

Structure of the C-terminal domain of transcription factor IIB from *Trypanosoma brucei*

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In trypanosomes, the production of mRNA relies on the synthesis of the spliced leader (SL) RNA. Expression of the SL RNA is initiated at the only known RNA polymerase II promoter in these parasites. In the pathogenic trypanosome, *Trypanosoma brucei*, transcription factor IIB (tTFIIB) is essential for SL RNA gene transcription and cell viability, but has a highly divergent primary sequence in comparison to TFIIB in well-studied eukaryotes. Here we describe the 2.3 Å resolution structure of the C-terminal domain of tTFIIB (tTFIIB_C). The tTFIIB_C structure consists of 2 closely packed helical modules followed by a C-terminal extension of 32 aa. Using the structure as a guide, alanine substitutions of basic residues in regions analogous to functionally important regions of the well-studied eukaryotic TFIIB support conservation of a general mechanism of TFIIB function in eukaryotes. Strikingly, tTFIIB_C contains additional loops and helices, and, in contrast to the highly basic DNA binding surface of human TFIIB, contains a neutral surface in the corresponding region. These attributes probably mediate trypanosome-specific interactions and have implications for the apparent bidirectional transcription by RNA polymerase II in protein-encoding gene expression in these organisms.

eukaryotic transcription | parasite | RNA polymerase II | trypanosome

Trypanosomes are flagellated protozoa, belonging to the order Kinetoplastida, that are exclusively parasitic (1). These organisms reside in insect vectors that transmit the parasites to mammals, birds, fish, and plants. *Trypanosoma brucei* and *Trypanosoma cruzi* are of particular medical concern because they cause debilitating and fatal diseases in millions of people annually in tropical regions of the world (2). Trypanosomes diverged from other eukaryotes early in evolution, and thus many biological processes that are well understood in metazoans are highly distinct in these parasites (3, 4). For example, little is known about how trypanosome RNA polymerase II (tRNAP-II) transcription is initiated. It is known that tRNAP-II transcribes most protein-encoding genes, which are arranged in tandem arrays, into polycistronic mRNAs (5). Thus far, the only tRNAP-II promoter that has been identified is the spliced leader (SL) RNA gene promoter (6). The SL RNA gene codes for the SL, which is capped and added in a *trans*-splicing reaction to the 5' end of each ORF contained in a polycistronic mRNA. The addition of an SL to every mRNA is a universal requirement in trypanosomes; therefore, understanding transcription initiation at the SL RNA gene promoter is a crucial step toward understanding the control of tRNAP-II transcription in these parasites (4, 7).

To date, components of the SL RNA transcriptional machinery have been studied by using genomics or biochemistry to identify candidate proteins followed by a combination of phenotypic analysis of protein function using either RNAi-silencing of endogenous protein in parasites or depletion/add back studies of in vitro transcription systems. These methods have identified several players in tRNAP-II-dependent SL RNA transcription,

including trypanosome TATA-box binding protein (tTBP), transcription factors IIB, -IIA, -IIH (tTFIIB, tTFIIA, tTFIIH), and tSNAPc (8–17). However, each trypanosome factor is only distantly related to its metazoan homolog and contains regions or subunits unique to trypanosomes. Indeed, a recent study of tTFIIH using single particle electron microscopy suggests that tTFIIH has a similar structure to human TFIH but contains trypanosome-specific subunits (18). Thus, the mechanism of transcription initiation in trypanosomes cannot be deduced from metazoan transcription systems. Our work presented here is the primary venture to understand the mechanics of the SL RNA transcriptional machinery using high-resolution structural analysis.

Trypanosome TFIIB is essential for SL RNA transcription in *T. brucei* (13, 14). tTFIIB associates with tTBP, tRNAP-II, and an SL RNA gene promoter fragment in nuclear extracts. In yeast and mammals, TFIIB binds specifically to TBP and DNA and recruits RNAP-II into a minimal preinitiation complex (reviewed in ref. 19). TFIIB consists of N- and C-terminal domains (20, 21). NMR analysis of the N-terminal domain from *Pyrococcus furiosus* TFIIB reveals a Zn ribbon motif and an extended B finger loop (22, 23). In the cocrystal structure of *Saccharomyces cerevisiae* TFIIB and RNAP-II, the TFIIB N-terminal domain binds RNAP-II and uses the B finger loop to finely position RNAP-II at the transcription start site (24). NMR analysis of the human TFIIB C-terminal domain reveals 2 helical repeats (25). In the human and *Pyrococcus woesei* TFIIB/TBP/DNA ternary complexes, the TFIIB C-terminal domain recognizes TBP and the promoter (26–29). The C-terminal domain of the trypanosome TFIIB (tTFIIB_C) contains 1 loosely conserved repeat module followed by a trypanosome-specific region (13, 14). Here, we report the 2.3 Å resolution structure of tTFIIB_C. The tTFIIB_C structure reveals 2, closely packed helical modules followed by a C-terminal extension of 32 aa. The trypanosome-specific region comprises the second helical module and the C-terminal extension. The overall tTFIIB_C structure is similar to other TFIIB_C structures (25–29), but contains additional loops and helices. In contrast to the highly basic DNA binding surface of human TFIIB, the structure reveals a neutral surface in the

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Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.pdb.org (PDB ID code 3H4C).

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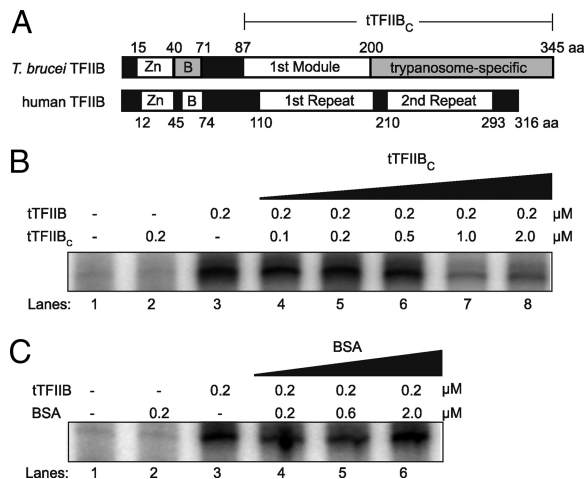


Fig. 1. Primary structure and function of tTFIIB and tTFIIB_c. (A) Schematic of *T. brucei* and human tTFIIB amino acid sequence alignment. The Zn ribbon and the first module/first repeat are well conserved between trypanosomes and humans. The B finger and the trypanosome-specific/second repeat are not well conserved (gray background). tTFIIB_c is 259 residues long, comprising the first module and the trypanosome-specific region (bracketed). (B) Activity of tTFIIB and tTFIIB_c in in vitro SL RNA transcription assays. Transcription activity in tTFIIB-depleted extracts (lane 1) could not be restored by 0.2 μM tTFIIB_c (lane 2), whereas tTFIIB restored transcription maximally at that concentration (lane 3). tTFIIB_c maximally inhibited restoration by tTFIIB when added in 5-fold molar excess (lanes 4–8). (C) In control assays, BSA did not restore transcription (lane 2) and did not inhibit restoration by tTFIIB (lanes 4–6).

corresponding region. These attributes are probably functionally important, mediating trypanosome-specific interactions in a preinitiation complex.

Results

tTFIIB and tTFIIB_c Are Stably Folded Monomers. We performed limited proteolysis on recombinant tTFIIB to identify a stable fragment amenable to crystallization, because the full-length protein was refractory to crystallization. The human tTFIIB C-terminal domain forms a protease-resistant core (20, 21). In contrast, tTFIIB was highly protease sensitive and did not yield any stable fragments (Fig. S1). Therefore, we defined the tTFIIB C-terminal domain (tTFIIB_c) by comparing the amino acid sequence of tTFIIB with those of crystal structures of human and archaeal tTFIIB C-terminal domains as shown schematically (Fig. 1A). tTFIIB_c, which comprises the first module and the trypanosome-specific region (residues 87–345), was expressed and purified.

Because the limited proteolysis results suggested that tTFIIB and tTFIIB_c might not be stably folded, we characterized both proteins biophysically. In gel filtration experiments, each protein eluted as a single peak at a position between its calculated dimer and monomer mass (Fig. S2A). This is consistent with the extended monomer conformation of tTFIIB and tTFIIB_c from other organisms (23–25). We found that the fluorescence spectrum of native tTFIIB was blue-shifted when compared to the spectrum under denaturing conditions, suggesting hydrophobic burial of Trp-260 in the native protein. Because Trp-260 is the only tryptophan in tTFIIB and resides in the C-terminal domain, unfolding of both tTFIIB and tTFIIB_c could be monitored by fluorescence. Denaturant-induced unfolding of tTFIIB and tTFIIB_c occurred in a cooperative and reversible fashion with a half-maximal C_m of unfolding between 2 and 3 M guanidinium chloride (Fig. S2B). tTFIIB_c appears to be slightly more stable than full-length tTFIIB (C_m = 2.9 M for tTFIIB_c versus 2.7 M for tTFIIB). Taken together, these results indicate that full-length tTFIIB and tTFIIB_c are stably folded monomers.

Activity of Purified tTFIIB and tTFIIB_c by In Vitro Transcription. We showed previously that tTFIIB expressed and purified from *Escherichia coli* restores transcription in vitro when added to tTFIIB-immunodepleted nuclear extracts of *T. brucei* (14). Using this system, tTFIIB restored transcription maximally at 0.2 μM (Fig. 1B, lane 3, and Fig. S3). By contrast, tTFIIB_c was unable to restore transcription (Fig. 1B, lane 2). Notably, tTFIIB_c had a *trans*-dominant negative effect on transcription. tTFIIB_c inhibited transcription in the presence of full-length tTFIIB when tTFIIB_c was added in 5-fold molar excess (Fig. 1B, lane 7). This effect is specific, as transcription by tTFIIB was unaffected by molar excess of BSA (Fig. 1C). We hypothesize that tTFIIB_c is able to bind to proteins and/or DNA in the preinitiation complex but requires the N-terminal domain to stimulate transcription, consistent with previous studies of human tTFIIB_c (21).

Structure of tTFIIB_c and Comparison to Human tTFIIB_c. We crystallized and solved the structure of tTFIIB_c at 2.3 Å resolution (R_{work} = 20.3%; R_{free} = 24.9%) (Table S1 and Table S2). The tTFIIB_c structure reveals 2, closely packed helical modules followed by a C-terminal extension of 32 aa (Fig. 2A). The trypanosome-specific region comprises the second helical module and the C-terminal extension. The first helical module contains 6 helices (H1–H6), while the second module has 8 helices (H1'–H6' and 2₃₁₀ helices, H2'A and H3'A). Both helical modules contain the canonical 5-helix cyclin fold characteristic of tTFIIB proteins (helices H1–H5 in the first module and H1'–H5' in the second module). The cyclin fold in each module aligns well to the cyclin folds of human tTFIIB_c (rmsd = 1.8 Å on 90 of 96 matched C_α atoms for the first module; rmsd = 2.3 Å on 86 of 100 matched C_α atoms for the second module). The cyclin fold of the second module has a slightly greater deviation from the human cyclin folds owing to divergent regions discussed below. The 32-residue C-terminal extension of tTFIIB_c (residues 314–345) is not visible in the crystal, suggesting that it is unstructured. The prevalence of Pro, Glu, Ser, and Lys residues in the C-terminal extension of the tTFIIB_c is consistent with this conclusion (30).

The overall tTFIIB_c structure is similar to other tTFIIB_c structures (25–29); however, the additional loops and helices between the modules and within the second module of tTFIIB_c explain why primary sequence analysis did not reveal the second cyclin fold (13, 14) (Fig. 2A and B). For instance, the first module has an additional helix H6 that links the 2 modules together by connecting H5 and H1'. Helix H6 essentially forms a bent, 24-residue helix with H1' of the second module (Fig. 2A, blue arrow). In human tTFIIB_c, the helical repeats are connected by a short, random-coil segment that interacts with TBP (26) (Fig. 2B, blue arrow). In tTFIIB_c, the presence of a helix at this position instead of a random coil creates a larger surface that would allow for additional contacts with tTBP.

The second module in tTFIIB_c contains 2 significant insertions within its cyclin fold that are not observed in other tTFIIB structures. The first insertion is the helix H2'A, which is a short 3₁₀ helix inserted into the loop connecting helices H2' and H3'. The second insertion is a much longer segment within the loop between helices H3' and H4'. This insertion consists of another 3₁₀ helix, H3'A, that is followed by a 13-residue linker (Fig. 2A, black arrow). In the tTFIIB_c structure, H3'A veers ≈90° from H3' and away from H4'. The linker (residues 162–174), which connects H3'A to H4', is not visible in the electron density, suggesting that it is unstructured. *T. cruzi* tTFIIB_c also contains a linker in this region of different length but with similar amino acid composition to *T. brucei* tTFIIB (Fig. 2C, black arrow). There appears to be no precedent for a long unstructured segment between helices H3' and H4' in either tTFIIB or cyclin structures in the Protein Data Bank, as assessed by a search using the DALI structural alignment server (31). In these well-studied proteins, the region connecting H3' and H4' is typically a

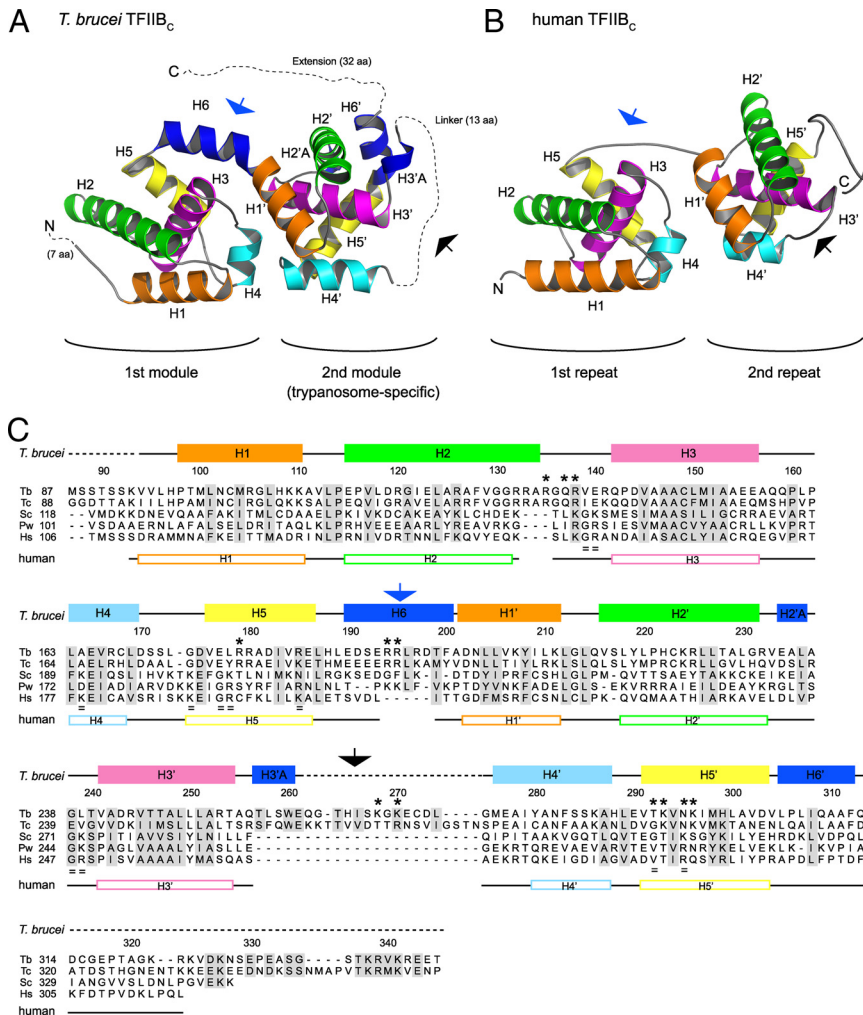


Fig. 2. Structure of the C-terminal domain of *T. brucei* TFIIB (tTFIIB_C) and comparison to human TFIIB_C. (A) Structure of tTFIIB_C (this work). Each helix of the 5-helix fold is colored identically in each of the 2 modules. Additional motifs distinct in tTFIIB (H6, H2'A, H3'A, and H6') are blue. H2'A is behind H2 in this view. Dashed lines denote amino acids not visible in the crystal, and the number of residues is in parentheses. Arrows denote regions for comparison between tTFIIB_C and human TFIIB_C in panel B and their position in the sequence in panel C as discussed in the text: Helix H6 (blue arrow) and the linker between H3'A and H4' (black arrow). (B) Structure of human TFIIB_C (PDB ID 1c9b) (29). (C) The amino acid sequence of tTFIIB_C (Tb; accession no. EAN76636) is aligned with that of *T. cruzi* (Tc; XP_806216), *S. cerevisiae* (Sc; P29055), and the sequences of known TFIIB_C structures from *P. woesei* (Pw; 1d3u), and *Homo sapiens* (Hs; 1c9b). Sequence gaps are denoted by dashes in panel C. tTFIIB_C amino acid numbers and secondary structure elements are indicated above the alignment, and human TFIIB_C secondary structure elements are indicated below the alignment. Helices are depicted as boxes, intervening segments as lines, gaps in the structural alignment are blank, and residues not visible in the electron density are denoted by dashes. Amino acids that were changed to alanine in tTFIIB variants are indicated by asterisks (see Figs. 3 and 4). Structurally equivalent residues in human TFIIB that affected function when mutated are denoted by equal signs (19). Residues Asp-229 to Thr-237 are omitted from the *S. cerevisiae* sequence for clarity.

structured loop of 5–8 residues (Fig. 2B, black arrow). Unstructured protein segments often contain short recognition motifs that mediate protein-protein or protein-nucleic acid interactions (32). We predict that the tTFIIB_C unstructured linker is functionally important, mediating trypanosome-specific interactions in the preinitiation complex.

The overall conformation of tTFIIB_C is closer to human TFIIB_C bound in the TFIIB_C/TBP/DNA complex rather than to the unbound protein (25, 29). This allows us to predict how tTFIIB_C might interact with DNA and tTBP. Strikingly, the electrostatic potential of the putative DNA binding surface of tTFIIB_C is markedly different from the DNA binding surface of human TFIIB_C (Fig. 3A). In the human TFIIB_C/TBP/DNA complex, TFIIB_C contacts DNA through a large positive surface (Fig. 3B). This surface stabilizes the TBP-induced deformation of the DNA by interacting both upstream and downstream of the TATA box. In the TFIIB_C/TBP/DNA complex, the DNA tracks

along the large positive surface of TFIIB_C (Fig. 3B). In stark contrast, the analogous surface of tTFIIB_C is relatively neutral and contains only 2 small patches of positive charge (Fig. 3A, Left). Thus the trypanosome TFIIB_C/TBP/DNA complex probably has a different conformation than the canonical ternary complex.

tTFIIB_C likely interacts with DNA through the 2 small positive patches found on the electrostatic surface of the structure (Fig. 3A, Left). The first patch corresponds to the loop between helices H2 and H3 (Fig. 3C, Left). In human TFIIB_C, the loop between helices H2 and H3, known as the recognition loop, binds to the DNA downstream of the TATA box at the downstream TFIIB recognition element (BRE^D) (Fig. 3A–C, Right) (26, 29). tTFIIB_C contains a similar loop that is preceded by an additional helical turn at the C-terminal end of H2 (Fig. 3C, Left, green arrow). This helical turn extends out from the first module into solvent, placing basic residues in the loop, namely Arg-135 and Arg-138, in position to interact

transcription, probably by stabilizing tTFIIB interactions at the SL RNA gene promoter.

Discussion

This work describes the high-resolution structure of a trypanosome transcription factor. The structure of trypanosome TFIIB_C shares important features with its human counterpart. tTFIIB_C contains 2 cyclin folds that align well to human TFIIB_C even though the second module was not apparent in tTFIIB_C from primary sequence information. In *T. brucei*, the loop between helices H2 and H3 is functionally important, as is the corresponding recognition loop in the human protein. Specifically, mutation of basic amino acids in this region completely compromised TFIIB function in the human and trypanosome proteins. Thus, our work supports a fundamental role of the loop between helices H2 and H3 in transcription.

Trypanosome TFIIB_C deviates in potentially significant ways from its human counterpart. The 2 cyclin folds are connected by a helix (H6) rather than a random coil, and in between helices H3' and H4', there is an additional helix (H3'A) followed by a disordered 13-residue linker. Importantly, mutations in these regions reduced transcriptional activity of tTFIIB, indicating that these regions play a role in tTFIIB function. Moreover, the presence of a potentially protease-sensitive linker between H3' and H4' may explain why tTFIIB lacks a protease-resistant core. In addition to the 13 disordered residues in the linker, the C-terminal 32 residues of the tTFIIB_C structure are also disordered. These regions are common to all trypanosome TFIIB proteins sequenced thus far, and while the sequences vary in length between each protein, they have similar amino acid composition. It is likely that these unstructured segments contain short, peptide motifs involved in recognizing DNA or other trypanosome-specific protein partners, possibly adopting secondary structure upon binding. These segments will be the object of future studies.

Trypanosome TFIIB lacks the conserved Gln and Arg residues in helices H4' and H5' that stabilize the interaction of HTH motifs with the DNA major groove. In addition, tTFIIB H4' has an additional helical turn that would hinder major groove binding. Mutations in H5' (Thr-292 and Asn-295) did not knock out function as they did in the corresponding residues (Val-283 and Arg-286) in human TFIIB (33). These traits in the trypanosome protein may be related to the lack of a BRE^U in the SL RNA gene promoter. In human TATA box-containing promoters, TFIIB binding to the BRE^U directs assembly of the preinitiation complex in the correct orientation. The majority of gene promoters do not contain a TATA box or an associated BRE^U, thus they rely on multiple interactions from transcription-associated factors for preinitiation complex orientation. In trypanosome TFIIB, the linker between H3' and H4', as well as the C-terminal extension, may recruit factors that help determine preinitiation orientation at the SL RNA gene promoter.

tTFIIB_C contains a large neutral surface in place of the highly basic surface on the human protein (Fig. 3). The large neutral surface might be required to stabilize TBP binding to DNA. Whereas human TBP has 4 phenylalanines that deform the TATA box upon binding, only 2 phenylalanines are present in trypanosome TBP (10, 34). Thus, tTFIIB may help stabilize weak tTBP-promoter interactions. In support of this argument, our previous studies showed that tTBP and tTFIIB tightly associate (14).

In well-studied eukaryotic transcription, the basic surface of TFIIB specifically binds at gene promoters to orient RNAP-II so that transcription proceeds in the correct direction. The lack of an extensive basic surface on trypanosome TFIIB may reflect a relaxed specificity of tRNAP-II orientation at promoters in trypanosomes. Most trypanosome protein-encoding genes are

transcribed in large arrays that appear to initiate bidirectionally from strand switch regions (35–37). Weak specificity of preinitiation complex orientation could cause bidirectional transcription from strand switch regions. Thus, trypanosome TFIIB would be relieved of the function to orient tRNAP-II for unidirectional transcription.

Methods

Protein Expression and Purification. Full-length *T. brucei* TFIIB (tTFIIB), the C-terminal domain (tTFIIB_C), and variants were recombinantly expressed and purified from *E. coli* as His₆-MBP fusion proteins following the protocol of Tropea et al. (38). TFIIB and variants were purified by sequential affinity and gel filtration chromatography. For tTFIIB_C, the latter step was replaced by ion exchange chromatography. The selenomethionine labeled tTFIIB_C (SeMet-tTFIIB_C) was expressed as described (39) and purified as described above for native tTFIIB_C. A detailed protocol is provided in the *SI Text*.

Limited Proteolysis. tTFIIB (20 μM) was digested with trypsin or subtilisin in 50 mM Tris-HCl, pH 8.0, 50 mM NaCl at 25 °C for 1 h in 50 μL with an enzyme: substrate molar ratio of 1:400. Aliquots (10 μL) were removed at 5-, 10-, 15-, 30-, 60-min intervals, and the reaction in the aliquot was stopped by addition of 4% SDS, 100 mM DTT, 20 mM EDTA, 2 mM PMSF, and flash-freezing. Samples were heated to 95 °C for 10 min and analyzed by SDS/PAGE.

Gel Filtration and Chemical Denaturation. Gel filtration was performed on a Sephacryl S-200 column equilibrated in 25 mM Tris-HCl, pH 8.0, 150 mM NaCl, 1 mM DTT, at 4 °C with 1 mg of tTFIIB and tTFIIB_C. Chemical denaturation experiments monitoring tryptophan fluorescence were performed at 25 °C (QuantaMaster spectrofluorometer; Photon Technology International). Samples of 3 μM tTFIIB or tTFIIB_C were prepared at each guanidinium chloride (GdmCl) concentration in 25 mM potassium phosphate, pH 7.5, 200 mM KCl, and 1 mM DTT and incubated overnight at room temperature. Fluorescence emission spectra were measured by exciting the samples at 265 nm and recording spectra from 300 to 400 nm (0.1-s integration time) and the center-of-mass of the spectral peak was calculated. The GdmCl concentration was determined by refractive index using a Bausch & Lomb refractometer. GdmCl denaturation of tTFIIB or tTFIIB_C was reversible as judged by the recovery of center-of-mass from dilution of the 4 M GdmCl sample to 2 M GdmCl. Curve fitting was performed in Kaleidagraph (Synergy Software).

Immunodepletion and in Vitro Transcription. *T. brucei* nuclear extracts were prepared as described (11, 40). Immunodepletion and in vitro transcription assays were performed as described (14) with the following exceptions. Transcription was monitored by detecting radiolabeled run-off transcript (172 bases) from a linearized plasmid rather than by primer extension. The plasmid pJP10, which contains the –125- to +120 region of the SL RNA gene was linearized with EcoRV. In the reaction, 250 μM CTP and GTP, 1 mM ATP, and 10 μCi alpha-³²P-labeled UTP were incubated at 28 °C with other reaction components for 10 min, after which cold UTP was added to a final concentration of 50 μM. The potassium glutamate concentration in the reaction was 130 mM.

Structure Determination of tTFIIB_C. Crystals of native tTFIIB_C were obtained by vapor diffusion against a reservoir containing 0.1 M Bis-Tris, pH 5.5, and 24% polyethylene glycol 3350. SeMet-tTFIIB_C protein crystallized in the same space group (P4₃2₁2) but required slightly higher concentrations of polyethylene glycol (32%) to form. The structure was solved using 3-wavelength multiple anomalous dispersion from data collected at National Synchrotron Light Source beamline X29 and refined against a native data set collected at the PHRI X-Ray Crystallography Core Facility. The final model consists of residues 94–261 and 275–313, 56 water molecules, and an ethylene glycol molecule, with values of R_{work} = 0.204 and R_{free} = 0.249. Details are provided in *SI Text*, and data collection and refinement statistics are given in *Table S1* and *Table S2*.

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