Nonsense mutations inhibit splicing of MVM RNA in cis when they interrupt the reading frame of either exon of the final spliced product

Lisa Kay Naeger, Robert V. Schoborg,1 Qihong Zhao, Gregory E. Tullis, and David J. Pintel 2

Department of Molecular Microbiology and Immunology, University of Missouri–Columbia, School of Medicine, Columbia, Missouri 65212 USA

mRNAs R1 and R2 of the autonomous parvovirus minute virus of mice (MVM), which encode the viral nonstructural proteins NS1 and NS2, respectively, are processed in an ordered splicing pathway in which R2 is generated from mature spliced R1. Introduction of translation termination signals into these genes alters the processing of these RNAs; there is a significant (up to fourfold) increase in the accumulated steady-state levels of R1 relative to R2, when compared with wild-type levels, although the total accumulated levels of R1 plus R2 remain the same. The increase in accumulated R1 relative to R2 in mutant infected or transfected murine cells is independent of RNA stability and transport and decreases, in a polar manner, with the distance of the inserted termination signal from the shared initiation codon for NS1 and NS2 at nucleotide 260. The increased ratio of R1 to R2 is a consequence of the artificially introduced translation termination signals acting in cis rather than in the absence of a functional viral gene product. These mutations have an effect when they interrupt previously open reading frames in either exon of the spliced product R2. Nonsense mutations that are located in the second exon of R2 inhibit splicing of R1 to R2 only when they interrupt an open reading frame (ORF) that has the potential, after normal splicing, to be joined in-frame with the initiating AUG. These results suggest that nonsense mutations inhibit splicing of R1 to R2 by influencing the mechanism by which exons are defined in murine cells.

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It has been known for some time that in various prokaryotic systems and in yeast, premature translation termination of certain mRNAs can result in their destabilization (Adhya and Gottesman 1978; Losson and Lacroute 1979; Kennell 1986; Nilsson et al. 1987; Brown 1989; Peltz et al. 1990). Mutations that terminate translation prematurely in higher eukaryotes have also been shown in some cases to act in cis to affect reduced cytoplasmic mRNA levels. Although the precise mechanism of this effect is still unclear, primary alterations in RNA synthesis and stability (Maquat et al. 1981; Graves et al. 1987; Arrigo and Beemon 1988; Peltz et al. 1990; Barker and Beemon 1991) and RNA processing and transport (Humphries et al. 1984; Takeshita et al. 1984; Baumann et al. 1985; Baserga and Benz 1988; Daar and Maquat 1988, Urlaub et al. 1989; Cheng et al. 1990; Peltz et al. 1990) have been proposed. Here, we report that introduction of translation termination signals into the open reading frames (ORFs) encoding the nonstructural genes NS1 and NS2 of the autonomous parvovirus minute virus of mice (MVM) act in cis to alter the processing of viral RNA in the nucleus of murine cells.

MVM, a member of the autonomously replicating subgroup of parvoviruses (for review, see Cotmore and Tattersall 1987, Berns 1990), is organized into two overlapping transcription units that produce three major transcript classes, all of which terminate near the right-hand end of the 5-kb viral genome [Pintel et al. 1983; Astell et al. 1986; Clemens and Pintel 1987] [Fig. 1A]. Transcripts R1 (4.8 kb) and R2 (3.3 kb) are generated from a promoter (P4) at viral map unit 4 [Pintel et al. 1983] and encode the viral nonstructural proteins NS1 (83 kD) and NS2 (24 kD), respectively [Cotmore and Tattersall 1986]. The R3 (3.0 kb) transcripts, generated from a promoter (P38) at map unit 38, encode the overlapping viral capsid proteins VP1 and VP2 from alternatively spliced mRNAs, using the ORF in the right half of the genome [Jongeneel 1986, Labieniec-Pintel and Pintel 1986]. Polyadenylation of MVM RNAs precedes splicing [Clemens and Pintel 1988]. Three splicing patterns are used to excise a small

1Present address: Division of Comparative Medicine, Johns Hopkins University School of Medicine, Baltimore, Maryland 21205 USA.
2Corresponding author.
introns at map units 44–46, which is common to all three transcript classes, resulting in nine different spliced MVM mRNAs species [Jongeneel et al. 1986; Morgan and Ward 1986; Clemens et al. 1990; Cotmore and Tattersall 1990]. The R2 transcripts are additionally spliced between map units 10 and 39 [Pintel et al. 1983; Cotmore and Tattersall 1986; Jongeneel et al. 1986; Clemens et al. 1990].

There is a regulated, temporal phasing of the accumulation of MVM RNA and protein: The products of the P4 promoter [the nonstructural genes] are produced before the products of the P38 promoter [the capsid genes] [Clemens and Pintel 1988; Cotmore and Tattersall 1990; Schoborg and Pintel 1991]. Later in infection, the products of the P38 promoter become predominant [Pintel et al. 1983; Clemens and Pintel 1988; Schoborg and Pintel 1991]. The steady-state levels of the nonstructural proteins, which are of critical importance for efficient viral replication, are also controlled at the level of RNA processing [Schoborg and Pintel 1991]. Analysis of the accumulation of MVM RNA in highly synchronized cells has shown that R2 accumulates faster and to a greater amount than R1 in both total and cytoplasmic RNA throughout infection, even although they are generated from the same promoter and have similar stabilities [Schoborg and Pintel 1991]. Detailed characterization of the accumulation of the various spliced and unspliced forms of MVM RNA has suggested a preferred [but not obligatory] order of splicing in which the R2 transcripts are generated from polyadenylated, spliced R1 RNAs [Clemens and Pintel 1988; Schoborg 1991; Schoborg and Pintel 1991; R.V. Schoborg, G. Tullis, Q. Zhao, and D. Pintel, unpubl.; R.V. Schoborg, Q. Zhao, and D.J. Pintel, in prep.] (Fig. 1B). Consistent with this model, cDNA constructs in which the small intron at map units 44–46 has been removed are impaired severely in further splicing of the large intron between map units 10 and 39 to form R2 [Schoborg 1991; R.V. Schoborg, G. Tullis, Q. Zhao, and D. Pintel, unpubl.; R.V. Schoborg, Q. Zhao, and D.J. Pintel, in prep.]. The ratio of accumulated R1 to R2 is dependent on the percentage of R1 that undergoes subsequent removal of the large intron [Clemens and Pintel 1988; Schoborg 1991; Schoborg and Pintel 1991; R.V. Schoborg, G. Tullis, Q. Zhao, and D. Pintel, unpubl.; R.V. Schoborg, Q. Zhao, and D.J. Pintel, in prep.]. The regulation of this event is critical in determining the steady-state levels of NS1 and NS2 [Schoborg and Pintel 1991] and, therefore, in determining the optimal balance between the essential roles of these proteins in viral replication and cytotoxicity [Cotmore and Tattersall 1987; Berns 1990].

The regulation of splicing of R1 to form R2 is mediated, at least in part, in cis by a nonconsensus donor (AA/GCAAG) at nucleotide 514 for the R2 large intron. When this donor is changed to consensus (AG/GTAAG), the majority of R1 is processed further to R2 [Schoborg 1991; R.V. Schoborg, G. Tullis, Q. Zhao, and D. Pintel, unpubl.; R.V. Schoborg, Q. Zhao, and D.J. Pintel, in prep.].

To determine whether any MVM gene products were involved in the regulation of the processing of R1 to R2, we examined a set of mutants in which translation termination signals were introduced into the coding regions of the nonstructural genes of MVM. We noticed that for several of these mutants the ratio of accumulated R1 relative to R2 was increased significantly compared with wild type, although the total accumulated levels of R1 plus R2 remained the same. The increase in accumulated R1 relative to R2, in mutant infected or transfected murine cells, is independent of RNA stability and transport, and the magnitude of the increase is correlated in a polar manner with the distance of the inserted termination codon from the shared initiation codon for NS1 and NS2 at nucleotide 260. The defect in processing of R1 to R2 is not complemented in trans by the viral nonstructural gene products. Mutational analysis has demonstrated that the increase in accumulated R1 relative to R2 is a consequence of cis-acting signals rather than the absence of a functional viral gene product. These cis-acting termination signals produce their effect when they interrupt previously open reading frames in either exon of R2. Nonsense mutations that are localized in the second exon of R2 inhibit splicing of R1 to R2 only when they interrupt an ORF that has the potential to be joined in-frame to the initiating AUG after normal splicing. These results suggest that nonsense mutations inhibit the splicing of R1 by influencing the mechanism by which exons are defined in murine cells. Because the processing of MVM RNAs is relatively simple and well defined, it provides a good model in which to study this association.

Results

Artificially introduced ochre termination codons in the nonstructural genes of MVM result in greater than wild-type accumulated ratios of R1 relative to R2

To assess the possible role of the MVM nonstructural proteins during the regulated accumulation of R1 and R2, a 16-nucleotide palindromic linker, containing ochre termination signals in all three reading frames, was inserted into pLLP-ΔH1 at nucleotide 385. This insertion terminates both nonstructural genes NS1 and NS2, which share ORF 3 in this region [Fig. 1A]. When the mutant plasmid [p385UTT] was transfected into murine A9 fibroblasts, very little P38-generated R3 RNA was detected, as expected [Fig. 2, lane 8], because the P38 promoter is dependent on NS1 for activation [Rhode 1985; Doering et al. 1988; Schoborg and Pintel 1991]. In addition, a significantly higher ratio of accumulated R1 to R2 was seen compared with parallel transfections of wild-type plasmids [Fig. 2A, lanes 2, 8; Table 1]. Extensive characterization of the rates and levels of accumulation of wild-type MVM RNAs during highly synchronous infection of murine A9 cells had demonstrated previously that MVM R2 accumulates faster and to a greater amount than R1 in both total and cytoplasmic RNA fractions throughout infection [Schoborg and Pintel 1991]. In contrast, p385UTT, in which both nonstructural proteins are prematurely terminated, generated approxi-
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![Diagram of MVM transcript classes and protein-encoding ORFs](image)

**Figure 1.** (A) Genetic map of MVM. The three major MVM transcript classes and protein-encoding ORFs are shown. Locations of most of the mutants used in this study are diagramed on the top line. The *bottom* line shows nucleotide locations and the three probes used for RNase protection assays as described in Materials and methods. There are no significant ORFs in MVM other than those indicated. (B) Splicing pathway of the P4-generated RNAs R1 and R2. The nonconsensus splice donor site for R2 at nucleotide 514 is indicated (ncD).

mately fourfold more viral R1 than R2 after transfection. In this and all subsequent analyses, all spliced and unspliced forms of R1 and R2 have been included in the quantitation.

Nonsense mutations were also introduced exclusively into the NS1 ORF at nucleotides 652 (p652[T4], p652UTT[T4], p652UTT[MB]), 762 (p762UTT), and 1081 (p1081UTT) (Fig. 1A). These mutant plasmids generated essentially wild-type ratios of accumulated R1 relative to R2 [Fig. 2A, lanes 3–7; Table 1], although transcription from P38 was abolished, as expected.

We then examined the relative RNA levels generated by an MVM mutant that exclusively affects NS2 (NS2-2018) [Naeger et al. 1990]. In this mutant, NS2 is truncated as a consequence of an ochre termination signal that was introduced by site-directed mutagenesis into the unique second exon of NS2 in ORF 2 at nucleotide 2018 [Fig. 1A] [Naeger et al. 1990]. Plasmids containing this mutant generated significantly increased levels of accumulated R1 relative to R2 when compared with wild type after transfection into A9 cells [Fig. 2, lane 9; Table 1], although the increase was not as great as that seen for p385UTT.

Virus containing nonsense mutations in NS2 display an increased ratio of accumulated R1 to R2 throughout synchronous infection, which was exhibited in a polar manner.

We have previously characterized a series of MVM NS2 mutants, including NS2-2018, in which ochre termination signals have been introduced at several positions in ORF 2 in the second exon of R2 [Naeger et al. 1990]. These mutants, in which NS1 is unaffected, are defective for single-strand progeny DNA accumulation and efficient virus production on the normal tissue culture host, murine A9 cells [Fig. 3]. They can, however, be propagated on a number of other nonmurine cell types [Naeger et al. 1990], permitting the generation of NS2 mutant virus stocks. NS2-2018 virus generated the same increased levels of R1 relative to R2 throughout infection of highly synchronous murine A9 cells, as seen after transfection of the parent mutant plasmid [Fig. 3]. These results confirm that the increased accumulated ratio of R1 to R2 described above for mutant plasmids is not unique to the transfection process. In addition, examination of the RNAs generated from four NS2 mutants that terminate at different points in the second exon of NS2 [NS2-2018, NS2-2159, NS2-2268, NS2-2381; see Fig. 1A] demonstrated that the increased R1 to R2 ratios detected are related, in a polar fashion, to the distance of the mutation from the initiating AUG for NS1 and NS2 at nucleotide 260 (and from the 5' end of R1 and R2) [Fig. 4]. The relative increase of accumulated ratios of R1 to R2, compared with wild type, seen during NS2-2018 infection, was greater than that for NS2-2159, whereas mutants NS2-2268 and NS2-2381 exhibited wild-type accumulated ratios of R1 to R2 [Fig. 4].

Although the accumulated levels of R1 relative to R2 after infection or transfection of NS2-2018 was significantly increased compared with wild type, the accumulated levels of total P4-generated RNA (R1 plus R2) remained the same, compared with P38-generated R3 [Fig. 3, data not shown].
The increase in the accumulation of R1 relative to R2 is not due to alterations in RNA stability

The altered ratio of accumulated R1 to R2 that we observed after transfection of mutant p385UTT and pNS2-2018 plasmids was not the result of altered stability of any of the viral RNAs. Characterization of the persistence of the individual viral RNAs in the presence of the transcription inhibitors actinomycin D (Fig. 5, lanes 2–16) or the adenosine analog 5,6-dichloro-1-β-D-ribofuranosylbenzimidazole.
Table 1. Ratios of accumulated R1/R2 following transfection of mutants

| Construct         | Relevant genotype | R1/R2a |
|-------------------|-------------------|--------|
| pLLPAH            | NS1 (+)/NS2 (+)   | 0.48 ± 0.04 |
| p385UTT           | NS1 (+)/NS2 (+)   | 1.83 ± 0.44 |
| p652UTT (T4)      | NS1 (+)/NS2 (+)   | 0.54 ± 0.32 |
| p652UTT (MB)      | NS1 (+)/NS2 (+)   | 0.50 ± 0.20 |
| pNS2-2018         | NS1 (+)/NS2 (+)   | 1.10 ± 0.19 |

aThe ratios presented are the average of the following number of RNase protection assays using probe A, as described in Materials and methods, and include 95% confidence limits. pLLPAH, 17; p385UTT, 7; p1081UTT, 5; p762UTT, 5; p652UTT (T4), 5; p652UTT (MB), 5; p652 [T4], 5; pNS2-2018, 10.

An increased ratio of accumulated R1 to R2 is seen in both nuclear and cytoplasmic RNAs

The availability of NS2-2018 mutant virus permitted efficient characterization of the accumulated ratios of R1 to R2 in both nuclear and cytoplasmic fractions of infected cells. The accumulated levels of R1 relative to R2 generated during NS2-2018 infection (Fig. 6, lanes 2–4) is significantly increased, compared with wild type (Fig. 6, lanes 6–8), in both nuclear and cytoplasmic fractions (Fig. 6, cf. lane 2 with lanes 3 and 4, and lane 6 with lanes 7 and 8; Table 2), although the total accumulated levels of R1 plus R2 remain the same, as determined by comparison to P38-generated R3 as an internal control (Table 2). This suggests that the altered accumulation of R1 to R2 is not primarily related to RNA transport of R1 or R2 to the cytoplasm and that R2 is not specifically degraded in the nucleus. This is also consistent with results demonstrating no decrease in the stability of R2 in total RNA. The amount of nuclear RNA is a significant portion of MVM-infected total RNA (Clemens and Pintel 1988; Schoborg and Pintel 1991); therefore, specific degradation of R2 in the nucleus would also be apparent in total RNA (see Fig. 5).

Because accumulated ratios of R1 to R2 generated during NS2-2018 infection are higher in the cytoplasmic fraction than in the nuclear fraction, we cannot exclude an additional effect on the transport of MVM RNA. In the wild-type infection, however, steady-state, accumulated ratios of R1 to R2 are lower in the cytoplasmic fraction than in the nuclear fraction. Taken together, these results are consistent with a model in which the rate of specific RNA transport is proportional to its concentration in the nucleus; therefore, such differences would be correspondingly amplified in cytoplasmic RNA compared with nuclear RNA.

The increased ratio of accumulated R1 to R2 is a consequence of a cis-acting signal in the RNA not the absence of a wild-type viral gene product

The analysis described up to this point demonstrates that the observed increased ratio of accumulated R1 to R2 was not the result of altered stability or transport of MVM RNA but, rather, the accumulated levels of R1 increased at the expense of R2. This suggested that the splicing of R1 to R2 was being inhibited. The data also suggested that perhaps the MVM NS2 protein might play a critical role in the regulation of processing of R1 to R2. Attempts to reverse the increased accumulated ratio of R1 to R2 generated from p385UTT, however, by complementation with NS2 in trans were unsuccessful. In these experiments, over a broad range of plasmid concentrations, pSVNS1/NS2, a plasmid that expresses the MVM nonstructural genes from the SV40 early promoter (Clemens et al. 1990), showed no effect on the accumulated ratio of R1 to R2 generated by the mutant p385UTT test plasmid (data not shown); even the P38 promoter of that plasmid was stimulated strongly by the nonstructural genes in trans. Attempts to complement p385UTT with constructs expressing an MVM NS2 cDNA (Clemens et al. 1990) were also unsuccessful [data not shown].

These results implied that perhaps the ochre termination signals, artificially introduced into MVM RNA, might act in cis, as has been described previously in

Figure 2. Artificially introduced termination signals interrupting MVM nonstructural genes affect relative accumulated RNA levels.

(A) RNase protection analysis of 40 μg of total RNA isolated from A9 cells 48 hr post-transfection with 20 μg of mutant plasmid DNA per dish is indicated. The final two lanes are RNase protection assays of 10 and 20 μg of total RNA taken from MVM-infected A9 cells, serving as a control to assure that protection assays were carried out in probe excess. Locations of mutations are indicated in Fig. 1A. RNase protection probe A (nucleotides 1854–2378) was used. The identities of the protected bands are diagramed as follows. (B) Unspliced and alternative splicing patterns of the R1, R2, and R3 transcripts are shown, and the corresponding RNase-protected regions, designated in bold, after hybridization to SP-6 generated antisense probe A (MVM nucleotides 1854–2378). The acceptor site for the large R2 splice lies at nucleotide 1990, and the initiation site for the R3 RNAs lies at approximately nucleotide 2005. Three splicing patterns are used to excise the small intron at map units 44–46 common to all three transcript classes. The major pattern (found in ~80% of molecules of all transcript classes) joins nucleotide 2280 to 2377. The minor pattern (found in~15% of molecules of all transcript classes) joins nucleotide 2317 to 2399. A rare pattern, present in <5% of MVM-spliced RNA joins nucleotide 2280 to 2399. M represents protection by those molecules that use the predominant splice donor at nucleotide 2280; m represents protection by those molecules that use the less frequently used donor at nucleotide 2317.

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Figure 3. Mutant NS2-2018 demonstrates increased accumulated levels of R1 relative to R2 throughout infection of murine fibroblasts. RNase protection analysis of total RNA isolated from NS2-2018 (left) and wild-type-infected A9 murine fibroblasts (right). Probe A-2018 (nucleotides 1854–2378) was used to protect 10 μg of total RNA isolated from highly synchronized A9 cells infected at m.o.i. 2 with NS2-2018 virus (left). Probe A (nucleotides 1854–2378) was used to protect 10 μg of total RNA isolated from highly synchronized A9 cells infected at m.o.i. 1 with wild-type virus (right). RNA was collected at 4, 8, 12, 16, and 24 hr postrelease, as indicated. Mock-infected RNA is protected in the lane indicated. The identities of the protected bands are as described in Fig. 2B. The average relative accumulated R1 versus R2 ratios taken from three separate infections with NS2-2018 and wild-type-infected A9 cells for each time point is shown at the bottom of each corresponding lane.

other systems. Therefore, we constructed mutants p260(T4) and p260UTT(T4) in which the initiating AUG codon for both NS1 and NS2 at nucleotide 260 was removed, thereby preventing their expression. These mutants, which express neither NS1 nor NS2, generated essentially wild-type levels of accumulated R1 relative to R2 [Fig. 7]. The absence of functional nonstructural proteins in these mutants was demonstrated by the absence of the P38-generated viral R3 RNA [Fig. 7], which in parallel experiments could be stimulated in trans by pS-VNS1/2 [data not shown]. These experiments established that neither NS1 nor NS2 participates in the regulation of the accumulated levels of R1 relative to R2 and suggested that the ochre termination signals acting in cis are responsible for inhibition of splicing of R1 to R2.

The increased ratio of accumulated R1 relative to R2 requires that the nonsense codon interrupt a previously open reading frame

Because the observed increase in the accumulated ratio of R1 to R2 (described above) was caused by a cis-acting effect, we expected that alteration of the processing of R1 to R2 might require that the termination signal interrupt a previously open reading frame. To examine this hypothesis and to test whether the interrupted ORF had to

Figure 4. Examination of RNAs generated from NS2 mutants demonstrates that the increased ratios of R1 to R2 are related in a polar fashion to the distance from the initiating AUG. Shown is RNase protection analysis of RNA isolated from murine A9 cells infected with NS2 mutants at a m.o.i. 1. Probe B (nucleotides 146–652) was used to protect 10 μg of total RNA taken at 22 hr postinfection with NS2-2018, NS2-2159, NS2-2268, NS2-2381, and wild type (lane 6), as indicated. Mock-infected RNA is protected in the first lane, and 20 μg of wild-type total RNA is protected in the lane indicated, as a probe excess control. The bands representing protection of probe B by R1 and R2 (451 and 313 nucleotides, respectively; see Fig. 1A) are indicated at left. Wild-type-infected RNA taken at 16 hr postinfection was also protected with probe A (nucleotides 1854–2378), as indicated, and the identities of these protected bands [right] are as described in Fig. 2B. The accumulated ratios of R1 to R2 averaged from three different infections are shown for each mutant at the bottom of the corresponding lane. (*) Average of three experiments.
be one that could potentially be joined in-frame to the initiating AUG, we constructed and analyzed the phenotype of a set of mutants based on mutant Tb60, which contains a 2-nucleotide insertion at the TaqI site at nucleotide 2073.

The Tb60 insertion shifts the NS2 reading frame in exon 2 of R2 from ORF 2 into ORF 3, which is normally used to encode the carboxyl terminus of NS1 (Fig. 8). The primary amino acid sequence of the NS2 gene product is therefore altered for most of exon 2 without introducing a nonsense codon until nucleotide 2280—beyond the point where nonsense codons inhibit the splicing of R2 [see Fig. 4]. After transfection into A9 cells, Tb60 had a ratio of R1 to R2 similar to wild type. These results support our conclusion from experiments described above that the NS2 gene product is not required to accumulate normal ratios of R1 to R2.

If the artificially introduced cis-acting signal responsible for the increased accumulated ratio of R1 to R2 must interrupt a previously open reading frame and if that reading frame must have the potential to become in-frame with the initiating AUG after splicing (Tb60-2087), the accumulated ratios of R1 to R2 would again be expected to be increased. The accumulated RNAs generated by these mutants are consistent with these predictions (Fig. 8) and strongly suggest that the mutant phenotype is dependent on the interruption of a previously open reading frame that must have the potential to become in-frame with the initiating AUG after splicing.

For Tb60, which generates a wild-type ratio of accumulated R1 relative to R2, the 2-nucleotide insertion at nucleotide 2073 also shifts ORF 3 to ORF 1, bringing a normally unused amber codon at nucleotides 2086–2088 into frame in NS1 [Fig. 9]. Although NS1 is terminated—as confirmed by the absence of P38 transcription from this mutant—the accumulated ratios of R1 to R2 remain like wild type, presumably because either the amber codon terminates the NS1 ORF too far downstream from the initiating AUG at nucleotide 260 or because nonsense codons that interrupt previously open reading frames in R1 have no effect. Mutant X27, a 4-nucleotide insertion at the Xhol site at nucleotide 2072, shifts ORF 2 to ORF 1, such that the same amber codon at nucleotides 2086–2088 now becomes in-frame in NS2 [Fig. 9]. Splicing of R2 also brings the NS2 in-frame amber codon much closer to the initiating AUG at nucleotide 260. In contrast to Tb60, X27 has a greatly increased accumulated R1 to R2 ratio [Fig. 9]. This experiment confirms that the position of the in-frame termination signal is of critical importance to the accumulated R1 to R2 ratio and demonstrates that interruption of a previously open reading frame by an amber codon can also affect MVM RNA processing in murine cells.

Results with the NS2 and Tb60 sets of mutants and mutant X27 demonstrate that the relative accumulation

Figure 5. Effects on RNA processing are independent of stability. Shown is RNase protection analysis using probe A (nucleotides 1854–2378), of 40 μg of total RNA isolated from A9 cells transfected with pLLP-ΔH, pNS2-2018, p385UTT, as indicated, mock-transfected [lanes 1, 5], or 10 μg of total RNA from A9 cells infected with MVM at m.o.i. 1 [lanes 12–14]. Forty-eight hours after transfection (or 9 hr after infection of highly synchronous cells), cells were treated with 40 μg/ml of actinomycin D and RNA was collected 0.5 hr [lanes 2, 6, 9, 12], 3.5 hr [lanes 3, 7, 10, 13], and 6.5 hr [4, 8, 11, 14] later. Alternatively, cells transfected with p385UTT were treated with 40 μg/ml of DRB and collected 0.5 hr [lane 17] or 3.5 hr [lane 18] later. Lanes 15 and 16 are protections with 10 and 20 μg of total RNA taken from untreated MVM-infected A9 cells serving as a control to ensure that assays were carried out in probe excess. The identities of the bands are as described in Fig. 2B. (*) Undigested probe. Unspliced R1 and unspliced R3 are processed to mature forms within the first 0.5 hr after drug addition (Schoborg and Pintel 1991).
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Figure 6. Increased ratios of accumulated R1 relative to R2 are seen in both nuclear and cytoplasmic RNA fractions. Shown are nuclear and cytoplasmic RNA fractions from NS2-2018 (lanes 1–4) and wild-type (lanes 5–8) infections of A9 murine fibroblasts analyzed by RNase protection analysis. Probe A-2018 (nucleotides 1854–2378) was used to protect 2.5 µg of nuclear RNA (lane 2), 10 µg of cytoplasmic RNA (lane 3), or 20 µg of cytoplasmic RNA (lane 4) [as a probe excess control], from NS2-2018-infected A9 cells or 20 µg of mock-infected RNA (lane 1). Probe A (nucleotides 1854–2378) was used to protect 2.5 µg of nuclear RNA (lane 6), 10 µg of cytoplasmic RNA (lane 7), or 20 µg of cytoplasmic RNA (lane 8) [as probe excess control] from wild-type-infected A9 cells, or 20 µg of mock-infected RNA (lane 5). The identities of the protected bands are as described in Fig. 2B. The average accumulated ratios of R1 versus R2 from nuclear and cytoplasmic fractions in this experiment are shown at the bottom of respective lanes. Nuclear RNA contains unspliced R1 and R3, which is absent from cytoplasmic RNA, however, there is no detectable unspliced R2 [molecules that have lost the large intron between nucleotides 514 and 1989 but not the small intron at map units 44–46] during MVM infection (Schoborg and Pintel 1991). Previous characterization of MVM RNA generated throughout infection has shown that MVM RNA in the nucleus comprises a significant percentage of total MVM RNA, as measured by the abundance of unspliced R1 and R3 (Clemens and Pintel 1988; Schoborg and Pintel 1991). In comparison with numerous other experiments, unspliced R1 is underrepresented in the nuclear fraction of this sample of NS2-2018-infected RNA. This was not a reproducible result and does not alter the total increased ratio of R1 to R2 seen for NS2-2018 infections.

Discussion

The 3.3-kb R2 mRNA of MVM is generated by further splicing of polyadenylated 4.8-kb R1 mRNA (Schoborg 1991, Schoborg and Pintel 1991, R.V. Schoborg, G. Tullis, Q. Zhao, and D. Pintel, unpubl.; R.V. Schoborg, Q. Zhao, and D.J. Pintel, in prep.) [Fig. 1B]. Regulation of the splicing of R1 to R2 is mediated, at least in part in cis, by the existence of a nonconsensus donor site for the large splice (nucleotides 514–1989) of R2 (Schoborg 1991, Schoborg and Pintel 1991; R.V. Schoborg, G. Tullis, Q. Zhao, and D. Pintel, unpubl.; R.V. Schoborg, Q. Zhao, and D.J. Pintel, in prep.). Throughout wild-type MVM infection, the relative accumulated levels of R1 and R2 are a critical parameter in determining the steady-state levels of NS1 and NS2 (Schoborg and Pintel 1991) and the optimal balance between the essential roles of these proteins in viral replication and cytotoxicity.

Our experiments show that nonsense codons artificially introduced into the nonstructural genes of MVM result in a significantly increased ratio of accumulated viral R1 relative to R2. This increase in accumulated R1 relative to R2 is independent of RNA stability and transport and is mediated by cis-acting nonsense codons rather than by the absence of a functional viral gene product.

Although termination signals that interrupt MVM ORFs significantly increase the accumulated ratio of R1 relative to R2, the total accumulated levels of R1 plus R2 remain the same. This is shown clearly for mutant NS2-2018, where the viral P38 products (R3 RNAs) can be compared as an internal control. In NS2-2018 infection or transfection of A9 cells, the accumulated levels of the P4 products (R1 plus R2) relative to R3 remain the same in both nuclear and cytoplasmic fractions; only the ratio of R1 to R2 has changed (Table 2). This suggests that nonsense codon mutations act in cis to inhibit the splicing of R1 to R2, rather than to cause a decrease in R2 stability or affect transport of R1 or R2 out of the nucleus.

Nonsense mutations in the second exon of R2 can inhibit splicing of R1 to R2 but only when they interrupt ORFs that have the potential to be joined in-frame with the initiating AUG after normal splicing. In wild-type virus, there are two ORFs in the second exon of R2. A nonsense mutation at nucleotide 2159 [NS2-2159], which interrupts the ORF that is joined in-frame to the initiating AUG at nucleotide 260, inhibits splicing of R1 to R2. In mutant Tb60-2159, in which a frameshift at nucleotide 2073 shifts the nonsense mutation at nucleotide 2159 into an ORF that can no longer be joined in-frame to the initiating AUG, splicing from R1 to R2 is unaffected. When a nonsense mutation is introduced into the ORF in Tb60, which now can become in-frame with the initiating AUG (Tb60-2087), splicing is again inhibited.

Similar to other observations of this type (Takeshita et al. 1984; Baumann et al. 1985; Baserga and Benz 1988; Urlaub et al. 1989; Cheng et al. 1990), the magnitude of the increased accumulation of R1 relative to R2 is re-
Table 2. Accumulated R1–R2 and \((R1 + R2)/R3\) ratios in total, nuclear and cytoplasmic RNA generated by NS2-2018 and wild-type MVM

|          | RNA       |        |        |
|----------|-----------|--------|--------|
|          | total      | nuclear | cytoplasmic |
| NS2-2018 | R1/R2     | 1.32 ± 0.25 | 1.30 ± 0.27 | 2.03 ± 0.36 |
|          | (R1 + R2)/R3 | 0.58 ± 0.14 | 0.55 ± 0.06 | 0.46 ± 0.14 |
| wild-type | R1/R2     | 0.48 ± 0.04 | 0.48*     | 0.16 ± 0.02 |
|          | (R1 + R2)/R3 | 0.51 ± 0.09 | 0.30 ± 0.25 | 0.55 ± 0.19 |

The ratios presented are the average of the following number of RNase protection assays using probe A (for wild-type RNA, see Fig. 2B) or probe A-2018 (for NS2-2018 RNA) as described in the text, and includes 95% confidence limits. Total [NS2-2018 R1/R2, (R1 + R2)/R3], 8; wild-type (R1/R2), 17; nuclear [NS2-2018 and wild-type], 3; cytoplasmic [NS2-2018 and wild-type], 8.

*Average of two experiments.
Figure 8. To effect RNA processing, termination signals must interrupt previously open reading frames that have the potential of being joined in-frame to the initiating AUG at nucleotide 260. The map shows the locations of the alterations in the Tb60 set of mutants. Relevant ORFs from the right half of the nonstructural genes [approximately nucleotides 1950-2400; see Fig. 1A] are shown. The frameshift mutation at nucleotide 2073, which shifts ORF 2 into ORF 3, is indicated by the vertical line. Sites of insertions of ochre termination signals and the reading frames that they interrupt are indicated. The column at right shows the relative accumulated ratio of R1 versus R2 that each mutant generates after transfection into A9 cells and is the average of five experiments, with 95% confidence limits included. RNase protection analyses were done as described in Materials and methods using either probe B or C (Fig. 1A).

Urlaub et al. (1989), proposes that translation of the 5' end of an mRNA molecule, on the cytoplasmic side of the nuclear envelope, facilitates transport, whereas splicing of the 3' part of that molecule is still taking place. If translation is terminated prematurely, transport to the cytoplasm is retarded and nuclear splicing is inhibited, presumably leading to increased degradation of that RNA in the nucleus.

Because termination codons that interrupt ORFs in the second exon of R2 and lie 3' to the splice that they affect [e.g., mutations in NS2-2018, NS2-2159, and X27] can inhibit the splicing of R1 to R2, our results are difficult to reconcile with the translation/translocation model as proposed previously (Urlaub et al. 1989). To conform to this model, ribosomes would need some direct influence on nuclear RNA processing before translation termination. Our results are more consistent with the nuclear exon scanning model also proposed by Urlaub et al. (1989), in which mRNAs are scanned for ORFs in the nucleus before splicing, thus avoiding the use of cryptic splice sites that result in nonproductive products. Such a model might be part of the process of exon definition. Recently, Robberson et al. (1990) have proposed a nuclear exon scanning model in which downstream exons are defined by interactions between splicing factors bound to adjacent splice sites across exons, and perhaps such a mechanism might also include scanning of ORFs. Because the first MVM exon lacks a 5' splice acceptor site, however, a different recognition point would be required to define this exon. As the effect of the termination signals interrupting an ORF is generally polar with respect to the initiating AUG of NS1 and NS2, such an effect on RNA processing might occur after the spliceosome shortens the distance between the initiating AUG (or the 5' end of the message) and the termination signal by looping out the intervening intron. Scanning of the first exon, however, apparently does not require recognition of the initiating AUG because p260[T4] and p260[T4]TTT, in which the initiating AUG has been removed, splice R1 to R2 normally. Experiments are currently under way to delineate further the role of the initiating AUG codon in this process.

Because nonsense mutations in the second exon of R2 inhibit splicing of R1 to R2 only if they interrupt an ORF
Nonsense mutations inhibit splicing in cis

Figure 9. The distance of the amber codon at nucleotide 2086 from the initiating AUG affects its role in RNA processing. The map displays the nonstructural genes of MVM from approximately nucleotides 200 to 2300. The frameshift in Tb60 shifts the NS1 in R1 from ORF 3 into ORF 1, where it is terminated by the amber codon at nucleotides 2086–2088, 1826 nucleotides from the initiating AUG at nucleotide 260. The frameshift in mutant X27 shifts the NS2 in R2 from ORF 2 into ORF 1, where it is terminated by the same amber codon at nucleotides 2086–2088, now 351 nucleotides from the initiating codon at 260. The column at right shows the relative accumulated ratio of R1 versus R2 that each mutant generates after transfection into A9 cells and is the average of five experiments, including 95% confidence limits. RNase protection analyses were done as described in Materials and methods using either probe B or C [Fig. 1A].

that has the potential to be joined in-frame to the first exon after splicing, there must be a functional cooperation between the exon scanning mechanism and the ability to link ORFs between the normal splice donor and acceptor. As the inhibition of splicing that we detect, however, is not complete, exon scanning is probably only one component of this complex process.

There is growing evidence that nonsense mutations that interrupt a previously open reading frame can effect nuclear RNA-processing events such as splicing and stability. Because the processing pathway for the parvovirus MVM is relatively simple and well defined, it offers an excellent opportunity to study the relationship between the definition of exons and RNA processing in murine cells.

Materials and methods

Cells, virus, infections, and transfections

Murine A9 cells were propagated as described previously (Pintel et al. 1983). Wild-type and NS2 mutant virus was titered and propagated, and infections were done as described previously.
RNA isolation and characterization

Total RNA was isolated as described previously [Schoborg and Pintel 1991]. Nuclear and cytoplasmic fractionation of infected cellular RNA was done as described [Miller et al. 1989]. Nuclear fractions were determined to be >95% pure, as monitored by the absence of mature rRNA and the presence of RNA precursors by ethidium bromide staining after gel electrophoresis (data not shown). RNase protection assays were done, using equal amounts of total, nuclear, or cytoplasmic RNA as indicated, exactly as described [Clemens and Pintel 1988], using an [α-32P]UTP-labeled, SP6-generated antisense MVM RNA probe from either MVM nucleotides 1854–2377, 146–652, or 385–652 (diagrammed as A, B, and C, respectively, in Fig. 1A), as indicated. Probe A [MVM nucleotides 1854–2378] extends from before the acceptor site of the large R2 splice to within the small intron common to all viral RNAs at map units 44–46 and distinguishes RNA species using either of the alternate small splice donors, designated M for the major splice 2280–2377 and m for the minor 2317–2399 [Fig. 2B, Clemens and Pintel 1988]. For analysis of RNA produced after transfection or infection of NS2-2087, probe A-2087 was used, which is identical to probe A except that the G→U mutation at nucleotide 2018 has been introduced. Probes B [nucleotides 146–652] and C [nucleotides 385–652] distinguish MVM R1 and R2, as indicated in the figures. Northern blot analysis [Pintel et al. 1983] and RNA stability assays using either 40 μg/ml of actinomycin D or 40 μg/ml of the adenosine analog DRB [Schoborg and Pintel 1991] were done exactly as described previously. RNase protection assay data were collected on a Betagen B-scanning image analyzer, and molar ratios of MVM RNA were determined by standardization to the number of uridines in each protected fragment. Total R1 and R2 [unspliced as well as spliced] RNA levels were included in the quantitation for each analysis.

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L K Naeger, R V Schoborg, Q Zhao, et al.

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