The Potential for Genotype-by-Environment Interactions to Maintain Genetic Variation in a Model Legume–Rhizobia Mutualism

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ABSTRACT

The maintenance of genetic variation in mutualism-related traits is key for understanding mutualism evolution, yet the mechanisms maintaining variation remain unclear. We asked whether genotype-by-environment (G×E) interaction is a potential mechanism maintaining variation in the model legume–rhizobia system, Medicago truncatula–Ensifer meliloti. We planted 50 legume genotypes in a greenhouse under ambient light and shade to reflect reduced carbon availability for plants. We found an expected reduction under shaded conditions for plant performance traits, such as leaf number, aboveground and belowground biomass, and a mutualism-related trait, nodule number. We also found G×E for nodule number, with ~83% of this interaction due to shifts in genotype fitness rank order across light environments, coupled with strong positive directional selection on nodule number regardless of light environment. Our results suggest that G×E can maintain genetic variation in a mutualism-related trait that is under consistent positive directional selection across light environments.

Key words: mutualism evolution, genetic variation, genotype-by-environment interaction, Medicago truncatula, light availability, partner quality

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INTRODUCTION

Characterizing the forces that maintain genetic variation in the face of ongoing selection is a long-standing, classic question in evolutionary biology (Walsh and Blows, 2009). For mutualisms—interspecific interactions that benefit both species—answering this question is especially challenging. By definition, the traits that promote mutualisms are positively related to fitness, and thus variation for fitness-related traits should be reduced (Charlesworth, 1987; Kruuk et al., 2000; Heath and Stinchcombe, 2014). Moreover, the mechanisms thought to promote stability and maintenance in mutualisms are themselves predicted to reduce variation in mutualist quality (Heath and Stinchcombe, 2014). One straightforward mechanism capable of maintaining variation in mutualisms, like all traits, is genotype-by-environment (G×E) interaction (Gillespie and Turelli, 1989), although we have few examples of whether G×E maintains variation in mutualism-related traits.

Genotype-by-environment interactions can occur either through changes in the rank order of genotype fitness or in the magnitude of genetic variance across environments (Bowman, 1972; Muir et al., 1992; Hühn et al., 1993; Des Marais et al., 2013). Only changes in the rank order of genotype fitness can contribute to the maintenance of genetic variation, because genotypes that perform well in one environment are not the same genotypes that perform well in another (Mitchell-Olids, 1992; Johnson, 2007). In contrast, G×E interactions that are mostly due to changes in genetic variance might affect the response to selection across environments (Fisher, 1958) but will not effectively contribute to the maintenance of genetic variation. Several studies have tested for G×E interactions in a variety of symbiotic systems, such as plant–fungi systems (Ahlholm et al., 2002; Werner et al., 2018), the wasp–Wolbachia symbiosis (Mouton et al., 2007), ant–plant mutualisms (Abdala-Roberts et al., 2012; Abdala-Roberts and Mooney, 2013), and legume–rhizobia mutualisms (Heath and Tiffin, 2007; Heath et al. 2010; Ossler and Heath, 2018; Batstone et al., 2020; Heath et al., 2020; Magnoli and Lau, 2020). Most of these studies concluded

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that G×E could contribute to genetic variation in mutualism-related traits and used crossing reaction norms as qualitative support. However, only two of these studies have explicitly tested the maintenance of genetic variation by including a quantitative assessment of the relative contribution of genotype rank order shifts versus changes in genetic variance across abiotic environments (e.g., Batstone et al., 2020; Heath et al., 2020).

Genetic variation in mutualisms could also be maintained through environmental heterogeneity in selection (Mitchell-Olms et al., 2007; Heath and Stinchcombe, 2014). Shifts in resource availability can alter the costs, benefits, and ultimately the strength of the mutualistic interaction, which could potentially cause changes in the magnitude and/or direction of selection on mutualism-related traits across environments (Thrall et al., 2007; Batstone et al., 2018). Changes in nutrient availability have been shown to reduce the net benefits for hosts in a variety of mutualistic systems, including coral, lichen, plant–fungi, and legume–rhizobia (Shantz et al., 2016). For example, in some legume–rhizobia systems, nitrogen enrichment decreased the net benefits provided by rhizobia to plant hosts, causing reduced investment by hosts in the mutualism (Regus et al., 2015, 2017; Weese et al., 2015). Such a reduction in mutualism investment could lead to relaxed selection by hosts for mutualistic partners, resulting in increased mutualism variation (or slowing the loss of variation) over time. Van Cauwenberghhe et al. (2015) showed that long-term nitrogen addition led to increased rhizobial genetic diversity in the legume–rhizobia system Vicia cracca–Rhizobium leguminosarum. In addition, Simonson and Stinchcombe (2014b) found a reduction in fitness costs for legumes that hosted ineffective rhizobia in the presence of herbivory, which could directly lead to reduced selection against ineffective partners and maintain mutualism variation. Batstone et al. (2020) explicitly tested for environmental heterogeneity in selection for a legume–rhizobia mutualism through a selection-by-environment (S×E) interaction analysis and found that the magnitude of selection for a mutualism-related trait varied across greenhouse and field plot conditions.

A common and useful model for studying the effects of genetics and the environment on the maintenance of mutualism variation is the legume–rhizobia mutualism. It is a resource-based mutualism, with leguminous plants providing carbon and shelter in root nodules, and mutualistic root-dwelling bacteria providing fixed nitrogen (Kiers et al., 2003; Gorton et al., 2012). There is a vast amount of information on the mechanisms regulating legume–rhizobia mutualisms—from genomic, cellular, to the individual level—making it a powerful system for studying mutualism evolution (Jones et al., 2007). In addition, abiotic and biotic factors can be effectively manipulated to shift the costs and benefits of the interaction due to resource availability (Bronstein, 1994; Heath and Stinchcombe, 2014). However, environmental manipulations of the legume–rhizobia mutualism tend to be largely one-sided, with most studies focusing on effects of soil nitrogen (N) availability due to its ecological and agricultural importance in artificial fertilizer use (e.g., Heath, 2010). The results of these studies tend to show variable outcomes for G×E, with some finding a significant G×E interaction with varying N availability (Heath and Tiffin, 2007; Heath, 2010), and others showing that plant hosts either consistently associated with (Grillo et al., 2016) or discriminated against (Regus et al., 2014) rhizobial strains regardless of N environment. The availability of carbon (C) resources is also equally important for understanding the relationship between legumes and rhizobia, but literature addressing this is scarce. Elevated CO₂ (eCO₂) has gained recent interest, but its effects on legume–rhizobia mutualisms also show inconsistent results depending on host species or genotype (Lüschers et al., 2000; West et al., 2005), rhizobial genotype (Bertrand et al., 2007), duration of eCO₂ exposure (Hungate et al., 2004; Rogers et al., 2006), or the surrounding environment (West et al., 2003). However, the results on eCO₂ do indicate that partner genotype and C availability have the potential to influence mutualism efficiency and genetic variation for mutualism-related traits.

An alternative way to test for the influence of C availability on mutualisms is through shifts in light availability. The physiological effects of light condition on plants are well documented (e.g., Evans, 1989; Cronin and Lodge, 2003), and individuals of most plant species show variation in their plastic responses to light (Winn and Evans, 1991; Sultan and Bazzaz, 1993). In addition, both G×E and adaptive plastic responses to light quality and quantity are well known (Donohue et al., 2000; Schmitt et al., 2003; Heschel et al., 2004; Stinchcombe et al., 2010). If legumes respond differently to varying light conditions like most plant species, it could affect how legume hosts allocate C resources and ultimately associate with rhizobia under varying light conditions (Schwartz and Hoeksema, 1998). A few studies have tested the effects of light availability on legume–rhizobia mutualisms using one-to-one pairings, but with variable results. Some of these studies suggested that maintaining the association becomes costly for plant hosts under low light, resulting in a shift from mutualism to parasitism (Lau et al., 2012; Ballhorn et al., 2016), while another suggested that the mutualistic association becomes commensal (Friesen and Friel, 2019). Heath et al. (2020) explicitly tested for G×E effects on legume–rhizobia mutualisms using legumes of a single genotype inoculated with one of 11 strains of rhizobia and two light environments. They showed a rhizobia genotype-by-light environment interaction for plant performance traits, but not for mutualism-related traits such as nodule number. Using several legume genotypes might more directly test for genetic variation in resource allocation and mutualism association under varying light conditions to effectively determine whether G×E interactions maintain genetic variation in mutualism-related traits.

We manipulated light availability, and hence the amount of C available for plants to provide to rhizobia, in an experiment testing for G×E in the model legume–rhizobia system, Medicago truncatula–Ensifer meliloti. Specifically, we asked the following questions: (1) Do legumes form fewer mutualistic associations with bacteria in low light (low C) environments? (2) Is there G×E for plant performance and mutualism-related traits in response to low light, and are these interactions capable of maintaining genetic variation? And (3) How does natural selection act on morphological and mutualistic traits in contrasting light environments? Here, we present the results of an experiment testing for G×E interactions in a plant–bacteria mutualism, quantitatively examining their potential to maintain variation in mutualism-related traits.
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Table 1. Summary Statistics for the Effects of Light Treatment and Plant Genotype on Plant Performance and Mutualism-Related Traits.

|                      | Leaf Number | Aboveground Biomass | Belowground Biomass | Aboveground:Belowground Ratio | Nodule number |
|----------------------|-------------|---------------------|---------------------|-------------------------------|--------------|
| **Fixed effects**    |             |                     |                     |                               |              |
| Light treatment      | $F(1,9.2944)$ | $F(1,13.209)$       | $F(1,13.281)$       | $F(1,9.0881)$                 | $F(1,11.67)$ |
|                      | 3.75        | 18.33***            | 19.16***            | 19.42**                       | 23.19**      |
| **Random effects**   |             |                     |                     |                               |              |
|                      | $\chi^2_{df=1}$ | $\chi^2_{df=1}$       | $\chi^2_{df=1}$       | $\chi^2_{df=1}$                 |              |
| Plant genotype       | 14.27***    | 5.16*               | 3.88*               | 2.76*                         | 8.13**       |
| GxE                  | 0.63        | 13.22***            | 32.45***            | 0.20                          | 5.42**       |

RESULTS

Poor germination and establishment led to a final sample of 666 plants that survived until the end of the experiment. While this germination and establishment fraction was lower than our past work with Medicago (and these lines), we verified that the total fraction of germinants was not correlated with the focal traits analyzed below ($| \rho | < 0.19$, $p > 0.17$), suggesting that germination and establishment variation did not bias our estimates of plant performance or nodulation. The experiment was terminated at 9 weeks to prevent nodule senescence (Puppo et al., 2005; El Msehli et al., 2019), at which point 409 (61%) plants had flowered. We found $25 \pm 6.50$ (mean $\pm$ S.E.) nodules in the uninoculated control plants, which suggests some rhizobial transfer among individuals within each block. However, this will not affect the results since all experimental plants received the same inoculation treatment. We excluded uninoculated control plants in the analysis of our data.

Plant Genotype and Light Environment Affect Plant Growth

The productivity and growth of plant hosts were estimated via leaf number, and aboveground and belowground biomass. Plants growing in shaded conditions produced marginally fewer number of leaves than ambient plants, showing a $\sim30\%$ decrease in leaf number from ambient to shade conditions (ambient $7.39 \pm 0.44$ versus shade $5.18 \pm 0.17$, Table 1 and Figure 1A). However, plant genotype had a significant effect on leaf number (Table 1), with the average number of leaves at harvest ranging from three to ten leaves across genotypes. There was no significant GxE for leaf number (Table 1 and Figure 1B).

Plants that received full light were significantly larger on average than plants that were shaded. Aboveground biomass decreased by $\sim61.5\%$ from ambient to shaded conditions (ambient $0.22 \pm 0.024$ g versus shade $0.085 \pm 0.0086$ g, Table 1 and Figure 1C), but was significantly affected by plant genotype and GxE (Table 1 and Figure 1D). Belowground biomass also decreased by $\sim72\%$ from ambient to shade (ambient $0.083 \pm 0.0084$ g versus shade $0.023 \pm 0.0012$ g, Table 1 and Figure 1E), and showed a significant genotype and GxE effect (Table 1 and Figure 1F). The relative contribution of genotype rank order shifts in the GxE interaction was $\sim31.6\%$ for aboveground biomass and $\sim12.5\%$ for belowground biomass (Supplemental Table 1), suggesting that a majority of the GxE was due to changes in the magnitude of genetic variance between environments; specifically, an increase in variance in the ambient treatment compared with the shade treatment (Figure 1D and 1F).

We also analyzed the ratio of aboveground to belowground biomass as an index of plant investment into different components of growth. Light availability significantly affected this ratio, but the response was in the opposite direction of the other traits. Aboveground:belowground ratio increased by $\sim50.3\%$ from ambient to shade (ambient $2.470.11$ g versus shade $3.710.24$ g, Table 1 and Figure 1G), indicating increased allocation to aboveground growth in the light-limited conditions. There was a marginal effect of plant genotype and no significant treatment-by-genotype interaction on aboveground:belowground biomass (Table 1 and Figure 1H).

Plant Genotype-Light Environment Interaction Affects Nodule Number

Nodule number showed a significant main effect of light treatment and genotype, as well as a significant GxE interaction effect (Table 1). Plants in the shade had about 20 fewer nodules on average than in full light ($\sim63\%$ decrease, ambient $32.05 \pm 2.51$ versus shade $11.85 \pm 0.73$, Figure 1I). Approximately 83.3% of the significant GxE interaction was explained by a change in genotype rank order between treatments (Supplemental Table 1). Most, but not all, plant genotypes showed higher average nodule number in the ambient treatment than shade, and the magnitude of this plastic response was variable among genotypes (Figure 1J).

Genetic Correlations between Traits and across Environments

There was a significant positive genetic correlation between most traits within each light treatment (Figure 2). Our focal mutualism-related trait, nodule number, was significantly positively correlated with all plant performance traits in both ambient and shade environments, except the aboveground:belowground ratio, which showed no significant correlation. When comparing across light treatments, all traits showed a significant positive correlation between the shade and ambient environments ranging from 0.34 for aboveground:belowground ratio to 0.76 for belowground biomass (Figure 2).
Selection for Nodule Number Does Not Differ between Light Environments

All three methods of our selection analyses (absolute, global, and local) revealed a significant $S \times E$ interaction for leaf number, while showing that selection for nodule number did not differ between light environments (Table 2 and Figure 3). The global and local analyses showed inconsistencies for belowground biomass, with a non-significant $S \times E$ interaction in the global analysis and significant $S \times E$ in the local analysis (Table 2 and Figure 3F and 3I, respectively). Estimates of the selection gradient ($\beta$) from the local analysis indicated positive directional selection for leaf number in both environments, but the magnitude of selection was significantly greater in the shade than ambient (Table 3 and Figure 3H). For nodule number, both the magnitude and direction of selection remained similar across light environments (Table 3 and Figure 3G).

DISCUSSION

Explaining the maintenance of variation in mutualisms remains a largely unresolved topic in evolutionary ecology, with several potential mechanisms that require testing (Heath and Stinchcombe, 2014). We used a manipulative experiment to test for $G \times E$ in mutualism-related traits, driven by changes in light and hence carbon availability. We found predicted effects of reduced plant performance (leaf number, and aboveground and belowground biomass) and mutualism-related traits (nodule number) under low light conditions, along with $G \times E$ interactions for nodule number driven by appreciable changes in rank order rather than variance. Our selection analyses suggest that nodule number is under strong positive directional selection in each environment, and this was consistent across all three methods of scaling: absolute, global (across environment), and local (within environment). There was little evidence of environmental heterogeneity in the magnitude or direction of selection for nodule number. Below, we discuss in detail two specific aspects of our main results. First, we evaluate the consequences of $G \times E$ being driven by changes in rank order and strong positive selection for nodule number on the evolution of mutualism-related traits. Second, we discuss whether plants produced fewer nodules in low light because they were smaller on a whole-plant level or as an active response against decreased carbon availability, and the implications of this result for the evolution of the mutualism.

$G \times E$ and Selection for Nodule Number

Our findings on nodule number illustrate how $G \times E$ can contribute to the maintenance of variation in mutualisms. Nodule number
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showed significant G×E, with the majority being driven by changes in rank order rather than variance. It is important to note, however, that this is not simply a function of the amount of G×E and the across environment correlation. Comparing nodule number and aboveground biomass, nodule number has a higher correlation across environments (i.e., less G×E: Cockerham, 1963), yet a much higher fraction of its G×E is due to changes in rank than variance (83% versus 31%). Nodule number was also under positive selection in each treatment, with little evidence of changes in selection depending on light availability. Consequently, genotypes that produce more nodules are strongly favored in each condition, although which genotypes are most strongly favored differs depending on light availability.

The contribution of genotype rank order shifts in nodule number variation across light environments suggests that some host genotypes invest more in the mutualism in one environment and reduce investment in the other. In addition, there was no significant shift in the magnitude or direction of selection for nodule number between light environments at both the global and local scale (Table 2); there was consistent positive directional selection in both light environments (Table 3 and Figure 3). A lack of S×E interaction coupled with a significant G×E due to genotype rank shifts can maintain genetic variation in nodule number because selection can favor plant genotypes with high nodule production in both light environments, but the genotypes that have high nodule numbers in one environment are not identical to those with high nodule numbers in the other. Furthermore, the genetic correlation for nodule number across environments was strongly positive (but less than 1: Figure 2), which indicates that the groups of genotypes that produce more or less nodules will be similar across environments, but that G×E with rank order shifts can affect which individual genotypes are selected for or against in each environment. Thus, a genotype rank order shift between light environments under positive directional selection could maintain genetic variation in the legume–rhizobia mutualism.

To precisely determine whether G×E is effective at maintaining genetic variation in the legume–rhizobia mutualism, it is also important to consider the frequency of selective environments experienced by natural populations (Gomulkiewicz and Kirkpatrick, 1992). Determining the natural frequency of selective environments proves to be challenging but it is critical for understanding whether the environmental context dependency we found is realistic. The evolutionary response of plastic traits can change drastically with the frequency of selective environments (McDonald and Ayala, 1974; Via and Lande, 1985; Gomulkiewicz and Kirkpatrick, 1992; Arnold and Peterson, 2002; Kelley et al., 2005; Stinchcombe et al., 2010). Our results indicate that plant hosts will experience positive directional selection for nodule number regardless of light environment, but the frequency of light environments will determine which genotypes will be selectively favored, and whether genetic variation will be maintained. For example, if individuals experience full light conditions more frequently than shade, the effects of G×E will no longer be a factor maintaining genetic variation; selection will act on genotypes only as they are ranked in the ambient treatment. Moreover, because there is more genetic variation expressed under ambient light than reduced light, we would expect larger responses to selection in those environments. Natural populations of *Medicago truncatula* experience a wide range of habitats across the Mediterranean (Batallion and Ronfort, 2006); however, information on the frequency of light conditions experienced by these populations is limited. Future studies must consider how frequently legume hosts experience varying light conditions in nature to determine whether G×E can effectively maintain genetic variation in the legume–rhizobia system.

### Reduced Nodule Number under Low Light: Causes and Implications

We found a significant average decrease in the number of nodules—the sites of resource exchange in the mutualism and our proxy for the prevalence of the mutualistic interaction (Heath and Tiffin, 2009). There was a ~63% reduction in nodule number from the ambient to shade treatment (Figure 1I), which could either be part of a whole-plant level response to low light

Table 2. Genetic Correlation Matrix between Traits and across Environments.
Correlations are of raw line means for each trait within the ambient (gold) and shade (green) treatments. Dark bordered boxes show between treatment correlations for each trait. Numbers present correlation coefficients, which are also represented by the color legend on the right. Insignificant correlations (p > 0.05) are indicated by the absence of an ellipse.
availability or an active response by plant hosts to reduce mutualistic interactions. A whole-plant level response is possible since we also found an average reduction in our plant performance traits from the ambient to shade treatment: leaf number, aboveground biomass, and belowground biomass, by approximately 30%, 61.5%, and 72%, respectively (Figure 1). Low light availability affects photosynthetic activity, respiration rates, and resource exchange and allocation, which can drastically influence whole-plant level traits, such as leaf number and biomass (Corrê, 1983; Givnish, 1988; Sultan and Bazzaz, 1993; Stuefer and Huber, 1998; Cronin and Lodge, 2003; Casal, 2012). However, our statistical models for nodule number included total biomass as a covariate, and we still found a significant average reduction under low light conditions, indicating that reduced allocation to rhizobia might be an active response to low light and low carbon availability.

A reduction in nodule number from ambient to shade could suggest that plant hosts invested less into the mutualistic interaction in low light conditions. The possibility of reduced mutualistic investment in low light is consistent with classic theoretical work that predicts shifts in mutualistic interactions based on the availability of traded resources (Trivers, 1971; Axelrod and Hamilton, 1981; Schwartz and Hoeksema, 1998; Sachs and Simms, 2006; Nathaniel Holland and DeAngelis, 2009). The carbon resources that plant hosts provide to their rhizobial partners may become less available in shaded conditions due to reduced photosynthetic activity and shifts in resource allocation from belowground to aboveground growth (e.g., Evans, 1989; Cronin and Lodge, 2003). Our experimental legumes showed a greater aboveground:belowground biomass ratio in the shade compared with full light conditions (Figure 1G), suggesting that resources in the shade may have been diverted toward aboveground growth as opposed to belowground, which could lead to reduced nodule production under light-limited conditions. A reduction in nodule number could also be a response by the host to minimize costs of the mutualistic association due to reduced plant performance at low light. Studies on plant–fungal (Zheng et al., 2015) and legume–rhizobial (Regus et al., 2015) systems at varying light conditions showed that, as the mutualistic benefits provided to hosts decreased at low light, so did host investment in the mutualism, which could weaken the interaction under low light availability.

A more precise assessment of whether light availability has a direct effect on the mutualism would require measuring the

| Model     | Term                  | Sum of Squares | F_{(1,92)} |
|-----------|-----------------------|----------------|------------|
| Absolute  | Intercept             | 0.018206       | 17.7636*** |
|           | Treatment             | 0.002129       | 2.0772     |
|           | Nodule number         | 0.006355       | 6.2011*    |
|           | Leaf number           | 0.014393       | 14.0437*** |
|           | Belowground biomass   | 0.065165       | 63.5825*** |
|           | Nodule × treatment    | 0.000000       | 0.0002     |
|           | Leaf × treatment      | 0.006106       | 5.9578*    |
|           | Belowground × treatment| 0.000700      | 0.6834     |
| Global    | (Intercept)           | 23.0933        | 539.2206***|
|           | Treatment             | 0.8351         | 19.4995*** |
|           | Nodule number         | 0.2656         | 6.2011*    |
|           | Leaf number           | 0.6015         | 14.0437*** |
|           | Belowground biomass   | 2.7231         | 63.5825*** |
|           | Nodule × treatment    | 0.0000         | 0.0002     |
|           | Leaf × treatment      | 0.2552         | 5.9578*    |
|           | Belowground × treatment| 0.0293       | 0.6834     |
| Local     | (Intercept)           | 100.000        | 1456.8018***|
|           | Treatment             | 0.0000         | 0.0000     |
|           | Nodule number         | 0.571          | 8.3197**   |
|           | Leaf number           | 0.982          | 14.3004*** |
|           | Belowground biomass   | 8.684          | 126.5117***|
|           | Nodule × treatment    | 0.019          | 0.2830     |
|           | Leaf × treatment      | 0.449          | 6.5364*    |
|           | Belowground × treatment| 0.854       | 12.4383*** |

Table 2. Selection-by-Environment Interactions Using Three Methods of Scaling Fitness (Aboveground Biomass) and Traits. ANCOVA F_{(num df, denom df)} are presented for three linear models: absolute fitness, globally (across light treatments), and locally (within light treatment) relativized fitness and standardized traits. Significant values are in bold and coded as: ***p < 0.001, **p < 0.01, *p < 0.05.
amount of costly photosynthetic carbon allocated to rhizobia and the relative exchange of carbon and nitrogen resources in shaded versus ambient environments. Nevertheless, our results suggest that legume hosts interact less with their rhizobial partners under low light, regardless of whether this is a direct or indirect response. The prevalence of the mutualism is reduced under low light and carbon conditions, but whether this response has any substantial effect on the evolution of the mutualism depends on how frequently and consistently the system experiences low light conditions.

METHODS

Study System

Medicago truncatula Gaertn. is an annual legume species native to the Mediterranean (Bataillon and Ronfort, 2006). It has a short generation
time, produces large uniform nodules, and a small genome size, with a growing availability of genomic tools and methods for studying legume–rhizobia mutualisms (Barker et al., 1990; Blondon et al., 1994; Cook, 1999; Stanton-Geddes et al., 2013). It is also a highly selfing species, which makes it possible to grow nearly genetically identical individuals of a single accession in multiple environments, facilitating studies of G×E. We used 50 M. truncatula genotypes supplied by the French National Institute for Agricultural Research in Montpellier, and the United States Department of Agriculture—Agricultural Research Services in Washington. We co-inoculated the legumes with two strains (Em1021 and Em1022) of the mutualistic rhizobial species, Ensifer meliloti. These strains differ in their ability to fix N (Batstone et al., 2017), making the rhizobial environment relatively more complex and realistic than a single strain inoculation. We used lab stocks of Em1021, while Em1022 was supplied by Batstone et al. (2017).

**Experimental Design**

We conducted a manipulative greenhouse experiment in the Earth Sciences Centre at the University of Toronto. We applied two light treatments: ambient and shade, where plants in the ambient treatment were exposed to normal greenhouse conditions (16:8 h light:dark cycle, 22°C day and 18°C night temperatures), and those in the shade treatment were covered with neutral shade cloth that blocked 70% of light. The experiment was set up as a split-plot randomized design, where each block contained both the ambient and shade treatments separated into two racks. We used one individual per genotype, placed in random locations in each rack, for a total of 50 experimental individuals per rack. To test for contamination among plants, an uninoculated control from an external source was included in each rack. The experiment was replicated two times, giving a total of 1020 plants (1000 experimental plants and 20 contamination control plants).

**Medicago truncatula Planting and Data Collection**

We prepared M. truncatula seedlings following standard protocols (Garcia et al., 2006; Simonsen and Stinchcombe, 2014a; Wood et al., 2018). We scarified seeds and sterilized them in bleach and 95% ethanol. We then imbibed seeds in distilled water for 30 min before placing them in 1% agar plates for stratification at 4°C for approximately 13 days.

Once stratified, we incubated the seeds in the dark at room temperature (~20°C) for approximately 24 h to allow for radial elongation, then exposed them to light for another hour to initiate chlorophyll production. The germinated seedlings were then moved to the greenhouse and planted into individual autoclaved 120 ml Conetainers plugged with polyester fiber (pillow stuffing) and filled with sand (Wood et al., 2018). Within 2 days of planting, the seedlings were co-inoculated with a mixture of Ensifer meliloti strains, Em1021 and Em1022.

**Statistical Analysis**

We used R (version 3.6.2, R Core Team, 2019) to conduct analyses for the effects of light environment and plant genotype on the plant performance traits, leaf number, aboveground and belowground biomass, and on the mutualism-related trait, nodule number. We used a linear mixed model (LMM) with log-transformed response variables using the lmer function from the lme4 package (Bates et al., 2015). Untransformed data did not meet parametric tests for normality, homoscedasticity, and linearity (Wood et al., 2018), and model diagnostics for generalized LMMs showed a misspecification with the data using the DHARMa package (version 0.2.7, Hartig, 2020). Our LMMs included light treatment as the fixed effect and random effects of genotype, block, and genotype-by-treatment and block-by-treatment interactions. We conducted significance tests using the Anova function from the car package (3rd edition, Fox and Weisberg, 2019) with mixed effects analyzed using the F statistic and type III sum of squares. We analyzed the significance of

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**Table 3. Summary Statistics of Local Scale (within Light Treatment) Selection Analysis.**

The summary presents selection gradient (β) estimates with standard errors (SE) and t values, as well as ANCOVA sum of squares and F_{(num df, denom df)} estimates with standard errors (SE) and t values, as well as ANCOVA sum of squares and F_{(num df, denom df)}.

Significant values are in bold and coded as: **“p < 0.001,” “p < 0.01,” “p < 0.05.”**
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random effects with log-likelihood ratio tests using the χ² test statistic to compare models with and without the effect of interest. We halved the p values for log-likelihood ratio tests because they are one-tailed tests of whether a variance is greater than zero. For analysis of nodule number, we included total biomass as a fixed effect covariate to account for con founding effects of plant size on nodule production. We also assessed belowground biomass as a covariate for nodule number to account for the effects of root size. The results for nodule number were consistent across both these models, so we report and discuss the analysis using total biomass as a covariate.

We analyzed genetic correlations among traits and across treatments using the rcorr function from the Hmisc package (version 4.4-0, Harrell, 2020) to calculate a matrix of Pearson correlation coefficients and corresponding p values. For this analysis, we used line means of raw data to obtain a correlation matrix within the two light environments. Correlation plots were made using the corplot function in the corplot package (version 0.84, Wei and Simko, 2017).

G×E interactions can be due to shifts in genotype rank order or in the magnitude of genetic variance across environments; the latter can change the magnitude of the response to selection (Fisher, 1958), but only the former can effectively maintain genetic variation. To assess whether significant G×E interactions were due to changes in rank order versus changes in the magnitude of genetic variance, we used an equation originally described by Cockerham (1963) and applied by Batstone et al. (2020):

$$V_{gij} = \frac{\sum_{i=1}^{g} \sum_{j=1}^{e} \left[ 2\sqrt{V_g} \sqrt{V_g} \left(1 - r_i\right) + \left(\sqrt{V_g} - \sqrt{V_g}\right)^2 \right]}{e(e-1)}$$

(Equation 1)

where $V_g$ is the genotypic variance component within environments $i$ and $j$, $r_i$ is the genetic correlation between environments $i$ and $j$, and $e$ is the number of environments.

The first half of (Equation 1) describes the G×E variance due to changes in genotype rank order, which can be isolated as:

$$V_{rank} = \frac{\sum_{i=1}^{g} \sum_{j=1}^{e} \left[ 2\sqrt{V_g} \sqrt{V_g} \left(1 - r_i\right) \right]}{e(e-1)}$$

(Equation 2)

The second half of (Equation 1) describes how changes in the magnitude of genetic variance contribute to G×E, which can be rewritten as:

$$V_{variance} = \frac{\sum_{i=1}^{g} \left[ \left(\sqrt{V_g} - \sqrt{V_g}\right)^2 \right]}{e(e-1)}$$

(Equation 3)

We obtained $V_{rank}$ and $V_{variance}$ using Equations 2 and 3, respectively, and estimated the relative portion of total G×E due to changes in rank ($V_{rank} / V_{total}$) and $V_{variance}$ ($V_{variance}$), following Batstone et al. (2020). To obtain genotypic variance components ($V_g$), we used LMMs of log-transformed data within each treatment with genotype as the main random effect, as well as a random effect of block. We obtained Pearson correlation coefficients between environments ($r_i$) using the cor function from the stats package (version 3.6.2, R Core Team, 2019), with raw line means that were also log-transformed to remain consistent across all G×E analyses. We determined genetic correlations across environments for all traits (see Supplemental Figure 1), but we only used $r_i$ from traits that showed significant G×E for the rank versus variance analysis.

Selection Analysis

To test whether there are any differences in how natural selection acts on mutualistic traits across environments (i.e., S×E), we conducted a selection analysis following the logic described by De Lisle and Svensson (2017) and Batstone et al. (2020). We used aboveground biomass as a proxy for plant fitness (most studies show a positive correlation between biomass and fitness components [Younginger et al., 2017]), and nodule number as our focal mutualistic trait. We also included leaf number and belowground biomass to analyze selection on plant performance traits, and to account for the high correlation of these traits with nodule number (see Results, Figure 2). We obtained raw line means to analyze ANCOVAs using linear models with the fitness proxy as a response variable, and focal traits and light treatment as covariate and interaction terms. We used three scaling methods for our S×E analysis: (1) absolute fitness, (2) relativizing fitness and standardizing traits on a global scale (across treatments), and (3) relativizing and standardizing on a local scale (within each treatment). We compared these methods to assess how differences in mean values and standard deviations across and within treatments might affect our S×E analysis (Batstone et al., 2020). We included an analysis using the absolute scale because it is free from any differences that might result from standardization and relativization in the other scaling methods, and it allows an assessment of the relationship between traits and fitness as they were measured. For the global analysis, we relativized aboveground biomass by dividing line means with a global mean across both treatments (line mean/global mean), and we standardized our focal traits by subtracting line means from the global mean and dividing this by a global SD (line mean – global mean)/global SD). The local analysis was similar except we relativized aboveground biomass using local means from within each treatment (line mean/local mean), and standardized the focal traits using local means and SD (line mean – local mean/local SD). Any difference in selection for our focal traits between light environments (i.e., S×E) was indicated by a significant trait-by-treatment interaction. If a significant S×E interaction from the global analysis is not present in the local analysis, it indicates that the interaction is mainly driven by differences in mean fitness because those differences are mathematically eliminated in the local analysis. We used the global and local S×E analyses to determine whether estimates of selection gradients ($\beta$) were significantly different between environments (i.e., significant S×E interaction). However, we used $\beta$ estimates from the local analysis to determine differences in the magnitude and direction of selection between environments, as these are more interpretable for making predictions of evolutionary change (De Lisle and Svensson, 2017).

SUPPLEMENTAL INFORMATION

Supplemental Information is available at Plant Communications Online.

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AUTHOR CONTRIBUTIONS

Conceptualization, P.V. and J.R.S.; Methodology, P.V. and J.R.S.; Investigation, P.V.; Formal Analysis, P.V.; Writing – Original Draft, P.V.; Writing – Review, Editing, & Revising, P.V. and J.R.S.; Resources & Supervision, J.R.S.

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REFERENCES

Abdala-Roberts, L., and Mooney, K.A. (2013). Environmental and plant genetic effects on tri-trophic interactions. Oikos 122:1157–1166.
Plant Communications

Abdala-Roberts, L., Agrawal, A.A., and Mooney, K.A. (2012). Ant-aphid interactions on Asclepias syriaca are mediated by plant genotype and caterpillar damage. Oikos 121:1005–1013.

Ahlholm, J.U., Helander, M., Henriksson, J., Metzler, M., and Saikkonen, K. (2002). Environmental conditions and host genotype directly genetic diversity of Venturia ditricha, a fungal endophyte of birch trees. Evolution 56:1566–1573.

Arnold, S.J., and Peterson, C.R. (2002). A model for optimal reaction norms: the case of the pregnant garter snake and her temperature-sensitive embryos. Am. Nat. 160:306–316.

Axelrod, R., and Hamilton, W.D. (1981). The evolution of cooperation. Science 211:1390–1396.

Ballhorn, D.J., Schadler, M., Elias, J.D., Millar, J.A., and Kauth, S. (2016). Friend or foe—light availability determines the relationship between mycorrhizal fungi, rhizobia and Lima Bean (Phaseolus lunatus L.). PLoS One 11. https://doi.org/10.1371/journal.pone.0154116.

Barker, D.G., Bianchi, S., Blondon, F., Dattée, Y., Duc, G., Essad, S., Flament, P., Gallusci, P., Génier, G., and Guy, P. (1990). Medicago truncatula, a model plant for studying the molecular genetics of the Rhizobium-legume symbiosis. Plant Mol. Biol. Rep. 8:40–49.

Bataillon, T., and Ronfort, J. (2006). Evolutionary and ecological genetics of Medicago truncatula. Medicago truncatula handbook, 1–12.

Bates, D., Maechler, M., Bolker, B., and Walker, S. (2015). Fitting linear mixed-effects models using lme4. J. Stat. Softw. 1:1–48.

Batstone, R.T., Dutton, E.M., Wang, D., Yang, M., and Frederickson, M.E. (2017). The evolution of symbiotic preference traits in the model legume Medicago truncatula. New Phytol. 213:1850–1861.

Batstone, R.T., Carscadden, K.A., Afkhami, M.E., and Frederickson, M.E. (2018). Using niche breadth theory to explain generalization in mutualisms. Ecology 99:1039–1050.

Batstone, R.T., Peters, M.A., Simonsen, A.K., Stinchcombe, J.R., and Frederickson, M.E. (2020). Environmental variation impacts trait expression and selection in the legume-rhizobium symbiosis. Am. J. Bot. 107:195–208.

Bertrand, A., Prévost, D., Bigras, F.J., Lalande, R., Tremblay, G.F., Castonguay, Y., and Bélanger, G. (2007). Alfalfa response to elevated atmospheric CO2 varies with the symbiotic rhizobial strain. Plant Soil 301:173–187.

Blondon, F., Marie, D., Brown, S., and Kondorosi, A. (1994). Genome size and base composition in Medicago sativa and M. truncatula species. Genome 37:264–270.

Bowman, J.C. (1972). Genotype X Environment Interactions (BioMed Central), p. 117.

Bronstein, J.L. (1994). Conditional outcomes in mutualistic interactions. Trends Ecol. Evol. 9:214–217.

Casal, J.J. (2012). Shade avoidance. Arabidopsis Book 10:e0157.

Charlesworth, B. (1987). The heritability of fitness. In Sexual Selection: Testing the Alternatives, J. Bradbury and M.B. Anderson, eds. (Oxford, United Kingdom: John Wiley & Sons, Ltd), pp. 21–40.

Cockerham, C.C. (1963). Estimation of genetic variances. In Statistical Genetics and Plant Breeding, W.D. Hanson and H.F. Robinson, eds. (Washington, DC, USA: National Research Council), pp. 53–94.

Cook, D.R. (1999). Medicago truncatula—a model in the making!. Curr. Opin. Plant Biol. 2:301–304.

Corré, W.J. (1983). Growth and morphogenesis of sun and shade plants I. The influence of light intensity. Acta Bot. Neerl. 32:49–62.

Cronin, G., and Lodge, D.M. (2003). Effects of light and nutrient availability on the growth, allocation, carbon/nitrogen balance, phenolic chemistry, and resistance to herbivory of two freshwater macrophytes. Oecologia 127:32–41.

De Lisle, S.P., and Svensson, E.I. (2017). On the standardization of fitness and traits in comparative studies of phenotypic selection. Evolution 71:2313–2326.

Des Marais, D.L., Hernandez, K.M., and Juenger, T.E. (2013). Genotype-by-environment interaction and plasticity: exploring genomic responses of plants to the abiotic environment. Annu. Rev. Ecol. E vol. Syst. 44:5–29.

Donohue, K., Messiqua, D., Pyle, E.H., Heschel, M.S., and Schmitt, J. (2000). Evidence of adaptive divergence in plasticity: density- and site-dependent selection on shade-avoidance responses in Impatiens capensis. Evolution 54:1956–1968.

El Mshehli, S., Lambert, A., Hopkins, J., Boncompagni, E., Smiti-Aschi, S., Héroaut, D., and Freno, P. (2019). Physiological and genetic changes during natural senescence of Medicago truncatula root nodules. J. Plant Nutr. Soil Sci. 182:385–392.

Evans, J.R. (1989). Partitioning of nitrogen between and within leaves grown under different irradiances. Funct. Plant Biol. 16:533–548.

Fisher, R.A. (1958). The Genetical Theory of Natural Selection, 2nd edn (New York: Dover Publications).

Fox, J., and Weisberg, S. (2019). An [R] Companion to Applied Regression, 3rd edn (Thousand Oaks CA: Sage).

Friesen, M.L., and Friel, C.A. (2019). Legumes modulate allocation to rhizobial nitrogen fixation in response to factorial light and nitrogen manipulation. Front. Plant Sci. 10:1316.

Garcia, J., Barker, D.G., and Journet, E.P. (2006). Seed Storage and Germination. The Medicago truncatula Handbook (Ardmore, OK: he Samuel Roberts Noble Foundation). http://www.noble.org/MedicagoHandbook.

Gillespie, J.H., and Turelli, M. (1989). Genotype-environment interactions and the maintenance of polygenic variation. Genetics 121:129–138.

Givnish, T.J. (1988). Adaptation to sun and shade: a whole-plant perspective. Funct. Plant Biol. 15:63–92.

Gomulkiewicz, R., and Kirkpatrick, M. (1992). Quantitative genetics and the evolution of reaction norms. Evolution 46:390–411.

Gorton, A.J., Heath, K.D., Pilet-Nayel, M.-L., Baronager, A., and Stinchcombe, J.R. (2012). Mapping the genetic basis of symbiotic variation in legume-rhizobium interactions in Medicago truncatula. G3: Genes Genomes Genet. 2:1291–1303.

Grillo, M.A., Stinchcombe, J.R., and Heath, K.D. (2016). Nitrogen addition does not influence pre-infection partner choice in the legume-rhizobium symbiosis. Am. J. Bot. 103:1763–1770.

Harrell, F. (2020). Hmisc: Harrell Miscellaneous. R Package Version 4.4-0 (The Comprehensive R Archive Network (CRAN)). https://biostat.org/R/Hmisc/.

Hartig, F. (2020). DHARMa: Residual Diagnostics for Hierarchical (Multi-Level/Mixed) Regression Models (The Comprehensive R Archive Network (CRAN)). http://florianhartig.github.io/DHARMa/.

Heath, K.D. (2010). Intergeneric epistasis and coevolutionary constraint in plants and rhizobia. Evol. Int. J. Org. Evol. 64:1446–1458.

Heath, K.D., Stock, A.J., and Stinchcombe, J.R. (2010). Mutualism variation in the nodulation response to nitrates. Journal of Evolutionary Biology 23:2494–2500.

Heath, K.D., and Tiffin, P. (2014). Explaining mutualism variation: a new evolutionary paradox? Evolution 68:309–317.

Heath, K.D., and Tiffin, P. (2007). Context dependence in the coevolution of plant and rhizobial mutualists. Proc. Biol. Sci. 274:1905–1912.

Heath, K.D., and Tiffin, P. (2009). Stabilizing mechanisms in a legume-rhizobium mutualism. Evol. Int. J. Org. Evol. 63:652–662.
Genetic Variation in Legume–Rhizobia Mutualism

Heath, K.D., Podowski, J.C., Heniff, S., Klingler, C.R., Burke, P.V., Weese, D.J., Yang, W.H., and Lau, J.A. (2020). Light availability and rhizobium variation interactively mediate the outcomes of legume-rhizobium symbiosis. Am. J. Bot. 107:229–238.

Heschel, M.S., Stinchcombe, J.R., Holsinger, K.E., and Schmitt, J. (2004). Natural selection on light response parameters in the herbaceous annual, Impatiens capensis. Oecologia 139:487–494.

Huhm, N., Lotito, S., and Piepho, H.P. (1993). Relationships between genotype x environment interactions and rank orders for a set of genotypes tested in different environments. Theor. Appl. Genet. 86:943–950.

Hungate, B.A., Stiling, P.D., Dijkstra, P., Johnson, D.W., Ketterer, H., Magnoli, S.M., and Lau, J.A. (2020). Novel plant–microbe interactions: evidence that symbiotic N2 fixation in fertile grassland is an important nutritional input. Am. J. Bot. 107:1299–1312.

Johnson, J.L., Stinchcombe, J.R., Weinig, C., and Schmitt, J. (2007). Genotype-by-environment interactions leads to variable selection on life-history strategy in Common Evening Primrose (Oenothera biennis). J. Evol. Biol. 20:190–200.

Jones, K.M., Kobayashi, H., Davies, B.W., Taga, M.E., and Walker, G.C. (2007). How rhizobial symbionts invade plants: the Sinorhizobium-Medicago model. Nat. Rev. Microbiol. 5:619–633.

Kelley, J.L., Stinchcombe, J.R., Weinig, C., and Schmitt, J. (2005). Soft and hard selection on plant defence traits in Arabidopsis thaliana. Evol. Ecol. Res. 7:287–302.

Kiers, E.T., Rousseau, R.A., West, S.A., and Denison, R.F. (2003). Host sanctions and the legume-rhizobium mutualism. Nature 425:78–81.

Kruuk, L.E.B., Clutton-Brock, T.H., Slate, J., Pemberton, J.M., Brotherstone, S., and Guinness, F.E. (2000). Heritability of fitness in a wild mammal population. Proc. Natl. Acad. Sci. U S A 97:698–703.

Lau, J.A., Bowling, E.J., Gentry, L.E., Glasser, P.A., Monarch, E.A., Olesen, W.M., Wam克斯ky, J., and Young, R.T. (2012). Direct and interactive effects of light and nutrients on the legume-rhizobia mutualism. Acta Oecol. 39:80–86.

Luscher, A., Hartwig, U.A., Suter, D., and Nosberger, J. (2000). Direct evidence that symbiotic N2 fixation in fertile grassland is an important trait for a strong response of plants to elevated atmospheric CO2. Glob. Change Biol. 6:655–662.

Magnoli, S.M., and Lau, J.A. (2020). Novel plant–microbe interactions: Rapid evolution of a legume–rhizobium mutualism in restored prairies. Journal of Ecology.

McDonald, J.F., and Ayala, F.J. (1974). Genetic response to environmental heterogeneity. Nature 250:572–574.

Mitchell-Olds, T. (1992). Does environmental variation maintain genetic variation? A question of scale. Trends Ecol. Evol. 7:397–400.

Mitchell-Olds, T., Willis, J.H., and Goldstein, D.B. (2007). Which evolutionary processes influence natural genetic variation for phenotypic traits? Nature Reviews Genetics 8 (11):845–856.

Mouton, L., Henri, H., Charif, D., Bouletreau, M., and Vavre, F. (2007). Interaction between host genotype and environmental conditions affects bacterial density in Wolbachia symbiosis. Biol. Lett. 3:210–213.

Muir, W., Nyquist, W.E., and Xu, S. (1992). Alternative partitioning of the genotype-by-environment interaction. Theor. Appl. Genet. 84:193–200.

Nathanial Holland, J., and DeAngelis, D.L. (2009). Consumer-resource theory predicts dynamic transitions between outcomes of interspecific interactions. Ecol. Lett. 12:1357–1366.

Osslter, J.N., and Heath, K.D. (2018). Shared genes but not shared genetic variation: legume colonization by two belowground symbionts. Am. Nat. 191:395–406.

Pupo, A., Groten, K., Bastian, F., Carzaniga, R., Soussis, M., Lucas, M.M., De Felipe, M.R., Harrison, J., Vanacker, H., and Foyer, C.H. (2005). Legume nodule senescence: roles for redox and hormone signalling in the orchestration of the natural aging process. New Phytol. 165:683–701.

Plant Communications

Regus, J.U., Gano, K.A., Hollowell, A.C., and Sachs, J.L. (2014). Efficiency of partner choice and sanctions in Lotus is not altered by nitrogen fertilization. J. Evol. Biol. 28:1:20132587.

Regus, J.U., Gano, K.A., Hollowell, A.C., Sofish, V., and Sachs, J.L. (2015). Lotus hosts delimit the mutualism-parasitism continuum of Bradyrhizobium. P. Roy. Soc. B Biol. Sci. 28:447–456.

Regus, J.U., Quides, K.W., O’Neill, M.R., Suzuki, R., Savory, E.A., Chang, J.H., and Sachs, J.L. (2017). Cell autonomous sanctions in legumes target ineffective rhizobia in nodules with mixed infections. Am. J. Bot. 104:1299–1312.

Rogers, A., Gibon, Y., Stitt, M., Morgan, P.B., Bernacchi, C.J., Ort, D.R., and Long, S.P. (2006). Increased C availability at elevated carbon dioxide concentration improves N assimilation in a legume. Plant Cell Environ. 29:1651–1658.

Sachs, J.L., and Simms, E.L. (2006). Pathways to mutualism breakdown. Trends Ecol. Evol. 21:585–592.

Schmitt, J., Stinchcombe, J.R., Heschel, M.S., and Huber, H. (2003). The adaptive evolution of plasticity: phytochrome-mediated shade avoidance responses. Integr. Comp. Biol. 43:459–469.

Schwartz, M.W., and Hoeksema, J.D. (1998). Specialization and resource trade: biological markets as a model of mutualisms. Ecology 79:1029–1038.

Shantz, A.A., Lemoine, N.P., and Burkepile, D.E. (2016). Nutrient loading alters the performance of key nutrient exchange mutualisms. Ecol. Lett. 19:20–28.

Simonsen, A.K., and Stinchcombe, J.R. (2014a). Herbivory eliminates fitness costs of mutualism exploiters. New Phytol. 202:651–661.

Simonsen, A.K., and Stinchcombe, J.R. (2014b). Standing genetic variation in host preference for mutualist microbial symbionts. P. Roy. Soc. B Biol. Sci. 281:20142036.

Stanton-Geddes, J., Pape, T., Epstein, B., Briskine, R., Yoder, J., Mudge, J., Bharot, A.K., Farmer, A.D., Zhou, P., and Denny, R. (2013). Candidate genes and genetic architecture of symbiotic and agronomic traits revealed by whole-genome, sequence-based association genetics in Medicago truncatula. PLoS One 8. https://doi.org/10.1371/journal.pone.0065688.

Stinchcombe, J.R., Izem, R., Heschel, M.S., McGoey, B.V., and Schmitt, J. (2010). Across-environment genetic correlations and the dynamics of selective environments shape the evolutionary dynamics of growth rate in Impatiens capensis. Evol. Int. J. Org. Evol. 64:2867–2903.

Stuefer, J.F., and Huber, H. (1998). Differential effects of light quantity and spectral light quality on growth, morphology and development of two stoloniferous Potentilla species. Oecologia 117:1–8.

Sultan, S.E., and Bazzaz, F.A. (1993). Phenotypic plasticity in Polygonum persicaria. I. Diversity and uniformity in genotypic norms of reaction to light. Evolution 47:1009–1031.

Thrall, P.H., Hochberg, M.E., Burdon, J.J., and Bever, J.D. (2007). Coevolution of symbiotic mutualists and parasites in a community context. Trends Ecol. Evol. 22:120–126.

Trivers, R.L. (1971). The evolution of reciprocal altruism. Q. Rev. Biol. 46:35–57.

Van Cauwenberghje, J., Michiels, J., and Honnay, O. (2015). Effects of local environmental variables and geographical location on the genetic diversity and composition of Rhizobium leguminosarum nodulating Vicia cracca populations. Soil Biol. Biochem. 90:71–79.
Plant Communications

Via, S., and Lande, R. (1985). Genotype-environment interaction and the evolution of phenotypic plasticity. Evolution 39:505–522.

Walsh, B., and Blows, M.W. (2009). Abundant genetic variation + strong selection = multivariate genetic constraints: a geometric view of adaptation. Annu. Rev. Ecol. Evol. Syst. 40:41–59.

Weese, D.J., Heath, K.D., Dentinger, B.T., and Lau, J.A. (2015). Long-term nitrogen addition causes the evolution of less-cooperative mutualists. Evolution 69:631–642.

Wei, T., and Simko, V. (2017). R Package “Corrplot”: Visualization of a Correlation Matrix, Version 0.84 (The Comprehensive R Archive Network (CRAN)). https://github.com/taiyun/corrplot.

Werner, G.D., Zhou, Y., Pieterse, C.M., and Kiers, E.T. (2018). Tracking plant preference for higher-quality mycorrhizal symbionts under varying CO₂ conditions over multiple generations. Ecol. Evol. 8:78–87.

Genetic Variation in Legume–Rhizobia Mutualism

West, J.B., HilleRisLambers, J., Lee, T.D., Hobbie, S.E., and Reich, P.B. (2005). Legume species identity and soil nitrogen supply determine symbiotic nitrogen-fixation responses to elevated atmospheric [CO₂]. New Phytol. 167:523–530.

Winn, A.A., and Evans, A.S. (1991). Variation among populations of Prunella vulgaris L. in plastic responses to light. Funct. Ecol. 5:562–571.

Wood, C.W., Pilkington, B.L., Vaidya, P., Biel, C., and Stinchcombe, J.R. (2018). Genetic conflict with a parasitic nematode disrupts the legume-rhizobia mutualism. Evol. Lett. 2:233–245.

Younginger, B.S., Sirová, D., Cruzan, M.B., and Ballhorn, D.J. (2017). Is biomass a reliable estimate of plant fitness? Appl. Plant Sci. 5:1600094.

Zheng, C., Ji, B., Zhang, J., Zhang, F., and Bever, J.D. (2015). Shading decreases plant carbon preferential allocation towards the most beneficial mycorrhizal mutualist. New Phytol. 205:361–368.