A long noncoding RNA promotes parasite differentiation in African trypanosomes

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The parasite Trypanosoma brucei causes African sleeping sickness that is fatal to patients if untreated. Parasite differentiation from a replicative slender form into a quiescent stumpy form promotes host survival and parasite transmission. Long noncoding RNAs (lncRNAs) are known to regulate cell differentiation in other eukaryotes. To determine whether lncRNAs are also involved in parasite differentiation, we used RNA sequencing to survey the T. brucei genome, identifying 1428 previously uncharacterized lncRNA genes. We find that grumpy lncRNA is a key regulator that promotes parasite differentiation into the quiescent stumpy form. This function is promoted by a small nuclear RNA encoded within the grumpy lncRNA. snoGRUMPY binds to messenger RNAs of at least two stumpy regulatory genes, promoting their expression. grumpy overexpression reduces parasitemia in infected mice. Our analyses suggest that T. brucei lncRNAs modulate parasite-host interactions and provide a mechanism by which grumpy regulates cell differentiation in trypanosomes.

INTRODUCTION

Trypanosoma brucei is a unicellular kinetoplastid parasite that cycles between an insect (tsetse fly) and a mammalian host. Once T. brucei reproduces to a critical density in mammalian blood, a quorum-sensing mechanism is activated, and parasites differentiate into a quiescent, nondividing stumpy form (1). This shift in parasite lifestyle limits parasite population size and extends host survival, making it a promising step for therapeutic intervention. The stumpy form also facilitates transmission to the tsetse fly vector and development into insect procyclic forms (2). In T. brucei, parasite density is sensed via the stumpy induction factor (SIF) (1) and the SIF signaling pathway, which promotes gene expression as well as morphological and metabolic changes associated with the stumpy form (3). To date, 43 genes have been shown to function in the SIF signaling pathway, playing roles that range from signal transduction to signal response (3). RNA-binding protein 7A (RBP7A) and RBP7B, proteins that are predicted to bind RNA, are necessary for SIF-induced stumpy formation (3). RBP7A/B-null mutant parasites are unresponsive to the SIF signal and are unable to differentiate into stumpy forms. RBP7 genes are therefore crucial regulators of parasite differentiation, yet their mode of action and target genes are unknown (4, 5).

In mammals, transcription is pervasive, with 80% of genomic DNA being actively transcribed in different cell types or physiological situations, yet only a small fraction (less than 3%) of the genome encodes protein-coding genes (6). The vast majority of the mammalian genome encodes noncoding DNA sequences such as introns, pseudo-genes, transposons, telomeres, or noncoding RNAs (ncRNAs). ncRNAs are divided into two groups, small ncRNAs (<200 nucleotides [nt]) and long ncRNAs (lncRNAs) (>200 nt). Small ncRNAs include RNAs that regulate splicing, small nuclear RNAs (snRNAs), tRNAs that help decode mRNA sequences into proteins, small interfering RNAs and microRNAs that regulate mRNA stability and translation, small nucleolar RNAs (snoRNAs) that regulate ribosomal RNA (rRNA) processing and modification, and others (7). T. brucei has all of the canonical small ncRNAs described above except for microRNAs (8).

In eukaryotes, lncRNAs resemble mRNAs as they harbor a 5′ cap and are polyadenylated (9). Their abundance is comparable to that of protein-coding genes (10). lncRNAs function in many cellular pathways (11–13), including cell differentiation (14, 15). This family of RNAs can regulate cell fate choice by either promoting or inhibiting differentiation. For example, skin stem cells express the ANCR (antidifferentiation ncRNA) and TINCR (terminal differentiation ncRNA) lncRNAs, which function antagonistically. While ANCR suppresses the epidermal differentiation pathway and maintains the stem cell compartment, TINCR promotes epidermal terminal differentiation (16, 17). lncRNAs also regulate antigenic variation of Plasmodium falciparum (18, 19) and the host cell response during Toxoplasma gondii infection (20), indicating that they are important players in parasite infection.

To date, only 95 putative lncRNA genes have been annotated in T. brucei, all with unknown functions (21). Compared to 9598 T. brucei protein-coding genes (22), this small number prompted us to analyze the noncoding repertoire of T. brucei. Here, we find 1428 additional T. brucei lncRNAs, including grumpy. We uncover a mechanism whereby grumpy encodes a snoRNA that regulates parasite differentiation.
RESULTS

We used a combination of strand-specific and paired-end RNA sequencing (RNA-seq), in silico analysis, and database integration to reannotate the lncRNA gene repertoire of *T. brucei* (Fig. 1A and figs. S1 to S5). We identified 1428 previously uncharacterized transcripts that are longer than 200 nt, have a low coding potential score, few ribosomal interactions, and which do not encode any unique peptides (tables S1 and S2 and figs. S2 to S5). These putative lncRNAs are scattered in a mostly intergenic fashion throughout the 11 chromosomes of the *T. brucei* genome (fig. S6). They are shorter, less expressed, and less GC-rich than *T. brucei* mRNAs (fig. S7), but they otherwise harbor regular mRNA trans-splicing/polyadenylation motifs (fig. S8). We confirmed the full-length sequences from the polyadenylate [poly(A)] tail to the splice leader sequence of 1016 of 1428 lncRNAs using Nanopore direct RNA-seq (Fig. 1B, table S3, and fig. S9). Nanopore sequencing also revealed that poly(A) tails of lncRNA tend to be longer (most frequently ~112 nt) than the poly(A) tails of other transcripts (~95 nt) (Fig. 1C). This opens up an avenue for future investigation of polyadenylation and processing of lncRNAs in trypanosomes. We detected some of these transcripts either in the nucleus or in the cytoplasm at various stages of the *T. brucei* life cycle (Fig. 1D). Notably, 25% of the newly identified lncRNA are differentially expressed between mammalian bloodstream form (BSF) and insect procyclic form (fig. S10 and table S4), a greater percentage than protein-coding transcripts (only 16% respond to parasite transition). The *T. brucei* lncRNA gene repertoire is substantial (11% of total genes) and shows a highly dynamic expression pattern during the parasite life cycle.

We analyzed output from an RNA interference (RNAi) screen (23) to test our hypothesis that lncRNAs are involved in parasite differentiation. We found a total of 399 lncRNA genes that appear to be required for differentiation to occur (Fig. 1E and table S5), consistent with our expectation that *T. brucei* lncRNAs regulate parasite transition and adaptation between mammalian and insect vector hosts. In other eukaryotes, lncRNAs have been reported to regulate cell differentiation by modulating the expression of their neighboring genes (14). We found 19 *T. brucei* lncRNAs genes located immediately upstream or downstream of 18 of the 43 SIF pathway genes (table S6). The lncRNA Ksplice-3137A, which we named grumpy (for “regulator of growth and stump formation”), is located upstream of RBP7A and RBP7B (Fig. 1A), which are both required for SIF-induced stump formation (3). *grumpy*’s pattern of expression is similar to that of RBP7, which is transcribed both in the BSF and the procyclic forms (PCF) of *T. brucei* (Fig. 2A). However, unlike RBP7 transcripts, grumpy does not stably interact with *T. brucei* ribosomes (Fig. 2A) and does not produce detectable peptides (table S2).

To further characterize the grumpy transcript, we used a circular reverse transcription polymerase chain reaction (cRT-PCR) assay, in which *T. brucei* RNAs are circularized via their 5’ to 3’ end junctions, amplified, and sequenced. We used gene-specific primers to confirm that grumpy is a trans-spliced and polyadenylated lncRNA transcript expressed as, at least, five different isoforms, including the smallest [359 base pairs (bp)], the major (397 bp), and the longest forms (432 bp) (Fig. 2B). These findings are consistent with the Ksplice in silico analysis, which revealed 1 splice-acceptor site and 10 alternative polyadenylation sites for grumpy (Fig. 2B). We also used Nanopore direct RNA-seq to confirm the full-length sequence of grumpy lncRNA in BSF and procyclic form (Fig. 2C).

RT quantitative PCR (qPCR) analysis reveals that RBP7 family genes and grumpy lncRNA expression differ during parasite differentiation. These different gene expression behaviors are independent of whether transition from the slender to stumpy form is induced by a chemical compound [pCPT-cAMP (chlorophenylthio)adenosine 3’,5’-cyclic monophosphate sodium salt)] or produced by cell density (Fig. 3A and fig. S11). While RBP7B mRNA decreases during stumpy formation, RBP7A and grumpy are up-regulated. Moreover, RNA–fluorescence in situ hybridization (FISH) analysis revealed changes in the subcellular localization of grumpy during stumpy formation (Fig. 3B). Whereas in slender forms, grumpy localizes in three distinct nuclear foci (one in the nucleolus and two in the nucleoplasm), in stumpy forms, grumpy localizes in a single nucleolar focus (Fig. 3B). Moreover, we observed that the signal intensity of nucleolar grumpy increases with time during stumpy formation (Fig. 3C). Contrary to our initial hypothesis, these changes in subcellular localization suggest that grumpy may act through a trans-acting mechanism, which means that grumpy acts in a nuclear localization different from its transcription locus.

Subcellular localization of lncRNAs is often regulated through recognition of RNA motifs and interactions with RNA binding proteins (24, 25). To study subcellular localization changes of grumpy during stumpy formation, pulldowns were performed in vitro using the stumpy form of nuclear lysates. Mass spectrometry (MS) analysis revealed seven RNA binding proteins that bind significantly to grumpy (Fig. 4A and table S7). The two proteins that showed the highest levels of association with grumpy are the heterogeneous nuclear ribonucleoprotein F/H (hnRNP F/H) and the double RNA binding domain protein 3 (DRBD3). hnRNP F/H and DRBD3 are two major factors that regulate RNA splicing and polyadenylation in *T. brucei* (26, 27). hnRNP F/H is a nuclear factor highly expressed in BSFs that regulates both mRNA stability and trans-splicing (26). hnRNP F/H can also repress trans-splicing of specific mRNAs (26). DRBD3 can be found both in the cytoplasm and nucleus of the parasite where it performs distinct functions (27). DRBD3 acts as stabilizer of a subset of mRNAs encoding for developmentally regulated membrane proteins (28), ribosomal proteins, and translation factors (29). DRBD3 moves to the nucleus in response to environmental cues (29). We validated the interaction of DRBD3 with grumpy in vivo using RNA immunoprecipitation and qPCR assays (or RIP-qPCR) (Fig. 4B).

Given that grumpy is recognized by and interacts with splicing factors involved both in trans-splicing and RNA stability, we examined whether grumpy RNA is further processed and matured in stumpy-stage parasites. A total of 88 nt of grumpy encode for a snoRNA gene (snoRNA), originally named TB10CS’2C1 and which we call here snoGRUMPY (fig. S12) (30, 31). Northern blot analysis using an antisense RNA probe targeting the snoRNA sequence reveals (i) that the grumpy lncRNA is processed into snoGRUMPY and (ii) that snoGRUMPY is more abundant in stumpy forms compared to the other life forms tested (Fig. 4C). Fractionation of whole-cell lysates in 10 to 30% sucrose gradients shows that snoGRUMPY is associated with smaller RNPs (fractions 1 to 3) and that grumpy lncRNA fractionates in a slightly heavier fraction (fraction 5) (fig. S13), suggesting that the lncRNA associates with protein(s) in addition to the core C/D small nucleolar RNP (snoRNP) that associates with snoGRUMPY (fig. S13).

snoGRUMPY maturation appears to depend on hnRNP F/H protein as parasites depleted of this splicing factor show lower levels of...
snoGRUMPY (Fig. 4D). DRBD3 depletion did not affect snoGRUMPY levels (fig. S14A). snoGRUMPY localizes mainly in the nucleolus of the cells (Fig. 4E), but it can also be found associated with the Cajal body (fig. S14B). snoGRUMPY, similar to grumpy, is up-regulated during stumpy formation induced both by cell density (Fig. 4F) and by CPT (fig. S15). snoGRUMPY is up-regulated two times more than grumpy during stumpy formation, suggesting that snoGRUMPY is processed from grumpy (Fig. 4F).

To identify the function of grumpy and snoGRUMPY, we started by overexpressing the full-length grumpy lncRNA (fig. S12). Upon

Fig. 1. Identification of 1428 lncRNAs in T. brucei. (A) Pipeline used for the identification of lncRNAs genes in T. brucei. (1) Strand-specific and paired-end RNA-seq. (2) Ksplice identified putative genes whose transcripts contained an SL sequence (SL) and a poly(A) tail (PA) at the extremities. Ksplice used LAST (48) to map RNA-seq reads to the T. brucei genome. (3) The noncoding nature of the putative lncRNAs was predicted from a low coding potential calculator (CPC) score, poor association with ribosomes, and no detectable peptides. grumpy lncRNA is intergenic and immediately upstream of RBP7 genes, previously shown to be involved in the SIF-dependent pathway. (B) The number of full-length lncRNAs [from the SL sequence to the poly(A) tail] sequenced with Nanopore in four RNA samples: two from BSF parasites (BSF) and two from procyclic forms (PCF). (C) Distribution of poly(A) tail lengths in lncRNA candidates versus other transcripts. (D) Subcellular localization of Ksplice IncRNA genes in slender forms (SL), stumpy forms (ST), and PCF of T. brucei, using RNA-FISH. (E) The number of Ksplice IncRNA genes that cause loss of parasite fitness upon down-regulation by RNAi [extracted from RIT-seq analysis (23)]. RNAi was induced in BSFs for 3 days (BSD3), 278 lncRNAs; in BSFs for 6 days (BSD6), 341 lncRNAs; during in vitro parasite differentiation from BSF to insect procyclic forms (DIF), 400 lncRNAs; in PCF, 402 lncRNAs. The total number of lncRNA genes essential for parasite fitness in this screen was 649.
induction, the levels of grumpy transcripts increased three- to fourfold, and that of snoGRUMPY increased eight- to ninefold (fig. S16), while the transcript levels of RBP7A and RBP7B remained unchanged (Fig. 5A). FISH probes spanning the full-length grumpy (fig. S12) show that the overexpressed RNAs retained the original nucleolar localization (Fig. 5B). We observed that exogenous expression of grumpy repressed T. brucei growth and increased life span in vitro (Fig. 5, C and D). We asked whether this reduction in parasite growth could be explained by a higher proportion of the stumpy forms in culture. Stumpy formation occurs only at high parasite density via the SIF-dependent quorum-sensing mechanism and can be quantified using flow cytometry (32, 33) by measuring the fraction of transgenic parasites expressing the fluorescent stumpy marker GFP::PAD1 (green fluorescent protein::protein associated with differentiation 1) 3′ UTR. After 2 days in culture, 60% of the grumpy-overexpressing parasites were in the stumpy form, compared to 7% in the parental line cultured for the same time period, suggesting that grumpy accelerates stumpy formation (Fig. 5E). This phenotype is not observed upon overexpression of other lncRNAs (fig. S17). grumpy overexpression also led to a lower parasite density via the SIF-dependent quorum-sensing mechanism and can be quantified using flow cytometry (32, 33) by measuring the fraction of transgenic parasites expressing the fluorescent stumpy marker GFP::PAD1 (green fluorescent protein::protein associated with differentiation 1) 3′ UTR. After 2 days in culture, 60% of the grumpy-overexpressing parasites were in the stumpy form, compared to 7% in the parental line cultured for the same time period, suggesting that grumpy accelerates stumpy formation (Fig. 5E). This phenotype is not observed upon overexpression of other lncRNAs (fig. S17). grumpy overexpression also led to a lower parasite density.
(<0.7 × 10^6 cells/ml) compared to the parental culture (1.4 × 10^6 cells/ml) (Fig. 5C). Parasites overexpressing *grumpy* displayed all the hallmarks of being in stumpy form, including PAD1 protein expression at the cell surface (Fig. 5F), arrest at the cell cycle G0-G1 phase (Fig. 5G), and preadaptation to differentiate into the insect procyclic stage (fig. S18).

To confirm these results in vivo, we induced mouse infections with parasites overexpressing *grumpy* and measured parasitemia, mouse survival, and stumpy formation, which were compared to infection by the parasite parental line. Mice infected with the parental cell line showed a typical infection profile characterized by successive waves of parasitemia (Fig. 6A) and an average survival of 43 days.
Fig. 4. grumpy lncRNA is processed into a snoRNA, called snoGRUMPY. (A) Protein partners of grumpy lncRNA identified using in vitro RNA-protein pulldown assay. Results are shown as the mean difference between pulldown assays performed with grumpy lncRNA and control lncRNA-5090a. Note that mitochondria proteins found in this in vitro RNA-protein pulldown assay could come from mitochondria contamination (see Materials and Methods). (B) RIP assay using anti-DRBD3 antiserum in BSFs and procyclic forms. Results are shown as the means (SEM, n = 3) and compared to RIP using control serum. Statistical test: two-sided t test, *P < 0.05, **P < 0.005, ***P < 0.0005, and ****P < 0.00005. (C to F) snoRNA encoded by grumpy lncRNA, named snoGRUMPY. (C) Northern blot analysis using probe against snoGRUMPY in procyclic forms, slender forms, and stumpy forms. tRNA<sup>ser</sup> serves as a loading control. The first lane shows the molecular marker (Mkr) and its different sizes annotated in nucleotides. (D) Northern blot analysis using probe against snoGRUMPY in RNAi hnRNP F/H cell line (in procyclic forms). 7SL RNA serves as a loading control for the Northern blot. Bottom: Western blot analysis showing the depletion of hnRNP F/H protein. ZC3H41 serves as a loading control for Western blot. (E) RNA-FISH analysis showing partial colocalization between snoGRUMPY and the nucleolar marker NHP2 (in procyclic forms). (F) grumpy and snoGRUMPY transcript level measured by RT-qPCR during the transition from slender to stumpy forms induced by the SIF signal (cell density). Tb927.2.2220 is used as a control to normalize transcript levels. Results are shown as means (SEM, n = 3), statistical test: two-way ANOVA (Dunnett’s multiple comparisons test), adjusted P values: *P < 0.05; ***P < 0.001; ****P < 0.0001.
By contrast, mice infected with parasites overexpressing _grumpy_ showed no detectable parasitemia and did not die from the infection (>100 days) (Fig. 6, A and B). When _grumpy_ overexpression was induced 4 days after infection, the parasites succeeded in establishing an infection (Fig. 6A), with three of four mice dying from the infection and mouse survival time increasing from approximately 43 to 72 days (Fig. 6B). Thus, _grumpy_ overexpression substantially reduces parasite virulence in mice.

Our in vivo analysis also recapitulated in vitro observations with respect to stumpy forms and density. Wild-type parasites started differentiating into stumpy forms (>20% stumpy forms in the blood) only at high parasitemia (>1.5 × 10^7 parasites/ml), whereas _grumpy_-overexpressing parasites differentiated into stumpy forms when parasitemia was as low as 1.1 × 10^6 parasites/ml (Fig. 6C). These results support the notion that _grumpy_ overexpression triggers premature _T. brucei_ differentiation into...
stumpy forms, which is associated with a reduction in parasite virulence.

Given that snoGRUMPY is processed from grumpy, we next studied the effect of overexpressing only snoGRUMPY in parasite differentiation (fig. S12). snoGRUMPY was overexpressed ~4 times from an exogenous genomic location (mini-chromosome) (Fig. 7A). Overexpressed snoGRUMPY localizes in the nucleolus (Fig. 7B), reduces parasite growth, and promotes differentiation into stumpy forms (Figs. 7C and 7D), recapitulating the grumpy lncRNA overexpression phenotype. Our results show that premature parasite differentiation into stumpy forms induced by grumpy lncRNA is likely mediated by the action of snoGRUMPY.

Next, we tried to study the function of grumpy or snoGRUMPY using loss-of-function strategies. All our attempts to knock out grumpy lncRNA were unsuccessful; therefore, we decided to use RNAi to strive for depletion of grumpy and/or snoGRUMPY (fig. S12). Although our RNAi targeting full-length grumpy successfully depleted ~80% of grumpy lncRNA at its 5’ end, the level of snoGRUMPY remained unchanged (fig. S19). Similarly, an RNAi construct that specifically targets the snoGRUMPY sequence did not succeed in reducing the levels of the snoRNA (figs. S12 and S20). Therefore, it appears that snoGRUMPY is protected from degradation, probably via interaction with snoRNPs (35, 36).

Next, we tried gymnastic delivery of locked nucleic acid (LNA) gapmeRs (fig. S21). While this strategy failed to deplete grumpy, this gapmeR induced the cleavage of the grumpy lncRNA at the LNA target site and promoted the release of snoGRUMPY (figs. S12 and S22A), which accumulated inside the nucleolus (fig. S22B), resulting

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**Fig. 6. Overexpression of grumpy promotes premature differentiation into stumpy forms in vivo and prolongs mouse survival.**

(A) Parasitemia in mice infected with the parental cell line (black line) or with a grumpy-overexpressing cell line. grumpy overexpression was either not induced or induced by adding doxycycline to drinking water either at day 0 (purple curve) or day 4 (yellow curve) of infection. Results are shown as the means (SEM, n = 4). Dunnett’s multiple comparisons test was used for statistical analysis using the parental cell line as the control (adjusted P values: **P < 0.01; ***P < 0.001). (B) Mouse survival rates according to the type of infection described in (A). Log-rank (Mantel-Cox) test for comparisons of Kaplan-Meier survival curves indicates a significant increase in the survival rates in mice infected with grumpy-overexpressing cell line parasites compared to mice infected with the parental cell line. ϕP = 0.0067 and ϕP = 0.0177. (C) Fraction of mice with at least 20% of parasites GFP::PAD1 positive (arbitrary threshold that we defined to compare the initiation of stumpy formation between the different conditions) in the blood as a function of parasitemia. Log-rank (Mantel-Cox) test for comparisons of Kaplan-Meier curves indicates significant premature parasite differentiation into stumpy forms in mice infected with grumpy-overexpressing cell line parasites compared to mice infected with the parental cell line. ϕP = 0.0067 and ϕP = 0.0177.
in a strong phenotype with premature parasite differentiation into stumpy forms (fig. S22, C and D). Thus, while our LNA gapmeR strategy failed as a loss-of-function strategy, it unexpectedly resulted in a gain of function of snoGRUMPY, confirming once again the phenotype of snoGRUMPY overexpression. This analysis confirmed that snoGRUMPY is a key player that causes premature parasite differentiation into stumpy forms upon grumpy lncRNA overexpression.

snoGRUMPY belongs to the C/D box snoRNA family, which guides 2′-O-methylation (Nm) in rRNA (30). To test whether snoGRUMPY plays a role in rRNA methylation, we used RiboMeth-seq (ribose methylations sequencing), which we previously showed that it can detect changes in Nm modification during the developmental cycle of the parasite (8, 37). Unexpectedly, analysis of the methylation profile of rRNA upon overexpression of grumpy lncRNA using RiboMeth-seq revealed no changes in the methylation site guided by snoGRUMPY (Am622). No changes were observed during the slender to stumpy transition or during overexpression of grumpy lncRNA (table S8). We conclude that the premature parasite differentiation phenotype induced by grumpy is not caused by increased methylation of rRNA mediated by snoGRUMPY.

snoRNAs have also recently been shown to also act on mRNA and regulate its translation efficacy (TE) in other eukaryotes (38). More recently, small antisense RNAs have been shown to either inhibit or enhance the translation of target mRNA in T. brucei (8). Next, we performed an in vivo cross-linking experiment to map the interaction between snoGRUMPY and its mRNA targets (8). To obtain enough material for analysis, this experiment was performed in procyclic forms. We found that snoGRUMPY interacts with 36 mRNAs and 22 other ncRNAs (including rRNA, tRNA, and others snoRNAs) (table S9). The two most remarkable mRNA targets are two hypothetical proteins: Tb927.8.2860 (HYP5) and Tb927.10.12080 (Fig. 8A, fig. S23, and table S9). HYP5 has previously been identified to be involved in driving stumpy formation (5). HYP5 has probable functions in ubiquitination and could be involved in protein degradation in slender forms. Tb927.10.12080 is the gene located just immediately upstream of grumpy lncRNA locus, but its function is unknown. When we overexpressed the coding sequence of Tb927.10.12080 alone or together with grumpy lncRNA, parasites showed a strong premature differentiation to stumpy forms (Fig. 8B and fig. S24). Parasite growth almost completely stops after 24 hours, and almost all parasites expressed GFP::PAD1 after 48 hours of Tb927.10.12080
Fig. 8. snoGRUMPY interacts with stumpy-related transcripts and regulates their expression. (A) Top 10 mRNA targets of snoGRUMPY identified using in vivo psoralen ultraviolet (UV) cross-linking. Results are shown as the total number of reads sequenced that match the specific target gene. (B to F) Tb927.10.12080 is an mRNA target of snoGRUMPY. (B) Growth curve of parasite overexpressing Tb927.10.12080 gene alone (light blue) or together with grumpy (dark blue). Results are shown as means (SEM, n = 3), statistical test: two-way ANOVA (Dunnett’s multiple comparisons test), adjusted P values: *P < 0.05; ****P < 0.0001. WT, wild-type. (C) Percentage of GFP::PAD1-positive parasites (stumpy forms) measured by FACS in parasite overexpressing Tb927.10.12080 gene alone (dark green) or together with grumpy (light green). Statistical test: two-way ANOVA (Tukey’s multiple comparisons test), adjusted P value: ****P < 0.0001. (D) RNA-FISH showing the localization of Tb927.10.12080 mRNA (green) and grumpy lncRNA (red), in parasite overexpressing Tb927.10.12080 together with grumpy. The white arrow shows the colocalization of signal between Tb927.10.12080 mRNA and grumpy lncRNA. (E) Change in Tb927.10.12080 protein expression in grumpy-overexpressing cell line is analyzed by Western blot and normalized by the expression of housekeeping protein HSP83 (in procyclic forms). (F) Change in HYP5 (Tb927.8.2860) protein expression in grumpy-overexpressing cell line is analyzed by Western blot and normalized by the expression of housekeeping protein HSP83 (in procyclic forms). Statistical test for (E) and (F): two-tailed paired t test (*P < 0.05; **P < 0.01).
overexpression (Fig. 8C). These data show that the two most remarkable targets of snoGRUMPY, HYP5 and Tb927.10.12080 mRNAs, are involved in stumpy formation in T. brucei.

RNA-FISH in parasites overexpressing both Tb927.10.12080 and grumpy from a single construct shows that grumpy partially colocalizes with Tb927.10.12080 mRNA (Fig. 8D, white arrow). This result not only confirms that grumpy IncRNA is independent from Tb927.10.12080 mRNA but also shows that these two transcripts can be found associated inside the nucleus, validating our previous cross-linking experiment. Together, these results suggest that snoGRUMPY can regulate the metabolism of Tb927.10.12080 mRNA. We tested this hypothesis by overexpressing grumpy and following Tb927.10.12080 and HYP5 protein levels. Western blotting showed that both proteins are up-regulated (Fig. 8, E and F), indicating that snoGRUMPY interacts with differentiation-related transcripts to promote their stability or translation, ultimately leading to stumpy formation.

DISCUSSION
Of the 1428 IncRNA genes that we have identified in T. brucei, 649 have been predicted, via an RNAi screen, to play a role in parasite fitness, including 399 IncRNA genes that appear to be involved in cell differentiation (Fig. 1C). Among them, we identified grumpy, an IncRNA located just upstream of RBP7 genes, a family involved in driving stumpy formation in T. brucei. We found that grumpy is the host gene of snoGRUMPY, a C/D box snoRNA. Overexpression of grumpy IncRNA and snoGRUMPY leads to premature parasite differentiation into stumpy forms and a reduction in parasite virulence in vivo.

Overexpression of snoGRUMPY recapitulates the premature parasite differentiation phenotype observed with grumpy overexpression. These results indicate that splicing and maturation of snoGRUMPY from grumpy IncRNA are likely to be the key molecular mechanism that governs parasite differentiation into stumpy forms. The antisense LNA gapmRNs, which lead to the strongest reduction in expression, were selected for the most intense GFP expression, which occurs in the nucleus in response to quorum-sensing signal. This reporter cell line was used as the genetic background to overexpress the grumpy IncRNA, snoGRUMPY, or grumpy RNAi construct. It was cultivated in HMI-11 at 37°C in 5% CO2 with G418 (2.5 μg/ml), hygromycin B (5 μg/ml), blasticidin S (5 μg/ml), and plethomycin (2.5 μg/ml). Procyclidine forms of T. brucei strain 29-13, which carries integrated genes for T7 polymerase and the tetracycline repressor, were grown in SDM-79 (45) supplemented with 10% fetal bovine serum in the presence of hygromycin B (50 μg/ml) and G418 (15 μg/ml). Transfected cells were cloned by serial dilution to obtain a clonal population. The cell lines used in this study to silence hnRNP F/H (26) and polyuridylic tract binding protein 1 (PTB1) were as previously described (27).

MATERIALS AND METHODS
Ethics statement
Male C57BL/6J (6- to 8-week-old) mice were purchased from Charles River Laboratories (Lyon, France). All animal care and experimental procedures were performed according to EU regulations (Directive 2010/63/EU9) and approved by the Animal Ethics Committee of Instituto de Medicina Molecular João Lobo Antunes (AWB2016_19FG_RNA).

T. brucei cell culture
A stumpy reporter cell line with a GFP:PAD1UTR construct (32) integrated into the tubulin locus was generated in T. brucei Antat1.1e (90:13) strain (44). The stumpy reporter cell line was selected for the most intense GFP expression, which occurs in the nucleus in response to quorum-sensing signal. This reporter cell line was used as the genetic background to overexpress the grumpy IncRNA, snoGRUMPY, or grumpy RNAi construct. It was cultivated in HMI-11 at 37°C in 5% CO2 with G418 (2.5 μg/ml), hygromycin B (5 μg/ml), blasticidin S (5 μg/ml), and plethomycin (2.5 μg/ml). Procyclidine forms of T. brucei strain 29-13, which carries integrated genes for T7 polymerase and the tetracycline repressor, were grown in SDM-79 (45) supplemented with 10% fetal bovine serum in the presence of hygromycin B (50 μg/ml) and G418 (15 μg/ml). Transfected cells were cloned by serial dilution to obtain a clonal population. The cell lines used in this study to silence hnRNP F/H (26) and polyuridylic tract binding protein 1 (PTB1) were as previously described (27).
RNA sequencing

*T. brucei* BSF and procyclic form parasites (strain Lister 427, antigenic type MiTat 1.2, clone 221a), from the PLIS cell line (46), were used to generate strand-specific libraries following the manufacturer’s instructions (Encore Complete RNA-Seq Library Systems, NuGen) for Illumina next-generation paired-end sequencing. RNA-seq was performed by the Genomics Core Facility, EMBL Heidelberg. The RNA-seq data from this study have been submitted to the European Nucleotide Archive—PRJEB38238.

Reconstruction of *T. brucei* transcriptome

In *T. brucei*, all mature mRNAs are trans-spliced and polyadenylated, which means that all mRNA transcripts start with a conserved spliced-leader (SL) sequence and finish with poly(A) tail sequence (47). We hypothesized that any new *T. brucei* transcripts, including ncRNAs, will bear these features. RNA-seq reads were assessed for quality using FastQC. To improve genome mapping, RNA-seq read size was increased, if possible, by merging the paired-end reads using PEAR software (Paired-End reAd mergeR; https://cme.h-its.org/exelixis/web/software/pear/). Merged and forward unmerged reads containing a minimum of 8 bp matching the SL sequence on their 5’ ends were extracted for 5’ splice-acceptor site detection, and the SL sequence was removed from the read. Reads containing stretches of at least nine A’s in the merged reads or nine T’s in the unmerged reverse reads were extracted for poly(A) site identification, and poly(A) tails were removed from the read.

SL and poly(A) reads were aligned to *T. brucei* genome (https://tritrypdb.org/tritrypdb/; genome annotation: version v5.1) using LAST (version 959) alignment tools (http://last.brc.jp/) (48). 5’ splice-acceptor sites were determined by the first position of all SL-containing reads mapping uniquely to the genome. Poly(A) sites were determined by the last position of all uniquely mapped poly(A)-containing reads. SL acceptor or poly(A) sites were considered for further analysis if a splice-acceptor or poly(A) site is supported by at least five reads. Putative *T. brucei* genes were defined by all genomic regions separated by at least one 5’ acceptor site and one 3’ poly(A) site occurring before the next downstream 5’ site. For each gene region, the longest transcript isoform was defined by the association of the most upstream SL acceptor site and the most downstream poly(A) site. In contrary, the major isoform of the *T. brucei* gene transcript was defined by the gene region bordered by the major SL acceptor and poly(A) sites (i.e., the ones with most reads aligned). This analysis identified 8831 genes in the *T. brucei* genome.

Identification of Ksplice putative new noncoding genes

A stringent selection pipeline was developed to systematically identify *T. brucei* ncRNAs. This pipeline aims to discard housekeeping (tRNAs, snRNAs, and snoRNAs) *T. brucei* ncRNAs and transcripts with protein-coding potential. First, only transcripts that do not overlap that annotated protein-coding and ncRNA genes from TriTrypdata (https://tritrypdb.org/tritrypdb/; genome annotation: version v5.1) were retained. Second, *T. brucei* transcripts with protein-coding potential were excluded. Protein-coding potential was determined using three different approaches. (i) The protein-coding potential for each transcript was calculated using coding potential calculator score (CPC2) (49). (ii) The association with *T. brucei* ribosomes and TE of each transcript was measured using the published ribosome profiling data from *T. brucei* (50) and reanalyzing it with our Ksplice gene annotation. (iii) The noncoding potential of each transcript was confirmed using proteomic data from three different life cycle stages of *T. brucei* (51). Each transcript with noncoding potential defined in parts (i) and (ii) and not encoding any peptides or encoding solely nonunique peptides in part (iii) was classified as a Ksplice ncRNA gene.

Coding potential calculator

The longest isoform of each Ksplice gene was used for CPC2 analysis. CPC2 (49) discriminates coding and noncoding DNA sequences based on four intrinsic features: Fickett TETSCODE score, open reading frame (ORF) length, ORF integrity, and isolectric point (pI). The Fickett TESTCODE score was calculated from the weighted nucleotide frequency of the full-length transcript, whereas the ORF length, ORF integrity, and pI were calculated from the longest putative ORF identified in each gene. A CPC2 score below 0.5 defined a transcript as a noncoding gene, whereas a CPC2 score of ≥0.5 was used to classify a transcript as encoded by a protein-coding gene.

Ribosome profiling

*T. brucei* ribosome profiling data (50) were reanalyzed using our merged genome annotation that consisted of the annotated protein-coding genes from TriTrypDB and our newly annotated Ksplice noncoding genes (major isoforms). Quantification and statistical analysis were performed as described by Vasquez et al. (50). A *T. brucei* transcript was defined as engaged in productive interaction with ribosomes if its TE score was ≥1 [TE = RPKM (reads per kilobase million) of ribosome profiling/RPKM of RNA-seq]. Conversely, a *T. brucei* transcript was defined to be not interacting with ribosomes if its TE score was ≤0.2857, meaning its transcript levels (RNA-seq data in RPKM) were 3.5× higher than its level of association with *T. brucei* ribosomes. A *T. brucei* transcript with a TE score in between (0.2857 < TE < 1) was defined to have low or few interactions with *T. brucei* ribosome. In addition, as in the work of Vasquez et al. (50), we investigated the 5′ end periodicity of mapped reads of both coding and putative noncoding genes. For fig. S6, for each gene, the number of reads mapping to each frame of translation (represented as +0, +1, and +2) was calculated, and the frame with the highest number of mapped reads was determined. A P value indicating the likelihood of periodicity was calculated by a binomial test on the frame with the highest number of mapped reads under the null hypothesis that this number should be equal to one-third of all reads mapped to that gene.

Proteomics

The MS proteomic data from Dejung et al. (51) were analyzed following the author’s methodology with some modifications using MaxQuant version 1.6.0.1 (52) and searching against our Ksplice protein database. Our Ksplice protein database is composed of three sets of proteins: protein-coding genes from TriTrypDB (version 33, 10,019 entries, excluding protein-coding genes with an internal codon stop), putative proteins that originate from the Ksplice new gene sequences (2003 Ksplice new genes + 72 Ksplice Kolev ncRNAs), and putative proteins originating from intergenic region sequences of *T. brucei* genome. Intergenic region sequences were selected to have, on average, the same size and number of sequences as Ksplice new genes. All putative protein sequences (enclosed by a start and stop codon) with a minimum of seven amino acids originating from Ksplice new genes or intergenic regions of *T. brucei* genome were extracted in order of the DNA sequence and from the...
three possible translation frames (excluding sequences without a start codon or/and containing an ambiguous base). A total of 14,261 proteins were extracted from Ksplice new genes and 28,750 proteins from the selected intergenic region of *T. brucei* genome.

**Full-length sequencing**

To investigate the presence of full-length transcripts in our RNA-seq dataset, read pairs containing the SL sequence on the forward read and a poly(A) tail on the reverse read were extracted and mapped to the *T. brucei* genome as described above (in the “Identification of Ksplice putative new noncoding genes” section). For all concordant alignments (both paired reads aligned), the boundaries of the transcripts were determined by the mapping positions of the two reads. Paired-end reads provide the accurate boundaries of *T. brucei* transcripts as reads are sequenced from the same RNA molecule.

**Nanopore direct RNA-seq**

Direct RNA-seq was performed as described by Bil ska et al. (53). Briefly, RNA libraries were prepared from 5 μg of total RNA isolated from the BSF and procyclic form, respectively, mixed with 50 to 200 ng of oligo(dt)25-enriched mRNA from *Saccharomyces cerevisiae* yeast with a Direct RNA Sequencing Kit (Oxford Nanopore Technologies, catalog no. SQK-RNA002) according to the manufacturer’s instructions. Sequencing was performed with RR9.4.1 RevD flow cells on a MinION device using MinKNOW software (Oxford Nanopore Technologies). The Nanopore direct RNA-seq data from this study have been submitted to the European Nucleotide Archive–PRJEB48655.

**Direct RNA-seq bioinformatic analysis**

Raw sequencing reads were base-called using Guppy v.4.4.1 (Oxford Nanopore Technologies). To find entirely sequenced reads (i.e., complete from 3′ to 5′ end), delimiting signals from both termini were assessed. The 3′ end was analyzed in the following manner: Obtained reads were mapped to custom references containing lincRNA candidates (provided by F.G.) using minimap2 v.2.17, with parameters -k 14 -ax map-ont --secondary=no. Supplemental alignments and reads mapping to the reverse strand were filtered out with SAMtools v.1.9 (SAMTools view -b -F 2320). The poly(A) tails of reads were assessed using the poly(A) estimation module of the program Nanopolish v.0.13.2 (54). Only reads fulfilling quality criteria (tagged as “PASS”) were taken into consideration in further analyses. 5′ ends were examined using two approaches. First, obtained reads were mapped to the reference genome (TriTrypDB:54_TBrucDetREU927_Genome.fasta) obtained from TriTryp database using minimap2 v.2.17 (55) with the following parameters: -k 14 -ax splice -uf --secondary=no. Second, mapped reads were merged and converted to BLAST databases with makeblastdb command using default settings for nucleotide sequences (for each sample separately). Subsequently, the BLASTn v.2.11.0 (56) algorithm was run with SL-RNA sequence used as a query against built databases with the following parameters: -evalue 1000 -num_alignments 1000000 -dust no -max_hsps 1 -word_size 4 -num_threads 10 -outfmt 6. The output data were filtered on the basis of the following criteria: (i) reads containing a >10-bp region of homology with SL-RNA query, (ii) reads aligned with an e value of <100, (iii) alignment starts at the termini of read (subject) sequence (value of blast tabular output columns qstart/qend == 1), and (iv) alignment starts at the termini of SL-RNA (query) sequence (value of blast tabular output columns qstart/qend == 1). Reads satisfying all of the above criteria were taken into account for further analysis. Second, reads mapping to the reference genome were filtered with cutadapt v.2.10 (57) with the following options: --discard-untrimmed -e 0.2 --action "lowercase,” based on the presence of the potential SL RNA sequence. Last, unique read IDs obtained with both 5′ end analysis approaches and 3′ analysis were combined and used to filter the genomic alignment files. Filtering was performed in R 4.0.2 using Rsamtools 2.6.0 from Bioconductor. The results were saved in BAM and FASTA formats. Gene body coverages were plotted using RseQC genebodycoverage.py script (58). The obtained sequence alignments were saved in BAM format. Read sequences were extracted in FASTA format as two variants: entirely (encompassing total base-called length) and with soft clip masking [soft clipping 5′ SL and miscalled bases from 3′ poly(A) tail were removed on the basis of cigar scores] using biostar84452.jar from jvarkit toolset. Two corresponding FASTA datasets were coerced, and 5′ SLs were identified. The coordinates of 3′ termini of transcripts were pointed out on the basis of sequence alignment and Nanopolish poly(A) function output.

**RNA-FISH**

Between 2.5 × 10^5 and 1 × 10^6 cells were harvested by centrifugation (1800g for 10 min), washed with 1× phosphate-buffered saline (PBS) or trypanosome dilution buffer (TDB), and resuspended in between 500 μl and 1 ml of fixation buffer [% 7.3% formaldehyde diluted in ribonuclease (RNase)–free PBS] for 10 min at room temperature. Fixed cells were washed with between 500 μl and 1 ml of RNase-free PBS and resuspended with 150 μl of RNase-free PBS. Cells were then settled on precoated polylysine culture dishes (35-mm glass bottom, MatTek) for at least 20 min. PBS was removed, and cells were permeabilized with 1 ml of 70% ethanol (EtOH; in RNase-free water) for at least 1 hour at +2° to +8°C. EtOH (70%) is discarded, and cells were washed with 200 μl of wash buffer A [10% (v/v) formamide in 1× wash buffer A; Biosearch Technologies, catalog no. SMF-WA1-60]. Cells were incubated with 100 μl of hybridization buffer containing 1.25 μM RNA-FISH probes in the dark at 37°C overnight (~16 hours). Cells were washed with 200 μl of wash buffer A and incubated with 200 μl of wash buffer A in the dark at 37°C for 30 min. Cells were stained with a solution of 4′,6-diamidino-2-phenylindole (DAPI; 1 μg/ml) (in wash buffer A) in the dark at 37°C for 30 min. Cells were washed with 200 μl of wash buffer B (Biosearch Technologies, catalog no. SMF-WB1-20) and incubated with it at room temperature for 2 to 5 min. VECTA-SHIELD (100 μl) was added to the dishes before analysis with the Zeiss cell observer widefield microscope.

RNA-FISH probes were designed using the online tools provided by LGC Biosearch Technologies (Stellars Probe Designer, www.biosearchtech.com/support/tools/design-software/stellars-probe-designer). A total of 17 probes were designed for grumpy lncRNA and 30 to 43 probes for Ksplice lncRNA223a, lncRNA1077a, lncRNA1735a, and lncRNA5090a.

**Differential expression of Ksplice new genes between BSF and procyclic form**

Differential expression analysis for Ksplice new genes between BSF and procyclic form was performed using our merged annotation of *T. brucei* genome (major isoform of Ksplice new genes + TriTryp...
protein-coding genes) and the DEseq2 package. To that end, we used our previously published transcriptomic data (59) containing 13 RNA interference target sequencing (RNA-seq) samples replicated for both BSF and procyclic form.

**RIT-seq analysis of Ksplice new genes**
The RNA interference target sequencing (RIT-seq) data from Alsford \textit{et al.} (23) were reanalyzed by aligning the sequence reads against our merged annotation of \textit{T. brucei} genome (major isoform of Ksplice new genes + TriTryp protein-coding genes). Quantification and statistical analysis were performed as described by Alsford \textit{et al.} (23).

**Transcript quantification and cRT-PCR**
Transcript quantification was performed by RT-qPCR, as described by Aresta-Branco \textit{et al.} (60), except that random hexamer primers were used to generate cDNA. The cRT-PCR protocol was performed essentially as described in (61). Briefly, parasites were harvested by centrifugation at 677 g for 10 min at 4°C and immediately resuspended in TRIzol (Life Technologies). Total RNA was isolated following the manufacturer’s instructions, and RNA was quantified in a NanoDrop 2000 (Thermo Fisher Scientific). The ideal RNA concentration to perform the cRT-PCR protocol is 0.5 to 1 mg/µl. The RNA cap and poly(A) tail were removed by oligonucleotide-directed RNase H cleavage using SL and oligo(dT) primers. After RNase H treatment, RNA was extracted with phenol/chloroform approach and precipitated using EtOH precipitation protocol. RNase H–treated RNA (3 to 5 µg) was circularized using T4 RNA ligase 1 [ssRNA Ligase, New England Biolabs (NEB)]. RNA was extracted using phenol/chloroform, and EtOH was precipitated. RNA was reverse-transcribed using the gene-specific primer R1 (100 nt from the 5′ end of the transcript or the RNase H cleavage site) and RT buffer and 5 mM magnesium from the SuperScript II kit (Life Technologies). The resulting cDNA molecules contained the juxtaposed 5′ and 3′ ends of circular RNA. PCR was performed on the produced cDNA using gene-specific primers R2 and forward F1. R2 primer is in a “nested” position relative to the R1 primer and contributes to the specificity of the PCR amplicon. PCR#1 product was purified using a MinElute PCR purification kit (QIAGEN), and a second round of PCR amplification was performed with gene-specific primers R2 and forward F2. The F2 primer is in a nested position compared to the F1 primer and contributes to PCR amplicon specificity. PCR#2 product was ligated to pGEM-T easy vector or TOPO vector following the manufacturer’s instructions (Promega). After transformation in bacteria and plasmid amplification, the subcloned PCR#2 fragments were amplified and sequenced using T7 and SP6 primers.

**Quantitative RT-PCR**
RNA was prepared with TRIzol reagent (Invitrogen) according to manufacturer’s instructions, and cDNA was synthesized with random primers and SuperScript II reverse transcriptase. qPCR was performed with AmpIiTaq Gold DNA Polymerase (Power SYBR Green Master Mix, Applied Biosystems) and gene-specific primers as follows: control of differentiation (Tb927.10.12970), CAGCTTCCTCAATCTC-CAG (forward) and GGCACAGTGGATAGCTTG (reverse); and GGAAAAGACCCAGAAACAGAAAG (reverse); \textit{grumpy} I (5′-CGAGGGTTTGGAAATGGTTTG -3′) and reverse (5′-TTATCGCACTGAAGCAGCTCC-3′) primers and inserted into pDEX577 (phleo) between the Hind III and Bam HI sites of the plasmid. pDEX577 vectors are highly modular expression vectors for inducible expression of transgenes, integrating in the mini-chromosome repeats, which was designed and constructed by Kelly \textit{et al.} (62). Moreover, two T7 terminator sequences were inserted between the Bam HI and Kpn I sites of the plasmid just downstream to the \textit{grumpy} lncRNA construct. For the overexpression of \textit{grumpy}, we used the same vector and cloning procedures as for overexpression of \textit{grumpy}, except that snoGRUMPY was amplified from \textit{T. brucei} Antat 1.1E genomic DNA with forward (5′-CCGTTTCGATCCTGTGTG-3′) and reverse (5′-TTATCGCACTGAAGCAGCTCC-3′) primers. For the RNAi construct of grumpy, lncRNA was designed as a long stem-loop double-stranded RNA and by following the experimental procedures described by Atayde \textit{et al.} (63). Briefly, a fragment of grumpy lncRNA sequence was amplified from \textit{T. brucei} Antat 1.1E genomic DNA with a forward primer (5′-CAAGGGCAGGAGAGGGGATGTGTAAC-3′) containing either Hind III or Bam HI restriction site in its 5′ end and a reverse primer (5′-GCGGGTTTGGAAATGGTTTG-3′) containing Eco RI restriction site followed by a 50-nt randomized sequence at its 5′ end. Two PCR amplicons were therefore obtained: one containing...
Inducible expression of RNAi and overexpression constructs

Cells were diluted at $5 \times 10^4$ parasites/ml and induced with tetracycline (1 µg/ml) for 3 days for grumpy RNAi and Tb927.10.12080 overexpression and for 6 days for grumpy overexpression cell lines. Cells were counted every day, live/dead cells were assessed by propidium iodide staining, GFP::PAD1 staining was assessed by measuring the GFP::PAD1 expression using flow cytometry (Accuri C6). BSFs were collected from culture, spun down, resuspended in differentiation trypanosome medium with 6 mM cis-aconitate at $1 \times 10^6$ parasites/ml, and incubated at 27°C. parasite differentiation was assessed by flow cytometry using anti-T. brucei procyclin antibody (0.5 mg; Cedarlane, clone TBRP1/247, CLP001AP) conjugated with Alexa Fluor 647 (protein labeling kit, Molecular Probes) (1:500 dilution in TDB).

PAD1 staining
BSF parasites ($5 \times 10^5$) were harvested by centrifugation (1800g for 10 min), washed with $1 \times$ PBS, and resuspended in 500 µl of fixation buffer (4% paraformaldehyde diluted in 1× PBS) for 10 min at room temperature. Fixed cells were washed with 500 µl of $1 \times$ PBS and resuspended with 100 µl of $1 \times$ PBS. Cells were then settled on precoated polyllysine culture dishes (35-mm glass bottom, MatTek) for at least 20 min. PBS was removed, and cells were permeabilized with 100 µl of 0.1% Triton X-100 in PBS for 2 min at room temperature. Permeabilized cells were washed five times with 200 µl of PBS and blocked with 2% bovine serum albumin (BSA) in PBS for 45 min at 37°C in a humidity chamber. Cells were incubated with 100 µl of the primary antibody anti-PAD1 (1:1000 in 2% BSA in PBS; antibody provided by K. Matthews) overnight at 4°C in a humidity chamber. Cells were washed five times with 200 µl of PBS and incubated with 100 µl of the secondary antibody anti-rabbit
Infections and sample collection

Four-week-old male c57BL/6 mice (Charles River Laboratories, France) were inoculated intraperitoneally with 2000 parasites. Mice were infected with either Antat1.1 90:13 GFP::PAD1 grumpy overexpression cell line or Antat1.1 90:13 GFP::PAD1 grumpy overexpression parasites were separated in three different cages: One cage of four mice received only water, one cage of four mice received water with doxycycline hyclate (1 mg/ml; Sigma-Aldrich) at day 4 after infection, and one cage of three mice received water with doxycycline hyclate (1 mg/ml) at the day of infection. Parasitemia was monitored by tail-vein bleeds every other day and counted using a hemocytometer with 1:150 blood dilution in TDB. The percentage of stumpy forms in the mouse blood was assessed by measuring GFP::PAD1 expression in blood-diluted sample using an Accuri C6 flow cytometer. Mouse survival was monitored every other day until 100 days after infections. Mice were euthanized at the first signs of severe disease distress, with all efforts to minimize animal suffering. Humane end points were used.

grumpy LNA treatment

Antisense LNA gapmeRs targeting grumpy lncRNA (LNA sequence: AGGATGCCGACGGGA; from QIAGEN) were resuspended at 50 μM with tris-EDTA buffer. The stumpy reporter cell line cultures (GFP::PAD1) were diluted at 5 × 10^6 parasites/ml and grown for 3 days with different concentrations of LNA gapmeRs (0 to 50 nM; diluted into culture medium). Cells were counted every day, live/dead cells were assessed by propidium iodide staining, GFP::PAD1-positive cells were scored, and all these parameters were quantified using an Accuri C6 flow cytometer. At day 1 after LNA exposure, RNA samples were collected by centrifugation of an equivalent number of cells and addition of TRIzol reagent (Invitrogen) to the cell pellets.

In vitro RNA-protein pulldown

Stumpy cells (350 × 10^6; GFP::PAD1-positive cells) were used for each condition of the RNA-protein pulldown. Cells were washed once with PBS and then kept on ice as a dry cell pellet (350 × 10^6 stumpy cells in ~30 μl of dry pellet). The nuclear proteins were then extracted from the dry cell pellets with the NE-PER kit according to the manufacturer’s instructions (Thermo Fisher Scientific). After phenol:chloroform extraction, the RNA baits were biotinylated using the Pierce RNA 3' End Desthiobiotinyllation Kit and following the manufacturer’s instructions (Thermo Fisher Scientific). RNA-protein pulldown assays were performed with the Pierce Magnetic RNA-Protein Pull-Down Kit according to the manufacturer’s instructions (Thermo Fisher Scientific). Nuclear proteins (~75 μg) were used under each condition of RNA-protein pulldown. The primers used to synthesize the RNA baits were the following:

grumpy lncRNA, taatagcactcactaggggCAAAAGGCACAGAT- TATAGGTTC (forward) and GATGCGCTCAACAGCAGAAG (reverse); lncRNA-5090a (lncRNA control), taatagcactcactagg- gagAAACCCCTCCACTCCTAC (forward) and TTGCCGTGTTTTC- GTGC (reverse).

Our grumpy RNAi vector was linearized using Bam HI restriction enzyme and used as a template for the RNA in vitro production of the grumpy RNAi bait. Note that we cannot exclude that the large proportion of mitochondria proteins found in our in vitro RNA-protein pulldown assay (Fig. 4A) could come from mitochondria contamination. The first lysis step of the NE-PER kit could be too mild to completely lyse mitochondria. Thus, mitochondria proteins will be extracted during the second lysis step together with the nuclear proteins. Under this condition, mitochondria and nuclear proteins will therefore be used together in our pulldown experiment.

MS sample preparation and measurement

Samples were loaded onto a Novex NuPage 4 to 12% precast gel (Thermo Fisher Scientific) and run in 1× Mops buffer for 10 min at 180 V. The Coomassie G250 (Roth)—stained gel lanes were minced and destained with 50% EtOH/25 mM ammonium bicarbonate (ABC). Subsequently, gel pieces were dehydrated with 100% acetonitrile (ACN) and dried for 5 min in a concentrator (Eppendorf). Samples were incubated with reduction buffer [10 mM dithiothreitol (DTT; Sigma-Aldrich)/50 mM ABC] for 30 min at 56°C and further alkylated for 30 min in the dark with iodoacetamide [50 mM indole-3-acetic acid (Sigma-Aldrich)/50 mM ABC]. Gel pieces were completely dehydrated with ACN and covered in MS-grade trypsin (Serva) solution (1 μg of trypsin per sample). Proteins were digested overnight at 37°C, and peptides were extracted twice by incubation with extraction buffer (3% trifluoroacetic acid and 30% ACN) for 15 min. The gel pieces were dehydrated with 100% ACN, and the extracted volume was reduced to approximately 150 μl in a concentrator (Eppendorf). Extracted peptides were desalted in StageTips using two layers of C18 material (Empore). Eluted peptides were injected via an autosampler into a uHPLC (EASY-nLC 1200, Thermo Fisher Scientific) and loaded on a 55-cm capillary (75-m inner diameter; New Objective) packed in-house with Reprosil C18-AQ 1.9-μm resin (Dr. Maisch GmbH) for reversed-phase chromatography. The EASY-nLC 1200 HPLC system was directly mounted to an Exporis 480 mass spectrometer (Thermo Fisher Scientific). Peptides were eluted from the column with a 73-min optimized gradient from 2 to 40% ACN with 0.1% formic acid at a flow rate of 250 nl/min. Chromatography was performed with a column oven setup operating at 55°C (Sonation). The ion transfer tube temperature was set to 250°C. Spray voltage ranged from 2.2 to 2.4 kV. The mass spectrometer was operated in data-dependent acquisition mode with one MS full scan and up to 15 triggered tandem MS (MS/MS) scans using higher energy collisional dissociation (HCD) fragmentation. MS full scans were obtained in the Orbitrap at 60,000 resolution, while MS/MS scan resolution was set to 15,000 resolution. Charge states 2 to 6 were included from MS/MS selection, and peptide match was preferred.

MS data analysis

Raw files were processed with MaxQuant (version 1.5.2.8) (52) and searched against T. brucei TREU927 database (version 8.1; 11,567 entries) downloaded from tritrypdb.org using the Andromeda search engine (31). All results were visualized using MaxQuant and Proteome Discoverer (version 1.4). In all cases, database search parameters were fixed as follows: trypsin cleavage, two missed cleavages were allowed, variable modifications were carbamidomethylation of Cys, oxidation of Met, and甲基化 of Lys and Arg. Raw files from technical replicates were averaged using the MaxQuant averaging function. For protein quantification, MS1 intensities of peptides with at least one unique sequence were used. Peptides were assigned to proteins with at least two unique peptides. Peptide assignments were filtered for a false discovery rate <0.01. Protein expression levels were calculated using the peak intensity of at least six peptides per protein.

The MS experiment was repeated twice. The experiments were statistically analyzed by Mann-Whitney U test. The error bars represent the mean ± standard deviation.
engine. Carbamidomethylation was set as a fixed modification, while acetyl (protein N terminus) and oxidation (Met) were considered as variable modifications. Trypsin (specific) was selected as a protease. Proteins were quantified with at least two quantification counts of which at least one of them was a unique peptide. Known contaminants and reverse hits were removed. For statistical analysis, the mean was calculated and Welch’s t test was performed between the conditions. Graphical data representation was done using the R environment (version 3.6.2).

RIP–DRBD3

*T. brucei* 449 procyclic cells (65) were cultured at 27°C in SDM-79 medium supplemented with 10% fetal bovine serum (66). For the RIP assays, 2 × 10^5 cells were washed in serum-free Ham’s H9 medium, resuspended in 1 ml of 10 mM tris-HCl (pH 7.4), 0.5% Igepal, 1 mM DTT, 2 mM ribonucleoside vanadyl complexes (Invitrogen), and protease inhibitors (Roche); and lysed by passing the cell suspension thrice on ice. Lysates were centrifuged at 16,000g for 10 min at 4°C, and NaCl was added to 0.9 ml of the supernatant to a final concentration of 150 mM. The supernatant was then split into two halves and rotated for 3 hours at 4°C in the presence of normal rabbit serum (NRS; Sigma-Aldrich) or anti-DRBD3 antisem (28) coupled to 1.5 mg of Dynabeads Protein G (previously blocked overnight in IPP*-150 [10 mM tris-HCl (pH 7.4), 150 mM NaCl, and 0.5% Igepal CA-630] containing 0.1% BSA). Beads were then washed four times in IPP*-150 and twice in IPP*-150 containing 1 M urea and incubated for 30 min at 50°C with 100 mg of protease K in 10 mM tris-HCl (pH 8), 100 mM NaCl, 0.5% SDS, and 1 mM EDTA. RNA was extracted with phenol:chloroform, precipitated with EtOH, and converted to cDNA using 0.5 µg of random hexamers (Invitrogen), 0.25 mM each dNTPs (deoxyribonucleotide triphosphates), 20 U of RiboLock (Thermo Fisher Scientific), and 200 U of Maxima reverse transcriptase (Thermo Fisher Scientific) in a final volume of 20 µl. Reactions were incubated for 5 min at room temperature, then incubated for 30 min at 50°C, and finally heated at 85°C for 5 min. RT-qPCR reactions were carried out in a Bio-Rad CFX96 thermal cycler. Reactions were set up in a final volume of 10 ml containing 0.1 to 0.5 µl of cDNA, 1× SYBR Green Master Mix (Thermo Fisher Scientific) and 0.5 µM each oligodeoxynucleotide. Fold enrichments of target transcripts relative to control serum (NRS) were calculated using actin mRNA as a reference according to the formula 2^ΔΔCt, where ΔΔCt = (ΔCtNRS_t.o.i. − ΔCtNRS_untreated) − (ΔCtDRBD3_t.o.i. − ΔCtDRBD3_untreated), and t.o.i. is the transcript of interest. Experiments were carried out in three biological replicates, and the data are represented as the means ± SEM.

Western blot analysis

Whole-cell lysates (10^7 cells) were fractionated by 10% SDS–polyacrylamide gel electrophoresis, transferred to PROTRAN membranes (Whatman), and reacted with antibodies. The bound antibodies were detected with goat anti-rabbit immunoglobulin G coupled to horseradish peroxidase and were visualized by ECL (Amersham Biosciences). The dilutions used for the antibodies are PTB1 (1:10,000) (27), hnRNP F/H (1:10,000) (26), cMYC (1:10,000; 9E10, Santa Cruz), HSP83 (1:10,000) (8), and ZC3H41 (1:10,000) (67).

Fractionation on sucrose gradient

Whole-cell extracts were prepared from 1010 *T. brucei* procyclic cells in a buffer containing 150 mM KCl, 20 mM tris (pH 7.6), 10 mM MgCl2, 0.5 M DTT, and 0.1% NP-40. In addition, 1 µl of leupeptin (10 mg/ml) protease inhibitor and 1 µl of RNasin (Thermo Fisher Scientific) were added to the lysate. Lysates were fractionated on a 10 to 30% (w/v) sucrose gradient by centrifugation for 3 hours at 35,000 rpm in a Beckman SW41 rotor at 4°C. Fractions (500 µl) were collected, EtOH was precipitated, and RNA was extracted using phenol-chloroform. RNA extracted from each fraction was separated on a 6 or 10% (w/v) polyacrylamide gel containing 7 M urea gel and transferred onto nitrocellulose membranes. The RNA blots were hybridized to a ^32^P-labeled antisense RNA probe and visualized by autoradiography (8).

Northern blot analysis

Total RNA was separated on a 6 or 10% (w/v) polyacrylamide gel containing 7 M urea or 1.2% agarose/formaldehyde gel and transferred onto nitrocellulose membranes. The RNA blots were hybridized to a ^32^P-labeled antisense RNA probe. The results were analyzed by autoradiography (8, 37). Primers used to amplify DNA templates for RNA probes synthesis (using T7 polymerase) were as follows: snoGRUMPY/7SL_F, 5′-GACGAATTCGAGATGATGGTTT3′; snoGRUMPY/7SL_R, 5′-TATAAGCTACTACATGAAATGGAGAGAGGCAATTCAGAAT3′; snoGRUMPY/27SL_F, 5′-TTGCCTGAAATTCGCTGACGAGTCTGG3′; snoGRUMPY/27SL_R, 5′-TATAAGCTACTACATGAAATGGAGAGGCAATTCAGAAT3′; snoGRUMPY/27SL_F, 5′-TATAAGCTACCTAAGGAGAAGATGATGGTTT3′; snoGRUMPY/27SL_R, 5′-TATAAGCTACTACATGAAATGGAGAGGCAATTCAGAAT3′.

In situ hybridization in procyclic cells

Procyclic *T. brucei* cells were fixed on circular coverslips using 1.6% formaldehyde and permeabilized using Triton X-100/Tween 20 (1:0.1%) in 1× PBS. In situ hybridization with specific CY3-labeled antisense RNA probes was performed as described (8). The nucleolus was visualized using anti-NHP2 antibody (0.7:1000 dilution) (68), the MTAP protein was visualized using an MTAP-YFP reporter (69), and the nucleus was stained with DAPI. Images were acquired using a Leica SP8 confocal microscope equipped with a white light laser and gated. Cells were oversampled both in the lateral and axial axis. The images were captured using an X100 HC Plan-Apo 1.4 numerical aperture objective at 512 by 512 pixels with Z slices taken every 200 nm. Excitation wavelengths used in this study were 405 nm for DAPI, 551 nm for detecting secondary anti-rabbit cy3-conjugated antibody, and 640 nm for detecting cy5-labeled antisense RNA (8 to 12 accumulations). The images were then deconvolved using Huygens Professional software with standard parameters (SVI, Laapersveld 6, 1213 VB Hilversum, The Netherlands).

RiboMeth-seq

Total RNA (5 µg) from *T. brucei* cells was initially denatured at 90°C for 2 min in a thermocycler, and an equal volume of buffer [NaHCO3/Na2CO3 (pH 9.9)] was added. The RNA samples were subsequently incubated at 90°C for 20 min and later kept on ice (37, 70). The hydrolyzed RNA was then used for library preparation. Briefly, 800 ng of the RNA was dephosphorylated with FastAP Thermosensitive Alkaline Phosphatase (Thermo Fisher Scientific) and cleaned by Agencourt RNA clean XP beads (Beckman Coulter).
The RNA was then ligated to 3’ linker using high-concentration T4 RNA Ligase 1 (NEB) in a buffer containing dimethyl sulfoxide (DMSO), adenosine 5’-triphosphate (ATP), polyethylene glycol (PEG) 8000, and RNase inhibitor (NEB) for 1.5 hours at 22°C. The ligated RNA was purified from excess linker using Dynabeads MyOne SILANE beads (Thermo Fisher Scientific), and first-strand cDNA was prepared using the AffinityScript Reverse Transcriptase enzyme (Agilent) at 55°C for 45 min. Next, the RNA was degraded using 2 μl of 1 M NaOH, and the cDNA was cleaned using Dynabeads MyOne SILANE beads (Thermo Fisher Scientific). The cDNA was further ligated to 3’ adapter using a high-concentration T4 RNA Ligase 1 (NEB) overnight at 22°C and cleaned of excess adapter using Dynabeads MyOne SILANE beads (Thermo Fisher Scientific). The adapter ligated cDNA was PCR-enriched using NEBNext High-Fidelity (NEB) polymerase (nine PCR cycles), separated on an E-Gel EX agarose gel (Invitrogen), and size-selected for the range of 150 to 300 bp (containing ~30 to 180 nt corresponding to RNA). The amplicons were gel-purified using a NucleoSpin Gel and PCR Clean-up kit (Macherey-Nagel) and sequenced in a NextSeq system (Illumina) in paired-end mode (20 million reads for each sample). BioProject ID for the RiboMeth-seq is the PRJNA776556.

Analysis of chimeric RNA molecules

The sequencing reads generated from in vivo psoralen UV cross-linking and ligation to generate chimeric RNA were retrieved from a previous study (8). The reads from the RNA-seq as described above were preprocessed using Trim Galore v0.5.0 (www.bioinformatics.babraham.ac.uk/projects/trim_galore/). Overlapping read pairs were merged using FLASH v1.2.11 (71). BWA-MEM v0.7.17-r11882 (72) was used to align the reads to a custom-made T. brucei transcriptome database. The SAM output was converted to BAM using SAMtools. The scripts from the SPLASH computational pipeline (73) were used to identify the chimeric RNA from the BAM files. The “find_chimeras.py” script was used to detect chimeric RNA. In the case of “intramolecular” chimeric RNA, the ligated RNAs were required to be spaced at least 15 nt apart in the same transcript.

Statistical analysis

For all graphs in Figs. 3A, 3C, 4F, 6A and 8B and figs. S11, S15, S22 (C and D) and S24, the results are shown as means (SEM, n = 3), and all statistical analyses are done with two-way analysis of variance (ANOVA; Dunnett’s multiple comparisons test). For all graphs in Figs. 5 (C to G) and 7 (A, C and D) and figs. S16A, S19A and S22A, the results are shown as means (SEM, n = 3), and all statistical analyses are done with two-way ANOVA (Tukey’s multiple comparisons test). For the graph in Fig. 8C, the results are shown as means (SEM, n = 3), and all statistical analyses are done with two-way ANOVA (Tukey’s multiple comparisons test). For the graph in Fig. 5A, the results are shown as means (SEM, n = 3), and all statistical analyses were done using multiple t test. For all graphs in Figs. 4B and 8 (E and F) and fig. S18, the results are shown as means (SEM, n = 3), and all statistical analyses were done with a two-tailed paired t test. For graphs in Fig. 6 (B and C), the results are shown as means (SEM, n = 4), and statistical analyses are done with log-rank (Mantel-Cox) test.
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