Evaluating the within-host fitness effects of mutations fixed during virus adaptation to different ecotypes of a new host

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The existence of genetic variation for resistance in host populations is assumed to be essential to the spread of an emerging virus. Models predict that the rate of spread slows down with the increasing frequency and higher diversity of resistance alleles in the host population. We have been using the experimental pathosystem *Arabidopsis thaliana*—tobacco etch potyvirus (TEV) to explore the interplay between genetic variation in host’s susceptibility and virus diversity. We have recently shown that TEV populations evolving in *A. thaliana* ecotypes that differ in susceptibility to infection gained within-host fitness, virulence and infectivity in a manner compatible with a gene-for-gene model of host–parasite interactions: hard-to-infect ecotypes were infected by generalist viruses, whereas easy-to-infect ecotypes were infected by every virus. We characterized the genomes of the evolved viruses and found cases of host-driven convergent mutations. To gain further insights in the mechanistic basis of this gene-for-gene model, we have generated all viral mutations individually as well as in specific combinations and tested their within-host fitness effects across ecotypes. Most of these mutations were deleterious or neutral in their local ecotype and only a very reduced number had a host-specific beneficial effect. We conclude that most of the mutations fixed during the evolution experiment were by drift or by selective sweeps along with the selected driver mutation. In addition, we evaluated the ruggedness of the underlying adaptive fitness landscape and found that mutational effects were mostly multiplicative, with few cases of significant epistasis.

1. Introduction

Characterizing the molecular basis of adaptation has been a central topic in modern evolutionary biology [1]. Compared with more complex organisms, this goal has proved to be relatively easy to achieve in RNA viruses owing to their small genomic sizes, which facilitates their full sequencing, the simplicity of the genotype-to-phenotype map, that allows performing functional analyses for many mutations, and the availability of reverse-genetic techniques to assess the effect of mutations individually or in given combinations. A common observation of such type of studies has been the identification of cases of convergent molecular evolution in independently evolved lineages [2–7]. This phenomenon is not only restricted to laboratory evolution experiments. It is also a relatively widespread observation among natural viral populations facing similar challenges, e.g. parallel changes are frequent in Human immunodeficiency virus type 1 clones isolated from patients treated with certain antiviral drugs, often following a common order of appearance [8–12]. Subsequent substitutions may confer increasing levels of drug resistance or, alternatively, may compensate for deleterious pleiotropic effects of earlier mutations [13–15]. Likewise, molecular convergences have been observed in the VPg cistron of
**Potato virus Y** populations that acquired the ability to break the resistance conferred by the *pop* 2 resistance gene of pepper [16–18]. This gene encodes for the ElF(iso)4G, whose correct interaction with VPg is essential for completing the virus replication cycle [16–18]. Convergent evolution at the molecular level is not controversial as long as it can be reconciled with the neutralist or the selectionist theories. The neutral theory suggests that convergences are simply accidents, whereas within the framework of selectionism, there are two qualifications for convergences. The first explanation considers convergences as being adaptive and the result of organisms facing the same or similar environments with a few alternative pathways of adaptation, as expected for highly compacted genomes. Second, given large enough population size and mutation rate (i.e. several beneficial mutations coexisting at any given time point) to make clonal interference an important evolutionary factor, the same mutations are expected to be fixed and in a more or less predictable order [19]. Another consequence of these pervasive genomic convergences is that although RNA viruses often respond quickly to strong selective pressures, their long-term evolutionary plasticity might be less impressive.

In recent years, we have been interested in exploring the molecular basis of virus adaptation to hosts that differ in their susceptibility to infection. Host populations with low genetic diversity in resistance-related loci show higher infection prevalence than populations with great diversity [20]. Since viruses evolve much faster than their multicellular hosts, the chances of generating escape mutants while replicating in a permissive host genotype are high, hence challenging the viability of host populations formed by individuals resistant only to the wild-type virus [21]. As a matter of fact, this process leads to local adaptation of parasites, where they have higher fitness in their current local host but lower in any foreign alternative host [22,23]. Local adaptation to a particular host genotype reduces the odds of successful transmission to a different one, thus minimizing the chances of epidemic spread. Despite this effect, too much diversity in susceptibility will allow infection by a wider range of parasites [24]. In consequence, the long-term outcome of the interplay between host and virus populations depends on the degree of genetic diversity of both contenders. The interaction between host genotypes and parasite genotypes has been modelled in the context of two different approaches. On the one hand, there is the gene-for-gene (GFG) model, in which a parasite genotype can infect all host genotypes and a universally susceptible host genotype exists [25]. Resistance occurs when a host ‘resistance’ gene is matched by at least one parasite ‘avirulence’ gene. Polymorphism in infectivity and resistance can be maintained only if virulence pays a cost. Infection matrices are, hence, nested. On the other hand, the matching-allele (MA) model is based on self- versus non-self-recognition systems in invertebrates. Infection is only possible if the parasite possess all alleles that match those of the host [26]. In this case, polymorphisms in infectivity and resistance are maintained by negative frequency-dependent selection and infection matrices are modular.

To tackle the effect of host genetic diversity for susceptibility to infection on the genetic composition and evolutionary dynamics of viral populations, we have chosen a model pathosystem formed by *Tobacco etch virus* (TEV; genus *Potyvirus*, family *Potyviridae*) and the experimental host *Arabidopsis thaliana* L. [7,27–30]. *A. thaliana* ecotypes vary in susceptibility to TEV [31]: some allow long-distance movement from inoculated to non-inoculated leaves while other support replication in inoculated leaves but do not allow for systemic movement. Susceptibility depends on the Restricted TEV Movement (RTM) multigenic system composed of the RTM1, RTM2 and RTM3 loci [31–37]. The presence of dominant alleles at all three loci is necessary for resistance; homozygous recessive mutations at any of the three loci result in systemic infection [34,35]. Aguadelo-Romero et al. [27] performed an evolution experiment in which TEV was adapted to the susceptible ecotype Ler-0 by serial passages. The ancestral TEV systemically infected Ler-0 plants, although the infection was asymptomatic. After 17 passages, the resulting strain, TEV-At17, fixed six point mutations, improved its accumulation ca 44-fold and induced severe symptoms. Comparative transcriptomics showed further differences between evolved and ancestral viruses: TEV-At17 downregulated developmental and metabolic processes, innate immunity and responses to abiotic stresses and to infection. Lalik et al. [29] showed that TEV-At17 systemically infected ecotypes that were resistant to the ancestral TEV. Furthermore, infectivity, viral load and severity of symptoms varied among ecotypes. Hillung et al. [30] compared the effect of TEV-At17b (a sequence variant of TEV-At17) infection on the transcriptome of the five ecotypes listed in table 1, finding differences in the way they perceived and responded to infection. The ecotypes could be classified into two groups according to their response to virus infection. In the first group, Ec-2 and Wt-1 upregulated genes involved in abiotic stresses and in the construction of new tissues: these ecotypes developed strong symptoms and, additionally, Wt-1 accumulated high viral loads. The second group (Ler-0, St-0 and Di-2) has mainly upregulated defence genes. Within this group, only Ler-0 developed severe symptoms as well as a very high viral load. St-0 and Di-2 developed mild symptoms, though Di-2 also accumulated high TEV-At17b loads. Finally, to evaluate the extent in which further evolution into a given plant ecotype (local host) conditioned the fitness of TEV-At17 derivatives into alternative ecotypes (foreign hosts), we performed the evolution experiments described by Hillung et al. [7] on the ecotypes.

### Table 1. List of *A. thaliana* ecotypes used in this study and the corresponding resistance phenotype according to the allelic combination at each RTM loci relative to the ancestral TEV.

<table>
<thead>
<tr>
<th>Ecotype</th>
<th>Origin</th>
<th>Genotype</th>
<th>Phenotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>Di-2</td>
<td>France</td>
<td>RTM1/RTM1 RTM2/RTM1 RTM3/RTM1</td>
<td>Resistant</td>
</tr>
<tr>
<td>Wt-1</td>
<td>Germany</td>
<td>RTM1/RTM1 RTM2/RTM1 RTM3/RTM1</td>
<td>Resistant</td>
</tr>
<tr>
<td>Ec-2</td>
<td>Germany</td>
<td>rtm1/rtm1 rtm2/rtm1 rtm3/rtm1</td>
<td>Sensitive</td>
</tr>
<tr>
<td>Ler-0</td>
<td>Germany</td>
<td>rtm1/rtm1 rtm2/rtm1 rtm3/rtm1</td>
<td>Sensitive</td>
</tr>
<tr>
<td>St-0</td>
<td>Sweden</td>
<td>RTM1/RTM1 RTM2/rtm3 RTM3/rtm3</td>
<td>Sensitive</td>
</tr>
</tbody>
</table>
Table 2. List of mutant TEV genotypes created for this study. All mutations were introduced in the TEV-Ar17b background [7].

<table>
<thead>
<tr>
<th>mutation</th>
<th>affected cistron</th>
<th>lineages in which mutations were observed</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1272U (synonymous)</td>
<td>HC-Pro</td>
<td>Di-2/1, Di-2/3, St-0/2, Wt-1/1</td>
</tr>
<tr>
<td>C8636U (G2831L)</td>
<td>CP</td>
<td>Di-2/1, Di-2/2, Di-2/3, Ler-0/1, Ler-0/3, Wt-1/3</td>
</tr>
<tr>
<td>A9240G (synonymous)</td>
<td>CP</td>
<td>Ler-0/1, St-0/2, Wt-1/2</td>
</tr>
<tr>
<td>G1272U/C8636U</td>
<td>HC-Pro/CP</td>
<td>Di-2/1, Di-2/2, Di-2/3</td>
</tr>
<tr>
<td>G1272U/A9240G</td>
<td>CP/CP</td>
<td>St-0/2</td>
</tr>
<tr>
<td>C8636U/A9240G</td>
<td>CP/CP</td>
<td>Ler-0/1</td>
</tr>
<tr>
<td>C2116U (synonymous)</td>
<td>HC-Pro</td>
<td></td>
</tr>
<tr>
<td>G6420A (synonymous)</td>
<td>CI</td>
<td></td>
</tr>
<tr>
<td>G1271U/G6420A</td>
<td>HC-Pro/CI</td>
<td></td>
</tr>
<tr>
<td>G3639A/G6420A</td>
<td>CI/NlaPro</td>
<td></td>
</tr>
<tr>
<td>C2116U/G3639A/G6420A</td>
<td>HC-Pro/CI/NlaPro</td>
<td>Ei-2/1, Ei-2/2, Ei-2/3</td>
</tr>
<tr>
<td>C795U (synonymous)</td>
<td>P1</td>
<td>St-0/1, St-0/3</td>
</tr>
</tbody>
</table>

listed in table 1. After 15 serial passages, we found that evolved viruses improved their within-host fitness, and showed higher infectivity and stronger virulence in their local hosts. We found that some ecotypes (e.g. St-0, Wt-1) were more permissive to infection than others (e.g. Ei-2, Di-2), and that viruses evolved in permissive hosts were more specialist than viruses evolved in more stringent hosts that became generalist, as predicted by a GFG model of infection.

The genome of evolved lineages was sequenced and we found that all lineages except Ler-0/2 contained at least one mutation, 43 being synonymous and 36 non-synonymous. Eight mutations appeared in multiple independent lineages (table 2). Three of these non-unique mutations were exclusive to Ei-2 lineages (C2116U, G3639A and G6420A). Mutation C795U was shared by St-0 lineages. Mutation G1272U was shared by Di-2 and St-0 lineages. Mutation A9240G was common to lineages Ler-0/1, St-0/1 and Wt-1/2. Very interestingly, these six convergent mutations were synonymous. Cases of convergent non-synonymous mutations were also observed: mutation C2912A was shared by lineages Di-2/1 and St-0/1, mutation C8636U was shared by all Di-2, all Ler-0 evolved lineages and by Wt-1/3, and lineages Ei-2/2 and Ei-2/3 each had a nucleotide substitution affecting the same codon at the CP cistron but resulting in different amino acid replacements. Convergent mutations are usually taken as an evidence of positive selection. Convergence at non-synonymous sites is usually explained as a consequence of identical selective pressures and the existence of limited accessible adaptive pathways. However, explaining convergences at synonymous sites is more problematic. Convergent synonymous mutations are a very common observation in evolving populations of RNA viruses [2,3,5,6,12] and studies analysing the mutational landscapes of RNA viruses have shown significant fitness effects associated to silent mutations [38–41]. Altogether, these observations support the notion that, at least for RNA genomes, equating synonymous substitution with neutral substitution is not always a valid premise. In our previous study, we provided some evidence supporting that selection for mRNA to protein translational efficiency might, in part, explain convergent synonymous mutations [7].

In this article, we focus our attention on the analysis of the adaptive value of convergent mutations observed during the TEV-Ar17b adaptation to ecotypes of A. thaliana that differ in their susceptibility to infection. More specifically, we want to evaluate the fitness effects in local and foreign hosts of convergent mutations and of their combinations. Are convergent mutations only beneficial in the local host wherein they arose or do they confer a host-independent benefit? Do mutations appear together in the same genotype because they contribute epistatically to the fitness benefit? Likewise, are mutations that do not appear together involved in reciprocal sign epistasis? To tackle these questions, we selected a set of six convergent mutations found by Hillung et al. [7] and previously classified as adaptive mutations (table 2). Five of these mutations were synonymous substitutions while only one was non-synonymous. These mutations were chosen on the basis of their coexistence in certain genotypes and apparent incompatibility with others. In total, we have created 14 genotypes, seven containing single mutations, six containing pairwise combinations of these mutations and one triple mutant. Some of these combinations were observed in the evolved lineages (G1272U/C8636U, G1271U/A9240G, C8638U/A9240G and G1271U/G6420A/G6420A; table 2), while others were not so, though they may have a transient existence in the evolving viral populations prior to the time at which they were sampled (G1271U/G3639A, C2116U/G6420A and G3639A/G6420A; table 2). In particular, we created these non-observed combinations to explore the local topography of the adaptive fitness landscape and its dependence on the host ecotype (i.e. environment). For each genotype, we have evaluated within-host fitness in the local host as well as in the four alternative foreign hosts.

2. Material and methods

(a) Plant ecotypes

The five A. thaliana ecotypes listed in table 1 were chosen for this study. According to their genetic makeup at the RTM loci, Ei-2, Ler-0 and St-0 are susceptible to infection with the ancestral TEV-At17b background [7].
Table 3. Relative within-host fitness of each viral genotype measured on every A. thaliana ecotype. Values are reported as average ± 1 s.e.m. (number of replicates). Asterisks indicate cases significantly different from TEV-Ar17b (Mann–Whitney test; FDR correction for multiple tests). Shadowed cells indicate the local host in which the mutation was detected. Last column shows the p-value of Kruskal–Wallis tests for differences in relative within-host fitness among ecotypes for each viral genotype.

<table>
<thead>
<tr>
<th>genotype label</th>
<th>Di-2</th>
<th>Wt-1</th>
<th>Ei-2</th>
<th>Ler-0</th>
<th>St-0</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>TEV-Ar17b</td>
<td>1.003 ± 0.008 (6)</td>
<td>0.999 ± 0.011 (4)</td>
<td>0.824 ± 0.109 (5)</td>
<td>0.996 ± 0.007 (9)</td>
<td>1.000 ± 0.2730 (2)</td>
<td>0.858</td>
</tr>
<tr>
<td>G1272U</td>
<td>0.831 ± 0.237 (2)</td>
<td>1.155 ± 0.000 (1)</td>
<td>0.969 ± 0.015 (7)</td>
<td>0.936 ± 0.060 (6)</td>
<td>0.702 ± 0.006* (5)</td>
<td>0.342</td>
</tr>
<tr>
<td>C8636U</td>
<td>1.062 ± 0.049 (2)</td>
<td>0.582 ± 0.013* (3)</td>
<td>1.028 ± 0.035 (3)</td>
<td>0.979 ± 0.006* (5)</td>
<td>0.722 ± 0.019* (2)</td>
<td>0.020</td>
</tr>
<tr>
<td>A9240G</td>
<td>0.995 ± 0.007 (4)</td>
<td>0.574 ± 0.007* (2)</td>
<td>0.984 ± 0.017 (10)</td>
<td>1.017 ± 0.018 (7)</td>
<td>1.198 ± 0.068* (8)</td>
<td>0.008</td>
</tr>
<tr>
<td>G1272U/C8636U</td>
<td>0.891 ± 0.161 (3)</td>
<td>0.552 ± 0.000* (2)</td>
<td>0.842 ± 0.073 (7)</td>
<td>1.020 ± 0.029 (5)</td>
<td>0.697 ± 0.000 (1)</td>
<td>0.046</td>
</tr>
<tr>
<td>G1272U/A9240G</td>
<td>0.604 ± 0.013* (4)</td>
<td>0.762 ± 0.214 (2)</td>
<td>0.575 ± 0.011* (4)</td>
<td>0.582 ± 0.011* (2)</td>
<td>0.750 ± 0.012* (3)</td>
<td>0.159</td>
</tr>
<tr>
<td>C8636U/A9240G</td>
<td>0.829 ± 0.197 (2)</td>
<td>n.d.</td>
<td>0.803 ± 0.077 (5)</td>
<td>0.760 ± 0.103 (5)</td>
<td>1.215 ± 0.014* (5)</td>
<td>0.017</td>
</tr>
<tr>
<td>C2116U</td>
<td>0.717 ± 0.109 (4)</td>
<td>0.797 ± 0.241 (2)</td>
<td>0.887 ± 0.054 (7)</td>
<td>1.000 ± 0.013 (5)</td>
<td>1.080 ± 0.095 (5)</td>
<td>0.066</td>
</tr>
<tr>
<td>G3639A</td>
<td>0.964 ± 0.079 (5)</td>
<td>0.678 ± 0.064* (9)</td>
<td>0.941 ± 0.017* (10)</td>
<td>0.985 ± 0.014 (10)</td>
<td>1.251 ± 0.012* (4)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>G6420A</td>
<td>0.992 ± 0.020 (3)</td>
<td>0.635 ± 0.072* (5)</td>
<td>0.763 ± 0.088 (6)</td>
<td>1.007 ± 0.011 (8)</td>
<td>0.918 ± 0.094 (5)</td>
<td>0.007</td>
</tr>
<tr>
<td>C2116U/G3639A</td>
<td>0.706 ± 0.103 (2)</td>
<td>0.917 ± 0.114 (4)</td>
<td>0.973 ± 0.011 (7)</td>
<td>0.914 ± 0.088 (4)</td>
<td>1.197 ± 0.013* (5)</td>
<td>0.009</td>
</tr>
<tr>
<td>C2116U/G6420A</td>
<td>0.742 ± 0.139 (3)</td>
<td>1.036 ± 0.021 (2)</td>
<td>0.790 ± 0.085 (5)</td>
<td>0.984 ± 0.009 (5)</td>
<td>0.985 ± 0.288 (2)</td>
<td>0.138</td>
</tr>
<tr>
<td>G3639A/G6420A</td>
<td>1.001 ± 0.014 (3)</td>
<td>1.032 ± 0.000 (1)</td>
<td>0.879 ± 0.086 (4)</td>
<td>0.955 ± 0.078 (6)</td>
<td>0.702 ± 0.004* (3)</td>
<td>0.079</td>
</tr>
<tr>
<td>C2116U/G3639A/G6420A</td>
<td>0.897 ± 0.124 (3)</td>
<td>0.555 ± 0.007* (3)</td>
<td>0.643 ± 0.031* (5)</td>
<td>0.886 ± 0.072 (6)</td>
<td>n.d.</td>
<td>0.015</td>
</tr>
<tr>
<td>C795U</td>
<td>0.927 ± 0.087 (5)</td>
<td>0.553 ± 0.005* (3)</td>
<td>0.948 ± 0.003* (3)</td>
<td>0.949 ± 0.047 (7)</td>
<td>1.260 ± 0.006* (4)</td>
<td>0.001</td>
</tr>
</tbody>
</table>

(b) Site-directed mutagenesis and production of infectious RNAs

The mutant genotypes listed in table 2 were generated by site-directed mutagenesis of a plasmid containing a cDNA of the TEV-Ar17b isolate, except synonymous substitution C6666U [7,30]. Mutagenesis was done using the PfuTurbo® DNA polymerase (Stratagene), as described in [42], and following the manufacturer’s instructions using the pairs of mutagenic primers listed in electronic supplementary material, table S1. The uniqueness of each mutation was confirmed by sequencing an 800 bp fragment encompassing the mutated nucleotide. Sequencing was done by GeneScreen (Lille, France) using BigDYE v. 3.1 in a 96-capillary ABI3730XL sequencing system (Applied Biosystems) and the most appropriated primers described in [27].

Infectious 5’-capped RNA of each genotype was obtained by in vitro transcription after BglII linearization of the corresponding plasmid as described in [42] and using the SP6 mMESSAGE mMACHINE kit (Ambion Inc.) following the manufacturer’s instructions. All RNAs were inoculated into Nicotiana benthamiana Dom plants to produce infectious viral particles. Infected N. benthamiana leaves were collected, homogenized and their viral load was quantified as described in §2c.

(c) Virus genomic RNA purification and quantification

Symptomatic whole plants were collected, ground into fine powder using a mortar and a pestle in liquid N₂ and stored at −80°C. RNA extraction from 100 mg tissue per plant was performed using InviTrap® Spin Plant RNA Mini Kit (Invitex) following the manufacturer’s instructions. The concentration of total plant RNA extracts was adjusted to 50 ng µl⁻¹ for each sample and the quantification of viral load was done by absolute real-time RT-qPCR using an external standard [43] as described elsewhere [29]. Amplifications were done using an ABI StepOnePlus™ real-time PCR System (Applied Biosystems) as follows: 5 min at 42°C, 10 s at 95°C following 40 cycles of 5 s at 95°C and 34 s at 60°C. Quantifications were performed in triplicate for each sample. Quantification results were examined using StepOne software v. 2.2.2 (Applied Biosystems).

(d) Evaluation of within-host fitness and epistasis

Previously quantified sap from infected N. benthamiana was diluted to the same viral concentration in all samples with buffer containing 50 mM sodium borate (pH 8.0) and 5 mM EDTA and was used for inoculation of A. thaliana plants. Two leaves from 21 day old A. thaliana plants were rub-inoculated with 10 µl of infectious sap and 10% carborundum (100 mg ml⁻¹). The inoculation of ecotypes was done in three temporal blocks, maintaining constant conditions of plant growth and inoculum. Between 1 and 10 (median 3) infected plants from each ecotype–mutant combination were collected 21 days post-inoculation (dpi) and processed as described in §2c. Total RNA was extracted and quantified by RT-qPCR as described in §2c.

A Malthusian growth rate per day was computed as

\[ m = \frac{1}{t} \log(Q_i/Q_0), \]

where \( Q_i \) is the number of TEV genomes/100 ng of total RNA quantified at t dpi by RT-qPCR. Relative within-host fitness was calculated as

\[ W_{xy} = \exp\left(m_{x,y} - m_{\text{TEV-Ar17b}}\right), \]

where \( m_{x,y} \) and \( m_{\text{TEV-Ar17b}} \) are the Malthusian growth rates of the viral isolate x and of the TEV-Ar17b isolate (estimated for the corresponding experimental block), respectively, evaluated in the same host h. Within-host fitness was evaluated for each mutant on the five different A. thaliana ecotypes.

Epistasis among a pair of mutations x and y was evaluated as

\[ e_{x,y} = W_{00h} - W_{xy} = W_{0y} - W_{xy} \]

where \( W_{00h} \), \( W_{xy} \), and \( W_{0y} \) are, respectively, the fitness of the ancestral TEV-Ar17b, of the double mutant, and of each single mutant [44], all evaluated in host ecotype h. In those cases for which epistasis turned out to be significant, we further proceeded to determine its type:
magnitude, sign or reciprocal sign. To do so, we have employed the inequality conditions derived by Poelwijk et al. [45]. All statistical analyses described in §3 have been done using IBM SPSS v. 21.

3. Results and discussion

(a) Exploration of within-host fitness data

Table 3 shows the within-host fitness values estimated for each genotype on each alternative host ecotype. For nine mutant genotypes, significant differences in within-host fitness have been observed among host ecotypes (Kruskal–Wallis test, \( p \leq 0.046 \)). However, by applying Fisher’s combined probability test we can conclude that, overall, differences among host types (table 2), only provided a significant beneficial effect in the non-permissive ecotype Ler-0. Mutation A9240G, identified in the most permissive host ecotypes (table 2), provided a significant beneficial effect only in St-0 and was significantly deleterious in Wt-1 (table 4). Mutation C795U, observed in two of the St-0 lineages (table 2), conferred a significant beneficial effect in St-0 and was significantly deleterious in Ei-2 and Wt-1 (table 3).

(b) Within-host fitness effects are greater in local than in foreign hosts

The above discussion (§3a) shows a complex interaction between the within-host fitness of each TEV genotype and the ecotype in which it is evaluated. If TEV genotypes were as well adapted to their local hosts as to the average foreign host, we would expect a positive correlation between these two measures of within-host fitness. Furthermore, in the hypothetical case of a perfect match in fitness among local and foreign hosts, the slope of the linear regression of fitness in the average foreign host on the local host(s) would be equal to unity. Deviations from the unity slope can be taken as indicative of fitness effects depending on the ecotype wherein they have been evaluated. A flatter-than-unity slope would indicate that the stronger the effects on the local host, the disproportionally weaker they may be in the foreign ones, thus suggesting that mutations contribute to specialization rather than to generalism. By contrast, a steeper-than-unity slope would indicate that the stronger the effect in the local host, the disproportionally stronger in the foreign ones they may be. To test this hypothesis, we first computed the median within-host fitness across all foreign hosts. In those cases where more than one local host existed, we also computed the median fitness. Figure 1 shows the relationship between within-host fitness in local and foreign hosts. The solid line represents the above null hypothesis of identical fitness among local and foreign hosts, the slope of the linear regression of fitness in the average foreign host on the local host(s) would be equal to unity. Deviations from the unity slope can be taken as indicative of fitness effects depending on the ecotype wherein they have been evaluated. A flatter-than-unity slope would indicate that the stronger the effects on the local host, the disproportionally weaker they may be in the foreign ones, thus suggesting that mutations contribute to specialization rather than to generalism.
For the eight genotypes above the line, fitness effects on the average foreign host are greater than expected given their effect on the local host ecotype. For the six genotypes below the line, fitness effects on the average foreign host are smaller than expected relative to their value on the local host ecotype. After thoughtful consideration of the properties of the genotypes at each side of the diagonal, we discovered that those below the diagonal were mostly obtained in the most permissive host ecotypes (as defined in [7]: Ler-0, St-0 and Wt-1; table 2: six out of eight instances), whereas the reverse is true for those genotypes above the diagonal (Di-2 and Ei-2; table 2: four out of 16 cases). Indeed, a Fisher’s exact test shows that this difference is significant ($p = 0.032$). Therefore, we provide further support to the conclusion of Hillung et al. [7] that more permissive hosts have selected for viral genotypes that are highly specialized for infecting their local host yet poorly capable of infecting more restrictive foreign hosts. By contrast, restrictive local hosts have selected for viral genotypes that are generalists successfully infecting every type of foreign host ecotype. This nested pattern of interactions is predicted by a GFG model of virus–plant interactions [25,46]. Nested infection networks are expected if a hierarchy of resistance among hosts and infection ability among viruses exists [46]. A GFG mechanism implies that mutations increasing fitness in the new local host exist that do not pay a fitness cost in Ler-0; thus the set of hosts that an isolate can infect are subsets of each other.

Very few experiments have tested whether the effects of adaptive mutations remain beneficial across all foreign hosts. In a pioneering study, Ostrowski et al. [47] found that mutations fixed in Escherichia coli lineages adapted to glucose as the only carbon source were also beneficial in five other carbon sources, concluding that positive pleiotropism was a norm for the bacterium and that local adaptation would not limit future evolution. In sharp contrast, evolution experiments with RNA viruses have shown that mutations expanding host range generally come with a cost in the original host [6,7,23,48–51], concluding that negative pleiotropism shall be the norm for these microparasites. The results reported here expand our previous observations [7,23,51] and give further support to this conclusion.

(c) Patterns of epistasis and the ruggedness of adaptive landscapes

Owing to their compact genomes and overlapping reading frames, epistasis is pervasive in RNA viruses (reviewed in [52]). The direction, magnitude and prevalence of epistasis is central to understanding many properties of virus genetic systems, such as reassortment of segments and recombination [53], complementation during co-infection [54], ploidy [55], phenotypic plasticity [56], robustness [57], the ruggedness of adaptive landscapes [45], or attempting to mechanistically explain dynamic processes such as the accumulation of mutations in finite populations upon transmission bottlenecks [58], and ultimately the origin of new viral species by an interruption of genetic exchange among populations [59].

In our previous work [7], we made two observations that pointed towards a possible role of epistasis and its dependence on the host ecotype. First, we found that mutations G1272U, C8636U and A9240G were common to lineages evolved in all ecotypes except Ei-2 (table 2), but their pairwise combinations were always exclusive to a given host ecotype (table 2): G1272U/C8636 in all Di-2-evolved lineages, G1272U/A9240G in lineage Ler-0/1. The second interesting observation involved mutations C2116U, G3639A and G6420A, their three pairwise combinations and the triple mutant. While the triple mutant was observed in all Ei-2-evolved lineages, the intermediate genotypes were observed neither in Ei-2 nor in other ecotypes. Henceforth, the observed sets of co-occurring mutations suggest that epistatic interactions played a prominent role in the genetic architecture of host adaptation. To assess this hypothesis, and more importantly the dependence of epistasis on the host ecotype, we constructed all the mutants shown in the genotypic network shown in figure 2. Table 4 shows all the epistasis values estimated. Looking at individual genotypes (rows in table 4), we found great variation in the direction, magnitude and prevalence of epistasis. Some mutations had positive or negative epistasis across all host ecotypes. On average, only C2116U/G6420A had a significant negative epistasis across host ecotypes. Moving to plant ecotypes (columns in table 4), the situation is similar: in all ecotypes we found cases of positive and negative epistasis.
in TEV. In Ei-2 and Lr-0, all but one case were of negative epistasis; by contrast, in Wt-1 positive epistasis was the most common. Finally, looking at individual combinations of mutations, only four cases have revealed significant epistasis. The combination of mutations G1272U/C8636U had a significantly negative epistasis coefficient in the foreign host Wt-1. The combination of mutations G1272U/A9240G was involved in significant negative epistatic interactions in the foreign hosts Ei-2 and Lr-0. Finally, the combination of mutations C3639A/G6420A was involved in significant positive epistasis in the foreign host Wt-1. In all other cases, we failed to detect a significant deviation from the null hypothesis of multiplicative fitness effects.

The significant negative interactions between genotypes G1272U/C8636U and C3639A/G6420A were of the sign epistasis type in Wt-1, thus suggesting the existence of local perturbations in the vicinity of these mutations in an otherwise smooth fitness landscape, although these perturbations are not strong enough as to create a deep valley. It is interesting that two out of two significant cases of sign epistasis reported are for the Wt-1 ecotype, suggesting that this host may impose some constraints that generate a certain degree of ruggedness in the adaptive landscape. The shape of the interaction between mutations G1272U/A9240G depends on the host ecotype where it is evaluated. It is of the reciprocal sign epistasis type in Ei-2, thus depicting a maximally rugged local surface where it is evaluated. It is of the reciprocal sign epistasis type in Wt-1, thus suggesting the existence of local fitness effects.

All other five mutations had a sign pleiotropic effect, i.e. the sign of a mutation’s effect on within-host fitness depends on the actual ecotype. Likewise, most of the multiple mutants analysed classify under the non-epistatic sign pleiotropy category, which always results in an obligatory cost of generalism [64]. Double-mutant C2116U/G6420A belongs to the non-epistatic no pleiotropy category, which always results in no-cost generalism [64]. Finally, mutations G1272U/A9240G and G1272U/C8636U are of the sign epistatic sign pleiotropy case. Under this scenario, populations may achieve either specialization or no-cost generalism, depending on the ecotype in which they evolved [64]. A scenario in which, depending on the ecotype wherein viral populations evolve, selection may favour genetic combination producing no-cost generalists, costly generalists or even specialists, explains the evolution of a GFG model of virus–plant interaction.

In conclusion, this study illustrates the necessity of performing reverse-genetic and functional analyses of mutations observed in evolving viral populations before concluding they are effectively beneficial and responsible for driving the observed phenotypic changes.

Data accessibility. All data necessary to reproduce the analyses reported in this article have been presented in tables 3 and 4.

Conflict of interest. We have no competing interests.

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