Molecular identification and prevalence of trypanosomes in cattle distributed within the Jebba axis of the River Niger, Kwara state, Nigeria

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Abstract
Background: Trypanosomiasis is a fatal disease that threatens the economy of at least 37 countries in sub-Saharan Africa, particularly with regard to livestock farming. In this study, we investigated the prevalence of trypanosome infection in cattle, and molecularly identified the species of trypanosomes in infected cattle and the spatial distribution of trypanosome-infected herds along the Jebba axis of the River Niger.

Methods: A randomized cross-sectional study was conducted along the Jebba axis of the River Niger by screening cattle from 36 herd clusters by nested PCR using ITS-1 generic primers. Data generated were analysed using the Chi-square test at a 95% confidence interval.

Results: Microscopic examination revealed three infected cattle out of 398 examined, representing 0.8% prevalence. Twelve animals (3.0%) were positive by PCR. Our results showed a decline in the packed cell volume of infected animals (24.7%). The infection rates were categorized as single infection in 11/12 (91.7%) and mixed infection in 1/12 (8.3%). Animals were most frequently infected by Trypanosoma congolense (50.0%), with T. congolense Savannah being the most prevalent subspecies (71.4%). Aside from the infection rate by age (10.0%) and relative distance of animals from the River Niger (56.2%), statistical differences in every other parameter tested were based on mere probabilistic chance. Spatial data showed that the disease was prevalent among herds located less than 3 km from the River Niger.

Conclusions: Six species of trypanosomes were identified in cattle herds along the Jebba axis of the River Niger, with T. congolense being the most prevalent. Age and relative distance of herds from the River Niger may be risk factors for trypanosome infection in cattle herds in this area.

Keywords: Trypanosomiasis, Cattle, ITS-1, Prevalence, Jebba, Geospatial distribution, River Niger

Background
African trypanosomiasis (AT) is a parasitic disease of public health concern affecting humans (human African trypanosomiasis) and animals (animal African trypanosomiasis), thereby limiting agricultural productivity in most developing countries in sub-Saharan Africa (sSA). The disease is caused by protozoan parasites belonging to the genus Trypanosoma. It is cyclically transmitted by tsetse flies and mechanically by other blood-feeding insects [1, 2]. Reports have shown that over 33% of Africa’s landmass has been pervaded by more than 30 Glossina species and subspecies [2, 3]. In Africa, the disease affects around 100 million head of cattle, with 6 million estimated in Nigeria alone out of a cattle population that...
is presently estimated at 20 million [4]. The World Health Organization (WHO) has estimated that 50–70 million individuals are at risk of tsetse bites in Africa, with about 30,000 reported cases per annum [3]. However, WHO-led intervention efforts targeting the elimination of HAT have resulted in a progressive decline in this incidence [5]. Nevertheless, no fewer than 1000 cases are being reported each year, despite continued active and passive screening efforts. The disease advances rapidly from a less virulent to a chronic form, and accounts for an economic loss of $4–5 billion in terms of gross domestic product [6, 7].

The major species responsible for animal African trypanosomiasis include *Trypanosoma congoense*, *T. brucei* and *T. vivax* [8]. In humans, *T. brucei gambiense* and *T. brucei rhodesiense* cause acute and chronic infections respectively [9]. However, several reports show the existence of atypical human infections caused by animal trypanosomes [10–14]. Therefore, identification of *Trypanosoma* species is fundamental to measuring the general threat posed by trypanosome species in animals and humans [15, 16].

There have been several recent developments in the diagnosis of AT towards understanding the disease transmission and distribution which is necessary for any successful treatment [17]. These advances have enabled accurate detection and identification of previously unidentified zoonotic *Trypanosoma* species [18]. While enzyme-based immuno sorbent assay (ELISA) was considered an improvement with regard to the sensitivity of pathogen determination, antigen detection using monoclonal antibody (mAb)-based ELISA is quite unreliable due to the presence of immune active agents in the blood which cannot distinguish between active and previous infections [19].

The advent of polymerase chain reaction (PCR) as a novel technique for direct parasite detection and identification offered greater sensitivity and reliability than methods previously used. The use of *Trypanosoma* species-specific DNA probes [20, 21] and PCR analysis [22–25] has significantly enhanced the efficiency of identification and understanding of trypanosome diversity, especially the high prevalence of mixed trypanosome infections which exist in the field [26–28]. Consequently, with specific PCR tests using generic primers [15], the identification of 11 known tsetse-transmitted trypanosome species/subspecies as well as unknown species has been made possible through amplification of the ribosomal RNA gene loci of the internal transcribed spacer (ITS-1) region of the trypanosome genome due to its variable inter-species length and high copy number [29, 30].

Several studies on trypanosomiasis transmission have been conducted in some parts of Nigeria [10, 11, 22]. However, to date, there is no published report on the prevalence of AAT in Kwara state. In this study, we investigated the prevalence of trypanosome infection in cattle and molecularly identified the species of trypanosomes in infected cattle and the spatial distribution of trypanosome-infected herds along the Jebba axis of the River Niger.

**Methods**

**Study area**

This study was conducted along the Jebba axis of the River Niger, Kwara state. Jebba is located at the geographical coordinates 9°9′14″N, 4°48′43″E, with views of the River Niger. In light of its area, it is known as the “midland” and the “door” between the southern and northern parts of Nigeria [31]. According to the 2006 census, the city’s population in 2006 was 22,411, and it is approximately 500 km from Abuja, and 306 km and 600 km from Lagos and Kaduna, respectively.

**Study population**

Our study population consisted of mostly transhumance cattle, having the possibility of mixing with sentinel animals. For parasite identification, animals with recent (≤ 2 weeks) administration of trypanocidal drugs were excluded from the study.

**Study design**

A cross-sectional study was undertaken to capture cattle distribution across the Jebba axis of the River Niger, Kwara state, and its tributaries to assess trypanosome distribution across the geographical area in June 2019.

**Sampling method and sample size**

A systematic random sampling technique was employed in this study. Herds of cattle in each coordinate were pooled and considered as a cluster from which animals were sampled by systematic randomization. The sampling frame was identified by listing herd locations across Jebba, and samples were obtained in each cluster based on proportion to size of herd; i.e. 6% of cattle in each herd. A systematic random sampling technique was used to select animals in each of the randomized clusters, whereby the sampling interval was generated by dividing the herd size by the sample size required for that herd. The first study subject was randomly selected from among the cattle, while others were selected at an interval based on herd size [32]. Due to the complete absence of previous records on the prevalence of trypanosomiasis in our study area, 50% prevalence was assumed and the sample size was estimated [33]. A minimum total sample size of 384 cattle was drawn from across the identified clusters. Sample size calculation was based on a 95%
Sample collection and parasitological analyses
Five millilitres of blood were collected from the jugular vein of each randomized animal using a sterile Vacutainer needle into tubes containing anticoagulant ethylenediaminetetraacetic acid (EDTA) [7]. Each sample was identified by a unique barcode system that corresponds to the name of the village, herd cluster and sample number. Samples were transported in an ice box to the laboratory and stored at 4 °C prior to laboratory analysis. Parasitological examination was done in the laboratory using the standard trypanosome detection methods, i.e., the Buffy coat method (BCM) [32], haematocrit centrifugation technique (HCT) [34], parasite load estimation [35] and Giemsa-stained thick and thin films [36]. The packed cell volume (PCV) of each animal was also determined and the parasites were identified [37, 38].

DNA extraction and PCR
The Quick-gDNA™ MiniPrep kit (Zymo Research Corporation, Irvine, CA, USA) was used for genomic DNA (gDNA) extraction from the blood as prescribed by the manufacturer. DNA yield and purity assessment were performed using a NanoDrop ND-100 UV spectrophotometer (NanoDrop Technologies, Inc./Thermo Fisher, Waltham, MA, USA) and the eluted DNA were stored at −20 °C until further use [32]. PCR amplification were carried out as described elsewhere [15], with slight modification. In the first round of the reaction, 3 µl of DNA was added to the PCR reaction mixture. PCR was performed in a total reaction volume of 25 µl containing 2.5 µl of standard Taq buffer (10x), 1.0 µl of dNTPs (10 mM), 1.0 µl of each primer (25 µM), 0.5 µl Taq DNA polymerase (5000 U/ml), 3.0 µl DNA template and 16.0 µl of nuclease-free water to make a final volume of 25 µl. The PCR runs were set as follows: initial denaturation at 95 °C for 2 min; 35 cycles of 30 s at 94 °C, 30 s at 54 °C and 60 s at 72 °C; final elongation for 5 min at 72 °C. In the first run of the nested PCR, Tryp 1 (forward, 5′AAGCGAAGTCATCCA TCG3′) and 2 (reverse, 5′TAGAGGAAGCAAAAG3′), which served as inner primers in the second reaction run. From the first run, 2.0 µl of the PCR products was added to 23 µl of the mixture in the second round of the reaction in a fresh PCR tube. Cycling conditions were the same as the standard PCR except for a 1 °C increase in annealing temperature in the second reaction. A positive control was used in the validation experiment, and a negative control was included in each run [39]. The positive control was DNA extracted from a blood sample of an animal positive by microscopy, whereas the negative control was DNA from a non-infected blood sample. The amplified DNA was resolved on 2% agarose gel, visualized under a UV transilluminator and photographed with a gel documentation apparatus (Bio-Rad Molecular Imager Gel Doc System 170-8170 v.3) for clear visualization and reference purposes.

DNA sequencing and phylogenetic analysis
Twenty microliters of samples positive by PCR were sequenced using the BigDye® Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Foster City, CA, USA), according to the manufacturer’s instructions. The sequences obtained were viewed on the FinchTV v.1.4.0 trace viewer (Applied Biosystems), while flanking regions of high signal-to-noise ratio were trimmed off the sequence to improve the accuracy and precision of sequence data obtained. Ambiguous nucleotides were edited and replaced with conventional ones based on the highest peaks recorded on the electropherogram. Each edited sequence was BLAST-searched against the DNA sequence database (NCBI) and/or the published databases for various trypanosome species (TriTrypDB), and isolates with GenBank hits of 80% or more were considered similar. Sequences of the ITS-1 region were aligned using ClustalW algorithm against known sequences in order to confirm species identity. MEGA (Molecular Evolutionary Genetics Analysis) version 7.0.2.6 was used to construct a phylogenetic tree using the maximum likelihood algorithm with 1000 bootstrap replicates to observe their evolutionary trend and variation over time.

Statistical analysis
The results obtained from this study were subjected to descriptive statistics to determine the frequency and distribution of trypanosome infection across the study area. The prevalence among localities, breeds of cattle, and age and sex of the animals was expressed as percentages of the total number of animals sampled. This was done by dividing the number of infected animals by the total number of animals examined and expressed as confidence level, 50% assumed prevalence and 0.05 tolerable error. In each cluster, animals were considered in randomization regardless of health status so as to give an overall current infection status of the herd. Animals aged one (1) year and younger were considered young calves, while those over one (1) year were regarded as adults. Dentition was used to determine the ages of animals, and body condition scores (BCS) were assessed and adequately scored. Other parameters including breed, sex, source and location of the cattle were recorded. Herd data were collected to include the residency, travel history, herd size, history of trypanocidal treatment and disease history.
percentages. Categorical values were evaluated using the Chi-square test to measure the strength between variables at a 95% confidence interval. All data obtained were analysed using SPSS statistical software version 20.0. Values of \( P < 0.05 \) were considered significant.

**Results**

Across the study area, 72 herd clusters were identified as a sample frame, from which 50% of the herds representing 36 clusters were ballotted by simple random sampling. In all, 398 blood samples were obtained from across the study area, three of which were screened positive by microscopy, representing 0.8% prevalence (Table 1), while 12 samples representing 3.0% tested positive by nested PCR, with distinct band size characteristics of the species involved (Fig. 1). Details on the herd location and number of samples obtained from each herd are contained in Additional file 1: Table S1.

Sequencing of the PCR products and bioinformatics analysis (http://blast.ncbi.nlm.nih.gov/Blast.cgi) further validated the species and subspecies of the trypanosomes (for details on analysed samples and sequences, see Additional file 3: Table S3 and Additional file 4: Dataset S1). To make an evolutionary inference, all isolates aligned with the salivarian group except JO6 (\( T. theileri \)), which fell within the stercorarian group. Small subunit RNA (ssrRNA) from JT11 (\( T. evansi \)), JAD7 (\( T. brucei \)) and JAG2 (\( T. brucei \)) fell within the same branch but different clades, which depicts a logical evolutionary event arising from within these species (Fig. 2). \( T. evansi \) is widely known to have evolved from \( T. brucei \), all of which were rooted on \( T. vivax \).

Within the study area, four different breeds of cattle were found. Table 2 shows the effect of trypanosome infection on the PCV in each of the cattle breeds. For infected animals, PCV estimates followed the order Red Bororo < White Fulani < Sokoto Gudali, while the PCV for the non-infected animals followed the order White Fulani > Sokoto Gudali > Red Bororo > Muturu.

Different trypanosoma species were identified in cattle distributed within the study area as shown in Table 3. Comparatively, the PCV values for non-infected animals were higher than for those infected with individual parasites. The animals infected with \( T. brucei \) were the least affected, with an average PCV of 30.3±0.92, while the animals infected with \( T. theileri \) were the most affected, with an average PCV of 19.2±1.12. Generally, the PCV in regard to species infection followed the order \( T. theileri < T. evansi < T. simiae < T. congolense < T. brucei \) (Table 3). Details on the average PCV obtained for each

| Cluster | Total cluster | Sampling point | Herd size | Total examined | +Ve | −Ve | Prevalence (%) | Species |
|---------|---------------|----------------|-----------|----------------|-----|-----|----------------|---------|
| Positive| 3             | JH             | 183       | 11             | 1   | 10  | 9.1            | T. c spp. |
|         |               | JAA            | 172       | 10             | 1   | 9   | 10             | T. b spp. |
|         |               | JAG            | 92        | 6              | 1   | 5   | 16.7           | T. b spp. |
| Negative| 33            | Others         | 6156      | 371            | 0   | 371 | 0              | Nil      |
| Total   | 36            | 6603           | 398       | 3              | 395 | 0.8 |                |          |

* \( T. c \) spp Trypanosoma congolense species, \( T. b \) spp Trypanosoma brucei species, JH Jebba, cluster H, JAA Jebba, cluster AA, JAG Jebba, cluster AG

**Table 1** Prevalence of trypanosome infection among cattle in Jebba by microscopic examination

**Fig. 1** PCR amplification of trypanosome from blood of cattle using ITS-1 generic primers. a Lane 1 (JA19a and JA19b): A mixed infection with \( T. congolense \) Kilifi and \( T. vivax \); lane 2 (JH4): \( T. congolense \) Savannah; lane 3 (JMB): \( T. congolense \) Savannah; lane 4 (JO6): \( T. theileri \); lane 5 (JO12): \( T. congolense \) Savannah; b lane 6 (JQ6): \( T. simiae \), lane 7 (JT4): \( T. congolense \) Savannah; lane 8 (JT11): \( T. evansi \); lane 9 (JT5): \( T. congolense \) Savannah; lane 10 (JAA7): \( T. brucei \) brucei; lane 11 (JAD7): \( T. congolense \) Forest; lane 12 (JAG2): \( T. brucei \) brucei. *Positive for microscopy. Lanes M: 100 bp superladder-mid (ABgene). Lanes N: negative control

**Fig. 2**
herd and the trypanosome species and subspecies infections are contained in Additional file 2: Table S2, Additional file 5: Table S4 and Additional file 6: Table S5, respectively.

Trypanosome infection rates varied with sex, age, cattle breed, residency/stay period, animal origin, disease and treatment history, travel history, herd size and location of herd relative to the River Niger. As shown

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**Fig. 2** Phylogenetic relationships of trypanosomes within the subgenus Nannomonas clade deduced from small subunit rRNA gene sequences. The phylogram was constructed by bootstrapped (1000 replicates) maximum likelihood (ML) analysis based on the Tamura–Nei model. The tree with the highest log likelihood (~1113.90) is shown. Bootstrap values for all major nodes are given and all branches receiving bootstrap support values > 50%. The tree was drawn to scale, with branch lengths measured in the number of substitutions per site. The analysis involved 39 nucleotide sequences and 77 positions in the final dataset. Evolutionary analyses were conducted in MEGA7.
Table 2 Comparison of mean PCV among cattle breeds infected with Trypanosoma species in the study location (June 2019)

| Cattle breed    | No. of animals infected | Average haematocrit (%) |
|-----------------|-------------------------|-------------------------|
|                 | Infected (mean±SEM)     | Not infected (mean±SEM) |
| Muturu          | 0                       | 32.8±1.23               |
| Red Bororo      | 1                       | 18.3±0.84               |
| Sokoto Gudali   | 3                       | 26.0±1.82               |
| White Fulani    | 8                       | 24.9±1.70               |

PCV values are means of three replicates of the same sample and are expressed as mean±SEM.

Table 3 Comparison of mean PCV of cattle infected with Trypanosoma species in the study location (June 2019)

| S/No. | Species infection | Number of animals infected | PCV (%) |
|-------|-------------------|---------------------------|---------|
| 1     | Non-infected      | 386                       | 36.2±3.71 |
| 2     | T. congolense     | 7                         | 23.8±1.15 |
| 3     | T. brucei         | 2                         | 30.3±0.92 |
| 4     | T. evansi         | 1                         | 20.0±0.32 |
| 5     | T. theileri       | 1                         | 19.2±0.12 |
| 6     | T. simiae         | 1                         | 22.0±0.63 |

PCV values are means of three replicates of the same sample and are expressed as mean±SEM.

Discussion

The low prevalence reported in this study by microscopic screening (Table 1) is not surprising considering the generally low sensitivity of the parasitological diagnostic method [40–42]. This is especially so for field animals characterized by low parasitaemia. The superiority of PCR over the micro-haematocrit centrifugation technique (MHCT) has been widely demonstrated in epidemiological studies of animal trypanosomiasis [3, 32, 43, 44]. These differences are due to the sensitivity thresholds of the techniques. Nested PCR gave an overall prevalence of 3.0%. Each species tested produced amplicons of between 200 and 700 base pairs (bp) in length (Fig. 1a, b). The ITS-1 PCR product size of T. evansi was similar to that of T. brucei, and sequencing analysis was key to differentiating between the two PCR products. Sample JT1 was further confirmed to be T. evansi (Additional file 3: Table S3), suggesting the possible role of cattle as reservoirs of T. evansi. Generally, bands obtained from the amplification results agreed with those of previous studies [15, 22, 45–51]. Samples JH4 and JM8 resulted in band sizes of approximately 700 bp, which were confirmed by sequence analysis to be T. congolense Savannah type. Molecular characterization of Trypanosoma species using ITS-1 generic primers and/or their slight modification has been reported to give an estimated range of ITS-1 band sizes with a maximum amplicon length of 640 bp [15], and it was noted that all species amplification using generic primers could lead to a band size of between 150 and 750 bp in length, as evidenced in this study and those previously reported [29, 45, 50, 52]. An amplicon length of 210 bp for T. vivax was not reported previously, which may be an indication that T. vivax 18S rRNA is more rapidly evolving than that of other non-salivarian trypanosomes and also evolving significantly faster than all other trypanosomes [53]. Further biological characterization will depend on isolation of a living specimen into culture. The ability to identify this trypanosome by the distinct size of the ITS-1 region has provided preliminary information on its distribution that should help track it in the field.

A drastic decrease in PCV has traditionally been considered a warning sign of trypanosomiasis [54, 55]. Classically, infection with trypanosome species that are pathogenic in local breeds of cattle result in retarded growth and anaemia, and nutritional status is a determining factor of infection [56, 57]. From our findings, PCV was decreased in trypanosome-positive animals (Table 3; Additional file 5: Table S4), possibly due to the effect of parasites on blood cells. Similarly, the average haematocrit values varied between cattle breeds (Table 2). However, the very low PCV presented by the Red Bororo breed may not be a true reflection of

in Table 4, the infection rate was higher in female animals (4.3%), young animals (10.0%), White Fulani cattle breeds (3.7%), animals that were recently domiciled ≤3 years (3.2%), transhumance animals (3.6%) and animals having a disease history (4.1%). Similarly, in Table 5, the infection rate was higher among herds closer to the River Niger (56.2%), larger herds (33.3%) and animals that had travelled to endemic zones (50.0%). Aside from the age of animals and location distance from the River Niger, the differences in the infection rates for all tested parameters were not statistically significant. In Fig. 3a, our results show a single infection rate of 91.7% as against 8.3% recorded for mixed infection, with T. congolense being the most prevalent (50.0%), followed by T. brucei brucei (16.7%), while other species had equal prevalence (8.3%). In the study area, the infection rate with T. congolense Savannah type was comparatively higher (71.4%) than other types, as shown in Fig. 3b (see also Additional file 6: Table S5). The spatial distribution of trypanosome-infected herds across the study area is shown in Fig. 4.
the PCV trend, as only one animal was screened positive. Trypanosome species had different effects on the PCV status of animals, with the average falling below the standard obtainable for cattle (24–46%), except for the animals infected with *T. brucei*, with an average PCV of 30.3 ± 0.92. The animal infected with *T. theileri*, a non-pathogenic trypanosome of cattle, had the lowest PCV value (19.2 ± 0.12), but it is difficult to conclude that this decrease was actually caused by *T. theileri* (Table 3). Co-infections or other possible causes of decreased PCV cannot be ruled out. However, it is possible that the parasite may have transited from a non-pathogenic to pathogenic form, hence the need for a controlled experiment aimed at monitoring the PCV in

| Table 4 | Parametric prevalence of trypanosome infection based on sex, age, breed, animal residency, origin, disease history and treatment history (June 2019) |
|---|---|---|---|---|---|
| Category | Subcategory | No. of cattle screened | No. of trypanosome infections | Prevalence (%) | $\chi^2$ | P-value |
| Sex | Male | 165 | 2 | 1.2 | 3.133 | 0.077 |
| | Female | 233 | 10 | 4.3 | 0.0 | 1.0 |
| | Total | 398 | 12 | 3.0 | 0.983 | 0.321 |
| Age (years) | ≤ 1 | 70 | 7 | 10.0 | 14.172 | 0.000 |
| | > 1 | 328 | 5 | 1.5 | 0.0 | 1.0 |
| | Total | 398 | 12 | 3.0 | 0.983 | 0.321 |
| Breed | W.F. | 214 | 8 | 3.7 | 1.172 | 0.760 |
| | S.G. | 130 | 3 | 2.3 | 0.0 | 1.0 |
| | R.B. | 36 | 1 | 2.8 | 0.0 | 1.0 |
| | MUT | 18 | 0 | 0.0 | 0.0 | 1.0 |
| | Total | 398 | 12 | 3.0 | 0.983 | 0.321 |
| Animal residency (years) | ≤ 3 | 126 | 4 | 3.2 | 0.016 | 0.899 |
| | > 3 | 272 | 8 | 2.9 | 0.0 | 1.0 |
| | Total | 398 | 12 | 3.0 | 0.983 | 0.321 |
| Animal origin | Sentinel | 174 | 4 | 2.3 | 0.542 | 0.461 |
| | Transhumance | 224 | 8 | 3.6 | 0.0 | 1.0 |
| | Total | 398 | 12 | 3.0 | 0.983 | 0.321 |
| Disease history | History | 148 | 6 | 4.1 | 0.870 | 0.351 |
| | No history | 250 | 6 | 2.4 | 0.0 | 1.0 |
| | Total | 398 | 12 | 3.0 | 0.983 | 0.321 |
| Treatment history | History | 366 | 11 | 3.0 | 0.001 | 0.970 |
| | No history | 32 | 1 | 3.1 | 0.0 | 1.0 |
| | Total | 398 | 12 | 3.0 | 0.983 | 0.321 |

*WF* White Fulani, *RB* Red Bororo, *SK* Sokoto Gudali, *MUT* Muturu. $\chi^2 = $ Chi-square test of association was carried out at a 95% confidence interval

| Table 5 | Parametric prevalence of trypanosome infection based on location, herd size and travel history to endemic zones (June 2019) |
|---|---|---|---|---|---|
| Category | Subcategory | No. of herds screened | No. of herd(s) infected | Prevalence (%) | $\chi^2$ | P-value |
| Location (Km) (distance from the River Niger) | Far (> 3 km) | 20 | 1 | 5.0 | 11.638 | 0.001 |
| | Close (≤ 3 km) | 16 | 9 | 56.2 | 0.0 | 1.0 |
| | Total | 36 | 10 | 27.8 | 0.0 | 1.0 |
| Herd size | Large ≥ 200 | 15 | 5 | 33.3 | 0.396 | 0.529 |
| | Small < 200 | 21 | 5 | 23.8 | 0.0 | 1.0 |
| | Total | 36 | 10 | 27.8 | 0.0 | 1.0 |
| Travel history to endemic areas | Endemic | 8 | 4 | 50.0 | 2.532 | 0.112 |
| | Not endemic | 28 | 6 | 21.4 | 0.0 | 1.0 |
| | Total | 36 | 10 | 27.8 | 0.0 | 1.0 |

*Travel history: Kaduna, Ogun, Oyo, Jos, Benue and Delta. $\chi^2 = $ Chi-square test of association was carried out at 95% confidence interval*
the face of *T. theileri* infection and other trypanosome species.

This study showed 3.0% overall prevalence, which compares well with the 4.3% national prevalence as reported by the European Economic Commission project of 1989 and 1996 [58], 3.9% in Ogbomosho [59], 4.7% in Oyo [60] and 9.4% in Kaduna [61], as against high prevalence of 53.4% in Kaura [62] and 46.8% in Jos [63]. These findings might reflect seasonal or local differences in tsetse populations, sample size and site, improved sensitization among nomads on grazing courses, better implementation of the use of trypanocidal drugs and urbanization.

Our results showed that females were more commonly infected (Table 4). This observed difference may be attributed to livestock management practices adopted in the farming community, wherein larger numbers of males are frequently sold off the herd at an early age while the rest are kept for breeding or animal traction. Also, females persist longer in herds for the purpose of breeding, thus allowing the chronic infection to be maintained for very long period. As a result, the remaining males are more
closely monitored while the females are readily exposed to hazard in the population vis-à-vis multiple copulation with limited male animals in the herd. Also, the larger population of females (59%) obtained in this study by simple random sampling may account for this difference. Previously, 199 males and 121 females were examined in the Ido Local Government Area (LGA), Oyo State, Nigeria, with no statistical difference in the infection prevalence [60]. Factors other than sex relating to the host or its environment could therefore have played a role in the susceptibility of animals to infection, which has been documented in several studies [8, 10, 64].

From our study, there is a decrease in disease prevalence as animals age, probably due to age-acquired immunity, which could represent a key positive factor, and bearing in mind that trypanocidal treatments are more frequently used on adults by local farmers. In addition, young animals are more vulnerable to tsetse bites due to their skin fragility. Moreover, they are not agile enough to ward off insects along the grazing route as are the adults. The tsetse flies also frequently target weak animals as a source of food in order to avoid being crushed by moving animals [65]. In this study, despite the very low number of young animals (17.6%) randomly sampled (Table 4), a prevalence of 10.0% (7/70) was recorded for younger animals and 1.5% (5/328) for adults, which is statistically significant ($\chi^2 = 14.172$, $df = 1$, $P < 0.001$). This indicates that the incidence rate was not similar between young and adult animals [8]. The prevalence of trypanosome infection was lowest in Sokoto Gudali (2.3%), a breed not known for being tolerant to trypanosome infection [66, 67]. The higher prevalence observed among the White Fulani breed may be attributed to their known susceptibility and perhaps due to their higher representation in the sampling (52.8%). Of the four cattle breeds studied, the White Fulani are usually raised under the nomadic system of management. This may be another possible explanation for the higher prevalence recorded in this cattle group [10].

The prevalence of trypanosome infection was higher in locations closer to the River Niger as compared to those further away (Table 5; Fig. 4). This difference is significant at a 95% confidence interval ($\chi^2 = 11.638$, $df = 1$, $P < 0.05$) and may be attributed to differences in herd management practices, grazing routes which predispose the herd to tsetse bites and herd composition which may differ in each herd. The river and the vegetation along its course could be a positive factor for the vector transmitting the disease as well as a source of water for grazing animals, which could expose them to risk of bite by riverine species of the flies [68].

The majority of the trypanosomes in cattle were \textit{T. congolense} and \textit{T. brucei}, with nearly half of the overall infection due to \textit{T. congolense} Savannah type (Fig. 3b), possibly as a result of the large host range or likely due to the fact that riverine species of tsetse are generally considered susceptible to \textit{T. congolense} infections [69]. The relatively low prevalence of \textit{T. brucei} as compared to \textit{T. congolense} may be associated with the poor efficiency of the generic ITS-1 primers in identifying members of the subgenus \textit{Trypanozoon} (e.g., \textit{T. brucei} subspecies) [70]. The high prevalence of \textit{T. congolense} infection is an indication of the dominance of the \textit{Glossina morsitans} tsetse fly [7, 8, 30, 32, 63, 71, 72] and could mean that its transmission is highly favoured by the obligate cyclical vector or that \textit{T. vivax} and \textit{T. brucei} respond better to the trypanocidal drugs, diminazene aceturate and homidium chloride, indiscriminately administered by farmers. A high prevalence of the Savannah subgroup in cattle may also indicate that the parasites were introduced recently into the tested herds, coupled with its reported virulence as compared to other subgroups [73]. The low prevalence of \textit{T. brucei} infection may relate to the reported resistance of indigenous West African cattle to the parasite [74]. However, the detection of \textit{T. brucei} and \textit{T. evansi} in Nigerian cattle might portend serious danger not only to cattle and other livestock but also to livestock owners and the host communities at large, as \textit{T. evansi} infection has been reported in cattle and humans [75, 76]. The low prevalence of \textit{T. simiae} infection is an indication of low transmission of the parasite, as animals infected with this species will probably not survive the acute and severe nature of the effects of this parasite [77, 78]. Double infections in animals are a normal occurrence in the field [79]. This study identified \textit{T. congolense} (Kilifi type) and \textit{T. vivax} mixed infection in only one of the tested herds. In Nigeria, previous surveys identified mainly \textit{T. congolense} and \textit{T. vivax} as animal-pathogenic trypanosomes [80], and co-circulation has been reported in studies conducted in northern Nigeria [22, 32, 81]. Co-infections with multiple \textit{Trypanosoma} species have also been documented previously due to bites from tsetse flies carrying more than one \textit{Trypanosoma} infection or successive bites from flies with different \textit{Trypanosoma} species [8, 82, 83].

**Conclusions**

Six species of trypanosomes were identified in cattle herds along the Jebba axis of the River Niger, with \textit{T. congolense} being the most prevalent. Age and relative distance of herds from the River Niger may be risk factors for trypanosome infection in cattle herds along the Jebba axis of the River Niger. According to the threshold of the European Economic Commission (i.e., 4.3%), the study area may not be classified as endemic, but our data suggest that the cattle population may play a role in the possible resurgence of the disease in this region.
Abbreviations
AT: African trypanosomiasis; AAT: African animal trypanosomiasis; HAT: Human African trypanosomiasis; PCR: Polymerase chain reaction; ITS-1: Internally transcribed spacers-1; HCT: Haematocrit centrifugation technique; BCVM: Buffy coat method; PCV: Packed cell volume.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s13071-021-05054-0.

Additional file 1: Table S1. Cluster data of herds screened for Trypanosoma infection in Jebba, Kwara State, Nigeria.
Additional file 2: Table S2. Cluster prevalence of Trypanosoma infection and average PCV of cattle in Jebba, Kwara State, Nigeria (June 2019).
Additional file 3: Table S3. Sequence similarity of ITS-1 DNA sequences of trypanosomes isolated from cattle with other sequences available in GenBank.
Additional file 4: Dataset S1: The DNA sequence data of Trypanosoma species obtained by BigDye Terminator Cycle Sequencing Kits v3.1.
Additional file 5: Table S4. Prevalence of Trypanosoma species infection among cattle in Jebba, Kwara State, Nigeria (June 2019).
Additional file 6: Table S5. Prevalence of Trypanosoma congolense infection (according to types: Kilifi, Savannah, and Forest), among cattle in Jebba, Kwara State, Nigeria (June 2019).

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Authors' contributions
IFH conceived and designed the study, collected samples, analysed samples by PCR and sequencing, analysed and interpreted the data, and drafted and reviewed the manuscript. GDC reviewed and edited the manuscript, facilitated the support of traditional and administrative authorities, and supervised the fieldwork and laboratory experiments. JKP designed the project, facilitated the support of traditional and administrative authorities, and supervised the fieldwork and laboratory experiments. All authors read and approved the final manuscript.

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Availability of data and materials
All data generated or analysed during this study are included in this published article and its supplementary data files.

Declarations
Ethics approval and consent to participate
Approval to collect blood from cattle was obtained from the community head, local cattle breeders and the Kwara State Ministry of Agriculture and Rural Development.

Consent for publication
Not applicable.

Competing interests
There are no competing interests in this research.

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