

Antigenic diversity is generated by distinct evolutionary mechanisms in African trypanosome species

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Antigenic variation enables pathogens to avoid the host immune response by continual switching of surface proteins. The protozoan blood parasite *Trypanosoma brucei* causes human African trypanosomiasis (“sleeping sickness”) across sub-Saharan Africa and is a model system for antigenic variation, surviving by periodically replacing a monolayer of variant surface glycoproteins (VSG) that covers its cell surface. We compared the genome of *Trypanosoma brucei* with two closely related parasites *Trypanosoma congolense* and *Trypanosoma vivax*, to reveal how the variant antigen repertoire has evolved and how it might affect contemporary antigenic diversity. We reconstruct VSG diversification showing that *Trypanosoma congolense* uses variant antigens derived from multiple ancestral VSG lineages, whereas in *Trypanosoma brucei* VSG have recent origins, and ancestral gene lineages have been repeatedly co-opted to novel functions. These historical differences are reflected in fundamental differences between species in the scale and mechanism of recombination. Using phylogenetic incompatibility as a metric for genetic exchange, we show that the frequency of recombination is comparable between *Trypanosoma congolense* and *Trypanosoma brucei* but is much lower in *Trypanosoma vivax*. Furthermore, in showing that the C-terminal domain of *Trypanosoma brucei* VSG plays a crucial role in facilitating exchange, we reveal substantial species differences in the mechanism of VSG diversification. Our results demonstrate how past VSG evolution indirectly determines the ability of contemporary parasites to generate novel variant antigens through recombination and suggest that the current model for antigenic variation in *Trypanosoma brucei* is only one means by which these parasites maintain chronic infections.

Antigenic variation enables pathogens to evade immune responses by continual switching of surface proteins (1, 2). The African trypanosomes (*Trypanosoma* spp.) are vector-borne protozoan blood parasites that survive in their hosts by antigenic variation, periodically replacing a monolayer of variant surface glycoproteins (VSG) (3) that shield the cell surface from immune effectors (4, 5). *Trypanosoma brucei* is the cause of human African trypanosomiasis (or “sleeping sickness”), and the mechanisms for expression and dynamic replacement of VSG in this species are a model system for antigenic variation (4) as well as a classic example of adaptive evolution at the host–pathogen interface. Two related veterinary parasites, *Trypanosoma congolense* and *Trypanosoma vivax*, also use antigenic variation to cause devastating diseases in domesticated animals. Through their detrimental effects on livestock productivity, these species arguably represent greater threats to socioeconomic well-being than *T. brucei* does in the agrarian societies in which they are endemic. Our understanding of how antigenic diversity is organized in *T. brucei* was greatly improved by the *T. brucei* 927 reference genome sequence (6). In this paper, we present draft genome sequences for *T. congolense* and *T. vivax*; we define their global VSG repertoires in a three-way comparative

analysis with *T. brucei*, revealing how antigenic diversity evolved in trypanosome genomes past and present.

The *T. brucei* genome includes many hundreds of VSG that encode a transcriptionally silent reservoir of variant antigens (6), and each cell expresses just a single gene from a specialized telomeric expression site at any time (4, 5). The parasite population collectively express multiple VSG; when the host becomes immune to the prevailing VSG, clones expressing alternative copies proliferate in a frequency-dependent manner, maintaining the infection and resulting in characteristic “waves of parasitaemia.” To survive long-term, *T. brucei* must generate novel VSG sequences through recombination; mechanisms may include domain shuffling (7) and gene conversion among silent, subtelomeric gene copies or possibly in situ within the expression site (8). Functional variant antigens in *T. brucei* consist of a- and b-type VSG (hereafter a-VSG and b-VSG), which share the cysteine-rich C-terminal domain (CTD) but are otherwise distantly related (9–11). Although VSG are known to occur in *T. congolense* and *T. vivax* (12–16), the repertoire of variant antigens in these species is uncharacterized. Consequently, the evolutionary diversification of the VSG gene family has not been examined, although it has been suggested that VSG are a source of novel genes. Two gene families, the expression site-associated genes (*ESAG6/7*) encoding transferrin receptor (TFR) and the VSG-related (*VR*) genes, are thought to have evolved from a-VSG (17, 18) and b-VSG (8, 11), respectively.

The antigenic variation phenotype is observed in all African trypanosomes, and it is assumed that this reflects a common physiological model, which has been defined in *T. brucei*. The aim of this study is to identify the evolutionary processes that have created contemporary VSG diversity and reveal any significant differences in how trypanosome species generate variant antigens.

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Data deposition: The draft genome sequences reported in this paper have been deposited in the European Molecular Biology Laboratory (EMBL) Nucleotide Sequence Database [accession nos. HE575314–HE575324 and CAEQ1000352–CAEQ1002824 (*Trypanosoma congolense*) and HE573017–HE573027 and CAEX01000001–CAEX01008277 (*Trypanosoma vivax*)]. The data can be examined via GeneDB (<http://www.genedb.org>) and TriTrypDB (<http://tritrypdb.org>). *T. vivax* transcriptome data have been submitted to the European Bioinformatics Institute Array Express Archive (accession no. E-MTAB-475). Sequence alignments and phylogenetic trees comprising the cell-surface phylome are contained in GeneDB (http://www.genedb.org/Page/trypanosoma_surface_phylome).

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Despite their shared phenotype, our results show that species differ in the organization of antigenic diversity at the genome level, and they provide a basis to better understand disease progression, pathology, and host range in all African trypanosomes.

Results

VSG Gene Repertoires of *T. congolense* and *T. vivax*. We have produced high-quality draft genome sequences for *T. congolense* IL3000, a sister species of *T. brucei*, and *T. vivax* Y486, a third species that branches close to the root of the African trypanosome lineage (19). These genome sequences are described in Table S1 and are accessible through GeneDB (<http://www.genedb.org>) or TriTrypDB (<http://tritrypdb.org>). Comparative analysis including the existing *T. brucei* 927 genome sequence shows that the principal differences in genome content relate to cell-surface architecture (Tables S2–S4). To define VSG repertoires, gene sequences with predicted cell-surface roles were extracted from all three genomes and sorted with BLASTx, resulting in 81 gene families (Materials and Methods and Table S5). Phylogenies of these families were estimated that we collectively termed the “cell-surface phylome,” (http://www.genedb.org/Page/trypanosoma_surface_phylome). The phylome contains VSG and related families already known in *T. brucei* but it also defines families that we believe encode the VSG repertoires of *T. congolense* and *T. vivax*.

The *T. congolense* VSG repertoire differs from that of *T. brucei* in three ways: First, there is no a-VSG subfamily of variant antigens; second, there are two b-VSG subfamilies, termed Fam13 ($n = 302$) and Fam16 ($n = 512$) by their phylome designations; and, third, unlike *T. brucei* VSG, which all share a relatively uniform CTD, *T. congolense* VSG have 15–20 different CTD types, each associated with a specific subset of Fam13 or Fam16 and none homologous to the *T. brucei* CTD. Hence, *T. congolense* b-VSG are more structurally heterogeneous than *T. brucei* b-VSG are. We know that both Fam13 and Fam16 contain functional variant antigens because each family encompasses both published *T. congolense* VSG (12–14) and VSG expressed sequence tags (15). Although there is no a-VSG variant antigen, there are homologs of the a-VSG-like TFR genes of *T. brucei*, i.e., *ESAG6/7* (Fam15; $n = 43$), as well as the procyclin-associated genes (*PAG*) (Fam14, $n = 22$), which also have an a-VSG-like structure (20).

VSG structural diversity is even greater in *T. vivax*. We have identified four VSG subfamilies (Fam23–26) that each possess definitive patterns of conserved cysteine residues (SI Results). Fam23 ($n = 540$) and Fam24 ($n = 279$) members possess sequence motifs homologous to a-VSG and b-VSG, respectively. Fam25 ($n = 227$) and Fam26 ($n = 87$) are two subfamilies unique to *T. vivax* but with low (~20%) protein sequence similarity to known VSG (Fig. S1). These specific subfamilies may have evolved in *T. vivax* or may represent ancestral lineages not inherited by *T. brucei* and *T. congolense*. Transcriptomic data show that multiple members of all four families are transcribed in bloodstream-stage cells (Table S6). We find no orthologs to the TFR-like genes of *T. brucei* and *T. congolense* among *T. vivax* VSG-like genes or indeed the numerous *T. vivax*-specific gene families.

Amino acid sequence homology with *T. brucei* VSG alone does not guarantee that putative *T. vivax* VSG function as variant antigens. To date, only one *T. vivax* VSG [ILDat 2.1 (16)] has been characterized, albeit from a dissimilar strain, and it is most closely related to Fam26. Therefore, we identified an expressed VSG in the genome strain Y486 by MS analysis of a protein specific to a relapsed infection population, peptide fragments of which are 100% identical to a predicted protein in Fam23 (TvY486_0027060; Fig. S2). Therefore, at least one a-VSG-like (i.e., Fam23) gene in *T. vivax* encodes a functional variant antigen.

Phylogeny of VSG Diversification. In total, the three genome sequences yielded 1,083 a-VSG-like and 1,537 b-VSG-like full-length genes (Table S7). We estimated Bayesian and maximum likelihood phylogenies from amino acid sequence alignments (Figs. S3 and S4); however, given the large number of sequences and to enable global visualization, we also estimated a similarity network from pairwise maximum likelihood protein distances that delivered a clearer picture of relationships within the a-VSG and b-VSG

lineages. The distance network includes examples of all VSG subfamilies and represents individual genes as spheres connected to others sharing identity above a threshold (Materials and Methods). The network and phylogenies are fully consistent. Fig. 1 shows the similarity network from two angles, and Movie S1 contains an animation of the 3D network; four principal features emerge.

First, the common CTD of *T. brucei* VSG must have evolved through horizontal transfer from one subfamily to the other. In Fig. 1, sequences cluster by lineage (a or b) rather than by species; for instance, *T. vivax* a-VSG (Fam23) is more similar to a-VSG-like subfamilies in *T. brucei* and *T. congolense* than to *T. vivax* b-VSG (Fam24). Therefore, VSG lineages are older than the genomes they occupy; indeed, they were present in the common ancestor of all African trypanosomes. The only above-threshold sequence connections occurring between a-VSG and b-VSG subfamilies (Fig. 1, *i*) concern *T. brucei* VSG and, in particular, their common CTD. This is a unique feature of *T. brucei* VSG and presents an interesting anomaly: despite belonging to ancient lineages separated in the ancestral trypanosome, a-VSG and b-VSG in *T. brucei* share a CTD that is species-specific, which can only be explained if the CTD evolved in one subfamily and was transposed to the other.

Second, b-VSG in *T. brucei* are derived from a single ancestral lineage, whereas *T. congolense* b-VSG are drawn from many lineages, suggesting that *T. brucei* b-VSG have passed through a “bottleneck.” In Fig. 1, all b-VSG in *T. brucei* (dark blue) form a cluster to the exclusion of other subfamilies. Hence, they share a recent common ancestor that evolved after the split from *T. congolense*. In contrast, *T. congolense* b-VSG comprise two lineages (Fam13 and Fam16) that originated in the *T. brucei*/*T. congolense* ancestor and form separate clusters in the network (Fig. 1, *ii*). We know that these lineages did not originate in *T. congolense* because their closest relatives are VSG-like genes in *T. brucei* (see below). In fact, Fam13 and Fam16 themselves split into multiple clusters in Fig. 1 (Fig. 1, *ii*), emphasizing the phylogenetic diversity of *T. congolense* VSG and relative homogeneity in *T. brucei*.

Third, VSG have repeatedly been a source of functional novelty on the cell surface. We know that VSG can be co-opted from variant antigen functions to novel roles, for example, the serum-resistance antigen [*SRA* (21)] and *Trypanosoma brucei gambiense*-specific glycoprotein [*TgsGP* (22)] proteins in *Trypanosoma brucei rhodesiense* and *T. b. gambiense*, respectively. However, SRA and tgsGP represent secondary loss of function among contemporary VSG. Fig. 1 shows that *ESAG2*, a gene family associated with the polycistronic VSG expression site in *T. brucei*, is a b-VSG-like gene, nested among *T. congolense* b-VSG (Fam13; Fig. 1, *iii*). Similarly, VR genes (purple in Fig. 1), rather than being derived from b-VSG in *T. brucei*, have an ancestral-type structure more akin to Fam16 in *T. congolense*. We have also identified another *T. brucei*-specific family (Fam1; pink in Fig. 1) that encodes proteins homologous to b-VSG, with a predicted GPI anchor but also a highly modified CTD. Fam1 (i.e., Tb927.6.1310) is preferentially expressed in bloodstream forms and localizes to the flagellar pocket and endosomal membranes (Fig. S5). Phylogenetic analysis clearly demonstrates that both *ESAG2* and VR gene subfamilies, for which the evidence is against a variant antigen function (8, 11), are not recent derivations from *T. brucei* VSG (like *SRA* and *TgsGP*) but belong to ancestral VSG lineages with representatives in *T. congolense* that still encode functional variant antigens (Fig. S4). Hence, some of the ancestral lineages in *T. congolense* identified above remain in *T. brucei* but have been co-opted to novel roles.

Finally, the network indicates that the TFR evolved from an a-VSG gene, as suggested previously (8, 23). However, the functional transition did not occur within the *T. brucei* VSG expression site but instead in the *T. brucei*/*T. congolense* ancestor. A tight cluster of TFR-like genes (i.e., *ESAG6/7* and *PAG*) from *T. brucei* as well as Fam14 and Fam15 sequences from *T. congolense* are distinct from other a-VSG subfamilies in Fig. 1 (Fig. 1, *iv*). The similarity network reflects their phylogeny, which shows that Fam14 and Fam15 are sister lineages to *PAG* and *ESAG6/7*, respectively, and their primary structures, which show that amino acid residues crucial for transferrin binding (18) are conserved in both species (Fig. S6). Given the absence of this entire family from *T. vivax*, we conclude that, rather than being loss from that species, the

a-VSG lineage

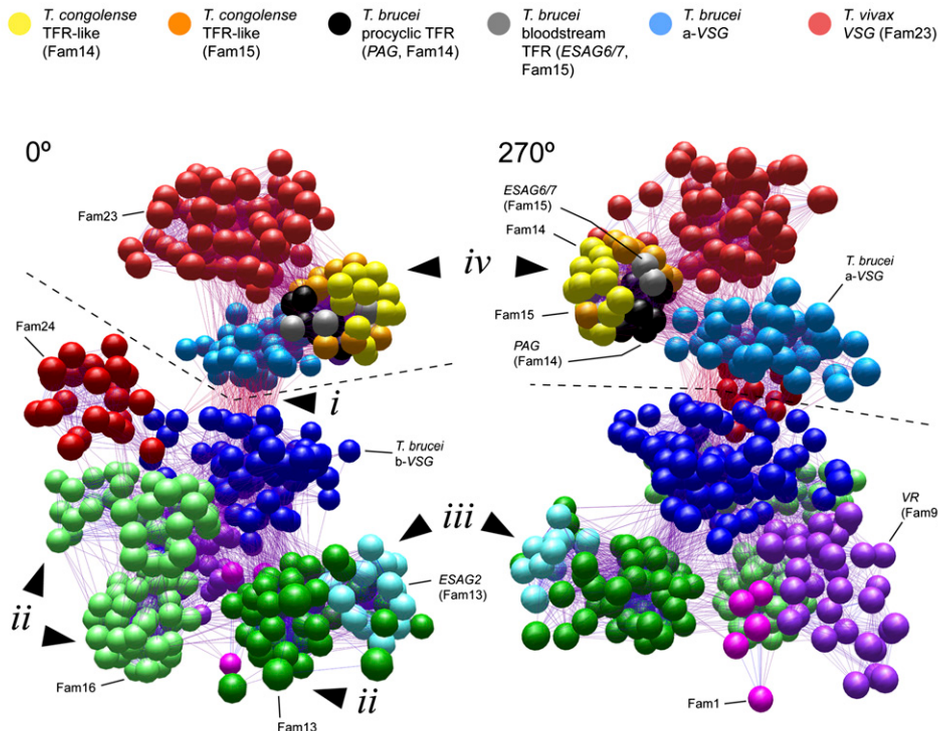


Fig. 1. A sequence-similarity network of VSG-like sequences from African trypanosome genomes, shown from 0° and 270° angles. A 3D rendering of the network is provided as [Movie S1](#). The network represents pairwise maximum likelihood protein distances, generated from multiple alignments of selected a-VSG-like (a-VSG, Fam23, TFR-like, and PAG-like proteins; $n = 174$) and b-VSG-like (b-VSG, Fam13, Fam16, Fam24, VR, ESAG2, and Fam1 proteins; $n = 339$) protein sequences, which are representative of global diversity. Spheres represent individual sequences shaded according to subfamily. A dashed line separates a-VSG subfamilies (above the line) and b-VSG subfamilies (below the line). Four significant features identified in the text are labeled: (i) sequence similarity between a-VSG and b-VSG lineages caused by the shared CTD of *T. brucei* VSG; (ii) diverse clusters of *T. congolense* b-VSG belonging to Fam13 and Fam16; (iii) the position of ESAG2 nested within Fam13; and (iv) tight cluster of TFR-like genes from both *T. brucei* and *T. congolense*.

b-VSG lineage

transferrin-receptor genes evolved before the separation of *T. brucei* and *T. congolense* but after their split from *T. vivax*. Our conclusion does not preclude other *T. vivax*-specific proteins performing a transferrin-binding function in that species.

These results are summarized in a model of VSG evolution (Fig. 2). The ancestral African trypanosome possessed a-VSG and b-VSG type genes that probably formed multigene families or functioned as variant antigens. Both VSG types were inherited by *T. vivax*, which retains an a-VSG family that includes functional variant antigens. The *T. brucei*/*T. congolense* ancestor inherited both a-VSG and b-VSG lineages, and, at this point, one a-VSG gene was co-opted to a transferrin-binding role differentiated between insect and vertebrate life stages, founding a lineage that was inherited by both daughter species. Another a-VSG lineage retained its variant antigen function in *T. brucei* but was lost from *T. congolense* (SI Results). Of the ancestral b-VSG repertoire, two different lineages have been inherited by both species: The first has produced ESAG2 and Fam13 in *T. brucei* and *T. congolense*, respectively, whereas the second has produced b-VSG and VR in *T. brucei* and Fam16 in *T. congolense*. There is no step in this deduced scheme where a trypanosome lacks variant antigen. Clearly, these two species have adapted their common legacy differently. *T. congolense* VSG are drawn from multiple ancestral lineages, whereas *T. brucei* has relegated corresponding genes (VR, ESAG2, and perhaps Fam1) to novel roles and derives its variant antigens from single lineages derived after speciation. This difference in the phylogenetic diversity of VSG repertoires is important because it could affect the ability of the parasites to present novel antigens to their hosts and therefore to maintain infection.

Tree Shape and Distribution of VSG Sequence Variation. We examined the phylogenies of VSG subfamilies within species for evidence that their distinct evolutionary legacies have affected

contemporary sequence evolution. Fig. 3 demonstrates how these trees have distinct topologies because of variation in the ratio of internal to terminal branches [described by “tree-ness” (24), T], which is low for *T. brucei* ($T = 0.282$ and 0.275), higher for *T. congolense* ($T = 0.376$ and 0.412), and highest for *T. vivax* ($T = 0.681$ and 0.763). *T. congolense* and *T. vivax* trees are more “tree-like” because they retain information about the past in basal nodes and internal branches, whereas the *T. brucei* tree consists mostly of long, terminal branches. Fig. 3 also compares the distribution of VSG sequence variation, showing that *T. brucei* distances have much narrower distributions than either *T. congolense* or *T. vivax* VSG do because both short, terminal branches and long, basal internodes are rare. Importantly, these patterns are genome-specific rather than lineage-specific effects, i.e., a-VSG and b-VSG in *T. brucei* display the same dynamic despite having greater identity with subfamilies in other species. They confirm that the mechanisms for antigenic variability vary between species now and likewise in the past.

Recombination is a principal evolutionary pressure affecting *T. brucei* VSG (5, 8), and exchange of the unique VSG CTD is well recorded (7, 11). Recombination is also the mechanism through which VSG are transposed from subtelomeric loci into the telomeric expression site (4, 5, 8). *T. brucei* VSG phylogenies in Fig. 3 are consistent with frequent recombination, but the cladistic structure of *T. congolense* and *T. vivax* VSG phylogenies could only persist if recombination between clades is rare. Furthermore, the incidence of pseudogenes, which result, at least partly, from gene conversion between VSG genes (5), is much lower in *T. congolense* (where only 21.1% of Fam13 and 29.7% of Fam16 are predicted pseudogenes) and *T. vivax* (15.5% and 27.2% of Fam23 and Fam24, respectively) than in *T. brucei* (69.2% of a-VSG and 72.2% of b-VSG) (6). Therefore, we

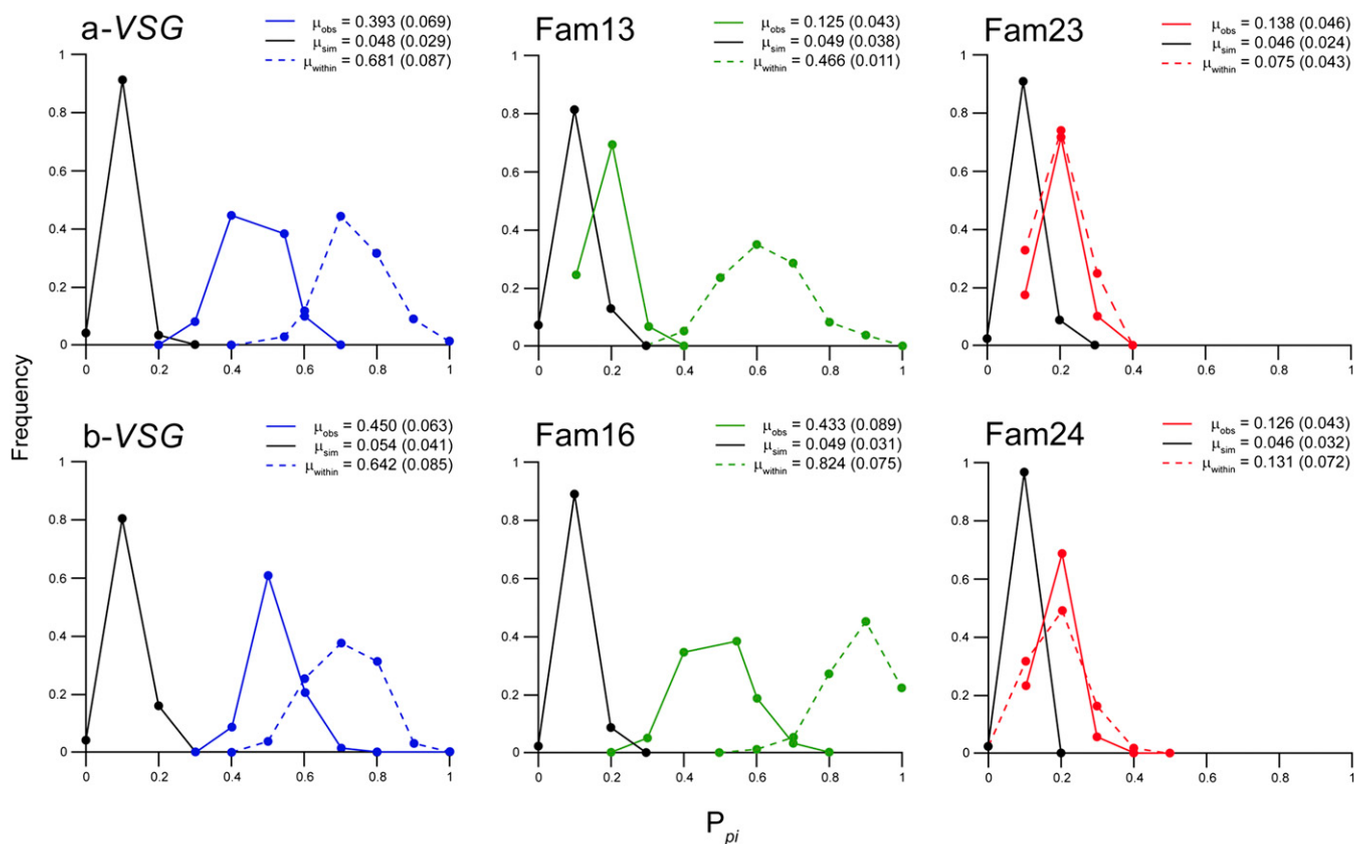


Fig. 4. Prevalence of significant PI within *VSG*-like sequence alignments. PI describes conflicting phylogenetic signals caused by recombination but also rate heterogeneity or other molecular homoplasy (25). Protein sequence alignments for six *VSG* subfamilies were examined for PI using the pairwise homoplasy index (26). Each alignment was randomly sampled 100 times, and the proportion displaying significant PI was counted (P_{pi}). A distribution for P_{pi} was generated by creating 100 bootstrapped alignments in each case (solid colored lines). To generate a null distribution, 100 alignments were simulated from the observed Bayesian phylogeny using a maximum likelihood substitution model (WAG+ Γ) that corrected for rate heterogeneity but did not consider recombination (black lines). To demonstrate the effect of genetic distance on PI, the analysis was repeated on smaller alignments of closely related sequences taken from crown clades (dashed colored lines; *Materials and Methods*). Mean average values, followed by SDs, are provided for observed (μ_{obs}), simulated (μ_{sim}), and within-clade (μ_{within}) sampling distributions.

Discussion

The past and present evolution of *VSG* can now be brought together. We have shown that the composition of contemporary *VSG* repertoires is determined by how each species has modified the common inheritance. *T. vivax* has the most structurally diverse repertoire, comprising a-VSG, b-VSG, and two additional types absent elsewhere; *T. congolense* combines multiple ancestral b-VSG lineages, each with a distinct CTD; and *T. brucei* a-VSG and b-VSG are recently derived single lineages with a common CTD. It is worth remembering that sequence mosaics generated in *T. brucei* infections are an additional source of diversity among expressed *VSG* (11); it is not known whether this dynamic assortment of *VSG* sequences occurs in other species. Nevertheless, as a result of compositional differences, the scale of recombination varies between species, being more frequent among *T. brucei* and *T. congolense* *VSG* than in *T. vivax*, and more prevalent among *T. brucei* *VSG* than in *T. congolense*. Opportunities for allelic recombination among *VSG* may be affected by species differences in mating system; recent work indicates that sexual reproduction is frequent in *T. brucei* and *T. congolense* populations (28–29), whereas *T. vivax* population structure is consistent with clonal reproduction (30). However, the importance of allelic recombination to *VSG* diversity is debatable; because the subtelomeres of homologous chromosomes in *T. brucei* are frequently dissimilar, subtelomeric *VSG* are effectively hemizygous, and so most recombination between *VSG* is ectopic (11). This study shows that PI in *T. brucei* *VSG* is attributable in large part to the CTD promoting exchange throughout the

repertoire, whereas the conservative CTDs of *T. congolense* *VSG* actually reduce the scale of PI and illustrate the lack of recombination between clades, emphasizing the role of gene structure in promoting ectopic recombination in mitosis rather than the frequency of allelic recombination during meiosis.

Differences in the role of the CTD indicate that, in addition to scale, the mechanism of recombination varies between species. The CTD is exchanged between *T. brucei* *VSG* but is not exposed to antibodies and therefore may not directly contribute to antigenic diversity (31). However, the CTD-encoding sequence may have a role in the duplicative transposition of *VSG*, which promotes the diversification of the silent archive (11) and is of paramount importance to antigenic variation because it is required to move *VSG* from silent, subtelomeric loci into the telomeric expression site (4, 8). It is thought that *VSG* transposition is facilitated by an upstream 70-bp repeat region and the conserved CTD (8), with the transposed region terminating around the 3' end of the coding sequence (32). Hence, in 37 recent such duplicative events among silent *VSG* in the *T. brucei* 927 genome (11), 16 terminated within the CTD region. In showing that the majority of PI in *T. brucei* *VSG* alignments concerns the CTD, our data demonstrate that a major recombination breakpoint occurs between the N-terminal domains and CTDs, supporting a principal role for the CTD in promoting *VSG* switching. Immediately, we can see that this mechanism cannot operate in *T. congolense*, where the CTDs are heterogeneous and have no role in promoting exchange. Hence, we propose that the preeminence of the CTD in PI among *T. brucei* *VSG* reflects the frequent transposition of N-terminal domains,

and, through its solitary CTD type, which originated uniquely through horizontal transfer between *VSG* lineages, *T. brucei* may have evolved a distinct mechanism for the *VSG* transpositions that are key for diversification and antigen switching.

Antigenic variation is central to the host–trypanosome relationship, intimately linked to the course and severity of disease, to parasite transmission and host range, and therefore to disease epidemiology. All African trypanosomes display antigenic variation, and, although the current *T. brucei*-based model might adequately describe the general phenomenon, this study shows that the genomic basis for antigenic variation has diverged among trypanosomes in a manner consistent with distinct mechanisms for generating antigenic variability. Consequently, we now have reason to expect substantial species differences beneath the general phenotype, a framework to dissect this variation, and so a basis for understanding how the enigmatic *VSG* connects with the wider disease.

Materials and Methods

Genome Sequencing and Annotation. *T. congolense* IL3000 and *T. vivax* Y486 genomes were capillary-sequenced with a whole-genome shotgun strategy as described previously (6).

Annotation of *VSG* Genes. *T. congolense* and *T. vivax* *VSG* were identified by BLASTp-based homology searches and hidden Markov models. The boundaries of all *VSG* ORFs were manually checked against global sequence alignments.

Comparison of Gene Content. OrthoMCL was used to examine putative gene gains and losses. All putative losses were confirmed by examining expected genomic position and by searching unassembled sequence reads for reciprocal sequence matches by tBLASTn/BLASTx.

***T. vivax* Transcriptome.** *T. vivax* Y486 was grown from stabilate in BALB/c mice immunosuppressed with cyclophosphamide and was amplified at patent parasitaemia in three immunosuppressed mice, from which whole blood

was collected. The blood was treated with the erythrocyte lysis buffer, and RNA was isolated from the pellet.

Analysis of *Fam1* Gene Expression. To determine mRNA expression levels of *Fam1* family members, quantitative real-time PCR (qRT-PCR) was carried out on total RNA.

Transfection and *Fam1* Protein Localization. Ectopic expression of HA epitope-tagged Tb927.6.1310 at the N terminus (after the predicted signal peptide sequence) was carried out with constitutive and inducible expression vectors.

***VSG* Purification and Sequencing.** *T. vivax* cell extracts were run in 1D SDS/PAGE, and three bands in the estimated size range were extracted from each, trypsinized, and subjected to liquid chromatography/tandem MS analysis.

Cell-Surface Phylome. Homologs to each *T. brucei* “surface” gene were identified among all *T. brucei*, *T. congolense*, *T. vivax*, and *Trypanosoma cruzi* predicted genes with wuBLAST. Gene family phylogenies were estimated with maximum likelihood and Bayesian methods.

Recombination Analysis. PI within *VSG* sequence alignments was used as a measure of recombination. The pairwise homoplasy index (26) returns a single probability value for PI, which was applied to amino acid sequence alignments for seven *VSG* subfamilies (Table S8), subsampled 1,000 times. The proportion of subalignments with significant PI, termed P_{pi} , was compared between species.

Additional information on methods is provided in *SI Materials and Methods*.

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