

# Phenotypic and genomic signatures of interspecies cooperation and conflict in naturally-occurring isolates of a model plant symbiont

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Given the need to predict the outcomes of (co)evolution in host-associated microbiomes, whether microbial and host fitnesses tend to trade off, generating conflict, remains a pressing question. Examining the relationships between host and microbe fitness proxies at both the phenotypic and genomic levels can illuminate the mechanisms underlying interspecies cooperation and conflict. We examined naturally-occurring genetic variation in 191 strains of the model microbial symbiont, *Ensifer meliloti*, paired with each of two host *Medicago truncatula* genotypes in single- or multi-strain experiments to determine how multiple proxies of microbial and host fitness were related to one another and test key predictions about mutualism evolution at the genomic scale, while also addressing the challenge of measuring microbial fitness. We found little evidence for interspecies fitness conflict; loci tended to have concordant effects on both microbe and host fitnesses, even in environments with multiple co-occurring strains. Our results emphasize the importance of quantifying microbial relative fitness for understanding microbiome evolution and thus harnessing microbiomes to improve host fitness. Additionally, we find that mutualistic coevolution between hosts and microbes acts to maintain, rather than erode, genetic diversity, potentially explaining why variation in mutualism traits persists in nature.

conflict | cooperation | GWAS | mutualism | pleiotropy

Recent advances in sequencing and microbiome research have revealed the ubiquity of microbial symbioses, meaning that many important host phenotypes, such as plant yield in agriculture or disease-related traits in humans, are actually **symbiotic extended phenotypes**, their variation being influenced by loci present within interacting microbial symbionts in addition to the host (1–6). When loci influence fitness-related traits of both host and symbiont, which we henceforth refer to as **symbiotic pleiotropy**, they determine the degree to which partners' fitnesses are aligned (i.e., same-sign or concordant effects) or in conflict (i.e., opposite-sign or discordant effects). Identifying the loci underlying symbiotic pleiotropy is therefore critical not only for illuminating the genetic basis of symbiotic extended phenotypes, but also for predicting how hosts and symbionts coevolve in nature.

In symbiotic mutualisms, wherein partners trade fitness benefits (7), whether fitness conflict or alignment drives the evolution of these interactions is hotly debated (8–13). Many models of mutualism rely on the key assumption that cooperation is inherently costly and thus that selection should favour less-cooperative, potentially 'cheating', partners that forgo paying the costs whilst continuing to receive benefits (14–17). At the phenotypic level, cheaters would be seen as symbiont genotypes that gain fitness at the host's expense (i.e., points in the bottom right quadrant of **Fig. 1A**), while fitness conflict would be seen as an overall negative correlation (grey line in **Fig. 1A**). In contrast, cooperators would be seen as symbiont genotypes whose increase in fitness is associated with an increase in host fitness (i.e., points in the top right quadrant of **Fig. 1A**), while fitness

20 alignment would be seen as an overall positive correlation (orange line in **Fig. 1A**). Evidence  
21 for fitness conflict within mutualism is mixed: although several studies have identified symbiont  
22 genotypes that gain fitness at their host's expense (e.g., 18, 19), recent experimental evolution  
23 studies instead found that microbial adaptation to particular host genotypes is associated with an  
24 increase, rather than a decrease, in host fitness (20–22). Yet, fitness alignment at the phenotypic  
25 level does not necessarily preclude fitness conflict at the genomic level: rather than dichotomous  
26 categories of "cooperator" or "cheater", mutualist genomes are best viewed as mosaics of loci (5),  
27 some underlying cooperation while others underlie conflict. Whether coevolution resulting from  
28 symbiosis leads to more beneficial interactions and greater mutualism stability, or alternatively, more  
29 antagonism and less stable interactions, therefore requires examining the relationships between host  
30 and symbiont fitness proxies at both the phenotypic and genomic levels.

31 Genome-wide association studies (i.e., GWAS) can be used to reveal the genes, as well as specific  
32 segregating mutations (i.e., variants), that underlie variation in symbiotic extended phenotypes  
33 in natural populations (20, 23–27). Because they provide an estimate of both the strength and  
34 direction of effects of particular alleles on the trait, henceforth referred to as **allelic effect sizes**,  
35 GWAS are especially useful for identifying loci underlying symbiotic pleiotropy, and thus, defining  
36 the mutational landscape of mutualism evolution in nature. For example, if symbiotic pleiotropy is  
37 extensive and its effects on fitness-related traits in the interacting partners tend to be discordant  
38 (grey quadrants in **Fig. 1B**), then conflict should underlie the evolution of mutualism, allowing for  
39 the possibility of cheating individuals that are competitively superior, as mutualism theory predicts  
40 (14–17). In contrast, if pleiotropic effects are overwhelmingly concordant (orange quadrants in **Fig.**  
41 **1B**), fitness alignment rather than conflict should be the null hypothesis in mutualism, and models  
42 relying on cheating genotypes as the main driver of mutualism evolution may not be suitable.

43 A longstanding mutualism paradox is that host-driven selection for the 'best' symbiont genotype  
44 should reduce overall symbiont diversity, yet diverse symbiont populations persist in nature (reviewed  
45 by 28, 29). Identifying patterns of selection acting on loci that determine fitness outcomes in natural  
46 populations could be key for resolving this paradox. Studies to date examining patterns of molecular  
47 variation have found stabilizing or purifying selection acting on candidate genes associated with  
48 partner recognition or quality, rather than patterns suggesting rapid turnover of alleles underlying  
49 conflict (30–33). In a recent *in silico* GWAS (5), conflict over mutualistic trait optima between

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50 hosts and microbes tended to increase genetic variance due to repeated sweeps, while alignment  
51 scenarios resulted in stabilizing selection and decreased genetic variance as trait optima were reached  
52 (5). Because GWAS has the power to reveal individual allelic effects, it can be used to identify loci  
53 underlying fundamental sources of conflict at the genomic level (e.g., grey quadrants, **Fig. 1B**) even  
54 if fitness alignment is realized at the phenotypic-level (e.g., orange line, **Fig. 1A**).

55 Over the past ~25 years, legume-rhizobium symbioses have been developed as models for un-  
56 derstanding mutualism evolution (34–37). This interaction is one of the most ecologically and  
57 economically important symbioses, contributing upwards of 20 million tonnes of nitrogen (N) to  
58 the global N-cycle, and saving billions of dollars that would have otherwise been spent on synthetic  
59 N fertilizer production (38). Legumes house rhizobia within specialized root nodules and supply  
60 them with photosynthate, while rhizobia within the nodules convert atmospheric N to a plant-usa-  
61 ble form, resulting in a beneficial exchange of resources. Key traits of this symbiosis, such as plant  
62 biomass and nodule number, are known to be influenced by variants in the genomes of both host  
63 and microbes, as well as the epistatic interactions between them (Genotype-by-Genotype or G x  
64 G interactions; e.g., 1, 26, 34, 39–42), making this symbiosis an excellent model for understanding  
65 mutualistic coevolution.

66 A quantitative comparison of single-strain and multi-strain proxies of rhizobium fitness, as well as  
67 their correlations with plant fitness, is needed to reveal interspecies conflict. Seed number, or close  
68 correlates such as aboveground (shoot) biomass and leaf chlorophyll A content, are well-established  
69 proxies for reproductive fitness in annual plants (23, 43, 44) (**Fig. 1C**). In contrast, estimating  
70 rhizobium fitness has been an empirical and conceptual challenge (8–10, 45, 46). Early attempts to  
71 estimate rhizobium fitness relied on single-strain experiments, whereby plants were inoculated with a  
72 single rhizobium strain, and fitness proxies including nodule number and nodule size were measured  
73 (**Fig. 1D**). Both measures reflect rhizobium fitness because a rhizobium that establishes a nodule  
74 in symbiosis can gain a fitness benefit on the order of  $10^5 - 10^7$  (47), larger nodules release more  
75 rhizobia (43, 46, 48), and intuitively, strains that produce more nodules will release more rhizobia  
76 (1, 37, 48). While rhizobium fitness proxies measured in single-strain experiments can be directly  
77 correlated with host benefit (e.g., shoot biomass), these measures have been criticized for producing  
78 spurious positive correlations due to stronger fitness feedbacks between host plants and rhizobia  
79 (9). In contrast, multi-strain experiments decouple individual strain fitness from host growth and  
80 better reflect rhizobia fitness in natural and agricultural soils where many strains coexist. Recent  
81 advances merging population genomics and metagenomics have enabled measuring **rhizobium**  
82 **relative fitness**, i.e., a strain's ability to compete for nodulation opportunities and extract host  
83 resources once in nodules when other strains are present (25, 49). Multi-strain experiments that have

84 estimated rhizobium relative fitness have so far hinted at a surprising lack of correlation between  
85 single-strain and multi-strain measures of rhizobium fitness (25), motivating our comprehensive  
86 analysis here.

87 Here we combine two datasets: first, GWAS that test for the association between rhizobium  
88 variants and both plant and rhizobium fitness proxies measured in single-strain experiments (Fig.  
89 1C), whereby 191 strains of *Ensifer meliloti* collected from natural populations were inoculated  
90 individually onto one of two genotypes of the host plant *Medicago truncatula*. Second, a GWAS based  
91 on a new dataset that measured rhizobium relative fitness in a multi-strain experiment (Fig. 1D),  
92 whereby 89 of the 191 *E. meliloti* strains were inoculated together onto the same two host genotypes.  
93 We combine these datasets to first ask: What are the relationships among rhizobium fitness proxies  
94 measured in both single-strain (nodule number, nodule size) and multi-strain (rhizobium relative  
95 fitness) experiments? Are these proxies genetically distinct and thus likely to evolve independently,  
96 or are they linked through pleiotropy, and thus, evolve together? We use this information to next  
97 address the potential for genomic conflict in this symbiosis, asking: Do variants tend to have aligned  
98 or conflicting effects on host and symbiont fitnesses? Finally, we ask whether there is any evidence  
99 that historical selection has differentially shaped loci underlying fitness alignment versus conflict, as  
100 we might predict under different models of mutualism evolution.

## 101 Materials and Methods

102 **Study system.** Full details are provided in Riley et al. (50) and **SI Methods**. *Ensifer* (formerly  
103 *Sinorhizobium*) *meliloti* is an Alphaproteobacteria that forms symbiosis with various *Medicago* spp.,  
104 fixing atmospheric N in return for plant photosynthate. All 191 strains used here were collected  
105 from the native range of the host *Medicago truncatula*, spanning Spain to France (as detailed in 50).  
106 *E. meliloti* has a multipartite genome, including a large (~3.7 Mbp) chromosome and two symbiosis  
107 plasmids (pSymA and pSymB, ~1.4 Mbp and ~1.7 Mbp, respectively); pSymA contains many of  
108 the canonical genes for nodulation and N-fixation (51, 52). We used two lines of *M. truncatula* DZA  
109 315.16 (hereafter DZA) and A17 in separate single-strain experiments and a multi-strain experiment  
110 detailed below.

111 **Single-strain experiments.** Full details are provided in Batstone et al. (42), and **SI Methods**.  
112 Briefly, we conducted two separate experiments in Sept and Nov 2018, one for each *M. truncatula* line,  
113 DZA and A17, respectively. At least three replicate seeds of each plant line were planted individually  
114 into pots (most treatments having four to five plant replicates) and were singly inoculated with one  
115 of 191 strains of *E. meliloti* described above. Within each experiment, we measured proxies for both

116 plant fitness (shoot biomass, leaf chlorophyll A content) and rhizobium fitness (nodule number, per  
117 nodule weight; **Fig. 1C,D**).

118 **Multi-strain experiment.** We grew A17 and DZA hosts with a multi-strain inoculum composed of 89  
119 of the 191 strains used in the single-strain experiments (described in more detail in **SI Methods**).  
120 Using the "select-and-resequence" approach (25), this experiment allowed us to generate new data  
121 on **rhizobium relative fitness**, which represents a strain's ability to both compete for nodulation  
122 opportunities and extract host resources once in nodules when 88 other strains were present (**Fig.**  
123 **1D**). This fitness proxy was obtained by sequencing pooled nodule samples from each plant at the  
124 end of the experiment and estimating each strain's frequency using a haplotype reconstruction  
125 method (53). We then obtained each strain's relative fitness by calculating the fold change in the  
126 frequency of a strain at the end of the experiment relative to its mean frequency at the beginning  
127 (see **SI Methods**). Because we wanted our rhizobium relative fitness metric to represent which  
128 strains will be present in the soil in subsequent generations, our method focuses on measuring  
129 strain frequencies of undifferentiated rhizobia in nodule pools; while differentiated bacteroids are  
130 responsible for fixing N, they are reproductively sterile, and thus, do not contribute to the next  
131 generation. Although the plant and microbe growth conditions used here differed slightly from those  
132 used in the single-strain experiments (due to these experiments being conducted in separate places  
133 and times), previous work has shown that strain frequencies are stable in response to environmental  
134 variation within an experiment, between experiments, and across host generations (49, 54). Thus,  
135 environmental variation due to differences in growth conditions is unlikely to significantly influence  
136 how strains behave across experiments.

137 **Phenotypic analyses.** As described in Batstone et al. (42), we calculated the estimated marginal  
138 means for each fitness proxy in each experiment (i.e., nodule number, nodule size, rhizobium relative  
139 fitness, shoot biomass, leaf chlorophyll A content), correcting for the effect of rack, using the emmeans  
140 package (v1.4.1, 55) in the R environment (56). We then conducted linear pairwise regressions (lm  
141 option in base R) for each fitness proxy against the other within each plant line.

142 **DNA isolation, whole-genome sequencing, and variant calling.** Detailed methods are provided in  
143 Riley et al. (50), and **SI Methods**. We obtained DNA from each of the 191 rhizobium isolates,  
144 sequenced their full genomes including the chromosome and two symbiosis plasmids, used a common  
145 reference-based assembly pipeline to align sequences and call single nucleotide polymorphisms  
146 (SNPs), and filtered the resulting SNPs based on sequence quality, depth, minor allele frequency,  
147 and missingness, resulting in a total of 36,526 filtered SNPs.

148 **Genome-wide association tests.** Detailed methods are provided in **SI Methods**. We conducted  
149 linear mixed models (LMMs) implemented in GEMMA (v0.98.1, [57](#)) to quantify **allelic effect sizes**  
150 that represent both the strength and direction of the association between variants and fitness proxies  
151 after correcting for rhizobium population structure. We ran ten separate association analyses in  
152 GEMMA, one for each of the four fitness proxies measured in single-strain experiments (**Fig. 1C**),  
153 plus rhizobium relative fitness measured in the multi-strain experiment, and for both host plant  
154 lines (DZA, A17; five proxies x two hosts = ten runs).

155 **Genomic analyses.** We first identified pleiotropic variants as those that were significantly associated  
156 with more than one trait on the same host line, significance determined using a permutation method  
157 (described in more detail in the **SI Methods**). Variants that were significantly associated with two  
158 or more rhizobium fitness proxies were categorized as underlying **rhizobium fitness pleiotropy**,  
159 whereas variants associated with at least one rhizobium AND plant fitness proxy were categorized  
160 as underlying **symbiotic pleiotropy**. We further categorized whether pleiotropic variants had  
161 discordant (opposite-sign; +,- or -,+) or concordant (same-sign; ++ or -,-) effects on pairwise fitness  
162 proxy combinations (**Fig. 1B**). Finally, to test whether selection acted on genes containing variants  
163 associated with rhizobium fitness and symbiotic pleiotropy, we used the R package PopGenome  
164 (v2.7.5, [58](#)) to compute several commonly used test statistics that can detect signatures of historical  
165 selection and/or departures of neutrality, namely, nucleotide diversity (i.e.,  $\pi$ ), Tajima's D, as well  
166 as Fu and Li's F and D ([59, 60](#)). Additional details appear in **SI Methods**.

167 **Data availability.** Strains and plant lines are available upon request. All raw data and analysis  
168 code are available on [GitHub](#) (see “Genetics\_conflict\_cooperation” folder). The Supplementary  
169 Information doc contains additional methods, results, figures and tables, as well as descriptions of  
170 datasets that have been uploaded along with this manuscript, and can be additionally accessed from  
171 [bioRxiv](#) and [GitHub](#). Once raw sequence reads and assemblies are archived and made accessible on  
172 NCBI, accession numbers will be added to this manuscript.

## 173 **Results**

174 **Relationships among rhizobium fitness proxies.** In single-strain experiments, when we regressed  
175 strain means for nodule number and nodule weight, we found significant negative correlations for  
176 both hosts (**Fig. 2A**, left), indicating that strains creating larger nodules tended to form fewer total  
177 nodules on both host genotypes. At the genomic-level, most variants had discordant effects (**Fig.**  
178 **2A**, right), similarly indicating a trade-off whereby variants that were positively associated with  
179 nodule weight tended to be negatively associated with nodule number, or vice-versa.

180 Comparing single- and multi-strain experiments, when we regressed strain means for rhizobium  
181 relative fitness and nodule number, we found a weak but significant negative relationship for host  
182 line DZA only (**Fig. 2B**, left), indicating that strains that were more common in the multi-strain  
183 experiment formed fewer nodules in the single-strain experiment. At the genomic-level on DZA,  
184 most variants had discordant effects (**Fig. 2B**, right), again indicating a trade-off between nodule  
185 number and rhizobium relative fitness. For host line A17, most pleiotropic variants (20/28) had  
186 discordant effects (**Fig. 2B**, right), despite no relationship between these fitness proxies at the  
187 phenotypic level (**Fig. 2B**, left).

188 We found a significant, positive relationship between strain means for rhizobium relative fitness  
189 and nodule weight, again for DZA only (**Fig. 2C**, left). This result indicates that strains that  
190 were more commonly found in nodules in the multi-strain experiment formed larger nodules in  
191 the single-strain experiment. At the genomic-level, all pleiotropic variants were concordant (i.e.,  
192 appearing in the top right or bottom left quadrants of **Fig. 2C**, right). For A17, while we found a  
193 lack of significant phenotypic correlation between nodule weight and rhizobium relative fitness (**Fig.**  
194 **2C**, left), at the genomic level, all pleiotropic variants had discordant effects on these fitness proxies  
195 (**Fig. 2C**, right), indicating a trade-off at this scale.

196 These results overall support trade-offs at the phenotypic level, plus underlying discordant  
197 pleiotropy at the genomic level, suggesting that strains for which plants form numerous small nodules  
198 in single-strain experiments are less able to proliferate within and compete for nodules in multi-strain  
199 experiments. We additionally found host-dependent relationships among rhizobium fitness proxies,  
200 especially at the genomic level when regressing rhizobium relative fitness and nodule weight (**Fig.**  
201 **2C**, right). However, we found fewer total pleiotropic variants associated with this relationship  
202 compared to the other two fitness proxy combinations (**SI Fig. S2A**, right), suggesting that nodule  
203 weight and rhizobium relative fitness are largely governed by different molecular mechanisms, and  
204 thus, likely to evolve independently.

205 **Relationships between plant and rhizobium fitness proxies.** We found no correlation, in either  
206 host, between strain means for nodule number and shoot biomass (**Fig. 3A** left), or nodule number  
207 and leaf chlorophyll A content on A17 (**SI Fig. S3A**, left). However we found a significantly negative  
208 correlation between nodule number and chlorophyll A for DZA (**SI Fig. 3A**, left). Assuming  
209 chlorophyll A is related to the N-fixation efficiency of each strain (61–63), this result suggests that  
210 strains forming more numerous (and smaller) nodules tended to fix less N. At the genomic level,  
211 most variants had concordant effects for nodule number and shoot biomass on DZA, whereas the  
212 opposite was true for the same proxy pair on A17 (**Fig. 3A**, right). Most variants had discordant

213 effects on nodule number and chlorophyll A on both hosts (**SI Fig. S3A**, right).

214 Strain means for nodule weight and shoot biomass were significantly positively correlated for both  
215 hosts (**Fig. 3B**, left). Nodule weight and chlorophyll A were significantly positively correlated  
216 on DZA, but uncorrelated on A17 (**SI Fig. S3B**, left). At the genomic level for both hosts, all  
217 pleiotropic variants had concordant effects for nodule weight and shoot biomass (**Fig. 3B**, right), as  
218 well as nodule weight and chlorophyll A (**SI Fig. S3B**, right).

219 Finally, rhizobium relative fitness and shoot biomass were significantly positively correlated for  
220 DZA but not for A17 (**Fig. 3C**, left), whereas rhizobium relative fitness and leaf chlorophyll A  
221 content were significantly positively correlated on both hosts (**SI Fig. S3C**, left). At the genomic  
222 level, most pleiotropic variants had concordant effects on rhizobium relative fitness and shoot biomass  
223 for both hosts (**Fig. 3C**, right), and this pattern was even stronger between rhizobium relative  
224 fitness and chlorophyll A on DZA with all but one pleiotropic variant having concordant effects  
225 (**SI Fig. S3C** right). Variant effects were mixed on A17 for rhizobium relative fitness and leaf  
226 chlorophyll A content (**SI Fig. S3C**, right).

227 Overall, we found mostly concordant relationships between rhizobium and plant fitnesses at  
228 both the phenotypic and genomic levels, suggesting a strong signal of fitness alignment in natural  
229 rhizobium populations. However, we note a lack of pleiotropic variants underlying the relationship  
230 between leaf chlorophyll A content and both rhizobium fitness proxies measured in single-strain  
231 experiments (i.e., nodule number and nodule weight; **SI Fig. S2C**, left & middle), suggesting these  
232 proxies are governed by different molecular mechanisms, and thus, should evolve independently.

233 **Selection acts differently on genes associated with alignment versus conflict.** Using multiple  
234 diversity and neutrality metrics, we found that rhizobium genes associated with fitness alignment  
235 exhibit higher nucleotide diversity and stronger signatures of balancing selection compared to any  
236 other gene category analyzed. For three of the four test statistics, genes associated with concordant  
237 symbiotic pleiotropy (i.e., solid orange lines in **Fig. 4**) had significantly elevated values relative  
238 to the "null" (i.e., distributions in **Fig. 4**, all genes containing significant variants identified by  
239 GWAS). We did not see any significant deviations from the null for genes associated with discordant  
240 symbiotic pleiotropy (i.e., dotted grey lines in **Fig. 4**), or for genes associated with both concordant  
241 and discordant rhizobium fitness pleiotropy (**SI Table S1**, **SI Fig. S4**, **SI Dataset S1**).

242 **Loci associated with symbiotic pleiotropy are host-dependent.** Comparing the identities and  
243 putative functions of variants associated with symbiotic pleiotropy on both host lines revealed little  
244 overlap – concordant variants giving rise to both high host and symbiont fitness (i.e., associated  
245 with fitness alignment) largely differed between the two host genotypes. Specifically, we identified a

total of 168 variants associated with symbiotic pleiotropy, corresponding to 128 coding-genes (see **SI Dataset S2** for variant-level and **SI Table S2 & Dataset S3** for gene-level summaries). 60 and 93 of these variants were uniquely associated with fitness proxies measured on plant lines A17 and DZA, respectively, while only 15 variants were shared between hosts. We highlight some of the noteworthy genes identified in our analysis for each pleiotropic category in **SI Results**.

## Discussion

Leveraging genomics to quantify the genetic architecture underlying symbiotic extended phenotypes gives us the power to address long-standing issues in mutualism evolution with genome-scale resolution. Our results overall suggest that: 1) fitness alignment between hosts and rhizobia is common at both the phenotypic and genomic levels, with genes associated with alignment showing elevated nucleotide diversity and signatures of balancing selection; and 2) the lack of a relationship or even trade-offs between rhizobium nodule number and rhizobium relative fitness mean that measures of rhizobium fitness in multi-strain experiments should be prioritized when we want to predict rhizobium evolution. We discuss these main points in turn below.

**Alignment of host and symbiont fitnesses.** In one-to-many symbioses, such as a single legume associating with a diverse population of rhizobia, less-beneficial symbionts are predicted to achieve higher relative fitness compared to more beneficial counterparts (14–16). While we find evidence for less beneficial rhizobia (i.e., points closer to zero along the y-axes of panels in **Fig. 3**), we find little evidence for "cheating" rhizobia genotypes or loci associated with an increase fitness at the host's expense (i.e., a lack of points in the bottom-right quadrants of panels in **Fig. 3**). Instead, we found strong fitness alignment even in environments where multiple strains occupy the same plant.

Fitness alignment is ultimately governed by the degree to which mutualistic trait optima are shared among partners (5), as well as the degree of fitness feedbacks that enforce alignment between partners (11, 14–17). For example, legume host plants have autoregulation of nodulation to limit the formation of costly nodules once sufficient N levels are achieved (64). No such constraint exists for rhizobia; every nodule formed is expected to lead to greater potential fitness benefits. Thus, a mismatch between the optimum nodule number for a plant versus rhizobium could generate conflict (5, 65). Indeed, the strongest evidence for conflict in our and other studies (e.g., 1, 44, 48) comes from regressing plant fitness proxies on nodule number, suggesting conflict over a host's total investment in nodulation.

In addition to controlling the total number of nodules formed, legumes can also allocate more carbon to nodules that fix more nitrogen (35, 66, 67), which acts to couple rhizobium quality and

278 fitness even when multiple strains are present. Such fitness coupling mechanisms can be disrupted  
279 by mismatches between legume and rhizobium genotypes, allowing strains that fix little to no N  
280 to proliferate within nodules (19, 68, 69). Our observations of abundant alignment between host  
281 fitness proxies, nodule size, and rhizobium relative fitness suggest that such mismatches may be rare  
282 in nature, although in our study, neither host line was collected from the same sites as the rhizobia  
283 strains, and thus, are unlikely to naturally co-occur. Overall, our results suggest that trait optima  
284 can be shared even in one-to-many interactions, and that fitness feedbacks operating to align host  
285 and symbiont fitness are present even in diverse communities irrespective of coevolutionary history.

286 **Genomic resolution of conflict and alignment in symbiosis.** Rather than to identify causal loci  
287 underlying symbiotic pleiotropy, the goal of our study was to examine broader patterns of fitness  
288 alignment or conflict across the genomes of numerous naturally occurring rhizobium strains. We  
289 found abundant concordant pleiotropic variants associated with both host and symbiont fitnesses,  
290 alongside evidence that selection has acted to maintain genetic variation within rhizobium genomes  
291 through time and/or space. A lack of discordant pleiotropic variants, like we have found, may have  
292 resulted from physiological constraints that make such variants impossible (i.e., alleles associated with  
293 larger nodules and small host biomass are rare or non-existent), or because of correlational selection  
294 that removes discordant variants from the population altogether (70, 71). In addition to direction,  
295 the extent of symbiotic pleiotropy (i.e., the number of pleiotropic variants) can inform whether  
296 traits are likely to evolve together or independently. Fitness proxy pairs with abundant pleiotropy  
297 (e.g., nodule size vs. shoot biomass) suggest a highly polygenic basis governed by many small-effect  
298 pleiotropic variants; other proxies with little pleiotropy (e.g., nodule number vs. chlorophyll) are  
299 largely governed by different sets of genetic mechanisms, and are thus likely to evolve independently.

300 One of our more critical findings, that the loci associated with both high host and symbiont fitness  
301 benefits (i.e., fitness alignment) largely differed across host genotypes, provides one solution to the  
302 mutualism paradox: if host genotypes act as distinct selective environments for rhizobia, meaning  
303 that the "best" symbiont genotype differs among host genotypes, then symbiont diversity could be  
304 maintained in the face of host selection. Host-dependent loci underlying fitness alignment were  
305 also found in experimentally evolved rhizobia isolates (20) and host-mediated balancing selection  
306 was previously proposed as a mechanism maintaining rhizobial diversity in native populations of  
307 *Bradyrhizobium* (72). Our results provide a solution to the mutualism paradox: symbiont diversity  
308 can be maintained via balancing selection acting on host-dependent rhizobium loci underlying fitness  
309 alignment.

310 We nonetheless found several instances of fitness conflict at the genomic level despite alignment at

311 the phenotypic level (e.g., shoot biomass vs rhizobium relative fitness on DZA). Linkage disequilibrium  
312 (LD) could lead to discordant alleles being “packaged” into multilocus genotypes that show fitness  
313 alignment. Such LD results from multiple, non-mutually exclusive factors including epistatic  
314 interactions among individual variants that render discordant variants effectively neutral, and/or  
315 past selection favouring allelic combinations that increase both host and symbiont fitness (and  
316 disfavours discordant combinations; [73](#), [74](#)).

317 Overall, these findings highlight the polygenic nature of symbiotic extended phenotype variation  
318 in nature — where the collective action of individual mutations, their additive and nonadditive  
319 effects ([42](#)), and a history of selection shapes the trait variation currently present in natural  
320 populations ([75](#), [76](#)). However, because of the limitations inherent within GWAS, including linkage  
321 disequilibrium that makes it difficult to pinpoint causal variants as well as false positive and negative  
322 associations, we acknowledge that the function of any specific locus identified here needs to be further  
323 validated in follow up experiments. Additionally, we only focus on allelic substitutions rather than  
324 presence/absence variation generated via gene gain and loss, the latter of which previous studies  
325 have found to be associated with exploitative traits in rhizobia (e.g., [68](#)).

326 **Trade-offs among rhizobium fitness proxies and the rhizobium competition problem.** Under-  
327 standing how microbial symbioses, which are ubiquitous and important in nature, evolve (or coevolve)  
328 requires accurate estimates of symbiont fitness in ecologically realistic settings. Given our evidence at  
329 both the phenotypic and genomic levels for trade-offs among rhizobium fitness proxies, and because  
330 diverse strains of rhizobia co-occur in nature, relative fitness proxies should be used whenever possible  
331 ([25](#)). Nevertheless these proxies are not replacements for those measured in single-strain experiments  
332 because they cannot be used to assign individual genetic means for whole-plant measures of host  
333 benefit (plant biomass and seed number) to individual strains (e.g., [1](#), [26](#), [39](#), [42](#)), necessitating that  
334 host benefit and rhizobium fitness be measured on separate individuals.

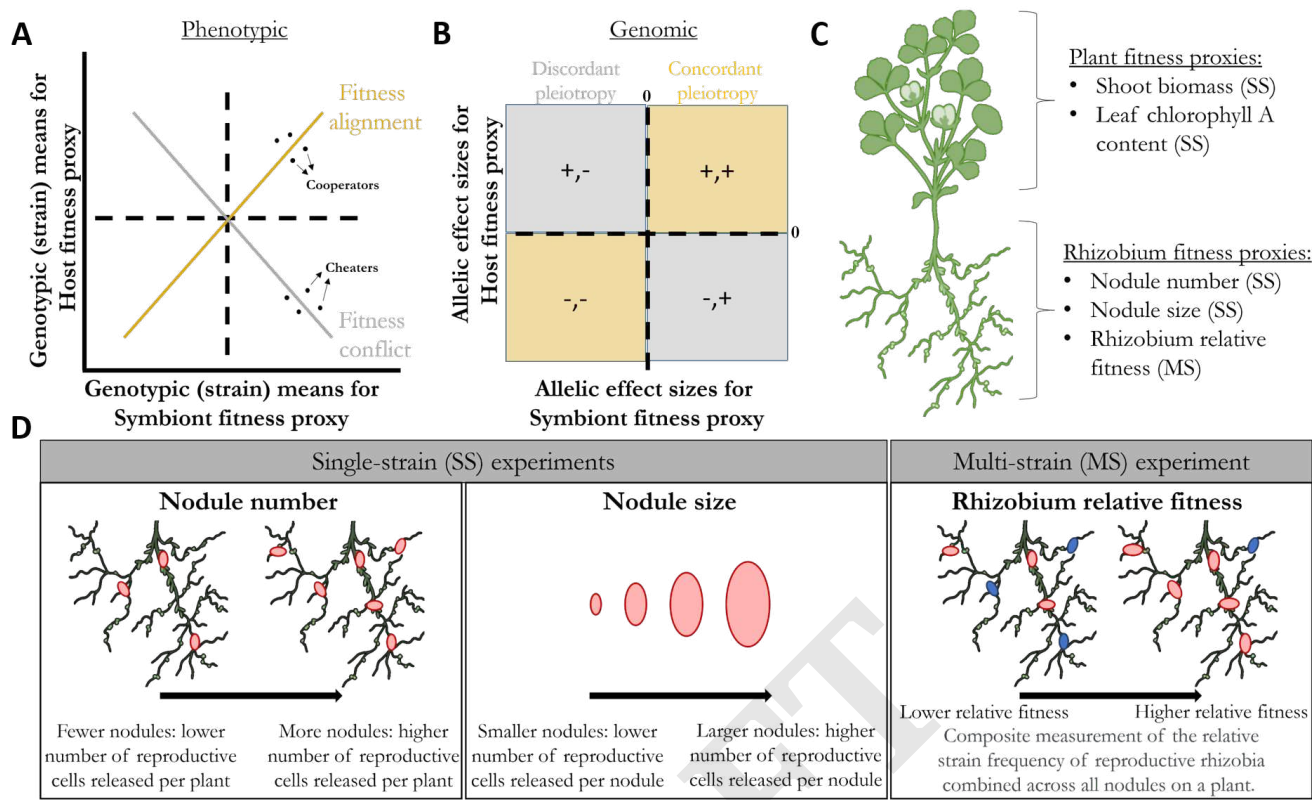
335 Together our results suggest that the genetic architectures associated with rhizobium fitness  
336 proxies, and their relationships, are host-dependent, and thus that their evolutionary trajectories are  
337 influenced not only by variation in the rhizobium’s genome, but also by variation in host traits. For  
338 example, host genotypes could differ in their ability to exert sanctions or partner choice, quantitative  
339 traits known to vary in host populations ([44](#), [77](#)). At the genomic-level, distinct variants underlying  
340 rhizobium fitness pleiotropy on each host genotype suggests that the genetic mechanisms (i.e.,  
341 genes, pathways, metabolic processes) governing the relationship between fitness proxies are largely  
342 non-overlapping when rhizobia interact with different hosts. Such host genotype-dependent shifts  
343 in the rhizobium genetic architecture of these symbiotic extended phenotypes are supported by

344 transcriptomics studies of G x G interactions (25, 40) and GWAS revealing distinct sets of candidate  
345 genes in different host backgrounds (26, 42). Similar genetic variation exists in hosts (78) and  
346 undoubtedly interacts with the variants we identify here, thus, should be accounted for if we want  
347 to uncover the multi-genomic basis of symbiotic extended phenotypes.

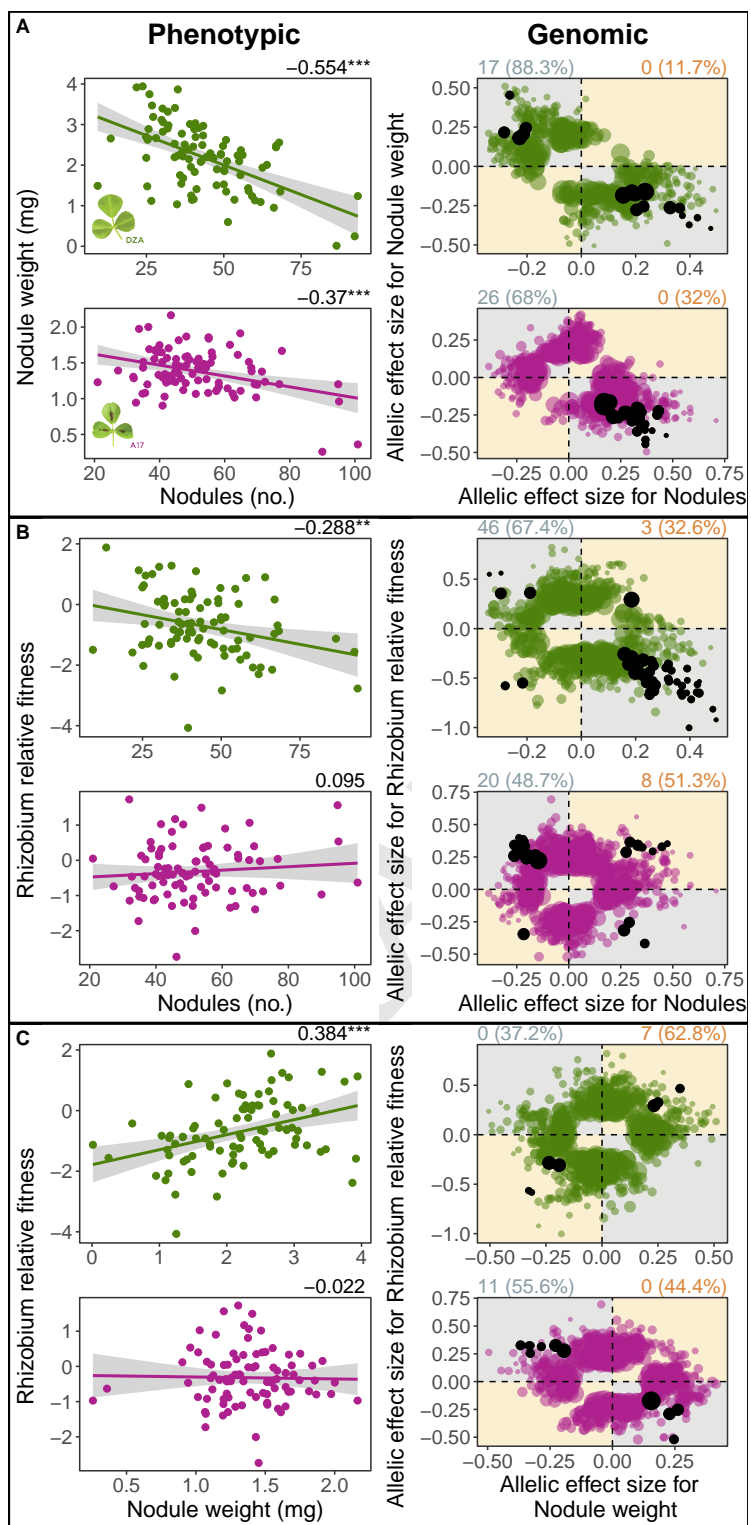
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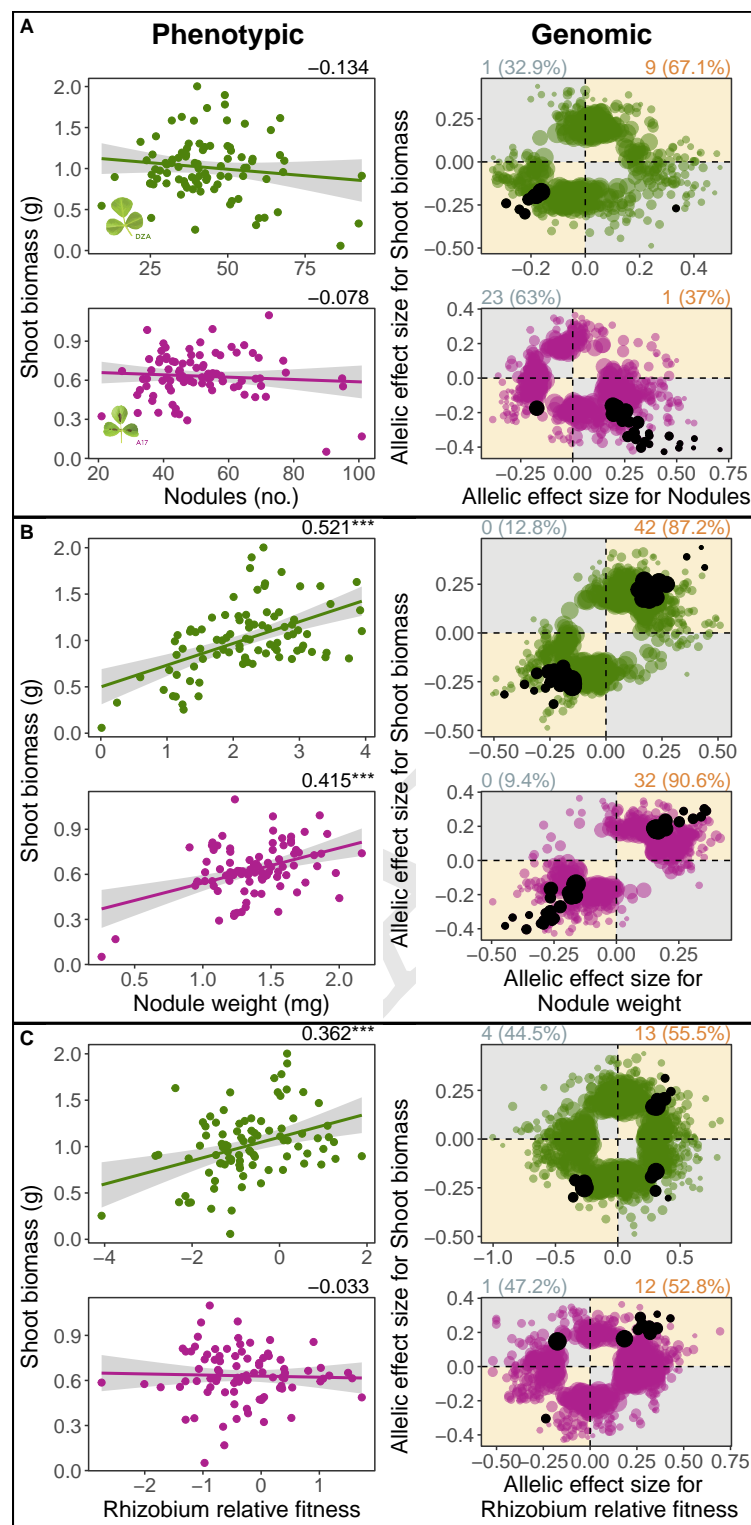
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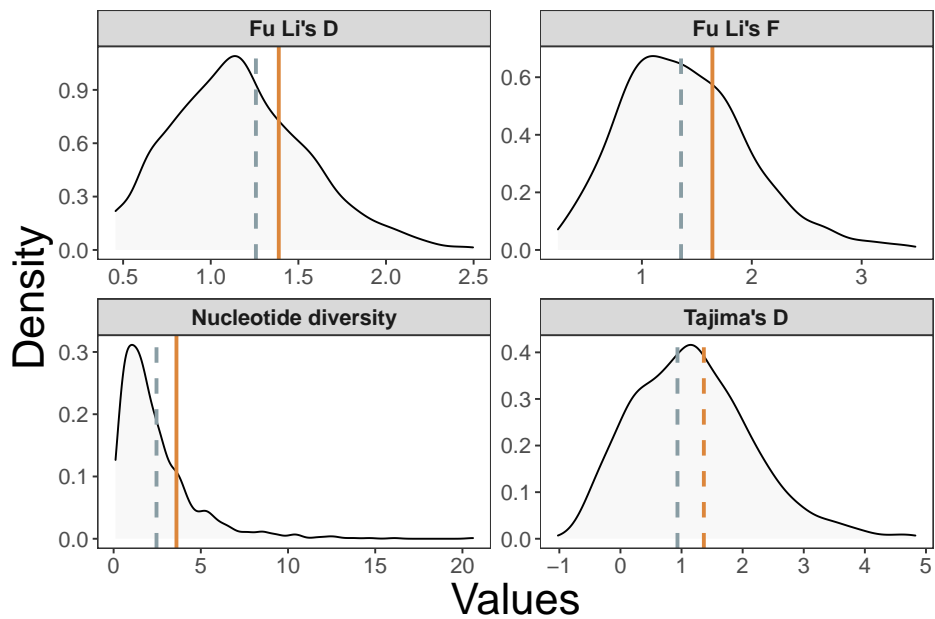
**Fig. 1. Interpreting patterns of fitness alignment and conflict at both the phenotypic and genomic levels.** **A)** Phenotypic-level correlations of genotypic (strain) means for rhizobium fitness proxies (e.g., nodule number, nodule size, rhizobium relative fitness) or plant fitness proxies (shoot biomass or leaf chlorophyll A content) on the x- and y-axes, respectively. The positive (orange) or negative (grey) correlations represent fitness alignment or conflict, respectively. **B)** Genomic-level correlations of allelic effect sizes determined in GWAS for rhizobium fitness proxies or plant fitness proxies on the x- and y-axes, respectively. Points appearing in orange or grey quadrants represent pleiotropic variants with concordant (same-sign) or discordant (opposite sign) effects, respectively, on both host and symbiont. **C)** Plant and rhizobium fitness proxies measured, parentheses indicating the experiment type for which proxies were measured (SS = single-strain; MS = multi-strain). **D)** Rhizobium fitness proxies corresponding to the pink rhizobium strain. In single-strain experiments, nodule number indicates the number of reproductive cells released per plant, while nodule size indicates the number of reproductive cells released per nodule. In the multi-strain experiment, rhizobium relative fitness is a composite metric that combines competition among strains for nodule occupancy and for host resources once in nodules. Nodules pink versus blue in colour are used here to illustrate two different rhizobium strains competing, however, a total of 89 strains were inoculated together onto plants in the multi-strain experiment.



**Fig. 2. Trade-offs among rhizobia (*Ensifer meliloti*) fitness proxies prevail at both the phenotypic (left panels) and genomic-levels (right panels).** **Phenotypic:** Genetic correlations between pairwise fitness proxies measured on plant lines DZA (green, top rows) or A17 (pink, bottom rows), based on 89 *Ensifer meliloti* strains. Dots represent estimated marginal strain means for nodule number and nodule weight, both being measured in single-strain experiments, or medians for rhizobium relative fitness measured in multi-strain experiments. Numbers at top right of each correlation represent Pearson correlation coefficients, while asterisks represent significance: \* =  $p < 0.05$ ; \*\* =  $p < 0.01$ ; \*\*\* =  $p < 0.001$ . **Genomic:** Dots represent allelic effect sizes (i.e., beta scores calculated in GEMMA), those falling along the diagonal (orange quadrants) or off-diagonal (grey quadrants) represent variants with concordant or discordant effects, respectively. Coloured dots represent variants that were significantly associated with one of the two fitness proxies, while black dots represent pleiotropic variants, i.e., significantly associated with both fitness proxies. Numbers outside of and percentages within parentheses at the top left and right of each plot represent the pleiotropic variant counts and the proportion of total significant variants, respectively, that are discordant (left, in grey) or concordant (right, in orange).



**Fig. 3. Fitness alignment between rhizobia (*Ensifer meliloti*) and host (*Medicago truncatula*) prevails at both the phenotypic (left panels) and genomic (right panels) levels. Phenotypic:** Genetic correlations between pairwise fitness proxies measured on plant lines DZA (green, top rows) or A17 (pink, bottom rows), based on 89 *Ensifer meliloti* strains. Dots represent estimated marginal strain means for shoot biomass, nodule number, and nodule weight, all being measured in single-strain experiments, or medians for rhizobium relative fitness measured in multi-strain experiments. Numbers at top right of each correlation represent Pearson correlation coefficients, while asterisks represent significance: \* =  $P < 0.05$ ; \*\* =  $P < 0.01$ ; \*\*\* =  $P < 0.001$ . **Genomic:** Dots represent allelic effect sizes (i.e., beta scores calculated in GEMMA), those falling along the diagonal (orange quadrants) or off-diagonal (grey quadrants) represent variants with concordant or discordant effects, respectively. Coloured dots represent variants that were significantly associated with one of the two fitness proxies, while black dots represent pleiotropic variants, i.e., significantly associated with both fitness proxies. Numbers outside of and percentages within parentheses at the top left and right of each plot represent the pleiotropic variant counts and the proportion of total significant variants, respectively, that are discordant (left, in grey) or concordant (right, in orange).



**Fig. 4. Neutrality statistics show elevated values for genes associated with fitness alignment.** Vertical lines represent the average values calculated for four separate statistics, grey for discordant and orange for concordant genes. Distributions represent the same statistics calculated for all genes containing significant variants based on GWAS. Dashed and solid lines represent non-significant ( $p > 0.1$ ) and significant ( $p < 0.1$ ) differences, respectively, between each focal gene category and all significant genes (i.e., distributions).