

37. Moon CP, Fleming KG. 2011. Side-chain hydrophobicity scale derived from transmembrane protein folding into lipid bilayers. Proc Natl Acad Sci U S A 108:10174–10177. https://doi.org/10.1073/pnas.1013979108.

38. Sharma V, Dahiya S, Jangra P, Kumar R, Sood S, Kapil A. 2013. Study of high-throughput genome sequencing data. J Clin Microbiol 52:139.

39. Hasman H, Saputra D, Sicheritz-Ponten T, Lund O, Svendsen CA, Frimodt-Møller N, Fields PI, Deng X. 2015. How high-throughput bacterial SNP typing identifies distinct clusters of Salmonella enterica serovar Typhimurium to fluoroquinolones and other antimicrobial. Antimicrob Agents Chemother 51:535–542.

40. Holt KE, Baker S, Dongol S, Basnyat B, Adhikari N, Thorson S, Pulickal AS, Whitehead S, Barrell BG. 2001. Complete genome sequence of a multiple drug-resistant Salmonella enterica serovar Typhi isolate related to a massive outbreak in Northern Taiwan attributable to production of CTX-M-4 and CMY-2 β-lactamases. J Clin Microbiol 43:3237–3243. https://doi.org/10.1128/JCM.43.13.3237-3243.2005.

41. Baker S, Duy PT, Nga TTV, Phan VT, Chau TT, Turner AK, Farrar J, Boni MF. 2013. Fitness benefits in fluoroquinolone-resistant Salmonella Typhi in the absence of antimicrobial pressure. Elife 2:e01229. https://doi.org/10.7554/eLife.01229.

42. Saha SK. 2018. Commentary: the new typhoid conjugate vaccine marks the dawn of a unique beginning. https://www.jhsph.edu/ivac/2018/03/27/the-new-typhoid-conjugate-vaccine-marks-the-dawn-of-a-unique-beginning/.

43. Levine MM, Simon R. 2018. The gathering storm: is untreated typhoid fever on the way? mBio 9:e00482-18. https://doi.org/10.1128/mBio.00482-18.

44. Saha SK, Baqui AH, Hanif M, Darmstadt GL, Nagatake T, Santoshm M, Black RE. 2001. Typhoid fever in Bangladesh: implications for vaccination policy. Pediatr Infect Dis J 20:521–524. https://doi.org/10.1097/00006454-200105000-00010.

45. European Committee on Antimicrobial Susceptibility Testing. 2018. Breakpoint tables for interpretation of MICs and zone diameters v8.0:5-9. http://www.eucast.org/fileadmin/src/media/PDFs/EUCAST_files/Breakpoint_tables/V8_0_Breakpoint_Tables.pdf.

46. Andrews S. 2010. FastQC: a quality control tool for high throughput sequence data. https://www.bioinformatics.babraham.ac.uk/projects/fastqc/.

47. Ewels P, Magnusson M, Lundin S, Käller M. 2016. MultiQC: summarize fastqc results for multiple tools and samples in a single report. Bioinformatics 32:3047–3048. https://doi.org/10.1093/bioinformatics/btw354.

48. Bolger AM, Lohse M, Usadel B. 2014. Trimomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 30:2114–2120. https://doi.org/10.1093/bioinformatics/btu170.

49. Larsen MV, CoSENTino S, Lukjancenko O, Saputra D, Rasmussen S, Hasman H, Sicheritz-Ponten T, Aarestrup FM, Ussery DW, Lund O. 2014. Benchmarking of methods for genomic taxonomy. J Clin Microbiol 52:1529–1539. https://doi.org/10.1128/JCM.02981-13.

50. Hasman H, Saputra D, Sicheritz-Ponten T, Lund O, Svendsen CA, Frimodt-Møller N, Aarestrup FM. 2014. Rapid whole genome sequencing for the detection and characterization of microorganisms directly from clinical samples. J Clin Microbiol 52:1529–1539. https://doi.org/10.1128/JCM.02981-13.

51. Letunic I, Bork P. 2016. Interactive tree of life (iTOL) v3: an online tool for displaying and annotation of phylogenetic and other trees. Nucleic Acids Res 44:W242–W245. https://doi.org/10.1093/nar/gkw290.