Deep kinetoplast genome analyses result in a novel molecular assay for detecting *Trypanosoma brucei gambiense*-specific minicircles

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ABSTRACT

The World Health Organization targeted *Trypanosoma brucei gambiense* (Tbg) human African trypanosomiasis for elimination of transmission by 2030. Sensitive molecular markers that specifically detect Tbg type 1 (Tbg1) parasites will be important tools to assist in reaching this goal. We aim at improving molecular diagnosis of Tbg1 infections by targeting the abundant mitochondrial minicircles within the kinetoplast of these parasites. Using Next-Generation Sequencing of total cellular DNA extracts, we assembled and annotated the kinetoplast genome and investigated minicircle sequence diversity in 38 animal- and human-infective trypanosome strains. Computational analyses recognized a total of 241 Minicircle Sequence Classes as Tbg1-specific, of which three were shared by the 18 studied Tbg1 strains. We developed a minicircle-based assay that is applicable on animals and as specific as the TgsGP-based assay, the current golden standard for molecular detection of Tbg1. The median copy number of the targeted minicircle was equal to eight, suggesting our minicircle-based assay may be used for the sensitive detection of Tbg1 parasites. Annotation of the targeted minicircle sequence indicated that it encodes genes essential for the survival of the parasite and will thus likely be preserved in natural Tbg1 populations, the latter ensuring the reliability of our novel diagnostic assay.

INTRODUCTION

Human African trypanosomiasis (HAT), also known as sleeping sickness, is a vector-borne disease caused by two *Trypanosoma brucei* (Tb) subspecies and transmitted by tsetse flies. *Trypanosoma brucei rhodesiense* (Tbr) causes acute infections in East Africa, whereas *Trypanosoma brucei gambiense* (Tbg) causes chronic infections in West and Central Africa (1). *Trypanosoma brucei rhodesiense* and Tbg type I (Tbg1) are defined by the presence of truncated variant surface glycoprotein (VSG) genes, respectively the serum-resistance-associated (SRA) gene (2) and the Tbg-specific glycoprotein (TgsGP) gene (3) that play a role in infectivity to humans (4). Unlike Tbr, Tbg1 parasites are genetically homogeneous and form a monophyletic group (5,6). They are responsible for the vast majority of the HAT-cases (85% of the 663 newly reported HAT-cases in 2020) (7). Infections by a third group of human-infective trypanosomes - Tbg type II (Tbg2) that lack the SRA and TgsGP genes - are extremely rare (8). Like the non-human-infective *T. b. brucei* (Tbb), Tbg2 is genetically and phenotypically highly diverse (8).

The WHO targeted Tbg-HAT (gHAT) for elimination as a public health problem by 2020 and for global elimination of transmission (EOT) to humans (i.e. zero reported cases) by 2030 (9). Elimination of gHAT as a public health problem has been reached in several countries and HAT foci, and recently Togo and Côte d’Ivoire have been validated as such by WHO (10,11). However, EOT remains challenging due to imperfect diagnostics and the risk of re-emergence from asymptomatic human infections and/or a possible animal reservoir (12–14). Current serological tests based on the Tbg-specific LiTat 1.3 and LiTat 1.5 VSG antigens (15–20) still require validation in different animal
species. The golden standard for molecular detection of \textit{Tbg1} involves the single-copy \textit{TgsGP} gene (21–23), but the analytical sensitivity of these tests is limited because they target a hemizygous single-copy gene (24). Among the molecular tests, several studies have proposed various genotyping techniques such as isoenzymes, ribosomal genes, VSGs, SNPs and microsatellites (25–32), but these have some disadvantages such as the requirement of multiple PCR reactions or high amounts of input material, without necessarily increasing the sensitivity of \textit{Tbg1} detection. To this end, alternative genetic markers are much needed to reliably and with improved sensitivity demonstrate \textit{Tbg1} infection in humans and in animals, including the tsetse fly vector.

Previous reports indicated that \textit{Tbg1}-specific minicircle sequences exist in the kinetoplast DNA (33,34). The kinetoplast DNA (kDNA), unique to the single mitochondrion of unicellular flagellates of the order \textit{Kinetoplastida}, is a giant network of dozens of homogenous maxicircles (20–30 kb) interlaced with hundreds to thousands of heterogeneous minicircles (0.5–2.5 kb) (35). Maxicircles are homologous to the mitochondrial genome of other eukaryotes and encode components of the respiratory chain complexes and the mitoribosome. Minicircles generally consist of a ∼100 bp conserved sequence region that contains hyperconserved sequences named Conserved Sequence Blocks (CSBs), and a variable region including genes encoding guide RNAs (gRNAs) that are responsible for directing post-transcriptional modification of the maxicircle-encoded messenger RNAs (35). A complete assembly and annotation of the kinetoplast genome of a lab-adapted \textit{Tbb} strain identified 391 minicircle classes, encoding ~1000 gRNA genes (36). Analysis of kDNA minicircles is already used for \textit{Leishmania} detection and differentiation (37,38) and allows subtypes of \textit{Trypanosoma evansi} to be distinguished (39). It has also been proposed for sensitive and specific detection of \textit{Tbg1} infection in humans (34) and animals (33). However, the exact nature of these sequences is unknown and their use for \textit{Tbg1} diagnosis is cumbersome, requiring both nested PCR and DNA hybridization.

Recent advances in the development of bioinformatic tools now facilitate the assembly and annotation of the structurally complex kinetoplast genome (36,40), allowing us to investigate minicircle sequence diversity in tens to hundreds of samples simultaneously (41). Therefore, we used next-generation sequencing of total cellular DNA extracts as a strategy to investigate minicircle sequence diversity in 38 animal- and human-infective trypanosome extracts as a strategy to investigate minicircle sequence diversity in tens to thousands of heterogeneous minicircles (20–30 kb) interlaced with homogenous maxicircles (0.5–2.5 kb) (35). Maxicircles are homologous to the mitochondrial genome of other eukaryotes and encode components of the respiratory chain complexes and the mitoribosome. Minicircles generally consist of a ∼100 bp conserved sequence region that contains hyperconserved sequences named Conserved Sequence Blocks (CSBs), and a variable region including genes encoding guide RNAs (gRNAs) that are responsible for directing post-transcriptional modification of the maxicircle-encoded messenger RNAs (35). A complete assembly and annotation of the kinetoplast genome of a lab-adapted \textit{Tbb} strain identified 391 minicircle classes, encoding ~1000 gRNA genes (36). Analysis of kDNA minicircles is already used for \textit{Leishmania} detection and differentiation (37,38) and allows subtypes of \textit{Trypanosoma evansi} to be distinguished (39). It has also been proposed for sensitive and specific detection of \textit{Tbg1} infection in humans (34) and animals (33). However, the exact nature of these sequences is unknown and their use for \textit{Tbg1} diagnosis is cumbersome, requiring both nested PCR and DNA hybridization.

Recent advances in the development of bioinformatic tools now facilitate the assembly and annotation of the structurally complex kinetoplast genome (36,40), allowing us to investigate minicircle sequence diversity in tens to hundreds of samples simultaneously (41). Therefore, we used next-generation sequencing of total cellular DNA extracts as a strategy to investigate minicircle sequence diversity in 38 animal- and human-infective trypanosome strains from diverse geographical origins. Following a series of computational analyses, we identify minicircles that were present in all \textit{Tbg1} strains and absent in all \textit{Tbb}, \textit{Tbg2} and \textit{Tbr} strains. Using a newly developed quantitative PCR (qPCR), we demonstrated that our minicircle-based assay reliably identifies \textit{Tbg1} within the Trypanozoon subgenus. Furthermore, we show that the copy number of the targeted minicircle was on average 4-fold higher compared to the hemizygous single-copy \textit{TgsGP} gene, suggesting that our minicircle-based assay may be used for sensitive detection of \textit{Tbg1} infections in humans and animals.

**MATERIALS AND METHODS**

**Ethics statement**

Expansion of bloodstream form trypanosome populations in mice received approval from the Animal Ethics Committee of the Institute of Tropical Medicine (DPU2017-1).

**DNA extraction and sequencing**

To cover a wide geographical range, we included a total of 18 \textit{Tbg1} strains isolated from humans between 1952 and 2008 from Cameroon (\(n = 2\)), the Democratic Republic of the Congo (\(n = 10\)), the Congo Brazzaville (\(n = 2\)), Côte d’Ivoire (\(n = 3\)) and South Sudan (\(n = 1\)) (Supplementary Table S1). For comparative purposes, we also included nine \textit{Tbb}, five \textit{Tbg2} and six \textit{Tbr} strains (Supplementary Table S1).

All 38 strains were propagated as bloodstream form populations in OF1 mice (Charles River, Belgium) and purified from the infected mouse blood via DEAE ion exchange chromatography (42). Purified trypanosomes were sedimented by centrifugation (17 000 × g, 10 min at 4°C). DNA of 50 μl pure trypanosome sediment was extracted using the standard phenol:chloroform method (43), aliquoted at 1 ng/μl and stored at –20°C. The concentration of extracted DNA was determined using a Qubit 4 Fluorometer (Invitrogen by Thermo Fisher Scientific). Paired-end 150 bp sequences were generated using the DNA nanoball sequencing technology (DNBSEQ™) at the Beijing Genomics Institute (BGI) in Hongkong, China.

**Genomic analyses**

Paired-end reads were aligned against the \textit{Tbb} TREU927 v4.6 reference genome (available on https://tritrypdb.org) using SMALT v0.7.6 (https://www.sanger.ac.uk/tool/smalt-0.0). A hash index of the reference genome was built using \texttt{H9262} and stored at –20°C. The concentration of extracted DNA was determined using a Qubit 4 Fluorometer (Invitrogen by Thermo Fisher Scientific). Paired-end 150 bp sequences were generated using the DNA nanoball sequencing technology (DNBSEQ™) at the Beijing Genomics Institute (BGI) in Hongkong, China.

Variant calling and filtering was performed using the Genome Analysis Toolkit (GATK) v4.1.4.1 (44). First, reads were assigned to a single read-group with \texttt{AddOrReplaceReadGroups} and duplicated reads were marked with \texttt{MarkDuplicates}. Variants were then called for each strain separately with \texttt{HaplotypeCaller} using default parameters. The resulting gVCF files for all strains were combined with \texttt{CombineGVCFs} to allow joint genotyping with \texttt{GenotypeGVCFs}. Single Nucleotide Polymorphisms (SNPs) were extracted from the resulting VCF file with \texttt{SelectVariants} and filtered with \texttt{VariantFiltration} using the following parameters: \texttt{QUAL < 500, DP < 5, EXQ < 2.0, FS > 60.0, MQ < 40.0, MQRankSum < -12.5, ReadPosRankSum < -8.0, -cluster_window_size 10 and -cluster_size 3}. Finally, we used BCFTools v1.10.2 (45) to extract bi-allelic SNP sites that were called in all \textit{Tb} strains. Using the resulting set of genome-wide SNPs, we
reconstructed a phylogenetic network with SplitsTree v4 (46) to infer the ancestral relationship among the 38 Tb strains.

The species identity of the Tbgl and Tbr strains was confirmed in silico by investigating the presence of the TgsGP and SRA genes, respectively. To this end, MEGAHit v1.2.9 (47) was used for de novo assembly of genomes of all 38 Tb strains using default parameters. The presence of TgsGP and SRA genes in the assembled contigs was then confirmed through a local BLAST search (48) using publicly available nucleotide sequences of TgsGP (21) (NCBI accession number: FN555993) and SRA (2) (NCBI accession number: Z37159) with the following parameters: minimum 90% identity, minimum e-value of 0.0001 and minimum alignment length of 500 bp.

Assembly of the kinoplast genome

Reads that did not align to the TbB TREU927 nuclear reference genome were extracted using SAMtools v1.9 (45) and converted to FASTQ format using GATK SamToFastq. These unmapped reads were aligned against the 23 kb maxicircle sequence of TbB Lister 427 (GenBank accession id M94286) using SMALT and the same parameters as described above except that the hash index was built with k-mer words of length six and the reads were mapped with a minimum identity threshold (−y) of 90% and a maximum insert size (−i) of 500. SNP calling was done using GATK as described above, and we extracted only those SNPs that passed the quality criteria (see above) and that were present within the maxicircle coding region (1.3–16.3 kb). Similar to the analyses above for the nuclear genome, a phylogenetic network analysis was done using maxicircle coding SNPs with SplitsTree.

Reads that did not align to the maxicircle sequence of TbB Lister 427 were extracted from the alignment file as described above, and used for the assembly of minicircle contigs. Before assembly, sequence reads were trimmed for high quality with fastp v0.20.0 (49) using the following parameters: allow for a maximum of 10% of bases per read with a maximum length of 100 and 155 bp after trimming, and cut the read on the average phred-scaled base quality at either end of the read when their phred-scaled quality is below 30, trim bases of clipped reads at either end of the assembled minicircles, and use a custom python script implemented in KOMICS. Following mapping with SMALT with exhaustive search (−x) and a percent identity of 97% (−y), we have calculated the following metrics using a bash script implemented in KOMICS: number of reads, number of mapped reads, number of properly paired reads, number of reads with mapping quality ≥20, number of CSB3-containing reads, number of mapped CSB3-containing reads and number of perfectly aligned CSB3-containing reads (i.e. alignments without any insertions or deletions). The proportion of perfect alignments of CSB3-containing reads serves as a proxy for the total number of minicircles that were initially present within the DNA sample. All metrics were processed and visualized using the R function msc.quality as implemented in the R package rKOMICS (41). In addition, the quality of the assembly was further verified by calling SNPs with BCFTools mpileup/call, retaining only SNPs with QUAL ≥60 and DP ≥30, and assuming that high-quality assemblies should yield relatively low number of homozygous SNPs.

Finally, using the rKOMICS function msc.depth, minicircle copy numbers (MCN) were estimated as the median read depth per minicircle contig divided by the median genome-wide read depth times two (assuming diploidy in all Tb strains).

Identification of minicircle sequences unique to Tbgl

The diversity and similarity of minicircle sequences within all Tb strains were examined with the R package rKOMICS (41). Following visual inspection of length distributions using msc.length, we used preprocess to retain minicircle sequences that had the expected length (800–1200 bp) and that were successfully circularized. Retained sequences of all samples were concatenated into a single FASTA file and clustered into Minicircle Sequence Classes (MSCs) based on a minimum percent identity (MPI) of 70, 80, 90 and 95–100 with VSEARCH. In order to choose an appropriate MPI for downstream analyses, we processed VSEARCH clustering results with msc.uc and inspected - at each MPI - the number of MSCs, the number of perfect alignments and the number of 2-nt and 3-nt gaps. In addition, VSEARCH clustering results were stored into a matrix using the rKOMICS function msc.matrix, which records the presence (1) or absence (0) of all MSCs (rows).
for each strain (columns). This matrix was subsequently used to document the number of MSCs per strain with *msc.richness*, to calculate the proportion of MSCs shared between the different *Tb* subspecies with *msc.similarity* and to investigate the ancestry among all *Tb* strains with *msc.pca*. Finally, we used the rKOMICS function *msc.subset* to find MSCs that were present in the *Tbg1* strains and absent in the strains belonging to the other *Tb* subspecies.

Development of minicircle-based quantitative PCR assays

For each of the common *Tbg1*-specific MSC, we extracted the assembled minicircle sequences for all 18 *Tbg1* strains from the alignment using the rKOMICS function *msc.seq*, generated consensus sequences with Jalview v2.11.1.4 (52) and designed primers and probes (Table 1) with the RealTimeDesign qPCR assay design software (LGC, Biosearch Technologies). Probes targeting *Tbg1*-specific MSCs were modified with a FAM dye label at the 5′ end and paired with BHQ-1 plus BHQ-2 at the 3′ end.

A 20 μL reaction mixture contained 1X PerfeCTa qPCR Toughmix (Quantabio), 100 nM of each primer (LGC, Biosearch Technologies), 300 μM of each probe (LGC, Biosearch Technologies) and 5 μL of template DNA. The thermal cycling profile consisted of an initial denaturation step at 95°C for 10 min followed by 40 cycles at 95°C for 15 s and 60°C for 1 min. qPCR was conducted on a Q-qPCR Instrument (Quantabio), and detection of the quantification cycle (Cq) was calculated using the Q-qPCR Instrument Software v1.0.2 with the automatic threshold enabled.

Each qPCR targeting *Tbg1*-specific MSCs was multiplexed with the *Trypanozoon*-specific q18S-assay targeting the multi-copy 18S rRNA gene (53), the *Trypanozoon*-specific qGPI-PLC-assay targeting the single-copy glycosylphosphatidylinositol-specific phospholipase C (GPI-PLC) gene (53) and a *Tbg1*-specific qTgsGP-assay, designed to target the single-copy TgsGP gene and avoid amplification of TgsGP-like genes (21). The multi-copy 18S rRNA gene was used as an internal standard for the sensitive detection of *Trypanozoon* DNA. The single-copy GPI-PLC gene was used as an internal standard to determine if sufficient *Trypanozoon* DNA was present to detect a single-copy sequence (54), and for the calculation of relative copy numbers (RCN) of each *Tbg1*-specific MSC (see below). The single-copy TgsGP gene was used as a golden standard for the specific detection of *Tbg1* DNA. The q18S-assay contains a CAL Fluor Orange 56 dye labeled probe paired with BHQ-1 plus BHQ-2 (53). The qGPI-PLC-assay contains a CAL Fluor Red 610 dye labeled probe paired with BHQ-2 plus BHQ-2 (53). The qTgsGP-assay contains a Quasar 670 dye labeled probe paired with BHQ-2 plus BHQ-2.

The qPCR efficiency and analytical sensitivity were calculated for each qPCR targeting *Tbg1*-specific MSCs in simplex and in quadruplex format. This was done using phenol:chloroform extracted DNA (see above) of two *Tbg1* strains. From these DNA extracts, ten-fold serial dilutions in DEPC-treated water, ranging from 100 pg/μL to 1 fg/μL, were prepared. Each qPCR was run in quadruplicate for each DNA dilution. A reaction was considered positive if at least three out of four replicates were positive.

The specificity of the quadruplex qPCR assays was assessed with the phenol:chloroform extracted genomic DNA of 34 *Tbb*, 49 *Tbg1*, 7 *Tbg2*, 15 *Tbr*, 2 *T. equiperdum* and 5 *T. evansi* strains (Supplementary Table S2). The 49 *Tbg1* samples originated from Burkina Faso (2), Cameroon (2), Côte d’Ivoire (6), Congo Brazzaville (3), Democratic Republic of the Congo (35) and South Sudan (1). Note that 26 strains from the Democratic Republic of the Congo were sampled within the context of a treatment outcome study in Mbuj-Maiyi (55), with 14 strains sampled from seven patients before (sample name include ‘BT’) and after (‘AT’) treatment. Quadruplex qPCR assays were run in duplicate for each DNA extract. The specificity of the assays was further assessed on DNA prepared from man, cattle, dog, goat, horse, sheep and tsetse (*Glossina fuscipes* from Kwamouth, Democratic Republic of the Congo, 2018), all known hosts of *Tbg1* (12).

Relative copy numbers (RCN) of each target were calculated using the ΔCq-method with qGPI-PLC as reference. This was done by subtracting the Cq-values obtained for each *Tbg1*-specific MSC from the Cq-value obtained for qGPI-PLC. The resulting ΔCq-value were averaged between replicates and transformed (2ΔCq) to yield RCNs for each target.

Annotation of minicircle sequences targeted by qPCR assays

Strain *Tbg1* 340AT (MHOM/CD/INRB/2006/21B) isolated in 2006 in Mbuj-Maiyi (DRC) (55) (Supplementary Table S1) was selected for representative minicircle annotation because of its comparatively high minicircle complexity (see results). DNA extraction, sequencing and alignment of sequence reads against the *Tbb* TREU9274 v4.6 reference genome was done as described above for the other 38 *Tb* isolates initially included in our study. Reads that did not align to the nuclear reference genome were used for the assembly of mitochondrial maxicircles and minicircles with KOMICS, as described above.

Due to the lack of transcriptomic data for *Tbg1* strain 340AT, edited mRNA sequences were predicted following a similar approach as in (40). Edited mRNA sequences for *Tbb* strains Lister 427, EATRO 164 and EATRO 1125 (36) were obtained from GenBank and manually corrected for changes in non-T residues based on alignment of the *Tbg1* 340AT maxicircle with the annotated *Tbb* EATRO 1125 maxicircle (36).

Guide RNA prediction and minicircle annotation were performed with python3 package for kDNA annotation (36). The alignments of the gRNAs encoded on the minicircles targeted by the diagnostic assay to their cognate mRNAs were carefully inspected to identify any ‘non-redundant’ gRNAs, i.e. gRNAs that direct editing events not covered by any other gRNAs. Furthermore, gRNAs 3’ of such non-redundant gRNAs were checked for potential premature truncations by the gRNA calling algorithm by carefully examining the editing capacity of their 3’ end extensions. Any gRNA genes that were confirmed to be non-redundant were considered essential, as were the minicircles that encoded them.
The genomes of 38 Tb strains were sequenced at a median depth (mean = 155, min = 126, max = 178) (Supplementary Table S3). On average 86.3% of the reads (min = 81.25%, max = 92.05%) aligned to the Tb TREU927 nuclear reference genome (Supplementary Table S3). Initial variant discovery with GATK identified a total of 1,558,963 SNPs across the 38 Tb strains. Strict quality filtering and the exclusion of multiallelic sites reduced the data set to 316,287 genome-wide bi-allelic SNPs, of which 310,701 SNPs (98.23%) were located within the 11 megabase chromosomes. In addition, joint genotyping identified a total of 150 SNPs within the maxicircle coding region.

Phylogenetic analyses based on genome-wide SNPs and SNPs from the maxicircle coding region confirmed that all 18 Tbgl parasites clustered together in a monophyletic group, a prerequisite for downstream analyses that aim at identifying Tbgl-specific minicircles (Supplementary Figure S1). Using a local BLAST search of assembled contigs, we also confirmed the presence of the Tbgl-specific TgsGP gene for the 18 Tbgl strains, and the Tbr-specific SRA gene for the six Tbr strains. These two genes were absent for the remaining nine Tbb strains and five Tbgl strains (Supplementary Table S4).

**Assembly and circularization of mitochondrial minicircles**

Mitochondrial minicircles were *de novo* assembled, circularized and reoriented for each of the 38 Tb strains using the Python package KOMICS. A total of 9,076 minicircle contigs were assembled across all 38 Tb strains, of which 7,156 (78.85%) were successfully circularized (Supplementary Table S5). The length of the majority of circularized minicircles (7,111 contigs, 99.37%) showed a unimodal distribution around ~1,000 bp (Supplementary Figure S2), which is comparable to the minicircle length found in Tbb (36). To verify whether the assembly process was impacted by excluding reads mapped on the nuclear genome or the maxicircle, we performed a BLAST search of the assembled minicircles against the reference genome, retaining only alignments with a minimum 90% identity, an e-value of 0.001 and a minimum length of 150 bp (the read length). This identified a total of 13 contigs that most likely originated from the nuclear genome, as they showed a length between 2,608 and 10,876 bp (which is much larger than the expect 1,000 bp length of minicircles). The length of one contig was 419 bp, which aligned with 96% identity and a length of 151 bp against chromosome 10. It is thus possible that this contig was not fully assembled because of sequence homology with the reference genome. However, this is only 1 contig in 1,409 minicircle contigs that were not successfully circularized. Hence, we believe that the impact of our filtering method has a negligible impact on the assembly process.

To validate the quality of the assembly process, sequence reads were aligned to the assembled minicircle contigs and several mapping and genotyping statistics were calculated. First, high-quality assemblies should result in relatively low numbers of homozygous SNPs when reads are aligned against the assembled contigs. Here, we identified a total of 302 homozygous SNPs within 127 minicircle contigs (Supplementary Table S5), which is only 1.4% of all assembled minicircle contigs. Only 48 homozygous SNPs were identified within 17 circularized contigs (0.2% of all circularized minicircles) (Supplementary Table S5). These results show that homozygous SNPs were found for only a fraction of the assembled contigs. Second, on average 96.99% of the sequence reads mapped in proper pairs.
and 94.46% aligned with a mapping quality larger than 20 (Supplementary Table S5), indicating that the large majority of mapped reads aligned with a high quality and with the expected orientation to the minicircle assemblies. Third, we calculated the number of aligned reads containing the CSB3 12-mer as a proxy for the total number of minicircles initially present within the DNA sample. This revealed that on average 93.24% of the CSB3-containing reads aligned against the assembled minicircles, and 88.77% aligned perfectly (Supplementary Table S5), suggesting that we were able to retrieve the vast majority of the minicircles. Note that for the assembly and circularization of minicircles, we only used sequencing reads that did not align against the nuclear genome or the coding region of the mitochondrial maxicircle. Therefore, the ~7% of unaligned CSB3-containing reads may have originated from minicircles that have not been (fully) assembled or circularized, the variable region of the minicircle or the minichromosomes.

**Estimation of maxicircle and minicircle copy numbers and network size**

To calculate the average number of maxicircles and minicircles per kinetoplast network, we used the coverage (i.e. median read depth) of the diploid nuclear genome. The average genome-wide coverage was 155 per diploid cell (two copies), and the average coverage per haploid sequence was equal to 78 (Supplementary Table S3). The coverage of the mitochondrial maxicircles (coding region only) and minicircles was estimated based on median read depths per 0.1 kb. Average coverage of the maxicircle was 1262 (median = 1325, min = 99, max = 2320) (Supplementary Table S6). This equaled an average copy number of 17 maxicircles (median = 18, min = 1, max = 28) per network (Supplementary Table S6), which is slightly lower compared with previous estimates of 20–50 copies per network (36,56–57). The average copy number of minicircles (MCN) ranged from 2.1 to 37.9 copies per network in *Tbg*1 strains, from 0.8 to 32.7 copies per network in *Tbb* strains, from 4 to 11.9 copies per network in *Tbg*2 strains and from 3.1 to 11.1 copies per network in *Tbb* strains (Supplementary Table S7). Thus, copy numbers for minicircles within each network varied substantially. The size of each kDNA network was estimated by adding up the estimated copy numbers for all minicircles. These calculations resulted in an estimated average of ~2,100 minicircles per network, ranging between 252 minicircles in the Nabe strain and 4,828 minicircles in the MSUS-CI-78-TSW-157 strain (Supplementary Table S7). These numbers are slightly lower compared to earlier estimates of 5000–10 000 minicircles per network (36,56–58), and future studies should attempt to confirm a lower amount of kDNA in *Tbg*1 by other methods, such as quantitation by microscopy.

**Sequence diversity and similarity of mitochondrial minicircles**

Minicircle sequence diversity was examined using a clustering approach, whereby minicircles were grouped into MSCs based on a minimum percent identity. This was done on the 7111 circularized minicircle contigs of the expected length, as these would produce the most robust alignments. At 100% identity, a total of 5883 unique MSCs were identified across the 38 *Tbg* parasites, leaving 1228 MSCs that are shared between two or more isolates. The number of MSCs decreased sharply with decreasing percent identities to a total of 719 MSCs at 70% identity (Supplementary Figure 3A). Regardless of the percent identity used, *Tbg*1 parasites contained an average of 103 MSCs per strain (median = 107), which is on average 2.48-fold lower compared to other taxa of the *Trypanosoma* subgenus (Figure 1). Most of the *Tbg*1 strains displayed a fairly similar number of MSCs, ranging between 89 and 122 MSCs (Figure 1), with the exception of LiTat 1.5 (50 MSCs) and Bosendja (76 MSCs). The composition of minicircle sequences was further investigated by quantifying the proportion of MSCs unique to *Tbg*1. At 98%-100% identity, the 18 *Tbg*1 parasites did not share any MSC with the non-*Tbg*1 subspecies (*Tbr, Tbg*2 and *Tbb*) (Supplementary Figure 3B). Below 98%, there was a steady increase in the proportion of shared MSCs between *Tbg*1 and non-*Tbg*1 subspecies (Supplementary Figure 3B). At the 98% identity threshold, the *Tbg*1 group contained 241 MSCs, none of which were found in the other *Tb* subspecies and three of which were found in all 18 *Tbg*1 strains (Figure 2). These three *Tbg*1-specific MSCs were retained as candidate markers for our new molecular test.

**Novel multiplex qPCRs including *Tbg*1-specific mitochondrial minicircles as target**

Using the RealTimeDesign qPCR assay design software, we successfully designed three simplex qPCR assays (here-after referred to as qMini1, qMini2 and qMini3) targeting two *Tbg*1-specific MSCs (here-after referred to as mO_078 and mO_104) (Table 1). Due to limitations such as ambiguous base count, low GC percentage and low melting temperatures, we were unable to design primers and probes for the third *Tbg*1-specific MSC. The *Tbg*1 specificity of the designed primers and probes was confirmed with a BLAST search (48) on TriTrypDB (https://tritrypdb.org), with a Primer BLAST search on NCBI (https://www.ncbi.nlm.nih.gov) and with the command line search tool grep in sequencing reads generated by this study. The three simplex qPCR assays qMini1, qMini2 and qMini3 were each multiplexed with q18S, qGPI-PLC and qTgsGP (Table 1) to produce three quadruplex reactions (here-after referred to as g-qPCR1, g-qPCR2 and g-qPCR3 when qMini1, qMini2 and qMini3 were included, respectively).

The efficiency and analytical sensitivity of the simplex and quadruplex qPCR assays were investigated using DNA from two *Tbg*1 strains (Supplementary Table S8), one with relatively low MCNs (Nabe; average MCN = 1) and one with relatively high MCNs (LOGRA; average MCN = 8). The lower detection limit of 0.05 pg DNA was reached in both strains for qMini1, qMini2 and qMini3 in their respective simplex reactions (Supplementary Figure S4), for q18S in all three g-qPCR assays (Supplementary Figure S4), and for qMini1 and qMini3 in their respective g-qPCR assays, with the exception of qMini3 that achieved the detection limit of 0.5 pg DNA in the Nabe strain (Supplementary Figure S4). This detection limit
of 0.5 pg DNA was also reached by qTgsGP and/or qGPI-PLC in all three g-qPCR assays with the LOGRA strain and in the g-qPCR2 and g-qPCR3 assays with the Nabe strain (Supplementary Figure S4). The analytical sensitivity of qMini2 was greatly reduced in the g-qPCR2 assay to 100 pg DNA with the Nabe strain and 0.5 pg DNA with the LOGRA strain (Supplementary Figure S4). The qPCR efficiency of g-qPCR1 and g-qPCR3 was estimated between 93% and 106% (Supplementary Figure S4), which is considered acceptable (https://www.thermofisher.com/content/dam/LifeTech/global/Forms/PDF/real-time-pcr-handbook.pdf).

The qMini3 assay displays a similar specificity as the qTgsGP assay

To assess the taxon-specificity of qMini1 and qMini3 within the Trypanozoon subgenus, a total of 118 DNA extracts were tested with the g-qPCR1 and g-qPCR3 assays (Supplementary Table S9). Here, qMini2 was excluded because of its low analytical sensitivity in the g-qPCR2 assay (see above) and because it targets the same minicircle as qMini3 (Table 1). Six of the 118 DNA extracts were excluded as they didn’t react in duplicate with qGPI-PLC, which was used as an internal standard to determine if sufficient Trypanozoon DNA was present to detect a single-copy sequence (Supplementary Table S10). The remaining 112 DNA extracts reacted with the Trypanozoon-specific assays q18S and qGPI-PLC (Supplementary Table S9). All 49 Tbg₁ DNA extracts reacted with qTgsGP, qMini1 and qMini3, with the exception of MSUS/Cl/82/TSW125_KP1_1cloneB (Tbg₁ isolated from a pig in Côte d’Ivoire), ALJO (Tbg₁ isolated from a human patient in DRC) and GUIWI-BOBO80-MURAZ18 (Tbg₁ isolated from a patient in Burkina Faso) that remained negative for qMini1. The qMini1 assay also showed one cross reaction with DNA extracted from the Tbr strain Etat 1.2 R, with a Cq-value 5× lower than that of the qGPI-PLC. The qMini3 assay showed no cross reactions or false negative results. In addition, the g-qPCR3 assay did not amplify DNA from human and non-human vertebrates that are known to be susceptible for Tbg₁ infection, in casu horse, cattle, goat, sheep, pig and dog, and DNA from Glossina fuscipes. Also, DNA from other livestock affecting trypanosomes like T. congolense, T. theileri and T. vivax was not amplified (Supplementary Table S9).

The qMini3 assay targets a minicircle with a relatively high, but variable copy number

Relative Copy Numbers (RCN) of the minicircle sequence targeted by qMini3 (RCNₘₙₜₜₜₜₜ), the target sequence of qTgsGP (RCNₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜₜ¢)

Figure 1. Number of Minicircle Sequence Classes (MSCs) for 38 Tb strains. Following de novo assembly and circularization, mitochondrial minicircle sequences were clustered into groups of sequences sharing a minimum percent identity (MSCs). Here, we summarized the number of MSCs identified within each Tb strain for a range of percent identities (70, 80, 90, 95 and 100). Strains were grouped as T. b. brucei (Tbb), T. b. gambiense type 1 (Tbg₁), T. b. gambiense type 2 (Tbg₂) and T. b. rhodesiense (Tbr).
The median RCNmO$_{0.078}$ (7.98) was 4.29x higher than the median RCNgTgsGP (1.86) and 2.06x higher than the median RCNg18S (3.88) (Figure 3A). However, RCNmO$_{0.078}$ also displayed a larger variation (SD = 9.01, min = 0.77, max = 40.58) compared to RCNgTgsGP (SD = 0.62, min = 0.94, max = 3.47) and RCNg18S (SD = 0.70, min = 1.51, max = 5.47). The RCNmO$_{0.078}$ was lower compared to RCNgTgsGP for 4/49 Tbg1 strains and to RCNg18S for 12/49 Tbg1 strains. There was no association between RCNmO$_{0.078}$ with the year of isolation (Pearson correlation test; cor = −0.1925748, t = −1.331, df = 46, P-value = 0.1897) or the geographical origin of the strain (Kruskal-Wallis test; country: chi-squared = 4.0896, df = 5, P-value = 0.5366; region: chi-squared = 3.4053, df = 2, P-value = 0.2181). High variation in RCNmO$_{0.078}$ was found within one group of Tbg1 strains isolated from humans between 2005 and 2009 in Mbuji-Mayi (Democratic Republic of the Congo), with a minimum RCNmO$_{0.078}$ of 0.8 in the 186BT strain and a maximum RCNmO$_{0.078}$ of 36.7 in the 93AT strain. Here, RCNmO$_{0.078}$ was significantly higher in strains sampled after treatment (mean RCNmO$_{0.078}$ = 14.9) compared to the RCNmO$_{0.078}$ in strains sampled before treatment (mean RCNmO$_{0.078}$ = 6.9), although this difference was not significant (Welch two sample t-test on all strains, t = 2.0305, df = 11.944, P = 0.06519 and paired t-test on paired strains, t = 1.5476, df = 5, P = 0.183). The RCNmO$_{0.078}$ as calculated using the ΔCq-method was strongly associated with MCNmO$_{0.078}$ as calculated using standardized read depths, with a coefficient of determination of 0.91, a slope of 0.32 and a y-intercept of 0.10 (Figure 3B).

The qMini3 assay targets a minicircle containing non-redundant guide RNA genes

Annotation of minicircles was done for strain 340AT. A local BLAST search of assembled contigs revealed the presence of the Tbg1-specific TgsGP gene, confirming that 340AT is a Tbg1 strain (Supplementary Table S3). The mitochondrial maxicircle and minicircles were assembled with KOMICS using sequence reads that did not align to the nuclear reference genome. This resulted in a maxicircle contig of 21 287 bp long (including the entire coding region) and a total of 143 minicircle contigs (including 129 circularized contigs). Hence, 340AT has the highest number of minicircles when compared to the 18 Tbg1 strains (max. 132 minicircles) initially sequenced in this study (Supplementary Table S7), which was the main motivation for using the 340AT data for representative minicircle annotation.

Annotation of the 143 minicircles revealed that the minicircle targeted by qMini3 encodes four gRNA genes (Figure 4). These gRNAs are involved in editing of the maxicircle genes cytochrome c oxidase subunit 3 (gCOX3(616–656) and gCOX3(341–369)), ATPase subunit 6 (gA6(415–452)) and NADH dehydrogenase subunit 7 (gND7(847–888)) (Figure 4). As expected, the minicircles contain the semi-conserved region characterized by conserved sequence blocks CSB1, CSB2 and CSB3 (36,50), and the gRNA genes are framed by imperfect 18bp inverted repeats (36,59–61).

Next, we investigated if any of the gRNAs encoded by minicircle mO$_{0.078}$ are non-redundant, i.e. whether they direct the editing of sites not covered by any of the other gRNAs encoded in this strain’s kDNA. Minicircles encoding only redundant gRNAs might be more prone to loss due to lack of selective pressure, which could make a diagnostic assay based on such minicircles less reliable. Our analyses showed that gRNAs gA6(415–452) and gND7(847–888) direct editing events not covered by any other gRNA (Figure 5), and are thus non-redundant. The two COX3 gRNAs are redundant with gRNAs encoded by other minicircles (results not shown).

DISCUSSION

This study presents a computational investigation of mitochondrial minicircle sequence diversity in trypanosome isolates, resulting in the development of a qPCR assay as a promising new tool for
sensitive diagnosis of *Tbg1* infections in humans and animals.

Computational and phylogenetic analyses using genome sequencing data revealed that the 18 sequenced *Tbg1* strains are monophyletic—contrary to opinions outlined by (62)—and contain the *TgsGP* gene, an essential precondition for identifying *Tbg1*-specific minicircles. The latter was achieved by grouping *de novo* assembled and circularized minicircles into MSCs according to sequence similarity. This uncovered a variable number of MSCs within the *Trypanozoon* subgenus, with a considerably lower number of MSCs in *Tbg1* compared to the other subspecies. The comparatively lower number of MSC within *Tbg1* may be the result of its asexual evolution (5) that results in the inevitable loss of redundant minicircles due to random genetic drift (63). Hence, the monophyletic origin and the asexual evolution of *Tbg1* may explain the less complex minicircle populations in isolates for this subspecies, with conserved presence of some minicircle classes, which facilitated the identification of *Tbg1*-specific minicircles. Identification of taxon-specific minicircles may prove more challenging for trypanosomatid parasites experiencing occasional recombination (64–66), leading to heterogeneous minicircle populations as a result of biparental inheritance of mitochondrial minicircles (40,67–69).

A total of 241 MSCs were recognized as *Tbg1*-specific, of which three were shared by the 18 studied *Tbg1* strains. For two of the three *Tbg1*-specific minicircles (mO\_078 and mO\_104), three molecular assays could be successfully developed and tested. While two qPCR assays were discontinued because of false-negative results or low analytical sensitivities, one qPCR assay targeting minicircle mO\_078 was fully specific (i.e. no false positives or false negatives) when tested on DNA of 112 different *Trypanosoma sp.* strains. These results show that the minicircle-based assay is as specific as the assay targeting the *TgsGP* gene, the current golden standard for molecular detection of *Tbg1* parasites (70–76), confirming the taxon-specificity of some kDNA minicircles in *T. b. gambiense* (33,34) and their exploitability in molecular tests as has been described for *Leishmania* (37,38) and *T. evansi* type A and B (39,77–79). Annotation of the mO\_078 minicircle demonstrated that it encodes two non-redundant gRNAs that are essential for completing the editing of ATP synthase subunit A6, a gene required for survival in both the bloodstream stage and the insect stage of *Tb* (80,81), and NADH dehydrogenase subunit 7, respectively. Hence, our results indicate that minicircle mO\_078 is essential for the survival of the parasite and will most likely be preserved in all natural *Tbg1* strains, ensuring the reliability of a diagnostic assay targeting this minicircle.

Molecular analyses revealed that the *Tbg1*-specific minicircle mO\_078 is a multicopy marker for the large majority of strains tested in this study, with a median copy number equal to eight and a maximum copy number of 41. Minicircle mO\_078 copy number exceeded that of the multicopy *18S* gene in 3/4 of the *Tbg1* strains and was as low

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**Figure 3.** Relative copy numbers of the qMini3 target sequence. (A) Relative copy numbers (RCN) were estimated for the qMini3, q18S and qTgsGP target sequences using the ΔCq-method with the qGPI-PLC as reference. Boxplots summarize the RCN estimates as calculated for 49 *Tbg1* strains. (B) Scatter plot showing the relationship between RCN (x-axis) and MCN (y-axis). To test whether RCN as calculated using the ΔCq-method is comparable to the Minicircle Copy Numbers (MCN) as calculated using standardized read depths, RCN and MCN were calculated for the qMini3 target sequence for 11 out of 18 *Tbg1* strains. For these 11 strains, there was sufficient DNA to allow both whole genome sequencing (for MCN calculation) and a qPCR run (for RCN calculation) on the same DNA extract. The remaining seven strains were excluded here as there was not sufficient DNA left for a qPCR run following whole genome sequencing.
as the TgsGP copy number in only four of the 49 tested Tbg1 strains. This finding confirmed the multicopy nature of mitochondrial minicircles in *Trypanosoma* and *Leishmania* parasites (36,82). However, our results also indicated that there is a relatively large variation in minicircle copy numbers across strains, suggesting that the detection limit of the minicircle-based assay may depend on the strain being investigated. The detection limit may also depend on the DNA extraction method, as DNA-extraction based on spin columns causes a random loss of small molecules like minicircles (36). Here, we avoided such biases by using the phenol-chloroform extraction method that captures all nucleic acids, although this method may be less amenable for high-throughput processing of human and animal specimens.

The high specificity and generally high copy number of minicircle mO_078 makes this a promising new marker for sensitive diagnosis of Tbg1 infections. Specifically, within the context of reaching EOT of gHAT by 2030 (9), our minicircle-based assay may prove valuable for studying the role of an animal reservoir in the epidemiology of gHAT (12). Therefore, we propose a new multiplex qPCR assay (g-qPCR3) that targets the Tbg1-specific minicircle mO_078 (serving as sensitive detection of Tbg1) in combination with the *Trypanozoon*-specific 18S (serving as sensitive detection of *Tb s.l.*) and the Tbg1-specific TgsGP gene (which confirms the trypanosome is human-infective and thus of epidemiological significance). We have shown that the g-qPCR3 assay is applicable on animals, as it does not amplify DNA from other livestock affecting trypanosomes like *T. congolense*, *T. theileri* and *T. vivax*, from the tsetse fly *Glossina fuscipes* and from six livestock species that are known to be susceptible to Tbg1 infection. Based on these data, we also trust that the specificity will not be compromised when testing blood from wild Bovidae and non-human primates (12), although we had no access to specimens from wild fauna to formally test this. The main limitation of our study is that the g-qPCR3 assay was tested on only one Tbg1 strain isolated from a host other than human, *in casu* a pig from Côte d’Ivoire (83), mainly because of the scarcity of such samples. However, given that minicircle mO_078 is non-redundant and universal (computational analyses confirmed that mO_078 was present in 193 sequenced Tbg1 strains sampled
Figure 5. Alignment of the non-redundant gRNAs from the qMini3 targeted minicircle mO078 to their target mRNAs. The alignment shows the non-redundant gRNAs in blue (A, gA6(415–452); B, gND7(847–888)) as well as the mRNA sequences immediately upstream and downstream, along with the neighbouring gRNAs encoded by other minicircles. Lines 1–3: mRNA position (hundreds, tens, ones); line 4: number of Us that have been deleted from the pre-edited mRNA at this position; line 5: edited mRNA sequence 5′ to 3′ (lowercase ‘u’s represent insertions). Note that the ‘anchor’ sequence at the 5′ end of each gRNA cannot direct editing events; line 6: protein sequence. For each gRNA: Line 1: name (mO_name(cassetteposition)mRNA(start-end of alignment on mRNA)), underscore characters denote the anchor; line 2: base-pairing: ‘|’: Watson-Crick basepair, ‘.’: GU basepair, ‘.’: mismatch basepair; line 3: gRNA sequence 3′ to 5′.

from humans in 11 different countries (data not shown), we are confident that the g-qPCR3 assay will be successful at amplifying Tbg1 DNA from livestock specimens.

In conclusion, this study exemplifies the power of genome assembly and annotation for identifying species-specific multicopy genetic markers. We developed a minicircle-based assay that is as specific as the current golden standard for molecular detection of Tbg1 infections, and argued that the g-qPCR3 has the diagnostic potential for assessing the importance of an animal reservoir in the epidemiology of gHAT.

DATA AVAILABILITY

Sequence reads generated within the context of this study have been deposited in the European Nucleotide Archive under accession number PRJEB49966 (https://www.ebi.ac.uk/ena/browser/view/PRJEB49966). The sequence of the Tbg1-specific minicircle mO078 was deposited to NCBI under accession number OM238297. Scripts used for processing WGS data in this study are available at https://github.com/GeertsManon/g-qPCR. All graphical analyses were performed using R v4.0.3 in RStudio v1.4.1103.

SUPPLEMENTARY DATA

Supplementary Data are available at NARGAB Online.

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