1	Genomic Diversity of Escherichia coli isolates from healthy children in rural Gambia
2	Ebenezer Foster-Nyarko ^{1,2} , Nabil-Fareed Alikhan ¹ , Usman Nurudeen Ikumapayi ² , Sarwar
3	Golam ² , M Jahangir Hossain ² , Catherine Okoi ² , Peggy-Estelle Tientcheu ² , Marianne
4	Defernez ¹ , Justin O'Grady ¹ , Martin Antonio ^{2,3} , Mark J. Pallen ^{1,4#}
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6	
7	¹ Quadram Institute Bioscience, Norwich Research Park, Norwich, Norfolk, United Kingdom
8	² Medical Research Council Unit The Gambia at the London School of Hygiene and Tropical
9	Medicine, Atlantic Boulevard Road, Fajara, the Gambia
10	³ Warwick Medical School, University of Warwick, Coventry, United Kingdom
11	⁴ School of Veterinary Medicine, University of Surrey, Guildford, Surrey, United Kingdom
12	
13	
14	[#] Corresponding author: Professor Mark J. Pallen, Quadram Institute Bioscience, Norwich
15	Research Park, Norwich, Norfolk, United Kingdom
16	Email: Mark.Pallen@quadram.ac.uk

18 Abstract

19	Little is known about the genomic diversity of Escherichia coli in healthy children from sub-
20	Saharan Africa, even though this is pertinent to understanding bacterial evolution and
21	ecology and their role in infection. We isolated and whole-genome sequenced up to five
22	colonies of faecal E. coli from 66 asymptomatic children aged three-to-five years in rural
23	Gambia (n=88 isolates from 21 positive stools). We identified 56 genotypes, with an average
24	of 2.7 genotypes per host. These were spread over 37 seven-allele sequence types and all
25	eight significant phylogroups of E. coli. Immigration events accounted for three quarters of
26	the diversity within our study population, while one quarter of variants appeared to have
27	arisen arose from within-host evolution. Several study strains were closely related to isolates
28	that caused disease in humans or originated from livestock. Our results suggest that within-
29	host evolution plays a minor role in the generation of diversity than independent immigration
30	and the establishment of strains among our study population. Also, this study adds
31	significantly to the number of commensal E. coli genomes, a group that has been traditionally
32	underrepresented in the sequencing of this species.
33	
34	Words: 184
35	Keywords: Escherichia coli, genomic diversity, within-host evolution.
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38 Introduction

39	Ease of culture and genetic tractability account for the unparalleled status of Escherichia coli
40	as "the biological rock star", driving advances in biotechnology (1), while also providing key
41	insights into biology and evolution (2). However, E. coli is also a widespread commensal, as
42	well as a versatile pathogen, linked to diarrhoea (particularly in the under-fives), urinary tract
43	infection, neonatal sepsis, bacteraemia and multi-drug resistant infection in hospitals (3-5).
44	Yet most of what we know about E. coli stems from investigation of laboratory strains, which
45	fail to capture the ecology and evolution of this key organism "in the wild" (6) What's more,
46	most studies of non-lab strains have focused on pathogenic strains or have been hampered by
47	low-resolution PCR methods, so we have relatively few genomic sequences from commensal
48	isolates, particularly from low- to middle-income countries (7-13).
49	We have a broad understanding of the population structure of <i>E. coli</i> , with eight major
50	phylogroups loosely linked to ecological niche and pathogenic potential (B2, D and F linked
51	to extraintestinal infection; A and B1 linked to severe intestinal infections such as
52	haemolytic-uraemic syndrome) (14-17), All phylogroups can colonise the human gut, but it
53	remains unclear how far commensals and pathogenic strains compete or collaborate with one
54	another—or engage in horizontal gene transfer—within this important niche (18, 19).
55	Although clinical microbiology typically relies on single-colony picks (which has the
56	potential to underestimate species diversity and transmission events), within-host diversity of
57	E. coli in the gut is crucial to our understanding of inter-strain competition and co-operation
58	and also for accurate diagnosis and epidemiological analyses. Pioneering efforts using
59	serotyping and molecular typing have shown that normal individuals typically harbour more
60	than one strain of <i>E. coli</i> (20-22), with one individual carrying 24 distinct clones (22-24).
61	More recently, whole-genome sequencing has illuminated molecular epidemiological

62 investigations (9), adaptation during and after infection (25, 26), as well as the intra-clonal

63 diversity in healthy hosts (27).

64	There are two plausible sources of within-host genomic diversity. Although, a
65	predominant strain usually colonises the host for extended periods (28), successful
66	immigration events mean that incoming strains can replace the dominant strain or co-exist
67	alongside it as minority populations (29). Strains originating from serial immigration events
68	are likely to differ by hundreds or thousands of single-nucleotide polymorphisms (SNPs).
69	Alternatively, within-host evolution can generate clouds of intra-clonal diversity, where
70	genotypes differ by just a handful of SNPs (20).
71	Most relevant studies have been limited to Western countries, with the exception of a
72	recent report from Tanzania (21), so little is known about the genomic diversity of E. coli in
73	sub-Saharan Africa. The Global Enteric Multicenter Study (GEMS) (30, 31) has documented
74	a high burden of diarrhoea attributable to E. coli (including Shigella) among children from
75	the Gambia, probably as a result of increased exposure to this organism through poor hygiene
76	and frequent contact with animals and the environment. In also facilitating access to stool
77	samples from healthy Gambian children, the GEMS study has given us a unique opportunity
78	to study within-host genomic diversity of commensal E. coli in this setting.
79	

80 Methods

81 **Study population**

We initially selected 76 faecal samples from three- to five-year-old asymptomatic Gambian
children, who had been recruited from Basse, Upper River Division, the Gambia, into the
GEMS study (30) as healthy controls from December 1, 2007 to March 3, 2011. Samples had
been collected according to a previously described sampling protocol (32). Archived stool
samples were retrieved from -80°C storage and allowed to thaw on ice. A 100-200 mg aliquot

from each sample was transferred aseptically into 1.8ml Nunc tubes for microbiological
processing below (Figure 1). Eleven of the original 76 samples proved unavailable for
processing in this study.

90

91 Bacterial growth and isolation

1 ml of physiological saline (0.85%) was added to each sample tube and vigorously vortexed
at 4200 rpm for at least 2 minutes. Next, the homogenised sample suspensions were taken

94 through four ten-fold dilution series. A100 µl aliquot from each dilution was then spread

- 95 evenly on a plate of tryptone-bile-X-glucuronide differential and selective agar. The
- 96 inoculated plates were incubated overnight at 37°C under aerobic conditions. Colony counts

97 were performed on the overnight cultures for each serial dilution for translucent colonies with

98 entire margins and blue-green pigmentation indicative of *E. coli*. Up to five representative

99 colonies were selected from each sample and sub-cultured on MacConkey agar overnight at

100 37°C before storing in 20% glycerol broth at -80°C. Individual isolates were assigned a

101 designation comprised of the subject ID followed by the colony number ("1-5").

102

103 Genomic DNA extraction and genome sequencing

104 Broth cultures were prepared from pure, fresh cultures of each colony-pick in 1 ml Luria-

105 Bertani broth and incubated overnight to attain between $10^9 - 10^{10}$ cfu per ml. Genomic DNA

106 was then extracted from the overnight broth cultures using the lysate method described in

107 (33). The eluted DNA was quantified by the Qubit high sensitivity DNA assay kit

108 (Invitrogen, MA, USA) and sequenced on the Illumina NextSeq 500 instrument (Illumina,

109 San Diego, CA) as described previously (34).

110 Following Dixit et al. (20), we sequenced a random selection of isolates twice, using DNA

111 obtained from independent cultures, to help in the determination of clones and the analysis of

within-host variants (Supplementary File 5). Bioinformatic analyses of the genome sequences
were carried out on the Cloud Infrastructure for Microbial Bioinformatics (CLIMB) platform
(35).

115

116 **Phylogenetic analysis**

117 The paired 150bp reads were quality checked and assembled, as previously described (34).

118 Snippy v4.3.2 (https://github.com/tseemann/snippy) was used for variant calling, using the

119 complete genome sequence of commensal E. coli str. K12 substr. MG1655 as a reference

120 strain (NCBI accession: NC_000913.3) and to generate a core-genome alignment, from

121 which a maximum-likelihood phylogeny with 1000 bootstrap replicates was reconstructed

122 using RAxML v8.2.4 (36), based on a general time-reversible nucleotide substitution model.

123 The phylogenetic tree was rooted using the genomic sequence of *E. fergusonii* as an outgroup

124 (NCBI accession: GCA_000026225.1). The phylogenetic tree was visualised in FigTree

125 v1.4.3 (https://github.com/rambaut/figtree/) and annotated in RStudio v3.5.1 and Adobe

126 Illustrator v 23.0.3 (Adobe Inc., San Jose, California). For visualisation, a single colony was

127 chosen to represent replicate colonies of the same strain (ST) with identical virulence,

128 plasmid and antimicrobial resistance profiles and a de-replicated phylogenetic tree

129 reconstructed using the representative isolates.

130

131 Multi-locus sequence typing, Clermont typing and SNPs

132 The merged reads were uploaded to EnteroBase (37), where *de novo* assembly and genome

133 annotation were carried out and *in-silico* multi-locus sequence types (STs) assigned based on

- 134 the Achtman scheme, allocating new sequence types (ST) if necessary. EnteroBase assigns
- 135 phylogroups using ClermontTyper and EzClermont (38, 39) and unique core-genome MLST
- 136 types based on 2, 513 core loci in *E. coli*. Publicly available *E. coli* sequences in EnteroBase

- 137 (http://enterobase.warwick.ac.uk/species/index/ecoli) (37) were included for comparative
- 138 analysis, including 23 previously sequenced isolates obtained from diarrhoeal cases recruited
- in the GEMS study in the Gambia (Supplementary File 1).
- 140 We computed pairwise single nucleotide polymorphism (SNP) distances between
- 141 genomes from the core-genome alignment using snp-dists v0.6
- 142 (https://github.com/tseemann/snp-dists). For the duplicate sequence reads of the same strains,
- 143 we used SPAdes v3.13.2 (40) to assemble each set of reads and map the raw sequences from
- 144 one sequencing run to the assembly of the other run and vice versa, as described previously
- 145 (20). SNPs were detected using the CSIPhylogeny tool
- 146 (https://cge.cbs.dtu.dk/services/CSIPhylogeny/) and compared between the two steps,
- 147 counting only those SNPs that were detected in both sets of reads as accurate.
- 148

149 Accessory gene content

- 150 We used ABRicate v0.9.8 (https://github.com/tseemann/abricate) to predict virulence factors,
- 151 acquired antimicrobial resistance genes and plasmid replicons by scanning the contigs against
- 152 the VFDB, ResFinder and PlasmidFinder databases respectively, using an identity threshold
- 153 of \ge 90% and a coverage of \ge 70%. Virulence factors and AMR genes were plotted next to the
- 154 phylogenetic tree using the ggtree, ggplot2 and phangorn packages in RStudio v3.5.1. We
- 155 calculated co-occurrence of AMR genes among study isolates and visualised this as a heat
- 156 map using RStudio v 3.5.1.
- 157

158 **Population structure and comparison of commensal and pathogenic strains**

- 159 We assessed the population structure using the hierarchical clustering algorithm in
- 160 EnteroBase. Briefly, the isolates were assigned stable population clusters at eleven levels
- 161 (from HC0 to HC 2350) based on pairwise cgMLST allelic differences. Hierarchical

162 clustering at 1100 alleles differences (HC1100) resolves populations into cgST complexes,

- 163 the equivalent of clonal complexes achieved with the legacy MLST clustering approaches
- 164 (37). We reconstructed neighbour-joining phylogenetic trees using NINJA (41), based on
- 165 clustering at HC1100 to display the population sub-clusters at this level as an indicator of the
- 166 genomic diversity within our study population and to infer the evolutionary relationship
- 167 among our strains and others in the public domain.
- 168 Next, we interrogated the HC1100 clusters that included both pathogenic and commensal
- 169 E. coli strains recovered from the GEMS study. For the clusters that encompassed commensal
- 170 and pathogenic strains belonging to the same ST, we reconstructed both neighbour-joining
- and SNP phylogenetic trees to display the genetic relationships among these strains. We
- 172 visualised the accessory genomes for the overlapping STs mentioned above to determine
- 173 genes associated with phages, virulence factors and AMR. The resulting phylogenetic trees
- 174 were annotated in Adobe Illustrator v 23.0.3 (Adobe Inc., San Jose, California).
- 175

176 Ethical statement

- 177 The study was approved by the joint Medical Research Council Unit The Gambia-Gambian178 Government ethical review board.
- 179
- 180 **Results**

181 **Population structure**

182 The study population included 27 females and 39 males (Table 1). All but one reported the

- 183 presence of a domestic animal within the household. 21 samples proved positive for growth
- 184 of *E. coli*, yielding 88 isolates. We detected 37 seven-allele sequence types (STs) among the
- isolates, with a fairly even distribution (Figure 2). Five STs were completely novel (ST9274,
- 186 ST9277, ST9278, ST9279 and ST9281). These study strains were scattered over all the eight

187	main phylogroups of E. coli (Table 2). Hierarchical clustering of core genomic STs revealed
188	twenty-seven cgST clonal complexes (Supplementary File 2).

189

190 Within-host diversity

- 191 Nine individuals were colonised by just a single ST, six carried two STs, four carried four
- 192 STs and two carried six STs. We found 56 distinct genotypes, which equates to an average of
- 193 2.7 genotypes per host. Two individuals (H-18 and H-2) shared an identical strain belonging
- 194 to ST9274 (zero SNP difference) (Supplementary File 4, yellow highlight), suggesting recent
- 195 transfer from one child to another or recent acquisition from a common source.
- 196 We observed thirteen cases where a single host harboured two or more variants within the
- 197 same SNP cloud (Table 2). Such within-host evolution accounted for around a quarter of the
- 198 observed variation, with immigration explaining the remaining three quarters. 22% of within-
- 199 host mutations represented synonymous changes. 43% were non-synonymous mutations,
- 200 while 31% occurred in non-coding regions and 4% represented stop-gained mutations
- 201 (Supplementary File 6). The average number of SNPs among variants within such SNP
- 202 clouds was 5 (range 0-18) (Table 3). However, in two subjects (H36 and H37), pairwise
- distances between genomes from the same ST (ST59 and ST5148) were as large as 14 and 18
- 204 SNPs respectively (Supplementary File 4, grey highlight).
- 205

206 Accessory gene content and relationships with other strains

207 A quarter of our isolates were most closely related to commensal strains from humans, with

- 208 smaller numbers most closely related to human pathogenic strains or strains from livestock,
- 209 poultry or the environment (Table 4). One isolate was most closely related to a canine isolate
- 210 from the UK. Three STs (ST38, ST10 and ST58) were shared by our study isolates and
- 211 diarrhoeal isolate from the GEMS study (Supplementary Figure 2), with just eight alleles

212 separating our commensal ST38 strain from a diarrhoeal isolate from the GEMS study

213 (Figure 5).

214	We detected 125 genes encoding putative virulence factors across the 88 study isolates
215	(Figure 2; Supplementary File 3). More than half of the isolates encoded resistance to three or
216	more clinically relevant classes of antibiotics (Figure 3; Supplementary Figure 1). The most
217	common resistance gene network was $-aph(6)-Id_1-sul2$ (41% of the isolates), followed by
218	<i>aph</i> (3")- <i>Ib_5-sul2</i> (27%) and <i>bla-TEM-aph</i> (3")- <i>Ib_5</i> (24%). Most isolates (67%) harboured
219	two or more plasmid types (Figure 4). Of the 24 plasmid types detected, IncFIB was the most
220	common (41%), followed by col156 (19%) and IncI_1-Alpha (15%). Nearly three quarters of
221	the multi-drug resistant isolates carried IncFIB (AP001918) plasmids, suggesting that these
222	large plasmids disseminate resistance genes within our study population.
223	

224 **Discussion**

225 This study provides an overview of the within-host genomic diversity of *E. coli* in healthy

226 children from a rural setting in the Gambia, West Africa. Surprisingly, we recovered a low

rate of colonisation than reported elsewhere among children of similar age groups (42), with

228 only a third of our study samples yielding growth of *E. coli*. This may reflect geographical

variation but might also be some hard-to-identify effect of the way the samples were handled,

even though they were kept frozen and thawed only just before culture.

231 Several studies have shown that sampling a single colony is insufficient to capture *E. coli*

strain diversity in stools (20, 21, 23). Lidin-Janson et al. (43) claim that sampling five

- 233 colonies provides a >99% chance of recovering dominant genotypes from single stool
- specimens, while Schlager et al. (24) calculate that sampling twenty-eight colonies provides a
- 235 >90% chance of recovering minor genotypes. Our results confirm the importance of multiple-

colony picks in faecal surveillance studies, as over half (57%) of our strains would have beenmissed by picking a single colony.

238	Although our strains encompassed all eight major phylotypes of E. coli, the majority fell
239	into the A and B1 phylogenetic groups, in line with previous reports that these phylogroups
240	dominate in stools from people in low- and middle-income countries (44, 45). The prevalence
241	of putative virulence genes in most of our isolates highlights the pathogenic potential of
242	commensal intestinal strains-regardless of their phylogroup-should they gain access to the
243	appropriate tissues, for example, the urinary tract. Our results complement previous studies
244	reporting genomic similarities between faecal E. coli isolates and those recovered from
245	urinary tract infection (25, 46).
246	We found that within-host evolution plays a minor role in generation of diversity, in line
247	with Dixit et al. (20), who reported that 83% of diversity originates from immigration events,
248	and with epidemiological data suggesting that the recurrent immigration events account for
249	the high faecal diversity of <i>E. coli</i> in the tropics (47). Co-colonising variants belonging to the
250	same ST tended to share an identical virulence, AMR and plasmid profile, signalling
251	similarities in their accessory gene content. The estimated mutation rate for E. coli lineages is
252	around one SNP per genome per year (48), so that two genomes with a most recent common
253	ancestor in the last five years would be expected to be around ten SNPs apart. However, in
254	two subjects, pairwise distances between genomes from the same ST (ST59 and ST5148)
255	were large enough (14 and 18 respectively) to suggest that they might have arisen from
256	independent immigration events, as insufficient time had elapsed in the child's life for such
257	divergence to occur within the host. However, it remains possible that the mutation rate was
258	higher than expected in these lineages, although we found no evidence of damage to DNA
259	repair genes. More than half of our isolates encode resistance to three or more classes of
260	antimicrobials echoing the high rate of MDR (65%; confirmed by phenotypic testing) in the

261	GEMS study. IncFIB (AP001918) was the most common plasmid Inc type from our study, in
262	line with the observation that IncF plasmids are frequently associated with the dissemination
263	of resistance (49). However, a limitation of our study is that we did not perform phenotypic
264	antimicrobial resistance testing, although Doyle et al. (50) reported that only a small
265	proportion of genotypic AMR predictions are discordant with phenotypic results.
266	Comparative analyses confirm the heterogeneous origins of the strains reported here,
267	documenting links to other human commensal strains or isolates sourced from livestock or
268	the environment. This is not surprising, as almost all study participants reported that animals
269	are kept in their homes and children in rural Gambia are often left to play on the ground,
270	close to domestic animals such as pets and poultry (51).
271	Our results show that the commensal E. coli population in the gut of healthy children in
272	rural Gambia is richly diverse, with the independent immigration and establishment of strains
273	contributing to the bulk of the observed diversity. In addition, this work has added
274	significantly to the number of commensal E. coli genomes, which are underrepresented in
275	public repositories. Although solely observational, our study paves the way for future studies
276	aimed at a mechanistic understanding of the factors driving the diversification of E. coli in
277	the human gut and what it takes to make a strain of E. coli successful in this habitat.
278	
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- 283 GEMS study in Basse Field Station and Fajara.
- 284

285 **Data summary**

- All genomic assemblies for the strains included in this study are freely available from
- 287 EnteroBase (http://enterobase.warwick.ac.uk/species/index/ecoli). The EnteroBase genome
- assembly barcodes are provided in Supplementary Files 1 and 2.
- 289 The raw genomic sequences have been deposited in the NCBI SRA, under the BioProject
- 290 ID PRJNA658685 and accession numbers SAMN15880286 to SAMN15880281.
- 291

292 **Conflicts of interest**

- 293 We declare no conflicts of interest.
- 294

295 Author contributions

- 296 Conceptualization: MA, MP; data curation, MP, NFA; formal analysis: EFN; analytical
- support: MD; funding: MA and MP; sample collection and storage: MJH, UNI, PET, CO;
- data management: SG; laboratory experiments, EFN, supervision, NFA, MP, JO, MA;
- 299 manuscript preparation original draft, EFN; review and editing, NFA, MP; review of final
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301

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- 466

467 **Tables and figure legends**

Sample ID	Lab ID	Age (months)	Gender	Bristol stool index	Domestic animal within household	Enrolment date
102135	H1	43	Female	Thick liquid	Thick liquid Goat, sheep	
102650	H2	45	Female	Soft	Goat, sheep, donkey	27-Jul-09
103296	Н3	44	Male	Soft	Goat, horse, donkey, rodent	27-Apr-10
103298	H4	44	Male	Formed	Sheep, fowl, horse, donkey, rodent	27-Apr-10
103621	Н5	37	Female	Soft	Sheep, fowl, rodent	01-Sep-10
103650	H6	48	Female	Soft	Fowl, donkey, rodent	29-Sep-10
103649	H7	45	Female	Soft	Goat, sheep, fowl, horse, rodent	29-Sep-10
103071	H8	53	Male	Formed	Goat, sheep, fowl	15-Jan-10
103622	Н9	39	Female	Soft	Goat, sheep	01-Sep-10
100167	H10	40	Female	Soft	Goat, sheep, fowl	01-Feb-08
100217	H11	57	Male	Formed	Cat, fowl, horse, rodent	21-Feb-08
100230	H12	51	Male	Soft	Goat, sheep, cat, fowl, rodent	28-Feb-08
100612	H13	55	Female	Formed	Goat, sheep, dog, fowl, horse, donkey, rodent	16-Aug-08
100162	H14	47	Female	Thick liquid	Sheep, horse, donkey, rodent	30-Jan-08
102255	H15	42	Male	Formed	Goat, sheep, fowl, horse, donkey, rodent	26-Mar-09
102250	H16	39	Male	Formed	Fowl	25-Mar-09
102114	H17	54	Male	Formed		
102123	H18	37	Female	Soft	Goat, sheep, fowl, rodent	
103282	H19	43	Male	Formed	formed Goat, sheep, dog, cat, cow, fowl,	
100817	H20	44	Male	Soft	Soft Dog, fowl	
100816	H21	40	Male	Soft	Goat, sheep, cow, fowl, horse, donkey, rodent	
102836	H22	47	Male	Thick liquid	Fowl, rodent	12-Oct-09
102837	H23	41	Male	Thick liquid	Sheep, fowl, rodent	12-Oct-09
102843	H24	44	Male	Soft	Fowl, rodent	13-Oct-09
102907	H25	36	Male	Soft	Goat, sheep, fowl	05-Nov-09
102905	H26	37	Male	Soft	Goat, sheep, fowl	
102262	H27	38	Male	Formed		
102728	H28	41	Male	Soft	Soft Goat, fowl	
102729	H29	41	Male	Soft Goat, dog, cat, fowl, donkey		24-Aug-09
100806	H30	55	Male	Soft Goat, sheep, dog, fowl 2		21-Nov-08
102053	H31	37	Female	Formed Cow, fowl, donkey, rodent 29-Jan-0		29-Jan-09
102052	H32	38	Female	Formed	Goat, sheep, cow, fowl, donkey, rodent	29-Jan-09
102511	H33	37	Male	Soft	Fowl, horse, donkey, rodent	19-Jun-09

Table 1: Characteristics of the study population

102649	H34	37	Male	Soft	Fowl, horse, donkey, rodent	27-Jul-09
102454	H35	52	Male	Soft	Sheep, fowl, donkey, rodent	02-Jun-09
102459	H36	51	Male	Formed	Goat, sheep, dog, cat, cow, horse, donkey, rodent	04-Jun-09
100303	H37	58	Male	Formed	Sheep, fowl	08-Apr-08
100320	H38	42	Female	Formed	Sheep, fowl, rodent	19-Apr-08
100319	H39	45	Female	Formed	Goat, sheep, fowl, rodent	17-Apr-08
103081	H40	39	Female	Thick liquid	Goat, sheep, fowl, horse, donkey, rodent	20-Jan-10
103082	H41	39	Female	Thick liquid	Goat, sheep, fowl, horse, donkey, rodent	20-Jan-10
100663	H42	36	Male	Thick liquid	Goat, sheep, fowl, donkey	10-Sep-08
100072	H43	51	Female	Formed	Goat, cow, fowl, rodent	03-Jan-08
103171	H44	36	Female	Soft	Goat, sheep, rodent, fowl, rodent	18-Feb-10
103172	H45	36	Female	Soft	Goat, sheep, fowl, rodent	18-Feb-10
103292	H46	39	Male	Soft	Goat, sheep, fowl	23-Apr-10
102952	H47	36	Male	Soft	Goat, sheep, fowl, rodent	20-Nov-09
102953	H48	37	Male	Soft	Goat, sheep, fowl, rodent	20-Nov-09
102964	H49	40	Female	Formed	Goat, fowl, rodent	26-Nov-09
102966	H50	37	Female	Formed	Goat, sheep, fowl, horse, donkey, rodent	22-Apr-10
103281	H51	44	Male	Formed	Goat, sheep, dog, cat, fowl	
100540	H52	43	Male	Soft	Goat, sheep, fowl, rodent	
103123	H53	38	Male	Soft	Sheep	
103124	H54	36	Male	Soft	Fowl	03-Feb-10
102089	H55	38	Female	Soft	Goat, cow, fowl, horse, donkey, rodent	05-Feb-09
103297	H56	38	Male	Soft	Goat, sheep, fowl, horse, donkey, rodent	27-Apr-10
102251	H57	39	Male	Formed	Fowl	25-Mar-09
103602	H58	38	Female	Formed	Goat, sheep, cow, fowl	26-Aug-10
103600	H59	39	Female	Formed	Goat, sheep, fowl	26-Aug-10
100026	H60	49	Female	Soft	Goat, sheep, cow, fowl	
102102	H61	47	Female	Opaque watery	None	11-Feb-09
102263	H62	38	Male	Formed	Horse, donkey, rodent	01-Apr-09
103070	H63	58	Male	Soft	Goat, sheep, fowl	15-Jan-10
103130	H64	40	Male	Soft	Sheep, fowl	03-Feb-10
102051	H65	36	Female	Formed	Goat, sheep, dog, cat, cow, fowl, donkey, rodent	29-Jan-09
102524	H66	36	Male	Soft	Goat, sheep, fowl, horse, donkey, rodent	24-Jun-09

	Genotype number						Migration events	Within-host evolution events
Host	1	2	3	4	5			
H-2	A (9274)	A (9274)	A (9274)	A (9274)	A (9274)	1	1	0
H-9	A (2705)	A (2705)	A (2705)	D (2914)	B1 (29)	3	3	0
H-15	B2 (9277)	B2 (9277)	B2 (9277)	Clade I (747)	Clade I (747)	3	2	1
H-18	D (38)	D (38)	B1 (9281)	A (9274)		4	3	1
H-21	B1 (58)	B1 (58)	B1 (223)	A (540)	D (1204)	4	4	0
H-22	B1 (316)	B1 (316)	B1 (316)	B1 (316)		2	1	1
H-25	A (181)	A (181)	A (181)	A (181)	B1 (337)	4	2	2
H-26	B1 (641)	B1 (2741)	A (10)	A (398)		4	4	0
H-28	B1 (469)	B1 (469)	B1 (469)	B1 (469)		2	1	1
H-32	B1 (101)	B1 (101)	B1 (101)	B1 (2175)	A (10)	3	3	0
H-34	B1 (603)	B1 (603)	B1 (603)	B1 (1727)	A (10)	4	3	1
H-35	A (226)					1	1	0
H-36	F (59)	F (59)	F (59)	F (59)	E (9278)	3	2	1
H-37	D (5148)	D (5148)	D (5148)	D (5148)	D (5148)	3	1	2
H-38	D (394)	D (394)	D (394)	D (394)	B1 (58)	4	2	2
H-39	B2 (452)	B2 (452)	B2 (452)	B2 (452)	B2 (452)	2	1	1
H-40	B1 (155)					1	1	0
H-41	A (43)	A (43)	A (43)	A (43)	B1 (9283)	2	2	0
H-48	Clade I (485)	Clade I (485)	Clade I (485)	Clade I (485)		1	1	0
H-50	C (410)	C (410)	C (410)	C (410)	B1 (515)	2	2	0
H-55	A (9279)					1	1	0

Table 2: Phylogroup and sequence types of the distinct clones isolated in each patient

Host	Sequence type (ST)	Colonies per ST	Pairwise SNP distances between multiple colonies of the same ST
H2	9274	5	0-9
H9	2705	3	0-1
H15	9277	3	0-1
H15	747	2	3
H18	38	2	3
H21	58	2	0
H22	316	4	0-3
H25	181	4	1-5
H28	469	4	0-3
H32	101	3	1-9
H34	603	3	2-8
H36	59	4	0-14
H37	5148	5	2-18
H38	394	4	1-3
H39	452	5	0-2
H41	43	4	0-1
H48	485	4	1-9
H50	410	4	0

Table 3: Pairwise SNP distances between variants arising from within-host evolution

Sample ID	7-gene ST	Neighbour host	Neighbour status	Neighbour's country of isolation	Allelic distance
H-32_5	10	Human	Unknown	UK	18
H-36_1	59	Human	Unknown	UK	18
H-39_1	452	Human	Commensal	UK	26
H-9_1	2705	Livestock		China	29
H-18_3	9274	Human	Commensal	Unknown	34
H-2_1	9274	Human	Commensal	Unknown	34
H-22_1	316	Human	Commensal	UK	35
H-38_1	394	Human	Pathogen (cystitis)	US	39
H-25_4	337	Human	Unknown	Mali	43
H-37_1	5148	Human	Pathogen (diarrhoea)	Ecuador	43
H-26_1	641	Livestock		US	46
H-26_5	398	Poultry		Kenya	47
H-48_2	485	Human	Commensal	Tanzania	57
H-15_1	9277	Human	Commensal	Zambia	68
H-15_2	747	Human	Commensal	Egypt	72
H-28_1	469	Human	Commensal	Kenya	77
H-21_2	1204	Avian		Kenya	81
H-34_2	10	Livestock		UK	83
H-38_2	58	Human	Pathogen (bloodstream infection)	Australia	87
H-34_4	1727	Unknown	Unknown	Unknown	89
H-35_1	226	Human	Commensal	China	93
H-21_1	58	Unknown	Unknown	Unknown	98
H-21_4	540	Human	Unknown	Belgium	100
H-32_2	2175	Livestock		UK	100
H-26_2	10	Livestock		US	111
H-32_1	101	Unknown	Unknown	Unknown	111
H-50_2	515	Environment		Canada	117
H-41_1	43	Unknown	Unknown	Unknown	120
H-26_4	2741	Human	Commensal	Germany	126
H-50_1	410	Livestock		US	140
H-18_1	38	Poultry		US	144
H-21_5	223	Unknown	Unknown	Unknown	145
H-40_1	155	Unknown	Unknown	US	146
– H-41_2	9283	Environment	Commensal	US	191
	9278	Avian		Kenya	208
H-9_3	2914	Canine		UK	272
H-9_5	29	Unknown	Unknown	Unknown	288
H-34_1	603	Laboratory		UK	325
H-55_1	9279	Environment		Unknown	333
H-18_2	9281	Unknown	Unknown	France	430
H-15_2 H-25_1	181	Human	Commensal	Tanzania	607

Table 4: Closest relatives to the study isolates

Legends to figures

Figure 2

A maximum-likelihood tree depicting the phylogenetic relationships among the study isolates. The tree was reconstructed with RAxML, using a general time-reversible nucleotide substitution model and 1,000 bootstrap replicates. The genome assembly of *E. coli* str. K12 substr. MG1655 was used s as the reference, and the tree rooted using the genomic assembly of *E. fergusonii* as an outgroup. The sample names are indicated at the tip, with the respective Achtman sequence types (ST) indicated beside the sample names. The respective phylogroups the isolates belong to are indicated with colour codes as displayed in the legend. *E. coli* reference genome is denoted in black. Asterisks (*) are used to indicate novel STs. The predicted antimicrobial resistance genes and putative virulence factors for each isolate are displayed next to the tree, with the virulence genes clustered according to their function. Multiple copies of the same strain (ST) isolated from a single host are not shown. Instead, we have shown only one representative isolate from each strain. Virulence and resistance factors were not detected in the reference strain either. We have provided a summary of the identified virulence factors and their known functions in Supplementary File 3.

Figure 3

A: The prevalence of antimicrobial-associated genes detected in the isolates. The y-axis shows the detected AMR-associated genes in the genomes, grouped by antimicrobial class.B: A histogram depicting the number of antimicrobial classes to which resistance genes were detected in the corresponding strains.

Figure 4

A: Plasmid replicons detected in the study isolates. B: A histogram depicting the number of plasmids co-harboured in a single strain.

Figure 5

A: A NINJA neighbour-joining tree showing the population structure of *E. coli* ST38, drawn using the genomes found in the core-genome MLST hierarchical cluster at HC1100, which corresponds to ST38 clonal complex. B: The closest neighbour to a pathogenic strain reported in GEMS ⁴ is shown to be a commensal isolate recovered from a healthy individual. C: The closest relatives to the commensal ST38 strain recovered from this study is shown (red highlights), with the number of core-genome MLST alleles separating the two genomes displayed. D: A maximum-likelihood phylogenetic tree reconstructed using the genomes found in the cluster in C above, comprising both pathogenic and commensal ST38 strains is presented, depicting the genetic relationship between strain 100415 (pathogenic) and 103709 (commensal) (red highlights). The nodes re coloured to depict the status of the strains as pathogenic (red) or commensal (blue). The geographical locations where isolates were recovered are displayed in Figures 4A-C; the genome counts shown in square brackets.

Supplementary material

Supplementary Figure 1

A co-occurrence matrix of acquired antimicrobial resistance genes detected in the study isolates. The diagonal values show how many isolates each individual gene was found in, while the intersections between the columns represent the number of isolates in which the corresponding antimicrobial resistance genes co-occurred.

Supplementary Figure 2

A Neighbour-joining phylogenetic tree depicting the genetic relationships among twenty-four strains isolated from diarrhoeal cases in the GEMS study ⁴. The Sequence types identified in

these isolates are shown in the legend, with the genome count displayed in square brackets next to the respective sequence types. Three STs (ST38, ST58 and ST10) overlapped with what was found among commensal strains from this study (see Figure 2).

Supplementary File 1

Sequencing statistics and characteristics of twenty-four previously sequenced GEMS cases included in this study.

Supplementary File 2

A summary of the sequencing statistics of the study isolates reported in this study.

Supplementary File 3

A summary of the virulence factors detected among the study isolates and their known functions.

Supplementary File 4

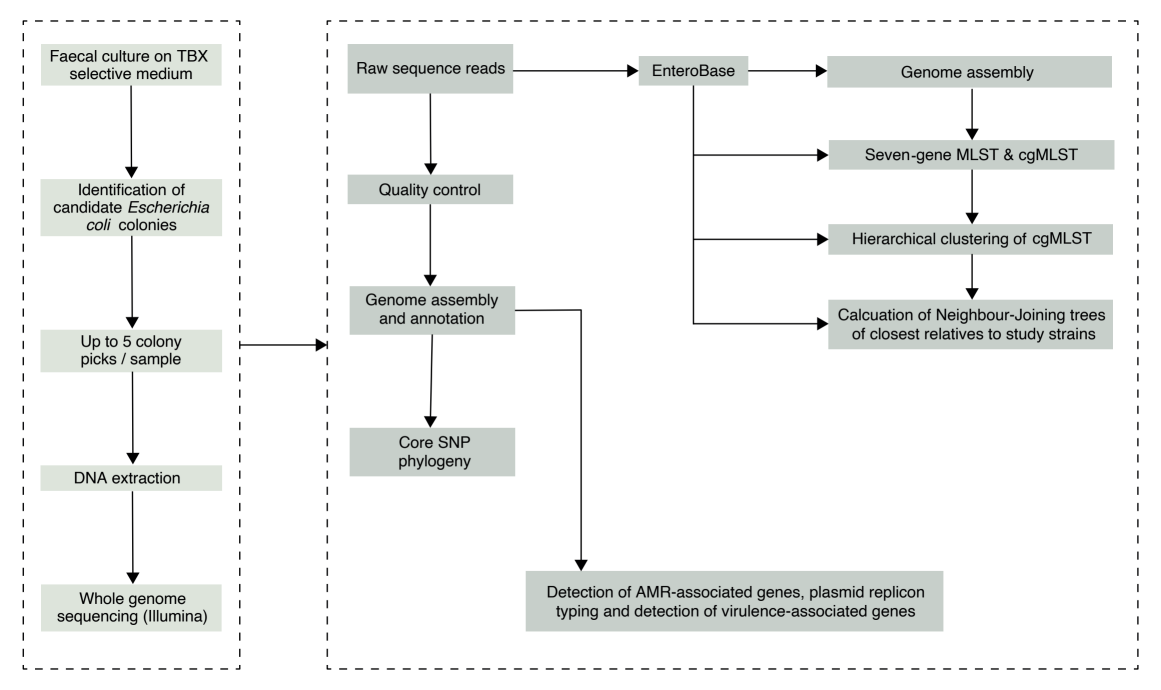
A pairwise single nucleotide polymorphism matrix showing the SNP distances between the study genomes.

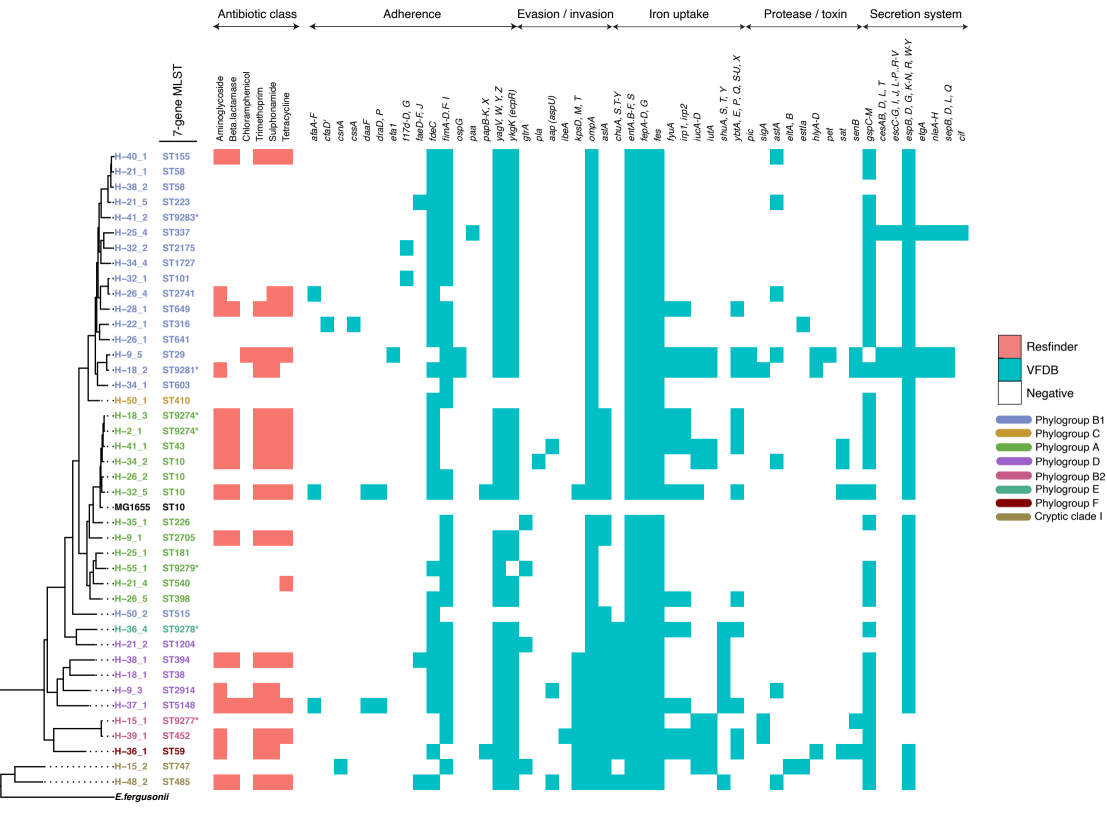
Supplementary File 5

Two independent cultures of the same clone of samples that were sequenced twice and used to find the SNPs between same clones.

Supplementary File 6

Mutations in variants inferred to have been derived from within-host evolution.





A

