Stage-specific transcription activator ESB1 regulates monoallelic antigen expression in Trypanosoma brucei

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Variant surface glycoprotein (VSG) coats bloodstream form Trypanosoma brucei parasites, and monoallelic VSG expression underpins the antigenic variation necessary for pathogenicity. One of thousands of VSG genes is transcribed by RNA polymerase I in a singular nuclear structure called the expression site body (ESB), but how monoallelic VSG transcription is achieved remains unclear. Using a localization screen of 153 proteins we found one, ESB-specific protein 1 (ESB1), that localized only to the ESB and is expressed only in VSG-expressing life cycle stages. ESB1 associates with DNA near the active VSG promoter and is necessary for VSG expression, with overexpression activating inactive VSG promoters. Mechanistically, ESB1 is necessary for recruitment of a subset of ESB components, including RNA polymerase I, revealing that the ESB has separately assembled subdomains. Because many trypanosomatid parasites have divergent ESB1 orthologues yet do not undergo antigenic variation, ESB1 probably represents an important class of transcription regulators.

Monoallelic expression of a single gene family member underpins a molecular ‘arms race’ between many pathogens and their host, through host monoallelic immunoglobulin and pathogen monoallelic antigen expression. The unicellular parasite Trypanosoma brucei is an archetypal example, achieving antigenic variation through monoallelic expression of one of a library of thousands of variant surface glycoproteins (VSGs). VSG covers the entire cell surface in life cycle stages that inhabit the host bloodstream or are preadapted for transmission to the host. The single active VSG gene is transcribed by RNA polymerase I (Pol I) from a specialized bloodstream form (BSF) telomeric expression site (BES), where it is co-transcribed along with four or more expression site (ES)-associated genes (ESAGs) using a single promoter. Switching of VSG is achieved by switching to transcription of one of several different telomeric BESs or replacement, by recombination, of the VSG in the active BES with one of the ~2,500 VSG gene and pseudogene variants elsewhere in the genome. The active BES is found in a specialized Pol I-containing, non-nucleolar, nuclear structure called the expression site body (ESB), from which inactive BESs are excluded. The ESB is present only in BSF parasites, despite procyclic forms (in tsetse fly) also employing Pol I-dependent transcription of their invariant surface coat (procyclic). Elegant biochemical candidate approaches and genetic screens of VSG expression have revealed the importance of epigenetic silencing, telomere and chromatin factors and SUMOylation. VEX proteins, required for exclusion of the inactive BESs, associate the single active BES with the spliced leader array chromosomal locations. These contain the repetitive genes encoding a sequence which, after transcription and processing, is added to every trypanosome messenger RNA. Hence, in addition to other properties, VEX proteins link an ESB-located exclusion phenomenon to an active VSG gene mRNA-processing capability. Notwithstanding these advances, bloodstream-specific factors remain elusive and the statement that “No ESB-specific factor has yet been identified” still holds true. Here we used a medium-throughput localization screen to identify ESB-specific protein 1 (ESB1), which is expressed only in mammalian infectious forms and is localized specifically to the ESB. ESB1 is required for VSG expression and is located near the active VSG promoter, with overexpression activating inactive VSG promoters. We show that ESB1 is required for recruitment of some, but not all, ESB components, revealing that the ESB has separately assembled subdomains. Many trypanosomatid parasites have a divergent ESB1 orthologue, and therefore ESB1 potentially represents an important class of trypanosome transcription regulators.

Results
We performed a candidate protein localization screen of proteins of unknown function upregulated in the BSF, and identified an ESB-specific protein, G1 BSF nuclei have one extranucleolar ESB. From 207 candidates, 153 were successfully localized and only one (Fig. 1b), Tb427.10.3800, exhibited this localization (Fig. 1c and Extended Data Fig. 1a) whilst endogenous tagging in the procyclic form gave no detectable signal (Extended Data Figs. 1a and 2a,b). We named this protein ESB1.

We used well-characterized ESB markers to confirm ESB1 localization. Pol I is the founding component of the ESB and localizes to both the nucleolus and ESB in BSFs. ESB1 lies extremely close to Pol I (RPA2) at the ESB (Fig. 1c), as confirmed by measurement of the distance between signal centre points (Fig. 1d). The ESB also has a VEX subcomplex involved in exclusion of inactive ESSs. ESB1 lies ~300 nm from the nearest VEX1 or VEX2 focus (Fig. 1c,d),
Fig. 1 | A protein localization screen identified ESB1. a, b. Degree of upregulation of T. brucei mRNAs in BSFs previously determined by RNA-seq49, highlighting known VSG monoallelic expression-associated factors (a) and candidates for tagging that we selected and successfully localized (b).

c–f. Fluorescence microscopy analysis of ESB1 subcellular localization relative to known ESB-associated proteins. c. Representative images from at least n = 3 independent sample preparations of G1 (1K1N) cells from cell lines expressing one mNG-tagged and one tdt-tagged ESB-associated protein. For cells expressing VEX1 or VEX2, examples with one nuclear focus are shown. d, Histograms of pairwise distance measurements between the ESB1 focus, RPA2 ESB focus and the nearest VEX1 or VEX2 focus. For each, n ≥ 45 cells from one clonal cell line and all distances are significantly different (P < 10−80, two-tailed Mann–Whitney U-test). Multicolour beads are a control for measurement accuracy (true distance from zero). e, Number of mNG-tagged ESB1, VEX1 or VEX2 foci per nucleus in different cell cycle stages; n indicates the number of cells counted from one clonal cell line. The number of ESB1 foci significantly differs from that of VEX1 or VEX2 in 1K1N cell nuclei (P < 10−9, χ2-test). f, ESB1 localization in mitotic nuclei representative of n = 3 independent sample preparations. RPK, reads per kilobase.

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ESB1 is necessary for active ES transcription. To determine ESB1 function, we generated a BSF ESB1 conditional knockout (cKO) cell line (Extended Data Fig. 3a–e). ESB1 cKO gave undetectable levels of ESB1 protein by 24 h (Extended Data Fig. 3a), which caused a profound proliferation defect due to failure of cytokinesis and further rounds of organelle duplication (Fig. 2a–c). To detect any effect on transcription we used RNA sequencing (RNA-seq) to profile mRNA levels, which showed that ESB1 cKO caused a marked decrease (~250-fold) in ESAG mRNAs, predominantly those transcribed from the active ([Fig. 2d and Extended Data Fig. 3e], associated with almost total loss of ESB1 transcript (Fig. 2e). mRNAs from the VSG gene in the active ES decreased ~eightfold (Fig. 2d), which we confirmed by quantitative PCR with reverse transcription (RT–qPCR) (Fig. 2f). The smaller decrease in VSG mRNAs is probably explained by the longer half-life of VSG mRNAs.

To understand at what stage ESB1 functions in VSG and ESAG mRNA production, we analysed changes to nascent mRNAs in the BSF ESB1 cKO. Co-transcriptional trans-splicing and polyadenylation generate mature mRNAs, enabling quantification of unprocessed transcript from RNA-seq reads spanning the spliced leader acceptor (SLAS) and polyadenylation (PAS) sites. Unprocessed ESAG and VSG mRNAs also dropped dramatically following ESB1 cKO (Fig. 2g), indicating that ESB1 cKO reduces active ES transcription rather than mRNA processing. Some low-processivity transcription of inactive ESs occurs; ESB1 cKO caused a smaller reduction in unprocessed transcript from inactive ESs (Fig. 2g) and mRNAs transcribed from specific inactive ESs (Fig. 2h), while mRNAs from promoter-proximal ESAG genes in the active ES tended to be less strongly reduced (Fig. 2i). Therefore, the highly processive active ES transcription is ESB1 dependent, with ESB1 cKO leaving a little residual transcription such as seen at silent ESs.

A specific transcription activator would be predicted to associate only with the promoter region of the active ES. Therefore, we carried out ESB1 chromatin immunoprecipitation sequencing (ChiP–seq). Across the genome, the highest peak in ESB1 ChiP/input DNA ratio (30-fold background signal) was in the active ES. Among the ESs the active ES had the highest average ChiP ratio (Fig. 2j and Extended Data Fig. 4), due to a large peak between ~5 and 15 kb upstream and a smaller peak ~5 kb downstream of the Pol I promoter (Fig. 2k). The former corresponds to the imperfect 50 base pair (bp) repeats found upstream of all ESs, but ESB1 associates with these repeats only at the active ES.

Procytic forms lack an active BES and an ESB and do not express ESB1, although they use Pol I for expression of their surface coat protein (procyclin) whose locus we refer to as a procyclin expression site. We tested ESB1 cryptic function in procytic forms by deletion of both ESB1 alleles, which resulted in no apparent growth or morphology defect. RNA-seq confirmed normal high expression of GPEET procyclin and no major changes to other mRNA transcripts (Fig. 3 and Extended Data Fig. 3f). ESB1 is therefore vital in BSFs for monoallelic VSG expression, but is dispensable in procytic forms.

For further experiments, the BSF cKO phenotype was recapitulated with the more experimentally amenable RNA interference knockdown of ESB1 (Extended Data Fig. 5). mRNA abundance changes correlated extremely well with cKO (Fig. 2 and Extended Data Fig. 5k), with the same ESAGs and VSGs mRNAs reduced, as were the same set of 11 upregulated mRNAs (probably endoplasmic reticulum stress-associated; Extended Data Fig. 5k). The rapid lethality of the RNAi phenotype naturally led to the appearance of RNAi escape subpopulations, therefore, we analysed only early RNAi time points.

ESB molecular composition depends on ESB1. We next determined whether ESB1, and thus active ES transcription, is required for the normal molecular composition of the ESB. We generated a panel of cell lines carrying the inducible ESB1 RNAi construct and tagged the following ESB-associated proteins: RPA2, SUMO (because the ESB is associated with a highly SUMOylated focus (HSF)) and VEX1 or VEX2 (Fig. 4a–h). As shown by others, the ESB focus of RPA2 was visible in 40% of G1 nuclei (that is, when not occluded by nucleolar RPA2) and the HSF in ~60% of G1 nuclei. After 24 h induction of ESB1 RNAi, RPA2 and SUMO were more dispersed through the nucleus and fewer nuclei had an ESB focus, in both morphologically normal and abnormal cells, while nucleolar RPA2 was unaffected (Fig. 4a–d). As seen previously, VEX1 and VEX2 localized to one or two foci in the nucleus. After 24 h induction of ESB1 RNAi, the localization pattern was unchanged, in both morphologically normal and abnormal cells (Fig. 4e–h). This indicates that ESB1 is necessary for both recruitment of Pol I to the ESB and higher local SUMOylation to form the HSF, but not for the formation of VEX foci.

The inverse, whether the ESB1 focus is VEX1 or VEX2 dependent, was analysed based on their deletion using RNAi and observing tagged ESB1 (Fig. 4i–l). VEX1 knockdown was confirmed using

**Fig. 2** | ESB1 is vital for BSFs and is required for transcription from the active VSG expression site. a–f. Cellular phenotype of BSF ESB1 cKO. Exogenous mNg-tagged ESB1 expression was maintained with 10 ng ml⁻¹ doxycycline (+Dox) in the BSF ESB1 cKO cell line validation shown in Extended Data Fig. 3a–d, and doxycycline washout (−Dox) induced the cKO phenotype. a. Culture growth (with subculture), mean ± s.d., n = 3 inductions. P value shown is at 48 h, two-tailed t-test, log cumulative growth. b. Counts of morphologically abnormal (‘Other’) cells following washout. n indicates number of cells counted, representative example from n = 3 inductions. P value derived from χ²-test. c. Representative images from n = 3 independent inductions showing mNg-tagged ESB1 signal before and after 24 h −Dox. d. Volcano plot of change in mRNA abundance as determined by RNA-seq after 48 h −Dox, n = 4 inductions (further time points shown in Extended Data Fig. 3e). P values derived from two-tailed t-test. A-BES and I-BES indicate active and inactive BESes, respectively. e. Change in ESB1 ORF read coverage for 48 h −Dox, mean of n = 4 inductions. f. RT–qPCR quantitation of TA-ES VSG mRNA (VSG221) −Dox, mean ± s.d. from n = 3 inductions. P ≤ 0.05 derived from two-tailed t-test. g. Profile of transcript abundance change following ESB1 loss in BSFs. h. Changes in total and unprocessed mRNA grouped into A-BES or I-BES VSg(s) and ESAgs for cKO ±Dox. Mean ± s.d. from n = 4 inductions. P ≤ 0.05 derived from two-tailed t-test. Average change in transcript abundance averaged per ES (h) and per gene (i) for active BES plotted by distance from the promoter after 48 h −Dox; n = 4 inductions, mean ± s.d. j. ESB1 ChiP–seq shown as the ratio of ChiP to input DNA, plotting mean ratio per BES (j) and mean ratio in 2 kb bins across the active BES (k) (extended in Extended Data Fig. 4). Non-analysable bins had insufficient uniquely mapped reads from the input DNA. n = 2 replicates.
Overexpressed ESB1 still localized to the ESB, although with more small growth reduction and some cytokinesis defects (Fig. 5a–c). Data Fig. 3b). In contrast to ESB1 cKO, overexpression yielded a 1284 mean ± s.d. 3 clonal ESB1 KO cell lines relative to the parental cell line, as determined by RNA-seq. Fourth bar shows change values derived from two-tailed t-test. n = 3 independent clonal cell lines (plotted individually in Extended Data Fig. 3f).

We have identified ESB1 as both an ESB-specific protein and/or supernumerary ESB formation. Overexpression was achieved using a cell line with an additional inducible tagged ESB1 locus (using 100 ng ml–1 doxycycline; Fig. 3f). ESB1 localization was unchanged following either VEX1 or VEX2 knockdown (Fig. 4k,l); therefore, formation of a singular ESB is not counterbalanced by strong repression of all other BESs, and processivity of transcription complex is additional machinery required for VSG transcript maturation, which could be regulated by ESB1. However, ESB1 was not strongly affected than promoter-distal ESs and VSGs (Extended Data Fig. 6d), unlike the phenotype of VEX1 knockdown (Extended Data Fig. 6e). ESB1 overexpression is therefore sufficient to cause activation of inactive BES transcription, although it may not be fully processive. All cells still expressed VSG221 (Extended Data Fig. 6f,g), therefore probably expressing multiple VSGs rather than switching to an alternative ES and VSG, whilst expression of procyclic form-specific surface proteins (procyclins) remained low (Extended Data Fig. 6a).

Finally, we forced expression of tagged ESB1 in procyclic form cells (Fig. 6a–d and Extended Data Fig. 6h). Expression produced no growth or cytokinesis defects (Fig. 6a,b) and tagko ESB1 was nuclear, but did not localize to a single extranucleolar ESB-like focus (Fig. 6c). RNA-seq analysis showed a large increase (up to ~200-fold) in mRNA levels for ESAGs, consistent with the activation by ESB1 of transcription initiation at ES promoters normally inactive in the procyclic form (Fig. 6d). This particular strain, we interrogated expression of the ESAGs and VSG from the sequenced BES37. Every ESAG transcribed from this BES was upregulated, typically ~three- to fivefold and up to ~80-fold (Fig. 6f,g). In contrast, VSG mRNAs (both published and from our de novo assembly of the transcriptome) were not strongly upregulated (Fig. 6d). We did not see a transcript from VSG 10.1, found in the sequenced BES, nor upregulation of any of the VSGs commonly expressed by this strain in BSFs during mouse infection38. This is despite ~50-fold overexpression of the ESB1 transcript relative to endogenous BSF expression (Figs. 5e and 6e). In regard to tagged ESB1 overexpression in the BSF, procyclin mRNA levels also remained unchanged (Fig. 6d). Hence ESB1 expression in procyclic forms activates BES transcription without formation of an ESB; however, transcription is either not fully processive to the most distal gene (VSG) or there is additional machinery required for VSG transcript maturation, processing and/or stability not expressed in the procyclic form—for example, CFB2 (ref. 39).

Discussion

Antigenic variation in T. brucei relies on monoallelic expression of the VSG gene in the active BES. Our results provide the basis for a model whereby strong transcription activation of the active BES is counterbalanced by strong repression of all other BESs, and provides insights into the ESB subdomains that orchestrate these different functions.

We have identified ESB1 as both an ESB-specific protein and an ES transcription activator enriched near the Pol I promoter. We show that ESB1 is necessary for the high level of transcription from the active BES and that its overexpression activates only VSG-containing ESs and not procyclin loci. Importantly, both BESs and metacyclic VSGs are upregulated, and previous transcriptomics showed that metacyclic forms have upregulated ESB1 (ref. 40), indicating that VSG expression in the earliest VSG-expressing

RNA-seq profiling of mRNA and, as previously described23, we saw derepression of inactive BESs (Extended Data Fig. 6e) with no growth defect (Fig. 4i,j). VEX2 knockdown was confirmed by carrying out knockdown in a cell line expressing tagged VEX2. ESB1 localization was unchanged following either VEX1 or VEX2 knockdown (Fig. 4k,l); therefore, formation of a singular ESB is not dependent on repression of inactive BESs by the VEX complex. The ESB1 and VEX2 compartments also have differing sensitivity to small molecule inhibitors. VEX2 foci became distributed following inhibition of Pol I transcription (BMH-21, an indirect Pol I inhibitor; Extended Data Fig. 6a). Expression produced no growth or cytokinesis defects (Fig. 4a,b) and tagko ESB1 was nuclear, but did not localize to a single extranucleolar ESB-like focus (Fig. 4c). RNA-seq analysis showed a large increase (up to ~200-fold) in mRNA levels for ESAGs, consistent with the activation by ESB1 of transcription initiation at ES promoters normally inactive in the procyclic form (Fig. 4d). This particular strain, we interrogated expression of the ESAGs and VSG from the sequenced BES37. Every ESAG transcribed from this BES was upregulated, typically ~three- to fivefold and up to ~80-fold (Fig. 4f,g). In contrast, VSG mRNAs (both published and from our de novo assembly of the transcriptome) were not strongly upregulated (Fig. 4d). We did not see a transcript from VSG 10.1, found in the sequenced BES, nor upregulation of any of the VSGs commonly expressed by this strain in BSFs during mouse infection38. This is despite ~50-fold overexpression of the ESB1 transcript relative to endogenous BSF expression (Figs. 4e and 4f). In regard to tagged ESB1 overexpression in the BSF, procyclin mRNA levels also remained unchanged (Fig. 4d). Hence ESB1 expression in procyclic forms activates BES transcription without formation of an ESB; however, transcription is either not fully processive to the most distal gene (VSG) or there is additional machinery required for VSG transcript maturation, processing and/or stability not expressed in the procyclic form—for example, CFB2 (ref. 39).

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Fig. 3 | ESB1 is dispensable in procyclic forms. a, c, RNA abundance phenotype of procyclic form ESB1 KO. a, Volcano plot of change in mRNA abundance; n = 3 independent clonal cell lines (plotted individually in Extended Data Fig. 3f). P values derived from two-tailed t-test. b, Change in ESB1 ORF read coverage, mean of n = 3 clonal ESB1 KO cell lines. c, Abundance of GPEET procyclin in n = 3 clonal ESB1 KO cell lines relative to the parental cell line, as determined by RNA-seq. Fourth bar shows mean ± s.d.
life cycle stage is ESB1 dependent. Ectopic expression of ESB1 in procyclic forms that never naturally express VSG was sufficient to activate BES promoter transcription, upregulating ESAGs located within a BES. However, ESB1 alone in procyclic forms was not sufficient for fully processive BES transcription and/or VSG mRNA processing. Interestingly, all trypanosomatid parasites, most of which do not undergo similar antigenic variation, have divergent ESB1 orthologues (Extended Data Fig. 7a). All orthologues have an N-terminal RING U-box domain while the weakly conserved C-terminal domain is not present in Trypanosoma cruzi and related Trypanosoma spp. and, when it is present, has very low sequence similarity to T. brucei (Extended Data Fig. 7b,c). This raises the prediction that PolI transcription of protein-coding genes and their regulation may occur in other trypanosomatid parasites.
**Fig. 5 | ESB1 overexpression in BSFs activates inactive BESs without affecting ESB formation.** a-f. Cellular phenotype of mNg-tagged ESB1 overexpression in BSFs induced with +Dox. a, Culture growth (with subculture), mean ± s.d., n = 3 inductions. P value shown at 48 h, two-tailed t-test, log cumulative growth. b, Counts of morphologically abnormal (Other) cells following washout. n indicates number of cells counted, representative example from n = 3 inductions. P value derived from χ²-test. c, Representative images from n = 3 independent inductions showing mNg-tagged ESB1 signal before and after 24 h +Dox. d, Volcano plot of change in mRNA abundance as determined by RNA-seq for 24 h –Dox, n = 4 inductions (further time points shown in Extended Data Fig. 6a). P value derived from two-tailed t-test. e, Change in ESB1 ORF read coverage as determined by RNA-seq 24 h after washout, mean ± s.d. from n = 3 inductions. P ≤ 0.05 derived from two-tailed t-test. f, RT-qPCR quantitation of A-ES VSg mRNA (VSg221) +Dox, mean ± s.d. from n = 3 inductions. χ²-test. g–i. Effect of mNg-tagged ESB1 overexpression on tdT-tagged RPA2 localization. Counts of morphologically abnormal cells (g) and number of cells with an RPA2-containing ESB focus (h); n indicates number of cells counted from one induction. P ≤ 0.05 derived from χ²-test. i, Representative images from n = 1 replicate showing tagged protein localization before induction and in morphologically normal and abnormal cells after ESB1 overexpression.

**Fig. 6 | ESB1 overexpression in PCFs activates BES transcription without formation of an ESB.** a-e, Cellular phenotype of mNg-tagged ESB1 overexpression in procyclic forms induced with 1 µg ml⁻¹ doxycycline. Culture growth (with subculture, a) and counts of morphologically abnormal cells (Other, b) after induction; n indicates number of cells counted from one induction, no changes in P ≤ 0.05 from χ²-test. c, Example fluorescence images from n = 1 induction of overexpressed mNg-tagged ESB1 in procyclic forms. d, Volcano plot of change in mRNA abundance, n = 4 inductions (further time points shown in Extended Data Fig. 6h); P values derived from two-tailed t-test. e, Change in ESB1 ORF read coverage as determined by RNA-seq 24 h after induction, mean of n = 4 inductions. f, ESP read coverage, as determined by RNA-seq 24 h after ESB1 overexpression, of the active and an example inactive BES in BSFs (f) (from Fig. 5), and an example inactive BES in procyclic forms (g); mean of n = 4 inductions. Further details given in Extended Data Fig. 8.
ESB1 alone was also not sufficient to support formation of the PolI and ESB1 focus, because overexpression of ESB1 did not give rise to an ESB-like body/bodies in the procyclic form, or to supernumerary ESB-like bodies in the BSF. Moreover, multiple active BESs in multiple ESBs do not represent a stable state: in cells forced to express two VSGs from two BESs, both were recruited to a single ESB15. Given this, our ESB1 overexpression results suggest that the reasons for ESB absence (procyclic forms) or singularity (BSFs) are likely to be more complex than a threshold level of ESB1 protein. Phase separation, common in nuclear compartment formation, is among potential mechanisms for ESB formation where singularity could be achieved by emergent properties (Otswald ripening); however, ESB1 appears strongly chromatin associated, perhaps acting as a single nucleation site. These are open hypotheses for future work, and may also have important implications for understanding of switching between BESs.

Our work, taken with that of others, shows that the ESB is a complex nuclear body with multiple subdomains. The defining subdomain is a focus of PolI around the active BES, which also contains basal PolI transcription factors20 and ESB1. This is associated with a HSF1. ESB1 is required for assembly of this subdomain. The BES is found in close proximity to one of the spliced leader array alleles26. PolII transcription of this array generates the spliced leader RNA necessary for processing of all transcripts into mRNA. Each spliced leader array allele is found in a PolII transcription focus1, and the proximity of one allelic copy to the ESB PolI subdomain provides a mechanism for efficient processing of the large amount of VSG mRNA. BES association with the ESB spliced leader array/PolII subdomain requires VEX2 (ref. 2) and ESB/BES/PolI subdomain overlaps, or is adjacent to one VEX1 and VEX2 nuclear focus2. We show that assembly of these foci is separable, with assembly of the VEX foci not dependent on ESB1 and vice versa. Importantly, we show that the PolI and ESB1 focus is strictly singular. This enhanced appreciation of the ESB in terms of spatially defined subdomains raises the possibility that it reflects an intrinsic functional architecture.

Methods

Parasite strains and cell culture. Trypanosoma brucei Lister 427 BSF was used because its expression sites are sequenced1 and assembled into contigs. BSFs were grown in HMI-9 (ref. 27) at 37 °C with 5% CO2 maintained under ~2 × 106 cells ml−1 by regular subculture. The active BES was BES1-containing VSG221 (also called VSG 427-2). T. brucei/TREU927 procyclic form (PCF), selected because it is the original genome strain with genome-wide PCF localization data28,45, was used as a wild-type control. The active BES was BES1-containing VSG221 (also called VSG 427-2). This strain was used in all experiments unless otherwise indicated. BSF927 was used for the localization screen (by limiting dilution cloning). Cultures were maintained, of the appropriate prewarmed medium for 6 h, then the necessary drugs added to achieve the expected growth rate defect and change in proportion of cells at different cell cycle stages.

PCR validation of endogenous locus ORF modification/loss. Key endogenous locus modifications were validated by PCR using template genomic DNA extracted using the DNeasy Blood & Tissue Kit (Qiagen). Primer pairs (Extended Data Table 2) spanned the expected genomic DNA sequence to integrated DNA: for deletions, the genomic DNA sequence (CDS) to the ESB1 CDS in PCFs.

Middle-throughput BSF localization screen for ESB proteins. Tagging candidates were selected using published mRNA abundance data (RNA-seq)22, taking those with significantly upregulated transcripts (P < 0.05, two-tailed t-test) in BSFs relative to PCFs and prioritizing those >2.5-fold upregulated (Fig. 1b).

Endogenous tagging. To tag genes at the endogenous gene loci, we used long primer PCR and the pPOT plasmid series as the template to generate tagging constructs and, for BSF tagging, PCR to generate DNA encoding single-guide RNA with a 17 promoter44. mNG was used for green fluorescent protein tagging, except for cell lines for ChlEP where e-yellow fluorescent protein (eYFP) was used. pPOTv7-blast-mNG was used for the medium-throughput BSF localization screen. pPOTv6-blast-3Ty::mNG-3Ty was used for other experiments and, for simplicity, we refer to this as a 6Ty::mNG construct. pPOTv7-hyg-Tomato was used for tagging with a red fluorescent protein. PCR confirmed correct fusion of the mNG tagging sequences (CDS) to the ESB1 CDS in PCFs.

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Inducible RNAi knockdown. For inducible ESB1, VEX1 or VEX2 RNAi knockdown we cloned a fragment (primer sequences shown in Extended Data Table 3) of the target gene ORF into a new doxycycline-inducible RNAi construct, pDRv0.5 (Supporting Information). This gives two copies of the fragment in reverse complement separated by a 150 nt stuffer. Two opposing T7 promoters under the control of doxycycline drive transcription of the resulting stem-loop. Cells were transfected with NotI linearized plasmid and selected using Hygromycin B. The construct integrates into the ribosomal RNA array. RNAi was induced using 1 µg ml−1 doxycycline.

To confirm effective knockdown, we introduced RNAi constructs into cell lines expressing endogenously tagged copy of the target protein whose knockdown was confirmed by light microscopy (Extended Data Figs. 5c and 41) and/or immunoblot (Extended Data Fig. 5d) and/or RNA-seq to determine the transcript abundance of the target gene (Fig. 4).

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Induction time series. RNAi and cKO cell lines were analysed as induction time series with paired induced and uninduced samples. Cells were subcultured to either 1 × 10^6 cells ml⁻¹ (BFSs) or 1 × 10^5 cells ml⁻¹ (PCFs), one sample without and one with the appropriate doxycycline concentration for induction. Each 24 h culture density was measured, samples taken and the remaining cells subcultured to either 1 × 10^6 cells ml⁻¹ (BFSs) or 1 × 10^5 cells ml⁻¹ (PCFs), with inclusion of doxycycline in the induced sample. For cultures with a strong growth defect, the culture was centrifuged at 1,200 × g for 5 min, the cell pellet resuspended in fresh medium and doxycycline added if needed, to maintain constant conditions. Growth defects were tested with two-tailed t-tests on log-transformed cumulative growth.

Microscopy. Unless otherwise noted, light microscopy was carried out on live cells adhered to glass, with DNA stained by Hoechst 33342 (ref. [7]), captured on a DM5500 B (Leica) wide-field epifluorescence microscope using a plan apo x63/1.4 numerical aperture phase contrast oil-immersion objective (Leica, no. 15506351) and a N.A. 1.45 (Andor) CMOS camera using Micro-Manager (v.1.4).45

Kinetoplasts (K, mitochondrial DNA) and nuclei (N) in cells were counted from micrographs as a measure of cell cycle stage. K division normally precedes N division, giving 1K1N, 2K1N then 2K2N cells before cytokinesis. Cells with abnormal K/N numbers were classified as ‘Other’. Change in cell cycle stage from micrographs as a measure of cell cycle. K division normally precedes N division.

For blinded counts, one researcher identified and cropped in-focus nuclei of 1K1N cells from a mixture of test and control samples and saved each with a randomized file name while generating an index. A second researcher classified the nuclei according to K+N unbinned or the defined index file.

For anti-VSG221 immunofluorescence, slides were prepared as for live-cell microscopy then cells were fixed with 2% formaldehyde for 5 min. Slides were then washed three times with PBS, incubated with 2:1:1 polyclonal rabbit anti-VSG246 for 1 h, washed three times with PBS, incubated anti-rabbit Alexa Fluor 647-conjugated secondary antibody for 1 h, washed three times with PBS and mounted with 50 mM phosphate-buffered 90% glycerol60.

Transcriptomic analysis. RNA samples for each experiment were purified simultaneously by inducing separate samples at appropriately staggered intervals. A paired uninduced sample, maintained by the same pattern of subculture, was generated for each induction time point. From this time series, a time of primary interest was identified and three further paired samples were prepared. For each, 10^6 cells were harvested by centrifugation at 3,200 × g for 90 s, the supernatant discarded by pipetting and the pellet flash-frozen in a dry ice/ethanol bath at −80 °C. Total RNA was diluted to 500 ng µl⁻¹ based on OD260 and RT–qPCR performed using the Quantitect SYBR Green RT-PCR Kit (Qiagen, no. 204423) with the manufacturer’s recommended reaction composition and thermocycle on an Mx3000p QPCR machine (Agilent).

For de novo transcriptome assembly we used Trinity (v.2.11.0) guided by Harvard PAS best practices. Sequencing errors were first corrected using USEARCH62 and trimmed (v.10.0) uncorrected reads were mapped using Trinity adapters and low-quality sections were trimmed with Trim Galore! (v.0.6.0) with flags −l 36, −q 5,–stringency 1 and −e 0.1. Finally, read sets that exactly matched four or more bases of the 3′ end of the T. brucei spliced leader sequence were trimmed. Trinity, using default settings, generated the assembly.

ChiP–seq. For ChiP–seq we used the following optimized protocol:5 × 10^5 cells were harvested by centrifugation at 3,200 × g for 90 s, the supernatant discarded, and the pellet flash-frozen in a dry ice/ethanol bath at −80 °C. Total RNA was diluted to 500 ng µl⁻¹ based on OD260, and RT–qPCR performed using the Quantitect SYBR Green RT-PCR Kit (Qiagen, no. 204423) with the manufacturer’s recommended reaction composition and thermocycle on an Mx3000p QPCR machine (Agilent).

Both input and ChIP DNA were sequenced by 50 bp single-end sequencing. To quantify transcript abundance from whole RNAs rather than CDS, we first mapped the 5′ and 3′ UTRs using all our BSF Lister 427 RNA-seq data. SLASs and spanned sites (1.00 if the only site, 0.05 if spanning a site used 5% of the genome) were identified and fitted and then normalized to a score per 1,000,000 reads (that is, reads per million like).

Mean read coverage was calculated from samtools depth, with flags -aa -d -t. Mean standard deviation of log fold change for transcripts binned by RPKM was calculated and fitted using a third-order polynomial for plotting. For time points with multiple replicates, mean and two-tailed t-tests were performed on log fold change were calculated for volcano plots.

Immature/nascent transcripts were quantified by filtering the alignments for reads spanning a SSAS or PAS, indicating that trans-splicing or polyadenylation, respectively, may not yet have occurred. Reads were scored by the sum frequency of reads spanning sites (1.00 if the only site, 0.05 if spanning a site used 5% of the genome) if spanning two sites used 63 and 34% of the time, respectively, and so on) then normalized to a score per 1,000,000 reads (that is, reads per million like).

Active VES (VSG221) RT–PCR was used a one-step protocol from total RNA, with β-tubulin as a control (primer sequences shown in Extended Data Table 4). Total RNA was diluted to 50 ng µl⁻¹ based on OD260, and RT–qPCR performed using the Quantitect SYBR Green RT-PCR Kit (Qiagen, no. 204423) with the manufacturer’s recommended reaction composition and thermocycle on an Mx3000p QPCR machine (Agilent). Specific PCR product was confirmed by gel electrophoresis and product melt curve analysis, with no template or primer contamination.

A six-step free-dead binding fluorescence assay (100 to 10^2 copies) of parental cell line RNA using a Neo v.5.5 (Andor) sCMOS camera using Micro-Manager (v.1.4).45

NATURE MICROBIOLOGY | VOLUME 7 | AUGUST 2022 | 1280-1290 | www.nature.com/naturemicrobiology
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**Author contributions**

Conceptualization was the responsibility of K.G., J.D.S., R.J.S. and S.D. Formal analysis was carried out by R.J.W. Funding was acquired by K.G., R.J.W., S.D. and J.D.S. L.E., B.H., C.H., J.D.S., R.W.J., S.D., B.A. and M.I. carried out investigations. Supervision was performed by K.G., J.D.S., R.J.W., S.D. and B.A. R.J.W. and L.E. carried out visualization. K.G., J.D.S., R.J.W. and S.D. wrote the paper.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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Extended Data Fig. 1 | Bloodstream form-upregulated nuclear T. brucei proteins. Screening epifluorescence images of cell lines expressing tagged proteins, each image representative of n = 1 non-clonal cell line. Images for each cell line are laid in the same format: Left, an overlay of the phase contrast (grey), mNG fluorescence (green) and Hoechst DNA stain (magenta). Middle, the DNA stain and the mNG fluorescence. Right, the mNG fluorescence in greyscale. Fold upregulation in bloodstream form cells is shown in the top right. A) Subcellular localisation of all 3 proteins >2.5x upregulated in bloodstream forms49 which localised to one or multiple points in the nucleus when tagged at the N terminus in bloodstream forms. B) Subcellular localisation of the 3 proteins in A) in equivalent procyclic form cell lines, shown at the same contrast levels. C) Subcellular localisation of the remaining 4 proteins >2.5x upregulated in bloodstream forms which localised to the wider nucleus when tagged at the N terminus in bloodstream forms. D) Subcellular localisation of the 4 proteins in C) in equivalent procyclic form cell lines, from TrypTag35, shown at the same contrast levels. We were unable to generate two cell lines.
Extended Data Fig. 2 | Tagging does not perturb ESB1 localisation or function. A) Clonal bloodstream form and procyclic form cell lines expressing Tb427.10.3800 or Tb927.10.3800 (ESB1) N terminally tagged with 6×Ty::mNg respectively were re-generated following the initial screen. Epifluorescence images representative of n = 1 clonal cell line of the localisation of the tagged protein by mNg fluorescence. B) Confirmation of the expected genetic modification of the cell lines in A) by PCR from genomic DNA using a forward mNg and a reverse ESB1 ORF primer. Schematic shows the primer binding sites, uncropped DNA gel shows the resulting PCR products from extracted genomic DNA from the tagged (Tag.) or parental (Par.) cell line. C) Epifluorescence images representative of n = 1 clonal cell line of bloodstream form cell lines expressing 6×Ty::mNg::ESB1 (N terminal tag), ESB1::mNg::6×tY (C terminal tag). D) Count of the number of points per nucleus at different stages of the cell cycle (1K1N, 2K1N and 2K2N) for N or C terminally tagged ESB1, from n = 1 replicate. E) Example field of view representative of n = 1 clonal cell line of cells expressing 6×Ty::mNg::ESB1 showing the strictly singular nature of the ESB1 nuclear focus. F) Epifluorescence image representative of n = 1 clonal cell line of bloodstream form cell line with one N terminally tagged ESB1 allele and the other deleted by replacement with a drug selectable marker. G) PCR validation of the sKO cell line. Schematics represent the deleted ESB1 ORF (top) and N terminally tagged locus (bottom) and primer binding sites, uncropped DNA gels shows the resulting PCR products from extracted genomic DNA. H) Count of the number of points per nucleus at different stages of the cell cycle for the sKO cell line, from n = 1 replicate. I) Proportion of cells at different stages of the cell cycle for the sKO in comparison to N or C terminal tagging, from n = 1 replicate, no changes p ≤ 0.05 from χ² test.
Extended Data Fig. 3 | Generation and validation of an ESB1 conditional knockout. A) Western blot validation of the cKO cell line and the BSF pDex577 tagged ESB1 exogenous expression cell line (the intermediate in cKO generation), both induced with 10 ng/ml doxycycline. Predicted molecular weights for ESB1 are: 108 kDa (untagged), 137 kDa (Ty::mNg tag) and 145 kDa (6×Ty::mNg tag). The uncropped Ponceau-stained membrane and anti-mNg blot are shown. B) Western blot validation of the BSF pDex577 tagged ESB1 exogenous expression cell line induced with 100 ng/ml doxycycline for overexpression. C) Validation of genetic modifications of the ESB1 conditional knockout (cKO). Schematics represent the deleted and tagged loci and primer binding sites and orientations, uncropped DNA gels shows the resulting PCR products from extracted genomic DNA. D) mRNA abundance in the BSF pDex577 tagged ESB1 exogenous expression cell line, plotting RPKM of uniquely mapped RNAseq reads. The exogenous expression prior to addition of doxycycline (0 h) is plotted relative to the parental pJ1339 cell line. Other plots are 12, 24 and 48 h after addition of 10 ng/ml doxycycline relative to the cell line grown without doxycycline. Each shows n = 1 induction replicate. E) mRNA abundance in the cKO, plotting RPKM of uniquely mapped RNAseq reads. The cKO prior to doxycycline washout (0 h) is plotted relative to the BSF pDex577 tagged ESB1 exogenous expression cell line induced with 10 ng/ml doxycycline. Other plots are 12, 24 and 48 h after doxycycline washout relative to the cell line grown with 10 ng/ml doxycycline. Each shows n = 1 induction replicate. F) mRNA abundance in the procyclic form KO, plotting RPKM of uniquely mapped RNAseq reads for three clonal KO cell lines relative to the parental cell line.
Extended Data Fig. 4 | Extended presentation of ChIP data showing the active and inactive BESs. ESB1 ChIP-seq in BSFs shown as the ratio of ChIP to input DNA in 2 kb bins across the active BES (BES1) and all inactive BESs. Unanalysable bins had insufficient uniquely-mapped reads from the input DNA. Extended version of Fig. 2K. n = 2 replicates, data points represent individual replicates and bars represent the mean.
Extended Data Fig. 5 | RNAi knockdown confirms ESB1 is vital and required for active BES transcription. Cellular phenotype of ESB1 RNAi knockdown. A) Growth curve of the ESB1 RNAi following doxycycline induction (+Dox) in comparison to uninduced (−Dox), using repeated subculture to maintain culture density under ~1×10⁶ cells/ml. Mean ± SD from n = 3 inductions. p shown at 48 h, two-tailed t test, log cumulative growth. B) Number of kinetoplasts (K) and nuclei (N) per cell, counted by light microscopy, at 24 h intervals following washout of doxycycline from the ESB1 cKO. Representative example from n = 3 inductions. p from χ² test. 1K1N, 2K1N and 2K2N are normal cell cycle stages. C) Representative images from n = 3 independent inductions of the ESB1 RNAi cell line before and after induction, showing a morphologically normal (1K1N) cell and a typical abnormal cell after 24 h induction. mNg signal is not detectable after 24 h induction. D) Uncropped anti-mNg Western blot validation of the ESB1 RNAi cell line. E) mRNA abundance in the BSF ESB1 RNAi cell line, plotting RPKM of uniquely mapped RNAseq reads. The uninduced cell line (0 h) is plotted relative to the parental 6×Ty::mNg::ESB1 cell line. Other plots are 12 and 24 after addition of 1 mg/ml doxycycline relative to the cell line grown without doxycycline. Each represents n = 1 induction replicate. F) Volcano plot of change mRNA abundance determined by RNAseq 24 h +Dox, n = 4 inductions. p from two-tailed t test. A-BES and I-BES indicate active and inactive BES respectively. G) qRT-PCR measurement of active BES VSG mRNA relative to the parental cell line, mean ± SD from n = 3 inductions, p ≤ 0.05 shown from two-tailed t test. H) Change in RNAseq read coverage over the ESB1 open reading frame shows reduced ESB1 transcript, mean from n = 4 inductions. I) Average change in transcript abundance averaged per BES and J) per gene for the active BES plotted by distance from the promoter 48 h −Dox, n = 4 inductions, mean ± SD from n = 4 inductions. K) Correlation of per mRNA Z scores for the bloodstream ESB1 RNAi cell line with the ESB1 cKO after 24 h induction. Upregulated genes are labelled, and are ER stress associated.
Extended Data Fig. 6 | Extended analysis of the bloodstream and procyclic form overexpression analysis. A) mRNA abundance in the bloodstream form ESB1 overexpression cell line, plotting RPKM of uniquely mapped RNAseq reads. The overexpression prior to doxycycline addition (0 h) is plotted relative to the parental pJ1339 cell line. Other plots are 12, 24 and 48 h after 100 ng/ml doxycycline addition relative to the cell line grown without doxycycline. Each represents n=1 induction replicate. B) Changes to total and unprocessed (not polyadenylated or not spliced) mRNA grouped into A-BES or I-BES VSg(s) and ESAgs for the overexpression +/−Dox. Mean ± SD from n=4 inductions. p ≤ 0.05 shown from two-tailed t test. C) Average change in transcript abundance averaged per BES and per gene for the inactive BESs plotted by distance from the promoter 12 h +Dox, n=4 inductions, mean ± SD. D) Correlation of per mRNA Z scores for the ESB1 overexpression, 12 h induction, with VEX1 RNAi, 72 h induction. Non-VSg and ESAg outliers are labelled. F,G) Anti-VSg221 immunofluorescence of the ESB1 overexpression line showing images of an example morphologically normal and abnormal cell after 48 h overexpression and G) counts of the proportion of VSg221-positive cells in comparison to the BSF and PCF cell lines. n numbers indicate number of cells counted from 1 induction replicate, no BSF changes p ≤ 0.05 from χ² test. H) mRNA abundance in the procyclic form ESB1 overexpression cell line, plotting RPKM of uniquely mapped RNAseq reads. The overexpression prior to doxycycline addition (0 h) is plotted relative to the parental pJ1339 cell line. Other plots are 12, 24 and 48 h after 1 mg/ml doxycycline addition relative to the cell line grown without doxycycline. Each represents n=1 induction replicate.
Extended Data Fig. 7 | ESB orthologs among kinetoplastid parasites. A) Fast approximately maximum-likelihood phylogenetic tree of ESB1 and its orthologs. Node values are SH-like support. B) ESB1 U-box RING finger domain compared to the canonical sequence showing a large insertion. C) Domain structure of ESB1 and its orthologs. From sequence alone, the C terminal domain could not be detected outside of trypanosomatids.
Extended Data Fig. 8 | Extended presentation of RNAseq data showing the active and inactive BESs. A,B) Read coverage of the active BES (BES1) in comparison to all sequenced inactive BESs, determined by RNAseq, for A) the BSF ESB1 cKO (characterised in Fig. 2A-F) 48 h after induction and B) the BSF ESB1 overexpression (characterised in Fig. 5A-F) 12 h after induction. The latter is an extended version of Fig. 6F. Mean of n = 4 inductions.
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| Study description | Briefly describe the study type including whether data are quantitative, qualitative, or mixed-methods (e.g. qualitative cross-sectional, quantitative experimental, mixed-methods case study). |
|-------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Research sample   | State the research sample (e.g. Harvard university undergraduates, villagers in rural India) and provide relevant demographic information (e.g. age, sex) and indicate whether the sample is representative. Provide a rationale for the study sample chosen. For studies involving existing datasets, please describe the dataset and source. |
| Sampling strategy | Describe the sampling procedure (e.g. random, snowball, stratified, convenience). Describe the statistical methods that were used to predetermine sample size OR if no sample-size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient. For qualitative data, please indicate whether data saturation was considered, and what criteria were used to decide that no further sampling was needed. |
| Data collection    | Provide details about the data collection procedure, including the instruments or devices used to record the data (e.g. pen and paper, computer, eye tracker, video or audio equipment) whether anyone was present besides the participant(s) and the researcher, and whether the researcher was blind to experimental condition and/or the study hypothesis during data collection. |
| Timing             | Indicate the start and stop dates of data collection. If there is a gap between collection periods, state the dates for each sample cohort. |
Data exclusions

If no data were excluded from the analyses, state so OR if data were excluded, provide the exact number of exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.

Non-participation

State how many participants dropped out/declined participation and the reason(s) given OR provide response rate OR state that no participants dropped out/declined participation.

Randomization

If participants were not allocated into experimental groups, state so OR describe how participants were allocated to groups, and if allocation was not random, describe how covariates were controlled.

Ecological, evolutionary & environmental sciences study design

All studies must disclose on these points even when the disclosure is negative.

Study description

Briefly describe the study. For quantitative data include treatment factors and interactions, design structure (e.g. factorial, nested, hierarchical), nature and number of experimental units and replicates.

Research sample

Describe the research sample (e.g. a group of tagged Passer domesticus, all Stenocereus thurberi within Organ Pipe Cactus National Monument), and provide a rationale for the sample choice. When relevant, describe the organism taxa, source, sex, age range and any manipulations. State what population the sample is meant to represent when applicable. For studies involving existing datasets, describe the data and its source.

Sampling strategy

Note the sampling procedure. Describe the statistical methods that were used to predetermine sample size OR if no sample-size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient.

Data collection

Describe the data collection procedure, including who recorded the data and how.

Timing and spatial scale

Indicate the start and stop dates of data collection, noting the frequency and periodicity of sampling and providing a rationale for these choices. If there is a gap between collection periods, state the dates for each sample cohort. Specify the spatial scale from which the data are taken.

Data exclusions

If no data were excluded from the analyses, state so OR if data were excluded, describe the exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.

Reproducibility

Describe the measures taken to verify the reproducibility of experimental findings. For each experiment, note whether any attempts to repeat the experiment failed OR state that all attempts to repeat the experiment were successful.

Randomization

Describe how samples/organisms/participants were allocated into groups. If allocation was not random, describe how covariates were controlled. If this is not relevant to your study, explain why.

Blinding

Describe the extent of blinding used during data acquisition and analysis. If blinding was not possible, describe why OR explain why blinding was not relevant to your study.

Field work, collection and transport

Field conditions

Describe the study conditions for field work, providing relevant parameters (e.g. temperature, rainfall).

Location

State the location of the sampling or experiment, providing relevant parameters (e.g. latitude and longitude, elevation, water depth).

Access & import/export

Describe the efforts you have made to access habitats and to collect and import/export your samples in a responsible manner and in compliance with local, national and international laws, noting any permits that were obtained (give the name of the issuing authority, the date of issue, and any identifying information).

Disturbance

Describe any disturbance caused by the study and how it was minimized.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.
Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| ☒ Antibodies |
| ☒ Eukaryotic cell lines |
| ☐ Palaeontology and archaeology |
| ☐ Animals and other organisms |
| ☐ Clinical data |
| ☐ Dual use research of concern |

Methods

| n/a | Involved in the study |
|-----|-----------------------|
| ☐ ChIP-seq |
| ☠ Flow cytometry |
| ☐ MRI-based neuroimaging |

Antibodies

| Antibodies used |
|----------------|
| Anti-mNG ChromoTek 32f6, anti-TY, anti-GFP Invitrogen A11122, anti-VSG221 |

Validation

- Anti-mNG: Validated by manufacturer with mNG-Actin and no cross interaction with other fluorescent proteins confirmed by the manufacturer.
- Anti-TY: Generated and validated previously in this research group, not a commercial antibody. Positive and negative controls through genetic modifications included in our western blot analysis in Extended Data Figure S3.
- Anti-GFP: Validated by manufacturer with H3-GFP, P65-GFPm and no cross interaction with mCh confirmed by the manufacturer.
- Anti-VSG221: Previously published polyclonal anti-VSG221 antibody, not a commercial antibody. Positive and negative controls through analysing VSG221 and non-VSG221 expressing life stages in Extended Data Fig S6.

Eukaryotic cell lines

Policy information about [cell lines and Sex and Gender in Research](#)

| Cell line source(s) |
|---------------------|
| Trypanosoma brucei brucei Lister 427 and TREU 927, laboratory long-term stocks |

| Authentication |
|----------------|
| RNA sequencing comparison to predicted CDSs of reference genomes on TriTrypDB.org |

| Mycoplasma contamination |
|--------------------------|
| No mycoplasma testing |

| Commonly misidentified lines |
|-------------------------------|
| N/A (See [CLAC register](#)) |

Palaeontology and Archaeology

| Specimen provenance |
|---------------------|
| Provide provenance information for specimens and describe permits that were obtained for the work (including the name of the issuing authority, the date of issue, and any identifying information). Permits should encompass collection and, where applicable, export. |

| Specimen deposition |
|---------------------|
| Indicate where the specimens have been deposited to permit free access by other researchers. |

| Dating methods |
|----------------|
| If new dates are provided, describe how they were obtained (e.g. collection, storage, sample pretreatment and measurement), where they were obtained (i.e. lab name), the calibration program and the protocol for quality assurance OR state that no new dates are provided. |

- Tick this box to confirm that the raw and calibrated dates are available in the paper or in Supplementary Information.

Ethics oversight

- Identify the organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or guidance was required and explain why not.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Animals and other research organisms

Policy information about [studies involving animals; ARRIVE guidelines](#) recommended for reporting animal research, and [Sex and Gender in Research](#)

| Laboratory animals |
|--------------------|
| For laboratory animals, report species, strain and age OR state that the study did not involve laboratory animals. |

| Wild animals |
|--------------|
| Provide details on animals observed in or captured in the field; report species and age where possible. Describe how animals were caught and transported and what happened to captive animals after the study (if killed, explain why and describe method; if released, say where and when) OR state that the study did not involve wild animals. |

| Reporting on sex |
|-----------------|
| Indicate if findings apply to only one sex; describe whether sex was considered in study design, methods used for assigning sex. Provide data disaggregated for sex where this information has been collected in the source data as appropriate; provide overall |
Field-collected samples: For laboratory work with field-collected samples, describe all relevant parameters such as housing, maintenance, temperature, photoperiod and end-of-experiment protocol OR state that the study did not involve samples collected from the field.

Ethics oversight: Identify the organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or guidance was required and explain why not.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

**Clinical data**

Policy information about [clinical studies](#). All manuscripts should comply with the ICMJE [guidelines for publication of clinical research](#) and a completed [CONSORT checklist](#) must be included with all submissions.

**Clinical trial registration**: Provide the trial registration number from ClinicalTrials.gov or an equivalent agency.

**Study protocol**: Note where the full trial protocol can be accessed OR if not available, explain why.

**Data collection**: Describe the settings and locales of data collection, noting the time periods of recruitment and data collection.

**Outcomes**: Describe how you pre-defined primary and secondary outcome measures and how you assessed these measures.

**Dual use research of concern**

Policy information about [dual use research of concern](#).

**Hazards**

Could the accidental, deliberate or reckless misuse of agents or technologies generated in the work, or the application of information presented in the manuscript, pose a threat to:

| Yes | No |
|-----|----|
| ☑   |    |
| Public health |
| National security |
| Crops and/or livestock |
| Ecosystems |
| Any other significant area |

**Experiments of concern**

Does the work involve any of these experiments of concern:

| Yes | No |
|-----|----|
| ☑   |    |
| Demonstrate how to render a vaccine ineffective |
| Confer resistance to therapeutically useful antibiotics or antiviral agents |
| Enhance the virulence of a pathogen or render a nonpathogen virulent |
| Increase transmissibility of a pathogen |
| Alter the host range of a pathogen |
| Enable evasion of diagnostic/detection modalities |
| Enable the weaponization of a biological agent or toxin |
| Any other potentially harmful combination of experiments and agents |

**ChIP-seq**

**Data deposition**

- Confirm that both raw and final processed data have been deposited in a public database such as [GEO](#).
- Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

**Data access links**

May remain private before publication.

NCBI BioProject PRJNA784098

Files in database submission

MiChIP3_1.fq.gz MiChIP4_1.fq.gz MiChIP19_1.fq.gz MiChIP20_1.fq.gz
**Methodology**

| Replicates                  | 2 biological replicates, carried out > 6 months apart, from clonal cell lines expressing eYFP-tagged target protein. |
|-----------------------------|-------------------------------------------------------------------------------------------------------------------|
| Sequencing depth            | ~100 million 50 bp single end reads for all experiments. ~1% of reads were uniquely mapped. Note that low mapping rate is expected due to the repetitive nature of the DNA encoding and flanking VSGs and ESAGs. |
| Antibodies                  | anti-GFP Invitrogen A11122 (no clone name available)                                                                |
| Peak calling parameters     | Peaks were not called. Standard tools are not applicable due to the repetitive nature of the enriched DNA. Analysis was of mapped reads binned into 2kb bins, carried out in Excel. See Figure 2J, K and Extended Data Figure S4 for custom analysis details. |
| Data quality                | Peaks were not called. Standard tools are not applicable due to the repetitive nature of the enriched DNA. Enrichment specificity is shown in Figure 2J and Extended Data S4. |
| Software                    | BWA-MEM 0.7.17, samtools 1.7, Excel                                                                                  |