

1 Antibodies against medically relevant arthropod-borne viruses  
2 in the ubiquitous African rodent *Mastomys natalensis*

3 Short title: Antibodies against arboviruses in *Mastomys natalensis* from Tanzania

4

5 Wim De Kesel <sup>1,2\*</sup>, Bram Vanden Broecke <sup>1,3</sup>, Benny Borremans <sup>1,4</sup>, Léa Fourchault <sup>5</sup>, Elisabeth Willems  
6 <sup>2</sup>, Ann Ceulemans <sup>2,6</sup>, Christopher Sabuni <sup>7</sup>, Apia Massawe <sup>7</sup>, Rhodes H. Makundi <sup>7</sup>, Herwig Leirs <sup>1</sup>,  
7 Martine Peeters <sup>8</sup>, Erik Verheyen <sup>1,5</sup>, Sophie Gryseels <sup>1,5</sup>, Joachim Mariën <sup>1,6</sup> and Kevin K. Ariën <sup>2,9\*</sup>

8

9 <sup>1</sup> Evolutionary Ecology Group, Department of Biology, Faculty of Science, University of Antwerp,  
10 Antwerp, Belgium

11 <sup>2</sup> Virology Unit, Department of Biomedical Sciences, Institute of Tropical Medicine, Antwerp, Belgium

12 <sup>3</sup> Terrestrial Ecology Unit, Department of Biology, Ghent University, Ghent, Belgium

13 <sup>4</sup> Wildlife Health Ecology Research Organization, San Diego, United States of America

14 <sup>5</sup> OD Taxonomy & Phylogeny, Royal Belgian Institute of Natural Sciences, Brussels, Belgium

15 <sup>6</sup> Virus Ecology Unit, Department of Biomedical Sciences, Institute of Tropical Medicine, Antwerp,  
16 Belgium

17 <sup>7</sup> Institute of Pest Management, Sokoine University of Agriculture, Morogoro, Tanzania

18 <sup>8</sup> TransVIHMI, University of Montpellier, Institut de Recherche pour le Développement (IRD), INSERM,  
19 Montpellier, France

20 <sup>9</sup> Department of Biomedical sciences, Faculty of Pharmaceutical, Biomedical and Veterinary Sciences,  
21 University of Antwerp, Antwerp, Belgium

22 \* Corresponding authors

23 E-mail: [wim.dekesel@uantwerpen.be](mailto:wim.dekesel@uantwerpen.be) (WDK); [karien@itg.be](mailto:karien@itg.be) (KKA)

## 24 Abstract

25 Over the past decades, the number of arthropod-borne virus (arbovirus) outbreaks  
26 has increased worldwide. Knowledge regarding the sylvatic cycle (i.e., non-human  
27 hosts/environment) of arboviruses is limited, particularly in Africa, and the main hosts for  
28 virus maintenance are unknown. Previous studies have shown the presence of antibodies  
29 against certain arboviruses (i.e., chikungunya-, dengue- and zika virus) in African non-human  
30 primates and bats. We hypothesize that small mammals, specifically rodents, may function as  
31 amplifying hosts in anthropogenic environments. The detection of RNA of most arboviruses  
32 is complicated by the virus's short viremic period within their hosts. An alternative to  
33 determine arbovirus hosts is by detecting antibodies, which can persist several months. We  
34 developed a high-throughput multiplex immunoassay to detect antibodies against 15  
35 medically relevant arboviruses. We used this assay to assess almost 1,300 blood samples of  
36 the multimammate mouse, *Mastomys natalensis* from Tanzania. In 24% of the samples, we  
37 detected antibodies against at least one of the tested arboviruses, with high seroprevalences  
38 of antibodies reacting against dengue virus serotype one (7.6%) and two (8.4%) and  
39 chikungunya virus (6%). Seroprevalence was higher in females and increased with age, which  
40 could be explained by inherent immunity and behavioral differences between sexes and the  
41 increased chance of exposure to an arbovirus with age. We evaluated whether antibodies  
42 against multiple arboviruses co-occur more often than randomly and found that this may be  
43 true for some members of the *Flaviviridae* and *Togaviridae*. In conclusion, the development  
44 of an assay against a wide diversity of medically relevant arboviruses enabled the analysis of  
45 a large sample collection of one of the most abundant African small mammals. Our findings  
46 suggest a role in the transmission of multiple arboviruses by this ubiquitous rodent and

47 provide a solid foundation for future molecular screening to elucidate the role in the arbovirus  
48 transmission cycle.

## 49 Author summary

50 One of the main causes of zoonotic related human morbidity and mortality is the  
51 transmission of arthropod-borne viruses such as dengue virus, Yellow Fever virus, and  
52 chikungunya virus. These viruses cannot only infect humans but also livestock, pets, and  
53 wildlife, though our understanding of their non-human hosts remains limited. Rodents are  
54 thought to be an interesting host for these viruses because they can be abundant, often live  
55 near humans and some are already known to be viral hosts. However, research has focused  
56 on non-human primates, neglecting other potential hosts. To address this gap, we have  
57 developed a high-throughput antibody test to screen rodent blood against 15 different  
58 arboviruses. Our findings reveal that a proportion of *Mastomys natalensis*, a common African  
59 rodent species, carry antibodies that (cross-)react against these viruses. We hypothesize that  
60 immunologically naïve juveniles may drive transmission, particularly during population  
61 outbreaks. These outbreaks coincide with environmental conditions that are favorable for  
62 mosquitoes, the vectors of these viruses. Thus, increasing the risk of spillover to humans,  
63 livestock, and wildlife. Understanding the role of rodents in arbovirus transmission dynamics  
64 is crucial for mitigating zoonotic disease risks.

65

66

## 67 Introduction

68           The African continent harbors a diverse array of infectious diseases with profound  
69 impacts on public health, economic development, and general well-being [1,2]. Diseases  
70 caused by arthropod-borne viruses, collectively known as arboviruses, are a growing threat  
71 for Africa and the rest of the world especially in relation to climate and environmental  
72 changes [3,4]. Arboviruses are a polyphyletic clade that includes several viral families, of  
73 which the most important are *Flaviviridae*, *Togaviridae*, *Bunyaviridae*, and *Reoviridae* [5].  
74 Some well-known arboviruses, notorious for their devastating effects on human health, are  
75 dengue virus, Yellow Fever virus, Zika virus, and chikungunya virus. Mosquitoes, ticks,  
76 sandflies, and midges are the primary vectors responsible for arbovirus transmission as they  
77 engage in hematophagy. These vectors do not only affect humans and livestock, but also a  
78 wide range of wildlife hosts [6–8]. Indeed, while for some arboviruses morbidity and mortality  
79 can be high in humans, similar impacts have been detected in other animals by arboviruses  
80 such as Rift Valley Fever virus in goats and sheep, West Nile virus in birds and horses and  
81 Japanese Encephalitis virus in birds and pigs [4,5,9–11]. The (re-)emergence of arboviruses  
82 appears, at least partially, to be caused by the increased urbanization and global connectivity,  
83 natural genetic evolution of viruses, and adaptations of the vectors to changing climate and  
84 environments [11,12]. Emerging arboviruses pose a threat for humans, livestock, as well as  
85 wildlife, therefore it needs to be approached from a One health perspective (i.e., including  
86 human, animal, and environment health) [13,14]. Nevertheless, our knowledge about the  
87 extent to which wild animals can serve as sylvatic hosts for human-infecting arboviruses and  
88 the natural diversity of arboviruses remains insufficient. This significantly limits our

89 understanding of arbovirus transmission dynamics, which is required to develop effective  
90 control measurements.

91 For decades, efforts have been made to identify natural reservoirs of arboviruses to  
92 monitor, prevent, and control sources of infection that pose a threat to human health [15–  
93 17]. Several studies have proposed non-human primates as significant potential reservoirs for  
94 arboviruses, as they have found arbovirus antibodies and viral RNA in this animal group [18–  
95 20]. However, other animal groups such as small mammals have often been neglected as  
96 arbovirus hosts [21]. Sporadic reports of arboviruses in small mammal species suggest that a  
97 more comprehensive investigation of their potential role as a host is needed [22–24].

98 Rodents have a number of characteristics that could make them potentially important  
99 hosts for several pathogens, including arboviruses [23]. Particularly the high species diversity,  
100 the fact that many species can reach high population abundances and turnover rates. The risk  
101 of pathogen spillover to humans increases with the role of some rodents as a pest species,  
102 due to their proximity to humans [25,26]. A notable example of such a pest species is the  
103 ubiquitous rodent *Mastomys natalensis*, commonly known as the multimammate mouse. This  
104 species inhabits many regions of sub-Saharan Africa, with a preference for crop fields, fallow  
105 land, and typically occurring within or at the fringes of urban settlements [27,28]. In east  
106 Africa, and especially in Tanzania, the reproductive cycle of *M. natalensis* is strongly  
107 correlated with seasonal rainfall which leads to strong seasonal fluctuations in density (20 -  
108 500 individuals/hectare), and occasionally even severe population outbreaks (>1000  
109 individuals/hectare) [29–32]. This has large ecological and societal impacts due to crop  
110 damage and influences seasonal transmission dynamics of different pathogens [33–35]. The  
111 multimammate mouse is a known host for several zoonotic pathogens such as *Lassa*  
112 *mammarenavirus*, *Yersinia pestis*, *Leptospira interrogans*, *Leishmania major* as well as

113 different ecto- and endoparasites [31,36–52]. No studies have investigated or reported on  
114 arboviruses in *M. natalensis*, except Diagne et al. (2019) who have detected Usutu virus RNA  
115 in *M. natalensis*. However, other studies have reported on sporadic arbovirus detections in  
116 other rodent species in sub-Saharan Africa [22,24,53,54]. These findings, along with the  
117 ecology of *M. natalensis* (i.e., high abundance during population outbreaks, proximity to  
118 humans, and its status as a proven pathogen host) may suggest that this species plays a role  
119 in the natural transmission cycle of arboviruses. Consequently, *M. natalensis* could thus pose  
120 a risk to humans in east Africa, particularly in Tanzania as an amplifying host.

121         The human population in Tanzania has experienced several outbreaks of chikungunya  
122 virus, Rift Valley fever virus, West Nile virus, and dengue virus in the past decades [55–59].  
123 Due to the symptomatic similarities between arbovirus and malaria infections, which has a  
124 prevalence of around 20% in Tanzania, it is probable that arbovirus cases are underreported  
125 [60,61]. While these studies confirm that the local human population is indeed exposed to  
126 arboviruses, the specific dynamics of arbovirus transmission in this region remains unclear.

127         The goal of this study was to investigate the potential of wild *M. natalensis* to serve as  
128 a host for arboviruses in their natural environment. To achieve this, we first developed a  
129 multiplex immune assay to detect immunoglobulin G (IgG) antibodies against 15 different  
130 arboviruses causing disease in humans and subsequently conducted a comprehensive  
131 screening of almost 1,300 blood samples obtained from *M. natalensis* from Morogoro,  
132 Tanzania.

## 133 Materials and Methods

### 134 Sample origin

135 The samples used in this study were collected during previous published and  
136 unpublished studies conducted by the University of Antwerp (UAntwerp) and Pest  
137 Management Center of the Sokoine University of Agriculture on *M. natalensis* in Morogoro,  
138 Tanzania, between 2010 and 2019 [31,62,63] (Fig 1). In this study, the samples were divided  
139 in two screening sessions. The first session consisted of approximately 500 dried blood spot  
140 (DBS) samples, from wild captured mice that were used in infection and behavioral  
141 experiments in six different years (i.e., 2010, 2011, 2015, 2017, 2018, and 2019) with an  
142 average of 80 samples per year. The second session consisted of 800 DBS samples from mice  
143 involved in capture mark recapture experiments in 2017 and 2019. All samples were randomly  
144 selected from the studies regardless of individual characteristics or trapping period.

145 During these studies, *M. natalensis* were live caught using Sherman traps (H.B.  
146 Sherman Traps, Tallahassee, USA) in a heterogeneous landscape (e.g., woodlands, maize  
147 fields, and fallow land) on the premises of the Sokoine University of Agriculture in Morogoro,  
148 Tanzania. Blood was collected from the retro-orbital plexus using a 50 $\mu$ L hematocrit capillary  
149 tube and preserved on filter paper (Serobuvar; LDA 22; Zoopole, France). The filter paper  
150 was dried for 12 hours at room temperature and archived at -20°C in envelopes with  
151 desiccant. Additional data related to characteristics such as sex, reproductive status, weight,  
152 and body measurements were recorded. More detailed Information pertaining to the  
153 trapping procedures and sampling methodology can be found in the primary research  
154 documents associated with these studies [31,62–64].

155



156

157 **Fig 1. African continent with a focus on Tanzania.** Samples were collected in the city of Morogoro (red triangle)  
158 which is located in the Morogoro region.

## 159 Analysis set up and protocol.

160 To assess the presence of arbovirus antibodies in DBS against a panel of arboviruses,  
161 we first developed a multiplex immune assay using Luminex technology [18,65]. Recombinant  
162 virus-derived proteins (Table 1) were covalently coupled to carboxyl-functionalized  
163 fluorescent magnetic beads ( $1\text{-}3\mu\text{g}/1.25 \times 10^6$  beads) (Luminex Corp. MagPlex®  
164 Microspheres; Bio-Rad; Temse, Belgium) employing the BioPlex amine coupling kit (Ref.:  
165 171406001; Bio-Rad; Temse, Belgium) following the manufacturer's instructions.

166 To obtain positive control samples we inoculated captive *M. natalensis* individuals  
167 (age: 5-12 months) from our breeding colony at UAntwerp with recombinant virus-derived  
168 proteins (Table 1) [66–68]. We subcutaneously injected  $4\mu\text{g}$  of the respective virus protein  
169 and  $1\mu\text{L}$  of vaccine adjuvant (Quil-A® adjuvant; InvivoGen; Toulouse, France), dissolved in  
170 autoclaved phosphate buffered saline (PBS) to achieve a final volume of 1mL. This inoculum  
171 was evenly divided, with 0.5mL administered into the scruff and 0.5mL into the hindlimb of  
172 the animal, using a 25-gauge, 12.5mm needle and a 0.5mL syringe. This inoculation was  
173 duplicated for each viral protein (i.e., performed in two mice) and repeated twice for each



174 mouse (i.e., inoculation on day 0 and day 20). We collected blood, according to the same  
 175 method as in the previous studies, every 10 days from day zero until day 30, at day 30 we also  
 176 collected whole blood from which serum was extracted. Serum from day 30 from individuals  
 177 were the antibody response increased over time were considered as positive samples. Day 30  
 178 had the highest antibody titer in our tests and is also a time point at which IgG antibody  
 179 development is anticipated to have reached a peak [69,70].

180 **Table 1. Recombinant arbovirus proteins for the inoculation of captive *Mastomys natalensis*.**

Viral family	Virus	Protein (reference)	Supplier
<i>Bunyaviridae</i>	Rift Valley Fever virus (RVFV)	Nucleoprotein (REC31640)	The native antigen company (Kidlington, United Kingdom)
<i>Flaviviridae</i>	Yellow Fever virus (YFV)	Nonstructural protein 1 (YFV-NS1)	The native antigen company (Kidlington, United Kingdom)
<i>Flaviviridae</i>	Zika virus (ZIKV)	Nonstructural protein 1 (40544-V07H)	Interchim (Montluçon Cedex, France)
<i>Flaviviridae</i>	Dengue virus serotype 1 (DENV1)	Nonstructural protein 1 (DEN-004)	Prospecbio (Rehovot, Israel)
<i>Flaviviridae</i>	Dengue virus serotype 2 (DENV2)	Nonstructural protein 1 (PIP048A)	BioRad (Temse, Belgium)
<i>Flaviviridae</i>	Dengue virus serotype 3 (DENV3)	Nonstructural protein 1 (DENV3-NS1)	The native antigen company (Kidlington, United Kingdom)
<i>Flaviviridae</i>	Dengue virus serotype 4 (DENV4)	Nonstructural protein 1 (DENV4-NS1)	The native antigen company (Kidlington, United Kingdom)
<i>Flaviviridae</i>	Usutu virus (USUV)	Nonstructural protein 1 (Ab218552)	The native antigen company (Kidlington, United Kingdom)
<i>Flaviviridae</i>	West Nile virus (WNV)	Nonstructural protein 1 (40346-V07H)	Sinobiological (Eschborn, Germany)
<i>Flaviviridae</i>	Tick-borne Encephalitis virus (TBEV)	Nonstructural protein 1 (TBEV-NS1)	The native antigen company (Kidlington, United Kingdom)
<i>Flaviviridae</i>	Wesselsbron virus (WSLV)	Nonstructural protein 1 (REC31698)	The native antigen company (Kidlington, United Kingdom)
<i>Nairoviridae</i>	Crimean Congo Hemorrhagic Fever virus (CCHFV)	Nucleoprotein (REC31639)	The native antigen company (Kidlington, United Kingdom)
<i>Togaviridae</i>	Chikungunya virus (CHIKV)	Envelope protein 2 (CHI-003)	Prospecbio (Rehovot, Israel)
<i>Togaviridae</i>	Mayaro virus (MAYV)	Envelope protein 2 (REC31644)	The native antigen company (Kidlington, United Kingdom)
<i>Togaviridae</i>	O'nyong nyong virus (ONNV)	Envelope protein 2 (B4TG40)	Interchim (Montluçon Cedex, France)

181

182            Screening was done in 96 flat-bottom well plates, each plate contained DBS samples  
183 of 80 wild *M. natalensis*, two background controls, eight negative controls and a six step  
184 dilution series (1:200 – 1:625,000) of a positive pool sample. Each well in the plate contained  
185 50µL of the corresponding sample type. The samples of the wild *M. natalensis* were acquired  
186 by placing a punched-out DBS (round, 0.5 cm diameter) in 200 µL of dilution buffer (1% bovine  
187 serum albumin, 0.2% Tween-20, 5% fetal calf serum, 45% distilled water, 50% Hypertonic PBS  
188 {0.08% NaH<sub>2</sub>PO<sub>4</sub>, 0.25% Na<sub>2</sub>HPO<sub>4</sub>, 8.8% NaCl}). One single DBS punch corresponds to  
189 approximately 10µL of blood [71]. The punched DBS were left to elute overnight, in a 1.5mL  
190 Eppendorf tube, maintained at a temperature of 4°C on a plate shaker. This elution was  
191 considered a 1:100 dilution and was diluted, with dilution buffer, to 1:200 prior to loading in  
192 the 96 well plate. This dilution gave the best signal to noise ratio in our preliminary tests and  
193 are in line with previous studies [18,72]. The background control was reading buffer (1%  
194 bovine serum albumin, 0.05% NaN<sub>3</sub>, 100% phosphate buffered saline). The eight negative  
195 controls were four DBS, treated the same as the wild *M. natalensis* DBS, and four serum  
196 samples in a 1:200 dilution. All negative controls originated from the breeding colony at  
197 UAntwerp. The positive pool sample was compiled from serum from the positive individuals  
198 acquired through the arbovirus protein inoculation experiment. Serum from 15 positive  
199 individuals (i.e., one for each arbovirus antigen) was pooled to create the positive pool  
200 sample, each individual serum had a final dilution in the pool of 1:200.

201            In each well of the 96 well plate, 25µL of bead mixture was added. The bead mixture  
202 consisted out of ~1000 protein-coated beads per arbovirus antigen suspended in reading  
203 buffer. The bead mixture of the first screening session did not contain ONNV beads.

204            Plates, containing 50µL of sample and 25µL of bead mixture per well, were incubated  
205 for one hour at room temperature, in the dark and on a plate shaker (Heidolph Titrimax 100;

206 VWR; Leuven, Belgium) at 400rpm/min. After incubation, plates underwent washing with  
207 dilution buffer using an automated plate washer (Tecan Hydroflex plate washer; Tecan  
208 Benelux; Mechelen, Belgium). Subsequently, we added 50 $\mu$ L Biotin anti-mouse IgG (4 $\mu$ g/mL)  
209 (Sigma-Aldrich B7022; Merck Life Science; Hoeilaart, Belgium) to each well and incubated for  
210 40 minutes. After another round of washing, we added 50 $\mu$ L of Streptavidin-R-phycoerythrin  
211 (1 $\mu$ g/mL) (10655783; Fisher Scientific; Brussel, Belgium) to each well, followed by a 10-minute  
212 incubation. The last wash step used reading buffer, and the final bead pellet was resuspended  
213 in 150 $\mu$ L of reading buffer. Beads were read on a Bio-Plex 200 System (Bio-Rad; Temse,  
214 Belgium). Results were quantified as the median fluorescent intensity (MFI) based on a  
215 minimum of 100 beads per antigen.

## 216 Data analysis and statistics

217 All data preparation, analysis and statistical procedures were conducted using R  
218 Statistical Software (R version 4.3.3) [73]. We used the body weight of the wild-caught *M.*  
219 *natalensis* individuals at the time of sample collection as a rough proxy for age, which we  
220 subdivided into three categories based on the 1/3 quantiles of weight; juvenile (5 - 26.7g),  
221 subadult (>26.7 - 42g) and adult (>42 - 91g). These weight classes coincide to the expected  
222 sexual maturity, with sexual maturity estimated to occur between 30 - 40g [29,32].

223 To control for variation between different assay plates and testing days, the MFI  
224 results were transformed to relative antibody units using the positive dilution series as a  
225 standard curve. The MFI result of the positive control starting dilution (i.e., 1:200) was  
226 equalized to 3,125 units and each following dilution step was adjusted proportionally (i.e., the  
227 final dilution step 1:625,000 corresponded to 1 unit). The results of the two sessions were  
228 combined by linear alignment adjustment. This alignment was based on 86 duplicate samples

229 encompassing the measurable range, allowing the adjustment of the results from the first  
230 session.

231 Finally, each sample was categorized as a binary value (i.e., 1= positive, 0= negative)  
232 for each of the tested arboviruses. This was done based on whether the unit value exceeded  
233 the mean cutoff value for that specific arbovirus antigen. Five cutoff values were determined  
234 for each arbovirus antigen: I) the mean plus three times the standard deviation of the negative  
235 controls (i.e. '*NegCtrl*') [18,65]; the change-point analysis, using R package '*changepoint*'  
236 (version: 2.2.4), calculated at most one changepoint based on the II) mean (i.e. '*CHP.m*'), III)  
237 variance (i.e. '*CHP.v*') and IV) a combination of mean and variance (i.e. '*CHP.mv*') of wild-  
238 caught samples [74,75] and V) the maximum value of an average antibody curve (i.e. '*Recap*').  
239 This curve was based on wild-caught individuals that were recaptured at least three times and  
240 showed seroconversion. Seroconversion of an individual was considered when the  
241 individual's maximum unit value was at least four-fold the minimum unit value. This four-fold  
242 increase is a standard seroconversion confirmation measure in human antibody studies [76].  
243 An average antibody curve, with days as the explanatory variable, was created for each  
244 antigen by aligning the maximum unit value of each recaptured seroconverted individual to  
245 the same day. The binary results were used to calculate the seroprevalence for each arbovirus  
246 along with a 95% confidence interval (CI), using the '*binom.exact*' from the package '*binom*'  
247 (version: 1.1.1.1) [77]. The seroprevalence according to the different cutoff methods was  
248 compared to the seroprevalence of the antibody curve cutoff using the '*chisq.test*' from the  
249 package '*stats*' (version 4.3.3) [73]

250 As an indication of cross-reactivity in antibody response between the tested  
251 arboviruses, pair-wise Pearson correlations were calculated on the binary results, according  
252 to the antibody curve cutoff, of all samples using the '*corr.test*' function of R package '*psych*'

253 (version: 2.4.1) [78]. The cross-reactivity in antibody response was visualized using the  
254 *'heatmap.2'* function of the R package *'gplots'* (version: 3.1.3) [79].

255 A generalized linear model (logit link function and binomial error distribution) was  
256 constructed with the package *'stat'* (version: 4.3.1), with the response variable being the  
257 binary serostatus of each sample [73]. Age (juvenile, subadult and adult), sex and their  
258 interaction were included as explanatory variables. The analysis of variance was performed  
259 using a likelihood ratio test, with p-values calculated assuming a chi-squared distribution.  
260 Pairwise comparison of the seroprevalence was performed between the six combinations of  
261 the explanatory variables (two levels of sex and three levels of age), using the *'emmeans'*  
262 package (version: 1.8.9) [80]. To prevent reporting statistical findings based on the reliance of  
263 an arbitrary p-value of 0.05, we instead present significance in terms of levels of statistical  
264 support based on p-values. P-values exceeding 0.1 are labeled as “no” support and values  
265 around 0.05 (range 0.1 -  $\geq$  0.01, symbol: \*) as “weak” support. “Moderate” support was  
266 assigned to p-values clearly below 0.05 (range  $<$  0.01 -  $\geq$  0.001, symbol: \*\*), while “strong”  
267 support is reserved for p-values lower than 0.05 ( $<$  0.001, symbol: \*\*\*). This representation  
268 in terms of statistical support is based on current statistical reporting practices [81].

## 269 Results

270 In total 1,280 DBS samples were tested of which 660 were female, consisting of 256  
271 juveniles, 172 subadults and 232 adults, 620 samples were male with 199 juveniles, 313  
272 subadults and 108 adults. Samples of recaptured individuals were considered as individual  
273 samples for all analysis.

## 274 Seroprevalence

275 The seroprevalences according to the different cutoff methods showed at least a weak  
276 statistical support for a different seroprevalence compared to the antibody curve  
277 seroprevalence for almost all arboviruses. Histograms of the data and seroprevalence for each  
278 tested arbovirus antigen according to the different cutoff methods is shown in S1 and S2 Figs.  
279 The cutoff value according to the antibody curve based on the recaptured seroconverted  
280 individuals was used as the main cutoff value for all further calculations.

281 The overall arbovirus seroprevalence, defined as at least positive for one of the tested  
282 arboviruses, except ONNV, was almost 24% (95% CI: 21.89 – 26.66%; N= 1280). ONNV was  
283 excluded since the samples of the first session were not screened for antibodies against the  
284 ONNV antigen. The seroprevalence for *Flaviviridae* was 20% (95% CI: 17.99 – 22.46%; N=  
285 1280) and for *Togaviridae*, excluding ONNV, almost 7% (95% CI: 5.48 – 8.32%; N= 1280).  
286 Overall, seroprevalences ranged from 0.62% for DENV3 (95% CI: 0.27 – 1.23%; N= 1280) and  
287 MAYV (95% CI: 0.27 – 1.23%; N= 1280) to 8.44% for DENV2 (95% CI: 6.97 – 10.10%; N= 1280),  
288 see Table 2.

289 **Table 2. Total seroprevalence of each arbovirus and virus family in the wild-caught *M. natalensis* sample set.**

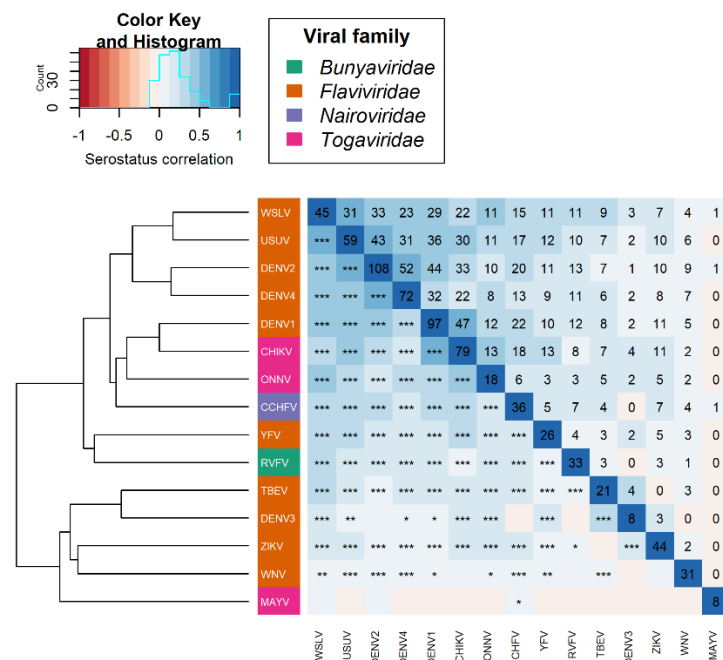
	Seroprevalence (%)	95% CI (%)	Nr. positive	Cutoff
Arbovirus <sup>a</sup>	24.22	21.89 – 26.66	310	
<i>Bunyaviridae</i>				
RVFV	2.58	1.78 – 3.60	66	37.94
<i>Flaviviridae</i>	20.16	17.99 – 22.46	258	
YFV	2.03	1.33 – 2.96	26	3.45
ZIKV	3.44	2.51 – 4.59	44	15.23
DENV1	7.58	6.19 – 9.17	97	42.11
DENV2	8.44	6.97 – 10.10	108	101.09
DENV3	0.62	0.27 – 1.23	8	32.27
DENV4	5.62	4.43 – 7.03	72	40.66
USUV	4.61	3.53 – 5.91	59	6.47
WNV	2.42	1.65 – 3.42	31	25.60
TBEV	1.64	1.02 – 2.50	21	50.85
WSLV	3.52	2.58 – 4.68	45	4.16

<i>Nairoviridae</i>				
CCHFV	2.81	1.98 – 3.87	36	5513.61
<i>Togaviridae</i> <sup>a</sup>	6.80	5.48 – 8.32	87	
CHIKV	6.17	4.92 – 7.63	79	43.95
MAYV	0.62	0.27 – 1.23	8	11.91
ONNV	2.18	1.30 – 3.42	18	77.10

290 A 95% confidence interval (CI) is provided and the calculated cutoff value is in units. Sample size was 1280 for  
 291 each tested arbovirus except for ONNV which had 826 samples. <sup>a</sup> Indicates that ONNV was not included for that  
 292 calculation.

## 293 Pairwise arbovirus serostatus correlation

294 The correlations in serostatus of samples between the tested arboviruses are  
 295 visualized in Fig 2. Correlation between two arboviruses is depicted in color scale with the  
 296 statistical symbol, lower triangle, and the number of positive samples in the upper triangle.  
 297 The matrix is accompanied by a dendrogram based on the hierarchical clustering of the  
 298 correlation coefficients. The branch lengths are a proxy for relative distance between  
 299 arboviruses based on the serostatus response of the samples.



300  
 301 **Fig 2. Correlation of the serostatus response between the tested arboviruses with a dendrogram of**  
 302 **hierarchical clustering.** Symbols in lower triangle represent significance of correlation, values in upper triangle,  
 303 including diagonal line, represents the number of positive individuals for the corresponding arboviruses.

304 The correlation in sample response between the tested arboviruses ranged from -  
305 2.44% for TBEV and ZIKV with no statistical support ( $p= 0.384$ ) to almost 59% between WSLV  
306 and USUV with a strong statistical support ( $p< 0.001$ ). The dendrogram based on the  
307 hierarchical clustering of the correlation showed that WSLV – USUV (correlation= 58.52%;  $p<$   
308 0.001), DENV2 – DENV4 (correlation= 56.03%;  $p< 0.001$ ) and DENV1 – CHIKV (correlation=  
309 50.31%;  $p< 0.001$ ) are relatively closer to each other than to other tested arboviruses.

## 310 Model analysis of antibody response

### 311 Sex, age, and interaction effects on serostatus

312 The generalized linear model indicated that there was a weak statistical interaction  
313 between the effects of sex and age on serostatus for DENV4 (Deviance [Df.= 2; Res.Df.= 1274]= 5.44;  
314  $p= 0.066$ ) and USUV (Deviance [Df.= 2; Res.Df.= 1274]= 7.72;  $p= 0.021$ ). For the other tested arboviruses,  
315 no support for a statistical interaction was detected, the interaction was thus removed from  
316 those models. In the case of RVFV, no statistical support was found for an effect of sex, age,  
317 or the interaction on the serostatus. All results from generalized linear model's analysis of  
318 variance are reported in S1 Table.

319 The analysis of the sex variable showed a moderate support for males having a lower  
320 seroprevalence compared to females for DENV2 (Est.  $_{\text{males}} \pm \text{SE}= -0.665 \pm 0.242$ ;  $p= 0.006$ ) and WSLV  
321 (Est.  $_{\text{males}} \pm \text{SE}= -1.446 \pm 0.483$ ;  $p= 0.003$ ). A weak statistical effect of a lower seroprevalence in males  
322 compared to females was detected in ZIKV (Est.  $_{\text{males}} \pm \text{SE}= -0.643 \pm 0.386$ ;  $p= 0.096$ ), TBEV (Est.  $_{\text{males}} \pm$   
323  $\text{SE}= -1.103 \pm 0.635$ ;  $p= 0.083$ ), CCHFV (Est.  $_{\text{males}} \pm \text{SE}= -0.821 \pm 0.460$ ;  $p= 0.074$ ) and CHIKV (Est.  $_{\text{males}} \pm \text{SE}= -$   
324  $0.633 \pm 0.296$ ;  $p= 0.032$ ). There was no support for a difference in seroprevalence between males  
325 and females for RVFV (Est.  $_{\text{males}} \pm \text{SE}= 0.097 \pm 0.381$ ;  $p= 0.798$ ), YFV (Est.  $_{\text{males}} \pm \text{SE}= 0.425 \pm 0.413$ ;  $p= 0.304$ ),  
326 DENV1 (Est.  $_{\text{males}} \pm \text{SE}= -0.377 \pm 0.248$ ;  $p= 0.129$ ), DENV3 (Est.  $_{\text{males}} \pm \text{SE}= -17.43 \pm 2021.76$ ;  $p= 0.993$ ), WNV



327 (Est.  $\text{males} \pm \text{SE} = -0.009 \pm 0.397$ ;  $p = 0.982$ ), MAYV (Est.  $\text{males} \pm \text{SE} = -1.222 \pm 1.080$ ;  $p = 0.258$ ) and ONNV (Est.  $\text{males}$   
 328  $\pm \text{SE} = -17.75 \pm 1663.70$ ;  $p = 0.991$ ).

329 The analysis of the age variable showed a strong statistical support for a higher  
 330 seroprevalence in subadults than in juveniles for DENV1 and DENV2, a moderate support for  
 331 CHIKV and a weak support for YFV, ZIKV, TBEV, WSLV and CCHFV. There was no support for a  
 332 difference in subadult and juvenile seroprevalence in the other tested arboviruses. A  
 333 significantly higher seroprevalence in adults compared to juveniles was shown for ZIKV,  
 334 DENV1, DENV2, WSLV and CHIKV with a strong support. A moderate support for a higher  
 335 seroprevalence in adults than in juveniles was detected for YFV and CCHFV. Adults showed a  
 336 weak statistical support for a higher seroprevalence in contrast to juveniles for WNV and  
 337 TBEV. All other tested arboviruses showed no support for a statistical difference between  
 338 adults and juveniles. The comparison between subadults and adults showed a strongly  
 339 supported statistical difference for DENV1 and CHIKV with a higher seroprevalence in adults.  
 340 A moderate support for a higher seroprevalence in adults compared to subadults was  
 341 detected for ZIKV and CCHFV. Yellow Fever virus, WNV, WSLV and ONNV showed a weak  
 342 support for a statistically higher seroprevalence in adults than in subadults. The other  
 343 arboviruses showed no statistically significant difference between adults and subadults. See  
 344 Table 3 for estimates, standard errors, and p-values.

345 **Table 3. Difference in coefficient estimate on logit scale between the age levels with standard error (SE).**

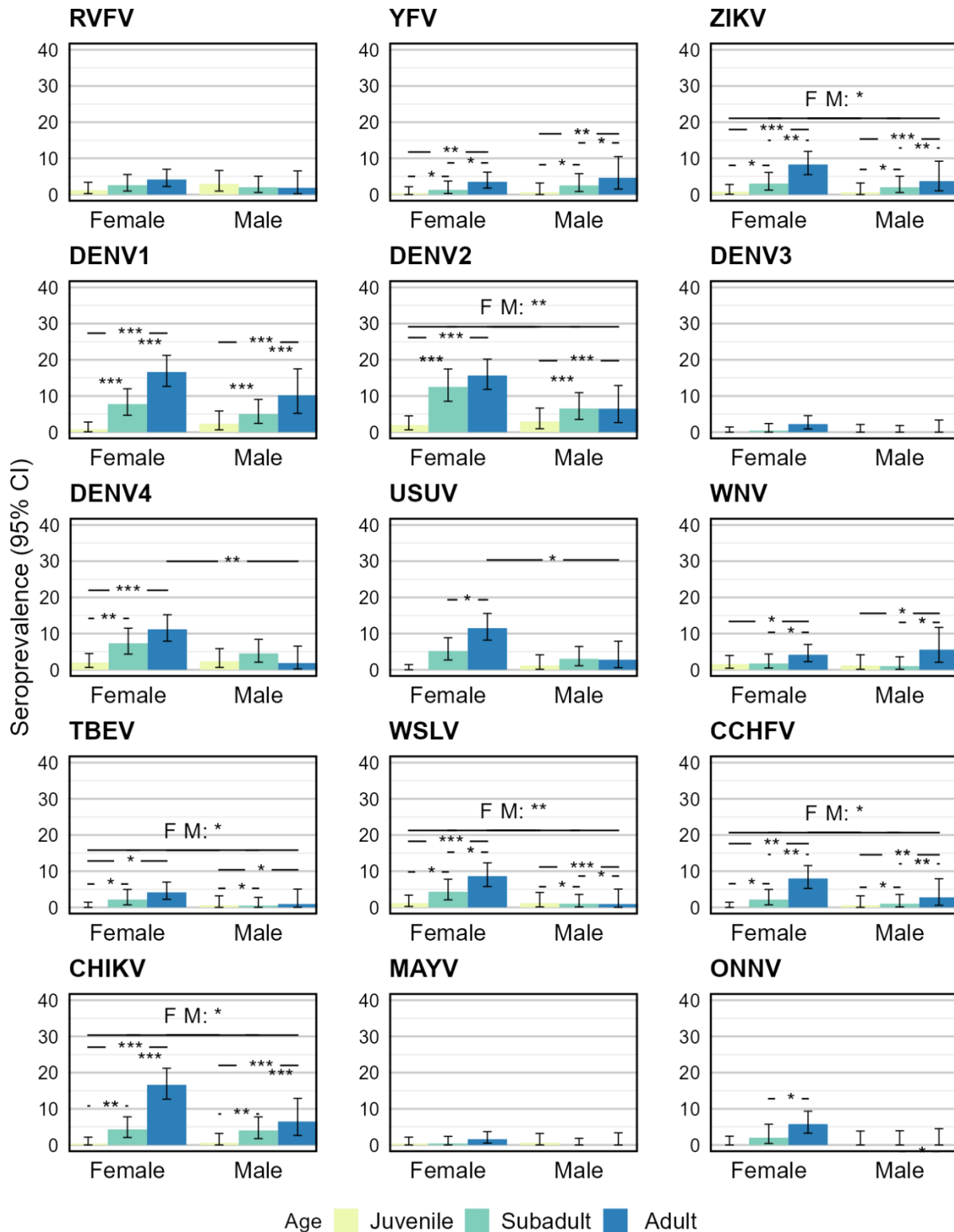
	Juvenile - Subadult		Juvenile - Adult		Subadult - Adult	
	Estimate $\pm$ SE	p-value	Estimate $\pm$ SE	p-value	Estimate $\pm$ SE	p-value
RVFV	-0.227 $\pm$ 0.480	0.637	-0.649 $\pm$ 0.446	0.146	-0.422 $\pm$ 0.421	0.316
YFV	-1.368 $\pm$ 0.794	<b>0.085</b>	-2.197 $\pm$ 0.757	<b>0.004</b>	-0.829 $\pm$ 0.449	<b>0.065</b>
ZIKV	-1.349 $\pm$ 0.656	<b>0.040</b>	-2.308 $\pm$ 0.611	<b>&lt; 0.001</b>	-0.959 $\pm$ 0.365	<b>0.009</b>
DENV1	-1.610 $\pm$ 0.456	<b>&lt; 0.001</b>	-2.468 $\pm$ 0.434	<b>&lt; 0.001</b>	-0.858 $\pm$ 0.243	<b>&lt; 0.001</b>
DENV2	-1.551 $\pm$ 0.356	<b>&lt; 0.001</b>	-1.780 $\pm$ 0.352	<b>&lt; 0.001</b>	-0.229 $\pm$ 0.221	0.301
DENV3	-16.47 $\pm$ 2176.65	0.994	-18.14 $\pm$ 2176.65	0.993	-1.665 $\pm$ 1.073	0.121
WNV	0.007 $\pm$ 0.582	0.991	-1.20 $\pm$ 0.477	<b>0.012</b>	-1.207 $\pm$ 0.480	<b>0.012</b>
TBEV	-1.855 $\pm$ 1.083	<b>0.087</b>	-2.566 $\pm$ 1.039	<b>0.014</b>	-0.711 $\pm$ 0.499	0.155

WSLV	-0.960 ± 0.539	<b>0.075</b>	-1.656 ± 0.492	<b>&lt; 0.001</b>	-0.697 ± 0.357	<b>0.051</b>
CCHFV	-1.999 ± 1.072	<b>0.062</b>	-3.321 ± 1.021	<b>0.001</b>	-1.321 ± 0.434	<b>0.002</b>
CHIKV	-2.267 ± 0.749	<b>0.002</b>	-3.474 ± 0.723	<b>&lt; 0.001</b>	-1.207 ± 0.283	<b>&lt; 0.001</b>
MAYV	0.642 ± 1.228	0.601	-0.807 ± 0.844	0.339	-1.449 ± 1.104	0.189
ONNV	-17.05 ± 1747.82	0.992	-18.15 ± 1747.82	0.992	-1.099 ± 0.641	<b>0.087</b>

346 Data originates from the pairwise comparison of the age class variables of the generalized linear model. P-values  
347 marked in bold have at least a weak statistical support ( $p < 0.1$ ).

348 In the case of DENV4, there was a weak support for an interaction, the analysis  
349 showed that there was a strong statistical support for a higher seroprevalence in female  
350 adults compared to female juveniles (Est.  $_{\text{female} - \text{adult}} \pm \text{SE} = 1.844 \pm 0.486$ ;  $p < 0.001$ ). A moderate  
351 support was shown for a higher seroprevalence in female subadults compared to female  
352 juveniles (Est.  $_{\text{female} - \text{subadult}} \pm \text{SE} = 1.379 \pm 0.517$ ;  $p = 0.008$ ) and a weak support for a higher  
353 seroprevalence in female adults compared to male adults (Est.  $_{\text{female} - \text{adult}} \pm \text{SE} = 1.898 \pm 0.736$ ;  $p =$   
354  $0.010$ ). For USUV the model analysis with a weak interaction, showed that there was weak  
355 statistical support for a higher seroprevalence in female adults compared to female subadults  
356 (Est.  $_{\text{female} - \text{adult}} \pm \text{SE} = 0.868 \pm 0.345$ ;  $p = 0.012$ ) and also a weak support for a higher seroprevalence in  
357 female adults compared to male adults (Est.  $_{\text{female} - \text{adult}} \pm \text{SE} = 1.515 \pm 0.612$ ;  $p = 0.013$ ).

358 Fig 3 displays the seroprevalence for the six distinct levels (two levels of sex and three  
359 levels of age) for all tested arboviruses, with statistical support lines based on the log odds.



360

361 **Fig 3. Seroprevalence according to sex and age combinations with 95% confidence error bars for each**  
 362 **arbovirus.** Statistical support on seroprevalence difference is indicated by asterisks in the horizontal lines.  
 363 Sample size: 660 females of which 256 juveniles, 172 subadults and 232 adult and 620 males with 199 juveniles,  
 364 313 subadults and 108 adults. Sample sizes for ONNV are 560 females: 150 juveniles, 150 subadults, 260 adults  
 365 and 266 males: 94 juveniles, 92 subadults and 80 adults.

## 366 Discussion

367 In this study we optimized a high-throughput multiplex immunoassay for the  
368 simultaneous detection of IgG antibodies against 15 medically relevant arboviruses and used  
369 it to investigate the potential of *M. natalensis* to serve as a host for arboviruses. We describe  
370 the screening results of a historic set of wild *M. natalensis* DBS samples. Our results revealed  
371 an overall seroprevalence of 24% against the entire panel of tested antigens. Virus family-  
372 specific seroprevalences were approximately 2.6%, 20%, 2.8% and 7% for respectively  
373 *Bunyaviridae*, *Flaviviridae*, *Nairoviridae* and *Togaviridae*. We further found that female  
374 rodents were more likely to be classified as antibody positive for eight of the 15 tested  
375 arboviruses. Additionally, positivity increased significantly with age for almost all tested  
376 arboviruses.

377 The lack of realistic natural positive controls limits the possibility to determine a true  
378 cutoff, we therefore used recaptured seroconverted individuals to determine a cutoff value.  
379 The use of antibody titers at multiple time points are a standard practice to determine  
380 antibody or pathogen development and (sero)conversions in human studies [76]. However,  
381 multiple samples of an individual animal across time are often impossible or very difficult in  
382 wildlife studies. Our study is unique in that regard that we have measurements of individual  
383 recaptured *M. natalensis*. We consider that our cutoff based on seroconverted individuals is  
384 a good proxy for the natural cutoff value, since it is based on similar methods as in human  
385 studies [76]. We tried to show in our analysis that the tested mathematical methods could  
386 approximate this calculated cutoff and thus provide a method for future studies that do not  
387 have access to recaptured seroconverted wildlife samples. Unfortunately, the tested cutoff  
388 methods did not significantly approximate the seroprevalence according to the cutoff using

389 samples from recaptures. The negative control-based cutoff (i.e., the mean plus three times  
390 the standard deviation of the negative control samples) gave unrealistic high seroprevalences.  
391 This can be explained by the fact that the negative control samples originate from a breeding  
392 colony. These animals have thus never been exposed to a natural environment and the  
393 pathogens that occur in the environment. The statistical methods vary in their seroprevalence  
394 with some methods approximating the estimated seroprevalence according to the recaptured  
395 cutoff. This high degree of variation makes it difficult to decide on one method that works for  
396 all the tested arboviruses. The cutoff value for CCHFV seems extremely high compared to the  
397 other arboviruses, but the unit values for CCHFV are also much higher than for the other  
398 arboviruses (see S1 Fig). The reason is that the unit values are calculated based on the positive  
399 dilution series and the positive controls for CCHFV were not of the same magnitude as for the  
400 other arboviruses. The value in determining a cutoff and the resulting seroprevalence is that  
401 it allows the comparison of results with previous and future studies on arbovirus  
402 seroprevalence in rodents or other wildlife. We are aware that the used cutoff and resulting  
403 seroprevalences could be an over- or underestimation and might not reflect the natural  
404 arbovirus seroprevalence. We therefore encourage future research to investigate and  
405 compare different cutoff methods for arbovirus (or pathogen) antibody detection in wildlife  
406 studies.

407         The detection of antibodies against each of 15 tested arbovirus antigens indicates that  
408 these arboviruses, or closely related viruses, are present in *M. natalensis*. The overall  
409 arbovirus seroprevalence of 24% suggests that this rodent species is commonly infected with  
410 one or more arboviruses and that it could thus play a significant role in virus transmission and  
411 persistence. Our results corroborate previous studies, which detected USUV and WNV RNA in  
412 respectively *M. natalensis* and *M. erythroleucus*, in Senegal [24,54]. Besides in this genus,

413 arboviral RNA has also been found in other rodents in Africa, such as *Rattus rattus* for USUV  
414 and WSLV and *Desmodillus auricularis* for WSLV [22,24,53]. The findings in our study thus  
415 further corroborate that arboviruses are likely present in rodents, and specifically in the  
416 ubiquitous *M. natalensis*. The demographic and ecological characteristics of *M. natalensis*  
417 may have particularly important implications for arbovirus transmission. The population  
418 densities of *M. natalensis* in Tanzania are strongly dependent on weather conditions. More  
419 specifically, early rainfall and elevated temperatures lead to an exponential growth in the  
420 population density, due to an increase influx of juveniles [29,30,32]. The rainfall and increased  
421 temperatures are also beneficial for the breeding of mosquitoes and the multiplication of  
422 arboviruses within these vectors [82]. Further, *M. natalensis* is highly abundant around  
423 houses and in the crop fields at the fringes of the villages. These factors increase the likelihood  
424 of arbovirus outbreaks in *M. natalensis* populations, with the possibility of spillover to  
425 humans.

426 Arboviruses that show the highest seroprevalence are DENV1, DENV2, DENV4 and  
427 CHIKV, with seroprevalences between five to nine percent. These seroprevalences could be  
428 caused by cross-reactivity due to antibodies of other dengue virus serotypes or other  
429 flaviviruses binding to the non-structural protein 1 (NS1 protein) of DENV1, DENV2 and  
430 DENV4. The same effect could also be true for alphaviruses binding to the envelope protein 2  
431 (E2 protein) of CHIKV. Whether these seroprevalences are indeed due to the presence of the  
432 arbovirus specific antibodies or a related arbovirus remains to be investigated. Nonetheless,  
433 it indicates that a part of the sampled *M. natalensis* population in Morogoro is exposed to  
434 dengue virus and CHIKV or respectively to a related flavivirus and alphavirus. This hypothesis  
435 is supported by the fact that flavi- and alphaviruses are the most prevalent arboviral genera

436 in humans, compared to other arbovirus genera, and potentially thus also in rodents involved  
437 in the sylvatic cycle [83,84].

438 A recent health survey has shown that, in the same region as where our rodent  
439 population was sampled, a high percentage of the human population is seropositive for CHIKV  
440 (9.83%) [55]. Another study in the same region reported acute infection of CHIKV in 1.28% of  
441 patients with fever and malaria-like symptoms [85]. Although these studies have not found  
442 any indication of dengue virus in humans, a large-scale cross-sectional study in Tanzania has  
443 found CHIKV and dengue virus antibodies in respectively 28.0% and 16.1% of the population  
444 [56]. These studies clearly indicate that the human population in Tanzania is exposed to  
445 arboviruses and then specifically to CHIKV and DENV.

446 The cross-reactivity analysis via the correlation matrix and hierarchical clustering (Fig  
447 2) showed an antibody response correlation between WSLV – USUV (59%), DENV2 – DENV4  
448 (56%) and DENV1 – CHIKV (50%). We expected that phylogenetically related arboviruses  
449 would show elevated levels of correlation due to cross-reactivity [86]. A remarkable result in  
450 this cross-reactivity analysis is that DENV1 – CHIKV cluster together with a correlation of 50%,  
451 based on the serostatus of the tested samples. The branch DENV1/CHIKV clusters also closer  
452 to ONNV than to the branches of WSLV/USUV and DENV2/DENV4. This is unexpected since  
453 CHIKV belongs to the *Togaviridae* and DENV1 to the *Flaviviridae* [87]. The proteins used for  
454 the antibody detection are also two different proteins, with the E2 protein used for the  
455 *Togaviridae* and NS1 protein for the *Flaviviridae*, thus limiting the possibility of cross-  
456 reactivity. Although we cannot exclude that there might be similar epitopes between the  
457 different proteins, but other studies have already indicated that cross-reactivity between the  
458 E2 protein of the *Togaviridae* and NS1 protein of the *Flaviviridae* is limited [88,89]. Given that  
459 both *Togaviridae* and *Flaviviridae* viruses are circulating in humans in East Africa, we

460 hypothesize that these viral families may also both be present in rodents [5,10]. More  
461 specifically, it is plausible that both viral families could be found in *M. natalensis*, where  
462 pathogen co-infections are common [49]. This hypothesis is further supported by the fact that  
463 some viruses in both families are transmitted by the same arthropod vectors, such as *Aedes*  
464 *aegypti* and *Aedes albopictus* for both dengue virus and CHIKV [4,82].

465 For some of the tested arboviruses we found statistical support for a higher  
466 seroprevalence in females than in males. This result is supported by previous studies where  
467 it is shown that female mice have a stronger innate immune response than male mice [90].  
468 Other animals (e.g., birds, fish, insects) as well as humans also display stronger immune  
469 responses in females [91–95]. The major driving forces behind these immune differences are  
470 genetic (i.e., X-chromosomes) and hormonal (i.e., different estrogen and testosterone levels)  
471 [96]. In the case of *M. natalensis*, behavioral differences could also be the cause for this  
472 divergence in seroprevalence. Previous studies have already shown that home range,  
473 behavior and pathogen presence differ between male and female *M. natalensis* [48,97].  
474 Besides the sex effects, we also found statistical support for a positive age effect on the  
475 presence of antibodies in some of the tested arboviruses. This increased seroprevalence with  
476 age corroborates previous findings for other pathogens (i.e., *Bartonella sp.*, *Anaplasma sp.*,  
477 helminths, and arenaviruses) [48,49,98]. This age effect further supports our hypothesis that  
478 *M. natalensis* is exposed to arboviruses and that individuals develop antibodies and gain  
479 immunity via repeated exposures throughout their life. To maintain the arbovirus  
480 transmission among the *M. natalensis* population, there needs to be a proportion of the  
481 population that is either chronically infected or immunologically naïve. Chronic infections in  
482 *M. natalensis* have already been documented for mammarenaviruses [31,99]. However, as  
483 far as we are aware naturally occurring chronic arbovirus infections have not been reported



484 in humans or non-human vertebrates. Therefore, the presumable driving factor in sustained  
485 transmission is the presence of immunologically naïve individuals. During the breeding  
486 season, which coincides with increased rainfall and temperature, there is an influx of  
487 immunologically naïve juveniles. This influx can reach high proportions during population  
488 outbreak periods [30,64]. We thus expect that it is mainly juveniles who are the major factor  
489 in sustaining the arbovirus transmission cycle. We predict that the prevalence of arboviral  
490 genetic material will be higher in juveniles than in adults, since juveniles do not possess the  
491 necessary antibodies to fight of the infection.

492 We conclude from our detected antibody responses that arboviruses, or related  
493 viruses, are present in *M. natalensis* in Morogoro, Tanzania. The higher seroprevalence we  
494 detect in females can be explained by genetic, hormonal, ecological and/or behavioral  
495 differences between sexes. Individuals are exposed to these viruses throughout their life and  
496 gain immunity as they age. We hypothesize that juvenile *M. natalensis* play an essential role  
497 in sustaining arbovirus transmission as they are immunologically naïve and can reach high  
498 densities in favorable climate conditions that coincide with optimal vector conditions. More  
499 extensive screening, such as virus neutralization tests and molecular screening of these  
500 viruses within *M. natalensis* are necessary to quantify the contribution of this rodent species  
501 in the arbovirus transmission cycle.

## 502 Acknowledgements

503 The Ethical Committee for Animal Testing at UAntwerp approved all experiments  
504 (ECD2021-79 and ECD2023-08). This study was funded by The Research Foundation – Flanders  
505 (FWO) through the Senior research project G054820N (to KKA, EV and MP) and PhD fellowship  
506 1171023N (to WDK).

## 507 References

- 508 [1] Boutayeb A. The Impact of Infectious Diseases on the Development of Africa.  
509 Handbook of Disease Burdens and Quality of Life Measures, 2010.  
510 [https://doi.org/10.1007/978-0-387-78665-0\\_66](https://doi.org/10.1007/978-0-387-78665-0_66).
- 511 [2] Nkengasong JN, Tessema SK. Africa Needs a New Public Health Order to Tackle  
512 Infectious Disease Threats. *Cell* 2020;183. <https://doi.org/10.1016/j.cell.2020.09.041>.
- 513 [3] Minakshi P, Brar B, Lambe UP, Ranjan K, Prasad G, Harimohan, et al. RNA viruses:  
514 Greatest global threat and one health solutions. *Virusdisease* 2019;30.
- 515 [4] Huang Y-JS, Higgs S, Vanlandingham DL. Emergence and re-emergence of mosquito-  
516 borne arboviruses. *Curr Opin Virol* 2019;34:104–9.
- 517 [5] Venter M. Assessing the zoonotic potential of arboviruses of African origin. *Curr Opin*  
518 *Virol* 2018;28:74–84.
- 519 [6] Omondi D, Masiga DK, Ajamma YU, Fielding BC, Njoroge L, Villinger J. Unraveling host-  
520 vector-arbovirus interactions by two-gene high resolution melting mosquito  
521 bloodmeal analysis in a Kenyan wildlife-livestock interface. *PLoS One*  
522 2015;10:e0134375.
- 523 [7] Musa AA, Muturi MW, Musyoki AM, Ouso DO, Oundo JW, Makhulu EE, et al.  
524 Arboviruses and blood meal sources in zoophilic mosquitoes at human-wildlife  
525 interfaces in Kenya. *Vector-Borne and Zoonotic Diseases* 2020;20:444–53.
- 526 [8] Thompson P, van Den Bergh C, Venter E, Schade M, Swanepoel R. Co-circulation of Rift  
527 Valley fever virus and other zoonotic arboviruses at the human-livestock-wildlife  
528 interface in KwaZulu-Natal, South Africa. *International Journal of Infectious Diseases*  
529 2020;101:535. <https://doi.org/https://doi.org/10.1016/j.ijid.2020.09.1387>.

- 530 [9] Pierson TC, Diamond MS. The continued threat of emerging flaviviruses. *Nat Microbiol*  
531 2020. <https://doi.org/10.1038/s41564-020-0714-0>.
- 532 [10] Mayer S V., Tesh RB, Vasilakis N. The emergence of arthropod-borne viral diseases: A  
533 global prospective on dengue, chikungunya and zika fevers. *Acta Trop* 2017.  
534 <https://doi.org/10.1016/j.actatropica.2016.11.020>.
- 535 [11] Gould E, Pettersson J, Higgs S, Charrel R, De Lamballerie X. Emerging arboviruses: why  
536 today? *One Health* 2017;4:1–13.
- 537 [12] Gould EA, Higgs S. Impact of climate change and other factors on emerging arbovirus  
538 diseases. *Trans R Soc Trop Med Hyg* 2009;103.  
539 <https://doi.org/10.1016/j.trstmh.2008.07.025>.
- 540 [13] Shahhosseini N, Wong G, Babuadze G, Camp JV, Ergonul O, Kobinger GP, et al. Crimean-  
541 Congo hemorrhagic fever virus in Asia, Africa and Europe. *Microorganisms*  
542 2021;9:1907.
- 543 [14] Akash S, Islam MdR, Rahman MdM. Rift Valley fever (RVF): a re-emerging zoonotic  
544 disease, pathogenesis, epidemiology, current status, and future perspective –  
545 correspondence. *International Journal of Surgery* 2023;109.
- 546 [15] Bernstein AS, Ando AW, Loch-Temzelides T, Vale MM, Li B V, Li H, et al. The costs and  
547 benefits of primary prevention of zoonotic pandemics. *Sci Adv* 2022;8:eabl4183.
- 548 [16] Vora NM, Hannah L, Lieberman S, Vale MM, Plowright RK, Bernstein AS. Want to  
549 prevent pandemics? Stop spillovers. *Nature* 2022;605:419–22.
- 550 [17] Gubler DJ. The global emergence/resurgence of arboviral diseases as public health  
551 problems. *Arch Med Res*, 2002. [https://doi.org/10.1016/S0188-4409\(02\)00378-8](https://doi.org/10.1016/S0188-4409(02)00378-8).
- 552 [18] Raulino R, Thaurignac G, Butel C, Villabona-Arenas CJ, Foe T, Loul S, et al. Multiplex  
553 detection of antibodies to Chikungunya, O'nyong-nyong, Zika, Dengue, West Nile and

- 554 Usutu viruses in diverse non-human primate species from Cameroon and the  
555 Democratic Republic of Congo. *PLoS Negl Trop Dis* 2021;15:e0009028-.
- 556 [19] Eastwood G, Sang RC, Guerbois M, Taracha ELN, Weaver SC. Enzootic circulation of  
557 chikungunya virus in East Africa: Serological evidence in non-human Kenyan primates.  
558 *Am J Trop Med Hyg* 2017;97:1399–404.
- 559 [20] Valentine MJ, Murdock CC, Kelly PJ. Sylvatic cycles of arboviruses in non-human  
560 primates. *Parasit Vectors* 2019;12:463.
- 561 [21] Olive M-M, Goodman SM, Reynes J-M. The role of wild mammals in the maintenance  
562 of Rift Valley fever virus. *J Wildl Dis* 2012;48:241–66.
- 563 [22] Diagne MM, Faye M, Faye O, Sow A, Balique F, Sembène M, et al. Emergence of  
564 Wesselsbron virus among black rat and humans in Eastern Senegal in 2013. *One Health*  
565 2017. <https://doi.org/10.1016/j.onehlt.2017.02.001>.
- 566 [23] Gora D, Yaya T, Jocelyn T, Didier F, Maoulouth D, Amadou S, et al. The potential role of  
567 rodents in the enzootic cycle of Rift Valley fever virus in Senegal. *Microbes Infect* 2000.  
568 [https://doi.org/10.1016/S1286-4579\(00\)00334-8](https://doi.org/10.1016/S1286-4579(00)00334-8).
- 569 [24] Diagne MM, Ndione MHD, Di Paola N, Fall G, Bedekelabou AP, Sembène PM, et al.  
570 Usutu virus isolated from rodents in Senegal. *Viruses* 2019;11:181.
- 571 [25] Dahmana H, Granjon L, Diagne C, Davoust B, Fenollar F, Mediannikov O. Rodents as  
572 hosts of pathogens and related zoonotic disease risk. *Pathogens* 2020;9:202.
- 573 [26] Fiedler LA. Rodent problems in Africa. *Rodent pest management*, CRC Press; 2018, p.  
574 35–65.
- 575 [27] Hánová A, Konečný A, Mikula O, Bryjová A, Šumbera R, Bryja J. Diversity, distribution,  
576 and evolutionary history of the most studied African rodents, multimammate mice of

- 577 the genus *Mastomys*: An overview after a quarter of century of using DNA sequencing.  
578 *Journal of Zoological Systematics and Evolutionary Research* 2021;59:2500–18.
- 579 [28] Chidodo S, Kilawe CJ, Mnyone LL, Broecke B Vanden, Mulungu LS. Factors affecting the  
580 composition of rodent assemblages in the North Uluguru Mountains, Tanzania. *J*  
581 *Vertebr Biol* 2020;69. <https://doi.org/10.25225/jvb.20047>.
- 582 [29] Leirs H, Verhagen R, Verheyen W. The basis of reproductive seasonality in *Mastomys*  
583 rats (Rodentia: Muridae) in Tanzania. *J Trop Ecol* 1994;10:55–66.
- 584 [30] Sluydts V, Crespín L, Davis S, Lima M, Leirs H. Survival and maturation rates of the  
585 African rodent, *Mastomys natalensis*: Density-dependence and rainfall. *Integr Zool*  
586 2007;2:220–32.
- 587 [31] Mariën J, Borremans B, Verhaeren C, Kirkpatrick L, Gryseels S, de Bellocq J, et al.  
588 Density dependence and persistence of Morogoro arenavirus transmission in a  
589 fluctuating population of its reservoir host. *Journal of Animal Ecology* 2020;89:506–18.  
590 <https://doi.org/10.1111/1365-2656.13107>.
- 591 [32] Leirs H, Stuyck J, Verhagen R, Verheyen W. Seasonal variation in growth of *Mastomys*  
592 *natalensis* (Rodentia: Muridae) in Morogoro, Tanzania. *Afr J Ecol* 1990;28:298–306.
- 593 [33] Mwanjabe PS, Sirima FB, Lusingu J. Crop losses due to outbreaks of *Mastomys*  
594 *natalensis* (Smith, 1834) Muridae, Rodentia, in the Lindi region of Tanzania. *Int*  
595 *Biodeterior Biodegradation* 2002;49. [https://doi.org/10.1016/S0964-8305\(01\)00113-](https://doi.org/10.1016/S0964-8305(01)00113-5)  
596 5.
- 597 [34] Stenseth NC, Leirs H, Skonhøft A, Davis SA, Pech RP, Andreassen HP, et al. Mice, rats,  
598 and people: The bio-economics of agricultural rodent pests. *Front Ecol Environ* 2003;1.  
599 [https://doi.org/10.1890/1540-9295\(2003\)001\[0367:MRAPTB\]2.0.CO;2](https://doi.org/10.1890/1540-9295(2003)001[0367:MRAPTB]2.0.CO;2).

- 600 [35] Mulungu LS. Control of rodent pests in maize cultivation: the case of Africa, 2017.  
601 <https://doi.org/10.19103/as.2016.0002.18>.
- 602 [36] Ziwa MH, Matee MI, Hang'Ombe BM, Lyamuya EF, Kilonzo BS. Plague in Tanzania: An  
603 overview. *Tanzan J Health Res* 2013;15. <https://doi.org/10.4314/thrb.v15i4.7>.
- 604 [37] Holt J, Davis S, Leirs H. A model of Leptospirosis infection in an African rodent to  
605 determine risk to humans: Seasonal fluctuations and the impact of rodent control. *Acta  
606 Trop* 2006. <https://doi.org/10.1016/j.actatropica.2006.08.003>.
- 607 [38] Sadlova J, Vojtkova B, Hrnčirova K, Lestínova T, Spitzova T, Becvar T, et al. Host  
608 competence of African rodents *Arvicanthis neumanni*, *A. niloticus* and *Mastomys  
609 natalensis* for *Leishmania major*. *Int J Parasitol Parasites Wildl* 2019;8.  
610 <https://doi.org/10.1016/j.ijppaw.2019.01.004>.
- 611 [39] Laudisoit A, Leirs H, Makundi R, Krasnov BR. Seasonal and habitat dependence of fleas  
612 parasitic on small mammals in Tanzania. *Integr Zool* 2009;4.  
613 <https://doi.org/10.1111/j.1749-4877.2009.00150.x>.
- 614 [40] Schwan TG, Lopez JE, Safronetz D, Anderson JM, Fischer RJ, Maïga O, et al. Fleas and  
615 trypanosomes of peridomestic small mammals in sub-Saharan Mali. *Parasit Vectors*  
616 2016;9. <https://doi.org/10.1186/s13071-016-1818-5>.
- 617 [41] Brouat C, Duplantier JM. Host habitat patchiness and the distance decay of similarity  
618 among gastro-intestinal nematode communities in two species of *Mastomys*  
619 (southeastern Senegal). *Oecologia* 2007;152. <https://doi.org/10.1007/s00442-007-0680-8>.
- 621 [42] Brouat C, Kane M, Diouf M, Bâ K, Sall-Dramé R, Duplantier JM. Host ecology and  
622 variation in helminth community structure in *Mastomys* rodents from Senegal.  
623 *Parasitology* 2007;134. <https://doi.org/10.1017/S003118200600151X>.

- 624 [43] Ribas A, López S, Makundi RH, Leirs H, De Bellocq JG. *Trichuris* spp. (Nematoda:  
625 Trichuridae) from two rodents, *Mastomys natalensis* and *Gerbilliscus vicinus* in  
626 Tanzania. *Journal of Parasitology* 2013;99. <https://doi.org/10.1645/12-151.1>.
- 627 [44] Ribas A, Diagne C, Tatard C, Diallo M, Poonlaphdecha S, Brouat C. Whipworm diversity  
628 in West African rodents: a molecular approach and the description of *Trichuris*  
629 *duplantieri* n. sp. (Nematoda: Trichuridae). *Parasitol Res* 2017;116.  
630 <https://doi.org/10.1007/s00436-017-5404-3>.
- 631 [45] Ribas A, Makundi RH, Bellocq JG De. *Paraconcinnum leirsi* n.sp. (Trematoda:  
632 Dicrocoeliidae) from rodents in Tanzania and its phylogenetic position within the  
633 dicrocoeliids. *Afr Zool* 2012;47. <https://doi.org/10.3377/004.047.0219>.
- 634 [46] Diouf M, Diagne CA, Quilichini Y, Dobigny G, Garba M, Marchand B. *Pterygodermatites*  
635 (mesopectines) *niameyensis* n. Sp. (nematoda: Rictulariidae), a parasite of *mastomys*  
636 *natalensis* (smith, 1834) (rodentia: Muridae) from Niger. *Journal of Parasitology*  
637 2013;99. <https://doi.org/10.1645/13-204.1>.
- 638 [47] Diagne CA, Charbonnel N, Henttonen H, Sironen T, Brouat C. Serological Survey of  
639 Zoonotic Viruses in Invasive and Native Commensal Rodents in Senegal, West Africa.  
640 *Vector-Borne and Zoonotic Diseases* 2017;17. <https://doi.org/10.1089/vbz.2017.2135>.
- 641 [48] Vanden Broecke B, Bernaerts L, Ribas A, Sluydts V, Mnyone L, Matthysen E, et al.  
642 Linking Behavior, Co-infection Patterns, and Viral Infection Risk With the Whole  
643 Gastrointestinal Helminth Community Structure in *Mastomys natalensis*. *Front Vet Sci*  
644 2021;8. <https://doi.org/10.3389/fvets.2021.669058>.
- 645 [49] Vanden Broecke B, Tafompa PJJ, Mwamundela BE, Bernaerts L, Ribas A, Mnyone LL, et  
646 al. Drivers behind co-occurrence patterns between pathogenic bacteria, protozoa, and

- 647 helminths in populations of the multimammate mouse, *Mastomys natalensis*. *Acta*  
648 *Trop* 2023;243. <https://doi.org/10.1016/j.actatropica.2023.106939>.
- 649 [50] Haikukutu L, Lyaku JR, Lyimo C, Kasanga CJ, Kandusi SE, Rahelinirina S, et al. Plague in  
650 Tanzania: first report of sylvatic plague in Morogoro region, persistence in Mbulu focus,  
651 and ongoing quiescence in Lushoto and Iringa foci. *IJID Regions* 2022;4:105–10.
- 652 [51] Calvignac-Spencer S, Kouadio L, Couacy-Hymann E, Sogoba N, Rosenke K, Davison AJ,  
653 et al. Multiple DNA viruses identified in multimammate mouse (*Mastomys natalensis*)  
654 populations from across regions of sub-Saharan Africa. *Arch Virol* 2020;165:2291–9.
- 655 [52] Oguge N, Rarieya M, Ondiaka P. A preliminary survey of macroparasite communities of  
656 rodents of Kahawa, Central Kenya. *Belgian Journal of Zoology (Belgium)* 1997;127.
- 657 [53] Weyer J, Thomas J, Leman PA, Grobbelaar AA, Kemp A, Paweska JT. Human cases of  
658 wesselsbron disease, South Africa 2010-2011. *Vector-Borne and Zoonotic Diseases*  
659 2013;13. <https://doi.org/10.1089/vbz.2012.1181>.
- 660 [54] Ndione MHD, Ndiaye EH, Faye M, Diagne MM, Diallo D, Diallo A, et al. Re-Introduction  
661 of West Nile Virus Lineage 1 in Senegal from Europe and Subsequent Circulation in  
662 Human and Mosquito Populations between 2012 and 2021. *Viruses* 2022;14.  
663 <https://doi.org/10.3390/v14122720>.
- 664 [55] Budodo RM, Horumpende PG, Mkumbaye SI, Mmbaga BT, Mwakapuja RS, Chilongola  
665 JO. Serological evidence of exposure to Rift Valley, Dengue and Chikungunya Viruses  
666 among agropastoral communities in Manyara and Morogoro regions in Tanzania: A  
667 community survey. *PLoS Negl Trop Dis* 2020;14:e0008061.
- 668 [56] Mwanyika GO, Sindato C, Rugarabamu S, Rumisha SF, Karimuribo ED, Misinzo G, et al.  
669 Seroprevalence and associated risk factors of chikungunya, dengue, and Zika in eight  
670 districts in Tanzania. *International Journal of Infectious Diseases* 2021;111:271–80.



- 671 [57] Chipwaza B, Mugasa JP, Selemani M, Amuri M, Mosha F. Dengue and Chikungunya  
672 Fever among Viral Diseases in Outpatient Febrile 2014.
- 673 [58] Mboera LEG, Mweya CN, Rumisha SF, Tungu PK, Stanley G, Makange MR, et al. The Risk  
674 of Dengue Virus Transmission in Dar es Salaam, Tanzania during an Epidemic Period of  
675 2014. *PLoS Negl Trop Dis* 2016. <https://doi.org/10.1371/journal.pntd.0004313>.
- 676 [59] Faustine NL, Sabuni EJ, Ndaró AJ, Paul E, Chilongola JO. Chikungunya, Dengue and West  
677 Nile virus Infections in Northern Tanzania. *J Adv Med Med Res* 2017:1–7.
- 678 [60] Mitchell CL, Ngasala B, Janko MM, Chacky F, Edwards JK, Pence BW, et al. Evaluating  
679 malaria prevalence and land cover across varying transmission intensity in Tanzania  
680 using a cross-sectional survey of school-aged children. *Malar J* 2022;21.  
681 <https://doi.org/10.1186/s12936-022-04107-8>.
- 682 [61] Chipwaza B, Mugasa JP, Selemani M, Amuri M, Mosha F, Ngatunga SD, et al. Dengue  
683 and Chikungunya fever among viral diseases in outpatient febrile children in Kilosa  
684 district hospital, Tanzania. *PLoS Negl Trop Dis* 2014;8:e3335.
- 685 [62] Borremans B, Hughes NK, Reijniers J, Sluydts V, Katakweba AAS, Mulungu LS, et al.  
686 Happily together forever: temporal variation in spatial patterns and complete lack of  
687 territoriality in a promiscuous rodent. *Popul Ecol* 2014;56:109–18.
- 688 [63] Vanden Broecke B, Mariën J, Sabuni CA, Mnyone L, Massawe AW, Matthysen E, et al.  
689 Relationship between population density and viral infection: A role for personality?  
690 *Ecol Evol* 2019;9:10213–24.
- 691 [64] Leirs H, Kirkpatrick L, Sluydts V, Sabuni C, Borremans B, Katakweba A, et al. Twenty-  
692 nine years of continuous monthly capture-mark-recapture data of multimammate  
693 mice (*Mastomys natalensis*) in Morogoro, Tanzania. *Sci Data* 2023;10:798.

- 694 [65] Raulino R, Thaurignac G, Keita AK, Esteban A, Goumou S, Diallo R, et al. Seroprevalence  
695 of IgG antibodies against multiple arboviruses in bats from Cameroon, Guinea, and the  
696 Democratic Republic of Congo. *Vector-Borne and Zoonotic Diseases* 2022;22:252–62.
- 697 [66] Amorim JH, Diniz MO, Cariri FAMO, Rodrigues JF, Bizerra RSP, Gonçalves AJS, et al.  
698 Protective immunity to DENV2 after immunization with a recombinant NS1 protein  
699 using a genetically detoxified heat-labile toxin as an adjuvant. *Vaccine* 2012;30:837–  
700 45.
- 701 [67] Li Y, Counor D, Lu P, Duong V, Yu Y, Deubel V. Protective immunity to Japanese  
702 encephalitis virus associated with anti-NS1 antibodies in a mouse model. *Virology*  
703 2012;9:1–13.
- 704 [68] Bailey MJ, Broecker F, Duehr J, Arumemi F, Krammer F, Palese P, et al. Antibodies  
705 elicited by an NS1-based vaccine protect mice against Zika virus. *MBio* 2019;10:10–  
706 1128.
- 707 [69] Easterbrook JD, Klein SL. Immunological mechanisms mediating hantavirus persistence  
708 in rodent reservoirs. *PLoS Pathog* 2008;4.  
709 <https://doi.org/10.1371/journal.ppat.1000172>.
- 710 [70] Spengler JR, Haddock E, Gardner D, Hjelle B, Feldmann H, Prescott J. Experimental  
711 Andes Virus Infection in Deer Mice: Characteristics of Infection and Clearance in a  
712 Heterologous Rodent Host. *PLoS One* 2013;8.  
713 <https://doi.org/10.1371/journal.pone.0055310>.
- 714 [71] Borremans B. Ammonium improves elution of fixed dried blood spots without affecting  
715 immunofluorescence assay quality. *Tropical Medicine and International Health*  
716 2014;19. <https://doi.org/10.1111/tmi.12259>.

- 717 [72] Ayouba A, Thaurignac G, Morquin D, Tuailon E, Raulino R, Nkuba A, et al. Multiplex  
718 detection and dynamics of IgG antibodies to SARS-CoV2 and the highly pathogenic  
719 human coronaviruses SARS-CoV and MERS-CoV. *Journal of Clinical Virology*  
720 2020;129:104521.
- 721 [73] R Core Team. R: A Language and Environment for Statistical Computing. R Foundation  
722 for Statistical Computing, Vienna, Austria 2024.
- 723 [74] Killick R, Eckley IA. Changepoint: An R package for changepoint analysis. *J Stat Softw*  
724 2014;58. <https://doi.org/10.18637/jss.v058.i03>.
- 725 [75] Donnelly Jr RA, Abdel-Raouf F. *Statistics*, 3E. Penguin; 2016.
- 726 [76] Piantadosi A, Kanjilal S. Diagnostic approach for arboviral infections in the united  
727 states. *J Clin Microbiol* 2020;58. <https://doi.org/10.1128/JCM.01926-19>.
- 728 [77] Dorai-Raj S. binom: Binomial confidence intervals for several parameterizations. R  
729 Package Version 2014;1.
- 730 [78] Revelle W. Package “psych” - Procedures for Psychological, Psychometric and  
731 Personality Research. R Package 2015.
- 732 [79] Warnes GR, Bolker B, Bonebakker L, Gentleman R, Huber W, Liaw WHA, et al. gplots:  
733 Various R Programming Tools for Plotting Data. R Package Version 313 2022.
- 734 [80] Lenth R V. emmeans: Estimated Marginal Means, aka Least-Squares Means. R Package  
735 Version 189 2023.
- 736 [81] Muff S, Nilsen EB, O’Hara RB, Nater CR. Rewriting results sections in the language of  
737 evidence. *Trends Ecol Evol* 2022;37. <https://doi.org/10.1016/j.tree.2021.10.009>.
- 738 [82] Chandra G, Mukherjee D. Effect of climate change on mosquito population and  
739 changing pattern of some diseases transmitted by them. *Advances in Animal*  
740 *Experimentation and Modeling*, Elsevier; 2022, p. 455–60.

- 741 [83] Mwanyika GO, Mboera LEG, Rugarabamu S, Ngingo B, Sindato C, Lutwama JJ, et al.  
742 Dengue virus infection and associated risk factors in africa: A systematic review and  
743 meta-analysis. *Viruses* 2021;13. <https://doi.org/10.3390/v13040536>.
- 744 [84] Madewell ZJ. Arboviruses and their vectors. *South Med J* 2020;113.  
745 <https://doi.org/10.14423/SMJ.0000000000001152>.
- 746 [85] Mboya LB, Nkya ES, Matemba LE, Kinimi E. Evidence of Chikungunya but not Dengue  
747 Virus Circulating among Febrile Patients during Low Transmission Period in Morogoro  
748 Municipality, Tanzania. *Int J Trop Dis Health* 2020.  
749 <https://doi.org/10.9734/ijtdh/2019/v40i430236>.
- 750 [86] Kasbergen LMR, Nieuwenhuijse DF, de Bruin E, Sikkema RS, Koopmans MPG. The  
751 increasing complexity of arbovirus serology: An in-depth systematic review on cross-  
752 reactivity. *PLoS Negl Trop Dis* 2023;17:e0011651.
- 753 [87] Lefkowitz EJ, Dempsey DM, Hendrickson RC, Orton RJ, Siddell SG, Smith DB. Virus  
754 taxonomy: The database of the International Committee on Taxonomy of Viruses  
755 (ICTV). *Nucleic Acids Res* 2018;46. <https://doi.org/10.1093/nar/gkx932>.
- 756 [88] Cho B, Jeon BY, Kim J, Noh J, Kim J, Park M, et al. Expression and evaluation of  
757 Chikungunya virus E1 and E2 envelope proteins for serodiagnosis of chikungunya virus  
758 infection. *Yonsei Med J* 2008;49. <https://doi.org/10.3349/ymj.2008.49.5.828>.
- 759 [89] Chang H-H, Huber RG, Bond PJ, Grad YH, Camerini D, Maurer-Stroh S, et al. Systematic  
760 analysis of protein identity between Zika virus and other arthropod-borne viruses. *Bull*  
761 *World Health Organ* 2017;95. <https://doi.org/10.2471/blt.16.182105>.
- 762 [90] Fink AL, Engle K, Ursin RL, Tang W-Y, Klein SL. Biological sex affects vaccine efficacy and  
763 protection against influenza in mice. *Proceedings of the National Academy of Sciences*  
764 2018;115:12477–82.

- 765 [91] Chrousos GP. Stress and sex versus immunity and inflammation. *Sci Signal*  
766 2010;3:pe36–pe36.
- 767 [92] Kelly CD, Stoehr AM, Nunn C, Smyth KN, Prokop ZM. Sexual dimorphism in immunity  
768 across animals: a meta-analysis. *Ecol Lett* 2018;21:1885–94.
- 769 [93] Vincze O, Vágási CI, Péntes J, Szabó K, Magonyi NM, Czirják GÁ, et al. Sexual  
770 dimorphism in immune function and oxidative physiology across birds: The role of  
771 sexual selection. *Ecol Lett* 2022;25:958–70.
- 772 [94] Jacobsen H, Klein SL. Sex differences in immunity to viral infections. *Front Immunol*.  
773 2021; 12: 720952 2021.
- 774 [95] Sciarra F, Campolo F, Franceschini E, Carlomagno F, Venneri MA. Gender-Specific  
775 Impact of Sex Hormones on the Immune System. *Int J Mol Sci* 2023;24:6302.
- 776 [96] Wilkinson NM, Chen H-C, Lechner MG, Su MA. Sex differences in immunity. *Annu Rev*  
777 *Immunol* 2022;40:75–94.
- 778 [97] Mlyashimbi ECM, Mlyashimbi ECM, Mariën J, Kimaro DN, Tarimo AJP, Machang’U RS,  
779 et al. Home ranges, sex ratio and recruitment of the multimammate rat (*Mastomys*  
780 *natalensis*) in semi-arid areas in Tanzania. *Mammalia* 2020;84.  
781 <https://doi.org/10.1515/mammalia-2019-0048>.
- 782 [98] Borremans B, Leirs H, Gryseels S, Günther S, Makundi R, De Bellocq JG. Presence of  
783 Mopeia virus, an African arenavirus, related to biotope and individual rodent host  
784 characteristics: Implications for virus transmission. *Vector-Borne and Zoonotic*  
785 *Diseases* 2011;11. <https://doi.org/10.1089/vbz.2010.0010>.
- 786 [99] Borremans B, Vossen R, Becker-Ziaja B, Gryseels S, Hughes N, Van Gestel M, et al.  
787 Shedding dynamics of Morogoro virus, an African arenavirus closely related to Lassa  
788 virus, in its natural reservoir host *Mastomys natalensis*. *Sci Rep* 2015;5:1–8.

789

## 790 Supporting information

791 **S1 Fig. Histograms of wild-caught *M. natalensis* for each tested arbovirus with on the x-axis the relative**  
792 **antibody units in a logarithmic scale.** The relative antibody units are calculated according to the positive control  
793 dilution series. The calculated cutoff values are represented by the colored vertical lines: 'CHP.m' is the  
794 changepoint mean, 'CHP.mv' is the changepoint mean-variance, 'CHP.v' is the changepoint variance, 'NegCtrl' is  
795 the mean plus three times the standard deviation of the negative control samples and 'Recap' is the maximum  
796 value of an antibody development curve based on recaptured seroconverted wild-caught *M. natalensis*.

797 **S2 Fig. Seroprevalence, according to the calculated cutoff methods, of the wild-caught *M. natalensis* with 95%**  
798 **confidence interval for each of the tested arboviruses.** The cutoff methods: 'CHP.m' is the changepoint mean,  
799 'CHP.mv' is the changepoint mean-variance, 'CHP.v' is the changepoint variance, 'NegCtrl' is the mean plus three  
800 times the standard deviation of the negative control samples and 'Recap' is the maximum value of an antibody  
801 development curve based on recaptured seroconverted wild-caught *M. natalensis*. Each calculated  
802 seroprevalence was compared to the 'Recap' seroprevalence using a Chi-square test, significant difference  
803 is depicted in asterisk (\*) symbols. P-values: \* 0.1 -  $\geq$  0.01; \*\* < 0.01 -  $\geq$  0.001; \*\*\* < 0.001.

804 **S1 Table. Analysis of variance from the generalized linear model (logit link function and binomial error**  
805 **distribution) with the response variable being the binary serostatus of each sample.** Sex and age and their  
806 interaction were included as explanatory variables. P values with at least a weak statistical support are marked  
807 in bold ( $p < 0.1$ ).