Cluster-specific gene marker enhance *Shigella* and Enteroinvasive *Escherichia coli* in silico serotyping

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Abstract

*Shigella* and enteroinvasive *Escherichia coli* (EIEC) cause human bacillary dysentery with similar invasion mechanisms and share similar physiological, biochemical and genetic characteristics. The ability to differentiate *Shigella* and EIEC from each other is important for clinical diagnostic and epidemiologic investigations. The existing genetic signatures may not discriminate between *Shigella* and EIEC. Phylogenetically, *Shigella* and EIEC strains composed of multiple clusters and are different forms of *E. coli*. In this study, we identified 10 *Shigella* clusters, 7 EIEC clusters and 53 sporadic types of EIEC by examining over 17,000 publicly available *Shigella*/EIEC genomes. We compared *Shigella* and EIEC accessory genomes to identify the cluster-specific gene markers or marker sets for the 17 clusters and 53 sporadic types. The gene markers showed 99.63% accuracy and more than 97.02% specificity. In addition, we developed a freely available *in silico* serotyping pipeline named *Shigella* EIEC Cluster Enhanced Serotype Finder (ShigEiFinder) by incorporating the cluster-specific gene markers and established *Shigella*/EIEC serotype specific O antigen genes and modification genes into typing. ShigEiFinder can process either paired end Illumina sequencing reads or assembled genomes. ShigEiFinder provided nearly perfect differentiation of *Shigella* from EIEC with 99.70% and 99.81% accuracy to assign isolates to the correct clusters for the assembled genomes and reads mapping respectively. ShigEiFinder was able to serotype over 59 *Shigella* serotypes and 22 EIEC serotypes and provided a high specificity with 99.40% for assembled genomes and 99.38% for reads mapping for serotyping. The cluster markers and our new serotyping tool, ShigEiFinder, will be useful for epidemiologic and diagnostic investigations.

Data summary

Sequencing data have been deposited at the National Center for Biotechnology Information under BioProject number PRJNA692536.
Introduction

*Shigella* is one of the most common etiologic agents of foodborne infections worldwide and can cause diarrhea with a very low infectious dose (1, 2). The infections can vary from mild diarrhea to severe bloody diarrhea referred to as bacillary dysentery. The estimated cases of *Shigella* infections are 190 million with at least 210,000 deaths annually, predominantly in children younger than 5 years old in developing countries (3-7). *Shigella* infections also have a significantly impact on public health in developed countries although most cases are travel-associated (8).

The *Shigella* genus consists of four species, *Shigella sonnei*, *Shigella flexneri*, *Shigella boydii* and *Shigella dysenteriae* (9). Serological testing further classifies *Shigella* species into more than 55 serotypes through the agglutination reaction of antisera to *Shigella* serotype specific O-antigens (10, 11). Up to 89.6% *Shigella* infections were caused by *S. flexneri* (65.9%) and *S. sonnei* (23.7%) globally (12, 13). The predominant serotype reported in *Shigella* infections has been *S. flexneri* serotype 2a while *S. dysenteriae* serotype 1 has caused the most severe disease (11, 14). Note that for brevity, in all references to *Shigella* serotypes below, *S. sonnei*, *S. flexneri*, *S. boydii* and *S. dysenteriae* are abbreviated as SS, SF, SB and SD respectively and a serotype is designated with abbreviated “species” name plus the serotype number e.g. *S. dysenteriae* serotype 1 is abbreviated as SD1.

Enteroinvasive *Escherichia coli* (EIEC) is a pathovar of *E. coli* that causes diarrhoea with less severe symptoms to *Shigella* infections in humans worldwide, particularly in developing countries (8, 13, 15-18). EIEC infections in developed countries are mainly imported (19). EIEC has more than 18 specific *E. coli* O-serotypes (19, 20). Although the incidence of EIEC is low (17), EIEC serotypes have been associated with outbreaks and sporadic cases of infections (20-22). In contrast to *Shigella*, EIEC infections are not notifiable in many countries (23, 24).

*Shigella* and EIEC have always been considered very closely related and share several characteristics (25-28). *Shigella* and EIEC are both non-motile and lack the ability of ferment lactose (24). Some of EIEC O antigens are identical or similar to *Shigella* O antigens (O112ac, O124, O136, O143, O152 and O164) (26, 29-31). Furthermore, *Shigella* and EIEC both carry the virulence plasmid pINV, which encodes virulence genes required for invasion (32, 33) and contains *ipaH* (invasion plasmid antigen H) genes with the exception of some SB13 isolates.
(10, 23, 24, 34, 35). Shigella and EIEC have arisen from *E. coli* in multiple independent events and should be regarded as a single pathovar of *E. coli* (25, 26, 28, 36-38). Previous phylogenetic studies suggested that *Shigella* isolates were divided into 3 clusters (C1, C2 and C3) with 5 outliers (SS, SB13, SD1, SD8 and SD10) (25, 38) whereas EIEC isolates were grouped into four clusters (C4, C5, C6 and C7) (26). The seven *Shigella/EIEC* clusters and 5 outliers of *Shigella* are within the broader non-enteroinvasive *E. coli* species except for SB13 which is closer to *Escherichia albertii* (39, 40). Alternative WGS-based phylogenomic studies have also defined multiple clusters of *Shigella* and EIEC (23, 28, 41).

The traditional biochemical test for motility and lysine decarboxylase (LDC) activity (42) and molecular test for the presence of *ipaH* gene have been used to differentiate *Shigella* and EIEC from non-enteroinvasive *E. coli* (24, 43-45). Agglutination with *Shigella/EIEC* associated antiserum further classify *Shigella* or EIEC to serotype level. However, cross-reactivity, strains not producing O antigens, and newly emerged *Shigella* serotypes may all prevent accurate serotyping (10, 46). Serotyping by antigenic agglutination is being replaced by molecular serotyping (47, 48), which can be achieved through examination of the sequences of O antigen biosynthesis and modification genes (8, 24, 49-52).

Recently, PCR-based molecular detection methods targeting the gene *lacY* were developed to distinguish *Shigella* from EIEC (53, 54). However, the ability of the primers described in these methods to accurately differentiate between *Shigella* and EIEC was later questioned (23, 28). With the uptake of whole-genome sequencing technology, several studies have identified phylogenetic clade specific markers, species specific markers and EIEC lineage-specific genes for discrimination between *Shigella* and EIEC and between *Shigella* species (23, 27, 28, 41, 55, 56). More recently, genetic markers *lacY, cadA, Ss_methylase* were used for identification of *Shigella* and EIEC (10). However, these markers failed to discriminate between *Shigella* and EIEC when a larger genetic diversity is considered (23, 28, 55). A Kmer-based approach can identify *Shigella* isolates to the species level but misidentification was also observed (56).

In this study, we aimed to i), identify phylogenetical clusters of *Shigella* and EIEC through large scale examination of publicly available genomes; ii), identify cluster-specific gene markers using comparative genomic analysis of *Shigella* and EIEC accessory genomes for differentiation of *Shigella* and EIEC; iii), develop a pipeline for *Shigella* and EIEC in silico serotyping based on the cluster-specific gene markers combined with *Shigella* and EIEC
serotype-specific O antigen and H antigen genes. We demonstrate that these cluster-specific

gene markers enhance in silico serotyping using genomic data. We also developed an
automated pipeline for cluster typing and serotyping of Shigella/EIEC from WGS data.

Materials and Methods

Identification of Shigella/EIEC isolates from NCBI database

E. coli/Shigella isolates from the NCBI SRA (National Center for Biotechnology Information
Sequence Read Archive) as May of 2019 were queried. Raw reads were retrieved from ENA
(European Nucleotide Archive). The ipaH gene (GenBank accession number M32063.1) was
used to screen E. coli/Shigella reads using Salmon v0.13.0 (57). Taxonomic classification for
E. coli/Shigella was confirmed by Kraken v1.1.1 (58). Molecular serotype prediction of ipaH
negative Shigella isolates was performed by ShigaTyper v1.0.6 (10). Isolates that were ipaH
positive and isolates with designation of SB13 by ShigaTyper were selected as Shigella/EIEC
database.

The sequence types (STs) and ribosomal STs (rSTs) of ipaH gene negative E. coli (non-
enteroinvasive E. coli) isolates were examined. STs and rSTs for these isolates were obtained
from the E. coli/Shigella database in the Enterobase (59) as of May 2019. For STs and rSTs
with only one isolate, the isolates were selected. For STs and rSTs with more than one isolates,
one representative isolate for each ST and rST were randomly selected. In total, 12,743 ipaH
negative E. coli isolates representing 3,800 STs and 11,463 rSTs were selected as non-
enteroinvasive E. coli control database.

Genome sequencing

Whole-genome sequencing (WGS) of 31 EIEC strains used in a previous study (26) was
performed by Illumina NextSeq (Illumina, Scoresby, VIC, Australia). DNA libraries were
constructed using Nextera XT Sample preparation kit (Illumina Inc., San Diego, CA, USA) and
sequenced using the NextSeq sequencer (Illumina Inc.). FASTQ sequences of the strains
sequenced in this study were deposited in the NCBI under the BioProject (PRJNA692536).

Genome assembly and data processing

Raw reads were de novo assembled using SPades v3.14.0 assembler with default settings
[http://bioinf.spbau.ru/spades] (60). The metrics of assembled genomes were obtained with
QUAST v5.0.0 (61). Three standard deviations (SD) from the mean for contig number, largest
contig, total length, GC, N50 and genes were used as quality filter for assembled genomes.

The STs for isolates in Shigella/EIEC database was checked by using mlst
(https://github.com/tseemann/mlst) with the E. coli scheme from PubMLST (62). rSTs were
extracted from the E. coli/Shigella rMLST database in Enterobase (59) as of May 2019.
Serotype prediction for isolates in Shigella/EIEC was performed by ShigaTyper v1.0.6 (10).
Serotyping of E. coli O and H antigens were predicted by using SerotypeFinder v2.0.1 (63).

Selection of isolates for Shigella/EIEC identification dataset

The selection of isolates for the identification dataset was based on the representative isolates
for each ST, rST and serotype of Shigella and EIEC in the Shigella/EIEC database. For STs
and rSTs with only one isolate, the isolate was selected. For STs and rSTs with more than one
isolates, one representative isolate for each ST, rST was randomly selected. A representative
experimentally confirmed isolate of each serotype of Shigella and EIEC was also randomly
selected. 72 ECOR strains downloaded from Enterobase (59) and 18 E. albertii strains were
used as controls for the identification dataset. The details of the identification dataset are listed
in Table S1. The remaining isolates in Shigella/EIEC database were referred as validation
dataset (Table S2).

The identification dataset was used for identification of phylogenetic relationships of Shigella
and EIEC. The identification dataset was also used for identification of cluster-specific genes.
The validation dataset was used to evaluate the performance of cluster-specific gene markers
using the in-silico serotyping pipeline.

Phylogeny of Shigella and EIEC based on WGS

Three phylogenetic trees including identification tree, confirmation tree and validation tree
were constructed by Quicktree v1.3 (64) with default parameters to identify and confirm the
phylogenetic clustering of Shigella and EIEC isolates. The phylogenetic trees were visualised
by Grapetree’s interactive mode and ITOL v5 (65, 66).

The identification phylogenetic tree was generated based on isolates in the identification
dataset for identification of clusters of Shigella and EIEC isolates (Fig. 1). A subset of 485
isolates known to represent each identified cluster from the identification dataset were then
selected. The subset of 485 isolates from the identification dataset and 1,872 non-enteroinvasive E. coli isolates from non-enteroinvasive E. coli control dataset (2,357 isolates total) were used to construct a confirmation tree. This tree was used for confirmation of the phylogenetic relationships between identified Shigella/EIEC clusters in the identification dataset and non-enteroinvasive E. coli isolates. The validation tree was generated based on 1,159 representative isolates from the validation dataset that were selected in the same way as the identification dataset and a subset of 485 isolates from the identification dataset to assign validation dataset isolates to clusters.

**Investigation of Shigella virulence plasmid pINV**

The presence of Shigella virulence plasmid pINV in isolates were investigated by using BWA-MEM v0.7.17 (Burrows-Wheeler Aligner) (67) to align isolate raw reads onto the reference sequence of pINV (68) (NC_024996.1). Mapped reads were sorted and indexed using Samtools v1.9 (69). The individual gene coverage from mapping was obtained using Bedtools coverage v2.27.1 (70).

**Identification of the cluster-specific gene markers**

Cluster-specific gene markers were identified from Shigella/EIEC accessory genomes. The genomes from the identification dataset were annotated using PROKKA v1.13.3 (71). Pan- and core-genomes were analysed by roary v3.12.0 (72) using an 80% sequence identity threshold. The genes specific to each cluster were identified from the accessory genes with an in-house python script. In this study, the number of genomes from a given cluster containing all specific genes for that cluster was termed true positives (TP), the number of genomes from the same cluster lacking any of those same genes was termed false negatives (FN). The number of genomes from other clusters containing all of those same genes was termed false positives (FP).

The sensitivity (True positive rate, TPR) of each cluster-specific gene marker was defined as TP/(TP+FN). The specificity (True negative rate, TNR) was defined as TN/(TN+FP).

**Validation of the cluster-specific gene markers**

The ability of cluster-specific gene markers to assign Shigella/EIEC isolates was examined by using BLASTN to search against the validation dataset (Table S2) and non-enteroinvasive E. coli control database for the presence of any of the cluster-specific gene marker or a set of
cluster-specific gene markers. The BLASTN thresholds were defined as 80% sequence identity and 50% gene length coverage.

**Development an automated pipeline for molecular serotyping of Shigella/EIEC**

The pipeline was developed using paired end illumina genome sequencing reads or assembled genomes identify cluster-specific gene markers combined with Shigella/EIEC serotype specific O antigen genes (wzx and wzy) and modification genes (Fig. 2, Data S1). We used the same signature O and H sequences from ShigaTyper and SerotypeFinder (Data S2) (10, 63). These includes Shigella serotype-specific wzx/wzy genes and modification genes from ShigaTyper and E. coli O antigen and fliC (H antigen) genes from SerotypeFinder. ipaH gene and 38 virulence genes used in analysis of virulence of 59 sporadic EIEC isolates were also included in the typing reference sequences database. Seven House Keeping (HK) genes - recA, purA, mdh, icd, gyrB, fumC and adk downloaded from NCBI were used for contamination checking.

Raw reads were aligned to the typing reference sequences by using BWA-MEM v0.7.17 (67). The mapping length percentage and the mean mapping depth for all genes were calculated using Samtools coverage v1.10 (69). To determine whether the genes present or absent, 50% of mapping length for all cluster-specific genes, virulence genes and O antigen genes and 10% for ipaH gene were used as cutoff value. The ratio of mean mapping depth to the mean mapping depth of the 7 HK genes was used to determine a contamination threshold with ratios less than 1% for ipaH gene and less than 10% for other genes assigned as contamination. Reads coverage mapped to particular regions of genes were checked by using samtools mpileup v1.10.

Assembled genomes were BLASTN v2.9.0 (73) searched against the typing reference sequences with 80% sequence identity and 50% gene length coverage for all genes with exception of ipaH gene which was defined as 10% gene length coverage.

The pipeline was tested with the identification dataset and validated with the Shigella/EIEC validation dataset and non-enteroinvasive E. coli control database. The specificity defined as (1 - the number of non-enteroinvasive E. coli isolates being detected / the total number of non-enteroinvasive E. coli isolates) * 100.
Results

Screening sequenced genomes for *Shigella/EIEC* isolates

We first screened available *E. coli* and *Shigella* genomes based on the presence of *ipaH* gene. We examined 122,361 isolates with the species annotation of *E. coli* (104,256) or *Shigella* (18,105) with paired end illumina sequencing reads available in NCBI SRA database. Of 122,361 isolates, 17,989 isolates were positive to the *ipaH* gene including 455 out of 104,256 *E. coli* isolates and 17,434 out of 18,105 *Shigella* isolates. The 17,989 *ipaH* positive *E. coli* and *Shigella* genomes and 571 *ipaH* negative “*Shigella*” genomes were checked for taxonomic classification and genome assembly quality. 17,320 *ipaH* positive *E. coli* and *Shigella* genomes and 246 *ipaH* negative “*Shigella*” genomes passed quality filters. Among 246 *ipaH* negative “*Shigella*” genomes, 11 isolates belonged to SB13 by using ShigaTyper (10) while the remaining 235 isolates were classified with taxonomic identifier of *E. coli* by Kraken v1.1.1 (58) and were removed from analysis. A total of 17,331 genomes including 17,320 *ipaH* positives and 11 SB13 genomes were selected to form the *Shigella/EIEC* database, which contained 429 genomes with species identifier of *E. coli* and 16,902 genomes with species identifier of *Shigella*.

Isolates in *Shigella/EIEC* database were typed using MLST, ShigaTyper and serotypeFinder. MLST and rMLST divided the 17,331 *Shigella/EIEC* isolates into 252 STs (73 isolates untypeable by MLST ) and 1,128 rSTs (3,513 isolates untypeable by rMLST). Of 16,902 genomes with species identifier of *Shigella*, 8,313 isolates and 8,189 isolates were typed as *Shigella* and EIEC respectively by ShigaTyper while 400 isolates were untypeable. ShigaTyper typed the majority of the 8,313 isolates as SF (66.82%) including 25.43% SF2a isolates, followed by SS (19.69%), SB (7.22%) and SD (6.27%).

SerotypeFinder typed 293 of the 429 *E. coli* genomes into 71 *E. coli* O/H antigen types. Among these 293 isolates with typable O/H antigen types, 190 isolates belonged to 22 known EIEC serotypes (O28ac:H-, O28ac:H7, O29:H4, O112ac:H26, O121:H30, O124:H30, O124:H24, O124:H7, O132:H4, O132:H21, O135:H30, O136:H7, O143:H26, O144:H25, O152:H-, O152:H30, O164:H-, O164:H30, O167:H26, O173:H7 and 2 newly emerged EIEC serotypes O96:H19 and O8:H19) (20-22). The remaining 136 of 429 genomes were O antigen untypable and typed to 15 H antigen types only by SerotypeFinder, of which H16 was the predominant H antigen type.
Identification of Shigella and EIEC clusters

Shigella and EIEC are known to have been derived from E. coli independently. To identify previously defined clusters (25, 26) and any new clusters from the 17,331 Shigella/EIEC genomes, we selected representative genomes to perform phylogenetic analysis as it was impractical to construct a tree with all genomes. The selection was based on ST, rST and serotype of the 17,331 Shigella/EIEC genomes. One isolate was selected to represent each ST, rST and serotype for a total of 1,830 isolates. The selection included 252 STs, 1,128 rSTs, 59 Shigella serotypes (21 SB serotypes, 20 SF serotypes, 17 SD serotypes and SS), 22 EIEC known serotypes and 31 other or partial antigen types. A further 31 in-house sequenced EIEC isolates, 18 EIEC isolates used in a previous typing study (41), 72 ECOR strains and 18 E. albertii strains were also included to form the identification dataset of 1,969 isolates. Details are listed in Table S1. A phylogenetic tree was constructed based on the identification dataset to identify the clusters (Fig. 1).

All known clusters were identified (Fig. 1) including 3 Shigella clusters (C1, C2, C3) and 5 outliers (SD1, SD8, SD10, SB13 and SS) as defined by Pupo et al (25) and 4 EIEC clusters (C4, C5, C6 and C7) defined by Lan et al. (26). Each of these clusters was supported by a bootstrap value of 80% or greater (Fig. S1). 1,789 isolates of the 1,879 Shigella/EIEC isolates (1,830 isolates from the Shigella/EIEC database, 31 in-house sequenced EIEC isolates and 18 EIEC isolates from Hazen et al.) fell within these clusters.

Of the remaining 90 Shigella/EIEC unclustered isolates, 31 belonged to 5 Shigella/EIEC serotypes including 5 SB13 isolates, 8 SB12 isolates, 2 EIEC O135:H30 isolates, 12 EIEC serotype O96:H19 isolates and 4 EIEC O8:H19 isolates, while 59 isolates were sporadic EIEC isolates which are described in detail in the separate section below. The 5 SB13 isolates were grouped into one lineage within E. coli and close to known Shigella/EIEC clusters rather than the established SB13 cluster outside E. coli which was within the E. albertii lineage. The former was previously named as atypical SB13 while the latter was previously named as typical SB13 (39). The 8 SB12 isolates formed one single cluster close to SD1 and atypical SB13 clusters. Two EIEC O135:H30 isolates were grouped as a separate cluster close to C5. Twelve isolates belonging to EIEC serotype O96:H19 and 4 isolates typed as O8:H19 were clustered into two separate clusters, both of which were more closely related to SD8 than other Shigella/EIEC clusters. Therefore, atypical SB13 and SB12 were defined as new clusters of
Shigella while EIEC O96:H19, EIEC O8:H19 and EIEC O135:H30 were defined as C8, C9 and C10 respectively. In total there were 10 Shigella clusters and 7 EIEC clusters (Table 1).

### Analysis of the 59 sporadic EIEC isolates

To determine the phylogenetic relationships of the above defined clusters and the remaining 59 sporadic EIEC isolates within the larger non-enteroinvasive E. coli population a confirmation tree was generated using 485 isolates representing the known clusters and 1,872 representative non-Shigella/EIEC isolates (Fig. S2). The 59 sporadic EIEC isolates including 2 EIEC isolates M2330 (O152:H51) and M2339 (O124:H7) sequenced in this study and 57 isolates were interspersed among non-Shigella/EIEC isolates and did not form large clusters. Groups of these isolates that were not previously identified were named as sporadic EIEC lineage followed by their serotype. For example, M2339 (O124:H7) grouped together with one other EIEC isolate with the same O and H antigens O124:H7 and were named ‘sporadic EIEC lineage O124:H7’.

There were 53 sporadic EIEC lineages including 5 lineages with 2 or more isolates and 48 lineages with only one isolate. The STs, rSTs and antigen types of these 59 isolates were listed in the Table S1.

Some of the sporadic EIEC isolates fell into STs containing *ipaH* negative isolates. We therefore examined the presence of the pINV virulence plasmid in the sporadic EIEC isolates. We selected 38 genes that are essential for virulence including 35 genes (12 *mxi* genes, 9 *spa* genes, 5 *ipaA-J* genes, 6 *ipgA-F* genes as well as *acp, virB, icsB*) in the conserved entry region encoding the Mxi-Spa-Ipa type III secretion system and its effectors and 3 regulator genes (*virF, virA* and *icsA/virG*) (24, 33, 68) and determined the presence of pINV in the 59 sporadic EIEC isolates by mapping the sequence reads onto a pINV reference sequence (68). Reads from 18 non-Shigella/EIEC isolates that shared the same ST as one of 58 sporadic isolates were positive for these genes.

The number of essential virulence genes with mapped reads in the 59 sporadic EIEC isolates were analysed (Fig. S3). Those isolates containing more than 25 of the 38 essential virulence genes were defined as virulence plasmid positive. While isolates containing between 13 and 25 were defined as intermediate and less than 13 were defined as virulence plasmid negative.

The 2 newly sequenced sporadic EIEC isolates (M2330 and M2339) were positive for the virulence plasmid and of the other 57 sporadic EIEC isolates, 39 isolates were positive, 9
isolates were negative and 9 isolates were intermediate (Table S1). The results were compared with those non-Shigella/EIEC isolates belonging to the same ST. The virulence plasmid was absent in all non-Shigella/EIEC isolates while all sporadic EIEC isolates in these STs were either positive or intermediate. Therefore, this analysis confirmed the sporadic isolates belonged to EIEC and the STs contained both EIEC and non-EIEC isolates.

**Identification of cluster-specific gene markers**

In this study, cluster-specific gene markers were either a single gene present in all isolates of a cluster and absent in all other isolates or a set of genes (two or more) that as a combination were only found in one cluster. For the marker sets, a subset of cluster-specific gene markers for a given cluster could be found in other clusters but the entire set was only found in the target cluster.

Comparative genomic analysis on 1,969 accessory genomes from the identification dataset was used to identify cluster-specific gene markers or marker sets. Multiple candidate cluster-specific gene markers or marker sets of markers for each of 17 Shigella/EIEC clusters and 53 sporadic EIEC lineages were identified through screening the accessory genes from 1,969 genomes. These gene markers or marker sets were 100% sensitive to clusters but with varying specificity. The cluster-specific gene markers or marker sets of markers with the lowest FP rates were then selected from candidate cluster-specific gene markers by BLASTN searches against genomes in the identification dataset using 80% sequence identity and 50% gene length coverage threshold.

Five single cluster-specific gene markers (C7, C10, SB12, SB13 and atypical SB13) and 12 sets of cluster-specific gene markers (C1, C2, C3, C4, C5, C6, C8, C9, SS, SD1, SD8 and SD10) were selected for Shigella/EIEC cluster typing. The sensitivity and specificity for each cluster-specific gene marker or a set of cluster-specific gene markers for the identification dataset were listed in Table 2. The cluster-specific gene markers or marker sets of markers were all 100% sensitive and 100% specific with exception of C1 (99.94%), C3 (99.91% specificity) and SS (99.8% specificity). A single specific gene for each of 53 sporadic EIEC lineages were also selected with the exception of one lineage which has a set of 2 genes. These genes were all 100% sensitive and specific for a given sporadic EIEC lineage.
All cluster-specific gene markers, 37 in total (5 single, 32 genes in 12 sets) and 54 sporadic EIEC lineages specific gene markers were located on chromosome but one of C4 gene markers and 5 sporadic EIEC lineages specific genes were located on plasmid. None of the cluster-specific gene markers were contiguous in the genomes. The location of these cluster-specific gene markers was determined by BLASTN against representative complete genomes of *Shigella*/EIEC containing gene features downloaded from NCBI GenBank. In those cluster or sporadic lineages with no representative complete genome specific gene markers were named using their cluster or sporadic EIEC lineage followed by the cluster or lineage number. For example, C7 specific gene marker was named “C7 specific gene”.

The functional characterization of these specific gene markers were identified from RAST annotation (74). For 37 cluster-specific gene markers, 22 had known functions and 15 encoded hypothetical proteins with unknown functions, while 11 sporadic EIEC lineages specific gene markers were identified with known functions and 43 were hypothetical proteins with unknown functions. The location and functions of specific gene markers are listed in Table S3.

**Validation of cluster-specific gene markers**

The ability of cluster-specific gene markers to correctly assign *Shigella*/EIEC isolates was evaluate with 15,501 *Shigella*/EIEC isolates in the validation dataset, 12,743 isolates from non-enteroinvasive *E. coli* control database.

Using cluster-specific gene markers, 15,443 of the 15,501 (99.63%) *Shigella*/EIEC isolates were correctly assigned to clusters which included 15,337 *Shigella* isolates, 102 EIEC isolates, 4 sporadic EIEC isolates, and 38 (0.24%) isolates with more than one clusters. Twenty of the 15,501 (0.13%) *Shigella*/EIEC isolates were not assigned to any of identified clusters.

To confirm the assignment of cluster-specific gene markers, we constructed a “validation” phylogenetic tree (Fig. S4) using 1,159 representative isolates from the validation dataset and a subset of 485 isolates from each cluster from the identification dataset. Isolates that grouped with known cluster isolates (from identification dataset) with strong bootstrap support were assigned to that cluster. All 1,159 isolates were grouped into known clusters on the validation phylogenetic tree. The cluster-specific gene markers assignments were entirely consistent with cluster assignments by phylogenetic tree.
We tested cluster-specific gene markers with the 12,743 non-enteroinvasive *E. coli* isolates. The *Shigella/EIEC* cluster-specific gene markers were highly specific with specificity varying from 98.8% to 100% for cluster-specific genes and 97.02% to 100% for sporadic EIEC specific genes. Details are listed in Table S4.

**Development an automated pipeline for molecular serotyping of *Shigella/EIEC***

Above results showed that cluster-specific gene markers were sensitive and specific and can distinguish *Shigella* and EIEC isolates. We therefore used these genes combined with established *Shigella/EIEC* serotype specific O antigen and H antigen genes to develop an automated pipeline for *in silico* serotyping of *Shigella/EIEC* (Fig. 2).

The pipeline is named *Shigella* EIEC Cluster Enhanced Serotype Finder (ShigEiFinder). ShigEiFinder can process either paired end Illumina sequencing reads or assembled genomes (https://github.com/LanLab/ShigEiFinder). ShigEiFinder classifies isolates into Non-*Shigella/EIEC*, *Shigella* or EIEC clusters based on the presence of *ipaH* gene, number of virulence genes, cluster specific genes. The “Not *Shigella/EIEC*” assignment was determined by the absence of *ipaH* gene, virulence genes (>25/38) and cluster-specific gene markers. The “*Shigella* or EIEC clusters” assignments were made based on the presence of *ipaH* gene, and/or more than 25 virulence genes together with the presence of any of cluster-specific gene markers or marker set, whereas the presence of *ipaH* gene and/or more than 25 virulence genes with absence of any of cluster-specific gene markers were assigned as “*Shigella/EIEC* unclustered”.

*Shigella* and EIEC isolates were differentiated and serotypes were assigned after cluster assignment. ShigEiFinder predicts a serotype through examining the presence of any of established *Shigella* serotype specific O antigen and modification genes and *E. coli* O and H antigen genes that differentiate the serotypes as ShigaTyper and SerotypeFinder (10, 63). A “novel serotype” is assigned if no match to known serotypes.

Two pairs of *Shigella* serotypes, SB1/SB20 and SB6/SB10, are known to be difficult to differentiate as they share identical O antigen genes (10, 46, 75). ShigaTyper used a heparinase gene for the differentiation of SB20 from SB1 and *wbaM* gene for the separation of SB6 from SB10. We found that fragments of the heparinase and *wbaM* genes may be present in other serotypes and cannot accurately differentiate SB1/SB20 and SB6/SB10. We found a SB20
specific gene which encoded hypothetical proteins with unknown functions and located on a plasmid by comparative genomic analysis of all isolates in C1 accessory genome. The SB20 specific gene can reliably differentiate SB20 from SB1 and also one SNP each in \textit{wzx} and \textit{wzy} genes that can differentiate SB6 from SB10. We used these differences (Data S1) in ShigEiFinder for the prediction of these serotypes.

**The accuracy and specificity of ShigEiFinder in cluster typing**

The accuracy of ShigEiFinder was tested with 1,969 isolates (1,969 assembled genomes and 1,951 Illumina reads [note no reads available for 18 EIEC isolates from NCBI] from the identification dataset and 15,501 isolates from the validation dataset. The results are listed in Table 3.

ShigEiFinder was able to assign 99.54% and 99.28% of the isolates in the identification dataset to clusters for assembled genomes and reads mapping respectively. The accuracy was 99.70% and 99.81% for assembled genomes and reads mapping respectively when applied to the validation dataset. Discrepancies were observed between assembled genomes and reads mapping (Table 3). There were more isolates assigned to “\textit{Shigella}/EIEC unclustered” in reads mapping, in contrast there were more isolates assigned to multiple clusters in genome assemblies. The specificity of ShigEiFinder was 99.40% for assembled genomes and 99.38% for reads mapping when evaluated with 12,743 non-\textit{Shigella}/EIEC \textit{E. coli} isolates. An additional 2 isolates were detected as sporadic EIEC lineages by reads mapping.

**Comparison of ShigEiFinder and ShigaTyper**

To demonstrate ShigEiFinder for differentiation of \textit{Shigella} from EIEC and enhancement of cluster based serotyping, the comparison of reads mapping results between ShigEiFinder and the existing \textit{in silico} \textit{Shigella} identification pipeline ShigaTyper (10) was performed with 488 isolates used in ShigaTyper and 15,501 isolates from \textit{Shigella}/EIEC validation dataset used in the present study.

The 488 isolates used in ShigaTyper consisted of 23 other species, 45 \textit{E. coli} isolates and 420 \textit{Shigella} isolates. ShigEiFinder identified 23 other species isolates and 453 out of 465 \textit{E. coli} and \textit{Shigella} isolates correctly, in agreement with ShigaTyper assignment. ShigEiFinder also assigned the remaining 12 \textit{E. coli} and \textit{Shigella} isolates including 3 EIEC isolates and 9
untypable (either multiple wzx or no wzx genes found) isolates by ShigaTyper to Shigella/EIEC clusters.

ShigEiFinder assigned 15,471 of 15,501 Shigella/EIEC isolates to Shigella or EIEC clusters and then to a serotype. The accuracy of ShigEiFinder to correctly assign isolates to Shigella or EIEC clusters was 99.81% (15,471/15,501). By contrast, ShigaTyper assigned 7,277 isolates (46.95%) to Shigella, 7,976 isolates (51.45%) to EIEC, 177 (1.14%) isolates to multiple wzx genes and failed to type 71 (0.46%) isolates.

The predicted serotype of 7,277 (46.96%) Shigella isolates by ShigaTyper agreed with the results of ShigEiFinder. For 8,224 isolates typed as EIEC or untypable by ShigaTyper, 99.73% (8,202/8,224) of the isolates were assigned to Shigella or EIEC clusters by ShigEiFinder (Table 4). Of these isolates, the majority belonged to SS, SD1 and SF which were erroneously predicted as EIEC by ShigaTyper.

**Discussion**

*Shigella* and EIEC cause human bacillary dysentery with similar invasion mechanisms, however the pathogenicity of these 2 groups varies (8, 43). The prevalence of each of the four *Shigella* “species” also varies (11-13). Differentiation of *Shigella* and EIEC from each other is important for epidemiologic and diagnostic investigations. However, their similar physiological, biochemical and genetic characteristics make this differentiation difficult.

**Determining phylogenetic clusters for better separation Shigella isolates from EIEC**

From phylogenetic perspective, *Shigella* and EIEC strains consisted of multiple phylogenetic lineages derived from commensal *E. coli*, which do not reflect the nomenclature of *Shigella* and EIEC (23, 25, 26, 28, 38, 41). In the present study, we identified all phylogenetic clusters of *Shigella* and EIEC through large scale examination of publicly available genomes.

Phylogenetic results demonstrated that *Shigella* isolates had at least 10 clusters while EIEC isolates had at least 7 clusters. The 10 *Shigella* clusters included the 7 previously defined lineages including 3 major clusters (C1, C2 and C3) and 5 outliers (SD1, SD8, SD10, SB13 and SS) (25) and 2 newly identified clusters (SB12 and SB13-atypical). The 7 EIEC clusters consisted of 4 previously defined EIEC clusters (C4, C5, C6 and C7) (26) and 3 newly identified EIEC clusters (C8 EIEC O96:H19, C9 EIEC O8:H19 and C10 EIEC O135:H30).
Our WGS-based phylogeny provided high resolution for assigning *Shigella* and EIEC isolates to clusters. Several serotypes that are currently increasing in frequency (SB19, SB20, SD14, SD15, SD provisional serotype 96-626) (76-79) were assigned to clusters and five new clusters/outliers were identified. SB13 isolates in this study formed two known lineages. One lineage was located outside of *Shigella/EIEC* clusters and represented the outlier SB13 which is in fact belonging to the newly defined species *E. albertii* (25, 26, 38, 39). The second lineage was with *E. coli*, and was defined as atypical SB13 previously (39). The newly identified *Shigella* outlier SB12 was previously grouped into C3 based on housekeeping gene trees (25, 38) but was seen as outliers in two other studies (28, 56).

Newly identified clusters C8 (EIEC O96:H19) and C9 (EIEC O8:H19) represented the emergence of novel EIEC serotypes. A recent study revealed that EIEC serotype O96:H19 (C8) could be the result of a recent acquisition of the invasion plasmid by commensal *E. coli* (80). The EIEC serotype O8:H19 (C9) had not been reported previously.

Apart from the 17 major and outlier clusters of *Shigella* and EIEC, the presence of 53 sporadic EIEC lineages indicated greater genetic diversity than has been observed previously. Isolates belonging to these sporadic EIEC groups were more closely related to non-enteroinvasive *E. coli* isolates than to major *Shigella/EIEC* lineages. However, 41 of the isolates representing 38 sporadic EIEC lineages that carried pINV. *Shigella* and EIEC both carry the *Shigella* virulence plasmid pINV which is vital for virulence and distinguishes *Shigella/EIEC* from other *E. coli* (24, 33, 68). Therefore, these isolates may represent recently formed EIEC lineages through acquisition of the pINV. The remaining 18 isolates contained the *ipaH* gene but may or may not carry pINV. It is possible that these strains carried very low copy number of pINV or the pINV plasmid was lost during culture.

**Highly sensitive and specific cluster-specific gene markers for differentiation of *Shigella* and EIEC isolates**

Several studies have identified phylogenetic related genomic markers for discrimination of *Shigella* and EIEC and between *Shigella* species (23, 27, 28, 41, 55, 56). However, these phylogenetic analyses were performed only with a small number of genomes (23, 28, 55). In addition, non-invasive *E. coli* isolates were included in some of the phylogenetic clusters identified (28) which led to non-invasive *E. coli* isolates being identified by the markers.
We identified cluster-specific gene markers for each respectively clusters which were only composed of Shigella or EIEC isolates. Sets of cluster-specific gene markers were identified for those clusters where no single suitable marker is present. The combination of genes enhances the specificity of cluster-specific gene markers as demonstrated by the 100% sensitivity and very high specificity in this analysis (Table 2). Genes specific to each of the 53 sporadic EIEC lineages were also identified and they were sensitive and specific, although it should be noted that these values are based on very small sample sizes.

The cluster-specific gene markers or marker sets can be used to differentiate Shigella/EIEC from non-enteroinvasive E. coli independent of ipaH gene. The ipaH gene as a molecular target has been used to differentiate Shigella and EIEC from non-enteroinvasive E. coli (24, 43-45). In our study, the cluster-specific gene markers were specific to Shigella/EIEC with 98.8% to 100% specificity when evaluated on non-enteroinvasive E. coli control database, giving us the confidence that the cluster-specific genes or sets are robust markers to identify Shigella/EIEC. 53 sporadic EIEC lineage specific gene markers also have very high specificity (97.02% to 100%) against non-enteroinvasive E. coli control database.

The cluster-specific gene markers or marker sets are able to assign Shigella/EIEC isolates correctly in 99.63% of cases and can clearly distinguish Shigella isolates from EIEC when applied to the validation dataset. While ShigaTyper assigned 46.95% isolates to Shigella and 51.45% isolates to EIEC in the same dataset we tested, leading to a large proportion of isolates incorrectly assigned. The majority of the isolates predicted as EIEC by ShigaTyper were SS or SD1 as they belonged to SS and SD1 specific STs and were positive to a set of SS or SD1 specific gene markers and grouped into SS or SD1 cluster on our phylogenetic tree. The genes used in ShigaTyper were SS specific marker Ss_methylase gene (81, 82) together with SS O antigen wzx gene. However, SS specific marker Ss_methylase gene was found in other Shigella serotypes and EIEC (10) and SS O antigen wzx gene were located on a plasmid which is frequently lost (83). Similarly, the SD1 O antigen genes used in ShigaTyper were plasmid-borne which may also lead to inconsistent detection (84, 85). A previous study identified 6 loci to distinguish EIEC from Shigella (23). We searched the 6 loci against our Shigella/EIEC database and found that some Shigella isolates were misidentified as EIEC isolates, such as SD8 isolates incorrectly identified as EIEC subtype 13. Our cluster-specific genes can differentiate SD8 isolates from EIEC with 100% accuracy. Therefore, the cluster-specific gene markers marker sets provided nearly perfect differentiation of Shigella from EIEC.
The cluster-specific gene markers or marker sets are able to differentiate SS and SF (with exception of SF6) from SB and SD. SF and SS are the major cause of *Shigella* infections, accounting for up to 89.6% annual cases (11-13). Differentiation of SS and SF isolates from SB and SD is also beneficial for diagnosis and surveillance. A recent study identified “species” specific markers for the detection of each of the four *Shigella* “species” and validated with only one isolate per species (55). A molecular algorithm based on *Shigella* O antigen genes can detect 85% of SF isolates (52). In contrast, a set of SF specific genes in our study can correctly identify SF isolates with 99.62% accuracy.

The cluster-specific gene markers or marker sets can also assign *Shigella/EIEC* isolates to serotype level if the cluster has single serotype such as SD1, SD8, SD10, SB13, SB12, EIEC O144:H25 (C7), EIEC O96:H19 (C8), EIEC O8:H19 (C9) and EIEC O135:H30 (C10). The remaining EIEC, SF, SB and SD serotypes were distributed over the major clusters C4-6, C3, C1 and C2 respectively. Cluster-specific gene markers combined with serotype associated O antigen and modification genes can further identify these isolates to serotype level.

**Cluster-specific gene marker based ShigEiFinder can accurately type *Shigella* and EIEC**

To facilitate the use of cluster-specific gene markers or marker sets for typing, we developed an automated pipeline, ShigEiFinder, for *in silico* molecular serotyping of *Shigella/EIEC*. ShigEiFinder provided *Shigella/ EIEC* differentiation as well as serotype prediction by yielding “presence or absence” of cluster-specific gene markers or marker sets combined with *Shigella/EIEC* O antigen genes and modification genes in a query isolate (either reads or assembled genomes). We showed 99.70% and 99.81% accuracy to assign isolates to the correct clusters from 15,501 *Shigella/EIEC* isolates in validation dataset for the assembled genomes and reads mapping respectively. In contrast, the existing *in silico* *Shigella* serotyping pipeline ShigaTyper had 46.95% accuracy for reads mapping when tested with the same validation dataset, with 51.45% of isolates in validation dataset being predicted as EIEC by ShigaTyper.

The genetic determinants used in ShigaTyper for differentiation of *Shigella* from EIEC and identification of SS were *lacY*, *cadA*, *Ss_methylase*, SS and SD1 O antigen *wzx* genes (10). As discussed above some of these genes were found to be non-specific in this study. Compared with ShigaTyper, the cluster-specific gene markers used in ShigEiFinder for identification of *Shigella* and EIEC provided higher discriminatory power than ShigaTyper. ShigEiFinder also
provided a high specificity with 99.40% for assembled genomes and 99.38% for reads mapping. ShigEiFinder can differentiate *Shigella* isolates from EIEC and distinguish SS and SF (with exception of SF6) isolates from SB and SD accurately. It also can identify SD1 isolates directly. ShigEiFinder was able to serotype over 59 *Shigella* serotypes and 22 EIEC serotypes. Therefore, ShigEiFinder will be useful for clinical, epidemiological and diagnostic investigations and the cluster-specific gene markers identified could be adapted for metagenomics or culture independent typing.

**Conclusion**

This study analysed over 17,000 publicly available *Shigella*/EIEC genomes and identified 10 clusters of *Shigella*, 7 clusters of EIEC and 53 sporadic types of EIEC. Cluster-specific gene markers or marker sets for the 17 major clusters and 53 sporadic types were identified and found to be valuable for *in silico* typing. We additionally developed a freely available *in silico* serotyping pipeline incorporating the cluster-specific gene markers to facilitate serotyping of *Shigella*/EIEC isolates using genome sequences with very high specificity and sensitivity.

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**Author contributions**

Conceptualization: R.L, M.P.; Investigation: X.Z., M.P., T.N., S.K.; Methodology: M.P., R.L. Writing – original draft: X.Z.; Writing – review and editing: M.P., R.L.

**Conflicts of interest**

The authors declare that there are no conflicts of interest.

**Data bibliography**
All the sequencing data generated within this study, NCBI BioProject number (PRJNA692536).

### Abbreviations

SS, *Shigella sonnei*; SF, *Shigella flexneri*; SB, *Shigella boydii*; SD, *Shigella dysenteriae*; EIEC, Enteroinvasive *Escherichia coli*; NCBI SRA, National Center for Biotechnology Information Sequence Read Archive; ST, sequence type; rST, ribosomal ST; MLST, Multilocus sequence typing; rMLST, Ribosomal MLST; ECOR, *Escherichia coli* reference collection; WGS, whole-genome sequencing; TP, true positive; FN, false negative; FP, false positive; HK, House Keeping.

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| Clusters (no of serotypes) | No of isolates | No. STs | No. rSTs | Serotypes |
|---------------------------|----------------|---------|----------|-----------|
| C1 (25)                   | 288            | 36      | 166      | SB1-4, SB6, SB8, SB10, SB14, SB18, SB11\textsuperscript{b}, SB19-20\textsuperscript{b}; SD3-7, SD9, SD11-13, SD14-15\textsuperscript{*}, SD-96-265\textsuperscript{*}; SF6 |
| C2 (9)                    | 101            | 19      | 56       | SB5, SB7, SB9, SB11, SB15, SB16, SB17; SD2, SD-E670-74\textsuperscript{b}; SD2 |
| C3 (20)                   | 744            | 81      | 437      | SF1a, SF1b, SF1c (7a), SF2a, SF2b, SF3a, SF3b, SF4a, SF4av, SF4b, SF4bv, SF5a, SF5b, SF7b, SFX, SFXv (4c), SFY, SFYv, SF novel serotype; SB-E1621-54\textsuperscript{*} |
| C4 (9)                    | 51             | 6       | 21       | O28ac:H-, O28ac:H7, O136:H7, O164:H-, O164:H7, O29:H4, O173:H7, O124:H7, O132:H7\textsuperscript{*} |
| C5 (6)                    | 62             | 4       | 15       | O121:H30, O124:H30, O164:H30, O132:H21, O152:H30, O152:H- |
| C6 (3)                    | 20             | 2       | 6        | O143:H26, O167:H26, O112ac:H26\textsuperscript{b} |
| C7                        | 10             | 1       | 3        | O144:H25 |
| C8\textsuperscript{a}     | 12             | 2       | 1        | O96:H19 |
| C9\textsuperscript{a}     | 4              | 1       | 2        | O8:H19 |
| C10 \textsuperscript{*}   | 2              | 1       | 1        | O135:H30 |
| CSS                       | 427            | 39      | 294      | |
| CSD1                      | 70             | 8       | 56       | SD1 |
| CSD8                      | 7              | 3       | 3        | SD8 |
| CSD10                     | 2              | 2       | 1        | SD10 |
| CSB12\textsuperscript{a}  | 8              | 2       | 6        | SB12 |
| CSB13                     | 7              | 3       | 3        | SB13 |
| Clusters (no of serotypes)* | No of isolates | No. STs | No. rSTs | Serotypes         |
|-----------------------------|----------------|---------|----------|------------------|
| CSB13-atypical*             | 5              | 3       | 3        | SB13             |
| Sporadic EIEC lineagesa (53)| 59             | 49      | 53       | 53 antigen types |

*Numbers in parentheses are the number of serotypes within that cluster.

a: Clusters identified as new clusters in this study.

b: Serotypes were inconsistent with previous analyses.
Table 2: The sensitivity and specificity of cluster-specific genes

| Clusters | Cluster-specific genes (Single/sets)b | Identification dataset (1969 isolates) |  |
|----------|--------------------------------------|----------------------------------------|---|
|          |                                      | No of isolates | Sensitivity | Specificity |
| C1       | Set of 4 genes                       | 288           | 100         | 99.94\(^a\) |
| C2       | Set of 3 genes                       | 101           | 100         | 100         |
| C3       | Set of 3 genes                       | 744           | 100         | 99.59\(^a\) |
| C4       | Set of 2 genes                       | 51            | 100         | 100         |
| C5       | Set of 3 genes                       | 62            | 100         | 100         |
| C6       | Set of 2 genes                       | 20            | 100         | 100         |
| C7       | Single gene                          | 10            | 100         | 100         |
| C8       | Set of 2 genes                       | 12            | 100         | 100         |
| C9       | Set of 2 genes                       | 4             | 100         | 100         |
| C10      | Single gene                          | 2             | 100         | 100         |
| CSS      | Set of 5 genes                       | 427           | 100         | 99.87\(^a\) |
| CSD1     | Set of 2 genes                       | 70            | 100         | 100         |
| CSD8     | Single gene                          | 7             | 100         | 100         |
| CSD10    | Single gene                          | 2             | 100         | 100         |
| CSB12    | Single gene                          | 8             | 100         | 100         |
| CSB13    | Single gene                          | 7             | 100         | 100         |
| CSB13-atypical | Single gene | 5 | 100 | 100 |
| 53 Sporadic EIEC lineages | Single gene / lineage | 59 | 100 | 100 |

\(^a\): The specificity of cluster-specific gene set less than 100% was due to at least one FP found in that set.

\(^b\): The sequences of these genes were listed in Data S1.
Table 3: The accuracy of ShigEiFinder with identification dataset and validation dataset

| ShigEiFinder assignments       | Identification Dataset (n=1,969) | Validation dataset (n=15,501) |
|-------------------------------|---------------------------------|--------------------------------|
|                               | Genomes | Reads mapping | Genomes | Reads mapping |
| Shigella/EIEC clusters        | 1871     | 1848          | 15,455  | 15,471        |
| Multiple Shigella/EIEC clusters | 9       | 6             | 33      | 7             |
| Shigella/EIEC unclustered     | 0        | 8             | 13      | 23            |
| Not Shigella/EIEC             | 89       | 89            | 0       | 0             |
| Accuracy<sup>b</sup>          | 99.54%   | 99.28%        | 99.70%  | 99.81%        |

<sup>a</sup>: Identification dataset has 90 non-Shigella/EIEC strains including 72 ECOR strains and 18 *E.albertii* strains. 1,969 assembled genomes and 1,951 reads (reads not available for 18 EIEC isolates downloaded from NCBI) in identification dataset. One of *E.albertii* strain was assigned as SB13 which was grouped into SB13 cluster on the phylogenetic tree.

<sup>b</sup>: The accuracy was defined as the number of Shigella/EIEC isolates being correctly assigned to cluster over the total number of tested.
### Table 4: Discrepant assignment of 8,224 isolates by ShigEiFinder and Shigatyper

| ShigEiFinder Assignment | ShigaTyper assignment | Total |
|-------------------------|-----------------------|-------|
|                         | EIEC                  | Multiple wzx | Non-prediction |     |
| SS                      | 7,465                 | 12          | 7              | 7,484 |
| SF                      | 117                   | 61          | 10             | 188   |
| C1 and C2 (SB/SD)       | 17                    | 99          | 51             | 167   |
| SB12                    | 0                     | 2           | 0              | 2     |
| SD1                     | 244                   | 1           | 1              | 246   |
| SD8                     | 1                     | 0           | 0              | 1     |
| SD10                    | 0                     | 0           | 2              | 2     |
| EIEC                    | 97                    | 0           | 0              | 97    |
| Sporadic EIEC lineages  | 15                    | 0           | 0              | 15    |
| Multiple clusters       | 5                     | 2           | 0              | 7     |
| Shigella/EIEC unclustered | 15               | 0           | 0              | 15    |
| Total                   | 7,976                 | 177         | 71             | 8,224 |
Figure legends:

Figure 1: *Shigella*/EIEC cluster **Identification phylogenetic tree**

Representative isolates from the identification dataset were used to construct the phylogenetic tree by Quicktree v1.3 (64) to identify *Shigella* and EIEC clusters and visualised by Grapetree’s interactive mode. The dendrogram tree shows the phylogenetic relationships of 1879 *Shigella* and EIEC isolates represented in the identification dataset. Branch lengths are log scale for clarity. The tree scales indicated the 0.2 substitutions per locus. *Shigella* and EIEC clusters are coloured. Numbers in square brackets indicate the number of isolates of each identified cluster. CSP is sporadic EIEC lineages.

Figure 2: *in silico* serotyping pipeline workflow

Schematic of *in silico* serotyping *Shigella* and EIEC by cluster-specific genes combined with the *ipaH* gene and O antigen and modification genes and H antigen genes, implemented in ShigEiFinder. Both assembled genomes and raw reads are accepted as data input.
**Supplementary Material**

**Figure S1: Identification phylogenetic tree**

An identification phylogenetic tree constructed by Quicktree v1.3 (64) and visualised by ITOL v5 shows the phylogenetic relationships of 1879 Shigella and EIEC isolates in identification dataset. The tree scales indicated the 0.01 substitutions per locus. Shigella and EIEC clusters are colored. The internal branches are colored to represent the bootstrap values. Green color indicates the maximum bootstrap value (1). The red color shows the minimum bootstrap value (0). Each of cluster is well supported by bootstrap value. CSP is sporadic EIEC lineages.

**Figure S2-A: Confirmation phylogenetic tree**

A confirmation phylogenetic tree was constructed by Quicktree v1.3 (64) based on 2375 isolates and visualised by Grapetree’s interactive mode. The tree shows the phylogenetic relationships between identified Shigella/EIEC clusters in identification dataset and non-enteroinvasive E. coli isolates. Branch lengths are log scale for clarity. The tree scales indicated the 0.1 substitutions per locus. Known Shigella and EIEC clusters from identification dataset are colored. Numbers in square brackets indicate the number of isolates of each identified cluster. CSP is sporadic EIEC lineages.

**Figure S2-B: Confirmation phylogenetic tree**

A confirmation phylogenetic tree constructed by Quicktree v1.3 (64) and visualised by ITOL v5 shows the phylogenetic relationships between identified Shigella/EIEC clusters in identification dataset and non-enteroinvasive E. coli isolates. The tree scales indicated the 0.01 substitutions per locus. Shigella and EIEC clusters are colored. The internal branches are colored to represent the bootstrap values. Green color indicates the maximum bootstrap value (1). The red color shows the minimum bootstrap value (0). Each of cluster is well supported by bootstrap value. CSP is sporadic EIEC lineages.

**Figure S3: Distribution of mapped 38 virulence genes in 58 sporadic isolates**

The presence of Shigella virulence plasmid pINV in 58 sporadic isolates in identification dataset was determined by the mapped 38 virulence genes. Detailed genes were described in Results “Investigation of Shigella virulence plasmid pINV in 58 sporadic isolates”. Three categories were defined based on the number of virulence genes mapped to isolate. Virulence plasmid positive: > 25 genes mapped to isolate; Intermediate: 13 to 25 genes mapped to isolate; Virulence plasmid negative: less than 13 genes mapped to isolate.
Figure S4 (A): Validation phylogenetic tree
A validation tree was generated by Quicktree v1.3 (64) and visualised by Grapetree’s interactive mode to assign representative isolates in validation dataset to clusters. Branch lengths are log scale for clarity. The tree scales indicated the 0.2 substitutions per locus. Known Shigella and EIEC clusters from identification dataset are colored. Numbers in square brackets indicate the number of isolates of each identified cluster. Isolates in validation dataset are colored white. The isolates are assigned to clusters if they grouped into known cluster isolates. CSP is sporadic EIEC lineages.

Figure S4 (B): Validation phylogenetic tree
A validation phylogenetic tree was constructed by Quicktree v1.3 (64) and visualised by ITOL v5 to assign representative isolates in validation dataset to clusters. The tree scales indicated the 0.01 substitutions per locus. Shigella and EIEC clusters are colored. The internal branches are colored to represent the bootstrap values. Green color indicates the maximum bootstrap value (1). The red color shows the minimum bootstrap value (0). Each of cluster is well supported by bootstrap value. Isolates that grouped with known cluster isolates (from identification dataset) with strong bootstrap support are categorised into that cluster. CSP is sporadic EIEC lineages.

Table S1: 1,969 isolates used in identification dataset
Table S2: 15,501 isolates used in validation dataset
Table S3: The location and function of cluster-specific genes
Table S4: The results of cluster-specific gene markers tested with 12,743 non-enteroinvasive E. coli isolates

Data S1: Algorithms incorporated into the ShigEiFinder
Data S2: Genetic signature O and H genes from ShigaTyper and SerotypeFinder

Data Availability Statement
Custom python scripts used in this study are available from the authors on request.
