Macrophage adaptation leads to parallel evolution of genetically diverse Escherichia coli small-colony variants with increased fitness in vivo and antibiotic collateral sensitivity

Ricardo S. Ramiro | Henrique Costa | Isabel Gordo

Abstract
Small-colony variants (SCVs) are commonly observed in evolution experiments and clinical isolates, being associated with antibiotic resistance and persistent infections. We recently observed the repeated emergence of Escherichia coli SCVs during adaptation to the interaction with macrophages. To identify the genetic targets underlying the emergence of this clinically relevant morphotype, we performed whole-genome sequencing of independently evolved SCV clones. We uncovered novel mutational targets, not previously associated with SCVs (e.g. cydA, pepP) and observed widespread functional parallelism. All SCV clones had mutations in genes related to the electron-transport chain. As SCVs emerged during adaptation to macrophages, and often show increased antibiotic resistance, we measured SCV fitness inside macrophages and measured their antibiotic resistance profiles. SCVs had a fitness advantage inside macrophages and showed increased aminoglycoside resistance in vitro, but had collateral sensitivity to other antibiotics (e.g. tetracycline). Importantly, we observed similar results in vivo. SCVs had a fitness advantage upon colonization of the mouse gut, which could be tuned by antibiotic treatment: kanamycin (aminoglycoside) increased SCV fitness, but tetracycline strongly reduced it. Our results highlight the power of using experimental evolution as the basis for identifying the causes and consequences of adaptation during host-microbe interactions.

KEYWORDS
antibiotic resistance, collateral sensitivity, experimental evolution, macrophages, mouse gut colonization, small-colony variants, whole-genome sequencing

1 | INTRODUCTION

Variation in colony morphology is often observed during in vitro bacterial adaptation (Miskinyte et al., 2013; Rainey & Travisano, 1998; Rozen & Lenski, 2000; Traverse, Mayo-Smith, Politak, & Cooper, 2013). One of the most commonly observed morphotypes are small-colony variants (SCVs) (e.g. Miskinyte et al., 2013; Rozen & Lenski, 2000; Traverse et al., 2013). SCVs emerge in response to multiple selective pressures, including adaptation to: single (Rozen & Lenski, 2000) or multiple carbon sources (Friesen, Saxer, Traviso, & Doebeli, 2004), biofilm conditions (Penterman et al., 2014; Traverse et al., 2013), the mouse gut (Lee et al., 2010), antibiotics (Musher, Baughn, Templeton, & Minuth, 1977; Yegian, Gallo, & Toll, 1959), antimicrobial peptides (Pranting & Andersson, 2010) or the intracellular environment of nonprofessional phagocytes (Cano, Pucciarelli, Martinez-Moya, Casadesus, & Garcia-del Portillo, 2003; Tuchscherr et al., 2011; Vesga...
et al., 1996). As could be expected, from such a diverse set of selective pressures, SCVs also present a diverse genetic basis (Cooper, Staples, Traverse, & Ellis, 2014; Herron & Doebeli, 2013; Pranting & Anders son, 2010; Proctor et al., 2006). The most common mutations are in genes related to the electron-transport chain (ETC), which are known to lead to the SCV phenotype in many species, both Gram-negative and Gram-positive (Cano et al., 2003; Dean, Olsen, Long, Rosato, & Musser, 2014; Kastbjerg, Hein-Kristensen, & Gram, 2014; Roggenkamp et al., 1998). Importantly, SCVs have been widely associated with antibiotic resistance, and for some of these species (e.g. S. aureus, E. coli, Pseudomonas aeruginosa), SCVs have been associated with chronic infections, such as infections of the lungs of cystic-fibrosis patients, urinary tract or prosthetics-association infections (Häussler, Tümmler, Weiβbrodt, Rohde, & Steinmetz, 1999; Proctor, van Langeveld, Kristjansson, Maslow, & Arbeit, 1995; Proctor et al., 2006; Roggenkamp et al., 1998; Rusthoven, Davies, & Lerner, 1979). In these natural infections, disentangling which selective pressures lead to particular phenotypes is often difficult. Interestingly, some of these SCVs show similar genetic/phenotypic bases to those which arise during evolution experiments (McDonald, Gehrig, Meintjes, Zhang, & Rainey, 2009; Proctor et al., 1995; Smith et al., 2006; Vesga et al., 1996), suggesting that experimental evolution may be useful to identify important selective pressures in vivo.

A recent aim in evolutionary biology is to understand whether evolution experiments can help understanding natural variation. A way to do this is to incorporate the selective pressures which are thought to be important in nature into evolution experiments (Bailey & Batail lon, 2016). In the context of infections and at the commensal-pathogen transition, a major selective pressure is the innate immune system, with macrophages (MΦs) being key in the response to microbial infections (Wynn, Chawla, & Pollard, 2013). Due to this role of MΦs, several studies have recently used experimental adaptation to understand how yeast or bacteria evolve under MΦ attack (Azevedo et al., 2016; Ensminger, Yassin, Miron, & Isberg, 2012; Miskinyte et al., 2013; Vázquez et al., 2014; Wartenberg et al., 2014). Recently, we used this approach to study the adaptation of a commensal E. coli to the interaction with MΦs (Miskinyte et al., 2013). In this experiment, replicate bacterial populations were allowed to evolve for 30 days in the presence of MΦs, while inhabiting both intracellular and extracellular environments (see Appendix S1 for summary of this experiment). Rapid and repeated emergence of SCVs occurred in independent bacterial populations, indicating that SCVs are an adaptive trait in the interaction with this key immune cell (Dettman et al., 2012; Elena & Lenski, 2003).

Here, we used whole-genome sequencing (WGS) of multiple clones to understand what is the genetic basis of SCVs and if the phenotypic parallelism is mediated by genetic parallelism. Our analysis shows evidence of parallelism from the gene to the pathway level and indicates widespread functional parallelism: all SCVs had mutations in genes related to the ETC, a common observation in clinically isolated SCVs (Proctor et al., 1995; Roggenkamp et al., 1998). Importantly, we identified several mutational targets that had not previously been associated with SCVs, including nuoG, hemY or ubiG. As is often observed for ETC-deficient SCVs (Proctor et al., 2006), we show that these SCV clones generally have a high reversion rate, are hemin auxotrophs, and generally reach a lower carrying capacity than the ancestral strain, both in poor and rich media. As SCVs emerged in the presence of MΦs, and such morphotypes have previously been shown to have increased antibiotic resistance, we tested for SCV fitness inside MΦs and for antibiotic resistance in vitro and upon colonization of the mouse gut. Our results show that SCVs can have an intracellular advantage inside MΦs, and that this morphotype has a pattern of antagonistic pleiotropy for antibiotic resistance in vitro: SCVs were more resistant to aminoglycosides but more sensitive to tetracycline, nalidixic acid and ciprofloxacin (quinolones), as well as cefotaxime (a β-lactam).

2 MATERIAL AND METHODS

2.1 Isolation of SCV clones, reference bacterial strains and MΦs

We isolated SCV clones from a previous experiment, in which E. coli was adapted to MΦs (see Appendix S1). These SCV clones were evolved from an ancestral strain of commensal E. coli (MC4100-CFP), which is streptomycin resistant (RpsL<sup>438</sup>) and constitutively expresses cyan-fluorescent protein (CFP; galK::CFP, Amp<sup>R</sup>). SCVs were identified in five of six independently evolving populations (Miskinyte et al., 2013; exception was population M4). Stocks of two randomly selected SCV clones (per population) were made and kept at −80°C. For population M4, two clones were also isolated. These clones produce colonies of similar size to the ancestral strain (see Fig. S1 for colony sizes) and were used as positive controls (i.e. clones that evolved with MΦs, but do not display a SCV phenotype) for the experiments below.

For competition assays, strain MC4100-YFP was used as reference strain. This differs from MC4100-CFP only in the expressed fluorescent protein. For the catalase activity test, a ΔµpO strain was used as a negative control (Al Mamun et al., 2012). The RAW 264.7 murine MΦ cell line was used to test for SCV intracellular survival (see Appendix S1 for MΦ-culture conditions).

2.2 Whole-genome sequencing and mutation prediction

Bacterial clones were grown in 10 ml RPMI at 37°C with 5% CO<sub>2</sub>. DNA was extracted using the procedure described by Wilson (2001). DNA library construction and sequencing was performed
by the IGC genomics facility. Each clone was paired-end sequenced on an Illumina MiSeq Benchtop-Sequencer. Standard procedures produced data sets of Illumina paired-end 250 bp read pairs, with mean coverage of 55× (32×–113× per clone). The genome sequencing data have been deposited in the NCBI Sequence Read Archive www.ncbi.nlm.nih.gov/sra (accession no.: SRP074322). bresq v0.20 was used to map reads and identify mutations, using the E. coli BW2952 genome as reference (NCBI accession: NC_012759.1; Deatherage & Barrick, 2014). Default settings were used, except for eliminating mutation predictions occurring in homopolymers of length >3. Mutations were determined by comparison with the ancestral (MC4100-CFP; previously sequenced for (Miskinyte et al., 2013). All predicted polymorphisms were manually inspected using IGV. Functional annotation clustering was performed using DAVID (Huang, Sherman, & Lempicki, 2008, 2009), with standard settings.

### 2.3 | Phenotypic traits of SCVs

Whole-genome sequencing revealed that SCVs had mutations in genes associated with the ETC. These SCVs often show the following phenotypes: (i) high reversion frequency to large colony phenotype; (ii) reduced catalase activity; (iii) slow growth in vitro; and (iv) auxotrophy for compounds involved in the ETC (Proctor et al., 2006). To test whether our SCVs had these phenotypes, we (i) measured reversion frequency (to large colony phenotype) and catalase activity as described in Miskinyte et al. (2013) and Roggenkamp et al. (1998) (see Appendix S1); (ii) estimated the relative growth rate and relative survival capacity of each SCV (relative to the ancestral) in rich (RPMI) and poor media (M9 mini medium); (iii) determined the auxotrophic requirements for eliminating mutation predictions occurring in homopolymers of length >3. Mutations were determined by comparison with the ancestral (MC4100-CFP; previously sequenced for (Miskinyte et al., 2013). All predicted polymorphisms were manually inspected using IGV. Functional annotation clustering was performed using DAVID (Huang, Sherman, & Lempicki, 2008, 2009), with standard settings.

### 2.4 | Bacterial survival inside MΦs

Three independently evolved clones carrying distinct mutations (28 and 41 as SCVs and 31 as a positive control; see Table 1) were competed against the YFP ancestral strain in the intracellular environment of MΦs. MΦs were seeded and activated, following a modification of Miskinsky & Gordo (2014) (see Appendix S1). On infection day, media were removed from each well and replaced by 1 ml RPMI with $10^7$ bacteria, a ~50:50 mixture of the YFP ancestral and one of the evolved (CFP) clones (three wells per clone-time point combination). Cultures were spun down (200 g, 5 min.) and plates incubated for 0.5 hr (37°C, 5% CO$_2$). After this period, media were removed from all wells and cells washed twice with RPMI to remove extracellular bacteria. RPMI with gentamicin was added to two sets of wells (100 μg/ml), to kill remaining extracellular bacteria and plates incubated for another 0.5 hr (i.e. total incubation time = 1 hr) or 5.5 hr (i.e. total = 6 hr). About 1 ml PBS was added to the remaining set of wells, and MΦs were detached and the contents of each well centrifuged to lyse MΦs (9400 g, 5 min). The pellet was then resuspended in 1 ml PBS. This allowed estimation of the number of bacteria that had been phagocytized by MΦs at 0.5 hr. At the 1 and 6 hr time-points, MΦs were washed, detached and centrifuged, as for the 0.5 hr timepoint. All timepoints were plated, and the number of YFP (ancestral) and CFP (SCVs) bacteria was used to calculate the competitive index (CI) as

$$CI = \log_{10} \left( \frac{\text{CFP}_{0.5hr}}{\text{YFP}_{0.5hr}} \right) \left( \frac{\text{YFP}_{xh}}{\text{CFP}_{xh}} \right)$$

where $xh$ represents 1- or 6-hr incubation, CFP (or YFP) represents the colony-forming units (cfu) at a particular time point. Six independent replicates were performed, per clone. We used gentamicin, a standard antibiotic for bacterial intracellular survival assays, due to its reduced ability to enter cells and kill intracellular bacteria (Cano et al., 2003; Carryn, Van Bambeke, Mingeot-Leclercq, & Tulkens, 2002; Nguyen et al., 2009). To ensure that the observed fitness effects are due to environmental conditions inside MΦs and not because gentamicin increases SCV relative fitness, we carried out in vitro competitions in the absence of MΦs. These mimic the competitions with MΦs: 50:50 mixtures ($10^7$ bacteria) of each SCV clone and the YFP ancestral were inoculated into RPMI (1 ml), with gentamicin at 100 μg/ml, at pH 5 (mimicking the MΦ intracellular environment) or 7.4 (standard for RPMI). Cultures were incubated for 6 hr and then plated. In three independent experiments (per SCV clone), SCV or ancestral colonies were only rarely observed, indicating that most bacteria (both SCVs and ancestral) cannot survive (Table S1). Moreover, we also carried competitions between isogenic CFP and YFP ancestral (Fig. S2) and observe that the CI was not significantly different from 0 at both 1 hr ($t_s = 0.43, p = .69$) and 6 hr ($t_s = 1.47, p = .22$), indicating that the fluorescent markers are neutral in this environment.

### 2.5 | In vitro antibiotic resistance assays

Antibiotic sensitivity was measured using disc diffusion assays (on LB-agar plates) for the following antibiotics: gentamicin (10 μg), kanamycin (30 μg), amikacin (30 μg), tobramycin (10 μg), netilmicin (10 μg), cefotaxime (30 μg), ciprofloxacin (5 μg), nalidixic acid (30 μg) and tetracycline (30 μg). The inhibition halo was measured 48 hr postincubation, at 37°C (3–8 independent replicates per clone-antibiotic combination).

### 2.6 | In vivo competition assays in the mouse gut

To test the competitive ability of SCVs in vivo, we used the streptomycin-treated mouse model of E. coli colonization (Conway, Kroghfeld, & Cohen, 2004). This model enables colonization by E. coli,
while maintaining a diverse microbiota (Barroso-Batista, Demengeot, & Gordo, 2015). Briefly, 8-week-old C57BL/6 female mice raised in specific pathogen-free conditions (ad libitum food), were given autoclaved water with streptomycin (5 g/L) for 7 days before and after colonization (total = 14 days). Mice were caged in pairs and gavaged with a suspension of $10^8$ bacteria (in 100 μl PBS; as in Barroso-Batista et al. (2014)), consisting of a mix of SCV-CFP and ancestral YFP (ancestral and SCV carry the streptomycin-resistance allele RpsL<sup>K43R</sup>; Trindade et al., 2009). The SCV population contained a mixture of equal numbers of the following genetically distinct clones: 1, 19, 26, 28, 41, 48 and 51, which cover all the mutations observed in our experiment (see Table 1). Groups of

### Table 1

Mutations identified in E. coli clones evolved in the presence of MΦs. Deletions (Δ), transposition of insertion sequences (IS) or small nucleotide insertions (+) are shown at the specific position of each mutation, within the gene. For SNPs, the particular nucleotide and corresponding amino acid replacement change is indicated. The deletion in clone 1 includes both intergenic and coding regions, starting at an intergenic position and ending within the cydA coding region. Genomic positions are indicated relative to the GenBank reference: NC_012759.1.

| Population | Day | Clone no. | Position | Gene | Mutation | Annotation | Product |
|------------|-----|-----------|----------|------|----------|------------|---------|
| M1         | 8   | 1         | 673,439  | mngB/ [cydA] Intergenic (+/−3) - coding (5/1,569 nt) | Δ6 bp | cydA: cytochrome bd-I terminal oxidase, subunit I; mngB: alpha-mannosidase |
| M1         | 8   | 61        | 174,223  | hemL  | A→C     | L220R (CTG→CGG) | Glutamate-1-semialdehyde aminotransferase |
|            |     |           | 3,077,939| yqiU  | Coding (562/630 nt) | IS5 | Predicted inner membrane protein |
|            |     |           | 4,205,688| yjbS  | Coding (193/204 nt) | IS1 | Hypothetical protein |
| M2         | 4   | 11        | 174,688  | hemL  | C→T     | G65D (GGG→GAC) | Glutamate-1-semialdehyde aminotransferase |
| M2         | 4   | 19        | 905,376  | ycbU  | Coding (297/543 nt) | IS2 | Predicted fimbrial-like adhesin protein |
|            |     |           | 2,223,615| ubiG  | Coding (222–231/723 nt) | Δ10 bp | Bifunctional 3-demethylubiquinone-8 3-O-methyltransferase and 2-octaprenyl-6-hydroxyphenol methylase |
| M3         | 5   | 26        | 174,223  | hemL  | A→C     | L220R (CTG→CGG) | Glutamate-1-semialdehyde aminotransferase |
|            |     |           | 3,077,939| yqiU  | Coding (562/630 nt) | IS5 | Predicted inner membrane protein |
| M3         | 5   | 28        | 174,223  | hemL  | A→C     | L220R (CTG→CGG) | Glutamate-1-semialdehyde aminotransferase |
|            |     |           | 3,077,939| yqiU  | Coding (562/630 nt) | IS5 | Predicted inner membrane protein |
|            |     |           | 4,205,688| yjbS  | Coding (193/204 nt) | IS1 | Hypothetical protein |
| M5         | 8   | 41        | 2,939,858| pepP  | Coding (153/1,326 nt) | IS1 | Proline aminopeptidase P II |
| M5         | 8   | 48        | 1,069,386| ndh   | C→T     | H52Y (CAC→TAC) | NADH:ubiquinone oxidoreductase II |
|            |     |           | 2,283,177| nuoG  | G→C     | Y272* (TAG→TAC) | NADH:ubiquinone oxidoreductase I, chain G |
| M6         | 4   | 51        | 2,927,004| ygfZ  | Coding (522/981 nt) | +GC | Folate-binding protein |
|            |     |           | 3,874,865| hemY  | T→A     | D237V (GAT→GTG) | Predicted protoheme IX synthesis protein |
| M6         | 4   | 52        | 2,927,004| ygfZ  | Coding (522/981 nt) | +GC | Folate-binding protein |
|            |     |           | 3,874,865| hemY  | T→A     | D237V (GAT→GTG) | Predicted protoheme IX synthesis protein |
| M4         | 5   | 31        | 4,324,488| mscM  | C→G     | W547C (TGG→TGC) | Mechanosensitive channel of miniconductance, monomer |
| M4         | 5   | 40        | 4,324,488| mscM  | C→G     | W547C (TGG→TGC) | Mechanosensitive channel of miniconductance, monomer |
four mice (caged in pairs) were used for each experimental group. Specifically, one group did not receive any additional treatment (‘no treatment’ in Fig. 3) and the other three groups were gavaged (between days 1 and 6) with the following antibiotics: kanamycin (1 mg), tetracycline (0.5 mg) or nalidixic acid (2 mg). Doses were based on per kg doses administered to human adults.

Mice in different cages were inoculated with independently grown bacteria and on different days. After *E. coli* gavage, faecal samples were obtained daily, homogenized and plated (LB-agar) for estimating CFP (SCV or revertant) and YFP (ancestral) cfu. As a control, four mice were also gavaged with a mix of isogenic CFP/YFP ancestral. For these mice, the YFP/CFP ratio did not significantly change over time ($\chi^2 = 0.82, p = .36$; Fig. S3), indicating that the fluorescent markers are neutral in this environment. Moreover, to ensure that the observed fitness effects are due to the environmental conditions in the gut and not because streptomycin increases the relative fitness of SCVs, in vitro competitions were carried, mimicking the in vivo competitions. This consisted of inoculating a 50:50 mixture of the SCV population and the YFP ancestral into RPMI at 100 $\mu$g/ml or 500 $\mu$g/ml of streptomycin ($n = 3$ replicates per concentration). Twenty-four hours post-incubation, the frequency of CFP/YFP bacteria was estimated, with SCVs always being deleterious (see Fig. S3D), in accordance with: (i) SCVs having a growth disadvantage in RPMI (Fig. S5); (ii) all clones being streptomycin resistant (RpsLA43R).

### 2.7 Data analysis

All analyses were carried in R version 3.2.1, using $t$-tests, Wilcoxon tests, ANOVA or linear mixed-effects models (http://www.r-project.org/). Where required, data were transformed to meet assumptions made by parametric statistics.

### 2.8 Ethics statement

All experiments involving animals were approved by the Institutional Ethics Committee at the Instituto Gulbenkian de Ciência (project nr. A009/2010 with approval date 2010/10/15), following the Portuguese legislation (PORT 1005/92) which complies with the European Directive 86/609/EEC of the European Council.

### 3 RESULTS AND DISCUSSION

#### 3.1 Novel genetic targets, associated with the ETC, mediate the SCV phenotype in *E. coli*

Whole-genome sequencing of 10 SCV clones evolved during interactions with Mfos and revealed a total of 19 mutations, with 1–3 mutations per clone (average: 1.9; Table 1). This is a large number of mutations, given that SCVs were isolated after only 60–120 generations of adaptation to Mfos. Interestingly, in half the populations (M1, M2 and M5), SCVs have distinct haplotypes and have therefore emerged independently. Thus, while we have not estimated the full genetic diversity within populations, it is clear that SCVs can be a polymorphic subpopulation, with different beneficial haplotypes in competition (a phenomenon known as clonal interference; Sniegoski & Gerrish, 2010).

The mutations in SCVs affected 11 different genetic targets, all in coding regions. To the best of our knowledge, 10 of these have not been previously associated with the SCV phenotype. The exception is *hemL*, the most common genetic target in our experiment: mutated in three of five independent populations. *hemL* has also been associated with SCVs in *S. enterica*, for spontaneous resistance to antimicrobial peptides (Pranting & Andersson, 2010) and during adaptation to the intracellular environment of fibroblasts (Canò et al., 2003). Additionally, we observed parallel mutations in two other genes (*yqJ* and *yjbS*: mutated twice), and at the haplotype level with clones carrying mutations in *hemL*, *yqJ* and *yjbS* being isolated from two different populations (M1 and M3). Importantly, functional annotation clustering (Table S2) and literature review on the function of the mutated genes revealed widespread functional parallelism, as all clones had at least one mutated gene associated with the ETC. Therefore, our WGS data show evidence for parallel adaptation at multiple levels.

Mutations in genes of the ETC are often associated with SCVs, however the diversity of mutational targets that can give rise to such SCVs is poorly understood. Our results (Table S2) indicate that the observed mutations target three classes of processes: first, oxidative phosphorylation, through mutations in subunit I of cytochrome bd oxidase (cydA; Green, Kranz, Lorence, & Gennis, 1984) and in both NADH:ubiquinone oxidoreductase I and II (nuoG and ndh, respectively, Young & Wallace, 1976; Weidner et al., 1993); second, heme and Fe-S cluster synthesis through mutations in: (i) two enzymes that are involved in heme synthesis (*hemL* and *hemY*), which is a key prosthetic group in several enzymes of the ETC (e.g. cytochromes; Alefounder, Abell, & Battersby, 1988; Iiag, Jahn, Eggertsson, & Söll, 1991); (ii) *ygfZ*, which is thought to be involved in the synthesis/repair of Fe-S clusters (present in enzymes as NADH:ubiquinone oxidoreductase I; Ote et al., 2006; Waller et al., 2010); and third, ubiquinone synthesis (the key electron acceptor for the ETC; Stroobant, Young, & Gibson, 1972), through mutations in an O-methyltransferase (*ubiG*) and in *pepP*. While the function of *pepP* (an aminopeptidase) is not directly related to ETC (Yoshimoto, Murayama, Honda, Tone, & Tsuru, 1988), transposition of an insertion sequence (IS) into *pepP* is known to have polar effects on *ubiH* activity (part of the same operon; Way, Salvistio, Maglizzo, & Goldber, 1999). We confirmed this result in our SCV clone, by demonstrating that (i) *ubiH* expression was strongly reduced in the *pepP* mutant (clone 41; Fig. S4A) and (ii) that this mutant was unable to grow in minimal media supplemented with succinate or glycerol, but not glucose (Fig. S4B and Appendix S1; this was previously known for *ubiH* mutants; Gupta, Mat-Jan, Latifi, & Clark, 2000; Joyce et al., 2006). Importantly, revertant clones, derived from clone 41 (i.e. producing a large colony morphotype), were able to grow on succinate and glycerol (Fig. S4C). As *ubiH* is involved in ubiquinone synthesis (Young, Stroobant, Macdonald, & Gibson, 1973), these data indicate that the SCV phenotype of the *pepP* mutant is also mediated by reduced activity of the ETC (rather than a direct effect of disrupting *pepP* function).
The two positive control clones, which do not show an SCV phenotype, (i.e. form large colonies; Fig. S1), carry a single mutation in mscM, which encodes a mechanosensitive ion channel of miniconductance. This is potentially important under hypo-osmotic shock (Edwards et al., 2012), but is not associated with the ETC. Therefore, these results generally suggest that the SCV phenotype that emerged during *E. coli* adaptation to MΦs is mediated by defects in the ETC, as has often been observed for SCVs isolated from experiments or clinical samples, across multiple species (Cano et al., 2003; Dean et al., 2014; Pranting & Andersson, 2010; Proctor et al., 2006; Roggenkamp et al., 1998).

### 3.2 E. coli SCVs have phenotypes similar to SCVs from clinical isolates

We measured several phenotypes that are typically observed in SCVs with mutations in the ETC (both in clinical and experimental isolates; results summarized in Table S3). We found that SCVs had a generally high (but variable) reversion frequency (to large colony phenotype), ranging from $10^{-6}$ to $10^{-9}$. In eight of 10 clones, the reversion frequency was $10^{-9}$, in line with previous observations (Häussler et al., 1999; Proctor et al., 2006; Roggenkamp et al., 1998; Trüdzsch et al., 2003). When tested for catalase activity, four SCV clones were catalase negative (Proctor et al., 2006), all of which carried mutations in hemL (clones 11, 26, 28, 61; Table S3). This is in agreement with the fact that heme is a co-factor in both *E. coli* catalases (Schellhorn, 1995).

Small-colony variants typically show poor growth under in vitro conditions, as a result of auxotrophy for specific compounds (Proctor et al., 2006; Wei et al., 2011). In agreement, our SCVs generally reached a lower carrying capacity than the ancestral strain, both in poor (M9glu0.4) and rich media (RPMI; Table S3; Fig. S5). However, the effect of the SCV phenotype on growth rate was media-dependent. In poor media, the relative growth rate was generally lower (8/10 SCV clones), but in rich media, this was strongly dependent on the SCV clone. Thus, it is important to understand how the within-host environment affects SCV growth in order to be able to predict the in vivo fitness of this morphotype (this could be done through indirect methods as in Barroso-Batista et al. 2015).

Two of the auxotrophies that are often observed in SCVs with mutations in the ETC are for heme and aminolevulinic acid (5-ALA; Proctor et al., 2006; Pranting & Andersson, 2010). Accordingly, supplementation of our SCV clones with either of these compounds improved their growth (Table S3; Fig. S6). However, while the benefit provided by heme was observed across all clones and growth parameters (lag phase, maximum growth rate and carrying capacity), 5-ALA specifically shortened the lag phase of only five clones, four of which carry mutations in hemL (~50% reduction). This is consistent with the described role of hemL in converting glutamate-1-semialdehyde to 5-ALA (Ilag et al., 1991), the first committed precursor of heme-biosynthesis, a key co-factor in several proteins of the ETC (Ilag et al., 1991).

These results highlight the pleiotropic consequences of the SCV phenotype. Moreover, while SCVs show qualitative convergence for slow growth and high reversion frequency, the phenotypes are quantitatively variable across SCV clones. The only clones that can be qualitatively distinguished are the hemL mutants due to their 5-ALA auxotrophy and negative catalase test.

### 3.3 SCVs have a fitness advantage in the intracellular environment of macrophages

As the SCVs used in this study emerged in the presence of MΦs, we carried out competition assays to test whether SCVs have a fitness advantage inside MΦs. Figure 1 shows that an SCV clone carrying mutations in hemL, yqiJ and yjbS had a fitness advantage (CI significantly different from 0) after both short (1 hr) and prolonged (6 hr) residence inside MΦs (at 1 hr: CI = 0.14 ± 0.06, mean ± 2 standard errors, $t_5 = 6.84$, $p < .01$; at 6 hr: CI = 0.22 ± 0.08, $t_5 = 8.25$, $p < .001$). A clone with an IS1 insertion in pepP displayed a transient fitness advantage, visible at 1 hr (CI = 0.08 ± 0.08, $t_5 = 2.9$, $p = .03$), but not at 6 hr. The ancestral strain of *E. coli* does not replicate inside MΦs (as most commensal strains; Koli, Sudan, Fitzgerald, Adhya, & Kar, 2011), the observed fitness advantage of these SCVs is likely mediated by an increased survival ability of SCVs inside MΦs. We also tested a clone with an mscM mutation that shows a colony size similar to the ancestral and observed no change in competitive ability. Overall, these data show that SCVs have increased intracellular survival in MΦs, providing an explanation for their repeated emergence during experimental adaptation of *E. coli* to MΦs (Miskine et al., 2013). This is also in agreement with studies showing that SCVs can emerge and have a fitness advantage in

![FIGURE 1 SCVs have a fitness advantage inside MΦs.](image-url)
the intracellular milieu of nonprofessional phagocytes (Cano et al., 2003; Tuchscherr et al., 2011; Vesga et al., 1996).

### 3.4 SCVs present antagonistic pleiotropy for antibiotic resistance in vitro

A common phenotype of SCVs is increased resistance to aminoglycoside antibiotics (von Eiff et al., 1997; Mush er et al., 1977). However, much less is known about the antibiotic resistance profile of SCVs against nonaminoglycoside antibiotics (Cano et al., 2003; Pranting & Andersson, 2010). In Fig. 2, we show that E. coli SCVs were generally more resistant to aminoglycosides, but were more susceptible to tetracycline, nalidixic acid and ciprofloxacin (quinolones), and cefotaxime (a β-lactam). All SCV clones had significantly increased resistance to at least three aminoglycosides (relative to the ancestral), but also increased susceptibility to at least three nonaminoglycoside antibiotics ($p < .05$ ANOVA contrasts; see Fig. S7 for mean ± 2SE). The positive control clones (mscM muta nts), whose colony size is similar to the ancestral, do not show such a pattern.

Recent work, using in vitro experimental evolution of E. coli in the presence of aminoglycosides, has shown that resistant mutants often have increased sensitivity to other antibiotics (including nalidixic acid, ciprofloxacin or tetracycline; Imamovic & Sommer, 2013; Pranting & Gordo, in press; Miskinyte & Gordo, 2013). In agreement with the in vitro results, SCV fitness in the mouse gut was dependent on the antibiotic treatment (time:treatment interaction: $\chi^2_1 = 11.4, p < .001$), with a positive selection coefficient (per day) of 0.14 (±0.05, standard error; SE) relative to the ancestral (Fig. 3A,B). Conversely, the frequency of revertant bacteria (forming large CFP colonies) decreased from ~0.12 on day 0 to below 0.14 (±0.05, SE) by day 7 post-colonization (Fig. 3B). These results show that SCVs can have an increased competitive ability in vivo that can be reversed with tetracycline treatment.

Remarkably, we observed that SCVs had a fitness advantage in the mouse gut (Fig. 3A,B): SCV frequency increased, and by day 7, SCVs constituted the majority of the population in three of four mice. Accordingly, SCV ratio significantly increased with time (Fig. 3A; $\chi^2_7 = 58.9, p < .0001$), with a positive selection coefficient (per day) of 0.14 (±0.05, standard error; SE); see methods; (Dykhuizen, 1990); see Fig. S3 for slopes). Interestingly, some of these studies recurrently observed mutations in genes with similar functions to those observed here (e.g. heme synthesis: hemA; NADH dehydrogenase I: nuoF and cytochrome bd-I terminal oxidase: cydA; Lázár et al., 2013; Oz et al., 2014; Suzuki, Horinouchi, & Furusawa, 2014), a phenomenon termed collateral sensitivity (Szybalski & Bryson, 1952). Interestingly, some of these studies recurrently observed mutations

### 3.5 SCVs have a fitness advantage in vivo that can be reversed with tetracycline treatment

While several recent papers have shown collateral sensitivity in vitro (Imamovic & Sommer, 2013; Lázár et al., 2013; Roemhild, Barbosa, Beardmore, Jansen, & Schu lenburg, 2015), whether this phenomenon also occurs in vivo is currently unknown. As the effect of resistance mutations is highly dependent on the environment (Miskinyte & Gordo, 2013; Trindade, Sousa, & Gordo, 2012), it is imperative to determine whether collateral sensitivity is also observed in vivo. Here, we used the streptomycin-treated mouse model of gut colonization to test for the competitive ability of SCVs against the ancestral, in the presence or absence of treatment with kanamycin, nalidixic acid or tetracycline.

In agreement with the in vitro results, SCV fitness in the mouse gut was dependent on the antibiotic treatment (time:treatment interaction: $\chi^2_2 = 58.9, p < .0001$). SCV fitness significantly increased with kanamycin treatment ($\chi^2_2 = 10.2, p < .01$), leading to a selection coefficient of 0.36 (±0.05, SE). On the other hand, tetracycline treatment,
significantly reduced SCV fitness within the mouse gut ($\chi^2 = 28.8$, $p < .0001$), leading to a fitness disadvantage (selection coefficient: $-0.23 \pm 0.05$, SE). Nalidixic acid also led to a reduction in the fitness advantage of SCVs (slope: $0.05 \pm 0.05$, SE). However, this difference was not significant ($\chi^2 = 1.9$, $p = .17$).

As far as we are aware, these results provide the first in vivo demonstration that the frequency of SCVs can be increased by aminoglycoside treatment, but decreased by tetracycline due to their collateral sensitivity to this antibiotic. Furthermore, while we observed that antibiotics strongly influence the composition of the *E. coli* population, the total population size remains relatively stable (Fig. S3), albeit lower than in mice drinking only streptomycin-treated water ($\chi^2 = 4.8$, $p = .03$). As commensal bacteria can acquire pathogenic traits through just a few mutations (Koli et al., 2011; Limoli et al., 2014; Miskinyte...
et al., 2013), these results suggest that it may be possible to reduce the frequency of bacteria with such traits (e.g. SCVs) without eliminating the commensals.

4 | CONCLUSIONS

Small-colony variants are an important bacterial morphotype often associated with chronic infections and antibiotic resistance. In this manuscript, we identified new mutational paths that confer an SCV phenotype and show that these mutations have multiple pleiotropic phenotypic consequences in vitro and in vivo (in the mouse gut). Thus, our results highlight that combining experimental evolution in the presence of key cells of the innate immune system, with WGS and phenotypic tests can reveal: (i) the genetic basis of bacterial traits important for within-host fitness and (ii) the selective pressures that lead to such traits.

Importantly, the SCV phenotype leads to a trade-off, both in vitro and in vivo, between increased aminoglycoside resistance and collateral sensitivity to other antibiotic classes. As SCVs carry multiple mutations in genes related to the ETC, collateral sensitivity is likely mediated by a reduction in proton-motive force (PMF) across the inner membrane. This is because resistance to other antibiotics is mediated by several PMF-dependent efflux pumps, thus reduced PMF can lead to increased sensitivity to some antibiotics (Lázár et al., 2013).

Collateral sensitivity has been suggested as an important principle for the design of treatment regimens that curb the evolution of antibiotic resistance (Imamovic & Sommer, 2013; Lázár et al., 2013). However, at the moment, we are aware of only a single report using this concept to treat bacteria in vivo (Gonzales et al., 2015). Thus, more work is currently needed to establish the in vivo effectiveness of therapies involving collateral sensitivity (Baym, Stone, & Kishony, 2016).

ACKNOWLEDGEMENTS

We would like to thank Daniela Güleresi for assistance in the laboratory, Jessica Thompson for editing the manuscript and all members of the Evolutionary Biology laboratory, two anonymous reviewers and editor for comments. This work was funded by European Research Council under the European Community’ Seventh Framework Programme (FP7/2007–2013)/ERC grant agreement no. 260421–ECOADAPT, German Science Foundation (DFG, G-410861, through SFB-680 collaborative research center) and from the Portuguese Science Foundation FCT (PTDC/BIA-EVF/118075/2010). IG acknowledges salary support of LAO/ITQB & FCT. We thank the IGC Animal House Unit which is partially funded through FCT grant RECI/IMI-IMU/0038/2012 and EU-FP7 infrastructure grants EMMA and Infracell3.

DATA ARCHIVING STATEMENT

Data available from the Dryad Digital Repository: http://dx.doi.org/10.5061/dryad.3h9f5

REFERENCES

Al Mamun, A. A. M., Lombardo, M.-J., Shee, C., Lisewski, A. M., Gonzalez, C., Lin, D., ... Nehring, R. B. (2012). Identity and function of a large gene network underlying mutagenic repair of DNA breaks. Science, 338, 1344–1348.

Alefounder, P. R., Abell, C., & Battersby, A. R. (1988). The sequence of hemC, hemD and two additional E. coli genes. Nucleic Acids Research, 16, 9871.

Azevedo, M., Sousa, A., Moura de Sousa, J., Thompson, J. A., Proença, J. T., & Gordo, I. (2016). Trade-offs of Escherichia coli adaptation to an intracellular lifestyle in macrophages. PLoS One, 11, e0146123.

Bailey, S. F., & Bataillon, T. (2016). Can the experimental evolution programme help us elucidate the genetic basis of adaptation in nature? Molecular Ecology, 25, 203–218.

Barroso-Batista, J., Demengeot, J., & Gordo, I. (2015). Adaptive immunity increases the pace and predictability of evolutionary change in commensal gut bacteria. Nature Communications, 6, 8945.

Barroso-Batista, J., Sousa, A., Lourenço, M., Bergman, M. L., Sobral, D., Demengeot, J., ... Gordo, I. (2014). The first steps of adaptation of Escherichia coli to the gut are dominated by soft sweeps. PLoS Genetics, 10, e1004182.

Baym, M., Stone, L. K., & Kishony, R. (2016). Multidrug evolutionary strategies to reverse antibiotic resistance. Science, 351, aad3292.

Cano, D. A., Pucciarelli, M. G., Martinez-Moya, M., Casadesús, J., & Garcia-del Portillo, F. (2003). Selection of small-colony variants of Salmonella enterica serovar typhimurium in nonphagocytic eucaryotic cells. Infection and Immunity, 71, 3690–3698.

Carryn, S., Van Bambke, F., Mingeot-Leclercq, M.-P., & Tulkens, P. M. (2002). Comparative intracellular (THP-1 macrophage) and extracellular activities of beta-lactams, azithromycin, gentamicin, and fluoroquinolones against Listeria monocytogenes at clinically relevant concentrations. Antimicrobial Agents and Chemotherapy, 46, 2095–2103.

Conway, T., Krogfelt, K., & Cohen, P. (2004). The life of commensal Escherichia coli in the mammalian intestine. EcoSal Plus, 1, 1–15.

Cooper, V. S., Staples, R. K., Traverse, C. C., & Ellis, C. N. (2014). Parallel evolution of small colony variants in Burkholderia cepacia biofilms. Genomics, 104, 447–452.

Dean, M. A., Olsen, R. J., Long, S. W., Rosato, A. E., & Musser, J. M. (2014). Identification of point mutations in clinical Staphylococcus aureus strains that produce small-colony variants auxotrophic for menadione. Infection and Immunity, 82, 1600–1605.

Deatherage, D. E., & Barrick, J. E. (2014). Identification of mutations in laboratory-evolved microbes from next-generation sequencing data using breseq. Methods in Molecular Biology, 1151, 165–188.

Dettman, J. R., Rodrigue, N., Melnyk, A. H., Wong, A., Bailey, S. F., & Kas-sen, R. (2012). Evolutionary insight from whole-genome sequencing of experimentally evolved microbes. Molecular Ecology, 21, 2058–2077.

Durão, P., Güleresi, D., Proença, J., & Gordo, I. (in press). Enhanced survival of Rfampicin and Streptomycin double resistant E. coli inside macrophages. Antimicrobial Agents and Chemotherapy.

Dykhuizen, D. E. (1990). Experimental studies of natural selection in bacteria. Annual Review of Ecology and Systematics, 21, 373–398.

Edwards, M. D., Black, S., Rasmussen, T., Rasmussen, A., Stokes, N. R., Stephen, T. L., ... Booth, I. R. (2012). Characterization of three novel mechanosensitive channel activities in Escherichia coli. Channels Austin Tex, 6, 272–281.

von Eiff, C., Bettin, D., Proctor, R. A., Rolauffs, B., Lindner, N., Winkelmann, W., & Peters, G. (1997). Recovery of small colony variants of Strepto-coccus aureus following gentamicin bead placement for osteomyelitis. Clinical Infectious Diseases, 25, 1250–1251.

Elena, S. F., & Lenski, R. E. (2003). Evolution experiments with microorganisms: The dynamics and genetic bases of adaptation. Nature Reviews Genetics, 4, 457–469.

Ensminger, A. W., Yassin, Y., Miron, A., & Isberg, R. R. (2012). Experimental evolution of Legionella pneumophila in mouse macrophages leads
to strains with altered determinants of environmental survival. PLoS Pathogens, 8, e1002731.
Friesen, M. L., Saxer, G., Travisano, M., & Doebeli, M. (2004). Experimental evidence for sympatric ecological diversification due to frequency-dependent competition in Escherichia coli. Evolution, 58, 245–260.
Gonzales, P. R., Pesesky, M. W., Bouley, R., Ballard, A., Biddy, B. A., Suckow, M. A., ... Dantas, G. (2015). Synergistic, collaterally sensitive β-lactam combinations suppress resistance in MRSA. Nature Chemical Biology, 11, 855–861.
Green, G. N., Kranz, R. G., Lorence, R. M., & Gennis, R. B. (1984). Identification of subunit I as the cytochrome b558 component of the cytochrome d terminal oxidase complex of Escherichia coli. Journal of Biological Chemistry, 259, 7994–7997.
Gupta, S., Mat-Jan, F., Latifi, M., & Clark, D. P. (2000). Acetalddehyde dehydrogenase activity of the AdhE protein of Escherichia coli is inhibited by intermediates in ubiquinone synthesis. FEMS Microbiology Letters, 182, 51–55.
Häussler, S., Tümmler, B., Weißbrodt, H., Rohde, M., & Steinmetz, I. (1999). - (1999). Small-colony variants of Pseudomonas aeruginosa in cystic fibrosis. Clinical Infectious Diseases, 29, 621–625.
Herron, M. D., & Doebeli, M. (2013). Parallel evolutionary dynamics of adaptive diversification in Escherichia coli. PLoS Biology, 11, e1001490.
Huang, D. W., Sherman, B. T., & Lempicki, R. A. (2008). Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. Nature Protocols, 4, 44–57.
Huang, D. W., Sherman, B. T., & Lempicki, R. A. (2009). Bioinformatics enrichment tools: Paths toward the comprehensive functional analysis of large gene lists. Nucleic Acids Research, 37, 1–13.
Ilag, L. L., Jahn, D., Eggertsson, G., & Söll, D. (1991). The Escherichia coli heml gene encodes glutamate 1-semialdehyde aminotransferase. Journal of Bacteriology, 173, 3408–3413.
Imamovic, L., & Sommer, M. O. A. (2013). Use of collateral sensitivity networks to design drug cycling protocols that avoid resistance development. Science Translational Medicine, 5, 204ra132–204ra132.
Joyce, A. R., Reed, J. L., White, A., Edwards, R., Osterman, A., Baba, T., ... Agarwala, S. (2006). Experimental and computational assessment of conditionally essential genes in Escherichia coli. Journal of Bacteriology, 188, 8259–8271.
Kastbjerg, V. G., Hein-Kristensen, L., & Gram, L. (2014). Triclosan induced aminoglycoside-tolerant Listeria monocytogenes isolates can appear as small-colony variants. Antimicrobial Agents Chemotherapy, doi: 10.1128/AAC.02666-13.
Koli, P., Sudan, S., Fitzgerald, D., Adhya, S., & Kar, S. (2011). Conversion of commensal Escherichia coli K-12 to an invasive form via expression of a mutant histone-like protein. mBio, 2, e00182–11.
Lázár, V., Singh, G. P., Spohn, R., Nagy, I., Horváth, B., Hrtyan, M., ... Pósfai, G. (2013). Bacterial evolution of antibiotic hypersensitivity. Molecular Systems Biology, 9, 700.
Lee, S. M., Wyse, A., Lesher, A., Everett, M. L., Lou, L., Holzknecht, Z. E., ... Parker, W. (2010). Adaptation in a mouse colony monoassociated with Escherichia coli K-12 for more than 1,000 days. Applied and Environment Microbiology, 76, 4655–4663.
Limoli, D. H., Rockel, A. B., Host, K. M., Jha, A., Kopp, B. T., Hollis, T., & Wozniak, D. J. (2014). Cationic antimicrobial peptides promote microbial mutagenesis and pathoadaptation in chronic infections. PLoS Pathogens, 10, e1004083.
McDonald, M. J., Gehrig, S. M., Meintjes, P. L., Zhang, X.-X., & Rainey, P. B. (2009). Adaptive divergence in experimental populations of Pseudomonas fluorescens. IV. Genetic constraints guide evolutionary trajectories in a parallel adaptive radiation. Genetics, 183, 1041–1053.
Miskinute, M., & Gordo, I. (2013). Increased survival of antibiotic-resistant Escherichia coli inside macrophages. Antimicrobial Agents and Chemistry, 57, 189–195.
Miskinute, M., & Gordo, I. (2014). Fitness measurements of evolved Escherichia coli. Bio-Protocol, 4, e1228.
