Establishment of an \textit{in vitro} trans-splicing system in \textit{Trypanosoma brucei} that requires endogenous spliced leader RNA

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Received October 22, 2009; Revised January 19, 2010; Accepted January 22, 2010

ABSTRACT

In trypanosomes a 39 nucleotide exon, the spliced leader (SL) is donated to all mRNAs from a small RNA, the SL RNA, by trans-splicing. Since the discovery of trans-splicing in trypanosomes two decades ago, numerous attempts failed to reconstitute the reaction \textit{in vitro}. In this study, a crude whole-cell extract utilizing the endogenous SL RNA and synthetic tubulin pre-mRNA were used to reconstitute the trans-splicing reaction. An RNase protection assay was used to detect the trans-spliced product. The reaction was optimized and shown to depend on ATP and intact U2 and U6 snRNPs. Mutations introduced at the polypyrimidine tract and the AG splice site reduced the reaction efficiency. To simplify the assay, RT–PCR and quantitative real-time PCR assays were established. The system was used to examine the structural requirements for SL RNA as a substrate in the reaction. Interestingly, synthetic SL RNA assembled poorly to its cognate particle and was not utilized in the reaction. However, SL RNA synthesized in cells lacking Sm proteins, which is defective in cap-4 modification, was active in the reaction. This study is the first step towards further elucidating the mechanism of trans-splicing, an essential reaction which determines the trypanosome transcriptome.

INTRODUCTION

mRNA maturation in trypanosomes differs from the process in most eukaryotes mainly because in trypanosomes, protein coding genes are transcribed into polycistronic RNAs. The polycistronic RNAs are dissected to individual mature mRNAs by the concerted action of trans-splicing and polyadenylation. Thus, trans-splicing functions to release mature capped mRNAs from the polycistronic RNA. In trans-splicing, a small RNA, the spliced leader RNA (SL RNA) harboring a special cap-4 and composed of a conserved exon (SL) and a more divergent intron, donates its exon to all mRNAs (1,2). Trans-splicing, like cis-splicing, proceeds through a two-step transesterification reaction, though a Y structure is formed instead of a lariat intermediate. Although first discovered in trypanosomes, trans-splicing was later found in nematodes, trematodes and even chordates (3), and more recently also in dinoflagellates (4). Cis-splicing also occurs in trypanosomes, but so far only three substrates were shown to undergo this process (5).

SL RNA is transcribed from a RNA polymerase II promoter (6,7). SL RNA undergoes modification to form the unique hypermethylated cap-4 (8). However, eliminating an individual cap modification does not interfere with the ability of the SL RNA to undergo trans-splicing (9). The SL RNA binds core Sm proteins, which are assisted by SMN and Gemin2 (10). SL RNA undergoes pseudouridylation by SLA1, which serves as a chaperone during SL RNA biogenesis. SL RNA transcription, pseudouridylation and assembly with core Sm proteins take place in a special nuclear location that we termed the SL RNP factory (11,12). The majority of SL RNA in the steady state is concentrated in this sub-domain (12,13). It is not currently known how the SL RNP is recruited to the trans-spliceosome.

Recently, much progress was made in identifying a variety of trypanosome splicing factors, and their functions were elucidated by down-regulation via RNAi (10,12,14–19). Splicing defects such as an increase in the
level of SL RNA, and changes in the level of the Y structure intermediate were observed following the depletion of each splicing factor (18, 19). Using this approach, the factors U2AF35, U2AF65 and SF1, which function to identify and specify the correct 3’ AG splice sites, were recently identified. The Trypanosoma brucei U2AF65 is larger than its mammalian counterpart, and surprisingly, unlike the protein’s interactions in metazoa, it interacts with SF1 but not with U2AF35 (20). An ~45 S spliceosomal complex was identified, suggesting the existence of a single-spliceosome complex that can potentially conduct both cis- and trans-splicing (19).

In vitro splicing reactions were established in mammalian and yeast extracts >25 years ago (21–23). A very efficient in vitro trans-splicing reaction was also established in nematodes using Ascaris lumbricoides whole-cell extracts derived from developing embryos. The reaction is very efficient and is based on a synthetic pre-mRNA acceptor and SL RNA derived from the extract, or synthetic radiolabeled SL RNA (24, 25). The in vitro cis-splicing systems in both mammals and yeast required different manipulations in order to optimize the reactions. Although whole-cell extracts were used in the mammalian system, nuclear extracts were later found to be more efficient (21). The in vitro reaction is more efficient using ‘tailored’ RNA molecules harboring introns which are derived from transcripts that are spliced in vivo with high efficiency. For instance, the first splicing substrate was constructed using exons and an intron from the adenovirus transcription unit (21, 23) and a better substrate was subsequently obtained by shortening the exon size (26). It was demonstrated in the mammalian system that the pre-mRNA does not require either a cap at its 5’-end or a poly-(A) tail for its splicing, but splicing is enhanced by the addition of a cap, probably due to stabilization of the pre-mRNA (27). In yeast, a cell-free splicing system was first established using a sensitive S1 nuclease assay monitoring the production of the spliced product, and later, radiolabeled pre-mRNA was used and the spliced product and intermediates were readily detected (22). Despite overcoming many hurdles in establishing in vitro splicing systems in mammals and yeast as mentioned above, the insights obtained did not prove to be useful for creating such an in vitro trans-splicing reaction in trypanosomes.

In this study, we established an in vitro trans-splicing system using substrates whose trans-splicing was first verified in vivo in transgenic parasites expressing these constructs. The model substrate contains the splicing signals of the β-tubulin gene fused to luciferase and the upstream polyadenylation signal of the z-tubulin gene. Three sensitive assays were established to monitor the production of the trans-spliced product, RNase protection, RT–PCR and a quantitative real-time PCR (qRT–PCR) assay. Trans-splicing requires ATP, is sensitive to heat-inactivation, and is dependent on U2 and U6 snRNAs. The reaction is inhibited when the AG splice site or the polypyrimidine tract are mutated. Mutant SL RNA, lacking the cap-4 +4 modification, can be utilized in trans-splicing, and synthetic cap-4 does not inhibit the reaction. Optimization of the reaction conditions indicates that SL RNA is the limiting component in the extract, but only endogenous and not m’G capped synthetic SL RNA is active in the reaction. This system can be further used to investigate in depth the mechanism and fidelity of 3’ splice site selection.

**MATERIALS AND METHODS**

**Cell growth, transfection and cloning**

All experiments were performed with the procyclic form of T. brucei strain 29–13. Cell culture and preparation of transgenic parasites was as previously described (14).

**Plasmids**

pNS21 was a kind gift from George Cross (Rockefeller University, USA). This plasmid contains the T. brucei procyclin promoter, followed by a linker containing a BglII restriction site, the procyclin (EP) 3’ splice site and synthetic 5’UTR, fused to the luciferase ORF that contains an XbaI restriction site 47-nt downstream from the ATG start codon. This vector contains the phleomycin resistance gene (28). To create plasmids carrying the splicing signals of β-tubulin or EP, the corresponding sequences were amplified from the genome and were cloned between the restriction sites BglII and XbaI (28). To generate the vector pNS21-TIR carrying the β-tubulin intergenic region, a PCR product was generated using forward primer (233-Tub-luc f1) and reverse primer (234-Tub-Luc.r1) to create the correct fusion. Similarly, the pNS21-PIR was generated by amplifying the EP-2-3 (Tb927.6.520; positions 226182-227114 on chromosome 6) with forward primer (246-EP-luc f1) and reverse primer (247-EP-luc r1) to create a fusion with the luciferase gene. The plasmids were linearized with NotI for integration into the rRNA locus. RNA transgenic parasites was used to prepare cDNA using the Reverse-iT (Thermo Scientific) kit with random primers. The cDNA was amplified using SL forward primer (220, SL RNA sense) and the luciferase reverse primer (575, T7 Luc 52-67 as). These cDNA clones were used to prepare the probes for the RNase protection assay described below. The oligonucleotide list is given in Supplementary Data 1.

**Mutants of pNS21-TIR**

The mutations that changed the polypyrimidine tract (PPTm), AG to AA, and the pNS21-TIRsub were generated by site directed mutagenesis using the Mega primer method. The mutants were generated using a forward primer and a reverse primer carrying the mutation, and a second pair composed of a sense primer carrying the mutation and a reverse primer containing the XbaI site (234). The two fragments carrying the mutation were amplified using the primer pair 233 and 234. The mutated fragments were cloned between the XbaI and BglII site of pNS21, and the plasmids were sequenced and used to prepare transgenic cell lines. cDNA from these cell lines were cloned and sequenced. The pairs of oligonucleotides used to generate the mutations are listed in Supplementary Data 1.
**In vitro trans-splicing extracts**

Extracts were prepared essentially as described (29), with minor changes. First, 21 of *T. brucei* 29-13 were grown in a 51 conical flask under constant stirring to a density of 1 x 10^7 cells/ml. Cells were harvested at 4°C, yielding a packed cell volume of around 1.5 ml. Cell pellets were washed twice in 10 ml of ice-cold wash solution (20 mM Tris–HCl (pH 7.4), 100 mM NaCl, 3 mM MgCl2, 1 mM EDTA) and once in 10 ml of ice-cold transcription buffer (150 mM sucrose, 20 mM potassium 1-glutamate, 3 mM MgCl2, 20 mM HEPES-KOH (pH 7.7), 2 mM dithiothreitol, and leupeptin (10 μg/ml)]. Finally, pellets were suspended in 3 ml of transcription buffer and incubated for 20 min on ice. Cells were broken in a 7-ml Dounce homogenizer with a type B pestle by applying rapid strokes continuously for ~5 min, until >75% of the cells were broken. The suspension was divided into 1 ml aliquots, frozen in liquid nitrogen and stored at -70°C. For a whole-cell extract preparation, an aliquot was thawed, 900 μl was mixed with 100 μl of transcription buffer containing 1.5 M KCl, and incubated for 20 min on ice with frequent mixing. Subsequently, the extract was centrifuged at 21 000g for 10 min at 4°C. The supernatant (~600 μl) was transferred to a new microtube, diluted with 0.5 volumes of transcription buffer, and centrifuged again for 2 min. The supernatant was loaded on an Amicon Ultra-4, 5KDa MWCO (Millipore) to concentrate the extract. When the volume reached ~500 μl, a second salt dilution to 40 mM KCl was performed by adding 750 μl transcription buffer, and the sample was further concentrated until a final volume of 150–200 μl was reached. The final cell extract was divided into ~50 μl aliquots, frozen, and stored at -70°C. Protein concentration of extracts varied between 40 and 50 mg/ml.

**In vitro trans-splicing**

The reaction buffer was essentially as described (30) with minor changes. Each 20 μl reaction mixture contained 8 μl cell extract (300–500 μg protein) in 20 mM potassium 1-glutamate, 20 mM KCl, 3 mM MgCl2, 20 mM HEPES–KOH, pH 7.7, 20 mM creatine phosphate, 0.48 mg/ml creatine kinase, 2.5% polyethylene glycol, 0.2 mM EDTA, 0.5 mM EGTA, 4 mM dithiothreitol, 10 μg/ml leupeptin, 10 μg/ml aprotinin and 25 μg/ml pre-mRNA substrate (0.5 μg). ATP (0.5 mM) was added after 15 min incubation on ice, and then the reaction was continued for 1 h at 28°C. When co-transcriptional trans-splicing was examined, 0.5 mM NTP mix was added to the reaction and 40 μg/ml of pNS21-TIR plasmid DNA was used. Total RNA was prepared and analyzed by RNase protection assay as previously described (14), except that RNA was extracted from the in vitro reaction using TriReagent (Sigma). RNase protection assay products were separated on 6% polyacrylamide/7 M urea gels and visualized by autoradiography.

**RNA preparation for the in vitro reaction**

Total RNA from the transgenic parasites was prepared using TriReagent (Sigma). To generate pre-mRNA for the trans-splicing reaction, PCR products were prepared using a sense primer containing the T7 promoter. The T7 in vitro transcribed pre-TIR and its derivate mutations were prepared with sense primer 259, and antisense primer 218 situated 86 nt downstream of the luciferase ATG. Synthetic SL RNA was prepared using primers 775 and 777, and plasmid coding for *T. brucei* SL RNA (14). RNA was synthesized using the MegaScript T7 Kit (Ambion) according to the manufacturer’s instructions. [α-32P]UTP-labeled antisense probes for RNase protection assay were prepared by using T7 RNA polymerase (Promega).

**Isolation of SL RNA transcribed in the cells**

The synthesis of [α-32P]UTP-labeled SL RNA in permeable cells was essentially as described (31). For isolation of native SL RNA, total RNA was prepared from 250 ml *T. brucei* parental strain (~1 mg) and separated on a 10% acrylamide denaturing gel. SL RNA co-migrates on such gels together with the 140 nt srRNA-3. RNA was visualized by UV lamp on a TLC plate. The ~140 nt RNA species was excised from the gel. SL RNA concentration was determined by primer extension using synthetically transcribed SL RNA as a standard (0.08 to 8 ng). Based on the primer extension calibration curve, the concentration of gel-extracted SL RNA was estimated to be 0.35 ng/μl.

**RNase H digestion of U snRNAs in the splicing extracts**

RNase H digestion was performed as follows: In vitro splicing extract (300–500 μg) was incubated with 200 pmol of primer, 0.4 mM ATP and 1U RNase H (New England Biolabs, USA) in a total reaction volume of 20 μl, for 90 min on ice. The reaction was then split. Half of the treated extract was used for an in vitro reaction and the other half was subjected to primer extension. The primers used for RNase H cleavage are listed in Supplementary Data 1.

**RT–PCR as a trans-splicing assay**

In vitro trans-splicing reaction was performed as described above and RNA was extracted by Proteinase K treatment, followed by phenol:chloroform extraction and ethanol precipitation. The cDNA was prepared from one standard 20 μl splicing reaction using random primer and the RevertAid™ First Strand cDNA synthesis kit (Fermentas) following the manufacturer’s instructions. PCR was performed on 0.1 μl of cDNA, 1 μM primers and ReadyMix™ Taq PCR Reaction Mix with MgCl2 (Sigma). The PCR conditions were as follows: 95°C for 2 min followed by 30 cycles of 95°C for 30 s, 60°C for 30 s and 72°C for 10 s (for pNS21-TIRsub mRNA amplification) or for 30 s (for pNS21-TIRsub pre-mRNA and Tubulin amplification).

**Quantitative real-time PCR**

Real-time PCR was performed in a two-step reaction. First, cDNA was prepared from either 10 ng of total RNA or a standard in vitro trans-splicing reaction using
random primer and the RevertAid™ First Strand cDNA synthesis kit (Fermentas) following the manufacturer’s instructions. Next, real-time PCR was performed using 1 μl of cDNA (diluted 1:100), 1 μM primers and Absolute Blue QPCR SYBR® Green ROX mix (Thermo Scientific). qRT–PCR was performed on a Chromo4 Real-Time PCR detection system (Bio-Rad), as follows: 95° for 2 min followed by 40 cycles of 95° for 30 s, 60° for 30 s and 72° for 10 s. A concentration curve of the trans-spliced product, purified using the QIAquick PCR purification kit (Qiagen), was determined using the Opticon Monitor3 software supplied with the Opticon4 apparatus. This concentration curve was used to determine the amount of PCR product produced in each sample. The reverse transcription could be performed using random-primers or primer specific to luciferase (216-Luc 66-68as). However, the random priming enabled simultaneous detection of the controls and the spliced product.

RESULTS

Constructing a pre-mRNA for in vitro trans-splicing

In vivo trans-splicing takes place on polycistronic pre-mRNA. A scheme depicting such a pre-mRNA is given in Figure 1A, highlighting the essential sequences required for trans-splicing including the 3' AG splice site, polypyrimidine tract (PPT), branch point and the polyadenylation signal of the upstream gene.

Plasmid pNS-21 was used in this study; this vector was previously used to study the optimal composition and distance between the AG splice and the PPT for trans-splicing (28). The sequences in the intergenic region of the vector are partially synthetic and lack the polyadenylation signal of an upstream gene. Because of the previously shown linkage between trans-splicing and polyadenylation (28,32–34), we replaced the intergenic region with authentic sequences. The sequences between the BglII site and the XbaI site were replaced by either the

| A | SL addition site | Polyadenylation site | BP | SL addition site | Polyadenylation site |
|---|------------------|----------------------|----|------------------|----------------------|
| PPT | AG | Gene 1 | A | PPT | AG | Gene 2 |

| B | pNS21-PIR pre-mRNA, 1019 nt |
|---|-----------------------------|
| polyA site | BP | PPT |
| Procyclin (EP-3) upstream intergenic sequence |
| probe | SL | PIR | Luciferase |
| in vitro trans-spliced product |
| endogenous mRNA |

| C (a) |
|---|
| M | 1 | 2 | 3 |
| 230 | 217 | 190 | 160 |
| 140 | 120 | 100 | 80 |
| 60 | 40 | 20 | 0 |

(b) | Luciferase mRNA in Transgenic Clones |
|---|
| Relative mRNA concentration (%)
| Transgenic clones |
| pNS21-PIR | pNS21-TIR |

Figure 1. (A) Schematic presentation of a trypanosome polycistronic pre-mRNA. The significant sequence elements essential for trans-splicing are indicated. (B) Pre-mRNA substrates used in this study. Schematic representation of the pre-mRNA substrates highlighting the significant sequence elements: polyadenylation site (polyA site), polypyrimidine tract (PPT) and AG splice site. The lengths of the PIR and TIR UTRs as well as the Luciferase ORF are indicated. Underneath each pre-mRNA, the respective probe, in vitro trans-spliced product and endogenous mRNA are shown, along with the expected size of the protected fragments after RNase protection. (C) RNase protection assay to monitor the expression of the TIR and PIR constructs. (a) Thirty micrograms of total RNA was hybridized with labeled antisense probe, complementary to the trans-spliced product, as described in 'Materials and Methods' section. Protected RNA products were separated on a 6% sequencing gel. M: DNA marker, labeled pBR322 DNA MspI digest. The size of the marker is indicated on the left. Lane contents are as follows: 1, RNA from parental strain; 2, RNA from transgenic parasites expressing the pNS21-PIR construct; 3, RNA from transgenic parasites expressing the pNS21-TIR construct. The scheme on the right hand side of the gel indicates the structure of the probe and the protected fragments. (b) Relative expression of the tubulin-luciferase or the EP-luciferase to the endogenous tubulin and EP transcripts.
326 bp intergenic region upstream to the β-tubulin locus (pNS21-TIR) or the 630 bp EP-3-2 intergenic region (pNS21-PIR) (Supplementary Figure S2). Transgenic parasites were obtained and the expression of the luciferase gene was determined by northern analysis. The results indicated that the presence of the authentic intergenic regions of either tubulin or EP enhanced the expression of the luciferase gene compared to the parental pNS21 (data not shown). Since these constructs differ at the 5′ flanking sequences, especially in the sequences upstream to the AG splice site, the difference in expression most probably reflects differences in trans-splicing efficiency.

To establish an assay that would directly measure the production of the trans-spliced product from pre-mRNA, we prepared an antisense RNA probe (carrying a small extension at the 5′-end to distinguish the probe from the protected fragments) using a cDNA clone derived from the transgenic parasites as a template. The RNase protection assay was used to examine the production of the trans-spliced products in parasites expressing either the TIR or PIR constructs. Schematic presentation of TIR and PIR constructs and the fragments that are generated from the RNase protection are depicted in Figure 1B. Three fragments were observed: (I) the residual undigested probe; (II) the trans-spliced product; and (III) the endogenous transcript. Based on the ratio between the trans-spliced product (II) and the endogenous transcript (III), it was evident that the PIR construct was more highly expressed and trans-spliced compared to the TIR transcript (Figure 1C-b compare lanes 2 to 3). As a control for the specificity of the assay, RNA from the parental strain was used, and the only protected fragment was of the endogenous tubulin transcript (lane 1). These results suggested that we established a chimeric construct that is properly trans-spliced in vivo and can be used as a substrate for establishing the in vitro trans-splicing reaction.

**In vitro reaction with the PIR and TIR pre-mRNA**

Several parameters were considered when establishing the in vitro reaction. Different protocols were used to prepare nuclear extracts (35,36) and a whole-cell extract (29). The efficiency of SL RNA transcription was examined in the extracts using an SL RNA gene plasmid harboring a 19-nt insertion mutation, and primer extension using a primer specific for the insertion (37). The results (data not shown) indicated that in our hands the most active transcription extract was the whole-cell extract prepared according to Laufer et al. (29). Note, that sucrose was used both for freezing the extract and in the reaction, rather than glycerol or sorbitol, which are used to stabilize the mammalian and yeast extracts, respectively. This change stems from the fact that the trypanosome extracts are highly enriched in glycolytic enzymes that require ATP, and hence, addition of glycerol depletes the ATP in the extract. It was previously reported that when the counter ion chloride is replaced by glutamate, the in vitro transcription in the trypanosome extracts is more efficient (35). Potassium glutamate was therefore used in the reaction. The first step was to determine if a trans-spliced product is generated in the reaction using T7 in vitro transcribed pre-TIR or pre-PIR mRNAs. The source of SL RNA was the endogenous SL RNP present in the extract. The reaction was terminated by extraction of the RNA, and the RNA was subjected to RNase protection with the corresponding probes. The RNase protection was performed in parallel on RNA extracted from transgenic parasites expressing the TIR and PIR constructs (Figure 2). Three fragments, whose identities are illustrated schematically on the right side of the gel, were

![Figure 2](https://academic.oup.com/nar/article/38/10/e114/2902780)
observed when pre-TIR and pre-PIR were used in the reaction: (i) residual undigested probe, which migrates like the probe (Figure 2A and B, lane 1); (ii) the trans-spliced product (Figure 2A and B, lane 3), seen as the protected fragments present in the RNA from the in vitro reactions. The protected fragments were identical to the fragments detected when RNA from transgenic parasites was used (compare lanes 2 to 3); and (iii) pre-mRNA; the intensity of the protected fragment indicates the extent of pre-mRNA that existed at the end of the reaction (present only in lane 3). Although the same amount (0.5 µg) of both pre-TIR and pre-PIR were added to the reaction, massive degradation of pre-PIR was observed in the extract. The trans-spliced product (lanes 3) was excised from the gel, amplified using the SL forward primer and the luciferase reverse primer (lanes 3) was excised from the gel, amplified using the SL forward primer and the luciferase reverse primer (described above) and sequenced. The sequence was identified to the product observed in the transgenic lines, suggesting that the reaction is faithful and generates a genuine trans-spliced product. The efficiency of the reaction was estimated by examining the fraction of pre-mRNA that was converted to trans-spliced product. The results were analyzed by densitometry and the intensity of the spliced product was divided by the intensity of the pre-mRNA. While this approach gives only a very rough estimation, our results indicate that between 0.5% and 2% of the pre-mRNA is trans-spliced. Note that the pre-mRNA is present in vast excess (see below).

The trans-splicing reaction is heat sensitive and requires ATP

To examine if the reaction shares known properties of in vitro splicing reactions (22–24), two requirements must be met. The reaction should be inactivated by heat and should depend on the presence of ATP. The results in Figure 3A demonstrate that heat inactivation (incubating the extract for 30 min at 55°C) abolished the production of the trans-spliced product (compare lanes 2 to 3). The position of the trans-spliced product was determined by the migration of trans-spliced product generated from RNA extracted from the transgenic line expressing the TIR plasmid (lane 1). Next, the dependence of the reaction on ATP was examined. If the ATP regenerating system was omitted from the reaction, no trans-spliced product was generated (Figure 3B, compare lanes 4 and 5). If the reaction lacked pre-mRNA, and instead the TIR plasmid was added to the reaction under transcription conditions, pre-mRNA was produced but no trans-spliced product was detected, suggesting that the amount of pre-mRNA produced in vitro did not enable the detection of the trans-spliced product (Figure 3B, lane 3). Next, we examined the time dependence of the reaction; the results demonstrate that the reaction was linear up to 60 minutes (Figure 3C and D). No increase in the trans-spliced product was observed thereafter, most probably because of consumption of all the SL RNA present in the extract. The degradation of the pre-mRNA most probably takes place immediately after addition of the pre-mRNA, since no change in the level of protected pre-mRNA was observed at longer incubation times, suggesting that the pre-mRNA is protected from further degradation once it is assembled into splicing complexes (Figure 3C; compare lanes 1–3).

Optimization of the reaction conditions

Salt concentration, the level of MgCl₂, the concentration of reactants and the identity of the acceptor and donor RNAs can all dramatically change the yield of the trans-splicing reaction. Thus, it was necessary to determine the optimal concentration of each of these constituents within the reaction. Previously, the type of counter ion used was shown to affect the in vitro transcription reaction. Changing KCl to K-glutamate increases the efficiency of SL RNA transcription (35). Thus, the in vitro reaction was incubated in the presence of elevated salt concentration, and the production of the trans-spliced product was monitored. The results shown in Figure 4A indicate that 40 mM K-glutamate was the optimal salt concentration. As expected, at higher salt concentrations, above 60 mM, the reaction was completely inhibited. Next, the effect of MgCl₂ concentration was examined and the results indicate that 3 mM MgCl₂ was optimal (results not shown). The amount of pre-mRNA was also examined, and the results (Figure 4B) indicate that a relatively high concentration (0.5 µg per reaction) was needed to enable production of the trans-spliced product. These results demonstrate that the RNAse protection assay reflects the increased amount of pre-mRNA present in the reaction. Note that concentrations higher than 0.5 µg poison the reaction. The results in the right panel present the quantification data from three independent experiments.

Addition of endogenous SL RNA but not in vitro transcribed SL RNA enhances the reaction efficiency

The trans-splicing reaction depends on the supply of endogenous SL RNA for the reaction to proceed. However, SL RNA might be a limiting factor in the extract. Exogenous SL RNA must first assemble into its cognate particle, the SL RNP, before it can participate in the reaction. The assembly of SL RNA into its particle was compared using in vitro transcribed SL RNA versus the assembly of native SL RNA that was synthesized in permeable cells. The results presented in Figure 5A-a, lanes 1 and 2, suggest that 100% of the native SL RNA prepared in permeable cells was incorporated into the SL RNP particle, whereas <1% of the in vitro transcribed SL RNA was assembled into the SL RNP (lanes 5 and 6). SL RNP assembly with the radiolabeled SL RNA was competed by the addition of cold-endogenous SL RNA (compare lanes 2 to 3) but not by tRNA (compare lanes 2 and 4). The assembly of SL RNA into the SL RNP was optimal when the reaction was performed with 4 mM MgCl₂ and 30 mM K-glutamate (Figure 5A-b), conditions very similar to those used in the in vitro reaction (Figure 4).

Next, we examined whether addition of native SL RNA or in vitro transcribed SL RNA to the in vitro splicing
reaction could increase the efficiency of producing the trans-spliced product. Addition of increasing amounts of endogenous SL RNA (up to 3 ng), increased the efficiency of the reaction, suggesting that the SL RNP is limiting in the extracts (Figure 5B). In contrast, addition of increasing amounts of synthetic SL RNA (up to 2 μg) had no effect on the production of trans-spliced product (Figure 5C-a); This result was expected based on the much more efficient assembly of endogenous SL RNA relative to the synthetic molecule. Next, it was determined if in vitro transcribed SL RNA containing an m7G cap could be used as a substrate for in vitro trans-splicing. Addition of up to 2 μg of synthetic m7G capped SL RNA did not increase the generation of the trans-spliced product (Figure 5C-b). This synthetic SL RNA also failed to efficiently assemble into SL RNP (data not shown), suggesting that the cap-4 modification might be necessary for SL RNP biogenesis. Therefore, increasing amounts of synthetic cap-4 (38) were added to the reaction. Figure 5D shows that adding cap-4 at concentrations up to 50 μM did not affect the reaction. Thus suggesting that cap-4 needs to be attached to the SL RNA in order to promote proper SL RNP assembly and that cap-4 provided in trans can not interfere with this process.

The absence of the +4 cap modification does not affect in vitro trans-splicing

To examine the dependence of trans-splicing in vitro on cap-4, we prepared an extract from SmD1-silenced cells. In these silenced cells, trans-splicing is inhibited, and SL RNA lacking modification at the +4 position of
cap-4 accumulates (14). The efficiency of knockdown is variable, and silencing is not very efficient if cells are silenced in large quantities. The silenced population also contains cells that are unaffected, and thus Sm proteins are nevertheless present to some extent in the extracts. When extracts are made from such cells, the majority of SL RNA however, is ‘defective’ SL RNA. First, we examined if SL RNA synthesized from SmD1-silenced cells can assemble efficiently into SL RNP particles. SL RNA was prepared from SmD1-silenced permeable cells. The SL RNA synthesized under these conditions lacks the +4 cap-4 modification and possesses a 5-nt tail (14). Defective SL RNA synthesized in permeable SmD1-silenced cells was able to assemble properly into SL RNP particles (Figure 6A compare lanes 2 and 4).

Next, we examined the capacity of extracts prepared from SmD1-silenced cells to carry out the in vitro trans-spooling reaction. Using these extracts, two distinct protected fragments were observed compared to a single protected fragment observed when wild-type splicing extract was used (Fig. 6B, compare lanes 1 and 2). In lane 2, the lower protected fragment represents splicing from the residual wild-type SL RNA present in the extract, and the upper fragment from splicing of defective SL RNA. The difference in size may reflect the fact that SL RNA lacking the cap-4 +4 nucleotide and harboring a 5-nt tail can serve as a bona-fide trans-spooling substrate.

Inactivation of U2 or U6 snRNA reduces the splicing efficiency

It was previously demonstrated that trans-spooling is inhibited in permeable trypanosome cells when either U2 or U6 snRNAs are inactivated by RNase H cleavage (39). To examine if trans-spooling in vitro relies on the presence of intact U2 and U6 snRNAs, extracts were incubated with U2 and U6 antisense oligonucleotides and RNase H. U3 cleavage was used as a control. The same oligonucleotides, U2SSC and U6-15, were previously used to inactivate these snRNAs in permeable cells (39). Cleavage reduced the level of U2 and U6 snRNA by 50% and 40%, respectively, as evident by primer extension that was performed on RNA extracted from the oligonucleotide-RNAse H treated extracts (Figure 7B and C). Indeed, in extracts depleted of these snRNAs, the production of the trans-spooled product was reduced by 60% and 45%, respectively, as evident by primer extension that was performed on RNA extracted from the oligonucleotide-RNAse H treated extracts (Figure 7A; lanes 4 and 5). These results suggest that, as expected, the reaction depends on the snRNAs that constitute the catalytic center of pre-mRNA splicing (40). The cleavage of U3 did not affect the level of U2 (Figure 7B, lane 2) or U6 (Figure...
7C; lane 2) snRNAs and did not affect splicing (Figure 7A, lane 3).

Mutations in the 3′ splice and polypyrimidine tract compromise the production of the trans-spliced product

The AG splice site and the PPT are sequences essential for trans-splicing (28,32,33). It was therefore essential to demonstrate that the production of the trans-spliced product is affected if these sequences are mutated. To this end, mutations were introduced to the AG splice site and the PPT, as illustrated in Figure 8A. Mutations were constructed and the mutated genes were cloned into the pNS21 vector. Transgenic parasites were obtained, and the production of the trans-spliced protected fragment was examined by RNase protection assay using RNA isolated from the transgenic parasites. The results demonstrate (Figure 8B) that the AG to AA mutation reduced...
the production of the trans-spliced product by 90% (compare lanes 1 and 2), whereas the PPT mutation reduced the production of the trans-spliced product by ~70% compared to the wild-type transcript (compare lanes 1 and 3). cDNA was prepared from the transgenic parasites and amplified with the SL forward primer and antisense luciferase primer. Although the RNase protection assay showed complete inhibition of the production of trans-spliced product using the RNase protection assay (Figure 8B), a PCR product was nevertheless obtained after PCR amplification. Sequence analysis indicated that SL was added exclusively 9-nt downstream to the authentic AG splice site, suggesting that, in the absence of a bonafide site, the splicing system ‘scans’ for an alternative AG splice site.

To investigate whether the splicing signal mutants are trans-spliced in vitro similarly to their splicing in vivo, pre-mRNAs were prepared from the mutants and the production of the trans-spliced product was examined. The results (Figure 8C) suggest that both mutations affected the ability to form the spliced product. The reduction in the formation of the trans-spliced product for the AA and PPT mutants was 60% and 70%, respectively (Figure 8C-b). Interestingly, the PPT mutation affected the in vitro reaction more strongly than its effect in vivo. The AA mutation, however, had a less severe effect in vitro compared to in vivo (compare Figure 8C-a, lanes 3 and 5 to Figure 8B-a, lanes 2 and 3).

Generating a TIR mutation downstream to the AG splice site to produce a model substrate for RT–PCR or real-time PCR assays

The RNase protection assay used to monitor the production of the trans-spliced product is a very laborious and time-consuming assay. However, it allows examination of the trans-spliced product without any amplification and is therefore less prone to artifacts compared to assays that involve amplification. Nevertheless, the technical difficulties in the RNase protection assay led us to develop an alternative RT–PCR assay that involves an amplification step. Performing RT–PCR on an in vitro reaction, using the pre-TIR as a substrate and the same oligonucleotides that were used to produce the cDNA in the transgenic parasites, produced a trans-spliced product whose production was insensitive to ATP depletion or to heat inactivation, suggesting that amplification can take place that is not dependent on trans-splicing. We therefore generated a 20-nt substitution mutation 2-nt downstream of the AG splice site (Figure 9A). The mutant was cloned into pNS-21, and cell lines were generated (pNS21-TIRsub). RNA was prepared from the cell line and used to amplify the cDNA using pairs of forward (SL) and antisense primers specific to the mutation (Figure 9C). No such product could be amplified using cDNA from cells expressing pNS21-TIR. The PCR product was sequenced and the results indicated that SL was added exclusively 9-nt downstream to the authentic AG splice site, suggesting that, in the absence of a bonafide site, the splicing system ‘scans’ for an alternative AG splice site.
prepared from the same reactions presented in (B), and the cDNA was amplified with three different pairs of oligonucleotides, a pair to amplify the pre-mRNA, the endogenous tubulin mRNA, and the transspliced product. The transspliced product was amplified with an antisense oligonucleotide complementary to the mutation and a forward SL oligonucleotide. The results indicate that identical amounts of pre-mRNA and extract (based on a tubulin control) were present in each reaction (Figure 10A, lanes 2 to 4), but the level of the transspliced product was sensitive to conditions that affected splicing. Thus, the RT–PCR results are in agreement with the RNase protection assay, and show 85% and 96% reduction in the formation of the transspliced product under ATP depletion or heat-inactivation, respectively (Figure 10).

Real-time PCR

In order to quantify the exact amount of PCR product that is amplified from the in vitro transsplicing reactions, and to show that the RT–PCR performed above was done while the reaction was in exponential phase, qRT–PCR was performed. A melting curve was performed to show that only a single transspliced product was amplified during the qRT–PCR, and the melting curve was identical for the fragment formed using the cDNA from the transgenic parasite and from the in vitro reaction (results not shown). Finally, a PCR product of known quantity was used to establish a concentration curve. This concentration curve was then used to determine the amount of PCR product amplified from an in vitro transsplicing reaction. As can be seen in Figure 10C, a standard in vitro transsplicing reaction, in which 4 pmol of pre-mRNA was added yielded 50 fmol of PCR product. This is comparable to the amount of PCR product amplified from 10 μg of total RNA from transgenic parasites expressing pNS21-TIRsub. Performing qRT–PCR on reactions in the absence of ATP or after heat inactivation yielded ~1 fmol and 0 fmol, respectively. The reduction in PCR amplification in the absence of ATP or upon heat inactivation was similar to the reduction as determined by RT–PCR and RNase protection. Thus, these experiments establish three different assays that faithfully monitor the production of transspliced product generated in vitro from synthetic pre-mRNA and endogenous SL RNA and demonstrate some of the requirements for these assays.

DISCUSSION

In this study, we describe for the first time a system that is able to faithfully monitor the formation of a chimeric transspliced RNA product in vitro using a T. brucei soluble cell extract and synthetic in vitro transcribed pre-mRNA. The ability to establish such an in vitro system stems from the reaction conditions and the sensitive assays used in this study. The study demonstrates that SL RNP is a limiting factor in the reaction, and only endogenous SL RNA but not m7G capped synthetic SL RNA can be utilized. Interestingly, SL RNA lacking the +4 cap modification and possessing a 5-nt longer tail can be utilized in vitro. The assembly of synthetic and
endogenous wild-type or SL RNA from SmD1-silenced cells suggests that the cap-4 modifications but not the cap-4 +4 modification are essential for SL RNP assembly. The performance of the in vitro trans-splicing reaction meets the requirements observed for other splicing systems, specifically the dependence on ATP, and on U2 and U6 snRNAs. Mutations in strategic positions such as the polypyrimidine tract and the AG splice site compromised the reaction in vivo and in vitro.

The most essential factors for establishing the in vitro system

The failure to date to establish an in vitro system in trypanosomes lies in the use of radiolabeled pre-mRNA substrates that were extensively degraded, and the deficiency in essential factor(s) in the extract. Massive degradation of pre-mRNA was also observed in our study. This degradation could not be eliminated even if the mRNA was capped at the 5'-end or at the 3'-end with pCp (our unpublished data). Our success in detecting the trans-spliced product in this study lies in three parameters: (i) the extract used is crude but contains all the factors necessary for the reaction. It is prepared very rapidly to eliminate the potential inactivation of essential factors. However, because the extract is not purified, it contains enzymes that may inactivate essential factors, thus reducing the reaction efficiency. (ii) A large amount of pre-mRNA is added to the reaction, leaving sufficient amounts of substrate (after its immediate degradation) to assemble into an active complex. The amount of pre-mRNA (4 pmol) is 50-fold higher than the amount of SL RNA in the extract (~80 fmol). Indeed, we demonstrate that increasing the amount of pre-mRNA increases degradation.
the reaction efficiency (Figure 4C). (iii) The assays used in this study. This is the most critical parameter that made it possible to detect the trans-spliced product. Three different assays were used. In all the assays, only the end-product but none of the intermediates are monitored. The reaction efficiency, roughly calculated by the RNase protection assay, is 1–2% of the input pre-mRNA utilizing almost 50% of the SL RNA present in the extract. Thus, the limiting factor in this system is the SL RNA.

Although the traditional assays to monitor splicing so far are based on the use of radiolabeled substrate, the real-time PCR assay represents a much desired substitute. It avoids the use of radiolabeled materials; it is quantitative, and is able to quantitatively detect the amount of trans-spliced product even at levels less than a few femtomoles. In addition, the assay is convenient relative to the RNase protection assay. Note that in all assays used in this study, the in vitro trans-spliced product was compared to the genuine trans-spliced product produced in vivo in transgenic parasites.

An additional parameter that was essential for establishing the in vitro system was the choice of pre-mRNA substrate. Our results indicate that the TIR pre-mRNA was a better substrate than the PIR pre-mRNA. The poor performance of the PIR substrate in vitro might stem from its size. The larger substrate appeared more vulnerable to degradation. In other splicing systems, it was demonstrated that shortening both the exons and the introns resulted in a substrate that is spliced more efficiently (26). In addition, microarray data of the transcriptome of cells silenced for splicing factors suggest that EP splicing requires a high level of SR proteins (Gupta and S.M., unpublished data). These factors, in the proper phosphorylation form, might be limited in the extracts used in this study. The choice of the β-tubulin gene over α-tubulin was based on the finding that efficient α-tubulin splicing in vivo requires exonic sequences present in the UTR. Such splicing might require additional factor(s) to enhance the protein binding to suboptimal sequences present in the vicinity of the AG splice site of the α-tubulin transcript (33).

The effect of pre-mRNA mutations on trans-splicing in vivo and in vitro

One of the interesting results of this study is the differential splicing of the AG and PPT mutants in vivo and in vitro. The mutation of AG to AA almost completely abolished the production of the trans-spliced product in vivo (Figure 8B). However, in vitro, only a 60% reduction...
alternative selection of this splice site took place when 6-nt downstream to the authentic site was utilized. Using a was observed (Figure 8C). RT–PCR of RNA extracted from the transgenic parasites revealed the production of a trans-spliced product where an alternative AG located 6-nt downstream to the authentic site was utilized. Using this cDNA in an RNase protection assay showed that alternative selection of this splice site took place when either wild-type transcript or the mutant AG to AA were used in vitro (our unpublished data), suggesting that in vitro, as opposed to in vivo, an alternative AG splice site can be chosen.

The difference between the in vivo and the in vitro utilization of the AG to AA mutant and the choice of an alternative AG splice site may be explained by the shortage in the extract of a factor that functions in selecting the optimal AG splice site. One such candidate is the U2AF65 homologue (20). Our recent study indicates that this large protein (90 kDa) is subjected to proteolysis in the extract. In addition, the active protein is phosphorylated, and the presence of phosphatases in the extract may render this factor inactive.

The PPT mutation, like the splice site mutation, had a different effect in the in vitro reaction. This mutation affected the in vitro reaction as strongly as the AG to AA mutation. In mammals, mutation or deletion of the AG splice site completely blocks ligation (step2), while the first step proceeds, albeit at reduced efficiency. In contrast, mutation of the PPT drastically reduces the efficiency of the first step; thus, the PPT sequence seems to be more important in splicing than the AG dinucleotide (41). In yeast, the 3’ splice site mutations activate a cryptic AG splice site even if this is not preceded by a PPT (42), while in mammals, the cryptic 3’ splice site is always preceded by a PPT (43). These results suggest that PPT recognition is more essential in mammalian than in yeast splicing. The PPT seems to also be very important for trans-splicing, especially in vitro. The trypanosome PPT was shown to be essential for trans-splicing and polyadenylation of the upstream gene (44–46). Mutations introduced in the PPT of trypanosome pre-mRNA suggest that although the size of the PPT does not have a large effect on the trans-splicing efficiency, its composition does. Replacing uridines with two or three consecutive purines severely compromises in vivo trans-splicing (28). Indeed, the mutation we introduced not only reduced the level of U’s within the tract but inserted consecutive purines into the PPT site (Figure 8A).

Endogenous SL RNA but not synthetic SL RNA can enhance the in vitro reaction

In nematodes, in vitro transcribed radiolabeled SL RNA is properly capped and efficiently utilized in the in vitro trans-splicing reaction (25). This is not the case with the trypanosome SL RNA. The special cap-4 is added to the SL RNA co-transcriptionally (47), and capping requires that the nascent SL RNA adopt a special conformation dictated by its interaction with SLA1 (11). Sm assembly is coupled with the cap-4 +4 modification (14); therefore, the in vitro transcribed SL RNA can not assemble efficiently with the Sm core proteins, and hence, the SL RNA is subject to major degradation. Indeed, in the reconstitution experiments presented in Figure 5A, we could not detect unassembled SL RNA; such RNA is most probably rapidly degraded in the extract. The capping of SL RNA by m7G neither improved its stability (our unpublished data), nor its ability to be utilized in the trans-splicing reaction (Figure 5C-b). The in vitro
transcribed SL RNA can not assemble properly into its cognate particle (results not shown) and therefore can not associate with the \textit{trans}-spliceosome. In contrast, SL RNA derived from SmD1-silenced cells that lack the +4 cap-4 modification can properly assemble and be utilized in the reaction (Figure 6B). Although the SmD1 SL RNA is defective compared to the wild-type RNA, it contains all the cap modifications with the exception of cap +4. This may suggest that these modifications are crucial for SL RNP assembly. Indeed, our results are in agreement with the \textit{in vivo} data demonstrating that an SL RNA mutant in the cap-4 +4 nucleotide has only a minor effect on its utilization compared to mutations in the +1 and +2 cap nucleotides, which severely affect \textit{trans}-splicing \textit{in vivo} (9).

In mammalian cells, it was demonstrated that splicing is enhanced by using capped pre-mRNA (48). In yeast, however, addition of cap analogues does not affect the reaction (22). In trypanosomes, the cap-4 may have additional biological functions during SL RNP assembly, other than just protecting the SL RNA from degradation. The main function of cap-4 might not be in the splicing reaction itself, but rather, in the assembly of functional SL RNP. The cap modification may serve as a landmark controlling the fidelity of SL RNA biogenesis; thus, only SL RNA containing the majority of modifications can properly assemble \textit{in vitro}. Indeed, it was shown that effector proteins distinguish the fully methylated form of the cap-4 from the incomplete cap structure. For instance, the \textit{T. brucei} nuclear cap-binding complex CBP20 binds with higher affinity to the mature cap-4 compared to uncapped SL RNA (49). In \textit{Leishmania}, homologues of the cap-dependent translation initiation factor eIF4E distinguish between different SL RNA caps (50). Interestingly, although cells can resist the loss of either modification at the cap-2 (51) or at positions 3 and 4 (51,52), it is not possible to derive viable cells with complete loss of cap ribose methylation (53). Indeed, \textit{in vitro} transcribed SL RNA lacking all the trypanosome-specific modifications can not assemble into SL RNP nor participate in the \textit{trans}-splicing reaction \textit{in vitro}. These modifications are therefore also essential for \textit{trans}-splicing.

One of future challenges will be to synthesize \textit{in vitro} SL RNA that can be utilized in the \textit{in vitro} \textit{trans}-splicing reaction. It will be interesting to use SL RNA from cells with knock-out of TbMTr1, TbMTr2, TbMTr3, or even a TbMTr-2 and 3 double knock-out, to examine the ability of undermethylated SL RNA to serve as a substrate in the \textit{in vitro} \textit{trans}-splicing reaction. Once the modifications that are most relevant for \textit{trans}-splicing are identified, it may be possible to synthesize \textit{in vitro} the properly capped SL RNA using the relevant recombinant methylating enzyme(s).

\textbf{What is the significance of an \textit{in vitro} \textit{trans}-splicing system and how can the system be improved?}

In this study, we utilized the \textit{in vitro} system to begin to assess the role of cap modification on the \textit{trans}-splicing reaction, and this can be further extended as outlined above. However, there are still many additional open questions related to our understanding of the mechanism and the machinery of trypanosome \textit{trans}-splicing. One major question is identifying the protein factor(s) that mediate the interactions between the SL RNP and the pre-mRNA machinery. In addition, relatively little is known about what protein factors and sequences are essential for determining the \textit{trans}-splicing efficiency and the selection of the proper 3' AG splice site. Transgenic parasites expressing reporter genes fused to upstream regulatory sequences were used to identify structural constraints within the 3' AG splice site, and the PPT was shown to be highly essential for \textit{trans}-splicing and its linkage to polyadenylation (28,32–34). However, such results may be influenced by mRNA stability, and the translatability of the reporter gene. An \textit{in vitro} system will therefore help to determine the exact role of sequences and factors in \textit{trans}-splicing. Recently, we described two polypyrimidine tract binding proteins in trypanosomes that affect both \textit{trans}-splicing and mRNA stability (54). However, in our studies, it was very difficult to dissect the contribution of these RNA binding proteins to \textit{trans}-splicing as compared to mRNA stability. The \textit{in vitro} system should assist in defining the factors that are essential for \textit{trans}-splicing of C rich polypyrimidine tract pre-mRNAs that bind PTB. Thus, \textit{in vitro} \textit{trans}-splicing could be used to study which factors and sequences are required for differential \textit{trans}-splicing, for example of the three phosphoglyceratekinase genes present in the same polycistrionic unit in \textit{T. brucei} (55). However, the immediate next goal is to identify factor(s) that might be missing or inactivated in the extracts, resulting in their reduced efficiency, especially for splicing genes that require additional factors. This can be achieved by either ‘spiking’ the reaction with fractionated extracts, or adding factors such as the U2AF65 homologue as recombinant proteins.

This study is only the first step towards developing a robust cell-free system that, together with data from \textit{in vivo} studies, should help to reveal the mechanism controlling the efficiency and fidelity of trypanosome \textit{trans}-splicing. \textit{Trans}-splicing is one of the most essential mRNA processing mechanisms in trypanosomes. This is the main biochemical process that differentiates the host from the parasite and therefore a better understanding of the mechanism may also identify novel targets for therapy.

\textbf{SUPPLEMENTARY DATA}

Supplementary Data are available at NAR Online.

\textbf{ACKNOWLEDGEMENTS}

We thank T. Nicolai Siegel and George Cross (Rockefeller University) for providing us with the pNS21 vector. We also thank Arthur Günl (University of Connecticut Health Center) for valuable advice on extract preparation.

\textbf{FUNDING}

United States-Israel Binational Science Foundation grant, by the Deutsche Forchshungsgemeinschaft (DFG);
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