Nuclease mapping and DNA sequence analysis of transcripts from the dihydrofolate reductase-thymidylate synthase (R) region of Leishmania major

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ABSTRACT

Trypanosomatid protozoan parasites utilize a number of nonstandard mechanisms in expressing their genes. To probe these phenomena in a genetically accessible system, we have mapped termini of eight transcripts arising from the amplified R region including the DHFR-TS gene of methotrexate-resistant Leishmania major. Poly(A)+ RNAs transcribed from the DHFR-TS-coding strand exhibit features similar to those observed around other trypanosomatid protein-coding genes. These include close spacing, the presence of a trans-spliced minilexon at the 5' termini, heterogeneity at both 5' and 3' ends, and in some cases S1 nuclease protection of intertranscript regions. Other than the splice acceptor site, no consensus sequence elements associated with either 5' or 3' ends were detected, although polydinucleotide tracts tended to be near intertranscript regions. Two poly(A)+ RNAs transcribed from the opposite strand of the upstream flanking regions lacked the minilexon. Sequencing of DNA encoding the overlapping 1.7 kb opposite strand transcripts (one bearing and one lacking the minilexon, both found on polysomes) revealed no reading frames likely to encode proteins, suggesting that at least some of these RNAs could be non-functional by-products of RNA processing.

INTRODUCTION

Trypanosomatid protozoa employ a variety of novel processes to express their genes, suggesting that the rules governing transcription are different from those of other eukaryotes (1–8). Transcription of trypanosomatid protein-coding genes yields a series of polyadenylated RNAs encoded in close proximity within the DNA, separated by short intertranscript regions. These RNAs bear the trans-spliced 39 nucleotide minilexon at their 5' end. In contrast to most other species intertranscript regions are generally transcribed at a rate similar to that of the regions encoding adjacent mature transcripts. This suggested the possibility that transcription occurs in a ‘polycistronic’ fashion, and that processing generates multiple mature mRNAs (7; 9–13). Consistent with the polycistronic model, conserved DNA sequences which commonly mark promoters in other organisms are usually absent in the immediate upstream regions of trypanosomatid genes (reviewed in 8). For some protein-coding genes, other transcripts arising from the putative polycistronic domain appear to lack significant open reading frames (14). These transcripts may be non-functional by-products analogous to introns, although these RNAs occur on polysomes (14; 15). Many questions remain concerning the specific mechanisms and results of polycistronic transcription in the loci studied thus far, and it is unknown whether polycistronic transcription is a universal feature of all trypanosomatid genes. Genetic tests of polycistronic vs. monocistronic transcription has not yet been obtained due to the very recent emergence of DNA transfection techniques (16–19).

We have utilized drug-resistant lines of Leishmania major for addressing a variety of molecular and biochemical questions in trypanosomatids. Methotrexate (MTX) resistant mutants often exhibit selective gene amplification of two separate regions of DNA, the R and H regions. The R region, encodes the bifunctional enzyme dihydrofolate reductase-thymidylate synthase (DHFR-TS; 20, 21), a normally single-copy gene which is amplified in the R1000-3 line as a 30 kilobase (kb) extrachromosomal DNA (22; Fig. 1). Previous analysis of the DNA sequence immediately upstream of the DHFR-TS mRNA did not identify conserved sequences present in other DHFR and TS genes which are known to promote transcription and bind trans-acting factors (23–27). Analysis of the transcripts of the R region revealed that approximately 95% of it is transcribed into 9–10 stable polyadenylated RNAs (Fig. 1), all of which are associated with polysomes (15). These include the ‘downstream’ RNAs, which are transcribed from the same DNA strand and overlap one another to varying extents, and two examples of opposite-strand ‘antisense’ RNAs localized to the
divergent RNA subregion, upstream of DHFR-TS (Fig. 1). R region transcripts were identical in structure in the R1000-3 and wild-type lines, indicating that all cis-acting elements necessary for transcription initiation and RNA processing reside within the amplified region (15; 19). In this report we have characterized transcripts of the amplified R region in more detail by nuclease protection studies and DNA sequencing.

METHODS AND MATERIALS

Parasites

Promastigotes (the insect stage of this digenetic parasite) of the R1000-3 line of Leishmania major (28) were grown in M199 medium as described (19) containing 1 mM MTX. Large scale cultures were grown in medium containing 100 μM MTX.

RNA isolation, S1 nuclease mapping and primer extension

Total and poly(A) RNA were prepared as described (29,30). S1 nuclease mapping was performed using 2 μg of total cellular poly(A) RNA (23). Primer extension was performed using specific oligonucleotide primers (at least 20 bases) whose 5' ends were located as described in the text and 2 μg of total cellular poly(A) RNA (15). Product sizes were estimated from multiple experiments using three different sets of molecular weight markers, and in all cases the standard deviation was less than 5 nucleotides.

Polymerase chain reaction (PCR) amplification of cDNA

cDNA synthesized from RNA-specific primers and 0.2 μg poly(A) RNA was subjected to PCR amplification as previously described (15,31), using the cDNA synthesis primer and either miniexon primer A (GGGAATTCCGATCC/AACGGTATATAAGTGTAC), which contains nucleotides (nt) 1 to 19 of the L. major miniexon (J. Miller, submitted for publication) and a 14 nt 5' extension, or miniexon primer B (TCAGTTTCTGTACTTTATTG) which contains nt 16-35 of the miniexon.

DNA sequence analysis

DNAs were cloned into M13mp18 and/or M13mp19 vectors (32) and sequenced by the dyeodeoxy method with modified T7 polymerase (Sequenase; U.S. Biochemical, Cleveland, Ohio). The complete sequence of both DNA strands was obtained. The sequence of the Sac I fragment containing the L. major DHFR-TS sequence (including Fig. 2B) was assigned accession number X51733 and the sequence of the divergent region (Fig. 5A) was assigned X51734.

RESULTS

RNA termini mapping

Downstream and DHFR-TS subregions. Northern blot mapping of the R region indicated that the subregion 3' to the DHFR-TS mRNA encodes at least five RNAs collectively termed the downstream RNAs (3.2, 2.8, 2.5 and 2.3 kb), which are all transcribed from the same strand as DHFR-TS and overlap to varying extents (Fig. 1; 15); the 5' ends of the 3.2 and 2.8 kb downstream RNAs mapped to a 0.6 kb SalI-SphI fragment of this region (Fig. 2B; 15). S1 nuclease analysis of this region produced a predominant protected fragment, mapping the 5' end of the 3.2 and 2.8 kb downstream RNAs 405 bp to the right of the SalI site; 3 additional minor 5' ends were mapped 280, 385 and 430 bp to the right of this site (Fig. 2B). Since Northern blot hybridization with this fragment indicated that the 3.2 and 2.8 kb downstream RNAs were equally abundant (15), we presume that both RNAs utilize the major 5' terminus identified here.

Thus, these data suggest that the 3.2 and 2.8 kb downstream RNAs have the miniexon at their 5' end. To confirm this, cDNA generated with the downstream RNA primer was subjected to PCR amplification with this primer in combination with either of two miniexon primers, A and B, which contain sequences from the 5' and 3' halves of the miniexon, respectively. If the miniexon were present, PCR amplification products would be present and be 30 nt larger when amplified with miniexon primer A than with primer B (miniexon primer A contains a 14 nt 5' extension, or miniexon primer B (TCAGTTTCTGTACTTTATTG) which contains nt 16-35 of the miniexon.

Aplification of cDNA synthesized from a downstream RNA-specific primer in conjunction with the A primer produced a major 385 bp product, while amplification with the B primer produced a major 360 bp product (Fig. 3; the size was determined more accurately from gels containing less DNA, not shown); other products may arise from alternative 5' ends bearing miniexons. The PCR products were specific, since they hybridized to a downstream RNA-specific probe and their appearance was dependent on the presence of both primers (data not shown). The sizes estimated for these PCR products were slightly smaller than expected, based on the S1 and primer extension products discussed above. We attribute this discrepancy to differences in gel electrophoresis conditions, since a similar discrepancy was observed in our analysis of the DHFR-TS mRNA (19) for which the cDNA sequence had been determined. Thus, these data suggest that the 3.2 and 2.8 kb downstream RNAs have the miniexon at their 5' end.

S1 nuclease mapping of the 3' end of the DHFR-TS mRNA (Fig. 2A) revealed a predominant protected fragment terminating 560 bp to the left of the PvuII site, in addition to less abundant 3' ends mapping 470, 590 and 710 bp to the left of this site (Fig. 2B). A small amount of full length protection was also observed. The 560 bp site agrees well with that of a previously sequenced cDNA (20).
These data identify a 440 bp intertranscript region between the major 5' downstream and 3' DHFR-TS-TS ends (Fig. 2C). This distance drops to 260 bp if minor ends are considered, and a small amount of RNA appears to span this region.

**DHFR-TS subregion.** Previous work has shown that the DHFR-TS mRNA originates within the 0.7 kb EcoRI-SalI fragment of this region (Figs. 1 and 2B; 15), and that the DHFR-TS mRNA has two major 5' ends mapping 253 and 388 bp to the right of the PvuII site and two minor ends mapping 208 and 308 bp to the right of this site (Fig. 2B; 23). A small amount of fully protected probe was also observed (23).

Primer extension of the DHFR-TS mRNA produced two major products with apparent 5' ends mapping 290 and 415 bp to the right of the PvuII site (Fig. 2A), approximately 30–35 bp beyond the major S1 sites (Fig. 2B, marked by dashed lines). PCR amplification of DHFR-TS cDNA with the DHFR-TS primer and the miniexon-A primer produced two products whose sizes (280 and 400 bp) were similar to those obtained in primer extension (Fig. 3). Previous DNA sequencing of the predominant 280 bp product showed that it contains a complete miniexon on its 5' end (19).

S1 nuclease mapping of the 3' end of the 13 kb RNA (Fig. 2A) identified three major protected products whose termini map 125, 130 and 135 bp to the left of the SalI site and two less abundant termini mapping 105 and 305 bp to the left of this site (Fig. 2B). These data identify a 445 bp intertranscript region between the most abundant 5' DHFR-TS and 3' 13 kb RNA ends (Fig. 2C). The intertranscript distance drops to 250 bp, if minor ends are considered, and a small amount of RNA may span this region (23).

**Leftwards-transcribed RNAs of the divergent RNA subregion.** The divergent RNA subregion (Fig. 1) contains the 5' end of the 13 kb RNA and encodes four additional RNAs (1.7-l, 1.7-r, 6.2-l and 6.2-r) which are transcribed from opposite strands and extensively overlap one another (Fig. 1; 15). The suffixes -l (leftward) and -r (rightward) denote the direction of transcription, with the leftward RNAs being transcribed in the same direction as DHFR-TS.

Previous Northern blotting showed that the 5' end of 13 kb RNA and 3' end of the 1.7-l RNA map within the 0.8 kb SalI-SphI fragment of this region (Fig. 4B; 15). S1 nuclease mapping of the 5' end of the 13 kb RNA (Fig. 4A) revealed considerable heterogeneity, with the most abundant protected
the 13 kb RNA bears the miniexon (Fig. 4B, dashed line), PCR products correlations among the less abundant products were however due to the large number of S1, primer extension and presence of both RNA-specific and miniexon primers (data not shown). The ability to amplify cDNAs mapping to the vicinity of larger products. PCR amplification with a primer whose 5' 3' 530 bp to the right of this site (Fig. 4B; 15). Primer extension of the 1.7-/ RNA (Fig. 4A) identified a predominant 3' end mapping 515 bp to the right of the Sall site, 55–60 bp longer than the S1 products. PCR amplification of the 1.7-/cDNA with the primer extension oligonucleotide in combination with miniexon primers A or B produced a major product of 205 bp with miniexon primer A and 175 bp with miniexon primer B (Fig. 3). These products hybridized with a probe specific for this region and their occurrence was dependent on having both RNA- and miniexon-specific primers in the reaction (data not shown). These data suggest that the 1.7-/ RNA has a miniexon on its 5' end.

Previous Northern blot analysis mapped the 3' end of the 6.2-/ RNA to the 0.9 kb Sall-Pstl fragment of this region (Fig. 4B; 15). S1 nuclease mapping of the 3' end of the 6.2-/ RNA (Fig. 4A) identified a major 3' end terminating 120 bp to the left of the Sphl site and minor termini mapping 165 and 280 bp to the left of this site (Fig. 4B). These analyses identified a 470 bp intertranscript region between the major 5' 1.7-/ and 3' 6.2-/ RNA termini, although if minor ends are considered this distance is 310 bp (Fig. 4C).

Rightwards RNAs of the divergent RNA region. Previous Northern blot analysis mapped the 5' end of the 1.7-r RNA to the 0.55 kb Sphl-Sphl site (Fig. 4D; 15). Multiple attempts to map the 5' end of the 1.7-r RNA by S1 nuclease protection with the 1.4 kb Sstl-Sall or 1.1 kb Sstl-Pstl fragments labeled on their 5' end at the Sall or Pstl sites (Fig. 4B) were unsuccessful, even though mixing experiments with a downstream RNA probe and the 1.7-r RNA Sstl-Pstl probe yielded a protected fragment corresponding to the major downstream RNA 5' end (Fig. 4A, lanes labeled Mixing). These results were perplexing, since S1 nuclease mapping of the 1.7-r RNA with a probe spanning the internal portion of this RNA (Pstl-HindIII, Fig. 4B) and overlapping the Sstl-Sall probe successfully yielded full length protection of the Leishmania sequences (Fig. 4D), as would be predicted for a contiguous 1.7-r RNA spanning this segment. This experiment also suggests that the formation of RNA-RNA duplexes cannot account for the failure to protect a fragment located at the 5' terminus. DNA sequence analysis of this region (below) revealed no unusual sequence organization or elements that could account for these discrepant S1 results. Previously, a probe labeled at the Sall site identified a predominant 3' end mapping 620–630 bp to the right of the Sall site (Fig. 4D; 15).

In contrast to the results obtained by S1 nuclease protection, primer extension mapping of the 5' end of the 1.7-/- RNA with a primer labeled at the Pstl site (Fig. 4B) yielded three comparably abundant products (Fig. 4D) with apparent 5' ends mapping 120, 135 and 140 bp to the left of the Pstl site. A second primer situated 95 bp to the right of the Pstl site
Figure 4. S1 nuclease and primer extension analysis of RNA termini within the divergent RNA subregion. Panel A. S1 nuclease and primer extension analysis. For locations of sites refer to the map shown in panel B. Symbols: +, 2 μg R1000-3 poly(A) RNA plus S1 nuclease; −, S1 nuclease alone; P, undigested probe. 5'S1, 3'S1, and P refer to S1 nuclease mapping of 5' and 3' termini of RNA and primer extension, respectively. DNA size markers (kb): MspI digested pHRR22 DNA (not all fragment sizes are indicated). Additional higher molecular weight markers (6X HindIII fragments) are indicated where appropriate. 13 kb: left panel, S1 nuclease mapping of the 5' end of the 13 kb RNA (lanes labeled 13 kb, 5'S1) with the 1.4 kb SalI-PstI fragment (positions -13.0 to -14.4) labeled on the 5' end of the SalI site; right panel, primer extension of the 13 kb RNA with primers labeled at the SalI site (lane labeled 13 kb, PE 1) or at the position 353 nt to the right of the SalI site (lane labeled 13 kb, PE 2); 1.7-f: left panel, S1 nuclease mapping of the 3' end of the 1.7-f RNA with a 3.75 kb SphI-BamHI fragment containing 0.75 kb of *Leishmania* DNA (SphI-SphI positions -16.0 to -16.0) and 3.0 kb of vector DNA labeled on the 3' end of the SphI site (lane labeled 1.7-f, SI); 6.2-f: left panel, S1 nuclease mapping of an internal segment of the 1.7-f RNA (lanes labeled 1.7-f, 3'S1) with a 5.0 kb PstI-BamHI fragment containing 1.0 kb of *Leishmania* DNA (PstI-HindIII positions -14.4 to -15.4) and 4.0 kb of vector DNA, labeled at the 3' end of the PstI site; right panel: primer extension of the 1.7-f RNA with a primer labeled on the 5' end at the PstI site (lane labeled 1.7-f, PE); 6.2-f: S1 nuclease mapping of the 3' end of the 6.2-f RNA with a 3.75 kb SphI-BamHI fragment containing 0.75 kb of *Leishmania* DNA (SphI-SphI, positions -15.2 to -16.0) and 3.0 kb of vector DNA labeled on the 3' end of the SphI site (lane labeled 6.2-f, SI); 1.7-r: left panel: S1 nuclease mapping of an internal segment of the 1.7-r RNA (lanes labeled 1.7-r, 3'S1) with a 5.0 kb PstI-BamHI fragment containing 1.0 kb of *Leishmania* DNA (PstI-HindIII positions -14.4 to -15.4) and 4.0 kb of vector DNA, labeled at the 3' end of the PstI site; right panel: primer extension of the 1.7-r RNA with a primer labeled on the 5' end at the PstI site (lane labeled 1.7-r, PE); 6.2-r: left panel, S1 nuclease mapping of the 5' end of the 6.2-r RNA with a 3.75 kb SphI-BamHI fragment containing 0.75 kb of *Leishmania* DNA (SphI-SphI positions -15.2 to -16.0) and 3.0 kb of vector DNA labeled on the 5' end of the SphI site (lane labeled 6.2-r, 5'S1); right panel, primer extension of the 6.2-r RNA with a primer labeled on the 5' end at the SphI site (lane labeled 6.2-r, PE); Mixing: Control S1 mapping experiment (lanes labeled mixing): 5' probes designed for the 5' end of the 1.7-r kb RNA probes (1.15 kb SalI-PstI fragment (positions -13.2 to -14.4) labeled on the 5' end of the SalI site and 5' end of the downstream RNAs (see Fig. 2B) were mixed and used for standard S1 nuclease digestion. The pattern shown is the same as that obtained with the downstream probe alone (see Fig. 2A). Panel B. Restriction map and summary of results obtained for leftwards RNAs. Results are shown using the conventions described in Fig. 2B. H, HindIII; I, SphI; P, PstI; S, SalI; T, SstI. Sites present more than once are distinguished with subscripts. 5' products of the 5' end of the 1.7-r RNA were previously determined (15). 5' and 3' refer to the ends that were mapped. Panel C. Summary of Transcript Organization. The RNAs of this portion of the divergent region are shown using the conventions described in Fig. 2C. Only the major RNAs are shown. Vertical tick marks represent termini. Panel D. Summary of results obtained for rightwards RNAs. Data are shown using the conventions described in Fig. 2B. S1 products of the 3' end of the 1.7-r RNA were previously determined (15).
produced three major primer extension products mapping to these same apparent positions (data not shown). The distance between the 5' and 3' ends of the 1.7-r RNA, mapped by primer extension and S1 nuclease, respectively, is approximately 1.6 kb, consistent with the size estimated by Northern blot analysis.

Previous Northern blots mapped the 5' end of the 6.2-r RNA to the 0.9 kb Sall-PstI fragment of this region (Fig. 4B; 15). S1 nuclease mapping of the 5' end of the 6.2-r RNA (Fig. 4A) mapped a major 5' end 560 bp to the left of the Sphl site (Fig. 4D), as well as a small amount of full length protection of the Leishmania sequences within this probe (750 nt fragment, Fig. 4A). Unlike the leftward transcribed RNAs which contain gaps immediately 3' of the AG marking the splice acceptor site for trans-splicing (Fig. 5). Most minor termini revealed by S1 protection analysis also mapped close to AG residues, although exceptions were noted (13K 5' ends located at positions -13447, -13540, -13550; downstream 5' end located at position +2900; Fig. 5).

Table 1 shows an alignment of the four miniexon addition sites of the leftwards RNAs and three other Leishmania genes. Most Leishmania splice acceptor sequences are flanked by pyrimidine-rich regions on both the 5' and 3' sides, whereas generally eukaryotic splice acceptor sequences show pyrimidine-richness only on the 5' side (33). This may reflect the fact that the splice acceptor sequence in trypanosomatids is invariably non-coding. Sequences related to the hexanucleotide CTTCC(T/C) were found in more than 20 positions, both 5' and 3' of the AG marking the splice acceptor site for all seven transcripts shown in Table 1.

One common sequence motif identified in the vicinity of the AG splice acceptor dinucleotide was the pentamer CGCAC. This or related sequences were found in more than 20 positions, both 5' and 3' of the AG dinucleotide. This sequence is complementary to the precursor RNA which donates the miniexon (termed the med-RNA) near the 5' splice donor site (Table 1). Layden and Eisen (34) have proposed that base-pairing interactions may serve to guide the site of the trans-splicing reaction, although the regions of complementarity that they identified were variable and frequently adjacent to the region complementary to CGCAC. However, the CGCAC sequence was found both upstream and downstream of the splice acceptor site as well as throughout the sequenced regions of Leishmania DNA, and other additional for shared or conserved sequences which could mark functionally important signals. All major 5' termini identified by S1 nuclease analysis mapped close to AG dinucleotides required for trans-splicing (Fig. 5).
Table 1. Sequence alignments of the 3' splice junction of *Leishmania* mRNAs

<table>
<thead>
<tr>
<th>mRNA</th>
<th>3' Splice Junction Alignment</th>
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<tr>
<td>DHFR-TS</td>
<td>GCTGGAATGA...CGCAATTCCAA</td>
</tr>
<tr>
<td>DWastr</td>
<td>AGCGGGCA...CTCTTCAGTCAG</td>
</tr>
<tr>
<td>1.7-l</td>
<td>CCTCTCTCTCACATCAAGAGAG</td>
</tr>
<tr>
<td>1.7-l</td>
<td>ACTCTCTCTCACATCAAGAGAG</td>
</tr>
<tr>
<td>13K</td>
<td>GCTGGAATGA...CGCAATTCCAA</td>
</tr>
<tr>
<td>hsp70</td>
<td>GGCGGGCGG...CGCAATTCCAA</td>
</tr>
<tr>
<td>a-TUB</td>
<td>AGCGGGCA...CTCTTCAGTCAG</td>
</tr>
<tr>
<td>B-TUB</td>
<td>GCTGGAATGA...CGCAATTCCAA</td>
</tr>
</tbody>
</table>

The splice junctions associated with the major termini of the RNAs listed are shown, with a '/' marking the splice site, pyrimidine rich regions marked by underlining, and the CGCAC motifs discussed in the text marked in bold type. Sequences are DHFR-TS, Dwastr (3.2 and 2.8 kb RNAs, Figs. 2B, 13K and 1.7-l (Fig. 4C), all from the R region of *L. major* (23; this work); HSP70, 70K heat shock protein homolog of *L. major* (53); α-TUB and β-TUB, α- and β-tubulins respectively of *L. enriettii* (42); med-RNA, the minicron precursor RNA sequence of *L. enriettii* (54). The '1' in this sequence denotes the splice donor site. The sequence shown of the med-RNA is identical in all *Leishmania* species studied thus far (55).

sequences potentially capable of base-pairing with the med-RNA were observed (not shown). These considerations suggest that the assignment of a role in trans-splicing to these sequences will require functional tests.

Analysis of the DNA flanking the 3' ends of the R region RNAs did not identify sequences which are conserved amongst these regions. These regions did not exhibit similarity with the prototypic polyadenylation signals of higher eukaryotes or yeast (35–37). The 3' end regions also tended to be flanked by pyrimidine rich segments, and short direct repeats specific to a given transcript were commonly found in the vicinity of the 3' ends.

Curiously, the predominant 3' ends of the 1.7-r and 6.2-r RNAs localized to similar positions, as did the 5' ends of the 1.7-l and 6.2-r RNAs. Using the method of Zucker (38), we examined these two regions for potential secondary structures which might exist in DNA or RNA. A potential hairpin structure with an estimated ΔG of −19.7 kcal/mol was observed only for the DNA surrounding the divergent 5' ends.

The intertranscript regions were searched for conserved or consensus sequence elements found in other species, however no similarities were identified which displayed a conserved position relative to the 5' or 3' RNA termini. Specific searches for elements such as TATAAT and CAAT boxes (39), binding sites for the transcription factor SP1 (GGGGCGG; 40), and the yeast splice acceptor branch site TACTAAC (41), among others, did not identify sequences which are conserved amongst these RNAs. Using the method of Zucker (38), we examined these two regions for potential secondary structures which might exist in DNA or RNA. A potential hairpin structure with an estimated ΔG of −19.7 kcal/mol was observed only for the DNA surrounding the divergent 5' ends.

Protein-coding potential of the 1.7 kb RNAs

The region encompassing the overlapping opposite-strand 1.7-l and 1.7-r RNAs was analyzed for open reading frames (ORFs), assuming initiation only at AUGs and the standard genetic code. The largest ORFs within the 1.7-l RNA were 210, 252 and 261 bp in length, with the most 5' of these beginning 900 nucleotides from the 5' end of the RNA and preceded by at least 6 AUGs and 4 smaller ORFs. Quantitative analysis of codon usage bias of these ORFs revealed them to be compositionally random and not in agreement with that of other *Leishmania* proteins (20; unpublished). The 1.7-r RNA exhibited one large 579 nt ORF, however this ORF began 650 bases from the 5' end of the mRNA, was proceeded by 10 AUGs and 5 smaller ORFs, and the predicted protein did not conform to the expected *Leishmania* protein codon bias. These results suggested that it was unlikely that either of the two 1.7 kb RNAs encoded proteins.

DISCUSSION

To further our understanding of the transcriptional organization of the DHFR-TS gene and surrounding R region, we have mapped the termini of 8 RNAs transcribed from the amplified R region of *Leishmania major* and sequenced the DNA encompassing 6 inter-transcript regions and two opposite strand transcripts. Heterogeneity at both 5' and 3' ends of the RNAs was commonly observed, the most extreme case being the 13 kb RNA which utilized at least eight 5' and four 3' termini. Whether the 5' and 3' ends are determined independently, as for the mouse DHFR gene (43), or in limited combinations has not been assessed. Our data show that the predominant termini of the leftwards RNAs contain minicexons at their 5' ends. Many of the minor termini appear to contain minicexons as well, and thus must be the result of alternative trans-splicing. Although alternative cis-splicing can play important functional roles in the expression of higher eukaryotic genes, this does not appear to be the case for the DHFR-TS mRNA since the 5' heterogeneity is invariant in promastigotes and amastigotes (23).

While many of the 5' ends of the R region RNAs contain minicexons, it is clear that some do not. Two of these are major RNAs, the rightwards-transcribed 1.7-r and 6.2-r RNAs, which partially overlap. Since one potential role of the minicexon is to provide the 5' cap structure required for translation, it would be expected that the RNAs lacking minicexons would be non-coding, and this appears to be the case for the 1.7-r RNA. However, the DNA sequence of the opposite-strand 1.7-l RNA, which bears the minicexon, also did not reveal clear evidence for an encoded protein. Both of these RNAs have previously been shown to occur on polysomes (15). Several mechanisms could account for these results: it is possible that certain termination codons are being suppressed, or that mechanisms such as nuclear RNA editing or RNA modification mediated by pairing of the antisense transcripts (44,45) could transform the genomic sequence into a protein-coding reading frame. If present, these modifications were not detected by S1 mapping using genomic DNA probes and have not been previously reported in trypanosomatids. In support of the sequencing results, two-dimensional gel analysis of proteins of R region-amplified *Leishmania* failed to reveal any over-expressed proteins other...
than DHFR-TS (46). Noncoding polysomal RNAs in trypanosomes have been reported (14), though precedents for non-translated polysomal RNAs are rare in other species (47–49).

In keeping with work from other trypanosome genes, other than limited similarity in the vicinity of the minixenon splice acceptor sites no conservation of universal sequence motifs at key positions around transcripts of the DHFR-TS region was observed. This is not surprising since one common eukaryotic signal, that marking poly-adenylation, has been shown to be non-functional in transfected derivatives of the R region (19). One feature of the intertranscript regions was the presence of poly(dC)-repeat sequences, most commonly poly-CA, which are also observed in other Leishmania genes. Poly-(CA) tracts, which are capable of adopting a lefthanded DNA conformation, can enhance transcription in mammalian cells and when transcribed form cytoplasmic ‘Z-RNA’ in trypanosomatids (50; 51; 52). It is possible that the dinucleotide tracts constitute loose signals for transcription or processing.

Mapping of the transcripts adjacent to the DHFR-TS mRNA has allowed us to consider these RNAs in the context of the current ‘polycistronic’ model for transcription in trypanosomatids. Transcripts of the R region exhibit several hallmarks associated with this model: 1) A series of closely juxtaposed RNAs transcribed from the same strand (6.2-, 1.7-, 13K, DHFR-TS, downstream RNAs), all bearing the trans-spliced miniexon. 2) Occasional protection of inter-transcript regions in S1 nuclease protection assays of total cellular poly-adenylated RNA preparations, not enriched for transcriptional intermediates. Had nuclear RNA been employed, the occurrence and amount of RNA apparently non-functional RNAs within this putative ‘polycistronic’ domain, minimally the 1.7- and 1.7-r RNAs and perhaps many of the others. At a structural level, these data are consistent with evidence obtained from other trypanosomatid loci thought to exhibit polycistronic transcription by these and additional criteria. In contrast, the transcripts proceeding in the rightwards direction lack many of these features, and may arise from an independent mechanism.

Although consistent with the polycistronic transcriptional model, our data do not rule out other models for transcription of the R region. Independent transcription of each RNA could occur, assuming that initiation was sufficiently far upstream to accommodate the trans-splicing mechanism as well as yielding at least some RNAs which span the inter-transcript regions. The emerging methods of DNA transfection of parasites promise to resolve this issue, as a polycistronic DHFR-TS transcription model predicts that transcription should be dependent upon specific promoters located at some considerable distance upstream of the mature DHFR-TS mRNA.

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REFERENCES
