**Arabidopsis thaliana** and the Robin Hood parasite: a chivalrous oomycete that steals fitness from fecund hosts and benefits the poorest one?

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Are parasites always harmful to their hosts? By definition, indeed, but in a few cases and particular environments, hosts experience higher fitness in the presence than in the absence of their parasites. Symbiotic associations form a continuum of interactions, from deleterious to beneficial effects on hosts. In this paper, we investigate the outcome of parasite infection of *Arabidopsis thaliana* by its natural pathogen *Hyaloperonospora arabidopsis*. This system exhibits a wide range of parasite impact on host fitness with, surprisingly, deleterious effects on high fecundity hosts and, at the opposite extreme, seemingly beneficial effects on the least fecund one. This phenomenon might result from varying levels of tolerance among host lines and even overcompensation for parasite damage analogous to what can be observed in plant–herbivore systems.

Keywords: host–parasite interaction; symbiosis; tolerance; evolved dependence

1. **INTRODUCTION**

Parasitism and mutualism are the two extreme types of interactions between associated species. Both imply the exploitation of resources of one organism by its interacting partner but they fundamentally differ in that mutualistic associations benefit both of them, whereas one partner gains at the expense of the other in parasitic ones. This distinction is not always clear, however, and numerous cases where mutualistic organisms occasionally cheat on their partner have been reported. Examples include well-known mutualisms such as mycorrhizae, yucca moths and cleaner fishes (Bronstein 2001). At the other extreme, rare reports exist of notorious parasites having a positive effect on their host in particular ecological contexts, when the negative impact of parasite exploitation is compensated by an indirect advantage of parasite presence, like protection against heavy metal toxicity (Thomas et al. 2000). Symbiotic associations thus form a continuum of interactions, ranging from deleterious to beneficial (Bronstein 1994) and the outcome of a given association may change depending on the individual partners or external factors. Elucidating the evolution of interspecies associations along this continuum will require identifying these factors and determining whether they are available to selection. Many studies investigating the fitness consequences of parasitic infections have found variation in virulence, i.e. reduction in host fitness, among parasite genotypes (e.g. Peever et al. 2000; Schulenburg & Ewbank 2004) and/or variation in tolerance among hosts (e.g. Peever et al. 2000; Kover & Schaal 2002). Such studies, however, usually focused on parasites with a large, and thus easily assessed, deleterious impact. On the other hand, mild to asymptomatic parasites represent an opportunity to investigate variation in infection effects from negative to positive. The model plant *Arabidopsis thaliana*, provides such a case, as recent experiments with different natural pathogens have revealed occasional cases where infection increase host fitness, albeit not significantly (Salvaudon et al. 2005; Goss & Bergelson 2007).

Here, we report the results of an experiment designed specifically to investigate whether host or parasite genotypes, or both, determine the fitness variation and advantage in *A. thaliana* when infected with its natural oomycete pathogen *Hyaloperonospora arabidopsis*, using several plant lines and parasite genotypes of different origins. The deleterious impact of parasite infection varied dramatically among plant genotypes, despite their similar susceptibility profile, from clear negative impacts to benevolence, and we confirm that this parasite can even increase reproductive success of its host.

2. **MATERIAL AND METHODS**

(a) Material

The oomycete *H. arabidopsis* is a natural pathogen of *A. thaliana* (Brassicaceae), its specific host. This parasite is biotrophic and produces asexual conidiophores on leaf surfaces a few days after infection, as well as sexual oospores that remain within leaves until host death. This pathogen is seldom lethal for its host, but nonetheless constitutes a selective pressure on *A. thaliana*, as more than 20 resistance genes targeted against *H. arabidopsis* have been reported (Slusarenko & Schlaich 2003). Six inbred lines of *A. thaliana* and seven parasite strains constituting a complete matrix of successful infections were selected from our collection. We used parasite strains of three different types of origin. Three strains, Emco, Emwa and Noco, were ‘laboratory strains’ provided by the Sainsbury Laboratory (John Innes Center, Norwich, UK) from isolates collected more than 10 years ago and subsequently maintained as asexual cultures on specific *A. thaliana* lines (Holub et al. 1994). ‘Orsay strains’, Ors3 and Ors5, were collected in spring 2004 from conidia spores on infected host plants from one population in Orsay, France. The ‘Fribourg strains’, Fr3 and Fr5, were obtained from oospores of two infected plants sampled in spring 2004 in a population in Fribourg, Switzerland. All strains were maintained on susceptible hosts for several asexual generations and thus are probably genetically homogeneous. The six *A. thaliana* inbred lines were issued from seeds collected in wild populations across Europe (Finland (Fin), England (Gh), Pyrenees (Pyr), Sweden (Sue) and Czech Republic (Tch)) or from the ecotype Tsu (Japan). Only the combination Fin/Emco unexpectedly failed to produce any successful infection.

(b) Methods

We inoculated all six host lines with the seven parasite strains and a water control, with five replicates for each of the 48 combinations. The plants were grown, inoculated and maintained under controlled conditions following the protocol described in Salvaudon et al. (2007), which reports on parasite transmission from this experiment. The life cycle of all plants was completed under greenhouse conditions (natural photoperiod: 23°C day–15°C night). Plants were watered ad libitum until they began...
fructification, and then only when pots were completely dry to hasten fruit production. Seeds were harvested progressively as they matured in order to collect all seeds before fruits opened. We estimated host fitness as the total weight of all seeds produced per plant, this parameter representing the lifetime investment in reproduction. Plants that died before flowering had a seed production of zero.

(c) Statistical analyses
All statistical analyses were performed with JMP software version 5.1.2 (SAS institute, Cary, NC). As the selected host and parasite genotypes did not represent a random sample of their respective populations but rather a subsample of intercompatible genotypes, we chose to treat them as fixed factors. Our purpose was not then to assess the existing variation for virulence in natural populations but to test whether host (or parasite) genotypes that have a similar susceptibility (or infectivity) profile can differ for virulence. The impact of inoculation treatment on host fitness was thus analysed with a nested ANOVA testing for the effect of host line, treatment type (water-inoculated controls or inoculation with laboratory, Fribourg or Orsay strains), parasite strain nested within the treatment type and interactions between host line and each of the two other factors.

We also tested whether virulence was linked to the intrinsic fecundity of host lines. By analogy with the type of analysis used to investigate costs of tolerance in plant–herbivore interactions, we investigated the relationship between the average fitness of attacked plants and the fitness in control plants of the same family (Strauss & Agrawal 1999). But instead of looking for a negative relationship that would indicate a cost, we tested whether the slope of this relationship was significantly different from one (Student’s t-test), as this value would be expected if virulence was completely unrelated to the intrinsic fecundity of host lines. This relationship was tested at all the levels of parasite treatment (mean of all pooled strains, parasite strains averaged over origins or each parasite strain separately), followed by Bonferroni correction of the critical α-value to account for multiple testing. Three plants were excluded from the analyses as there were doubts about their possible contamination or errors during inoculation.

3. RESULTS
Of the 240 plants, six died before flowering. The weight of total seed production varied significantly among the six host lines ($F_{5,189} = 31.007, p < 0.0001$) but not among the treatment types (controls, laboratory, Fribourg or Orsay strains) or parasite strains within the treatment types. However, there was a significant, albeit small, interaction between treatment type and host line ($F_{15,189} = 1.733; p = 0.0476$), which was due to a difference in seed production between control and inoculated plants in some host lines: seed yield was lower in infected plants of Gb and Tch lines, unchanged in Fin, Sue and Tsu lines and higher in the Pyr line. This small interaction term was nonetheless robust to the type of analysis performed, and remained significant with a mixed model (Hocking formulation; Strauss & Agrawal 1999) incorporating only host lines as random factor, the type of treatment and their interaction ($F_{15,213} = 1.758, p = 0.0424$).

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This interaction was due to significant differences between control and laboratory strains that were both positive (contrast on Pyr, $T$-ratio = 2.56, $p = 0.011$) and negative (Gb, $T$-ratio = −2.097, $p = 0.037$; Tch, $T$-ratio = −2.51, $p = 0.025$). Orsay and Fribourg strains showed little significance effect (figure 1). Furthermore, these changes in seed production covaried with the
fitness (Strauss & Agrawal 1999). Indeed, genetic that reduces the negative effect of infection on host growth. But they could reflect variation in tolerance levels of resistance that reduces parasite infectivity or et al. (Salvaudon et al. 2005) has previously been demonstrated in Arabidopsis. 

Table 1. Slope of the relationship between average seed production of inoculated and control plants of the same host line. The Student *t*-test (5 d.f.) tests whether the slope equals one. Since we test the same hypothesis multiple times we mark in italics only those tests significant after Bonferroni correction of the critical *a*-value.

<table>
<thead>
<tr>
<th>mean by origin</th>
<th>slope coefficient</th>
<th>Student <em>t</em></th>
<th><em>p</em>-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory</td>
<td>0.364</td>
<td>7.074</td>
<td>0.0009</td>
</tr>
<tr>
<td>Fribourg</td>
<td>0.571</td>
<td>4.074</td>
<td>0.0096</td>
</tr>
<tr>
<td>Orsay</td>
<td>0.671</td>
<td>2.022</td>
<td>0.0991</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>mean by strain</th>
<th>slope coefficient</th>
<th>Student <em>t</em></th>
<th><em>p</em>-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emco</td>
<td>0.226</td>
<td>5.308</td>
<td>0.0032</td>
</tr>
<tr>
<td>Emwa</td>
<td>0.197</td>
<td>5.395</td>
<td>0.0030</td>
</tr>
<tr>
<td>Noco</td>
<td>0.669</td>
<td>4.057</td>
<td>0.0098</td>
</tr>
<tr>
<td>Fri3</td>
<td>0.488</td>
<td>5.312</td>
<td>0.0032</td>
</tr>
<tr>
<td>Fri5</td>
<td>0.655</td>
<td>2.571</td>
<td>0.0500</td>
</tr>
<tr>
<td>Ors3</td>
<td>0.587</td>
<td>2.832</td>
<td>0.0366</td>
</tr>
<tr>
<td>Ors5</td>
<td>0.755</td>
<td>1.175</td>
<td>0.2828</td>
</tr>
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The more striking aspect of our results, which cannot be explained by tolerance alone (either to stress or parasites), is the fact that the less fecund host line benefited from the infection. Here again it is double that H. arabidopsis behaved generously by providing some advantage. However, such an increase in host fitness has sometimes been reported for plants damaged by herbivores and a similar phenomenon could possibly have occurred with our parasite. This increase in plant reproduction following damage, or ‘overcompensation’, has been explained as an adaptation of plants to recurrent and predictable herbivory, with plants delaying inflorescence development until grazers have passed. The corollary is that such plants then become dependent on herbivores to trigger the initiation of reproduction and perform less well in their absence. Overcompensation is thus an artefact of the plants’ evolved dependence on their herbivores (Agrawal 2000; de Mazancourt et al. 2005): their being unprepared to live without them. This phenomenon is less documented for parasites (but see Pannebakker et al. 2007), but there is no reason why a predictable attack by parasites would not select for the same kind of dependence. We do not exclude that other factors could also be invoked (like other aspects of the experimental conditions for which this line was maladapted, or even a real, yet undetermined, advantage given by the parasite). However, our hypothesis provides testable predictions. We expect that the gradient observed among the host lines tested reflects a gradient in parasite pressure in their population of origin, with low tolerance in populations with low parasite pressure and high tolerance even associated with evolved dependence in those undergoing regular pathogen epidemics. In any case, the dramatic variation in parasite impact we observed among host lines in our experiment points towards an important role in host–parasite coevolution played by host variation in compensating for infection.

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