Mutagenesis-Induced, Large Fitness Variations with an Invariant Arenavirus Consensus Genomic Nucleotide Sequence

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Enhanced mutagenesis may result in RNA virus extinction, but the molecular events underlying this process are not well understood. Here we show that 5-fluorouracil (FU)-induced mutagenesis of the arenavirus lymphocytic choriomeningitis virus (LCMV) resulted in preextinction populations whose consensus genomic nucleotide sequence remained unaltered. Furthermore, fitness recovery passages in the absence of FU, or alternate virus passages in the presence and absence of FU, led to profound differences in the capacity of LCMV to produce progeny, without modification of the consensus genomic sequence. Molecular genetic analysis failed to produce evidence of hypermutated LCMV genomes. The results suggest that low-level mutagenesis to enrich the viral population with defector, interfering genomes harboring limited numbers of mutations may mediate the loss of infectivity that accompanies viral extinction.

Mutagenic agents administered alone or in combination with antiviral inhibitors can drive RNA virus populations to extinction (4, 18, 20, 21, 41, 42, 46, 48, 50, 51, 53, 65, 67, 72, 77, 79; reviewed in references 6, 27, 28, 31, and 39). Loss of infectivity has been interpreted as an irreversible transition that occurs when template copying fidelity falls below a critical value termed the error threshold. Such a transition has been termed virus entry into error catastrophe, or lethal mutagenesis (50), and its existence has been supported by several theoretical studies (5, 31, 60, 83). In agreement with this concept, analysis of the mutant spectra of virus populations on their way to extinction has shown 2- to 17-fold increases in mutation frequencies, calculated among components of the mutant spectra, as well as increases in Shannon entropy (a measure of the diversity of the different types of sequences present in a mutant spectrum) to nearly maximal values (that is, each component of the mutant spectrum showed a unique sequence) (reviewed in references 27 and 28).

The prototypic arenavirus lymphocytic choriomeningitis virus (LCMV) (12, 63, 64) showed extreme sensitivity to 5-fluorouracil (FU)-induced mutagenesis (41, 72) compared with other RNA viruses subjected to comparable doses of the mutagen (79). The extreme sensitivity of LCMV to FU offered an opportunity to analyze the capacity of LCMV to regain infectivity following FU mutagenesis as well as the modification of genomic nucleotide sequence variations as the virus moves toward or away from the error threshold and to explore the possible dominance of hypermutated genomes in the transition to extinction. The results reveal a remarkable capacity of LCMV populations to modify their fitness level while maintaining an invariant consensus sequence. Multiple molecular clones were analyzed to define a sequence at nucleotides 470 to 550 within the intergenic region of genomic segment L. A number of standard and mutated oligonucleotide primers have failed to produce evidence of hypermutated viral sequences in the L polymerase gene. The results suggest that limited mutagenesis may be sufficient to drive LCMV close to the error threshold and that LCMV can rapidly regain fitness in the absence of a mutagen. The invariance of the consensus genomic sequence, associated with the movements toward and away from the error threshold, suggests a decisive participation of the mutant spectrum in determining infectivity levels and supports a lethal deflection model for virus extinction through enhanced mutagenesis.

MATERIALS AND METHODS

Cells and virus. Baby hamster kidney cells (BHK-21) were grown in Dulbecco's modified Eagle's medium (DMEM) with 5% fetal calf serum (FCS) as described previously (79). Vero cells were maintained in DMEM supplemented with 3% FCS and 2% L-glutamine. LCMV strain Armstrong (ARM) 53b is a triple plaque-purified clone of Armstrong CA 1371, passaged four times in BHK-21 cells (10, 23, 30, 62, 73, 80, 82). For the present experiments, a stock virus (LCMV p0) was prepared by infecting 1 × 10^7 BHK-21 cells with LCMV at a multiplicity of infection (MOI) of 0.01 PFU per cell (41) (LCMV ARM 53b was kindly supplied by P. Borrow).

Virus infections. Semiconfluent (2.8 × 10^6 cells in 100-mm-diameter dishes [Falcon]) monolayers of BHK-21 were infected with 0.01 PFU of LCMV ARM 53b per cell in 5 ml of DMEM supplemented with 10% FCS, 10% tryptose phosphate broth, 2% l-glutamine, 0.52% glucose, and 50 µg/ml gentamicin. Viral supernatants were harvested 48 h postinfection, clarified by centrifugation at 2,500 rpm for 30 min at 4°C, and stored at −80°C. Supernatants were titrated on Vero cell monolayers under a semisolid agar medium, as described elsewhere (3). Viral yields usually were ≥10^9 PFU/ml.

Serial passages of LCMV ARM 53b were carried out by infecting control (D) or FU-treated (F) cell monolayers with the corresponding supernatants of the previous infection. For infections carried out with a virus obtained following a D passage, 2.8 × 10^6 BHK-21 cells were infected with about 3 × 10^4 PFU of LCMV, whereas for infections with a virus derived from an F passage virus, the input was ≤10^4 PFU of LCMV. Virus supernatants and cells were harvested 48 h postinfection. To obtain preextinction virus populations, semiconfluent BHK-21 monolayers (15 182-cm^2 flasks with 1.6 × 10^6 cells per flask) were infected at an

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MOI of 0.01 PFU/cell and incubated for 48 h in 10 ml of infection medium (with 100 µg/ml of FU) per flask. No specific LCMV sequences could be amplified and no infectivity (<10 PFU/ml) obtained from cell culture supernatants after a second passage in 100 µg/ml FU, even after 25-fold concentration with a Centricon Plus-20 Biomax-100 (Millipore) filter. Virus titers are means of three titrations; standard deviations never exceeded 10% of the mean.

Drug treatment. Preparation of FU stock solutions, BHK-21 cell viability assays, and infections in the presence of the mutagenic analogue were performed as previously described (41, 79). LCMV was considered extinguished when no reverse transcriptase PCR (RT-PCR)-amplifiable material and no infectivity (<10 PFU/ml) could be detected after three blind passages of the undiluted viral population in BHK-21 cells in standard culture medium in the absence of mutagens.

RNA extraction, RT-PCR, nucleotide sequencing, molecular cloning, and calculation of mutan-i spectrum complexity. RNA was extracted from supernatants of infected cell cultures by using Trizol (Invitrogen) according to the manufacturer's protocol. RNAs were amplified by two-step RT-PCR using ThermoScript reverse transcriptase (Promega) at 37°C for 45 min and then at 94°C for 2 min, followed by 20 cycles at 95°C for 1 min, 60°C for 1 min, and 72°C for 1 min with 16 pmol of reverse and forward primers. The reaction conditions were as follows: in the S genomic RNA, the glycoprotein-coding region between residues 268 and 738 was amplified using primers GP424F (forward) and GP57RF (reverse); in the nucleoprotein-coding region between nucleotides 2500 and 4240 was amplified with primers L2929F (forward) and L4260R (reverse) (this set amplified the region corresponding to amino acids 978 to 1422, which encompasses premotif A to motif E, including the catalytic domain of the viral polymerase [24, 52, 87]). Consensus nucleotide sequences were determined using entire viral RNA populations as templates, except for the L intergenic region, for which consensus sequences were derived from the alignment of sequences from multiple individual molecular clones. Most nucleotide sequences were determined two or more times using different sense and antisense primers (described at http://www.cbm.uam.es/fkfactory.esdomain/webs/CBMSO/pult_LinesInvestigacion.aspx?idObjeto=17). Nucleotide positions are given in the viral (genomic) sense and refer to the consensus genomic sequence determined in the present study. LCMV genomic nucleotide sequences were retrieved from GenBank (74–76) (GenBank accession numbers BANKIT 691408 and AY 894816). Sequences were compared using SEQMAN from the DNASTAR software package (Lasergene), and alignments were done with the GeneDoc program (K. B. Nicholas and H. B. Nicholas, Jr., Pittsburgh Supercomputing Center, Pittsburgh, Pa.).

Computer-based predictions of RNA secondary structure were carried out using the MFOLD program (92) with default folding.

The entire LCMV genome was sequenced using the primers and the strategy given in the supplementary material (http://www.cbm.uam.es/fkfactory.esdomain/webs/CBMSO/pult_LinesInvestigacion.aspx?idObjeto=17). For sampling of partial genomic sequences, three LCMV genomic regions were amplified by RT-PCR and sequenced, as follows: in the S genomic RNA, the glycoprotein-coding region between residues 268 and 738 was amplified using primers GP424F (forward) and GP57RF (reverse); in the nucleoprotein-coding region, residues 2244 to 2723 were amplified with primers NP2223F (forward) and NP2743R (reverse); in the L RNA, the polymerase-coding region between nucleotides 2500 and 4240 was amplified with primers L2929F (forward) and L4260R (reverse) (this set amplified the region corresponding to amino acids 978 to 1422, which encompasses premotif A to motif E, including the catalytic domain of the viral polymerase [24, 52, 87]). Consensus nucleotide sequences were determined using entire viral RNA populations as templates, except for the L intergenic region, for which consensus sequences were derived from the alignment of sequences from multiple individual molecular clones. Most nucleotide sequences were determined two or more times using different sense and antisense primers (described at http://www.cbm.uam.es/fkfactory.esdomain/webs/CBMSO/pult_LinesInvestigacion.aspx?idObjeto=17). Nucleotide positions are given in the viral (genomic) sense and refer to the consensus genomic sequence determined in the present study. LCMV genomic nucleotide sequences were retrieved from GenBank (74–76) (GenBank accession numbers BANKIT 691408 and AY 894816). Sequences were compared using SEQMAN from the DNASTAR software package (Lasergene), and alignments were done with the GeneDoc program (K. B. Nicholas and H. B. Nicholas, Jr., Pittsburgh Supercomputing Center, Pittsburgh, Pa.). Computer-based predictions of RNA secondary structure were carried out using the MFOLD program (92) with default folding.

Detection of hypermutated molecules. Conditions for reverse transcription and PCR were set to increase the probability of amplification of putative hypermutated molecules. The RT reaction was done with avian myeloblastosis virus (AMV) reverse transcriptase (Promega) at 37°C for 45 min and then at 94°C for 5 min. This incubation temperature is that recommended for the synthesis of first-strand cDNA using random oligonucleotide primers. One microliter of RNA was thawed at 70°C for 3 min, and after cooling on ice, 8 pmol of reverse primer were added and the mixture incubated at 65°C for 5 min. A mixture of 2.5 U of AMV and 7.5 U of RNasin RNA inhibitor (Promega) was added in a final reaction volume of 20 µl. Five microliters of the first-strand reaction was used in a touchdown PCR in a volume of 50 µl with 1.25 U of Pfu DNA polymerase and 16 pmol of reverse and forward primers. The reaction conditions were as follows: denaturation hold at 95°C for 2 min, followed by 20 cycles at 95°C for 1 min, 60°C
for 30 s (starting at an annealing temperature of 60°C, with a decrease of 0.5°C per cycle), and 72°C for 2 min, and finally 10 additional cycles at an annealing temperature of 47°C. Assuming that the more-frequent mutations induced by FU are transitions of the types A→G and U→C (41, 65, 67, 72, 79), oligonucleotides with different degrees of degeneracy (which should favor hybridization with mutated templates) were synthesized. Oligonucleotides L3662F (forward) and L4260R (reverse) were modified to produce a battery of degenerate primers to amplify residues 3662 to 4260 of the L polymerase-coding region (Table 1). The cDNAs synthesized were purified with a Wizard PCR purification kit (Promega), ligated into the pGEM-T Easy vector (Promega), and used to transform E. coli DH5α; positive bacterial clones were amplified with the TempliPhi amplification kit (Amersham). Purified cDNA and molecular clones were subjected to cycle sequencing. Standard RNA consisted of an SP6 RNA polymerase runoff transcript of a molecular clone that contained the polymerase region amplified with primers L3662F and L4260R in the genomic sense. Quantification of RNAs of control and FU-treated virus populations was done by two-step RT-PCR with ThermoScript RT (Invitrogen) at 60°C for 45 min, followed by a hold at 85°C for 5 min; 2 μl of the first-strand reaction product was used in a 20-μl reaction mixture with the Fast Start DNA Master SYBR Green I kit (Roche) in a Light Cycler instrument (Roche). A standard curve employed serial 10-fold dilutions of the standard RNA from 10^{10} to 10^{3} molecules. Mutation frequencies were calculated as described above.

**RESULTS**

**Rapid FU-mediated decrease of LCMV infectivity and rapid recovery in the absence of mutagens.** Serial population passages of RNA viruses under a defined set of stable conditions generally result in fitness gain (32, 58), and lethal mutagenesis is less effective with high-fitness virus (67, 79). To investigate whether loss of infectivity of LCMV upon replication in the presence of FU was dependent on the passage number of the virus in BHK-21 cells, LCMV was serially passaged in BHK-21 cells at an MOI of 0.01 PFU/cell in the absence of FU, as detailed in Materials and Methods. Using virus from passage 1, 2, 4, 5, 6, 7, 8, or 9, an infection in the presence of 100 μg/ml FU resulted in 10^{-2} to 10^{-9}-fold decreases in infection progeny production, and the decrease did not show any trend with the passage number of LCMV in BHK-21 cells (Fig. 1A).

To study whether mutagenized populations could regain the capacity to produce progeny in the absence of FU, several FU-treated populations were allowed to replicate in the absence of the mutagen. In the first passage without FU, the capacity to produce progeny, which is taken as an estimate of relative viral fitness (33, 47), was impaired. However, two to three passages in the absence of mutagens allowed LCMV to regain the capacity to produce progeny at levels comparable to those of LCMV not subjected to mutagenic treatment (Fig. 1B). Two passages in the presence of FU led to virus extinction, as defined in Materials and Methods. In one case, extinction occurred despite an intervening recovery passage (Fig. 1B). A recovered preextinction population was driven to extinction by reiteration of the mutagenic process, and the preextinction population that resulted from this second round of mutagenesis could again attain the standard capacity to produce progeny (Fig. 1B). Thus, LCMV is very sensitive to FU-induced lethal mutagenesis, but preextinction populations maintain the potential to produce normal infectious progeny levels in BHK-21 cells upon replication in the absence of mutagen.

**Fitness variations with an invariant consensus sequence.** To study whether large fluctuations in replication capacity were associated with modification of the consensus LCMV genomic sequence, the entire genomic nucleotide sequences of the virus after one passage in the absence of drug (D1), the virus subjected to one passage in the presence of FU (F1), and the virus subjected to three subsequent recovery passages in the absence of FU (F1D3) were determined (the nomenclature of LCMV populations that defines the passage history in the presence or absence of FU is described in the legend for Fig. 1B). No differences in the consensus nucleotide sequence of the three
viruses were observed. Although some differences were seen relative to the previously published LCMV ARM 53b sequence (73–76), our sequence is in good agreement with the LCMV genomic sequence deposited in GenBank with accession number AY894816. Under the standard amplification conditions, it was not possible to determine a defined nucleotide sequence (i.e., the peak pattern suggested the presence of insertions and/or deletions) for the intergenic region of segment L (nucleotides 470 to 550). RT-PCR at a higher temperature (25, 87) in 5% dimethyl sulfoxide was used to obtain cDNA that was cloned into the pGEM-T Easy vector (Promega). The nucleotide sequences of 13 clones from viral populations D1, F1, and F1D3 indicated a remarkable genetic heterogeneity with G-rich and C-rich sequences. Despite this heterogeneity, no evidence of dominance of different subpopulations was obtained for viruses D1, F1, and F1D3 (Fig. 2A). Two of the clones harbored large deletions which likely correspond to stem-loop structures deleted during the amplification or during the sequencing reaction by polymerase slippage (71, 89); however, heterogeneity in the viral RNA cannot be excluded. We have not subjected in vitro-transcribed RNA containing this region to RT-PCR to assess the contribution to the observed heterogeneity of errors during the amplification procedure. Previous studies have reported deletions in the intergenic regions of LCMV (75) and Lassa fever virus (87).

To further ascertain that variations in fitness were not accompanied by changes in the consensus genomic nucleotide sequence, three genomic regions (nucleotides 2950 to 4240 of the L segment, encoding amino acids 985 to 1415 of protein L; nucleotides 268 to 738 of the S segment, encoding amino acids 198 to 357 of NP) were sequenced for viruses D1, F1, and F1D3 (Fig. 3). For primers without degeneracy, lowering the temperature during first-strand synthesis, followed by touchdown amplification, allowed detection of genomes with a number of mutations that remained undetected under standard, more stringent conditions of amplification. Under these new amplification conditions, the mutation frequency for the F population was 4.7-fold higher than the value obtained under standard amplification conditions whereas for the D population the mutation frequency was 2.0-fold lower (Fig. 3). Also, for the F population, the mutation frequencies scored were lower when any combination of degenerate oligonucleotides was used than with standard oligonucleotides representing the wild-type LCMV consensus sequence. Control amplifications employing similar numbers of molecules of L RNA runoff transcripts as those in the D and F populations indicate that the differences in the mutation frequency values were not due to the amplification conditions (Fig. 3). Thus, the results do not support our initial hypothesis that hypermutated molecules may be dominant during the transition into error catastrophe (41). Rather, the results suggest that mutant spectra, composed of genomes harboring modest numbers of fitness-decreasing mutations, dominate the molecular landscape in the LCMV transition into error catastrophe, supporting a lethal deflection model for LCMV extinction (40).

**DISCUSSION**

Lethal mutagenesis has been approached with a number of RNA viruses (4, 18, 19, 21, 41, 42, 46, 48, 50, 51, 53, 65, 67, 72, 77, 79) and is currently being explored as a possible new antiviral strategy to provide an alternative to the use of inhibitors of viral replication (26, 35, 38). The molecular events that mediate the critical transition into viral extinction, also termed virus entry into error catastrophe (5, 31, 60, 83), are still poorly understood, but such understanding may be crucial for finding new virus-specific mutagenic agents that, in combination with inhibitors, could lead to a clinical application. The previous studies with LCMV documented a very high sensitivity of the virus to FU in cell culture (41, 72) and prevention by FU of the establishment of a persistent LCMV infection of RAG−/− (deficient in the adaptive immune response) mice (72). This latter study constituted a proof-of-principle of the potential of mutagenesis to prevent a viral infection in vivo. Moreover, there is evidence that the antiviral activity of the nucleoside analogue ribavirin, currently licensed for the treatment of several viral diseases (68, 81), might be mediated in some cases by its mutagenic action (4, 18–21, 39, 45, 53, 66, 88).
FIG. 2. Nucleotide sequence and predicted secondary structure of an LCMV genomic region belonging to the L intergenic region. (A) Nucleotide sequence of residues 470 to 550, corresponding to the intergenic L region of clones from populations D1, F1, and F1D3. Sequences of clones are aligned under our consensus LCMV ARM 53b; at the bottom, the sequence previously published (termed here ARM 53b*) (76) is given. Our sequence is in agreement with a recent sequence (deposited in GenBank with accession number AY894816), determined independently of ours. Deletions introduced for best alignment of heteropolymeric stretches are indicated by dashes. (B) Predicted secondary structure of intergenic L residues 365 to 466 of Lassa fever virus and predicted secondary structures of residues 470 to 550 of LCMV ARM 53b, clones D1-2, F1-4, and F1D3-4 analyzed in the present study. Procedures for nucleotide sequence determination and secondary-structure prediction are described in Materials and Methods.
LCMV shows higher sensitivity to FU mutagenesis than foot-and-mouth disease virus (FMDV) in cell culture when the two viruses are subjected to similar FU doses (41, 65, 67, 72, 79). The basis for this difference is not known, but it may relate either to the size of the essential polymerase gene, which occupies 62.6% and 17.4% of the total genetic information of LCMV and FMDV, respectively; to a larger number of essential genomic sites susceptible to FU-induced mutations in LCMV than in FMDV; or to other replicative differences between the two viruses still to be elucidated (different affinities of the two polymerases for FU-triphosphate, etc.).

The comparison of complete genomic LCMV sequences reported here has indicated that neither an abrupt decrease in infectivity associated with FU treatment nor the ensuing recovery of the capacity to produce infectious progeny upon replication in the absence of mutagen modified the consensus nucleotide sequence of the virus. Likewise, no variations were detected among LCMV populations subjected to one or several rounds of replication in the absence or presence of FU, or among populations that underwent alternations between the two passage regimens. In these cases the consensus sequences analyzed involved three genomic regions, one of which was a part of the polymerase (L) that included premotif A and motifs A to E (11, 24, 52, 70, 86, 87). The possibility that FU, in addition to its mutagenic activity documented by increases in mutant-spectrum complexity, could exert some mutagenesis-independent, inhibitory activity (i.e., directly on the viral polymerase or indirectly through alteration of nucleotide pools, etc.) on LCMV replication cannot be excluded. This would mean that FU could contribute to decreases in viral load, thereby favoring mutagenesis-induced extinction, as documented with FMDV (65–67).

It could be argued that fitness variations despite an invariant consensus sequence may occur without modification of the mutant spectrum, simply by an increase of dominance of a preexisting subset of genomes with the same consensus sequence. A static mutant spectrum is highly unlikely given the number of LCMV progeny per cell (Fig. 1), since it would imply mutation rates much lower than those that have been calculated for RNA viruses (29). Low mutation rates are not supported by the estimates of the frequencies of cytotoxic T-lymphocyte (CTL)- and antibody-escape mutants in LCMV populations (1, 15, 43, 49; reviewed in reference 78). Fitness variations without modification of consensus genomic sequences have been reported for vesicular stomatitis virus clones in the process of fitness recovery after bottleneck passages (59).

The L RNA intergenic region may function as a transcription termination signal, as previously suggested for LCMV (69, 74), other arenaviruses (25, 34, 44), and the stem-loop present in the S RNA intergenic region of LCMV (55), Pichinde virus (7), and Lassa fever virus (8, 16). RNA secondary-structure predictions suggest that the L RNA intergenic region of LCMV ARM 53b forms a stem-loop stabilized by 15 GC pairs (Fig. 3B), similar to a loop predicted for the L RNA intergenic regions of Tacaribe and Lassa fever virus (25, 34, 44). We have not distinguished whether the nucleotide sequence heterogeneity seen in the L intergenic region (Fig. 3A) corresponds to a heterogeneity present in the LCMV used (perhaps with func-
tional implications) or is due to the production of altered copies during RT-PCR, or both.

Although it cannot be excluded that highly mutated or hypermutated RNA molecules derived from LCMV RNA may be present during the transition to error catastrophe, attempts using standard and multiply substituted oligonucleotide primers have failed to reveal the dominance of such genomes (Table 1 and Fig. 3). In the amplification with primers L3662F and L4260 RY4, one sequence was found that had nine mutations, of which six were U→C transitions and the others were A→G, C→A, and G→A. The mutation frequency for this molecule relative to the consensus sequence (1.6 × 10⁻²) would lie within the lower limit in the range of 3.3 × 10⁻² to 1.0 × 10⁻¹ mutations per nucleotide for the U→C (or A→G) hypermutation, previously characterized in genomes of measles virus, human parainfluenza virus 3, vesicular stomatitis virus, and Rous-associated virus, which has been related to posttranscriptional unwinding and mutagenic activities (13, 36, 54, 56, 57, 61). Hypermutated genomes have not been identified, either, among multiple molecular clones copied from preextinction populations of FMDV (4, 67, 79). In the FU-mediated transition to extinction of LCMV in the course of a persistent infection of BHK-21 cells, decreases in viral infectivity preceded decreases in viral RNA levels (40). The specific infectivity of LCMV reached values 10⁻² to 10⁻³-fold lower than those for LCMV from parallel cultures not subjected to FU mutagenesis (40). Similar decreases in specific infectivity were noted for mutagenized FMDV during persistent or cytolytic infections (4, 37, 65, 66). These quantifications of specific infectivity, together with the complexity of mutant spectra of mutagenized LCMV and FMDV populations (4, 37, 40, 41, 65–67, 72, 79), suggest that rather than requiring extensive mutagenesis, virus extinction may be mediated by the presence of defective genomes (which have been termed “defectors” in several studies of RNA virus evolution [85, 90]). Therefore, the present evidence suggests that low-activity mutagens capable of being incorporated specifically by viral polymerases may be sufficient to enrich the defector population to a level at which it may interfere with standard genome replication. A likely molecular mechanism mediating this type of interference is the effect of multiple, low-frequency dominant-negative mutants generated in mutagenized virus populations, an extension to an intrapopulation level of the mechanisms of interference between different viruses, well documented in classical viral genetics (reviewed in references 2, 17, and 91). Because such interfering genomes—due to their partial or complete defectiveness—cannot dominate the population, no alteration of consensus sequences is observed despite profound fitness losses. Thus, the results reported here support a lethal defection model for the mutagenesis-induced genetic meltdown that results in viral extinction (40). Treatment with the mutagenic nucleoside analogue ribavirin of cell cultures infected with poliovirus or Hantaan virus resulted in decreases in infectivity that preceded decreases in viral RNA levels (19–21, 77), again suggesting a role of mutagenesis-induced defective genomes that maintain their replication competence prior to detectable losses of infectivity.

The results presented here underscore the relevance of mutant spectra in determining virus behavior, illustrated here by the capacity of LCMV to either decrease or increase its fitness without any alteration of the consensus nucleotide sequence. This observation must be added to results with several other virus-host systems showing suppressive or modulating effects of mutant spectra on specific types of variants with superior fitness (9, 14, 22, 37, 84).

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