

Open access • Posted Content • DOI:10.1101/078428

Quantifying the evolutionary potential and constraints of a drug-targeted viral protein — Source link [2]

Lei Dai, Yushen Du, Hangfei Qi, Nicholas C. Wu ...+2 more authors Institutions: University of California, Los Angeles Published on: 30 Sep 2016 - bioRxiv (Cold Spring Harbor Laboratory) Topics: Robustness (evolution), Mutation rate, Fitness landscape, Selection coefficient and Viral protein

Related papers:

- Adaptive potential of a drug-targeted viral protein as a function of positive selection
- · Adaptive potential of a drug-targeted viral protein as a function of environmental stress
- Assessing in vivo mutation frequencies and creating a high-resolution genome-wide map of fitness costs of
 Hepatitis C virus
- Evolutionary dynamics of viral escape under antibodies stress: A biophysical model.
- Costs and benefits of mutational robustness in RNA viruses.



Adaptive potential of a drug-targeted viral protein as a function of environmental stress

- 2
- Lei Dai^{1, 2*}, Yushen Du^{1*}, Hangfei Qi¹, Christian D. Huber², Nicholas C. Wu³, Ergang Wang¹, James O.
- 4 Lloyd-Smith², Ren Sun¹
- 5
- 6 ¹Department of Molecular and Medical Pharmacology
- ⁷ ²Department of Ecology and Evolutionary Biology, University of California Los Angeles, Los Angeles, United
- 8 States
- ⁹ ³Department of Integrative Structural and Computational Biology, The Scripps Research Institute, La Jolla,
- 10 United States
- 11
- ¹² *These authors contributed equally to this work.
- 13
- 14 Corresponding author:
- 15 Lei Dai, Ph.D.
- 16 Email: leidai@ucla.edu
- 17 Ren Sun, Ph.D.
- 18 Email: rsun@mednet.ucla.edu

19 Abstract

RNA viruses are notorious for their ability to evolve rapidly under selection in novel environments. It is known 20 that the high mutation rate of RNA viruses can generate huge genetic diversity to facilitate viral adaptation. 21 However, less attention has been paid to the underlying fitness landscape that represents the selection 22 forces on viral genomes. Here we systematically guantified the distribution of fitness effects (DFE) of about 23 1,600 single amino acid substitutions in the drug-targeted region of NS5A protein of Hepatitis C Virus (HCV). 24 We found that the majority of non-synonymous substitutions incur large fitness costs, suggesting that NS5A 25 protein is highly optimized in natural conditions. We characterized the adaptive potential of HCV by 26 subjecting the mutant viruses to selection by the antiviral drug Daclatasvir. Both the selection coefficient and 27 28 the number of beneficial mutations are found to increase with the level of environmental stress, which is modulated by the concentration of Daclatasvir. The changes in the spectrum of beneficial mutations in NS5A 29 protein can be explained by a pharmacodynamics model describing viral fitness as a function of drug 30 concentration. We test theoretical predictions regarding the distribution of beneficial fitness effects of mutations. 31 32 We also interpret the data in the context of Fisher's Geometric Model and find an increased distance to optimum as a function of environmental stress. Finally, we show that replication fitness of viruses is correlated with the 33 pattern of sequence conservation in nature and viral evolution is constrained by the need to maintain protein 34 stability. 35

37 Introduction

In our evolutionary battles with microbial pathogens, RNA viruses are among the most formidable foes. 38 HIV-1 and Hepatitis C Virus acquire drug resistance in patients under antiviral therapy. Influenza and Ebola 39 virus cross the species barrier to infect human hosts. Understanding the evolution of RNA viruses is 40 therefore of paramount importance for developing antivirals and vaccines and assessing the risk of future 41 emergence events (Goldberg et al. 2012; Domingo et al. 2012; Metcalf et al. 2015). Comprehensive 42 characterization of viral fitness landscapes, and the principles underpinning them, will provide us with a map 43 of evolutionary pathways accessible to viruses and guide our design of effective strategies to limit antiviral 44 resistance, immune escape and cross-species transmission (Turner and Elena 2000; Ke et al. 2015; Barton 45 46 et al. 2016).

Although the concept of fitness landscapes has been around for a long time (Wright 1932), we still know 47 little about their properties in real biological systems. Previous empirical studies of fitness landscapes have 48 been constrained by very limited sampling of sequence space. In a typical study, mutants are generated by 49 site-directed mutagenesis and assayed for growth rate individually. We and others have recently developed 50 a high-throughput technique, often referred to as "deep mutational scanning" or "quantitative high-resolution" 51 genetics", to profile the fitness effect of mutations by integrating deep sequencing with selection experiments 52 in vitro or in vivo (Hietpas et al. 2011; Wu et al. 2013; Thyagarajan and Bloom 2014; Qi et al. 2014; Fowler 53 and Fields 2014). This novel application of next generation sequencing has raised an exciting prospect of 54 large-scale fitness measurements (Olson et al. 2014; Puchta et al. 2015; Li et al. 2016; Wu et al. 2016) and a 55 revolution in our understanding of molecular evolution (He and Liu 2016). 56

The distribution of fitness effects (DFE) of mutations is a fundamental entity in genetics and reveals the local structure of a fitness landscape (Burch and Chao 2000; Eyre-Walker and Keightley 2007; Hietpas *et al.* 2011; Desai 2013; Jacquier *et al.* 2013; Bataillon and Bailey 2014; Chevereau *et al.* 2015; Bank *et al.* 2015). Deleterious mutations are usually abundant and impose severe constraints on the accessibility of fitness landscapes. In contrast, beneficial mutations are rare and provide the raw materials of adaptation. Quantifying the DFE of viruses is crucial for understanding how these pathogens evolve to acquire drug resistance and surmount other evolutionary challenges.

Most empirical studies of the DFE have been performed in a single, static environment (Eyre-Walker and Keightley 2007; Bataillon and Bailey 2014). A central challenge is to characterize the DFE, and its determinants, in fluctuating or heterogeneous environments where evolution typically occurs (e.g. fluctuating

67 drug concentrations or a gradient across space). Previous studies on yeast have investigated the change in DFE across different levels of temperature and salinity (Hietpas et al. 2013; Bank et al. 2014). For bacteria, 68 the fitness effects of mutations at different drug concentrations have been studied (Firnberg et al. 2014). 69 One recent study has demonstrated that drug concentration modulates the shape of the DFE and 70 determines the evolvability under new environments (Stiffler et al. 2015). In another study, the implications of 71 differing drug concentrations on the adaptive landscape have been examined in the context of resistance 72 evolution (Ogbunugafor et al. 2016). For viruses, the fitness effects of mutations have been measured 73 across different hosts (Lalić et al. 2011; Vale et al. 2012). The shape of DFE of viruses has been inferred 74 from experimentally passaged populations (Foll et al. 2014) and from patient data (Renzette et al. 2017). 75

In this study, we profile the DFE of ~1.600 single amino acid substitutions in a drug-targeted viral protein 76 by combining selection experiment of a mutant library and deep sequencing. We examine the changes in 77 DFE under varying levels of environmental stress by tuning the concentration of an antiviral drug. We test 78 theoretical predictions regarding the distribution of beneficial fitness effects of mutations (Orr 2003). We also 79 interpret the data in the context of Fisher's Geometric Model (Martin and Lenormand 2006b) and find an 80 increased distance to optimum as a function of environmental stress. Finally, we show that replication fitness 81 of viruses is correlated with the pattern of sequence conservation in nature and viral evolution is constrained 82 by the need to maintain protein stability. 83

84

85 **Results**

86 Profiling the fitness landscape of the drug-interacting domain of HCV NS5A protein

The system used in our study is Hepatitis C Virus (HCV), a positive sense single-stranded RNA virus with a genome of ~9.6 kb. HCV has been studied extensively in the past two decades in patients and in laboratory and provides an excellent model system to study viral evolution. We applied high-throughput assays to map the fitness effects of all single amino acid substitutions in domain IA (amino acid 18-103) of HCV NS5A protein (Methods). This domain is the target of several directly-acting antiviral drugs, including the potent HCV NS5A inhibitor Daclatasvir (DCV) (Gao *et al.* 2010).

To study the DFE of mutations of HCV NS5A protein, we conducted new selection experiments using a previously constructed saturation mutagenesis library of mutant viruses (Qi *et al.* 2014). Briefly, each codon in the mutated region was randomized to cover all possible single amino acid substitutions. We observed 2520 non-synonymous mutations in the plasmid library, as well as 105 synonymous mutations. After transfection to reconstitute mutant viruses, we performed selection in an HCV cell culture system

98 (Lindenbach *et al.* 2005; Wakita *et al.* 2005). The relative fitness of a mutant virus to the wild-type virus was 99 calculated based on the changes in frequency of the mutant virus and the wild-type virus after one round of 100 selection in cell culture (Supplementary Figure 1). In our selection experiment, we grew 5 small sub-libraries 101 (~500 mutants each) separately to reduce the noise in fitness measurements (Methods). The fitness data 102 reported in this study is highly correlated to an independent experiment using the same plasmid library 103 (Supplementary Figure 2) (Qi *et al.* 2014).

Our experiment provides a comprehensive profiling of the fitness effect of single amino acid substitutions 104 (1565 out of 1634 possible substitutions, after filtering out low frequency mutants in the plasmid library). We 105 grouped together non-synonymous mutations leading to the same amino acid substitution. As expected, the 106 fitness effects of synonymous mutations were nearly neutral, while most non-synonymous mutations were 107 deleterious (Figure 1). We found that the majority of single amino acid mutations had fitness costs and more 108 than half of them were found to be significantly deleterious, or "lethal" (Methods). The fraction of lethal 109 mutations (not shown explicitly in Figure 1) is 57.0% (932/1634) for single amino acid substitutions, 1.0% 110 (1/105) for synonymous mutations and 90.6% (77/85) for nonsense mutations. The low tolerance of 111 non-synonymous mutations in HCV NS5A, which is an essential protein for viral replication, is consistent with 112 previous small-scale mutagenesis studies of RNA viruses (Sanjuan et al. 2004). Our data support the view 113 that RNA viruses are very sensitive to the effect of deleterious mutations, possibly due to the compactness of 114 their genomes (Elena et al. 2006; Rihn et al. 2013). 115

Using the distribution of fitness effects of synonymous mutations as a benchmark for neutrality, we identified that only 2.3% (37/1634) of single amino acid mutations are beneficial (Methods). The estimated fraction of beneficial mutations is consistent with previous small-scale mutagenesis studies in viruses including bacteriophages, vesicular stomatitis virus, etc. (Sanjuan *et al.* 2004; Burch *et al.* 2007; Silander *et al.* 2007; Eyre-Walker and Keightley 2007). Our results indicate that HCV NS5A protein is under strong purifying selection, suggesting that viral proteins are highly optimized in their natural conditions.

122 Adaptive potential as a function of environmental stress

Beneficial mutations are the raw materials of protein adaptation (Eyre-Walker and Keightley 2007). In this study, we aimed to study the role of environmental stress in modulating the adaptive potential of drug-targeted viral proteins. In an independent study (Qi *et al.* 2014), the mutant library of HCV NS5A protein was selected under a single drug concentration ([DCV]=20 pM) to profile the effects of mutations on drug resistance. In this study, we selected the mutant library at 10, 40 and 100 pM of DCV. The drug concentrations were chosen based on in vitro IC₅₀ of wild type HCV virus (~20 pM) to represent different

levels of environmental stress (mild, intermediate and strong).

By tuning the concentration of DCV, we observed a change in the DFE (Supplementary Table 1&2), 130 particularly of beneficial mutations (Figure 2A). At higher drug concentrations, we observed an increase in 131 the median selection coefficient (Figure 2B) as well as the total number of beneficial mutations (Figure 2C, 132 Supplementary Table 3). We further tested whether the shape of this distribution changed under drug 133 selection. Previous empirical studies supported the hypothesis that the DFE of beneficial mutations is 134 exponential (Orr 1998, 2003, 2006; Imhof and Schlötterer 2001; Sanjuan et al. 2004; Rokyta et al. 2005; 135 Cowperthwaite et al. 2005; Kassen and Bataillon 2006; Burch et al. 2007; Carrasco et al. 2007; MacLean 136 and Buckling 2009; Peris et al. 2010; Bataillon et al. 2011). Following a maximum likelihood approach, we fit 137 the DFE of beneficial mutations to the Generalized Pareto Distribution (Supplementary Figure 3, Methods). 138 The fitted distribution is described by two parameters: a scale parameter (τ), and a shape parameter (κ) that 139 determines the behavior of the distribution's tail. Using a likelihood-ratio test (Beisel et al. 2007), we found 140 that our data are consistent with the null hypothesis that the DFE of beneficial mutations is exponential ($\kappa = 0$) 141 (Supplementary Table 4). 142

Furthermore, we used a maximum-likelihood approach to fit a displaced-gamma distribution to the DFE 143 to estimate the distance to the phenotypic optimum in Fisher's Geometric Model (FGM) (Martin and 144 Lenormand 2006b; Bank et al. 2014) (Supplementary Figure 4). The displaced-gamma distribution has the 145 shape of a negative gamma distribution, shifted by a parameter s_0 that indicates the distance of the initial 146 genotype (i.e. wild-type) to the optimum (Methods). Estimated distances to the optimum under different 147 conditions are summarized in Supplementary Table 5. In accordance with theoretical expectations, we found 148 that the distance to the optimum increased as the level of environmental stress increased (i.e. increasing 149 drug concentration). 150

151 The effects of mutations on drug resistance and replication fitness

Our results show that the adaptive potential of proteins is modulated by the strength of environmental stress. The changing spectra of beneficial mutations upon drug treatment can be explained by a pharmacodynamics model describing viral fitness as a function of drug concentration (i.e. phenotype-fitness mapping) (Figure 3A).

156
$$f = f_0 \frac{IC_{50}}{IC_{50} + [drug]}$$

where f_0 is the fitness without drug selection and IC_{50} is the half inhibitory concentration. The absolute fitness f decreases with drug concentration [drug]. In this paper, we define a drug-resistant mutant as any viral variant that is less inhibited than the wild type for some drug concentration, i.e. higher IC_{50} than wild-type (Rosenbloom *et al.* 2012).

Mutations that reduce a protein's binding affinity to drug molecules (i.e. less inhibited by the drug) may 161 come with a fitness cost (i.e. smaller f_0 than wild-type). Thus, a drug-resistant mutant that is deleterious in 162 the absence of drug may become beneficial under drug selection, leading to an increase in the number of 163 beneficial mutations. Moreover, the relative fitness of the drug-resistant mutant is expected to increase with 164 stronger selection pressure (Figure 3A, dashed line). The dose response curves were previously measured 165 for a set of mutants constructed by site-directed mutagenesis (Supplementary Figure 5) (Qi et al. 2014). 166 Indeed, we found that the relative fitness of validated drug-resistant mutants increased at higher drug 167 concentration (Figure 3B); in contrast, drug-sensitive mutants became less fit under drug selection. 168

169 Furthermore, we showed that the effects of mutations on drug resistance can be estimated from the fitness data and the results were generally consistent with estimates based on the dose response curves 170 (Supplementary Figure 6, Methods). Among all the non-lethal single amino acid substitutions profiled in our 171 HCV NS5A protein library, we found that roughly half of the mutations increased resistance to DCV (i.e. 172 improved new function) at the expense of replication fitness without drug (Figure 3C, Spearman's p=-0.13, 173 p=8.3×10⁻⁴). This group of resistance mutations (lower right section in Figure 3C) can become beneficial 174 when the positive selection imposed by the antiviral drug is strong, leading to an increase in the supply of 175 beneficial mutations at higher drug concentrations. We found no association between drug resistance and 176 fitness cost (Fisher's exact test, p=0.26), suggesting that there is no or very weak tradeoff in adaptation of 177 NS5A protein under the two different environments (i.e. with and without DCV selection). 178

179 Deleterious mutations as evolutionary constraints

While beneficial mutations open up adaptive pathways to genotypes with higher fitness, mutations that severely reduce replication fitness impose constraints on the evolution of viruses and are less likely to contribute to adaptation through gain of function. We analyzed sequence diversity of HCV sequences identified in patients from the HCV sequence database of Los Alamos National Lab (Methods). As expected, we found that amino acid sites with high fitness costs are often highly conserved (Figure 4A). The sequence diversity at each site was highly correlated to the replication fitness (the median fitness of observed mutants

at each site) measured in our study (Spearman's ρ =0.82, p=1.8×10⁻²¹).

To understand the biophysical basis of mutational effects (Liberles et al. 2012), we took advantage of 187 the available structural information (Supplementary Figure 7A). The crystal structure of NS5A domain I is 188 available excluding the amphipathic helix at N-terminus (Tellinghuisen et al. 2005; Love et al. 2009). We 189 found that the fitness effects of deleterious mutations at buried sites (i.e. with lower solvent accessibility) 190 were more pronounced than those at surface exposed sites (Figure 4B, Spearman's p=0.51, p=5.1×10⁻⁶; 191 Supplementary Figure 8A) (Ramsey et al. 2011). Moreover, we performed simulations of protein stability for 192 individual mutants using PyRosetta (Methods) (Das and Baker 2008; Chaudhury et al. 2010). A mutation 193 with $\Delta\Delta G>0$, i.e. shifting the free energy difference to favor the unfolded state, is expected to destabilize the 194 protein. We found that mutations that decreased protein stability led to reduced viral fitness (Figure 4C, 195 Spearman's $\rho = -0.57$, $p = 1.5 \times 10^{-7}$). For example, mutations at a stretch of highly conserved residues 196 (F88-N91) that run through the core of NS5A protein tended to destabilize the protein and significantly 197 reduced the viral fitness. Mutations that increase $\Delta\Delta G$ beyond a threshold (~5 Rosetta Energy Unit) were 198 mostly lethal (Supplementary Figure 8B). This is consistent with the threshold robustness model, which 199 predicts that proteins become unfolded after using up the stability margin (Bloom et al. 2005; Wylie and 200 Shakhnovich 2011; Olson et al. 2014). Also, we note that mutations can be deleterious because they impair 201 protein function rather than destabilize the protein, so the correlation between protein stability and fitness is 202 not expected to be perfect. The level of correlation between $\Delta\Delta G$ and fitness that we observed is similar to 203 previous studies in other proteins (Firnberg et al. 2014; Wu et al. 2015). 204

205

206 **Discussion**

Site-directed mutagenesis and experimental evolution are traditional approaches to examine the DFE 207 (Domingo-Calap et al. 2009; Sanjuán 2010; Levy et al. 2015; Visher et al. 2016). Both methods provide 208 pivotal insights into the shape of the DFE, yet with limitations. The site-directed mutagenesis approach 209 requires fitness assays for each individual mutant and can only provide a sparse sampling of mutations. In 210 experimental evolution, the sampling of sequence space via de novo mutations is biased towards 211 large-effect beneficial mutations, as they are more likely to fix in the population. In contrast, the deep 212 mutational scanning approach (Fowler and Fields 2014), which utilizes high-throughput sequencing to 213 simultaneously assay the fitness or phenotype of a library of mutants, allows for unbiased and large-scale 214 sampling of fitness landscapes and thus is ideal for studying the characteristics of empirical DFE. The 215 downside of this high-throughput approach is that the fitness measurements can be noisy, especially for 216

large mutant libraries (Matuszewski et al. 2016). In our experiment, we divided the mutant library into smaller 217 sub-libraries (~500 mutants) in selection experiments. We compared the data to an independent experiment 218 and found that the fitness estimates were largely reproducible (Supplementary Figure 2). We also showed 219 that the observed change in the DFE under different conditions was consistent with validation experiments 220 (Figure 3). Since this study is focused on the properties of the entire distribution of mutations rather than the 221 effects of specific mutations, our findings on the general patterns of DFE are robust to the errors in fitness 222 estimates. Our study quantified the fitness effects of single amino acid substitutions in the drug-targeted 223 region of an essential viral protein. In general, the empirical DFE of HCV NS5A was consistent with previous 224 findings that viral proteins were highly optimized in the natural condition and very sensitive to the effects of 225 deleterious mutations. 226

One crucial but often overlooked point is that DFE will vary as a function of the environment (Martin and 227 Lenormand 2006a; Lalić et al. 2011; Stiffler et al. 2015). In the study by Stiffler et al. 2015, the level of 228 environmental stress is controlled by ampicillin concentration. Because TEM-1's function is to degrade 229 ampicillin, deleterious mutations that impair the enzyme function ("loss-of-function") would become more 230 deleterious at higher dose of ampicillin. In our system, we do not expect the dose of Daclatasvir to alter the 231 strength of purifying selection on maintaining HCV NS5A protein's function in viral replication. Indeed, we do 232 not find much difference on the deleterious side of DFE across different environments. Instead, we have 233 observed significant changes on the beneficial side of DFE as a function of the drug dose. Because HCV 234 NS5A protein is not well adapted in the novel environment of Daclatasvir selection, the effect of drug 235 resistance mutations ("gain-of-function") becomes more beneficial at higher drug dose. Moreover, due 236 to the pleiotropic effect of mutations on drug resistance and replication fitness (Figure 3), there is an 237 238 increasing supply of beneficial mutations at higher drug dose.

Although different systems have distinct protein-drug interactions that lead to different resistance 239 profiles (Robinson et al. 2011), the results in our study provide a general framework to study DFE of 240 drug-targeted proteins. Future studies along this line will further our understanding of how proteins evolve 241 new functions under the constraint of maintaining their original function (Soskine and Tawfik 2010), as 242 exemplified in the evolution of resistance to directly-acting antiviral drugs (Rosenbloom et al. 2012). 243 244 Quantifying the characteristics of DFE of drug-targeted proteins under different environments (e.g. varying levels of environmental stress, or conflicting selection pressures), would allow us to assess repeatability in 245 the outcomes of viral evolution (de Visser and Krug 2014) and guide the design of therapies to minimize 246 247 drug resistance (Ogbunugafor et al. 2016).

248

249 Conclusions

Many viruses adapt rapidly to novel selection pressures, such as antiviral drugs. Understanding how 250 pathogens evolve under drug selection is critical for the success of antiviral therapy against human 251 pathogens. By combining deep sequencing with selection experiments in cell culture, we have guantified the 252 distribution of fitness effects of mutations in the drug-targeted domain of Hepatitis C Virus NS5A protein. Our 253 results indicate that the majority of single amino acid substitutions in NS5A protein incur large fitness costs. 254 By subjecting the mutant viruses to selection under an antiviral drug, we find that the adaptive potential of 255 viral proteins in a novel environment is modulated by the level of environmental stress. We test theoretical 256 predictions regarding the distribution of fitness effects of mutations. Finally, we show that viral evolution is 257 constrained by the need to maintain protein stability. 258

259

260 Materials and Methods

261 Mutagenesis

The mutant library of HCV NS5A protein domain IA (86 amino acids) was constructed using saturation mutagenesis as previously described (Qi *et al.* 2014). In brief, the entire region was divided into five sub-libraries each containing 17-18 amino acids (~500 mutants in each sub-library). NNK (N: A/T/C/G, K: T/G) was used to replace each amino acid. The oligos, each of which contains one random codon, were synthesized by IDT. The mutated region was ligated to the flanking constant regions, subcloned into the pFNX-HCV plasmid and then transformed into bacteria. The pFNX-HCV plasmid carrying the viral genome was synthesized in Dr. Ren Sun's lab based on the chimeric sequence of genotype 2a HCV strains J6/JFH1.

269 Cell culture

The human hepatoma cell line (Huh-7.5.1) was provided by Dr. Francis Chisari from the Scripps Research Institute, La Jolla. The cells were cultured in T-75 tissue culture flasks (Genesee Scientific) at 37 °C with 5% CO₂. The complete growth medium contained Dulbecco's Modified Eagle's Medium (Corning Cellgro), 10% heat-inactivated Fetal Bovine Serum (Omega Scientific), 10 mM HEPES (Life Technologies), 1x MEM Non-Essential Amino Acids Solution (Life Technologies) and 1x Penicillin-Streptomycin-Glutamine (Life Technologies).

276 Selection of mutant viruses

Plasmid mutant library was transcribed *in vitro* using T7 RiboMAX Express Large Scale RNA Production
System (Promega) and purified by PureLink RNA Mini Kit (Life Technologies). 10 µg of *in vitro* transcribed

RNA was used to transfect 4 million Huh-7.5.1 cells via electroporation by Bio-Rad Gene Pulser (246 V, 950 μ F). The supernatant was collected 6 days post transfection and virus titer was determined by immunofluorescence assay. The viruses collected after transfection were used to infect ~2 million Huh-7.5.1 cells with an MOI at around 0.1-0.2. The five sub-libraries were passaged for selection separately. For the three different levels of selection pressure, the growth media was supplemented with 10 pM, 40 pM and 100 pM HCV NS5A inhibitor Daclatasvir (BMS-790052), respectively. The supernatant was collected at 6 days post infection.

286 **Preparation of Illumina sequencing samples**

For each sample, viral RNA was extracted from 700 µl supernatant collected after transfection and after 287 selection using QIAamp Viral RNA Mini Kit (Qiagen). Extracted RNA was reverse transcribed into cDNA by 288 SuperScript III Reverse Transcriptase Kit (Life Technologies). The targeted region in NS5A (51-54 nt) was 289 PCR amplified using KOD Hot Start DNA polymerase (Novagen). The Eppendorf thermocycler was set as 290 following: 2 min at 95 °C; 25 to 35 three-step cycles of 20 s at 95 °C, 15 s at 52-56 °C (sub-library #1, 52 °C; 291 #2, 52 °C; #3, 52 °C; #4, 56 °C; #5, 54 °C) and 25s at 68 °C; 1 min at 68 °C. The number of PCR cycles are 292 chosen based on the copy number of cDNA templates as determined by gPCR (Bio-Rad). The PCR primers 293 are listed in Supplementary Table 6. The PCR products were purified using PureLink PCR Purification Kit 294 (Life Technologies) and prepared for Illumina HiSeg 2000 sequencing (paired-end 100 bp) following 295 5'-phosphorylation using T4 Polynucleotide Kinase (New England BioLabs), 3' dA-tailing using dA-tailing 296 module (New England BioLabs), and TA ligation of the adapter using T4 DNA ligase (Life Technologies). 297 Each sample was tagged with a unique 3-bp customized barcodes, which were part of the adapter sequence 298 and were sequenced as the first three nucleotides in both the forward and reverse reads (Wu et al. 2015) 299 300 (Supplementary Table 7).

301 Analysis of Illumina sequencing data

The sequencing data were parsed by SegIO function of BioPython. The reads from different samples were 302 de-multiplexed by the barcodes and mapped to the entire mutated region in NS5A by allowing at maximum 5 303 mismatches with the reference genome (Supplementary Data 3) (Qi et al. 2014). Since both forward and 304 305 reverse reads cover the whole amplicon, we used paired reads to correct for sequencing errors. A mutation was called only if it was observed in both reads and the quality score at the corresponding position was at 306 least 30. Sequencing reads containing mutations not supposed to appear in our single-codon mutant library 307 were excluded from downstream analysis. The sequencing depth for each sub-library is at least ~10⁵ and 308 309 two orders of magnitude higher than the library complexity.

310 Calculation of relative fitness

For each condition of selection experiments (i.e. different concentration of Daclatasvir [DCV]), the relative fitness (RF) of a mutant virus to the wild-type virus is calculated by the relative changes in frequency after selection,

314
$$RF_{mut}([DCV]) = \left(\frac{f_{mut}^{T=2}}{f_{mut}^{T=1}}\right) / \left(\frac{f_{WT}^{T=2}}{f_{WT}^{T=1}}\right)$$

where $f_{mut}^{T=round}$ and $f_{WT}^{T=round}$ is the frequency of the mutant virus and the wild-type virus at round 1 (after 315 transfection) or round 2 (after infection). The fitness of wild-type virus is normalized to 1. The fitness values 316 estimated from one round (round 1 to round 2) have been shown to be highly consistent to estimated based 317 round 0 to round 1 (Supplementary Figure 2), and estimates from multiple rounds of selection (Qi et al. 318 2014). A mutant was labeled as "missing" if the mutant's frequency in the plasmid library was less than 319 0.0005 (RF=NaN, see Supplementary Data 1 and 2). A mutant was labeled as "lethal" if the mutant's 320 frequency after transfection was less than 0.0005, or its frequency after infection was 0 (RF=0) (Qi et al. 321 2014). 322

323 The selection coefficient is defined in the context of discrete generations (Chevin 2010)

$$324 \qquad s_{mut} = \log(RF_{mut})$$

The threshold for beneficial mutations is chosen as $2\sigma_{silent}$, where σ_{silent} is the standard deviation of the selection coefficients of synonymous mutations (Figure 1). The fitness effects of non-synonymous mutations leading to the same amino acid substitution were averaged to estimate the fitness effect of the given single amino acid substitution.

329 Fitting the distribution of fitness effects of beneficial mutations

The distribution of selection coefficients of beneficial mutations were fitted to a Generalized Pareto Distribution following a maximum likelihood approach (Beisel *et al.* 2007),

$$F(x|\kappa,\tau) = \begin{cases} 1 - (1 + \frac{\kappa}{\tau}x)^{-\frac{1}{\kappa}}, x \ge 0, & \text{if } \kappa > 0 \\ 1 - (1 + \frac{\kappa}{\tau}x)^{-\frac{1}{\kappa}}, 0 \le x < -\frac{\tau}{\kappa}, & \text{if } \kappa < 0 \\ 1 - e^{-\frac{\kappa}{\tau}}, x \ge 0, & \text{if } \kappa = 0 \end{cases}$$
(Weibull)
(Gumbel)

Only mutations with selection coefficients higher than the beneficial threshold $2\sigma_{silent}$ were included in the distribution of beneficial mutations. The selection coefficients were normalized to the beneficial threshold. The shape parameter κ determines the tail behavior of the distribution, which can be divided into three domains of attraction: Gumbel domain (exponential tail, $\kappa = 0$), Weibull domain (truncated tail, $\kappa < 0$) and Fréchet domain (heavy tail, $\kappa > 0$). For each selection condition, a likelihood ratio test is performed to evaluate whether the null hypothesis $\kappa = 0$ (exponential distribution) can be rejected.

339 Fitting the distribution of fitness effects to Fisher's Geometrical model

Fisher's Geometrical Model predicts that the distribution of fitness effects of mutations is distributed 340 according to a negative displaced gamma distribution (Martin and Lenormand 2006a, Bank et al. 2014). This 341 distribution has a shape parameter (α), a scale parameter (β), and a displacement parameter (s_0). We 342 assume that selection coefficients are measured with a normally distributed measurement error with 343 standard deviation σ_{silent} . Thus, the observed distribution of selection coefficients is modeled as the sum of a 344 gamma and normally distributed random variable. We use the NormalGamma package in R to numerically 345 compute the normal-gamma density function (Plancade et al. 2012). Maximum likelihood estimates of the 346 parameters of the negative displaced gamma distribution are obtained with L-BFGS-B optimization 347 implemented in the R function optim. 348

349 Inferring drug resistance from fitness data

We can quantify the drug resistance of each mutant in the library by computing its fold change in relative fitness,

352
$$W([DCV]) = \frac{RF_{mut}([DCV])}{RF_{mut}}$$

Here RF_{mut} is the relative fitness of a mutant under the natural condition (i.e. no drug). W is the fold change in relative fitness and represents the level of drug resistance relative to the wild type. W > 1 indicates drug resistance, and W < 1 indicates drug sensitivity.

This empirical measure of drug resistance can be directly linked to a simple pharmacodynamics model (Rosenbloom *et al.* 2012), where the viral replicative fitness is modeled as a function of drug dose,

358
$$W_{predict}([DCV]) = \left(\frac{IC_{mut}}{[DCV] + IC_{mut}}\right) / \left(\frac{IC_{wt}}{[DCV] + IC_{wt}}\right)$$

Here IC denotes the half-inhibitory concentration. The Hill coefficient describing the sigmoidal shape of the dose response curve is fixed to 1, as used in fitting the dose response curves of wild-type virus and validated

mutant viruses (Supplementary Figure 5). The drug resistance score *W* inferred from fitness data is consistent with the drug resistance score $W_{predict}$ predicted from dose response curves of validated mutants (Supplementary Figure 6).

364 Calculation of relative solvent accessibility

DSSP (http://www.cmbi.ru.nl/dssp.html) was used to compute the Solvent Accessible Surface Area (SASA)
(Kabsch and Sander 1983) from the HCV NS5A protein structure (PDB: 3FQM) (Love *et al.* 2009). SASA
was then normalized to Relative Solvent Accessibility (RSA) using the empirical scale reported in (Tien *et al.*2013).

369 **Predictions of protein stability**

 $\Delta\Delta G$ (in Rosetta Energy Unit) of HCV NS5A mutants was predicted by PyRosetta (version: "monolith.ubuntu.release-104") as the difference in scores between the monomer structure of mutants (single amino acid mutations from site 32 to 103) and the reference (PDB: 3FQM). The score is designed to capture the change in thermodynamic stability caused by the mutation ($\Delta\Delta G$) (Das and Baker 2008). The reference sequence of NS5A in the PDB file (PDB: 3FQM) is different from the WT sequence in our experiment by 20 amino acid substitutions. Thus instead of directly comparing $\Delta\Delta G$ to fitness effects of individual mutations, we used the median $\Delta\Delta G$ caused by amino acid substitutions at each site.

The PDB file of NS5A dimer was cleaned and trimmed to a monomer (chain A). Next, all side chains were repacked (sampling from the 2010 Dunbrack rotamer library (Shapovalov and Dunbrack 2011)) and minimized for the reference structure using the talaris2014 scoring function. After an amino acid mutation was introduced, the mutated residue was repacked, followed by quasi-Newton minimization of the backbone and all side chains (algorithm: "lbfgs_armijo_nonmonotone"). This procedure was performed 50 times, and the predicted ΔG of a mutant structure is the average of the three lowest scoring structures.

We note that predictions based on NS5A monomer structure were only meant to provide a crude profile of how mutations at each site may impact protein stability. Potential structural constraints at the dimer interface have been ignored, which is further complicated by the observations of two different NS5A dimer structures (Tellinghuisen *et al.* 2005; Love *et al.* 2009).

387 Diversity of HCV sequences identified in patients

Aligned nucleotide sequences of HCV NS5A protein were downloaded from Los Alamos National Lab database (Kuiken *et al.* 2005) (all HCV genotypes, ~2600 sequences total) and clipped to the region of

interest (amino acid 18-103 of NS5A). Sequences that caused gaps in the alignment of H77 reference genome were manually removed. After translation to amino acid sequences, sequences with ambiguous amino acids were removed (~2300 amino acid sequences after filtering). The sequence diversity at each amino acid site was guantified by Shannon entropy.

394 Data and reagent availability

All research materials are available upon request. Raw sequencing data have been submitted to the NIH Short Read Archive (SRA) under accession number: BioProject PRJNA395730. All scripts have been deposited to https://github.com/leidai-evolution/DFE-HCV.

398 Ethics Statement

- The use of human cell lines and infectious agents in this paper is approved by Institutional Biosafety Committee at University of California, Los Angeles (IBC #40.10.2-f).
- 401

402 Acknowledgements

We thank Daniel Weinreich and two anonymous reviewers for constructive comments on the manuscript.
L.D. was supported by HHMI Postdoctoral Fellowship from Jane Coffin Childs Memorial Fund for Medical
Research. N.C.W. was supported by Croucher Foundation Fellowship. R.S. was supported by NSFC
81172314, NIH DE023591 and NIH CA177322.

407

408 Author contributions

L.D., Y.D., H.Q. and R.S. designed the experiments. L.D., H.Q. and Y.D. performed the experiments. L.D.
and Y.D. analyzed the experimental data. L.D., E.W. and Y.D. performed the bioinformatics analyses. C.D.H.
and L.D. performed the analysis on FGM. L.D. wrote the first draft of the manuscript, with revisions from Y.D.,
H.Q., N.C.W., J.O.L-S., and R.S.. All authors discussed the results and commented on the manuscript.

413 **References**

- Bank C., Hietpas R. T., Wong A., Bolon D. N., Jensen J. D., 2014 A Bayesian MCMC Approach to Assess
- the Complete Distribution of Fitness Effects of New Mutations: Uncovering the Potential for Adaptive
 Walks in Challenging Environments. Genetics 196.
- Bank C., Hietpas R. T., Jensen J. D., Bolon D. N. A., 2015 A systematic survey of an intragenic epistatic
 landscape. Mol. Biol. Evol. 32: 229–38.
- Barton J. P., Goonetilleke N., Butler T. C., Walker B. D., McMichael A. J., et al., 2016 Relative rate and
- location of intra-host HIV evolution to evade cellular immunity are predictable. Nat. Commun. 7: 11660.
- Bataillon T., Zhang T., Kassen R., 2011 Cost of adaptation and fitness effects of beneficial mutations in
 Pseudomonas fluorescens. Genetics 189: 939–949.
- Bataillon T., Bailey S., 2014 Effects of new mutations on fitness: insights from models and data. Ann. N. Y.
 Acad. Sci. 1320: 76–92.
- Beisel C. J., Rokyta D. R., Wichman H. A., Joyce P., 2007 Testing the extreme value domain of attraction for
 distributions of beneficial fitness effects. Genetics 176: 2441–9.
- Bloom J. D., Silberg J. J., Wilke C. O., Drummond D. A., Adami C., *et al.*, 2005 Thermodynamic prediction of protein neutrality. Proc. Natl. Acad. Sci. U. S. A. 102: 606–11.
- Burch C. L., Chao L., 2000 Evolvability of an RNA virus is determined by its mutational neighbourhood.
 Nature 406: 625–8.
- Burch C., Guyader S., Samarov D., Shen H., 2007 Experimental estimate of the abundance and effects of
 nearly neutral mutations in the RNA virus φ6. Genetics 476: 467–476.
- Carrasco P., Iglesia F. de Ia, Elena S., 2007 Distribution of fitness and virulence effects caused by
 single-nucleotide substitutions in Tobacco etch virus. J. Virol. 81: 12979–12984.
- Chaudhury S., Lyskov S., Gray J. J., 2010 PyRosetta: a script-based interface for implementing molecular
 modeling algorithms using Rosetta. Bioinformatics 26: 689–91.
- Chevereau G., Dravecká M., Batur T., Guvenek A., Ayhan D. H., *et al.*, 2015 Quantifying the Determinants of
 Evolutionary Dynamics Leading to Drug Resistance. PLoS Biol. 13: e1002299.
- 439 Chevin L.-M., 2010 On measuring selection in experimental evolution. Biol. Lett.
- Cowperthwaite M. C., Bull J. J., Meyers L. A., 2005 Distributions of beneficial fitness effects in RNA.
 Genetics 170: 1449–57.
- 442 Das R., Baker D., 2008 Macromolecular Modeling with Rosetta. Annu. Rev. Biochem. 77: 363–382.
- 443 Desai M. M., 2013 Statistical questions in experimental evolution. J. Stat. Mech. Theory Exp. 2013: P01003.

- 444 Domingo E., Sheldon J., Perales C., 2012 Viral guasispecies evolution. Microbiol. Mol. Biol. Rev. 76: 159–
- 445 **216**.
- Domingo-Calap P., Cuevas J. M., Sanjuán R., 2009 The Fitness Effects of Random Mutations in
 Single-Stranded DNA and RNA Bacteriophages (DJ Begun, Ed.). PLoS Genet. 5: e1000742.
- Elena S. F., Carrasco P., Daròs J.-A., Sanjuán R., 2006 Mechanisms of genetic robustness in RNA viruses.
 EMBO Rep. 7: 168–73.
- 450 Eyre-Walker A., Keightley P. D., 2007 The distribution of fitness effects of new mutations. Nat. Rev. Genet. 8:
- 451 610–8.
- 452 Firnberg E., Labonte J. W., Gray J. J., Ostermeier M., 2014 A comprehensive, high-resolution map of a 453 gene's fitness landscape. Mol. Biol. Evol. 31: 1581–92.
- Foll M., Poh Y.-P., Renzette N., Ferrer-Admetlla A., Bank C., *et al.*, 2014 Influenza virus drug resistance: a
 time-sampled population genetics perspective. PLoS Genet. 10: e1004185.
- Fowler D. M., Fields S., 2014 Deep mutational scanning: a new style of protein science. Nat. Methods 11:
 801–807.
- Gao M., Nettles R. E., Belema M., Snyder L. B., Nguyen V. N., *et al.*, 2010 Chemical genetics strategy identifies an HCV NS5A inhibitor with a potent clinical effect. Nature 465: 96–100.
- Goldberg D. E., Siliciano R. F., Jacobs W. R., 2012 Outwitting evolution: fighting drug-resistant TB, malaria,
 and HIV. Cell 148: 1271–83.
- He X., Liu L., 2016 Toward a prospective molecular evolution. Science 352: 769–70.
- Hietpas R. T., Jensen J. D., Bolon D. N. A., 2011 Experimental illumination of a fitness landscape. Proc. Natl.
 Acad. Sci. U. S. A. 108: 7896–901.
- Hietpas R. T., Bank C., Jensen J. D., Bolon D. N. A., 2013 SHIFTING FITNESS LANDSCAPES IN
 RESPONSE TO ALTERED ENVIRONMENTS. Evolution (N. Y). 67: 3512–3522.
- Imhof M., Schlötterer C., 2001 Fitness effects of advantageous mutations in evolving Escherichia coli
 populations. Proc. Natl. Acad. Sci. U. S. A. 98: 1113–1117.
- Jacquier H., Birgy A., Nagard H. Le, Mechulam Y., Schmitt E., *et al.*, 2013 Capturing the mutational landscape of the beta-lactamase TEM-1. Proc. Natl. Acad. Sci. U. S. A. 110: 13067–72.
- Kabsch W., Sander C., 1983 Dictionary of protein secondary structure: Pattern recognition of
 hydrogen-bonded and geometrical features. Biopolymers 22: 2577–2637.
- Kassen R., Bataillon T., 2006 Distribution of fitness effects among beneficial mutations before selection in
 experimental populations of bacteria. Nat. Genet. 38: 484–8.

- Ke R., Loverdo C., Qi H., Sun R., Lloyd-Smith J. O., 2015 Rational Design and Adaptive Management of
- 476 Combination Therapies for Hepatitis C Virus Infection. PLoS Comput. Biol. 11: e1004040.
- Kuiken C., Yusim K., Boykin L., Richardson R., 2005 The Los Alamos hepatitis C sequence database.
 Bioinformatics 21: 379–384.
- Lalić J., Cuevas J. M., Elena S. F., 2011 Effect of host species on the distribution of mutational fitness effects
 for an RNA virus. PLoS Genet. 7: e1002378.
- Levy S. F., Blundell J. R., Venkataram S., Petrov D. A., Fisher D. S., *et al.*, 2015 Quantitative evolutionary
 dynamics using high-resolution lineage tracking. Nature 519: 181–6.
- Li C., Qian W., Maclean C. J., Zhang J., 2016 The fitness landscape of a tRNA gene. Science 352: 837–840.
- Liberles D. A., Teichmann S. A., Bahar I., Bastolla U., Bloom J., *et al.*, 2012 The interface of protein structure,
 protein biophysics, and molecular evolution. Protein Sci. 21: 769–785.
- Lindenbach B. D., Evans M. J., Syder A. J., Wölk B., Tellinghuisen T. L., *et al.*, 2005 Complete Replication of
 Hepatitis C Virus in Cell Culture. Science 309: 623–626.
- Love R. A., Brodsky O., Hickey M. J., Wells P. A., Cronin C. N., 2009 Crystal structure of a novel dimeric form of NS5A domain I protein from hepatitis C virus. J. Virol. 83: 4395–403.
- MacLean R. C., Buckling A., 2009 The distribution of fitness effects of beneficial mutations in Pseudomonas
 aeruginosa. PLoS Genet. 5: e1000406.
- Martin G., Lenormand T., 2006a The fitness effect of mutations across environments: a survey in light of
 fitness landscape models. Evolution 60: 2413–2427.
- 494 Martin G., Lenormand T., 2006b A general multivariate extension of Fisher's geometrical model and the 495 distribution of mutation fitness effects across species. Evolution 60: 893–907.
- Matuszewski S., Hildebrandt M. E., Ghenu A.-H., Jensen J. D., Bank C., 2016 A Statistical Guide to the
 Design of Deep Mutational Scanning Experiments. Genetics.
- Metcalf C. J. E., Birger R. B., Funk S., Kouyos R. D., Lloyd-Smith J. O., *et al.*, 2015 Five challenges in
 evolution and infectious diseases. Epidemics 10: 40–44.
- Ogbunugafor C. B., Wylie C. S., Diakite I., Weinreich D. M., Hartl D. L., 2016 Adaptive Landscape by
 Environment Interactions Dictate Evolutionary Dynamics in Models of Drug Resistance. PLoS Comput.
 Biol. 12: e1004710.
- 503 Olson C. A., Wu N. C., Sun R., 2014 A comprehensive biophysical description of pairwise epistasis 504 throughout an entire protein domain. Curr. Biol. 24: 2643–51.
- 505 Orr H. A., 1998 The population genetics of adaptation: the distribution of factors fixed during adaptive

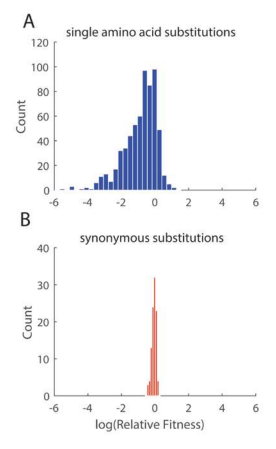
- 506 evolution. Evolution (N. Y). 52: 935–949.
- 507 Orr H., 2003 The distribution of fitness effects among beneficial mutations. Genetics 1526: 1519–1526.
- 508 Orr H., 2006 The distribution of fitness effects among beneficial mutations in Fisher's geometric model of 509 adaptation. J. Theor. Biol. 238: 279–285.
- Peris J. B., Davis P., Cuevas J. M., Nebot M. R., Sanjuán R., 2010 Distribution of fitness effects caused by
 single-nucleotide substitutions in bacteriophage f1. Genetics 185: 603–9.
- Plancade S., Rozenholc Y., Lund E., 2012 Generalization of the normal-exponential model: exploration of a
 more accurate parametrisation for the signal distribution on Illumina BeadArrays. BMC Bioinformatics
 13: 329.
- Puchta O., Cseke B., Czaja H., Tollervey D., Sanguinetti G., *et al.*, 2015 Network of epistatic interactions
 within a yeast snoRNA. Science 352: 840–844.
- Qi H., Olson C. A., Wu N. C., Ke R., Loverdo C., *et al.*, 2014 A quantitative high-resolution genetic profile
 rapidly identifies sequence determinants of hepatitis C viral fitness and drug sensitivity. PLoS Pathog.
 10: e1004064.
- Ramsey D. C., Scherrer M. P., Zhou T., Wilke C. O., 2011 The Relationship Between Relative Solvent
 Accessibility and Evolutionary Rate in Protein Evolution. Genetics 188: 479–488.
- Renzette N., Pfeifer S. P., Matuszewski S., Kowalik T. F., Jensen J. D., 2017 On the Analysis of Intrahost and
- Interhost Viral Populations: Human Cytomegalovirus as a Case Study of Pitfalls and Expectations. J.
 Virol. 91: e01976-16.
- Rihn S. J., Wilson S. J., Loman N. J., Alim M., Bakker S. E., *et al.*, 2013 Extreme genetic fragility of the HIV-1
 capsid. PLoS Pathog. 9: e1003461.
- Robinson M., Tian Y., Delaney W. E., Greenstein A. E., 2011 Preexisting drug-resistance mutations reveal
 unique barriers to resistance for distinct antivirals. Proc. Natl. Acad. Sci. U. S. A. 108: 10290–5.
- Rokyta D., Joyce P., Caudle S., Wichman H., 2005 An empirical test of the mutational landscape model of
 adaptation using a single-stranded DNA virus. Nat. Genet. 37: 441–444.
- Rosenbloom D. I. S., Hill A. L., Rabi S. A., Siliciano R. F., Nowak M. A., 2012 Antiretroviral dynamics
 determines HIV evolution and predicts therapy outcome. Nat. Med. 18: 1378–85.
- Sanjuan R., Moya A., Elena S. F., 2004 The distribution of fitness effects caused by single-nucleotide
 substitutions in an RNA virus. Proc. Natl. Acad. Sci. 101: 8396–8401.
- Sanjuán R., 2010 Mutational fitness effects in RNA and single-stranded DNA viruses: common patterns
 revealed by site-directed mutagenesis studies. Philos. Trans. R. Soc. Lond. B. Biol. Sci. 365: 1975–82.

- 537 Shapovalov M. V, Dunbrack R. L., 2011 A smoothed backbone-dependent rotamer library for proteins
- derived from adaptive kernel density estimates and regressions. Structure 19: 844–58.
- Silander O. K., Tenaillon O., Chao L., 2007 Understanding the Evolutionary Fate of Finite Populations: The
 Dynamics of Mutational Effects. PLoS Biol. 5: e94.
- Soskine M., Tawfik D. S., 2010 Mutational effects and the evolution of new protein functions. Nat. Rev. Genet.
 11: 572–82.
- 543 Stiffler M. A., Hekstra D. R., Ranganathan R., 2015 Evolvability as a Function of Purifying Selection in 544 TEM-1 β-Lactamase. Cell 160: 882–892.
- Tellinghuisen T. L., Marcotrigiano J., Rice C. M., 2005 Structure of the zinc-binding domain of an essential component of the hepatitis C virus replicase. Nature 435: 374–9.
- Thyagarajan B., Bloom J. D., 2014 The inherent mutational tolerance and antigenic evolvability of influenza
 hemagglutinin. Elife 3: e03300.
- Tien M. Z., Meyer A. G., Sydykova D. K., Spielman S. J., Wilke C. O., 2013 Maximum allowed solvent accessibilites of residues in proteins. PLoS One 8: e80635.
- 551 Turner P. E., Elena S. F., 2000 Cost of Host Radiation in an RNA Virus. Genetics 156: 1465–1470.
- Vale P. F., Choisy M., Froissart R., Sanjuán R., Gandon S., 2012 THE DISTRIBUTION OF MUTATIONAL
- 553 FITNESS EFFECTS OF PHAGE φX174 ON DIFFERENT HOSTS. Evolution (N. Y). 66: 3495–3507.
- Visher E., Whitefield S. E., McCrone J. T., Fitzsimmons W., Lauring A. S., 2016 The Mutational Robustness
 of Influenza A Virus. PLOS Pathog. 12: e1005856.
- Visser J. A. G. M. de, Krug J., 2014 Empirical fitness landscapes and the predictability of evolution. Nat. Rev.
 Genet. 15: 480–90.
- 558 Wakita T., Pietschmann T., Kato T., Date T., Miyamoto M., *et al.*, 2005 Production of infectious hepatitis C 559 virus in tissue culture from a cloned viral genome. Nat. Med. 11: 791–796.
- 560 Wright S., 1932 The Roles of Mutation, Inbreeding, Crossbreeding and Selection in Evolution. 1: 356–366.

Wu N. C., Young A. P., Dandekar S., Wijersuriya H., Al-Mawsawi L. Q., *et al.*, 2013 Systematic identification

- of H274Y compensatory mutations in influenza A virus neuraminidase by high-throughput screening. J.
 Virol. 87: 1193–9.
- Wu N. C., Olson C. A., Du Y., Le S., Tran K., *et al.*, 2015 Functional Constraint Profiling of a Viral Protein
 Reveals Discordance of Evolutionary Conservation and Functionality. PLOS Genet. 11: e1005310.
- 566 Wu N. C., Dai L., Olson C. A., Lloyd-Smith J. O., Sun R., 2016 Adaptation in protein fitness landscapes is
- 567 facilitated by indirect paths. Elife 5: e16965.

- 568 Wylie C. S., Shakhnovich E. I., 2011 A biophysical protein folding model accounts for most mutational fitness
- effects in viruses. Proc. Natl. Acad. Sci. 108: 9916–9921.
- 570



571

572 Figure 1. Distribution of fitness effects (DFE) of single amino acid substitutions in domain IA of HCV

573 **NS5A protein without drug selection.** DFE of single amino acid substitutions (A) and synonymous 574 substitutions (B). Lethal mutations are not shown in the histogram.

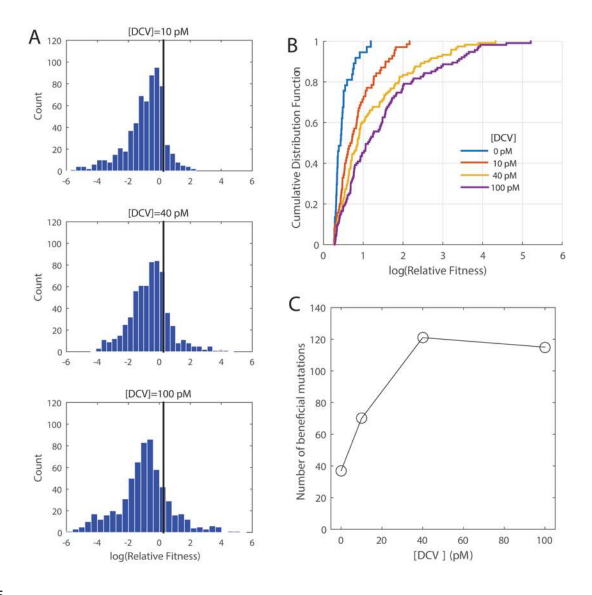




Figure 2. The spectrum of beneficial mutations changes under increasing environmental stress imposed by the antiviral drug Daclatasvir. (A) DFE of single amino acid substitutions in domain IA of HCV NS5A protein under increasing environmental stress by Daclatasvir. The black line indicates the threshold used for classifying beneficial mutations (Methods). (B) The cumulative distribution function of the fitness effect of beneficial mutations. (C) The number of beneficial mutations as a function of environmental stress imposed by Daclatasvir.

bioRxiv preprint doi: https://doi.org/10.1101/078428; this version posted January 9, 2018. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

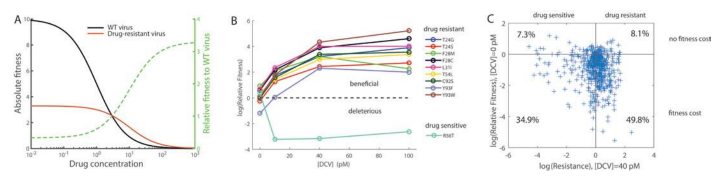


Figure 3. The adaptive potential under drug selection is determined by the effects of mutations on replication fitness and drug resistance. (A) Hypothetical dose response curves of the wild-type virus and a drug-resistant mutant virus. The absolute fitness f decreases with drug concentration [drug]

582

586
$$f = f_0 \frac{IC_{50}}{IC_{50} + [drug]}$$
, where f_0 is the fitness without drug selection and IC_{50} is the half inhibitory

concentration. Compared to the wild-type virus, the hypothetical drug-resistant mutant carries a fitness cost (smaller f_0) but is less sensitive to drug inhibition (larger IC_{50}). Relative fitness of the drug-resistant mutant is expected to increase with drug concentration. (B) Relative fitness of validated drug-resistant and drug-sensitive mutants (Supplementary Figure 5) as a function of [DCV]. (C) The effects of mutations on replication fitness (i.e. fitness without drug) and drug resistance score W at [DCV]=40 pM (Methods).

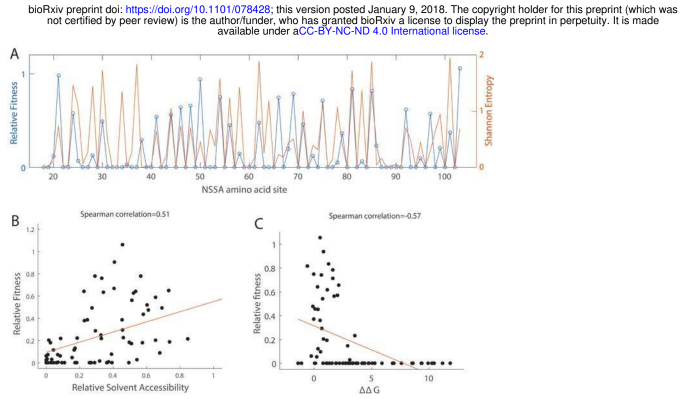


Figure 4. Mutations with deleterious fitness effects reveal constraints of protein evolution. (A) The 593 pattern of sequence conservation observed in patient sequences is highly correlated to the replication 594 fitness measured in cell culture. (B) Mutations at amino acid sites with lower solvent accessibility tend to 595 incur larger fitness costs. (C) Mutations at amino acid sites with larger effects on destabilizing protein 596 597 stability ($\Delta\Delta$ G>0) tend to reduce the viral replication fitness. Changes in folding free energy $\Delta\Delta$ G (Rosetta Energy Unit) of NS5A monomer were predicted by PyRosetta. The median ΔΔG at each amino acid site is 598 shown. In (A-C), the median fitness of observed mutants at each amino acid site is shown. In (B) and (C), 599 red lines represent the fits by linear regression and are only used to guide the eye. 600