1 Inducible Cooperation in a Synthetic Gut Bacterial Consortium Introduces

2 **Population Balance and Stability**

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22 Abstract

Commensal microbes in the gut do not act alone but instead as cooperative consortia to conduct their myriad functions. Cooperative interactions and feedback mechanisms are key to consortia performance, yet are often ignored in current synthetic biology efforts to engineer the microbiota. To this end, we engineered mutual metabolic dependencies between four heterogeneous gut-dwelling bacterial species. Each species was made auxotrophic for three amino acids and an overproducer for one amino acid to share with the other species. By performing dynamical systems inference from time-series measurements, we show that our engineering introduced positive interactions that either reversed or neutralized pre-existing competitive interactions and improved stability of the consortium. We further demonstrate that we can induce population balance in the engineered consortia, both *in vitro* and in the mouse gut, through nutrient and dietary manipulations. Our findings indicate that induced cooperation can introduce evenness and stability in a synthetic microbial ecosystem, and have implications for development of synthetic approaches to manipulate the microbiome.

53 Introduction

54 In nature, microbes occur as conglomerates of various species with diverse sets of 55 genomes and metabolic capabilities, allowing for division of labor and increased 56 robustness (Hays et al., 2015a). For example, microbial consortia have been shown to 57 withstand external perturbations such as invasion of other species, toxic compounds 58 and nutrient sparseness (Burmølle et al., 2006; Lapara et al., 2002). A major driver for 59 microbial consortia robustness is cooperative behavior through production of public 60 goods and metabolic cross-feeding (Cavaliere et al., 2017). Consortia robustness is 61 correlated with population balance among the microbes and an extensive network of 62 interactions between species (Stelling et al., 2004; Stenuit and Agathos, 2015). 63

Metagenomic analyses reveal nutrient auxotrophies as a prevalent feature of microbial 64 65 communities, suggesting that cross-feeding might be a common mode of interaction in 66 natural consortia (Mee et al., 2014; Pande et al., 2014). Examples of such natural 67 microbial consortia include metabolically interacting communities in soil (Venail and 68 Vives, 2013) and in the mammalian gut (Rakoff-Nahoum et al., 2016). Distributing 69 metabolic capabilities over multiple species, a form of functional complementarity, can 70 increase productivity of the consortium through more efficient resource utilization 71 ((Pande et al., 2014; Savage et al., 2007).

72

73 Amino acid cross-feeding is an attractive means to introduce cooperation into synthetic 74 microbial consortia. These metabolites are more readily secreted than others, e.g., E. 75 coli secretes certain amino acids upon starvation (Burkovski et al., 1995; Kaderbhai et 76 al., 2003; Valle et al., 2008). Indeed, amino acids have been shown to play important roles in inter-species communication in natural systems (McCutcheon and Von Dohlen, 77 78 2011). For instance, amino acids are used by S. cerevisiae to regulate nitrogen 79 overflow, which leads to natural cross-feeding to lactic acid bacteria (Ponomarova et al., 80 2017). Numerous studies have engineered pairwise amino acid cross-feeding in E. coli 81 producing normal amino acid levels and generated quantitative models to describe their 82 behavior (Estrela and Gudelj, 2010; Kerner et al., 2012; Pande et al., 2014; Stolyar et al., 2007; Wintermute and Silver, 2010a). 83

84

85 Engineering cooperative microbial consortia has been of longstanding interest in 86 synthetic biology; studies were performed to gain basic scientific insights, or as 87 engineering proofs-of-principles (Mee et al., 2014; Wintermute and Silver, 2010b). With 88 recent advances in our understanding of the human microbiota, there is increasing 89 interest in applying synthetic biology approaches to construct a well-defined gut 90 microbiome, living bacterial diagnostics and therapeutics (Riglar and Silver, 2018). 91 Creating such diagnostics and treatments in the context of cooperative consortia has 92 numerous potential advantages, including the aforementioned capability of consortia to 93 carry out more complex functions in a more stable manner (Cavaliere et al., 2017), with 94 stability in this context defined as the ability to withstand external disturbances. 95 Additionally, consortia can potentially allow for greater safety and control. For instance, 96 a consortium that maintains population balance through cooperativity could be used to 97 control dosing of a therapeutic compound in the gut. 98 99 However, almost all prior synthetic biology studies that have engineered cooperativity in 100 bacterial communities have used a single species (Kong et al., 2018; Mee et al., 2014; 101 Wintermute and Silver, 2010b). However, natural microbial ecosystems contain a 102 diversity of interacting species. In advancing synthetic biology to real applications in 103 complex environments, it will be essential to expand engineering capabilities to diverse, 104 multi-species consortia. Importantly, bacteria from naturally occurring ecosystems are 105 likely to have pre-existing interactions, which are often competitive. These interactions 106 must be considered and often overcome to achieve cooperative consortia. 107

As a step toward developing multi-species consortia, we have constructed a synthetic consortium of four different bacterial species, each derived from the mammalian gut, and engineered mutual interactions by cross-feeding of four amino acids. Using several experimental approaches, combined with statistical inference from data and computational modeling, we demonstrate our ability to engineer cooperativity in the consortium that overcomes pre-existing competitive interactions. Further, we show that this cooperativity is inducible through nutrient or dietary manipulations, and that the

- 115 engineered consortium exhibits population balance that is stable when subjected to
- 116 perturbations *in vitro* and when introduced into the mammalian gut.
- 117

118 **Results**

119 Cooperative Consortia Design and Engineering

- 120 To gain intuition into synthetic consortia designs, we simulated behavior of non-
- 121 interacting collections of four bacterial species versus consortia linked by
- 122 positive/cooperative interactions (Figure 1A). Our simulations demonstrate that while a
- 123 collection of non-interacting bacteria can exhibit population balance, it is highly
- 124 susceptible to external disturbances that can drastically change the composition of the
- 125 community. In particular, disturbances can readily cause a species to die out in the
- 126 community. However, when we linked bacterial species through cooperative
- 127 interactions, the resulting consortia can withstand much higher levels of external
- 128 disturbances without dramatically altering its composition. Thus, our simulations
- 129 suggest that engineering a network of positive interactions within a bacterial consortium
- 130 could introduce stability towards environmental disturbances.
- 131

132 To construct our synthetic consortium, we selected four bacterial species, *Escherichia* 133 coli NGF-1, Salmonella enterica subsp enterica serovar Typhimurium LT2, Bacteroides 134 thetaiotaomicron VPI-5482, and Bacteroides fragilis 638R. These species are not only 135 genetically tractable, but also able to survive in the mammalian gut in diverse niches 136 and have varied abundances within the total microbiota. These characteristics allow us 137 to investigate key synthetic biology engineering principles in a controlled, but more 138 realistic context, and also maximize potential for downstream applications, for example 139 for bacterial therapeutics and diagnostics in the gut.

140

141 *E. coli* NGF-1 was isolated from BALB/c mice, has been shown to stably colonize the

- mouse gut, and can be engineered with standard genetic tools (Kotula et al., 2014;
- 143 Riglar et al., 2017). S. Typhimurium LT2 was further attenuated by removing the
- pathogenicity islands SPI1 and SPI2, and thus did not cause any disease phenotype
- 145 when administered to mice. The two Bacteroides species, *Bacteroides thetaiotaomicron*

and *Bacteroides fragilis,* are human commensals that can achieve high abundance in
 the mammalian gut, and are also genetically tractable.

148

149 We engineered each of the constituent species to depend on the other three by cross-

150 feeding of the four metabolites L-methionine, L-histidine, L-tryptophan and L-arginine

151 (hereafter referred to as Met, His, Trp and Arg) (Figure 1B). Auxotrophies for three of

- 152 these amino acids were generated in each strain (*E. coli*: His, Trp and Arg;
- 153 S. Typhimurium: Met, Trp, Arg; B. theta: Met, His, Arg; B. fragilis: Met, His, Trp), along
- 154 with the ability to overproduce one amino acid in each strain (*E. coli*: Met; *S.*

155 *Typhimurium*: His; *B. theta*: Trp; *B. fragilis*: Arg) (Table 1). *E. coli* and *S. Typhimurium*

156 were engineered by sequential phage transduction from three single auxotroph strains.

157 *E. coli* was transduced with genome fragments from BW25113 that contained insertions

158 in argA, trpC, hisA (see Methods) and S. *Typhimurium* with genome fragments of the

same parent strain with insertions in argA, trpC, metA. *Bacteroides spp*. triple knockout

- 160 generation utilized the pExchange-tdk vector to precisely delete metA, hisG and argF in
- 161 *B. theta* and *metA*, *hisG* and *trpC* in *B. fragilis*. To engineer overproduction of amino

acids, we selected for bacterial strains that showed resistance to specific

163 antimetabolites.

164

165 Characterization of Auxotrophies and Overproduction

166 To assess the auxotrophic strains' amino acid requirements, we measured growth on 167 varying concentrations of each metabolite in the presence of non-limiting concentrations 168 of all the other metabolites (Figure 2A). Each strain had a requirement for specific and 169 differing levels of the amino acids. Overproduction of metabolites was measured in 170 comparison to a defined amino acid standard using LC-MS (Figure 2A, horizontal bars). 171 In order to compare overproduction with each species' amino acid requirements, we fit a 172 sigmoidal curve to the growth response data (Figure 2A), to produce an expected 173 concentration (OD600) for the species for a given overproduction rate. This comparison 174 of requirements and overproduction levels provides information about the expected 175 relative strengths of the engineered interactions.

177 *B. fragilis* was the highest overproducer (Arg at 362 uM). Corresponding

- supplementation would allow growth of *E. coli* to OD 0.144, *S. Typhimurium* to OD
- 179 0.166 and *B. theta* to OD 0.154. *B. theta* overproduced Trp at 34 uM, allowing expected
- 180 growth for *E. coli* to OD 0.411, *S. Typhimurium* to OD 0.156 and *B. fragilis* to OD 0.128.
- 181 E. coli overproduced Met at 5.3 uM, allowing expected growth of S. Typhimurium to OD
- 182 0.032, *B. theta* to 0.017 and *B. fragilis* to OD 0.027. *S. Typhimurium* overproduces His
- at 16 uM, allowing for expected growth of E. coli to OD 0.028, but not supporting
- 184 *Bacteroides spp.* growth, which required concentrations higher than 100 uM.
- 185 Interestingly, the unengineered *Bacteroides spp*. also produced detectable amounts of
- 186 some amino acids, whereas the other wild-type species did not. In the case of *B. theta*,
- 187 the detected levels of Trp would, in principle, be high enough to support growth of other
- 188 consortium members. Overall, our findings suggest relatively strong engineered
- 189 cooperation from *Bacteroides spp* to other strains, moderate cooperation from *E. coli* to
- 190 other strains, and the weakest cooperation from *S. Typhimurium* to other strains.
- 191

192 Amino Acid Overproducers can Rescue Growth Defects of Corresponding

193 Auxotrophs

194 Having established amino acid requirements and overproduction levels for each strain, 195 we assessed pairwise cross-feeding using culture supernatants from the overproducing 196 and wild-type strains to test for growth of the corresponding auxotrophs (Figure 2b). 197 Cells were grown for 24 hr before supernatant was collected (Figure 2c). Notably, three 198 out of the four engineered species did not show any growth defect compared to the WT; 199 B. theta growth was decreased by 3-fold. Extent of rescue was determined by OD600 200 values after 24 hr of growth in supernatant that was diluted 1:1 with fresh media lacking 201 the tested amino acid (Figure 2d). As another comparator, we grew the auxotrophic 202 strains without amino acid supplementation and with full supplementation (1 mM of each 203 amino acid). As expected, we detected no growth in any of the auxotrophs when no 204 amino acid was supplied, and growth with full supplementation. 205

206 Consistent with our design, and our amino acid requirement and overproduction data, *E.* 207 *coli* grew well in supernatant from engineered *B. theta* (180% of fully supplemented growth) and *B. fragilis* (130%), and somewhat in supernatant from engineered *S.*

- 209 Typhimurium (13%). Interestingly, E. coli grew better in Bacteroides spp. supernatant
- than in fully supplemented media, suggesting that these *Bacteroides spp.* may produce
- other beneficial metabolites for *E. coli*. *S*. Typhimurium grew relatively well in *E. coli*
- supernatant (88%), and also showed enhanced growth in the Bacteroides spp.
- supernatants (330% in *B. theta* supernatant, and 227% in *B. fragilis* supernatant). As
- 214 expected from our requirement and overproduction data, *B. theta* is only marginally
- 215 rescued by engineered *E. coli* (3%). Overall, these experiments indicate cross-feeding
- 216 between the engineered bacterial strains.
- 217

218 However, in some cases the supernatants did not perform as well as predicted or in 219 comparison to co-cultures. B. fragilis rescued B. theta growth much less than expected 220 (12%) from our overproduction data. This finding suggests that *B. fragilis* may secrete 221 factors that inhibit B. theta growth, but not E. coli or S. Typhimurium. Indeed, 222 competitive interactions among more closely related species have previously been 223 reported, possibly due to competition for similar niches (Bauer et al., 2018). No growth 224 by B. theta in S. Typhimurium supernatant was evident. B. fragilis was not rescued well 225 by any of the strains (Figure 2d, blue panel). Taken together, these results may reflect 226 production of toxic compounds that are enriched in supernatants of grown cultures but 227 might play a lesser role in co-cultures.

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230 Cooperation and Population Balance of the Consortium *in vitro* is Inducible

- 231 Based on Amino Acid Abundance
- Having investigated pairwise interactions in our consortium, we next sought to
- 233 characterize properties of the entire consortium versus individual members. Using a
- 234 medium that we specifically designed to accommodate the four bacterial species in a
- single batch culture, and without amino acid supplementation, we grew monocultures
- and co-cultures of WT and the engineered consortia and estimated bacterial abundance
- 237 (cfu/mL) via qPCR after 24 hr (Figure 3A,B).
- 238

239 By comparing growth of strains in monoculture to the full consortium co-culture, we can 240 assess the degree of cooperativity in the bacterial community. We quantify this behavior 241 using a cooperation factor, defined as the total concentration of the co-culture divided 242 by the sum of the concentrations of the monocultures. According to this definition, a 243 cooperation factor <1 indicates competitive behavior, whereas a cooperation factor of 244 >1 indicates cooperation. For the WT consortium, each of the strains grew better in 245 monoculture than in co-culture (Figure 3B), with a cooperation factor of 0.14. This 246 finding suggests pre-existing negative interactions among the WT species, e.g., 247 competition for nutrients or production of compounds toxic to the other species. For the 248 engineered consortia, the cooperation factor is 1.18, indicating that our engineering 249 strategy has led to a net growth improvement when the complete consortium is able to 250 interact. Note that much of the growth improvement is due to *E. coli* and *S.* 251 Typhimurium growth, whereas B. theta and B. fragilis growth is essentially unchanged. 252 This suggests that our engineering introduced net positive interactions for *E. coli* and *S.* 253 Typhimurium, while neutralizing competitive effects on *B. theta* and *B. fragilis* present in 254 the WT consortium.

255

256 Since our engineering design is based on amino-acid cross-feeding, we hypothesized 257 that by varying the concentrations of amino acids in the medium, we could control the 258 degree of cooperativity among the consortium members. To test this hypothesis, we 259 subjected the engineered bacteria in monoculture and co-culture to different 260 concentrations of relative amino acid supplementation (Figure 3C). We measured 261 bacterial abundance (cfu/mL) via gPCR after 24 hr and calculated cooperation factors 262 for each condition. Overall, bacterial growth decreases for both monoculture and co-263 culture conditions with decreasing supplementation, as expected. Interestingly, we were 264 effectively able to ablate cooperativity with high amino acid supplementation; in this 265 regime, the engineered consortium behaves like the WT consortium, with monoculture 266 growth exceeding co-culture growth. As supplementation decreased, we found that the 267 cooperativity factor consistently increased, with the factor exceeding 1 at a 268 supplementation level of 3 uM. These findings are consistent with cross-feeding

behavior in naturally occurring microbial consortia, in which cooperativity only occursduring nutrient scarcity (Carlson et al., 2018).

271

272 Our simulation studies showed that our engineering design could result in a consortium 273 with a balanced population less susceptible to environmental changes. Thus, we were 274 interested in how varying amino acid supplementation would affect population balance. 275 To assess this, we measured relative abundances of each species in the engineered 276 consortium (Figure 3C, lower panel) via strain-specific gPCR. We quantitated population 277 balance using the normalized entropy measure (also called the evenness index) for the 278 consortium. With this measure, a completely even (balanced) community would have a 279 relative entropy of one.

280

We found that the highest population evenness occurred with the highest

supplementation (1000 uM.) This is likely because all the required amino acids are

supplied, which reduces competition that would lower population evenness. When

supplementation is decreased to intermediate levels (30 and 100 uM), the relative

abundances of bacteria become less even, i.e. entropy decreases. Specifically,

286 Bacteroides spp. abundance decreases dramatically, and E. coli and S. Typhimurium

dominate the culture. This level of supplementation represents a "mismatched" regime.

in which amino acid concentrations are high enough to support some of the species (S.

289 Typhimurium and E. coli, which have the lowest amino acid requirements), but not the

290 others (the *Bacteroides spp*., which have higher requirements.) When amino acid

supplementation is reduced further (particularly below 20 uM), the ecosystem enters a

low nutrient regime characterized by increased cooperativity as described above, and

293 population evenness increases again, almost to levels seen with the highest levels of

supplementation.

295

296 The Engineered Consortium Exhibits Greater Stability

297 Our amino acid supplementation experiments demonstrated that a microbial consortium

with low cooperativity can exhibit high population evenness, but can also be less stable.

299 One measure of stability is the extent to which a system tends to reach the same end-

300 point or steady-state, even if it starts in a different initial state. Indeed, this behavior is a 301 necessary condition for common microbial dynamical systems models to exhibit 302 asymptotic stability (Gibson et al., 2017). To assess the stability of our consortium 303 according to this criteria, we inoculated both WT and engineered consortia at five 304 different starting concentrations each (Figure 4). In condition 1, all bacterial species 305 were inoculated at the same ratios, and in conditions 2-5, we reduced one of the 306 species' inocula by a factor of 10. We then assessed growth of each strain in co-culture 307 over 12 hours (Figure 4A, B) via strain-specfic gPCR. Both consortia reached consistent total concentrations at 12 hrs (approximately 5 x 10⁷ cfu/mL, 1 x 10⁶ cfu/mL for 308 309 engineered.) However, the end-point abundances of the WT consortium members 310 differed markedly, depending on the starting condition. In particular, for conditions 2-4, 311 the low inoculum species remained low at 12 hrs. In contrast, the engineered 312 consortium exhibited significantly greater consistency (p-value 0.0355) in end-point 313 concentrations of the consortium members, regardless of the starting condition. These 314 results demonstrate increased stability, a dynamical systems property, of the 315 engineered consortium.

316

317 Computational Analysis of *in vitro* Growth Dynamics of the Engineered Consortia 318 over Time Elucidate a Net Positive Interaction Network

319 Our design created a mutually coupled bacterial consortium designed to function

- 320 together, suggesting that consortium behavior is best assessed in the full assemblage,
- 321 rather than through pairwise co-culture experiments. In order to investigate such
- 322 behavior, we analyzed the densely sampled time-series data from our experiments
- 323 using the entire consortium with different initial starting conditions (Figures 4A,4B;
- 324 described in the previous section) with a dynamical systems inference approach.
- 325
- 326 Our dynamical systems approach uses a tailored model based on stochastic
- 327 generalized Lotka-Volterra (gLV) dynamics and an associated fully Bayesian machine-
- 328 learning/statistical inference algorithm (Methods, Gibson and Gerber, 2018). Continuous
- 329 time stochastic generalized Lotka-Volterra (gLV) dynamics can be expressed as:

330
$$d\boldsymbol{x}_{t,i,\ell} = \left(\boldsymbol{r}_i \boldsymbol{x}_{t,i,\ell} + \sum_{j=1}^n \boldsymbol{a}_{i,j} \boldsymbol{x}_{t,i,\ell} \boldsymbol{x}_{t,j,\ell}\right) dt + d\boldsymbol{w}_{t,i,\ell}$$

where $x_{t,i,\ell}$ is the abundance of microbe *i* at time *t* in experiment ℓ . The parameter r_i denotes the growth rate of microbe *i*, and $a_{i,j}$ is the effect that microbe *j* has on microbe *i*. When i = j the expression $a_{i,i}$ is a self-limiting term and together with r_i determines the carrying capacity, $-\frac{r_i}{a_{i,i}}$, of microbe *i* if no other microbes were present. Finally, *w* is the process disturbance term, which we assume is a Brownian motion. For inference, we discretize the continuous dynamics as described fully in Methods.

337

338 We used our method to infer growth rates and microbe-microbe interaction strengths 339 from our time-series data (Figure 4E,F). As we described above, the WT consortium 340 achieves an overall higher concentration of approximately 5 x 10⁷ versus 1 x 10⁶ cfu/mL 341 for the engineered consortium. This results in different scales for the self-interaction and 342 interaction parameters across the two consortia, so we normalized the interaction 343 matrices by steady-state dynamics (see Methods) to render the two consortia 344 comparable. We see that the WT consortium has several strong aggregate negative 345 interactions, e.g., mutual negative interactions between *E. coli* and *S. Typhimurium*. In 346 contrast, the engineered consortium has aggregate neutral interactions, aside from one 347 strong positive interaction from *B. fragilis* to *S. Typhimurium*.

348

349 We can gain insight into the *net changes* in the guantitative structure of the synthetic 350 microbial interaction network introduced by engineering, by subtracting the normalized 351 WT network from the normalized engineered network, and keeping only interactions 352 deemed significant with our inference method in at least one network. Using this 353 analysis, we found 5 net positive and 7 net neutral interactions, confirming the ability of 354 our engineering approach to promote cooperation in the consortium. Interestingly, since 355 the WT interaction network shows strong competitive interactions, our model suggests 356 that our engineering approach mostly promotes cooperatively by significantly weakening 357 the naturally occurring competitive interactions.

359 The inferred network is generally consistent with our mono-culture and co-culture 360 results. For instance, the strongest positive interaction in the engineered consortium is 361 from *B. fragilis* to *S. Typhimurium*. Concordantly, *B. fragilis* overproduces the highest 362 amount of its crossfed metabolite Arg, and S. Typhimurium is the strain that benefits 363 most in co-culture. As another example, we did not infer any incoming positive 364 interactions for *B. fragilis*, which is consistent with our finding that *B. fragilis* does not 365 show improved growth in co-culture compared to monoculture. Our model inferred 366 mutual net positive interactions between E. coli to S. Typhimurium, which are consistent 367 with our supernatant complementation experiments.

368

369 Some aspects of the inferred model are inconsistent with the mono- and co-culture 370 experiments, however. For instance, these experiments suggest a positive interaction 371 from *B. fragilis* to *E. coli*, but this interaction does not appear in the inferred network. 372 Because our model is inferred from longitudinal data, which is relatively sparse and 373 noisy, it is not surprising that some interactions may not be detected. Further, our 374 approach is fully Bayesian and takes into account uncertainty in both the model and 375 measurements. This approach is by design conservative, meaning that it requires 376 strong evidence from the data to formally detect an interaction. Thus, our approach will 377 tend to report the strongest pre-existing or engineering induced interactions, and may 378 miss weaker but still present interactions. Although weak interactions were enhanced by 379 removing any amino acid supplementation in our experiments, these interactions may 380 still fall below our threshold of detection.

381

382 Simulations Based on Realistic Design Constraints Reveal Principles for Stability 383 in Metabolically Cooperative Microbial Consortia

We used information from our experiments and data-derived microbial interaction networks to systematically study consortia stability, and gain insight into general design principles. We present these simulations with increasingly realistic design constraints and demonstrate how such constraints lead to different cooperativity regimes. For simplicity of exposition, we assume the four species have identical growth rates ρ , selfinteraction terms δ (which is always negative), and identical interactions coefficients α 390 (Figure 5A). In our first, and least realistic design (Figure 5B), the growth rate and self-391 interaction terms are kept constant, and only the positive interaction strength is 392 increased. Under these constraints, the carrying capacity and stability of the system 393 increases as the interaction strengths increase, up to the point of system instability. 394 However, such a scenario is unrealistic, because on theoretical grounds, it would allow 395 for the overall carrying capacity of the ecosystem to be arbitrarily increased. Moreover, 396 it is inconsistent with our experimental evidence and data-derived model, which both 397 show a lower overall carrying capacity for the engineered consortia relative to WT. This 398 lowered carrying capacity is driven by both lower growth rates as well as increased 399 negative autoregulation in each species, likely due to the dual burden of auxotrophy and 400 overproduction.

401

402 To match these realistic design constrains, we ran simulations in which increases in 403 cooperative interactions were always accompanied by decreases in intrinsic growth 404 rates and increases in the magnitude of negative autoregulation (Figure 5C.) We further 405 assumed no interactions prior to engineering. In this scenario, the stability margin 406 cannot be arbitrarily increased, and an engineering design arises, with intermediate 407 cooperation strengths, that optimizes consortium robustness. If cooperation strengths 408 are increased beyond this level, the consortium becomes less stable. Thus, we see that 409 under realistic constraints, there is a trade-off between cooperativity and self-interest in 410 the consortium, with an optimal intermediate.

411

412 We next investigated the impact of pre-existing interactions, an important feature of 413 naturally occurring heterogeneous bacterial species, on stability (Figure 5D.) In this 414 case, the robustness of the consortium again cannot be arbitrarily increased, and an 415 engineering design that optimizes consortium stability arises. However, in contrast to 416 the case with no interactions prior to engineering (Figure 5C), the optimal cooperativity 417 strength is higher. This reflects the fact that engineering must first push pre-existing 418 competitive interactions toward neutrality before pushing interactions into the optimal 419 regime for consortium stability. These results suggest gualitative design principles for

420 engineering cooperative bacterial consortia, and provide tools for future analyses of

- 421 specific designs.
- 422

423 Consortia Engineering Increases Population Calance in the Mammalian Gut in a

424 Diet Dependent Manner

425 We investigated the behavior of our consortium in the mammalian gut, using gnotobiotic 426 mice as a controlled yet sufficiently complex environment for evaluation. To investigate 427 the role of amino acid cross-feeding *in vivo*, we altered amino acid levels in the gut by 428 changing the animal's diet (Ravindran et al., 2016). Groups of five germfree mice were 429 fed standard or low protein (3%) chow and gavaged with either the WT or engineered 430 consortium (Figure 6). The consortia were allowed to colonize for 10 days, and then 431 stool samples were collected and interrogated via qPCR with species-specific primers. 432 The engineered consortium consistently exhibited greater population evenness in mice 433 that were fed low protein diet compared to the three other groups (Mann-Whitney test: 434 *p*-values: 0.02, 0.03, 0.02).

435

436 Our results show that diet influences total bacterial concentrations in both engineered 437 and WT consortia, and each species in the consortium is affected to a different extent. 438 For the engineered consortium, species abundances were higher by a factor of 439 approximately 3 for *E. coli*, 8 for *S. Typhimurium*, 16 for *B. theta*, and 11 for *B. fragilis* in 440 mice fed a standard versus a low protein diet. In the case of the WT consortium, S. 441 Typhimurium, B. theta and B. fragilis concentrations were similarly higher on standard 442 chow (fold changes of approximately: 10, 22, 13 respectively). WT E. coli 443 concentrations were dramatically higher (fold change of approximately 51), and partially 444 account for the greater population imbalance in the WT consortium. Interestingly, in the 445 mice on a low protein diet, S. Typhimurium grew about 8-fold better in the engineered 446 consortium compared to the WT consortium. This finding is consistent with our *in vitro* 447 results, which indicated that engineered S. Typhimurium benefits most from growing in 448 co-culture. The same trend can be observed in mice that were fed standard diet, albeit 449 to a lesser extent.

451

452 **Discussion**

453 We have engineered a heterogeneous synthetic bacterial consortium from four different 454 gut-derived species, and demonstrated that this consortium exhibits inducible 455 cooperativity, with increased stability and population balance. Using data-driven 456 dynamical systems models, we have elucidated the interaction network among 457 consortia members and shown that our engineering strategy acts to increase 458 cooperativity largely by neutralizing pre-existing competitive interactions. Simulations 459 based on the derived model provide further insights into general synthetic design 460 strategies for these systems, showing a regime of optimal cooperativity that maximizes 461 consortium stability. Finally, we demonstrate that our engineered consortium exhibits 462 increased population balance in the complex mammalian gut environment when 463 induced to cooperate, with this behavior alterable by the host diet. 464 465 This work differs from previous approaches in two key aspects. First, while there have been reports of engineered multi-strain consortia (Minty et al., 2013; Zhou et al., 2015), 466 467 we have engineered four different species that have relevance to gut applications. 468 Second, previous efforts to engineer interactions via metabolite cross-feeding have 469 relied on metabolite auxotrophies without corresponding overproduction (Mee et al., 470 2014; Wintermute and Silver, 2010a). Here, we selected for overproducing strains for 471 each of the four species, applying knowledge from industrial amino acid overproduction 472 (Becker and Wittmann, 2012).

473

474 We found that there are strong pre-existing negative interactions between WT strains 475 (Figure 3b), which is not altogether surprising given the natural history of gut 476 commensal bacteria. Negative interactions may include acidification (Ratzke et al., 477 2018), scavenging of metals or other micronutrients (Hider and Kong, 2010), and 478 competition for carbohydrate sources, among others. In some instances, we achieved 479 measurable positive interactions, whereas in other cases the negative interactions were 480 neutralized. Of note, our engineered consortia growth is reduced by about 100-fold 481 compared to the WT consortia (Figure 3 a,b). In essence, we have introduced improved

482 cooperativity into the consortium with concomitant gains in stability and population 483 balance, at the expense of strains' individual fitness. Loss of fitness in this case is likely 484 due to insufficient complementation by the overproducers as shown in our initial system 485 characterization (Figure 2). Complementation could be improved in two ways: first 486 directed evolution approaches could render a more efficient consortium, and second, 487 we could apply rational engineering (e.g. introducing transporters or increasing 488 overproduction) to improve cross-feeding. Genome sequencing of our mutated strains 489 could aid in such approaches in future work.

490

491 We have created a model ecosystem that allows us to study the effects that cooperation 492 has on microbial consortia. While microbial cooperation is found in natural habitats 493 (Hays et al., 2015b; Ponomarova et al., 2017; Rakoff-Nahoum et al., 2016), and there 494 are many ecological theories that attempt to explain its evolution (Nowak, 2006; West et 495 al., 2006; Zomorrodi and Segrè, 2016), there is a dearth of experimental systems to test 496 such hypotheses. In our system, cooperation leads to improved growth of the overall 497 synthetic consortia, but it also promotes continuing survival of each single species 498 through improved population evenness. These two characteristics of our system make it 499 an attractive test-bed to address questions about the evolution of cooperation, which 500 serves both the consortia as a whole and each single species' survival.

501

502 We chose cross-feeding of amino acids as a model of cooperativity, because this approach has been well established and could readily be applied to disparate bacterial 503 504 species. However, amino acids are also abundant in the animal gut and other 505 environments, raising the possibility that background amino acids levels would simply 506 saturate our synthetic consortium. Remarkably, our consortium demonstrated inducible 507 cooperativity with increased population balance in our gnotobiotic mouse model (Figure 508 6), despite the fact that the germ-free mouse gut contains amino acids from both diet 509 and host sources. This experimental system provides a relatively controlled, but still 510 complex environment, and allows us to readily interpret behavior of the consortia alone 511 without having to consider interactions with a pre-existing microbiota. Of note, all of the 512 strains we used have been shown to individually colonize conventional mice, in some

513 cases without the need for prior antibiotic treatment (Kotula et al., 2014; Riglar et al.,

514 2017). However, our approach using amino acid crossfeeding may not translate to a

515 mammalian gut colonized with a pre-existing microbiota. Thus, when extending our

516 results to more complex systems, more orthologous approaches to cooperativity are

517 likely to be necessary to avoid host interference.

518

519 Our computational approach is data-driven and phenomenological, abstracting various 520 types of possible biological interactions (e.g., competition for nutrients, bacteriocin 521 production, syntrophy, etc.) into quantitative pairwise interaction coefficients. This is a 522 different modeling approach than that of many prior auxotroph-overproducer studies, 523 which built detailed metabolic models. These models have not been shown to be 524 reliable for diverse bacteria, such as the *Bacteroides spp*. in our consortium, in part due 525 to limited knowledge of bacterial metabolism outside of a small number of model 526 organisms. Moreover, as discussed, in a consortium with heterogeneous commensal 527 gut bacteria as members, we expect there to be pre-existing interactions, many of which 528 may not be metabolic. Thus, to gain insights into our consortium, rather than build a 529 bottom-up metabolic model, we infer a phenomenological model from data. Our model 530 is based on stochastic generalized Volterra-Lotka (gLV) dynamics, which is a relatively 531 simple model. More sophisticated models incorporate higher-order interactions or 532 nonlinearities such as saturation effects. However, gLV models have been shown to 533 accurately forecast dynamics in complex host-microbial ecosystems (Bucci et al., 2016), 534 suggesting that the relatively simple assumption of pairwise guadratic interactions may 535 dominate higher-order and more nonlinear effects. Moreover, reliable inference from 536 data for more complicated models is not possible given the amount of densely sampled 537 time-series data that we could feasibly collect for the present study. 538

539 Overall, we have demonstrated a design, build and test cycle, applicable to engineering 540 microbial consortia that function in the mammalian gut. There is currently interest in 541 developing living bacterial diagnostics and therapeutics for human diseases. The first 542 approach has been to attempt to transfer an uncharacterized microbiota, i.e., fecal 543 microbiota transplants (Boyle, 2015). Subsequently, there have been efforts to

544	assemble defined	collections of na	aturally occurring	commensal bacteria	(Atarashi et al.).
					(**************************************

545 While these approaches may work for some diseases, we could anticipate a need for

- 546 precise and controllable behavior of therapeutics, which will require synthetic biology.
- 547 Our results help to close this gap, providing new insights into the design principles
- 548 needed to engineer robust and heterogeneous bacterial consortia.
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567 Author Contributions

- 568 Conceptualization, J.C.W, P.A.S, G.K.G; Methodology, M.Z., T.G., J.O., G.K.G.;
- 569 Software, T.G.; Formal Analysis, M.Z., T.G.; Investigation, M.Z., J.O., A.S., T.G., N.D.,
- 570 K.L.; Resources, L.B., P.A.S., G.G.; Writing Original Draft, M.Z., T.G.; Writing –
- 571 Review & Editing, B.B.H., D.T.R, P.A.S., G.K.G.; Visualization, D.T.R., B.B.H.;
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- 574

- 575 Financial Interests
- 576 G.K.G. is a shareholder and member of the Strategic Advisory Board of Kaleido
- 577 Biosciences, and shareholder and member of the Scientific Advisory Board of Consortia
- 578 Rx; neither company provided funding for this work.
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- 581
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- 584
- 585 Methods

586 Auxotroph engineering

587 For auxotroph generation in the *E. coli* NGF-1 strain we introduced multiple knockouts 588 using sequential P1 transduction (Thomason et al., 2007) from the Keio knockout 589 collection (Baba et al., 2006). Flip-out of kanamycin cassettes was done using pCP20 590 (Cherepanov and Wackernagel, 1995). In brief, for P1 transduction we prepared phage 591 by diluting an overnight culture of the donor strain 1:100 LB with 0.2% glucose, 5 mM 592 CaCl₂ and 25 mM MgCl₂ and incubated for 1-2 hours at 37 °C until slightly turbid. We 593 then added 40 µL P1 lysate and continued growth for 1-3 h at 37 °C while shaking until 594 lysed. Lysate was then filtered with a 20 µm sterile filter and stored in the fridge. For 595 transduction, we harvested 2 mL overnight culture of recipient strain and re-suspended 596 in 2 mL LB with 5 mM CaCl₂ and 100 mM MgSO₄. We then mixed 100 µL donor lysate 597 with 100 µL recipient, incubated 30 min at 37 °C and added 200 µL sodium citrate (1 M, 598 pH 5.5) and 1 mL LB and incubated for another 1 hr at 37 °C. Cells were harvested, re-599 suspended in 100 µL LB with 100 mM sodium citrate and plated on LB Kan plates (75 600 µg/mL). The transduced kanamycin cassette was then removed using pCP20 according 601 to protocol. We transformed pCP20 via electroporation and transformants were selected 602 on LB agar plates supplemented with 100 µg/mL carbenicillin grown at 30 °C. Single 603 colonies were re-streaked on LB without drugs and incubated for 10 hours at 42 °C. 604 From there, single colonies were re-streaked on LB plates without drugs and grown 605 overnight at 37 °C. Colonies were checked for Carbenicillin, and Kanamycin sensitivity

and further confirmed via PCR at respective loci. This procedure was repeated until allknockouts were introduced.

608

609 Engineering of S. Typhimurium LT2 required generation of single knockout strains that 610 contained pKD46 integrated into the genome, which allowed for linear DNA integration 611 using lambda red recombination (Cherepanov and Wackernagel, 1995). We then 612 introduced the knockouts into the S. Typhimurium strain through sequential P22 613 transduction and pCP20 flipout analogous to *E. coli* engineering. Single knockout 614 strains were generated by PCR amplifying a Kanamycin resistance cassette from 615 pKD13 generating linear fragments that contained upstream and downstream homology 616 to the gene of interest and the kanamycin cassette with FRT sequences. Fragments 617 were introduced via electroporation and selected on LB agar plates supplemented with 618 50 µg/mL Kanamycin. Sequential P22 transduction and pCP20 flip-out was essentially 619 performed as described above for P1 transduction but lysis was done overnight.

620

621 For knockout generation of both *B. theta* and *B. fragilis*, we used pExchange KO vectors 622 as described (Mimee et al., 2015). Briefly, we introduced 750 bp flanking regions for 623 genes of interest adjacent to each other into the vector. The vector contains an 624 erythromycin resistance positive marker and a thymidine kinase as counter selection 625 marker. Cloning was done in pir⁺ E. coli strains and vectors were transferred to MFDpir 626 for conjugation (Ferrières et al., 2010). Conjugation was done according to protocol with 627 minor changes. In brief, five drops of overnight culture of *E. coli* donor was inoculated in 628 LB supplemented with 300 µM Diamino pimelic acid (DAP) and five drops of recipient 629 overnight culture was inoculated in 50 mL basal media. Both cultures were grown for 630 about 2 hr (E. coli aerobically, Bacteroides spp. anaerobically) until E. coli culture was 631 well turbid and *Bacteroides* culture just slightly turbid. Subsequently, 9 mL recipient and 632 3 mL donor were combined and spun down for 10 min at 4000 rpm together. The pellet 633 was re-suspended in 100 µL fresh basal media with 300 µM DAP and pipetted on basal 634 media agar plates without cysteine and supplemented with 300 µM DAP. The cells were 635 incubated at 37 °C aerobically face up for up to 20 hr, scraped off and re-suspended in 636 10% glycerol. Dilutions were plated on basal-agar plates supplemented with 10 µg/mL

637 erythromycin and incubated at 37 °C anaerobically for 2-3 days. Single colonies were 638 re-streaked in the presence of erythromycin and grown for another 2 days. 10 single 639 colonies were inoculated in basal media without drug and grown overnight. 500 µL of 640 each culture was mixed, spun down and re-suspended in 10% glycerol. We then plated 641 different dilutions on basal media plates supplemented with 5-fluoro-2-deoxy-uridine 642 (FuDR) (200 µg/mL) and incubated at 37 °C anaerobically for 3 days. Knockouts were 643 verified via PCR. This procedure was repeated multiple times to obtain the multiple 644 auxotroph strains.

645

646 **Overproducer selection**

647 Overproducers were generated by selecting for mutants that could grow on minimal media agar plates supplemented with anti-metabolites (E. coli: 5 mg/mL Norleucine for 648 649 Met overproduction; S. Typhimurium: > 0.7 mg/mL beta-(2-thiazolyl)-DL-alanine for His 650 overproduction: B. theta: 50 µg/mL 4-methyl tryptophan for Trp overproduction; B. 651 fragilis: 80 µg/mL Canavinine for Arg overproduction). Single colonies that showed 652 halos were re-streaked and overproduction was measured using a bioassay. In brief, for 653 screening of overproducing mutants the isolated strains were grown overnight at 37 °C 654 shaking aerobically (for E. coli and S. typhimurium) or anaerobically without agitation 655 (for *Bacteroides spp.*). Supernatant was harvested, diluted 1:1 with fresh media, and E. 656 coli auxotrophs were inoculated and their growth was recorded after 24 hr. For E. coli NGF-1 overproducers, we used a S. Typhimurium auxotroph instead, since its colicin 657 658 production prevented the *E. coli* biosensor from growing. Confirmed overproducers 659 were further quantified using LC-MS.

660

661 LC-MS for Overproduction Measurements

To quantitate amino acid levels in overproducer supernatants, a standard curve was obained using freshly prepared amino acid standards dissolved in growth media (1mM, 500 uM, 100 uM, 50 uM, 10 uM of L. Methionine, L/Histidine, L-Tryptophan, L-Arginine each). To prepare for HPLC-MS analysis, 0.5 mL sample or standard were added to 1.5 mL ice-cold methanol and incubated on ice for 10 min. The mixture was centrifuged for 5 min at 15,000 rpm and 500 µL supernatant was vacuum concentrated and resuspended in 50 µL methanol. Samples were kept on ice or at 4°C. HPLC-MS analysis

- 669 of standards and extracts was carried out using an Agilent 1260 Infinity HPLC system
- equipped with an Agilent Eclipse Plus C18 (100 × 4.6 mm, particle size 3.5 mm, flow
- rate: 0.3 mL/min, solvent A: dd.H₂O/0.1% (v/v) formic acid, solvent B: acetonitrile,
- 672 injection volume: 4 mL) connected to an Agilent 6530 Accurate-Mass Q-TOF
- 673 instrument. The following gradient was used (time/min, %B): 0, 0; 0.5, 0; 14, 100; 19,
- 100; 20, 0, 25, 0. The mass spectrometer was operated in positive mode and the
- autosampler was kept at 4°C. After HPLC-MS analysis, extracted ion current (EIC)
- 676 peaks were automatically integrated using the MassHunter Workstation Software
- 677 (version: B.07.00). A plot of peak area versus amino acid concentration was used to
- 678 generate a linear fit.
- 679

680 Sequencing

- 681 Bacterial cultures were prepared in rich media (basal for *Bacteroides spp.* and LB for *E.* 682 coli and S. Typhimurium). Genomic DNA (gDNA) extraction was performed using the 683 Wizard Genomic DNA Purification Kit (Promega) according to protocol. The extracted 684 gDNA was sheared using Covaris DNA Shearing, and the library was prepared using 685 Kapa Biosystem DNA Hyper Prep NGS Library (Dana Faber Core MBCFL Genomics). 686 Sequencing was performed on the Illumina MiSeg instrument, with the 150 bp paired 687 End (PE150) reagents. Sequences were analyzed for SNPs using Geneious software 688 and published genome sequences (*E. coli*: CP016007.1; *S. typhimurium*: NC 003197; 689 B. theta: AE015928; B. fragilis: NC 016776) (Table S1).
- 690

691 Growth and Media Conditions

All basal media and co-culture media was pre-incubated for at least 24 hr anaerobically

- 693 before use. *Bacteroides spp*. were inoculated from glycerol stock into basal media,
- 694 grown overnight and 400 μL was inoculated in 5 mL basal and grown 2 hr anaerobically.
- 695 Cells were spun down, washed twice in PBS and diluted in growth media as described
- 696 for each experiment in Results. *E. coli* and *S. Typhimurium* were inoculated from
- $\,$ 697 $\,$ glycerol stock into LB and grown overnight at 37 °C while shaking. 100 μL of culture
- 698 was then inoculated into pre-incubated LB and grown anaerobically for 2 hr, diluted,

- 699 washed in PBS and diluted into co-culture media as described. Co-culture media
- consisted of modified M9 salts (0.2 g/L Na₂HPO₄, 90 mg/L KH₂PO₄, 30 mg/L NH₄Cl, 15
- 701 mg/L NaCl), 1 mM MgSO₄, 10 μg/mL heme, 0.1 mM CaCl₂, 1 μg/mL Niacinamide,
- vitamin B12 and thiamine, 400 µg/mL L-cysteine, 0.3% bicarbonate buffer, 2.5 ng/mL
- vitamin K, 2 μ g/mL FeSO₄*7H₂O and carbon sources and amino acid supplementation
- as described in Results.
- 705

706 Multiplex qPCR

- 707 We designed strain specific primer/probe-fluorophore pairs according to IDT protocol
- 708 (Table S2). We chose strain specific genes by multiple genome alignment between the
- strain of interest, the other consortia members and closely related strains using Mauve
- 710 (Darling et al., 2004). Multiplex qPCR was used to quantify each strain in co-culture by
- vising a standard curve obtained by plating late log phase cultures grown in rich media.
- In brief, each strain was grown from overnight culture for ~5 hours until about OD of 1.
- 713 Cells were then counted by plating. Cultures were mixed, diluted and frozen at -80 °C
- for use as standard curve. Samples were diluted 1:10 in ddH₂O and snap-frozen in
- 715 liquid nitrogen and stored at least overnight at -80 °C. Growth curve and sample were
- both thawed together and prepared in a 5 μ L Primetime Mastermix (IDT) with 1 μ L
- 717 Primer/Probe mixture (final concentrations: 100 nM for primers and 50 nM for probes).
- The qPCR was run with the following program: 20 min at 98 μ C (to boil the cells and
- denature gDNA), followed by 40 cycles of 60 °C and 98 °C.
- 720

721 Calculation of Normalized Entropy (Pielou's Evenness) and Hellinger Distance

Normalized entropy (Pielou's evenness) was calculated according to the given formula:

Pielou's Evenness Index =
$$\frac{-\sum_{i=1}^{4} p_i * \ln(p_i)}{\ln(S)}$$

725

724

Where p_i refers to the population ratio of a given strain in the consortium of four strains.

727 S is the number of species.

- Hellinger distance was used as a metric for comparing consortia relative abundances.
- 729 We computed Hellinger distances between different starting conditions in WT and
- rain engineered strains using the formula:
- 731

732 *Hellinger distance* =
$$\frac{1}{\sqrt{2}} \sqrt{\sum_{i=1}^{4} (\sqrt{p_i} - \sqrt{q_i})^2}$$

- 733 Where p_i and q_i are the ratios of each strain in the consortium in two different conditions.
- 734
- 735

736 Pairwise Supernatant Experiment

- 737 Overproducers and WT strains were grown overnight anaerobically at 37 °C in co-
- culture media and supernatant was harvested and sterile filtered. Auxotrophs were
- prepared as described in **Media and Growth conditions** and inoculated in media that
- contained 50% fresh co-culture media and 50% spent supernatant. OD600 was
- 741 measured after 24 hr growth anaerobically at 37 °C.
- 742

743 In vivo Experiments

- Adult (6-8 weeks) male Swiss Webster germ free mice bred in house at the
- 745 Massachusetts Host-Microbiome Center were used. Animals were fed either on
- standard chow in the facility for the entire experiment, or on low-protein diet (3% custom
- 747 diet, envigo, doubly irradiated) beginning 10 days prior to the experiment and continuing
- for its duration. To prepare bacteria for gavage, we grew each strain to mid-log phase,
- plated for counting and snap-froze aliquots. For gavage, aliquots were thawed, spun
- down, combined to achieve concentrations of approximately 10⁷ per bacteria per
- 751 gavage, and re-suspended in 200 μL 1x PBS with 0.05% L-cysteine for gavage. After
- gavage, mice were transferred to Optimice cages and maintained gnotobiotic for 10
- days. Fecal samples were collected prior gavage and at 10 days, and snap-frozen for
- storage at -80 °C.
- 755

756 Molecular Analysis of *in vivo* Samples

757 DNA was extracted from fecal samples using the Zymobiomic 96 DNA Kit with the

- following modification: we omitted the silicon-ATM-HRC wash. Cells were lysed in a
- bead beater at speed 20 for 10 min, plates were turned and lysed for another 10 min at
- the same speed. We added an additional 3 min incubation step for Binding Buffer and
- additional 5 min incubation steps when transferring to Zymo-Spin-I-96-Z plates. Elution
- 762 was done in 50 µL ZymoBIOMIC DNase/RNase Free Water.
- 763
- 764 Direct multiplex probe-based qPCR was done on extracted DNA samples as described
- above. For standard curves, we used plated overnight cultures spiked into germfree
- ⁷⁶⁶ fecal samples and extracted them as described above.
- 767

768 Synthetic simulations of deterministic microbial dynamics in the presence of a

- 769 disturbance
- For the synthetic results in Figure 1 and Figure 5, we used the following (deterministic)
- generalized Lotka-Volterra (gLV) dynamics with a constant disturbance d

772
$$\frac{\mathrm{d}x_{k+1,i}}{\mathrm{d}t} = x_{k,i} \left(r_i + \sum_{j=1}^4 a_{i,j} x_{k,j} \right) + d_i$$

The corresponding first order Euler integration of these dynamics with step size Δ_k is as follows

775
$$x_{k+1,i} = x_{k,i} + x_{k,i} \left(r_i + \sum_{j=1}^4 a_{i,j} x_{k,j} \right) \Delta_k + d_i \Delta_k.$$

- For the simulations to generate Figure 1, the growth rate vector, interaction matrix, and
- 777 disturbance vector for the non-interacting consortia are

778
$$r = \begin{bmatrix} .3\\ 6\\ 9\\ 12 \end{bmatrix}, \quad A = \begin{bmatrix} -2 & 0 & 0 & 0\\ 0 & -2 & 0 & 0\\ 0 & 0 & -2 & 0\\ 0 & 0 & 0 & -2 \end{bmatrix}, \text{ and } d = \begin{bmatrix} 0\\ -5.3\\ 0\\ 0 \end{bmatrix}.$$

The growth rate vector, interaction matrix, and disturbance vector for the cooperative

780 community are then as follows

781
$$r = \begin{bmatrix} .3\\ 6\\ 9\\ 12 \end{bmatrix}, A = \begin{bmatrix} -2\ 0.5\ 0\ 0.5\\ 0\ -2\ 0\ 0.5\\ 0\ 0.2\ -2\ 0\\ 0.2\ 0\ 0.2\ -2 \end{bmatrix}, \text{ and } d = \begin{bmatrix} 0\\ -5.3\\ 0\\ 0 \end{bmatrix}.$$

Two simulation were performed for both the non-interacting and the cooperative

community: one simulation without the disturbance present, and then the subsequent

simulation with the disturbance present.

785

For the simulations to generate Figure 5, the parameterization and simulation

parameters are as specified in the figure. The robustness margin was calculated as the

788 largest value of κ for which the disturbance vector

789 $d = \begin{bmatrix} 0 \\ -\kappa \\ 0 \\ 0 \end{bmatrix}.$

can be applied to the dynamics without any of the x_i becoming <0.01 in abundance (i.e.,

without any of the species going extinct during the simulation). For all synthetic results

shown in Figures 1 and 5, the step size was $\Delta_k = 0.005$ for a total of 3000 steps. For

generating Figure 5, the initial condition for all the species was an abundance of 2.

794

795 Bayesian dynamical systems inference

Our dynamical systems model and associated inference algorithm is a version of our previously published method (Gibson and Gerber, 2018) that we have customized for this study. Briefly, our model is based on continuous time stochastic gLV dynamics as described in Results. We approximate these dynamics with a first order Euler integration of step size $\Delta_{k,\ell}$

801
$$x_{k+1,i,\ell} = x_{k,i,\ell} + x_{k,i,\ell} \left(r_i + \sum_{j=1}^4 a_{i,j} x_{k,j,\ell} \right) \Delta_{k,\ell} + \sqrt{\Delta_{k,\ell}} \left(w_{k+1,i,\ell} - w_{k,i,\ell} \right)$$

802 Our model is fully Bayesian, with the inference goal being to learn the posterior 803 probability distribution over both the model parameters as well as the qualitative 804 interaction structure graph. To model the interaction structure graph, we use indicator 805 variables $z_{i,j}$, which indicate the presence or absence of the edge from species *i* to *j* in 806 the graph. The resulting conditional distribution for dynamics is then:

807
$$x_{k+1,i,\ell} | x_{k,[n],\ell}, r, a, z, v_i^w$$

808

~ Normal
$$\left(\boldsymbol{x}_{k,i,\ell} + \boldsymbol{x}_{k,i,\ell} \left(\boldsymbol{r}_i + \boldsymbol{a}_{i,i} \boldsymbol{x}_{k,i,\ell} + \sum_{j \neq i} \boldsymbol{a}_{i,j} \boldsymbol{z}_{i,j} \boldsymbol{x}_{k,j,\ell} \right) \Delta_{k,\ell}, \Delta_{k,\ell} \boldsymbol{v}_{i,\ell}^{\boldsymbol{w}} \right)$$

809 Where $v_{i,\ell}^w$ is the variance for the Brownian motion. Our measurements of microbial 810 abundances are captured by the observed variable y. We introduce an auxiliary variable 811 q that enables efficient inference of a relaxed system (see Gibson and Gerber, 2018 for 812 details):

813
$$\boldsymbol{q}_{k,i,\ell} \mid \boldsymbol{x}_{k,i,\ell} \sim \operatorname{Normal}(\boldsymbol{x}_{k,i,\ell}, \boldsymbol{v}^{\boldsymbol{q}})$$

814
$$\boldsymbol{q}_{k,i,\ell} \sim \text{Uniform}[0,L)$$

815
$$\mathbf{y}_{k,i,\ell} \mid \mathbf{q}_{k,i,\ell} \sim \operatorname{Normal}_{\geq 0}(\mathbf{q}_{k,i,\ell}, \mathbf{v}_{k,i,\ell}^{\mathbf{y}})$$

We estimate the measurement variance $v_{k,i,\ell}^{y}$ directly from qPCR technical replicates, obtaining $L = 10^{10}$, and $v^{q} = 10^{6}$. The prior probability distributions for the indicator variables, the growth rates, and the interactions terms as defined below:

- 819 $\mathbf{z}_{i,i} \sim \text{Bernouli}(0.5)$
- 820 $r_i | v^r \sim \text{Normal}(0, v^r)$

821
$$\boldsymbol{a}_{i,j} \mid \boldsymbol{\nu}^{\boldsymbol{a}} \sim \operatorname{Normal}(0, \boldsymbol{\nu}^{\boldsymbol{a}})$$

Note that our prior for *z*, the indicator variables for presence or absence of interaction edges is set to indicate maximum uncertainty, e.g., no *a priori* assumption about the presence or absence of an interaction. The variance parameters in our model that are not *a-priori* fixed or estimated from technical replicates have the following conjugate priors

827 $\boldsymbol{\nu}^{\boldsymbol{r}} \sim \operatorname{Inv} - \mathbf{X}^2(\boldsymbol{\eta}_{\boldsymbol{r}}, \boldsymbol{\theta}_{\boldsymbol{r}})$

828
$$\boldsymbol{\nu}^{\boldsymbol{a}} \sim \operatorname{Inv} - \mathbf{X}^2(\eta_{\boldsymbol{a}}, \theta_{\boldsymbol{a}})$$

829
$$\boldsymbol{\nu}^{\boldsymbol{w}} \sim \operatorname{Inv} - \mathbf{X}^2(\eta_{\boldsymbol{w}}, \theta_{\boldsymbol{w}})$$

830 Our parameterization of the $Inv - X^2$ is as follows

831
$$f(x;\eta,\theta) \triangleq \frac{(\theta\eta/2)^{\eta/2}}{\Gamma(\eta/2)} \frac{\exp\left[-\frac{\eta\theta}{2x}\right]}{x^{1+\eta/2}}$$

832 which is sometimes referred to as the Scale – $Inv - X^2$ distribution, because it has two 833 parameters, number of degrees of freedom η , and scale parameter θ . In our model 834 $\eta_r, \eta_r, \eta_r = 1$ with $\theta_r = 1$, $\theta_a = 10^{-10}$, and $\theta_w = 10^6$, specifying relatively diffuse priors. A 835 compact representation of our model is shown in Supplemental Figure 1a with an 836 accompanying graphical model with plate notation in Supplemental Figure 1b

837

838 We perform model inference using a custom Markov Chain Monte Carlo (MCMC)

algorithm. Almost all variables in the model can be updated with Gibbs or collapsed

Gibbs sampling, with the exception of *q* and *x*. Sampling of the auxiliary variables and

841 latent trajectories require Metropolis-Hastings (MH) steps. For q, the MH proposal is

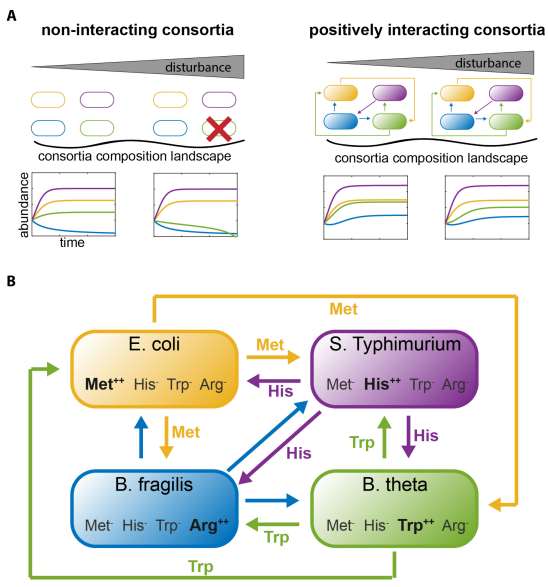
based on a Generalized-Linear Model approximation. For x, we use a one time-step

ahead proposal that is essentially the forward pass of a Kalman filter, see Supplemental

Figure 1c and (Gibson and Gerber, 2018). Inference was performed with 6,000 MCMC

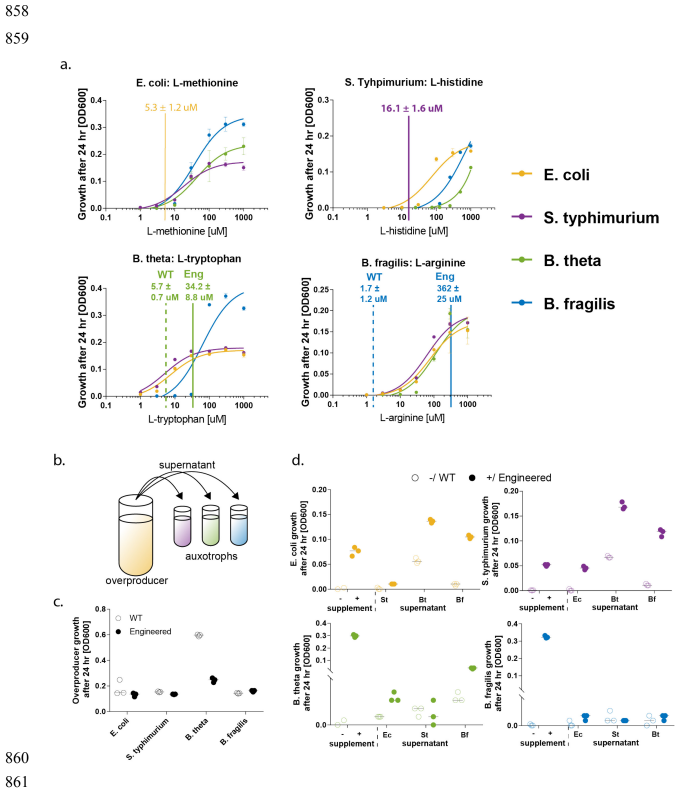
steps where the first 1,000 steps are discarded (burn in).

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848 Figure 1 Engineering positive interactions in a microbial consortia introduces 849 resilience to disturbances. (A) Simulations show increased stability in consortia with 850 positive interactions. In a non-interacting consortia, disturbances can lead to dramatic 851 changes in consortia composition; in this example, the disturbance causes one of the 852 strains to die out. Introducing positive interactions among the consortia strains 853 increases stability in the presence of disturbances. (B) Our engineering design, which 854 introduces mutual positive interactions by cross-feeding metabolites. Each strain was 855 knocked out for three amino acid biosynthesis pathways and mutated to overproduce 856 one amino acid. Thus, four amino acids L-methionine (Met), L-histidine (His), L-857 tryptophan (Trp) and L-arginine (Arg) are cross-fed between the four strains.

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862 Figure 2 Characterization of auxotroph requirements and overproduction

863 capabilities in a 4-species consortium. a. Growth response with indicated

overproduction. Each auxotroph was grown in media supplemented with varying
concentrations of one amino acid and saturating concentration of the two others.
Depicted is the average of three biological replicates; error bars indicate standard
deviation. A sigmoidal curve was fit using GraphPad Prism 7. Overproduction
(horizontal lines) falls largely within requirement values. **b.** Cross-feeding capabilities of

868 (nonzontal lines) rais largely within requirement values. **b.** Cross-recting capabilities of
 869 each strain were assessed by testing for rescue of auxotrophs in supernatants obtained

870 from overproducers. **c.** Growth of overproducers after 24 hr, at which point supernatant

871 was collected for cross-feeding experiment. Overproduction does not affect growth with

the exception of *B. theta* (3-fold reduction). Shown are three biological replicates with

873 median indicated as horizontal line. **d.** Growth of auxotrophs with and without amino

acid supplementation and in supernatant of engineered overproducers and WT

equivalents. Shown are three biological replicates with median indicated as horizontal

876 lines. Except for *B. fragilis*, all auxotrophs can be rescued to varying degrees.

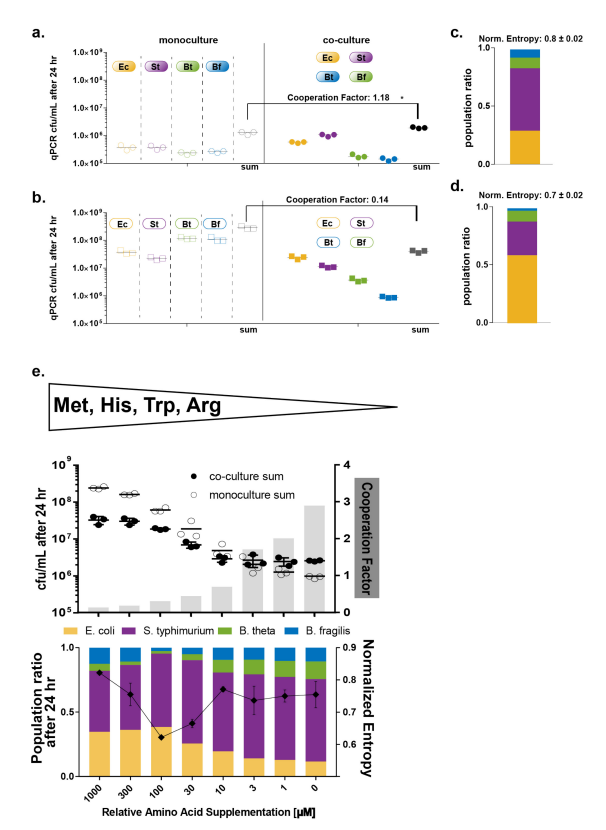
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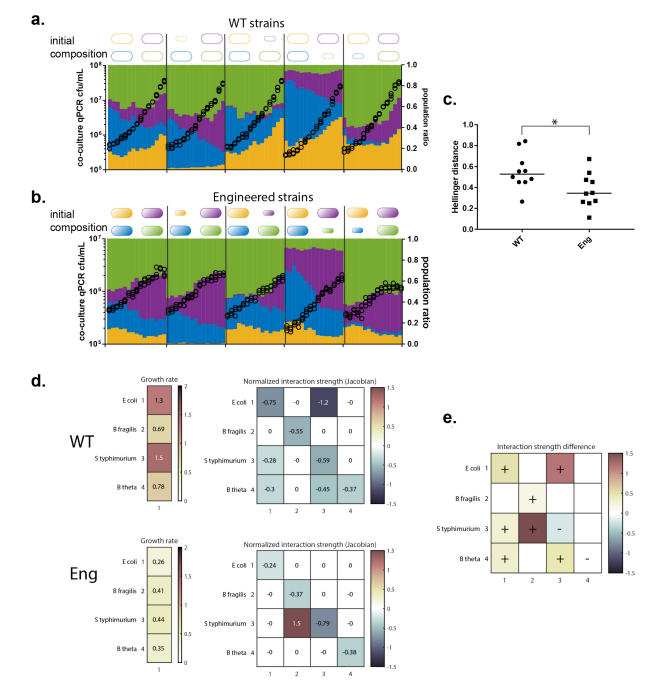




balance. a./b. Growth of engineered (a.) and WT (b.) mono cultures and co-culture

886	grown without supplementation was recorded after 24 hr as qPCR cfu/mL estimates.
887	Shown are three replicates with median indicated. For engineered strains co-culture
888	reaches higher cfu/mL than sum of monocultures quantified by cooperation factor. *sum
889	in co-culture in engineered strains is significantly larger than sum in monocultures
890	(Mann-Whitney test; <i>p</i> -value: 0.0071) c./d. Population ratios and calculated balance
891	factor for engineered and WT co-cultures. Balance for engineered strains is higher than
892	for WT strains. e. Supplementation titration experiment. Engineered consortia co-culture
893	and mono cultures were subjected to a range of amino acid supplementation and was
894	analyzed after 24 hr via qPCR. Upper panel: sum of monocultures growth (empty
895	circles) and co-culture growth (filled circles) and calculated cooperation factor (grey
896	bars). Shown are three replicas with median indicated as black line. Cooperation
897	increases with decreasing supplementation. Lower Panel: Population ratios of co-
898	cultures as function of supplementation. Shown is the mean of three replicas. At high
899	supplementation, evenness is high, then drops at intermediate levels and rises again at
900	lower levels. Cooperation and evenness co-vary.
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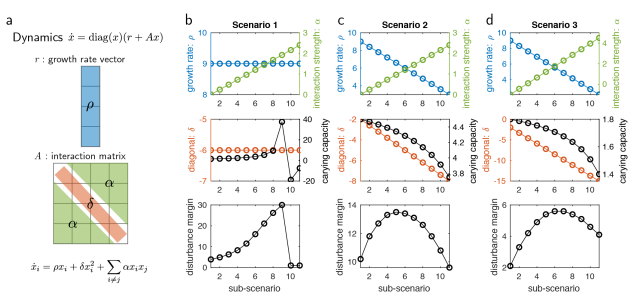
912 Fig 4 Engineered consortia exhibits net positive interaction structure and

913 increased stability. a. We inoculated WT and engineered consortia in media without

- amino acid supplementation and followed growth over time. Starting inocula varied to
- 915 mimic external perturbations. Each strain was inoculated at the same ratio (condition 1)
- and then each one dropped down by 1:10 (condition 2-5). Total bacterial abundance
- 917 (black dots) and relative abundances (colored bars) for each starting condition for WT
- and engineered strains in sequence. Shown are data from one representative

919 experiment. WT consortia grow to about 100-fold higher cfu/mL. c. Hellinger distance 920 between population ratios of all conditions at 12 hr time point calculated for WT and 921 engineered consortia. The Hellinger distance for engineered consortia is lower than for 922 WT consortia (Mann-Whitney *p*-value: 0.0355). **d.** Inferred growth rates and Jacobians 923 of the inferred dynamics for both WT and engineered consortia (only interactions with 924 Bayes factors greater than 1 shown). The Jacobians provide normalized measures of 925 interaction strengths. e. Difference between engineered and WT Jacobians identifies 5 926 net positive, 7 neutral interactions.







929 Figure 5 Simulations investigating consortia stability with different design

930 constraints. a. Baseline model of Lotka-Volterra dynamical system parameterized by

931 growth rate ρ , self-interaction δ (which is negative), and identical interactions 932 coefficients α , for all species. **b.** Scenario 1: as the interactions coefficient is increased,

933 the system has an increased carrying capacity and robustness up to the point of system

934 instability. **c.** Scenario 2: interactions coefficient is increased as the magnitude of the

935 self-limiting term is increased, and the growth rate is decreased. This results in a more

936 realistic scenario and a trade-off between cooperativity and self-interest emerges. d.

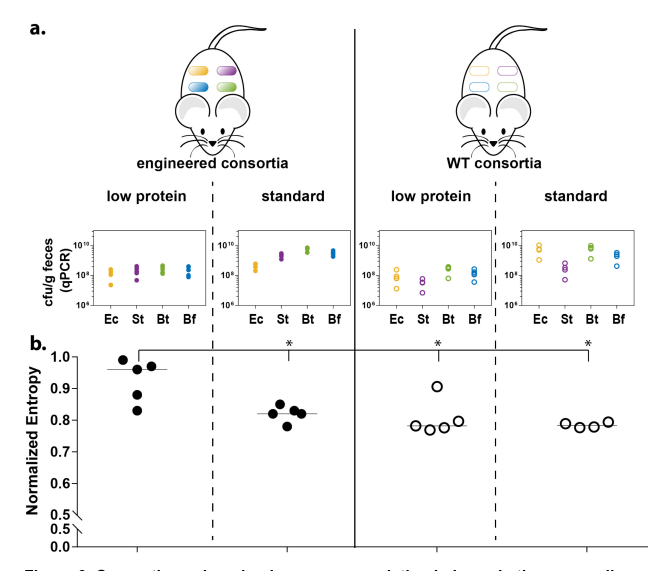
- 937 Scenario 3: similar to Scenario 2, but instead of the interactions coefficient starting at
- 238 zero, it begins with a negative value (pre-existing competitive interactions), mimicking

939 what we observed in our engineered consortia. This also results in a trade-off between

940 cooperativity and self-interest, with a higher optimal interactions coefficient than **c**.

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943 Figure 6. Consortia engineering increases population balance in the mammalian 944 gut in a diet dependent manner. a. Four groups of 5 germfree mice were either fed 945 low protein diet or standard diet, and inoculated with either the engineered or WT 946 bacterial consortia. Fecal samples 10 days post-inoculation were analyzed via strain-947 specific qPCR to assess concentrations of each consortia species. b. Population 948 balance or evenness of consortia, expressed as normalized entropy of the population 949 proportions. Bar indicates Median. Mann-Whitney test showed significantly increased 950 population of the engineered consortia in mice that were fed low protein diet compared 951 to the consortia in the three other groups (p-values: 0.024, 0.032, 0.015). Ec: E. coli; St: 952 S. Typhimurium; Bt: B. theta; Bf: B. fragilis.

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956 **Table 1 Engineered strains and genotypes.** ^{a)}prevents feedback inhibition (Veeravalli

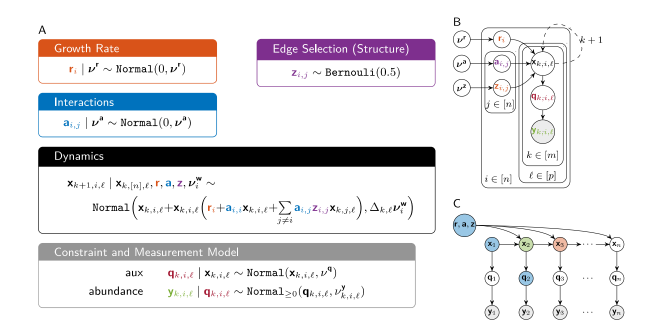
957 et al., 2014), ^{b)}decouples from histidine feedback inhibition (Malykh et al., 2018); ^{c)}trpE,

- 958 removes feedback inhibition (Fang et al., 2015); ^{d)}arginine repressor, nonfunctional
- 959 (Ginesy et al., 2015)

Species	Strain	Auxotroph Genotype	Other Genotype	Overproduction Mutations
E. coli	NGF	$\Delta argA$, $\Delta trpC$, $\Delta hisA$	∆thiE	metA(I296S) ^{a)}
S. Tyhpimurium	LT2	$\Delta argA$, $\Delta trpC$, $\Delta metA$	Δ thiE, Δ SPI1, Δ SPI2	hisG(E271K) ^{b)}
B theta	VPI5482	$\Delta metA$, $\Delta hisG$, $\Delta argF$,	Δ thiSEG, Δ tdk	BT_0532 (A306V; N63D) ^{c)}
B. fragilis	368R	$\Delta metA$, $\Delta hisG$, $\Delta trpC$	Δ thiSEG, Δ tdk	BF638R_0532 (L26R) ^{d)}
			1	= `,

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964 Supplemental Figure 1. a. Description of key components of our Bayesian dynamical 965 systems model for the consortia. Higher level priors not depicted for simplicity. b. 966 Graphical Model depiction with plate notation. c. Portion of the Graphical Model 967 unraveled in time, depicting a single time series experiment, and color coded to 968 illustrate our inference method, using Metropolis-Hastings proposals, for filtering the 969 latent state. The proposal uses information from the blue nodes to propose for the green

970	node, but does not use future time information highlighted in red. The future information
971	is however accounted for in the target distribution and thus the algorithm samples from
972	the true posterior.
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999	References
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