Chemotaxis and autoinducer-2 signalling mediate colonization and contribute to co-existence of Escherichia coli strains in the murine gut

**Journal Article** 

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Chemotaxis and autoinducer-2 signalling enhance gut colonization and contribute to 1 2 niche segregation of Escherichia coli strains in the mammalian gut Leanid Laganenka<sup>1</sup>, Jae-Woo Lee<sup>2</sup>, Lukas Malfertheiner<sup>3</sup>, Cora Lisbeth Dieterich<sup>1</sup>, Lea Fuchs<sup>1</sup>, 3 Jörn Piel<sup>1</sup>, Christian von Mering<sup>3</sup>, Victor Sourjik<sup>2</sup>, Wolf-Dietrich Hardt<sup>1</sup> 4 5 <sup>1</sup>Institute of Microbiology, D-BIOL, ETH Zurich, Zurich, Switzerland 6 <sup>2</sup>Max Planck Institute for Terrestrial Microbiology and Center for Synthetic Microbiology (SYNMIKRO), 7 Marburg, Germany 8 <sup>3</sup>Department of Molecular Life Sciences and SIB Swiss Institute of Bioinformatics, University of Zurich, 9 Zurich, Switzerland Corresponding author: Wolf-Dietrich Hardt, hardt@micro.biol.ethz.ch 10 11 12 Abstract 13 Bacteria communicate and coordinate their behavior by producing and sensing 14 extracellular small molecules called autoinducers. The astounding structural diversity 15 of these molecules allows bacteria to synchronize their behavior on both intra- and 16 interspecies levels. Autoinducer 2 (AI-2) is produced and detected by a variety of 17 bacteria, thus principally allowing interspecies communication. Although AI-2 is a major 18 autoinducer molecule present in the mammalian gut, its role in bacteria-bacteria and 19 20 bacteria-host interactions during gut colonization remains elusive. Here, we show that 21 chemotaxis and AI-2 signalling promote gut colonization by Escherichia coli, which is in turn connected to the ability of the bacteria to utilize fructoselysine. We further show 22 23 that the genomic diversity of *E. coli* strains with respect to AI-2 signaling allows ecological niche segregation and stable co-existence of different E. coli strains in the 24 25 mammalian gut. 26 27 Introduction 28 Chemotaxis allows motile bacteria to navigate in chemical gradients. Although being a 29 costly cellular behaviour, chemotaxis provides bacteria with physiological advantage by 30 enhancing access to nutrient and energy sources<sup>1–3</sup>. Furthermore, it facilitates detection and 31 colonization of beneficial niches by free-living and host-associated bacteria, including plant 32

and human pathogens<sup>4</sup>. In the latter case, some sensed compounds might serve as orientation
 cues within the host, even if they lack a direct metabolic value. These include hormones,

neurotransmitters, acids, and other compounds<sup>4-7</sup>. However, despite the ever-growing 35 mechanistic understanding of chemotaxis systems and the range of molecules sensed by them 36 in diverse bacteria, the ecological role of chemotaxis has received much less attention<sup>8–10</sup>. The 37 38 physiological importance of chemotaxis has remained incompletely understood even for well-39 established model organisms such as E. coli. To our knowledge, only one study addressed 40 this question, finding that neither motility nor chemotaxis are required for gut colonization of E. 41 *coli* F-18<sup>11</sup>. However, this strain was shown to lose motility during growth in the mouse large intestine, which was associated with mutations in the regulatory region of *flhDC* operon<sup>12</sup>. 42 Besides leading to loss of flagella, these mutations further resulted in beneficial pleiotropic 43 44 metabolic effects, making it impossible to draw any conclusions about the role of chemotaxis 45 in motile *E. coli*.

46 Recent ex vivo studies have shown that collective behaviours of E. coli such as 47 autoaggregation and biofilm formation are dependent on chemotaxis towards the interspecies quorum sensing signal autoinducer-2 (AI-2)<sup>13–15</sup>. Self-produced AI-2 attracts bacteria towards 48 the growing aggregates, and it further enhances mature biofilm formation in a chemotaxis-49 dependent manner. AI-2 is produced and sensed by a vast number of bacterial species, and 50 AI-2 mimics were reported to be synthesized by eukaryotic cells<sup>16–18</sup>. The chemotactic 51 response to AI-2 has been clearly shown for several bacterial species, and it is apparently not 52 restricted to AI-2-producing bacteria, suggesting its important role in establishing complex 53 multispecies communities<sup>19,20</sup>. Indeed, AI-2 seems to affect bacterial community structure of 54 55 the mammalian gut after antibiotic-induced dysbiosis and to promote colonization resistance to certain enteric pathogens<sup>21,22</sup>. However, there are still fundamental mechanistic gaps in our 56 57 knowledge of how bacteria might benefit from AI-2 signalling under physiologically relevant 58 conditions.

Here we studied the roles of chemotaxis and AI-2 signalling in *E. coli* gut colonization. We show that chemotaxis towards self-produced AI-2 provides *E. coli* with a fitness advantage during gut colonization, and that this hinges on fructoselysine metabolism. We further report the novel role of AI-2 chemotaxis in contributing to niche segregation and thus to co-existence of different *E. coli* strains in the gut based on their ability to perform AI-2 chemotaxis. These findings might be relevant for other AI-2 chemotactic bacteria in their natural habitats.

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# Results

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68 **Chemotaxis provides** *E. coli* with a fitness advantage during gut colonization. To 69 assess the role of chemotaxis in gut colonization by *E. coli*, we infected ampicillin-pretreated 70 specific pathogen-free (SPF) mice with a 1:1 mixture of *E. coli* Z1331 wild-type strain, a motile 71 stool isolate from a healthy human volunteer<sup>23</sup>, and its non-chemotactic  $\Delta cheY$  derivative (Fig.

1a). A wild-type *E. coli* isolate was chosen over the classical K-12 laboratory strain since the 72 accumulation of lab cultivation-derived mutations might result in overall loss of fitness during 73 the gut colonization and thus compromise the physiological relevance of the study<sup>24–27</sup>. Since 74 SPF mice are normally resistant to E. coli colonization, antibiotic pretreatment is required to 75 76 transiently suppress resident gut microbiota, thus allowing for gut luminal E. coli colonization<sup>28</sup>. 77 Levels of *E. coli* colonization and relative fitness of the WT and  $\Delta cheY$  strains were determined 78 by differential plating of faeces collected at several time points within 72 h.p.i.. E. coli Z1331 colonized the gut of the mice within 8 h.p.i. at densities of  $\approx 10^9$  c.f.u./g stool and remained at 79 carrying capacity throughout the course of the experiment (Extended Data Fig. 1a). In 80 81 competitive infections with a 1:1 inoculum,  $\Delta cheY$  knockout cells were consistently 82 outcompeted by up to 50-fold by the wild-type strain, indicating that chemotaxis is required for successful gut colonization by E. coli Z1331 (Fig. 1b, see Materials and Methods for the 83 rationale for using competitive infections). This phenotype was observed along the whole 84 length of the small and the large intestine at 72 h.p.i. (Fig. 1c), indicating that the competitive 85 fitness advantage provided by chemotaxis is not limited to a particular region of the mouse gut. 86 Interestingly, although in Salmonella enterica serovar Typhimurium, a close relative of E. coli, 87 the beneficial role of chemotaxis only becomes apparent at high levels of gut inflammation<sup>29,30</sup>, 88 this was not the case for E. coli Z1331 in our experiments. The Lipocalin-2 level, a marker of 89 gut inflammation, remained within the range of concentrations characteristic for unperturbed 90 91 mice during the entire E. coli colonization experiment (Extended Data Fig. 1b). Notably, a 92 similar loss of fitness was observed for a  $\Delta cheY$  knockout mutant of E. coli K-12 W3110 as well as for several other *E. coli* isolates from different phylogroups (Extended Data Fig. 1c). 93 94 These results suggest that the competitive colonization benefit provided by chemotaxis might 95 be not strictly strain-specific, but rather of general importance for motile *E. coli* strains.

96 To visualize the spatial distribution of the WT and  $\Delta cheY$  cells in the gut by confocal 97 microscopy, we infected mice with the WT and  $\Delta cheY$  cells that constitutively express mCherry and GFP, respectively. At 72 h.p.i., ileal, caecal and proximal colon tissues were excised and 98 99 fixed in 4% paraformaldehyde. Subsequently, 10 µm tissue sections were additionally stained with DAPI and phalloidin to visualize the host tissue. As seen in Fig. 1d, WT cells appeared to 100 form clusters reminiscent of the previously described aggregates<sup>14</sup>, whereas much less 101 aggregate formation was observed for  $\Delta cheY$  cells (Fig. 1d, e; Extended Data Fig. 2). Less 102 aggregation of  $\Delta cheY$  knockout was also observed in single-strain infections, where no 103 colonization defect of  $\Delta cheY$  was detected (Extended Data Fig. 3). Our results suggest that 104 chemotaxis contributes to gut colonization and spatial organization of *E. coli* cells in the gut. 105

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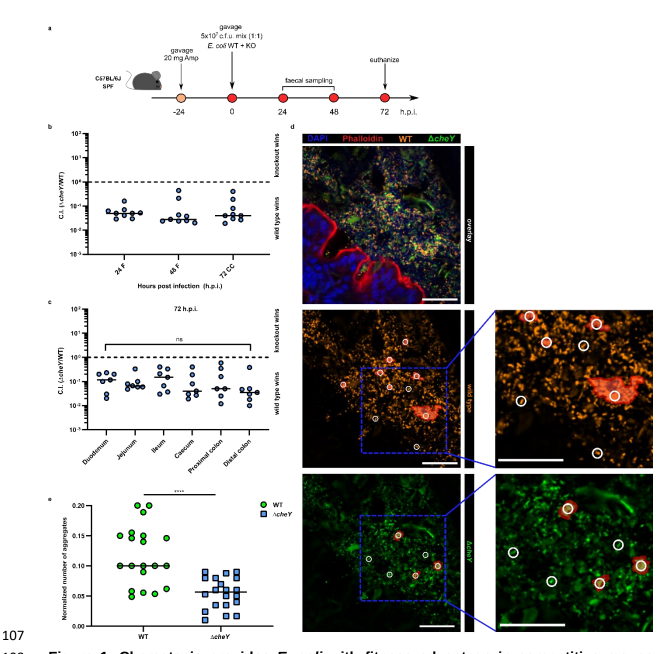


Figure 1. Chemotaxis provides *E. coli* with fitness advantage in competitive mouse 108 infections. a, Experimental scheme of competitive infection. C57BL/6J specific pathogen-free 109 (SPF) mice were pretreated with 20 mg ampicillin by oral gavage 24 h prior to infection with E. 110 coli (1:1 mix WT and KO strain). Faeces were collected at 24, 48 h.p.i., unless stated otherwise, 111 and mice were euthanized at 72 h.p.i. **b**, Competitive index (C.I.) of non-chemotactic  $\Delta cheY$ 112 (Z7741, unless stated otherwise) mutant in  $\Delta cheY/WT$  competitive infection. F, faeces, CC, 113 caecal content. Lines indicate median values (n=9, at least two independent replicates). c, C.I. 114 values of  $\Delta cheY$  in  $\Delta cheY/WT$  competitive infection along the gut at 72 h.p.i. Lines indicate 115 median values (n=7, at least two independent replicates). P values were analyzed using one-116 way ANOVA test (ns, not significant). d, Caecal tissue sections of mice infected with E. coli 117 WT (mCherry-positive, shown in orange) and  $\Delta cheY$  (Z7730, GFP-positive, shown in green) 118 at 72 h.p.i (\(\Delta che Y/WT C.I.=0.16)\). Actin filaments (red) and DNA (blue) were stained with 119 120 phalloidin and DAPI, respectively. White circles indicate the threshold area ( $\sim$ 50 px<sup>2</sup>) for

adjacent cells (~10 px<sup>2</sup> in size) to be considered an aggregate. Examples of such aggregates are highlighted in red. Scale bars, 50  $\mu$ m. **e**, Number of aggregates formed by WT and  $\Delta cheY$ cells normalized to the number of detected cells in a tissue section (Mann-Whitney test, \*\*\*\**P*<0.0001). Lines indicate median values (n=20, tissues sections from two independent experiments were analyzed).

- 126
- 127 Chemotaxis towards self-produced AI-2 enhances gut colonization by E. coli. As 128 stated above, the clusters of *E. coli* cells observed in the gut tissue sections were similar to the aggregates formed by swimming E. coli cells in liquid medium. Since aggregation ex vivo was 129 shown to be dependent on chemotaxis towards self-produced AI-2<sup>14</sup>, we hypothesized that AI-130 131 2 chemotaxis might as well play a role during gut colonization. AI-2 is known to be produced and sensed by variety of both Gram-positive and Gram-negative bacteria<sup>16</sup>, and a substantial 132 number of gut-associated bacteria encode LuxS, the AI-2 synthase enzyme, potentially 133 rendering AI-2 the most abundant interspecies guorum-sensing molecule in the gut<sup>31</sup>. Ex vivo, 134 135 E. coli, chemotaxis towards AI-2 has been previously shown to control such group behaviours as autoaggregation and biofilm formation<sup>14,15,32,33</sup>. In contrast, its role *in vivo* remained unclear. 136
- When extracellular AI-2 exceeds a certain threshold ex vivo, E. coli strains activate 137 expression of the Isr operon, which contains genes required for AI-2 import (via an ABC 138 139 transporter) and degradation. The LsrB protein binds AI-2 in the periplasm and directs its import via the LsrACD ABC transporter<sup>34,35</sup>. Additionally, AI-2-bound LsrB elicits a chemotactic 140 141 response of *E. coli* cells to AI-2 by binding to the Tsr chemoreceptor<sup>36</sup>. As no chemotaxisindependent AI-2-related phenotypes have been observed in *E. coli*<sup>13</sup>, deletion of the *lsrB* gene 142 alone should be sufficient to abolish chemotaxis to AI-2 and AI-2-mediated phenotypes in vitro. 143 Interestingly, ampicillin pretreatment of the SPF mice resulted in the transient increase of 144 luminal AI-2 levels at 24 h post treatment, possibly due to the lysis of the resident microbial 145 146 cells or shifts in the microbiota composition towards AI-2 producing bacteria (Extended Data Fig. 4a, b). As we could further detect *lsr* operon expression by *E. coli* Z1331 in the mouse gut 147 lumen (Fig. 2a), we decided to explore the function of LsrB in gut colonization. Similar to the 148 149  $\Delta cheY$  knockout, WT cells consistently outcompeted the isogenic  $\Delta lsrB$  mutant in SPF mice already after 24 h.p.i. (Fig. 2b). In contrast, no fitness defect was observed for the  $\Delta$ *lsrC* and 150  $\Delta$ *lsrD* knockouts, indicating that the  $\Delta$ *lsrB* phenotype is indeed attributable to the lack of 151 chemotaxis towards AI-2, rather than impaired AI-2 import. Loss of chemotaxis towards AI-2 152 similarly affected the fitness of E. coli K-12 W3110 and 8850, other Isr operon-encoding 153 154 isolates (Extended Data Fig. 5).

To further test our hypothesis, we took advantage of a well-established Al-2 overproducing *E. coli* strain (ARO071)<sup>21</sup>. Since Al-2 is sensed indirectly via an Al-2 binding protein LsrB, a narrow sensitivity range would be expected due to saturation of the receptors

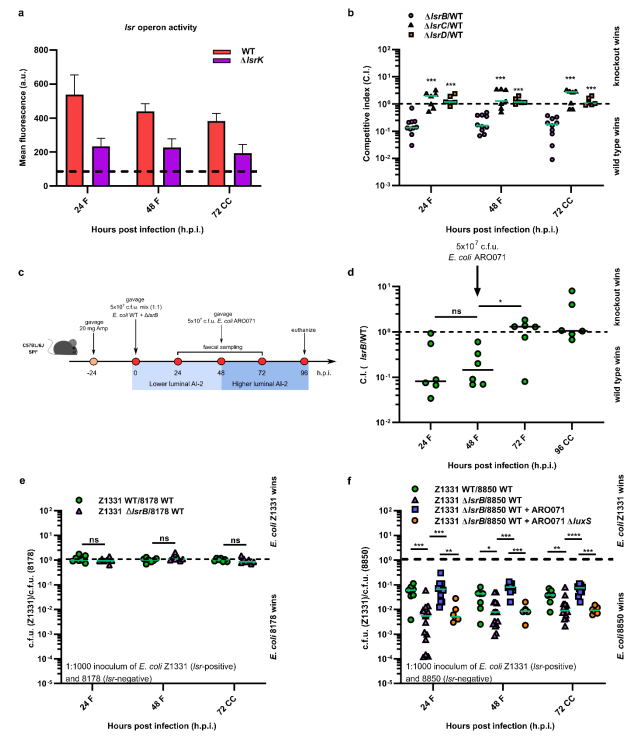
at high background stimulation<sup>37</sup>. Increasing luminal AI-2 concentrations by introducing E. coli 158 ARO071 should thus saturate the chemotactic response and eliminate the advantage of the 159 160 WT in a competitive infection. To test this, we again infected mice with a 1:1 mix of the WT E. 161 coli Z1331 and its isogenic  $\Delta lsrB$  mutant. Although  $\Delta lsrB$  was again stably outcompeted by the WT at 24 and 48 h.p.i. (C.I.≈10<sup>-1</sup>), introducing the AI-2-overproducing strain at 48 h.p.i. 162 abolished the competitive advantage of the WT strain within one day (C.I.≈1; Fig. 2c, d; 163 164 Extended Data Fig. 4b, c). This was apparently related to a slight (though not significant) rise in the stool density of the  $\Delta$ *lsrB* mutant that may go along with a very slight decrease in WT *E*. 165 166 coli Z1331 densities (which is again not significant in our experiment). Regardless, we 167 hypothesize that the observed shift in the C.I. values is attributable to the dynamic nature of 168 the favourable niches in the gut. Any E. coli-occupied niche, in the mucus layer or in the gut lumen, is constantly washed out and renewed. Therefore, such niches must be constantly re-169 occupied by the respective strains. Abolishing AI-2 chemotaxis by saturating luminal AI-2 170 concentrations after two days of the competitive infection results in both wild-type and  $\Delta lsrB$ 171 strains having the same chance of establishing themselves in the newly opened niches by 172 means of random motility. Furthermore, the dispersal of the wild-type E. coli from the existing 173 niches (like the aggregates and biofilms of E. coli disperse upon addition of saturating AI-2 174 concentrations in vitro<sup>14</sup>) might also contribute to this process. Upon addition, *E. coli* ARO071 175 might also compete transiently with E. coli Z1331 WT for a niche that was previously accessible 176 177 to this strain, and might thereby contribute to  $\Delta IsrB$  reaching WT *E. coli* Z1331 densities.

178 Our further analysis showed that  $\Delta cheY$  or  $\Delta cheY \Delta lsrB$  featured the same competitive 179 defect against the wild-type strain, while  $\Delta lsrB$  or  $\Delta cheY$  were about as competitive as  $\Delta cheY$ 180  $\Delta lsrB$  (Extended Data Fig. 6). These results strongly suggest that chemotaxis towards AI-2 181 enhances gut colonization by *E. coli*. Interestingly, we noticed a minor fitness defect of  $\Delta cheY$ 182 in the  $\Delta lsrB$  vs.  $\Delta cheY \Delta lsrB$  infection (Extended Data Fig. 6). Altogether, our observations 183 suggest that AI-2 is the major chemoeffector, albeit not the only one, contributing to 184 chemotaxis-driven gut colonization of ampicillin-pretreated SPF mice by *E. coli*.

185 It has been recently shown that AI-2 mimics can be produced by eukaryotic cells: by Saccharomyces cerevisiae and, more intriguingly, by intestinal epithelial cells<sup>17,18</sup>. This 186 187 suggests that there are three potential sources of AI-2-type molecules in the gut which might 188 affect *E. coli* colonization in our experiments: epithelium-, microbiota-, or self-produced signals. We therefore aimed at pinpointing the major source of AI-2 or AI-2 mimics sensed by the E. 189 coli Z1331 cells. To distinguish between host- and self-produced molecules, we investigated 190 the role of LsrB during gut colonization of germ-free (GF) mice in WT E. coli Z1331 and an 191 isogenic strain incapable of AI-2 production (E. coli Z1331 \DeltaluxS). As in the case of ampicillin-192 193 pretreated SPF mice,  $\Delta$ *lsrB* was outcompeted by the WT strain in GF mice (without antibiotic 194 pretreatment; Extended Data Fig. 7a, b). However, no advantage of LsrB was observed when both strains were lacking *luxS*. Thus, it is not epithelium-derived, but self-produced AI-2 that is
involved in colonization. In agreement with this explanation, no accumulation of the WT cells
was observed close the epithelial tissue (Fig. 1d, e).

Although antibiotic treatment leads to suppression of resident microbiota and stochastic 198 199 loss of bacterial phyla in SPF mice, the microbiota is never completely cleared by the antibiotic and regrows within several days after the treatment<sup>38</sup>. It was thus interesting to see whether 200 AI-2 produced by residual microbiota may contribute to the LsrB-mediated phenotype in E. coli 201 202 Z1331. To answer this question, we repeated the experiment described above in ampicillin-203 pretreated SPF mice. Again, no fitness loss was observed for  $\Delta lsrB\Delta luxS$  vs  $\Delta luxS$ , suggesting 204 that self-produced AI-2 that is sensed by *E. coli* during gut colonization confers the competitive 205 advantage in all our initial experiments (Extended Data Fig. 7c, d). 206

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Figure 2. Chemotaxis towards AI-2 promotes colonization and drives ecological niche 209 segregation of E. coli strains in the gut. a, In vivo Isr promoter activity in wild-type E. coli 210 cells transformed with pUA66::P/sr-gfp, as measured by flow cytometry and expressed in 211 212 arbitrary units (a.u.).  $\Delta$ *lsrK* strain (no activation of *lsr* operon) was used as a negative control. Error bars indicate s.d. (n=5, at least two independent replicates). Dashed line indicates mean 213 214 fluorescence of E. coli cells harbouring the promoterless pUA66 plasmid as measured at 8 215 h.p.i. F, faeces. **b**, Competitive infection experiments. C.I. values for  $\Delta IsrB$ ,  $\Delta IsrC$  and  $\Delta IsrD$  in competitive infections against the WT strain. F, faeces, CC, caecal content. Lines indicate 216 median values (minimum n=6, from at least two independent experiments). P values were 217

calculated using the Mann-Whitney test (\*\*\*P<0.0005). c, Experimental scheme for 218 competitive infection experiments with subsequent introduction of the AI-2- overproducing 219 strain E. coli strain ARO071. SPF mice were pretreated with 20 mg ampicillin by oral gavage 220 221 24 h prior to inoculation with *E. coli* (1:1 mix WT vs Δ*lsrB*). At 48 h.p.i., mice were additionally inoculated with 5x10<sup>7</sup> c.f.u. of *E. coli* ARO071 by oral gavage, resulting in increased luminal 222 concentration of AI-2. d, C.I. values for ΔlsrB in Z1331 ΔlsrB vs WT E. coli Z1331 competitive 223 224 infection before and after addition of the AI-2-overproducing E. coli ARO071. F, faeces, CC, caecal content. Lines indicate median values (n=6, from at least two independent 225 226 experiments). P values were calculated using the Mann-Whitney test (\*P<0.05; ns, not 227 significant). e, Competitive infection experiments of E. coli Z1331 vs E. coli 8178 (Isr operon-228 negative) or E. coli Z1331 ΔlsrB vs E. coli 8178. SPF mice were infected with E. coli Z1331 WT or  $\Delta$  lsrB and E. coli 8178 WT (5x10<sup>7</sup> c.f.u. by gavage; 1:1000 ratio). F, faeces, CC, caecal 229 content. Lines indicate median values (n=7, from at least two independent experiments). P 230 values were calculated using the Mann-Whitney test (ns, not significant). f, Competitive 231 232 infection experiments with or without AI-2 overproduction. SPF mice were infected with E. coli Z1331 WT or  $\Delta$ IsrB and E. coli 8850 WT (Isr operon-positive; 5x10<sup>7</sup> c.f.u. by gavage; 1:1000 233 ratio). To probe the effect of AI-2 overproduction, mice were inoculated with Z1331 vs 8850 as 234 above, supplemented with *E. coli* ARO071 (5x10<sup>7</sup> c.f.u.; by gavage; at 1:1 ratio). F, faeces, 235 CC, caecal content. Lines indicate median values (minimum n=6, from at least two 236 independent experiments). P values were calculated using the Mann-Whitney test 237 (\*\*\*\**P*<0.0001; \*\*\**P*<0.0005; \*\**P*<0.005; \**P*<0.05). 238

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Genomic diversity of Al-2 sensing contributes to niche segregation of *E. coli* strains in the gut. The mammalian gut is a complex and dynamic environment with a fine scale spatial structure. Heterogenous spatial distribution of available nutrients, microorganisms, signalling and other host-derived molecules allows niche segregation and can thereby permit the stable co-existence of several strains of a given bacterial species based on their metabolic preferences<sup>39–41</sup>. The relevant metabolic pathways are thought to differ from case to case, and they were unknown for *E. coli* Z1331.

247 E. coliniche segregation was shown to depend at least partially on the differential ability of individual strains to utilize certain compounds as carbon or nitrogen source<sup>42,43</sup>. Intriguingly, 248 the *lsr* operon (and thus the ability to sense and chemotactically respond to AI-2) is found in 249 some, but absent in other *E. coli* strains<sup>44</sup>. We hypothesized that AI-2 sensing might contribute 250 to niche segregation of *E. coli* strains in the gut. To test this, we analysed the competition of 251 E. coli Z1331 against two other E. coli mouse isolates, E. coli 8178 which naturally lacks the 252 253 Isr-operon and is therefore devoid of AI-2 signaling<sup>45</sup>; and *E. coli* 8850, which is naturally Isr positive<sup>45</sup>. The latter two strains were applied in a 1000-fold surplus (compared to *E. coli* 254

Z1331). The overabundance of *E. coli* 8178 or 8850 in the inoculum would allow these strains 255 to rapidly occupy their respective niches. If chemotaxis towards AI-2 indeed allows E. coli 256 257 Z1331 to reach a distinct niche and thus avoid direct competition with E. coli 8178, it should 258 be able to grow up to reach a 1:1 ratio (at least in the absence of other competitive effects like bacteriocins production<sup>46</sup>). If true, *E. coli* Z1331 should not be able to catch up in growth with 259 260 Isr positive E. coli 8850 strain. As seen in Fig. 2e and Extended Data Fig. 8d, E. coli Z1331 261 was indeed able to reach a 1:1 ratio compared to E. coli 8178 within 24 h.p.i. This phenotype was independent of LsrB, suggesting that E. coli Z1331 and 8178 do not compete for the LsrB-262 263 dependent niche. In this case, LsrB-dependent chemotaxis may therefore contribute to niche 264 segregation in competitive infections.

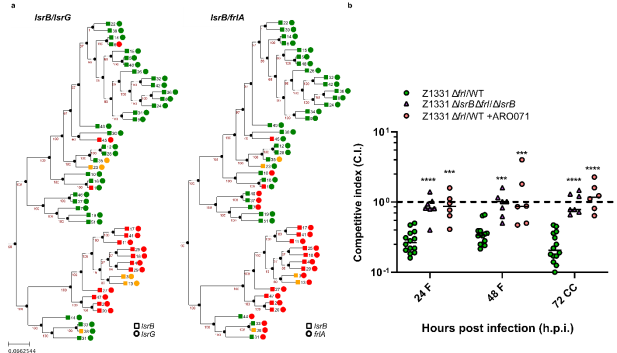
265 In contrast, the E. coli 8850 outnumbered E. coli Z1331 by 10-50 fold throughout the course of the experiment (Fig. 2f). Deletion of IsrB further decreased fitness of Z1331 to 100-266 1000 fold, and this fitness defect was alleviated in presence of the AI-2-overproducing E. coli 267 ARO071 (Fig. 2f and Extended Data Fig. 8e). As expected, no inflammation was observed 268 upon infection of mice with E. coli 8178 and 8850 (Extended Data Fig. 8a-c). Additionally, 269 colonization levels of E. coli 8850 remained unchanged in co-infection experiments with 270 ARO071, suggesting that the regain of fitness by E. coli Z1331 \Delta\lsrB was not due to the 271 competition between E. coli 8850 and ARO071 (Extended Data Fig. 8e). In agreement with 272 this observation, the AI-2-deficient mutant of ARO071 was incapable of rescuing the E. coli 273 274 Z1331  $\Delta$ /srB phenotype (Fig. 2f). These findings suggest that the differential ability to sense 275 and chemotactically respond to AI-2 can contribute to niche segregation of *lsr*-expressing and non-expressing *E. coli* strains in the gut. 276

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278 E. coli Z1331 benefits from fructoselysine utilization in a LsrB-dependent 279 manner. Although degradation of AI-2 in E. coli cells yields acetyl-CoA and dihydroxyacetone 280 phosphate, which in turn can be fed into glycolysis and the citric acid cycle<sup>47</sup>, *E. coli* strains show generally poor or no growth with AI-2 as a sole carbon source<sup>47,48</sup>. Although the observed 281 aggregation of *E. coli* Z1331 cells in the gut might be per se beneficial<sup>14,49</sup>, we aimed at further 282 deciphering how E. coli benefits from AI-2 chemotaxis in vivo. We reasoned that Isr positive E. 283 284 coli strains, by reaching their respective ecological niche, might benefit from utilizing niche-285 specific nutrients. Therefore, we reasoned that the lsr operon may show a pattern of cooccurrence with genes responsible for the utilization of niche-specific nutrients when 286 comparing E. coli genomes. To explore this, 10146 E. coli and Shigella genomes and their 287 genes were downloaded from a high-quality genome collection<sup>50</sup>. In order to identify known 288 genes and pathways that may be connected to the *lsr* operon, this collection was reprocessed 289 290 to obtain gene frequencies of all genes with a known function in 47 different E. coli and Shigella 291 lineages. The analysis resulted in a list of 168 genes whose presence correlated with IsrB (Pearson correlation coefficient ≥0.5; Supplementary Table 2). These included structural,
 metabolic as well as regulatory genes and operons. Interestingly, among these correlated
 genes were those belonging to the *frl* operon, which is required for fructoselysine utilization<sup>51</sup>.
 Fructoselysine, an Amadori product of the non-enzymatic reaction of glucose with primary
 amines, is highly abundant in thermally processed foods including mouse chow<sup>52,53</sup>.

We compared the phylogenetic relatedness, measured by extracting the tree branch 297 298 length from a maximum likelihood phylogenetic tree of representative genomes, as well as the 299 functional similarity, based on the fraction of shared annotated genes with a known function, 300 of all 47 lineages in an all-against-all manner. In addition to the previously observed correlation 301 of IsrB and frIA genes, we could thereby show that lineages which possess both IsrB and frIA 302 are closely related to one another, both in evolutionary distance as well as functional similarity. When comparing these lineages to others which encode neither of the two genes, the opposite 303 trend was apparent, e.g., they were considerably more dissimilar in the phylogenetic 304 relatedness and functional similarity (Fig. 3a, Extended Data Fig. 9). Consistent with a role in 305 306 gut colonization by particular E. coli strains, mutations in the frl operon repressor FrIR were detected in long-term colonization experiments with an *frl*-positive *E. coli* strain<sup>54,55</sup>. 307 Accordingly, the deletion of the frl operon attenuated E. coli Z1331 gut luminal growth in 308 309 competitive infection experiments (Fig. 3b). In order to assess if fructoselysine utilization requires LsrB-dependent chemotaxis, we created an equivalent pair of isogenic mutants in a 310 *IsrB*-deficient background. In competitive infections, *E. coli* Z1331  $\Delta$ *IsrB*  $\Delta$ *frI* colonized the gut 311 312 as well as *E. coli* Z1331 ∆*lsrB* (C.I.≈1; Fig. 3b). Moreover, increasing the luminal AI-2 levels using E. coli ARO071 abolished the fitness advantage of the WT E. coli Z1331 over E. coli 313 314 Z1331  $\Delta frl$ , suggesting a direct connection between AI-2 chemotaxis and fructoselysine 315 utilization (C.I.≈1; Fig. 3b). We therefore conclude that *E. coli* Z1331 requires *lsrB* in order to 316 benefit from fructoselysine. We further selected and tested several other genes from the 317 Supplementary Table 2, but failed to identify those potentially connected to AI-2 chemotaxis (Supplementary Table 3). 318

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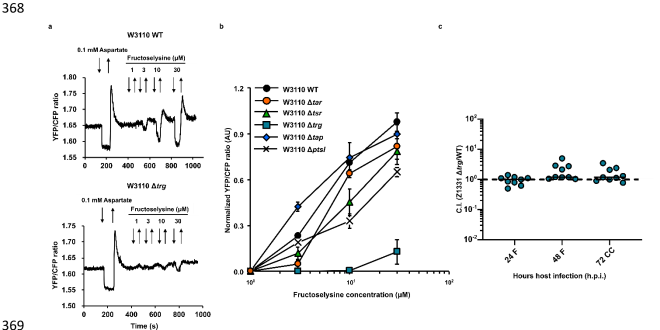
Figure 3. Co-occurrence analysis and competitive infection experiments demonstrating 322 323 a functional link between fructoselysine utilization and LsrB in E. coli Z1331. a, Maximum likelihood phylogenetic tree of representative genomes from the 47 investigated E. coli 324 lineages. We annotated with the gene frequencies of IsrB and IsrG (positive control) and for 325 326 frIA and IsrB genes. Bootstrap values are shown in red. A green dot indicates the respective gene is present in the majority of genomes belonging to the respective lineage (>95% of all 327 genomes possess the gene), orange means partly present (5%-95%) and red means it is not 328 present (<5%). b, C.I. values for *E. coli* Z1331 *Afrl* knockout during competitive infection 329 against the WT strain in the WT and the  $\Delta IsrB$  strain background in the absence or the 330 331 presence of the AI-2 overproducing *E. coli* strain ARO071. F, faeces, CC, caecal content. Lines 332 indicate median values (minimum n=6, from at least two independent experiments). P values were calculated using Mann-Whitney test (\*\*\*\**P*<0.0001; \*\*\**P*<0.0005). 333

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Fructoselysine is an attractant sensed by the chemoreceptor Trg. To further 335 investigate the interplay between AI-2 chemotaxis and fructoselysine metabolism, we analysed 336 337 the chemotactic response of E. coli to fructoselysine using a well-established Förster resonance energy transfer (FRET) assay<sup>56</sup>. This assay allows investigating the response of 338 the chemotaxis pathway to its ligands by monitoring the phosphorylation-dependent interaction 339 between fluorescent CheY-YFP and CheZ-CFP fusion proteins<sup>57,58</sup>. Since the chemotaxis 340 signalling pathway is highly conserved between different *E. coli* strains, the FRET assay could 341 be performed in the E. coli K-12 derivative strain W3110. As wild type E. coli isolates like those 342 employed in our study do often fail to express motility and chemotaxis genes under the FRET 343 assay conditions<sup>59</sup>, we have used *E. coli* W3110 for FRET experiments. In the wild-type *E. coli* 344

K12 cells, we observed an attractant response to 1-30 µM fructoselysine, as reflected by the 345 rapid drop in the YFP/CFP fluorescence ratio. (Fig. 4a, b, Extended Data Fig. 10a). E. coli 346 strains typically possess up to 5 different types of chemoreceptors responsible for sensing a 347 large repertoire of molecules<sup>1</sup>. To pinpoint the chemoreceptor involved in the chemotactic 348 response to fructoselysine, we performed FRET assays using E. coli K12 strains deleted for 349 each of the four receptors (Tar, Tsr, Trg and Tap) that mediate responses to chemical ligands. 350 351 The response to fructoselysine was severely reduced in  $\Delta trg$  knockout cells, suggesting that chemotaxis to fructoselysine is mediated by Trg (Fig. 4a, b, Extended Data Fig. 10b). In 352 353 contrast, a specific response to fructoselysine was retained in the knockouts of pts/ 354 (phosphotransferase enzyme of the phosphotransferase system which is involved in chemotactic response to some sugars<sup>60</sup>) and other receptor genes (Fig. 4a, Extended Data 355 Fig. 10c-q), including the AI-2-specific receptor Tsr and also dipeptide receptor Tap that 356 showed genome correlation with *lsr* operon (Extended Data Fig. 10e, Supplementary Table 2). 357

However, although Trg-mediated chemotaxis apparently represents the primary 358 mechanism of cell attraction towards fructoselysine, no effect of  $\Delta trg$  deletion in *E. coli* Z1331 359 was observed in *in vivo* experiments (Fig 4c). This might be due to the pleiotropic nature of the 360  $\Delta trg$  deletion, since Trg is known to mediate chemotactic responses to several sugars<sup>61</sup>, which 361 may have different effects on the cell fitness in the gut context. A similar situation was observed 362 for strain lacking Tsr, that mediates the response to AI-2 but also to a number of other stimuli, 363 including amino acids, pH and redox potential<sup>61</sup>. Despite sharing the same signalling pathway 364 365 of AI-2 sensing,  $\Delta IsrB$  and  $\Delta tsr$  strains show opposite competitive indexes in mouse experiments (Fig. 2b, Extended Data Fig. 11), again likely due to the impact of the  $\Delta tsr$  deletion 366 367 on responses other than AI-2. Further studies are needed to understand this complexity.



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370 Figure 4. Fructoselysine is an attractant sensed by Trg chemoreceptor. a, Examples of FRET measurements of *E. coli* W3110 wild-type and  $\Delta trg$  responses to fructoselysine. The 371 efficiency of FRET and thus the activity of chemotaxis pathway is reflected by the ratio of 372 373 YFP/CFP fluorescence. Stimulation with an attractant results in pathway inactivation and thus 374 decrease in the FRET ratio. Buffer-adapted cells were stimulated with step-like addition and 375 subsequent removal of compounds, indicated by downward and upward arrows, respectively. 376 Stimulation with a saturating concentration of strong attractant aspartate was used as a positive control. **b**, Dose-response curves of *E. coli* W3110 wild-type, chemoreceptor ( $\Delta tar$ , 377 378  $\Delta tsr$ ,  $\Delta trg$ ,  $\Delta tap$ ) and PTS ( $\Delta ptsl$ ) knockout strains to fructoselysine. Means of three 379 independent values of the pathway response upon step-like stimulation with indicated 380 concentrations of fructoselysine measured by FRET as in (a) and in Extended Data Fig. 10, normalized to the maximal response to control attractants aspartate or serine are shown. Direct 381 effect of fructoselysine on the YFP/CFP fluorescence ratio, measured in the negative control 382 strain, was subtracted. Error bars indicate s.d., c, Competitive infection experiment. C.I. values 383 for *E. coli* Z1331 Δ*trg* knockout during competitive infection against the WT strain. F, faeces, 384 CC, caecal content. Lines indicate median values (n=9, from at least two independent 385 386 experiments).

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Fructoselysine utilization represses Isr operon expression, resulting in 388 389 accumulation of extracellular AI-2. Since the chemotactic responses to fructoselysine and 390 AI-2 are directly related, we further aimed at understanding the connection between selfproduced AI-2 and fructoselysine metabolism. We hypothesized that fructoselysine uptake and 391 392 metabolism might affect *lsr* operon activity. Addition of 1% fructoselysine to exponentially 393 growing E. coli Z1331 cultures resulted in decreased Isr operon expression, and the observed 394 inhibition was independent of fructoselysine ( $\Delta frIA$ ) or AI-2 ( $\Delta IsrB$ ) import (Fig. 5a). 395 Interestingly, no repression of *lsr* operon activity was detected in a phosphotransferase system (PTS)-deficient *Aptsl* knockout strain, albeit at reduced background levels of *lsr* expression. *lsr* 396 397 operon expression is known to be regulated by catabolite repression, resulting in inhibition of its expression upon import of sugars through the PTS mediated by cAMP and its receptor 398 CRP<sup>62,63</sup>. This suggests that although the *frl* operon encodes a fructoselysine importer<sup>51</sup>, 399 fructoselysine might be partially imported via PTS or its uptake affects the activity of PTS 400 indirectly, similar to several other non-sugar carbon sources (Extended Data Fig. 12)<sup>60</sup>. Indeed, 401 de-repression of frl operon expression caused by FrIR inactivation upon addition of 402 fructoselysine<sup>64</sup> was only detected in  $\Delta ptsl$  background (Fig. 5b). This suggests a rather 403 unconventional double regulation of the *frl* operon activity by fructoselysine interactions with 404 405 PTS and FrIR. Consistent with CRP-mediated regulation of frl operon activity, we observed a decreased frIA promoter activity in presence of glucose, a known PTS substrate, and 406

fructoselysine in both WT and fructoselysine import-deficient  $\Delta frlA$  strains<sup>64</sup>. At this point, further studies are needed to clarify the mode of interaction between fructoselysine and PTS. Apart from CRP-mediated catabolite repression of the *frl* operon, direct inhibition of fructoselysine kinase FrIR by the unphosphorylated form of the PTS phosphocarrier protein HPr might as well be possible. This has been previously shown for the *lsr* operon, where unphosphorylated HPr inhibited AI-2 kinase LsrK<sup>63</sup>.

Upon its activation, the *lsr* operon induces rapid import and degradation of extracellular AI-2<sup>34</sup>. Decreased *lsr* operon activity as a response to fructoselysine metabolism might result in increased extracellular AI-2 levels. To test this hypothesis, we incubated *E. coli* Z1331 cells with or without 1% fructoselysine, followed by measurements of extracellular AI-2. As expected, fructoselysine-dependent inhibition of *lsr* operon activity resulted in elevated extracellular AI-2 levels (Fig. 5c). By generating more AI-2, fructoselysine-metabolizing *E. coli* might thus recruit additional cells to the source of fructoselysine in the gut.

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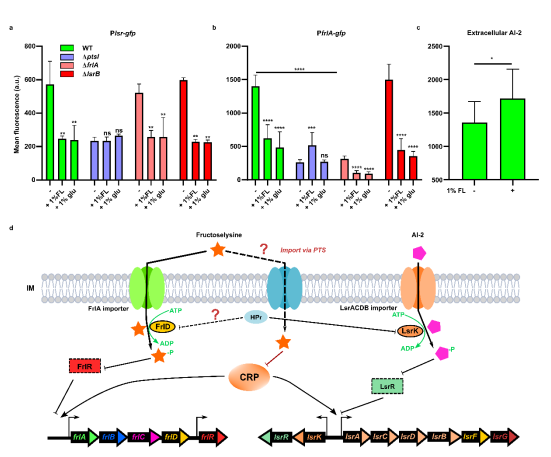


Figure 5. Effect of fructoselysine on *Isr* and *frl* operon expression in *E. coli* Z1331. a, *Isr* and b, *frl* operon activity in absence and presence of 1% fructoselysine (FL) in *E. coli* WT and its PTS-, fructoselysine and AI-2 import-deficient knockouts ( $\Delta ptsl$ ,  $\Delta frlA$  and  $\Delta IsrB$ , respectively). FL was added to exponentially growing *E. coli* cells (TB medium) containing

plasmid based Plsr-gfp or PfrIA-gfp fluorescent reporter. Glucose (glu), was used as a positive 428 control for PTS system effects. Fluorescence was measured 2 h after incubation with FL or 429 glucose with flow cytometry and expressed in arbitrary units (a.u.). Error bars indicate s.d. 430 431 (n=6, from at least two independent experiments). c, Levels of extracellular AI-2 produced by 432 E. coli cells quantified in supernatants using fluorescence reporter strains as described in 433 Methods. Reporter fluorescence was measured with flow cytometry and expressed in arbitrary 434 units (a.u.). Error bars indicate s.d. (minimum n=7, from at least two independent experiments). P values were calculated using the Mann-Whitney test \*\*P<0.005; \*P<0.05; ns, not significant). 435 d, Working model of FL- and AI-2-mediated regulation of *frl* and *lsr* operons, respectively. 436 437 Import of AI-2 via the LsrACDB importer results in de-repression of the Isr operon. Note that AI-2 internalization in *E. coli* is not solely dependent on the Lsr system<sup>34,65</sup>. Our data indicates 438 that FL appears to interact with a PTS system (dashed line, is potentially imported via an 439 unidentified PTS), leading to catabolite repression of both frl and lsr operons via CRP. In case 440 of the *frl* operon, FrIA-mediated FL import can slightly upregulate transcription, presumably by 441 alleviating FrIR-mediated repression (that is induction; as seen in the *ptsI* mutant). *frI* operon 442 repression by PTS might however also occur via direct inhibition of the fructoselysine kinase 443 FrID by the PTS phosphocarrier protein HPr (dashed line), as previously shown for the *lsr* 444 operon. Fructoselysine-dependent inhibition of *lsr* operon expression results in less AI-2 import 445 and therefore higher levels of extracellular AI-2, as seen in panel c. IM, inner membrane. 446 447 Question marks indicated potential interactions that are yet to be experimentally addressed.

448

## 449 Discussion

450 Chemotactic bacteria are found in various environments, ranging from the rhizosphere 451 and aquatic habitats to the mammalian gut. The role of chemotaxis in nutrient acquisition, 452 biofilm formation and host-microbe interactions has been clearly shown for several bacterial 453 species<sup>8</sup>. However, despite our detailed knowledge of its underlying molecular machinery, the importance of chemotactic behaviour in bacteria under physiologically relevant conditions 454 455 remains poorly studied<sup>9</sup>. E. coli, a common mammalian gut inhabitant, has been shown to benefit from chemotaxis on both individual and population levels. In a series of *in vitro* studies, 456 457 the role of chemotaxis in its foraging behaviour, expansion of the population range, autoaggregation and biofilm formation has been well documented<sup>13–15,66–68</sup>. Surprisingly, 458 although E. coli has been a preferred model for ex vivo chemotaxis studies for decades, the 459 460 question of how it might benefit from chemotaxis in vivo has not yet been fully addressed. In this study, we combined molecular and bioinformatics approaches to understand the role of E. 461 coli chemotaxis system during gut colonization. 462

By competing the non-chemotactic  $\Delta cheY$  knockout against the WT strain in ampicillinpretreated SPF mice, we could clearly show the fitness advantage of chemotaxis for *E. coli* 

Z1331 *in vivo*. Interestingly, although a similar phenotype was previously observed for  $\Delta cheY$ 465 mutants during S. Tm infection, this latter phenotype was strongly associated with gut 466 inflammation<sup>30</sup>. In contrast, no inflammation was observed during our colonization experiments 467 468 with E. coli, suggesting that even in closely related enteric organisms like E. coli and S. Tm, 469 chemotaxis can be adopted for different strategies of proliferation and survival in the gut environment. It is important to note that *E. coli* is as well capable of adapting another strategy 470 471 during gut colonization, namely inactivating motility and chemotaxis. Building and maintaining motility and chemotaxis machinery represents one of the most energetically costly behaviours 472 473 for the cell<sup>1,2</sup>, and mutations that inhibit flagella synthesis might prove beneficial under certain 474 conditions. However, our data suggests that chemotaxis indeed provides fitness advantages for motile *E. coli* strains across different phylogroups<sup>12,69–71</sup>. 475

We further show that the fitness advantage provided by chemotaxis is largely dependent on the response to the self-produced interspecies quorum sensing signal AI-2. Multiple roles of AI-2 signalling in collective behaviour of bacteria, phage-bacteria interactions and gut community structure have been reported<sup>16,21,48,72</sup>. Chemotaxis towards AI-2 promotes autoaggregation and biofilm formation in *E. coli* and biofilm dispersal in *Helicobacter pylori*<sup>14,15,73,74</sup>. Moreover, the chemotactic response to AI-2 is apparently not limited to AI-2 producing bacteria, suggesting its broad function in host-associated communities<sup>19</sup>.

In E. coli, AI-2 is indirectly sensed by the Tsr chemoreceptor via AI-2 binding LsrB 483 protein<sup>36</sup>. LsrB is in turn encoded by the AI-2-responsive *lsr* operon required for rapid signal 484 internalization and degradation<sup>16</sup>. The ability of an *E. coli* strain to chemotactically respond to 485 AI-2 is thus tightly linked to the presence of the *lsr* operon, which may be different in other 486 487 bacteria<sup>44</sup>. One of the competitive fitness advantages provided by AI-2 in the gut is related to 488 the niche segregation between E. coli strains that differ in their ability to chemotactically 489 respond to AI-2. Given that any gut ecosystem might contain several strains of the same 490 bacterial species that must stably coexist over extended period of time<sup>39–41</sup>, their segregation 491 due to different tactic preferences might generally facilitate such co-existence, either on its own 492 or in combination with different metabolic preferences or utilization of the same nutrients in distinct niches (niche segregation) as proposed by a niche theory<sup>41,75</sup>. The former case might 493 apply to AI-2, since it is not directly used by *E. coli* as a nutrient source<sup>47,48</sup>, thus expanding 494 495 the nutrient niche theory to molecules with no apparent nutritional value.

The inability of *E. coli* to use AI-2 as sole nutrient, however, poses a question of how exactly *E. coli* benefits from AI-2 chemotaxis during gut colonization. To address this question, we analysed 10146 *E. coli* and *Shigella* genomes to find genes correlated with the presence of the *Isr* operon. We found some level of correlation between *Isr* and the *frl* operon, which is required for fructoselysine utilization. This is of particular interest, since fructoselysine is widely found in thermally processed foods<sup>53</sup>. Indeed, we could show that *E. coli* cells benefit from

fructoselysine utilization in a AI-2 chemotaxis-dependent manner. Intriguingly, although our 502 experiments revealed that fructoselysine could itself act as an attractant sensed via the 503 504 chemoreceptor Trg, its primary mode of signalling might rather rely on the regulation of local 505 AI-2 levels in the gut, mediated by the inhibitory effect of fructoselysine on lsr operon 506 expression and therefore leading to increased extracellular AI-2 levels. We hypothesize that fructoselysine chemotaxis and metabolism, by increasing AI-2 levels surrounding 507 508 fructoselysine-utilizing cells, attracts AI-2 chemotactic E. coli to the source of fructoselysine. Consistently, a positive loop in AI-2-mediated cell recruitment was previously reported for 509 510 growing aggregates of *E. coli* cells *in vitro*<sup>14</sup>. Furthermore, it has been proposed that such 511 integration of two independent signalling pathways enhances chemotaxis of E. coli cells 512 towards the nutrient source<sup>76</sup>.

AI-2 is produced and sensed by a wide range of bacterial species, with AI-2 mimics 513 being synthesized by eukaryotic cells<sup>16–18</sup>. Given the apparently ubiquitous presence of this 514 signal, it seems rather counter-intuitive that E. coli cells rely on it to occupy specific niche within 515 516 the gut. A possible explanation for that would be that AI-2 chemotaxis is only relevant under antibiotics- or inflammation-induced dysbiosis conditions, which are known to result in 517 microbiota suppression and expansion of Enterobacteriaceae<sup>77–80</sup>. In absence of other AI-2 518 producers, no interference with self-produced AI-2 would occur. Additionally, our analysis 519 shows that E. coli lineages that possess both Isr and frl operons show high levels of similarity 520 521 to each other both in terms of their phylogenetic relatedness as well as in their functional 522 potential, ensuring that closely related E. coli have higher chances of co-accumulation at the fructoselysine source than more distantly related strains. Collectively, our study provides first 523 524 evidence for a causal link between AI-2 chemotaxis, gut colonization and niche segregation of 525 E. coli strains in the mammalian gut. It further establishes a link between AI-2 chemotaxis and 526 fructoselysine metabolism in vivo. We suggest that similar mechanisms of AI-2-mediated host 527 colonization might exist in other chemotactic bacteria.

528

## 529 Materials and Methods

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## Bacterial strains and growth conditions

The strains and plasmids used in this study are listed in Supplementary table 1. *E. coli* cells were routinely grown either on 1.5% Lysogeny Broth (LB) agar or in liquid LB or tryptone broth (TB) medium (10 g tryptone, 5 g NaCl per litre) supplemented with ampicillin (100  $\mu$ g/ml), kanamycin (50  $\mu$ g/ml), streptomycin (50  $\mu$ g/ml) or chloramphenicol (35  $\mu$ g/ml), where necessary. For fructoselysine utilization experiments, M9 minimal medium (6.7 g Na<sub>2</sub>HPO<sub>4</sub>, 3 g KH<sub>2</sub>PO<sub>4</sub>, 0.5 g NaCl, 1 g NH<sub>4</sub>Cl per litre, 2 mM MgSO<sub>4</sub>, 0.1 mM CaCl<sub>2</sub>) was used. Gene deletions were obtained via PCR-based inactivation<sup>81</sup>, and Km<sup>r</sup> cassettes were eliminated via
 FLP recombination<sup>82</sup>.

540

#### 541 Animals

542 C57BL/6 (JAX:000664, The Jackson Laboratory) mice were held under specific 543 pathogen-free (SPF) conditions at the EPIC facility at ETH Zürich. Germ-free (GF) mice were 544 bred in flexible film isolators at the isolator facility (EPIC, ETH Zürich). All animal experiments 545 were reviewed and approved by Kantonales Veterinäramt Zürich under license ZH158/2019, 546 complying with the cantonal and Swiss legislation. 8-12 week old mice of both sexes were 547 randomly assigned to experimental groups.

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## 549

## Mouse infection experiments

8-12 week old SPF mice were orally pretreated with ampicillin (20 mg) 24 h prior 550 infection. No pretreatment was required prior to infection of GF mice. E. coli cultures were 551 grown overnight in TB at 37 °C with shaking, diluted 1:100 in fresh TB and incubated at 37 °C 552 with shaking to the OD<sub>600</sub>=0.5-0.6 was reached. The cells were then washed in PBS (137 mM 553 NaCl, 2.7 mM KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, 1.8 mM KH<sub>2</sub>PO<sub>4</sub>), and mice were orally gavaged with 554  $5x10^7$  c.f.u. in 50 µl of single culture or a 1:1 mixture of strains, unless specified otherwise. 555 Faecal samples were collected every 24 h, unless stated otherwise, and animals were 556 557 sacrificed at 72 or 96 h.p.i. by CO<sub>2</sub> asphyxiation. Fresh faecal pellets and intestinal contents 558 were harvested and suspended in 500 µI PBS, followed by homogenization in a Tissue Lyser (Qiagen). Bacteria were plated on MacConkey (Oxoid) or LB agar plates with appropriate 559 560 antibiotics.

561 Competitive index (C.I.) of a wild-type strain (WT) and respective knockouts (KO) in a 562 competitive infection was determined as a ratio between c.f.u. (KO) and c.f.u. (WT) divided by the ratio of both strains in the inoculum. In most cases, competitive infection assays can 563 resolve gut lumen colonization phenotypes of Enterobacteriaceae much better than a 564 565 comparison between data from animals that are infected with either the wild-type or the mutant strain alone. This is likely attributable to the fact that these Enterobacteriaceae strains 566 567 generally do not reach their fastest possible growth rate in the gut lumen, at least in the 568 absence of massive bacteriotoxic events (like massive gut inflammation). Thus, slightly reduced efficiency of one mechanism to access a particular fraction of the nutrients can be 569 compensated by utilizing alternative, even if less efficient, mechanisms. As a result, 570 competitive infection assays in the non-inflamed gut are not accompanied by a significant 571 change in the overall E. coli density, while competitions between a wild-type strain and an 572 573 attenuated mutant would still yield a significant competitive index.

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# 575 Immunofluorescence microscopy and image analysis

SPF ampicillin-pretreated mice were infected with 5x10<sup>7</sup> c.f.u. (single infection or 1:1 576 mix) of *E. coli* Z1331 WT and Δ*che* Y carrying pFPV25.5 (mCherry under control of constitutive 577 578 rpsM promoter) and pFPV25.1 (GFP under control of constitutive rpsM promoter) plasmids, respectively<sup>83,84</sup>. At 72 h.p.i., mice were sacrificed by CO<sub>2</sub> asphyxiation. Caecal contents were 579 collected and plated to confirm the expected C.I. ( $\Delta che Y/WT$ ). Small intestine (lleum), caecal 580 581 and proximal colon tissue was excised, fixed in 4% paraformaldehyde (w/v in PBS) for 48 h at 4 °C, followed by 4 h in 20% sucrose solution (w/v in PBS) at 4 °C. The samples were then 582 583 embedded and frozen in Tissue-Tek OCT medium (Sysmex). 10 µm sections of the resulting 584 cryoblocks were cut and stained using DAPI and Cy5-Phalloidin. Images of fixed tissues were 585 visualized using a confocal Zeiss Axiovert 200 m microscope equipped with two evolve 512 EMCCD cameras (Photometrics) and a 40x oil objective. Images were analysed using 586 Particles Analysis Tool (ImageJ, https://imagej.nih.gov/) to determine the number of 587 aggregated formed by WT and  $\Delta cheY$  cells. Aggregates were defined as objects with the size 588 at least 50 px<sup>2</sup>, with single cells being  $\sim$ 10 px<sup>2</sup>. The number of aggregates formed by WT and 589 ΔcheY cells was then normalized to the total amount of detected particles in mCherry and GFP 590 channels. Fluorescent particles of non-bacterial origin (i.e., food particles) were manually 591 592 excluded from the analysis.

593

## 594 Flow cytometry

Activity of the *lsr* promoter was analyzed *in vivo* in SPF ampicillin-pretreated mice using a plasmid-based P*lsr-gfp* reporter<sup>14</sup>. Fresh faecal pellets were collected at 8, 24 and 48 h.p.i., incubated for 1 h at room temperature with 2  $\mu$ g/ml chloramphenicol to inhibit protein synthesis and to allow GFP proteins to fully maturate. Maximum of 20% reporter plasmid loss was observed during the experiment as determined by differential plating of faeces. Fluorescence was measured with Cytoflex flow cytometer (Beckman Coulter). *In vitro* culture of *E. coli* cells was used as a control for a forward and side scatter gate to exclude debris.

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# Extracellular AI-2 measurements

Extracellular AI-2 in faeces was measured by the same plasmid-based P*lsr-gfp* reporter introduced into *E. coli* W3110  $\Delta luxS$  strain<sup>14</sup>. Faecal suspensions were centrifuged for 10 min at 14 000 r. p.m. 20 µl aliquots of the reporter strain (OD<sub>600</sub>=0.3-0.5) were added to debris-free supernatants. Fluorescence of the reporter was measured after 1 h of incubation at 37 °C with shaking.

To measure AI-2 levels in *E. coli* supernatants treated with fructoselysine, exponentially growing *E. coli* Z1331 cells were incubated for 2 h with or without 1% fructoselysine. The cells

- 611 were subsequently washed twice in PBS and incubated for additional hour in fresh TB to allow
- for AI-2 production. AI-2 levels in cell-free supernatants were measured as described above.
- 613

## 614 Lipocalin-2 ELISA

- Lipocalin-2 was measured in faeces and caecal contents homogenized in 500 μl PBS
   by ELISA DuoSet Lipocalin ELISA kit (DY1857, R&D Systems).
- 617

618 Histopathology

619 Caecal tissue samples were embedded and frozen in Tissue-Tek OCT medium 620 (Sysmex). Cryosections (10  $\mu$ m) were stained with haematoxylin and eosin (H&E). 621 Pathological analysis (submucosal edema, goblet cells numbers, epithelial integrity and 622 polymorphonuclear granulocytes infiltration into the lamina propria) was performed as 623 described previously<sup>85</sup>.

624

## 625 Fructoselysine synthesis

626 Fructoselysine synthesis was performed as previously described<sup>86</sup>.

627

# 628 FRET measurements

The FRET measurements were performed as previously described<sup>57,58</sup>. Briefly, *E. coli* 629 630 cells were grown in TB supplemented with the antibiotics (100 mg/mL ampicillin) and inducer (50 mM IPTG) at 34 °C and 275 r.p.m. Cells were harvested at OD<sub>600</sub>=0.6 by centrifugation 631 (4000 r.p.m. for 5 min), washed twice with tethering buffer (10 mM KPO<sub>4</sub>, 0.1 mM EDTA, 1 µM 632 633 methionine, 10 mM lactic acid, pH 7), and stored at 4 °C for 30 min. The sample was attached 634 to a polylysine-coated coverslip, placed in a flow chamber under constant flow (300 µl/min) of 635 tethering buffer using a syringe pump (Harvard Apparatus, Massachusetts, United States), which was used for stimulation with compounds of interest. Measurements were performed on 636 an upright fluorescence microscope (Zeiss AxioImager.Z1) equipped with photon counters 637 638 (Hamamatsu) connected to a computer with custom written LabView7 software (National Instruments). CFP fluorescence was excited at 436/20 nm through a 455 nm dichroic mirror by 639 640 a 75 W Xenon lamp. To detect CFP and YFP emissions, 480/40 nm band pass and 520 nm 641 long pass emission filters were used, respectively. Fluorescence of a monolayer of 300-500 cells was continuously recorded in the cyan and yellow channels using photon counters with a 642 643 1.0 s integration time.

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## 645 Bioinformatic analysis

The genome collection provided by Horesh *et al.* was used as a base for our bioinformatic analysis<sup>50</sup>. Briefly summarizing their work, the authors downloaded 18156 *E. coli*  and *Shigella* genomes from human hosts and extensively curated them, resulting in a highquality dataset of 10146 genomes. Coding sequences were identified with Prodigal<sup>87</sup> and annotated with PROKKA<sup>88</sup> and genomes were assigned to lineages with a k-mer based wholegenome comparison approach using popPUNK<sup>89</sup>.

652 We downloaded the supplementary files from the publicly available collection 653 (<u>https://microbiology.figshare.com/articles/dataset/A\_comprehensive\_and\_high-</u>

guality\_collection\_of\_E\_coli\_genomes\_and\_their\_genes/13270073, 654 assessed on 16.08.2021) and used the Supplementary file F4 to obtain gene annotations and lineage 655 656 assignments for each genome. Selected annotations of interest, such as the Isr operon and 657 associated genes, were manually verified by blasting the representative sequence with 658 blastx<sup>90</sup>. We excluded the gene family "IsrB\_1", which mapped best to "LacI family transcriptional regulator [Escherichia coli]", hence most likely not corresponding to a functional 659 660 IsrB protein. Some gene clusters encoding the same function were split up in the original analysis, for example *frIA\_1* and *frIA\_2*. We used the short names to merge gene clusters with 661 the same known function in every genome with a custom script in Python 3.7.6., generating a 662 663 presence/absence matrix resulting in a total amount of 3964 annotated genes (https://github.com/lukasmalfi/E Coli). This matrix was combined with the lineage assignment 664 of each genome to compute gene frequencies (percentage of genomes within one lineage in 665 which the respective gene is present) for all lineages containing >20 genomes, dividing the 666 667 sum of genomes encoding a respective gene by the total amount of genomes. This process 668 was repeated for each gene and lineage, resulting in a gene frequency table of all 47 lineages 669 and 3964 genes.

Pearson correlation values of the *IsrB* frequencies against all other gene frequencies were computed from the gene frequency table using the DataFrame.corr function of pandas  $1.0.3^{91}$ . The resulting coefficients were sorted in descending order and descriptions for gene short names were obtained from the *E. coli* K12 MG1655 genome in the STRING Database Version 11.5<sup>92</sup>.

675 The maximum likelihood phylogenetic tree "tree 50.nwk", based on single nucleotide polymorphisms in the core-gene alignment of one representative genome per lineage, was 676 677 obtained from the previously mentioned data collection and edited using the ete3 toolkit version 678 3.1.2<sup>93</sup>. The representative genomes were chosen in the genome collection by Horesh et al., 679 with Treemer v 3.0 to ensure a sub-sample that is representative of the original diversity<sup>94</sup>. We subsequently removed the three lineages (21,43,49) that contained less than 20 genomes from 680 the tree and colored the nodes of the remaining lineages according to the gene frequencies of 681 both *IsrB* and *frIA*, as well as *IsrB* and *IsrG* to serve as a positive control. 682

To compare the similarity of lineages containing both *IsrB* and *frIA* in comparison to lineages that possess only one or none of the genes of interest, we used two parameters, their

evolutionary distance and their functional similarity. When comparing two lineages, the tree 685 branch length extracted from the phylogenetic tree with the get\_distance function of the ete3 686 687 toolkit version 3.1.2 was used as a proxy for their evolutionary distance. To obtain a value for 688 the functional similarity of two lineages, the fraction of shared annotated gene frequencies was 689 calculated. For each gene with a known function, the lower frequency was identified, and the 690 sum of all lower frequencies was divided by the sum of all higher frequencies, resulting in a 691 value ranging from 1 (all gene frequencies are exactly the same in both lineages) to 0 (none of the genes are present in both lineages). This was done using the NumPy package version 692 693 1.18.1<sup>95</sup>.

All lineages (n=47) were assigned into one of three groups: Containing both IsrB as well as frIA (gene frequency of both genes >0.1, n=24), containing neither of the two genes (gene frequency of both genes <0.1, n=13) or containing only one of the two genes (n=10). All lineages were then compared with one another and a scatter plot illustrating the comparison of functional and genomic relatedness of all lineages, colored according to their assignment to the three groups, was generated with bokeh version 2.2.3 (<u>https://bokeh.org/</u>).

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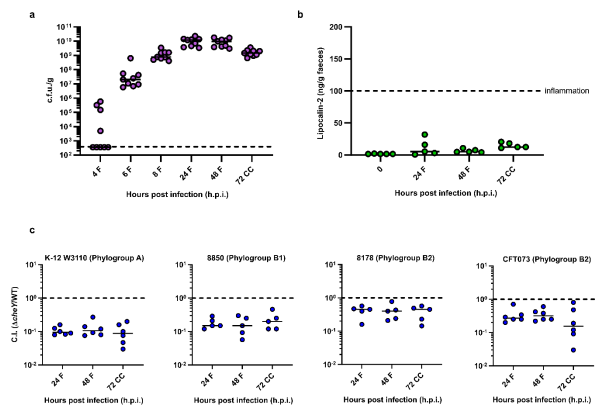
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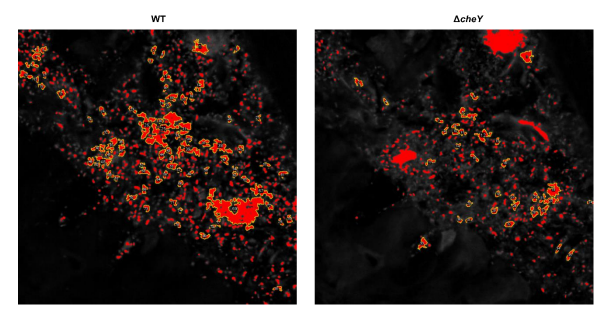
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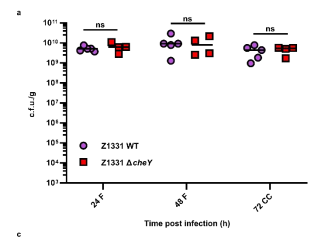
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962	Contributions
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964	performed the experiments, L.M. and C. v. M. performed bioinformatic analysis, C.L.D., L.F.
965	and J.P. synthesized fructoselysine. All authors contributed to data analysis and writing of the
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969	Ethics declarations
970	Competing interests
971	The authors declare no competing interests



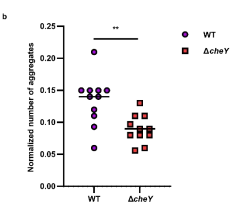
**Extended Data Figure 1.** *E. coli* **Z1331 colonizes ampicillin-pretreated SPF mice without causing inflammation. a**, c.f.u. of *E. coli* Z1331 WT (*yidX-bla*, amp<sup>1</sup>) detected in faeces (F) and caecal content (CC) of ampicillin-pretreated SPF mice at different time points of a 72 h infection. Lines indicate median values (n=9, from ≥2 independent animal experiments). The slight drop of fecal *E. coli* densities between 48 h and 72 h.p.i. is likely due to the regrowth of microbiota. The dashed line indicates the detection limit. **b**, Lipocalin-2 levels in faeces (F) and caecal content (CC) of *E. coli*-infected mice as measured by ELISA. Lines represent median values (n=5, from ≥2 independent animal experiments). Dashed line indicates approximate threshold of lipocalin-2 concentration marking a shift from non-inflamed to the inflamed gut, as observed in the streptomycin mouse model for *Salmonella* diarrhea<sup>1,2</sup>. Note that gut colonization by wild type S. Typhimurium yields lipocalin-2 levels of 10<sup>4</sup> ng/g faeces during fullblown gut inflammation<sup>1</sup>. **c**, Competitive indices (C.I.) for chemotaxis-deficient  $\Delta cheY$  strains from different phylogroups in competition against the respective WT strains in SPF ampicillinpretreated mice. F, faeces. CC, caecal content. Lines indicate median values (min n=5, at least two independent replicates).

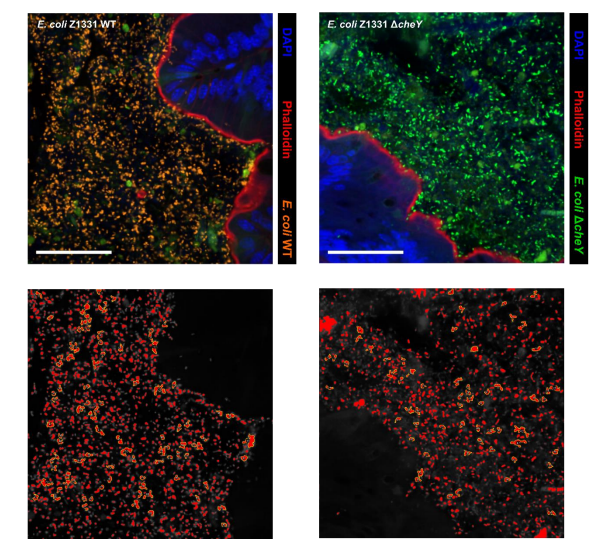


Extended Data Figure 2. An example of image segmentation and analysis of bacterial aggregates (as seen in Fig. 1d) using ImageJ. Detected particles are indicated in red, with aggregates (at least 50 px<sup>2</sup> in size) outlined in yellow. Particles of non-bacterial origin (food fibers etc, as seen in  $\Delta cheY$  panel) were manually excluded from analysis.

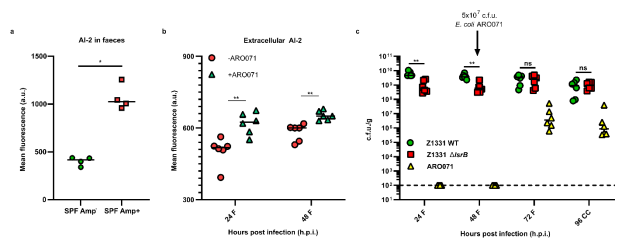


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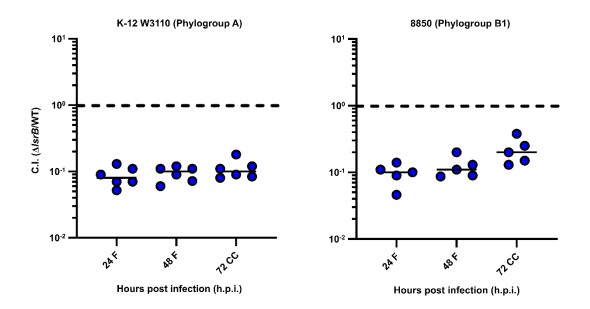




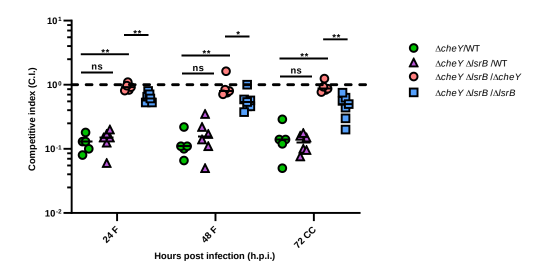
**Extended Data Figure 3.** *E. coli* Z1331  $\Delta cheY$  has no colonization defect in singlestrain infection. **a**, , c.f.u. of *E. coli* Z1331 WT and  $\Delta cheY$  detected in faeces (F) and caecal content (CC) of ampicillin-pretreated SPF mice at different time points of a 72 h infection. Lines indicate median values (n=4, 2 independent replicates). **b**, Number of aggregates formed by WT and  $\Delta cheY$  cells in a single-strain infection normalized to the number of detected cells in a tissue section as seen below (Mann-Whitney test, \*\**P*<0.005). Lines indicate median values (n=11, tissues sections from two independent experiments were analyzed). **c**, Caecal tissue sections of mice infected either with *E. coli* WT (mCherry-positive, shown in orange) or  $\Delta cheY$  (GFP-positive, shown in green) at 72 h.p.i. Actin filaments (red) and DNA (blue) were stained with phalloidin and DAPI, respectively. Scale bars, 50 µm. **d**, An example of image segmentation and analysis of bacterial aggregates (as seen above) using ImageJ. Detected particles are indicated in red, with aggregates (at least 50 px<sup>2</sup> in size) outlined in yellow. Particles of non-bacterial origin (food fibers etc, as seen in  $\Delta cheY$  panel) were manually excluded from analysis.



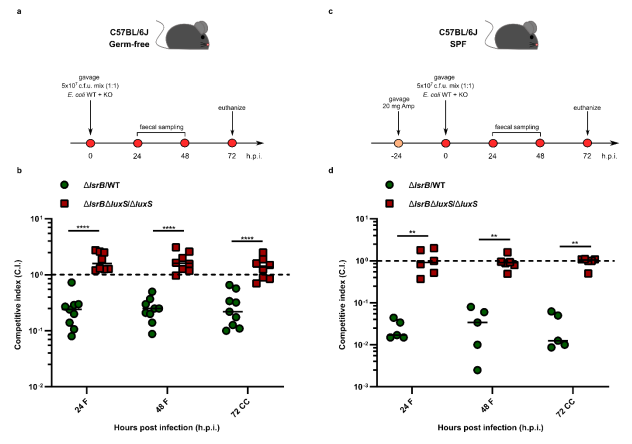
**Extended Data Figure 4. Increased luminal AI-2 levels abolish fitness advantage of wild-type** *E. coli* in  $\Delta$ *IsrB*/WT competitive infection. a, AI-2 levels of AI-2 in faeces of SPF mice before (SPF Amp<sup>-</sup>) and 24 h after (SPF Amp<sup>+</sup>) treatment with 20 mg ampicillin. Mean fluorescence of a plasmid-based AI-2 reporter strain was measured by flow cytometry and plotted in arbitrary units (a.u.). Lines indicate median values (n=4, at least two independent replicates). *P* values were calculated using the Mann-Whitney test (\**P*<0.05). **b**, AI-2 levels of AI-2 in faeces (F) of SPF ampicillin-pretreated mice infected with *E. coli* Z1331 WT (-ARO071) or with 1:1 mix of *E. coli* Z1331 WT and *E. coli* ARO071. Mean fluorescence of a plasmid-based AI-2 reporter strain was measured by flow cytometry and plotted in arbitrary units (a.u.). Lines indicate median values (n=6, at least two independent replicates). *P* values were calculated using the Mann-Whitney test (\*\**P*<0.005). **c**, c.f.u. data for the experiment shown in Fig. 2d. F, faeces, CC, caecal content. Lines indicate median values (n=6, at least two independent replicates). *P* values were calculated using Mann-Whitney test (\*\**P*<0.005; ns, not significant). The dashed line indicates the detection limit. Note that the total c.f.u. loads can differ between caecum and faeces due to yet unidentified reasons.



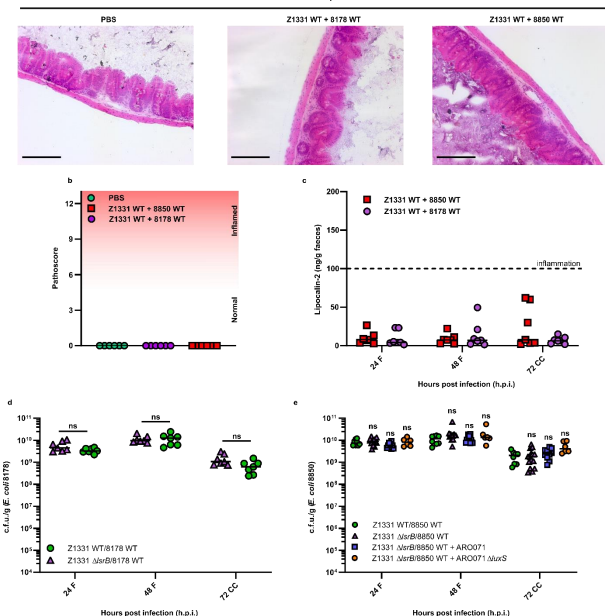
**Extended Data Figure 5.** Competitive indices (C.I.) for  $\Delta IsrB$  strains of *Isr*-positive *E. coli* W3110 and 8550 in competition against the respective WT strains in SPF ampicillinpretreated mice. F, faeces. CC, caecal content. Lines indicate median values (min n=5, at least two independent replicates).



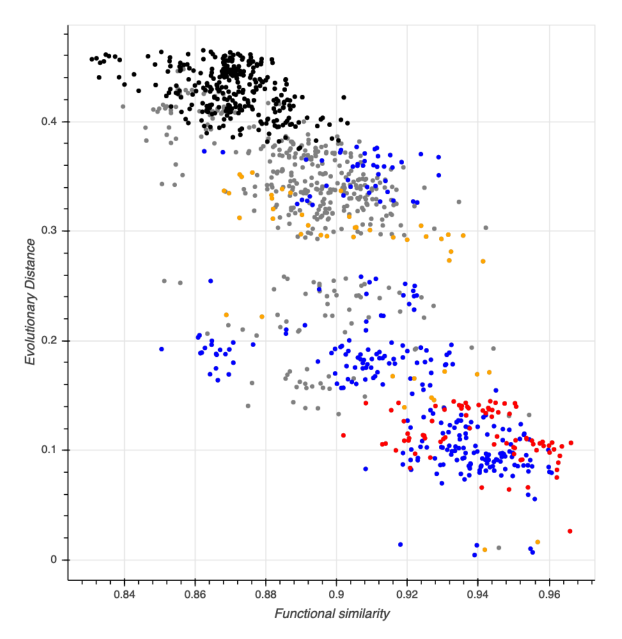
Extended Data Figure 6. CheY and LsrB belong to the same regulatory pathway. *E. coli* Z1331  $\Delta$ *cheY* and  $\Delta$ *cheY*  $\Delta$ *lsrB* knockout strains were competed against the wild-type strain. Additionally competitive indices (C.I.) of  $\Delta$ *lsrB* and  $\Delta$ *cheY* mutants were analyzed in  $\Delta$ *cheY* and  $\Delta$ *lsrB* backgrounds, respectively. F, faeces, CC, caecal content. Lines indicate median values (min n=5, from at least two independent infection experiments). *P* values were analyzed using the Mann-Whitney test (\*\**P*<0.005; \**P*<0.05; ns, not significant)



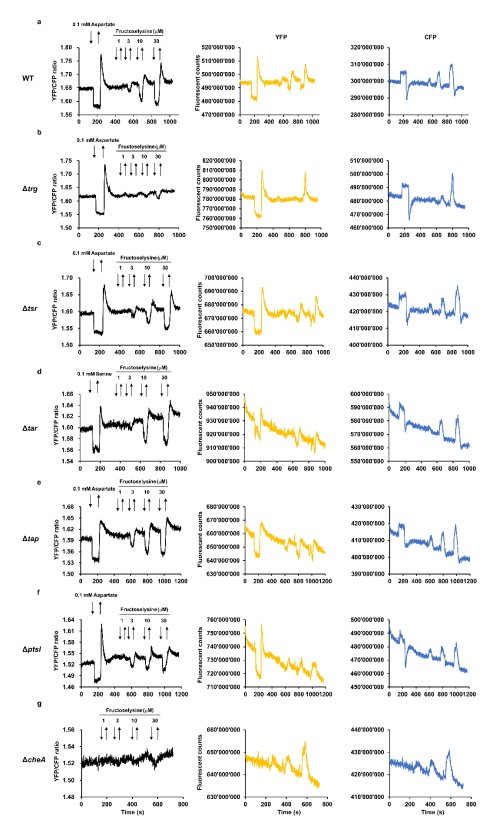
**Extended Data Figure 7. Self-produced Al-2 enhances gut colonization by** *E. coli.* **a**, Experimental scheme of competitive infection in germ-free (GF) mice. C57BL/6J GF mice were orally infected with  $5\times10^7$  c.f.u. *E. coli* W3110 WT and  $\Delta lsrB$  or  $\Delta lsrB \Delta luxS$  and  $\Delta luxS$  at a 1:1 ratio. Faeces were collected 24, 48 h.p.i. and mice were euthanized at 72 h.p.i. **b**, C.I. of non-Al-2 chemotactic  $\Delta lsrB$  mutant in WT and  $\Delta luxS$  background strains in the GF mouse infection model. F, faeces, CC, caecal content. Lines indicate median values (n=9, from least two independent experiments). *P* values were calculated using the Mann-Whitney test (\*\*\*\**P*<0.0001). **c**, Experimental scheme of competitive infection in SPF mice. C57BL/6J SPF mice were pretreated with 20 mg ampicillin by oral gavage 24 h prior to infection with *E. coli* W3110 WT and  $\Delta lsrB$  or  $\Delta lsrB \Delta luxS$  and  $\Delta luxS$  at 1:1 ratio. Faeces were collected at 24, 48 h.p.i and mice were euthanized at 72 h.p.i. **d**, C.I. of non-Al-2 chemotactic  $\Delta lsrB$  mutant in WT and  $\Delta luxS$  background strains in SPF ampicillin-pretreated mouse infection model. F, faeces, CC, caecal content. Lines indicate median values (min n=5, from at least two independent experiments). *P* values were calculated using the Mann-Whitney test (\*\**P*<0.005).



**Extended Data Figure 8. Infection of SPF ampicillin-pretreated mice with** *E. coli* 8178 and 8850 does not cause inflammation. a, H&E staining of caecal tissue of uninfected mice (PBS) and mice infected with 5x10<sup>7</sup> c.f.u. of *E. coli* Z1331 WT + 8178 WT and *E. coli* Z1331 WT + 8850 WT (1:1000 ratio) at 72 h.p.i (as seen in Fig. 4). Scale bar, 50 µm. b, Histopathology analysis of the caecal tissue section as seen above. 3 sections from 2 mice per group were analyzed. c, Lipocalin-2 levels in faeces (F) and caecal content (CC) of *E. coli*-infected mice as measured by ELISA. Lines represent median values (n=7, at least two independent animal experiments). Dashed line indicates approximate threshold of lipocalin-2 concentration marking a shift from non-inflamed to the inflamed gut. d, Colonization levels of *E. coli* 8178 in competition experiments with *E. coli* Z1331 as seen in Fig. 2. Lines indicate median values (n=7, at least two independent replicates). *P* values were calculated using the Mann-Whitney test (ns, not significant). e, Colonization levels of *E. coli* 8850 in competition experiments with *E. coli* 21331 as seen in Fig. 2. Lines indicate median values (min n=6, at least two independent replicates). *P* values were calculated using the Mann-Whitney test (ns, not significant). *P* values were calculated using the Mann-Whitney test (ns, not significant). *P* values were calculated using the Mann-Whitney test (ns, not significant). *P* values were calculated using the Mann-Whitney test (ns, not significant). *P* values were calculated using the Mann-Whitney test (ns, not significant). *P* values were calculated using the Mann-Whitney test (ns, not significant). *P* values were calculated using the Mann-Whitney test (ns, not significant).

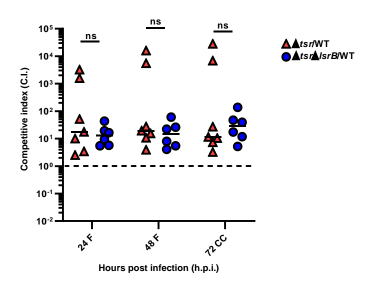


**Extended Data Figure 9.** *E. coli* genomes containing both *IsrB* and *frIA* are more closely related to each other than the average *E. coli* genomes. *E. coli* lineages were split up into three groups. Group 1 contains both *frIA* and *IsrB*, group 2 contains neither *frIA* nor *IsrB* and group 3 contains either *frIA* or *IsrB*. Phylogenetic distance (based on tree branch lengths, Y-axis) was plotted against the fraction of shared annotated genes (x-axis) of the lineages in an all-against all manner and colored according to their groups. Blue dots indicate group 1 compared with group 1, red dots indicate group 2 compared with group 2, orange dots indicate group 3 compared against group 3, black dots indicate group 1 compared with group 2, and grey dots indicate group 3 against group 1 and group 2, respectively.

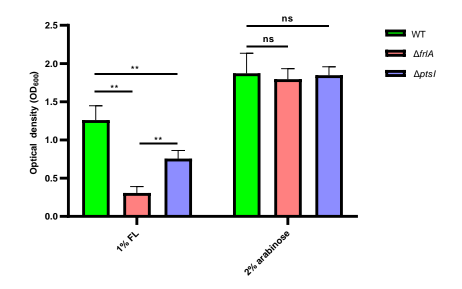


Extended Data Figure 10. Fructoselysine is an attractant sensed by the Trg chemoreceptor. Examples of FRET measurements of the response to fructoselysine by *E. coli* W3110 **a**, wild-type, **b**,  $\Delta trg$ , **c**,  $\Delta tsr$ , **d**,  $\Delta tar$ , **e**,  $\Delta tap$ , **f**,  $\Delta ptsI$  and **g**,  $\Delta cheA$  (negative control) knockout strains. Buffer-adapted cells were stimulated with step-like addition and removal of compounds (indicated by downward and upward arrows, respectively). Stimulation with saturating concentration of aspartate or serine, two strong attractants, was used as a positive control. Time traces of fluorescence intensity in the YFP and CFP channels are shown in the right. Opposite changes in two channels indicate specific FRET response. Note that

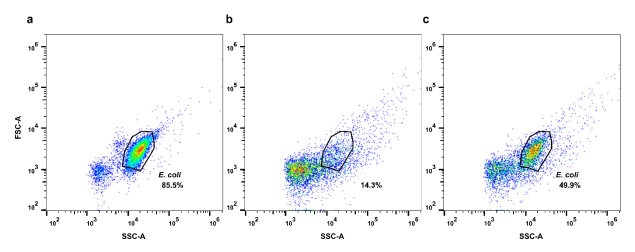
higher concentrations of fructoselysine solution have unspecific effect on fluorescence in both YFP and CFP channels, particularly visible in  $\Delta$ *cheA* negative control, but little effect on the YFP/CFP ratio. Residual effect on the YFP/CFP ratio in the negative control was subtracted from all dose-response curves in Figure 4b.



**Extended Data Figure 11. LsrB and Tsr belong to the same regulatory pathway.** Competitive indices (C.I.) of *E. coli*  $\Delta tsr$  and  $\Delta tsr \Delta lsrB$  mutant strains vs the wild-type strain *E. coli* Z1331 in SPF ampicillin-pretreated mice. F, faeces, CC, caecal content. Lines indicate median values (min n=6, at least two independent replicates). *P* values were analyzed using the Mann-Whitney test (ns, not significant).



# **Extended Data Figure 12.** *E. coli* Z1331 utilizes fructoselysine as a sole carbon source. *E. coli* Z1331 WT, $\Delta frlA$ and $\Delta ptsl$ strains were grown aerobically for 24 h in M9 minimal medium supplemented with either 1% fructoselysine (FL) or 2% arabinose (non-PTS sugar, used as a control for $\Delta ptsl$ growth) and NH<sub>4</sub>Cl as a nitrogen source. Error bars indicate s.d. (n=6, from at least two independent experiments). *P* values were calculated using the Mann-



**Extended Data Figure 13. Gating strategy for measuring E. coli gene expression in vitro and in vivo (as seen in Fig. 2a and Fig. 5a-c).** Representative plots of forward versus side scatter gating of **a**, *in vitro E. coli* culture grown in TB, **b**, faeces sample of uninfected SPF mice 24 h after ampicillin treatment (negative control), **c**, *E. coli* in faeces of ampicillinpretreated mice 24 h.p.i.

## **Supplementary References**

Whitney test (\*\*P<0.005; ns, not significant).

1. Maier, L. *et al.* Microbiota-derived hydrogen fuels Salmonella typhimurium invasion of the gut ecosystem. *Cell Host Microbe* **14**, 641–651 (2013).

2. Nguyen, B. D. *et al.* Import of Aspartate and Malate by DcuABC Drives H2/Fumarate Respiration to Promote Initial Salmonella Gut-Lumen Colonization in Mice. *Cell Host Microbe* **27**, 922-936.e6 (2020).