

Recurrent Reverse Evolution Maintains Polymorphism After Strong Bottlenecks in Commensal Gut Bacteria

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**Recurrent Reverse Evolution Maintains Polymorphism After Strong
Bottlenecks in Commensal Gut Bacteria**

Authors: A. Sousa^{1,2}, R. S. Ramiro¹, J. Barroso-Batista¹, D. Güleresi¹, M. Lourenço^{1,3}, I. Gordo^{1*}

Affiliations:

¹Instituto Gulbenkian de Ciência, Rua da Quinta Grande 6, Oeiras, Portugal.

²Institute for Biomedicine, Department of Medical Sciences, University of Aveiro, Aveiro, Portugal

³Current address: Department of Microbiology, Institute Pasteur, Paris.

*Correspondence to: igordo@igc.gulbenkian.pt

Abstract

The evolution of new strains within the gut ecosystem is poorly understood. We used a natural but controlled system to follow the emergence of intra-species diversity of commensal *Escherichia coli*, during three rounds of adaptation to the mouse gut (~1300 generations). We previously showed that, in the first round, a strongly beneficial phenotype (loss-of-function for galactitol consumption; *gat*-negative) spread to >90% frequency in all colonized mice. Here, we show that this loss-of-function is repeatedly reversed when a *gat*-negative clone colonizes new mice. The re-gain of function occurs via compensatory mutation and reversion, the latter leaving no trace of past adaptation. We further show that loss-of-function adaptive mutants re-evolve, after colonization with an evolved *gat*-positive clone. Thus, even under strong bottlenecks a regime of strong-mutation-strong-selection dominates adaptation. Coupling experiments and modeling, we establish that reverse evolution recurrently generates two coexisting phenotypes within the microbiota that can or

not consume galactitol (*gat*-positive and *gat*-negative, respectively). While the abundance of the dominant strain, the *gat*-negative, depends on the microbiota composition, *gat*-positive abundance is independent of the microbiota composition and can be precisely manipulated by supplementing the diet with galactitol. These results show that a specific diet is able to change the abundance of specific strains. Importantly, we find polymorphism for these phenotypes in indigenous Enterobacteria of mice and man. Our results demonstrate that natural selection can greatly overwhelm genetic drift at structuring the strain diversity of gut commensals and that competition for limiting resources may be a key mechanism for maintaining polymorphism in the gut.

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1 **Introduction**

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3 The mammalian gut is inhabited by a highly diverse microbial community (including
4 multiple phyla of bacteria, archaea, eukaryotes and viruses). Current data suggests that
5 maintaining a species rich gut bacterial community is important for multiple aspects of host
6 health (Clemente et al. 2012), including: resistance to pathogens (van Nood et al. 2013),
7 development of the immune system (Faith et al. 2014) and even behavior (Dinan and Cryan
8 2017). Maintenance of species diversity in the gut may be influenced by several ecological
9 factors: spatial structure (transversal and longitudinal), temporal fluctuations in the
10 environment (e.g. food intake, (David et al. 2013) and competitive interactions amongst
11 colonizing species (Coyte et al. 2015). Much less studied is the diversity within each
12 species of the microbiota (Caugant et al. 1981) and the extent to which such intra-species
13 variation has functional consequences for both bacteria and their hosts (Schloissnig et al.
14 2012; Greenblum et al. 2015). In particular the dynamics of emergence and spread of
15 adaptive mutations in commensal gut bacteria are poorly known (Waldor et al. 2015).
16 Different evolutionary mechanisms can promote genetic diversity within species (Chesson
17 2000). The recurrent emergence of new adaptive mutations in different clones that compete
18 for fixation - clonal interference - leads to unstable levels of polymorphism (Gerrish and
19 Lenski 1998). On the other hand resource partitioning and negative-frequency dependent
20 selection leads to stable polymorphisms (Chesson 2000). Evolution by *de novo* mutation in
21 the gut has recently started to be studied using a mouse model of colonization, the
22 streptomycin-treated mouse. In this model the antibiotic (streptomycin) is used to deplete
23 some of the microbiota members (the facultative anaerobes) so that specific strains of

commensal or pathogenic bacteria can colonize the gut (Krogfelt et al. 2004) . Clonal interference (Barroso-Batista et al. 2014; Lourenço et al. 2016), trade-offs between stress resistance and nutritional competence (De Paepe et al. 2011) and negative frequency-dependent selection (Lourenço et al. 2016), have all been shown to affect the adaptive dynamics of newly emerging mutations in the guts of mice.

The long-term success of bacterial lineages also depends on their ability to transmit between hosts. Such transmission can be vertical, e.g. from mother to offspring, or horizontal, e.g. in the case of fecal transplantation or migration of microbes between hosts inhabiting the same environment. The process of transmission can entail strong population bottlenecks that may reduce genetic diversity, potentially breaking interactions between strains and/or species that were created within the gut of a given host (Coyte et al. 2015).

Genetic diversity can then be regained during population re-establishment. However, it is currently poorly understood whether the recovery in diversity involves mutations that re-establish previously existing polymorphisms at the same or at different loci/traits.

Escherichia coli generally associates with humans as a commensal, being one of the first colonizers of our guts. In particular, the strain K12 constitutes one of the best-studied model organisms. Recently, we began to uncover the nature of the adaptive process that *E. coli* experiences in the gut, by following the evolutionary dynamics of a lineage colonizing mice. Using the streptomycin-treated mouse model of colonization, described above, and a fluorescently labeled strain of *E. coli*, we unraveled the spread of adaptive mutations, by the signature it leaves in the dynamics of neutral fluorescent markers (Barroso-Batista et al. 2014). In this mouse model, the *E. coli* that naturally inhabits the mouse gut, is killed by the antibiotic whereas the fluorescently labeled *E. coli* (which is streptomycin resistant) is able

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to stably colonize and evolve in the gut. We found that the first step of adaptation to the gut, arising as rapidly as three days post-colonization, consisted in the selective inactivation of the operon that allows *E. coli* to metabolize galactitol (i.e. *gat*-negative mutant; (Barroso-Batista et al. 2014)), a sugar alcohol derived from galactose. Distinct, but phenotypically equivalent, knock-out alleles of this operon bearing a similar selective effect (7.5% benefit (Barroso-Batista et al. 2015)) were recurrently observed to emerge across all populations adapting to the gut of independent mice, reaching 95-99% frequency after 24 days (~430 generations). Thus, the fitness effects of mutations comprising the first step of adaptation to the gut could be described by a simple distribution, where all mutations have similar effect (Hegreness 2006).

Here, we use an evolved *gat*-negative single mutant to colonize a new set of mice (mimicking a strong transmission bottleneck), and ask if the rate of adaptation slows down in this complex ecosystem. A deceleration in the rate of adaptation is expected under simple fitness landscape models (Couce and Tenaillon 2015) and is typically found in bacterial populations adapting to laboratory environments (Chou et al. 2011; Khan et al. 2011; Kryazhimskiy et al. 2014). If the selective benefit of emerging mutations decreases during adaptive walks, then diversity might be maintained for longer periods of time. We find that the rate of adaptation does not seem to decelerate for at least two consecutive bouts of adaptation. Unexpectedly, whole-genome sequencing of the evolving populations of the second round of adaptation, revealed that evolution took a step back, with the emergence of multiple mutations that re-enabled galactitol metabolism (*gat*-positive) by *E. coli*. These phenotypic reversions occurred by compensatory mutation and genetic reversion. Subsequently, upon a third round of adaptation, using an evolved *gat*-positive

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3 70 strain to colonize new mice, we find *de novo* loss-of-function mutations. The results
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5 71 demonstrate that in this system, irrespectively of the starting genotype and transmission
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8 72 involving the strongest of bottlenecks, natural selection can maintain polymorphism
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10 73 through recurrent reverse evolution. By assaying microbiota composition and *in vivo*
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12 74 competitions, we then show that the recurrent reverse evolution leads to the coexistence of
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14 75 two phenotypes whose niches appear to be independent, as they can consume different
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16 76 resources. Furthermore, while the total size of the *E. coli* species is dependent on
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18 77 microbiota composition, – with a more diverse microbiota being associated with a smaller
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20 78 load of *E. coli* – its diversification is not. Indeed, the abundance of the strain that can
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22 79 consume the resource galactitol (*gat*-positive) is relatively constant across mice with
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24 80 distinct microbiota compositions and can be increased by galactitol supplementation, in a
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26 81 concentration dependent manner. Conversely, the abundance of the clones that cannot grow
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28 82 on galactitol (*gat*-negative) is influenced by microbiota composition. Finally, we find that
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30 83 natural isolates of *E. coli*, inhabiting the guts of healthy humans show variation for
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32 84 galactitol consumption – a trait that we find to be under strong selection in laboratory mice.
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41 **Results and Discussion**

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45 **Rate of adaptation to the mouse gut during the second steps of adaptation**

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48 90 We colonized 15 mice with a clonal population of *E. coli* exhibiting the first beneficial
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50 91 phenotype – inability to uptake galactitol- conferred by a single base pair insertion in the
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52 92 coding region of the *gatC* gene, which encodes a subunit of the galactitol transporter. The
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54 93 colonizing population was made dimorphic for a neutral fluorescent marker to allow
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56 94 determination of the spread of further adaptive changes (Hegreness 2006). The time series
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dynamics of the fluorescent marker frequency resulting from the second steps of adaptation are shown in fig. 1A. These dynamics can be summarized in two effective evolutionary parameters (Hegreness 2006; Barrick et al. 2010; Barroso-Batista et al. 2014), the mutation rate (U_e) and the fitness effects of adaptive mutations (S_e) that are estimated to be $U_e = 7.1 \times 10^{-7}$, $S_e = 9\%$ (see Methods). These quantitatively describe the simplest adaptive scenario where new beneficial mutations emerge at a constant rate and have a constant effect (Hegreness 2006). The estimates of the evolutionary parameters for the second steps of adaptation are remarkably similar to those previously estimated for the first step ($U_e = 7 \times 10^{-7}$, $s_e = 7.5\%$) (Barroso-Batista et al. 2014). Direct *in vivo* competitions between samples of evolved clones and their ancestor (a *gat*-negative strain) further corroborate that the fitness benefit of the accumulated mutations is large, ranging from 5 to 14% fitness increase (mean fitness increase of $9 \pm 4\%$ (2SE), fig. 1B), which is similar to what was observed for the first steps of adaptation where evolved populations were competed against the ancestral *gat*-positive strain ($8 \pm 1\%$ (2SE)). Thus the strength of selection *in vivo* does not seem to decrease much after the first step of adaptation.

Widespread reversion of a strongly beneficial mutation upon transmission

To determine the genetic basis of adaptation underlying the rapid changes in neutral marker frequency (fig. 1A) we performed whole genome sequencing (WGS) of samples of two evolving populations along time in two lineages (2.14 and 2.15 from fig. 1A-B). Remarkably, we uncovered the rapid emergence of several mutations in the galactitol operon (fig. 2A-B, see also Table S1), particularly in the previously pseudogenized *gatC* (fig. 2C). These mutations restored the open reading frame of *gatC* suggesting that an

unexpected re-gain of function could have evolved. Thus, we tested hundreds of evolved clones from these populations for the ability to metabolize galactitol and found that the new mutations in *gatC*, spreading through the populations (fig. 2A-B), were indeed gain of function mutations. In face of our previous findings (Barroso-Batista et al. 2014), where *gat*-positive *E. coli* evolved a *gat*-negative phenotype that reached more than 75% frequency in all the mice after 24 days of colonization (fig. 3A), the re-emergence of the *gat*-positive phenotype was completely unexpected. Indeed, we had assumed that the *gat*-negative phenotype would eventually fix in all populations and therefore started a second colonization with a single adaptive *gat*-negative mutant. This design, mimicking a strong bottleneck or transmission event, serendipitously revealed that evolution could reverse.

To query whether the two populations studied reflect widespread reversion (Porter and Crandall 2003), we tested large samples of clones from all the evolved populations for a *gat*-positive phenotype. We found that phenotypic reversion was highly probable, as in 80% of the 15 independently colonized mice the bacteria re-gained the ability to consume galactitol, with this phenotype reaching a detectable frequency of at least 0.3% and up to 77% in the evolving populations (fig. 3B). Thus, under a transmission event involving a single clone, re-acquisition of polymorphism at the *gat* locus constituted a highly predictable adaptive event. Interestingly, the power of selection on this re-gain of function is clearly observed in population 2.14 (fig. 2A), where an allele (involving a 1bp deletion, the only mutation detected by WGS of the population as indicated in Table S1) emerges in this population at day 11, on the YFP background, but will then compete with a CFP clone which acquired another mutation causing the same phenotype. Thus the emergence of a *gat*-positive phenotype in both fluorescent backgrounds within the same evolving

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population (fig. 2A) and the parallelism across independent replicate populations (fig. 3B) strongly supports the adaptive nature of the re-gain of function in this ecosystem (de Visser and Krug 2014). To provide further evidence for selection acting on mutations conferring the *gat*-positive phenotype, in contrast to being hitchhiker alleles, we sequenced two *gat*-positive clones. Genome sequencing of the two clones (isolated from population 2.15 at day 6) revealed that: a back-mutation (“rev allele” in fig. 2C) had occurred in one clone, with no other mutations detected in its genome; a compensatory mutation (“comp3 allele” in fig. 2C), that redoes the open reading frame of *gatC* had occurred in the other clone, whose genome carried no further mutations. These results suggest that the *gat*-positive phenotype is intrinsically beneficial. Moreover, the rare event of gain-of-function through back-mutation could even be observed in one of the assayed populations (2.15, fig. 2B-C), supporting the conclusion that the ancestral gene can invade a population where loss of function at this locus had previously evolved. Such an event is particularly important since it erases the signature of the strong past selection that had occurred.

Natural selection maintains polymorphism during multiple rounds of transmission

To understand the extent to which selection can overwhelm genetic drift (associated with extreme population bottlenecks at host-to-host transmission) in driving the emergence of intra-species diversity at this locus, we performed a third colonization. This was initiated with an evolved *gat*-positive clone (isolated from population 2.14 – fig. 2A), which carries a 2bp insertion that redoes the *gatC* reading frame and two mutations at other genomic loci (see Materials and Methods for the exact genotype). In all 6 colonized mice, emergence of the *gat*-negative phenotype, consistent with inactivation of the *gat* operon, was seen (fig.

3C). In this new genetic background the emergence of polymorphism at detectable frequencies occurred slightly later than in the first colonization: the median time for *gat*-negative phenotype to reach 5% frequency was 7 days (6.25-7.75, Interquartile range), whereas for the first colonization it was 3 days (2-4, Interquartile range, Mann-Whitney-Wilcoxon Test, $W=1.5$, $P=0.0006$). The difference in the time of emergence of the polymorphism could be related with the two additional mutations whose phenotype was not characterized.

The three colonizations together (fig. 3A-C) suggest a model of cycling evolution, where a *E. coli* population can loose genetic diversity due to intense drift, caused by strong bottlenecks during transmission events between hosts, but can readily restore polymorphism at this locus due to strong selection in this ecosystem and high mutation rate (fluctuation tests show mutation rates of: 1.0×10^{-5} per generation (95% CI, [6.7×10^{-6} , 4.0×10^{-4}]) (Lourenço et al. 2016) for *gat*-positive to *gat*-negative and 9.7×10^{-9} (95% CI, [3×10^{-9} , 1.9×10^{-8}]) for *gat*-negative to *gat*-positive (see Materials and Methods)). Such strong bottlenecks may occur during vertical transmission from parent to offspring (Nayfach et al. 2016) and horizontal transmission of pathogenic strains in a hospital setting.

Competition for limiting resources can help explaining the maintenance of polymorphism at the *gat* operon

Competition for limiting resources is one of the mechanisms likely to be important in the gut microbial ecosystem (Coyte et al. 2015). Such a mechanism can lead to strong selection for evolving polymorphisms (Chesson 2000). Given the observed polymorphism for

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galactitol metabolism, we hypothesized that this sugar alcohol could be a key limiting resource for *E. coli* in the mouse gut. Though we did not measure directly the levels of galactitol in the gut, we have explored the potential implications of this being a limiting resource. We first explored the conditions under which a simple resource competition model (Hansen and Hubbell 1980; van Opheusden et al. 2015) could explain the invasions observed in the independent colonizations of fig. 3A-C (see Materials and Methods and fig. S1) and then extended the model to understand the conditions under which a mutant for galactitol metabolism could invade a resident population of bacteria which possesses a regulated operon (see below and fig S2). In the simplest model we assume that the *gat*-negative and the *gat*-positive clones consume two different resources. Given that the *gat*-positive strain (*E. coli* K-12 MG1655) constitutively expresses the *gat* operon (it carries an IS1 insertion in the operon repressor, *gatR*), we further make the simplifying assumption that this strain only consumes one of the resources (galactitol). As expected under these simple assumptions the theoretical model can reproduce quantitatively the dynamics of emergence of polymorphisms observed in the experiments (fig. S1A). Under these conditions we expect that supplementation of the mouse diet with galactitol should lead to an increase in frequency of *gat*-positive clones (fig. S1B). To test for this we colonized new mice with a co-culture of *gat*-positive and *gat*-negative bacteria (at a ratio 1 to 10) while supplementing the mice diet with different concentrations of galactitol. Indeed we observed that the frequency of *gat*-positive bacteria increased in mice drinking water with galactitol ($\chi^2_2=31.1$, $p<0.001$, from day 0 to day 2), and more so when the concentration of galactitol was higher (fig. 4A, also compare with fig. S1B). After day 3 the frequency of *gat*-positive bacteria stabilizes at different levels with different galactitol amounts, with all treatments being significantly different from each other (Tukey tests, $p<0.0001$). Furthermore the

absolute abundance of the *gat*-positive strain increases with galactitol concentration ($\chi^2_1=6.4$, $p=0.01$; log10 of CFUs: 7.2 ± 0.1 , 7.5 ± 0.1 and 8.4 ± 0.03 , for 0%, 0.01% and 0.1% galactitol, respectively, with all treatments being significantly different from each other $p<0.05$). The total load of *E. coli* was not significantly different across diets ($\chi^2_2=4.9$, $p=0.09$), consistent with the main effect of galactitol supplementation being the increase of *gat*-positive abundance.

We then assayed if the composition of the other species in the mouse gut could be altered when adding galactitol to the diet, by 16S rDNA metagenomics of the mouse fecal samples. Figure 4C shows that the microbiota of the mice drinking galactitol-supplemented water do not cluster separately from those that undergo a normal diet. The same conclusion was reached after performing a PCoA analysis on the data shown in fig. 4C, either considering days 3 through 7 (fig. S6) or each of the three days separately (fig. S7 – day 3, fig. S8 – day 4 and fig. S9 – day 7). Overall, these results suggest that galactitol can increase the frequency of specific strains of *E. coli* (fig. 4A) without significantly changing the load of this species (fig. 4B) and the relative abundance of other species of the microbiota (fig. 4C - within the limits of 16S profiling which is unable to detect differences below the genus level). Nevertheless, we cannot rule out the possibility that galactitol affects the background microbiota in the absence of *E. coli*.

The results so far presented provide a possible explanation for both the emergence and maintenance of polymorphism for galactitol consumption based on the hypothesis of non-overlapping niches. Yet, they do not exclude other hypotheses underlying the recurrent inactivation of the galactitol operon. The *gat*-positive strains which seeded the 1st and 3rd

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3 237 colonization constitutively express the *gat* operon (due to the IS element insertion in the
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5 238 *gatR*) and thus lack regulation. We hypothesized that a strain that could regulate galactitol
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10 240 *gat* operon would not be selected. To test for this, we replaced the pseudogene *gatR*, in the
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13 241 original strain, by a functional version, which does not impair the ability of consuming
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15 242 galactitol but regulates the operon expression. We then colonized new mice with this strain
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17 243 and followed its evolution in the mouse gut for 19 days ($n=6$). In contrast to the
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20 244 colonization initiated with *gat*-positive (constitutive), we did not observe the emergence of
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22 245 *gat*-negative mutants. Therefore this experiment suggests that the ability to regulate the *gat*
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24 246 operon impedes (or strongly delays) the loss of function previously observed in this
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27 247 ecosystem. Remarkably though, we did observe the emergence of *gat*-constitutive bacteria
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29 248 in all mice colonized with a regulated strain (fig. 5). Furthermore targeted PCR of the
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31 249 evolved clones showed that the phenotype could be caused by *de novo* IS insertions in the
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34 250 coding region of *gatR*. Interestingly, these emerging mutants are phenotypically equivalent
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36 251 to the starting *E. coli* clone, which suggests that constitutive mutants might be indeed
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38 252 segregating in natural populations. These results are predicted by the simple model of
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41 253 resource competition (fig. S2). The observed loss of regulation *in vivo* is also consistent
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43 254 with previous experiments of adaptation to limiting nutrient concentrations *in vitro*, in
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45 255 which the repressors of operons for sugar metabolism were inactivated (Horiuchi et al.
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47 256 1962; Zhong et al. 2004; Zhong et al. 2009).
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51 257 Overall the colonization experiments (fig. 3 and 5) imply that irrespectively of the founder
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53 258 clone being constitutive or regulated, in the long run *gat*-negative and *gat*-positive
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55 259 polymorphism is likely to emerge and be maintained in the gut.
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Microbiota composition affects *E. coli* abundance but not *E. coli* reverse evolution

The success of *E. coli* colonization, in terms of population size, is variable among mice. On the second colonization, approximately half of the animals are colonized with $\sim 10^8$ CFU/g of feces whereas the other half are colonized with $\sim 10^9$ CFU/g of feces (fig 6A). This variation could be mediated by the composition of the other members of mouse gut microbiota. We thus enquired whether we could observe an association between the loads of *E. coli* and the microbiota community composition in the gut of the colonized mice (fig. 6A-B). Remarkably, mice with low loads of *E. coli* (fig. 6C) exhibit a microbiota community composition significantly different from mice with high loads of *E. coli* (Fig. 6D). Because the great majority of the *E. coli* population is *gat*-negative the same applies to this fraction of the population. In contrast a remarkable constancy is observed for the abundance of *gat*-positive *E. coli* clones, whose numbers remain around 10^7 CFUs/g irrespectively of microbiota composition. We observed two main microbiota compositions (UPGMA clusters) that we term cluster I and cluster II (I in red and II in blue; fig. 6B, C-E). Cluster I shows significantly higher species richness (Shannon index= 4.7 ± 0.2 (2SE); $\chi^2_1=13.5$, $p<0.001$; fig. 6E), being enriched for several OTUs (two Firmicutes and one OTU from Bacteroidetes, Actinobacteria and Deferibacteres; fig. S10). Cluster II shows a lower Shannon index of $3.3 (\pm 0.3)$, being enriched for a single OTU belonging to Actinobacteria (Coriobacteriaceae), Proteobacteria (Entobacteriaceae) and Bacteroidetes (*Bacteroides*). Importantly, cluster I is associated with an abundance of *gat*-negative *E. coli* that decreases overtime, tending to $<10^8$ (per feces gram), which is significantly lower than that observed in cluster II (with average abundance of 10^9 per feces gram; $\chi^2_1=33.6$, $p<0.001$). These results suggest that *gat*-negative *E. coli* may compete with other members

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284 of the microbiota for resources and that the *gat*-positive *E. coli*, which acquired mutations
285 leading to increase expression of the *gat* operon and increased growth on galactitol, may be
286 selected for because they avoid such competition. As galactitol supplementation and the
287 microbiota independently affect the abundances of *gat*-positive and *gat*-negative *E. coli*
288 (respectively), these results appear to suggest that the two phenotypes occupy different
289 niches.

290 The theoretical model suggests that the equilibrium ratio of *gat*-positive to *gat*-
291 negative is dependent on the relative abundance of galactitol to other undefined resources
292 (see Materials and Methods) and that the abundance of these two phenotypes can vary
293 independently as these consume different resources. Interestingly, in the galactitol
294 supplementation experiment (fig. 4A-C), we see that increasing galactitol concentration
295 leads to an increase in the abundance of *gat*-positive bacteria ($\chi^2_1=6.4, p=0.01$), without
296 affecting the abundance of *E. coli* ($p>0.05$). Conversely, in the colonization experiment
297 (fig. 3B) it is the abundance of *gat*-negative bacteria that varies across mice, while the
298 abundance of *gat*-positive bacteria remains relatively constant after their emergence (fig.
299 6A). This suggests that there is variation in the concentrations of the unidentified resources
300 that are consumed by the *gat*-negative *E. coli*.

301
302 **Natural polymorphism for galactitol consumption in indigenous Enterobacteriaceae of**
303 **mice and humans**

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305 The findings reported here suggest that the gut ecology may favor natural
306 polymorphism for galactitol metabolism in its microbiota. To test for this we sampled

Enterobacteriaceae clones from the natural microbiota of our host model organism (fecal samples of mice untreated with antibiotics in the IGC animal house ($n=20$)) and from the gut microbiota of healthy humans with no controlled diet ($n=9$). Variation for this trait was found to be segregating across mice and man (Table 1). In 1 out of 9 human hosts, polymorphism for the ability to consume galactitol was detected, with 58% of the clones being *gat*-negative. In 5 humans the *gat*-negative phenotype was present at frequencies above 90%, with some showing a frequency of ~99%. Only 3 humans showed no *gat*-negative Enterobacteriaceae. From 20 mice assayed, one was monomorphic for the *gat*-negative phenotype, one showed an intermediate frequency for this phenotype and the rest showed only *gat*-positive clones.. These results indicate that the level of polymorphism at this locus may be influenced by antibiotic treatment or the strain background used for gut colonization. Nevertheless, in more natural conditions trait polymorphisms can also be found. Interestingly, in a comparative genomics survey of genes under positive selection in *E. coli*, *gatC* was found to be a target of repeated replacement substitution driven by positive selection (Chattopadhyay et al. 2009).

Conclusions

Overall our results highlight how evolutionary and ecological processes interact to shape the genetic and phenotypic composition of commensal bacteria within a natural ecosystem. At the evolutionary level, we find that a strong-mutation-strong-selection-strong-drift regime of adaptation can lead to the maintenance of polymorphism through frequent episodes of reverse evolution, which defy Dollo's law of evolutionary irreversibility (Gould 1970; Marshall et al. 1994). This regime of adaptation has been poorly studied, at both the theoretical and experimental levels, but may be important for host-associated microbes as

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330 their population dynamics often involves transmission bottlenecks (strong-drift) followed
331 by expansion to large population sizes (strong-mutation-strong-selection). Interestingly,
332 reversion has also been shown to rapidly occur for immune escape mutations in HIV
333 (Friedrich et al. 2004). Our study reveals that phenotypic and genetic reversals may be
334 common in bacteria colonizing the mammalian intestine. Such scenario may have been
335 difficult to predict, as bacteria could be expected to respond to changing environments by
336 gene regulation and not through *de novo* mutation, as we showed here. One context where
337 reversion is pivotal regards antibiotic resistance. Reversion of antibiotic resistant bacteria to
338 sensitivity has been more rarely observed than what one would desire (Andersson and
339 Hughes 2010). However, since most studies have been done *in vitro*, it would be important
340 to determine if the rate of reversion of resistance *in vivo* could be as high as the one we
341 found here for a metabolic trait (Björkman et al. 2000).

342 At the ecological level, our results indicate that resource specialization may allow new
343 emerging strains to avoid competition with other members of the microbiota (Conway and
344 Cohen 2015), even if to stably colonize the gut at low abundances. Our results further
345 highlight that precise dietary supplementations may allow for controlling the colonization
346 level of specific strains.

347 In sum, the evolutionary invasion and ecological stability of *de novo* emerging clones in
348 genetic identical hosts, nurtured in identical conditions, was highly reproducible, despite
349 being accompanied by significant differences in the level of microbiota species richness
350 between hosts.

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Material and Methods

Bacterial strains and culture conditions

All strains used in this study were derived from MG1655, a K12 commensal strain of *Escherichia coli* (Blattner et al. 1997).

Ancestors of the first colonization: the two ancestors of the first colonization (Fig. 3A), described in a previous experiment (Barroso-Batista et al. 2014), were DM08-YFP and DM09-CFP (MG1655, *galK::YFP/CFP* amp^R , *str*^R (*rpsL150*), *ΔlacIZYA*)).

Ancestors of the second colonization: strains JB19-YFP and JB18-CFP (MG1655, *galK::YFP/CFP* cm^R , *str*^R (*rpsL150*), *ΔlacIZYA*, Ins (1bp) *gatC*) were used for the second colonization (Fig. 3B) and were previously described in (Lourenço et al. 2016). These strains differ from the ancestral MG1655 fluorescent strains DM08-YFP and DM09-CFP by a mutation in the *gatC* gene (1bp insertion in the coding region), rendering it unable to metabolize galactitol. To construct these strains the ampicillin resistance cassette in the ancestral strains DM08-YFP and DM09-CFP was replaced with a chloramphenicol resistant cassette using the Datsenko and Wanner method (Datsenko and Wanner 2000). The yellow (*yfp*) and cyan (*cfp*) fluorescent genes linked to cm^R were then transferred by P1 transduction to a derivative of clone 12YFP (Barroso-Batista et al. 2015), an evolved clone of DM08-YFP, isolated after 24 days of adaptation in the gut of WT mice, that carried an insertion of 1bp in *gatC* and a large duplication. During the genetic manipulations the large duplication was lost, confirmed by whole genome sequencing, leaving this clone with a single mutation in *gatC* (Lourenço et al. 2016).

Ancestor of the third colonization: the strain used to start the third colonization was isolated from population 2.4 of the second colonization at day 24 post-colonization. This strain (named 19CFP (Lourenço et al. 2016) has a 2bp insertion (+TC) in *gatC* which

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reconstructed the open reading frame allowing it to recover the ability to consume galactitol (see below details about the phenotype confirmation). Additionally, this strain has the following mutations: an IS2 insertion in the coding region of *ykgB* and a 6790bp deletion from gene *intZ* to *eutA*.

Reconstruction of the transcriptional regulator of the galactitol metabolism (*gatR*): the *gatR* coding region is interrupted by an IS element in *E. coli* MG1655, and therefore in all strains used in this study. This insertion inactivates *gatR* leading to the constitutive expression of the *gat*-operon. To restore the function of the gene, and therefore the ability to regulate the operon, a copy of the *gatR* native sequence (obtained from the strain *E. coli* HS) was used to replace the non-functional copy. This was accomplished by the *SacB-TetA* counter selection method as previously described (Li et al. 2013)). Briefly, the *SacB-TetA* fragment with the overhangs for recombination was obtained by target PCR using the primers *gatR_SacB_F* / *gatR_SacB_R* (Table S2) and genomic DNA of *E. coli* XTL298 as a template. The resulting PCR product was then used to replace the non-functional copy of *gatR* present in strains DM08 and DM09 by the Wanner method (Datsenko and Wanner 2000). The DNA fragment containing the native sequence of *gatR* was obtained by target PCR using the primers *yegS_F* / *gatD_R* and *E. coli* HS as a template. Finally, this fragment was used to replace the counter selection cassette in the *gatR* locus of the recipient strains DM08 and DM09 by the Wanner method (Datsenko and Wanner 2000) and plated on double-counter-selective medium. Candidates for successful restoration of *gatR* were screened by PCR and tested for antibiotic resistances to confirm the loss of the cassette and the plasmid before sequencing.

To distinguish between *gat*-negative and *gat*-positive bacteria we used the differential medium MacConkey agar supplemented with galactitol 1% and streptomycin (100µg/mL).

Plates were incubated at 30°C. The frequency of *gat*-negative mutants was estimated by counting the number of white (auxotrophic for galactitol) and red colonies. On average around 1000 colonies per time point were plated to estimate the frequency of *gat*-negative mutants, therefore the minimal frequency detected by this method is ~0.5%.

***In vivo* Experimental Evolution**

In order to study *E. coli*'s adaptation to the gut we used the classical streptomycin-treated mouse model of colonization (Krogfelt et al. 2013) and performed the evolution experiments using the same conditions as before (Barroso-Batista et al. 2014; Barroso-Batista et al. 2015; Lourenço et al. 2016). Briefly, 6- to 8-week old C57BL/6 male mice raised in specific pathogen free conditions were given autoclaved drinking water containing streptomycin (5g/L) for one day. After 4 hours of starvation for water and food, the animals were gavaged with 100µL of a suspension of 10⁸ colony forming units of a mixture of YFP- and CFP-labeled bacteria (ratio 1:1) grown at 37°C in brain heart infusion medium to OD₆₀₀ of 2. After the gavage, all animals were housed separately and both water with streptomycin and food were returned. Mice fecal samples were collected for 24 days and diluted in PBS, from which a sample was stored in 15% glycerol at -80°C and the remaining was plated in Luria Broth agar supplemented with streptomycin (LB plates). Plates were incubated overnight at 37°C after which fluorescent colonies were counted using a fluorescent stereoscope (SteREO Lumar, Carl Zeiss) to assess the frequencies of CFP- and YFP-labelled bacteria. These fluorescent proteins are used as neutral markers with which we can follow to detect the emergence of beneficial mutations, since these markers hitchhike with the beneficial mutations that spread in the populations (Hegreness 2006).

***In vivo* competitive assays**

To test the *in vivo* advantage of the evolving populations at the last time point of the evolution experiment (Fig. 1A) samples of either YFP or CFP clones isolated from day 24 (sub-populations) were competed against the respective ancestor labeled with the opposite fluorescent marker (n=2 for each population). These sub-populations were composed of mixtures of approximately 30 colonies with the same fluorescent marker isolated after plating the appropriate dilution of mice fecal pellets. The mixtures of clones were then grown in 10 mL of LB supplemented with chloramphenicol (100µg/mL) and streptomycin (100µg/mL) and stored in 15% glycerol at -80°C. *In vivo* competitions of evolved sub-populations against the ancestral were performed at a ratio of 1 to 1, following the same procedure described above for the evolution experiment. Mice fecal pellets were collected for 3 days, diluted in PBS and frozen in 15% glycerol at -80°C. Frequencies of CFP- and YFP-labelled bacteria were estimated using a fluorescent stereoscope (SteREO Lumar, Carl Zeiss). The selective coefficient (fitness gain) of these mixtures of clones *in vivo* (presented in Fig. 1B) was estimated as: $s_b = \ln \left(\frac{Rf_{ev/anc}}{Ri_{ev/anc}} \right) / t$, where s_b is the selective advantage of the evolved clone, $Rf_{ev/anc}$ and $Ri_{ev/anc}$ are the ratios of evolved to ancestral bacteria in the end (f) or in the beginning (i) of the competition and t is the number of generation per day. We assumed $t=18$ generations, in accordance with the 80 minute generation time estimated in previous studies on *E. coli* colonization of streptomycin-treated mouse (Poulsen et al. 1995; Rang et al. 1999).

Whole genome re-sequencing and mutation prediction

Clone analysis: Two *gat*-positive clones (revertant) isolated from the second colonization (day 6 of population 2.15) were sequenced to test for the presence of additional mutations, besides the one reverting the *gat*-phenotype. Two independent colonies were used to inoculate 10 mL of LB and incubated at 37°C with agitation for DNA extraction (following a previously described protocol (Wilson 2001)). The DNA library construction and sequencing was carried out at the IGC sequencing facility. Each sample was paired-end sequenced on an Illumina Mi Seq Benchtop Sequencer. Standard procedures produced data sets of Illumina paired-end 250 bp read pairs. Genome sequencing data have been deposited in the NCBI Read Archive, <https://www.ncbi.nlm.nih.gov/bioproject/PRJNA295680>. Mutations were identified using the BRESEQ pipeline (Deatherage and Barrick 2014). To detect potential duplication events we used ssaha2 (Ning et al. 2001) with the paired end information. This is a stringent analysis that maps reads only to their unique match (with less than 3 mismatches) on the reference genome. Sequence coverage along the genome was assessed with a 250 bp window and corrected for GC% composition by normalizing by the mean coverage of regions with the same GC%. We then looked for regions with high differences (>1.4) in coverage. Large deletions were identified based on the absence of coverage. For additional verification of mutations predicted by BRESEQ, we also used the software IGV (version 2.1) (Robinson et al. 2011).

Population analysis: DNA isolation, library construction and sequencing were carried as described above for the clone analysis except that now it derived from a mixture of >1000 clones per population grown in LB agar. Two populations, from the evolution experiment, were sequenced: 2.14 and 2.15. Those were sequenced for three time points during the adaptive period (generation 198 (day 11), generation 306 (day 17) and

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3 471 generation 432 (day 24)). Mean coverage per sample was between ~100x and ~300x for
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5 472 population 2.14 and between ~70x and ~115x for population 2.15. Mutations were
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7 473 identified using the BRESEQ pipeline (version 0.26) with the polymorphism option on.
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9 474 The default settings were used except for: a) requirement of a minimum coverage of 3 reads
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11 475 on each strand per polymorphism; b) eliminating polymorphism predictions occurring in
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13 476 homopolymers of length greater than 3; c) polymorphism predictions with significant
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15 477 ($P=0.05$) strand or base quality score bias were discarded. Data presented in Table S1.
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23 479 **Diet supplementation experiment**

24 480 To test the selective pressure exerted by galactitol, when present in the gut, we
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26 481 performed *in vivo* competitions between *gat*-positive (DM08) and *gat*-negative (JB18)
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28 482 bacteria, while supplementing the diet with this sugar. Competition experiments were
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30 483 performed as described above except that the initial frequency of *gat*-positive bacteria was
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32 484 ~10% and the drinking water was supplemented with 0%, 0.01% or 0.1% of galactitol (n=3
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34 485 for each galactitol concentration) together with streptomycin (5g/L). Fecal samples were
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36 486 collected daily and plated to access the frequencies of both phenotypes during four days.
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43 488 **Estimate of gain-of-function mutation rate from *gat*-negative to *gat*-positive phenotype**

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45 489 Strains JB19 (*gatC* 1bp Ins) and 4YFP (*gatZ* IS Ins) were grown overnight in 10 mL
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47 490 of LB at 37°C with aeration. After growth, the total number of cells in the cultures was
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49 491 measured by flow cytometry (using BD LSR Fortessa; BD Biosciences) and approximately
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51 492 1000 cells were used to inoculate 1 mL of LB (10 replicates of each strain) and incubated
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53 493 overnight. Aliquots of each replicate tube were plated in LB agar and MM agar
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55 494 supplemented galactitol 1% and incubated overnight at 37°C. The number of spontaneous
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gat-positive mutants and total number of cells grown on LB were used to estimate the mutation rate using the maximum likelihood approach as implemented in FALCOR (Hall et al. 2009).

We measured the mutation rate to re-acquire the ability to consume galactitol, since this locus was previously demonstrated to be a hotspot for loss of function mutations (Lourenço et al. 2016). The fluctuation test revealed that the spontaneous rate at which the ancestor of the second colonization (*gat*-negative) recovers the ability to consume galactitol is 9.7×10^{-9} (95% CI, $[3 \times 10^{-9}, 1.9 \times 10^{-8}]$) per generation. Considering that the estimated spontaneous rate of small indels in *E. coli* is $\sim 2 \times 10^{-11}$ (Lee et al. 2012), and that about one third of such mutations in a region of ~ 30 aminoacids of *gatC* could restore the open reading frame in the ancestral clone of the second colonization, the expected rate of mutation towards a *gat*-positive phenotype is $\sim 6 \times 10^{-10}$. This estimate is between 5 to 15 times lower than that observed in the fluctuation assay, suggesting that a high mutation rate may have also contributed to the frequent re-emergence of *gat*-positive bacteria in the evolution experiment.

Identification of adaptive mutations and estimate of haplotype frequencies in selected populations of the evolution experiment

In order to estimate the haplotype frequencies depicted in Fig. 2A-B two complementary strategies were employed. In addition to the WGS of the populations, targeted PCR of the identified parallel mutations was performed. For the targeted PCR, 20 to 80 clones from different time points were screened (from populations 2.14 and 2.15) using the same primers and PCR conditions as in (Barroso-Batista et al. 2014; Lourenço et al. 2016). Because all target mutations correspond to IS insertions an increase in size of the

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519 PCR band is indicative of the presence of an IS. Frequency of *gat*-revertant was estimated
520 by plating in differential media (MacConkey supplemented with 1% galactitol).
521 Frequencies are depicted in Table S1.

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523 **Microbiota analysis**

524 We extracted DNA from mice fecal samples from two experiments: the second
525 colonization (Fig. 3B) and the galactitol supplementation experiment (Fig. 4A-B). For the
526 second colonization we extracted DNA from mice where *gat*-positive *E. coli* could be
527 detected (12 out of 15), at days 12 and 17 or 18 post-colonization (*i.e.* 2 samples per
528 mouse). For the galactitol supplementation experiment, we extracted DNA at days 1, 2, 3, 4
529 and 7 post-colonization, for the nine mice involved in this experiment. Moreover, we
530 extracted and sequenced a negative control (extraction without feces).

531 Fecal DNA was extracted with a QIAamp DNA Stool Mini Kit (Qiagen), according to
532 the manufacturer's instructions and with an additional step of mechanical disruption
533 (Thompson et al. 2015). 16S rRNA gene amplification and sequencing was carried out at
534 the Gene Expression Unit from Instituto Gulbenkian de Ciência, following the service
535 protocol. For each sample, the V4 region of the 16S rRNA gene was amplified in triplicate,
536 using the primer pair F515/R806, under the following PCR cycling conditions: 94°C for 3
537 min., 35 cycles of 94°C for 60s, 50°C for 60s and 72°C for 105s, with an extension step of
538 72°C for 10min (Caporaso et al. 2011; Caporaso et al. 2012). Samples were then pair-end
539 sequenced on an Illumina MiSeq Benchtop Sequencer, following Illumina
540 recommendations.

541 QIIME (Caporaso et al. 2010) was used to analyze the 16S rRNA sequences by first
542 quality filtering the samples with a phred quality threshold of 20 (with script

split_libraries_fastq) and then OTU picking by assigning operational taxonomic units at 97% similarity against the Greengenes database (DeSantis et al. 2006); using script pick_open_reference_otus with default settings, but enabling reverse strand matches). The OTU table generated by the latter script was then filtered in two steps. First, with script remove_low_confidence_OTUs, from the Microbiome helper repository (Comeau et al. 2017), to remove potentially spurious OTUs. Second, to remove contaminants (Salter et al. 2014) identified in the negative control, we filtered out any OTUs that were present in both the negative control and 2 or less samples at a frequency <0.01 . The OTUs that were kept in the final OTU table and that were shared between negative control and samples had a maximum frequency of 0.004 in the negative control, were present at a frequency of >0.01 in at least 5 samples and had an average frequency in samples that was at least one order of magnitude higher than the observed in the negative control. The OTU table obtained after these two steps was then used for the analysis below. To understand the clustering pattern of the mouse gut microbiota in each experiment, we used the script jackknifed_beta_diversity to build rarefied OTU tables (at 75% of the minimum read depth in each experiment, as advised on QIIME's website). This built 100 rarefied OTU tables, computed weighted UniFrac (Lozupone and Knight 2005) and generated a consensus tree (per experiment) with jackknife support of tree nodes. Consensus trees were then plotted with R package ggtree together with the OTU relative abundance (Yu et al. 2017). To calculate alpha diversity, the OTU tables for each experiment were rarefied 100 times to even depth (multiple_rarefactions_even_depth), alpha diversity was estimated and collated (scripts alpha_diversity and collate_alpha) and the mean for each sample-metric combination was calculated (plotted in fig. S5 and S11). To test whether there was an effect

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of cluster (second colonization) or galactitol concentration (galactitol supplementation experiment), we used linear mixed effects models with mouse as a random effect.

For the second colonization, UPGMA clustering identified two main clusters (Fig. 6). Thus, we then used linear discriminant analysis effect size (LefSe) to identify if there were any taxa that were differentially abundant between the two clusters (Segata et al. 2011). The input data for LefSe (fig. S10) was obtained after filtering the 97% OTUs for those that were present in at least 6 samples (*i.e.* at least 3 mice) and at a frequency that was above 0.5%, which corresponds to ~100 reads.

Furthermore, for the galactitol supplementation experiment, we used the R package phyloseq (McMurdie and Holmes 2013) to compute the following beta-diversity metrics: Jaccard index, Bray-Curtis, unweighted and weighted UniFrac. We then tested for an effect of galactitol concentration on each beta-diversity matrix with PERMANOVA by using the adonis function with the vegan R package, with 1000 permutations (Anderson 2001; Oksanen et al. 2009). These procedures were applied to data from day 3, 4 and 7 post-colonization or each of the last three days separately (as shown in fig. S6-9).

Natural fecal isolates collection and phenotyping

Fecal samples were collected from 9 specific pathogen free mice of different litters (strain C57BC-67) and from 9 healthy humans. Samples were weighted and re-suspended in PBS with 15% glycerol before storage at -80°C. To isolate lactose fermenting Enterobacteriaceae (red colonies) appropriate dilutions of the fecal samples were plated on MacConkey plates supplemented with 0.4% lactose and incubated overnight at 37°C. The frequency of *gat*-positive bacteria among the lactose fermenting Enterobacteriaceae was estimated by replica-plating ~96 isolated clones on M9 minimal medium supplemented

with 0.4% galactitol. The M9 agar plates were incubated at 30°C and bacterial growth was scored after 24h, 48h, 72h and 96h. The ability to metabolize galactitol was tested by three independent trials.

Statistical Analysis

To analyze temporal dynamics data, we used linear mixed models, with mouse as a random effect. Where required, data were transformed to meet assumptions made by parametric statistics. All analysis were performed in R (R Core Team 2016) .

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55 791 J.B., M. L., D.G. performed the experiments. A.S., R.R., M.L., I.G. analyzed the
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57 792 data and wrote the manuscript.
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793 **Figures:**

794 **Fig. 1. Rapid adaptation characterizes the evolution of colonizing the mouse gut. (A)**

795 Change in frequency of a neutral fluorescent marker (YFP) along 24 days of colonization.

796 **(B)** Fitness increase (per generation) measured by direct *in vivo* competitive assays of
797 evolved clones against the ancestor (n=2).

799 **Fig. 2. Reverse evolution leads to the recurrent regain of a previously lost phenotype,**

800 **by compensatory and back-mutation. (A and B)** Muller Plots showing the emergence of

801 polymorphism in two evolving populations (pop 2.14 (A) and pop 2.15 (B)), as numbered

802 in fig 1A) where regain of function (ability to metabolize galactitol) is detected - *gat*-rev.

803 Yellow and blue shaded areas are proportional to the frequencies of YFP- and CFP-labelled

804 bacteria. Darker tones of yellow and blue denote the accumulation of adaptive mutations in

805 the respective background. Days shown below the graph depict the temporal points where

806 haplotype frequencies were estimated (see Table S1) through a combination of different

807 approaches. Numbers in red represent temporal points where frequencies were estimated by

808 WGS of the population and numbers in black represent time points where frequencies were

809 obtained by target PCR of collections of clones. Dotted areas represent the proportion of

810 the population that reverted the ancestral *gat*-negative phenotype and was determined by

811 phenotypic testing. **(C)** The genetic basis of the *gat*-reversion phenotype (*gat*-rev) involves

812 both compensatory (represented by alleles comp1-6) and back-mutation (allele rev). Six

813 different alleles were found in pop 2.14 at day 11 (comp1 to comp6). After 6 days (day17)

814 only alleles comp1 (~1% frequency) and comp2 (~13% frequency) could be detected by

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3 815 WGS; we note however that the frequency of *gat*-rev phenotype in the same time-point was
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5 816 considerably higher (~50%). This suggests that the polymorphism in the *gatC* locus
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8 817 estimated by WGS was considerably underestimated in this time-point. On day 24, comp2
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10 818 was the only allele detected. Days 6 and 11 of pop 2.15 revealed two different alleles
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12 819 (comp3 and rev). On day 17 and 24 only the rev allele was found.
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17 821 **Fig. 3. Reacquisition of the ancestral metabolic ability is pervasive across**
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19 822 **independently evolving populations. (A-C)** The ancestral genotype and phenotype of
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21 823 each colonization is indicated above the respective graph. (A) Massive selection to
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23 824 inactivate the *gat*-operon characterized the first colonization, yet in half of the populations
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25 825 the frequency of *gat*-positive bacteria was kept above 1%. (B) In the second colonization,
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27 826 initiated with a *gat*-negative strain, reacquisition of the ancestral metabolic ability was
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29 827 pervasive across independently evolving populations re-establishing the polymorphism for
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31 828 the *gat* phenotype. (C) The third colonization, initiated with a *gat*-positive clone isolated
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33 829 from the second colonization, exhibited the re-emergence of *gat*-negative mutants, as in the
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35 830 first colonization.
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43 832 **Fig. 4. Diet supplementation with galactitol increases the frequency and abundance of**
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45 833 ***gat*-positive bacteria, without affecting total *E. coli* density and microbiota**
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47 834 **composition.** Temporal dynamics for the frequency of *gat*-positive *E. coli* (A). Temporal
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49 835 dynamics for the density of *gat*-positive (solid lines) and total *E. coli* (dashed lines); error
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51 836 bars correspond to standard error (B). Analysis of the mouse gut microbiota shows no
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53 837 evidence for microbiota clustering being influenced by galactitol supplementation (C).
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55 838 Dendrogram on the left represents UPGMA clustering of weighted Unifrac (beta diversity),
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with * indicating when jackknife node support was above 0.9 (100 subsamples of the OTU table). Consistently, alpha diversity is also not influenced by galactitol supplementation (fig. S5). Bars on the right indicate the relative frequency of the different operational taxonomic units (OTUs; at 97% identity; full legend is provided in fig. S3). Data from days 3, 4 and 7 was used in this analysis, as after day 3 both the *gat*-positive frequency and the mouse microbiota had started to stabilize (fig. S4 shows the temporal dynamics of gut microbiota per mouse). Colors correspond to: blue – no galactitol supplementation; yellow – 0.01% galactitol; red – 0.1% galactitol (galactitol was diluted in the mice drinking water). Different symbols correspond to different mice.

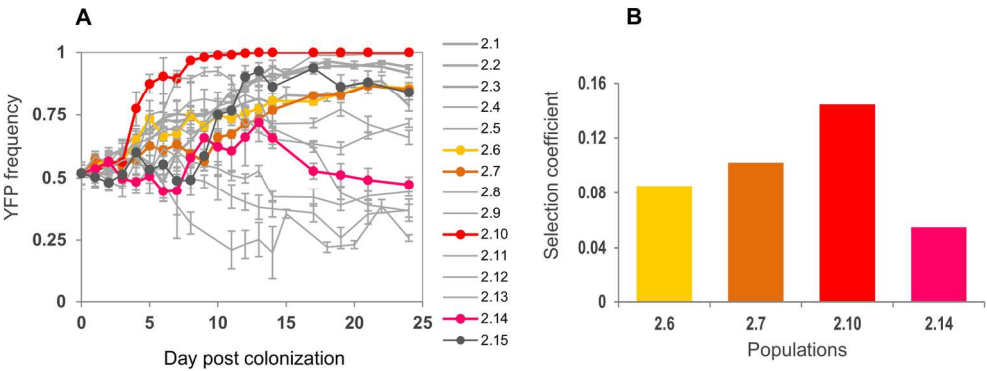
Fig. 5. Emergence of *gat*-constitutive strains in the guts of mice colonized with a clonal population of bacteria where the repressor of the galactitol operon (*gatR*) has been restored. (A) Frequency of *gat*-constitutive mutants that emerged *de novo* in all colonized mice (n=6) during the first 5 days of colonization (A). (B) Frequency of *gat*-constitutive mutants at day 19 in the same mice.

Fig. 6. Microbiota composition affects the density of *gat*-negative, but not *gat*-positive *E. coli*. Microbiome analysis was carried for mice from the second colonization (fig. 1 and 3B) in which *gat*-positive revertants had emerged. Moreover, the densities of *gat*-positive and *gat*-negative *E. coli* were estimated. The gut microbiota affects the densities of *gat*-negative but not *gat*-positive *E. coli* (A). Dot plots indicate the densities of *E. coli* with each phenotype for the corresponding mouse-day combination. Line and shading indicate the mean and standard error. For *gat*-negative, two lines are shown as the bacterial densities differ significantly between red and blue clusters. Conversely, for *gat*-positive, only one

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3 863 line is shown as there is no significant difference. (B) Analysis of the mouse gut microbiota
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5 864 divides mice into two main clusters, I (red) and II (blue). Dendrogram on the right
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8 865 represents UPGMA clustering of weighted Unifrac (beta diversity), with * indicating when
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10 866 jackknife node support was above 0.9 (100 subsamples of the OTU table). Bars on the left
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12 867 indicate the relative frequency of the different operational taxonomic units (OTUs; at 97%
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15 868 identity). Labeled taxa are enriched in either cluster I (red labels) or II (blue labels), as
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17 869 identified by LEfSe (full legend is provided in fig. S3 and LefSe results in fig. S10). (C-D)
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19 870 Temporal dynamics of *gat*-negative and *gat*-positive *E. coli* showing that the densities of
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21 871 *gat*-negative *E. coli* are lower in mice with the cluster I microbiota (C), than in mice with
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23 872 the cluster II microbiota (D) (cluster:time interaction: $\chi^2_1=33.6, p<0.001$). Conversely, the
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25 873 densities of *gat*-positive clones remain constant across all mice (cluster:time interaction:
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27 874 $\chi^2_1=3.3, p=0.07$ and cluster single effect: $\chi^2_1=1.0, p=0.31$). (E) Microbiota diversity
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29 875 (estimated as Shannon Index) is significantly lower in cluster II than in cluster I (similar
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31 876 observations are made for other diversity indices; see fig. S11).
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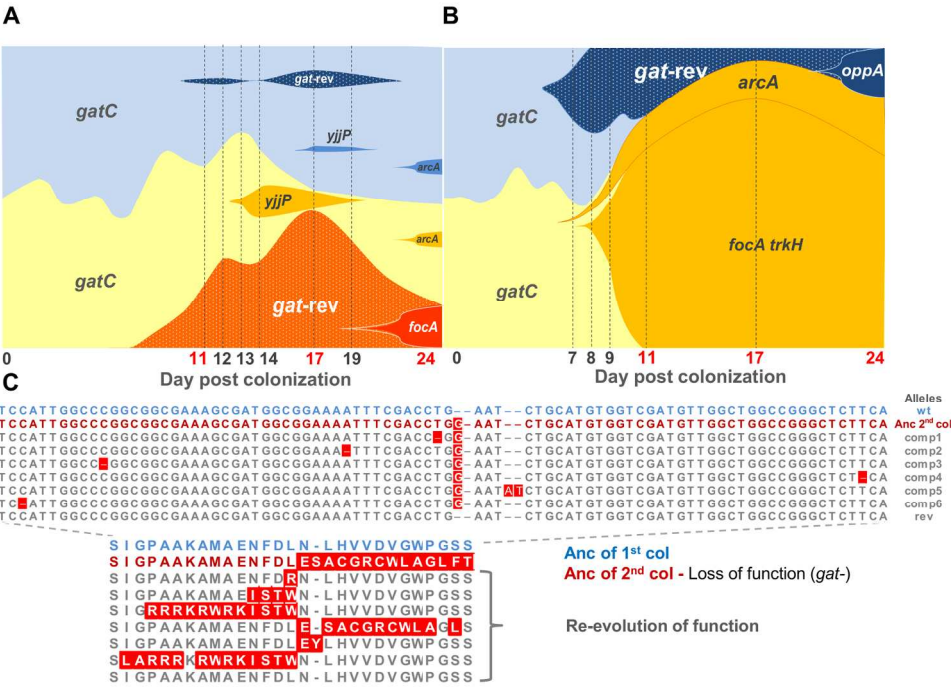
Table 1. Polymorphism for galactitol metabolism in natural fecal isolates.

Sample #	<i>gat</i> -pos	<i>gat</i> -neg	Total	Frequency of <i>gat</i> -neg	2SE
Mouse 1	1	143	144	0.99	0.01
Mouse 2	125	19	144	0.13	0.06
Mouse 3-20	144	0	144	0.00	0.01
Human 1	189	3	192	0.02	0.02
Human 2	12	180	192	0.94	0.03
Human 3	192	0	192	0.00	0.01
Human 4	2	190	192	0.99	0.01
Human 5	11	181	192	0.94	0.03
Human 6	81	111	192	0.58	0.07
Human 7	1	191	192	0.99	0.01
Human 8	19	173	192	0.90	0.04
Human 9	192	0	192	0.00	0.01



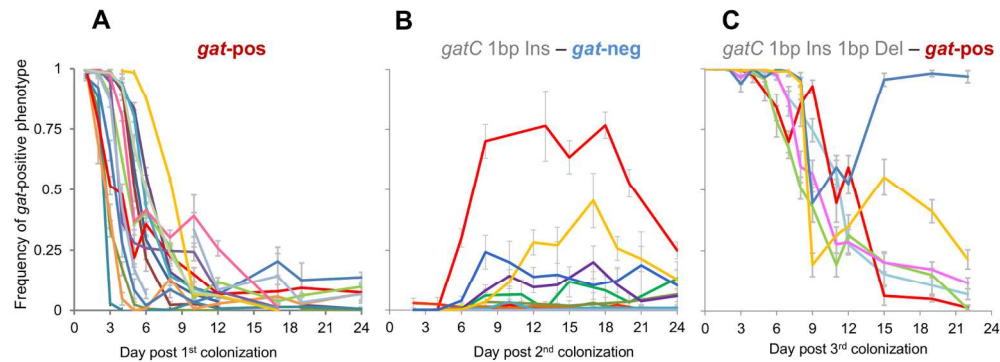
Rapid adaptation characterizes the evolution of colonizing the mouse gut.

168x64mm (300 x 300 DPI)



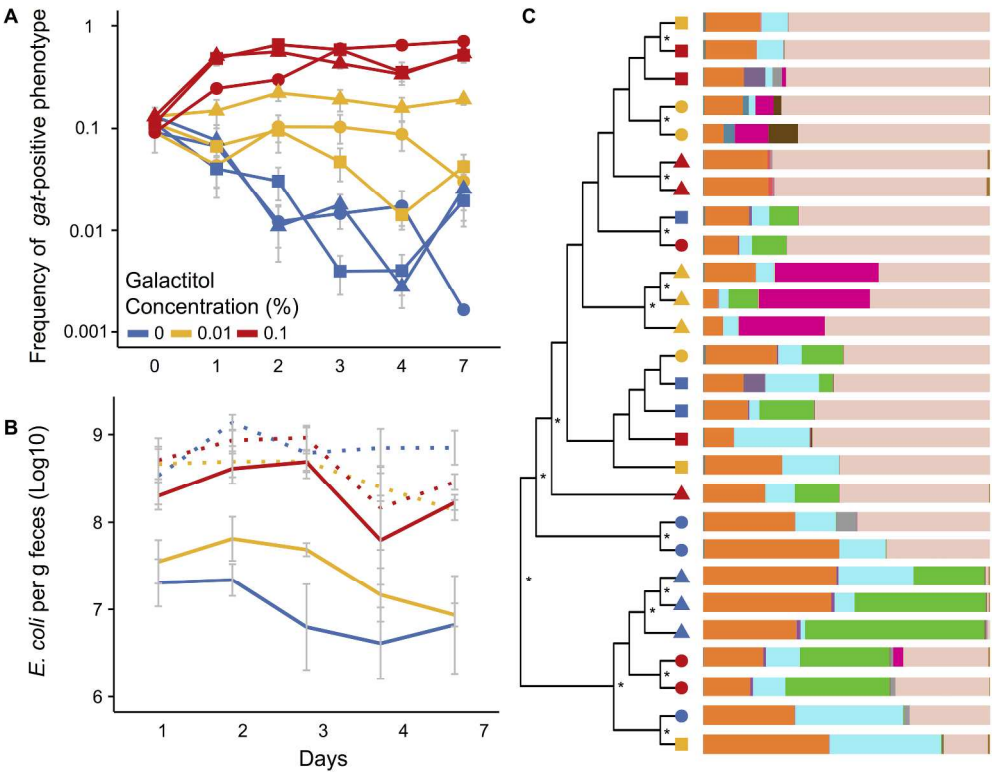
Reverse evolution leads to the recurrent regain of a previously lost phenotype, by compensatory and back-mutation.

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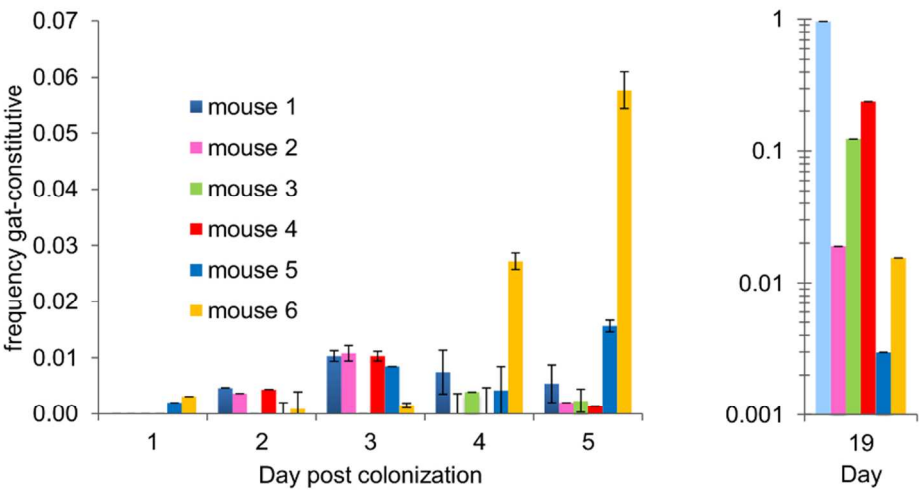
Reacquisition of the ancestral metabolic ability is pervasive across independently evolving populations.

168x63mm (300 x 300 DPI)



Diet supplementation with galactitol increases the frequency and abundance of *gat*-positive bacteria, without affecting total *E. coli* density and microbiota

131x101mm (600 x 600 DPI)



Emergence of gat-constitutive strains in the guts of mice colonized with a clonal population of bacteria where the repressor of the galactitol operon (gatR) has been restored.

82x43mm (300 x 300 DPI)

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Microbiota composition affects the density of gat-negative, but not gat-positive *E. coli*.