

# Genome-wide analysis in *Escherichia coli* unravels a high level of genetic homoplasmy associated with cefotaxime resistance

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## Abstract

Cefotaxime (CTX) is a third-generation cephalosporin (3GC) commonly used to treat infections caused by *Escherichia coli*. Two genetic mechanisms have been associated with 3GC resistance in *E. coli*. The first is the conjugative transfer of a plasmid harbouring antibiotic-resistance genes. The second is the introduction of mutations in the promoter region of the *ampC*  $\beta$ -lactamase gene that cause chromosome-encoded  $\beta$ -lactamase hyperproduction. A wide variety of promoter mutations related to AmpC hyperproduction have been described. However, their link to CTX resistance has not been reported. We recultured 172 cefoxitin-resistant *E. coli* isolates with known CTX minimum inhibitory concentrations and performed genome-wide analysis of homoplastic mutations associated with CTX resistance by comparing Illumina whole-genome sequencing data of all isolates to a PacBio sequenced reference chromosome. We mapped the mutations on the reference chromosome and determined their occurrence in the phylogeny, revealing extreme homoplasmy at the -42 position of the *ampC* promoter. The 24 occurrences of a T at the -42 position rather than the wild-type C, resulted from 18 independent C>T mutations in five phylogroups. The -42 C>T mutation was only observed in *E. coli* lacking a plasmid-encoded *ampC* gene. The association of the -42 C>T mutation with CTX resistance was confirmed to be significant (false discovery rate <0.05). To conclude, genome-wide analysis of homoplasmy in combination with CTX resistance identifies the -42 C>T mutation of the *ampC* promoter as significantly associated with CTX resistance and underlines the role of recurrent mutations in the spread of antibiotic resistance.

## DATA SUMMARY

All data is available from the National Center for Biotechnology Information (NCBI) under BioProject number PRJNA592140. Raw Illumina sequencing data and metadata for all 171 *Escherichia coli* isolates used in this study is available from the NCBI Sequence Read Archive database under accession numbers SAMN15052485 to SAMN15052655. The full reference chromosome of *ampC\_0069* is available via GenBank accession number CP046396.1 and NCBI Reference Sequence NZ\_CP046396.1. The

scripts used to calculate homoplasmy-based association analysis are available from GitHub (<https://github.com/JordyCoolen/hombaampC>) under MIT license.

## INTRODUCTION

*Escherichia coli* is an important pathogen in both community and healthcare-associated infections [1, 2]. In the past decades, a substantial increase in resistance to third-generation cephalosporin

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**Abbreviations:** *campC*, chromosome-mediated *ampC*; CTX, cefotaxime; ESBL, extended-spectrum  $\beta$ -lactamase; EUCAST, European Committee on Antimicrobial Susceptibility Testing; FDR, false discovery rate; FOX, cefoxitin; 3GC, third-generation cephalosporin; MIC, minimum inhibitory concentration; MLST, multilocus sequence typing; NCBI, National Center for Biotechnology Information; *pampC*, plasmid-mediated *ampC*; qRT-PCR, quantitative reverse-transcriptase PCR; SMRT, single-molecule real-time; ST, sequence type; VCF, variant calling file; WGS, whole-genome sequencing.

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**Data statement:** All supporting data, code and protocols have been provided within the article or through supplementary data files. Four supplementary tables and three supplementary figures are available with the online version of this article.

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(3GC) antibiotics in *E. coli* has been observed worldwide, mainly caused by the production of extended-spectrum  $\beta$ -lactamases (ESBLs) and AmpC  $\beta$ -lactamases, restricting available treatment options for common infections [3]. AmpC  $\beta$ -lactamases differ from ESBL as they hydrolyse not only broad-spectrum penicillins and cephalosporins, but also cephamycins. Moreover, AmpC  $\beta$ -lactamases are not inhibited by ESBL-inhibitors like clavulanic acid [3], limiting antibiotic treatment options even further. A widely used screening method for AmpC production is the use of susceptibility to ceftiofloxacin (FOX), a member of the cephamycins [4].

Although *ampC*  $\beta$ -lactamase genes can be plasmid-encoded (plasmid-mediated *ampC*, *pampC*), they are also encoded on the chromosomes of numerous *Enterobacteriales*. *E. coli* naturally carries a chromosome-mediated *ampC* (*campC*) gene but, unlike most other *Enterobacteriales*, this gene is non-inducible due to the absence of the *ampR* regulator gene [3]. Chromosomal AmpC production in *E. coli* is exclusively regulated by promoter and attenuator mechanisms. This results in constitutive low-level *campC* expression that still allows the use of 3GC antibiotics, such as cefotaxime (CTX), to treat *E. coli* infections [3]. However, various mutations in the promoter/attenuator region of *E. coli* may cause constitutive hyperexpression of *campC* [5, 6], thereby increasing the minimum inhibitory concentrations (MICs) for broad-spectrum penicillins and cephalosporins and limiting appropriate treatment options.

A wide variety of promoter and attenuator mutations have been related to AmpC hyperproduction [6]. AmpC hyperproduction is primarily caused by alterations of the *ampC* promoter region, leading to a promoter sequence that more closely resembles the *E. coli* consensus  $\sigma^{70}$  promoter with a TTGACA  $-35$  box separated by 17bp from a TATAAT  $-10$  box. These alterations can be divided into different variants associated with, for example, an alternate displaced promoter box, a promoter box mutation or an alternate spacer length due to insertions [6]. Furthermore, mutations of the attenuator sequence can lead to changes in the hairpin structure that strengthen the effect of promoter alterations on AmpC hyperproduction. In the study by Tracz *et al.* on FOX-resistant *E. coli* isolated from Canadian hospitals, 52 variants of the promoter and attenuator region were described [6]. Tracz *et al.* used a two-step quantitative reverse-transcriptase PCR (qRT-PCR) to determine the effect of promoter/attenuator variants on *ampC* expression. Various mutations were related to different delta–delta cycle threshold values in the qRT-PCR and corresponding variations in FOX resistance. An interesting observation that emerged from this study was that the  $-32T>A$  and the  $-42C>T$  mutation were the major alterations that strengthened the *ampC* promoter. Both result in a consensus  $-35$  box. Although it is known that AmpC hyperproduction leads to FOX resistance, as studied by Tracz *et al.*, the effects of various mutations on resistance to a 3GC antibiotic such as CTX have not been explored. This is relevant because CTX is commonly used in the treatment of patients with severe *E. coli* infections, often in combination with selective digestive tract decontamination in intensive care units [7, 8].

### Impact Statement

In the past decades, the worldwide spread of extended-spectrum  $\beta$ -lactamases (ESBLs) has led to a substantial increase in the prevalence of resistant common pathogens, thereby restricting available treatment options. Although acquired resistance genes, e.g. ESBLs, get most attention, chromosome-encoded resistance mechanisms may play an important role as well. In *Escherichia coli*, chromosome-encoded  $\beta$ -lactam resistance can be caused by alterations in the promoter region of the *ampC* gene. To improve our understanding of how frequently these alterations occur, a comprehensive interpretation of the evolution of these mutations is essential. In the current study, we apply genome-wide homoplasmy analysis to better perceive adaptation of the *E. coli* genome to antibiotics. Thereby, this study grants insights into how chromosome-encoded antibiotic resistance evolves and, by combining genome-wide association studies with homoplasmy analyses, provides potential strategies for future association studies into the causes of antibiotics resistance.

While previous research mainly focused on the chromosomal AmpC resistance mechanism and the impact of AmpC hyperproduction, there is a lack in knowledge and understanding of the evolutionary origin of these promoter/attenuator variants. Notably, it is unexplored how the two most prominent promoter mutations,  $-32T>A$  and  $-42C>T$ , are distributed over the *E. coli* phylogeny and therewith how often they occur. More precisely, literature shows selective pressure can lead to convergent evolution that results in the reoccurrence of a mutation in multiple isolates independently and in separate lineages [9]. This phenomenon is named homoplasmy [10]. A consistency index can be calculated to quantify homoplasmy by dividing the minimum number of changes on the phylogeny by the number of different nucleotides observed at that site minus one [11], effectively quantifying how often the same mutation occurred in a phylogenetic tree. One can use the consistency index to recognize genomic locations subjected to homoplasmy, and relate the SNP positions that are inconsistent with the phylogeny to antibiotic resistance, as has been done, for example, in multiple studies on *Mycobacteria* spp. [12–15].

In the present study, we hypothesize that some of the mutations in the *ampC* promoter/attenuator region are homoplastic and are associated with CTX resistance. To test our hypothesis, we performed genome-wide homoplasmy analysis and combined it with a genome-wide analysis of polymorphisms associated with CTX resistance by constructing an *E. coli* reference chromosome and combining it with whole-genome sequencing (WGS) data of 172 both FOX resistant and ESBL-negative *E. coli* isolates from human and animal origin previously collected by our research group [16].

## METHODS

### Isolate selection, DNA isolation, library preparation and DNA sequencing

One hundred and seventy-two both FOX resistant (MIC >8 mg l<sup>-1</sup>) and ESBL-negative *E. coli* isolates previously used by our study group [16] were selected in the present study (Table S1, available with the online version of this article). To summarize the methods, DNA isolation was performed as previously described [16], library preparations were performed using an Illumina Nextera XT library preparation kit, and DNA sequencing was performed using an Illumina NextSeq 500 to generate 2×150 bp paired-end reads or 2×300 bp reads on an Illumina MiSeq. *De novo* assembly was also performed identically to the method as described previously [16] using SPAdes version 3.11.1 [17].

### Phylogroup and multilocus sequence typing (MLST)

Phylogroup stratification was performed using ClermonTyping version 1.4.0 [18]. MLST sequence types (STs) were derived from the contigs using Mlst version 2.5 PubMLST (31 October 2017) [19, 20].

### Obtaining the *ampC* promoter/attenuator region

To detect the promoter/attenuator region, a custom BLAST database [21] was created using the 271 bp fragment as described by Peter-Getzlaff *et al.* [22] using *E. coli* K-12 strain ER3413 (accession no. CP009789.1). ABRicate version 0.8.9 [23] was used to locate matching regions per sample and these were extracted and converted into multi-FASTA format using a custom Python script. Strains were labelled AmpC putative hyperproducer when promoter mutations were found, as previously identified by Caroff *et al.* [24], Siu *et al.* [25] and Tracz *et al.* [6].

### *pampC* detection

Detection of *pampC* genes was performed by using ABRicate version 0.5 [23] and ResFinder database (16/02/2018) as described by Coolen *et al.* [16].

### PacBio single-molecule real-time (SMRT) sequencing of an *E. coli* isolate

To enable an accurate SNP analysis, a reference chromosome of an *E. coli* isolate from our collection (*ampC*\_0069) was constructed using PacBio SMRT sequencing. For sequencing, genomic DNA (gDNA) was extracted using a bacterial gDNA isolation kit (Norgen Biotek). A single *E. coli* isolate was subjected to DNA shearing using Covaris g-TUBEs for 30 s at 11000 r.p.m. Each DNA sample was separated into two aliquots. Size selection was performed using a 0.75% agarose cassette and marker S1 on the BluePippin system (Sage Science) to obtain either 4–8 kb or 4–12 kb DNA fragments. This size selection was chosen to maintain all DNA fragments, including those originating from plasmids (data not used in this study). Library preparation was performed using the PacBio SMRTbell template prep kit 1.0 (Pacific Biosciences). For cost-effectiveness, samples were barcoded

and pooled with other samples that are not relevant for this study. Sequencing was conducted using the PacBio Sequel I (Pacific Biosciences) on a Sequel SMRT Cell 1M v2 (Pacific Biosciences) with a movie time of 10 h (and 186 min pre-extension time). Subreads per sample were obtained by extracting the BAM files using SMRT Link version 5.1.0.26412 (Pacific Biosciences).

### Chromosomal reconstruction using *de novo* hybrid assembly

To obtain a full-length chromosome, Unicycler version 0.4.7 [26] (settings: --mode bold) was used, combining Illumina NextSeq 500 2×150 bp paired-end reads with PacBio Sequel SMRT subreads. Because Unicycler requires FASTA reads as input, the subreads in BAM format were converted to FASTA by using bam2fasta version 1.1.1 from pbbioconda (<https://github.com/PacificBiosciences/pbbioconda>) prior to *de novo* hybrid assembly. The full circular chromosome was uploaded to the National Center for Biotechnology Information (NCBI) database and annotated using NCBI Prokaryotic Genome Annotation Pipeline (PGAP) version 4.10 [27, 28].

### SNP analysis using *E. coli* reference *ampC*\_0069

Alignment of Illumina reads and SNP calling was performed for all isolates to the reference chromosome of *E. coli* isolate *ampC*\_0069 using Snippy version 4.3.6 (<https://github.com/tseemann/snippy>). A full-length alignment (fullSNP) and a coreSNP alignment containing SNP positions shared among all isolates were generated by using snippy-core version 4.3.6 (<https://github.com/tseemann/snippy>).

### Inferring of phylogeny

A phylogenetic tree was inferred by using the coreSNP alignment as input for FastTree(MP) version 2.1.3 SSE3 (settings: -nt -gtr) [29].

### Detection of homoplasy

The consistency index for all nucleotide positions on the chromosome was calculated using HomoplasyFinder version 0.0.0.9000 [10]. The coreSNP phylogeny was used as true phylogeny and the consistency index was calculated using the multiple sequence alignments fullSNP alignment as input.

### Relating mutations to CTX resistance

To assess whether certain mutations were linked to CTX resistance, all non-plasmid-harboursing *ampC* *E. coli* isolates were used. CTX resistance was defined using European Committee on Antimicrobial Susceptibility Testing (EUCAST) guideline standards of CTX MIC >2 mg l<sup>-1</sup> [30]. CTX MIC results were obtained from our previous study [16]. For each nucleotide position on the reference chromosome, the numbers of resistant and sensitive isolates were counted and tested for adenine versus all other nucleotides, thymine versus all other nucleotides, cytosine versus all other nucleotides, and guanine versus all other nucleotides, creating a contingency table and performing a Fisher's exact test in R 3.6.1 [31]. To correct for

multiple testing, *P* values were adjusted using false discovery rate (FDR) [32].

### Selection of genomic positions of interest

Genomic positions with potential roles in CTX resistance were identified based on FDR  $\leq 0.05$  for CTX and a consistency index of  $\leq 0.05882353$ . Annotation of mutation positions was obtained by using the genome annotation of the reference chromosome (GenBank accession no. CP046396) and applying snpEff (version 4.3 t) [33]. The EnteroBase core-genome MLST and whole-genome MLST schemes were used to distinguish core and accessory genes [34].

### Recombination analysis

Gubbins version 2.4.1. (settings: -f 30) was used to detect recombination regions with coreSNP alignment and tree as input [35].

### Pyseer analysis

To compare our homoplasy-associated analysis method to an alternative method, we used Pyseer (version 1.3.6) [36]. In short, variant calling files (VCFs) were obtained from snippy, and bcftools (version 1.11) was used to merge and filter the VCFs from all samples to a single VCF [37]. A phylogenetic distance file was calculated by using the phylogeny\_distance.py included in Pyseer on the corrected for recombination phylogeny of Gubbins. Finally, the distance, trait and VCF file was used to run Pyseer (default fixed effects) with settings --min-af 0.01, --max-af 0.99.

### Visualization of data

The interactive tree of life web-based tool (iTOL) version 5.3 was used to visualize the phylogenetic tree [38]. Information about CTX resistance, presence of the *pampC* gene, *campC* hyperproduction as defined, MLST and phylogroup, as well as alignments of promoter and attenuator region, were incorporated into the visualization. The sequence logo of the promoter and the attenuator alignment were generated using the web-based application WebLogo, version 3.7 [39] (<http://weblogo.threeplusone.com>). A chromosome ideogram of the *E. coli* isolate ampC\_0069 reference chromosome was visualized using the CIRCOS software package, version 0.69-8 [40]. Consistency index scores and significant mutations associated with CTX resistance were plotted in the ideogram. Gubbins results were displayed by using Phandango [41].

### Overview of the method

A workflow graph of the method is visualized in Fig. 1, using the web-based application yEd Live version 4.4.2 (<https://www.yworks.com/yed-live/>).

## RESULTS

### *E. coli* collection

To study genetic homoplasy events in suspected AmpC-producing *E. coli*, FOX-resistant ESBL phenotype negative

*E. coli* isolates ( $n=172$ ) were selected, as previously described by Coolen et al. [16] (see Table S1). The entire collection was subjected to WGS, followed by *de novo* assembly of the sequence reads to obtain contigs.

### MLST and phylogroup variants

To access the genetic diversity of our *E. coli* collection, we identified both MLST and phylogroup variants of each of the 172 *E. coli* isolates. A total of 75 different STs were identified, of which ST131 (8.1%,  $n=14$ ), ST38 (7.0%,  $n=12$ ) and ST73 (7.0%,  $n=12$ ) were the most prevalent. The STs of 13 isolates are unknown. Phylogroup stratification revealed that the isolates belonged to eight different phylogroups (Table 1). Phylogroup B1, B2 and D were the most prevalent. One isolate belonged to *E. coli* clade IV (strain no. ampC\_0128).

### *ampC* promoter and attenuator variants

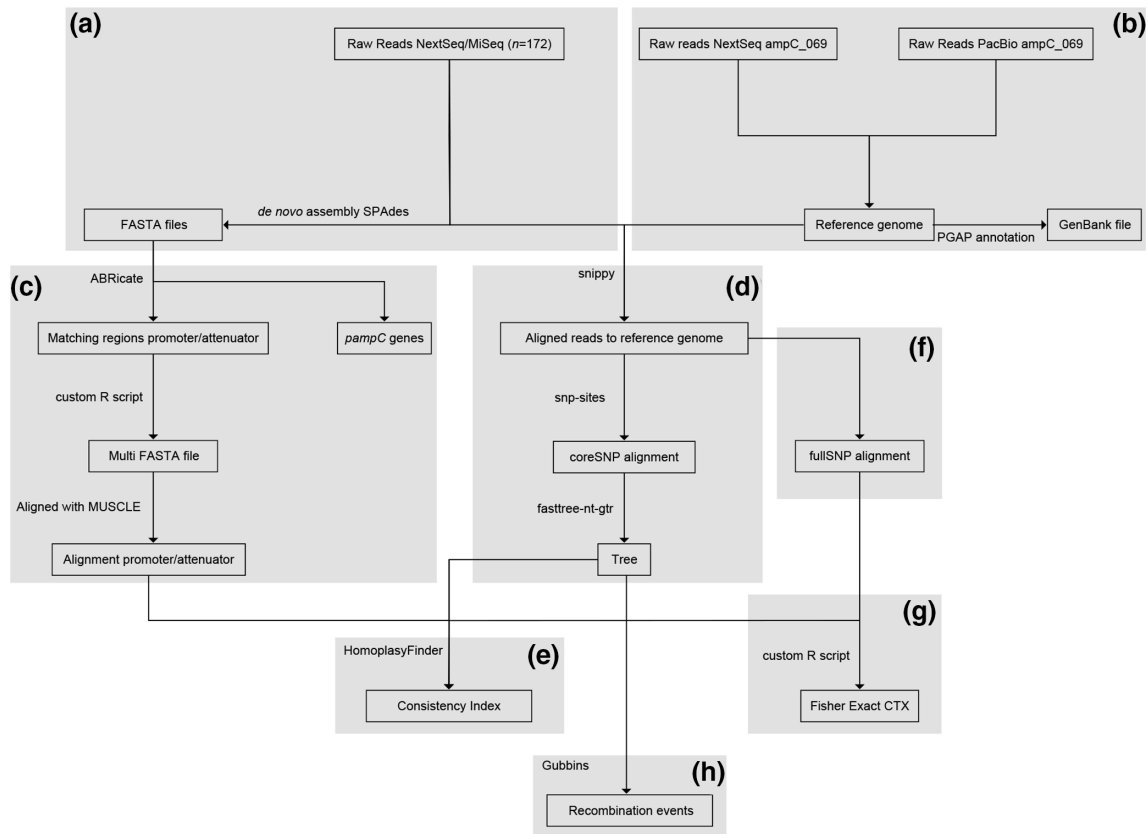
We examined the whole *E. coli* genome. However, we firstly focused on mutations in the *ampC* promoter and attenuator region. Previously described mutations in the *ampC* promoter region that according to described literature lead to 'hyperproduction' of AmpC were detected in 61 (35.5%) of the isolates [6, 24, 25] (see Tables S1 and S2). These isolates were, therefore, labelled as putative hyperproducers. Analysis of the promoter area (-42 to -8) resulted in 20 different variants and the wild-type (see Table 1). In the attenuator region (+17 to +37), 18 different variants were identified (see Table 1). One isolate (ampC\_0128) showed an unusual promoter variant, a four nucleotide deletion (-45\_-42delATCC). Moreover, an insertion (21\_22insG) of unknown function was detected in the attenuator of this isolate (ampC\_0128) as well (see Tables S1 and S2).

### *pampC* variants

As we aimed to associate chromosomal mutations with CTX resistance, differentiation of *pampC*-harbouring isolates from non-*pampC*-harbouring isolates was required. Genomic analysis showed that 90 (52.3%) of the isolates harboured a *pampC* gene of which *bla*<sub>CMY-2</sub> was the most prevalent ( $n=78$ ). One isolate harboured two different *pampC* genes (*bla*<sub>CMY-4</sub> and *bla*<sub>DHA-1</sub>) (ampC\_0119). One isolate contained a *bla*<sub>CTX-M-27</sub> gene combined with a *bla*<sub>CMY-2</sub> gene (ampC\_0114), but was ESBL disc test negative (see Table S1). In 21 (12.2%) of the isolates, neither *pampC* nor described mutations related to AmpC hyperproduction were detected and are noted as putative low-level AmpC producers.

### Reference chromosome

To be able to reconstruct an accurate phylogeny, we selected *E. coli* isolate ampC\_0069, one of the strains of the study, to use as self-constructed reference chromosome for SNP calling. The reference chromosome was constructed through a hybrid assembly of  $n=4423109 \times 150$  bp Illumina NextSeq 500 paired-end reads together with  $n=218475$  PacBio Sequel SMRT subreads (median 5640 bp). This resulted in a high-quality full circular chromosome of *E. coli* isolate ampC\_0069, with a size of 5056572 bp. This isolate belongs to ST648 and



**Fig. 1.** Schematic of the workflow used to perform the homoplasmy-based association analysis. Starting from the top, (a) the *de novo* assembly of the NextSeq/MiSeq reads and (b) the hybrid assembly of the reference chromosome ampC\_069. On the left side, (c) the alignment of promoter/attenuator region. In the middle, (d) the coreSNP analysis for the phylogeny used in (e) the homoplasmy analysis combined with (f) the fullSNP data, on the right, which was also used for (g) the statistics (Fisher's exact test and FDR) to relate CTX resistance to SNP positions. (h) Inferring recombination events using Gubbins.

contains a plasmid-encoded *bla*<sub>CMY-42</sub>. The full circular chromosome was uploaded to GenBank under accession number CP046396 and was used for further analysis. Genome annotation with the NCBI Prokaryotic Genome Annotation Pipeline (PGAP) identified 4720 coding sequences.

### SNP calling

The mapping of the reads of all isolates to the reference chromosome *E. coli* ampC\_0069 (accession no. CP046396) resulted in a coreSNP alignment containing 314200 variable core SNP positions. For further details per isolate see Table S3. To validate our SNP calling method, we compared the ampC\_0069 Illumina NextSeq 500 paired-end reads to the reference chromosome of ampC\_0069, resulting in 0 SNPs detected, supporting that the SNP calling data and method produce no false positives.

### Phylogenetic tree based on coreSNP

The coreSNP alignment was used for further analysis. Fig. 2 illustrates the approximately maximum-likelihood phylogenetic tree of all 172 isolates based on the coreSNP alignment. The tree has a robust topology as indicated by the

SH-like (Shimodaira–Hasegawa like) branch support, only three positions have a value  $\leq 60\%$  [29]. When focusing on the *ampC* promoter mutations, they were most prevalent in phylogroups B1, B2 and C, although they were present in all phylogroups except phylogroup E that lacked mutations in either the promoter or attenuator region. Interestingly, two positions previously highlighted by Tracz *et al.*,  $-42$  and  $-32$ , are only mutated in the absence of a *pampC* gene, even in isolates with a similar MLST ST (ST12, ST88 and ST131). The  $-42C>T$  mutation, which results in an alternate displaced promoter box and, therefore, leads to increased resistance [6], is present in 24 isolates in five distinct phylogroups and in 17 separate phylogenetic branches, indicating that this mutation is homoplastic. Additionally, the  $-32T>A$  mutation in the promoter, previously also associated with resistance [6], is present in 20 isolates in three distinct phylogroups and in 14 separate phylogenetic branches. To quantify the level of homoplasmy, we calculated the consistency index.

### Genomic homoplastic mutations

We calculated the consistency index for all SNP positions on the *E. coli* reference chromosome. A low consistency

**Table 1.** Table of the distribution of the different AmpC promoter and attenuator variants, as well as the numbers of different MLST STs and phylogroups per grouped genotype (*pampC*, putative hyperproducers and putative low-level AmpC producers)

Genotype	Isolate	Promoter variant	Attenuator variant	MLST	Phylogroup
<i>pampC</i>	<i>n</i> =90	3 variants	6 variants	44 STs and 4 unknown	
Putative hyperproducers	<i>n</i> =61	13 variants	14 variants	30 STs and 5 unknown	
Putative low-level AmpC producers	<i>n</i> =21	8 variants	5 variants	14 STs and 4 unknown	
Total	<i>n</i> =172	21 variants	18 variants	75 STs and 13 unknown	

phylogroup ■ A ■ B1 ■ B2 ■ C ■ clade IV ■ F

index value for a position indicates a high degree of inconsistency with the chromosomal phylogeny and can be calculated by HomoplasyFinder as described in earlier studies [10, 42, 43]. As can be observed in Fig. 3, results clearly indicate position -42 (4470140) and -32 (4470150) are the lowest scoring positions on the consistency index concerning the promoter and attenuator region, respectively, 0.05882353 and 0.07142857 (see also Fig. S1). To access how extreme these values are, we calculated the consistency index for all SNP positions in the chromosome (see Fig. S2). All consistency indexes <1.0 are plotted in the outer ring (ring A) of Fig. 4. Results show that only 9640 out of 5056572 positions (0.19%) had a consistency index  $\leq 0.07142857$  (see Fig. 4, ring a, cut-off is indicated by a black circle). This clearly indicates that positions with low consistency indexes are rare, but not unique. Although these 9640 positions have a low consistency index, we do not yet know their relation to CTX resistance.

### CTX-resistance measurements

CTX MIC measurements from Coolen *et al.* [16] in relation to the genotype of the *E. coli* isolates are shown in Table S1. Eighty-four of ninety (93.3%) *pampC*-harbouring *E. coli* were CTX resistant (MIC >2 mg l<sup>-1</sup>) based on EUCAST clinical breakpoints. Twenty-two of sixty-one (36.1%) isolates categorized as putative hyperproducers based on Tracz *et al.* were CTX resistant, primarily isolates with the -42 (*n*=15,

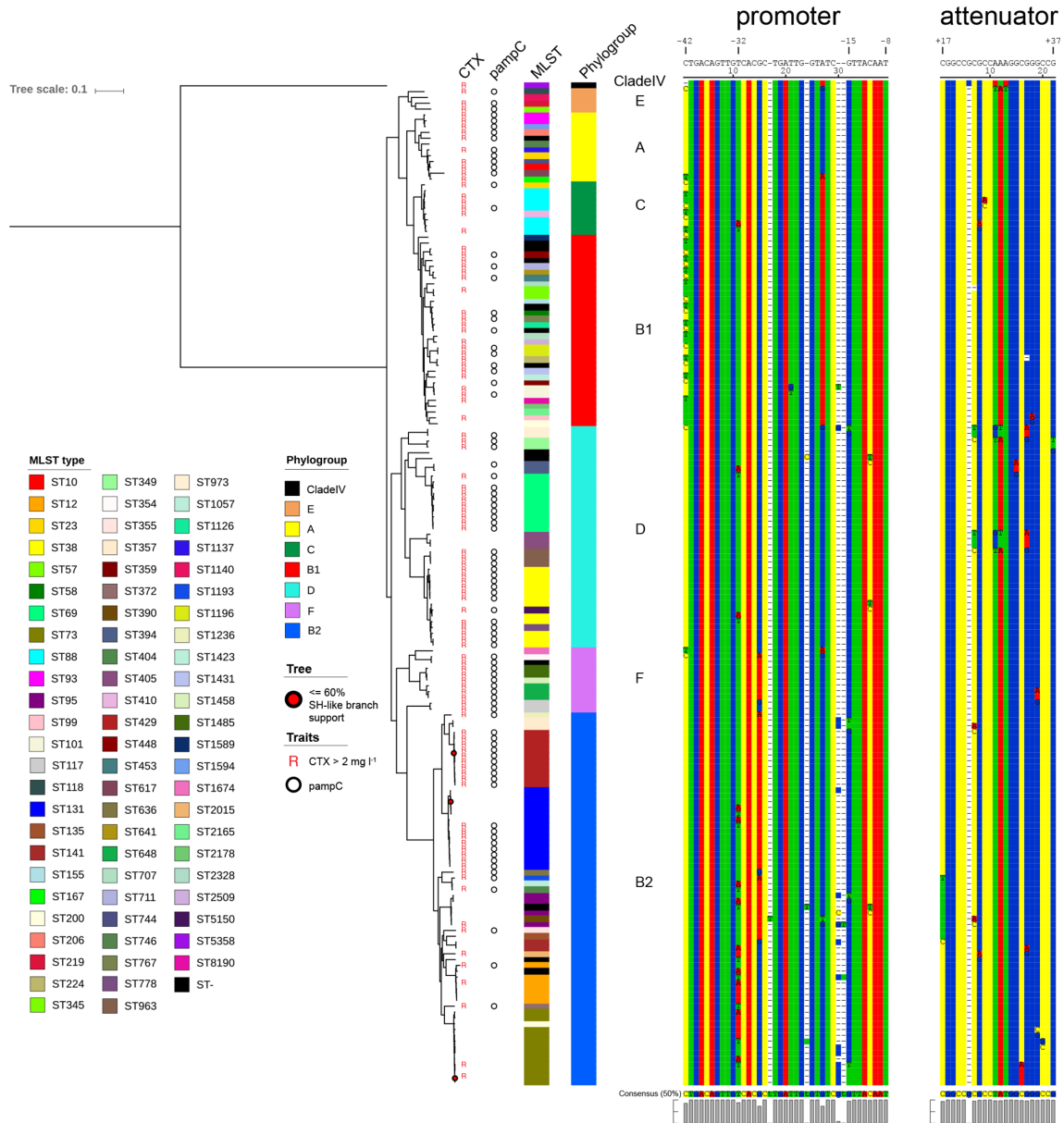
62.5% of isolates with -42 mutation) or -32 mutation (*n*=2, 10.0% of isolates with -32 mutation). The *pampC* genes never occurred simultaneously with the -42 or -32 mutations in any of these isolates. As depicted in Fig. 2, the non-*pampC* strains with a phenotype of CTX >2 mg l<sup>-1</sup> were present in all phylogroups, although CTX-resistant isolates with the -42 or -32 mutation were predominantly present in phylogroups B1, B2 and C.

### Genotype to phenotype

To be able to link *E. coli* chromosomal mutations to CTX resistance, we excluded all *E. coli* isolates with a plasmid containing an *ampC*  $\beta$ -lactamase gene. The association of SNPs with the CTX-resistance phenotype (MIC >2 mg l<sup>-1</sup>) was tested in the remaining 82 isolates using Fisher's exact test. After FDR correction to 0.05, 45998 significant positions were found (see Fig. 4, ring b). Mutation C>T on position -42 of the *ampC* promoter was found to be significantly associated with CTX resistance (FDR=0.034). However, position -32 A>T was not significantly associated with CTX resistance (FDR=1).

### Homoplasy-based association analysis

Combining the outcome of the homoplasy analysis with the significant CTX-resistance-associated positions results in genomic positions associated with CTX resistance that have



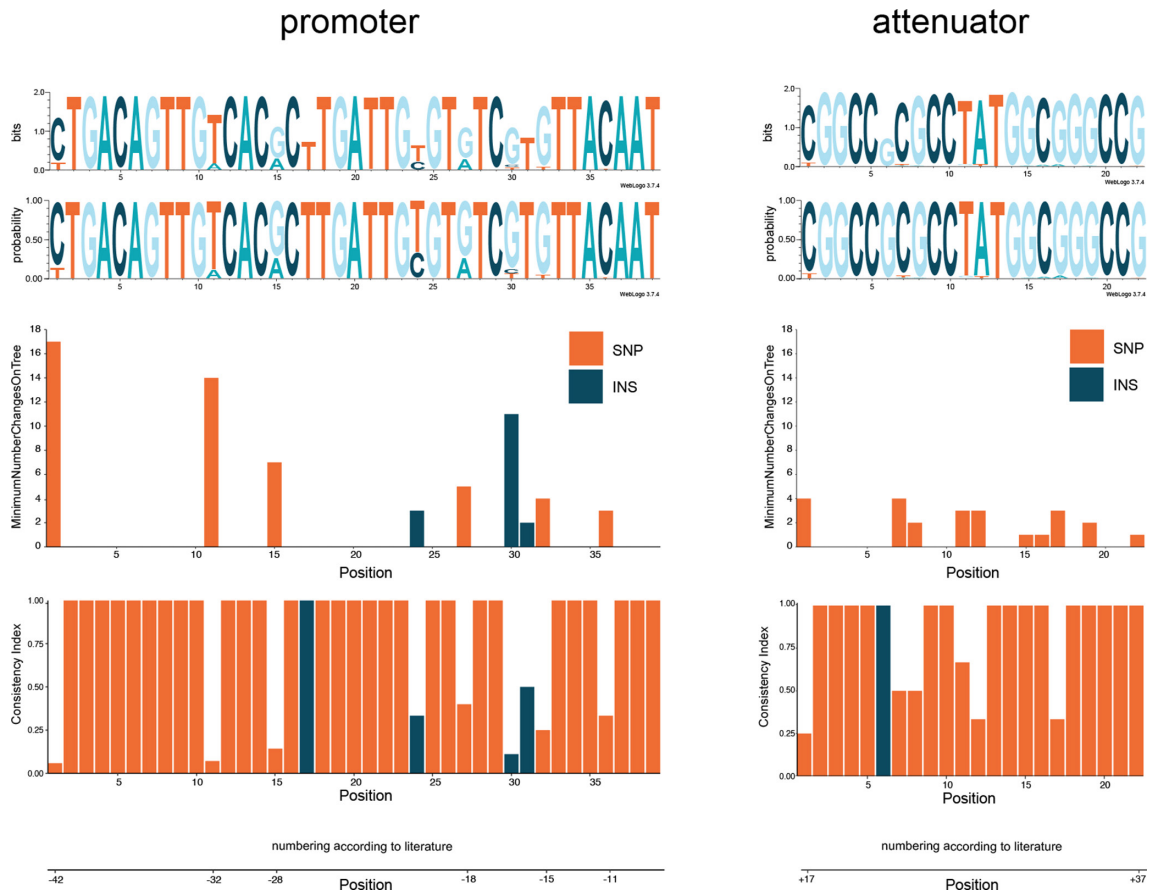
**Fig. 2.** Approximately maximum-likelihood phylogenetic tree of all 172 *E. coli* isolates based on the coreSNP alignment with the resistance for CTX, *pampC* gene presence, MLST STs, phylogroups, and the alignments of the promoter and the attenuator region. Positions with a SH-like (Shimodaira–Hasegawa like) branch support  $\leq 60\%$  are indicated as red dots. Scale bar indicates branch length calculated by FastTree.

evolved multiple times independently. After selecting the lowest scoring consistency index positions,  $\leq 0.05882353$ , 24 relevant genomic positions were identified that had both a low consistency index and a significant association with CTX resistance. Most notably, 1 of these 24 positions is position  $-42$ . Only 2 mutations of those 24 that were located in genes were non-synonymous: a (conservative) missense mutation in the type II secretion system protein L (*gspL*) gene leading to a Ser330Thr alteration, and a mutation in the hydroxyethylthiazole kinase (*thiM*) gene resulting in a Thr122Ala alteration according to the annotation of

*E. coli* strain ampC\_0069 (accession no. CP046396). In addition to the non-synonymous mutation found on the *gspL* gene, eight synonymous mutations are also located in genes annotated as being part of the type II secretion system. A complete overview is presented in Tables 2 and S4.

### Recombination analysis

To verify whether the level of homoplasmy could be a result of recombination, we used the Genealogies Unbiased By



**Fig. 3.** WebLogo sequence logo with both bits and probability score for the promoter and the attenuator region. Consistency index and the minimum number of changes on the tree per position are represented in the bar charts below the sequence logos. Orange bars are positions containing SNPs that are tested for homoplasy. Blue bars are positions containing insertions that are tested for homoplasy.

recombinations in Nucleotide Sequences (Gubbins) algorithm to predict recombination events in our isolate collection [35]. This analysis showed frequent recombination events in our 172 *E. coli* isolates (see Fig. S3). Results illustrate that recombination blocks cover the region of the *gspL* and the *thiM* gene, and their high homoplasy levels could, thus, be due to recurrent recombination rather than independent mutations. Nonetheless, position  $-42$  in the *ampC* promoter is not located in a region affected by recombination, as shown in Fig. S3. Moreover, when inferring the phylogenetic tree corrected for recombination events as obtained from the Gubbins analysis, the  $-42$  C>T mutation actually occurred in 18 independent branches rather than the 17 branches in the uncorrected tree. This supports our previous result that this mutation is homoplastic, and not the result of a recurrent recombination event.

### Pyseer analysis

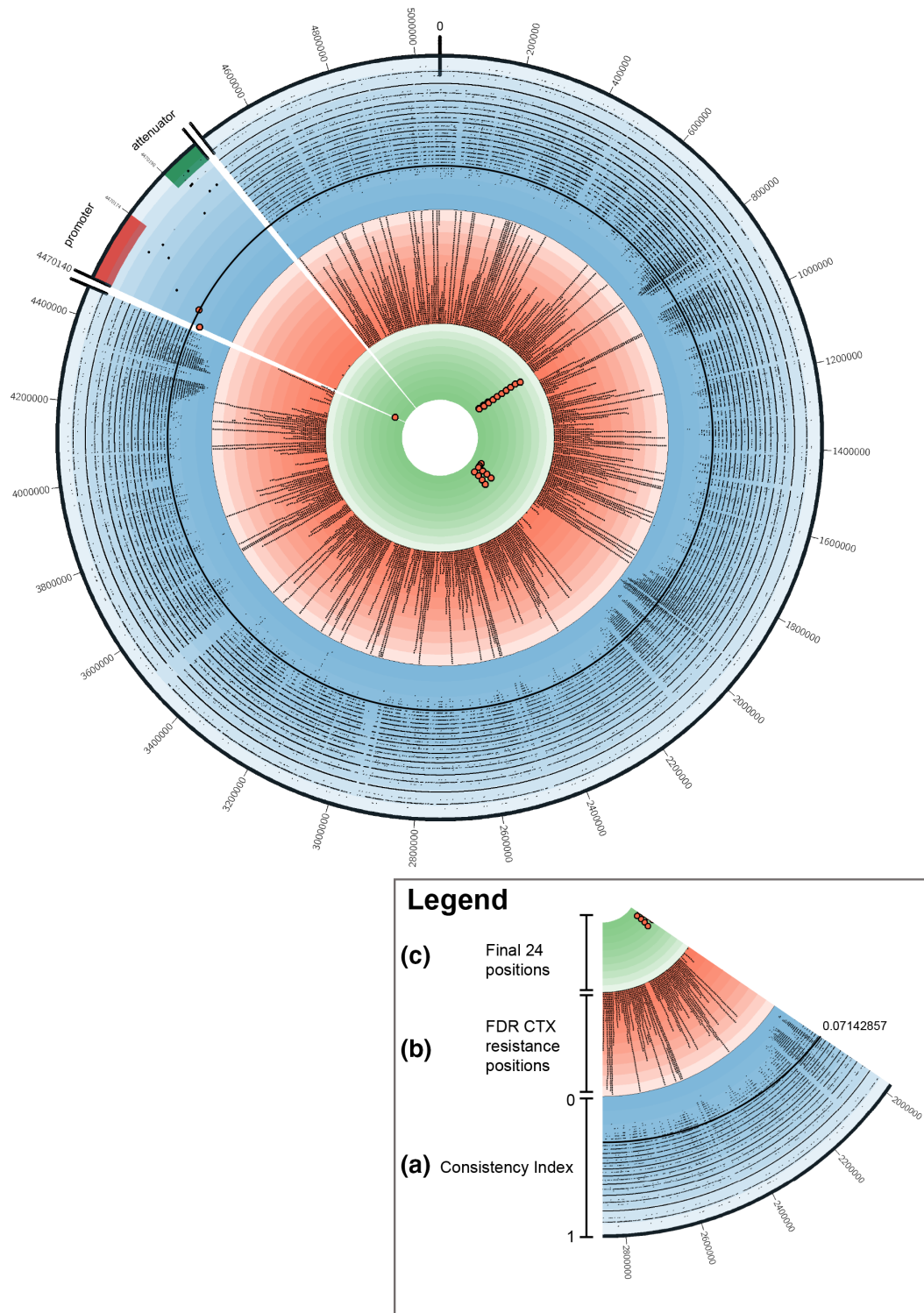
To add additional support to our findings, we used Pyseer as an alternative method to compare the 24 positions identified with the homoplasy-based association analysis (see Table 2). Pyseer identified 65501 unique significant

mutations associated with CTX resistance with filter  $P$  value  $\leq 0.05$  and 1097 unique positions with filter and likelihood ratio test (lrt)  $P$  value  $\leq 0.05$ . Of the 24 positions identified with the homoplasy-based association analyses, we identified 8 positions also reported by the Pyseer method (see Table 2). Furthermore, the Pyseer method identified 6 complex mutation variants and a total of 14 positions that have multiple mutation variants overlapping the same genomic positions as found by the homoplasy-based method. The most dominant position associated with CTX resistance is the  $-42$  C>T promoter mutation as indicated by both methods. No further positions on the promoter or the attenuator were found significantly associated with CTX resistance.

### DISCUSSION

We present a genome-wide analysis in which homoplastic mutations are associated with antibiotic resistance in *E. coli*. By comparing WGS data of 172 *E. coli* isolates to a reference chromosome, we were able to reconstruct the evolution of the genomes and therewith map recurrent events, allowing us to detect homoplasy associated with CTX resistance.





**Fig. 4.** CIRCOS plot for the full chromosome of *ampC\_0069* (accession no. CP046396) with, per position, the various metrics used. (a) The blue coloured ring represents the consistency index results per genomic position. The two red dots indicate the  $-42$  and  $-32$  position on the promoter. The black circle line indicates the  $0.07142857$  consistency index value. (b) The ring with a red background shows all positions that were significantly associated with CTX resistance in all non-*pampC*-harbouring *E. coli* isolates. Larger bars pointing outwards indicate multiple significant associated positions in a small genomic region. (c) The ring with the green background shows all 24 positions that have a low consistency index of  $\leq 0.05882353$  and are significantly associated with CTX resistance in all non-*pampC*-harbouring *E. coli* isolates.

**Table 2.** Details of the 24 positions with a significant association with CTX resistance (FDR  $\leq 0.05$ ) and with a consistency index  $\leq 0.05882353$ 

Homoplasy-based association analysis											Pyscer			
Genomic position	Mutation	ACGT counts	Min changes	CI	Crosstab	AF	Fisher P-value	FDR	Gene name	Genomic position	Mutation	AF	Filter P-value	Irt P-value
810581	A>G	41:0:122:0	18	0.056	0:22:26:34	6.83E-01	8.11E-05	0.039	<i>glcE</i>	810581	A>G	0.683	1.87E-04	2.09E-02
810791	G>T	0:0:54:109	17	0.059	0:22:28:32	6.59E-01	1.68E-05	0.035	<i>glcE</i>	810791*	GGGT>TGGC	0.646	4.08E-04	1.94E-02
815680	C>T	0:69:0:93	18	0.056	2:20:34:26	5.61E-01	1.06E-04	0.042	<i>glcA</i>	815680	C>T	0.561	1.20E-04	2.07E-02
824522	T>C	0:82:0:73	18	0.056	2:20:36:24	5.37E-01	3.58E-05	0.036	<i>gspC</i>	824522*	T>C	0.378	8.08E-06	1.00E+00
828695	A>G	79:0:74:0	18	0.056	3:19:41:19	4.63E-01	1.14E-05	0.034	<i>gspF</i>	828695	A>G	0.463	1.08E-05	3.26E-02
830684	T>C	0:80:0:73	18	0.056	2:20:37:23	5.24E-01	1.61E-05	0.034	<i>gspJ</i>	830684*	T>C	0.488	3.74E-05	1.06E-01
830708	A>G	68:0:85:0	19	0.053	2:20:36:24	5.37E-01	3.58E-05	0.036	<i>gspJ</i>	830708*	A>G	0.111	6.41E-01	5.02E-01
830732	T>C	0:82:0:71	17	0.059	2:20:36:24	5.37E-01	3.58E-05	0.036	<i>gspJ</i>	830732	T>C	0.537	4.20E-05	4.19E-02
831564	G>A	71:0:83:0	17	0.059	5:17:45:15	3.90E-01	2.76E-05	0.036	<i>gspK</i>	831564	G>A	0.39	1.71E-05	2.37E-01
832152	T>C	0:86:0:72	18	0.056	2:20:38:22	5.83E-01	1.08E-05	0.034	<i>gspK</i>	832152*	CGTGTG>TGTCA	0.195	6.94E-03	2.29E-01
832287	A>T	81:0:0:79	17	0.059	2:20:37:23	5.24E-01	1.61E-05	0.034	<i>gspK</i>	832287*	T>A	0.232	6.73E-02	5.73E-01
833451	A>T	64:0:0:98	20	0.05	1:21:34:26	5.73E-01	1.03E-05	0.034	<i>gspL</i>	833451*	T>A	0.378	1.69E-04	9.04E-02
843887	G>A	64:0:40:0	19	0.053	21:1:29:31	3.90E-01	7.00E-05	0.036	Unnamed	843884*	GTTTGGC>TCGACGT	0.354	4.08E-04	1.08E-01
1863004	A>G	126:0:46:0	18	0.056	9:13:53:7	2.44E-01	3.37E-05	0.036	<i>flrM</i>	1862989*	CTGTCTCATAAGCCCAACCGCC > TTATCTTTTAAACCGGCAGCG	0.232	4.55E-05	3.30E-02
1911551	T>C	0:116:0:44	17	0.059	16:6:14:46	6.34E-01	7.02E-05	0.036	Unnamed	1911551*	C>T*	0.28	1.22E-03	5.28E-01
1946016	A>G	58:0:91:0	23	0.043	17:5:17:43	5.85E-01	1.03E-04	0.041	<i>ugd</i>	1946016*	A>G*	0.402	5.02E-02	4.92E-01
1946067	G>T	0:0:65:84	22	0.045	18:4:20:40	5.37E-01	1.25E-04	0.05	Non-coding region	1946067	G>T	0.537	9.58E-05	2.26E-01
1946072	T>A	84:0:0:65	22	0.045	18:4:20:40	5.37E-01	1.25E-04	0.05	Non-coding region	1946072	T>A	0.537	9.58E-05	2.26E-01
1952745	A>G	131:0:40:0	28	0.036	9:13:52:8	2.56E-01	7.67E-05	0.037	<i>hisD</i>	1952745	G>A	0.732	6.54E-05	7.96E-03
2051214	T>A	137:0:0:34	19	0.053	11:11:55:5	8.05E-01	1.00E-04	0.041	Unnamed	2051211*	CACT>TACA	0.659	3.92E-03	4.87E-01
2051220	T>A	137:0:0:34	19	0.053	11:11:55:5	8.05E-01	1.00E-04	0.041	Unnamed	2051220*	T>A	0.793	4.79E-06	5.72E-02
2057518	A>G	111:0:60:0	19	0.053	7:15:49:11	3.17E-01	3.89E-05	0.036	<i>dem</i>	2057506*	ATGTTTCCCTGGCAGCGAGT > CTGCTATCCGGCAACGTTAT	0.0122	9.66E-02	1.00E+00
2068593	G>A	45:0:125:0	17	0.059	9:13:52:8	2.56E-01	7.67E-05	0.037	<i>flrM</i>	2068593	G>A	0.256	2.60E-05	1.28E-01
4470140	C>T	0:147:0:24	17	0.059	7:15:51:9	2.93E-01	8.35E-06	0.034	<i>ampC</i> promoter	4470140	C>T	0.293	2.74E-06	5.42E-03

\*pyscer identified multiple mutation variants, affecting the outcome, only the position with the highest allele frequency (AF) is reported which overlaps the result of the homoplasy-based association analysis. Sequences in the pyscer mutation of the pyscer outcome that are the corresponding position with the tested genomic position. crosstab:R(other:R/mutation:S/other:S/mutation). AF, allele frequency; CI, consistency index; FDR, false discovery rate; Irt, likelihood ratio test.

Our foremost finding is the significant association of the  $-42C>T$  mutation, in the *ampC* promoter, with CTX resistance that evolved independently at least 17 times in five distinct phylogroups. The  $-42C>T$  mutation has been confirmed in former studies to result in AmpC hyperproduction in *E. coli* [6, 24, 44]. Nelson *et al.* demonstrated an 8 to 18 times increase in activity of AmpC when cloning the promoter upstream a *lac* operon [44]. Conversely, Caroff *et al.* found a decrease in expression of AmpC when cloning the promoter with a  $-42T>C$  mutation in a pKK232-8 reporter plasmid with a chloramphenicol acetyltransferase gene [24]. Tracz *et al.* confirmed that the  $-42C>T$  mutation has the strongest effect on the *ampC* promoter, resulting in a high expression of the *ampC* gene as detected by qRT-PCR [6]. Despite the fact that the  $-42C>T$  mutation has such a strong effect on AmpC production, the effect of the mutation on CTX MICs had not been confirmed. Moreover, the contribution of convergent evolution on this position relative to the role of the expansion of a clone with a beneficial mutation at this position has not been determined. That being the case, this study provides evidence that this  $-42 C>T$  mutation is not a result of a recombination event and most likely evolved many times independently.

In the current study, we see a strong correlation between the  $-42C>T$  mutation and CTX resistance, even though there were exceptions, as not all isolates with this mutation were considered resistant according to EUCAST guidelines. As described by Coolen *et al.*, the MIC for CTX in putative AmpC hyperproducers was generally higher than in the putative low-level AmpC producers, though the range in CTX MICs overlapped [16]. Yet, the lowest MIC measured in the isolates with the  $-42C>T$  mutation was  $0.75 \text{ mg l}^{-1}$ , which is at the higher end of the EUCAST epidemiological cut-off values (ECOFF) distribution. The variation in phenotypical testing could be an explanation, although an interplay of AmpC hyperproduction and other strain-specific factors as previously described by Tracz *et al.* may also be considered [6].

In order to avoid biasing towards a single method, in our case the homoplasy-based association method, we performed the analysis using Pyseer on the same data set. The outcome of the Pyseer analysis provided a similar number of CTX-resistance-associated mutations and also confirmed the strong association of the  $-42 C>T$  mutation. Nonetheless, when zooming in on the identified positions by the homoplasy-based method, not all 24 mutations were significantly associated with CTX by Pyseer. An explanation for this discrepancy is that the combination of snippy and Pyseer identifies various mutation variants on the same overlapping genomic region by stratifying it as complex, indels and/or SNP. This greatly affects the power of the test on certain positions. An example of the differences in mutation variants is illustrated in Table 2.

Remarkably, we observed that the  $-42C>T$  mutation never occurs in the presence of a *pampC* gene (0 out of 24 cases). This was even noticed in isolates with the same MLST, i.e. ST88  $-42C>T$  ( $n=3$ ) and *pampC* ( $n=1$ ), suggesting preferred exclusivity for one of the resistance mechanisms. One study

mentioned the co-occurrence of the  $-42C>T$  mutation and a *pampC* gene in only 1 out of 36 strains [45]. One could speculate that the exclusivity is a matter of what arrives first, the plasmid or the mutation, after which there is no selective advantage for the second mechanism, or that there is actually a fitness cost to having both the mutation and the plasmid relative to having only the mutation or the plasmid. This hypothesis might be a start for future studies to determine the relative fitness and resistance provided by the  $-42C>T$  mutation relative to isolates harbouring a *pampC* gene.

The study performed by Tracz *et al.* showed that position  $-32T>A$  on the promoter of *ampC* associates with AmpC hyperproduction that results in elevated MIC levels for FOX [6]. Surprisingly, in the current study, no significant association of  $-32T>A$  with CTX resistance was noticed despite its low consistency index. Only 2 out of 20 isolates with the  $-32T>A$  were CTX resistant, 4 out of 20 showed an intermediate elevated CTX MIC, and 14 were susceptible to CTX. Although we do not know under which conditions this mutation did arise, it can be speculated that the high level of homoplasy at the  $-32$  position is associated with a different trait, e.g. resistance against another antibiotic.

Prior studies discovered the importance of mutations in the promoter elements. Although an existing promoter is often copied upstream to the gene, a *de novo* promoter can also evolve out of an existing sequence region. Random sequences can even evolve expression comparable to the wild-type promoter elements after only a single mutation [46]. This means that the *de novo* creation of a promoter region within the *E. coli* may be much more often the result of mutation rather than a rearrangement. Furthermore, these promoter elements evolve to only a few forms, indicating convergent evolution [47], as also observed in the present study. All encountered variants seem to result in a sequence that resembles the *E. coli* consensus  $\sigma^{70}$  promoter more than the wild-type sequence they are derived from [6].

Next to the  $-42C>T$  promoter mutation, we detected 23 other positions in our analysis that are associated with CTX resistance and have extremely high levels of homoplasy. Most of these are synonymous mutations, with only two missense mutations (*thiM* and *gspL*) found. It is remarkable that one missense mutation (p.Ser330Thr) is located in *gspL* that encodes a protein of the type II secretion system. The type II secretion system is used by many Gram-negative bacteria to translocate folded proteins from the periplasm, through the outer membrane, into the extracellular milieu [48]. The system is composed of 12–15 different general secretory pathway (Gsp) proteins and is related to virulence of various pathogenic *E. coli*, e.g. EHEC (enterohaemorrhagic *E. coli*) and UPEC (uropathogenic *E. coli*) [49–51]. It could be that in our selection of mainly clinical samples a certain predilection has occurred towards isolates with particular virulence traits and not based on mechanistic benefits. The *gspL* gene has been described as being part of the accessory genome of *E. coli* [52]. Our study supports this finding as some strains did not harbour this gene. Additionally, we found evidence that

recombination events in the type II secretion system could be the underlying cause of the extreme homoplasy levels. Still, it is remarkable that missense mutation p.Ser330Thr in the *gspL* gene correlates with the CTX-resistance trait even though it is most likely caused by a recombination event. To the best of our knowledge, no relationship between the type II secretion system and CTX resistance has been observed before. One could hypothesize that the mutation is a secondary adaptation needed to cope with the elevated AmpC production, as the peptidoglycan (PG) layer is affected by AmpC hyperproduction and the type II secretion system contains proteins that are partly localized in the periplasm [53, 54].

The use of genomic data to detect homoplasy events is an accepted scientific technique [55–57]. In *Mycobacterium tuberculosis*, it is a well-known method for identifying advantageous mutations, as they are likely to be associated with phenotypes such as drug resistance, heightened transmissibility or host adaptation [12–15]. In other species, e.g. *Staphylococcus aureus* and *Burkholderia pseudomallei*, the method has been effectively applied to identify mutations associated with antibiotic resistance or virulence-associated genes [58, 59]. Homoplasy-based association analysis limits phylogenetic bias by correcting for genetic relatedness of strains with the same phenotype, thereby increasing statistical power to find true associations [14]. Taking this into account, the use of homoplasy-based association analysis seems viable to relate polymorphic sites to phenotypic traits in bacteria. Still, studies on other genera than mycobacteria are scarce. To our knowledge, no homoplasy studies have used this method on *E. coli*.

The increase of 3GC resistance imposes a clinical threat by restricting treatment options and it is essential to understand the underlying resistance mechanisms. To be able to explore these mechanisms, we selected primarily clinical *E. coli* strains. The current study is directed on exploring AmpC-mediated CTX resistance. Therefore, we included isolates that are already suspected for increased AmpC production based on elevated FOX resistance. Since a random sample of *E. coli* would limit finding homoplasy-based associated promoter mutations with CTX resistance. A downside of these selection criteria might be that we over-estimated certain genetic variants associated with the trait, as we do not know the frequency of these variants in the general population. Despite the fact that the spontaneous mutation rate in *E. coli* is relatively low [60], it is still likely that this particular mutation occurs often in the general population, given the vast amounts of *E. coli* in the environment [61], providing ample opportunities for adaptation to antibiotics and arguing for antibiotics of which genomic adaptation requires multiple mutations in order to develop resistance.

The findings of this study have a number of implications for future practice. This study not only grants insights into how chromosome-encoded antibiotic resistance evolves, but also provides potential strategies for future homoplasy-based association studies. Furthermore, the use of genome-wide homoplasy-based analysis could be applied to optimize

outbreak analysis. Prior studies have optimized outbreak analysis by removing recombinant regions [62, 63]. Homoplasy events disturbs the true phylogeny; hence, removing genomic positions that are heavily affected by homoplasy could improve tree topology, thereby refining outbreak analysis, although this strategy is still under debate [64].

## Conclusions

To conclude, our method demonstrates extreme levels of homoplasy in *E. coli* that are significantly associated with CTX resistance. Greater access to WGS data provides new opportunities to perform large-scale genome-wide analysis. Homoplasy-based methods can have a potential role in future studies as they constitute an effective strategy to relate phenotypic traits to variable genomic positions.

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## Author contributions

H. F. L. W., J. A. J. W. K. and M. A. H. conceived and supervised the study. J. P. M. C., E. P. M. D., E. K., J. A. S., J. J. V., W. J. G. M. and K. N. performed the data acquisition. J. P. M. C. and E. P. M. D. performed the data analysis. J. P. M. C. performed bioinformatic analysis. J. P. M. C., E. P. M. D. and M. A. H. performed the data interpretation and wrote the manuscript. All authors read and approved the final manuscript.

## Conflicts of interest

The authors declare that there are no conflicts of interest.

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