Host–Parasite Interactions and the Evolution of Gene Expression

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Interactions between hosts and parasites provide an ongoing source of selection that promotes the evolution of a variety of features in the interacting species. Here, we use a genetically explicit mathematical model to explore how patterns of gene expression evolve at genetic loci responsible for host resistance and parasite infection. Our results reveal the striking yet intuitive conclusion that gene expression should evolve along very different trajectories in the two interacting species. Specifically, host resistance loci should frequently evolve to co-express alleles, whereas parasite infection loci should evolve to express only a single allele. This result arises because hosts that co-express resistance alleles are able to recognize and clear a greater diversity of parasite genotypes. By the same token, parasites that co-express antigen or elicitor alleles are more likely to be recognized and cleared by the host, and this favours the expression of only a single allele. Our model predicts testable predictions that can help interpret accumulating data on expression levels for genes relevant to host–parasite interactions.

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Introduction

Hosts and parasites are locked in a continual co-evolutionary race, which generates persistent selection for resistant hosts and infectious parasites. Understanding the direct effects of this process on spatial patterns of local adaptation [1–5], the evolution of virulence/pathogenicity [6–8], and the spread of infectious disease [9–11] has been a central focus of research into host–parasite interactions. Yet host–parasite interactions also generate indirect selection on a variety of other features of the interacting species. The classical example of indirect selection imposed by host–parasite interactions is on the mode of reproduction [12–16]. Host–parasite interactions can select for sexual rather than asexual reproduction, although they tend to do so only when selection is strong and sex is rare [17]. Recently, we have shown that indirect selection also acts on genome size (ploidy level), with selection favouring diploidy more often among host species and haploidy more often among parasite species [18]. There are a variety of other genomic features besides ploidy level that should experience indirect selection in response to host–parasite interactions. Here, we examine the evolution of expression levels using a model that is structurally similar to models of the evolution of dominance (as in the classic papers by Fisher [19,20], Wright [21–23], and Haldane [24,25], and more recent papers reviewed in Otto and Bourgouet [26]).

The Model

To explore the evolution of expression levels, we assumed that infection/resistance was determined by a single gene in the host with alleles A and a, and a single gene in the parasite with alleles B and b. We then tracked changes in allele frequency at a single modifier locus, whose alleles (M and m) altered the pattern of expression in heterozygotes at the A locus (if in hosts) or B locus (if in parasites). Thus, we refer to this modifier locus, M, as a regulatory locus. To simplify the analysis and interpretation, we allowed expression levels to evolve in only one species (the “focal species”) at a time.

Determining how expression patterns evolve during the course of host–parasite co-evolution requires that we relate expression patterns to the phenotype expressed by heterozygous genotypes. We assumed that a heterozygous individual of species j could express the phenotype of homogygotes carrying allele A (or B) with probability \( p_{1j} \), and A (or B) and a (or b) with probability \( p_{2j} \), and A (or B) with probability \( p_{3j} \), where the terms in parentheses are appropriate when the focal species is the parasite. These probabilities were assumed to sum to one (\( p_{1j} + p_{2j} + p_{3j} = 1 \)), for both hosts \( (j = h) \) and parasites \( (j = p) \). This constraint prevents heterozygotes from having fitness greater than the best homozygous genotype in any given encounter between host and parasite genotypes. An implicit assumption of this mapping between genotype and phenotype is that heterozygotes can, if \( p_{2j} = 1 \), co-express both alleles without decreasing the function of either allele. To take a concrete example, our mapping of phenotype onto genotype assumes that AA hosts could express receptor A as effectively as AA hosts and also express receptor a as effectively as aa hosts. The model is easily generalized, however, to relax this assumption (results available upon request). Alleles at the regulatory locus, M, were allowed to alter the pattern of expression in heterozygotes by altering the probabilities, \( p_{i(j)} \). Because an individual's genotype at the regulatory locus determines these probabilities, we specify

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Abbreviations: GFG, gene-for-gene; IMA, inverse-matching-alleles; MA, matching-alleles

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the genotype in square brackets (e.g., \(p_{ij}[MM]\)). When exposed to selection induced by the interacting species, alleles at the regulatory locus might evolve to upregulate one allele over the other or to express both alleles equally (co-expression), as illustrated in Figure 1.

We incorporated host–parasite co-evolution into the modifier framework described above by considering the following well-studied genetic interactions. In the gene-for-gene (GFG) model [27], avirulence alleles in the parasite produce signal molecules that elicit a defence response in resistant hosts, whereas parasites carrying virulence alleles fail to produce the signal molecule and cannot be detected by any host. GFG interactions are considered to be prevalent in plant–pathogen interactions [28]. Costs of resistance and virulence alleles have been demonstrated in some GFG systems [29,30], so we let \(C_g\) be the fitness cost of expressing only the resistant allele in hosts, and \(C_p\) be the fitness cost of expressing only the virulence allele in parasites. Co-expressing both alleles might reduce these costs, particularly when the susceptible allele in the host or the avirulent allele in the parasite performs a beneficial function. The fitness costs experienced by heterozygotes expressing both alleles were thus set to \(C_s\) in hosts and \(C_p\) in parasites. The matching-alleles (MA) model is predicated upon a system of self/non-self recognition. Hosts can successfully defend against attack by a parasite whose genotype does not match their own. Such recognition systems have been observed in invertebrates [31] and vertebrates [32]. Finally, in the inverse-matching-alleles (IMA) model, host defence involves an array of recognition molecules (e.g., antibodies) that are able to recognize specific antigens and resist attack by parasites carrying these antigens [32]. Following the rules imposed by each of these modes of co-evolution allowed us to create a matrix that describes the outcome of an interaction between any two phenotypes (Table 1). In all cases, we assumed that infection results in a loss of host fitness but an increase in parasite fitness.

We assumed a life cycle where selection due to interactions between host and parasite was followed by sexual reproduction. Species interactions are assumed to depend on loci: a regulatory locus with alleles \(M\) and \(m\) and an interaction locus with alleles \(A\) and \(a\) if the focal species is the host, or \(B\) and \(b\) if the focal species is the parasite. Thus, there are four chromosome types in each species: \(MA\) (MB), \(Ma\) (Mb), \(mA\) (mB), and \(ma\) (mb), where the terms in parentheses correspond to cases where the focal species is the parasite. We track evolution at the regulatory locus in only one species at a time and assume that the regulatory locus is fixed on \(M\) in the non-focal species. The non-focal species is assumed to be diploid, although results derived with a haploid non-focal species were similar. Species \(j\) is assumed to undergo sexual reproduction with random mating with probability \(s_{xy}\) and to reproduce asexually with

### Table 1. An Interaction between a Host and a Parasite Results in Either Infection or Resistance, Depending on the Phenotype of the Interacting Species

| Parasite Phenotype | Host Phenotype | A | A and a | a |
|--------------------|---------------|---|---------|---|
| \(B\)              | \{UR\}        | \{LR\} | \{UJ\} |
| \(B\) and \(b\)    | \{RR\}        | \{LR\} | \{UR\} |
| \(b\)              | \{RR\}        | \{LR\} | \{UR\} |

Each vector represents the outcome of a species interaction, either infection (I) or resistance (R), under the following three models: GFG (first number in each set), MA (second number in each set), and IMA (third number in each set). In GFG interactions, infection reduces host fitness by \(\gamma_h\), resistance reduces parasite fitness by \(\gamma_p\). In MA interactions, infection reduces host fitness by \(\gamma_h\), resistance reduces parasite fitness by \(\gamma_p\).

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probability \((1 - sex)\). During sexual reproduction, the two loci are allowed to recombine at rate \(r_{ij}\).

Genotype frequencies after one round of selection can be determined using standard population genetic equations once the fitnesses of genotypes have been determined. We assume that encounters between species occur at random and that at most one interaction occurs per generation per individual. When interacting with genotype \(k\) in species \(j\), the fitness of genotype \(i\) in species \(j\) is denoted by \(W_{ij} = \sum_{genotypes\ k} W_{ij-kj}\). Thus, we assume that fitness depends upon genotype frequencies but is independent of the population sizes of the interacting species (e.g., [33,34]). The mean fitness in species \(j\) is calculated as the weighted sum of equation 1 over all genotypes in species \(j\):

\[
W_j = \sum_{genotypes\ i} X_{ij} W_{ij}.
\]

\(W_{ij-kj}\) can be calculated using Table 1 and the probabilities, \(p_{ij}\), that heterozygous hosts (parasites) express a particular phenotype. The encounter rate between hosts and parasites is implicitly incorporated in \(W_{ij-kj}\); when hosts and parasites rarely encounter one another, the fitnesses will be more similar to one another, all else being equal.

Assuming an infinite population size and ignoring mutation, we can write down recursions for the frequency, \(X_{ij}\) of each diplodid genotype (e.g., \(i = MA/MA\) or \(MB/MB\)) in species \(j\) after one round of selection followed by reproduction. For example, the first four recursions for host genotypes are given by

\[
X'_{MA/MA,k} = \frac{1 - sex}{2} X'_{MA/MA,k} + \frac{sex}{2} (p'_{M,k})^2
\]

\[
X'_{MA/MA,h} = \frac{1 - sex}{2} X'_{MA/MA,h} + \frac{sex}{2} 2p'_{M,h}p'_{M,h}
\]

\[
X'_{MA/MA,K} = \frac{1 - sex}{2} X'_{MA/MA,K} + \frac{sex}{2} 2p'_{M,h}p'_{M,h}
\]

\[
X'_{MA/MA,H} = \frac{1 - sex}{2} X'_{MA/MA,H} + \frac{sex}{2} 2p'_{M,h}p'_{M,h},
\]

where primes indicate post-selection genotype \((X'_{ij})\) and gamete \((p'_{ij})\) frequencies. Specifically, the frequency of genotype \(i\) after selection is given by \(X'_{ij} = X_{ij} W_{ij}/W_p\), and the gametes produced by the surviving hosts are in the following frequencies:

\[
p'_{M,h} = X'_{MA/MA,h} + \frac{X'_{MA/MA,h}}{2} + \frac{X'_{MA/MA,h}}{2} (1 - r_{ij})
\]

\[
p'_{M,h} = X'_{MA/MA,h} + \frac{X'_{MA/MA,h}}{2} + \frac{X'_{MA/MA,h}}{2} (1 - r_{ij})
\]

Recursions for the parasite species are identical, with the exceptions of the subscripts \(A, a, p, h\) which are replaced by \(B, b, p, h\), respectively.

**Results**

To analyze the model, we assumed that selection was weak relative to the rate of recombination between the modifier locus and the locus determining infection/resistance. This allowed us to derive very general conditions for the evolution of expression levels in the focal species using quasi-linkage equilibrium approximations [35,36]. In short, the frequency of sex and recombination are assumed to be high enough relative to the strength of selection that the disequilibrium between the regulatory and interaction locus \((D_h = \text{freq}(MA), \text{freq}(MA_h) - \text{freq}(MA) \text{freq}(MA_h)\) in hosts) reaches a steady-state value that depends on the current allele frequencies in the host and parasite. Solving for this disequilibrium then allows us to calculate the rate of allele frequency change at the regulatory locus to leading order in the selection coefficients (Protocol S1).

When the host was the focal species, the frequency of allele \(M\) at the regulatory locus changed at a per-generation rate of:

\[
\Delta p_{M,h} = 2p_{A,h}p_{B,h}p_{M,h}p_{M,h} s_{M,h},
\]

where \(p_{ij}\) is the frequency of allele \(i\) in species \(j\) and where \(s_{M,h}\) depends on the model of host–parasite interactions and, for the GFG, MA, and IMA models, is given by

\[
s_{GFG,h} = -\gamma_h (\Delta p_{A,h} (p_{A,h}^2 + 2p_{A,h}p_B(p_{A,h} + p_{B,h})))
\]

\[
- \Delta p_{A,h} (C_h - \alpha) + \Delta p_{A,h} \alpha
\]

\[
s_{MA,h} = \frac{\xi_h \Delta p_{A,h} (p_{A,h}^2 + 2p_{A,h}p_B(p_{A,h} + p_{B,h}))}{\Delta p_{A,h} (p_{A,h}^2 + 2p_{A,h}p_B(p_{A,h} + p_{B,h})))}
\]

\[
+ \Delta p_{A,h} (p_{A,h}^2 + 2p_{A,h}p_B(p_{A,h} + p_{B,h})))
\]

In equations 12–14, \(\gamma_h, \xi_h, \alpha, \alpha_h\) measure the strength of selection acting on species \(i\) due to GFG interactions, MA interactions, and IMA interactions, respectively (see Table 1), and \(\Delta p_{ij}\) represents the average effect of allele \(M\) on the probability of expression pattern \(i\) in species \(j\):

\[
\Delta p_{ij} = \langle p_{ij}[MM] - p_{ij}[Mm] \rangle p_{M,j} + \langle p_{ij}[Mm] - p_{ij}[mm] \rangle p_{m,j}.
\]

Assuming weak selection, equation 11 is equivalent to the allele frequency change in a standard one-locus model with a selection coefficient given by the selection term \(s_{M,h}\) multiplied by the frequency of \(AA\) hosts (2\(p_{A,h}p_{A,h}\). As
discussed in greater detail in Protocol S1, the selection coefficient \( s_{\text{model}M} \) can be readily interpreted on the basis of the change in fitness expected if a heterozygous host will be infected.

Similarly, when the parasite was the focal species, the frequency of allele \( M \) at the regulatory locus changed at a per-generation rate of

\[
\Delta p_{M,p} = 2p_Bp_Hp_{A,H}p_{M,H}p_{M,P}^{\text{model}},
\]

where now

\[
s_{\text{GFG},p} = \gamma_p(\Delta p_{A,p}(p_{A,H}^2 + 2p_{A,H}p_{A,H}(\rho_{1,H} + \rho_{2,H})))
\]

\[
- \Delta p_{A,p}(C_p - c_p) + \Delta p_{A,p}c_p
\]

\[
s_{\text{IMA},p} = \gamma_p(\Delta p_{A,p}(p_{A,H}^2 + 2p_{A,H}p_{A,H}(\rho_{1,H})))
\]

\[
+ \Delta p_{A,p}(p_{A,H}^2 + 2p_{A,H}p_{A,H}(\rho_{2,H}))
\]

\[
s_{\text{IMA},p} = \gamma_p(\Delta p_{A,p}(p_{A,H}^2 + 2p_{A,H}p_{A,H}(\rho_{1,H})))
\]

\[
+ \Delta p_{A,p}(p_{A,H}^2 + 2p_{A,H}p_{A,H}(\rho_{2,H})).
\]

To the order of these approximations, genetic associations (\( D_h, D_q \)) had no influence on the frequency of the alleles at the regulatory locus, \( M \). Instead, frequency change at the regulatory locus resulted from the direct effect of altered expression levels on fitness. Indeed, to leading order in the selection coefficients, equations 12–14 in hosts and 17–19 in parasites describe the change in fitness expected if a randomly chosen \( m \) allele were replaced by an \( M \) allele within an \( A \) heterozygote.

Examining the signs of equations 12–14 and 17–19 allows us to predict the directions in which expression levels should evolve in heterozygotes. These results are summarized in Figure 1A for the host and Figure 1B for the parasite. As is clear from Figure 1, selection typically favours the evolution of co-expression among hosts but rarely favours co-expression among parasites. These results are conceptually similar to recent findings on the evolution of ploidy levels [18]. In order to recognize and clear a wide array of parasites, selection favours hosts with a broader arsenal of recognition molecules, thus favouring diploid life cycles and the co-expression of alleles in heterozygotes. In contrast, in order to evade a host’s immune system or defence response, selection favours parasitic individuals that express a narrow array of antigens and elicitors, thus favouring haploid life cycles or expression of only one allele in heterozygotes. Exceptions to these general rules arise when selection acts in ways other than recognition and evasion. In the MA model, hosts are more likely to survive if they are difficult to mimic, which selects for a narrow expression pattern of only one allele. Furthermore, when costs are added to the GFG model, there are periods of time when selection favours expression of only the least costly allele (i.e., expression of the susceptible allele in hosts when virulence is common among parasites [see equation 12] or the expression of the avirulent allele in parasites when resistance is rare among hosts [see equation 17]).

To evaluate whether our analytical results are robust to violations of the assumption that recombination is frequent and selection is weak, we numerically iterated the exact recursions. For each genetic model of co-evolution, we considered both focal hosts and focal parasites, and modifiers that altered the expression probabilities \( \rho_1, \rho_2, \) and \( \rho_3 \) (Protocol S1). In each case, we considered all combinations of the following selection intensities (0.005, 0.05, and 0.50) and recombination rates (0.005, 0.05, and 0.50) and ran five simulations with randomly chosen initial allele frequencies. In the GFG model, we considered six levels of the costs of expressing only the resistance allele \( (C_h) \) or only the virulence allele \( (C_p) \): 15%, 30%, 45%, 60%, 75%, or 90% of the value of the fitness cost of infection in hosts, \( \gamma_p \), and the fitness cost of resistance in parasites, \( \gamma_p \), respectively. In addition, the costs of co-expression \( (c_h \) or \( c_p \)) were set to 33%, 66%, or 100% of the total costs of resistance or virulence \( (C_h \) or \( C_p) \). In all simulations, the modifier was introduced at an initial frequency of 0.5 after a 1,000 generation burn-in period had elapsed. All simulations were then run for an additional 4,000 generations, and the modifier was considered to have changed in frequency if its final frequency differed from its initial frequency by an amount greater than \( 10^{-15} \). This minimum threshold was set to eliminate false positives due to numerical imprecision and was based upon the maximum change in frequency observed for a modifier with no effect. The simulation results always coincided with the analytical predictions (Figure 1).

Taken together, our analytical and simulation results suggest that heterozygous hosts should generally evolve to co-express resistance alleles but heterozygous parasites should evolve to express only a single infection allele (Figure 1). It is not clear from the analytical results, however, which allele \( (B \) or \( b ) \), will ultimately be expressed in heterozygous parasites. Specifically, our analytical results suggest that expression of the \( B \) allele is favoured at some host allele frequencies, whereas expression of the \( b \) allele is favoured at others (see equations 17–19). Thus, the potential exists for patterns of parasite gene expression in heterozygotes to cycle over evolutionary time. Results from numerical simulations demonstrate that this is indeed the case. Cycles in parasite gene expression, where allele \( B \) was expressed during some periods of time and allele \( b \) at others, were frequently observed in IMA interactions and occasionally in GFG interactions with a cost of resistance (Figure 2). In contrast, cyclical patterns are less likely to persist in host species over long periods of evolutionary time because modifiers that increase co-expression generally spread to fixation (see Figure 1A). Only in the MA model do we expect long-term cycles in levels of dominance to potentially occur in both host and parasite.

**Discussion**

Our results demonstrate that co-evolution between hosts and parasites favours co-expression of alleles more often in hosts than in parasites. This predicted pattern is particularly striking among the models with the greatest empirical support (GFG and IMA) and helps explain observed patterns of expression of loci governing infection/resistance in hosts and parasites. Co-expression of resistance alleles has been observed in both the \( R \) gene family in plant hosts [37,38] and the major histocompatibility complex and immunoglobulin gene families in animal hosts [39]. In contrast, many parasites typically express only one of many antigen alleles encoded by large gene families. For instance, trypanosomes typically express only one of thousands of variant surface glycoprotein
In both panels, the frequency of a modifier allele that increases the expression of the B allele in the parasite population is shown in orange. The frequency of the B allele in the parasite population is shown in blue, and the frequency of the A allele in the host population is shown in maroon. Both panels considered an expression modifier introduced at an initial frequency of 0.5 with the following effects: \( p_1[M] = 0.75, p_2[M] = 0.50, p_3[M] = 0.25, p_4[M] = 0.25, p_5[M] = 0.5, p_6[M] = 0.75 \), and \( p_7[M] = 1 - p_1[M] - p_2[M] - p_3[M] \). Parameters for the IMA model (A) were \( q_1 = 0.15, q_2 = 0.20, q_3 = 0.25, \) and \( q_4 = 0.25 \). Parameters for the GFG model (B) were \( q_1 = 0.15, q_2 = 0.20, C_p = 0.075, C_r = 0.10, C_m = 0.0075, C_p = 0.01, f_m = 0.25, \) and \( f_r = 0.25 \). In both panels, the initial frequency of the A allele and the B allele was 0.55.

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Figure 2. Co-Evolutionary Dynamics of Parasite Gene Expression

In both panels, the frequency of a modifier allele that increases the expression of the B allele in the parasite population is shown in orange. The frequency of the B allele in the parasite population is shown in blue, and the frequency of the A allele in the host population is shown in maroon. Both panels considered an expression modifier introduced at an initial frequency of 0.5 with the following effects: \( p_1[M] = 0.75, p_2[M] = 0.50, p_3[M] = 0.25, p_4[M] = 0.25, p_5[M] = 0.5, p_6[M] = 0.75 \), and \( p_7[M] = 1 - p_1[M] - p_2[M] - p_3[M] \). Parameters for the IMA model (A) were \( q_1 = 0.15, q_2 = 0.20, q_3 = 0.25, \) and \( q_4 = 0.25 \). Parameters for the GFG model (B) were \( q_1 = 0.15, q_2 = 0.20, C_p = 0.075, C_r = 0.10, C_m = 0.0075, C_p = 0.01, f_m = 0.25, \) and \( f_r = 0.25 \). In both panels, the initial frequency of the A allele and the B allele was 0.55.

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