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Competition-colonization dynamics in an RNA virus

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During replication, RNA viruses rapidly generate diverse mutant progeny which differ in their ability to kill host cells. We report that the progeny of a single RNA viral genome diversified during hundreds of passages in cell culture and self-organized into two genetically distinct subpopulations that exhibited the competition-colonization dynamics previously recognized in many classical ecological systems. Viral colonizers alone were more efficient in killing cells than competitors in culture. In cells coinfecting with both competitors and colonizers, viral interference resulted in reduced cell killing, and competitors replaced colonizers. Mathematical modeling of this coinfection dynamics predicted selection to be density dependent, which was confirmed experimentally. Thus, as is known for other ecological systems, biodiversity and even cell killing of virus populations can be shaped by a tradeoff between competition and colonization. Our results suggest a model for the evolution of virulence in viruses based on internal interactions within mutant spectra of viral quasispecies.

evolution | quasispecies | self-organization | virulence

RNA viruses replicate as complex mutant distributions termed viral quasispecies (1–4). Viral clones diversify upon replication because of mutation rates in the range of 10^{-3} to 10^{-5} substitutions per nucleotide copied, due to absence (or low efficiency) of proof-reading-repair functions in viral RNA dependent RNA polymerases (5–7). Little is known of the evolution of virulence when a viral clone diversifies to produce a broad mutant distribution. Large population passages result in fitness gain (8, 9). However, fitness and virulence are not necessarily correlated traits, as shown with clones of the important picornaviral pathogen foot-and-mouth disease virus (FMDV) using cell killing as a marker for virulence (10), or using Tobacco Etch virus and reduction in seed production in planta as a marker for virulence (11).

FMDV has been used as a model system to study quasispecies dynamics (12). A biological clone of FMDV of serotype C termed C-S8c1 was extensively passaged in BHK-21, resulting in genetic diversification and fitness increase (13, 14). At passage 143, a monoclonal antibody (MAb) SD6-resistant mutant termed MARLS was isolated (14). This mutant differed in 32 mutations from its parental C-S8c1 virus and displayed high fitness and a 10^3 -fold greater ability to kill cells than C-S8c1 (10, 15). In the present study we have examined the mutant composition of the population that resulted after 240 serial passages of FMDV C-S8c1 (Fig. 1). The clonal population diversified into two genotypically and phenotypically distinct classes of FMDV genomes that correspond to competitors and colonizers, as previously recognized in ecological systems (16, 17). We provide evidence that cell killing is the result of a compromise between the two phenotypes that coexist in the populations and develop a mathematical model for the competition-colonization dynamics.

Results

High Fitness FMDV Clones Were Suppressed by the Dominant Population. A biological clone of FMDV was passaged in BHK-21 cells at a multiplicity of infection (MOI) of 1–20 plaque-forming-units per cell (PFU/cell) for a total of 240 passages (see *SI Appendix*). The

populations at passages 200, 219, 225 and 240 were subjected to three low-MOI passages (0.006–0.02 PFU/cell) to obtain the corresponding populations termed 200p3d, 219p3d, 225p3d, and 240p3d, respectively, as detailed in *Materials and Methods*. The four populations and their 3d derivatives were analyzed phylogenetically (Fig. 1A). The results show that the sequence of the initially dominant populations is similar to the consensus sequence of passage 200 (p200), although the sequence of the p3d derivatives is similar to MARLS, a MAb-escape mutant isolated from the same viral lineage (14), and previously characterized as a high-fitness and high-virulence variant (10, 15). To ascertain that the viral population included individual infectious particles that were either p200 or MARLS, a total of 16 biological clones derived from the populations at passages 219, 225, and 240 were isolated and their genomic RNA analyzed by nucleotide sequencing. The phylogenetic analysis showed that indeed two subclasses of genomes were present and that they segregated into p200 and MARLS sequences (Fig. 1B). These phylogenetic comparisons suggest that MARLS clones might be suppressed by p200 variants replicating in the same population and that the MARLS variants could become dominant after the three low-MOI passages (Fig. 1). Therefore, limiting coinfection of cells by MARLS and p200 genomes permitted the former to attain dominance in the viral population.

Competitor and Colonizer FMDV Subpopulations. To further analyze the interaction between p200 and MARLS, the two viral subpopulations were characterized phenotypically. The MARLS population and its derived clones killed BHK-21 cells faster than the p200 population and its derived clones in a cell killing assay that compared the time needed to kill 10^4 BHK-21 cells as a function of the initial number of PFU (10) (Fig. 2A and B). To further evaluate this difference, viruses were tested in cell killing-interference assays (see *Materials and Methods*). Cells coinfecting by fast-killing MARLS and slow-killing p200 viruses died at a similar rate as cells infected only by slow p200 viruses, either using whole populations or individual clones (Fig. 2C–D). A possible dose effect was excluded because no alteration of the cell killing time was observed when cells were infected by twice the amount of fast viruses. The delay of cell killing could be attributed specifically to slow (p200) viruses because cells infected by two fast (MARLS) viruses experienced no delay (Fig. 2E). A tradeoff between higher cell killing and lower progeny production was also rejected because MARLS clone 240c2 production was statistically indistinguishable from p200 clones

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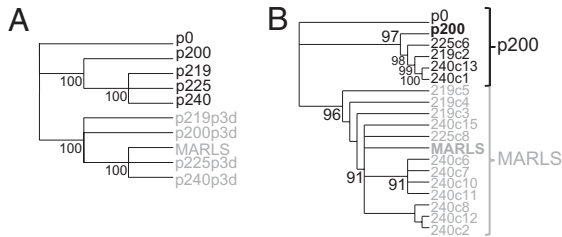


Fig. 1. Genetic characterization of p200 and MARLS viruses. (A) Maximum likelihood reconstruction of consensus nucleotide sequences (nucleotides 1033 through 1154 and 1570 through 3853, see *Materials and Methods*) of populations before and after three sequential low-MOI infections (indicated by "p3d"). (B) Maximum likelihood reconstruction [nucleotides 1033 through 3853; residue numbering is as described in (10)] of biological clones. Clones are identified by passage number and clone number (e.g., 225c6). Consensus reference sequences at passages 0, 200 (p0, p200), and of MARLS virus are included. Confidence values higher than 80% are indicated.

240c1 and 240c13 (KS-test, $P > 0.05$, *SI Appendix*). MARLS clone 240c12 production was slightly higher than that of both p200 clones (KS-test, $P < 0.05$). These results demonstrate that slow p200 viruses directly interfere with the replication of fast

MARLS viruses when coinfecting the same cell. We refer to p200 viruses as competitors because of their intracellular competitive advantage and to MARLS viruses as colonizers because of their higher cell killing capacity, which entails a faster dispersal upon completion of the infectious cycle.

Mathematical Model of Viral Coinfection Dynamics. To pinpoint the coevolutionary dynamics of competitors and colonizers, and to generate testable hypotheses, we developed a mathematical model of virus dynamics (18, 19) in cell culture that accounts for intracellular interactions. Two different viruses, whose abundances are denoted by v_1 and v_2 , compete for an uninfected cell pool of size x and give rise to singly infected and coinfecting cell populations of size y_1 , y_2 , and y_{12} , respectively. The dynamics are given by the ordinary differential equations (ODE)

$$\begin{aligned}\dot{x} &= -\beta xv_1 - \beta xv_2 \\ \dot{y}_1 &= \beta xv_1 - \beta y_1 v_2 - a_1 y_1 \\ \dot{y}_{12} &= \beta y_1 v_2 + \beta y_2 v_1 - a_1 y_{12} \\ \dot{y}_2 &= \beta xv_2 - \beta y_2 v_1 - a_2 y_2 \\ \dot{v}_1 &= k_1 y_1 + c k_1 y_{12} - uv_1 \\ \dot{v}_2 &= k_2 y_2 + (1 - c) k_1 y_{12} - uv_2\end{aligned}$$

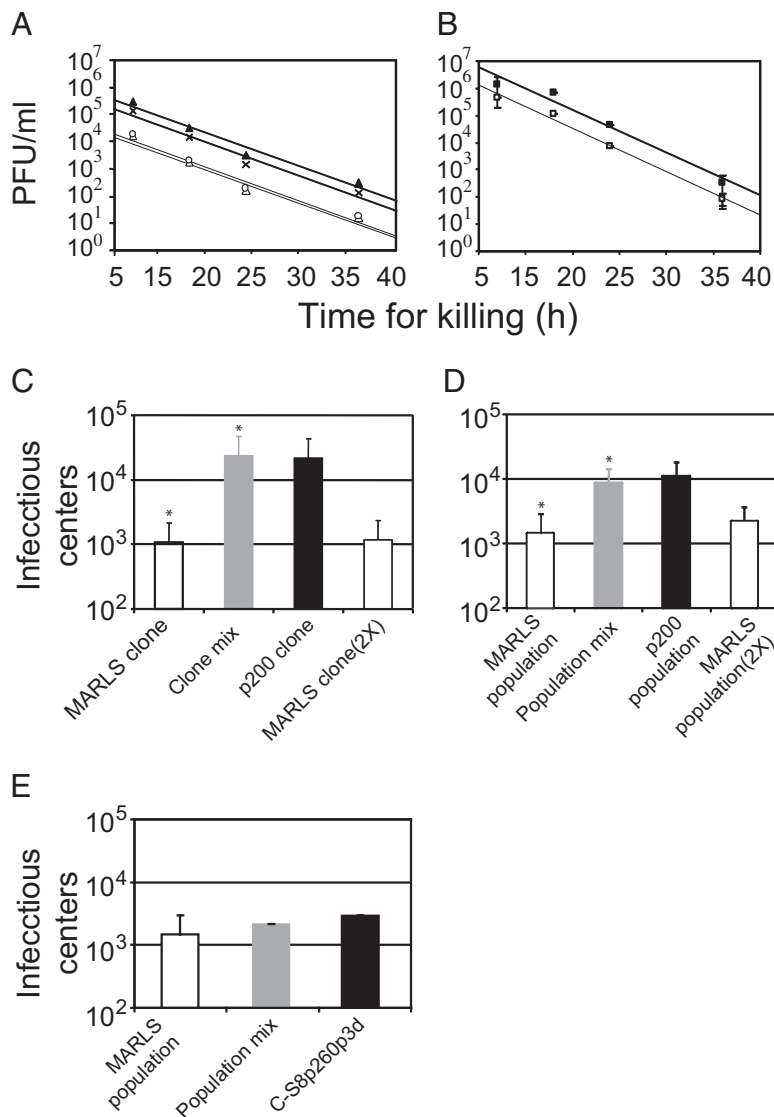


Fig. 2. Cell killing and cell killing-interference assays. (A and B) Cell killing capacity of viral populations and clones, measured as the time needed to kill 10^4 BHK-21 cells as a function of the initial number of PFU. (A) MARLS clones: 240c2 (\circ) and 240c12 (Δ); p200 clones: 240c1 (\blacktriangle) and 240c13 (\times). (B) Population p200 (\blacksquare), population p200p5d, MARLS population (\square). For each point, the average and standard deviation from three independent determinations are indicated. (C–E) Cell killing-interference assays. Number of infectious centers required by p200, MARLS, or the mixture of both viruses to kill 10^4 BHK-21 cells in 9.5h; 2 \times , double dose of MARLS clone or population. Each bar represents the mean and standard deviation from triplicate assays. (C) p200 clone and MARLS clone correspond to 240c1 and 240c12, respectively; *The mean of the "Clone mix" is higher than the mean of the "MARLS clone" with marginal significance ($P < 0.08$, T-Student). (D) population mix indicates a mixture of MARLS population and p200 population. *The mean of the "population mix" is significantly higher ($P < 0.05$, T-Student) than the mean of the MARLS population. (E) C-S8p260p3d is the virus recovered by passage of C-S8c1 after 260 transfers in cell culture and 3 passages at low-MOI; this virus has a virulence similar to MARLS (10).

This ODE system describes uninfected cells being infected with efficiency β , infected cells dying and releasing viral offspring at rate a , and free virus being produced at rate k and inactivated at rate u . Because cell monolayers are confluent, no additional parameters describing any external supply of cells or cell division are required. The model parameters a and k are indexed by virus type, and the intracellular competition parameter c denotes the probability that viral offspring of a coinfecting cell is of type 1. Virus 1 is the competitor and its parameters have been measured from p200 viruses, whereas virus 2 is the colonizer, characterized by parameters obtained from MARLS viruses (Table 1). Specifically, competitor viruses are more likely to be produced by coinfecting cells ($c > 1/2$) as determined by high-MOI infection experiments (see *SI Appendix*). Interference is reflected by the condition $a_{12} = a_1$, which expresses the delay of colonizers in cells coinfecting by competitors and which was demonstrated in cell killing-interference assays (Fig. 2 C–E). As shown in the cell killing experiments (Fig. 2 A and B), colonizers are more efficient in cell killing ($a_2 > a_1$), and because the burst sizes ($K_i = k_i/a_i$) of the two viruses are equal, colonizers also replicate faster ($k_2 > k_1$). By introducing a linear change of coordinates the parameter space of the ODE system can be seen to be 2D (see *SI Appendix*). It is given by the rescaled parameters $a = a_2/a_1$, the ratio of cell death rates, and c , the intracellular competition parameter. The competitors p200 and the colonizers MARLS differ only in precisely these two parameters (Table 1).

We have solved the ODE system numerically for different initial viral densities to assess the winner of the competition as the virus type produced with the highest total abundance. For the competitor-colonizer region of the parameter space, i.e., for $c > 1/2$ and $a > 1$, the winner of a competition can be either the competitor or the colonizer, depending on the initial MOI (see *SI Appendix*). The advantage of the colonizer decreases with the initial density of viruses, to the point where the competitor can outcompete the colonizer under high-density conditions. In the limit of high MOI, we have also found an analytical solution of the ODE system which confirms this observation. Competitors outcompete colonizers in total numbers if their intracellular advantage c is larger than the critical value

$$c^* = \frac{a + v_0}{1 + a + 2v_0}$$

In the competitor-colonizer regime ($a > 1$), this threshold is always greater than $1/2$. Increasing initial viral densities v_0 shift the fitness advantage from colonizers to competitors because $dc^*/dv_0 < 0$. Thus, the model predicts a density-dependent outcome of the coinfection.

Experimental Test of Model Prediction. To test the density-dependent outcome of the infection predicted by the model, we determined the final abundances of viruses for different initial viral densities both computationally, using the parameters measured from p200

and MARLS (Table 1; an estimate of the values of the parameters is included in the *SI Appendix*), and experimentally. The p200 and MARLS clones and populations were subjected to standard virus competition assays (see *Materials and Methods*) at different initial MOI (see *SI Appendix*). The average fitness of MARLS relative to p200 was found to be dependent on the initial density: the higher the MOI, the lower the relative fitness of MARLS viruses (Fig. 3). The observed fitness values are in good agreement with the predicted fitness values and they confirm the predicted power law dependence of fitness on MOI, although the effect of the space (2D nature of cell monolayers) may be responsible for the lack of exact match. In particular, the experimental data validate the main prediction of the model, namely that fitness is density-dependent and that both competitors and colonizers can be winners of this coevolutionary process.

Discussion

We have described rapid self-organization in the progeny mutant spectrum of a single viral clone into two different ecological niche specialists. The coexistence of the two strategies relies on intracellular reproductive success due to interference (competition) and on intercellular spread due to increased virulence (colonization) (17, 20). The rapid quasispecies rearrangement was probably facilitated by a high mutation rate (21–23) and the fact that cell killing in RNA viruses can be modulated by a small subset of mutations at potentially many different genomic sites (10, 11). The suppression of colonizers by competitors at high viral density can lead to the attenuation of a viral population. This mechanism could also explain the previous observation of mutant suppression or density-dependent selection in dissimilar viral systems such as vesicular stomatitis virus (24, 25), FMDV (26), and bacteriophages $\phi 6$ and $\phi X174$ (27, 28). In these studies, a subset of clones or subpopulations that showed high fitness in low-MOI infections was outcompeted under high-MOI conditions. The faster replicators under low-MOI conditions probably acted as colonizers, whereas the suppressors at high-MOI acted as competitors. Defective interfering particles maintenance in high-MOI infections strongly attenuates viral infections (29), whereas serial bottleneck (low-density) transfers maintained the cell killing capacity of an FMDV clone despite its reduced replicative fitness (10). Fitness determinants of FMDV are scattered along the viral genome (9, 10) whereas BHK-21 cell-killing determinants mapped mainly in the nonstructural protein-coding region (10).

Mutants with a high cell-killing efficiency that replicate in independent cells may have a selective advantage because they spread faster (30–32). However, viruses share gene products inside coinfecting cells allowing the progression of dominant-negative mutants (24, 33–35) during processes such as pathogen-derived resistance (36), lethal defection (34, 37), or pseudotype formation (38). Therefore, unlike in bacteria or protozoa (39, 40), coinfection would tend to attenuate virus populations. Our model might also explain the attenuation of a clone of FMDV

Table 1. Parameters and estimated values of the model of virus competition in cell culture

Parameter*	Value	Description	Method of measurement†
a_1	0.14 h^{-1}	Cell killing rate of virus 1	Counting of live cells
a_2	0.25 h^{-1}	Cell killing rate of virus 2	Calculated from the basic reproductive ratio
u	0.28 h^{-1}	Viral inactivation rate	Slope of exponential decay of infectivity
β	$7.8 \cdot 10^{-8}$ ml h^{-1}	Infectivity rate‡	Calculated from the basic reproductive ratio
c	0.62	Probability of a coinfecting cell to produce virus 1	Specific RNA quantification in high-MOI infections
K	250	Burst size	Titration of viral progeny after complete cell lysis

*See *SI Appendix* for details.

†Virus 1 and virus 2 correspond to p200 (competitor) and MARLS (colonizer) viruses, respectively.

‡The initial number of susceptible cells was $x_0 = 2 \times 10^6$ cells in a volume of 2 ml.

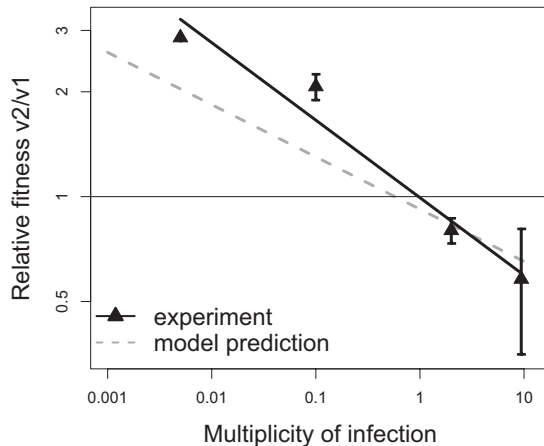


Fig. 3. Model predictions and experimental results of virus competition experiments. The basic model of virus competition in cell culture was implemented and solved for the parameters listed in Table 1, to obtain relative fitness of MARLS compared to p200 which are plotted as a function of MOI (●). Experimental fitness values were obtained from virus competition experiments. Two representative clones each of MARLS and p200 subpopulations were mixed at equal PFU, and challenged in competitions with variable initial MOI. Reference p200 and MARLS populations (p200p5d) were also mixed in direct competition at variable initial MOI. Relative fitness values obtained in each competition for both clones and populations are plotted against MOI (▲). Experimental measurements with standard errors (solid line) and model predictions (dashed line) were fitted by linear regression. Procedures for fitness value measurements and competition experiments are detailed in *Materials and Methods* and the *SI Appendix*.

after mice-to-mice transfers because attenuated strains were only isolated in organs in which the virus spreads locally thereby increasing the probability of coinfection (41). Coinfection of cells by two or more viral particles must be abundant *in vivo*, as judged by the frequency of viral recombinants that are described as epidemiologically relevant for many viral systems (42–44). Recombination could be enhanced by the fact that double infections of cells might be more frequent than expected from dual hits occurring at random as documented for some virus-host systems (45, 46). Coinfection of cells has been described in HIV and Dengue infections (47, 48). The analyses with FMDV reported here document that mutants within a viral quasispecies behave as ecological entities and follow competition-colonization dynamics. This compromise, which ensues from intrapopulation self-organization, can modulate the cell killing capacity of the entire viral population. In addition to implications for virulence as a trait that can be modulated in complex viral populations, the results reinforce the concept that mutant spectra can act as a unit of selection (4, 22, 33, 34).

Materials and Methods

Cells, Viruses, and Infections. Infections of BHK-21 cell monolayers with FMDV have been previously described (10, 15, 26). The FMDVs used in the present study are the initial clone C-58c1 (p0, GenBank NC 002554) (49) and viral populations derived by passage of p0 at high MOI (1–10 PFU/cell). They are labeled by “p” followed by the passage number (e.g., p200, GenBank FJ824812). MARLS (GenBank AF274010) is a monoclonal antibody-escape mutant of FMDV C-58c1, selected as a minority component from p213 (15). Biological clones were obtained by dilution and plating in semisolid agar

medium (50). Clones are labeled with the number of the population, followed by “c” and a clone number (e.g., p240c1). Specifically, p240c1 and p240c13 have a MARLS sequence, and p240c2 and p240c12 have a p200 sequence, as shown by phylogenetic analysis (see Fig. 1). Serial low-MOI passages (MOI = 0.006–0.02 PFU/cell) were carried with p200, p219, p225, and p240 to obtain populations p3d after three low-MOI passages, and p5d after five low-MOI passages (see *SI Appendix*). Population p200p5d was employed throughout the study as the reference MARLS population. C-58p260p3d derives from C-58c1 after 263 passages at high-MOI (1–10PFU/cell), and its cell killing capacity is similar to that of MARLS (10).

Competition Between Viruses. Competition between viruses was carried out as described previously (10). Briefly, known numbers of PFU of the two viruses were adsorbed onto BHK-21 cell monolayers for 1 h at 37 °C; then the monolayers were washed to remove unadsorbed virions, and the infection was allowed to proceed until complete cytopathology. Viral RNA specific for each virus, present in the cell culture supernatants was quantitated by real-time RT-PCR. The primers that specifically amplify MARLS and p200 sequences are CACGCTACTATTTTCTGATTG and CACGCTACTCTTTTCTGATCTG, respectively. Relative fitness values were calculated as described in the *SI Appendix*.

Cell-Killing Assay. The capacity of FMDV to kill BHK-21 cells was measured as previously described (10). The assay consists in determining the minimum number of PFU required to kill 10^4 BHK-21 cells after variable times of infection. The experiments were performed using multiwell M96 plates seeded with 10^4 BHK-21 cells per well and then infected by serial dilutions of the virus to be tested, following the standard infection protocol. At different times postinfection the cells were fixed with 2% formaldehyde. The results are expressed as the logarithm of the number of PFU (PFU/mL of the virus used for the infection) as a function of the time needed to kill the 10^4 BHK-21 cells. Control viruses with previously measured cell killing capacities were included in all of the experiments. Cell viability was measured by cell counting after trypan blue staining (37).

Cell Killing-Interference Assay. The cell killing assay was adapted to measure the interference that FMDV variants exerted on the killing of BHK-21 by other variants, as follows. Monolayers of $5 \cdot 10^5$ cells were infected at high-MOI (> 20 PFU/cell) with either individual variants or with an equal number of PFUs of the two variants (killing and interfering) to be tested. After a 1 h adsorption period, cells were detached by trypsin-EDTA treatment, washed, serially diluted, and applied onto fresh monolayers of 10^4 cells. The monolayers were overlaid with semisolid agar. At 9.5 h postapplication, the minimum number of infected cells required to kill 10^4 BHK-21 cells was measured.

Viral Phylogenies. Phylogenetic trees were inferred by maximum likelihood, using the Tamura-Nei substitution model with Gamma distributed heterogeneous rates (TN-81) (51). The genomic regions chosen for sequence comparison included the capsid-coding region and provided a sufficient average number of mutations in the consensus sequence and among individual clones to achieve the required resolution in the phylogenetic analyses (9, 10, 15, 52, 53). Nucleotides 1154 through 1570 were not considered in the analysis presented in Fig. 1A because some populations contain a proportion of genomes harboring internal deletions (52).

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Parameter	Description	Virus 1	Virus 2	Unit
		(competitor)	(colonizer)	
r_0	initial viral growth rate	1.43	1.94	h^{-1}
R_0	basic reproductive number	69.5	69.5	1
a	death rate of infected cells	0.14	0.25	h^{-1}
u	clearance rate of free virus	0.28	0.28	h^{-1}
$K = k/a$	burst size	250	250	1
k	production rate of free virus	34.6	62.5	h^{-1}
β	rate of infection	$7.8 \cdot 10^{-8}$	$7.8 \cdot 10^{-8}$	$ml \cdot h^{-1}$
c	prob. of type 1 viral offspring	0.62	$1 - c$	

Table S1: Parameters of the basic model of virus dynamics in cell culture.

- equal infection efficiencies, $\beta_1 = \beta_2 = \beta_{12} = \beta_{21} =: \beta$,
- equal viral clearance rates, $u_1 = u_2 =: u$.

Our experimental results (Table S1) further specify the parameters to

- equal death rates of cells infected by virus 1 and those doubly infected, $a_1 = a_{12}$,
- equal burst sizes, $K := k_1/a_1 = k_2/a_2 = k_{12}/a_{12}$.

These two constraints also imply the equality $k_{12} = k_1$ of viral production rates. In other words, the experimental findings summarized in Table S1 assert that a coinfecting cell behaves like a cell infected only by virus 1 with the exception that it produces both types of viruses.

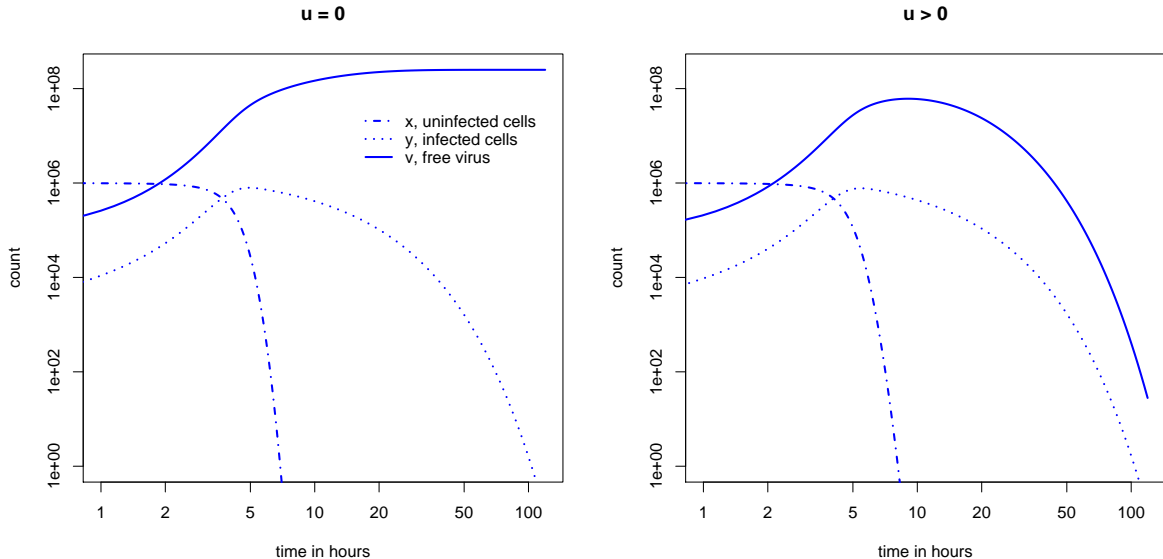


Figure S1: Dynamics of the basic cell culture model (S1) assuming no clearance of free virus ($u = 0$, left) and a positive death rate of free viruses ($u > 0$, right). Shown are the abundances of uninfected cells (x , dash-dot line), infected cells (y , dotted line), and free virus (v , solid line).

1.3.1 Initial conditions

To control the amount of virus that is added to a cell culture, the first stages of infection are carried out as in a virus titration assay. A volume of medium containing a known amount of virus is added to the culture. One hour later, the virus inoculum is removed and fresh medium is added to the culture. For this technical reason, in the first infection we deal with infected cells in absence of free virus. Thus, we calculate the number of infected cells in this single round of infection and use this value, rather than the initial number of viruses, as the starting solution of the ODE system.

Let us first consider a single virus at initial multiplicity of infection $\lambda = v/x_0$. The probability of finding a k -fold infected cell, denoted $X = k$, after one round of infection is given by the Poisson distribution [2]

$$\text{Prob}(X = k) = \frac{\lambda^k e^{-\lambda}}{k!},$$

if we assume a large number of cells and independent viruses. Thus, the probability of a cell being infected by at least one virus particle is

$$\text{Prob}(X \geq 1) = 1 - \text{Prob}(X = 0) = 1 - e^{-\lambda}.$$

For two viruses with $\lambda_1 = v_1/x_0$ and $\lambda_2 = v_2/x_0$, let $X_i = k$ denote a k -fold infection with virus i , $i = 1, 2$. Assuming that infections are independent, we find

$$\begin{aligned} p_{00} &= \text{Prob}(X_1 = 0, X_2 = 0) = e^{-(\lambda_1 + \lambda_2)} \\ p_{01} &= \text{Prob}(X_1 = 0, X_2 \geq 1) = e^{-\lambda_1} (1 - e^{-\lambda_2}) \\ p_{10} &= \text{Prob}(X_1 \geq 1, X_2 = 0) = (1 - e^{-\lambda_1}) e^{-\lambda_2} \\ p_{11} &= \text{Prob}(X_1 \geq 1, X_2 \geq 1) = (1 - e^{-\lambda_1}) (1 - e^{-\lambda_2}) \end{aligned}$$

and use the following initial conditions for the ODE system:

$$x(0) = p_{00}x_0, \quad y_1(0) = p_{10}x_0, \quad y_2(0) = p_{01}x_0, \quad y_{12}(0) = p_{11}x_0, \quad v_1(0) = v_2(0) = 0 \quad (\text{S4})$$

1.3.2 Counting viruses

As a comparative measure of success of two competing viruses in cell culture, we quantify their abundances at the end of the infection experiment. Because the experimental read-out can not distinguish infectious from non-infectious virus particles, in the mathematical model, we also have to account for inactivated viruses. The abundances of inactivated viruses, w_1 and w_2 , follow the additional equations

$$\begin{aligned} \dot{w}_1 &= uv_1 \\ \dot{w}_2 &= uv_2. \end{aligned}$$

We define t^* as the earliest time point at which the number of infected cells is less than one,

$$t^* = \inf\{t \geq 0 \mid y_1(t) < 1, y_2(t) < 1, y_{12}(t) < 1\},$$

and write $v(t^*) = v^*$, etc. The computational read-out is the relative fitness advantage of virus 2, the colonizer, over virus 1, the competitor,

$$f = \frac{v_2^* + w_2^*}{v_1^* + w_1^*}.$$

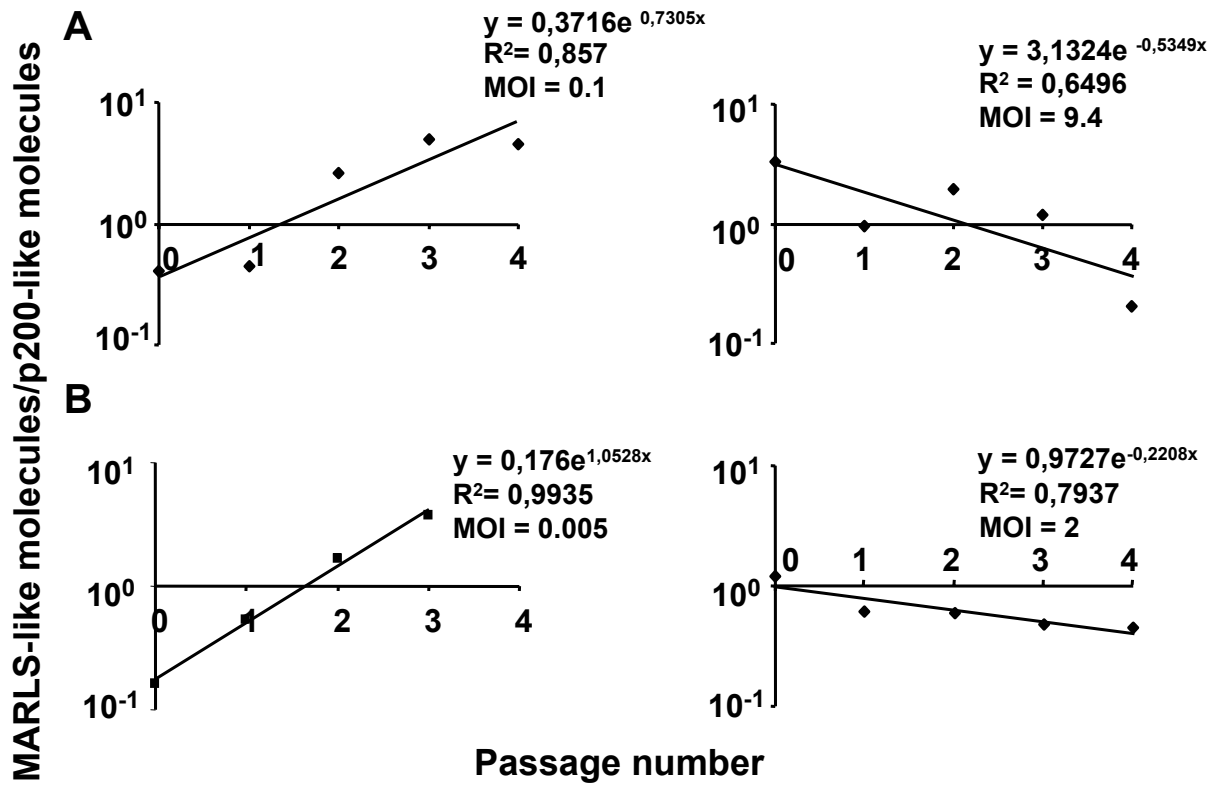


Figure S3: Competition experiments between MARLS and p200 viruses at different MOI. The competitions and quantifications of viral RNA were carried out as specified in the Methods section. Each point represents the number of MARLS genomic RNA molecules divided by the number of p200 genomic RNA molecules. Data have been fitted by linear regression; Regression equation, R^2 value, and MOI used are indicated in each plot. A) Competition between populations p200 and p200p5d (MARLS). B) Competition between two p200 representative clones (240c1, 240c13), and two MARLS representative clones (p240c2, p240c12). All viruses were mixed using equal number of PFU. As a relative fitness measure we used the slope of each graph which is the increase in frequency of each genotype per passage.

The experimental assessment of the final relative abundances of both viruses was performed in competition assays (Figure S3) for different initial viral densities

$$\lambda = \frac{v_{0,1} + v_{0,2}}{x_0}. \quad (S5)$$

Figure S2 illustrates that, according to the model, the winner of the competition can be either the competitor or the colonizer, depending on the initial multiplicity of infection. In Figure 2 of the main text, this model prediction is validated by comparing predicted and observed relative viral abundances.

2 Experimental measurements

2.1 Origin of viral strains

We first describe the experimental procedures that were employed to obtain the two different viral subpopulations.

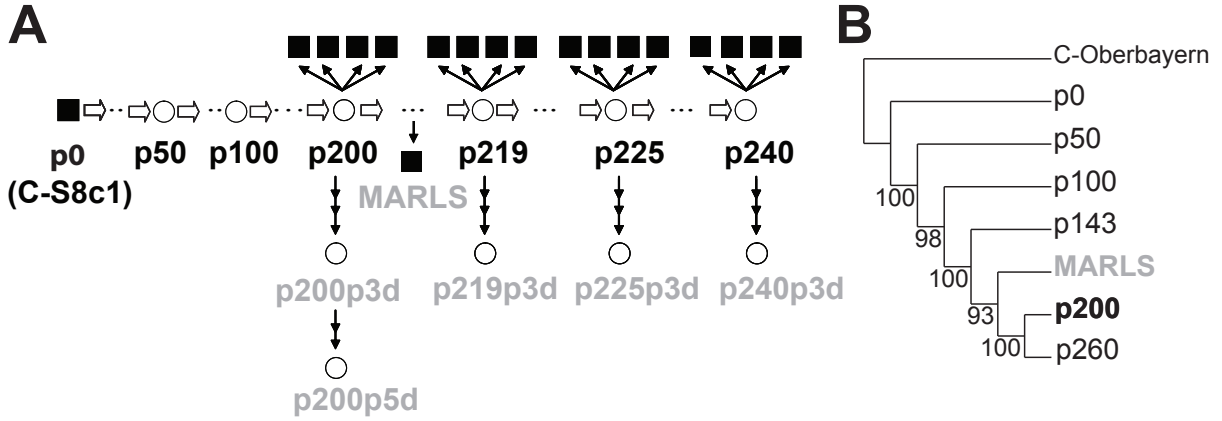


Figure S4: Genetic diversification of a biological clone of FMDV in cell culture. (A) Schematic representation of passages of clone C-S8c1 (p0) in BHK-21 cells. Biological clones are depicted as filled squares and populations as empty circles. Thick arrows indicate high-MOI passages (1 to 20 PFU/cell), sequential thin arrows indicate low-MOI infections (0.006 to 0.02 PFU/cell), and single thin arrows represent isolation of biological clones. Viral populations are labelled by passage number (e.g., p200 at passage 200). Populations with suffix p3d and p5d were derived by three and five low-MOI passages, respectively. MARLS is a monoclonal antibody-escape clone isolated at passage 213. Viruses resembling MARLS and p200 are labelled in grey and black, respectively. (B) Phylogenetic reconstruction based on whole genome consensus nucleotide sequences of populations derived from biological clone p0. The sequence of C-Oberbayern is included as outgroup. The tree was constructed by maximum likelihood using the Tamura-Nei substitution model with Gamma distributed rates (TN-8 Γ). Confidence values higher than 80% are shown at the corresponding branching points.

Serial infections of BHK-21 cells at high MOI were carried out with the biologically pure clone of FMDV C-S8c1 (Figure S4A). Evolutionary history of the consensus sequences at consecutive passages was reconstructed by maximum likelihood phylogenetic analysis (Figure S4B). The phylogenetic tree reflected the successive evolution of the virus and showed that C-S8c1 diversified into MARLS between passage 143 and passage 200.

2.2 Model parameters

We have performed several experiments and employed theoretical considerations in order to obtain the parameter values displayed in Table S1.

Basic reproductive number, R_0 . The basic reproductive number of a virus, R_0 , is defined as the number of secondary infections, i.e., the number of infections that result from a single infected cell, when all cells are uninfected. If $R_0 > 1$, each cell produces on average more than one virus and the virus population will initially grow exponentially as $v(t) \propto e^{r_0 t}$. The growth rate, r_0 , is the largest root of the equation

$$r_0^2 + (a + u)r_0 + au(1 - R_0) = 0 \quad (S6)$$

[3]. From this equation, we can derive the basic reproductive number, R_0 . The initial growth rate of each virus, $r_{0,i}$, has, in turn, been determined experimentally by averaging the slope of the virus growth curves depicted in Figure S5

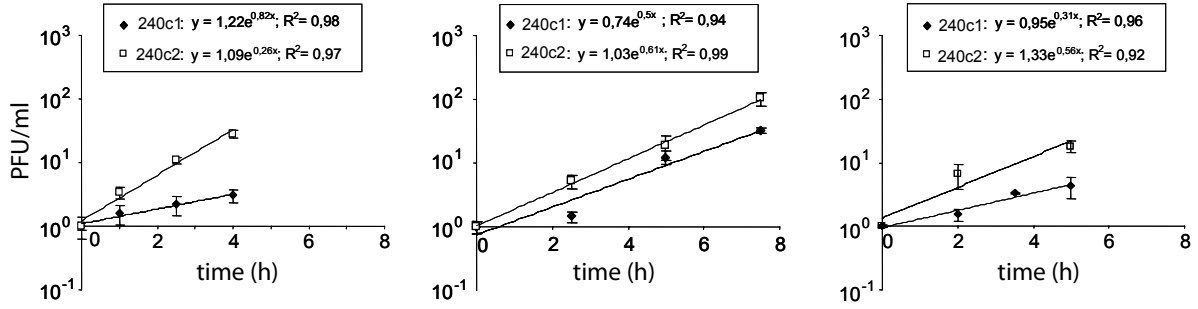


Figure S5: Viral growth curves. The initial exponential growth of the MARLS clone, 240c2, and the p200 clone, 240c1, was monitored by titration at several time points of the supernatant of cells infected independently by each clone. The growth rates $r_{0,i}$ correspond to the average slopes of each experiment represented in the individual panels.

Virus ^a	Subpopulation	Progeny production (PFU/ml) ^b
240c1	p200, competitor	$2.95 \times 10^7 \pm 9.84 \times 10^6$
240c13	p200, competitor	$3.43 \times 10^7 \pm 4.41 \times 10^6$
240c2	MARLS, colonizer	$3.74 \times 10^7 \pm 5.49 \times 10^6$
240c12	MARLS, colonizer	$6.90 \times 10^7 \pm 2.78 \times 10^7$

^a The origin of clones is described in the Materials and Methods Section, in Section 2.1, and in Figure S4

^b Each titration was determined at least in triplicate. Two independent infections were carried out with each clone to determine the virus yield.

Table S2: Progeny production of MARLS and p200 clones. BHK-21 cells were infected with equal amounts of specified virus clones. After 1h of adsorption to the cells, the inoculum was removed, cells were washed, and fresh medium was added. When complete cytopathic effect was reached, infectivity was determined by titration of the cell culture supernatant.

Death rate of infected cells, a . To obtain the differences in a , the time that the slow virus takes to kill one cell (a_2) was measured by monitoring cell viability with the trypan blue staining technique (see Methods section in the main text). This procedure does not allow to determine, with sufficient accuracy, the death rate of fast viruses that kill host cells in a few hours. For this reason, we use Equation S6 to derive a_1 in a precise fashion.

Clearance rate of free virus, u . FMDV infectivity decays exponentially with time [1]. We assume uniform inactivation rates $u_1 = u_2 = u$ (cf. Sec 1.2) and take the mean value of $u = 0.28 \text{ h}^{-1}$, which corresponds to the standard observation in our laboratory of the inactivation rate of FMDV at 37°C.

Burst size, $K = k/a$. Production of MARLS and p200 viruses was very similar as shown by titration of individual viral clones and reported in Table S2. Dividing these virus titers by the number of infected cells, one obtains $K = k_1/a_1 = k_2/a_2 = k_{12}/a_{12}$.

Efficiency of infection, β . We assume uniform infection efficiencies $\beta_1 = \beta_2 = \beta_{12} = \beta_{21} = \beta$ (cf. Sec 1.2) and obtain a numerical estimate from the equation $R_0 = \beta k x_0 / a u$ for the basic reproductive number [3].

Probability of producing type 1 viral offspring, c . As described for the calculation of the burst size, the viral progeny production in a viral infection at high multiplicity of infection (MOI) represents an average of the production by each individual cell. The advantage of p200 viruses in coinfecting cells was calculated by averaging the results of the serial competitive infections used to determine relative fitness and confirmed by three additional individual infections carried out at high MOI.

3 Mathematical analysis

For our mathematical analysis of the ODE system describing two competing viruses in cell culture, we rewrite the equations using $K = k_1/a_1 = k_2/a_2$,

$$\begin{aligned}\dot{x} &= -\beta x v_1 - \beta x v_2 \\ \dot{y}_1 &= \beta x v_1 - \beta y_1 v_2 - a_1 y_1 \\ \dot{y}_{12} &= \beta y_1 v_2 + \beta y_2 v_1 - a_1 y_{12} \\ \dot{y}_2 &= \beta x v_2 - \beta y_2 v_1 - a_2 y_2 \\ \dot{v}_1 &= K a_1 y_1 + c K a_1 y_{12} - u v_1 \\ \dot{v}_2 &= K a_2 y_2 + (1 - c) K a_1 y_{12} - u v_2 \\ \dot{w}_1 &= u v_1 \\ \dot{w}_2 &= u v_2.\end{aligned}$$

3.1 Conservation law

Slightly abusing notation we set, for $i = 1, 2$, $v_i^* = v_i(\infty)$ and $w_i^* = w_i(\infty)$. The sum

$$\phi = x + y_1 + y_{12} + y_2 + \frac{v_1 + w_1 + v_2 + w_2}{K}$$

is a conserved quantity of the system, i.e., $\dot{\phi} = 0$. For the initial conditions (S4), we obtain from $\phi(0) = \phi(\infty)$ the equation

$$v_1^* + w_1^* + v_2^* + w_2^* = K x_0.$$

This relation does not determine the final numbers of both viruses, but defines a constraint on them. This restriction does not depend on β (even if we would not have assumed that all infection rates are equal). Similarly, for the initial conditions

$$x(0) = x_0, \quad y_1(0) = y_2(0) = y_{12}(0) = 0, \quad v_1(0) = v_{1,0}, \quad v_2(0) = v_{2,0}$$

we have

$$v_1^* + w_1^* - v_{1,0} + v_2^* + w_2^* - v_{2,0} = K x_0,$$

Using Eq. S5 we obtain an expression similar to Eq. S2,

$$v_1^* + w_1^* + v_2^* + w_2^* = (K + \lambda) x_0.$$

3.2 Parameter space

We introduce the following linear change of coordinates,

$$\begin{aligned}\hat{t} &= a_1 t, \quad \hat{x} = \frac{\beta K}{a_1} x, \quad \hat{y}_1 = \frac{\beta K}{a_1} y_1, \quad \hat{y}_{12} = \frac{\beta K}{a_1} y_{12}, \quad \hat{y}_2 = \frac{\beta K}{a_1} y_2, \\ \hat{v}_1 &= \frac{\beta}{a_1} v_1, \quad \hat{v}_2 = \frac{\beta}{a_1} v_2, \quad \hat{w}_1 = \frac{\beta}{a_1} w_1, \quad \hat{w}_2 = \frac{\beta}{a_1} w_2,\end{aligned}$$

to obtain the simplified ODE system (where we drop the hats)

$$\begin{aligned}\dot{x} &= -xv_1 - xv_2 \\ \dot{y}_1 &= xv_1 - y_1v_2 - y_1 \\ \dot{y}_{12} &= y_1v_2 + y_2v_1 - y_{12} \\ \dot{y}_2 &= xv_2 - y_2v_1 - ay_2 \\ \dot{v}_1 &= y_1 + cy_{12} - \mu v_1 \\ \dot{v}_2 &= ay_2 + (1 - c)y_{12} - \mu v_2 \\ \dot{w}_1 &= \mu v_1 \\ \dot{w}_2 &= \mu v_2\end{aligned}$$

This system has only three parameters, namely $a := a_2/a_1$, c , and $\mu := u/a$. We have eliminated the direct dependencies on β , K , and a_1 . The competitor and the colonizer described in Table S1 differ only in two of the three remaining parameters, namely a and c .

In order to explore the complete parameter space of the model it is enough to consider the case where $\beta = 1$, $K = 1$, and $a_1 = 1$. We are interested whether the relative fitness

$$f = \frac{v_2^* + w_2^*}{v_1^* + w_1^*} = \frac{(1 + \lambda)x_0}{v_1^* + w_1^*} - 1$$

is greater or less than 1 for any given values of a , c , and μ .

Figure S6 explores this quantity by plotting the sign of its logarithm in the a - c plane for three different multiplicities of infection and three different values of μ . Positive values ($f > 1$) indicate that colonizers win the competition and are displayed in red, whereas at negative values ($f < 1$) competitors win.

For values of c below $1/2$, colonizers will always win, whereas for $c > 1/2$ and small values of a , competitors will always outcompete colonizers. However, for $c > 1/2$ and $a > 1$, the outcome of the competition depends on the initial multiplicity of infection: colonizers win under low-density conditions, but competitors win under high-density conditions.

The rows of Figure S6 show that this effect is largely independent of the viral inactivation rate μ . We therefore assume $\mu = 0$ for the following analytical analysis of the model. The ODE system then simplifies to

$$\begin{aligned}\dot{x} &= -xv_1 - xv_2 \\ \dot{y}_1 &= xv_1 - y_1v_2 - y_1 \\ \dot{y}_{12} &= y_1v_2 + y_2v_1 - y_{12} \\ \dot{y}_2 &= xv_2 - y_2v_1 - ay_2 \\ \dot{v}_1 &= y_1 + cy_{12} \\ \dot{v}_2 &= ay_2 + (1 - c)y_{12}\end{aligned}\tag{S7}$$

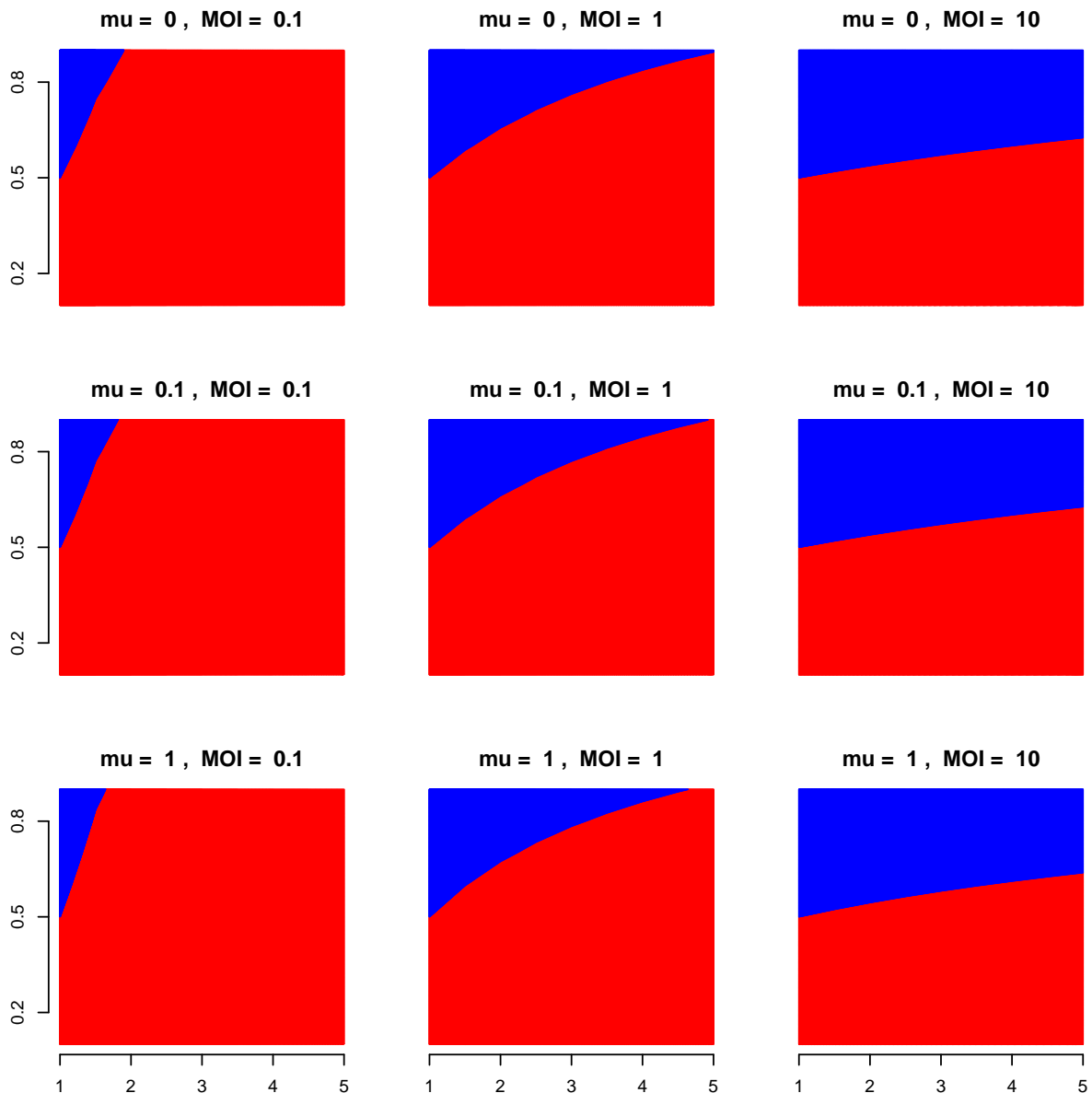


Figure S6: Viral competition outcome. Each plot indicates the winner of the competition experiment (blue: competitor wins; red: colonizer wins) over the parameter subspace defined by the cell killing ratio $a = a_2/a_1$ (x -axis) and the intracellular advantage c (y -axis). Subfigures correspond, from left to right, to increasing multiplicities of infection (MOI), λ , and, from top to bottom, to increasing viral inactivation rates, μ .

and it depends exactly on the two parameters a and c in which the competitors and colonizers of Table S1 differ.

3.3 Large initial virus load limit

Analytic treatment of the nonlinear ODE system (S7) is quite challenging. In order to obtain the final virus densities, we need to find the complete time dependent solution; the stationary solution is not enough. We can find a solution of (S7) in the limit of large initial virus density (MOI). Since the rate of infection is proportional to the virus load, for large virus loads the slow effect of virus inactivation can be neglected. We assume

that $v_1(0), v_2(0) \gg u/\beta$, and hence we take $u = 0$ as in (S7).

If the initial virus density is large, its relative change will be small during the entire process. Hence we will assume that the virus densities are constant during the process, $v_1 = v_1(0)$ and $v_2 = v_2(0)$, which makes the first four equations of (S7) linear. Then by letting the virus densities depend on time again, we can integrate the fifth and sixth equations of (S7) to obtain the total virus production. Let us now follow this outline.

We concentrate on the initial value problem $x(0) = x_0$ and $y_1(0) = y_{12}(0) = y_2(0) = 0$. The first four equations of (S7) remain unchanged if we measure all densities x , y_1 , y_{12} , and y_2 in units of x_0 . Hence the final virus production will be proportional to x_0 and, for simplicity, we can set $x(0) = x_0 = 1$.

For constant v_1 and v_2 the time dependent solution of the densities can be obtained by integrating (S7). We find

$$\begin{aligned} x(t) &= e^{-t(v_1+v_2)} \\ y_1(t) &= \frac{v_1}{v_1-1} [e^{-t(1+v_2)} - e^{-t(v_1+v_2)}] \\ y_2(t) &= \frac{v_2}{v_2-a} [e^{-t(a+v_1)} - e^{-t(v_1+v_2)}] \end{aligned} \quad (\text{S8})$$

The solution for y_{12} can be obtained by integrating the third equation of (S7), but the result is too cumbersome to include here. Now we can obtain the change in virus densities during the process by integrating the last two equations of (S7),

$$\begin{aligned} \Delta v_1 &= \int_0^\infty y_1(t) + cy_{12}(t) dt = I_1 + cI_{12} \\ \Delta v_2 &= \int_0^\infty ay_2(t) + (1-c)y_{12}(t) dt = aI_2 + (1-c)I_{12} \end{aligned}$$

where

$$\begin{aligned} I_1 &= \int_0^\infty y_1(t) dt = \frac{v_1}{(1+v_2)(v_1+v_2)} \\ I_2 &= \int_0^\infty y_2(t) dt = \frac{v_2}{(a+v_1)(v_1+v_2)} \\ I_{12} &= \int_0^\infty y_{12}(t) dt = \frac{v_1v_2(1+a+v_1+v_2)}{(a+v_1)(1+v_2)(v_1+v_2)}. \end{aligned} \quad (\text{S9})$$

We are interested in the difference of the final viral densities

$$\begin{aligned} \Delta &= v_2^* - v_1^* \\ &= v_2 + \Delta v_2 - v_1 - \Delta v_1 \\ &= v_2 - v_1 + aI_2 - I_1 + (1-2c)I_{12} \end{aligned}$$

where I_1 , I_2 , and I_{12} are given above by (S9), and in particular in the case of symmetric initial virus load, i.e., $v := v_1 = v_2$. In that case we obtain

$$\Delta = 1 - 2c - \frac{1-c}{1+v} + \frac{ac}{a+v} \quad (\text{S10})$$

which is a linear function of c . If both viruses equal in cell killing ($a = 1$) and in offspring production ($c = 1/2$), then of course $\Delta = 0$. Also, for any other value of x_0 , the difference Δ simply becomes x_0 times larger.

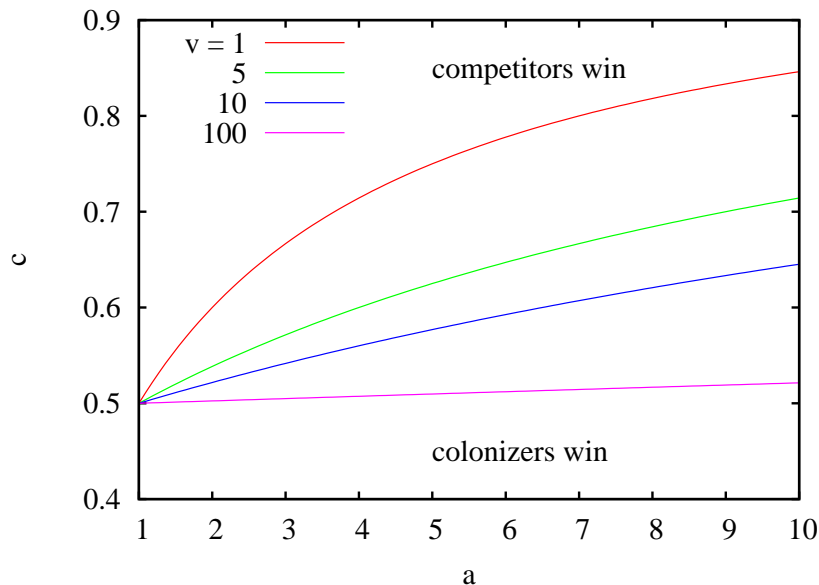


Figure S7: Boundary between regions where colonizers or competitors are favored at different values of initial virus density (v). The lines correspond to the exact solution in the large virus density limit (S11). Low viral density v favors the colonizers (second virus), and larger v favors the competitors (first virus).

Our main question is when the second virus (the competitor) becomes more successful than the first virus (the colonizer), that is when $\Delta > 0$. We find from (S10) that $\Delta > 0$ for $c < c^*$, where the threshold value is

$$c^* = \frac{a + v}{1 + a + 2v} \quad (\text{S11})$$

Competitors are more successful within coinfecting cells ($c > 1/2$), whereas colonizers replicate faster ($a = a_2/a_1 > 1$). In the regime of $a > 1$, we indeed have $c^* > 1/2$. For $a > 1$, the critical intracellular competition parameter c^* decreases as v increases, because its rate of change is always negative,

$$\frac{dc^*}{dv} = \frac{1 - a}{(1 + a + 2v)^2} < 0.$$

Hence, we have shown analytically that smaller viral density v favors colonizers, while larger viral density (MOI) favors competitors (Figure S7).

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