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Import of aspartate and malate by DcuABC drives H₂/fumarate respiration to promote initial *Salmonella* gut-lumen colonization in mice

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Abstract

Initial enteropathogen growth in the microbiota-colonized gut is poorly understood. *Salmonella* Typhimurium is metabolically adaptable and can harvest energy by anaerobic respiration using microbiota-derived hydrogen (H₂) as an electron donor and fumarate as an electron acceptor. As fumarate is scarce in the gut, the source of this electron acceptor is unclear. Here, transposon sequencing analysis along the colonization-trajectory of *S.*Typhimurium implicates the C₄-dicarboxylate antiporter DcuABC in early murine gut colonization. In competitive colonization assays, DcuABC and enzymes that convert the C₄-dicarboxylates aspartate and malate into fumarate (AspA, FumABC), are required for fumarate/H₂-dependent initial growth. Thus, *S.*Typhimurium obtains fumarate by DcuABC-mediated import and conversion of L-malate and L-aspartate. Fumarate reduction yields succinate, which is exported by DcuABC in exchange for L-aspartate and L-malate. This cycle allows *S.*Typhimurium to harvest energy by H₂/fumarate respiration in the microbiota-colonized gut. This strategy may also be relevant for commensal *E. coli* diminishing the *S.*Typhimurium infection.

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Author contributions

Conceived and designed the experiments: BDN, WDH, MC, BC, JH, EG. Performed the experiments: BDN, JH, RB, SM, JR, CM, LH, FB, MF, CF, PPE, EG, LL, PK, LF. Analyzed the data: BDN, MCV, JH, PPE, BC, MC, SS, EG, PK, WDH. Contributed reagents/materials/analysis tools: MM, SP, JAV, US. Computational analysis: MCV, MC, BC, SS. Wrote the paper: BDN, WDH.

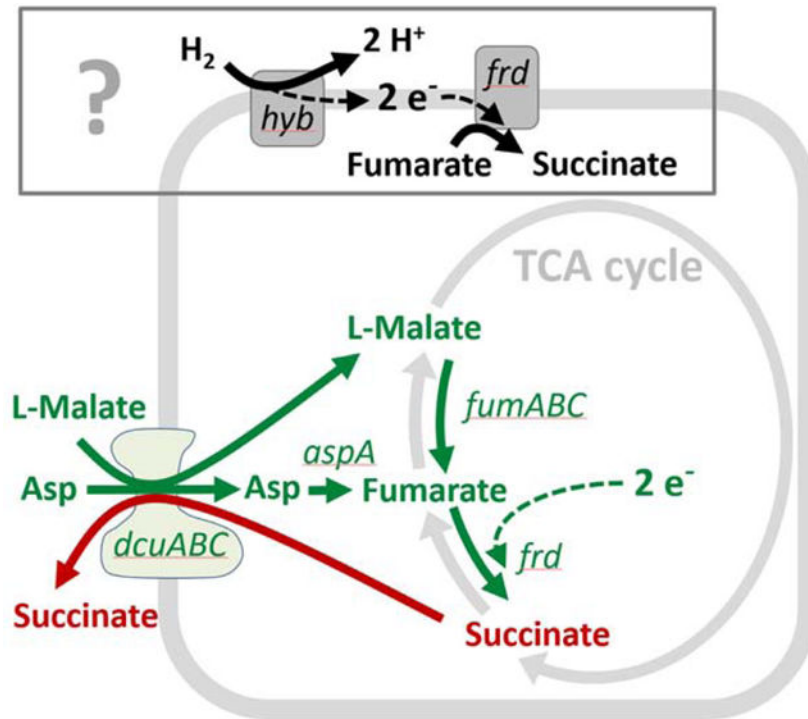
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Declaration of Interests

The authors have no conflicting financial interests.

Graphical Abstract



eTOC Blurp

Bidong Nguyen and colleagues show how *Salmonella* Typhimurium grows after arrival in the gut. *Salmonella* Typhimurium imports aspartate and malate using the DcuABC transporters and converts these substrates into fumarate to fuel anaerobic H₂/fumarate respiration. Thereby, *Salmonella* can grow in face of an intact gut microbiota to establish the infection.

Introduction:

The mammalian gut limits microbial growth via chemical defenses, physical barriers and by depleting nutrients. Most microbial growth is observed in the large intestine, which harbors the bulk of the microbiota. This dense microbial consortium utilizes nutrients that are not depleted by the host, e.g. by fermenting complex carbohydrates. Importantly, the microbiota can limit pathogen growth in the large intestinal lumen (Stecher and Hardt, 2011, Buffie and Pamer, 2013). This colonization resistance (CR) is attributable to diverse mechanisms like microbiota-mediated nutrient depletion, release of pathogen-inhibitors (bacteriocins, lantibiotics, type VI secretion systems), blocking of binding sites, depletion of terminal electron acceptors, or vitamin B6 production and microbiota-mediated modulation of enterocyte metabolism (Fabich et al., 2008, Freter et al., 1983, Sonnenburg et al., 2005, Stecher, 2015, Stecher et al., 2013, Miki et al., 2017, Buffie and Pamer, 2013, Hecht et al., 2016, Byndloss et al., 2017). CR reduces the incidence of enteropathogen colonization in humans and mouse models, alike. Nevertheless, in some cases, enteropathogens succeed in colonizing the gut lumen in face of the microbiota. The underlying mechanisms of how

enteropathogens can grow before the onset of overt enteric disease are not completely understood.

Salmonella enterica serovar Typhimurium (*S. Typhimurium*) is a common enteropathogenic bacterium, which can infect a broad range of hosts, including livestock, humans and mice. While CR suppresses *S. Typhimurium* growth in the intestine of most exposed hosts, in some cases the pathogen can initiate gut luminal growth (Simonsen et al., 2009) (Stecher and Hardt, 2011, Buffie and Pamer, 2013). Studies from mouse infections and human outbreaks suggest that there is an initial phase of pathogen growth (termed “initial growth”) before the mucosa responds with overt enteropathy, characterized by pronounced phagocyte infiltration, re-modeled mucosal architecture and an altered gut luminal milieu (Wotzka et al., 2017) (Organization and Nations, 2002) (Maier et al., 2013). To fuel initial growth, *S. Typhimurium* can harvest energy by anaerobic hydrogen (H₂)/fumarate respiration (Maier et al., 2013). H₂ is a product of anaerobic microbiota metabolism and serves as an electron donor for the *hyb* hydrogenase (Maier et al., 2013) (Maier et al., 2014a) (Dubini et al., 2002) (Maier, 2005). The resulting protons feed the proton gradient to support ATP production (Unden et al., 2016) and the electrons are transferred to fumarate reductase (Frd) which reduces fumarate to succinate (Maier et al., 2013). However, it remained unclear how *S. Typhimurium* disposes of succinate and how it obtains fumarate. The free fumarate concentration in the gut lumen might be too low to support this mechanism of energy conservation (Spiga et al., 2017, Cummings et al., 1987).

Here, we employ an unbiased mutagenesis approach, gene deletion epistasis experiments and metabolite analyses to identify DcuABC-dependent uptake of malate and aspartate and their conversion into fumarate as the principle mechanism fueling anaerobic hydrogen (H₂)/fumarate respiration during initial growth.

Results:

RB-TnSeq screen implicates C4-dicarboxylate transport in initial growth.

To gain insights into the initial growth physiology of *S. Typhimurium* in the gut lumen, we have used strain SL1344, a well-established model for *Salmonella* gut colonization (Barthel et al., 2003) (Maier et al., 2013) (Wotzka et al., 2019) (Sana et al., 2016) and employed a random-barcoded transposon sequencing approach (RB-TnSeq; (Price et al., 2018, Wetmore et al., 2015)). Via transposome transformation (Langridge et al., 2009) (Canals et al., 2012), we generated a library of \approx 2000 randomly barcoded transposon insertion mutants of *S. Typhimurium* SL1344 (RB-TnSeq library; Fig. 1A, Fig. S1A). This number of mutants is close to the upper limit for parallel analysis in the same mouse (Maier et al., 2014b) (Maier et al., 2013). The RB-TnSeq mutations were mapped by sequencing of their unique barcodes in conjunction with their transposon’s insertion sites (Fig. S1B). To facilitate quality control, the inoculum pool was spiked with several controls (Fig. 1A; Key Resources Table). Wild type isogenic tagged strains (WITS, Key Resources Table; (Grant et al., 2008)) were used to flag cases where gut inflammation might confound mutant fitness analysis by inflicting bottlenecks in the gut-luminal *S. Typhimurium* population (Maier et al., 2014b) (Fig. 1A, Fig. S1C). An ampicillin resistant *hyb* mutant was included to verify H₂-dependent *S. Typhimurium* growth in the gut lumen. Titration series of RB-TnSeq-barcoded wild type *S.*

Typhimurium, and isogenic mutants (*hyb*, *ttss-1/ttss-2*, *cheY*) were also spiked in (Fig. 1A, Key Resources Table). These control strains featured unique RB-TnSeq-barcode sequences, which served as internal standards during the transposon barcode amplification and quantification procedure by tracking fitness of the RB-TnSeq mutants along the colonization trajectory.

The screen was performed in gnotobiotic mice associated with a low-complexity microbiota consortium (LCM mice; Star Methods (Maier et al., 2013)). LCM mice provide a well-defined and highly reproducible gut colonization system where the initial H₂-dependent growth of *S. Typhimurium* lasts for 2 days and which permits the fitness analysis of 2000 strains in parallel during this colonization process ((Maier et al., 2013); Fig. S1C). We orally infected 6 LCM mice with our inoculum pool composed of ≈2000 strains (Fig. 1A, 5 × 10⁶ cfu total) and subsequently recovered mutant mixtures from feces collected at days 1–4 (Fig. 1B). We reasoned that the frequencies of mutants impaired in colonization are reduced along the infection trajectory, resulting in decreased frequencies of the corresponding barcodes from feces samples compared to the inoculum pool. Plating on ampicillin agar verified *hyb*-dependent gut colonization in all mice (competitive index C.I. = 0.1; Fig. 1C,D). Fecal levels of the gut inflammation marker lipocalin-2 verified that initial growth in the absence of overt gut inflammation lasted for 2 days and that pronounced enteropathy commenced by days 3 and 4 post infection (Fig. 1E; Fig. S1D). DNA was prepared from the inoculum pool and from fecal bacteria (see Materials and Methods) to measure fitness phenotypes of mutants and control strains. WITS- and RB-TnSeq controls verified the absence of bottlenecks during days 1 and 2 (Shannon evenness score E_H > 0.95, Fig. S1E; linearity of dilution series R > 0.93; Fig. S1F). Using stringent quality controls (Materials and Methods), we could determine the fitness of 1305 mapped RB-TnSeq mutants during initial growth (days 1 and 2 post infection (pi); Tab. S1). This covered 732 genes and 205 intergenic regions. Linearity was reduced in 3/6 mice by day 3 and in 5/6 mice by day 4 pi (R = 0.3; Fig. S1F). Therefore, we limited our mutant phenotype analysis to days 1 and 2 pi.

By comparing fitness values across the profiled RB-TnSeq mutants along the infection trajectory, we mapped a set of 44 Tn5 mutants impaired or enhanced in colonization at day 1 and/or 2 pi (p<0.05; Fig. 1F, red dots; Tab. S1). Functional classification revealed that the majority of the identified genes comprise cellular functions such as metabolism, virulence, lipopolysaccharide biosynthesis and gene regulation. The identified gene-set included well-characterized pathogenicity factors, including attenuating mutations in LPS O-sidechain biosynthesis (Ilg et al., 2009) and *hypD* (Maier et al., 2013) (Fig. 1G; Tab. S1). The latter encodes a maturation factor required for [NiFE] hydrogenases like Hyb, which is used for H₂/fumarate respiration (Stripp et al., 2013). We also observed the expected growth acceleration of “cheater mutants” including *hilC* and *hilD* defective in the costly expression of the TTSS-1 type III secretion system ((Diard et al., 2013) (Bakkeren et al., 2017); Fig. 1F,G; Tab. S1). Moreover, mutants lacking the two component system DcuR/S, involved in dicarboxylate transport regulation in the closely related bacterium *E. coli* (Uden et al., 2016), were attenuated (Fig. 1F,G; Tab. S1).

To validate the importance of the identified colonization factors, we constructed a panel of 29 barcoded in-frame deletion mutants (Key Resources Table). We included 11 genes that

had yielded significantly enhanced or reduced C.I.s (C.I. < 0.68 or C.I. > 2.06; $p < 0.05$, day 1 pi; Tab. S1) in the RB-TnSeq follow-up experiment. They represented the identified gene classes of interest (Tab. 1). In addition, we included 18 controls (Tab. S2). Competitive infections of LCM mice confirmed the reduced C.I.s for 9/9 mutants with reduced and for 2/2 mutants with enhanced C.I.s (C.I. > 2.06); Tab. 1 and Tab. S2) identified by our screen during initial growth. Among the identified colonization factors, the DcuR/S two-component system attracted our particular attention, as two independent hits were identified in our screen, the in-frame deletion verified attenuation (C.I._(*dcuR*) = 0.34, $p = 0.002$; day 1 pi) and as DcuR/S is linked to fumarate respiration. If C4-dicarboxylates are available under anaerobic conditions, the DcuR/S signaling system induces the expression of genes involved in the uptake and metabolism of C4-dicarboxylates. Specifically, DcuR/S regulates expression of the C4-dicarboxylate transporter DcuB, of FumB, an enzyme converting L-malate into fumarate, and of fumarate reductase (Kleefeld et al., 2009) (Golby et al., 1999) (Unden et al., 2016). Therefore, the identification of DcuR/S in our RB-TnSeq screen suggested that C4-dicarboxylate transport and the conversion of C4-dicarboxylates into fumarate may contribute to initial growth in the gut lumen.

The DcuABC C4-dicarboxylate transporters are required for initial growth in the gut lumen.

S. Typhimurium encodes three Dcu transporters (DcuA, DcuB, DcuC). In *E. coli*, they can functionally complement each other to support anaerobic fumarate-dependent growth on C4-dicarboxylates in vitro (Fig. 2A; (Unden et al., 2016, Zientz et al., 1999, Zientz et al., 1996)). While Dcu transporters have been implicated in *S. Typhimurium* growth in the inflamed gut (Spiga et al., 2017) (Tchawa Yimga et al., 2006, Mercado-Lubo et al., 2009, Rivera-Chavez et al., 2016) (Wilson et al., 2019), their putative role in the initial gut luminal growth of *S. Typhimurium* by H₂/fumarate respiration has remained unclear. Therefore, we decided to decipher their function. As Dcu transporter genes were not represented in our RB-TnSeq library, we created the respective *S. Typhimurium* mutants by site-directed mutagenesis (Key Resources Table) and analyzed gut colonization during initial growth at days 1 and 2 of infection, as well as in the overtly inflamed gut (days 3–4 pi, in case of wild type *S. Typhimurium* infections; Fig. S2).

DcuB is an antiporter, importing external C4-dicarboxylates (fumarate, malate, aspartate, and D-tartrate) in exchange for cellular succinate and is expressed under anaerobic conditions (Engel et al., 1992) (Engel et al., 1994) (Six et al., 1994) (Bin Kim et al., 2007, Kroger et al., 2013) (Fig. 2A). In competitive infections of LCM mice, a *dcuB* mutant (*S.Tm^{dcuB}*; Key Resources Table) had a mild gut colonization defect (C.I. = 0.32 at day 2 pi; Fig. 2B, yellow symbols; Fig. S2A–D; Tab. S3). This mild attenuation is similar to that of *dcuS* and *dcuR* mutants in our RB-TnSeq screen and the verification experiment (Fig. 1F, Tab. 1), but much smaller than the attenuation of *S. Typhimurium* mutants lacking the H₂-hydrogenase *hyb* and the fumarate reductase *frd* (*S.Tm^{frd}*, *S.Tm^{hyb}*; Fig. 2B; blue and grey symbols). Genetic redundancy might mask fitness consequences of individual transporter mutants, as suggested by in vitro studies on *E. coli* (Zientz et al., 1996, Zientz et al., 1999). Therefore, we assessed the colonization phenotype of double and triple *dcu* mutants.

Indeed, while *S.Tm^{dcuAC}* and *S.Tm^{dcuB}* were only mildly attenuated, *S.Tm^{dcuABC}* exhibited ≈ 100 to 1000-fold attenuated initial growth compared to wild type *S. Typhimurium* (C.I. = 0.01 – 0.001; Fig. 2B, red symbols; Tab. S3). This colonization defect could be complemented (Fig. 2C, blue symbols). Deleting *hyb* did not further attenuate *S.Tm^{dcuABC}* at days 1–4 pi (compare to *S.Tm^{dcuABC}* (red symbols) and *S.Tm^{dcuABC hyb}* mutants (green symbols) in Fig. 2B), indicating that *dcu* genes are epistatic to *hyb*, at least during initial growth.

The data presented above indicated that DcuABC transporters are required for the H₂/fumarate-dependent initial growth in the gut lumen of LCM mice. Accordingly, competitive infections with a 1:1 mixture of *S.Tm^{frd}* and *S.Tm^{dcuABC frd}* verified that both strains have similar gut luminal growth defects (C.I. ≈ 1 ; Fig. 2D, blue symbols; Fig. S2E–H). Compared to mice infected with strain mixtures that include wild type *S. Typhimurium* (e.g. *S.Tm* and *S.Tm^{dcuAB}*, Fig. 2D and Fig. S2E–H, red symbols) mice infected with a mixture of *S.Tm^{frd}* and *S.Tm^{dcuABC frd}* showed a delayed rise of gut luminal pathogen densities and delayed gut inflammation kinetics (Fig. S2E,G,H, blue symbols). This prolonged the phase of initial growth and delayed progression towards the next stage of the gut infection process. Thus, DcuABC promote gut colonization and accelerate mucosal inflammation in LCM mice.

The LCM gut microbiota composition shows little change during initial *S. Typhimurium* growth

While overt gut inflammation is known for its dramatic effects on the microbiota, it remained to be established how the microbiota composition is affected during initial growth. Therefore, we infected LCM mice with wild type *S. Typhimurium*. To assess the effects of C4-dicarboxylate-fuelled pathogen growth, we infected a second group with *S.Tm^{dcuABC}*. In line with the data presented above, wild type *S. Typhimurium* featured faster initial growth and elicited gut inflammation about 1 day earlier than *S.Tm^{dcuABC}*, i.e. at day 3 vs. day 4 pi (Fig. 3A,B,C).

At days 1 and 2 pi, we observed only little change in the relative abundance of the different microbiota phylotypes (Fig. 3D,E). This pertained to wild type *S.Tm* and *S.Tm^{dcuABC}* infected mice, as indicated by the relative abundance of the different detectable OTUs and by the low Bray-Curtis dissimilarity scores (Fig. 3D,E). Please note that the Bray-Curtis dissimilarity scores of the feces before the infection (day 0) were defined as the reference (Bray-Curtis dissimilarity score at day 0 = 0). The scores of ≈ 0.2 at days 1 and 2 pi are likely attributable to the normal variability of the microbiota community composition that is also observed in unperturbed LCM mice (see below in Fig. S6B; approximately 20% variation in water-control group). The newly introduced pathogen (which is not present at day 0) may also contribute to some extent. Similarly, low-grade mucosal responses observed at days 1–2 pi might contribute a little (lipocalin 10⁰-10² ng/g; Fig. 3B, Fig. 1E, Fig. S2C, Fig. S3). These low-grade responses are characterized by ≈ 10 -fold induction of *Il6*, *Il1b*, *nos2*, and a mucosal morphology equivalent to uninfected controls (Fig. S3D,E, days 1 and 2 pi). However, these mucosal responses and inflammation-boosted pathogen growth (e.g. via *cheY*-driven chemotaxis, NO₃⁻ and O₂-respiration) are lower than those observed after the onset of overt gut inflammation (i.e. days 3–4 pi with wild type *S.Tm*, Fig. S3B). Thus,

during the first 1–2 days of infection in LCM mice, the initial growth of *S. Typhimurium* proceeds while mucosal responses remain sub-acute and the microbiota shows little change.

The end of the initial growth phase is defined functionally by the onset of overt gut inflammation, as indicated by histopathological analysis and fecal lipocalin-2 concentrations above 3×10^2 ng/g (Fig. 3B,C). In the case of wild type *S. Typhimurium*, this occurs typically by day 3 pi. At the same time, many strains of the microbiota are suppressed, as indicated by the elevated Bray-Curtis dissimilarity scores (Fig. 3D). Suppression is particularly pronounced in the cases of the *Parabacteroides*, *Lachnoclostridium*, *Blautia* and *Flavonifractor* spp. (Fig. 3E). In line with the delayed onset of overt gut inflammation, microbiota disruption occurs later (day 4 pi) and is less pronounced in the case of *S.Tm^{dcuABC}* (Fig. 3D,E). Thus, DcuABC accelerates the infection by promoting initial growth in face of the microbiota. However, the nature of the relevant C4-dicarboxylate substrates of DcuABC remained unclear.

Gut luminal availability of malate, aspartate, citrate and succinate

DcuABC transporters can import several C4-dicarboxylates, namely L-malate, L-aspartate, fumarate or D-tartrate and exhibit affinities in the micromolar range (DcuAB, $K_m(\text{fumarate})$ 100 μM ; (Unden et al., 2016, Six et al., 1994, Janausch et al., 2002, Engel et al., 1994, Bin Kim et al., 2007)). To gain further insights into the relevant substrates fueling the DcuABC-dependent initial growth, we focused on the cecum lumen. This is the main site of *S. Typhimurium* growth in the murine gut. We measured the extracellular metabolites by targeted liquid chromatography/mass spectrometry (Materials and Methods). In line with previous work (Spiga et al., 2017, Cummings et al., 1987), we observed low amounts of fumarate (1.27–3.75 nmol/g cecum content; Fig. 4A) in the cecum lumen of LCM mice and mice harboring a complex pathogen-free microbiota (CON^E). This residual fumarate might originate from bacterial lysis (see lysis control experiment in Fig. S4B) or malate degradation, as germ free mice featured even lower amounts of free fumarate (<0.8 nmol/g cecum content). Overall, we conclude that extracellular fumarate in the cecum lumen is likely insufficient to support anaerobic H₂/fumarate respiration.

In contrast to fumarate, our metabolomics measurements revealed that aspartate (100–500 nmol/g cecum content) and malate (10–20 nmol/g cecum content) were highly abundant in LCM and CON^E mice (Fig. 4A). Assuming that particulate matter and cells occupy 90% of the cecum lumen, the aspartate and malate concentrations in the extracellular space (1–5 mM or 100–200 μM respectively) are close to or above the reported K_M -values of DcuA and DcuB ($K_M = 30\text{--}50$ μM ; (Engel et al., 1994, Six et al., 1994)). A second approach confirmed the abundance of malate and aspartate and established that free citrate is below the detection limit (Fig. S4A). Our assays did not provide data for free L/D-tartrate. The amounts of free succinate in the cecum lumen of LCM and CON^E mice ranged between those of aspartate and malate (≈ 100 nmol/g cecum content; Fig. 4A). These succinate values are 6–30 fold lower than previously reported for gut contents sampled from humans and mice (Spiga et al., 2017, Cummings et al., 1987) (Meijer-Severs and van Santen, 1987). We speculate that these differences are attributable to differences in food, intestinal resorption rates, microbiota composition or bacterial lysis during sample preparation (as in Fig. S4B). Regardless,

because succinate is the reduction end-product of fumarate respiration, it cannot serve as substrate for further anaerobic respiration during initial growth and must be secreted by the DcuABC antiporters (Unden et al., 2016). Collectively, these findings show that aspartate and malate are present in the cecum lumen and that their concentrations suffice to support H₂/fumarate respiration during initial growth in the murine gut.

L-malate and L-aspartate serve as important nutrients for gut luminal H₂/fumarate respiration.

To further deduce the role of malate and aspartate metabolism during the initial growth, we performed a series of competitive infection experiments in LCM mice and profiled *aspA*, *fumA*, *fumB* and *fumC* mutants for colonization defects. The *aspA* gene encodes for aspartate-ammonia lyase, which converts aspartate to fumarate and ammonia (Fig. 4B; (Williams and Lartigue, 1967)). The genome of *S. Typhimurium* also encodes three fumarases (encoded by *fumA*, *fumB*, or *fumC*), which are hydro-lyases catalyzing inter-conversion of L-malate and fumarate (Mercado-Lubo et al., 2009). In competitive infection assays, *S.Tm^{fumAC fumB}* and *S.Tm^{aspA}* both exhibited impaired initial growth as compared to wild type *S. Typhimurium* (C.I. = 0.119 or 0.052, respectively; day 2 pi; Fig. 4B; Tab. S3; for day 1 pi data see Fig. S4C) suggesting that catabolism of malate and aspartate is important for fueling gut luminal H₂/fumarate respiration. Accordingly, the quadruple mutant *S.Tm^{fumAC fumB aspA}*, which can neither convert L-malate nor L-aspartate, showed severe attenuation (C.I. = 0.006 at day 2 pi; Fig. 4B; Tab. S3) to a similar level as observed for *S.Tm^{frd}*, *S.Tm^{hyb}*, or *S.Tm^{hyb frd}* (C.I. = 0.010, 0.013, or 0.002 respectively; day 2 pi; Fig. 4B; Tab. S3). This genetic evidence suggests that malate and aspartate are the main sources to drive H₂/fumarate respiration during the initial growth of *S. Typhimurium*. By inference, we conclude that the import of fumarate from the gut lumen does not contribute significantly as initial growth in the absence of *aspA*, *fumA*, *fumB* and *fumC* was severely impaired. This is in line with our metabolomics measurements establishing low availability of free fumarate in the cecum lumen (Fig. 4A).

While anaerobic H₂/fumarate respiration is well established for *E. coli* growth in vitro (Unden et al., 2016, Zientz et al., 1999, Zientz et al., 1996), it remained to be verified for *S. Typhimurium*. We performed anaerobic (4% H₂, 10% CO₂, 86% N₂) *S. Typhimurium* growth experiments to verify the conversion of L-malate and aspartate to succinate. In the presence of C4-dicarboxylates, wild type *S. Typhimurium* grew anaerobically in minimal medium containing pyruvate, while much slower growth was observed in the absence of C4-dicarboxylates (Fig. S4D). Supplementing the minimal medium with ¹³C C4-dicarboxylates resulted in the consumption of all the available C4-dicarboxylate and the release of 50–70% mole equivalents of ¹³C succinate, as measured by mass spectrometry of the spent medium (Fig. 4C). Thus, similar to *E. coli* (Unden et al., 2016, Zientz et al., 1999, Zientz et al., 1996), *S. Typhimurium* can grow by anaerobic H₂/fumarate respiration, if C4-dicarboxylates are available.

Besides DcuABC-mediated uptake, *S. Typhimurium* encodes several alternative transporters and enzymes which may replenish cellular malate and aspartate pools under anaerobic conditions (Fig. 4B). However, these do not appear to affect initial growth in the gut, as

shown by a series of competitive infection experiments, which are summarized in Fig. 4B, Fig. S4C and Tab. S3 and discussed in the supplement (Supplemental Discussion).

In addition to transport of aspartate by DcuABC, the genome of *S. Typhimurium* encodes a H⁺/glutamate-aspartate transporter GltP (Deguchi et al., 1989). However, *S.Tm^{gltP}* colonized the gut about as efficiently as wild type *S. Typhimurium* (C.I. = 1.2 at day 2 pi, Fig. 4B) and disrupting *gltP* did not further attenuate the *dcuA dcuB dcuC* mutant (compare *S.Tm^{dcuABC}* vs. *S.Tm^{dcuABC gltP}*) suggesting that the DcuABC is the main import route for aspartate. Collectively, these findings suggest that Dcu transporters promote initial growth in the gut lumen by importing mainly malate and aspartate to drive H₂/fumarate respiration.

A fully operating TCA cycle is not required for initial gut lumen colonization.

Next, we asked whether the observed phenotypes (Fig. 4B, Fig. S4C, Tab. S3) are attributed solely to defects in fumarate respiration or, alternatively, were due to other disturbances of the TCA cycle. The latter was conceivable, as several recent studies have reported full expression of the TCA cycle enzymes in antibiotic pretreated or inflamed guts (Ng et al., 2013) (Ali et al., 2014) (Wilson et al., 2019, Rivera-Chavez et al., 2016, Spiga et al., 2017). Some of the TCA enzymes and substrates are shared with H₂/fumarate respiration. However, deletions in genes exclusively involved in the TCA cycle but not required for H₂/fumarate respiration exhibited only minor effects on initial growth compared to *frd*, *hyb* or *dcuABC* mutants. Among them was an *sdh* mutant lacking succinate dehydrogenase activity, which colonized LCM mice almost as well as wild type *S. Typhimurium* (*S.Tm^{sdh}* C.I. = 0.66, day 2 pi; Fig. 4B, Tab. S3). Similarly, deletion of *sucCD* lacking succinyl-CoA synthetase activity only mildly affected initial growth (*S.Tm^{sucCD}* C.I. = 0.68, day 2 pi; Fig. 4B, Tab. S3). This was in line with earlier work analyzing the gut-luminal growth of a *S. Typhimurium* TTSS-1/2 mutant incapable of eliciting mucosal inflammation (Spiga et al., 2017). In sum, our findings suggested that a functioning TCA cycle is not a prerequisite for initial growth in the murine gut.

DcuABC promotes *S. Typhimurium* growth in the face of a complex microbiota

To assess the role of *dcuABC*-mediated C4-dicarboxylate transport for initial growth in face of a complex microbiota, we performed competitive infection experiments with barcoded strains in CON^E mice (Fig. 4D–G; Star Methods; no pretreatment). In line with the pronounced colonization resistance conferred by complex microbiotas (Barthel et al., 2003, Stecher et al., 2010, Wotzka et al., 2019), we detected wild type and/or *Salmonella* cells in feces of only 16/30 animals by day 1 pi (Fig. 4D). The pathogen loads declined further by day 2 pi, and we observed no signs of overt gut inflammation (Fig. 4F,G). The low colonization efficiency in CON^E mice explains why we only detected robust signals for the barcodes of all 7 strains in the 6 mice carrying the highest fecal pathogen loads at day 1 pi (Fig. 4D,E; highlighted with colors; >10⁵ cfu/g). This data is noisier than that from LCM mice. Nevertheless, in these 6 mice *S.Tm^{frd}*, *S.Tm^{hyb}*, *S.Tm^{dcuABC frd}*, *S.Tm^{dcuABC}*, *S.Tm^{dcuABC hyb}* and *S.Tm^{fumAC fumB aspA}* were significantly attenuated compared to wild type *S. Typhimurium* (median attenuation: 10–100-fold). Subsequent experiments further substantiated this (see below; unperturbed CON^E controls in Fig. S5C). In sum, these

findings suggest that *dcuABC*-mediated C4-dicarboxylate transport promotes initial gut luminal growth in mice harboring different microbiotas.

Aspartate and Malate supplementation promotes gut luminal growth of *S. Typhimurium* in face of a complex microbiota.

Our data suggested that gut luminal levels of malate and aspartate are critical determinants promoting gut colonization. To assess whether colonization resistance is affected by malate or aspartate availability, we supplemented CON^E mice with water (= negative control) or L-aspartate and L-malate (= experimental group) and infected the animals for 4 days with an equal mix of WITS-barcoded wild type *S. Tm*, *S. Tm^{dcuABC}*, *S. Tm^{fumABC aspA}*, *S. Tm^{hyb}*, *S. Tm^{aspA}*, *S. Tm^{fumABC}*, and *S. Tm^{frd}*. We found that the total fecal *S. Typhimurium* load was 10–100 fold higher and more animals developed mucosal inflammation when aspartate and malate was supplemented (3/9 mice) as compared to controls (0/10 mice; Fig. 5A,B; Fig. S5A). Systemic colonization might also be slightly elevated in the L-aspartate/L-malate treated group. However, this effect was not significant (p > 0.05; Fig. S5B). These data indicated that aspartate and L-malate promote initial colonization and progression to disease.

In line with the data presented above (Fig. 4E, Fig. 2B, Tab. S3), *S. Tm^{dcuABC}*, *S. Tm^{fumABC aspA}*, *S. Tm^{hyb}*, *S. Tm^{aspA}*, and *S. Tm^{frd}* were attenuated in the control CON^E mice upon aspartate and malate supplementation (Fig. S5C). However, as total pathogen densities were low, these data were again quite noisy. This may also explain why we could not detect significant growth defects for some of the strains like *S. Tm^{fumABC}* in this experiment.

We also assessed if malate- and aspartate-inflicted changes in microbiota composition might contribute to pathogen growth. Malate and aspartate supplementation shifted the microbiota composition (Fig. S6A,B). However, this shift was mild compared to those inflicted by the onset of gut inflammation (LCM mice, Fig. 3D) or by high fat diet exposure of CON^E mice (Fig. S6A,B, right side), a stimulus which can boost *S. Typhimurium* gut colonization by 1000-fold (Wotzka et al., 2019). Fumarate supplementation had an even milder effect on the microbiota composition than malate and aspartate supplementation (Fig. S6A,B). We speculate that this may be attributable to fumarate resorption by the host, as food contains much more fumarate than the cecum lumen of germ-free mice (Fig. 4A). Importantly, the fumarate supplementation did not enhance initial pathogen growth in the gut lumen or enteropathy (Fig. S6C,D). Thus, the observed small changes in microbiota composition alone appear insufficient for boosting initial growth of *S. Typhimurium*. However, we cannot exclude that they contribute in malate and aspartate supplemented mice.

Pathogen blooms in the aspartate and malate supplemented CON^E mice still required *dcuABC*, *aspA*, *hyb* and *frd* (Fig. S5C). This supported the idea that pathogen growth still relied on H₂/fumarate respiration. We cannot exclude that additional effects attributable to altered microbiota metabolism or community structure might further promote the pathogen's growth. Collectively, our findings underscore the importance of C4-dicarboxylic acid levels in the gut lumen to promote initial growth of *S. Typhimurium* in CON^E mice.

***E. coli* 8178 requires DcuAB for competitive gut luminal growth**

Genome analyses indicated that DcuABC homologs and genes implicated in H₂/fumarate respiration are encoded by several other enteric pathogens and by commensal *E. coli* strains (Tab. S4), including *E. coli* 8178. *E. coli* 8178 had been isolated from mice and is capable of limiting gut luminal *S. Typhimurium* growth after a shift to Western-type diet and in the inflamed gut (Stecher et al., 2012) (Wotzka et al., 2019). We hypothesized that Dcu-mediated C4-dicarboxylate import might contribute to the competitive capacity of *E. coli* 8178. Based on our *S. Typhimurium* data, we focused on DcuA and DcuB, as these should show the strongest phenotypes. To test our hypothesis, we performed a competitive infection experiment in mice harboring a complex microbiota that lacks commensal *E. coli*. We used CON^X mice (Star Methods), as these animals allow sufficient *E. coli* and *S. Typhimurium* densities for our experiment (approx. 10⁵-10⁷ cfu/g stool at day 1 p.i.). The animals were inoculated with a mix of wild type *S. Typhimurium*, wild type *E. coli* 8178 and an isogenic mutant lacking DcuAB (*E. coli*^{Δ*dcuAB*}; Key Resources Table). Control mice were inoculated with wild type *S. Typhimurium* alone (Fig. 6). Lipocalin-2 levels of CON^X mice were slightly higher than those of uninfected CON^E mice (Fig. 6D; compare to Fig. 3B, Fig. S1D). However, tissue sections revealed no enteropathy and Lipocalin-2 levels did not change over time (Fig. 6D,E), suggesting that initial growth lasted throughout the course of the 4 days of infection. In presence of *E. coli*, *S. Typhimurium* had a reduced initial growth in the gut lumen (Fig. 6A). We also observed a trend towards reduced spread to systemic sites (Fig. 6C). However, this trend was not significant and more animals would be necessary to verify this effect. *E. coli*^{Δ*dcuAB*} was 10–100-fold attenuated in colonization compared to wild type *E. coli* (Fig. 6A,B). Wild type *E. coli* diminished initial *S. Typhimurium* growth in CON^X mice more efficiently than *E. coli*^{Δ*dcuAB*} (Fig. 6F) and *E. coli*^{Δ*frd*} was attenuated in competitive infections (Fig. 6G). These data confirm that commensal *E. coli* can fortify colonization resistance against *S. Typhimurium* (Velazquez et al., 2019) and suggest that DcuAB-dependent fumarate respiration promotes its' competitive capacity in the gut lumen.

Discussion

The initial growth of *S. Typhimurium* in the anaerobic milieu of the microbiota-colonized gut is fueled in part by H₂/fumarate respiration where fumarate serves as an alternative electron acceptor for electrons derived from the *hyb* H₂-hydrogenase reaction to sustain anaerobic growth in the gut lumen (Maier et al., 2013). Here, we found that fumarate is obtained via conversion of L-aspartate and/or L-malate, which are imported in exchange for succinate, the reduction product of fumarate. This DcuABC-dependent antiporter process allows *S. Typhimurium* to maintain respiration in the absence of suitable alternative electron acceptors. This metabolic strategy employs just one or two steps of the reverse TCA cycle, i.e. the fumarate reductase (*frd*) and fumarase (*fumABC*).

What is the source of the L-aspartate and the L-malate? Aspartate is a typical food constituent. Our mouse diet contains about 1–2 μmol/g (≈0.01% w/w; Fig. 4A) free aspartate, and much additional aspartate is bound in proteins (18.5% w/w protein content). Most proteins are hydrolyzed, but most aspartate is resorbed before the digesta leave the

small intestine. In pigs, aspartate resorption occurs with apparent transporter affinities of $K_m=5\text{--}20$ mM (Buddington et al., 2001). If this holds true for mice, much of the free aspartate detected in the cecum lumen (100–500 nmol/g cecum content; corresponding to $\approx 1\text{--}5$ mM, as discussed above) may originate directly from the food. This free aspartate concentration is sufficient for DcuABC-mediated C4-dicarboxylate uptake by *S. Typhimurium*, as the pathogen's transporters feature 20–100 fold higher substrate affinity than the host's transporters (DcuAB, $K_m(\text{fumarate})$ 0.1 mM; (Six et al., 1994, Jausch et al., 2002, Engel et al., 1994)). As germ-free mice, LCM mice and CON^E mice have similar amounts of free aspartate in the cecum lumen (0.1–0.4 mM), the microbiota may not contribute much. Thus, most of the aspartate fueling *S. Typhimurium* growth in the cecum lumen may originate from food.

Malate is present in mouse food at 8 $\mu\text{mol/g}$ ($\approx 0.1\%$ w/w). Rat feeding experiments with ¹⁴C-labeled D/L-malic acid suggest that $\approx 0.3\%$ of this malate will be excreted with the feces (Daniel, 1969). Based on these data, we would expect about 24 nmol/g free malate in the cecum lumen. While germ-free mice had only 2 nmol/g, we observed 10–20 nmol/g free malate in the cecum content of LCM and CON^E mice. Therefore, this malate may derive from both, food and the microbiota. Our data indicates that the malate and the aspartate concentrations are in the same range as the C4-dicarboxylate K_m values of the DcuABC transporters. This supports our hypothesis, that Dcu-mediated C4-dicarboxylate uptake fuels *S. Typhimurium*'s initial growth by H₂/fumarate respiration. Interestingly, the substrates fueling this initial growth are co-fed from two distinct sources, the food (L-aspartate, some L-malate) and the microbiota (H₂ (Maier et al., 2013), some L-malate).

In addition to the use of C4-dicarboxylates for H₂/fumarate respiration, *E. coli* and *S. Typhimurium* can also use C4-dicarboxylates during aerobic growth (Unden et al., 2016). In this case, C4-dicarboxylates are mainly imported via DctA and serve as electron donors, which are oxidized to CO₂. This occurs during a later phase of *S. Typhimurium* gut colonization, i.e. when gut inflammation commences and creates an oxidative gut-luminal milieu (Spiga et al., 2017). In most LCM animals, this occurs at days 3–4 pi when mucosal responses are induced by 10–100 fold (e.g. *I16*, *I11b*, *nos2*; Fig. S3D,E). The central carbon metabolism of *S. Typhimurium* adapts accordingly by switching away from H₂/fumarate respiration to utilizing inflammation-elicited terminal electron acceptors (O₂, NO⁻₃, tetrathionate; (Winter et al., 2010) (Rivera-Chavez et al., 2016) (Spiga et al., 2017); see also Fig. S3) and diverse electron donors, including microbiota-released carbohydrates, lactate released by enterocytes (Gillis et al., 2018), 1,2-propanediol (Faber et al., 2017), galactarate and glucarate (Faber et al., 2016), formate (Hughes et al., 2017), ethanolamine (Thiennimitr et al., 2011) and succinate (Spiga et al., 2017). As a consequence, the full TCA cycle is engaged in the *forward* direction. This is confirmed by our data for days 3 and 4 pi in LCM mice (C.I. of *S.Tm*^{*sdh*} = 0.28; C.I. of *S.Tm*^{*sucCD*} = 0.11; Tab. S3). In this phase of the infection, C4-dicarboxylate importers like DctA support *S. Typhimurium* blooms during gut inflammation by importing microbiota-derived succinate (Spiga et al., 2017). Thus, *S. Typhimurium* utilizes C4-dicarboxylates in distinct ways during initial growth vs. inflammation-promoted pathogen blooms.

Gut luminal growth of commensal *E. coli* 8178 and its competitive growth vs. *S. Typhimurium* also relies on DcuAB and fumarate reductase. This is in line with our *S. Typhimurium* data and earlier work on the growth of *E. coli ftd* mutants in mice treated with low doses of streptomycin (Jones et al., 2011). Sequence data hints that other Enterobacteriaceae might also employ aspartate or malate to gain energy from H₂/fumarate respiration. For example, orthologs of *dcuABC*, *fumABC*, *aspA*, *frd* and diverse H₂-hydrogenases are observed in additional *Salmonella enterica* spp., commensal and pathogenic *E. coli* strains, *Helicobacter pylori*, *Shigella flexneri* and *Yersinia enterocolitica*, but not in other commensal microbiota strains analyzed (Tab. S4). Along these lines, it is also tempting to speculate that some microbiota members may inhibit infections by depleting, or fuel infections by releasing L-aspartate, L-malate or other C4-dicarboxylates, as suggested in the Restaurant hypothesis (Conway and Cohen, 2015) (Maltby et al., 2013). It would be interesting for future work to test these hypotheses.

Star Methods

LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Wolf-Dietrich Hardt (hardt@micro.biol.ethz.ch). Mice can be obtained from Jackson laboratories. Gnotobiotic mice are available upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mouse lines.—We used male and female 8–12 week old mice and assigned animals of either sex randomly to the experimental groups. All lines lack *E. coli*, as determined by plating on MacConkey agar. All mice originate from C57BL/6 or 129SvEv breeders originally obtained from Jackson laboratories. The mice with a normal complex microbiota (CON^E, CON^X) were specific pathogen-free and bred under full barrier conditions in individually ventilated cage systems in the EPIC mouse facility of ETH Zurich, Switzerland. CON^E mice have a C57BL/6 genetic background, CON^X mice have the 129SvEv genetic background. **LCM mice** (Stecher et al., 2010) (Maier et al., 2013) are ex-germ free C57BL/6 animals that have been associated with the strains of the altered Schädler flora. They are bred in flexible film isolators under strict exclusion of microbial contamination at the isolator facility of the EPIC mouse facility of ETH Zurich, Switzerland. **Germ-free** C57BL/6 mice were bred in flexible film isolators under strict exclusion of microbial contamination at the isolator facility of the EPIC mouse facility of ETH Zurich, Switzerland.

All studies were performed in accordance with ethical and legal requirements and were reviewed and approved by the Kantonales Veterinäramt Zürich under the licenses 222/2013, 193/2016 and ZH158/19.

Bacteria strains and culture conditions—All *Salmonella* strains are isogenic to *S. Tm* SB300, a re-isolate of SL1344. The *E. coli* strains are derivatives of *E. coli* 8178, an isolate from the stool of a SPF C57BL/6 mouse colony. They are listed in the KEY RESOURCES TABLE. All strains were routinely grown overnight at 37°C in Lysogeny broth (LB) with agitation. Strains were stored at – 80°C in peptone glycerol broth.

METHOD DETAILS

Construction of bacteria strains and plasmids

Transposon Mutagenesis.: We constructed the DNA-barcoded EZ:Tn5 transposome by PCR from a linearized plasmid pZ922 template using primers that contained a random 17 nucleotide sequence flanking the RB-TnSeq PCR priming sites and the terminal inverted repeats of Tn5 (Fig. S1A; KEY RESOURCES TABLE). Specifically, we used primer tn5+ME+17N-F3 and primer tn5+ME-R2 to amplify a kanamycin resistance cassette from pZ922, which has an Illumina primer binding sequence at the 5' end and a transcriptional terminator and a T7 promoter downstream. We performed a PCR in 100 µl total volume with 0.1 µmol of each primer, 25 ng of pZ922 template (NotI digested), and 0.5 µl Phusion DNA polymerase (NEB 1.0U/µl) with the following cycling program: initial denaturation step at 94 °C for 1 min followed by 30 cycles of 94 °C for 30 sec, 55 °C for 30 sec, 72 °C for 1 min, and a terminal extension step at 72 °C for 5 min. The PCR product was purified by gel electrophoresis in the absence of DNA intercalating dyes, excised from the gel for column-based DNA extraction (Macherey&Nagel), and eluted in 10 mM Tris/HCl, 1 mM EDTA (TE buffer). We prepared the transposome by adding 1 µl of EZ-Tn5 Transposon DNA (100µg/ml in TE) (Epicentre; 1.0 U/µl), and 1 µl of glycerol, mixed by vortexing, and incubated for 30 min at room temperature. Excess salts were removed by drop dialysis on a Type-VS Millipore membrane (0.025 µm pore size, Millipore). We electroporated 1 µl of the DNA-barcoded EZ:Tn5 transposome into 50 µl of electrocompetent wild type *S. Typhimurium* (prepared from mid log cultures; ~10¹⁰-10¹¹ cfu/ml). Cells were immediately recovered in SOC medium, incubated for 45 min with agitation at 37 °C, and plated in batches on LB agar (50 µg/ml kanamycin). The transposon mutants were recovered and aliquots of a pool of 2000 colonies were stored in peptone glycerol at -80 °C.

RB-TnSeq Control Strains.: We constructed the RB-TnSeq control strains using the lambda red recombination system (Datsenko and Wanner, 2000) in order to insert the barcoded transposon into a fitness neutral locus. 40-nucleotide long sequences that are homologous to the intergenic region between the pseudogenes, *malX* and *malY* were attached to the ends of the barcoded transposons (as generated from above) by PCR using the primers, ydgA tn5-aphT-R2 and ydgA tn5-aphT-F2. PCR products were electroporated into a wild type *S. Typhimurium* carrying the pKD46 plasmid. Recombinant strains were selected on LB agar (50 µg/ml kanamycin). The transposon insertion was confirmed by PCR (primers ydgAtn5aphTcheckR and ydgAtn5aphTcheckF) and the associated 17 nucleotide sequence was determined by Sanger sequencing. To generate RB-TnSeq barcoded control strains with known DNA barcodes, we transduced from these strains five barcoded transposons each via P22 phage into the mutant strains, *hybABChypO*, *cheY*, and *ssaV invG* as well as wild type *S. Typhimurium* (KEY RESOURCES TABLE).

Mouse infection experiments.—8 to 12 week old mice were inoculated with *S. Typhimurium* cells prepared from a culture grown for 12 h in LB medium containing the appropriate antibiotic(s), diluted 1:20 and sub-cultured for 4 h in the same medium w/o antibiotics. Bacteria were washed twice with PBS and mice were infected with bacteria by gavage (5·10⁶ cfu in 50 µl, if not stated otherwise). Mice were kept in grid-floor cages to avoid transmission between mice. Animals were sacrificed at day 4 pi by cervical

dislocation. Fresh fecal pellets, whole cecum content, a piece of the liver, the mesenteric lymph nodes and the spleens were harvested and suspended in PBS (500 μ l) using a Tissue Lyser device (Qiagen) before plating to determine the total bacterial population size. HE-staining of cyro-embedded tissues and subsequent pathoscore for granulocyte infiltration was performed as described previously (Barthel et al., 2003).

Lipocalin-2 ELISA.—Lipocalin 2 was detected in feces homogenized in 500 μ l sterile PBS by ELISA (DuoSet Lipocalin ELISA kit (DY1857, R&D Systems, Minneapolis, MN, USA).

Sequencing of transposon insertion sites and their associated barcodes.

Sequencing library preparation.: Transposon sequencing and DNA library preparation was performed as previously described (Yung et al., 2015, Christen et al., 2011). In brief, we used semi-arbitrarily primed (semi-AP) and nested PCR reactions to amplify the transposon junctions bearing the unique DNA barcodes as well as the adjacent genomic DNA and to add terminal adapter sequences compatible to the Illumina sequencing platform (Fig. S1B). To prepare the PCR template, we diluted the transposon library in sterile water to an OD of 1.0 and subjected the diluted sample to six freeze-thaw-heat cycles (submersion in liquid nitrogen for 1 min, followed by incubation at room temperature for 1 min and at 95 °C for 1 min). We then performed separate semi-AP PCRs with the crude bacterial lysate as template and the transposon specific primer (ARB-Tn5-R1) and one of the indexed semi-arbitrary primers (ARB3A_PE, ARB3C_PE, ARB3D_PE, ARB3F_PE; KEY RESOURCES TABLE), using the following program: (1) 94 °C for 3 min, (2) 94 °C for 30 s, (3) 42 °C for 30 s, slope -1 °C per cycle, (4) 72 °C for 1 min, (5) go 6 times to step 2; (6) 94 °C for 30 s, (7) 58 °C for 30 s, (8) 72 °C for 1 min, (9) go 25 times to step 6; (10) 72 °C for 3 min. We enriched and prepared the PCR products for Illumina sequencing by performing a second PCR with nested primers that carry terminal Illumina sequencing adapters (PE_PCR_Primer_1.0, PE_PCR_Primer_2.0; KEY RESOURCES TABLE) and the following program: (1) 94 °C for 3 min, (2) 94 °C for 30 s, (3) 64 °C for 30 s, (4) 72 °C for 1 min, (5) go 30 times to step 2; (6) 72 °C for 3 min. We size-selected 200–500 bp fragments by gel electrophoresis (2% agarose) prior to Illumina sequencing.

Illumina sequencing.: Cluster-generation was performed within standard Illumina paired-end flow-cells (Illumina cluster chemistry v2) and settings according to the Illumina cluster generation protocol. The DNA sample concentration (25 ng/ μ l) of each transposon junction library was titrated to produce approximately 1.3×10^5 clusters per tile on the flow cell. Paired-end sequencing with 150 bp read length from both ends of a given DNA fragment was performed on an Illumina HiSeq instrument using sequencing chemistry (v4) and standard paired-end sequencing primers (PE 1.0 and PE 2.0, Illumina).

Mapping of the transposon insertion sites.: Raw transposon read processing and mapping of genomic insertion sites was performed using previously described data analysis routines (Christen et al., 2016, Christen et al., 2011). In brief, reads with at least a 15 bp long perfect match to the Tn5 transposon end sequence [‘GTGTATAAGAGACAG’] and bearing genomic inserts larger than 15 base pairs were selected. Transposon internal RB-TnSeq-

barcode sequences were extracted and sequencing strings corresponding to the illumina adapters, transposon end or arbitrary PCR primers sequences were trimmed. Using bwa-07.12 (Li and Durbin, 2010), filtered transposon reads were aligned onto the *S. Typhimurium* NCBI reference genome (FQ312003). Reads with correct paired-end alignment, no mismatches within the first 15 bases of both paired-end reads, and with an insert size smaller than 700 bases, were selected for subsequent analysis. Transposon insertion sites were defined according to the first reference base detected adjacent to the transposon end sequence.

RB-TnSeq barcode assignment to genomic insertion sites.: Barcode sequences were unambiguously assigned to genomic insertion locations selecting the most abundant barcode sequence associated with a specific insertion site and excluding secondary barcode occurrences at alternative insertion sites with lower read counts. (Fig. S1B; Fig. S4). Cumulatively, we could unambiguously associate 1305 unique RB-TnSeq-barcodes with specific transposon insertion sites, corresponding to 732 genes.

Barcode annotation.: Each barcode was assigned to specific gene if the insertion locations occurred inside the protein coding sequence according to the reference genome coordinates (Tab. S3).

Analysis of RB-TnSeq mutant gut colonization phenotypes.

Sample preparation.: Fecal *S. Typhimurium* cells were enriched in 10 ml LB supplemented with 50 µg/ml of streptomycin or 3.0 ml LB supplemented with 12.5 µg/ml of tetracycline for 4 hours to select and enrich for living salmonellae. Bacterial cells were pelleted, resuspended in TE buffer, and stored at -20 °C. DNA was extracted from thawed pellets using commercial kits (Qiagen Mini DNA) according to manufacturer's instructions. To PCR-amplify the barcode sequences, we added 200 ng of extracted DNA, 100 nmol of each primer (PE PCR Primer 1.0 and indexed PE2.0-ME, Table S1), 10 µmol dNTPs and standard amounts of reaction buffer and Taq (0.2 µl, 1 U/µL) in a 50 µl total volume. We ran the reaction using the following cycling program: initial denaturation step at 94 °C for 3.0 min followed by 23 cycles of 94 °C for 30 sec, 60 °C for 30 sec, 72 °C for 20 sec, and a terminal extension step at 72 °C for 2.0 min. PCR products were column-purified, measured for DNA concentration, pooled (450 ng per sample), and then gel purified. Purified PCR products were diluted to a final concentration 25 ng/µl in TE buffer.

Illumina sequencing.: Sequencing was performed as described, above (see sequencing of RB-TnSeq transposon insertion sites).

Barcode counting.: In order to analyze barcode sequence abundance in the inoculum and the output pools, we extracted random barcode sequences and index sequences used to track individual sample and experimental conditions from raw read files. Barcodes were demultiplexed and corresponding occurrences were computed using shell scripts (Fig. S3B). We only considered 100% matches.

Sample QC.: For each sample, a linear regression is made with the dilution series of wild type control strains to verify that $r^2 > 0.8$. If this is not the case, two more successive linear regressions are made using only 4 or 3 control wild type RB-TnSeq strains with the higher titers. If in neither of these cases the sample maintains linearity $r^2 > 0.8$, the sample is discarded completely. In order to avoid noise originating from RB-TnSeq mutants that were poorly represented in the inoculum pool, we only retained RB-TnSeq mutants with equal or higher number of reads than the lowest wild type RB-TnSeq strain that maintains linearity.

Rarefaction.: All of the samples were rarefied to the same value, equivalent to the shallowest sample that passed QC. This was done with the vegan R package (v 2.5-1)

Set detection limit.: Linear regressions were calculated for all the samples in a similar procedure to the Sample QC. All barcodes less abundant than the lowest linear wt control are deemed below detection limit and set to the corresponding value

Calculate fitness.: Fitness values were calculated as the number of observed barcode reads at a particular time point divided by the number of barcode reads observed in the inoculum. In the cases where there is more than one barcode per gene, the median value was selected. To test for significance, a Wilcox unpaired exact test was performed by comparing each RB-TnSeq gene fitness value and the corresponding mean fitness for the fitness values of the 3/2000, the 1/2000 and the 1/6000 dilutions of the RB-TnSeq-barcoded *ssaV invG* control strains, and then FDR adjusted.

Calculate CI.: The CI (Competitive index) is calculated by dividing the fitness values of each RB-TnSeq mutant to the mean fitness value of the 3/2000, the 1/2000 and the 1/6000 dilutions of the RB-TnSeq-barcoded wild type *S. Typhimurium* control strains. To test for significance, a Wilcox unpaired exact test is performed for each gene CI and the corresponding mean CI for the 3/2000, the 1/2000 and the 1/6000 dilutions of the RB-TnSeq-barcoded *ssaV invG* control strains, and then FDR adjusted. The decision to use the RB-TnSeq-barcoded *ssaV invG* strains for comparison (and not the RB-TnSeq-barcoded wild type strains) is that the RB-TnSeq-barcoded wild type strains had already been used for calculating the CI. During co-infections with wild type *S. Typhimurium*, the *ssaV invG* strains are known to colonize the gut lumen just as well as the isogenic wild type strain (Stecher et al., 2007). Therefore, we could use the RB-TnSeq-barcoded *ssaV invG* strains for comparison (Fig. S3A)

Analysis of WITS mutant gut colonization phenotypes.—*Inoculum mixtures* were prepared by mixing equal ratios of the indicated WITS-barcoded strains including a WITS21-barcoded wild type *S. Typhimurium* strain (KEY RESOURCES TABLE). An equivalent proportion (50% of the total cfu) of non-barcoded wild type *S. Typhimurium* was added to ensure equivalent disease kinetics between different experiments. If two independent clones were available from a given mutant (KEY RESOURCES TABLE), we used the different clones in the different independent replicates of each experiment and verified that both clones yielded equivalent results. This strategy avoided misinterpretation of our data that might be caused by non-neutral second-site mutations.

Sample preparation for WITS-distribution analysis.: Samples were prepared and DNA was recovered, as described above (Analysis of RB-TnSeq mutant phenotypes). rtqPCR analysis was performed using the FastStart Universal SYBR Green Master Mix (Roche) and primers listed in KEY RESOURCES TABLE. The temperature profiles were described previously (Maier et al., 2014b) (Grant et al., 2008) (Kaiser et al., 2013). The population size of each tagged strain was calculated by multiplying the number of kanamycin resistant colonies/g feces with the ratio of WITS determined by rtqPCR.

To calculate Shannon Evenness (E_H), we used the formula: $E_H = \frac{H}{\ln S}$, where $H = -\sum_{p_i} p_i \ln p_i$, \ln , with H =Shannon's diversity index, S =total number of strains, p_i =proportion of S made up of the i th strain.

16S rRNA gene sequencing—Microbial DNA was extracted from feces as described (Wotzka et al., 2019). Feces were collected before the infection (day 0) or at days 1, 2, 3, or 4 as indicated. The samples were immediately frozen in liquid nitrogen for storage at -80°C . The AllPrep DNA/RNA Kit (Qiagen) was used to extract DNA, using 600 μl of RLT buffer. One unit of 3 mm metal beads (Berani Kugellager AG) and 600 mg of 0.1 mm zirconia beads (OPS Diagnostics) were added to the tube containing the feces, which were bead-beaten two times at 30 Hz for 5 min with a 5 minute incubation time in between, using a Retsch M400 apparatus. The lysates were transferred into the corresponding columns which were then washed as described in the manufacturer's protocol. Finally, the bacterial DNA was eluted in 100 μl elution buffer (EB).

The NEXTflex® 16S V4 Amplicon-Seq Kit 2.0 (Barcodes 1–96; Bio Scientific) was employed to produce the 16S libraries. Genomic DNA (30 ng/ μl) was used as template and the library preparation was performed according to the manufacturer's protocol (25 μl reaction volumes). A modified primer pair was used for the first PCR reaction, replacing the primer pair 515f-806r from the kit with the degenerate primers 515F (5'-GTGYCAGCMGCCGCGGTAA-3') and 806rB (5'-GGACTACNVGGGTWTCTAAT-3') as described^{52,53}. Q5 High-Fidelity DNA Polymerase (BioConcept (NEB)) was used for the first PCR reaction: 1) first denaturation: 95°C for 4 min; 2) denaturation in each PCR cycle: 95°C for 30 sec; 3) annealing (each PCR cycle): 56°C for 30 sec; 4) extension (each PCR cycle): 72°C for 90 sec; 5) final extension at the end of the reaction: 72°C for 4 min. The cycles 2–4 were repeated 8 times. The PCR products were purified using AMPure XP magnetic beads (Beckman Coulter SA), eluted in 20 μl of resuspension buffer and served as templates in the next PCR reaction, which employed barcoded primers for multiplexing. The quantity of the amplicons was measured using a Qubit fluorometer and quality was verified using a Fragment Analyzer (Advanced Analytical). DNA concentrations were measured using a Qubit fluorometer. PCR products (approx. 450 bp length) were diluted to a final concentration of 60 ng DNA per 20 μl multiplexed sample. Illumina MiSeq paired end read sequencing was performed at the Functional Genomics Center Zürich.

The raw sequencing data were processed with custom scripts to execute the following commands of the USEARCH software (v9.1.13): the fastq_mergepairs command was used with default settings to merge and quality-filter paired reads. The merged reads were

subsequently filtered using the `fastq_filter` command (`-fastq_maxee 0.1`). We only merged reads with perfect primer matches and we only selected reads of 100 bp length. The `fastx_uniques` command was used for de-replication. Then, data were clustered into operational taxonomic units (OTUs; 97% threshold) using the `cluster_otus` command (`-minsize 2`). Thereby, chimeric sequences were also removed. The `usearch_global` command (`-strand both; -id 0.97`) was used to quantify the OTU abundances for each sample. Taxonomic annotation was done by querying OTU sequences against the SILVA database (v 128). For this purpose, the `usearch_global` command (`-id 0.90; -maxaccepts 20; -maxrejects 500; -strand both; -top_hits_only`) was performed. The scripts used for these steps are available in https://github.com/SushiLab/Amplicon_Recipes.git and the raw data is available from ENA with the accession number PRJEB35643

Downstream analyses were done in RStudio (version 1.0.143 based on [R] version 3.3.3) using the libraries `vegan` and `ggplot2`. Read counts were rarefied to 55,000 (CON^E experiments) or 100,000 (LCM experiments) reads per sample. To analyze changes in the microbiota composition between samples, time points or treatments, the Bray-Curtis dissimilarity was analyzed.

Anaerobic culture of *S. Typhimurium*.—*S. Typhimurium* cells were cultured in minimal M9 medium (12.8 g/L Na₂HPO₄·7H₂O, 3 g/L KH₂PO₄, 0.5 g/L NaCl, 1 g/L NH₄Cl, 100 μM CaCl₂, 2 mM MgSO₄) supplemented with Wolin's trace elements (13.4 μM EDTA, 3.1 μM FeCl₃·6H₂O, 0.62 μM ZnCl₂, 76 nM CuCl₂·2H₂O, 42 nM CoCl₂·2H₂O, 162 nM H₃BO₃, 8.1 nM MnCl₂·4H₂O), 250 mg/L ¹²C Histidine-HCl and 45 mM ¹²C pyruvate. 10 mM ¹³C C4-dicarboxylate (aspartate, fumarate, malate) was added, as indicated. Growth was analyzed under anaerobic conditions (4% H₂, 10% CO₂, 86% N₂) at 37 °C. Supernatant was sampled at the transition from exponential to stationary phase and centrifuged for 2.0 min at 11,000 × g at 4 °C to pellet and remove bacterial cells. Samples were diluted (1:1000 v/v) in a buffer corresponding to the starting conditions of the mobile phase used for liquid chromatography (see below: mass spectrometry, metabolite measurements) containing unlabeled standards.

Mass spectrometry.

Materials.: Stable isotope labeled (SIL) standards [¹³C₄]-fumaric acid (99%), [¹³C₄, ¹⁵N]-L-aspartic acid (99%), [1, 4-¹³C₂]-succinic acid (99%) and [¹³C₄]-L-malic acid (98%) were purchased from Cambridge Isotope Laboratories.

Sample preparation.: Cecum contents were recovered in pre-weighed 2 ml microfuge tubes. 0.5 ml of ice cold PBS was added per 1 g cecum content. Samples were homogenized using a tissuelyzer (Qiagen) at 25 Hz for 2.0 min, chilled for 1 min and homogenized again for another 2 min. Samples were centrifuged for 5.0 min at 20,000 × g at 4 °C to pellet cells. Supernatants were collected and centrifuged for 30 min to pellet remaining suspended particulate matter. Clear supernatants were syringe-filtered (0.22 μm). Samples were centrifuged for 2 min at 20,000 × g. Mouse chow (MC) pellets (1 pellet per sample) were coarsely grounded and collected in 2 ml microfuge tubes. MilliQ water was added to achieve a concentration of 0.033 g mouse chow /ml and cleared supernatants were prepared, as

described above. Samples were diluted (1:10 v/v for cecum content samples, 1:1 v/v for extracted mouse chow) in dH₂O containing a mix of SIL standards (aspartic acid, fumaric acid, malic acid, succinic acid). Prior to acquisition, samples were further diluted 1:10 (v/v) in a buffer corresponding to the starting conditions of the mobile phase used for liquid chromatography (see below).

Metabolite measurements. Microscale ultrahigh performance liquid chromatography mass spectrometry (UPLC-MS) was performed with a Dionex Ultimate 3000 (Thermo Scientific) hyphenated to a Q Exactive Plus mass spectrometer (Thermo Scientific) with a heated electrospray ionization (HESI) source. Separation of metabolites was achieved with a C18 column (Kinetex®, 2.6 µm Polar C18, with dimensions 100×2.1 mm) as the stationary phase, and LC-MS grade solvents A/B/C/D as a mobile phase. Solvent A (1.5% formic acid in dH₂O), and C (methanol) were kept constant at 10% and 25% respectively. For elution, solvent B (acetonitrile) was used against solvent D (dH₂O) with the following gradient: 0 min, 0%; 0.4 min, 0%; 1.4 min, 50%; 2.4 min, 50%; 2.9 min, 0%; 7.0 min, 0%. Flow rate was 500 µl min⁻¹; the injection volume was 2 µl. The mass spectrometer was operating in negative FTMS mode (m/z range 80 to 1000); resolution was set to 70000 (at m/z 200). Ionization was performed using a HESI probe with a sheath and aux gas flow rate of 60 and 20 respectively. Spray voltage was 3.0 kV, the S-lens RF level 50.0, capillary temperature 275°C and the aux gas heater was set to 400°C. Between samples, blank measurements were included.

Metabolites were identified based on exact monoisotopic mass (± 0.002 Da), matching retention time and predicted mass shifts with corresponding SIL analytical standards.

Metabolites were quantified using unlabeled or SIL compounds as internal standards and custom written scripts employing the Python-based platform eMZed2 (Kiefer et al., 2013) (Hartl et al., 2017) (<http://emzed.ethz.ch/>). Briefly, corresponding monoisotopic (unlabeled) and respective (labeled) SIL peaks for each compound were determined by targeted extraction. Peak areas were calculated by integration using the trapezoidal rule. Background signals as determined by the median response in blank measurements were subtracted. For Fig. 4A, a linear response was confirmed by calibration curves of isotope labeled standards spiked into a mix (n=6) of the respective sample matrices (i.e. cecum content from CON^E, LCM, or GF mice). Average metabolite concentrations were estimated for each sample type, and approximately equimolar concentrations of respective SIL standards were spiked into individual samples. As the obtained peak areas from SIL standards and analyzed samples were accordingly in the same range, metabolite concentrations were directly determined by calculating the ratio of peak areas A_{unlabeled} (sample) over A_{labeled} (SIL standard) and multiplication with the known concentration of the corresponding spiked SIL standards.

Limits of detection (LODs) were determined for the data shown in Fig. 4A using a statistical method (Armbruster and Pry, 2008). Briefly, samples were prepared as described above, and low concentration SIL standards were spiked into a mix (n=6) of respective matrices and repeatedly measured (n=8 per matrix). LODs were then calculated based on the obtained standard deviation (SD) between repeatedly measured samples.

$$LOD = Z_score * SD(\text{concentration})$$

The Z-score was based on a confidence interval of 95% and corrected for low sample number with the corresponding t-distribution. Obtained LODs were subsequently confirmed by manual inspection of calibration curves.

Additional quantifications of C4-dicarboxylates and other free metabolites in the mouse cecum in Fig. S4A: Chromatographic separation was performed via hydrophilic interaction (HILIC) on a AdvanceBio MS Spent Media column (100 × 2.1 mm, Agilent technologies) using a mobile phase A: H₂O, 10 mM ammonium acetate titrated at pH 9.0 and B: Acetonitrile, 10 mM ammonium acetate titrated at pH 9.0. Samples were prepared as above and diluted 10 times in B phase. 2 µl of sample were injected at 100% B and 0.250 ml/min followed by the following gradient: 0.0 min 100%, 2.0 min 100%, 10.0 min 60%, 10.50 min 100%, 18.0 min 100%. The qTOF (Agilent 6520) was operated in negative mode at 2 GHz extended dynamic range and with a mass/ charge (m/z) range of 50–1000 and the following source parameters: VCap: 3500 C, nozzle voltage: 2000 V, gas temp: 325°C m drying gas 5 l/min, nebulizer 30 psig. Online mass calibration was performed using a second ionization source and a constant flow (5 ul/min) of reference solution (119.0363, 301.9981 and 980.0163). Compounds were identified based on the retention time of chemical standards and their accurate mass (tolerance 20 ppm) The MassHunter Quantitative Analysis Software (Agilent, version 7.0) was used for peak integration. Quantification was based on dilution series of chemical standards. Limits of detection for each metabolite was determined as mean +3.3x standard deviation of the signal detected in the lowest detected standard concentration. Obtained LODs were subsequently confirmed by manual inspection of calibration curves.

DNA sequence analyses for Tab. S4.—We identified *Salmonella enterica* Typhimurium SL1344 genes of interest (*dcuABC*, *aspA*, *fumABC*, *frdABCD*, *sdh* and H₂-hydrogenases) in genomes from representative enteropathogens and microbiota using a reciprocal best BLAST hit approach. We used parameters previously recommended for the most sensitive and precise detection of orthologous genes in pairwise comparisons of genomes (Moreno-Hagelsieb and Latimer, 2008). Specifically, we used BLASTp of the NCBI BLAST+ (version 2.8.1) suite (Camacho et al., 2009) with soft filtering and Smith-Waterman alignment (“-seg yes - soft_masking true -use_sw_tback”) to reciprocally align the protein sequences of *S. enterica* Typhimurium with those of other genomes. We subsequently parsed the results for the best reciprocal hits using a custom python script and considered two genes as orthologs if the respective alignment score exceeded 60 bits. Protein sequences were annotated to Clusters of Orthologous Groups (Galperin et al., 2015) with EggNOG-mapper 2.0.0 (Huerta-Cepas et al., 2017) [4] in diamond mode (-m diamond), targetting the ‘bact’ database version 5.0.0 (Huerta-Cepas et al., 2019) (-d bact), using 8 threads per annotation (--cpu 8).

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical analysis—The Mann Whitney test was performed using Graphpad Prism Version 6.0 for Windows (GraphPad Software, La Jolla California USA, www.graphpad.com). P values of < 0.05 (two-tailed) were considered as statistically significant. * P < 0.05, ** P < 0.01, *** P < 0.001.

Downstream analysis of the RB-TnSeq and 16s data were performed with custom R scripts including the packages vegan (2.5–5), ggplot2 (2.2.1), dplyr (0.8.0), tidyr (0.8.2), ggrepel (0.7.0) and readr (1.1.1).

For analyzing mass spectrometry data, we used the MassHunter Quantitative Analysis Software (Agilent, version 7.0), as detailed, above.

For analyzing the C.I. data in Fig. S3B, the Multiple Comparisons Test with Sidak's correction was used.

For analyzing the attenuation of *E. coli*^{dcuAB} (Fig. 6A), Wilcoxon's matched-pairs signed rank test (nonparametric, paired) was performed.

DATA AND CODE AVAILABILITY

The published article includes all datasets generated or analyzed during this study. All code is freely available upon request (except for the commercial data analysis programs listed). The RB-TnSeq data are listed in Tab. S1, the gut colonization phenotypes of the analyzed mutants are listed in Tab. S2. The 16S sequencing data is available at <http://www.ebi.ac.uk/ena/data/view/PRJEB35643>.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

A high-throughput screen identifies key *Salmonella* genes used in initial gut colonization
Host food and microbiota provide luminal aspartate and malate for *Salmonella*
respiration The DcuABC transporters pump aspartate and malate into the *Salmonella* cell
Aspartate and malate conversion into fumarate fuels growth by H₂/fumarate respiration

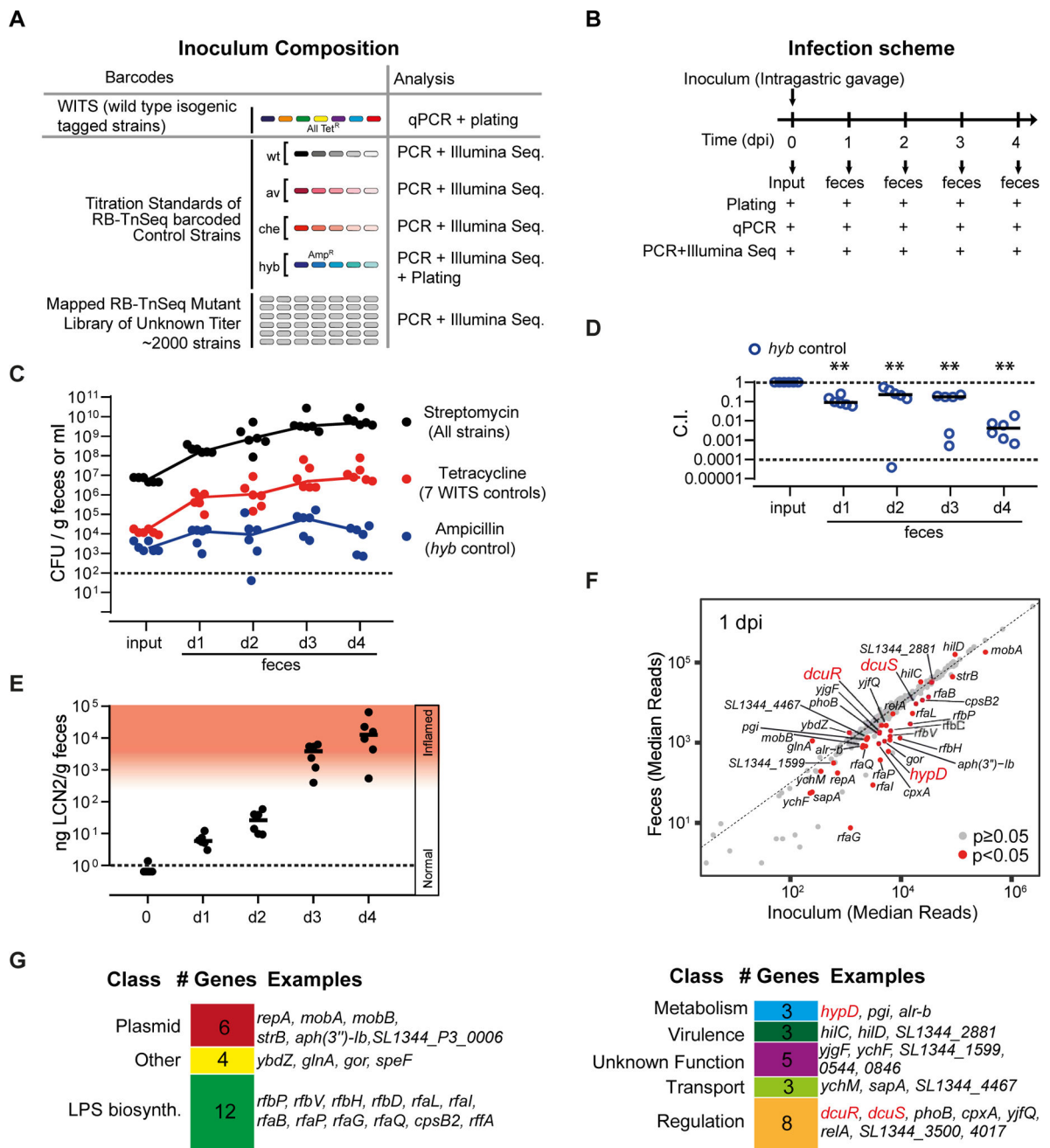


Figure 1. RB-TnSeq screen to identify *Salmonella* genes involved in growth and colonization of the murine gut.

A. Composition of the inoculum pool. WITS-barcoded wild type controls were spiked in at densities of 1/2000 and titration standards of RB-TnSeq barcoded wild type and mutant controls as detailed in Star Methods and the Key Resource Table. B. Mouse infection protocol and sampling procedures. C. *S. Typhimurium* loads in the inoculum and the feces at days 1–4 pi as determined by plating. Black: Total *S. Typhimurium* loads. Red: total loads of WITS-barcoded wild type controls. Blue: Loads of the *hyb* mutant. Dotted line: detection limit. D. Attenuation of the *hyb* control strain (C.I.). E. Gut inflammation as measured by Lipocalin-2 ELISA. F. Fitness defects of RB-TnSeq mutants. We compared the relative

barcode read abundances from the inoculum and the feces (day 1 pi; full data shown in Tab. S3). Red dots: mutants with significantly reduced or enhanced fitness compared to the wild type control strains ($p < 0.05$). G. Gene classes represented among the 43 RB-TnSeq hits implicated in initial growth.

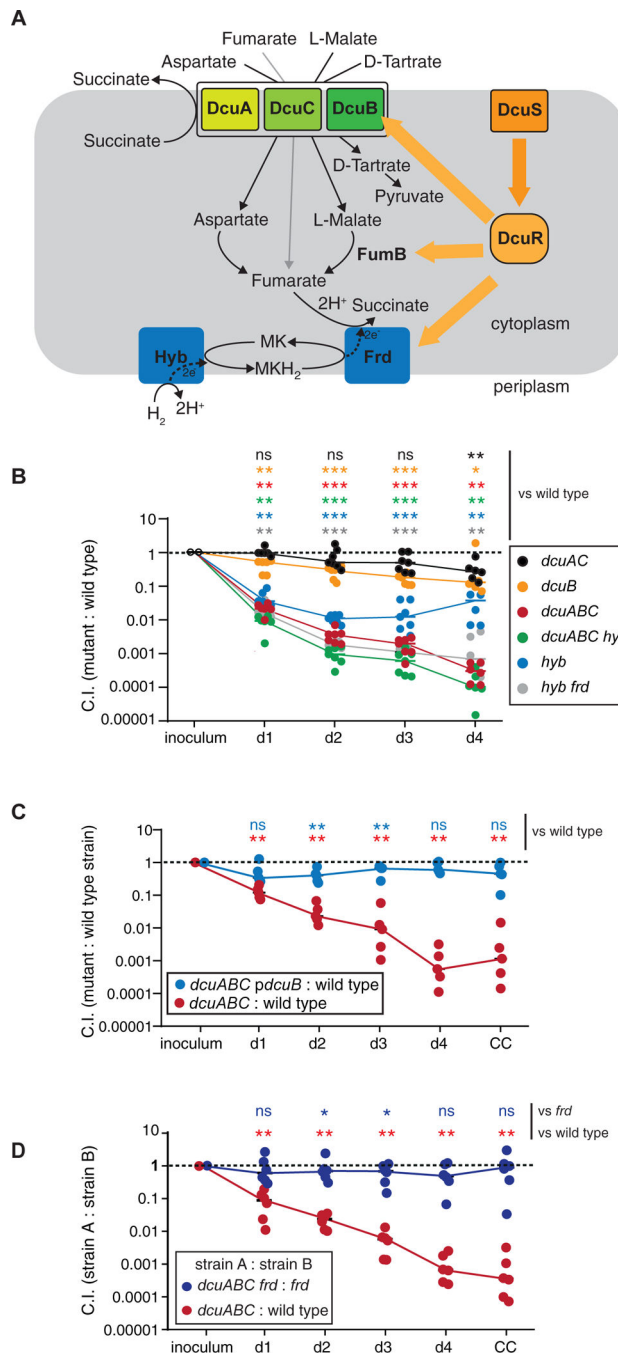


Figure 2. Dcu transporters are contributing to anaerobic respiration during initial gut luminal growth.

A. Scheme illustrating the regulation by DcuRS and the role of Dcu transporters in C4-dicarboxylate-fueled anaerobic respiration by *E. coli*. Under anaerobic conditions, and in the presence of suitable substrates, DcuRS activates expression of the indicated proteins. C4-Dicarboxylates, namely aspartate, fumarate, L-malate and L-tartrate, can be imported by Dcu transporters. Under certain conditions, they can be subsequently converted into fumarate to fuel H₂/fumarate respiration. The reduction product succinate is secreted in exchange for new aspartate, fumarate, L-malate or L-tartrate. B. Competitive infection with

seven different strains. LCM mice were infected with a mixture of untagged wild type *S. Typhimurium*, WITS-tagged wild type *S. Typhimurium* and the indicated 6 *S. Typhimurium* mutants (5×10^6 cfu in total; by gavage). Strain abundance in the inoculum and the feces was analyzed by qPCR. C. Complementation of *S.Tm^{dcuABC}*. LCM mice were infected with a mixture of wild type *S. Typhimurium*, *S.Tm^{dcuABC}* and *S.Tm^{dcuABC} (pdcuB)*, which carries a plasmid bearing a *dcuB* gene on a native promoter (5×10^6 cfu in total; by gavage). Fecal salmonellae were plated on selective media to distinguish between the three strains. D. *dcuABC* fueled growth requires the fumarate reductase. LCM mice were infected with 1:1 mixtures of wild type *S. Typhimurium* and *S.Tm^{dcuABC}* (red symbols) or *S.Tm^{dcuABC frd}* and *S.Tm^{frd}* (5×10^6 cfu in total; by gavage). Fecal salmonellae were plated on selective media to distinguish between the three strains. The Mann-Whitney U-test was used for statistical analysis; *: $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$.

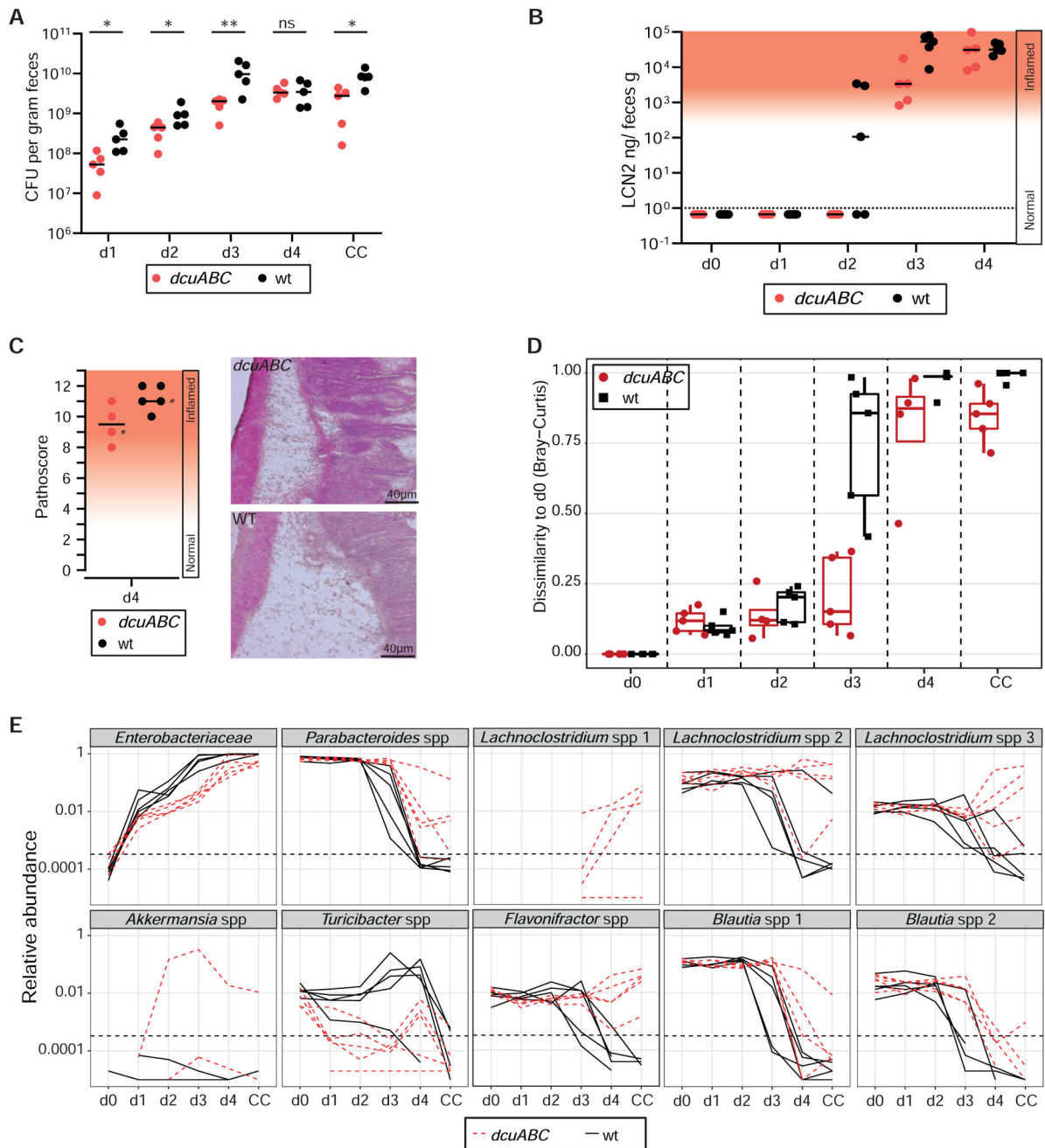


Figure 3. Dcu transporters promote gut colonization and accelerate the kinetics of mucosal inflammation and of microbiota perturbation.

A. Fecal pathogen loads in LCM mice infected with wt *S. Typhimurium* (black) or *S.Tm^{dcuABC}* (red; n=5 per group; 5×10^6 cfu; by gavage) as determined by plating. B. Gut inflammation was measured by Lipocalin-2 ELISA. C. Histopathology of the infected cecum tissue at day 4 pi. D,E. 16S rDNA analysis of the fecal microbiota composition before the infection and at days 1,2,3,4 pi. D. Bray-Curtis dissimilarity scores (black = wild type *S. Typhimurium* infection; red: *S.Tm^{dcuABC}* infection). E. Relative abundance of the identified phylotypes (rarefied to 100,000 16S sequences). Solid lines = wild type *S.Tm* infection; dashed lines = *S.Tm^{dcuABC}* infection.

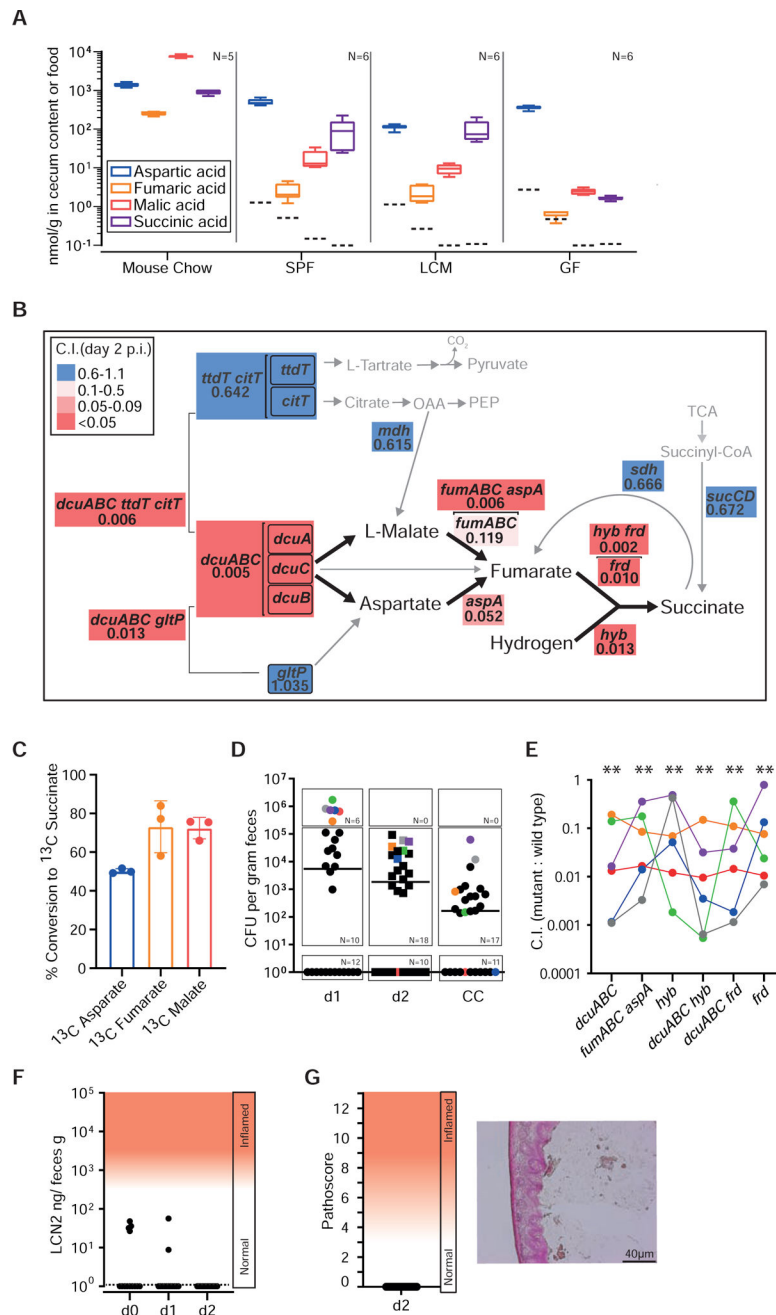


Figure 4. Aspartate and malate are important nutrients used to fuel anaerobic respiration.
 A. C4-Dicarboxylate levels in the cecum lumen. Cecum contents (without particulate matter or cells) was recovered from mouse chow, germ free (GF), specific pathogen free (CON^E), and LCM mice and analyzed by liquid chromatography mass spectrometry (LCMS). Independent measurements of n = 5 (mouse chow) and n = 6 (cecum samples) biological replicates. Dashed lines indicate the limits of detection; box plots extend from the 25th to 75th percentiles with whiskers going down to the smallest value and up to the largest. Center lines indicate median values. B. Competitive indices of site-directed mutants vs. wild type *S. Typhimurium*. We infected LCM mice with WITS-barcoded *S. Typhimurium* strain mixtures

(5×10^6 cfu in total, by gavage; at least 6 independent experiments for each mutant) and analyzed the C.I. at days 1–4 pi by qPCR (Materials and Methods). All data are summarized in Tab. S3. Panel B shows data for day 2 pi. Gene symbols denote the mutants, analyzed. Arrows: metabolic conversions; Brackets indicate pathways that were collectively deleted. Numbers: median C.I. at day 2 pi; Colored boxes indicate the degree of attenuation, as indicated in the legend. Grey: Genes that are not essential for C4-dicarboxylate driven $H_2/fumarate$ respiration (mutant C.I. values 0.6–1.1). C. Conversion of ^{13}C C4-dicarboxylates by *S. Typhimurium*. Wild type *S. Typhimurium* was grown anaerobically (37°C, 4% H_2 , 10% CO_2 , 86% N_2) in minimal medium containing 45 mM ^{12}C pyruvate and 10 mM of ^{13}C aspartate, ^{13}C fumarate or ^{13}C L-malate (Star Methods). After reaching stationary phase, cell free culture supernatants were harvested and the amounts of released ^{13}C succinate were analyzed by LC-MS (n = 3 independent cultures). D-G. Competitive infection experiment in CON^E mice. N = 30 animals were infected for two days with wild type *S. Typhimurium*, *S.Tm^{dcuABC}*, *S.Tm^{fumAC fumB aspA}*, *S.Tm^{hyb}*, *S.Tm^{dcuABC hyb}*, *S.Tm^{dcuABC frd}*, or *S.Tm^{frd}* (1:1:1:1:1:1 mix; 5×10^7 cfu in total, by gavage). D. Total fecal *Salmonella* loads as determined by plating. E. Competitive index, as determined by qPCR analysis of the strains' barcodes. F. Gut inflammation as measured in feces by Lipocalin-2 ELISA. G. Histopathology of the infected cecum tissue at day 2 pi.

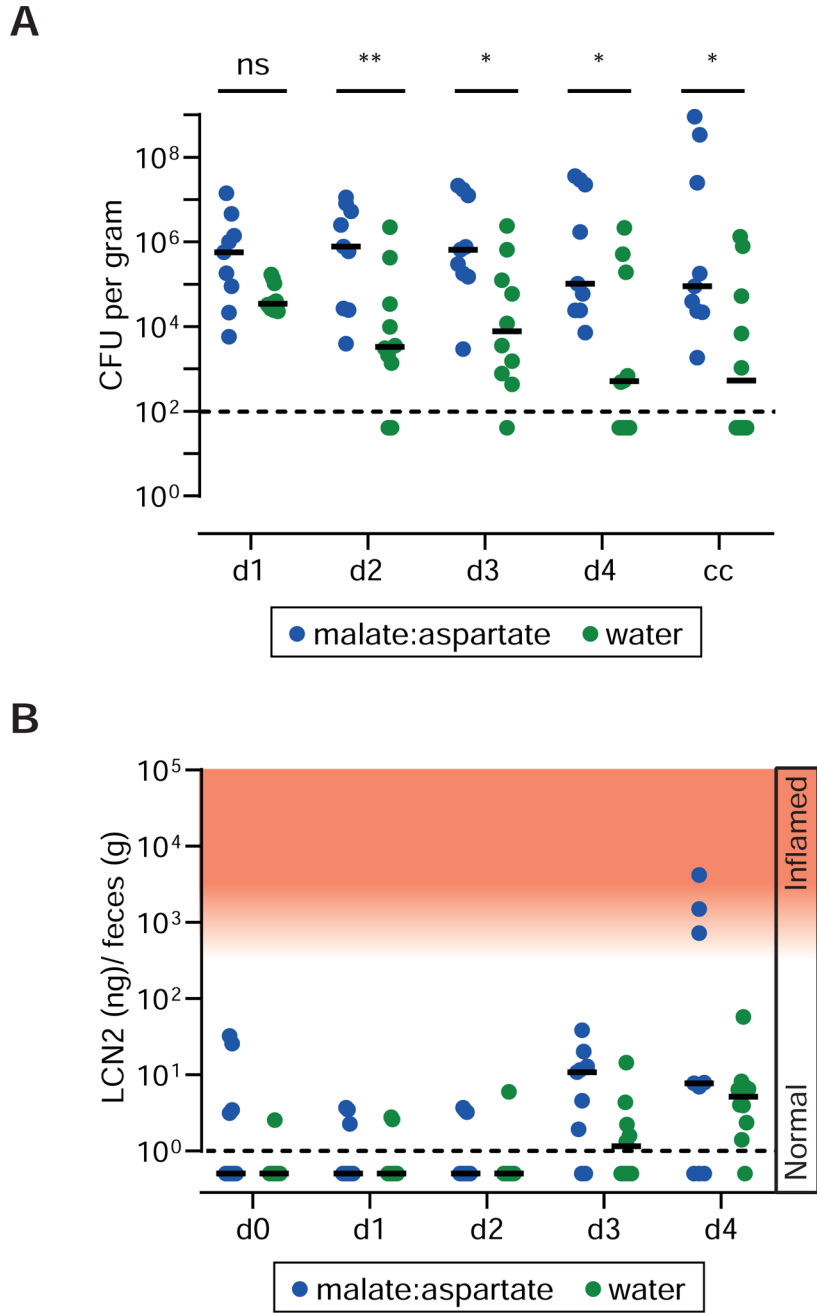


Figure 5. Oral administration of malate and aspartate promotes initial growth of *S. Typhimurium* in the gut lumen.

A. CON^E mice (N=9) were orally gavaged with 20 mg of an equal mixture (w/w) of L-aspartate and L-malate four hours after infection with a mixed inoculum composed of WITS-barcoded *S.Tm*^{wt}, *S.Tm*^{dcuABC}, *S.Tm*^{fumABC aspA}, *S.Tm*^{hyb}, *S.Tm*^{aspA}, *S.Tm*^{fumABC}, *S.Tm*^{frd}, and untagged wild type *S. Typhimurium* (5×10^7 cfu; mixed inoculum of 8 strains) and once per day thereafter. Control mice were treated with water (N=10). A. Total *S. Typhimurium* loads in the feces collected at days 1–4 pi. B. Lipocalin-2 ELISA.

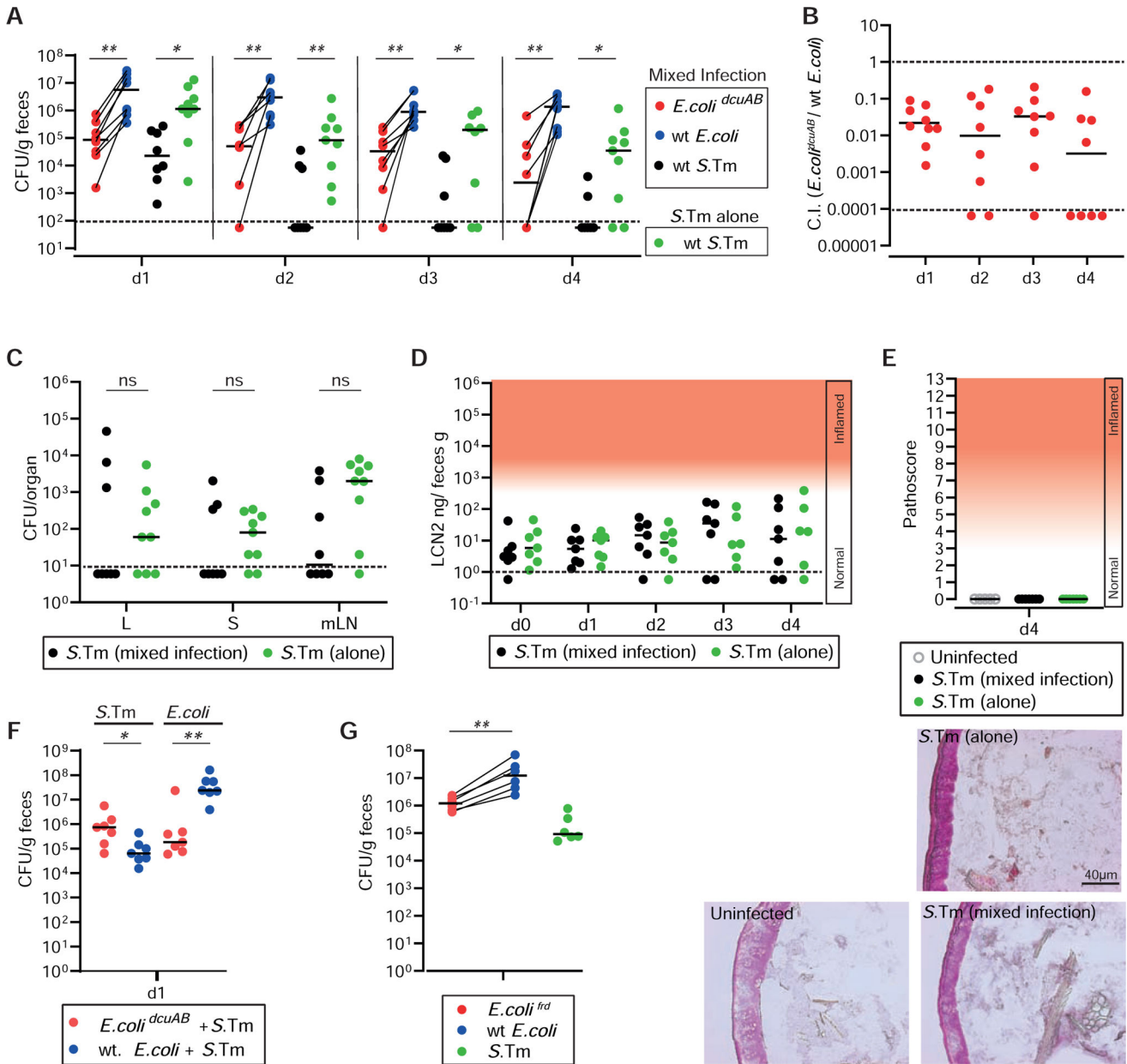


Figure 6. Competitive growth of *E. coli* 8178 requires DcuAB.

A. CON^X mice were infected for 4 days with a 1:1:1 mix of wild type *E. coli* (Cm^R), *E. coli*^{*dcuAB*} (Km^R) and wild type *S. Typhimurium* (Sm^R; N = 8) or with *S. Typhimurium* alone (N = 9; 10⁸ cfu each; by gavage). A. Fecal loads of the indicated *E. coli* and *S. Typhimurium* strains as determined by plating on MacConkey agar. B. C.I. values for *E. coli*^{*dcuAB*} vs. wild type *E. coli*. C. *S. Typhimurium* loads in livers, spleens and mesenteric lymph nodes. D. Gut inflammation as measured by Lipocalin-2 ELISA. E. Representative images of hematoxylin and eosin (H&E) stained cecum tissue from uninfected mice or animals infected for 4 days with the mix, or wild type *S. Typhimurium*, alone. F. CON^X mice were infected for 1 day with a 1:1 mix of wild type *E. coli* (Cm^R) and wild type *S. Typhimurium* (Sm^R) or with a 1:1 mix of *E. coli*^{*dcuAB*} (Km^R) and wild type *S. Typhimurium* (N = 7; 10⁸ cfu each; by

gavage). We determined fecal loads of the indicated *E. coli* and *S. Typhimurium* strains by plating on MacConkey agar. G. CON^X mice were infected for 1 day with a 1:1:1 mix of wt *E. coli* (Cm^R), *E. coli*^{fitd} (Km^R) and wild type *S. Typhimurium* (Sm^R; N = 6). Fecal loads of the indicated *E. coli* and *S. Typhimurium* strains were determined by plating on MacConkey agar.

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Table 1.

Gut colonization of deletion mutants in LCM mice (C.I. values for hits identified in Fig.1)*.

gene	function	gene #	day 1 p.i.		day 2 p.i.		day 3 p.i.		day 4 p.i.	
			median C.I.	p value	median C.I.	p value	median C.I.	p value	median C.I.	p value
<i>rfaP</i>	LPS synth.	3687	0.10	0.002	0.11	1E-03	0.06	1E-03	0.00	0.016
<i>gor</i>	other	3562	0.11	0.002	0.13	1E-03	0.07	1E-03	0.00	0.016
<i>rfbP</i>	LPS synth.	2059	0.62	0.027	0.33	1E-03	0.21	2E-03	0.43	0.016
<i>cpxA</i>	regulation	4007	0.29	0.002	0.11	1E-03	0.09	1E-03	0.00	0.016
<i>rfaB</i>	LPS synth.	3685	0.21	0.002	0.13	1E-03	0.06	1E-03	0.01	0.016
<i>phoB</i>	regulation	392	0.28	0.002	0.23	1E-03	0.26	1E-03	0.49	0.016
<i>yjgF</i>	unknown	4389	0.28	0.002	0.21	1E-03	0.23	1E-03	0.40	0.016
<i>yjfQ</i>	regulation	4314	0.30	0.002	0.16	1E-03	0.13	1E-03	0.00	0.016
<i>dcuR</i>	regulation	4240	0.34	0.002	0.27	1E-03	0.25	1E-03	0.01	0.016
<i>hilC</i>	virulence	2847	1.14	0.014	1.25	1E-02	2.63	2E-01	6.13	0.047
<i>hilD</i>	virulence	2855	1.42	0.049	1.42	4E-01	2.76	7E-02	6.97	0.016

* we selected the hits with C.I.s < 0.68 or >2.06 in RB-TnSeq screen and analyzed 10 (or 9) mice (day 1), 11 mice (day 2, day 3), or 7 mice (day 4) respectively; The full dataset is shown in Tab. S2

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Bacterial and Virus Strains		
See Supplemental Table S5		
Biological Samples		
Chemicals, Peptides, and Recombinant Proteins		
Isotopes: [13C4]-fumaric acid (99%), [13C4, 15N]-L-aspartic acid (99%), [1, 4-13C2]-succinic acid (99%) and [13C4]-L-malic acid (98%)	Cambridge Isotope Laboratories	CLM-1529, CNLM-544-H-0.25, CLM-1084-PK CLM-8065-PK, respectively
Critical Commercial Assays		
DuoSet Lipocalin ELISA kit	R&D Systems, Minneapolis, MN, USA	DY1857
AllPrep DNA/RNA Kit	Qiagen	80204
NEXTflex® 16S V4 Amplicon-Seq Kit 2.0	Bio Scientific	NOVA-4203-04
Deposited Data		
16S sequencing data of microbiome in mice	this manuscript	http://www.ebi.ac.uk/ena/data/view/PRJEB35643 .
Experimental Models: Cell Lines		
Experimental Models: Organisms/Strains		
Mouse: LCM mice (C57BL/6 genetic background)	(Stecher et al., 2010) (Maier et al., 2013)	N/A
<i>Mouse: 129SvEv</i>	Jackson Laboratories; bred at EPIC mouse facility of ETH Zurich, Switzerland	N/A
Mouse: C57BL/6	Jackson Laboratories; bred at EPIC mouse facility of ETH Zurich, Switzerland	N/A
Oligonucleotides		
See Supplemental Table S6		

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Recombinant DNA		
Software and Algorithms		
Graphpad Prism Version 6.0 for Windows	GraphPad Software, La Jolla California USA	N/A
eMZed2	(Kiefer et al., 2013) (Hartl et al., 2017)	http://emzed.ethz.ch/
BLASTp of the NCBI BLAST+ (version 2.8.1)	(Camacho et al., 2009)	N/A
Clusters of Orthologous Groups with EggNOG-mapper 2.0.0	(Galperin et al., 2015) (Huerta-Cepas et al., 2017)	N/A
MassHunter Quantitative Analysis Software	(Agilent, version 7.0)	N/A
Other		

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