

Short Research Article

Phylogeographical Analysis and Visualization of Data of *Trypanosoma* Isolates from different Foci in Nigeria Using Microreact and Phandango (2003 – 2018)

Abstract

Introduction: A combination of phylogenetic tree of trypanosome isolates and the geographical information of the epidemiologic foci can acquaint us with the occurrences of African trypanosomiasis outbreak necessitous for educating the spatial distribution of the parasites, vectors and drawing inferences for planning strategic control programs. Microreact and Phandango are free online tools that enables the visual representation and study of genomic epidemiology of infectious diseases. This study elicits the phylogenetic data of trypanosome isolates and the specific host along with geographical information

Methods: A total of 46 isolates. Protein sequence data of trypanosomes isolated in Nigeria between 2003 and 2018 were extracted from National Center for Biotechnology Information (NCBI) database. Multiple sequence alignment and tree construction were carried out on sequence data files and Newick output was downloaded. Metadata .csv files were prepared accordingly and loaded for visualization in microreact and phandango.

Results The preponderance of *Glossina palpalis* was reported in old Oyo National Park and they were majorly incriminated in the transmission of *Trypanosoma gravi* in *Glossina palpalis*. *Trypanosoma theileri* was isolated from the midgut of *Glossina morsitans* reported to be found in Kainji Lake National Park only. *Trypanosoma congolense* was reportedly isolated from cattle at Zaria, *Glossina morsitans* at Kainji Lake National Park, *Glossina tachinoid* at old Oyo National Park at Oyo and, *Glossina palpalis* at Yankari Game Reserve Bauchi. *Trypanosoma brucei brucei* was isolated from a rat in April 2018 at Federe and Vom, Plateau state. In Federe and Vom, *Trypanosoma brucei brucei* was found in rat in April 2018 while *Trypanosoma evansi* was obtained from animal in Vom December 2008. In Bida and Tsuaa, *Trypanosoma brucei gambiense* was isolated from unidentified animals (probably cattles) in October 2009. The analyses also revealed 7 protein sequence which includes 7 protein sequenced including cathepsin L-like protein (CPL) and glycosomal glyceraldehyde-3-phosphate dehydrogenase (G-GAPDH) from *Trypanosoma vivi* and *Trypanosoma congolense*, glycoprotein (GP) from *Trypanosoma brucei gambiense*; trans-sialidase (TSS) from *Trypanosoma evansi*; phospholipase A2 (PLA2) and hexokinase 1 (HK1) from *Trypanosoma brucei brucei*; and also glycosomal glyceraldehyde-3-phosphate dehydrogenase (G-GAPDH) from *Trypanosoma theileri*.

Comment [u1]: No mention of *Trypanosoma sius/simiae*, which I believe is a major species affect the porcine species both domestic and wild in Nigeria.

Conclusion: The study generated a concise visual representation and trajectory of trypanosome isolates in Nigeria along with their genetic metadata and geographic location on a phylogenetic tree which will serve as a reference for information on trypanosomiasis outbreaks in Nigeria, provide blue prints for future outbreak predictions and guide decision-making on strategic control programs.

Keywords: Phylogenetics, Geographical Information, Epidemiological foci, Hosts, Trypanosome

Introduction

Trypanosomes are members of protozoans of the genus *Trypanosoma* which proliferate in the body fluids mostly blood of the vertebrate hosts such as man, domestic and wild animals. They reside in the salivary glands or alimentary canal of the invertebrate hosts, precisely tsetse flies and other biting insects (Bouyer et al 2012). Trypanosomes infection often vary with hosts (Odeniran et al 2019). Notable trypanosomes species such as *T. congolense*, *T. vivax*, *T. brucei*, *T. grayi*, *T. theileri*, *T. simiae*, etc. infect both vertebrate and invertebrate hosts (Isaac et al., 2016; Odeniran et al., 2019; Weber et al., 2019; Omonona et al., 2020).

Wildlife is an important natural resource and it plays an essential role in sustaining the livelihoods of communities in sub-Saharan Africa (Odeniran et al., 2018b). Consequent to the suitable habitat provided by the national park for the flies and the pathogenic trypanosome, the wild animal in the park and domestic livestock grazing around are steadily exposed to the risk of trypanosomiasis. Cases of tsetse flies has been documented in the national parks and game reserves where diverse species of living organisms are protected (Ajibade and Agbede, 2008; Okoh et al., 2011; OIE, 2013; Isaac et al., 2016; Shaida et al., 2018; Wama et al., 2018; Omonona et al., 2020).

Factors militating the health and productivity of wild animals constitute hindrance to a developed and stable ecosystem (Omonona et al., 2020). Morbidity and occasional mortality in wild animals have been reported to be a major result of trypanosomiasis transmitted by tsetse flies (OIE, 2013). While wild animals are usually the reservoir hosts, some species of tsetse fly have been attributed to certain species of wildlife (Omonona et al., 2020). A combination of phylogenetic tree of trypanosome isolates and the geographical information of the epidemiologic foci can serve as a tool that provides critical information regarding the occurrences of African trypanosomiasis outbreak necessitous for educating the spatial distribution of the parasites, vectors as well as drawing inferences for planning strategic control programs. Locally and globally, Phylogenetics serve as an essential tool for identifying disease outbreaks and monitoring the spread of a pathogen. This tool is capable of monitoring the spread or distribution of isolates as it can infer relationship between samples collected at various times, sources and places.

The combination of high-resolution information on the phenotypic and genotypic relationships of isolates with corresponding data, can be valuable in understanding the distribution of the vectors, parasites, history of outbreaks; secondarily, provide inferences towards identification of enabling conditions contributing to the occurrences and distribution of the disease over a specific period.

Microreact is a free online tool that will enable you to visualize phylogenetic relationships of isolates linked to geographic locations. This tool allows for dynamic visualization of the data with an interactive map using your own phylogenetic tree and metadata (Argimón *et al.*, 2016).

Phandango is an interactive visualisation tool for phylogenetics trees and can be viewed with other information that would benefit from a visual output such as metadata (e.g. resistance, virulence, serotyping and multi locus sequence typing or MLST data) and genomic structure information (e.g pangenome, recombination blocks, genome wide association studies or GWAS) (Hadfield *et al.*, 2017).

This project is aimed at generating visual chart for the phylogenetic data combined with geographical information to navigate the vectors/hosts and *Trypanosoma* species in specific existing foci using microreact and phandango using protein sequence of *Trypanosoma* species found in Nigeria between 2003 – 2018 datasets obtained from National Center for Biotechnology Information (NCBI) website.

Materials and Methods

Data analysis

Protein sequence of Trypanosome isolates (represented by accession number) found in Nigeria between 2003 and 2018 were downloaded from National Center for Biotechnology Information (NCBI) database in 15th, September, 2020.

Multiple sequence alignment was performed using Multiple Alignment using Fast Fourier Transform (MAFFT). Phylogenetic tree was constructed and estimated by Phylogeny Maximum Likelihood (PhyML option in Seaview version 2.0), with blossom62 model and bootstrap set to 100 replicates as described by Guindon *et al.* 2010. The output was saved in NEWICK version (.nwk format).

Geographical Information of isolate origin

Geographic coordinates of epidemiological foci of the trypanosome isolates were generated with the web-based Google Earth application (Google Inc., California, USA).

Metadata File for Phandango preparation

A metadata file (.csv format) was prepared (in excel readable by phandango server <https://jameshadfield.github.io/phandango/#>) from the protein sequenced data using information such as accession number, years, month and of collection, location of collection, amino acids sequenced and identity of protein sequenced. Protocols for entry data found in https://en.wikipedia.org/wiki/Newick_format were adhere to as describe by Hadfield *et al.*, 2017. The .csv file was saved as comma delimited format.

Metadata File for Microreact preparation

A metadata file (.csv format) prepared in excel readable by microreact server (<https://microreact.org/>) from the protein sequenced data which contains details such as accession number, years, month and of collection, location of collection, amino acids sequenced

Phandango chart Figure 1.2 consist of 46 isolates with 7 different sequence proteins, 7 different *Trypanosoma* species which were obtained from 3 isolation sources (blood, gut and proboscis) from 7 vectors/hosts found in 10 foci across the Nigeria.

Comparing the genetic relationship of isolates (represented by accession numbers) and the hosts/*Trypanosoma*, isolates that are in group in the tree, have similar grouping in organism name (orgname).

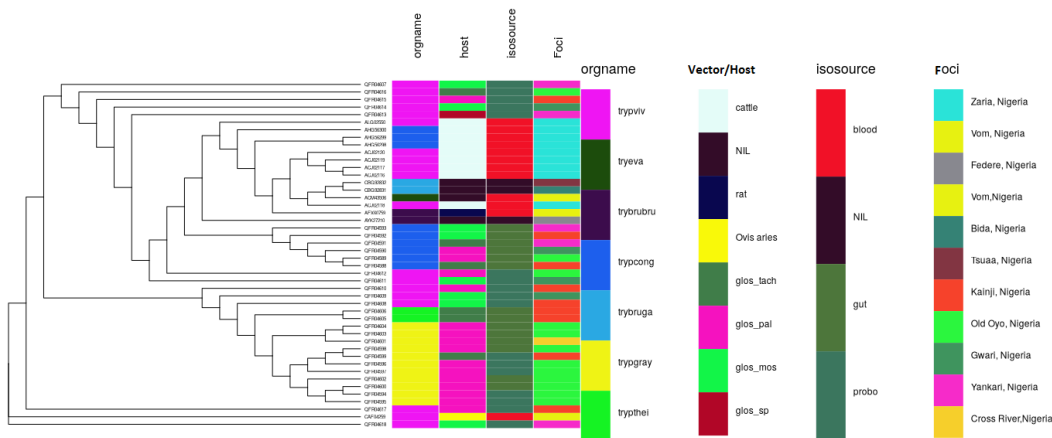


Figure 1.2: Phandango chart of phylogenetic tree of isolates in relationship with *Trypanosoma* species, hosts, foci and isolation source.

Figure 1.3 contain the visualized chart of phylogenetic tree in relation to protein sequenced from *Trypanosoma* species and we can deduced that cathepsin L-like protein (CPL) and glycosomal glyceraldehyde-3-phosphate dehydrogenase (G-GAPDH) are sequenced from *Trypanosoma vivi* and *Trypanosoma congolensis*; glycoprotein (GP) is sequenced from *Trypanosoma brucei gambiensis*; trans-sialidase (TSs) from *Trypanosoma evansi*; phospholipase A2 (PLA2) and hexokinase 1 (HK1) from *Trypanosoma brucei brucei*; and also glycosomal glyceraldehyde-3-phosphate dehydrogenase (G-GAPDH) from *Trypanosoma theileri*.

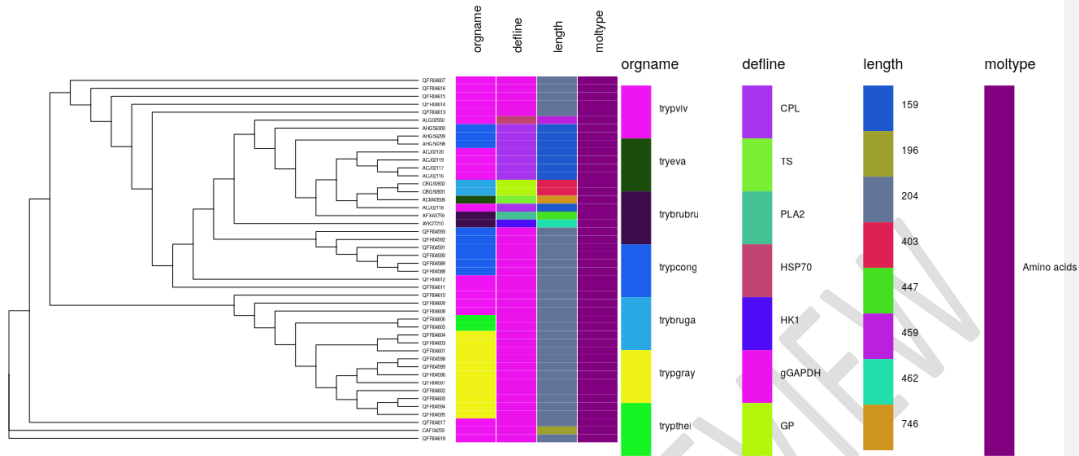
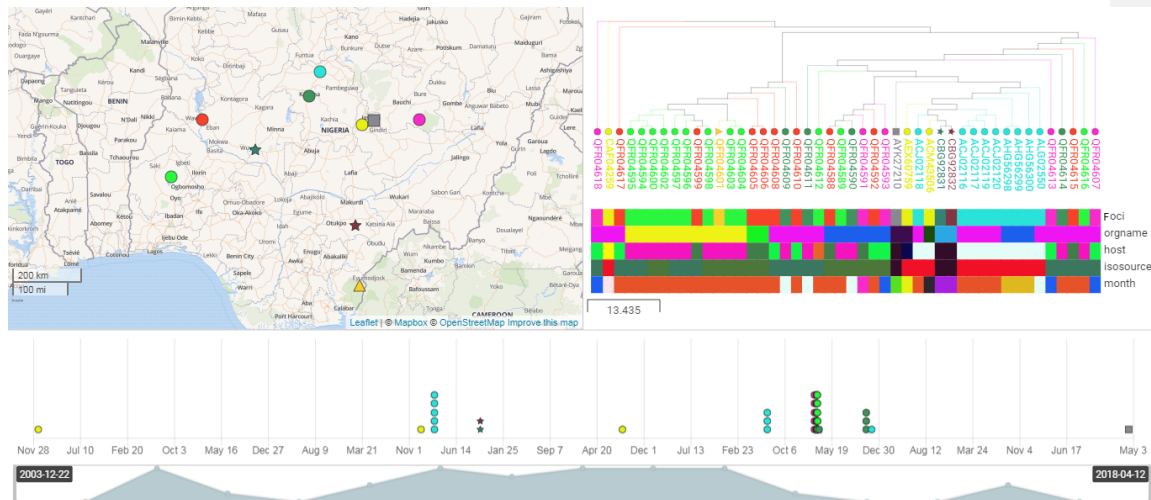


Figure 1.3: Phandango chart of phylogenetic tree of isolates in relationship with *Trypanosoma* species, protein sequenced and others.

Phylogenetic and Microreact Visualization

The chart consists of four components including the map panel on the left, the phylogenetic tree on the right panel, the legend and the timeline at the bottom panel.



Abbreviations: organism name (orgname); *Trypanosoma congolense* (trypcong), *Trypanosoma grayi* (trypgray), *Trypanosoma brucei brucei* (trybrubu), *Trypanosoma evansi* (tryeva), *Trypanosoma theileri* (trypthei), *Trypanosoma vivax* (trypviv), *Glossina* sp. (glos_sp), *Trypanosoma brucei gambiense* (trybruga); **Host**– *Glossina tachenoid* (glos_tach), *Glossina palpalis* (glos_pal), *Glossina morsitans* (glos_mos), *Glossina* sp. (glos_sp); (isolation source) (isosource), proboscis (probo)

<https://microreact.org/project/4SM4tVyb4FHfAo6PsWqKfZ/87dcfa8d>

Figure 1.4: Microreact chart embedded with phandango metadata information.

The microreact and phandango chart show the relationship of phylogeny with various species of *Trypanosoma* isolated from sources in diverse hosts at different foci of unique geographical information and date as shown in Figure 1.4. *Trypanosoma vivi* (purple) are from isolates (ACJ02116, ACJ02117, ACJ02118, ACJ02119, ACJ02120, ALG02550, CAF04259, QFR04607, QFR04608, QFR04609, QFR04610, QFR04611, QFR04612, QFR04613, QFR04614, QFR04615, QFR04616, QFR04617, QFR04618) found in diverse hosts and different foci. They are isolated from the blood of cattle in Zaria in march 2009; proboscis of *Glossina morsitan* in Yankari Game Reserve, Ijah Gwari and Kainji Lake National Park in 2014; proboscis of *Glossina palpalis* in the Old Oyo National Park and Kainji Lake National Park in 2014; and in the blood of *Ovis aries* (sheep) in Vom, Nigeria in 2003.

Trypanosoma evansi (deep green) from isolate (ACM43506) is obtained from the blood of unidentified host in Vom, Nigeria in December, 2008. *Trypanosoma brucei brucei* (dark blue) (AEX60759, AYK27210) are identified in the blood of rat in Vom in august, 2011; and in unidentified host in Federe, Nigeria in April, 2018. *Trypanosoma congolensis* (blue) (AHG56298, AHG56299, AHG56300, QFR04588, QFR04589, QFR04590, QFR04591, QFR04592, QFR04593) are as well isolated from the blood of cattle in Zaria in July, 2013; guts of *Glossina morsitan* and *Glossina tachnoid* in Yankari Game Reserve and Kainji Lake National Park in 2014; guts of *Glossina palpalis* in Ijah Gwari and Old Oyo National Park, 2014. *Trypanosoma brucei gambiense* (light blue) (CBG92831, CBG92832) are obtained from unidentified host in Tsuaa and Bida, Nigeria October, 2009. *Trypanosoma grayi* (yellow) (QFR04594, QFR04595, QFR04596, QFR04597, QFR04598, QFR04599, QFR04600, QFR04601, QFR04602, QFR04603, QFR04604) are isolated from the guts of *Glossina palpalis* in Old Oyo National Park and Cross River Park in March, 2014; proboscis of *Glossina palpalis* in Old Oyo National Park, 2014; and also proboscis of *Glossina tachnoid* in Kainji Lake National Park in March 2014. *Trypanosoma theileri* (green) (QFR04605, QFR04606) are only obtained from the guts of *Glossina tachnoid* in Kainji Lake National Park in March 2014.

Glossina palpalis are readily abundant in Old Oyo National Park and they transmitted mostly *Trypanosoma grayi*. The large grazing activities of cattle in Zaria could be responsible for the large number of *Trypanosoma vivi* parasites recorded. *Trypanosoma theileri* isolated from *Glossina morsitan* are only found in Kainji Lake National Park. *Trypanosoma congolensis* are identified to be readily isolated in the blood of cattle, *Glossina morsitan*, *Glossina tachnoides* and *Glossina palpalis* in Zaria, Old Oyo National Park and Yankari Game Reserve respectively.

Relationship of Vector/host Distribution to the Isolates and the Location with respect to Chart Phylogeny, Timeline and Map.

Vectors are sensitive to climate change and other environmental factors which together affect Trypanosomiasis by influencing their distribution. Increased understanding in the population structure of *Glossina* species and other hosts in the different vegetation zones of Nigeria will aid decisions on appropriate control strategies (Shaida *et al.*, 2018).

In figure 1.4, the chart consist of three major informative panel. The phylogenetic tree panel describes the isolate phylogeny and coloured based on the location/foci of isolation. The timeline panel at the bottom describe the (months, days and years of isolation) season of prevalence of vectors and the map panel gives the geographical information of the foci of vectors and isolates.

The chart in figure 1.4 mainly used the information on phylogeny isolates and foci to explain vector distribution through accurate interpretation of the data displayed on the panels.

The chart shown phylogeny isolates including (QFR04607, QFR04613, QFR04593, QFR04618) and (QFR04592, QFR04610, QFR04608, QFR04617) found in Yankari Game Reserve and Kainji Lake National Park foci respectively showed that *Glossina morsitan* vector was geographically restrained to the foci forming cluster on the timeline (February, 2014 – March 2014). The clustered explained that the vectors are in the same foci or region. Yankari Game Reserve is situated in Bauchi State within the Northern Guinea/Sudan savannah vegetation zone and is located between latitude 9° 45' N and longitude 10° 30' E. Kainji Lake National Park covers an area of 5340 km² across Niger and Kwara States (10° 22' N and 4° 33' E), within the southern guinea savannah vegetation zone. The morsitans tsetse fly formed the primary vector of trypanosomiasis that thrive mostly in savannah. The species identified in an earlier study (Isaac *et al.*, 2016; Shaida *et al.*, 2018) was *Glossina morsitan submorsitans*, which corroborates our results.

The chart also displayed phylogenetic isolates including (QFR04614, QFR04590, QFR04611, QFR04609), (QFR04616, QFR04589, QFR04604, QFR04603, QFR04598, QFR04596, QFR0497, QFR04602, QFR04600, QFR04594, QFR04595), and (QFR04601) found in Ijah Gwari, Old Oyo National Park, and Cross River National Park respectively showed that *Glossina palpalis* vector was geographically restricted to specific foci forming clusters in the timeline (2013-2014 and 2014-2015). Old Oyo National Park is located in the northern part of Oyo State, South Western Nigeria. It has a total land area of 2512 km². The Park lies within the derived savannah vegetation zone between latitudes (8° 10' - 9° 05' N) and longitudes (3° 00' - 4° 02' E). Cross River National Park is located in the rainforest ecological zone in the extreme South East of Nigeria on the border with Republic of Cameroon. The park occupies a total land area of about 4000 km² of tropical rain forest ecosystem and lies between latitude (5° 05' - 6° 29' N) and longitude (8° 15' - 9° 30' E) (NPN, 2004). Ijah Gwari (near Suleja) is located between latitude 9° 12' N and 9° 24' N and longitude 7° 12' E and 7° 20' E in Tafa Local Government Area of Niger State. It is characterized by several small streams traverse the area and the vegetation is riverine fringing forest forming a dense two-storey canopy. Timeline information explained that *Glossina palpalis* vector was readily found in Old Oyo National Park, and Cross River National Park around March, 2014, but however was found abundant in November, 2014 in riverine fringing forest of Ijah Gwari due to seasonal fluctuation. The fact that *Glossina palpalis* vector is readily found in different habitat in distinct timeline may be attributed to variations in microclimatic conditions at the sampling spots (Shaida *et al.*, 2018).

The information in the chart shown the phylogeny isolates (QFR04615, QFR04588, QFR04606, QFR04605, QFR04599) and (QFR04591) found in the Kainji Lake National Park and Yankari Game Reserve respectively explained that the *Glossina tachinoides* vectors were geographically

restrained as shown by the clustered in the timeline March, 2014. Kainji Lake National Park is characterized by thick riverine vegetation, gallery forest and game animals which formed suitable habitat for *Glossina tachinoides* vectors. The coexistence of more than one vectors in the same spot could be attributed to variations in microclimatic conditions at the sampling spots (Shaida *et al.*, 2018).

The chart also displayed phylogeny isolates (ALG02550, AHG56298, AHG56299, AHG56300, ACJ02120, ACJ02119, ACJ02118, ACJ02117, ACJ02116), (ACM43506, AEX60759, CAF04259), (CBG92832), (CBG92831) and (AYK27201) found in Zaria, Vom, Tsua, Bida and Federe respectively which shown that the isolates from hosts such as cattle, sheep, etc were geographically spread across foci in timeline (December, 2003 – April, 2018), hence they were seen in different location in Nigeria due to seasonal variation of green pasture.

Conclusion

This study develop chart that can be used by field researchers to navigate (identify) the existing *Trypanosoma* foci in Nigeria and predict isolation source in vectors/hosts and parasites which can be found in diverse foci. The study generated a concise visual representation and trajectory of trypanosome isolates in Nigeria along with their genetic metadata and geographic location on a phylogenetic tree which will serve as a reference for information on trypanosomiasis outbreaks in Nigeria, provide blue prints for future outbreak predictions and guide decision-making on strategic control programs.

Limitation in this study

The exact geographical locations were not known as the latitudes and longitudes of the foci were not parts of the information deposited in NCBI websites.

Recommendation

Geographical information system of isolation should be included among the information imputed in the NCBI website.

<https://microreact.org/project/4SM4tVy4b4FHfAo6PsWqKfZ/87dcfa8d>

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