The role of deadenylation in the degradation of unstable mRNAs in trypanosomes

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ABSTRACT

Removal of the poly(A) tail is the first step in the degradation of many eukaryotic mRNAs. In metazoa and yeast, the Ccr4/Caf1/Not complex has the predominant deadenylase activity, while the Pan2/Pan3 complex may trim poly(A) tails to the correct size, or initiate deadenylation. In trypanosomes, turnover of several constitutively-expressed or long-lived mRNAs is not affected by depletion of the 5′–3′ exoribonuclease XRNA, but is almost completely inhibited by depletion of the deadenylase CAF1. In contrast, two highly unstable mRNAs, encoding EP procyclin and a phosphoglycerate kinase, PGKB, accumulate when XRNA levels are reduced. We here show that degradation of EP mRNA was partially inhibited after CAF1 depletion. RNAi-targeting trypanosome PAN2 had a mild effect on global deadenylation, and on degradation of a few mRNAs including EP. By amplifying and sequencing degradation intermediates, we demonstrated that a reduction in XRNA had no effect on degradation of a stable mRNA encoding a ribosomal protein, but caused accumulation of EP mRNA fragments that had lost substantial portions of the 5′ and 3′ ends. The results support a model in which trypanosome mRNAs can be degraded by at least two different, partially independent, cytoplasmic degradation pathways attacking both ends of the mRNA.

INTRODUCTION

Messenger RNA degradation is an important component of the control of gene expression in eukaryotes. Messenger RNAs are generally protected from degradation by the interaction of poly(A)-binding protein (PABP) with both the poly(A) tail at the 3′-end, and the cap-binding complex at the 5′-end (1). In the simplest model, mRNA degradation is initiated by digestion of the poly(A) tail; once the tail has been shortened below a certain threshold length, poly(A)-binding protein drops off, the interaction with the 5′-end is disrupted and the RNA ends become available for digestion (2,3). Three major deadenylases are known: PARN, Pan2/Pan3 and the Ccr4/Caf1/Not complex.

The poly(A) ribonuclease PARN has specialized functions, such as deadenylation of stored mRNAs during the maturation of oocytes (4,5) and regulation of embryogenesis in plants (6). It is stimulated by PABP (7) and by binding to the 5′ cap (5).

The Ccr4/Caf1/Not complex has the major deadenylation activity in animal cells and yeast. (8,9). In some organisms, including yeast and mammals, it contains two catalytic subunits, Ccr4 and Caf1. Ccr4 is bound to Caf1, which in turn interacts with the scaffold protein Not1. Several other subunits, with stimulatory or other functions, are also associated with Not1 (10–12). Although Ccr4 is the most active deadenylase in the Saccharomyces cerevisiae Ccr4/Caf1/Not complex (13), Caf1 is more important in animal cells (8,14–16). Indeed, whereas Caf1 is conserved throughout eukaryotic evolution, Ccr4 is absent in many species. In some mammalian in vitro extracts, degradation of an mRNA substrate is stimulated by addition of poly(A) oligoribonucleotides, suggesting that the major deadenylation activity is inhibited by PABP (3,17). This would correlate with the known inhibition of Ccr4/Caf1/Not by PABP (18).

Pan2 is the catalytic subunit of the Pan2/Pan3 deadenylase and is a member of the DEDDh subfamily of exonuclease III enzymes. In contrast with Caf1, the activity of the complex is stimulated by PABP in vitro (19,20).

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Pan3 has stimulatory function: the C-terminal part interacts with Pan2, while the N-termimnus interacts with PABP (20,21). Pan3 also has a central kinase domain, the role of which is unknown. Pan2/3 is found in the yeast nucleus and cytoplasm. In the nucleus, it trims the poly(A) tails of newly synthesized mRNAs to specific lengths (22) while the cytoplasmic fraction plays a role in mature mRNA deadenylation (12). Neither Pan2 nor Pan3 is essential for yeast growth (23,24) but their absence results in longer bulk poly(A) tails in vivo. In Drosophila Schneider cells Pan2 contributes to the deadenylation of the hsp70 mRNA during recovery from heat shock (25). There is evidence that in mammalian cells, the Pan2/Pan3 deadenylase initiates cytoplasmic deadenylation (26).

After deadenylation, mRNAs can be degraded either unidirectionally from either end, or from both ends; the nature of the mRNA and the species determine which pathway predominates. Results from work in S. cerevisiae showed that the major pathway for degradation of some unstable deadenylated mRNAs is decapping, by a complex containing the MutT hydrolase Dcp2, then degradation by the 5′→3′ exonuclease Xrn1 (27–29). Alternatively, or in addition, deadenylated transcripts can be degraded by 3′→5′ exonucleases associated with the exosome (30,31); in this case the residual cap structure is degraded by the scavenger decapping enzyme DcpS (32).

Computer modelling of mRNA degradation in yeast indicated that modulation of deadenylation has a stronger effect on mRNA half-life and levels than changes in decapping or the 3′→5′ and 5′→3′ decay pathways (33). Mammalian mRNAs that are destabilized by AU-rich elements (ARE) in the 3′-untranslated region (UTR) are subject to accelerated deadenylation (34–36) followed by digestion by 5′→3′ and 3′→5′ pathways (37,38). 5′ degradation is also important in Drosophila cells (25). In contrast, the stability of the alpha globin mRNA depends on binding of a complex that inhibits 3′→5′ decay (39). The 3′ pathway has also been shown to be important during degradation of mammalian mRNAs with premature termination codons (nonsense-mediated mRNA decay, or NMD) (40–42). In mammalian cells, the decapping enzyme, Xrn1, Dhl1, Pan2 and Caf1 co-localize in cytoplasmic processing bodies (P-bodies) (15) which are thought to be a site of mRNA decay.

Despite the importance of deadenylation, other deadenylation-independent mRNA degradation pathways clearly exist. First, degradation may be initiated by an endonuclease: examples include mammalian mRNAs subject to iron regulation (43), the alpha globin mRNA in mammalian erythroid cells (44) and the albumin RNA in Xenopus hepatocytes (45). In Drosophila and mammalian cells some mRNAs with premature termination codons (PTCs) are also attacked by an endonuclease (46–48). In yeast, RNAs with PTCs (49) and the mRNAs encoding the ribosomal protein Rps28b (50) and the decapping enhancer Edc1 (51) are subject to deadenylation-independent decapping. In contrast, degradation of mRNAs lacking a stop codon depends primarily on the exosome (52). Finally, degradation of mammalian histone mRNAs, which are not polyadenylated, is initiated by addition of poly(U) (53).

Trypanosomes are unicellular parasites which branched from the Opisthomont (fungi/animal) line very early in eukaryotic evolution (54). In trypanosomes, transcription by RNA polymerase II is polycistronic (55), individual mRNAs being generated by processing (trans splicing and polyadenylation) (56,57). Control of polymerase II initiation—even for the polycistronic units—has not been documented. Nevertheless, gene expression is regulated in order to allow trypanosomes to survive and grow in the mammalian host (the ‘bloodstream form’) and in the digestive tract of the Tsetse fly (the ‘procyclic form’) (58). To regulate gene expression, trypanosomes are therefore heavily dependent on mRNA degradation. The stability of trypanosome mRNAs is—as in other eukaryotes—determined mainly by sequences in the 3′-UTR (59). We have previously shown that degradation of relatively stable, constitutively expressed mRNAs depends on CAF1 activity (14). (Trypanosomes lack CCR4.) Trypanosome extracts possess decapping activities (60), but the proteins involved have not been identified. Of four XRN homologues, only XRNA has so far been shown to play a role in mRNA decay (61). Trypanosome XRNA and DHH1 are found in P-bodies (62).

Our previous work on degradation of highly unstable, developmentally regulated mRNAs has focussed mainly on two mRNAs that are highly expressed in the procycolic form trypanosomes, EP and PGKB. The EP mRNAs encode the major surface proteins of procyclic cells, the EP procyclins. Pairs of procyclin genes are arranged in a tandem repeat which is transcribed by RNA polymerase I (63). EP transcription is approximately 10-fold less efficient in bloodstream forms than in procyclic forms, most likely a consequence of epigenetic control (64). The EP mRNAs have a half-life of about 5 min in bloodstream-form trypanosomes, and over an hour in procyclic forms (65). Regulation of procyclin protein expression is completed by blocks to translation and protein processing (66). PGKB encodes a cytosolic phosphoglycerate kinase that is specific to procyclic forms; it is co-transcribed with the PGKC mRNA, which encodes a phosphoglycerate kinase that is abundant in bloodstream forms (56). The results of mathematical modelling indicate that mRNA stability is the major determinant of PGKB and PGKC expression (67).

Chloramphenicol acetyltransferase (CAT) reporter mRNAs with 3′-UTRs of either EP procyclin mRNA (CAT-EP) or PGKB mRNA (CAT-PGKB) are very unstable in bloodstream forms, with half-lives of around 5 min (68–70). The CAT-EP reporter thus has the same half-life as endogenous EP mRNAs. The half-life of PGKB mRNA has not been measured accurately because of its low abundance (less than one molecule per cell), but available data suggests a value of around 5 min (67). Mutational analyses on both reporters identified a single-stranded poly(U) tract in the 3′-UTR that was required for mRNA instability (69–73). Analysis of the kinetics of CAT-EP mRNA degradation indicated that degradation occurs without accumulation of deadenylated intermediates, and that the RNA is destroyed by both 5′→3′ and 3′→5′ exoribonuclease activities (68).
Correspondingly, depletion of either the exosome (74) or the 5′-3′ exonuclease XRNA (61) delays degradation of the CAT-EP and CAT-PGKB mRNAs. The results of XRNA depletion are more dramatic, with general disruption of PGKB and PGKC regulation: steady-state PGKB mRNA increases in bloodstream forms, while PGKC mRNA appears in procyclins (61). Analysis of CAT-EP degradation intermediates also suggested action of both deadenylation-dependent and -independent mRNA degradation pathways. In this article we investigate the roles of trypanosome CAF1 and PAN2 in rapid mRNA degradation.

MATERIALS AND METHODS

Sequence alignments

To find PAN2 and PAN3 genes, we searched various genomes using BLASTp, with the S. cerevisiae PAN2 or PAN3 protein sequences as the query, and then used the top match (or matches) as queries for the S. cerevisiae genome. Protein sequences that yielded S. cerevisiae PAN2 or PAN3 as the best match were categorized as potential homologues. Sequence alignments were done using ClustalW, with DNASTar software. Functional motifs were found using Prosite and NCBI searches.

Trypanosome culture, RNAi and plasmids

Bloodstream form Trypanosoma brucei stably expressing the tetracycline represor (pHD 1313) and T7 RNA polymerase (pHD 514) were grown, transfected and clonally selected as described (75). The plasmids used in this study are described in Table S1. For CAF1 and PAN3 RNAi the vector p2T7 TA blue (75) was used to make dsRNA from opposing tetracycline-inducible T7 promoters. In the PAN2 and MEX67 RNAi constructs, an inducible RNA polymerase I promoter drove expression of stem-loops (76). The plasmids were linearized with Not I and transfected into bloodstream-form trypanosomes expressing the represor and T7 RNA polymerase (77). Transfectants were selected with 10 µg/ml hygromycin. The RNAi was induced by adding 1 µg/ml tetracycline to the medium. Samples were taken after 18.5 h of CAF1 RNAi induction or after 2-3 days after PAN2 RNAi induction.

In situ V5 tagging was done using the V5 plasmid described in (78), while over-expression of open reading frames with two myc tags was done as in (79).

RNA preparation and blotting

To analyse mRNA decay the cells were treated with Sinefungin (final concentration 2 µg/ml) for 5 min then Actinomycin D was added (final concentration 10 µg/ml) (80). Total RNA was isolated using peqGold Trifast (peqLab, Germany). To remove poly(A) tails, 170 pmol oligo d(T)20 was incubated with 10–12 µg total RNA at 42°C for 10 min, 5 U RNase H (New England Biolabs) and RNase H buffer were added and the mixture incubated for 40 min at 37°C.

To select polyadenylated RNA, oligo d(T) cellulose (GE Healthcare) was treated with 0.1 M NaOH and washed in RNA-binding buffer (20 mM HEPES pH 7.4, 5 mM EDTA, 0.4% SDS, 500 mM NaCl). A total of 20 µg RNA was dissolved in RNA denaturation buffer (20 mM HEPES pH 7.4, 10 mM EDTA, 1% SDS). The mixture was heated at 65°C for 10 min then put on ice. A 1.5x volume of RNA dilution buffer (20 mM HEPES pH 7.4, 800 mM NaCl) was added. The RNA was transferred to a tube containing 10 µg resuspended oligo d(T) cellulose and incubated for 10 min at room temperature. After centrifugation the supernatant contained the poly(A) fraction. The oligo d(T) cellulose was washed three times with washing buffer (20 mM HEPES, pH 7.4, 5 mM EDTA, 0.5% SDS). Poly(A)+ RNA was eluted with water.

RNA was analysed by formaldehyde gel electrophoresis, blotting onto Nitran membranes (GE Healthcare) and hybridization with radioactive DNA (Prime-IT RmT Random Primer Labelling Kit, Stratagene) or antisense RNA (MAXiscriptT7/T3, Ambion) probes. Signals were measured using a phosphorimager and normalized relative to the signal from a 7SL (signal recognition particle) probe. Northern was stripped by brief boiling in 0.1% SDS; the efficacy of stripping assessed by phosphorimager, and any residual signals left to decay as necessary before re-use.

RNA half-lives were measured by plotting relative signal intensities using Kaleidograph (Synergy software), including only the segments of the time course that gave exponential decay curves (fitted with a linear correlation coefficient generally exceeding 0.9). To mathematically describe the half-life of the EP transcript, we used Microsoft Excel to build models, assuming the presence of one or two exponential decay components. Since XRNA depletion results in simple exponential kinetics (half-life about 12 min) we varied one half-life around this value. The other half-life was varied from 1.5 min upwards; the proportions of mRNA affected by each pathway were also varied. The root mean square deviation of the model from the data was calculated and the model with minimum deviation adopted.

Poly(A) tail lengths were analysed as described previously (14). For pooled statistical analysis of time points 5 min, 15 min and 30 min, a Wilcoxon signed rank test was used (81).

Analysis of circular RNAs

RNA samples (60 µg, 400 µl) were treated with 40 units of DNase I (Invitrogen) with 240 units of RNaseOUT (Invitrogen) for 30 mins at room temperature. This and all following reactions were terminated by phenol-chloroform extraction and ethanol precipitation. To remove the terminal cap, ~10 µg of DNase-treated RNA was incubated with 2.5 units of Tobacco Acid Pyrophosphatase (TAP; Epicentre Biotechnologies) and 40 units of RNaseOUT (Invitrogen) in a 20 µl reaction for 1h at 37°C. To remove the cap with the modified 5′ nt, 10 µg of DNase-treated RNA was incubated with 100 pmol of an oligonucleotide complementary to the
first 15 nt of the spliced leader (5′ TCTAATAATAGCGT T 3′) at 37°C in 33 μl of water. We then added 5 μl (1× RNase H buffer) of RNase H buffer, 1 μl (5 U) of RNase H and incubated for 1 h at 37°C. The RNA was extracted with phenol/chloroform and ethanol precipitated. Circularization of 10 μg RNA was done in a reaction volume of 400 μl, using 40 units of T4 RNA ligase (New England Biolabs) and 80 units of RNaseOUT (Invitrogen), for 16 h at 16°C. For reverse transcription ~2 μg of RNA were incubated with 50 pmol of gene-specific reverse primer, 200 units SuperScript III RT (Invitrogen) and 40 units RNaseOUT (Invitrogen) in a 20 μl reaction. PCR amplification was performed in a 30 μl reaction [1 μl of cDNA (5%)], 10 pmol each of forward and reverse primers, 2.5 units Taq DNA polymerase (New England Biolabs) (30 s at 95°C, 30 s at 52°C and 45 s at 72°C). For the RPL37A mRNA, we used 38 cycles; for EP, each PCR was for 30 cycles. PCR products were purified and cloned into pGEMT-easy vector and randomly selected clones sequenced. Details of the oligonucleotides are in the Supplementary Data.

In situ hybridization

In situ hybridization followed approximately the protocols published previously (62,82). Gene-frames (Thermo scientific) were applied to glass slides, which were then coated with poly-l-lysine. For each slide chamber, 10° to 10° cells were pelleted, washed with PBS, and fixed with 500 μl of freshly prepared 4% (w/v) paraformaldehyde for 18 min. Fixed cells were pelleted, washed three times in PBS, resuspended 200 μl PBS and allowed to settle on the slides overnight at 4°C. Cells were permeabilized with an 0.2% (v/v) TritonX-100 in PBS (30 min), and washed three times with PBS. Total 50 μl hybridization buffer was added per chamber: 10× Denhardt's solution, 4× SSC, 1 mM EDTA, 35% deionized formamide, 0.5 μg/ml tRNA, 2 μM/ml RNase OUT. After 30 min 60 ng of 5′Cy3-oligo d(T)30 was mixed with 10 μl of hybridization buffer and added to the hybridization mix. The chamber was covered with a coverslip, slides were heated to 65°C for 3 min., then incubated overnight at room temperature in a humid box. Coverslips and gene frames were removed and the slides were washed twice with 2× SSC (15 min), twice with 1× SSC (15 min) and twice with PBS (5 min). Cells were stained with DAPI solution (100 ng/ml, 15 min), washed twice with PBS (5 min) and allowed to dry for 1 h. Mounted preparations (Vectashield, Vector laboratories, H-1000) were viewed within 24 h.

Immunofluorescence

Totally 10° cells were fixed in 4% paraformaldehyde/PBS (w/v) for 20 min. The cells were allowed to settle on a poly-lysine coated glass slide (25 min), permeabilized with 0.2% (v/v) Triton X-100/PBS (room temperature, 20 min). Slides were blocked with 0.5% (w/v) gelatine (20 min), incubated with a 1:100 dilution of mouse α-V5 (Invitrogen) for 1 h, then a 1:500 dilution of the secondary antibody Alexa Fluor 488 goat α-mouse IgG (Molecular probes) for 40 min before repeated washing. The kinetoplast and the nuclear DNA were stained with 100 ng/ml DAPI/PBS for 10 min.

Cell fractionation

The fractionation method was taken from (83). A total of 2 × 10° cells were washed in PBS and resuspended in buffer A [150 mM sucrose, 20 mM KCl, 3 mM MgCl2, 20 mM HEPES–KOH (pH 7.9), 1 mM DTT, complete EDTA-free protease inhibitor (Roche)]. Igepal CA-630 was added to a final concentration of 0.2%. The cells were passed three times through a 27 g needle. After a centrifugation (13 000 rpm, minifuge, 10 min) the supernatant (cytosolic fraction) was kept and purified with an additional centrifugation step, while the nuclear pellet was resuspended in buffer A and passed 15 times through a 27 g needle. The nuclear fraction was pelleted again (13 000 rpm, 10 min) and washed once more before use.

RESULTS

The trypanosome genome encodes potential homologues of PAN2 and PAN3

The trypanosome genome sequence encodes a single potential PAN2: the locus, Tb927.6.1670, is designated as a pseudogene in the 927 strain and is absent from the (incomplete) genome sequence of the 427 strain, but the gene is intact in other Kinetoplastid genomes (Figure S1). Trypanosoma gambiense and T. brucei PAN2 genes are identical in all but 19 (out of 3256) nt, but a T at position 2697 in the T. gambiense sequence is missing in the corresponding T. brucei 927 sequence. To analyse the protein sequence we therefore restored this residue; sequencing of DNA from the 427 strain indeed revealed an intact open reading frame (M. Carrington and L. Ellis, University of Cambridge, personal communication).

Trypanosome PAN2, like the human and yeast proteins, has a C-terminal exonuclease III domain of the DEDDh RNase family (84), and this is the most conserved part of the protein: it is 38% identical to the equivalent portion of human PAN2. We compared the sequence with that of other, diverse eukaryotes (Figures 1; Figure S1 shows the complete alignment). The PAN2 of Monosiga brevicollis, the nearest unicellular relative to animals, retains all three domains, as do the PAN2s of Chlamydomonas and Dictyostelium discoideum. The Chlamydomonas sequence has mutations in putative active site and catalytic residues (20,84); for example, Asp 1083 was shown to be essential for activity of human PAN2 (20), but is missing in Chlamydomonas PAN2. Aureococcus anophagefferens (a chromalveolate alga), Naegleria (an amoeba), Micromonas pusilla (an alga related to green plants) and Thalassiosira pseudonana (a marine diatom) have truncated versions containing only the exonuclease domain (Figures 1 and S1), but in all of these apart from Thalassiosira, N-terminal active site residues are missing. We could not find PAN2 in Arabidopsis, Plasmodium, Entamoeba, Trichomonas, or Giardia: in these organisms the best match to PAN2 is another 3′–5′ exoribonuclease of S. cerevisiae, Rnh70, which is involved in maturation of 5S rRNA and tRNAs.
Figure 1. Cartoon maps of PAN2 and PAN3 from various species, as indicated on the left. Domains recognized by Prosite are also shown. Sc: Saccharomyces cerevisiae; Hs: Homo sapiens; Tb: T. brucei; Clamy: Chlamydomonas reinhardtii; Thp: Thalassiosira pseudonana; Neig: Nagelea fowleri; Aure: Aureococcus anophagefferens; Micp: Micromonas pusilla; EXOe is the exoribonuclease domain; ‘Kin’ is the putative kinase domain. The downward arrows indicate the approximate positions of residues thought to be required for catalytic activity according to an NCBI domain search and specifications for the cd06143 sequence cluster [e.g. (114,115)]. Complete sequence alignments are provided as Figures S1 (PAN2) and S2 (PAN3).

The WD40 repeat of the yeast and human PAN2s is missing from the trypanosome homologue. The central peptidase-C19 like domain—similar to ubiquitin hydrolases—was 27% identical to the human PAN2 peptidase domain and was recognized using default settings of the NCBI domain recognition algorithm, but the score was low (E = 5 × 10⁻⁹) and the domain was not recognized by Prosite. Moreover, in the putative TbPAN2 peptidase domain, two of the three conserved active site residues (85) are mutated (C→K, H→D). Thus T. brucei PAN2 is unlikely to have ubiquitin hydrolase activity.

A search of the trypanosome database for PAN3 yielded Tb11.01.5540. For the potential PAN3 sequences found in a multi-organism search, sequence identities varied between 20% and 30% (Figures 1 and S2). A central kinase-like domain (SSF56112) was recognized in some proteins but not others; the role of this domain is unknown. The PABP-binding regions of yeast and human PAN3 are at different positions in the proteins, and are not sufficiently similar to enable us to determine whether such a region is conserved in the trypanosome protein. Interestingly, three organisms with truncated versions of PAN2 had a candidate PAN3; but we could not find it in Thalassiosira, Monosiga, Chlamydomonas or Dictyostelium, or in the organisms that lacked PAN2. Although some searches may have failed because the genomes were not 100% complete, and alternative algorithms might yield additional PAN2 and PAN3 sequences, the results so far indicate that both PAN2 and PAN3 have been lost several times in eukaryotic evolution.

Trypanosome PAN2 is in the cytoplasm and is required for normal growth

To find the location of PAN2, we integrated a sequence encoding a V5 epitope tag immediately upstream of the gene, to create an N-terminal fusion. The resulting V5-tagged protein was mostly in the cytoplasm by both immunofluorescence (Figure 2A) and cell fractionation (Figure 2B). We inducibly expressed PAN3 with a C-terminal myc tag; this was also in the cytoplasm (data not shown). Because the signals were weak in comparison with that from DHH1, we were unable to check whether PAN2 or PAN3 were in P-bodies. Preliminary results using co-immunoprecipitation suggested that a small proportion of the PAN3-myc was associated with V5-PAN2 (data not shown). Unfortunately, expression of PAN3-myc was rapidly lost during cultivation of the cell line, precluding further analysis. Attempts to tag the PAN2 gene in situ with a TAP tag failed, and we have so far been unable to express PAN2 in E. coli.

We made several different trypanosome lines with tetracycline-inducible RNA interference targeting PAN2 or PAN3. Induction of PAN3 RNAi did not affect growth (data not shown). In contrast, cells with PAN2 RNAi showed variable levels of growth inhibition. Figure 2C shows cultures that ceased growth completely after PAN2 depletion, whereas the line illustrated in Figure 2D were less affected; another line (data not shown) had a division time of 6.7 h without tetracycline and 8.2 h after RNAi induction. All lines rapidly lost RNAi upon either storage or propagation, and some grew slowly in the absence of tetracycline, suggesting leakage of the RNAi. These problems, which are common in trypanosome clones with detrimental inducible RNAi, prevented us from conducting extensive series of experiments with individual inducible cell lines.

From these experiments, we concluded that PAN2 is in the trypanosome cytoplasm, is probably required for optimal growth, and might be essential. It is not clear whether, in trypanosomes, PAN2 is stably associated with PAN3, and we have no evidence that PAN3 is required for PAN2 function.

Depletion of PAN2 decreases deadenylation

To find out whether PAN2 is important for mRNA deadenylation in trypanosomes, we measured bulk poly(A) tail lengths in cells with or without RNAi, and after inhibition of transcription. Cells with PAN2 RNAi reproducibly had more poly(A), as judged by the signals on the autoradiograms (Figure 3A). Although equal amounts of starting RNA were used for each reaction, this effect cannot be quantitated accurately because the design of the assay precludes inclusion of a loading control. Instead, to measure the effect of RNAi, we analysed the distributions of poly(A) tail lengths within each sample (Figure 3B). At steady state (time = 0), there was no significant difference in poly(A) tail lengths between the normal and PAN2-depleted cells, but after transcription inhibition, deadenylation was delayed. Quantitation of the relative intensities of the signals for short (0–15), intermediate (51–100) and long (101–200) poly (A) tails revealed that after 5, 15 and 30 min of transcription inhibition, the samples from PAN2-depleted cells had longer poly(A) tails than samples from the normal cells (Figure 3B). The effect was weaker than that seen upon depletion of CAF1 (14). This could indicate that PAN2 has a minor role in deadenylation of all mRNAs, or that it attacks only a subset of mRNAs. It is also possible that the weakness
of the effect is caused by persistence of PAN2 protein after RNAi induction.

We previously showed that CAF1 RNAi strongly inhibits degradation of constitutively-expressed trypanosome mRNAs, such as those encoding glycosomal PGKC, histone H4 (HISH4), tubulin (TUB) or actin (ACT) [(14) and see also Figure 4A]. PAN2 depletion, like CAF1 depletion, did not cause an increase in steady-state levels of these mRNAs (Figure 3D and E, and data not shown). PAN2 depletion had no significant effect on HISH4 (Figures 3D) or TUB (Figure S3) mRNA decay. In Figure S3 we show the mRNA degradation curves normalized to wild-type, so that any effects of RNAi on in steady-state mRNA levels are apparent. ACT RNA was slightly more stable in PAN2-depleted cells (Figure 3C and E). This effect was clearest 30 min after RNA synthesis inhibition: at this time point, in five out of five experiments, there was more mRNA left in the RNAi-induced cells than in the controls. At 60 min, ACT mRNA was undetectable in the control samples but detectable in the RNAi samples. We concluded that PAN2 depletion affected decay of the ACT mRNA.

**Deadenylation plays a role in degradation of the unstable EP mRNA**

We previously showed that RNAi targeting the 5′–3′ exonuclease XRNA in bloodstream forms caused accumulation of the unstable EP procyclin and PGKB
mRNAs. In addition, a role for deadenylation was indicated by the existence of deadenylated EP and CAT-EP mRNA (61,74). We therefore analysed the effects of both CAF1 and PAN2 depletion on EP mRNA turnover. HISH4 mRNA was included as a control for deadenylation inhibition in the CAF1-depleted cells.

After inhibition of transcription, in cells without RNAi induction, EP procyclin transcripts showed an initial rapid decrease, followed by a slower phase (Figure 4A). The kinetics could be explained by the existence of two exponential decay components: about half of the RNA had a half-life of around 2 min and the rest, a half-life of 14 min. After CAF1 RNAi, the abundance of EP transcripts...
increased 1.2- to 3.1-fold, and their degradation was inhibited (Figures 4A and S3B). After CAF1 depletion, the degradation kinetics were best explained by a model in which half of the RNA was degraded with a half-life of 12 min and the remainder had a half-life of over 8 h: even after 90 min, 45% of the EP mRNA persisted. From these kinetics, it appeared that half of the EP mRNA is degraded by a pathway that depends on CAF1.

In the PAN2 RNAi cell line, EP mRNA had a half-life of about 7 min in the absence of RNAi induction, suggesting some inhibition of degradation because of RNAi ‘leakage’. With PAN2 RNAi, degradation was further inhibited and 10% appeared to be stable (Figures 4B and S3A). These results suggest that PAN2 is required for optimal EP mRNA degradation. We cannot make any judgements concerning the relative importance of the two deadenylases in EP mRNA degradation since we do not know the extents of PAN2 and CAF1 depletion and—even if we did—we do not know the threshold below which depletion might have an effect. Digestion with oligo d(T) and RNaseH increased the mobility of EP mRNA, suggesting the presence of a poly(A) tail of at least 50 nt (Figure 4A and B).

We had previously shown that XRNA RNAi inhibited the degradation of a CAT-EP reporter mRNA. We now examined the effect of XRNA RNAi on EP mRNA degradation (Figure 4C). The graph quantitates the ~900-nt mature EP mRNA. Degradation in cells without the RNAi construct was biphasic as expected. Uninduced XRNA RNAi cells showed a slight retardation of the degradation, relative to wild-type cells, and XRNA RNAi eliminated the fast degradation component.
Modelling using our previous data for the CAT-EP mRNA (61) yielded almost identical results to those for EP (Figure S4A).

Surprisingly, northern blots of RNA from cells containing the XRNA RNAi construct had, in addition to the expected band at 0.9 kb, two additional bands: one at 1.3 kb and one at about 0.4 kb. These were faint, but detectable, in RNA from cells grown without tetracycline (data not shown) and clearly visible after RNAi induction (Figure 4C). Both hybridized with an EP probe containing only the coding region; the shorter product was not detected in the poly(A)+ fraction (data not shown) and could be a coding region fragment. The longer product was polyadenylated; a band of similar size hybridized to an intergenic region probe (data not shown). This RNA is too long to be a polyadenylated primary transcript, but could be a previously detected minor alternatively spliced version of EP2 or EP3 mRNA, extended at the 5' end (86,87). Thus XRNA seems to be implicated in destruction of abnormal, probably non-functional mRNAs from the procyclin locus.

These results indicate that extremely rapid degradation of EP mRNA requires XRNA, but some EP mRNA is degraded by a pathway that depends on the presence of CAF1.

Degradation of the CAT-PGKB and PGKB mRNAs

The endogenous PGKB mRNA, produced by pol II, is virtually undetectable by Northern blotting. Even using real-time PCR, the levels were only just above the detection limit, making accurate quantitation impossible (data not shown). The only conclusion we could make was that as for other polymerase II transcripts, the steady-state level of PGKB mRNA did not appear to have been changed by CAF1 depletion. The mRNA level dropped below the detection limit 5–15 min after inhibition of mRNA synthesis so the degradation rate could not be measured. In order to obtain sufficient mRNA for analysis, we therefore investigated the effects of deadenylase depletion in cell lines expressing the CAT-PGKB reporter mRNA from an RNA polymerase I promoter. The CAT-PGKB mRNA has a half-life of 5–7 min (70,74). Depletion of CAF1 resulted in modest stabilization (Figures 5A and S3B). In two experiments with a PAN2 RNAi cell line, we also saw stabilization of CAT-PGKB mRNA (see for example upper panel of Figure 5B) but this was not seen in two further experiments. Thus the role of PAN2 in CAT-PGKB mRNA degradation is uncertain. From these results we concluded that deadenylation plays a minor role in degradation of CAT-PGKB mRNA.

TbCAF1 depletion partially inhibits 3'→5' degradation of CAT-EP mRNA

To examine the various degradation pathways for CAT-EP in more detail, we depleted CAF1 in a cell line expressing a CAT-GC-EP reporter, which has a C30G30 sequence between the CAT ORF and the EP 3'-UTR (61,68,74). The C30G30 causes stalling of exoribonucleases.
Figure 6. Degradation of the CAT-GC-EP reporter mRNA in trypanosomes. (A) Possible pathways of degradation. The CAT-GC-EP mRNA has, from 5′ to 3′, a cap (small circle), a short 5′UTR (white fill), a CAT open reading frame (black), a G30C30 sequence, then the EP1 3′-UTR (white, with destabilising 26-mer in grey) and poly(A) tail (AAAA). (1) deadenylation as the first step in degradation (product ‘A’); (2) degradation of the deadenylated RNA from the 3′-end, passing at the G30C30 sequence to produce the CAT-GC intermediate (product ‘B’); (3) decapping of the deadenylated mRNA; product ‘C’, indistinguishable from ‘A’ by northern blotting; (4) degradation of product ‘C’ from the 5′-end, passing at the G30C30 sequence to give the deadenylated 3′ GC-EP RNA (product ‘E’); (5) decapping as the first step in degradation, to give product ‘F’, indistinguishable from the initial RNA by northern blotting; (6) degradation of product ‘F’ from the 5′-end, passing G30C30 to give a polyadenylated 3′ GC-EP RNA (product ‘G’); (7) deadenylation of RNA ‘D’ to give RNA ‘E’. (B) The effect of inducible CAF1 RNAi on CAT-GC-EP mRNA degradation. CAF1+ cells without tetracycline; CAF1 with downshock: cells with tetracycline (19–21.5h). Total RNA (T) was prepared and part of it was separated into poly(A)+ (A+) and poly(A)− (A−) fractions. The identities of the bands are shown on the left and sizes on the right. The upper panels were hybridized with an EP 3′-UTR riboprobe. This reproducibly cross-hybridizes with the smallest rRNA fragment. Although this signal has been cropped from the picture, it smears a little into the upper portions of the total and poly(A)− lanes. The lower panels are hybridizations with a CAT probe. The ratio of the CAT-GC fragment to the full-length mRNA fragment was measured in three experiments: for cells without tetracycline the average ratio was 21% (range 15–26%) and for CAF1-depleted cells 14% (range 12–16%). The degradation fragments were detected readily only if we probed for them first: the upper and lower panels therefore originate from different experiments.

The pathways that could be responsible for the various possible degradation intermediates [previously identified in (61)] are depicted in Figure 6A. A CAT probe detected the full-length CAT-GC-EP mRNA (Figure 6B, lane 1), which was recovered in the poly(A)+ fraction (Figure 6B, lane 3). Based on published analyses (88) we estimate that this fraction included RNAs with tails of greater than 20 nt. The CAT probe also detected a fragment containing the 5′-UTR joined to the CAT gene and terminating at the C30G30 sequence (CAT-GC); this was in the poly(A)− fraction as expected (Figure 6B, lanes 1 and 2). A probe for the EP 3′-UTR detected the polyadenylated CAT-GC-EP and EP procyclin mRNAs (Figure 6B, lanes 1 and 3). In addition, the EP probe detected two GC-EP 3′-end fragments, one of which was deadenylated (Figure 6B, lane 2) and one of which was, on average, about 100-nt longer and bound to oligo (dT) (Figure 6B, lane 3). The presence of the latter fragment suggests that 5′→3′ exonuclease is able to target an mRNA with a poly(A) tail longer than 50 nt (61). Meanwhile, the deadenylated GC-EP fragment could arise either from deadenylation of the complete CAT-GC-EP mRNA, followed by 5′→3′ degradation, or from deadenylation of pre-made GC-EP RNA fragments. Thirty minutes after addition of Actinomycin D, only shadows of deadenylated EP, CAT-GC and GC-EP RNAs remained (Figure 6B, lanes 7 and 8); the poly(A)+ lane (lane 9) was reproducibly empty.

We had previously shown that after XRNA depletion, a smear of CAT-GC-EP and EP mRNAs becomes detectable in the steady-state poly(A)− fraction, indicating stabilization of deadenylated mRNA (61) and Figure S4B. Exosome depletion had no effect on the pattern of intermediates at steady state, but partially stabilized deadenylated full-length CAT-GC-EP mRNA (74) and Figure S4B. Depletion of PAN2 did not result in any obvious change in the pattern of CAT-GC-EP mRNA degradation (data not shown). After depletion of CAF1, the only effect at steady state was in a slight decrease in the abundance of the CAT-GC fragment relative to the amount of CAT-GC-EP RNA (Figure 6B, lower panel, lanes 4 and 5; see figure legend for quantitation). This would be expected if the production of CAT-GC were initiated by deadenylation. 30 min after Actinomycin D addition full-length mRNA was more abundant after CAF1 depletion (Figure 6B, visible in lanes 7 and 10 hybridized with the CAT probe). More strikingly, the poly(A)+ lane (Figure 6B, lane 12) contained some CAT-EP mRNA, EP mRNA and the polyadenylated 3′ GC-EP fragment; in contrast, for CAF1+ samples, this lane was completely empty (Figure 6B, lane 9).

These results demonstrate the role of CAF1 in deadenylating the CAT-GC-EP and EP mRNAs, and in initiating 3′→5′ degradation.

**Sequencing of uncapped mRNA reveals multiple degradation pathways**

To more characterize endogenous mRNA degradation intermediates, we used an RNA circularization and sequencing strategy (89). In this procedure (Figure 7A), RNAs with a 5′-phosphate and 3′-OH group are diluted to a low concentration, then incubated with T4 RNA ligase to join the ends, creating circles. After that, the 5′–3′ junction is reverse transcribed, then amplified by PCR (Figure 7A). The products are subsequently cloned and sequenced. To sequence full-length mRNAs, the cap must be removed since capped 5′-ends cannot be ligated; the methylated nucleotides just downstream of the cap must also be removed because they inhibit reverse transcriptase (90–94). In order to obtain ligation products
A Method

<table>
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<th>RNA ligase</th>
<th>RT-PCR</th>
<th>5' cap + methylations</th>
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<td>lanes 1 and 3</td>
<td>Products from 5' oligo RNA ligase ++</td>
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B RPL37A

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<th>RT-PCR</th>
<th>5' cap + methylations</th>
</tr>
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<td>Agarose gel</td>
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<td>Products from 5' oligo RNA ligase ++</td>
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<tr>
<td>Primers b</td>
<td>M 1 2 3 4 5</td>
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</table>

C EP

<table>
<thead>
<tr>
<th>Primer Set</th>
<th>RNA ligase</th>
<th>RT-PCR</th>
<th>5' cap + methylations</th>
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<td>Primers a</td>
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<td>lanes 1 and 3</td>
<td>Products from 5' oligo RNA ligase ++</td>
</tr>
<tr>
<td>Primers b</td>
<td>M 1 2 3 4 5</td>
<td></td>
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</table>

Figure 7. RT–PCR of circularized mRNAs. (A) Illustration of the method for intact, capped RNA (right) and a degradation intermediate (left). The open reading frame is filled grey, other symbols are shown on the figure. For detailed description see text. (B) Ethidium-bromide stained gels of amplified products for RPL37A. A cartoon of the mRNA, not to scale, showing the positions of the primers, is above the gels. Products from the first PCR (primers ‘a’) are shown on the gel containing most DNA was excised for purification and cloning. In each case, the area of the gel containing most DNA was excised for purification and cloning of the PCR products, and clones were selected randomly for sequencing. Gels for amplifications with 3' primers 33 or 34, with 5' primer 51, looked very similar to the lower panel for primers 51 and 32; the same procedure was followed but only clones with the longest inserts were sequenced.

from mRNAs with intact 5'-ends and cap structures, we therefore pre-incubated samples with RNase H and an oligonucleotide complementary to the 5'-end of the spliced leader. When considering the results, it is important to note that if an mRNA had been decapped in vivo, but still retained the 5' methylated nucleotides, it would be detected only in RNase H treated samples. Such decapped RNAs are expected to accumulate after XRNA depletion.

Tandemly repeated precursor RNAs should be substrates for RNA-ligase-independent amplification, and are expected for both HISH4 and TUB. We therefore chose, as a control, the ribosomal protein L37a mRNA (RPL37A), which is encoded by a single locus, has a predicted length of about 400 nt, and has a half-life exceeding 90 min in bloodstream-form trypanosomes (Figure 5B). RNA circle results for RPL37A are shown in Figure 7B, and sequencing results for a random selection of cloned products are in Tables 1 and S2, and Figure S5. The mRNA is trans spliced at a single site, but there are several alternative polyadenylation sites. Figure 7B shows the products of the RT–PCRs. The first PCR showed a ladder in all lanes, except for the negative control without ligase (lane 5). In lanes 1 and 3, showing products from 5' RNase H-treated RNA, there was a broad band around 300 nt, which was absent in the products from RNA that had not been RNase H treated. The predicted length for amplification products without the poly(A) tail was 159–210 bp, so the size of this band suggested average poly(A) tail lengths of about 100 nt. Cloning, however, instead gave poly(A/T) tracts of 36–37 bp. In the original article describing the method (89), the poly(A) tails of four rat mRNAs were measured by gel electrophoresis, without cloning. Amplification of three of them indicated poly(A) tails of 20–50 nt; for one mRNA there were two populations, corresponding to 20–50 nt and 120–200 nt. Since mammalian poly(A) tails are expected to be 100–200 nt long, these results hint that the circle–PCR method may have an intrinsic bias against longer poly(A) tails. Our results show that the bias is exacerbated if the products are cloned.

RPL37A products from RNA that had not been treated with RNase H were expected to correspond to mRNAs that had already been decapped and had begun to be degraded from the 5'-end: they formed a smear in the agarose gel (Figure 7B). In order to reduce contamination with irrelevant cDNAs prior to cloning, we performed a nested PCR. Surprisingly, the 5'-ends of the degradation intermediates were not random. Instead, 25 out of 46 clones had lost the first 27–31 nt of the spliced leader. About half of the degradation intermediates had lost the poly(A) tail and parts of the 3'-UTR as well. In contrast to observations for mammalian histone RNAs (53), here was no evidence for addition of U residues at the 3'-ends of the degrading mRNAs. Depletion of XRNA did not affect the patterns of degradation products. If poly(A) tails were present they were almost always in the 30–40 nt size range. Because of the bias noted earlier, this result gives us the lower limit of poly(A) lengths for deadenylation-dependent decapping, but does not tell us the upper threshold. Just 2 of the 43 cloned intermediates had much...
longer poly(A) tails (93 nt and 112 nt), illustrating the minor role of deadenylation-independent decay.

We next repeated the experiment for EP mRNA. In bloodstream forms, at steady state, EP mRNAs have a poly(A) tail of at least 50 nt, added about 300-nt downstream of the stop codon (e.g. Figures 4 and 6B, lanes 2 and 3). Primers 52 and 31 (Figures 7C and S6) were designed to allow detection of a broad range of degradation products, including those that had lost long segments of the 5' and 3'-ends. First, we examined RNase H-treated mRNA, in order to see the full-length mRNA. The corresponding amplified product should have been 430 bp plus poly(A). To our surprise, a band of this size was not seen and no clones corresponding to full-length mRNA were obtained. Instead, the RNase H-treated RNA yielded EP products that were predominantly in the 200 bp range (Figure 7C, lanes 1 and 3). Randomly selected clones nearly all retained SL sequence, but either lacked a poly(A) tail, or had (A)3–11 at positions over 100-nt upstream of the usual site (Tables 1 and S3, and Figure S7).

We next amplified EP cDNAs using RNaseH-treated RNA from XRNA-depleted cells with primers 52 and 31. The average size of the products was shorter than before (Figure 7B, compare lane 1 with lane 3). Although all eight cloned products had retained the expected portions of the spliced leader, only one of them had a poly(A) tail (Table S3 and Figure S7). This illustrates stabilization of deadenylated CAT-EP mRNA after XRNA depletion (61).

To find out whether the procedure could amplify full-length EP mRNA from bloodstream forms at all, we repeated amplification with primers 33 and 34 and selected only the longest inserts for sequencing (Table S4 and Figure S8). The product distributions on agarose gels were similar to those obtained with the first primer sets (data not shown). Even with primer 34, no clones had poly(A) in the expected position, although one was polyadenylated a few nucleotides further downstream (Figure S8). This result is strange, since it is inconsistent with the northern blot results. It will not be discussed further here, because a considerable number of further experiments is required to determine the origin of these products. In procyclic forms, the EP mRNA is very stable and polyadenylated. In preliminary experiments with RNaseH-treated mRNA from this form, using primers 33 and 34, 6 out of 12 clones were polyadenylated in the expected position at +297, with an average poly(A) length of 29 nt (maximum 58 nt). This shows that the procedure can indeed amplify the full-length polyadenylated RNA, but probably biases towards shorter products and truncates the tails.

Returning to the bloodstream-form trypanosomes, we next analysed circularized EP mRNAs without prior RNAase treatment. Using RNA from cells with normal XRNA levels, approximately half of the amplified products using 3' primer 32 had an oligo(A) tail, well upstream of the expected position; the intermediates without poly(A) had lost variable portions of the 3'-UTR.

It was obvious on the stained agarose gel that products from cells with XRNA RNAi (Figure 7C, lane 2) were smaller than those obtained from cells with a normal amount of XRNA (Figure 7C, lane 4). The clones from the XRNA-depleted cells mostly lacked poly(A) and had been digested significantly further at the 5'-end than those from the normal cells: 56% had lost part of the coding region, and only 1/18 had retained part of the spliced leader, while in XRNA positive cells, 14% had lost coding sequence and 64% retained part of the spliced leader. Surprisingly although the coding region primers were designed to hybridize equally to EP1, EP2 and EP3 sequences, all clones originated from EP1. Two clones (no RNase H, normal XRNA) originated from the primary transcript, enabling precise mapping of the transcription start site at position –115 relative to the start codon.

---

Table 1. Summary of results for circularized RNAs

<table>
<thead>
<tr>
<th>Gene</th>
<th>RNaseH</th>
<th>XRNA</th>
<th>#</th>
<th>Nt 5'</th>
<th>nt 3'</th>
<th>pA (%)</th>
<th>(A)n</th>
</tr>
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<tbody>
<tr>
<td>RPL37A full</td>
<td>+</td>
<td>+</td>
<td>5</td>
<td>55</td>
<td>108–136</td>
<td>36±4</td>
<td></td>
</tr>
<tr>
<td>RPL37A circles</td>
<td>-</td>
<td>+</td>
<td>23</td>
<td>38±4</td>
<td>108, 127, 136</td>
<td>43</td>
<td>12–28</td>
</tr>
<tr>
<td></td>
<td>+</td>
<td>-</td>
<td>5</td>
<td>27±8</td>
<td>78±42</td>
<td>44 (5–93)</td>
<td>36±5</td>
</tr>
<tr>
<td></td>
<td>-</td>
<td>-</td>
<td>20</td>
<td>39±4</td>
<td>108,136</td>
<td>100</td>
<td>34 (5–112)</td>
</tr>
<tr>
<td>EP full</td>
<td></td>
<td></td>
<td></td>
<td>20±15</td>
<td>97±37</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td>EP circles</td>
<td>+</td>
<td>+</td>
<td>8</td>
<td>70</td>
<td>312</td>
<td>62</td>
<td>7 (3–11)</td>
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<td>-</td>
<td>+</td>
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<td>49±25</td>
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<td>5</td>
<td>38±33**</td>
<td>29±35</td>
<td>59</td>
<td>12 (3-28)</td>
</tr>
<tr>
<td></td>
<td>-</td>
<td>-</td>
<td>8</td>
<td>57±1</td>
<td>28±22</td>
<td>12</td>
<td>10</td>
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<td></td>
<td></td>
<td></td>
<td>16</td>
<td>−11±28*</td>
<td>76±98</td>
<td>19</td>
<td>5 (4,5)</td>
</tr>
</tbody>
</table>

Results are shown for products of RPL37A amplified with the ‘b’ primers (Figure 7B) and for EP mRNAs amplified with 5’ primers 51 or 52, and 3'-primer 32 (Figure 7C). 'RNase H'—removal of 5'-end of SL by RNase H digestion; 'XRNA'—presence of normal XRNA (+) or depleted XRNA (−). #: denotes number of clones sequenced; in each case, clones with various insert lengths were selected. Results for 5' and 3' ends (nt 5', nt 3') are the numbers of nucleotides beyond the ATG start codon or the stop codon, respectively, expressed as arithmetic mean±SD, including clones with and without poly(A). The lengths for full-lengths RNA are given for reference (‘full’) and are shown in italics. 'pA' shows the percentage of clones that had a poly(A) tail. The 'p(A)n' lane indicates the arithmetic mean lengths of poly(A) tails, where present, with either the standard deviation or the range.

**Not including the primary transcripts.

*Significantly different, P < 0.001 (t-test).
Finally, results in Figure 4C had suggested that XRNA depletion stabilized an intermediate of about 400 nt. Since this fragment had not been detected from CAT-EP mRNAs after XRNA depletion (61), we suspected that it might depend on the EP coding region. We looked for such RNAs using primers 52 and 35; the latter hybridizes within the repetitive region encoding the EP repeats. The result was a clean ladder of products which was unaffected by either RNase H decapping or XRNA depletion (Figure 5C). All clones obtained, (with or without XRNA depletion) started at position +51 downstream of the ATG, and ended at position 10 in the 3′-UTR. This was paradoxical since no clones obtained with other primers had these boundaries.

Two novel findings emerged from this approach. First, we saw hitherto unsuspected heterogeneity in polyadenylation of EP mRNAs in bloodstream forms. Second, XRNA depletion caused accumulation of EP mRNA degradation intermediates that lacked not only the poly(A) tail, but also portions of the 5′-end. We were disappointed that we could not establish an upper poly(A) length threshold for deadenylation-dependent decapping; nevertheless, it was interesting that cloned EP mRNAs had much shorter poly(A)s than the RPL37A clones.

**RNAi targeting a MEX67 homologue does not accelerate EP mRNA decay**

XRNA is present in both the nucleus and the cytoplasm, so could theoretically attack EP and PGKB mRNA immediately after synthesis, prior to nuclear export. To address this possibility, we searched for homologues of the export factor MEX67 (human TAP). The protein predicted from one, Tb927.5.1270, is only 216 amino acids long, and RNAi had no effect on growth (data not shown). The protein encoded by Tb11.22.0004 is, in contrast, closer to the expected length, has two NTF2 homology domains (superfamily SSF54427), and is 30% identical (49% similar) to mouse TAP/MEX67 over the first 158 amino acids. RNAi targeting Tb11.22.0004 inhibited growth (Figure 8A). We used in situ hybridization to examine the distribution of poly(A) after RNAi targeting Tb11.22.0004. In normal cells, or RNAi cells without tetracycline addition, poly(A) was always distributed uniformly throughout the cytoplasm and nucleus (Figure 8B).
After RNAi, the cells showed a range of phenotypes, varying from normal poly(A) staining to strong retention of poly(A) in the nucleus (Figure 8C). These results, seen with several independent RNAi lines, suggest that the Tb11.22.0004 locus indeed encodes a MEX67-like protein that is required for mRNA export, so we have named the gene TbMEX67.

If the XRNA pathway were in the nucleus, one would expect a decrease in EP and PGKB mRNAs after MEX67 RNAi. Instead, MEX67 RNAi reproducibly caused an increase in both mRNAs (Figure 8D and E). At the same time, PGKC mRNA decreased. In normal bloodstream forms, the PGKC:PGKB ratio was greater than 10:1; after MEX67 RNAi, the ratio changed to about 4:1. Results of a preliminary cell fractionation experiment suggested that even after 48 h tetracycline treatment, most of the EP mRNA was in the cytoplasm (data not shown). This is consistent with the in situ hybridization results suggesting that mRNA export from the nucleus had slowed down, but not stopped. These results lent no support to the notion that degradation of either EP or PGKB mRNA occurs in the nucleus.

**DISCUSSION**

**Roles of PAN2 and PAN3 in trypanosomes**

In yeast and animals, the roles of the PAN2/PAN3 complex may be mainly in quality control—trimming poly(A) tails in the nucleus and cytoplasm. The complex is not essential in yeast, and appears to have been lost many times in eukaryotic evolution. In addition, some organisms have truncated or mutant versions of PAN2 which have lost active site residues. PAN3 was present in most organisms that had PAN2, but absent in all that lacked it. Even in *Plasmodium*, which shows translational regulation (95) and has few potential regulatory transcription factors (96), we could find no PAN2/3.

Homologues of PAN2 and PAN3 are present in the trypanosome genome. Results from RNAi suggested that PAN2 is required for optimal trypanosome growth. We found no indication that PAN3 is required for PAN2 activity, and evidence for a PAN2–PAN3 interaction was weak. The effects of RNAi targeting PAN2 were difficult to analyse since the RNAi effect was rapidly lost during culture. Nevertheless, it was clear that even when PAN2 depletion was sufficient to stop trypanosome growth, the degradation kinetics of several mRNAs were unchanged. In contrast, there was clear stabilization of the regulated EP mRNA and mild inhibition of ACT and CAT-PGKB mRNA decay. These results suggest that PAN2 may have a specialized role in degrading some, but not all mRNAs.

**Decapping in trypanosomes**

Decapping has been seen in trypanosome extracts (60), and was also measured indirectly in *Leishmania* (97). RNAi targeting the only reasonable candidate for a trypanosome DCP2-like decapping enzyme, Tb10.70.2530, did not affect cell growth (for details see Supplementary Data). We therefore wondered whether trypanosome mRNAs were decapped by some other mechanism. Since all mRNAs have a 5′ SL, sequence-specific endonuclease digestion within the SL could substitute for decapping. To analyse this we used RNA ligation and PCR to clone mRNA degradation intermediates. When we examined decay intermediates of the mRNA encoding ribosomal protein L37A, there was a favoured 5′-end, at positions 28–31 of the SL. However, the frequency of this was not increased after XRNA depletion, as would be expected for an endonuclease product. Moreover, some L37A RNAs had more SL sequence, and a similar pattern was not seen for EP mRNA. The result for L37A could be an artefact of the method. If it is not, there might be an SL-specific endonuclease, affecting some, but not all, mRNAs. Alternatively, XRNA (or another unidentified 5′–3′ exoribonuclease) may pause in vivo at position 28–31 because of secondary structure or the presence of bound protein.

**CAF1 depletion increased the steady-state abundance of EP mRNA**

Depletion of CAF1 almost completely stopped degradation of *HISH4, TUB, ACT* and *PGKC* mRNAs but the steady-state abundances were unaffected (14). We previously suggested that when mRNA degradation is stopped, a feedback mechanism might inhibit RNA polymerase II transcription. In contrast, depletion of CAF1 selectively increased the abundances of EP mRNA, which is made by RNA polymerase I. It is therefore possible that if a feedback mechanism exists, RNA polymerase I is less susceptible to it than RNA polymerase II. Also, it has recently been shown in *S. cerevisiae* that the Rpb4 and Rpb7 subunits of RNA polymerase II play roles in mRNA degradation (98,99). It is conceivable that these subunits also influence mRNA decay in trypanosomes (100).

**Deadenylation and decapping of the L37A mRNA**

In mammalian cells, decapping occurs on mRNAs that have reduced poly(A) tail lengths; the threshold may correspond to a minimum number of 25-nt-binding sites for PABP (101,102). In studies of budding yeast, the deadenylation limit is indeed 10–20 residues (103,104) though not many transcripts have been analysed in detail. A study of four mammalian liver mRNAs, using circularization, found that for one, decapped mRNAs had poly(A) tails of 20–60 nt, but the three others had shorter tails; there was no obvious relationship between the minimum tail length and the mRNA half-life (89). The stable betaglobin mRNA exhibits synchronous deadenylation until the poly(A) tail has a length of about 110 nt; after that, an asynchronous reduction to A(20) is observed before the mRNA is degraded (26). In contrast, completely deadenylated cmyc mRNA has been detected (105).

Our previous experiments had suggested that the poly(A) tails on histone H4 mRNA are reduced to about 50 nt before the mRNA body is degraded (14,61). Our attempts to analyse overall poly(A) lengths of the L37A mRNA by sequencing RT–PCR amplified, circularized mRNAs were thwarted by methodological bias towards shorter products. Nevertheless, the majority of
products were consistent with deadenylation-dependent decay. XRNA depletion had no effect on the overall pattern, and we can suggest a lower limit of \(A_{30-40}\) for the threshold for decapping. The presence of short tails in nearly half of the decapped intermediates implies a temporal and mechanistic discontinuity in the decay process: after deadenylation reaches the threshold, the mRNA is decapped, but there is then a pause while the short-tailed, decapped mRNA is ‘handed over’ to the 5′–3′ exonuclease and/or the exosome.

**Polyadenylation of EP mRNA**

All evidence from northern blotting indicates that most EP mRNAs in bloodstream-form trypanosomes are polyadenylated at approximately the same position as in procylic forms, about 300 nt downstream of the stop codon, and that the poly(A) tail is at least 50-nt long (see e.g. Figure 4). We were therefore surprised that after circle-PCR, almost all of the cloned polyadenylated EP RNAs had shorter 3′-UTRs. If such mRNAs were as abundant as the cloning results imply, the northern would have a smear and no full-length band at all. These mRNAs are nevertheless probably present, albeit at much lower abundance than the cloning results suggest. They could arise by use of alternative processing signals: this has been seen for EP before, but only in reporters with deletions of the wild-type sequence (87,106). Alternatively, oligo(A) tails might have been added to degradation intermediates in the cytoplasm. The exosome from some Archaea has the ability to add nucleotides to RNA; adenosine is added preferentially because ATP is most abundant in the cytoplasm (107). The trypanosomal exosome lacks RNase PH activity (108), so is unlikely to be able to catalyse the polyadenylation reaction, but some other cytoplasmic protein might conceivably have this function.

**Pathways of EP mRNA degradation**

We previously suggested that in bloodstream-form trypanosomes, CAT-EP, EP and PGKB mRNAs are destroyed by two pathways, one dependent on XRNA and the other on deadenylation. The results presented here suggest a more complicated picture. The presence of very active, and at least partially deadenylation-independent, 5′–3′ pathway for EP mRNA degradation is supported by the presence of polyadenylated CAT-GC-EP degradation intermediates, and by the inhibition of degradation by depletion of XRNA. However, we also found deadenylated CAT-GC-EP intermediates, and 3′–5′ exonuclease products from both CAT-GC-EP and EP. The EP mRNA degradation kinetics included a fast, XRNA-dependent component and a slower one dependent on CAF1. Even the rapid degradation pathway, had a 3′-exonucleolytic component: it was made slower by CAF1 depletion, and was delayed by RNAi targeting the exosome (74).

Yeast XRN1 is processive on a poly(A) substrate (109). If XRNA were processive on mRNA in vivo, a reduction in the XRNA level should cause accumulation of full-length, but decapped EP mRNA, but should not change the overall distribution of mRNAs in which are ‘caught in the act’ of 5′ degradation at the time of RNA isolation. If XRNA were distributive in vivo, RNAi should cause accumulation of many different 5′ degradation products; the same picture would be obtained if, after XRNA depletion, decapped mRNAs were degraded by a different, distributive, 5′–3′ exoribonuclease. Inhibition by secondary structures, bound proteins and ribosomes could also affect the pattern. We found that the cloned intermediates from XRNA-depleted cells were significantly shorter at the 5′-end than those from normal cells. There are multiple possible explanations, and we present just one, highly speculative, scenario here. Suppose that in normal cells, XRNA is responsible for most 5′–3′ degradation: its activity is transiently delayed by the exon junction complex or other proteins bound to the spliced leader; but once over this barrier, digestion is processive. This would explain the predominance of intermediates that retain parts of the spliced leader in normal cells. In the absence of XRNA, decapped mRNA accumulates, and is either attacked from the 3′-end, or degraded from the 5′-end by other, distributive exoribonucleases, or by endonucleases.

When CAF1 was depleted, half of the EP mRNA was degraded within 15 min, but the rest appeared to be very stable, and so apparently not susceptible to degradation by other enzymes. This suggests mechanistic separation of the CAF1-dependent and -independent pathways. Temporal separation, with deadenylation-independent degradation only within the first 15 min after synthesis, would be easily understood if the XRNA pathway were located in the nucleus; but this is unlikely, since EP mRNA accumulated after depletion of MEX67. CAF1 RNAi might also result in abnormal sequestering of mRNAs in stress granules (62) or other structures (110). We looked for this, but 24 h after induction of RNAi against either CAF1 or NOT1 there was little, if any, effect on the distribution of the helicase DHH1 (data not shown), which is a stress granule marker (62).

We suggest that the choice of mRNA degradation pathway is dictated by binding of different combinations of regulatory RNA-binding proteins. The RPL37A, HISH4 and other relatively stable mRNAs are bound by combinations of RNA-binding proteins that actively prevent degradation (111); deadenylation is slow, but at a threshold (minimally, \(A_{30-40}\)) there is a pause before subsequent attack at the 5′ and 3′ ends. Protein combinations specific for EP, PGKB and related unstable mRNAs, in contrast, can actively recruit either XRNA and the elusive decapping enzyme, or deadenylases; this causes either very rapid degradation initiated from the 5′-end, or a rapid entry to the deadenylation-dependent pathway. If XRNA is not available, the mRNA defaults to the deadenylation-dependent pathway. Possibly, the RNA-binding proteins route the highly unstable mRNAs directly to P-bodies.

This scenario is entirely consistent with results from other eukaryotes. For example, the alpha1 tubulin mRNA of *Chlamydomonas reinhardtii* is normally degraded via deadenylation followed by decapping and 5′–3′ digestion. After deflagellation, synthesis of the mRNAs is induced, but their half-lives decrease to 10–15 min, and 5′ degradation occurs without deadenylation (112). Mammalian mRNAs with different AU-rich
elements also show different deadenylation kinetics (113). The next challenge, for trypanosomes, consists in identifying the RNA-binding proteins that guide these choices.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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