RESEARCH ARTICLE

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Effects of a previously selected antibiotic resistance on mutations acquired during development of a second resistance in *Escherichia coli*



Marloes Hoeksema¹, Martijs J. Jonker², Stanley Brul¹ and Benno H. ter Kuile^{1,3*}

Abstract

Background: The effect of mutations conferring antibiotic resistance can depend on the genetic background. To determine if a previously de novo acquired antibiotic resistance influences the adaptation to a second antibiotic, antibiotic resistance was selected for by exposure to stepwise increasing sublethal levels of amoxicillin, enrofloxacin, kanamycin, or tetracycline. *E. coli* populations adapted to either a single or two antibiotics sequentially were characterized using whole genome population sequencing and MIC measurements.

Results: In a wild-type background, adaptation to any of the antibiotics resulted in the appearance of well-known mutations, as well as a number of mutated genes not known to be associated with antibiotic resistance. Development of a second resistance in a strain with an earlier acquired resistance to a different antibiotic did not always result in the appearance of all mutations associated with resistance in a wild-type background. In general, a more varied set of mutations was acquired during secondary adaptation. The ability of *E. coli* to maintain the first resistance during this process depended on the combination of antibiotics used. The maintenance of mutations associated with resistance to the first antibiotic did not always predict the residual MIC for that compound.

Conclusions: In general, the data presented here indicate that adaptation to each antibiotic is unique and independent. The mutational trajectories available in already resistant cells appear more varied than in wild-type cells, indicating that the genetic background of *E. coli* influences resistance development. The observed mutations cannot always fully explain the resistance pattern observed, indicating a crucial role for adaptation on the gene expression level in de novo acquisition of antibiotic resistance.

Keywords: Bactericidal antibiotics, de novo resistance, Mutations, Whole genome sequencing

Background

To be able to predict, prevent, or slow down development of antibiotic resistance, the molecular mechanisms that drive development of antibiotic resistance need to be understood. Antibiotic resistance can develop in three distinct ways: through horizontal gene transfer, chromosomal mutations, or phenotypic adaptation. In *Escherichia coli*,

the first stages of de novo development of antibiotic resistance occur on a phenotypic level, controlled by adaptation on the gene expression level [1]. After this initial stage, mutations appear that, in most cases, confer a reduction in bacterial fitness [1-3]. Often, this is followed by the appearance of compensatory mutations that reduce the loss in fitness without reducing the acquired resistance [4].

Antibiotic resistance can be selected for by exposure to stepwise increasing sublethal levels of the antibiotic [5]. Although each antibiotic class has a specific cellular target, the radical-based theory suggests that as a secondary effect increased levels of cellular reactive oxygen species (ROS) occur as a result of a pathway common to

Full list of author information is available at the end of the article



^{*} Correspondence: b.h.terkuile@uva.nl

¹Laboratory for Molecular Biology and Microbial Food Safety, Swammerdam Institute for Life Sciences, University of Amsterdam, Amsterdam, The Netherlands

³Netherlands Food and Consumer Product Safety Authority, Office for Risk Assessment, Utrecht, The Netherlands

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all bactericidal antibiotics [6, 7]. Following this logic, similar mutations might occur in cells with acquired resistance to different bactericidal antibiotics. During evolution experiments exposing *E. coli* to steadily increasing levels of antibiotics, cells with an earlier acquired resistance to a bactericidal antibiotic adapt to a second bactericidal, but not bacteriostatic, antibiotic at a more rapid rate, corroborating this theory [8].

The effect of resistance associated mutations varies depending on the genetic background, such as the presence of other resistance mutations, a phenomenon known as epistasis [9, 10]. The fitness cost incurred by mutations determines the evolutionary pathways available [11], suggesting that the evolutionary history of bacteria may influence the type or number of mutations that are acquired upon exposure to an antibiotic. In this study, we investigate if a previously de novo acquired antibiotic resistance influences the adaptation to a second antibiotic. Whole genome population sequencing was applied to E. coli strains with de novo acquired resistance to either one or to two antibiotics sequentially. We previously reported on larger genomic changes that occur during adaptation to a single antibiotic, or two antibiotics successively [12]. In this study, we provide a comprehensive overview of the different mutations that are acquired when wild-type or antibiotic-resistant E. coli is exposed to amoxicillin, enrofloxacin, kanamycin, or tetracycline.

Results

This study addressed three questions: 1) which mutations are associated with the development of resistance by *E. coli* wild-type against specific antibiotics? 2) Are the same mutations observed when cells made resistant against one antibiotic become resistant to a second one? 3) Are initial mutations lost when cells resistant to one antibiotic are made resistant to another one? To answer these questions, strains exposed to only amoxicillin, enrofloxacin, or kanamycin, all bactericidal antibiotics from different classes, or the bacteriostatic antibiotic tetracycline were compared to cells made resistant to two of these antibiotics successively. Resistant strains were generated by exposing wild-type E. coli or a strain with an earlier acquired resistance to increasing but sublethal concentrations of any of these antibiotics [8]. The initial resistance was selected for in duplicate strains. Two replicates of each of these strains were made resistant to a second antibiotic, resulting in four strains with the same exposure history (Fig. 1). Genomic DNA was isolated from the entire population at selected time points for whole genome sequencing at an average read depth of 226 to identify genetic changes associated with acquired resistance to a single and to subsequent antibiotics.

Adaptation of the wild-type to amoxicillin (Table 1) resulted in two well-scrutinized mutations in the ampC promoter [1, 13], and in mutations in ompR, citF, and *rpoD*, all with a population frequency of 1. One of the replicate strains acquired a mutation in waaB in addition to these mutations. In the wild-type strain adapted to enrofloxacin, either ten or eleven genes were mutated, and a low-frequency mutation occurred in mutL. Mutations in gyrA, parC, parE, soxR, and acrR, commonly associated with resistance to fluoroquinolones [14-16] were accompanied by less frequently observed mutations in the genes *proY* and *yciO*. In addition, mutations in the 5' UTR of rpsJ, ybjL or ybjM, and gtlB were detected. When wild-type cells were exposed to kanamycin, mutations accumulated in sbmA, sapF, cpxA, npr, fusA, and the 5' UTR of uidA. Interestingly, a mutation in gyrB, most often associated with resistance to fluoroquinolones [17], was identified with a population frequency of 1 in both wild-type-adapted strains. Development of resistance to tetracycline in wild-type E. coli resulted in mutations in acrR and acrB, ompF, mlaA. A mutation in rpoC was only detected in a strain with partial resistance to tetracycline.

Development of secondary resistance does not always result in the appearance of all mutations associated with resistance to that specific antibiotic in a wild-type strain (Fig. 2, Tables 2, 3, 4 and 5). This is most apparent in the strains adapted to kanamycin (Fig. 2c), where only mutations in *sbmA* and *fusA* are acquired by all strains. A high degree of variability can be observed between the replicates, as indicated by the frequencies displayed in Tables 2, 3, 4 and 5(column headed "Strains"), suggesting that secondary adaptation allows for more flexibility than primary development of resistance. Moreover, selected resistance to a second antibiotic results in the appearance of mutations that are acquired by multiple strains already resistant to another antibiotic but not by wild-type strains, such as rpoA during secondary amoxicillin resistance development (Fig. 2a), caiA, ahpC, rph, and spoT during adaptation to enrofloxacin (Fig. 2b), kdpD and acrB during development of resistance to kanamycin as a second antibiotic (Fig. 2c), or envZ when already resistant cells adapt to tetracycline (Fig. 2d). Within different antibiotic-resistant populations, a *mutL* mutation was identified in several samples, with a population frequency varying from 0.09 to 0.20.

All genes containing a mutation were functionally annotated using clusters of orthologous groups (COG) analysis and clustered according to the four different categories (Fig. 3). The majority of mutations occur in genes involved in information storage and processing in general (145/268) and transcription (65/268) (Fig. 3a). For development of resistance to amoxicillin, enrofloxacin, and kanamycin, no major shift in the function of the

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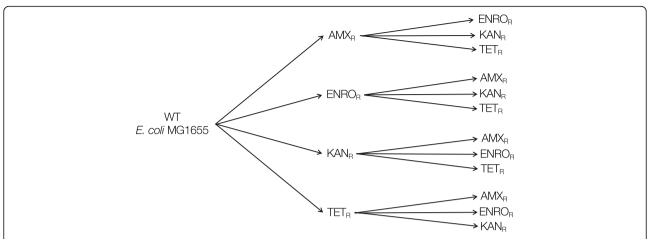


Fig. 1 Set-up of evolution experiments inducing resistance. Duplicate strains of wild-type *E. coli* MG1655 were made resistant by exposure to increasing non-lethal concentrations of amoxicillin (AMX), enrofloxacin (ENRO), kanamycin (KAN), or tetracycline (TET). Two replicates of each of the primary resistant strains were subsequently made resistant to one of the three other antibiotics using the same approach. As a result 4 replicates were obtained of all double exposed strains

mutated genes could be detected (Fig. 3b). For tetracycline, the total number of mutations is significantly lower than the numbers observed with the other three antibiotics, making the comparison slightly problematic.

The minimal inhibitory concentration (MIC) of the first antibiotic was measured in all strains exposed to two antibiotics sequentially, to determine if the original resistance could be maintained during development of resistance to a second antibiotic (Fig. 4). In general, the ability of *E. coli* to maintain its original resistance appears to depend on the combination of antibiotics used. Amoxicillin resistance is compatible with kanamycin and tetracycline exposure, but not with enrofloxacin. For enrofloxacin resistance, the same applies. Acquired resistance to kanamycin resistance can co-exist with enrofloxacin resistance, but not with resistance to other antibiotics. Tetracycline resistance persists after exposure to kanamycin, but not to enrofloxacin. Exposure to amoxicillin results in a mixed pattern.

The percentage of residual mutations was determined by dividing the number of original mutations still present in the strain at the end of the experiment by the number of mutations acquired by the parental strain and multiplying this number by 100 (Fig. 4). When comparing the residual MIC with the number of original mutations still present for most antibiotic combinations, loss of resistance can be correlated with loss of mutations and vice versa. However, a clear correlation cannot always be established. For amoxicillin and enrofloxacin resistance, the absence of mutations correlates to the loss of the original resistance. Resistance to kanamycin and tetracycline, in contrast, is partially or fully maintained, but this does not coincide with the maintenance of the original resistance mutations, suggesting that either the

additionally acquired mutations also contribute to the original resistance, or that subsequent adaptation on a gene expression level maintains the level of resistance to the first antibiotic.

Discussion

Taken together, the data presented in this study indicate that the mutations involved in de novo acquisition of resistance to each antibiotic are unique for each specific antibiotic. The shared killing mechanism proposed for all bactericidal antibiotics [6, 7] and the reduction of ROS levels measured in resistant strains [8] suggest that common mutations may occur during adaptation. However, no common mutation shared by all populations resistant to one of the bactericidal antibiotics could be identified. The one common mutation identified throughout the dataset is a mutL W390 L mutation (Tables 2, 3, 4 and 5). In E. coli, MutL is essential for mismatch repair [18]. Defects in mismatch repair are often observed in antibiotic-resistant strains [19, 20] and are known to increase mutation rates over 200-fold [21]. Considering its role in DNA mismatch repair, the mutation frequency varying between 0.09-0.20 most likely provides the population with a source of increased variation without causing deleterious effects on the entire population.

In all resistant strains, most shared mutations that are acquired by both the wild-type and strains with acquired resistance mostly have a frequency of (close to) 1, indicating fixation in the population. For each antibiotic, mutations previously associated with that antibiotic resistance were observed (Table 1). The occurrence of the same mutations in replicates concurs with the earlier observation that while the initial mutations during the

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Table 1 Mutations associated with resistance to amoxicillin, enrofloxacin, kanamycin, or tetracycline in wild-type *E. coli*

Exposure to amoxicillin					
			WT		WT
Gene product	Gene	Mutation	40 μg/ml	1280 μg/ml	1280 μg/n
B-lactamase	атрС	G23 T (promoter)	1	1	1
		C-11 T (promoter)	1	1	1
Transcriptional regulator	ompR	E88A		1	1
Citrate lyase	citF	G432A		1	0.97
Sigma 70	rpoD	D445V		0.96	1
UDP-D-galactose	waaB	Del <u>C3803246</u> (FS297)			1
Exposure to enrofloxacin					
				WT	WT
Gene product	Gene	Mutation		1024 μg/ml	1024 μg/n
DNA gyrase subunit A	gyrA	D87G		1	
		S83 L		1	1
		D87N			1
DNA gyrase subunit B	gyrB	S463F			1
DNA topoisomerase IV subunit A	parC	S80R		1	1
DNA topoisomerase IV subunit B	parE	Ins Q458		0.94	1
Cryptic proline/histidine transporter	proY	A235S		1	1
Transcriptional regulator	acrR	Del <u>485,885–11</u> (FS42)		1	1
Transcriptional regulator	soxR	T133S, del <u>4,277,899–903</u> (FS144)		1	1
S10 subunit of 30S	rpsJ	5' UTR (<u>G3453306 T</u>)		1	1
Putative transport protein/Putative inner membrane protein	ybjL/ybjM*	5' UTR (<u>A889923G</u>)		1	1
Glutamate synthase subunit	gltB	5' UTR (<u>C3354487T</u>)		1	1
Putative RNA binding protein	yciO	N64Y		1	1
Mismatch repair protein	mutL	W390 L			0.12
Exposure to kanamycin					
_	_		WT		WT
Gene product	Gene	Mutation	32 μg/mL	1024 μg/mL	1024 μg/m
Peptide antibiotic transporter	sbmA	Del <u>397,306–27</u> (FS 222)		1	1
ATP binding protein of putrescine ABC exporter	sapF	L181Q		1	1
Sensor protein of Cpx TCS	срхА	Q242L		1	1
DNA gyrase subunit B	gyrB	S464Y		1	1
Phosphorelay protein	npr	L23R		1	1
Elongation factor G	fusA	Q242L		1	
		A608V	0.19		
		F05 L	0.74		1
β -D-glucuronidase	uidA	5' UTR (<u>T1696220C</u>)			1
Exposure to tetracycline					
			WT		
Gene product	Gene	Mutation	16 μg/mL	64 μg/mL	
Multidrug efflux pump RND permease	астВ	145L	1	1	
Transcriptional regulator	acrR	P85Q	1	1	
Outer membrane porin F	ompF	T71S	0.65	0.04	
		Ins 7 nt after <u>G986771</u> (FS71)		0.60	
Outer membrane lipoprotein	mlaA	Del N41/F42		0.9	
RNA polymerase subunit β'	rpoC	G367C	0.34		

Numbers shown indicate frequency of mutation in population. *Del* = deletion, *Ins* = insertion, underlined letters and numbers indicate nucleotides and their genomic position, * indicates that mutation could affect either gene. For amoxicillin, enrofloxacin, and kanamycin, two independent strains were sequenced

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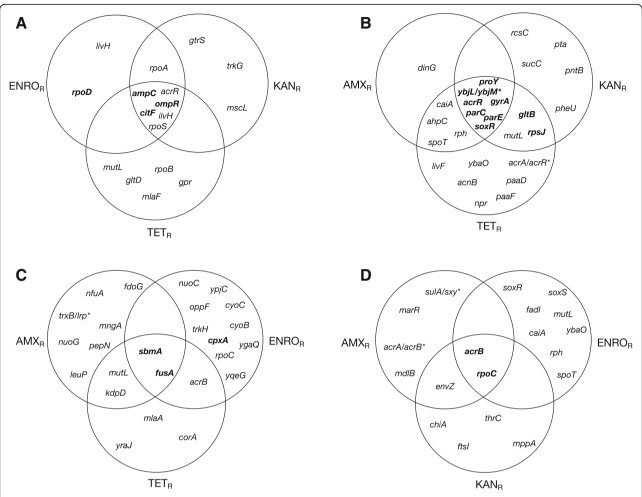


Fig. 2 Overlap of mutations associated with resistance to amoxicillin (**a**), enrofloxacin (**b**), kanamycin (**c**), or tetracycline (**d**) in strains with a previously acquired resistance to a different antibiotic. AMX_R: amoxicillin resistant, ENRO_R: enrofloxacin resistant, KAN_R: kanamycin resistant, TET_R: tetracycline resistant. Mutations associated with resistance in a wild-type background are shown in bold. Mutations associated with the previous resistance have been omitted from the figure

development of resistance may vary, at the higher concentrations a single mutation or set of mutations becomes dominant [1]. From a mechanistic point of view, acquired antibiotic resistance can be classified into three groups: 1) prevention of access to target by reduction of permeability (ompR for amoxicillin, soxR for enrofloxacin, sbmA and sapF for kanamycin, ompF for tetracycline) or increased efflux (acrR and soxR for enrofloxacin, acrB and acrR for tetracycline), modification of target (gyrA, parC, and parE for enrofloxacin), or 3) antibiotic modification (ampC for amoxicillin). The effect of other mutations identified in this dataset is not always clear and could vary. Some mutations appear to have a non-structural function, such as mutations found in the 5' UTR region of rpsJ, gltB, and ybjL or ybjM in strains with acquired resistance to enrofloxacin, by potentially influencing gene expression or translation [22, 23]. Mutations in sigma factors, for example *rpoD* in amoxicillin resistance, *rpoC* in tetracycline resistance, or *rpoS* during secondary adaptation to amoxicillin or tetracycline, can redirect expression of an entire set of genes [24].

Protein moonlighting is a phenomenon in which proteins, often metabolic proteins or molecular chaperones, fulfil more than one biological function [25]. Metabolic proteins from the glycolytic or glyoxylate cycle often function as moonlighting enzymes [26]. *CitF*, encoding isocitrate lyase, is a candidate for such a role.

Interestingly, a number of synonymous mutations were also identified (Tables 2, 3, 4 and 5). Synonymous mutations are now recognized to have a range of, mostly deleterious, effects on cells, including altering mRNA structure [27, 28], rate of translation [29], protein folding [30], and fitness [31]. So far, synonymous mutations

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Table 2 Mutations associated with resistance to amoxicillin in strains with a previously acquired resistance to enrofloxacin, kanamycin, or tetracycline

Gene product	Gene	Mutation	Pop. frequency	Strains
Transcriptional regulator	acrR	L204R, Del C205	0.98–1	E _{1/4} , K _{2/4} T _{2/4}
B-lactamase	ampC	C35A (promoter)	1	E _{4/4,} K _{3/4,} T _{1/4}
		C-11 T (promoter)	1	E _{4/4,} K _{3/4,} T _{1/4}
		Ins <u>T</u> after <u>C4379837</u> (promoter)	1	K _{1/4}
		Ins <u>T</u> after <u>A4379015</u> (promoter)	1	T _{1/4}
Citrate lyase	citF	G432A	1	E _{4/4} , K _{3/4} , T _{1/}
				4
Glutamate synthase subunit	gltD	A80A	0.18	T _{1/4}
L-glyceraldehyde 3-phosphate reductase	gpr	L180F	1	T _{2/4}
CPS-53 glucosyl transferase	gtrS	V351 V	0.96	K _{1/4}
Acetolactate synthase/acetohydroxybutanoate synthase subunit	ilvH	G52C	0.60–1	E _{2/4} , K _{2/4} , T _{1/}
Branched chain amino acid/phenylalanine ABC transporter membrane subunit	livH	G239R	0.18	E _{1/4}
Intermembrane phospholipid transport system subunit	mlaF	Ins AIT between T38-A39	0.65-0.66	T _{2/4}
Large conductance mechanosensitive channel	mscL	L45Q	0.98	K _{1/4}
Mismatch repair protein	mutL	W390 L	0.11-0.2	T _{2/4}
Transcriptional regulator	ompR	E88A	1	E _{4/4} , K _{3/4} , T _{1/}
RNA polymerase subunit α	rpoA	C269S	1	E _{1/4} , K _{1/4}
RNA polymerase subunit β	гроВ	E564A	0.15	T _{1/4}
Sigma 70	rpoD	D445V	1	E _{4/4}
Alternative sigma factor σS	rpoS	E16Stop	0.96-1	K _{2/4} , T _{1/4}
		Del Q59-E330	0.91-1	T _{2/4}
Potassium ion transporter	trkG	G159D	1	K _{1/4}

Del = deletion, Ins = insertion, underlined letters and numbers indicate nucleotides and their genomic position. Population frequency indicates the mutational frequency in the sequenced population. Strains indicates the prevalence of a mutation within different replicates. E: population with previously acquired resistance to enrofloxacin, K: population with previously acquired resistance to tetracycline. X/Y: X indicates number of replicates with particular mutation, Y indicates total number of replicates that were sequenced. Mutations associated with resistance in a wild-type background are shown in **bold**. Mutations associated with the previous resistance have been omitted from the table

contributing to antibiotic resistance development have been described in TEM-1 β -lactamase [32], gyrA, and rpoB [33].

Not all mutations identified in the resistant strains are necessarily functionally relevant. Through genetic hitch-hiking, mutations can increase in frequency because they are genetically linked to mutations that do provide a survival advantage under antibiotic pressure [34, 35].

Most resistance mutations in bacteria confer a fitness cost [36], which can be reduced by the appearance of compensatory mutations [37] or additional resistance mutations [38]. *E. coli* with acquired resistance to amoxicillin, enrofloxacin, or tetracycline is able to maintain the level of resistance when cultured under the same conditions in the absence of antibiotics [5]. It is therefore likely that some of the mutations observed in this dataset do not

confer resistance themselves, but compensate for the fitness cost associated with resistance mutations. In general, mutations observed in this data set either interfere directly with the target-drug interaction, or they facilitate development of resistance by adjusting other cellular processes and thereby compensating for metabolic costs.

The observation that not all mutations associated with a particular resistance in wild-type *E. coli* appeared during secondary adaptation to the same antibiotic confirms that the effect of a single mutation on phenotype or fitness can depend on the genetic background [11, 39]. For some mutations that do not appear during secondary adaptation, the associated fitness cost could be too high in that particular genetic background. Alternatively, the mutation

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Table 3 Mutations associated with resistance to enrofloxacin in strains with a previously acquired resistance to amoxicillin, kanamycin, or tetracycline

Gene product	Gene	Mutation	Pop. frequency	Strains
Multidrug efflux pump membrane fusion lipoprotein/transcriptional regulator	acrA/acrR*	5' UTR (Del <u>485,695–711</u>)	1	T _{1/3}
Transcriptional regulator	acrR	Del <u>485,885–11</u> (FS42)	1	A _{3/4} , K _{1/4} , T _{2/3}
Aconitate hydratase B/2-methylisocitrate dehydratase	acnB	G362 V, Del L363-V865	1	T _{1/3}
Alkyl hydroperoxide reductase	ahpC	Del F38	0.40-1	A _{2/4} , T _{1/3}
Crotonobetainyl-CoA reductase	caiA	C154A, F155P, I156T, T157P, S158R, S159 T, A160S, Y161W, Del T162-R380	1	A _{2/4} , T _{1/4}
ATP-dependent helicase	dinG	Del Q641-R716	0.89	A _{1/4}
Glutamate synthase subunit	gltB	5' UTR (<u>C3354487T</u>)	1	$K_{1/4}, T_{2/3}$
DNA gyrase subunit A	gyrA	S83 L	1	A _{4/4} , E _{1/4} , T _{3/3}
		D87G	1	A _{3/4} , K _{2/4} , T _{1/3}
		D87N	1	$K_{2/4}, T_{2/3}$
Branched chain amino acid/phenylalanine ABC transporter membrane subunit	livF	G239F	0.87	T _{1/4}
Mismatch repair protein	mutL	W390 L	0.09-0.14	K _{1/4} , T _{2/3}
Phosphorelay protein	npr	L23R	1	T _{1/3}
Phenylacetate degradation protein	paaD	A8S	1	T _{1/3}
Putative 2,3-dehydroadipyl-CoA hydratase	рааҒ	V35 L	1	T _{1/3}
DNA topoisomerase IV subunit A	parC	S80R	0.94-1	A _{3/4} , K _{4/4} , T _{3/3}
DNA topoisomerase IV subunit B	parE	Ins Q458	0.97-1	A _{3/4} , K _{2/4} , T _{2/3}
		E459Q	0.03	A _{1/4}
		S425P	0.81	K _{2/4}
		Ins S after A457	0.95	T _{1/3}
tRNA-Phe	pheU	5' UTR (<u>G4362633 T</u>)	1	K _{1/4}
Cryptic proline/histidine transporter	proY	A235S	1	A _{3/4} , K _{1/4} , T _{2/3}
Pyridine nucleotide transhydrogenase subunit β	ptnB	V434 V	0.57	K _{1/4}
Phosphate acetyltransferase	pta	Del T431-Q714	0.56	K _{1/4}
Sensory histidine kinase	rcsC	E44K	1	K _{2/4}
Truncated RNase PH	rph	Del R676	1	$A_{2/4}, T_{1/3}$
S10 subunit of 30S	rpsJ	5' UTR (<u>G3453306 T</u>)	1	$K_{1/4}, T_{2/3}$
Bifunctional (p) ppGpp synthase/hydrolase	spoT	Ins C after <u>T3815879</u> (FS223)	1	$A_{2/4}, T_{1/4}$
Transcriptional regulator	soxR	T113S, del 4,277,899-903 (FS14)	1	A _{3/4} , K _{1/4} , T _{1/3}
		Ins G136	0.15	K _{1/4}
		L139F, del T after <u>C4277882</u>	0.85	K _{1/4}
		A146E	1	K _{1/4}
		Del R139-N154	1	K _{1/4}
Succinyl-CoA synthetase subunit β	sucC	Del A218-N221	0.51	K _{1/4}
DNA-binding transcriptional activator	ybaO	T33P	1	T _{1/4}
Putative RNA binding protein	yciO	N64Y	1	A _{1/4}
Putative transport protein/Putative inner membrane protein	ybjL/ybjM*	5' UTR (A889923G)	1	A _{3/4} , K _{1/4} , T _{2/3}

Del = deletion, Ins = insertion, underlined letters and numbers indicate nucleotides and their genomic position, * indicates that mutation could affect either gene. Population frequency indicates the mutational frequency in the sequenced population. Strains indicates the prevalence of a mutation within different replicates. E: population with previously acquired resistance to enrofloxacin, K: population with previously acquired resistance to kanamycin, T: population with previously acquired resistance to tetracycline. X/Y: X indicates number of replicates with particular mutation, Y indicates total number of replicates that were sequenced. Mutations associated with resistance in a wild-type background are shown in **bold**. Mutations associated with the previous resistance have been omitted from the table

might not be necessary for resistance. Although no cross-resistance between resistant strains from this study has been found [8], the different genetic background itself could contribute to the ability to

become resistant to a second antibiotic. Another explanation could be that the particular mutation is a compensatory mutation that is no longer necessary because of the different genetic background [38].

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Table 4 Mutations associated with resistance to kanamycin in strains with a previously acquired resistance to amoxicillin, enrofloxacin, or tetracycline

Gene product	Gene	Mutation	Pop. frequency	Strains
Multidrug efflux pump RND permease	астВ	V139F	0.73–1	E _{2/3}
Inner membrane magnesium ion transporter	corA	Del V264-F266	0.93	T _{1/4}
Sensor protein of Cpx two-component system	срхА	Q242L	1	E2/3
Cytochrome bo3 ubiquinol oxidase subunit 1	суоВ	G662C	0.86-0.93	E _{2/3}
		Del G448658 (FS661)	0.13	E _{1/3}
		Start1L	0.12	E _{1/3}
Cytochrome bo3 ubiquinol oxidase subunit 3	cyoC	Start1I	0.86-0.93	E _{2/3}
		Del C448658 (FS1)	0.13	E _{1/3}
		N661Stop	0.12	E _{1/3}
Formate dehydrogenase O subunit α	fdoG	V226 L	0.36-0.58	A _{2/3}
Elongation factor G	fusA	T393I	1	A1/3
		F605 L	1	E3/3, T3/4
		P610L	0.92-0.97	A2/3
		P610T	0.96	T1/4
sensory histidine kinase	kdpD	Q728P	0.36-0.61	A _{1/3} , T _{1/4}
tRNA-Leu	leuP	V16 V	0.67	A _{1/3}
Outer membrane lipoprotein	mlaA	Q728P	0.88	T _{1/4}
		Del F42-N43	0.92	T _{1/4}
2-O-α-mannosyl-D-glycerate specific PTS permease	mngA	F431 L	0.28-0.59	A _{2/3}
Mismatch repair protein	mutL	W390 L	0.11-0.13	A _{1/4} , T _{2/4}
Iron-sulfur cluster carrier protein	nfuA	Ins 8 nt after <u>G3546085</u> (FS154)	0.23	A _{1/3}
NADH:quinone oxidoreductase subunit CD	nuoC	Del R471-L474	0.93	E _{1/3}
NADH:quinone oxidoreductase subunit G	nuoG	Ins C after A2399396 (FS257)	0.4	A _{1/3}
Murein/oligopeptide ABC transporter subunit	oppF	L888Stop	0.74-1	E _{2/3}
		Ins SIQ after L187	0.14	E _{1/3}
Aminopeptidase N	pepN	V809E	0.57	A _{1/3}
RNA polymerase subunit β^\prime	rpoC	G367C	0.16	E _{1/3}
ATP binding protein of putrescine ABC exporter	sapF	L181Q	1	T2/4
Peptide antibiotic transporter	sbmA	Ins <u>G</u> after <u>T393897</u> (FS88)	0.99	E _{1/3}
		W98Stop	0.44	A _{1/3}
		Y162Stop	0.11	T _{1/4}
		W179Stop	0.53-0.60	T _{2/4}
		S250Stop	0.44	A _{1/3}
		L369 L	0.14	T _{2/4}
		1370L	0.14	T _{2/4}
Uncharacterized protein	ygaQ	Q417K	1	E _{2/3}
Uncharacterized protein	урјС	5' UTR (<u>T2785462C</u>)	1	E _{2/3}
Putative transporter	yqeG	P282P	0.22	E _{1/3}
Putative fimbrial usher protein	yraJ	Q614L	0.18	T _{1/4}
		H615H	0.19	T _{1/4}

Del = deletion, Ins = insertion, underlined letters and numbers indicate nucleotides and their genomic position. Population frequency indicates the mutational frequency in the sequenced population. Strains indicates the prevalence of a mutation within different replicates. A: population with previously acquired resistance to amoxicillin, E: population with previously acquired resistance to enrofloxacin, T: population with previously acquired resistance to tetracycline. X/Y: X indicates number of replicates with particular mutation, Y indicates total number of replicates that were sequenced. Mutations associated with resistance in a wild-type background are shown in **bold**. Mutations associated with the previous resistance have been omitted from the table

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Table 5 Mutations associated with resistance to tetracycline in strains with a previously acquired resistance to amoxicillin, enrofloxacin, or kanamycin

Gene product	Gene	Mutation	Pop. frequency	Strains
Multidrug efflux pump membrane fusion lipoprotein/multidrug efflux pump RND permease		5' UTR (Del <u>485,695–711)</u>	0.77	A _{1/4}
Multidrug efflux pump RND permease	acrB	V139F	1	A _{1/4} , E _{4/4}
		S665A	1	A _{1/4}
Transcriptional regulator	acrR	145L	0.97	K _{1/1}
		P85Q	1	A _{3/4} , K _{1/4}
		E130K	1	K _{1/1}
		R442Q	1	K _{1/1}
Crotonobetainyl-CoA reductase	caiA	C154A, F155P, I156T, T157P, S158R, S159 T, A160S, Y161W, Del T162-R380	1	E _{1/4}
Endochitinase	chiA	W702R	1	K _{1/1}
Membrane associated sensor kinase	envZ	P247S	0.24-0.95	A _{1/4} , K _{1/1}
3-ketoacyl-CoA thiolase	fadl	L246 L	1	E _{2/4}
Peptidoglycan DD-transpeptidase	ftsI	A513S	1	K _{1/1}
DNA-binding transcriptional repressor	marR	Ins <u>C</u> after <u>A1619472</u> (FS118)	1	A _{1/4}
ABC transporter family protein	mdlB	A16A	1	A _{2/4}
	mlaA	Del F42-N43	0.93	K _{1/1}
Murein tripeptide ABC transporter periplasmic binding protein	тррА	F244F	1	K _{1/1}
Mismatch repair protein	mutL	W390 L	0.1	E _{1/4}
Outer membrane porin F	ompF	Ins 7 nt after <u>G986771</u> (FS71)	0.04	E _{1/4}
Truncated RNase PH	rph	Del R676	1	E _{1/4}
RNA polymerase subunit β^\prime	rpoC	Del T208-K213	1	E _{1/4}
DNA-binding transcriptional dual regulator	soxR	T145S, G146T, A147L, R148A, L149G, L150R, Del E151-N155	1	E _{1/4}
DNA-binding transcriptional dual regulator	soxS	S2F	0.98-1	E _{3/4}
Bifunctional (p) ppGpp synthase/hydrolase	spoT	Ins <u>C</u> after <u>T3815879</u> (FS223)	1	E _{1/4}
DNA-binding transcriptional activator	ybaO	T33P	1	E _{1/4}

Numbers shown indicate frequency of mutation in population. *Del* = deletion, *Ins* = insertion, underlined letters and numbers indicate nucleotides and their genomic position. Population frequency indicates the mutational frequency in the sequenced population. Strains indicates the prevalence of a mutation within different replicates. A: population with previously acquired resistance to amoxicillin, E: population with previously acquired resistance to enrofloxacin, K: population with previously acquired resistance to kanamycin. X/Y: X indicates number of replicates with particular mutation, Y indicates total number of replicates that were sequenced. Mutations associated with resistance in a wild-type background are shown in **bold**. Mutations associated with the previous resistance have been omitted from the table

The mutations identified in strains exposed to a second antibiotic do not appear to be random, as the population frequency is high (Tables 2, 3, 4 and 5) and mutations appear in replicate strains (Fig. 2). The higher number of mutations could be necessary to compensate for the mutations already present in the resistant strains, or could represent alternative pathways to adaptation to the second antibiotic. In addition, a much higher degree of variability was displayed during secondary adaptation, both in the population frequencies of the mutations (Tables 2, 3, 4 and 5), as well as in the incidence of mutations among replicates (Tables 2, 3, 4 and 5). This confirms that, in addition to the different selective strength [40, 41], the genetic background can influence the mutational trajectories available [11, 42].

In an antibiotic-free environment, strains with acquired resistance to amoxicillin, enrofloxacin, and tetracycline maintain this resistance for at least two weeks after daily passaging [5]. When transferred to an environment with a different antibiotic, depending on the antibiotic combination, the original resistance is lost, partially maintained or fully preserved (Fig. 4). This suggests that the cellular adjustments necessary for adaptation to a new antibiotic are not always compatible with the earlier acquired mutations, which can have implications for the de novo development of multidrug resistance.

During resistance development, adaptation to one antibiotic can result in an increased or decreased susceptibility to other antibiotics, a concept known as collateral Hoeksema et al. BMC Genomics (2019) 20:284 Page 10 of 14

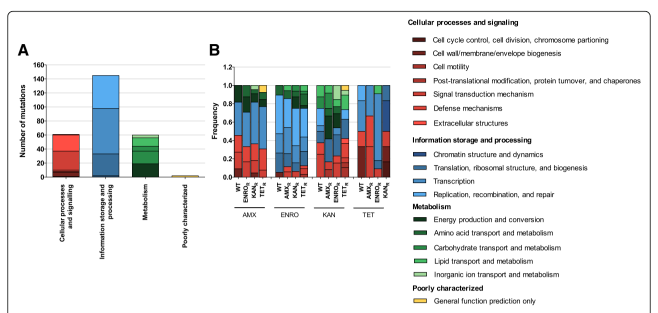


Fig. 3 Functional annotation of mutated genes associated with antibiotic resistance. All genes containing a mutations were classified using cluster of orthologous groups (COG) classifications and grouped by the four main categories (**a**) or by resistant strain (**b**). The genes were classified into COG categories using IMG/M (https://img.jgi.doe.gov)

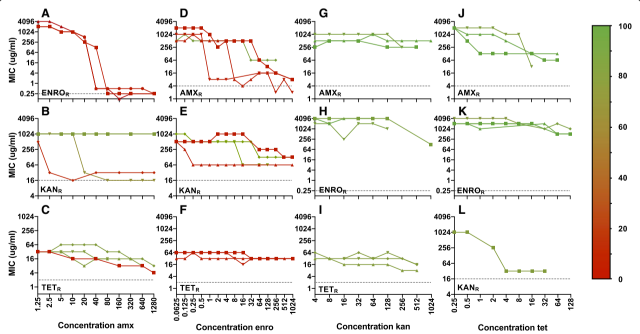


Fig. 4 MIC values over time in resistant strains exposed to a second antibiotic. Plots show MIC (ug/ml) of the first de novo acquired resistance, indicated on the X-axes, when exposed to amoxicillin (amx, a-c), enrofloxacin (enro, d-f), kanamycin (kan, g-i), or tetracycline (tet, j-l). The dotted black lines indicate the MICs for wild-type *E. coli* MG1655. Different lines in each plot indicate the different replicates. The percentage of residual mutations was determined by dividing the number of original mutations still present at the end of the experiment by the number of mutations observed in the beginning and multiplying this number by 100. The percentages are indicated in the plots using colors based on the gradient shown on the right

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sensitivity or collateral resistance. This is a concept often employed to develop strategies to limit resistance development [43–47], but is not observed in the resistant strains generated in this study [8] and can therefore not explain the drug-specific effect on loss or maintenance of resistance. Similar experiments in *Pseudomonas aeruginosa* show a very similar pattern on resensitization of antibiotic resistant strains, where the effect clearly seems to depend on the combination of the antibiotics, but no explanation is available as of yet [48] .

The loss of resistance is not always correlated with the loss of the associated mutations. Resistance to kanamycin and tetracycline (Fig. 4) appears to be partially maintained, but no mutations associated with resistance could be identified in those samples. The residual resistance could be explained by the genetic background contributing to resistance, such as *gyrB* during adaptation to kanamycin and enrofloxacin, or *acrR* during adaptation to enrofloxacin or tetracycline, although no cross-resistance between resistant strains has been observed [8]. Alternatively, adaptation on a gene expression level could contribute to residual levels of resistance.

Conclusion

The acquisition of drug resistance involves changes at various levels of cellular organization, including the genome, transcripts, and metabolites, highlighting that a complex interaction network is involved [49]. The data presented in this study support this notion. Development of resistance in strains with a previously acquired resistance does not result in the accumulation of the same mutations observed during primary adaptation in wild-type *E. coli*. This observation suggests that the genetic background of the strain itself plays an important role in resistance development. Moreover, mutations alone cannot always fully explain the resistance pattern observed, supporting the earlier suggested role for adaptation on the gene expression level in de novo acquisition of antibiotic resistance [50].

Methods

Bacterial strains, growth media, antibiotics, and MIC measurement

In all experimental tracks inducing resistance by exposing cells to step-wise increasing concentrations of the antibiotic [5], *E. coli* MG1655 wild-type strains were used as starting point. The choice of starting concentrations was based on MIC measurements.

Evolution experiments were performed as follows: wild-type *E. coli* was exposed to either $1.25 \,\mu\text{g/ml}$ amoxicillin, $0.0625 \,\mu\text{g/ml}$ enrofloxacin, $4 \,\mu\text{g/ml}$ kanamycin, or $0.5 \,\mu\text{g/ml}$ tetracycline. Cells were grown overnight at $37 \,^{\circ}\text{C}$, and OD_{600} were measured after $24 \,\text{h}$. When the OD_{600} of the exposed culture was at least

75% of the ${\rm OD_{600}}$ of a reference culture that was not exposed to antibiotics, the population was considered adapted and the antibiotic concentration was doubled. Using the exposed population, new flasks were re-inoculated to a final ${\rm OD_{600}}$ of 0.1, with the flasks with the lower concentration now serving as the reference culture. The initial adaptation to each antibiotic was performed in duplicate.

After development of resistance to one antibiotic, two duplicates of each duplicate strain were used for exposure to a second antibiotic, resulting in four strains with an identical exposure history, as depicted in Fig. 1. To re-start the evolution experiments, flasks were re-inoculated with one of the resistant strains to a final ${\rm OD}_{600}$ of 0.1. Strains with acquired resistance to one or two antibiotics were compared [8]. When a population was considered adapted, glycerol stocks were made and stored at -80 °C. When relevant, cells from glycerol stocks were plated on LB agar and grown in Evans medium with the appropriate antibiotic.

Stock solutions of amoxicillin, enrofloxacin, kanamycin, or tetracycline (10 mg/mL) were filter sterilized and stored at $4\,^{\circ}\mathrm{C}$ for maximally 2 weeks. Minimal inhibitory concentrations (MICs) were determined as described before [51]. Measurements were performed in 96-well ThermoScientific Multiskan FC spectrophotometer plate readers, shaken and at 37 °C in a final volume of 150 μ l with a starting OD $_{595}$ of 0.05. Antibiotic concentrations increasing by a factor of 2 and ranging from 0.0625 to 4096 μ g/ml were applied. The lowest concentrations that limited final OD after 23 h to 0.2 or less was reported as MIC.

Whole genome sequencing

Glycerol stocks of selected strains were plated on LB agar, and over 100 single colonies were combined to grow a liquid culture (Evans minimal medium) in the presence of the antibiotic. The culture was still growing when samples were taken to freeze at -80 °C. We then used roughly 109 cells (corresponding to 2 mL of a culture with an OD600 of 1) for genomic DNA isolation. Genomic DNA was isolated using the DNeasy blood and tissue kit (Qiagen). DNA was isolated from the wild-type, the duplicates of the single exposure strains and all four replicates of the double exposure strains. A wild-type E. coli MG1655 was submitted to the same protocol and used as control. In this control two point mutations were observed when compared to the MG1655 reference strain. These point mutations were observed in all strains and are not reported.

gDNA libraries were generated according to the manufacturers' protocols using the Ion Xpress™ Plus gDNA Fragment Library Preparations (Thermo Fisher Scientific). Bar-coded libraries were prepared according to the

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Ion Plus fragment library kit (Thermo Fisher Scientific) and the Ion Xpress DNA bar-coding kit (Thermo Fisher Scientific) according to the 200-base-read Ion Proton libraries instructions of the manufacturer. The size distribution and yield of the barcoded libraries were assessed using the 2200 TapeStation System with Agilent High Sensitivity D1000 ScreenTapes (Agilent Technologies). Sequencing templates were prepared on the Ion Chef System using the Ion PI Hi-Q Chef Kit (Thermo Fisher Scientific). Sequencing was performed on an Ion Proton System using a Ion PI v3 chip (Thermo Fisher Scientific) according to the instructions of the manufacturer.

For quality control procedures, the FASTQ files of individual samples were assessed with fastqc (http://www. bioinformatics.babraham.ac.uk/projects/fastqc/). Several quality metrics (sequencing depth, read length distribution, read quality distribution, mean read quality along the read, base frequency at each read position) were compared across samples in relation to the experimental factors using in-house software based on samtools and R (https://www.r-project.org/). To map all accepted reads to the E. coli K-12 MG1655 reference genome, Tmap was applied (https://github.com/iontorrent/TMAP/blob/ master/doc/tmap-book.pdf). The Ion Proton system generates sequencing reads of variable lengths, and Tmap combines a short read algorithm [52] and long read algorithms [52, 53] in a multistage mapping approach. The genes were classified into COG categories using IMG/M (https://img.jgi.doe.gov).

The average sequencing depth was 226. Torrent Variant Caller (Thermo Fisher Scientific) was used to identify deviations, such as single-nucleotide polymorphisms, insertions, and deletions from the reference genome. The Torrent Variant Caller is based on Freebayes [54] and capable of somatic variant calling. Overall, parameters such as the minimum phred-scaled call quality, the minimum coverage, and the maximum strand bias, were set such that variants were called with relatively high reliability, at the cost of sensitivity. On the other hand, the minimum observed allele frequency required for a non-reference variant call was set relatively low (5%) to enable detection of low frequency events in the bacterial populations. Mutation frequency was calculated as the ratio of the number of reads containing a genetic variation to the overall read number. To differentiate between sequencing errors and true mutations, genetic deviations were excluded if either of the following conditions applied: 1) the mutations appeared in a homopolymer region, or 2) the Phred quality score was < 30 and the same mutation was not identified in a different sample with a Phred quality score of ≥30. Mutations also present in the wild-type were excluded from the analysis.

Sequencing reads were mapped directly to the MG1655 genome to enable detection of mutations and short indels. For the detection of long inserts the single nucleotide polymorphism (SNP) calling algorithm was used. Locations with a high number of called mutations were assumed to be alignment artifacts due to inserts in the genome under study and were detected with the cn.mops package [55]. The regions containing these alignment artefacts were subsequently amplified with PCR and sequenced by Sanger sequencing to achieve a more reliable analysis than the alternative, de novo assembly of the entire genome, would allow [12].

Abbreviations

COG: clusters of orthologous groups; MIC: minimal inhibitory concentration; ROS: reactive oxygen species; SNP: single nucleotide polymorphism

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Availability of data and materials

The binary alignment/map (bam) files of the sequenced strains have been deposited in the NCBI database and can accessed at https://www.ncbi.nlm.nih.gov/Traces/study/?acc=PRJNA489357

Authors' contributions

MH and BtK conceived the project. MH performed experiments. MJ performed the bioinformatic analysis. MH and BtK wrote the manuscript. SB contributed to the final manuscript. All authors critically reviewed the manuscript and approved the final version.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Author details

¹Laboratory for Molecular Biology and Microbial Food Safety, Swammerdam Institute for Life Sciences, University of Amsterdam, Amsterdam, The Netherlands. ²RNA Biology & Applied Bioinformatics, Swammerdam Institute for Life Sciences, University of Amsterdam, Amsterdam, The Netherlands. ³Netherlands Food and Consumer Product Safety Authority, Office for Risk Assessment, Utrecht, The Netherlands.

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References

 Handel N, Schuurmans JM, Feng Y, Brul S, ter Kuile BH. Interaction between mutations and regulation of gene expression during development of de novo antibiotic resistance. Antimicrob Agents Chemother. 2014;58(8):4371–9. Hoeksema et al. BMC Genomics (2019) 20:284 Page 13 of 14

- Bagel S, Hullen V, Wiedemann B, Heisig P. Impact of gyrA and parC mutations on quinolone resistance, doubling time, and supercoiling degree of Escherichia coli. Antimicrob Agents Chemother. 1999;43(4):868–75.
- Vogwill T, MacLean RC. The genetic basis of the fitness costs of antimicrobial resistance: a meta-analysis approach. Evol Appl. 2015;8(3):284–95.
- Maisnier-Patin S, Andersson DI. Adaptation to the deleterious effects of antimicrobial drug resistance mutations by compensatory evolution. Res Microbiol. 2004;155(5):360–9.
- van der Horst MA, Schuurmans JM, Smid MC, Koenders BB, ter Kuile BH.
 De novo acquisition of resistance to three antibiotics by Escherichia coli.
 Microb Drug Resist. 2011;17(2):141–7.
- Kohanski MA, Dwyer DJ, Hayete B, Lawrence CA, Collins JJ. A common mechanism of cellular death induced by bactericidal antibiotics. Cell. 2007;130(5):797–810.
- Dwyer DJ, Belenky PA, Yang JH, MacDonald IC, Martell JD, Takahashi N, et al. Antibiotics induce redox-related physiological alterations as part of their lethality. Proc Natl Acad Sci. 2014;111(20):F2100–9.
- Hoeksema M, Brul S, Ter Kuile BH. Influence of reactive oxygen species on De novo Acquisition of Resistance to bactericidal antibiotics. Antimicrob Agents Chemother. 2018;62(6):e02354–17.
- Knopp M, Andersson DI. Predictable phenotypes of antibiotic resistance mutations. mBio. 2018;9(3):e00770–18.
- Durao P, Trindade S, Sousa A, Gordo I. Multiple resistance at no cost: rifampicin and streptomycin a dangerous liaison in the spread of antibiotic resistance. Mol Biol Evol. 2015;32(10):2675–80.
- Weinreich DM, Watson RA, Chao L. Perspective: sign epistasis and genetic constraint on evolutionary trajectories. Evolution. 2005;59(6):1165–74.
- Hoeksema M, Jonker MJ, Bel K, Brul S, Ter Kuile BH. Genome rearrangements in Escherichia coli during de novo acquisition of resistance to a single antibiotic or two antibiotics successively. BMC Genomics. 2018;19(1):973.
- Corvec S, Caroff N, Espaze E, Marraillac J, Reynaud A. 11 mutation in the ampC promoter increasing resistance to beta-lactams in a clinical Escherichia coli strain. Antimicrob Agents Chemother. 2002;46(10):3265–7.
- Ruiz J. Mechanisms of resistance to quinolones: target alterations, decreased accumulation and DNA gyrase protection. J Antimicrob Chemother. 2003;51(5):1109–17.
- Koutsolioutsou A, Pena-Llopis S, Demple B. Constitutive soxR mutations contribute to multiple-antibiotic resistance in clinical Escherichia coli isolates. Antimicrob Agents Chemother. 2005;49(7):2746–52.
- Vinue L, Hooper DC, Jacoby GA. Chromosomal mutations that accompany qnr in clinical isolates of Escherichia coli. Int J Antimicrob Agents. 2018;51(3):479–83.
- Karczmarczyk M, Martins M, Quinn T, Leonard N, Fanning S. Mechanisms of fluoroquinolone resistance in Escherichia coli isolates from food-producing animals. Appl Environ Microbiol. 2011;77(20):7113–20.
- Acharya S, Foster PL, Brooks P, Fishel R. The coordinated functions of the E. Coli MutS and MutL proteins in mismatch repair. Mol Cell. 2003;12(1):233–46.
- Taddei F, Matic I, Godelle B, Radman M. To be a mutator, or how pathogenic and commensal bacteria can evolve rapidly. Trends Microbiol. 1997;5(11):427–8 discussion 8-9.
- LeClerc JE, Li B, Payne WL, Cebula TA. High mutation frequencies among Escherichia coli and Salmonella pathogens. Science (New York, NY). 1996;274(5290):1208–11.
- 21. Marinus MG. DNA methylation and mutator genes in Escherichia coli K-12. Mutat Res. 2010;705(2):71–6.
- Caroff N, Espaze E, Gautreau D, Richet H, Reynaud A. Analysis of the effects of -42 and -32 ampC promoter mutations in clinical isolates of Escherichia coli hyperproducing ampC. J Antimicrob Chemother. 2000; 45(6):783-8.
- Rinder H, Thomschke A, Rusch-Gerdes S, Bretzel G, Feldmann K, Rifai M, et al. Significance of ahpC promoter mutations for the prediction of isoniazid resistance in mycobacterium tuberculosis. Eur J Clin Microbiol Infect Dis. 1998;17(7):508–11.
- Gruber TM, Gross CA. Multiple sigma subunits and the partitioning of bacterial transcription space. Annu Rev Microbiol. 2003;57:441–66.
- 25. Jeffery CJ. An introduction to protein moonlighting. Biochem Soc Trans. 2014;42(6):1679–83.
- Henderson B, Martin A. Bacterial virulence in the moonlight: multitasking bacterial moonlighting proteins are virulence determinants in infectious disease. Infect Immun. 2011;79(9):3476–91.

- 27. Goodman DB, Church GM, Kosuri S. Causes and effects of N-terminal codon bias in bacterial genes. Science (New York, NY). 2013;342(6157):475–9.
- Kudla G, Murray AW, Tollervey D, Plotkin JB. Coding-sequence determinants of gene expression in Escherichia coli. Science (New York, NY). 2009; 324(5924):255–8.
- Sorensen MA, Pedersen S. Absolute in vivo translation rates of individual codons in Escherichia coli. The two glutamic acid codons GAA and GAG are translated with a threefold difference in rate. J Mol Biol 1991;222(2):265–280.
- Zhang G, Hubalewska M, Ignatova Z. Transient ribosomal attenuation coordinates protein synthesis and co-translational folding. Nat Struct Mol Biol. 2009;16(3):274–80.
- 31. Firnberg E, Labonte JW, Gray JJ, Ostermeier M. A comprehensive, high-resolution map of a gene's fitness landscape. Mol Biol Evol. 2014;31(6):1581–92.
- Schenk MF, Szendro IG, Krug J, de Visser JA. Quantifying the adaptive potential of an antibiotic resistance enzyme. PLoS Genet. 2012;8(6): e1002783.
- Lai CC, Chen CC, Lu YC, Chuang YC, Tang HJ. The clinical significance of silent mutations with respect to ciprofloxacin resistance in MRSA. Infect Drug Resist. 2018;11:681–7.
- 34. Kim Y, Stephan W. Detecting a local signature of genetic hitchhiking along a recombining chromosome. Genetics. 2002;160(2):765–77.
- Lang GI, Rice DP, Hickman MJ, Sodergren E, Weinstock GM, Botstein D, et al. Pervasive genetic hitchhiking and clonal interference in forty evolving yeast populations. Nature. 2013;500(7464):571–4.
- Melnyk AH, Wong A, Kassen R. The fitness costs of antibiotic resistance mutations. Evol Appl. 2015;8(3):273–83.
- Maisnier-Patin S, Berg OG, Liljas L, Andersson Dl. Compensatory adaptation to the deleterious effect of antibiotic resistance in Salmonella typhimurium. Mol Microbiol. 2002;46(2):355–66.
- Trindade S, Sousa A, Xavier KB, Dionisio F, Ferreira MG, Gordo I. Positive epistasis drives the acquisition of multidrug resistance. PLoS Genet. 2009;5(7):e1000578.
- Wong A. Epistasis and the evolution of antimicrobial resistance. Front Microbiol. 2017;8:246.
- Wistrand-Yuen E, Knopp M, Hjort K, Koskiniemi S, Berg OG, Andersson DI. Evolution of high-level resistance during low-level antibiotic exposure. Nat Commun. 2018;9(1):1599.
- Mira PM, Meza JC, Nandipati A, Barlow M. Adaptive landscapes of resistance genes change as antibiotic concentrations change. Mol Biol Evol. 2015;32(10):2707–15.
- Borrell S, Teo Y, Giardina F, Streicher EM, Klopper M, Feldmann J, et al. Epistasis between antibiotic resistance mutations drives the evolution of extensively drug-resistant tuberculosis. Evol Med Public Health. 2013;2013(1):65–74.
- 43. Barbosa C, Trebosc V, Kemmer C, Rosenstiel P, Beardmore R, Schulenburg H, et al. Alternative evolutionary paths to bacterial antibiotic resistance cause distinct collateral effects. Mol Biol Evol. 2017;34(9):2229–44.
- Imamovic L, Sommer MO. Use of collateral sensitivity networks to design drug cycling protocols that avoid resistance development. Sci Transl Med. 2013;5(204):204ra132.
- Lazar V, Pal Singh G, Spohn R, Nagy I, Horvath B, Hrtyan M, et al. Bacterial evolution of antibiotic hypersensitivity. Mol Syst Biol. 2013;9:700.
- Munck C, Gumpert HK, Wallin AI, Wang HH, Sommer MO. Prediction of resistance development against drug combinations by collateral responses to component drugs. Sci Transl Med. 2014;6(262):262ra156.
- Rodriguez de Evgrafov M, Gumpert H, Munck C, Thomsen TT, Sommer MO. Collateral resistance and sensitivity modulate evolution of high-level resistance to drug combination treatment in Staphylococcus aureus. Mol Biol Evol. 2015;32(5):1175–85.
- Yen P, Papin JA. History of antibiotic adaptation influences microbial evolutionary dynamics during subsequent treatment. PLoS Biol. 2017;15(8):e2001586.
- Suzuki S, Horinouchi T, Furusawa C. Prediction of antibiotic resistance by gene expression profiles. Nat Commun. 2014;5:5792.
- Handel N, Schuurmans JM, Brul S, ter Kuile BH. Compensation of the metabolic costs of antibiotic resistance by physiological adaptation in Escherichia coli. Antimicrob Agents Chemother. 2013;57(8):3752–62.
- Schuurmans JM, Nuri Hayali AS, Koenders BB, ter Kuile BH. Variations in MIC value caused by differences in experimental protocol. J Microbiol Methods. 2009;79(1):44–7.

Hoeksema et al. BMC Genomics (2019) 20:284 Page 14 of 14

- 52. Li H, Durbin R. Fast and accurate short read alignment with burrows-wheeler transform. Bioinformatics. 2009;25(14):1754–60.
- 53. Ning Z, Cox AJ, Mullikin JC. SSAHA: a fast search method for large DNA databases. Genome Res. 2001;11(10):1725–9.
- 54. Garrison EM, Gabor. Haplotype-based variant detection from short-read sequencing. eprint arXiv. 2012:1207.3907v2.
- Klambauer G, Schwarzbauer K, Mayr A, Clevert DA, Mitterecker A, Bodenhofer U, et al. Cn. MOPS: mixture of Poissons for discovering copy number variations in next-generation sequencing data with a low false discovery rate. Nucleic Acids Res. 2012;40(9):e69.

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