Costs and Benefits of Mutational Robustness in RNA Viruses

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SUMMARY

The accumulation of mutations in RNA viruses is thought to facilitate rapid adaptation to changes in the environment. However, most mutations have deleterious effects on fitness, especially for viruses. Thus, tolerance to mutations should determine the nature and extent of genetic diversity that can be maintained in the population. Here, we combine population genetics theory, computer simulation, and experimental evolution to examine the advantages and disadvantages of tolerance to mutations, also known as mutational robustness. We find that mutational robustness increases neutral diversity and, as expected, can facilitate adaptation to a new environment. Surprisingly, under certain conditions, robustness may also be an impediment for viral adaptation, if a highly diverse population contains a large proportion of previously neutral mutations that are deleterious in the new environment. These findings may inform therapeutic strategies that cause extinction of otherwise robust viral populations.

INTRODUCTION

Mutational robustness provides a population with the ability to maintain a given phenotype despite mutational perturbation (de Visser et al., 2003; Wagner, 2005). Previous theoretical studies have suggested that mutational robustness can enhance adaptation (Draghi et al., 2010; Masel and Trotter, 2010; Wagner, 2008). These studies proposed that in asexual organisms, robustness might allow higher connectivity of the neutral network of genotypes in a population, allowing better exploration of the sequence space and future access to potentially beneficial genotypes. In sexual organisms, it is expected that robustness will directly increase the amount of hidden (“cryptic”) genetic variation within a population (Gibson and Dworkin, 2004). Thus, robust populations should harbor substantial neutral genetic variation that may provide “stepping stones” to future adaptation (Masel, 2006; Waddington and Kacser, 1957).

The concept of mutational robustness has gained much interest in the context of RNA viruses (Elena, 2012; Lauring et al., 2013; O’Dea et al., 2010; Sanjuán et al., 2007), which display exceptionally high mutation rates, roughly on the order of $10^{-4}$–$10^{-6}$ errors per base per replication round (Sanjuán et al., 2010). Given the compact genome size of viruses, this level of replication fidelity results in 0.01–1 errors per genome per replication. Furthermore, because viruses typically generate enormous population sizes in the infected host (e.g., Haase et al., 1996), it has been estimated that all possible single mutations are created each day in a host (Coffin, 1995). Several recent studies have shown that a large proportion of these mutants will be either lethal or deleterious (reviewed in Sanjuán, 2010), suggesting that viruses suffer from an exceptionally high mutation load. Indeed, it has been shown that viral populations can be pushed to near extinction both through mutational meltdown (e.g., Crotty et al., 2001) and as deleterious mutations accumulate irreversibly in the genome (Chao, 1990; Duarte et al., 1992; Jaramillo et al., 2013) following the “Müller’s ratchet” effect (Smith, 1978).

Given this extreme selection regime, it is puzzling how viruses preserve functionality. Accordingly, it has been proposed that RNA viruses maintain different mechanisms of robustness that allow them to buffer the negative effects of mutations (Lauring et al., 2013). Large population size is an efficient strategy employed by viruses to purge and tolerate high mutation load (Elena, 2012). Cellular mechanisms like mutation “buffering” by molecular chaperones may play an important role in mitigating the negative effects of mutant viral proteins (Geller et al., 2007; R. Geller, C. Parrot, R. White Anderson, R.A., and J. Frydman, unpublished data). Another, highly prevalent, mechanism of robustness in viruses involves coinfection, when two or more viruses coinfect the same cell, and low-fit genotypes may be rescued by complementation with high-fit genotypes (Froissart et al., 2004; Montville et al., 2005).

Viral populations constantly experience new environmental challenges: they may experience different temperatures, and
infect different tissues, different hosts of the same species, and different hosts of different species. However, the central question whether mutational robustness is beneficial for evolvability in new environments (Draghi et al., 2010; Wagner, 2008) has remained experimentally unanswered and challenged. Some studies have shown that robust viruses were better prepared to evolve than their brittle counterparts in response to heat adaptation (McBride et al., 2008). Furthermore, cryptic neutral networks have also been implied in the adaptability of influenza virus (van Nimwegen, 2006). In contrast, a recent report showed that a more brittle viral strain adapted better to a host switch as compared to a robust strain (Cuevas et al., 2009).

Here, we examined the importance of mutational robustness (hereby termed robustness) as viruses establish an infection in a novel environment. We began by developing a theoretical framework that defines robustness through the distribution of mutational fitness effects (DMFEs) of a viral population. Although the model we use is general, in this study, we focused on robustness as manifested during viral coinfec tion. We first experimentally demonstrated that coinfection indeed buffers detrimental mutations, and then used these data to guide our theoretical modeling.

Most evolutionary studies have focused on mutations that are beneficial in a novel environment (Draghi et al., 2010; Gagneux et al., 2006; Hermisson and Pennings, 2005; Otto and Whitlock, 1997; Pennings and Hermisson, 2006). However, there has been little attention on the impact of novel deleterious changes. One exception is the study by Masel (2006), which examined how fixation of beneficial mutations (arising from cryptic genetic variation) is affected by a background of deleterious variants. Cryptic genetic variation was found to most likely allow more rapid adaptation.

Here, we set about to explore the effects of cryptic genetic variation in the unique settings affecting viruses: (1) rapid rate of mutation, (2) relatively frequent and often drastic environmental changes, and (3) large population size coupled with frequent bottlenecks. Importantly, in contrast to the assumptions in the Masel (2006) study, we consider environmental changes where alleles may completely change their fitness effects. This type of scenario has been described for drug-resistance and immune-evasion mutations (Agranovich et al., 2011; Andersson, 2006), and modeling these changes is important for understanding viral adaptation. Our model considers that, following an environmental change, alleles might become either beneficial or deleterious in the new environment. We note that due to the high rate of viral mutation, these changes will encompass both variants previously segregating in the population, as well as emerging mutations that occur after the environmental shift. We evaluated two types of environmental change in viruses: transmission to a susceptible host, and encountering a host response (e.g., drug) during infection. By computationally varying the multiplicity of infection (moi) of different populations, we were able to modulate levels of robustness of a population in realistic scenarios of viral infection and environmental change. We then were able to estimate the conditions under which viral populations have a higher probability of establishing a successful infection. Our results suggest that populations evolving at low moi, and hence lower robustness, may be more likely to successfully adapt to new environments.

RESULTS
Defining Mutational Robustness
During their life cycle, RNA viruses encounter a number of different environments and challenges. It is generally accepted that the genetic diversity of RNA virus populations affords them greater capacity for adaptation to novel environments. To test this hypothesis and gain further insight into the role of robustness in virus adaptation, we began by formulating a precise evolutionary definition of mutational robustness (r). We considered the distribution of mutational fitness (w) effects at each site of a genome (Figures 1A and 1B). We propose that mean fitness across all nonbeneficial mutants in the genome, \( r = \bar{w} \), defines the mutational robustness of a population. Robustness can then be estimated as \( r = q + p_{\text{del}} \bar{w}_{\text{del}} \), where q is the fraction of neutral mutations, \( p_{\text{del}} \) is the fraction of deleterious mutations, and \( \bar{w}_{\text{del}} \) is the mean fitness effect of a deleterious mutation. Essentially, r measures the mean fitness of new mutations. Hence, higher values of r correspond to more robust populations. We note that robustness is here defined as a property of the environment in which the population is replicating.

Mutational Robustness Is Mediated by Viral moi
To provide an experimental context in which robustness can be modulated, we focused on the condition under which viral populations are produced with respect to the virus-to-cell ratio. During higher moi, a given cell supports replication of two or more different genomes. In this scenario, genetic complementation can increase the robustness of the viral population. Although a mutated viral protein can limit or even halt viral replication at a low moi, this defect will be masked by “wild-type” (WT) versions of the same gene product in cells infected with more than one genome.

To determine whether indeed higher levels of coinfec tion correspond to higher robustness, we passed oral polio vaccine (OPV) type 2 (OPV2) strain under low moi (0.1) or high moi (>10) (Experimental Procedures). We then used a highly accurate sequencing method to determine the genetic landscape of the population (Acevedo et al., 2014). This approach enables us to detect rare mutational events with unprecedented accuracy at the single-nucleotide level. Our approach allowed estimation of the DMFEs of the virus populations growing at low moi and high moi. OPV2 grown at high moi showed a DMFE consistent with higher mutational robustness (r = 0.49) as compared to low moi (r = 0.37) (p < 1 × 10^{-15}, t test; p < 1 × 10^{-15}, Kolmogorov-Smirnov test). Indeed, we observed that populations that replicated at high moi bear a larger proportion of neutral alleles and a lower proportion of lethal alleles than those populations replicating at low moi (Figures 1C and 1D). These results are consistent with previous observations suggesting that coinfec tion buffers the effects of deleterious alleles via complementation (Froissart et al., 2004). Thus, buffering allows alleles to be present at higher frequencies in the population and to present as less deleterious (more neutral) than under low coinfection conditions. Notably, we evolved the OPV2 strain in an environment where it is not at equilibrium. This strain was originally produced in nonhuman cells, whereas our experiment was carried out in HeLa cells (Experimental Procedures). Under these conditions,
we detected beneficial alleles, which we have not included in our approximate estimation of robustness ($r$). The role of robustness in beneficial mutations and adaptation is discussed in Supplemental Information section 5.

**Modeling a Transmission to a Susceptible Host**

We next carried out numerical simulations to determine how robust and brittle virus populations adapt following transmission between hosts. Our model compares two different situations: one in which the transmitting virus population derives from the focal point of infection, where moi is presumable high, and a second scenario in which virus derives from a more disperse infection, where moi is assumed to be lower. Upon transmission to a novel host, viral populations experience a population bottleneck (e.g., Betancourt et al., 2008), i.e., reduction in population size, and thus the initial replication cycles will likely occur at low moi (Abrahams et al., 2009). Our aim was to compare the effect on population fitness upon infection of a new host of viruses produced at low moi (brittle) to those produced at high moi (robust).

To model transmission between hosts, we computationally imposed an environmental change on populations of viruses evolving at equilibrium (Experimental Procedures). Such a change was introduced through reassigning selection coefficients to a random number of sites in the genome: some sites were reassigned with a negative (deleterious) selection coefficient, whereas some sites were assigned with a positive (adaptive) selection coefficient (Figure 2A). In order to consider a realistic case of transmission, we further modeled a strong reduction in population size during the environmental change, followed by logistic growth (see also Supplemental Information section 1 and Figure S1). Notably, because we reassign selection coefficients randomly, robustness in our study is related directly to revelation of cryptic genetic variation (see also Richardson et al., 2013 on the complex relationship between robustness and revelation of cryptic genetic variation).

We used simulation and theory to estimate mean waiting times as a function of robustness until (1) a novel beneficial allele takes over a population (adaptation), and (2) a novel deleterious allele is purged from the population (clearance) following transmission (Experimental Procedures). Our results show that mean waiting time for adaptation decreases as robustness increases (Figure 2). This occurs because robustness shifts the DMFE to include more sites where mutations have neutral fitness effects, and thus, robust populations harbor a larger number of neutral
genetic variants segregating at appreciable frequencies as compared to brittle populations. When such higher-frequency “hidden” alleles become beneficial, the rate of adaptation will be faster (Hermisson and Pennings, 2005). However, the higher frequency of neutral alleles has an additional consequence. When measuring the rate at which deleterious mutations are removed, we observed that robust populations purge the novel deleterious mutations more slowly (Figure 2; Supplemental Information section 1), and they appear to be at a disadvantage as compared to more brittle populations.

We next examined which of these two opposite effects dominate during an environmental change. Although the effect of purging appears to be less significant than the effect of adaptation (Figure 2), the first few generations of viral replication are critical for establishing a novel infection. To this end, we determined the mean fitness of the population as it adapted to the new environment, which affects multiple sites and their fitness, over time. This allowed us to compare a robust \( r = 0.97; q = 0.8 \) with a non-robust \( r = 0.9; q = 0.2 \) population and determine which population has a higher probability to establish a productive infection. We first derived a simple analytic expression to estimate the difference in log mean fitness of a robust versus brittle population, given the effect of an environmental change on numerous alleles in the previous environment (Supplemental Information section 2). Under these conditions, the difference in log mean fitness of a robust versus brittle population is approximated by \( \Delta w = \Delta q \cdot \sum \log((1 + s_i^{\text{new}})^T - 1) \cdot X_0 + 1 \), where \( \Delta w \) is the difference in log mean fitness between a robust and brittle population over \( T \) generations, \( s_i^{\text{new}} \) is the selection coefficient at site \( i \) in the new environment, and \( X_0 \) is the mean frequency of a neutral allele in the first environment. This term can be positive or negative depending on the number of positive and negative selection coefficients in the new environment.

The analytic term above relies on the assumption of deterministic selection. Yet, fluctuating viral population sizes may allow a powerful role for genetic drift. We thus used stochastic simulations to explore the adaptation of viral populations to a new environment (Experimental Procedures) (Figure 3). Notably, it has been shown that exponential microbial growth leads to a loss of beneficial mutations due to stochastic effects (Wahl and Gerrish, 2001; Wahl et al., 2002), and this might cause deviations from the analytic approach described above.

Initially, we explored two extreme cases. In one scenario, the environmental change results in a larger proportion of beneficial alleles in the new environment \( \sum s_i^{\text{new}} = 100, \sum s_i^{\text{old}} = 5 \), Figure 4A). In the other case, only a few alleles became beneficial, whereas the majority of them became deleterious \( \sum s_i^{\text{new}} = 5, \sum s_i^{\text{old}} = 100 \), Figure 4B). In order to test which population would outperform its counterpart, we calculated the log mean fitness of the robust and brittle population during the first generations following the environmental change. The difference in fitness between the two populations hence corresponds to the shaded differences between the robust (gray) and brittle (black) curves (Figures 4A and 4B). We observed that whereas the originally robust populations adapt more rapidly when many alleles become beneficial in the new environment, they are also significantly less fit during the initial rounds of replication in the new environment if a large proportion of neutral alleles becomes detrimental. Indeed, under this scenario, the originally robust population suffers from a severe drop in fitness following the environmental change, due to the sudden influx of deleterious variants in the population segregating at a higher frequency (Figure 4B).

To examine the larger spectrum of possible scenarios, we next simulated a series of 400 environmental changes, which encompassed relatively widespread changes of selection coefficients in the genome defined by randomly drawing \( N_r \) and \( N_d \) sites, whose selection coefficient was reassigned after the environmental...
1% of the sites in the genome), and all combinations of deleterious sites (blue region in Figure 4C). Under the latter scenario, the brittle population outperforms the robust population, namely when the magnitude of adaptive sites is much smaller than the magnitude of deleterious sites (blue region in Figure 4C). We also identify a range of conditions where the brittle populations have an advantage for adaptation to some new environmental conditions, but in the majority of new environments, brittle populations have a larger advantage as compared to the WT fitness in the native host (caption in Figure 4C; Experimental Procedures). This set of results suggests that a switch to a new environment may result in a predominantly higher load of deleterious alleles in the new environment. Under this scenario, we can roughly extrapolate that the brittle virus populations would outperform their more robust counterparts (Figure 4C, dashed circle; Supplemental Information section 3; Table S1).

Acute versus Persistent Modes of Viral Infection

Our analytic results coupled with the simulations showed that mean fitness of a virus population depends on the level of neutral diversity in the population prior to the environmental change. Theoretical models often assume that allele frequencies are at mutation-selection balance. However, for acute virus infections, this balance is unlikely to be attained given that the environment and other features of infection frequently fluctuate. Of note, the current and previous studies showed that some acute viral infections accumulate limited diversity during infection of a host (Acevedo et al., 2014; Thai et al., 2012). In contrast, during persistent infections, population diversity should increase with time as the virus replicates within the infected individual, akin to a population accumulating genetic charge (see Le Rouzic and Carlborg, 2008). Accordingly, we hypothesized that the effect of robustness will be more pronounced in persistent infections.

To test this hypothesis, we examined whether or not acute versus persistent viruses are equally affected by the level of robustness during transmission events. We simulated two possible scenarios: (1) for acute viruses, we allowed shorter waiting times between environmental changes (corresponding to transmission events) \( (T = 20) \) and larger maximal population size \( (N_{max} = 10^8) \); and (2) for persistent viruses, we allowed longer waiting times between transmission events \( (T = 100) \) yet smaller maximal population sizes \( (N_{max} = 10^7) \) (Experimental Procedures). These parameters were selected based on evidence suggesting that acute infections produce large population sizes in a short period of time, whereas persistent infections often are characterized by smaller size populations (Grenfell et al., 2004). As a consequence of the increased time for replication, under persistent infection conditions, we observed a significant increase in genetic diversity (data not shown). Importantly, our simulations showed that robust populations have an advantage for adaptation to some new environmental conditions, but in the majority of new environments, brittle populations have a larger advantage as compared to the acute infection viruses. This effect is also observed in acute infections, but the effect is less pronounced (Figure 5). We conclude that robustness determines diversity, diversity is increased in robust populations that replicate for a longer period of time in the same environment (persistent infections), and diversity can be either beneficial or detrimental in a new environment.

**DISCUSSION**

The role of mutational robustness in evolution has remained controversial. In this study, we begin with a simple quantitative...
definition of mutational robustness (see also Elena et al., 2007; Krakauer and Plotkin, 2002). This definition is highly amenable to direct measurement in populations of evolved RNA viruses, and we show that viral coinfection is a relevant model for studying the effects of mutational robustness. We then construct a theoretical model that is guided by our experimental results (Figure 1). This process allows us to predict how viral populations, modulated by different levels of moi, would adapt to new different conditions.

Mutational robustness has been shown to be prevalent in many organisms. In several cellular organisms, it is present in the form of gene redundancy, modularity, and alternative regulatory pathways. In fact, for many organisms, most gene knockouts are viable (Costanzo et al., 2011). In principle, it would be reasonable to expect selection to maximize mutational robustness because this serves to decrease mutational load and increase diversity. However, experimental evidence suggests that virus populations are mostly composed of lethal or deleterious mutations (Acevedo et al., 2014; Rihn et al., 2013; Sanjuañ, 2010). This has been explained largely by the small genome sizes of viruses, which limit redundancy. However, an alternative explanation is that brittle populations may also have an advantage under certain conditions. This advantage of brittle populations manifests as lowered sensitivity to novel deleterious mutations and, subsequently, higher fitness in a new environment. Thus, brittle populations suffer less from the transient decrease in fitness in a novel environment (Figure 4B).

Limitations of our study are in the decoupling of the mechanisms that generate robustness in the first environment and the proportion of beneficial-to-deleterious mutations in the new environment. Supplemental Information section 4 and Table S2 discuss different mechanisms of robustness and their potential effects on fitness. In the study herein, we find that high moi buffers deleterious mutations and masks the effects of beneficial mutations, as evident from the changed distributions in Figure 1C versus Figure 1D (see also Supplemental Information text 5 and Figure S2). However, other mechanisms of robustness may well directly affect the distribution of fitness effects in the new environment. The authors conclude that the study provides insights into the evolution of mutational robustness and its impact on viral fitness.
environment, and an explicit model, which ties the effects of robustness with the effects of the new environment, is warranted.

Our current model, for simplicity, does not take into account linkage among sites, entailing infinite recombination. Background selection, genetic hitchhiking, and genetic interference are all manifestations of the Hill-Robertson (HR) effect (Felsenstein, 1974; Hill and Robertson, 1966), which occurs when selection and linkage combine. Some of the important derivations of the HR complex effect are that the effective population size is reduced, genetic variance is reduced, and selection is less effective (e.g., McVean and Charlesworth, 2000; Roze and Barton, 2006). Multilocus HR interference has been shown to be prevalent in simulations and, in fact, may lead to a selective advantage for recombination (Barton and Otto, 2005; Keightley and Otto, 2006). Interestingly, negative disequilibrium (associations between deleterious and beneficial alleles) is expected to prevail over positive disequilibrium (Barton and Otto, 2005). Even more complex effects may take place when rates of recombination are low, such as clonal expansion of a beneficial genotype. Furthermore, viral colonization may involve structured populations replicating autonomously, different degrees of coinfection and genetic complementation, effects of migration, and strong competition for resources (Turner and Chao, 1999). It remains to be studied how these different phenomena affect the relationship between robustness and adaptation. This will be facilitated by developing explicit models of viral evolution, which take into account the complex processes occurring during viral establishment of an infection.

The framework proposed in this study may help resolve conflicting evidence on the role of mutational robustness in promoting evolvability: whereas McBride et al. (2008) showed that robust phage clones adapted more successfully to a new thermal niche, Cuevas et al. (2009) showed that brittle vesicular stomatitis virus populations were more adaptable to a new host cell type than their robust counterparts. The results of our study support the notion that, depending on the conditions, either robust or brittle populations can successfully establish an infection. We conclude that the change in the distribution of fitness effects in the novel environment as compared to the previous environment will determine the outcome of an infection.

Previous studies by Masel (2006) concluded that robustness should promote evolvability even in the presence of cryptic alleles that are deleterious in a new environment. Our results on the other hand identify a range of conditions where a brittle population would prevail, namely when many neutral (or near neutral) mutations drastically change their selection coefficient into a deleterious one. These conditions will largely depend on whether mutation effect distributions are correlated or not across environments. Although Masel (2006) assumed that mutational fitness values are correlated between environments, we relaxed this assumption and considered that fitness values of mutations do not necessarily correlate between two given environments.
Another difference between these studies is that the Masel (2006) study takes into account conditionally lethal mutations, whereas we focus only on conditionally deleterious mutations. However, we argue that considering conditionally lethal mutations would not change the conclusions of our study, given that they increase the negative effects on fitness of a population in the new environment.

We thus propose that different modes of virus replication and different types of environment conditions will most likely dictate whether fitness effects are indeed correlated. For instance, some viruses that infect multiple hosts (such as vector-borne viruses) are expected to regularly alternate between very different environments, whereas other viruses may experience relatively similar environments, and hence, the mutation effect distributions might not change dramatically. Interestingly, it has been shown that vector-borne RNA viruses experience more purifying selection than nonvector-borne viruses (Woelk and Holmes, 2002). Because evolutionary rates and mutation rates are highly correlated in viruses (Sanjuan, 2012), increased purifying selection implies more deleterious and lethal mutations. Because vector-borne viruses experience frequent dramatic environment changes and, hence, a higher proportion of deleterious mutations in the new host (i.e., the blue region in Figure 4C), this suggests that selection may have favored such brittle populations over the course of evolution of vector-borne viruses.

Finally, our results suggest an interesting relationship among the rate of environmental change, the accumulated diversity, and mutational robustness. The time to accumulate neutral mutations in a relatively constant environment is significantly longer during persistent infection of robust populations. As a result, robust populations, replicating under persistent infection, should accumulate a considerably larger diversity than brittle viruses. Indeed, we show that persistent infection exacerbates the effects of robustness (Figure 5), and depending on the conditions of the new environment, the degree of robustness in the population would be a critical factor in the establishment of infection in the new environment. Considering thousands of years of evolution, brittle populations of persistent viruses would have an increased probability of establishing infection if a large proportion of neutral mutations become detrimental in the new environment. We thus speculate that under these conditions, brittle viruses that use a persistent mode of infection are more likely than robust viruses to successfully infect in diverse environments. On the other hand, acute infection viruses, which undergo rapid transmission and generate less genetic diversity, will be less affected by robustness. This conclusion seems counterintuitive in light of theory and experiments, which show an increased rate of extinction as the rate of environmental change increases (e.g., Lindsey et al., 2013). However, this may be explained given that our framework assumes a random process of environmental change and, hence, lack of correlation between selection coefficients before and after an environment changes. Although this requires further empirical study, it is likely that successive viral transmission events do not adhere to a gradual process of unidirectional change (e.g., climate change). Under the latter scenario, populations would gradually climb to a new fitness peak, and we would not expect an abundance of novel deleterious mutations as the unidirectional change proceeded.

The ability to precisely measure the DMFEs of viral populations as they adapt to different environmental changes will be central to establishing the constraints affecting viral evolution and will allow a better understanding of the link among standing genetic variation, mutational robustness, and capacity to evolve.

**EXPERIMENTAL PROCEDURES**

**Experimental Evolution of Viruses and Fitness Measurements**

To understand population diversity and robustness dynamics under different pressures, we passaged OPV2 in HeLa S3 (ATCC; CCL22) cells at low and high moi. For low moi passages, 10^7 HeLa S3 cells seeded the day before the experiment were infected with OPV2 at a moi of 0.1 at 33°C for 1 hr to allow virus adsorption, and then replaced with virus culture medium. The infected cells were maintained at 33°C for 10 hr and then harvested by freezing at –80°C. After three freeze-thaw cycles, virus suspension was clarified by centrifugation at 3,500 x g for 10 min at 4°C, and stored at –80°C for future passages. Plaque assays were performed to determine virus titer of each passage for a subsequent seven passages.

For high moi passages, all passages were obtained by infecting HeLa S3 cells at a moi of 0.1 as described above but were harvested at 24 hr postinfection. Thus, viruses experienced one replication cycle at a low moi of 0.1 and an additional cycle at a higher moi (>10; inferred assuming a burst size of >100 infectious particles per infected cell, based on low moi titers). Plaque assay was also performed to determine virus titer of each passage for subsequent passages. All viral populations were amplified once in HeLa S3 to increase the percentage of viral RNA for sequencing. Library preparation, base calling, and mutational fitness were calculated based on time-series allele frequency measurements as described in Acevedo et al. (2014) for CirSeq. Briefly, we assume deterministic selection, which is approximated by

\[
P(t+1) = wP(t)(1 - \mu) + (1 - P(t))\mu. \tag{1}
\]

where \(P(t)\) is the frequency at generation \(t\). Assuming linear behavior at small time intervals, we obtain the following ordinary differential equation:

\[
\frac{dP}{dt} = \frac{P(t + \Delta t) - P(t)}{\Delta t} = \frac{P(t)(a + \mu). \tag{2}
\]

where \(a = (w(1 - \mu) - \mu - 1)\). Solving this equation yields

\[
P(t) = ce^{\mu t} - \frac{\mu}{a} \tag{3}
\]

The mutation rate \(\mu\) is estimated based on the median frequencies of lethal mutations as described in Acevedo et al. (2014). Thus, Equation 3 allows us to infer the constant c and fitness \(w\) for any allele, given a pair of frequency estimates \((P(t_i)\) and \(P(t_j))\) at any two passages. For multiple passages, we use linear regression to estimate a unique \(w\) value. We note that this approach will attribute all changes in allele frequencies to the combined effect of mutation and selection, while ignoring the effect of genetic drift and sampling. For rare alleles, with small allele counts, the effect will be to overestimate the effect of selection. There are a number of approaches for incorporating genetic drift and sampling variance into the estimation of selection coefficients (Boilback et al., 2008; Mathieson and McVean, 2013; Song and Steinrueck, 2012). However, these methods are highly computationally demanding and are not developed for analyses of thousands of sites. Furthermore, our results rely on comparing distributions rather than comparing estimates for single sites. For that reason, we use the deterministic approximation and interpret the results accordingly.

**Virus Evolution Model**

We assume a haploid population of \(N\) individuals, with a genome length \(L\) of 10,000 bp. Mutations were modeled assuming two allelic states in each site with an equal rate of mutation between the two states of 10^{-5} mutations/replication/base. Each population is assigned a DMFE, from which a selection
coefficient is drawn for one of the alleles at each position (labeled the alternative allele), whereas the fitness of the other allele (labeled the WT allele) is set to 1. The total fitness was calculated assuming multiplicative effects across sites (see below). We assume that the DMFE follows a mixture of a log-normal distribution of nonlethal deleterious selection coefficients ($s < 0$) with probability 1, and a point mass at $s = 0$ with probability $q$. Robustness of a population of viruses is defined as $r = \pi$. A survey across several DMFEs of various RNA and DNA viruses has shown a highly similar distribution for nonlethal deleterious selection coefficients by Sanjuán et al. (2004). Hence, here, $q$ effectively defines the robustness of the population, and values of 0.2 and 0.8 are explored, corresponding to $r = 0.896$ and 0.974, respectively. We note that $q$ will also capture any near-neutral mutations that “behave” neutrally when $s < 1/N_0$ (where $N_0$ is the effective population size).

To simplify our calculations, we made several assumptions: (1) we assume absence of linkage disequilibrium, and (2) we considered only nonlethal mutations across both environments modeled. The latter assumption allows us to focus on the “effective” portion of the genome where genetic variability exists. The former assumption on lack of linkage entails infinite recombination. Although this is not a realistic assumption, for many RNA viruses, the rate of recombination is exceptionally high (e.g., Kirkegaard and Baltimore, 1986; Runce et al., 2013; Tromas et al., 2014). This high rate is especially true for poliovirus studied herein. We have addressed this important limitation of our model in the Discussion and in Supplemental Information section 6, where we show that our results may likely hold when alleviating the assumption of lack of linkage.

**Simulations**

We perform Wright-Fisher simulations with selection, where binomial sampling probabilities are weighted by the fitness of the allele:

$$Pr(X_t = x_t|X_{t-1} = x_{t-1}, s, \mu, N_t) = \frac{N_0^N}{(N_0 - x_{t-1})! (N - x_{t-1})!} \left(1 - P_0\right)^{N_0 - x_{t-1}} \left(P_0\right)^{x_{t-1}}$$

(Equation 4)

where $X_t$ is the frequency of the minor allele at time $t$, $s$ is its selection coefficient, $\mu$ is the mutation rate, and $N_t$ is the population size. $P_0$ is the probability of sampling an allele in the next generation and is determined by its relative fitness as compared to the WT: $P_0 = \left(1 + s\right)X_{t-1}/\left(1 - \mu\right) + \mu(1 - X_{t-1})$ (as in Equation 1).

Each site in the population of genomes is simulated independently. To initialize the first environment, each site is first simulated with the WT allele frequency set to 1, and evolved for 150,000 generations with a population size $N = 10^5$. Then, an environmental change is imposed, which changes the selection coefficients at a varying number of random sites in the genome. Because selection coefficients are always relative to the WT allele (which is assigned a value of 1), changes are only made to the alternative allele. We note that this implicitly captures all types of changes in selection, both to the WT and mutant alleles, because we allow a change in the magnitude and sign of the novel selection coefficient. Thus, for example, a situation where the WT allele becomes deleterious in the new environment would be captured in our simulations as the alternative allele becoming beneficial. The population is then simulated for several generations in the new environment. For the “transmission scenario,” a severe bottleneck is assumed, and the population size is reduced to $N_0 = 100$ as the environmental change takes place. This is followed by population growth modeled as a logistic density-dependent growth of the population (Acevedo et al., 2014; Sanjuán et al., 2004; $N_t = K(N_t/N_0 + e^{-r}(K - N_0)$), where $K$ (maximal carrying capacity) was set to $10^5$ for the “transmission scenario,” and $r$ (intrinsic growth rate) was set to 2. This allowed explosive growth, as observed during the acute infection phase of several viruses (Otto and Whitlock, 1997). Different values of $r$ (0.5, 1, and 2) yield essentially the same results (data not shown). For the “within-host scenario,” population size remained large ($N = 10^5$) before and after the environmental change. When a buffering effect is modeled, the beneficial fitness effect is reduced from 1.1 to 1.04 (~6% decrease), and the deleterious fitness effect is increased (selection coefficient is decreased) from 0.9 to 0.93 (~4% increase), in line with the experimental results determined on the effects of high moi.

**Mean Fitness of the Digital Viral Populations**

Assuming sites are independent, log mean population fitness at a given generation of simulated populations was calculated as $\log(W) = \sum_{i=1}^{n} \log(1 + s_i f_a + 1(1 - f_a))$, where $s_i$ and $f_a$ are the selection coefficient and frequency of the alternative allele at position $i$ of the genome, respectively. The Supplemental Information elaborates on calculating mean fitness over time analytically.

**Mapping Experimental DMFE Measurements**

For the TEV measurements (Lalai et al., 2011), values of $w$ (fitness) were measured across eight hosts: the native host and seven additional hosts of varying phylogenetic relatedness to the native host. Selection coefficients in the novel hosts (corresponding to the novel environment) were calculated as $w_{\text{novel}}/w_{\text{native}} - 1$, where each $w$ was obtained by averaging across the six replicates performed in the study. The $\Delta$X174 study (Vale et al., 2012) directly reported selection coefficients in the single novel host tested. For each host shift, we calculated the extrapolated difference in log fitness $\sum_{i=1}^{n} \log((1 + s_i f_a - 1)\cdot x_i + 1)$ for those selection coefficients that had changed from neutral in the natural host to either positive or deleterious, with $T = 10$ and $N_0 = 0.5$. As per the $\Delta$X174 study (Vale et al., 2012), mutations were classified as nonneutral based on a t test when mean $s$ values significantly deviated from zero. Because both studies only measured selection coefficients for a sample of n sites in a genome of length $L$, we corrected for this by multiplying the result by $L/n$. Accordingly, for all but one of the novel hosts, the results showed a higher extrapolated fitness for a brittle host had it been competed against a robust one during a host shift (see Supplemental Information text part 3 for more details).

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, two figures, and two tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2014.07.011.

**AUTHOR CONTRIBUTIONS**

A.S. and R.A. designed the project. A.S. and S.B. performed all of the theoretical, simulation, and data analysis work. M.T.Y., C.W., and K.B. performed the laboratory experimental work. R.N. guided the theoretical framework. A.S. and R.A. wrote the manuscript. All authors commented on the manuscript.

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