The contribution of epistasis to the architecture of fitness in an RNA virus

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The tendency for genetic architectures to exhibit epistasis among mutations plays a central role in the modern synthesis of evolutionary biology and in theoretical descriptions of many evolutionary processes. Nevertheless, few studies unambiguously show whether, and how, mutations typically interact. Beneficial mutations are especially difficult to identify because of their scarcity. Consequently, epistasis among pairs of this important class of mutations has, to our knowledge, never before been explored. Interactions among genome components should be of special relevance in compacted genomes such as those of RNA viruses. To tackle these issues, we first generated 47 genotypes of vesicular stomatitis virus carrying pairs of nucleotide substitution mutations whose separated and combined deleterious effects on fitness were determined. Several pairs exhibited significant interactions for fitness, including antagonistic and synergistic epistasis. Synthetic lethals represented 50% of the latter. In a second set of experiments, 15 genotypes carrying pairs of beneficial mutations were also created. In this case, all significant interactions were antagonistic. Our results show that the architecture of the fitness depends on complex interactions among genome components.

The nature, amount, and intensity of epistasis among alleles that affect fitness are barely known but are of extreme relevance for many evolutionary theories seeking to explain the origin and maintenance of genetic systems such as sex, recombination, diploidy, genetic canalization, and reproductive isolation; they are also an important component of S. Wright’s shifting-balance theory (1). Much effort has gone into finding and characterizing epistasis among genes by using different experimental approaches. Quantitative genetics experiments, in principle, are able to detect epistasis as a component of genetic variance in quantitative traits (2, 3). However, this approach often generates ambiguous results because of low statistical power. Other studies have shown compensatory evolution (4, 5), but these studies do not determine whether epistatic effects are rare or common in absolute terms, because compensatory mutations are favored by selection among a large set of possible mutations. Mutation-accumulation experiments have also been used to show deviations from additive fitness declines (6, 7), but this kind of experiment suffers from two limitations: (i) a lack of knowledge of the number of mutations accumulated and (ii) the fact that deviations from a linear fitness decline may indicate an absence of epistasis or widespread epistasis in which antagonistic and synergistic interactions are equally common (8). So far, the only straightforward approach that avoids all of the above problems relies on constructing genotypes containing exactly the same mutations alone and in combination, then measuring their individual and combined fitness effects and, finally, comparing the results with predictions generated under the null hypothesis of nonepistatic interactions. This approach has been applied to the bacterium Escherichia coli (8) and the fungus Aspergillus niger (9). Both studies reached a similar conclusion: epistases were widespread, but synergistic and antagonistic types were equally common. In addition, all of the above studies explored the interaction among deleterious or quasineutral alleles. However, to our knowledge, no study has yet explored the extent and sign of epistasis among those mutations responsible for adaptive evolution: beneficial mutations.

RNA viruses are characterized as having small compacted genomes, often with overlapping genes and multifunctional components (10). Therefore, epistatic effects on fitness should be especially important for these organisms (10). However, current models of viral evolution do not explicitly incorporate the existence of epistasis (11). Until now, only two studies have sought epistasis for fitness in RNA viruses. The first of them, with foot-and-mouth disease virus, failed to find such epistasis (12). The second, with the segmented bacteriophage φ6, found antagonistic epistasis (13). Nonetheless, these were mutation-accumulation studies, and therefore their conclusions have to be understood with caution for the reasons given above.

Here, we took a direct approach for characterizing the distribution of epistatic effects (8, 9). The starting point for our experiments was a collection of 91 single-nucleotide substitution mutants of vesicular stomatitis virus (VSV) created by site-directed mutagenesis (14). In a first set of experiments, we chose 28 of these genotypes that fulfilled the following two conditions (14): (i) the genomic position to be mutated and the nucleotide incorporated were both randomly chosen, and (ii) mutations had deleterious (although nonlethal) fitness effects. We randomly picked 47 pairs of these mutants and constructed the corresponding double-mutant genotypes. In a second set of experiments, we chose six genotypes for which the mutation incorporated had a beneficial fitness effect (14) and created all of the 15 possible double mutants resulting from combining these single mutations. Table 1, which is published as supporting information on the PNAS web site, contains information about each single and double mutant.

Materials and Methods

Site-Directed Mutagenesis. A full-length infectious VSV cDNA clone was used as template for creating the collection of single and double mutants (15). Site-directed mutageneses were performed as described in the QuikChange XL kit (Stratagene) user’s manual. It is important to mention that the method minimizes the chance of the appearance of undesired mutations by using the high-fidelity Pfu DNA polymerase. Sequencing of the cDNAs was done to confirm that each desired mutation was successfully incorporated.

As a first step, we introduced the substitution A-3853→C in the plus strand (Asp-259→Ala substitution in the G surface glycoprotein), which confers the ability of growing in presence of the I1 mAb [mAb-resistant mutant (MARM) phenotype] at a concentration that inhibits wild-type growth (16). This cDNA, named MARM R5V, was used as template for the rest of the mutagenesis.

For the set of randomly chosen mutations, the mutations were

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distributed in the genome as follows: four mutations were introduced in the nucleocapside N gene (one of them was synonymous); two mutations were generated in the P gene, the small component of the RNA polymerase; one mutant was created in the matrix gene, M; seven mutations were introduced in the envelope glycoprotein G gene (including two synonymous changes); and 12 mutations (including two synonymous substitutions) were introduced in the L gene, the major component of the RNA polymerase. In addition, two mutants were created containing one change in the M-G intercistronic noncoding region. For the set of beneficial mutations, all changes introduced were nonsynonymous: two mutants contained one change in the N gene, three in the M gene, and one in the G gene. More details about each mutant can be found in Table 1.

**Virus Recovery from cDNA Clones.** Approximately $10^5$ (90–95% confluent) baby hamster kidney (BHK21) cells (obtained from the American Type Culture Collection) were infected with a recombinant vaccinia virus that expressed T7 RNA polymerase. After incubation, cells were cotransfected with the mutant cDNA clone plus three support plasmids encoding the P, L, and N proteins of VSV (15). Transfections were done by using Lipofectamine Plus Reagent (Invitrogen) and adding 25 μg/ml 1-B-D-arabinofuranosylcytosine 6 h postinoculation (hpi) to inhibit vaccinia replication. The supernatant was harvested 96 hpi, and residual vaccinia was removed by filtering the supernatant through 0.2-μm membranes (Millipore). Dilutions (100- to 10^{-6}-fold) were plated on a fresh monolayer with 0.4% agarose in the overlay DMEM (supplemented with 5% calf serum). The presence of plaque-forming units (pfu) 24 hpi indicated the successful recovery of infectious VSV particles. Posttransfection titers, which ranged from $10^4$ to $10^6$ pfu/ml, were equalized to $\sim 5 \times 10^6$ pfu/ml before the competition assays to avoid any possible density effect on fitness (14). Failed transfection experiments were repeated until a positive result was obtained, with a maximum of five trials. The assignation of a mutant as putative lethal was done, taking into account our chosen pairs of mutations. For each double mutant, we ran 5 independent blocks for random mutations and in 10 blocks when beneficial mutations were combined. Each block contained each pair of singles as well as three replicates of MARM RSV. Doubles were assayed in different blocks that also contained MARM RSV. Fitness estimates of each mutant genotype ($W$) were adjusted by dividing $\omega$ by the grand-mean fitness of MARM RSV estimated in the same block. MARM RSV was effectively neutral relative to the wild type ($\omega = 0.997 \pm 0.009; t_{50} = 0.405, P = 0.687$).

**Quantifying the Strength and Direction of Epistasis.** Fitness was determined for each double mutant ($W_{ij}$) as well as for their corresponding single mutants ($W_i$ and $W_j$). Under the null hypothesis of nonepistatic effects, the expected fitness for the double mutant equals the product of the fitness estimated for each single mutation (i.e., $W_{ij} = W_iW_j$). If deleterious mutations were to interact in a synergistic way, then the observed fitness for the double mutant would be lower than expected by multiplying the fitness of both single mutations, and hence the difference between observed and expected fitnesses would become negative. By contrast, if deleterious mutations were to interact in an antagonistic way, then the observed fitness would be larger than expected under the null hypothesis of multiplicative fitness effects, and thus the difference between observed and expected fitnesses would become positive. From this argument, a convenient way of detecting the existence (and sign) of epistasis is by computing the index $e_{ij} = W_{ij} - W_iW_j$. For deleterious alleles ($W_i < 1$ and $W_j < 1$), synergistic epistasis is defined by $e_{ij} < 0$, whereas antagonistic epistasis is defined by $e_{ij} > 0$. For beneficial alleles ($W_i > 1$ and $W_j > 1$), the signs of $e_{ij}$ must be inverted.

**Results**

**Epistasis Among Pairs of Deleterious Mutations.** Fig. 1 shows the relationship between observed and expected fitness for each of the 62 double-mutant VSV genotypes. The solid line indicates the result expected under the null hypothesis of nonepistatic interactions. Let us first focus our attention on the 47 randomly chosen pairs of mutations. For each double mutant, we ran t tests assessing the significance of $e_{ij}$ (Table 1). Ten cases showed significant antagonistic epistasis, whereas three cases had synergistic epistasis (all $P = 0.042$). If we apply the Bonferroni method (17) to take into account the multiplicity of tests for interactions, which is a highly conservative approach, only three antagonistic interactions remained significant. Interestingly, in three cases, we were unable to replicate genotypes combining two viable mutations. These can be considered as putative synthetic lethals (18), which represent an extreme form of synergistic interaction. If putative synthetic lethals were removed (originally having single nonsynonymous mutations), however, presented one additional synonymous change with no presumably fitness effect (14). In conclusion, compensatory mutations occurring before competition experiments do not take place at a noticeable rate.

**Relative Fitness Assays.** The fitness of each mutant relative to the wild type was assessed by seeding $\sim 2.5 \times 10^3$ plaque-forming units (pfu) of each genotype into $\sim 10^5$ cells. To minimize the probability of fixation of new mutations during competition experiments, they were run for only 12 hpi. Preliminary assays showed that exponential growth occurred during this interval. During competition, the titers of both genotypes were determined by plating the appropriate dilution in the presence and absence of $I_1$ mAb. Intrinsice growth rates, $r$, were calculated as the slope of the log-titer against time (hpi) during the period of exponential growth. Relative fitness can hence be calculated as $\omega = \exp(r_{\text{mutant}} - r_{\text{wild type}})$. We used this fitness definition because it allows straightforward testing for epistasis in a multiplicative scale (see below). Assays were replicated in 5 independent blocks for random mutations and in 10 blocks when beneficial mutations were combined. Each block contained each pair of singles as well as three replicates of MARM RSV. Doubles were assayed in different blocks that also contained MARM RSV. Fitness estimates of each mutant genotype ($W$) were adjusted by dividing $\omega$ by the grand-mean fitness of MARM RSV estimated in the same block. MARM RSV was effectively neutral relative to the wild type ($\omega = 0.997 \pm 0.009; t_{50} = 0.405, P = 0.687$).
interactions was effectively symmetrical (Fig. 2; \( g \) abundant in both cases. Antagonistic interactions are more expected values were calculated from the fitness values of the corresponding mutations (after excluding synthetic lethals). The lower distribution was obtained from the 44 mutants carrying pairs of deleterious mutations (Fig. 2; \( t_{ij} \)). The conclusion was completely robust to changes in the sample sizes used during the bootstrap resampling, because 14 of 15 cases had \( \bar{e} < 0 \). This result gives even more strength to our conclusion of a predominance of antagonistic epistasis among beneficial mutations.

After studying each genotype separately, eight cases showed significant antagonistic epistasis (\( t \) tests, \( P < 0.027 \)) but none showed synergistic epistasis. Six antagonistic cases remained significant even after correcting the significance level with Bonferroni’s sequential method (17). Actually, in five of these instances, the fitness of the double mutant was even less than that of either single mutant. This particular case of antagonism between mutational fitness effects is known as decompensatory epistasis (1). Therefore, on average, a viral genotype carrying two beneficial mutations does not get the entire benefit individually associated with each mutation. Indeed, when epistasis is decompensatory, both beneficial alleles involved in the interaction cannot spread to fixation in the population, because the double mutant is less fit than each single mutant (1). As a consequence, lineages bearing alternative beneficial mutations should compete with each other on their way to fixation and, as a consequence of asexuality and clonal interference (19, 20), only the best competitor will eventually become fixed in the population.

**Discussion**

In this work, the distribution of epistatic interactions on fitness for an RNA virus has been estimated by using explicit pairs of single-nucleotide substitutions, each having a deleterious fitness effect. Among pairs of deleterious mutations, although both synergistic and antagonistic epistases have been detected, interactions were predominantly antagonistic, such that their combined effect is significantly smaller than expected under a multiplicative model. This result is in good agreement with very recent observations made with bacteriophage \( \phi \delta \) (15). A significant fraction of these deleterious, but still viable, mutations create synthetic lethals when combined. We also explored the distribution of epistatic interactions among beneficial mutations. Imaging the results obtained for deleterious mutations, antagonistic epistasis represents the most abundant type of interaction among beneficial mutations, with several cases showing decompensatory epistasis.

Previous studies with *E. coli* (8) and *A. niger* (9) found that both synergistic and antagonistic epistases were more or less equally common among deleterious mutations. The results we obtained for deleterious mutations are similar (i.e., both types of interactions exist) but not identical, because antagonistic epistasis are more abundant in VSV. This result might be a reflection of the differences in genome complexity and mutation rate. Evolution under the high mutation rate characteristic of RNA viruses should have been pushing VSV toward regions in sequence space that are more robust to the action of deleterious mutations (21). In this sense, because it involves masking the interaction among deleterious alleles, antagonistic epistasis might be seen as a sort of genetic mutational robustness (22).
The results reported here have two important implications for theories seeking explanations for the evolutionary advantage of recombination and sexual reproduction. First, according to the Fisher–Müller argument, sex and recombination are advantageous because they combine into a common genotype beneficial mutations that arose in different ones,speeding up the rate of adaptation (23,24). However, if the genetic architecture of RNA viruses determines that, in general, antagonistic epistasis and, in particular, decompensatory epistasis among beneficial mutations is the norm, then recombination would not necessarily imply a benefit in terms of adaptive evolution. Second, sex might still be beneficial for RNA viruses as an efficient mechanism for purging deleterious mutations (25,26). However, according with the Mutational Deterministic Hypothesis (27), if this is the case, an excess of synergistic epistasis among deleterious mutations is required to compensate the 2-fold advantage of clonal reproduction. Our first data set shows that synergistic interactions among random mutations are neither stronger nor more common than antagonistic interactions. Indeed, the existence of variability among loci in the sign and strength of epistasis, and especially the dominance of antagonistic epistasis, decreases the parameter space over which sex may evolve (28). Taken together, these two observations might explain why VSV, as many other RNA viruses, has evolved asexuality.

Despite the often invoked limitless adaptability of RNA viruses, the existence of abundant antagonistic epistasis among different genome components might impose a strong burden on viral adaptation. As stated by Holmes (10), “Although we can continue to marvel at the adaptability of RNA viruses . . . rather than thinking about what RNA viruses can do in their evolution, we should concentrate on their limitations. RNA viruses might be more at the mercy of their mutation rates than we think.” The existence of decompensatory epistasis among beneficial mutations might constrain adaptability of RNA viruses.

Finally, we would like to hint that the above findings prompt the necessity of considering nonmultiplicative fitness effects in mathematical descriptions of viral evolution. So far, the only mathematical model put forward to specifically explain viral evolution that explicitly incorporates epistasis (29) considers only the existence of compensatory mutations (i.e., antagonistic epistasis). Although this is a very good starting point, the results we present here suggest that more realistic models must incorporate variance in the type and strength of epistasis among mutations.

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