The genome of minute virus of mice, an autonomous parvovirus, encodes two overlapping transcription units

David Pintel, Dinshaw Dadachanji, Caroline R. Astell* and David C. Ward

Departments of Human Genetics and Molecular Biophysics-Biochemistry, Yale University School of Medicine, New Haven, CT 06510, USA and *Department of Biochemistry, Faculty of Medicine, University of British Columbia, Vancouver, British Columbia, V6T 1WF, Canada

Received 1 December 1982; Revised and Accepted 25 January 1983

ABSTRACT

Four virus-specific transcripts have been identified in murine cells infected with Minute-Virus-of-Mice (MVM). These RNAs, 4.8, 3.3, 3.0 and 1.8 kilobases in length, designated R1 to R4 respectively, are all transcribed from the virion (-) strand of DNA and they are all polyadenylated and spliced. The R1 transcript is derived from sequences that reside on the genome between 4.0 and 95 map units (mu). Transcript R2 is composed of exon sequences derived from mu coordinates 4.0-10.0, 40-46 and 48-95. The most abundant RNA, R3, is transcribed from sequences mapping between 40 and 95 mu. All three of these RNAs have a short intron sequence between 46-48 mu removed. The least abundant transcript, R4, has not been mapped precisely, however it hybridizes with all three EcoRI fragments which span the entire 5 kb genome. In vitro transcription of cloned restriction fragments of MVM DNA confirm the existence of functional promoters at map coordinates 4.0 and 39 and sequence analysis of these regions of the viral DNA reveal the characteristic features of RNA polymerase II promoters. These results indicate that MVM DNA encodes two overlapping transcription units with separate promoters near the left end (4.0 mu) and middle (39 mu) of the genome.

INTRODUCTION

Mammalian paroviruses possess linear, single-stranded DNA genomes that are only 4500-5200 nucleotides in length (1). The limited genetic complexity of these genomes makes them attractive model systems for studying features of eukaryotic gene expression. To date the majority of such studies have been done with Adeno-Associated Virus, type 2 [AAV-2] (2-6), a prototype of the defective parovirus subgroup that require Adenovirus (7) or Herpes simplex virus (8) functions to initiate their own replication. The emerging picture of AAV-transcription (9) is that there are six species of polyadenylated cytoplasmic RNA synthesized in infected cells. These are 4.2, 3.9, 3.6, 3.3, 2.6 and 2.3 kilobases (kb) in length, respectively. The 4.2 and 3.9 kb transcripts are initiated from a promoter at map position (mu) 5.0 and differ only in that the 3.9 kb species has genomic sequences between 40 and 48 mu spliced out (5,9). The same region of the
genome also has been spliced out in the 3.3 and 2.3 kb RNAs. These latter transcripts and their unspliced (3.6 and 2.6 kb) counterparts are initiated from promoters at 19.0 and 39.0 µm, respectively (5,9, E. Lusby and K. Berns, personal communication). Since each of the six RNAs share an extensive block of common sequence, derived from the genome between 48 and 96 µm, it is apparent that AAV DNA encodes three overlapping transcription units which coterminate at µm 96.

Initial studies on the transcription of the DNA from H-1 virus and Minute Virus of Mice (MVM), two members of the autonomous parvovirus sub-group that replicate without the aid of a helper virus (1), suggested that these viral genomes encode only a single transcription unit with a promoter near µm 4.0 (10,11). To further characterize the transcriptional organization of an autonomous parvovirus genome we undertook an analysis of the RNA species produced in MVM-infected mouse cells. The results presented here demonstrate the presence of four spliced viral RNAs, all transcribed from the virion strand of DNA. The three major transcripts have been shown to arise from two overlapping transcription units with separate promoters positioned near the left end (4.0 µm) and the middle (39 µm) of the genome. Additional features of the transcriptional organization of MVM DNA are discussed in the following paper, which presents the complete nucleotide sequence of the MVM genome (12). A preliminary report of some of these data has been presented (13).

**MATERIALS AND METHODS**

**Virus and Cells**

Plaque-purified MVM(p), the prototype strain of MVM, was grown in the A-9 strain of mouse L cells (14). The maintenance of cell cultures and virus stocks was as described (15).

To obtain 32P-labeled viral DNA, carrier-free 32P-orthophosphoric acid (New England Nuclear) was added at 80 µCi/ml to culture media, containing only 2-5% of the normal inorganic phosphate, 7-9 hours post infection. Cells were harvested 26 hours post infection, and viral DNA prepared from purified virions as described (16). This labeling condition yielded DNA of approximately 6-8x10^5 cpm/µg.

**RNA extraction and cell fractionation**

RNA was extracted from whole cells by the guanidinium chloride method described by Cox (17) and modified by Strohman et al. (18).

Cells were divided into nuclear and cytoplasmic fractions with Nonidet
P-40 (NP-40) as described by Borun et al. (19) and modified by Carter and Rose (20). RNA was extracted from the nuclear and cytoplasmic fractions of infected and mock infected cells exactly as described by Carter and Rose (20). Poly A⁺ RNA was selected on an oligo(dT)-cellulose column (Collaborative Research).

RNA-DNA Hybridization in Solution.

a) In aqueous solution.

RNA-DNA hybridizations were carried out in 2xSSC at 67°C for 23 hours (21). Each assay mix (100 μl), containing 2-5x10³ cpm of ³²P-labeled MVM DNA (2x10⁵ cpm/μg) and varying concentrations of unlabeled RNA from infected or mock-infected cells, was heat-denatured before annealing. At the end of the hybridization incubation, 25 μl of each mix was TCA-precipitated onto GF/A filters and counted, while 50 μl of the mix was digested with SI nuclease prior to TCA precipitation.

b) In 80% formamide.

Hybridizations under conditions favoring DNA-RNA annealing were carried out as described (22,23). The hybridization buffer consisted of purified (24) 80% formamide (Macalaster-Bicknell or Fisher), 0.4 M NaCl, 0.04 M PIPES-Na, pH 6.4, and 1 mM EDTA. The hybridization mix (100 μl) usually contained between 30-100 ng MVM [³²P]-DNA, and between 3-36 μg cellular RNA (as specified in the figure legends). Each assay mix was first incubated at 80°C for 5 min to denature the nucleic acids and then placed at the hybridization temperature. The optimum temperature for selective hybridization was found to be 57°C, and the incubation time ranged from 3-18 hours. At the end of each incubation, aliquots were precipitated with TCA before and after treatment with either SI or mung bean nuclease.

SI Nuclease Digestion.

The digestion buffer consisted of 0.03 M NaOAc, pH 4.5, 0.25 M NaCl, 1 mM ZnSO₄. Samples obtained from RNA-DNA hybridization assays were diluted 10-fold in the above digestion buffer, and incubated with 65 U/ml SI nuclease (Sigma) in the presence of sonicated, heat-denatured calf thymus DNA (20 μg/ml) at 48°C for 75 min and the TCA-precipitable counts were then determined in the presence of carrier RNA. (1 unit of SI nuclease is defined as the amount of enzyme that converts 1 μg of nucleic acid to a TCA-soluble form in 1 min at 37°C, at pH 4.6).

Mung Bean Nuclease Digestion.

Samples obtained from RNA-DNA hybridization assays were diluted 10-fold.
with buffer (0.05 M NaOAc/HOAc, pH 5.0, 0.05 M NaCl, 1 mM ZnCl\textsubscript{2}). Both native and heat-denatured, sonicated calf thymus DNA, each at a final concentration of 10μg/ml, were added and each assay mix was incubated with 20 U/ml mung bean nuclease (PL Biochemicals), at 45°C for 45 min (unless otherwise indicated). Samples were either tested for TCA-precipitable counts or alcohol-precipitated, and redissolved in TE 8.3 buffer, and analyzed by gel electrophoresis. (1 unit of mung bean nuclease is defined as the amount of enzyme that produces 1 μg TCA-soluble material in 1 min at 37°C, at pH 4.6).

**RESULTS**

Initially, unlabelled RNA was prepared from MVM infected and mock infected A9 cells 14 hours post infection and the RNA was extracted from nuclear, cytoplasmic or whole cell fractions. These RNA preparations were annealed in aqueous solution to \textsuperscript{32}P-vDNA over a range of RNA/DNA ratios, and assayed for the extent of hybridization by determining the proportion of \textsuperscript{32}P-radioactivity found resistant to the action of SI nuclease. The results of these analyses are shown in Figure 1. It can be seen that less than 10% of the viral DNA is resistant to SI nuclease when it is annealed to itself or to RNA from uninfected cells. This background level of annea-
MVM specific RNA is transcribed using only the viral strand as template; we were unable to detect RNA which hybridized to viral C-strand probes. When MVM $^{32}P$-cDNA was purified as described (29), hybridized in solution with cytoplasmic fractions of MVM infected A9 cells, and treated with SI nuclease the percent of nuclease resistant cDNA did not rise above the background obtained using mock infected cell RNA, even at RNA/DNA ratios up to 60,000:1. At this ratio, had transcripts complementary to cDNA been represented as $5 \times 10^{-5}$% of the cellular RNA tested, the proportion of cDNA resistant to SI would have risen to at least twice the background level. MVM $^{32}P$-cDNA was also used to probe "Northern" blots from MVM infected cells. Exposure times 50X longer than those necessary to visualize RNA hybridizing to $^{32}P$-vDNA probes detected a very weak pattern. However, the pattern obtained was identical to that seen using $^{32}P$-vDNA probes, and most probably reflects hybridization with a trace amount of
contaminating vDNA in the cDNA preparations. A search for symmetrical transcripts, first described by Aloni and Attardi (30), which might have prevented detection of low abundance RNA complementary to cDNA (due to the hybridization of transcripts of opposite polarity to each other) was also negative. There was no increase of double stranded RNA species, as detected by pancreatic and T1 nuclease resistance, in MVM infected cells compared to mock infected cells, and neither pool of duplex RNA hybridized in solution with excess cDNA probes (data not shown).

To determine the number and size of viral specific transcripts, total cell RNA was extracted from MVM infected A9 cells 23 hours post infection and fractionated by electrophoresis in a formaldehyde-agarose gel. The RNA was transferred to nitrocellulose and hybridized with $^{32}$P-labeled, nick-translated, MVM RF DNA. Four RNA species, 4.8, 3.3, 3.0 and 1.8 kilobases (kb) in length, can be detected by this type of analysis (Figure 2a). Similar experiments using poly A selected RNA and $^{32}$P-labeled vDNA yield identical results (data not shown).

To determine the proper annealing conditions for analyzing hybrids by the nuclease procedure of Berk and Sharp, $^{32}$P-labelled vDNA was annealed, in 80% formamide, at various temperatures, to total intracellular RNA taken from MVM infected A9 cells 26 hours post infection. The hybridization mix was digested with mung bean nuclease, and run on a 1.4% agarose gel under neutral pH conditions. For each temperature tested both DNA/DNA hybridization of separate viral restriction fragments and DNA hybridization to total RNA from mock infected cells were run in parallel. The results shown in Fig. 2b indicate that the optimum temperature for hybridization is 56°C-58°C (lanes 2-7). The gels also demonstrate the existence of four major hybrid bands when $^{32}$P-labelled vDNA was hybridized with RNA extracted from MVM infected cells. The three largest of these bands 4.8, 3.3 and 3.0 kb correspond both in size and relative intensity to the three largest transcripts seen by Northern analysis. In addition, a fourth band appears at approximately 2.6 kb. This band is smaller than any of the major transcripts seen by Northern analysis and, as discussed below, is the size of the major shared exon of the three largest and most abundant RNA species. We believe that this band represents an artifact of the nuclease digestion, resulting from RNA strand scission at the point of a splice junction. Similar artifactual cleavages have been reported previously by others (10,34).

Since the three largest transcripts represent over 95% of the viral RNA
Figure 2A. Hybridization of cloned MVM RF DNA $^{32}$P-labeled by nick-translation to RNA extracted from MVM-infected (lane 1) and uninfected (lane 2) A9 cells 26 hours post infection. The methods of RNA extraction, blotting and hybridization are described in Materials and Methods. The relative abundance of the four viral transcripts are: 4.8 kb band (10-15%), 3.3 kb band (15-20%), 3.0 kb band (65-70%), and 1.8 kb band (2-3%). B. Effect of temperature on the specific formation of DNA/RNA hybrids. Total RNA (17 µg), prepared from infected A9 cells 26 hours post infection, was hybridized in 80% formamide buffer to MVM $[^{32}P]$-vDNA (12,000 cpm, 21 ng) at 56° (lane 3), 58° (lane 6), 60° (lane 9) or 62° (lane 12), then treated with mung bean nuclease and electrophoresed on a 1.4% agarose gel for 4 hours at 100V. (At this RNA/DNA ratio, less than 20% of the DNA probe becomes SI resistant). RNA (18 µg) from uninfected A9 cells was hybridized at these temperatures with the $^{32}$P-MVM DNA probe (21 ng), nuclease treated and electrophoresed in lane 4 (56°), lane 7 (58°), lane 10 (60°) and lane 13 (62°). To control for DNA:DNA reannealing, $^{32}$P-MVM RF DNA, synthesized in vitro (16), was subjected to partial digestion with EcoRI, heat denatured and annealed at the indicated temperatures for 2 hours in 80% formamide buffer. The three bands observed represent fragments which spontaneously renature since the V and C strands are covalently linked. The EcoRI A,B and AB fragments (see Figure 4), in which V and C strands are not covalently associated, do not reanneal at any of the four hybridization temperatures used.

in infected cells, total cytoplasmic RNA (extracted 26 hr post infection) was hybridized to $^{32}$P-labelled vDNA using the optimal hybridization conditions described above, treated with mung bean nuclease, and run on
Figure 3. Alkaline gel analysis of MVM RNA-DNA hybrids obtained by the "Berk and Sharp" technique. A. Equal aliquots of MVM [*32P]-vDNA (20,000 cpm, 0.03 µg) were annealed with 36 µg of cytoplasmic RNA from MVM-infected A9 cells taken 24 hours post infection (lane 1) or from mock-infected cells (lane 2). Hybridization were performed in 80% formamide hybridization buffer, at 57°C for 4 hours. Samples were then digested with mung bean nuclease and run on an alkaline 1.4% agarose gel. Lane 3: partial digest of MVM [*32P]-RF DNA. B. Autoradiogram of an alkaline 1.4% agarose gel, analyzing the three major RNA-DNA hybrid bands individually isolated from a neutral gel. MVM [*32P]-vDNA was hybridized to total RNA from MVM-infected A9 cells (extracted at 25 h p.i.), in 80% formamide hybridization buffer at 57°C for 20 h (at an RNA/DNA ratio of 515:1). The sample was digested with mung bean nuclease, electrophoresed through a preparative 1.4% agarose gel, pH 8.3, and each of the three largest hybrid bands (see text) was sliced out, dissolved in saturated KI solution, and recovered by chromatography on hydroxylapatite. The RNA-DNA hybrids derived from the 4.8, 3.3 and 3.0 kb transcripts are shown in lanes 2 through 4, respectively. Lane 1: partial EcoRI digest of MVM [*32P]-RF DNA. C. The effect of varying mung bean nuclease concentrations on RNA/DNA hybrids. An aliquot of [*32P]-vDNA (72,000 cpm, 168 ng) was annealed with total RNA from MVM infected A9 cells extracted 26 hours post infection, at an RNA/DNA ratio of 200:1, in 80% formamide hybridization buffer, at 57°C for 13 h. The sample was then divided into 4 equal aliquots, digested with increasing concentrations of mung bean nuclease: 3.6 U/ml (track 3), 7.2 U/ml (track 4), 14.4 U/ml (track 5) and 28.8 U/ml (track 6), and electrophoresed on an alkaline 1.4% agarose gel. Lane 1: partial EcoRI digest of MVM [*32P]-RF DNA. Lane 2: blank.
alkaline 1.4% agarose gels. As seen in Fig. 3a, five bands are generated under denaturing conditions, whereas four bands were observed under the neutral gel conditions discussed above and shown in Fig. 2b. The very prominent band at 2.6 kb in addition to the bands at 2.1 and 0.3 kb represent exons which constitute the major RNA species as detailed below. The origin of the minor bands at 3.0 and 1.8 kb are also discussed below.

When the three largest hybrid bands (4.8, 3.3 and 3.0 kb) illustrated in Fig. 2b were individually eluted, and separately run on alkaline 1.4% agarose gels the bands shown in Fig. 3b were observed. It can be seen immediately that all three of the major transcripts have an exon fragment 2.6 kb in length. The 4.8 kb transcript (Fig. 3b, lane 2) can be seen to contain the 2.1 kb exon fragment, previously noted in Fig. 3a. The 4.8 kb transcript also protects DNA sequences on the alkaline gel which migrate at 4.8 kb and 2.8 kb. These bands could represent either incomplete digestion products or minor unspliced species (see below). The 3.3 kb transcript can be seen to contain the 2.6 kb exon, the 2.8 kb band, and, upon longer exposure, a 0.3K exon band. The major 3.0 kb transcript generates the 2.6 kb exon band, a 0.3 kb exon band, the 2.8 kb band, the 3.0 kb band (previously noted in Fig. 3a) and a minor band at 3.2 kb - larger than the original 3.0 kb transcript.

The presence of a 2.8 kb band in each of these samples was unexpected since no such band was observed in Fig. 3a, which displays the hybrid protected DNA fragments generated by the total viral RNA pool after alkalai treatment. To account for this apparent discrepancy we next analyzed the effect of varying mung bean nuclease concentration on the DNA/RNA hybrids generated during such analysis. Fig. 3c shows that at low nuclease concentration bands at 2.8 kb as well as at 4.9 and 5.1 kb are seen; these bands disappear as the nuclease concentration increases. In contrast the bands at 2.1, 2.6 and 3.0 kb in these digests remain unaffected by increasing the nuclease level. (The 0.3 kb exon was run off the gel in this experiment in order to get better resolution of higher molecular weight bands). We interpret the 2.8 kb band, seen in Fig. 3b, as a partial digestion product which is not present in the sample illustrated in Fig. 3a. Restriction fragment mapping data presented below shows that the major 2.6 kb exon RNA extends very close to the 5' hairpin duplex. We conclude that the partial digestion band at 2.8 kb represents incomplete cutting at this site, near the 5'-termini, which results in fragments approximately 250 nucleotides longer than expected. The minor 3.2 kb fragment observed in the analysis
of the 3.0 kb transcripts (lane 4, Fig. 3b) is also interpreted to be due to incomplete removal of the 5' terminal hairpin sequence. The 5.1 and 4.9 kb bands obtained at low nuclease concentration (lane 1 of Fig. 3c) most likely results from incomplete nuclease digestion at one or both of the terminal hairpin sequences.

Although no band of 1.8 kb was observed in the nuclease protection studies illustrated in Fig. 2b, where the protected DNA fragments were resolved on neutral agarose gels, a minor 1.8 kb band is observed on the alkaline gel shown in Fig. 3a. The band could reflect an unspliced form of the 1.8 kb transcript seen by Northern analysis (Fig. 2a). Unfortunately, since this RNA species appears in such low abundance in MVM infected cells we have not been able to analyze this RNA further. However, the 1.8 kb transcript does hybridize to all three EcoRI fragments of MVM RF DNA, which span the entire 5 kb genome of MVM, suggesting that the majority of the molecules in the RNA species are spliced.

Although Exonuclease VII has been used widely in nuclease protection experiments to determine the size of intron sequences, such studies, using whole MVM genomic DNA as probe, have proven extremely difficult in our hands since the hairpin duplexes at the genomic termini render the MVM-vDNA probe extremely resistant to this enzyme. Therefore, the sites and sizes of intron sequences and the map coordinates of the splice junctions were determined directly by the method of Berk and Sharp (23) using mung bean nuclease. Total cellular RNA from MVM infected A9 cells, taken at 26 hours post infection, was hybridized in solution with isolated restriction fragments of 32P-labelled MVM RF DNA. After treatment with mung bean nuclease the RNA protected DNA fragments were analyzed under neutral and denaturing gel conditions. The Hind III B and Hind III C fragments, which map between 54-80.5 (nucleotide 2650-3993) and 80.5-100 (nucleotide 3993-5081) respectively on the viral genome, are wholly protected from nuclease in the above analysis. Nucleotide numbers are taken from the complete nucleotide sequence of MVM, (12). The alkaline agarose gel profile shown in Figure 4c indicates that the rightward half of the genome is transcribed without interruption, and since the Hind III B and C fragments span a region approximately 2440 nucleotides in length and the two largest exons have been determined to be 2.6 and 2.1 kb, this region must encode the large exon of 2.6 kb, common to the three most abundant transcripts, demonstrated in Figure 3a. There is also a diffuse band generated from the Hind III C hybrid on the alkaline gel which migrates at approximately
Figure 4. Analysis of hybrids between $^{32}$P-labeled restriction fragments of MVM RF DNA and total cell RNA which had been extracted from MVM infected A9 cells 26 h post infection. Hybridization were performed in 80% formaldehyde hybridization buffer at 57°C for 15-18 h at an RNA/DNA ratio of 100:1 (at which ratio only 5% [above background] of the genome is SI resistant). The samples were then digested to completion with mung bean nuclease and run on either an alkaline 1.4% agarose gel, Fig. 4D; 4C; neutral 1.4% agarose gel, Fig. 4B; or an 8M urea - 8% polyacrylamide gel, Fig. 4A. A map of the restriction fragments appears on top. A: analysis of nuclease treated hybrids obtained with Hinf I A (956-2664), lane 3; Hinf I C (534-956), lane 5; and Hinf I E (225-534), lane 7. Lanes 2, 4, 6: unhybridized, purified $^{32}$P-labeled Hinf I A, C and E fragments respectively. Lane 1: Hhal digest of φX174 $^{32}$P-RF DNA. B: Neutral agarose gel analysis of nuclease treated hybrids obtained with the EcoRI A (1086-3521) fragment (lane 2). Lane 1: $^{32}$P-labeled purified EcoRI A fragment before hybridization. Lane 3: EcoRI digest of MVM $^{32}$P-RF DNA. C: Alkaline agarose gel analysis of the same samples as in B. D: Alkaline agarose analysis of nuclease treated hybrids with the Hind III B (2652-3996) fragment (lane 3) and Hind III C (3996-5081) fragment (lane 5). Lanes 2 and 4 are $^{32}$P-labeled purified Hind III B and C fragments respectively, before hybridization. Lane 1: EcoRI digest of MVM $^{32}$P-RF DNA.

220-240 nucleotides (Figure 4d, lane 5). This is the size of the 5' terminal hairpin which is most likely released intact after nuclease digestion. Although four additional minor bands are seen in the HindIII B protection experiment, (Fig. 4d, lane 3), these bands are also present in
the unhybridized DNA fragment control (lane 2), and most likely represent degradation products of the restriction fragment probe. Hybrids with the EcoRI A fragment (21-70 μm, nucleotide 1084-3518) help delineate the boundaries of the two major introns in the genome. Neutral gel analysis demonstrates three bands (Figure 4b, lane 2). The largest (2430 nucleotide) band seen in Figure 4b, lane 2, migrates identically to the unhybridized EcoRI A fragment and represents hybrids with the RNA subpopulation that spans the boundaries of the fragment. However, the major band observed is approximately 1475 nucleotides in length. Since the Hind III B fragment is transcribed co-linearly and the EcoRI A fragment overlaps approximately 870 nucleotides of Hind III B, the 1475 bp hybrid must be from a transcript that initiates 1475 nucleotides from the right end boundary of this fragment, i.e., at approximately nucleotide 2000 (μm 40). This hybrid determines the left hand boundary of the major 3.0 hybrid (seen in the neutral gels of Figure 2b), and from the right hand boundary determined above, we conclude that the 3.0 kb transcript lies between map units 40 and 100 μm. The minor band of 1150 nucleotides also on the gel is the size of the largest exon segment protected in the hybrid (see below) and constitutes another example of the nuclease digestion artifact described above. On alkaline gels (Figure 4c) the major band migrates at 1150 nucleotides and, in addition, there is a less intense band at 0.3 kb. This indicates that the transcript which initiates at μm 40 continues for approximately 300 nucleotides, contains a small splice between μm 46-48, and then continues uninterrupted through the remainder of the EcoRI A fragment and through the Hind III B and C fragments as well. Together these results demonstrate that the common 2.6 kb exon maps between μm 48 and 96 on the genome.

The Hinf I A fragment (μm 17.1 to 49.4, nucleotide 954-2662) overlaps the majority of the EcoRI A fragment and is useful in confirming the results described above. Neutral gel analysis shows that approximately 650-690 nucleotides of the Hinf I A fragment are protected by MVM RNA (data not shown); this provides further evidence that the 3.0 kb transcript starts at μm 40. The hybrids protected by the Hinf I A fragment were also run on alkaline acrylamide gels as shown in Figure 4a. It can be seen that the 650-690 hybrid forms multiple bands centering around 280 nucleotides (Figure 4a, lane 3). The sizes of these fragments corroborate the existence of two exons of approximately 265-330 nucleotides, which are separated by a small splice at μm 46-48. The multiple band pattern seen
may indicate heterogeneity in these RNA species. As expected, there is also a minor hybrid fragment that spans the entire Hinf I fragment, indicative of the large 4.8 kb transcript that is unspliced. Alkaline gel analysis of hybrids with HinfI C fragment (mu 9.6-171, nucleotide 531-954) shows no interruption of the coding sequence in this region (Fig. 4a, lane 5). Analysis of hybrids with the Hinf I E fragment (mu 4.3-9.6, nucleotide 224-531), however, gives evidence for two differentially spliced molecules coded for by this region. Neutral gel analysis shows that all RNA hybridizing with this fragment span the entire fragment (data not shown) while the alkaline acrylamide gel displayed in Fig. 4a, lane 7 demonstrates that while one species remains unspliced through the region, another species is truncated approximately 25 nucleotides from the 3' end of the fragment, or approximately nucleotide 510, map position 10. This leaves a leader sequence 300 nucleotides long between mu 4.5-10. Since the 3.3 kb message appears to be comprised of two 0.3 kb leader sequences attached to the common 2.6 kb exon, the smaller region of HinfI E which is protected encodes the leftmost of these two leaders segments.

To corroborate the emerging map of viral transcripts, we next probed four identical samples of total RNA from MVM infected cells which had been electrophoresed through formaldehyde-agarose and transferred to nitrocellulose with four 32P-labeled cloned MVM DNA fragments from different areas across the genome. As expected the cloned Hind III C fragment (nucleotide 3993-5081) and the cloned EcoRI A (nucleotide 1084-3518) fragment hybridized to each of the 4.8, 3.3, and 3.0 kb messages. Cloned DNA from nucleotide 1084-1661, which falls totally within the large intron hybridizes with only the 4.8 kb message, and the cloned PvuII C fragment (nucleotide 1-760) hybridizes with only the 4.8 and 3.3 kb messages (data not shown).

The map coordinates of the 3 major transcripts, deduced from the data presented above, are summarized in Figure 5. The largest message, R1, is initiated at the left end of the genome at ~mu 4 and extends the length of the genome to ~mu 96. Most of these molecules have a small region between mu 46-48 spliced out, although there apparently does exist a subpopulation that lack the small splice. The next largest transcript, R2, which also begins at ~mu 4.0, extends to ~mu 10 at which point it is spliced to ~mu 40 on the genome. The RNA continues ~300 more nucleotides downstream at which point a small region between mu 46-48 is spliced out; it then continues uninterrupted to ~mu 96. This message lacks two blocks of
Figure 5. Map displaying the organization of the three major transcripts of MVM (see text for details).

genomic sequences, one large block between 10-40 mu and one small block between 46-48 mu. The map coordinates of the R2 transcript deduced by the nuclease protection study agree extremely well with those determined by Tal et al. (11) using electron microscopic measurements of DNA-RNA hybrids. However, the small splice in this transcript at 46-48 mu was not observed by Tal et al., presumably because the resultant intron loop was too small to be detected by electron microscopy. Since only hybrid molecules with detectable intron loops were analyzed in the EM study, the map coordinates of the most abundant 3.0 kb transcript were not reported. The data presented above indicates that the major R3 transcript is initiated from a promoter near mu 40 and possesses sequences complementary to genomic DNA mapping between 40-46 and 48-96 mu. It remained possible however, that the above analyses might have missed a very smaller leader sequence on the R3 transcript encoded by the left end of the genome around mu 4.

To corroborate the existence of two promoters in the MVM genome we have examined restriction fragments from viral RF DNA and cloned MVM DNA segments for promoter activity in the standard in vitro transcription system developed by Manley et al., (27). The cloning of the complete molecule of MVM or subgenomic fragments of the viral DNA into pBR322 and the characterization of the resultant clones will be presented elsewhere (Merchlinsky et al., manuscript in preparation).

In our hands the in vitro HeLa cell transcription system neither splices nor polyadenylates transcripts, however it is very useful for identifying and mapping promoter sites onto isolated DNA. When full length MVM RF DNA cloned into pBR322 is used to prime the in vitro transcription system two transcripts approximately 4.8 and 3.2 kb in length are produced (Fig. 6, part A). The 3.2 kb band is a doublet because the template pool contains a minor population of molecules which contain a deletion of approximately 85 base pairs within the 5' hairpin sequences of MVM which is generated during propagation in bacteria (Merchlinsky et al., unpublished).
Figure 6. In vitro transcription of cloned MVM RF DNA in Hela cell lysates. Incubation conditions were exactly as described by Manley et al. (27). RNA products were treated with glyoxal at 50°C for 1 hour, electrophoresed on 1.4% agarose gels and autoradiographed. Lanes 1 and 6 are 32P-labeled lambda DNA cut with Hind III. A. Transcription of cloned MVM RF DNA. Templates are: lane 2, cloned total RF DNA excised from pBR322 vector; lane 3, linearized pBR322 vector alone. B. Transcription of EcoRI C fragment (nucleotide 0-1086). Templates are: lane 4, purified EcoRI C fragment; lane 5, purified EcoRI C fragment plus 2 µg/ml of α-amanitin. C. Transcription of cloned EcoRI A (nucleotide 1086-3521) and EcoRI B (nucleotide 3521-5081) fragments. Templates are: lane 7, linearized pBR322 alone; lane 8, EcoRI A fragment purified from the plasmid vector; lane 9, purified EcoRI A fragment cut with PVUI (nuc. 2842) truncating 679 bp from the right hand end; lane 10, purified EcoRI A fragment cut with Hind III (nuc. 2652) truncating 869 bp from the right hand end; lane 11, EcoRI B fragment excised from the plasmid vector. D. Templates are: lane 12, clone EcoRI A fragment excised from pBR322 vector; lane 13, the same as lane 12 above plus 2 µg/ml α-amanitin.

observation). To demonstrate the activity of the left end promoter in vitro we chose to transcribe the EcoRI C fragment, purified from MVM RF DNA, which terminates at nucleotide 1084 (c.f. Fig. 4). This fragment primes the synthesis of a discreet runoff transcript approximately 870 nucleotides in length (Fig. 6, lane 4) which is not produced in the presence of 2 µg/ml α-amanitin, characteristic of authentic polymerase II
transcripts. The size of this runoff transcript places the initiation of transcription in vitro at approximately nucleotide 205-225 or mu 4. The EcoRI A fragment (nucleotide 1084-3518; c.f. Fig. 4) was used to demonstrate the activity in vitro of the internal promoter (Fig. 6, part C). This fragment generates a runoff transcript of approximately 1475 nucleotides (Fig. 6, lane 8), consistent in size with the nuclease protected hybrid obtained previously between the EcoRI A fragment and RNA from MVM infected cells as seen in Fig. 4, part B. When the EcoRI A fragment, truncated at nucleotide 2840 by PvuII (removing 679 nucleotides), or at nucleotide 2650 by Hind III (removing 869 nucleotides), is used to prime the in vitro system, transcripts 800 nucleotides (shorter by ~ 675 nucleotides) and 600 nucleotides (shorter by ~ 875 nucleotides) in length are generated (Fig. 6, lanes 9 and 10). These results map the initiation of transcription to approximately nucleotide 2000, i.e. mu 39-40. Recent experiments using the primer extension method of Weaver and Weissman (35), has precisely mapped the 5' end of the 3.0 kb message, revealing heterogeneous initiation at nucleotides 2003 (major start site), 2006 and 2010 (C. Astell, unpublished results). The 1475 nucleotide transcript generated in response to the EcoRI A fragment is also sensitive to low levels of α-amanitin as shown in Fig. 6, part D. In vitro transcription of cloned EcoRI B fragment (nucleotide 3518-5081, c.f. Fig. 4) produced no specific transcript (Fig. 6, lane 11). Our in vitro system generates numerous minor bands which are DNA dependent but α-amanitin insensitive. Such bands, presumed not to be RNA polymerase II products, have been widely observed by others (27,28) and they were not investigated further.

The identification of the regions at 4 and 39 map units as functional RNA promoter regions both in vivo and in vitro led us to an examination of the nucleotide sequences in these areas. At 3.5 μm, nucleotide 175-181, there is a canonical TATA box (36,37), "TATATAA", on the viral complementary strand (Fig. 7a). These sequences reside approximately 60 nucleotides downstream from the 3' hairpin of MVM. At 39 μm, nucleotide 1976-1982, there is also a 7 nucleotide canonical TATA box, "TATAAAT", which is preceded 87 nucleotides upstream by a canonical CAT box (37,38), "CCATTCT", at nucleotides 1889-1895 (Fig. 7b).

DISCUSSION

We have identified and mapped the three major RNA transcripts, which together constitute greater than 97% of the total viral specific RNA.
Figure 7. Nucleotide sequence of MVM RF in the vicinity of map unit 4 (7A), and 39 (7B). The complete nucleotide sequence of MVM DNA is described in the accompanying paper (12).

produced by the autonomous parvovirus MVM during productive infection of murine A9 cells. These transcripts, which are 4.8, 3.3 and 3.0 kb respectively, are organized into two overlapping transcriptional units as illustrated in Fig. 5. Each of these transcripts is spliced in the region of mu 46 and each contains a similar exon of 2.6 kb between mu 48-96. Minor unspliced species of R1 and R3 have been detected at very low levels in total RNA preparations. These results suggest a more complex pattern of autonomous parvovirus transcription than has been previously described (10,11) and resembles in many ways the transcriptional topography described for the defective parvoviruses. The transcription organization of the defective parvovirus AAV-2 is that of three overlapping transcriptional units, encoding six RNA species, with each promoter (located at mu 5, 19 and 39) generating both a spliced and unspliced species of RNA (9). All the spliced species of AAV-2 RNA, although initiated at three different sites, share a similar exon which comprises the right half of the genome. The spliced RNAs also share a similarly sized and placed splice at mu 40-48. The size of this intron (~350 nucleotides) is, however, considerably larger than
for the analogous common intron at mu 46 that is spliced out of the MVM transcripts. In addition, while AAV-2 encodes two significant transcripts initiated at mu 19, we found no evidence for a third such promoter in MVM. No in vivo RNAs that mapped to that initiation site were detected, and in vitro transcription experiments with cloned DNA representing 6-25 mu of MVM RF DNA failed to demonstrate efficient in vitro promotion from that region of the genome (data not shown). The emerging picture of MVM transcription, while resembling that of the defective paroviruses, is still distinctly different.

There are a number of still unanswered questions raised by the data presented here. Foremost of these is the map position(s) of the low abundance 1.8 kb transcript R4. In addition, the results of Fig. 4A, lane 3 indicates that the small leader sequence between mu 40-46 is heterogeneous. Although this may be an artifact of the procedures used, it may reflect, as in SV40 (39,40) and polyoma (41) transcription, subtle differences in splice junctions either among individual members of the same transcript class or differences between the different class species. Additional discussion of the splice junctions in this area based on nucleotide analysis of MVM DNA is presented in the accompanying paper (12).

Previous work on the autonomous parovirus H1 (10) had revealed 5 viral RNA species, 4.7, 3.0, 2.8, 1.45 and 1.3 kb in length. The 4.7 kb transcript, although spliced, was found only in the nucleus, leading to the speculation that it was a nuclear precursor to the smaller transcripts found in the cytoplasm. We have detected the MVM 4.8 kb transcript both in the nuclear and cytoplasmic fractions. In addition, in vitro translation of R1, which had been selected by hybridization to cloned MVM DNA from nucleotide 1086-1661 (totally within the large intron), has recently been shown to generate at least one previously unidentified non structural protein (S. Cotmore and P. Tattersall, personal communication). We conclude from this that the 4.8 kb message appears in a functional form in the cytoplasm of infected cells.

We have presented here a more comprehensive map of MVM transcription products than previously available, however, precise definition of the map coordinates of all RNA species require further analysis. Such studies are currently in progress.

ACKNOWLEDGEMENTS

This work was supported by grants from the United States Public Health
Service (GM-20124 and CA-16038 to D.C.W) and by the Medical Research Council of Canada and British Columbia Health Care Research Foundation (to C.R.A.). We thank Peter Tattersall, Michael Merchlinsky, Jeffry Leary and Susan Cotmore for providing cloned stocks of MVM DNA and for many helpful comments and suggestions. D.P. was a Fellow in Cancer Research Supported by Grant DRG-374-F of the Damon Runyon-Walter Winchell Cancer Fund.

REFERENCES

Nucleic Acids Research