Novel artificial selection method improves function of simulated microbial communities

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There is increasing interest in artificially selecting or breeding 38 microbial communities, but experiments have reported mod- 39 2 est success and it remains unclear how to best design such a selection experiment. Here, we develop computational models to simulate two previously known selection methods and compare them to a new "disassembly" method that we have devel-6 oped. Our method relies on repeatedly competing different communities of known species combinations against one another, 8 and sometimes changing the species combinations. Our approach significantly outperformed previous methods that could ⁴⁶ 10 not maintain enough between-community diversity for selection 47 11 to act on. Instead, the disassembly method allowed many species 48 12 combinations to be explored throughout a single selection ex- 49 13 periment. Nevertheless, selection at the community level in our 50 14 simulations did not counteract selection at the individual level. 51 15 Species in our model can mutate, and we found that they evolved 52 16 to invest less into community function and more into growth. 17 Increased growth compensated for reduced investment, how-18 ever, and overall community performance was barely affected 19 55 by within-species evolution. Our work provides important in-20 sights that will help design community selection experiments. 21

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23 Introduction

Humans have been breeding plants and animals for centuries 24 by allowing individuals with the most desirable traits to se-25 lectively produce offspring. Also known as "artificial selec-26 tion" or "directed evolution", breeding has altered traits such 27 as the size of fruits or the enzymatic activity of proteins used 28 in biotechnology (1). More recently, we have started to ap- $_{66}$ 29 preciate that microbes — often multi-species communities of 67 30 microbes — play an important role for health and the envi- 68 31 ronment. One way to improve or optimize the functions and 69 32 services that these microbes provide is to select for their traits 70 33 in the same way as traditional breeding. 34

³⁵ However, breeding microbial communities is less straightfor-⁷²

- ³⁶ ward than individual organisms (2, 3), mainly because the ⁷³
- 37 breeder selects whole groups of organisms rather than indi-74

vidual plants, animals or proteins. According to evolutionary theory, group-level selection suffers from reduced heritability, one of the main requirements for evolution by natural selection (4). The problem arises when a single communitylevel "generation", which we will call a "round of selection" to avoid confusion, can comprise several generations of cells, each belonging to different genotypes (i.e. species and strains), (Fig. 1A). Since the genotypes all reproduce at varying rates, their relative abundances can change during one round of community growth and over subsequent rounds. Because community traits depend on the traits of all of its genetically distinct constituent members and their proportions, an "offspring" community may not resemble its "parent" (4-6). Another issue with group-level selection is that withinand between-species selection continue to operate within a round. If there are trade-offs between growth and contribution to the community trait, cheaters that contribute less can emerge and sweep to fixation (2, 7). A third challenge is to find a good constellation of different community members and their proportions that can best achieve the desired function. Generating different constellations of member species at each round of selection is also important to have enough variability for selection to act on (4). The major challenges for community-level selection then, are (i) ensuring that community functions are heritable, (ii) that within-community selection does not dominate over between-community selection, and (iii) ensuring variability, that communities differ in phenotype.

In the earliest community breeding experiments, Swenson *et al.* selected microbial communities to yield plants with high and low biomass and to control pH (8). In two out of three experiments, the communities selected for high vs. low function differed significantly from each other, but were not significantly different from the starting communities. The results were also noisy and inconsistent across experimental systems (8, 9). Many attempts have been made since, aiming to optimize several microbial community traits, includ-

ing increased microbial biomass production (10), the stim- 123 75 ulation of various plant properties (10-15), chitin degrada- 124 76 tion (16), the stimulation of fruit fly development (17), to re- 125 77 duce wastewater CO_2 emissions (18), and to hydrolyze starch 126 78 (19). Some of these studies have managed to significantly im- 127 79 prove the average community function over several rounds 128 80 of selection, but sometimes only as an effect of time with- 129 81 out any significant differences between selection treatments 130 82 (8, 16, 17). Overall, community breeding experiments have 131 83 shown mixed success (3, 20), but computer simulations have 132 84 provided some clues on how to improve them (5, 6, 21-23). ¹³³ 85 All previous experiments have followed one of two methods 134 86 to propagate the communities with the highest scores to the 135 87 next round: in the "propagule" selection method (PS), a frac- 136 88 tion of the cells in the highest-scoring communities are se-137 89 lected and transferred by dilution (Fig. 1B), while in "mi-138 90 grant pool" selection (MS), all populations of the selected 139 91 communities are mixed in a pool before they are diluted in 140 92 equal proportions to the new tubes (Fig. 1C). While both 141 93 selection methods have achieved some success, they suffer 142 94 from a rapid decrease in between-community variability (24), 143 95 such that selection has little to act on. Intuitively, the loss of 96 variability arises firstly because only a fraction of commu-144 97 nity members are selected and replicated for the next round. 98 Second, species composition can only change through loss 99 of members when the communities are diluted, meaning that 100 the communities evaluated throughout the whole experiment 101 can only be sub-communities of the initial ones. Given that 102 finding the right species composition is one of the goals of 150 103 community-level selection, this suggests that we need novel 151 104 selection methods that can better explore the search space of 152 105 species combinations (23). 106 153

In this manuscript, we propose a new selection method that 154 107 we call "disassembly selection" (DS), that is designed to 155 108 maintain heritability as well as between-community variabil-109 ity. After each round, we disassemble the selected commu-110 nities by isolating the constituent species before recombin-158 111 ing them into new communities for the next round of growth 159 112 (Fig. 1D). We construct two computational models of mi-113 crobes in a well-mixed liquid culture, one individual-based 161 114 and one based on differential equations, to systematically 162 115 compare our new approach to the classical propagule selec-163 116 tion (PS) and migrant pool selection (MS) methods. 117

Inspired by a four-species community that degrades an in- 165 dustrial pollutant (25), we aim to select for microbial com- 166 munities with improved degradation capabilities. Based on 167 this experimental system, the microbes in our models face a 168 dilemma: whether to invest consumed nutrients into growth 169 or into degradation of toxic compounds that would otherwise cause cell death. The populations evolve by random mutations to this relative investment. We evaluate the selection methods by comparing how the degradation scores change over several rounds of growth and selection starting from the same initial communities. We simulate community selection in both models separately, to test whether our results depend on the choice of model framework.

Our results confirm our intuition that propagule and migrant pool selection do not maintain enough variability to explore many different species combinations, which means that the communities can only improve by mutation. In contrast, our new disassembly approach maintains variability between communities, allowing it to find some of the best possible species combinations. Nevertheless, disassembly selection still suffers from an important problem in group selection: competition within species leads to the dominance of strains that invest less into the function and more into growth. Our work thereby suggests a new method to find species combinations whose community function is high, but in which between-individual competition may be inevitable.

Results

Simulating community-level selection. In either model (see Methods for details), each species is described by its growth and uptake rates for each of 4 available nutrients, and its death and degradation rates for each of 10 toxic compounds. We assume that interactions between cells occur only via nutrients and toxic compounds, as cells of type *i* invest a fraction f_{ik} into degradation of the toxic compounds and the rest into growth. Cells of the same species differentiate by accumulating "mutations" as they grow and divide, that alter the total investment $f_{i} = \sum_k f_{ik}$. All other species properties remain unchanged throughout the simulations.

The simulations start with 21 communities of 4 species each, chosen at random with replacement from a set of 15 initial species, that are described by randomly drawn model parameters. The 21 communities are grown in simulated batch cultures containing defined initial concentrations of nutrients and toxic compounds for a fixed number of time-steps (Fig. 1F). At the end of each round, the 21 communities are scored based on degradation of the ten toxic compounds. The best 7 communities are then selected and propagated to the next round, depending on the selection method: communities are diluted in propagule and migrant pool, whereas they are re-inoculated to a defined population size with equal proportions in the disassembly method (Fig. 1B-D). In disassembly, communities are penalized by species extinctions, and

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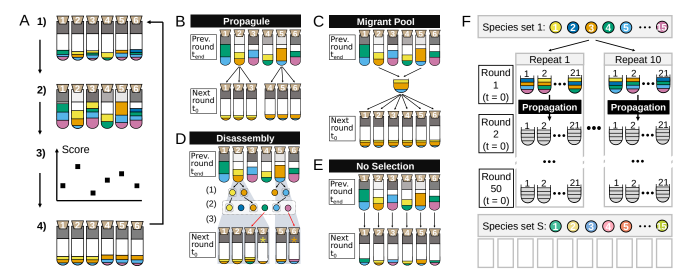


Fig. 1. (A) Overall method for artificial selection of microbial communities. Communities are illustrated as test tubes with bacterial "species" in different colors (white represents empty "space"). The concentration of toxic compounds is shown in shades of gray in the upper part of each tube (darker is more toxic). The inoculated communities (1) grow until the measurement (2) of toxic compound concentration, from which we (3) calculate a score for each community. (4) The highest-scoring communities are selected for propagation into offspring communities and the process is repeated. (B) Propagule: each selected community from the previous round is diluted to form the same number of communities for the next round. (C) Migrant pool: selected communities are merged before dilution. (D) Disassembly: Microbes are (1) isolated from the chosen communities and (2) saved in a repository (dotted rectangle). Each selected community contributes offspring communities in proportion to their degradation score (3). A fraction of the new communities receive new species (red arrows) or lose members from the previous round (asterisk in color of removed species). (E) No-selection control: each community is diluted into a new tube. Propagule, migrant pool and disassembly have selection treatments (PS, MS and DS) and random treatments (PR, MR and DR), where community scores are ignored (see Methods). (F) A "species set" consists of 15 randomly generated species. From this set, we draw 21 initial communities of 4 randomly chosen species each and for each of five species sets, simulate 10 repeats from different initial communities over 50 rounds of selection under each of the propagation methods (B-E).

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communities are randomly chosen to receive or lose species 194 170 (Fig. 1D). We compare each method to a corresponding ran- 195 171 dom control line (e.g. random propagule: PR) where 7 com- 196 172 munities are chosen at random instead of according to their 197 173 score, and to a no-selection control (NS) where every com- 198 174 munity is diluted without selection (Fig. 1E). This last control 199 175 forms a baseline for how communities change due to species 200 176 interactions (23, 26). To achieve statistical power, 5 species 201 177 sets were generated, each with a new set of 15 species. From 202 178 each species set, we then sampled the 21 communities 10 203 179 times to run 10 replicate simulations, which were all sub-204 180 jected to 50 rounds of selection. The same initial conditions 205 181 were used for the different selection methods to allow for a 206 182 fair comparison (Fig. 1F). 207 183

²⁰⁹ Disassembly finds communities whose degradation ²⁰⁹ ranks in the top percentile of all possible communi-²¹⁰

ties. All simulated selection methods succeeded in improv-

ing the median degradation score across the 21 communities 211
between round 0 and 50 (Fig. S1), which is consistent with 212
previous work (3, 23). However, DS was the only propa-213
gation method to significantly and consistently improve the 214
maximum degradation score, meaning that on average, the 215
best community in round 50 degraded significantly better 216
than the best community in round 0 (one-sided Wilcoxon 217

signed rank-test n = 50, 10 repeated runs of 5 species sets, $p < 10^{-9}$ for both IBM and ODE, Fig. 2A, C). The increase in maximum score in DS $(0.22 \pm 0.06, 0.14 \pm 0.08$ for IBM, ODE), was also significantly different from the classical selection methods (-0.03 ± 0.06 and -0.12 ± 0.09 for PS and MS in the IBM, and -0.1 ± 0.08 for PS in the ODE), from its own random control (DR), and from NS (all two-sided Wilcoxon tests of diff. in max. degradation between DS and other methods, n = 50, $p < 10^{-9}$, for IBM and ODE). For comparison, we computed the degradation scores of all $2^{15} - 1 = 32767$ possible communities consisting of 1 up to 15 species for each species set and sorted them from best to worst. The communities found by DS ranked among the best few hundred in both our models, finding the very best community out of 32767 (Fig. 2D, F) in 17 out of 50 runs in the IBM and 23 out of 50 in the ODE. We next investigate what distinguishes these high-ranking communities.

Communities selected by disassembly invest more into degradation and are composed of diverse species with complementary phenotypes. In our model, community performance depends on (a) the overall investment into degradation of toxic compounds relative to growth, and (b) how well community members complement each other. Community members will compete less if they take up dif-

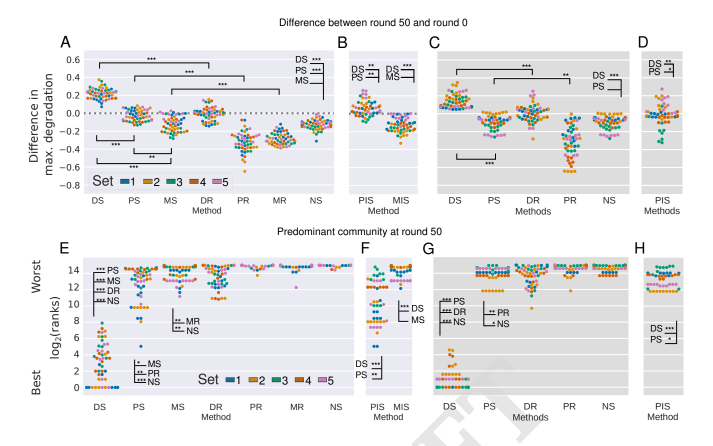


Fig. 2. Degradation scores and ranking of selected communities. Panels A, B, E, F with lighter background show results from the IBM, while panels C, D, G, H with darker background show results from the ODE model. Asterisks show the significance of a Wilcoxon signed-rank test for difference in degradation between methods (*: $p < 10^{-3}$, **: $p < 10^{-6}$, ***: $p < 10^{-6}$, ***: $p < 10^{-9}$). (A-D) The difference in maximum degradation score between round 50 and round 0 over the 21 communities is shown as one dot for each of 10 repeated runs, colored by species set, with 50 dots in total. As each run starts from identical communities for all methods, we have compared pairs of runs between the selection methods. (E-H) The rank of the predominant community (the most common combination of species among the 21 communities in the last round of selection, not counting sub-communities) in terms of its degradation score compared to all of the 32767 possible combinations of 1, 2, ..., 15 ancestral species. As above, each of the 50 dots marks 1 out of 10 repeated runs of 1 out of 5 sets of species.

²¹⁸ ferent nutrients while the degradation score of a community ²³⁷
²¹⁹ can increase if its members specialize on degrading different ²³⁸
²²⁰ toxic compounds (Eq. 1). ²³⁹

To understand how these two properties changed over time, 221 we first quantified the "total investment", i.e. the fraction ²⁴¹ 222 $\sum_{k=1}^{10} f_{ik} < 1$ of resources invested into degradation of all²⁴² 223 toxic compounds k, averaged over the species in each com-²⁴³ 224 munity. Starting from an average investment of 0.5, DS²⁴⁴ 225 finds communities that invest significantly more resources 245 226 into degradation at round 50 than in the first round (one-sided ²⁴⁶ 227 Wilcoxon test of average total investment, all $p < 10^{-9}$, $n = {}^{247}$ 228 50 for both IBM and ODE, Fig. 3A, C). This is not due to any ²⁴⁸ 229 single species with unusually high degradation capabilities,²⁴⁹ 230 but rather because DS finds a combination of species with ²⁵⁰ 231 high investment. The average within-community species di-²⁵¹ 232 versity increases over the 50 rounds (Fig. 3E), which means²⁵² 233 that the communities consist of an increasing number of ²⁵³ 234 species and/or that the communities are increasingly even. 254 235 Accordingly, in DS, the effective number of consumed nu- 255 236

trients and toxic compounds increases over the 50 rounds (Fig. 3F). This increase in coverage and community diversity was not observed for the other selection methods (Fig. 3E, Fig. S2).

Given the complementarity in nutrient uptake and toxic compound degradation, one might expect species to grow and degrade better together compared to when they are alone, as they may be facilitated by other species that degrade compounds that they themselves cannot. We use "synergy" to quantify whether a community property (e.g. degradation) is greater than that of its member species together (Fig. 3G). Against a baseline of all possible species combinations for a given community size – richness in our models increases niche overlap and competition for resources, which decreases synergy – communities selected by DS have significantly higher synergy, for both degradation and cumulative biomass (Kruskal–Wallis *H* test, $p < 10^{-9}$ in either case, Fig. 3G).

In sum, communities selected by DS invest more into degradation compared to communities from other methods. These

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communities are diverse in composition, consist of species 303
with minimal niche overlap, and cover the toxic compounds 304
evenly (Fig. 3F). 305

306 Disassembly can explore more species combinations 259 by diversifying the selected communities. Seeing that $_{_{308}}$ 260 communities selected by DS are diverse and efficient de-261 graders, we now investigate how the method finds these com-262 munities. First, DS explores more species combinations than 263 the other methods (Fig. 4A, B, Fig. S3). The classical propag-264 ule method (PS) can only find sub-communities of the species 265 combinations present in round 0. Similarly, while migrant 266 pool (MS) is in principle able to search all sub-communities 267 of the first set of selected communities, they are in practice 268 limited to a smaller subset as species tend to go extinct due 269 to the toxic compounds, inter-species competition and/or the 270 dilution bottleneck at each round. Accordingly, most com-271 318 munities available for selection by PS or MS resemble one 272 another, seen as a rapid drop in between-community (or beta)³¹⁹ 273 diversity (Fig. 4C, D, Fig. S4). In contrast, changing the 274 321 species composition of some selected communities by insert-275 ing or removing species at random, DS can search a larger ³²² 276 323 number of communities and the resulting drop in beta diver-277 324 sity is not as steep. The beta diversity of the no-selection 278 325 control depends on the diversity of the initial communities. 279

Propagule selection —but not migrant pool— per-327 280 forms better by periodically adding species to se-328 281 lected communities. In the disassembly method, more and 329 282 better communities can be found by randomly adding and 330 283 removing species in some of the communities. To explore 331 284 whether species introduction could improve PS and MS in 332 285 our models (previously shown for PS (23)), we implemented 333 286 two new versions (PIS and MIS), where in each round, a ³³⁴ 287 fixed number of communities chosen at random will re-335 288 ceive one or more "invader" species (also chosen at ran-336 289 dom) with a defined initial population size. With this modi- 337 290 fication, PIS increases the maximum degradation (one-sided 338 291 Wilcoxon signed-rank test of degradation scores in round 50 339 292 versus 0, $p < 10^{-3}$, n = 50, Fig. 2B) and improves upon ³⁴⁰ 293 the standard PS method (two-sided Wilcoxon signed-rank 341 294 test, $p < 10^{-6}$, n = 50) in the IBM. The results are how-³⁴² 295 ever model-dependent. While the PIS method still improved 343 296 upon the PS method in the ODE model (two-sided Wilcoxon 344 297 signed-rank test, $p < 10^{-3}$, n = 50), we did not find any ³⁴⁵ 298 significant improvements in the maximum degradation score 346 299 compared to round 0 (p = 0.9, n = 50, Fig. 2D). Further, ³⁴⁷ 300 PIS finds higher-ranking communities than PS in both the 348 301 IBM (two-sided Wilcoxon signed-rank test for differences in 349 302

ranks between PIS and PS, $p < 10^{-6}$, n = 50, Fig. 2F) and the ODE model ($p < 10^{-3}$, n = 50, Fig. 2H) over the 50 rounds. PIS can explore more combinations than the regular PS, and the initial drop in beta diversity is less severe in both models (Fig. 4A-D), indicating that there is more variability for selection to act on. In contrast, MIS does not improve significantly on MS, either in terms of degradation, ranks or investment. Even though MIS explores more species combinations than MS, the beta diversity rapidly drops (Fig. 4C), and the introduced species do not contribute much to diversity or degradation of the resulting communities.

Mutation and selection can decrease per-species investment, but this increases biomass, maintaining community degradation. We have shown that DS can improve degradation by exploring many different species combinations and find ones that rank highly. Shuffling species around is, however, not the only way to improve degradation scores. Our models allow for mutations to the parameter f_{ik} that determines the trade-off between investment into degradation and biomass production for a cell. If a mutant is more competitive than its parent, it can replace the original type in future rounds, even as other species come and go around it. To investigate the effect of mutations, we now compare the investment into degradation of species at round 50 to that of their ancestors from round 0, and analyze how these changes affect degradation at the community level.

In DS, the total per-species investment $\sum_{k=1}^{10} f_{ik}$ into degradation was significantly lower after 50 rounds of selection than that of the corresponding ancestral species (one-sided Wilcoxon signed-rank test of total investment in initial vs final round of selection, $p < 10^{-6}$, n = 50, Fig. 5A, Fig. S5). Given the trade-off between investing into growth versus degradation, the communities made up of evolved species had greater total biomass than communities composed of the corresponding ancestral species (one-sided Wilcoxon signedrank test of total AUC in communities, initial vs final round, $p < 10^{-9}$, n = 50, Fig. 5B), such that overall, the degradation of the evolved communities was marginally but significantly higher $(+6 \times 10^{-3} \text{ units}, \text{ averaged over all species sets}, \text{ one$ sided Wilcoxon signed-rank test $p < 10^{-3}$, n = 50) than that of communities made up of their ancestors (Fig. 5C, Fig. S6). Compared to the improvement in degradation due to finding better species combinations, the improvement due to species evolution is very small and is not likely to have a large effect on the outcome of selection.

In summary, the disassembly method improved the degradation scores over the 50 rounds of selection by finding better

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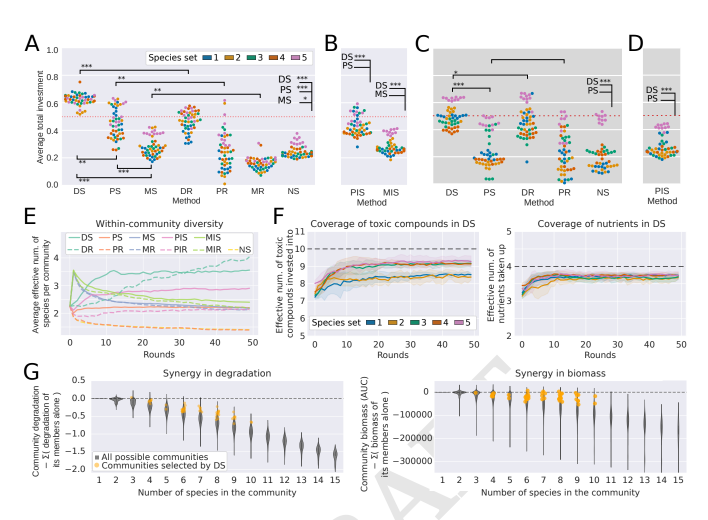


Fig. 3. Total investment into degradation, diversity, coverage and synergy in degradation. Data from the IBM shown on lighter background everywhere except for panels CD, which has data from the ODE model on darker background. **(A-D)** Average total investment in a community at round 50, averaged for the 21 communities in a run. For each species we calculate the average investment weighted by strain population and then we do the unweighted average of all the species in the community. The red dotted line at 0.5 indicates the theoretical mean investment at round 0. IBM results in panels A, B and from the ODE in C and D. One dot for each out of 10 repeated runs, colored by the 5 species sets, with 50 dots in total. The asterisks indicate the results of a Wilcoxon signed-rank test with n = 50 (*: $p < 10^{-3}$, **: $p < 10^{-6}$, ***: $p < 10^{-9}$). **(E)** Within-community species diversity measured as the effective number of species Eq. (13), averaged over all 21 communities in a run. The line shows the average over the 10 repeats and the 5 sets of species, with error bars per set of species in Fig. S2. **(F)** The coverage of toxic compounds and nutrients in communities selected by DS, measured as the effective number invested into (f_{ik}) or taken up (n_{ij}) respectively within a community (mean \pm s.d. over the 10 repeated runs for a given set of species), Table 1 (see Methods). Results from the IBM. **(G)** Synergy in communities selected by DS at round 50, grouped by community richness. Synergy is the difference in degradation scores (left panel) or biomass (right) between a co-culture and the sum of the values of the corresponding monocultures. The violin plot shows the distribution of synergy for each possible community of that richness level after one round of growth. The dots show the average synergy per repeated run in the last round for the 5 species sets. The average species richness per repeated run is rounded to obtain an exact value. For visibility, we have plotted all species sets in the same

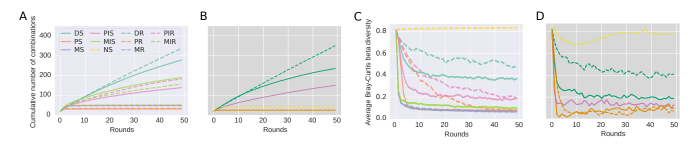


Fig. 4. Cumulative number of communities found by the selection methods and between-community diversity explain how DS can find better communities. We show the mean over the 10 repeats and 5 species sets for each propagation method and refer to Fig. S3-S4 for the full results. Results for the IBM and ODE model are shown against a light or a dark background, respectively. (A, B) Time series of cumulative number of unique communities for each selection method. (C, D) Between-community or beta diversity, calculated as the average Bray–Curtis distance of each pair of communities.

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species combinations. Within those communities, individual ³⁹⁷
species evolved to invest less into degradation and more into ³⁹⁸
biomass production. As an effect of the trade-off between ³⁹⁹
degradation and growth, the communities still maintain their ⁴⁰⁰
degradation capabilities and the most efficient communities ⁴⁰¹
are the species combinations found in round 50, composed of ⁴⁰²
either their ancestral or evolved genotypes. ⁴⁰³

Communities selected by DS are less stable than 357 those selected by PS and MS. The disassembly method 358 has features to ensure heritability and promote within-359 community diversity: we re-inoculate species in fixed and 360 equal abundances, punish extinctions and re-inoculate ex-361 tinct species. Controlling the ecological dynamics so tightly $_{_{410}}$ 362 means that if we were to simply transfer these communities 363 without adjusting relative abundances and without selection, $_{_{412}}$ 364 as in the no-selection treatment, they could drift towards a $_{\scriptscriptstyle 413}$ 365 different equilibrium with a lower degradation score. To as-366 sess the ecological stability of the selected communities, we $_{_{415}}$ 367 transferred the communities from round 50 for an additional $_{_{416}}$ 368 25 rounds of growth and dilution, this time without selec-369 tion (Fig. 6A) and found that the degradation scores of com-370 munities selected by DS dropped by -0.21 ± 0.14 on aver-371 age when left to their natural dynamics, close to how much $_{\rm _{420}}$ 372 the selection method increased the degradation (0.22 \pm 0.06). 373 This indicates that the high performance of these commu-374 nities relied on controlling the ecological dynamics. This 375 means that the communities converge, once ecologically sta-376 ble, to a degradation score that is not significantly different to 377 the average of the initial communities (one-sided Wilcoxon 378 signed-rank test, p = 0.24, n = 50, Fig. 2A, B). 379 427 In contrast, the degradation does not drop as much in commu-380 nities selected by the classical methods PS and MS ($-0.02 \pm$ 381 0.03 and -0.03 ± 0.03 in max degradation, respectively, $_{_{\rm 429}}$ 382 Fig. 6A). The methods are stable in the sense that the com-383 munities do not change much after the first few rounds of 430 384 selection, either in terms of composition (Fig. 3E) or degra-431 385 dation (Fig. S7). The methods with invasion, PIS and MIS, 432 386 show an intermediate drop in degradation (-0.07 ± 0.07 and 433 387 -0.07 ± 0.04) indicating that the invasion step has an effect ⁴³⁴ 388 on community stability. In order to remain effective, the com-435 389 munities found by DS should be grown in the same condi-436 390 tions as they were selected, i.e. (i) from equal abundance, (ii) 437 391 without any intermediate rounds of growth in between rounds 438 392 of selection. The latter has been suggested to stabilize the dy-439 393

³⁹⁴ namics and improve the community selection (23).

395 Varying experimental parameters to decrease the size 442

³⁹⁶ of the experiment. Our model shows that DS can outper- ⁴⁴³

form other propagation methods, as long as the ecological dynamics of the communities are controlled. However, DS is more cumbersome than the other methods from an experimental perspective: constantly dis- and re-assembling communities and having to adjust the population sizes of each species at every round could cost a lot of time and resources. We now investigate how four experimental parameters impact the degradation scores in DS, and affect experiment size. We focus on DS but also compare it to the other methods (Fig. 6B-E, Fig. S8-S11).

The parameter with the strongest effect on experiment size and the maximum degradation score is the number of species in the initial set (Spearman's rank correlation coefficient $\rho = 0.67, p < 10^{-9}$, Fig. S9). This means that the metacommunity needs to be as rich as possible to efficiently improve degradation, and the main effort should be invested into managing a larger number of species, ideally by adding species that have positive effects on degradation or the growth of others. In contrast, the number of communities clearly affects experiment size, but it had a weaker correlation to degradation for DS ($\rho = 0.35$, $p < 10^{-9}$), meaning that the number of communities could be decreased, which would reduce effort with a limited effect on community performance. Next, we turn to two parameters that affect the degradation scores but not size of the experiment. The number of communities receiving an invading species is negatively correlated with degradation ($\rho = -0.29$, $p < 10^{-9}$). Introducing species to a smaller number of communities should improve the final degradation score (Fig. S8). Finally, the dilution factor (i.e. how large a fraction of the culture to re-inoculate for the next round of growth) is positively correlated to degradation scores ($\rho = 0.52, p < 10^{-9}$).

Discussion

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The major challenges for community breeding are ensuring (i) that the community function is heritable, (ii) that withincommunity selection does not dominate over betweencommunity selection and (iii) that communities differ sufficiently in phenotype. While other theoretical studies have investigated heritability and the balance between within- and between-community selection (6, 21–23), our "disassembly" method contributes to improving the third point: how to maintain variability between communities.

We have shown that disassembly can improve significantly upon the maximum degradation scores of simulated synthetic communities, compared to a random line and a noselection control. The method further outperformed the classical propagule and migrant pool methods, which could only

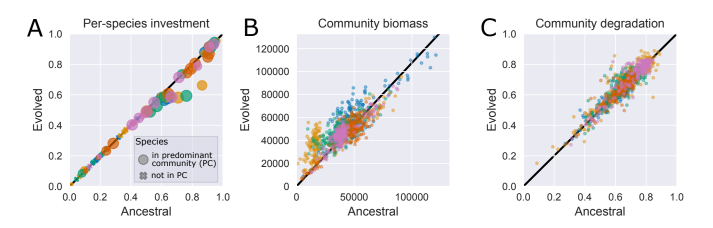


Fig. 5. Change in degradation investment per species and the effect of such change in the biomass and degradation of the communities in the last round, colored by species sets. All results from the IBM. **(A)** Total investment $f_i = \sum_k f_{ik}$ for each species, i = 1, 2, ..., 15. There are 75 dots: 15 species, colored by the species sets 1-5. For evolved condition, the markers show the average total investment of the species, weighted by population size for each occurrence in communities from the last round of selection where the species is present and not weighted between repeated runs; while ancestral condition corresponds to the total investment of each ancestral species prior to any growth or evolution. Species that were present in a predominant community (P.C., see definition in the caption of Fig. 2) in the last round are shown as circles, where the radius is proportional to the number of repeated runs where the species appear in a P.C. Species that were never present in a P.C. are represented by crosses. We summarize the p-values of Wilcoxon tests of whether the investment is different in the last round of selection compared to the first in Fig. **S5. (B)** Total biomass per community, measured as the sum of the area under the growth curves (AUC) for species in the community. The initial AUC is calculated from one round of growth, where the community is composed of ancestral strains of the same species in the same proportions as the last community. There are 1050 dots: 21 communities per 10 repeated runs, for each of the 5 species sets. **(C)** Degradation scores of the same 1050 communities in (B). The degradation of the initial community is calculated over one round of growth when the community is composed of ancestral strains of the same species.

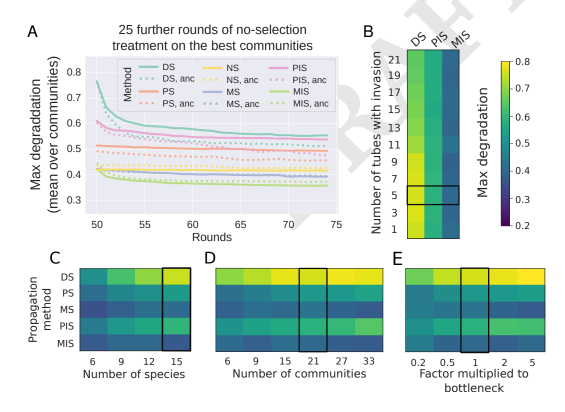


Fig. 6. (A) Stability in degradation over 25 additional rounds after releasing the selection pressure, calculated for the highest-scoring community for each selection method in the IBM. Results show the average degradation for each repeat. The dotted lines show the corresponding degradation when the community is composed of ancestral species. (B-E) Max. degradation score in the last round of selection as a function of the experimental parameters: (B) the number of tubes to receive or lose a species, (C) the number of species in the initial set, (D) the number of tubes or communities and (E) a scaling factor for the dilution bottleneck. The heat maps show the median degradation score over all species sets and repeated runs (in (C) also over sub-samples) for each selection method. Fig. S9-S11 show the full data set. The color bar is the same for panels B-E. The black outline marks the parameter value used throughout the rest of the paper. All data from the IBM.

improve the maximum function for some initial combination 492
 of species, confirming previous findings (3, 23).

The problem with the classical methods is that they rapidly 494 446 lose between-community diversity, which selection acts upon 495 447 to improve community function. In these methods, diversity 496 448 only arises through mutations, or through loss of species due 497 449 to competition and dilutions between rounds of growth, while 498 450 directed selection reduces between-community variability by 499 451 only propagating a small subset of high-performing commu- 500 452 nities. When communities become increasingly similar, fit- 501 453 ness differences become increasingly random which makes 502 454 selection less effective (23). By removing and introducing 503 455 species, the disassembly method reshuffles the species com- 504 456 position to access new communities that sometimes outper- 505 457 form the original best community. As proposed in (23), we 506 458 show that the classical methods can be improved by periodi- 507 459 cally invading them with new species, which allows them to 508 460 maintain some variability. This approach still under-performs 509 461 compared to disassembly, however. 510 462

In a sense, the disassembly method is inspired by the 511 463 crossover operator used in genetic algorithms in the field of 512 464 evolutionary computing (27–29), whereby building blocks 513 465 are recombined between digital individuals and generate vari- 514 466 ability for selection to act on. Of course, crossover in ge- 515 467 netic algorithms is itself inspired by recombination in sex- 516 468 ual organisms (30, 31). While interesting, these parallels do ₅₁₇ 469 not map directly to species exchange in community breeding, 518 470 where the units that are subject to exchange are well-defined 519 471 and have their own ecology and within-species evolution. 520 472 Next, heritability is crucial for evolution (32) and a major 521 473 challenge for community selection (6). In the disassembly 522 474 method, as in (6), we sidestep the issue of heritability and 523 475 ecological stability by re-assembling communities in a fixed 524 476 abundance and equal species proportions. In our models, this 525 477 allows community dynamics to unfold in almost the same 526 478 way after each transfer. The disadvantage of this approach is 527 479 that we cannot guarantee that communities selected by disas- 528 480 sembly would maintain either their community composition 529 481 or function when propagated by regular dilution. One way to 530 482 overcome this would be to include a few rounds of transfer-531 483 ring without selection to allow the communities to equilibrate 532 484 before each selection step (23). This would, however, make 533 485 disassembly quite inefficient, in which case, propagule with 534 486 invasion can be more stable than disassembly (Fig. 6A). 487 535 Another factor that we do not explore in detail here is the 536 488 relationship between community stability and the timing of 537 489 selection. While our model is robust to timing errors, a com- 538 490 munity that degrades the toxic compounds too quickly can in 539 491

principle be invaded by cheater cells that profit from the initial degradation and out-compete degraders in the absence of toxic compounds (6). Such ecological succession was found in a chitin-degrading community (16, 33).

While we have focused on increasing community performance from an applied perspective, our study provides important insights into discussions on the levels of selection and whether group selection can dominate over individual-level selection. Our choice to implement mutations (unlike other theoretical studies (23)) means that selection will also act at the level of individual genotypes. We found that competition between individuals is indeed strong in our simulated communities, leading individual species to reduce their investment into degradation in favor of growth. Interestingly, though, the faster growth rate of these evolved species compensates for the reduced investment by producing larger population sizes of degraders, such that the difference in degradation between ancestral and evolved communities was negligible. Our approach does not overcome within-community selection for growth, but this may not impact community function as much as one might expect when the community function is indirectly coupled to growth. Going back to the practical perspective, in a lab setting, it may be worth investigating whether the best selected species composition is more efficient when assembled from ancestral or evolved strains.

Ultimately, the goal of our investigation is to help design experiments rather than just computer simulations. While we found that our method can efficiently search for better species combinations, its scope is limited to communities that can be disassembled in the lab, and the effort needed to find out how to isolate species from the communities should not be underestimated. We have, however, confirmed that a periodic introduction of species (which is possible independently of whether we can remove species or disassemble the communities) improves the propagule method (23) to balance experimental feasibility and improvement of community functions. From an experimenter's perspective, it is also useful to consider where one can reduce the size and complexity of the experiment. We find that the parameter with the biggest effect on the method is the size of the initial species set and that the disassembly method is more efficient for larger sets of species (Fig. 6). Disassembling rich communities may however prove quite challenging in practice, and future experimental work will aim to make this step more efficient. We also find that for disassembly, a higher number of communities grown in each selection round can compensate for the number of selection rounds needed since we are then able to search more distinct communities per round of selection.

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We have made a number of assumptions for our models. 587 540 First, we assume a well-mixed liquid culture, where in re- 588 541 ality, clumps may affect species interactions and community 589 542 function. We also assumed a trade-off between degradation 590 543 and growth in both models and that the toxic compounds can- 591 544 not be used as nutrients. Both assumptions serve to decou- 592 545 ple the community phenotype from population growth and 593 546 while they are not completely independent, a smaller con- 594 547 tribution from growth on the community phenotype should 595 548 make artificial selection more difficult. In a sense, this is 596 549 the more interesting problem to explore, since all the meth- 597 550 ods we explored are expected to improve a community phe-598 551 notype that is aligned with population growth. Further, we 599 552 have assumed that species diversity is key to functional di- 600 553 versity: each species can only degrade a subset of the toxic 601 554 compounds in our model and complete removal of the com- 602 555 pounds depends on finding other species with complemen- 603 556 tary degradation capabilities. Within-community diversity is 604 557 in this way fundamental for community success, and also de- 605 558 creases competition, as each species only uses a subset of the 606 559 nutrients. In our simulations, it is therefore unlikely that any 607 560 mono-culture scores higher than a multi-species community, 608 561 and the median size of a selected community was $6.9 \pm 1.8_{\text{ 609}}$ 562 species after 50 rounds of selection by disassembly (in the 610 563 IBM model). This optimum will, however, differ for each 611 564 system (25). 565

Taken together, we have introduced a new approach to com-566 munity selection, where species composition is shuffled be-567 tween competing communities, allowing for a greater explo-568 ration of the space of possible communities to find the best 569 performing ones. In doing so, we have been able to im-570 prove community function with respect to randomly assem-571 bled communities, but show that genetic mutation can con-615 572 tribute to reduced investment by individual strains into com-573 munity function. We are testing this approach experimentally 574 in parallel work. 575

576 Materials and methods

Growth and community function. We separately imple-577 ment an individual-based model (IBM) of individual cell 578 growth and a system of ordinary differential equations 623 579 (ODEs) that model population-level dynamics. Both mod-624 580 els simulate different microbial species growing together in 625 581 batch culture, in a medium containing 4 types of nutrients 626 582 that allow cells to grow, and 10 toxic compounds that cause 627 583 cell death. Species are described by how quickly they grow, 628 584 the rate at which they convert nutrients into biomass and how 629 585 they are affected by toxic compounds. We describe the details 630 586

of both models in the following sections. All interspecies interactions are due to the consumption of nutrients and degradation of the toxic compounds. The concentration of each nutrient j = 1, 2, 3, 4 is denoted by N_j and the concentration of each toxic compound k = 1, 2, ..., 10 is denoted by T_k . The removal of toxic compounds in either model is determined by the parameters f_{ik} that denote the fraction of resources that a cell (IBM) or population (ODE) of type *i* invests into the degradation of toxic compound *k*. The remaining fraction $1 - \sum_k f_{ik}$ is invested into growth. As we describe in the following section, f_{ik} is the only parameter that is subject to mutation and therefore gives rise to different strains of the same species.

Communities are created in two steps: First we draw a library of 15 species, each with unique (random) combinations of model parameters sampled as described in Table 1 and 2. These species are then randomly assigned (with replacement) to 21 communities of four species each, such that each species is present in at least one community.

In each round, each community grows and degrades the toxic compounds independently of the other communities, for 80 time-steps in the IBM and 100 in the ODE model. The root-mean-square decrease in T_k from the initial time point t_0 to the last t_{end} forms the basis of our community function, the degradation score D:

$$D = 1 - \sqrt{\frac{1}{10} \sum_{k=1}^{10} \left(\frac{T_k(t_{end})}{T_k(t_0)}\right)^2}.$$
 (1)

These scores are used to rank the communities, so that we can propagate the best ones by the different selection methods (elaborated below). To compare the selection methods, we simulate 50 rounds of growth, degradation and selection with the different methods. Upon propagating communities to fresh medium, we only transfer the cells and no leftover media. To reduce bias due to initial conditions, we run 10 simulations with different initial communities for each out of 5 sets of 15 species (Fig. 1F). The same exact set of initial conditions is used for all selection methods to make comparisons fair.

Within-species evolution. In both models, a strain *i* of the ancestral species l_i ("I" as in lineage) invests a fraction $0 \le f_{ik} \le 1$ of the resources that they take up into degradation of toxic compound *k*. These fractions can mutate, creating new strains of the same species that invest different amounts into degradation and the rest, $1 - \sum_k f_{ik}$, into population growth. The mutations take place at cell division in the IBM and during community replication in the ODE. The dis-

tinction between "species" of a fixed set of 15 and "strains", 673
that emerge from an ancestral strain of each species as mu- 674
tations arise, is important. For example, in the disassembly 675
method described below, we inoculate a fixed concentration 676
of each *species* in each round, which can consist of different 677 *strains* of the species, depending on their frequency in the 678
previous round.

To evaluate the evolution of the total investment $f_i = \sum_k f_{ik}$ 680 (Fig. 5A) of species l, we compare the investment of the an- 681 cestral strain i to that in the last round of selection (or as late 682 as possible in case the species went extinct). The average 683 per-species investment is weighted by the population size S_i 684 of the different strains i of species l,

$$\tilde{f}_{l.} = \sum_{i \text{ of species } l} S_i f_{i.} / \sum_i S_i$$
(2)

to emphasize the investment of abundant strains instead of small recent mutants that have not contributed as much to the community function. The value \tilde{f}_l is further averaged over the 10 repeated runs (recall: each run starts from the same set of species) for each of the 15 species, without weighting by population size. For the statistical comparison, we hence have 75 data points: 15 species per each of the 5 sets.

Individual-based model (IBM). We simulate well-mixed 693 651 batch cultures that initially contain 10 cells per species in 694 652 the communities. All cells are initially equally exposed to 695 653 $T_k(t_0) = 700$ units of each of the 10 toxic compounds, which ⁶⁹⁶ 654 can cause cell death (see below), and have equal access to 697 655 $N_i(t_0) = 2000$ units of each of the 4 nutrients that allow the ⁶⁹⁸ 656 cells to divide and reduce the concentrations of toxic com-699 657 pounds (see below). These processes occur according to the 700 658 parameters of each strain *i* (Alg. 2, Table 1, Fig. S12). 701 659 Cell division consists of two steps, here called "activation" 702 660 and "replication", that respectively involve the acquisition of 703 661 some nutrients by the cell and their utilization for cell repro-704 662 duction (34). Cells of strain *i* share their parameters (Table 1) 705 663 and can be in either of two states: the initial "inactivated" 706 664 with population size p_{i0} , or the "activated" with size p_{i1} . Ev-⁷⁰⁷ 665 ery activation, replication, mutation and death event occurs 708 666 with a given probability by random sampling from a Poisson 709 667 distribution with rate $population \cdot probability$ (22). 710 668 Degradation and activation are costly and are carried out first. 711 669

670 At each time step, a cell of type i can take up an amount n_{ij} 712

of the nutrient j with current concentration N_j , with

$$max_uptake = \sum_{j} (n_{ij} \text{ if } N_j > n_{ij}).$$
 (3) ₇₁₅

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⁶⁷² The *max_uptake* scales down the amount of degradation ⁷¹⁷

or the probability to activate when not all the nutrients consumed by the cell are present.

Regardless of their activation state, at each time step, all the cells of a strain (i.e. the total population size $S_i = p_{i0} + p_{i1}$) degrade the amount $f_{ik} \cdot max_uptake$ units of each toxic compound k, consuming f_i . units of nutrients in total. If there are not enough nutrients, a smaller fraction of S_i degrades, determined by the amount of nutrients available. When a toxic compound is depleted, its degradation and thus the corresponding nutrient consumption does not occur.

Following degradation, an inactive cell can activate with probability

$$a_i \cdot (1 - \sum_k f_{ik}) \cdot max_uptake \cdot \sum_j \left(\hat{n}_{ij} \frac{N_j(t)}{N_j(t_0)} \right), \quad \textbf{(4)}$$

where \hat{n}_{ij} is a re-scaled version of n_{ij} : if some nutrients are depleted, their n_{ij} are set to 0 and the n_{ij} of the remaining nutrients are re-scaled such that $\sum_j n_{ij} = 1$. To activate, a cell of type *i* consumes in total $1 - f_i$. units of nutrients and if the it does not replicate, it needs the same amount of nutrients in subsequent time steps to remain activated.

The amount of nutrient of type j that is consumed for degradation and activation depends on the parameters n_{ij} and on the amount of each nutrient that is available. When one type of nutrient gets depleted, cells will take up more of the other available nutrients that they require. At every time step we check how many cells of strain i can degrade and activate based on the scarcest nutrient. Thus, when a nutrient jis nearly depleted, fewer cells that require this nutrient degrade and have the chance to activate in the current time step. In following time steps, when nutrient j is depleted, cells consume the remaining required nutrient types. So, although now cells would use nutrients less efficiently, if these nutrients are sufficiently abundant, a greater population can degrade and activate. This models a pause in degradation and division caused by a metabolic shift towards consuming fewer nutrients. These events are stochastic and lead to noise between runs of the model with the same starting conditions. When we calculate the growth and degradation of specific communities such as the 32767 possible combinations of species (Fig. 2 E-F, Fig. 3 G), we average the results over three replicates of the simulations. We use the same seed for the random number generators for consistency.

At each time step, activated cells divide with probability $r_i \cdot (1 - f_{i\cdot})$ without additional cost, resulting in two inactivated daughter cells: one daughter maintains the parameter values, and the other is susceptible to mutation with probability $\mu = 0.01$. Upon mutation, the previous value of at least one f_{ik}

Parameter	Description	Randomly sampled from
l_i	Species ID of strain <i>i</i>	
a_i	Activation probability	Beta(2,2)
r_i	Replication probability	Beta(2,2)
n_{ij}	Consumption rate of nutrient j	Uni(0,1), Sparse,
		Rescaled so that
		$\sum_{j} n_{ij} = 1$
f_{ik}	Fraction of consumed nutrients invested	Uni(0,1), Sparse,
	into degradation of toxic compound k	Rescaled so that
		$\sum_{k} f_{ik} = Uni(0,1)$
m_{ik}	Death rate of strain i due to toxic compound k	Uni(0.001,0.02), Sparse

Table 1. Parameters defining a microbial strain *i* in the IBM. Growth rates, death rates and degradation investment vectors r_{ij} , m_{ik} and f_{ik} are made sparse by multiplying them by a vector drawn from Bernoulli(0.5). Each species can this way only take up a random fraction of nutrients, be affected by a random fraction of the toxic compounds and degrade another fraction of the toxic compounds. Despite changing by mutation, the total investment f_i . is limited to the interval [0, 1].

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is multiplied by a random number from the lognormal ($\mu =$

719 $0, \sigma^2 = 0.4$) distribution, making sure the total investment f_i .

falls in the [0, 1] interval. As a result, a new strain of the

same species with population $p_{i0} = 1$ is introduced.

722 At each time step, activated and inactivated cells may die with

⁷²³ a probability determined by the following Hill function:

$$\sum_{k} m_{ik} \frac{T_k^2}{T_k^2 + K^2} \tag{5}_{727}^{726}$$

Where T_k is the current concentration of toxic compound k^{729} and the constant K = 700.

Population-level model. As the IBM above, the ODE ⁷³³ model simulates well-mixed batch cultures with nutrients and ⁷³⁴ toxic compounds, extending a previous model (25). In the ⁷³⁵ model, the population size S_i of each strain *i* in a community ⁷³⁶ grows in relation to the concentrations of nutrients N_j and ⁷³⁷ decline by the toxic compounds T_k (Fig. S13) by the model ⁷³⁸ parameters in Table 2. Growth, death, nutrient uptake and ⁷³⁹ degradation is described by the following ODE system: ⁷⁴⁰

$$\frac{dS_i}{dt} = \left((1 - \sum_k f_{ik})\rho_i(\mathbf{N}) - \mu_i(\mathbf{T}) \right) S_i \tag{6}$$

$$\frac{dT_k}{dt} = -T_k \sum_i f_{ik} \delta_i \rho_i(\mathbf{N}) S_i$$
(8)⁷⁴⁷
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The bold-face N, T denote the vectors of all nutrients and $_{750}$ toxic compounds, respectively. We assume Monod and Hill $_{751}$

functions for the per-capita growth and death rates ρ_i , μ_i .

$$\rho_i(\mathbf{N}) = \sum_j r_{ij} \frac{N_j}{N_j + K_N} \tag{9}$$

$$\mu_i(\mathbf{T}) = \sum_k m_{ik} \frac{T_k^2}{T_k^2 + K_T^2}$$
(10)

The system of equations Eq. (6)–Eq. (8) is solved with a standard ODE solver (dopri5, (35, 36)) for 100 time steps with initial conditions $S_i(t_0) = 100$, $N_j(t_0) = 100$ and $T_k(t_0) =$ 100 for all i, j, k.

The investment f_{ik} can mutate to form different strains of the same species. When this happens, we add a new population equation of the type Eq. (6) to the ODE system, with the same parameters r_{ij} , m_{ik} , Y_i and d_i as the ancestor but with the modified f_{ik} . To not make the system of equations too large, we have limited the number of strains to 28 per community. We estimate this to be enough since we expect mutants to rapidly replace their ancestral strains if their growth rate is higher, and otherwise disappear rapidly. If there are already 28 strains in a community, then no more mutants are allowed. Otherwise, when communities are propagated to the next round of growth, any surviving strain can have a mutant with probability 0.05. Having chosen which strains i to mutate, we pick one or more traits f_{ik} at random and multiply them by numbers drawn at random from lognormal(0, 0.4)and ensure that both the mutated traits f_{ik} and the total investment f_i . falls in the [0, 1] interval. The mutant receives the same r_{ii} , m_{ik} , Y_i and d_i parameters as its ancestor and is introduced with population size 100, the same as the initial population before the first round of growth. This population size is chosen relatively high, in order to speed up the competition between ancestor and mutant strain.

Parameter	Description	Sampled from
l_i	Species ID of strain <i>i</i>	
r_{ij}	Maximum growth rate	Uni(0.01, 0.1)
	with respect to nutrient j	Sparse
K_N	Half-saturation constant	$K_N = 10$
	for nutrients	(fixed)
m_{ik}	Maximum death rate	Uni $(10^{-4}, 10^{-3})$
	with respect to toxic compound k	Sparse
K_T	Half-saturation constant	$K_T = 10$
	for toxic compounds	(fixed)
f_{ik}	Fraction of amassed nutrients	Uni(0,1), Sparse
	that are invested into	Rescaled so that
	degradation of toxic compound k	$\sum_k f_{ik} = \operatorname{Uni}(0,1)$
Y_i	(Average) biomass yield	$\log \operatorname{lognormal}(\log(10^{-3}), \log(5))$
	with respect to the nutrients	
δ_i	(Average) degradation efficiency	$\log \operatorname{normal}(\log(10^{-4}), \log(5))$
	with respect to the toxic compounds	

Table 2. Parameters defining a microbial strain *i* in the ODE model. Growth and degradation parameters in relation to nutrients and toxic compounds N_j and T_k . All parameters are assumed to be positive, and the investment f_{ik} is limited to the interval [0, 1]. The matrices of growth rates, death rates and degradation investment r_{ij} , m_{ik} and f_{ik} are made sparse by multiplying them by matrices drawn from Bernoulli(0.5), i.e. flipping a coin for each entry. In this way, each species takes up approximately half of the nutrients, is affected by half of the toxic compounds and degrades half of the toxic compounds.

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Artificial selection methods. After scoring all communi-777 752 ties in a given round (see above), a fraction of these "par-778 753 ent" communities is propagated to "offspring" for the next 779 754 round of growth (Fig. 1A, Alg. 1). Here we implement 780 755 several methods of propagating the selected communities 781 756 as described below (Fig. 1B, C), and compare them to a 782 757 no-selection control (NS) where each community is propa-783 758 gated by 100-fold or approximately 20-fold (stochastic pro-784 759 cess based on Poisson distribution) dilution, respectively for 785 760 the ODE model or IBM (Fig. 1E). NS shows the baseline 761 change in community function due to interspecies interac-762 786

⁷⁶³ tions and changes to the species composition (23).

Propagule selection. In the propagule method, the 7 commu-789 764 nities with the highest scores are propagated to the next round 790 765 by dilution (10, 24) (Fig. 1C, Alg. 7, Alg. 8). The commu-791 766 nities are populated uniformly such that each selected par-792 767 ent contributes 3 offspring communities for the next round. 793 768 The important design parameters are the dilution factor and 794 769 the fraction of communities to propagate, i.e. the selection 795 770 bottleneck. In previous experimental studies, dilution factors 796 771 between 5 and 30 have been used for bottlenecks between 797 772 1/10 and 1/3 of parent communities (8–10, 17, 19). We keep 798 773 a wide bottleneck, selecting 7 out of 21 communities, before 799 774 diluting them by a factor 100 (ODE model) or approximately 800 775 20 in the IBM (where population sizes are smaller and we 801 776

sample the cells to propagate at random, according to a Poisson distribution) for the next round. See also simulation studies in (5, 6, 22, 23, 37). We compare PS to the random control PR, where the communities are selected at random without regards to their degradation scores. We also compare PS to a version that we call propagule with invasion (PIS) and its corresponding random control (PIR). In this version, we introduce at least one species to 5 out of the 21 offspring communities (chosen at random with uniform probability) (23).

Migrant pool selection. Here, selected parent communities are mixed in a *migrant pool* before new offspring communities are formed by taking samples from the pool (10, 24) (Fig. 1D, Alg. 9). Previous experiments have used microbial communities from wastewater (18), soil and rhizospheres (8, 10–14), marine environments (16) and other strain collections (38), selecting between 1/10 and 2/7 of communities and diluting them by factors between 1/100 and 1/2. The method has also been subject to at least one simulation study (5). We select 7 communities out of 21, merge them in a pool and create 21 new communities by sampling without replacement cells from the pool with an approximately 20-fold dilution (stochastic process according to Poisson distribution). We compare MS to a random control (MR), where we select communities with uniform probability without regards to their degradation scores. We also implement a version of mi-

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grant pool selection where we introduce one or more species 844 802 to 5 out of the 21 offspring communities (chosen at random 845 803 with uniform probability), and call this migrant pool with in-846 804

vasion (MIS and the random control MIR). 805

Disassembly selection. Our proposed method is intended for 806 synthetic communities, where each species can be grown sep- $_{_{850}}$ 807 arately and isolated from a multispecies community, such that 808 the communities can be disassembled between transfers. We 809 select 7 out of 21 parent communities by degradation scores 810 (Fig. 1D, Alg. 10). By disassembling these communities, 854 811 we maintain a record of samples of each species that were 812 present in at least one selected community in each round. If 813 a species is present in more than one selected community, we 856 814 sample from the highest-scoring community that this species 857 815 was part of. In this way, we are able to re-introduce any 858 816 species that went extinct. 817 859

To select against extinctions and communities whose mem-860 818 bers out-compete one another, we scale the degradation score 861

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D by the fraction of surviving species at the end of a round 862 820 of growth as follows:

$$\hat{D} = D \times \frac{\text{number of surviving species}}{\text{number of species in the community}}.$$
 (11)

For example, if a 5-species community loses one member 822

species, its degradation score is scaled by 0.8. Next, we draw 865 823

21 offspring communities from the 7 (n = 1, ..., 7) selected 824

parent communities for the next round of growth, in propor-825

tion to their scores \hat{D} with probability: 826

$$\frac{1}{\sum_k \hat{D}_k} \hat{D}_n,$$
 (12) 868

for each selected community. In this way, parent communi- 870 827 ties with (i) high degradation scores and (ii) low or no extinc-871 828 tions will have more offspring. The offspring communities 872 829 have the same species composition as their parents, but we 873 830 reset the initial abundance of each species to a specified pop-874 831 ulation size (100 in the ODE, around 10 cells in the IBM by 875 832 a random sampling with replacement from a Poisson distri-876 833 bution) to standardize the growth conditions between rounds, 877 834 i.e. to maintain heritability. 835

To introduce variability between communities, we change the 879 836 species composition of a few of the 21 offspring communi-880 837 ties. First, we choose 5 offspring communities at random 881 838 and remove one or more species, always one species plus an 882 839 additional number drawn from Poisson(0.5). If the drawn 883 840 number is equal to or higher than the number of species cur- 884 841 rently in the community, we leave one species to avoid emp-885 842 tying or completely changing the community composition. 886 843

Having found a number of species to remove, we choose the species to remove with uniform probability, but avoid removing any species that is present in only this community. Next, we introduce one or more invader species -as above, 1 + Poisson(0.5) — chosen with uniform probability from the frozen stock, to 5 randomly chosen communities. These 5 are chosen anew and could be the same communities that we just removed species from, or not. In order to maintain diversity, we ensure that all species appear in at least one community by preferentially introducing species that are not currently present in any offspring community. See Alg. 10 for more details.

Statistical and other analyses. Correlations are evaluated by the Spearman's rank correlation coefficient ρ . We compare selected communities to the set of all possible communities by a Kruskal-Wallis H test for differences in median. We use the scipy (36) implementations for all three methods. We quantify species diversity within a community (Fig. 3E, Fig. S2) as the Hill number of order 1 or the average effective number of species present in the community (39, 40), which is based on the Shannon index H':

$$\exp(H') = \exp(-\sum_{l=1} p_l \log p_l), \tag{13}$$

which in turn depends on the species' relative abundances

$$p_l = S_l / S_{tot} \tag{14}$$

where we divide the population size S_l of each species by the total population size in the community $S_{tot} = \sum_l S_l$. If more than one strain is present, we sum up their population sizes to find the species' total population size. We then average $\exp(H')$ over all communities to find the average effective number of species. The measure falls between 0 and 15 effective species in an average community.

Beta diversity (Fig. 4 C, D, Fig. S4) is calculated by considering each community as a vector of the population sizes of the 15 species. Species absence from the community is marked by zero. We find the beta diversity as the average Bray-Curtis dissimilarity of each of the 210 possible pairs in the 21 communities.

The community coverage of nutrients and toxic compounds (Fig. 3F) is quantified similarly to species diversity, as the effective number of toxic compounds invested into or nutrients taken up. Toxic compound coverage is calculated from the vector $\tilde{f}_{k} = \sum_{l} \tilde{f}_{lk}$ of total investment into degrading toxic compound k in a given community. The 'tilde' indicates that we have scaled the f_{lk} for each strain of a species by the corresponding population sizes of the strains in the community

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as in Eq. (2). In this way, we emphasize the most relevant $_{930}$ strain of each species and do not bias the result to the number of competing strains within a species. Note that we do not $_{932}$ scale $\tilde{f}_{\cdot k}$ by species abundance in the community. Once each $_{934}^{933}$ investment $\tilde{f}_{\cdot k}$ is rescaled so that $\sum_k \tilde{f}_{\cdot k} = 1$, we calculate $_{935}^{936}$ the effective number of toxic compounds invested into as

$$\exp(H') = \exp\left(-\sum_{k=1} \tilde{f}_{\cdot k} \log \tilde{f}_{\cdot k}\right), \qquad (15)_{940}^{939}$$

and average this value over all 21 communities. The measure ⁹⁴³ falls between 0 (no toxic compounds are invested into) and 10 ⁹⁴⁴ (all compounds). For the nutrient coverage, we use the same ⁹⁴⁶ calculation using the nutrient uptake rates n_{ij} , but without ⁹⁴⁷ scaling between different strains as this parameter does not ⁹⁴⁹ mutate. The effective number of nutrients taken up takes val-⁹⁵⁰ ues between 0 and 4, the number of different nutrients. ⁹⁵²

953 To evaluate the stability of the selection methods (Fig. 6A), $\frac{1}{954}$ 900 we choose the highest-scoring community that each method 955 901 found after 50 rounds of selection (one community for each 957 902 repeated run of each species set), and seed 10 replicates with 958 903 its identical initial composition. Then we grow and dilute $_{960}$ 904 them for a further 25 rounds, as we would do for the no-961 905 selection treatment. We do not allow mutations in these 963 906 rounds, to focus only on the ecological stability of the found 964 907 communities. For the analysis of sensitivity to the number of $_{966}$ 908 species in the initial species pool (Fig. 6C, Fig. S9), we sam-909 ple 5 random subsets of 6, 9, 12 from the original set of 15 969 910 species, and run for each of them 5 simulations with differ-⁹⁷⁰ 911 ent 21 initial communities. Drawing new species sets of the 972 912 corresponding size would introduce further variance, which 973 913 we would rather avoid. For the effect of the dilution factor 975 914 (Fig. 6E, Fig. S11), we multiply the 10-cell inoculum by a ⁹⁷⁶ 915 factor 0.2, 0.5, 1, 2 or 5 for the disassembly method and scale $_{978}$ 916 the 5% dilution fraction for the other methods by the same ⁹⁷⁹ 917 factor. 918 981

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BV, FAS and SM conceived the project. PGF and BV devel-986 920 oped, implemented and simulated the models and selection. 988 921 PGF and BV analyzed the data. BV wrote the first draft. BV, 989 922 PGF and SM wrote the manuscript. FAS commented on the 991 923 manuscript. BV, FAS, PGF and SM were funded by Euro-992 924 pean Research Council Starting Grant 715097, and SM by the 994 925 NCCR Microbiomes (Swiss National Science Foundation). 995 926 We thank the Mitri lab at the University of Lausanne for valu- 997 927 able discussions, in particular Afra Salazar de Dios and Shota 998 928 999 Shibasaki for detailed comments on the manuscript. 929 1000

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Supplementary tables

Method 1	Method 2	p (IBM)	p (ODE)
DS	Ø	4×10^{-10}	4×10^{-10}
DS	PS	8×10^{-10}	8×10^{-10}
DS	MS	8×10^{-10}	-
DS	PIS	1×10^{-9}	2×10^{-8}
DS	MIS	8×10^{-10}	-
DS	DR	8×10^{-10}	8×10^{-10}
DS	NS	8×10^{-10}	8×10^{-10}
PS	Ø	1.0	1.0
PS	MS	1×10^{-7}	-
PS	PIS	4×10^{-9}	7×10^{-5}
PS	PR	8×10^{-10}	3×10^{-9}
PS	NS	8×10^{-10}	0.1
MS	Ø	1.0	-
MS	MIS	0.67	-
MS	MR	8×10^{-10}	-
MS	NS	0.3	-
PIS	Ø	9×10^{-6}	0.8
MIS	Ø	1.0	-

 Table S1. P-values for Fig.2A-D a Wilcoxon signed-rank test of difference in maximum degradation between methods.

Method 1	Method 2	<i>p</i> (IBM)	p (ODE)
DS	PS	8×10^{-10}	7×10^{-10}
DS	MS	8×10^{-10}	-
DS	PIS	8×10^{-10}	7×10^{-10}
DS	MIS	8×10^{-10}	-
DS	DR	8×10^{-10}	8×10^{-10}
DS	NS	8×10^{-10}	7×10^{-10}
PS	MS	1×10^{-4}	-
PS	PIS	3×10^{-8}	7×10^{-4}
PS	PR	2×10^{-9}	7×10^{-7}
PS	NS	8×10^{-10}	3×10^{-6}
MS	MIS	8×10^{-2}	-
MS	MR	2×10^{-8}	-
MS	NS	2×10^{-9}	

Table S2. P-values for Fig.2E-H a Wilcoxon signed-rank test of difference in degradation ranks between methods.

Method 1	Method 2	p (IBM)	p (ODE)
DS	PS	6×10^{-9}	8×10^{-10}
DS	MS	8×10^{-10}	-
DS	PIS	8×10^{-10}	8×10^{-10}
DS	MIS	8×10^{-10}	
DS	DR	8×10^{-10}	6×10^{-4}
DS	NS	8×10^{-10}	8×10^{-10}
PS	MS	1×10^{-9}	-
PS	PIS	0.1	0.01
PS	PR	2×10^{-8}	0.5
PS	NS	8×10^{-10}	0.3
MS	MIS	0.13	-
MS	MR	1×10^{-9}	-
MS	NS	1×10^{-5}	-

Table S3. P-values from a Wilcoxon signed-rank test of difference in total investment in communities, between methods for Fig.3A-D.

Supplementary figures

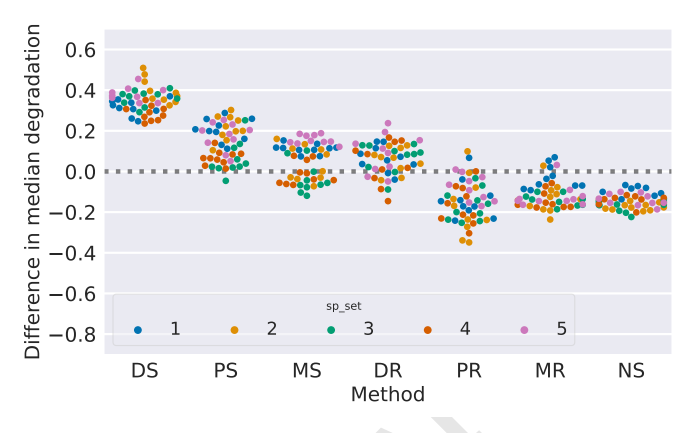


Fig. S1. The difference in median degradation between round 50 and round 0 for each propagation method, corresponding to Fig. 2. The two-sided Wilcoxon test for difference in degradation against the no-selection control is significant for the selection methods DS, PS and MS. Data generated by the IBM.

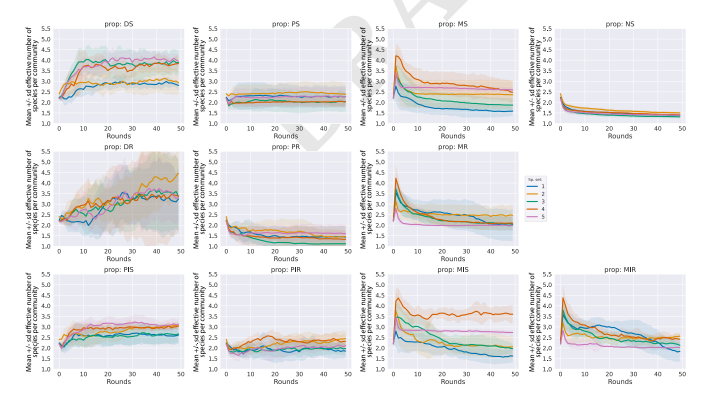


Fig. S2. Time-series of the species diversity (effective number of species per community) corresponding to Fig. 4D. Each panel shows the mean \pm standard deviation over the 10 repeated runs, for each species set 1-5, for one propagation method. Data generated by the IBM.

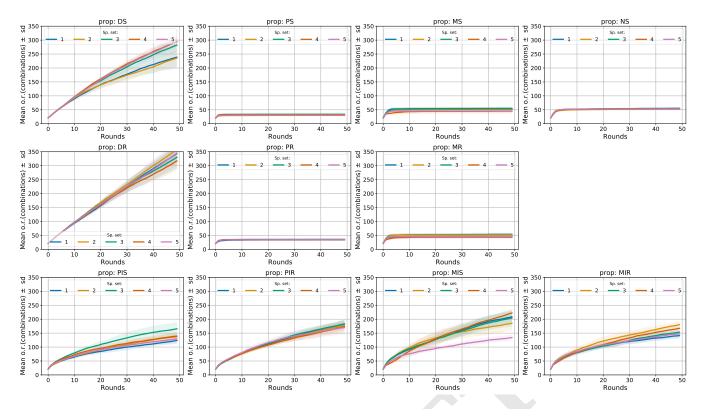


Fig. S3. Time-series of the number of explored communities, corresponding to Fig. 4B. Each panel shows the mean \pm standard deviation over the 10 repeated runs, for each species set 1-5, for one propagation method. Data generated by the IBM.

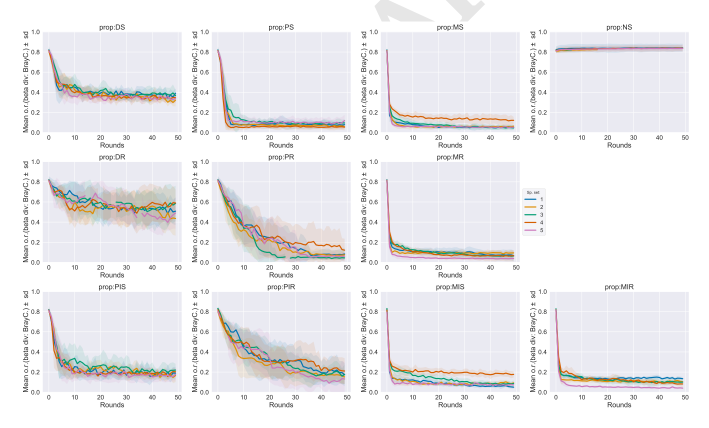


Fig. S4. Time-series of the beta diversity corresponding to Fig. 4F. Each panel shows the mean \pm standard deviation over the 10 repeated runs, for each species set 1-5, for one propagation method. Data generated by the IBM.

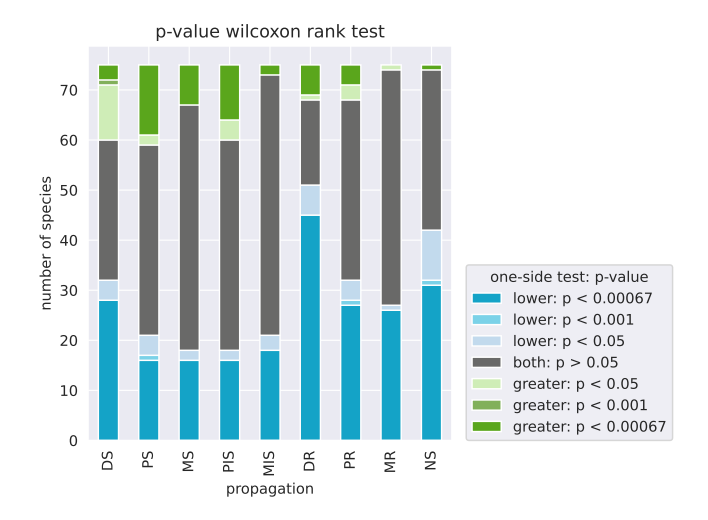


Fig. S5. Distribution of p-values from a one-sided Wilcoxon signed-rank test of whether the total investment $f_{l.}$ of a species is larger/smaller in the last round where a species survived, than the investment of the ancestral species. There is one bar for each selection method, with 15 species x 5 sets of species for each bar. The alternative hypothesis is that difference in investment (ancestral-evolved) is greater (green) or less (blue) than zero. Data generated by the IBM. Data for DS is shown in Fig. 5.

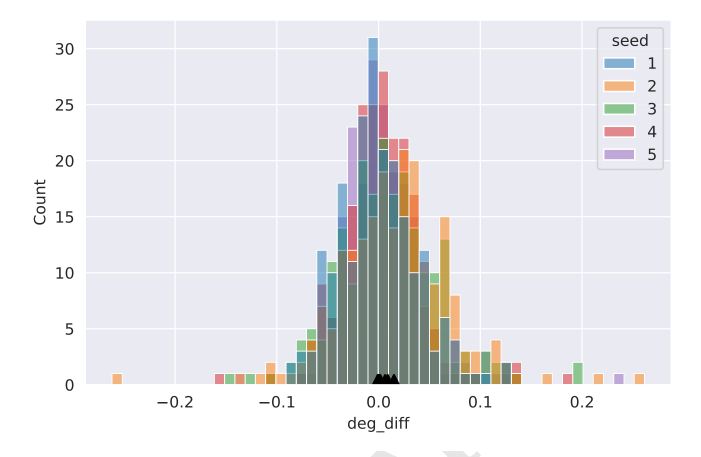


Fig. S6. Histogram of difference in max degradation between evolved and ancestral communities. Triangles indicate the mean values for each species set. Data generated by the IBM.

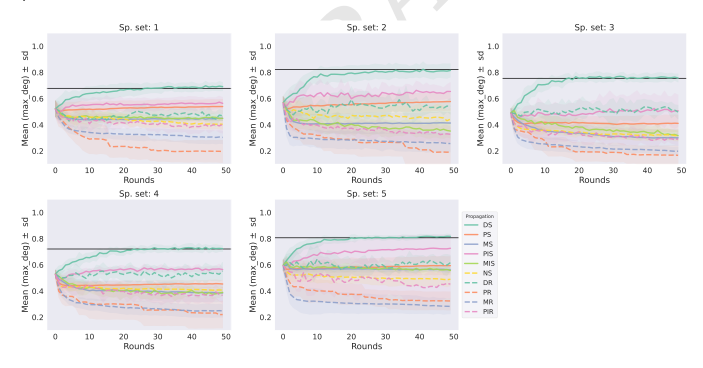


Fig. S7. Time-series of max. degradation over 50 rounds of selection, for the different propagation methods. Each plot corresponds to one species set and shows the maximum degradation in the meta-community averaged over repeats, with the standard deviation in shades of the corresponding color. For each species set, each repeat etc, the degradation score at transfer 50 forms the swarms in Fig. 2. The black line shows the degradation score of the best ancestral community out of the 32767 combinations. Data generated by the IBM.

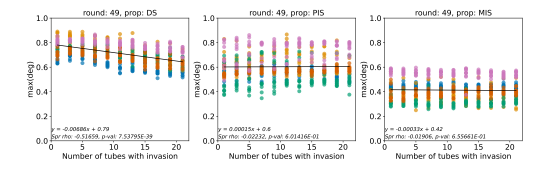


Fig. S8. Effect on max community degradation score from changing the number of communities to receive a migrating species. Data generated by the IBM.

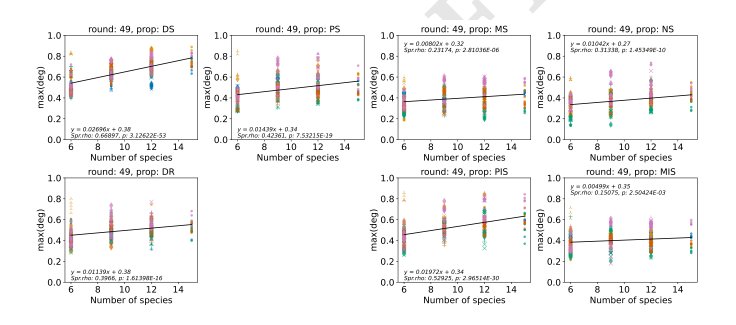


Fig. S9. Effect on max community degradation score from changing the number of species in the ancestral community. Different marker shape indicates sub-sample (1-5) for each species group of size 6, 9, 12. For 15 species we keep the original species sets. Data generated by the IBM.

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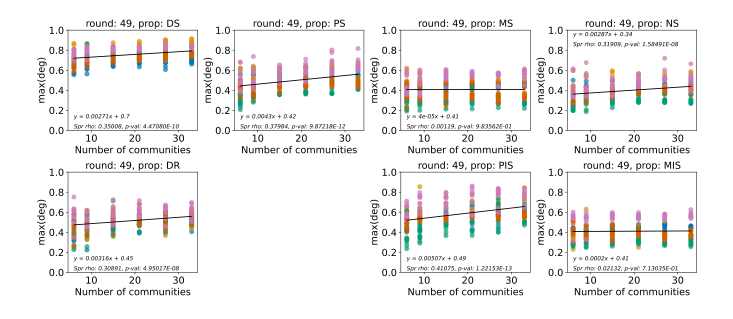


Fig. S10. Effect on max community degradation score from changing the number of communities. Data generated by the IBM.

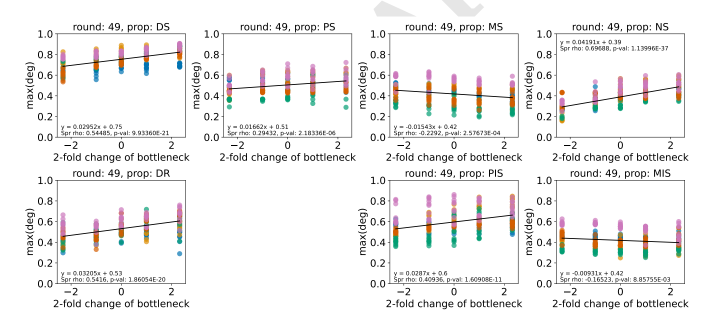


Fig. S11. Effect on max community degradation score from scaling the dilution factor or inoculum size. Data generated by the IBM.

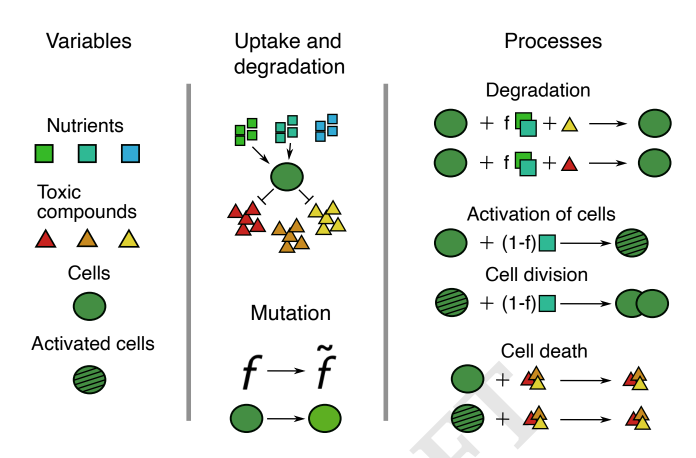


Fig. S12. Illustration of the variables and processes in the individual-based model. Cells of a certain species vary in their preferences for nutrients and degradation capabilities. Cells use the available nutrients to degrade the toxic compounds and for cell division.

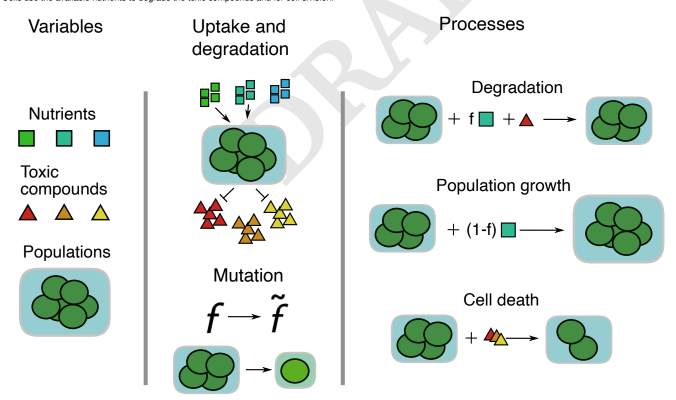


Fig. S13. Illustration of the variables and processes in the ODE model. Populations of cells vary in their preferences for nutrients and degradation capabilities. The populations use the available nutrients to degrade the toxic compounds and to grow.

¹⁰³⁷ Pseudo-code for implementation of models and selection methods

Input: A set of 15 species, defined by model parameters.

Input: Experimental parameters: Number of communities, time span $[t_0, t_{end}]$ of growth in batch. Community bottleneck $\beta = 1/3$, dilution ratio $d \in [0, 1]$. Initial conditions $S_i(t_0), N_j(t_0), T_k(t_0)$.

Assemble 21 communities by randomly drawing 4 species with replacement from the species set. Ensure that each species is present in at least one initial community.

for Each round of selection do

// Population growth, interspecies competition and invasion of mutants

for Each community do

Grow the communities for a time span $[t_0, t_{end}]$. (IBM implementation: Alg. 2, ODE implementation: Alg. 6).

Save the population sizes $S_i(t_{end})$ for each strain *i* in the community

Save the end-state concentrations $T_k(t_{end})$ for each toxic compound k

Compute the degradation score D from $T_k(t_{end})$ by Eq. (1)

end

// Propagate the communities by the chosen selection method

// Required parameters: the community bottleneck β , dilution ratio d

// Required variables: degradation scores D for each community

// For propagule method, follow Algs. 7 and 8

// For migrant pool method, IBM only, follow Alg. 9

// For disassembly method, follow Alg. 10

// Replenish the substrates

Set $N_j(t_0) = N_0$ and $T_k(t_0) = T_0$ for all j, k

end

Algorithm 1: Overall flow of the selection simulations from population growth, community dynamics and mutations to propagation by the different selection methods.

Input: A community where each microbial strain *i* is defined by the parameters in Tab. 1. Size of the inactive and active sub-populations p_{i0} , p_{i1} and total population size $S_i = p_{i0} + p_{i1}$. Nutrient concentrations N_j , toxic compound concentrations T_k .

Input: Mutation parameters: mutation rate μ_{mut} , trait deviation σ_m .

for Each time step do

```
for Each community 1, ..., 21 do
    for Each strain i do
        // Maximum uptake of nutrients of this strain, used to scale the growth and degradation according to the
          concentration of nutrients
        max\_uptake := \sum_{j} (n_{ij} \text{ if } N_j > n_{ij})
        // If some nutrients are depleted we re-scale n_{ij} to consume the remaining nutrients in higher amounts
        if N_j < n_{ij} then
         | n_{ij} := 0
        end
        \hat{n}_{ij} := \frac{n_{ij}}{\sum_j (n_{ij} \text{ if } N_j > n_{ij})}
        // The largest number of cells that can consume the scarcest nutrient at this time step, we just take its integer part
         S_i^{max} := int(min(N_j/\hat{n}_{ij} \text{ if } N_j > \hat{n}_{ij}))
        // Toxic compound degradation
        if S_i^{max} > 0 then
             II Maximal population that can degrade
             P_{i,tot} := \min(S_i^{max}, S_i)
             for Each compound T_k do
                 if T_k cannot be completely degraded in this time step then
                      T_k := T_k - P_{i,tot} \cdot max\_uptake \cdot f_{ik}
                      for Each nutrient N_j do
                       N_j := N_j - \hat{n}_{ij} \cdot P_{i,tot} \cdot f_{ik}
                      end
                 end
                 else
                  Degrade the remaining toxic compounds and consume the corresponding nutrients
                 end
             end
        end
        // Cell division step 1: Costly activation
        Alg. 3.
    end
end
// Cell division step 2: Replication
Alg. 4.
// Cell death
Alg. 5.
```

Algorithm 2: Implementation of population growth, competition and mutations in the IBM model described in the section *Individual-based model*.

end

Input: Communities where each strain *i* is defined by parameters in Tab. 1. Inactive and active sub-populations p_{i0} , p_{i1} . The maximal population S_i^{max} that can afford to consume nutrients, based on their current availability. Re-scaled nutrient consumption rates \hat{n}_{ij} . Current nutrient concentrations N_j .

Input: Parameters: Initial nutrient concentration N_0 .

// Cell activation if $S_{i}^{max} > 0$ then *II Already activated cells consume nutrients* if $S_i^{max} \ge p_{i1}$ then $N_j := N_j - \hat{n}_{ij} \cdot p_{i1} \cdot (1 - \sum_k f_{ik})$ $S_i^{max} := S_i^{max} - p_{i1}$ end else Deactivate cells that cannot afford to stay activated, consume the corresponding nutrients for cells that remain activated, set $S_i^{max} := 0$ end // Newly activated cells $cells_activate := \text{Poisson}(a_i \cdot p_{i0} \cdot max_uptake \cdot (1 - \sum_k f_{ik}) \cdot \sum_j (\hat{n}_{ij} \cdot N_j / N_0))$ if $cells_activate > p_{i0}$ then $| cells_activate := p_{i0}$ end if $cells_activate > S_i^{max}$ then $cells_activate := S_i^{max}$ end $p_{i0} := p_{i0} - cells_activate$ $p_{i1} := p_{i1} + cells_activate$ for Each nutrient N_i do $| N_j := N_j - \hat{n}_{ij} \cdot cells_activate \cdot (1 - \sum_k f_{ik})$ end

end

else | Deactivate all the activated cells end

return Populations p_{i0} , p_{i1} for strain *i*, current nutrient concentrations N_j

Algorithm 3: Activation of cells, the first step of cell division in the IBM described in Alg. 2.

Input: Communities where each strain i is defined by parameters in Tab. 1. Inactive and active sub-populations p_{i0} , p_{i1} . for Each community **do**

for Each strain *i* in the community **do** // Calculate the number of new cells appearing due to division $new_cells :=$ Poisson $(p_{i1} \cdot r_i \cdot (1 - \sum_k f_{ik}))$ if $new_cells > p_{i1}$ then $| new_cells := p_{i1}$ end // Calculate how many new_cells will carry mutations $mutants := Poisson (new cells \cdot \mu_{mut})$ if $mutants > new_cells$ then $mutants := new_cells$ end $p_{i1} := p_{i1} - new_cells$ $p_{i0} := p_{i0} + new_cells \cdot 2 - mutants$ **I**Mutation for Each new mutant do Add a new strain i to the community, with the model parameters of the ancestor and set $p_{i0} := 1$ and $p_{i1} := 0$ Decide which f_{ik} to mutate by drawing from Bernoulli $\left(\frac{1}{N_{tox}}\right)$ for each f_{ik} ; ensure that at least one f_{ik} mutates for Each successful draw do Multiply the chosen f_{ik} , by a factor $x \sim lognormal(0.0, \sigma_m)$ Re-scale so $\sum_k f_{ik} \leq 1$, if needed end end end end **return** Populations p_{i0} , p_{i1} for each strain *i*, including the new ones resulted from mutation.

Algorithm 4: Replication and mutation, second step of cell division of the IBM described in Alg. 2.

Input: Communities where each strain i is defined by parameters in Tab. 1. Inactive and active sub-populations p_{i0} , p_{i1} .

Current concentrations T_k of toxic compounds. Death rates m_{ik} and K constant for the Hill function.

for Each community do

for Each strain *i*, looping over the community in reverse order do

$$p_{i0} := p_{i0} - \operatorname{Poisson}\left(p_{i0} \cdot \sum_{k} (m_{ik} \cdot \frac{T_k^2}{T_k^2 + K^2})\right); \text{ ensure that } p_{i_0} \ge 0$$

$$p_{i1} := p_{i1} - \operatorname{Poisson}\left(p_{i1} \cdot \sum_{k} (m_{ik} \cdot \frac{T_k^2}{T_k^2 + K^2})\right); \text{ ensure that } p_{i_1} \ge 0$$

$$if p_{i0} + p_{i1} = 0 \text{ then}$$

$$remove strain i from the community$$

remove strain *i* from the commun

end

end

Randomly shuffle strains in community

end

return Populations p_{i0} , p_{i1} for each strain *i*.

Algorithm 5: Cell death of the IBM described in Alg. 2.

Input: Strains with model parameters from 2 for each population *i*.

Input: Mutation parameters: rate μ_{mut} , trait deviation σ_m .

Input: Experimental parameters: Number of toxic compounds N_{tox} , initial concentrations N_0 , T_0 of nutrients and toxic compounds, initial population size S_0 . Time span for growth $[t_0, t_{end}]$

for Each community do

// Growth and competition within one round

Solve the equations Eq. (6)–Eq. (8) for a time span $[t_0, t_{end}]$.

Save the end states $S_i(t_{end})$, $N_j(t_{end})$, $T_k(t_{end})$.

// Mutations, ODE model

for Each strain *i*, with probability μ_{mut} do

Copy the species parameters to an empty place in the list of populations

Choose a f_{ik} at random by drawing from Bernoulli $(1/N_{tox})$ for each k = 1, ..., 10

For each chosen f_{ik} , multiply by a factor $x_k \sim lognormal(0.0, \sigma_m)$

Set the inoculum size to S_0

end

end

return $S_i(t_{end})$, $N_j(t_{end})$, $T_k(t_{end})$, f_{ik}

Algorithm 6: Implementation of population growth, competition and mutations in the ODE model described in the section Population-level model. To solve the equations, we use dopri5 from the SciPy library (35, 36).

Input: Communities with populations S_i and degradation scores D. End states $T_k(t_{end})$.

Input: Experimental parameters: selection bottleneck $\beta = 1/3$, dilution ratio d.

Rank the communities by degradation D

Select the top $N_{\beta} = 7$ of communities with the highest ranks.

// Re-populate the new set of tubes

Allocate $1/\beta$ new tubes for each selected community

for Each selected community 1, 2, ..., 7 do

for Each population S_i in the selected community do

Dilute $S_i(t_0) := d \cdot S_i(t_{end})$

if
$$S_i(t_0) < 1.0$$
 then

 $S_i(t_0) < 1.0$ then // The population is extinct

Set
$$S_i(t_0) := 0$$
.

Remove all species parameters from the community

end

Copy model parameters and population sizes $S_i(t_0)$ of each strain in the parent communities to each of the $1/\beta$

offspring communities

end end

Algorithm 7: Implementation of the propagule selection method for the ODE model.

Input: Communities with degradation scores D and strains S_i with total population $S_i = p_{i0} + p_{i1}$. End-state concentration of toxic compounds $T_k(t_{end})$.

Input: Experimental parameters: selection bottleneck $\beta = 1/3$, dilution ratio d.

Rank the communities by D

Select the top $N_{\beta} = 7$ of communities with the highest ranks

// Re-populate the new set of tubes

Allocate $1/\beta$ new tubes for each selected community

for Each selected community $1, 2, \ldots, 7$ do

for Each strain *i* in the community **do**

```
 \begin{array}{l} \textit{ // Deactivate cells} \\ p_{i0}(t_{end}) = S_i(t_{end}) \\ p_{i1}(t_{end}) = 0 \\ \textit{ // Dilute the population, new cells will be inactivated} \\ p_{i0}(t_0) := \operatorname{Poisson}(d \cdot S_i(t_{end})) \\ \text{ if } p_{i0}(t_0) > S_i(t_{end}) \text{ then} \\ | p_{i0}(t_0) = S_i(t_{end}) \\ \text{ end} \\ \text{ if } p_{i0}(t_0) > 0 \text{ then} \\ | \text{ Append strain } i \text{ with population } p_0(t_0) \text{ to the new tube} \\ \text{ end} \\ \textit{ // Delete selected cells to not chose them again} \\ S_i(t_{end}) = S_i(t_{end}) - p_{i0}(t_0) \\ \text{ end} \end{array}
```

end

Algorithm 8: Implementation of the propagule selection method for the IBM.

Input: Communities with degradation scores D and strains S_i with total population $S_i = p_{i0} + p_{i1}$. End-state concentration of toxic compounds $T_k(t_{end})$.

Input: Experimental parameters: selection bottleneck $\beta = 1/3$, dilution ratio d.

Rank the communities by D

Select the top $N_{\beta} = 7$ communities with the highest ranks, and pool their populations.

// Re-populate the new set of tubes

Allocate 21 new tubes

for Each offspring community 1, 2, ..., 21 do

```
for Each strain i in the pool do

// Deactivate cells

p_{i0}(t_{end}) = S_i(t_{end})

p_{i1}(t_{end}) = 0

// Dilute the population, new cells will be inactivated

Draw p_{i0}(t_0) from Poisson (\frac{d}{N_{\beta}} \cdot S_i(t_{end}))

if p_{i0}(t_0) > S_i(t_{end}) then

| p_{i0}(t_0) = S_i(t_{end})

end

if p_{i0}(t_0) > 0 then

| Append strain i with population p_{i0}(t_0) to the new tube

end

// Delete selected cells to not chose them again

S_i(t_{end}) = S_i(t_{end}) - p_{i0}(t_0)
```

end

end

Algorithm 9: Implementation of the migrant pool selection method for the IBM

- **Input:** Communities with populations S_i (in the IBM $S_i = p_{i0} + p_{i1}$) with model parameters from (Tab. 1, Tab. 2) and degradation scores D.
- Input: Experimental parameters: selection bottleneck $\beta = 1/3$, initial population size S_0 and number of new communities to emigrate species from $N_{emi} = 5$, and immigrate species to $N_{immi} = 5$.

// Rank the communities

Rank the communities by the degradation score D

// Update the fossil record with the top communities in this round

for Each selected community $1, 2, \ldots, 7$ do

for Each species *l* in the selected community do

if The record of species *l* is not yet updated in this round of selection then

Add all strains i of species l to the fossil record, including the corresponding parameters and population sizes end

end

end

// Propose new communities in proportion to their degradation scores and survival

Follow Alg. 11

// Emigration

Draw $N_{emi} = 5$ communities with uniform probability

for Each chosen community $1, \ldots, N_{emi}$ do

// Find emigrating species

 $Number_of_emigrants = 1 + Poisson(0.5)$

Verify that at least one species will remain

for Each emigrant do

Choose the emigrant at random, with priority for species that occur in more than one community

Remove all strains of this species from the community

end

end

// Immigration

Draw $N_{immi} = 5$ communities with uniform probability

```
for Each chosen community 1, \ldots, N_{immi} do
```

```
// Find immigrating species
```

 $Number_of_immigrants = 1 + Poisson(0.5)$

for Each immigrant do

if There are species that do not feature in any community then

```
Choose one of them at random
```

end

else

| Choose a species that is not already in the community with uniform probability

end

Take all strains of the species from the species record, add them to the offspring community

Set the population size to S_0 (approximately S_0 in the IBM, see Alg. 11), in proportion to strain relative abundance end

end

Algorithm 10: Implementation of the disassembly method.

Input: The subset of selected communities, their degradation scores D. Population sizes S_i of all strains (in the IBM,

 $S_i = p_{i0} + p_{i1}$).

Input: Parameters: Number of species N_{spc} that were inoculated in the different communities. Inoculum size per species S_0 . // Scale degradation scores by species extinctions

for Each selected community 1, 2, ..., 7 do

// Count the number of surviving species in this community

Set the extinction counter E = 0

for Each species *l* in the community **do**

if $S_i = 0$ for all strains of species l then

Set E := E + 1

// Re-introduce species l from the record

Take all strains of species l from the most recent record

end

end

// Scale the degradation score D by the fraction of surviving species

Set $\hat{D} := D \cdot (N_{spc} - E) / N_{spc}$

end

// Calculate a probability distribution based on degradation scores Set $p_n := \hat{D}_n / \sum_m \hat{D}_m$ for each community n// Propose new communities randomly in proportion to p_n

for Each offspring community do

Choose a parental community at random, by the probability distribution p_n

for Each species l in the parental community do

Copy the growth parameters and f_{ik} of all strains i to the offspring community

// ODE: Set the population size of each species to S_0 in total, in proportion to the relative abundance of the strains.

// IBM: Sample approximately S_0 cells with replacement as follows:

for Each strain i (with population S_i) of species l (with population S_l) do

//Deactivate cells

$$p_{i0}(t_{end}) = S_i(t_{end})$$

 $p_{i1}(t_{end}) = 0$

// Draw new population

Draw $p_{i0}(t_0)$ from Poisson $(S_0 \cdot \frac{S_i}{S_i})$

if
$$p_{i0}(t_0) > 0$$
 then

Add strain *i* with population $p_{i0}(t_0)$ to the new community

```
end
```

end

end

Algorithm 11: Method to propose new communities based on their degradation scores from the previous round. Called by Alg. 10