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Detection and genetic diversity of Mopeia virus in *Mastomys natalensis* from different habitats in the Limpopo National Park, Mozambique

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**ABSTRACT**

Mammarenaviruses have been a growing concern for public health in Africa since the 1970s when Lassa virus cases in humans were first described in west Africa. In southern Africa, a single outbreak of Lujo virus was reported to date in South Africa in 2008 with a case fatality rate of 80%. The natural reservoir of Lassa virus is *Mastomys natalensis* while for the Lujo virus the natural host has yet to be identified. Mopeia virus was described for the first time in *M. natalensis* in the central Mozambique in 1977 but few studies have been conducted in the region. In this study, rodents were trapped between March and November 2019 in villages, croplands fields and mopane woodland forest. The aim was to assess the potential circulation and to evaluate the genetic diversity of mammarenaviruses in *M. natalensis* trapped in the Limpopo National Park and its buffer zone in Massingir district, Mozambique. A total of 534 *M. natalensis* were screened by RT-PCR and the overall proportion of positive individuals was 16.9%. No significant differences were detected between the sampled habitats ($\chi^2 = 0.018; DF = 1; p = 0.893$). The Mopeia virus (bootstrap value 91%) was the Mammarenavirus circulating in the study area sites, forming a specific sub-clade with eight different sub-clusters. We concluded that Mopeia virus circulates in all habitats investigated and it forms a different sub-clade to the one reported in central Mozambique in 1977.

1. Introduction

Mammarenaviruses are enveloped, ambisense or negative single-strand RNA microorganisms and their genome (~10.5 kb) consists in two or three RNA segments named large (L), medium (M) and small (S). Currently, there are four genera in the Arenaviridae family (*Mammarenavirus*, *Antennavirus*, *Hamartnavirus* and *Reptararenavirus*). The genus *Mammarenavirus* infects mainly mammals including rodents (*Radoski et al., 2019*). In Africa the viruses of the *Mammarenavirus* genus have been a growing concern for public health since the 1970s when the first cases of human disease associated with haemorrhagic fevers caused by the *Lassa virus* were described in west Africa (*Buckley et al., 1970; Frame et al., 1970*). Based on geographical occurrence, genetic and serological features, mammarenaviruses are divided into old world group (OWG) and new world group (NWG) (*Bowen et al., 1997; Albarino et al., 1998*). The OWG is restricted to the Eastern Hemisphere while the NWG occur in the Western Hemisphere. The *Lympohycytic chorioniometitis virus* an OWG *Mammarenavirus* has a worldwide...
distribution (Gratz, 2006; Fornůsková et al., 2021). To date, 16 species of Mammarenavirus have been isolated from African mainland and all belong to the OWG (Burrell et al., 2017; Radoshitzky et al., 2019). Two of them, Lassa virus and Lujo virus, are associated with human lethal diseases. Lassa virus is responsible for 300,000 to 500,000 infection cases resulting in about 5000 deaths each year in west Africa (Ogbu et al., 2007) while a single outbreak of Lujo virus has been reported to date in South Africa in 2008 where four out of the five infected patients died (Paweska et al., 2009). The natural reservoir of Lassa virus is the rodent M. natalensis (Monath et al., 1974) while for the Lujo virus the natural host is yet to be identified (Simulundu et al., 2016). M. natalensis is largely distributed in sub-Saharan African (IUCN, 2016a), with a wide ecological range and can be found in savannas, grasslands, agricultural fields and houses (Coetzee, 1975; Monadjem et al., 2015). It hosts a number of non-pathogenic mammarenaviruses for humans such as Mopeia virus, Luna virus, Link virus, Morogoro virus, Gairo virus (Wulff et al., 1977; Taylor et al., 1981; Günther et al., 2009; Ishii et al., 2011; Gryseels et al., 2015; Cuypers et al., 2020).

The Mopeia virus, identified for the very first time in Mopeia district, Zambézia province (Fig. 1A), central Mozambique (Wulff et al., 1977), was the second African Mammarenavirus described in M. natalensis after the Lassa virus. No human cases of Mopeia virus diseases have been reported so far. However, this virus has been foreseen as an useful model to understand Lassa virus transmission ecology and immune responses (Borremans et al., 2011; Russier et al., 2012; Schaeffer et al., 2019). Grobbelaar et al. (2021), highlighted the importance of information on the occurrence and diversity of mammarenaviruses in Africa to understand possible risks to human health. A new Mammarenavirus non-pathogenic for humans (Dhati Welel virus) has been recently described in M. natalensis in eastern Africa (Gouy de Bellocq et al., 2020); which suggests that with further studies new mammarenaviruses could be identified in Africa and that a systematic mapping of Mastomys and Mammarenavirus deserves more attention. Indeed, apart from an early study from the 1970s (Wulff et al., 1977), there is a lack of data about Mammarenavirus circulation in Mozambique. Thus, this study was set out to investigate the diversity of mammarenaviruses circulating in M. natalensis trapped in villages, cropland fields and mopane woodland forest in the Limpopo National Park (LNP) and its buffer zone.

2. Materials and methods

2.1. Study sites

The study was carried out in the Massingir district counterpart of the LNP in Gaza province, south Mozambique (Fig. 1A). The rodents were trapped between March and November 2019 in three different habitats (villages, croplands fields and mopane woodland forest) from localities inside the LNP (Bingo, Macavene and Mavoze) and its buffer zone (Chibotane, Machavule and Madingane). (Fig. 1B). The Massingir district has 5893 km² of surface, 37,664 inhabitants, is dominated by dry semi-arid type zones with average annual temperatures of 30 °C and rainfall 600 mm. The study area falls within the Great Limpopo Trans-frontier Conservation Area adjacent to the South African Kruger National Park (KNP) and the Zimbabwe’s Gonarezhou National Park. The area supports subsistence farming dominated by rain-fed cropping in elevated land and with small irrigation schemes close to rivers and to Massingir dam. There are an extensive livestock production. Four localities are located inside the LNP with no physical barrier to separate wildlife and human. The wildlife mammal community includes large, medium and small sized terrestrial animals, such as elephants, buffaloes, lions, wild dogs, wild pigs and zebras.
2.2. Rodent trapping and sampling

The following experimental protocol was approved by the Ethical Committee of VetAgro-Sup in Lyon, France (Comité d’Éthique de VetAgro Sup n°18, Avis 1905) and the credential for rodent capture in the LNP was obtained from the Mozambican National Administration for Conservation Areas (Credential Nr. 1/02/2021).

Two types of sites were included during this study: capture mark and recapture (CMR) sites and removal sites. In CMR sites trapped rodents were marked and released for potential recapture for rodent ecology studies (results not shown here). Only rodents found dead in the traps in the CMR sites were samples for Mammarenavirus screening. By contrast in the removal sites, all M. natalensis rodents captured were sacrificed and the samples were screened for Mammarenavirus RNA detection. In villages, rodents were trapped inside houses and granaries. The captures in cropland fields included irrigated and rain-fed sites. The sites were selected based on the information collected from the local population about the presence of rodents and we also checked for signs of rodent’s presence before setting the traps. During the sampling, crops (mainly beans and maize) were abundant in irrigated cropland fields while in the rain-fed cropland field, the sampling occurred after peanut harvesting.

In houses and granaries, the rodents were trapped using mesh traps baited with chorizo and Sherman (small and medium sizes) baited with toasted coconuts, peanut butter and oats. In cropland fields and mopane woodland only Sherman traps were set up. The number of traps deployed in each site was adjusted according to the house, granary and field size. In each house or granary, the number of traps varied between two and five while in cropland fields and mopane woodland the minimum number was 24 and the maximum 144 traps. Due to logistic and safety reasons (i.e., presence of dangerous wildlife) a single mopane woodland was sampled four times inside the LNP. The traps were armed at the end of the day and inspected the following morning. Traps were set for one to five consecutive nights, depending on the trapping success. Captured rodents were transported to the laboratory located in the LNP headquarters where the team was based during the trapping sessions. Rodents were euthanised with isoflurane, dissected and tissues (lung, heart, liver, spleen, kidney, cerebral, tail and ear samples) as well as blood (Dried Blood Spot, DBS) were collected. All tissue samples were conserved in homemade RNAlater solution (http://www.protocol-onlin e.org/prot/Protocols/RNAlater-3999.html) and tail and ear for rodent species genetic identification were conserved in 70% ethanol (Herbreteau et al., 2011).

2.3. Rodent species genetic identification

Initially, morphological characteristics were used for rodent species in-field identification (Herbreteau et al., 2011). Because species in some African rodent genera are difficult to recognize and the taxonomy of many groups is still not well resolved (Monadjem et al., 2015), at least 10% of rodents morphologically identified as M. natalensis were selected for genetic identification at each capture site. We applied the DNA-barcoding approach to amplify and sequence (by the Sanger method) the complete mitochondrial cytochrome b gene (CYTB; 1140 bp) of selected individuals. We followed the protocols described by Bryja (2014) for DNA extraction, polymerase chain reaction (PCR) and sequencing. Species and intraspecific lineages were identified by a maximum likelihood (ML) analysis in RAxML 8.2.8 (Stamatakis, 2014), using GTRCAT substitution model recommended by the author. The robustness of the nodes was evaluated by the default bootstrap procedure with 1000 replications and new sequences were considered as part of previously defined taxa/lineages if the bootstrap support was >95%. For this study, we specifically compared the new sequences with those from the phylogenetic studies of the genus Mastomys (Colangelo et al., 2013; Martynov et al., 2020).

2.4. Mammarenavirus detection and phylogeny

Only M. natalensis individuals were considered for Mammarenavirus screening. RNA was extracted from lung tissues using Qiagen RNeasy Mini kit (Hilden, Germany). Briefly, 30 mg of lung tissue samples were disrupted for RNA release by adding 20 μL of proteinase K, one 2 mm inox bead and 500 μL of lysis buffer in a 2 μl tube and vortexed using a Qiagen TissueLyser LT system for 5 min at 50 Hz. The homogeneous lystate in the 2 μl tube from the above step was centrifuged at 11,000 × g for 3 min and then the supernatant was collected and used for viral RNA extraction following the manufacturer’s instructions.

Two pairs of primers P1-LSF/R and P2-LSF/R (Li et al., 2015) targeted to conserved regions of the L segment of mammarenaviruses were used to amplify a 611 bp fragment. The total volume of the reaction mix for the first-strand cDNA synthesis was 20 μL containing 5 μL of template RNA, 20 pmol of Mammarenavirus genus-specific primer (P1-LSF), 0.5 mM of each dNTP, 4 μL of 5× RT Buffer, 20 units of Ribolock RNase Inhibitor and 100 units of Maxima H Minus Reverse Transcriptase. The thermal profile conditions were those described in the manufacture protocol (Maxima H minus Reverse Transcriptase, Thermo Scientific). The pair P1-LSF/R was used on the first amplification and the P2-LSF/R on the second PCR. The final volume of PCR mix for the two reactions was 50 μL and the volume or concentration of mix components were: 5 μL of cDNA, 1.25 units of Dream Taq Hot Start DNA Polymerase (Thermo Scientific, city and country), 1× Dream Taq Buffer (Thermo Scientific) with 2 mM MgCl2, 0.2 mM of each dNTP (Thermo Scientific) and 3 pmol of each primer. The same thermal profile was used for each primer pair: initial denaturation at 95 °C for 3 min, followed by 40 cycles of denaturation at 95 °C for 30 s, annealing at 50 °C for 30 s and extension at 72 °C for 1 min. The final extension was at 72 °C during 5 min. The expected RT-PCR products were visualized on 1% agarose gel.

Positive RT-PCR amplicons including the positive control were purified using a Wizard SV Gel and PCR Clean-up system (Promega, Wisconsin, USA). Only purified amplicons with sufficient concentration were sent to LGC Genomics (Berlin, Germany) using the primer pair P2-LSF/R.

The final ML analysis of Mammarenavirus included 67 sequences of which 50 generated in this study, one Mopeia virus used in this study as positive control, 15 Mammarenavirus partial L gene fragment of representatives of the OWG and one representative of the NWG as outgroup from GenBank. Mammarenavirus nucleic acid sequences were aligned using MEGA 7 (Kumar et al., 2016), with minor manual adjustments. Sites that could not be unambiguously aligned were excluded and divergent regions were excluded from subsequent analysis. Thus, the final size of our alignment used for the phylogenetic analysis was of 468 bp. Phylogenies were inferred using ML method implemented in PhyML (Guindon et al., 2010). The reliability of branching orders was tested using the bootstrap approach (1000 replicates). The suited evolution model (GTR + I + G) was selected by Akaike’s Information criterion (AIC) using Topali software (Milne et al., 2009). Mopeia virus nucleic acid sequences generated in this study were deposited in GenBank under accession numbers MZ512094 to MZ512143.

2.5. Statistical analysis

The proportion of Mopeia virus RNA positive individuals were calculated by habitat and locality with 95% confidence intervals (CI) using Wilson score test (Agresti and Coull, 1998). The χ² test of independence was performed to evaluate differences in proportions of positive individuals among the habitat types as well as among the localities.

3. Results

3.1. Rodent trapping and sampling

During a total of 7290 trap nights; 1235 rodents were captured and
82.5% (n = 1019) were morphologically identified as *M. natalensis*. In CMR sites we caught 541 *M. natalensis* individuals out of which 56 were found died in the traps and in removal sites we trapped 478 individuals. A total of 534 lung tissue samples were obtained from all *M. natalensis* trapped and sacrificed in removal sites and from all individuals found died in the traps deployed in CMR sites h. *M. natalensis* were captured in all the three types of habitats and across all the localities trapped. The highest number of *M. natalensis* was trapped from cropland fields and the lowest number of individuals was caught in the mopane woodland (Table 1 and Table S1).

### 3.2. Rodent species genetic identification

Mitochondrial CYTB genotyping of selected 101 out of 534 specimens (Table S2) confirmed all of them to be *M. natalensis* and all belonging to phylogroup B-VI (Colangelo et al., 2013).

### 3.3. Mammarenavirus molecular detection

The overall proportion of positive individuals for *Mammarenavirus* RNA detection was 16.9% (95% CI: 13.9–20.3). Neither croplands nor villages were sampled in Macavene locality because the entire population that lived there was resettled outside the park. *Mammarenavirus* RNA was detected in 16.9% (95% IC: 13.8–20.5) and 16.3% (95% IC: 8.1–30.0) of specimens, respectively from cropland fields and villages. A single mopane woodland forest was sampled in the Macavene locality where 16.7% (95% CI: 3.0–56.4) of individuals tested positive. Nevertheless, the proportions of *Mopeia virus* positive rodents did not differ significantly among the habitats (χ² = 0.018; DF = 1; p = 0.893). Across the localities, Machavule (only two individuals were caught) had the highest proportion 100% (95% CI: 15.2–32.5) and the lowest proportion 2.9% (95% CI: 0.8–9.8) was found in Madingane locality (Table 1).

Concerning the differences among habitats (or localities), five samples that had doubtful RT-PCR results were excluded from the chi-square test. Besides the mopane woodland forest habitat and two localities (Macavene and Machavule) were excluded from the analysis because of low expected values (McHugh, 2013). The proportion of positive individuals differed significantly among the localities (χ² = 12.512; DF = 3; p = 0.0058).

### 3.4. Phylogenetic analysis

A total of 50 out of 90 purified amplicons had sufficient concentration for sequencing, thus the phylogenetic analysis was performed for all the positive samples from villages (n = 7) and mopane woodland forest (n = 1) and for 51% (42/82) of the positive samples from the cropland fields.

All our *Mammarenavirus* sequences clustered together with *Mopeia virus* (bootstrap value 91%) (Fig. 2). However, all the *Mopeia virus* identified in this study formed a specific *Mopeia virus* sub-clade with eight different sub-clusters. The sub-clusters I, III, IV, VI, VII and VIII were well sustained with bootstrap value >70%, whereas for sub-clusters II and IV the tree topology seemed unresolved with bootstrap values of 48% and 56%, respectively. The nodes of divergence between the sub-clusters I and II; and the sub-clusters IV, V, VI, VII were also not well supported with bootstrap value <70% (Fig. 2).

### 4. Discussion

In the present study, we screened *M. natalensis* by RT-PCR and we detected *Mopeia virus* in all habitats and localities across the study area at the interface between LNP and communal land in its buffer zone. We reported the presence of *Mopeia virus* in 16.9% (90/534) of tested *M. natalensis* with no significant differences between habitats. Phylogenetic analysis indicated that the viruses detected formed a specific *Mopeia virus* sub-clade with eight sub-clusters.

Several villages and cropland sites were surveyed inside the LNP and its buffer zone while a single mopane woodland was sampled four times inside the LNP. *M. natalensis* has a very similar external morphology with *Mastomys coucha* which is largely endemic in southern Africa (IUCN, 2016b). Sympathric occurrence of *M. natalensis* and *M. coucha* was reported in some regions of South Africa including the KNP (Kneidinger et al., 2014) which is adjacent to the LNP in our study area. The mitochondrial phylogenetic analysis split *M. natalensis* into six matrilineage phylogroups: A-I, A-II, A-III, B-I, B—V, B-VI (Colangelo et al., 2013) and all were identified as natural reservoirs of mammarenaviruses (Gouy de Bellaco et al., 2020). The CYTB mitochondrial gene sequence phylogenetic analysis indicated that all captured *Mastomys* in our study area were *M. natalensis* belonging to the B-VI lineage. Previous studies reported that two lineages of *M. natalensis* (B—V and B-VI) occur in parapaternity in Mozambique with the former being geographically restricted to the northernmost part of the country (Colangelo et al., 2013; Gouy de Bellaco et al., 2020).

*Mammarenavirus* detection in Mozambique dates back to 1977 when the *Mopeia virus* was first identified in *M. natalensis* population from Mopeia district in central Mozambique (Wulff et al., 1977). Studies have been carried out in southern Africa and indeed detected non-pathogenic *Mammarenavirus* such as *Mopeia virus*, *Morogoro virus*, *Gairo virus* and *Luna virus* in *M. natalensis* (Wulff et al., 1977; Günther et al., 2009; Ishii et al., 2011; Gryseels et al., 2015). Here, we detected the *Mopeia virus* in 16.9% (90/534) of tested *M. natalensis* samples by the viral RNA molecular detection system (RT-PCR). The proportion of positives is below the ones detected by previous serological studies: 21% in Mozambique (Wulff et al., 1977), 20% in Zimbabwe (Johnson et al., 1981) and the highest of 31.8% from retrospective samples collected South Africa and Zimbabwe (Grobbelaar et al., 2021). In Tanzania Borremans et al. (Borremans et al., 2014) which is adjacent to the LNP in our study area. The mitochondrial genetic analysis indicated that the viruses detected formed a specific *Mopeia virus* sub-clade with eight sub-clusters.

#### Table 1

<table>
<thead>
<tr>
<th>Locality</th>
<th>Habitat</th>
<th>Cropland fields</th>
<th>Villages</th>
<th>Mopane woodland</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nr. of positives (n)</td>
<td>Proportion [CI, 95%]</td>
<td>Nr. of positives (n)</td>
<td>Proportion [CI, 95%]</td>
<td>Nr. of positives (n)</td>
</tr>
<tr>
<td>Bingo</td>
<td>17 (78)</td>
<td>21.8 [14.1–32.2]</td>
<td>3 (10)</td>
<td>30.0 [10.8–60.3]</td>
<td>/</td>
</tr>
<tr>
<td>Chibotane</td>
<td>28 (144)</td>
<td>19.4 [13.8–26.7]</td>
<td>0 (19)</td>
<td>0.0 [0.0–16.8]</td>
<td>/</td>
</tr>
<tr>
<td>Macavene</td>
<td>/</td>
<td>/</td>
<td>2 (2)</td>
<td>100.0</td>
<td>/</td>
</tr>
<tr>
<td>Machavule</td>
<td>/</td>
<td>/</td>
<td>[34.2–100.0]</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>Madingane</td>
<td>1 (62)</td>
<td>1.6 [0.3–8.6]</td>
<td>1 (8)</td>
<td>12.5 [2.2–47.1]</td>
<td>/</td>
</tr>
<tr>
<td>Mavoze</td>
<td>26 (201)</td>
<td>17.9 [12.2–23.8]</td>
<td>1 (4)</td>
<td>25.0 [4.6–69.9]</td>
<td>/</td>
</tr>
</tbody>
</table>

Five individuals had dubious RT-PCR results in Bingo (n = 1), Chibotane (n = 2) and Mavoze (n = 2).
Mopeia virus

Fig. 2. Maximum Likelihood analysis, implemented in PhyML, of a partial nucleotide sequence of Mammarenavirus L gene fragment from the samples, our positive control genotyped, OWG Mammarenavirus reference sequences previously identified in Africa and available in the GenBank (Table S4). One Mammarenavirus sequence of the NWG (Table S4) were used as outgroup. The asterisks at nodes represent the bootstrap values ≥70% calculated from 1000 bootstrap replicates. Scale bars indicate the number of base substitutions per site. The sample reference number is followed by three letters indicating the collection site as following: bgc = Bingo cropland; bgv = Bingo village; cbc = Chibotane cropland; cnc = Macavene mopan woodland; mvc = Mavoze cropland; mvc = Madoine village; mlv = Machavule village; mgc = Madingane cropland and mgv = Madingane village. Village. The colors differentiate the types of habitats: black = cropland fields, blue = villages and red = mopane woodland. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

We sampled three different habitats (only two individuals were caught in the mopane woodland forest) and no significant differences were observed on the proportion of individuals positive for Mopeia virus RNA across the habitats. This observation suggests that the level of Mopeia virus circulation is the same among M. natalensis populations living in these three habitats. The possible explanation for this homogeneity could be that the studied habitats offer similar environmental conditions for viral viability and transmission. Borremans et al. (2011) also reported a great homogeneity in Mopeia virus prevalence across different habitat types in Tanzania and suggested that the abiotic environment may not be an important determinant of virus prevalence.

The proportion of positive individuals differed significantly among the localities; however, the 100% [95% CI: 99.9–100.0] found in Machavule is less informative because of the very small number of rodents (N = 2) captured in this locality. Thus, the 22.7% [95% CI: 15.2–32.5] detected in Bingo is referenced as the highest proportion of positive rodents across the localities in our study area. On the other hand, our statistical inference within the localities was influenced by the lowest proportion of Mopeia virus positive rodents detected in Madingane 2.9% [95% CI: 0.8–9.8]. The differences observed within localities suggests that in some localities humans are relatively more exposed to potentially zoonotic Mammarenaviruses. Nonetheless, the localities were sampled in different periods and this could possibly explain the differences of proportion of positive samples detected in Madingane locality. Rodent-host population dynamics is an important factor of the epidemiology of mammarenaviruses, whereby seasonal peaks of virus circulation may be associated with increased rodent populations and could explain prevalence differences between different months (Jay et al., 2005; Akhmetzhanov et al., 2019). The level of Mammarenavirus circulation in rodent population depend on the interaction between the virus, rodent-hosts and ecological variables (Childs and Peters, 1993). Therefore, a study incorporating an analysis of epidemiological and ecological parameters in our study area is needed to understand the homogeneity of virus circulation in habitats, the heterogeneity of prevalence in the localities and possibly during the different sampling months.

The mammarenaviruses detected in three different habitats (villages, croplands and mopane woodland forest) clustered with Mopeia virus, the unique Mammarenavirus species described so far in Mozambique (Wulff et al., 1977). The mammarenaviruses from the study area formed a specific sub-clade that did not include the reference Mopeia virus isolate from Mozambique (GenBank accession Nr. AY772169.1) as well as the Mopeia virus positive control isolate ("Mopeia virus, strain UVE/MOPV/UNK/MZ/Mozambique 20410 | EVAg"). The mammarenaviruses detected in the study area formed eight sub-clusters and the sub-cluster I is the one most widespread with viruses from the three habitats and from five out of six localities included in the study. However, the sub-clusters II and V were not well sustained. Likewise, the bootstrap values at the roots of sub-clusters I and II as well as for sub-clusters IV to VII were low suggesting an unresolved tree topology. Consequently, it is probable that...
the viruses could change the place between the sub-clusters I and II as well as among the sub-clusters IV to VII. This is likely due to the small size (468 bp) of the fragment analyzed. Therefore, we think that the analysis of the whole L segment would give better insight of the genetic diversity of the mammareviruses in our study area. Besides, the basal position of sample 765 from Bingo cropland field may suggest that in fact there is some degree of genetic variability of Mopeia virus in the study area. The detection of these probable different sub-clusters in the different habitats may indicate wide circulation of viruses of different sub-clusters within the study area. Evidence of Mammarenavirus circulation in other species than M. natalensis had been reported. Mopeia virus antibodies were detected in Aethomys chrysophilus in Zimbabwe (Johnson et al., 1981). Mus minutoides carries Lank virus in Zambia and Kodoko virus in West Africa (Lecompte et al., 2007; Ishii et al., 2012). Recently, two novel mammarenaviruses were detected in Angola: the first one, Bitu virus, related to Okahandja virus, was found in Micaelamus namaquensis; the second one, Kwanza virus, related to Mobala virus, was detected in Mus triton (Těšíková et al., 2021). Results obtained in samples from South Africa and Zimbabwe indicated presence of mammarenaviruses in 14 rodent species that included Aethomys, chrysophilus, Mastomys, coucha, Rattus norvegicus, Rattus rattus, Saccostomus campestris and Gerbilliscus leucogaster (Grobelnlar et al., 2021). The assessment of the potential circulation of mammarenaviruses in other rodent species from our study area is therefore important.

Mammarenaviruses transmission from rodent to humans occur through contaminated rodent excreta and secretions and crop harvesting or hunting activities. Poor hygiene in households and rodent consumption increase the risk of transmission (Paweska, 2014; Akhmetzhanov et al., 2019) The human population in our study area practices agriculture, hunting (although prohibited by law), some consume rodents and like in other parts of the country the environment sanitation is deficient. Here we reported that Mopeia virus circulates in M. natalensis rodents, thus we hypothesize that humans in the study area may be exposed to mammarenaviruses. Unpublished data cited by Grobelnlar et al. (2021), indicated that antibodies to Mopeia virus were detected in humans’ sera but with no association with clinical disease resembling mammarenaviruses. Serological and molecular investigation of Mopeia virus in humans will inform about viral exposure and the viral genetic profile in the study area.

5. Conclusions

Mopeia virus was detected in M. natalensis trapped in villages, crop-land fields and mopane woodland from all the six surveyed localities. The Mopeia virus we detected formed a specific Mopeia virus sub-clade than the previously described in Mozambique. Habitat type does not have influence on the proportion of positive individuals in the study area. The genus Mammarenavirus contain zoonotic species and possibly others potentially zoonotic could emerge, thus, our results suggest that humans could be exposed to mammarenaviruses in all habitats, so awareness campaigns could be designed to sensitize local citizens about the potential risks of zoonotic disease spill over from rodents as well as further risk-based studies to explore which group of stakeholders are more exposed than others. We have expanded and updated the data on Mopeia virus in Mozambique and made more sequences available for this virus.

Supplementary data to this article can be found online at https://doi.org/10.1016/j.meegid.2022.105204.

Data availability

The authors declare that all the data analyzed here will be available without any restrictions.