

Environmental heterogeneity drives tsetse fly population dynamics

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Abstract

A spatially and temporally heterogeneous environment may lead to unexpected population dynamics, and knowledge still is needed on which of the local environment properties favour population maintenance at larger scale. As regards pathogen vectors, such as tsetse flies transmitting human and animal African trypanosomosis, such a knowledge is crucial for proposing relevant management strategy. We developed an original mechanistic spatio-temporal model of tsetse fly population dynamics, accounting for combined effects of spatial complexity, density-dependence, and temperature on the age-structured population, and parametrized with field and laboratory data. We confirmed the strong impact of temperature and adult mortality on tsetse populations. We showed that patches with the lowest mean temperatures and lowest variations act as refuges when adult mortality is homogeneously increased. Our results highlighted the importance of baseline data collection to characterize the targeted ecosystem before any control measure is implemented to maximize its efficiency.

1 **1. Introduction**

2 Environmental spatial heterogeneity is a key driver of population dynamics (Tilman & Kareiva 1997;
3 Vinatier et al. 2011), inducing movements from source to sink patches possibly enhancing population
4 persistence in unsuitable patches (Holt 1985; Pulliam 1988). In addition, environmental suitability
5 varies over time both at local scale, due to microclimate variations as related to vegetation growth
6 (Keppel et al. 2017), and at large scale, due to a seasonal occurrence of unfavourable periods.
7 Confounding the role of spatial and temporal environmental heterogeneity potentially gives rise to
8 erroneous predictions of ecological processes (Clark 2005). However, relating such a complex time-
9 and space-varying habitat with population dynamics still is a challenge in ecology (Sutherland et al.
10 2013; Crone 2016; Griffith et al. 2016). Therefore, illustrative examples about the complex interplay
11 between spatio-temporal environmental variability and population dynamics are welcome to feed
12 theory and assess which patch properties (co)contribute to define sources and sinks in heterogeneous
13 environments.

14 This is particularly true when it comes to controlling infectious diseases, given that vector-borne
15 disease dynamics is largely determined by those of vector populations (Hartemink et al. 2015). First,
16 spatial heterogeneity is expected to favour vector persistence thanks to the rescue effect, especially if
17 control is not area-wide, i.e. targeting an entire insect pest population within a circumscribed area
18 (Reichard 2002; Hendrichs et al. 2007). Second, such populations and associated pathogens face
19 seasonal variations of habitat suitability (Charron et al. 2013). Environmental suitability varying in
20 space and time could induce unexpected population dynamics, potentially impairing its management,
21 whereas control strategies are nonetheless often elaborated without considering local environmental
22 specificities.

23 Tsetse flies (*Glossina* spp.) are vectors of African trypanosomes, widely recognized as a major
24 pathological constraint for efficient livestock species and agricultural development in sub-Saharan
25 Africa (Alsan 2015). *Trypanosoma* spp. parasites both cause Human African Trypanosomosis and
26 African Animal Trypanosomosis. Widely distributed, they occur in 38 countries and infest 10 million

27 km² (Vreysen et al. 2013), with over 60 million people continuously exposed to the risk of infection
28 of this neurological, potentially lethal disease, mainly in remote rural areas with limited access to
29 health services. Besides, farms in tsetse-infested areas suffer a 20% to 40% loss in livestock
30 productivity, adding up to an estimated \$4500 million loss annually for producers and consumers in
31 sub-Saharan Africa (Budd 1999). Among the 31 species and subspecies known of tsetse flies, a third
32 is of economic (agricultural and veterinary) and sanitary importance (Solano et al. 2010a). Efforts to
33 manage the vector and the disease in Africa are on-going for decades but largely fail to create
34 sustainable tsetse free areas, resulting in only a reduction of less than 2% of tsetse distribution
35 (Allsopp 2001; Bouyer et al. 2013a). Although tsetse flies turned out to be extremely complex species,
36 their very low rate of reproduction would make them a relevant target to eradicate, making crucial to
37 better apprehend their spatio-temporal population dynamics (Peck & Bouyer 2012).

38 Mathematical models have proved to be relevant tools in ecology, to better understand the dynamics
39 of populations (Hasting 2012) and to predict such dynamics under modified conditions (Evans et al.
40 2012). Process-based models incorporate at minimal costs sparse and heterogeneous knowledge from
41 various areas, species, and fields of expertise. Simulations are complementary to field observations
42 and experiments (Restif et al. 2012), enabling the fast acquisition of quantitative predictions which
43 can in turn emphasize the need for further biological investigations. Moreover, the range of
44 behaviours of complex systems can be scanned using mechanistic models, and scenarios are tested
45 easily (Cailly et al. 2012). Provided hypotheses and limits are clearly stated (Getz et al. 2018), models
46 can guide decision-making (Sutherland & Freckleton 2012).

47 As regards tsetse biology and population dynamics, entomologists quickly realized how useful
48 models could be (Rogers 1988, 1990; and more recently: Vale & Torr 2005; Lin et al. 2015), and
49 encouraged their use when designing management decisions (Hargrove 2003; Childs 2011; Meyer et
50 al. 2018). However, most models have failed to predict the persistence of target populations leading
51 to misleading guidelines for control programs (Peck & Bouyer 2012; Bouyer et al. 2013b). Most of
52 these programs were not implemented following area-wide principles (Klassen 2005) and their failure

53 could be imputed to population resurgence in non-eradicated patches or re-invasion of the targeted
54 zone by neighbouring populations (Meyer et al. 2016; Lord et al. 2017). However, it is still unclear
55 what the relevant patch properties are and how they combine to define sources and sinks in a hostile
56 environment created by eradication efforts. To address such an issue, the spatial complexity of the
57 environment has to be accounted for. While omitted in most models until recently, it has been shown
58 to considerably influence predictions once incorporated (Peck 2012; Barclay & Vreysen 2013; Lord
59 et al. 2017). Indeed, population dynamics is expected to vary locally among patches of variable
60 suitability, possibly affecting population dynamics at large metapopulation scale. To better assess
61 how large scale tsetse fly population dynamics are affected by local dynamics, there is a need for an
62 integrated spatio-temporal model thoroughly evaluated against field and experimental data, and fed
63 by environmental data to account for landscape heterogeneity.

64 To assess if spatial and temporal environmental heterogeneity drives tsetse fly population dynamics
65 at large scale, we developed an original mechanistic spatio-temporal model of tsetse fly population
66 dynamics and incorporated environmental heterogeneity through a data-driven approach. The model
67 was applied to *Glossina palpalis gambiensis* in the Niayes (Senegal), a region with an ongoing
68 eradication project (Dicko et al. 2014). In this area, less than 4% of the habitat is suitable (Bouyer et
69 al. 2010), and tsetse flies harbour a metapopulation structure (Solano et al. 2010b). This knowledge
70 was incorporated in the model, accounting for combined effects of spatial complexity, density-
71 dependence, and temperature on the age-structured population.

72 **2. Material and methods**

73 **Key knowledge on tsetse biology**

74 Meteorological variables influence the abundance and spatial distribution of arthropod disease
75 vectors (Hay et al. 1996). For tsetse flies, effect magnitude depends on species (Rogers & Randolph
76 1991; Rogers et al. 1996; Hargrove 2001), but average temperature is the most influential

77 meteorological variable on life cycle (Hargrove 2004). However, its influence compared to or
78 combined with demographic processes is barely known.

79 The tsetse fly is adenotrophic viviparous: the egg hatches in the female and the larva is nourished by
80 dedicated organs until larviposition. A temperature decrease lengthens the time between
81 larvipositions (Harley 1968). Similarly, the colder it gets in breeding sites, the longer the pupa
82 development (Glasgow 1963; Phelps & Burrows 1969a,b). After pupa emergence, the newly emerged
83 fly (teneral) takes its first blood meal to strengthen its musculature and reproduce. The first oocyte
84 maturation into pupa takes around 18 days, making the first larviposition longer than subsequent ones
85 (10 days) depending on species and temperature (Hargrove 2004).

86 Extreme temperatures, cold or warm, increase fly mortality (Hargrove 2001). Mortality, related to
87 predation and feeding success, is density-dependent (Rogers & Randolph 1984) and age-dependent
88 (Hargrove 1990), with remarkably high losses in tenerals partly due to starvation risk (Phelps &
89 Clarke 1974; Hargrove 2004). Learning capacities of older flies make them return on their first host,
90 increasing their hunting efficiency with age (Bouyer et al. 2007).

91 Tsetse flies are classified into three groups of different behaviours and distributions: forest (subgenus
92 *Fusca*), savannah (subgenus *Morsitans*), and riverine flies (subgenus *Palpalis*). Most of previous
93 model concerned *Glossina pallidipes* and *G. morsitans*, both of the savannah group. We focused on
94 *G. p. gambiensis*, a riverine fly living in forest galleries and riparian thickets (Bouyer et al. 2005).
95 Due to habitat characteristics, this species is known to mostly disperse in one dimension (along
96 rivers). However, climate changes induce the disappearance of rivers and associated vegetation as
97 evidenced in our study area (Niayes, Senegal, Fig. 1). *G. p. gambiensis* adapted to patchy vegetation
98 mainly associated to human watering activities (Bouyer et al. 2010), dispersing in two dimensions.
99 Furthermore, isolated populations in fragmented habitats are preferential targets for area-wide
100 integrated pest management programs (Hendrichs et al. 2007; Bouyer et al. 2015). Hence, our case
101 study is of broad relevance for better understanding and predicting tsetse fly spatio-temporal

102 population dynamics in rapidly changing ecosystems that are gradually becoming the norm (Guerrini
103 et al. 2008).

104 **Data on tsetse biology**

105 Variations in mortality and fecundity with temperature were measured for the studied strain under
106 experimental conditions (Pagabeleguem et al. 2016). We used data on the first larval period (time
107 between emergence and first pupa production) and on subsequent inter-larval periods (time between
108 reproductive cycles). As the colony was maintained at 24°C with only temperatures above 24°C tested
109 to assess the maximum critical temperature for flies, most data used to estimate female mortality were
110 obtained at 24°C and none at a lower temperature. In addition, the effect of temperature on *G. p.*
111 *gambiensis* pupal duration was measured under experimental conditions (Centre International de
112 Recherche-Développement sur l'Élevage en zones Subhumides, CIRDES, Bobo-Dioulasso, Burkina
113 Faso, 2009). One hundred and twenty 20-day old pupae were held in climate controlled rooms until
114 emergence. The experiment was replicated three times for each temperature tested (Table S1).

115 Dispersing abilities of *G. p. gambiensis* were assessed from release-recapture data of marked sterile
116 males (Oct. 2010 to Dec. 2012; Pagabeleguem 2012). Flies were mass-reared in CIRDES Burkina
117 Faso and shipped as irradiated pupae to Senegal (Pagabeleguem et al. 2015). Flies were released twice
118 a month in four locations (Parc de Hann in Dakar, Diacksaw Peul, Pout, and Kayar; Fig. 1). Two
119 release points were selected per location (in suitable vs. unsuitable habitats). Released flies were
120 trapped using Vavoua traps (Laveissière & Grébaud 1990) up to 2kms from release points. Distance
121 between traps varied between 100m and 300m. Traps were set in the morning before 9:00 and
122 collected in the afternoon after 16:00, every 3 days. The monitoring of a release stopped when less
123 than 2 marked males were recaptured.

124 In another study, natural abortion rate was monitored in Hann, Diacksaw, Sebikotane, and Pout (Fig.
125 1). Ten traps per site were deployed monthly from March 2008 to February 2009, and then every
126 three months until September 2010 (Hann, Diacksaw) or December 2011 (Pout, Sebikotane). Flies
127 were collected at least once a day. Fresh flies were dissected to estimate their ovarian sequence. This

128 female dataset was used to calculate the population age structure, to be compared to simulation results
129 for partial validation.

130 **Environmental data**

131 The spatio-temporal heterogeneity of the environment was realistically represented using an original
132 data-driven approach. The environmental carrying capacity and the local daily temperatures were
133 incorporated in the model.

134 The carrying capacity was defined as the maximum sustainable number of individuals for a given
135 area and was estimated as (Eq. 1):

$$136 \quad k = \frac{SI \times ADT}{\sigma} \quad (\text{Eq. 1})$$

137 with *SI* the suitability index, estimated with a species distribution model (Dicko et al. 2014) based on
138 the maximum entropy (Maxent) (Supporting Information 2.1), σ the trap efficiency, i.e. the
139 probability that a trap catches a fly within 1km² within a day (Barclay and Hargrove 2005), and *ADT*
140 the apparent density of flies per trap per day (Dicko et al. 2015). All available data from catches
141 obtained between 2007 and 2010 in the Niayes before the start of the eradication campaign were used
142 to estimate local carrying capacities (Supporting Information 2.1).

143 Air temperatures measured in weather stations are not those experienced by flies in resting places.
144 Indeed, flies prefer microenvironments that are 2-6°C lower than the ambient temperature (Hargrove
145 & Coates 1990). In addition, temperature largely varies from the centre of a gallery forest towards its
146 edges (Bouyer 2006). Therefore, micro-climate and approximated local temperatures truly perceived
147 by tsetse flies were explicitly modelled using input data from weather stations transformed using a
148 spatio-temporal geostatistical model (Kilibarda et al. 2014). Available temperature data recorded in
149 selected suitable patches were used to correct the bias present in moderate resolution imaging
150 spectroradiometer (MODIS) Land-Surface Temperature (macro-climate; Supporting Information
151 2.2). High resolution macro-climate data were available only for 2011. Approximated temperatures

152 were used as model inputs in a zone known as suitable for tsetse to check if the simulated population
 153 persisted as expected.

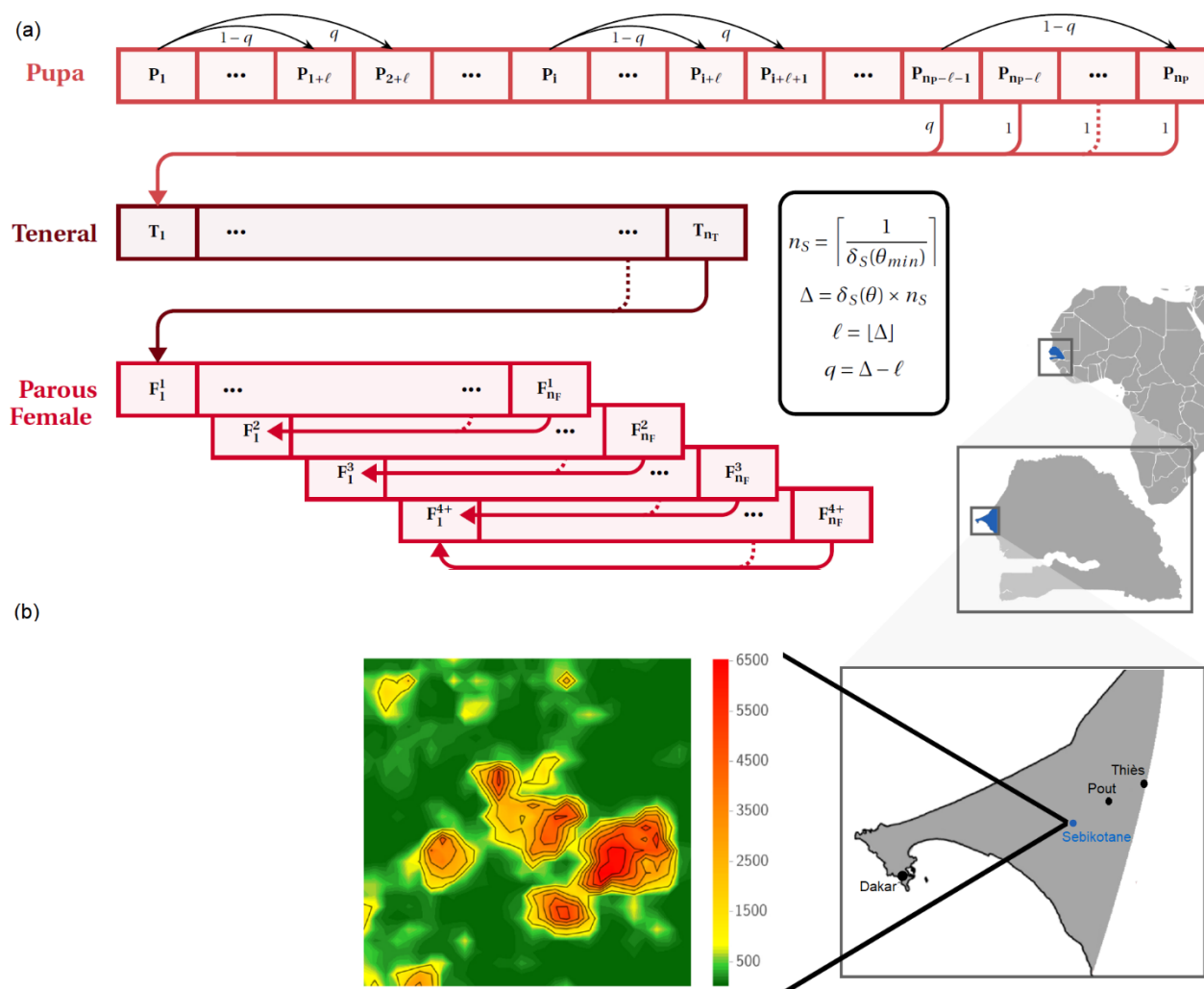


Figure 1. Local and general tsetse fly population dynamics applied to the Niayes in Senegal. (a) within-cell model diagram of tsetse fly populations dynamics (time unit is a day). All transitions between stages except P to T triggers the birth of a new pupa P_1 . Transitions occur at a development rate δ_S for stage S according to temperature $\theta_{t,c}$ at time t in cell c , giving rise daily to a minimum jump of l states from each state i of stage S , with $(1-q)S_{t,c,i}$ individuals going from state $S_{t,c,i}$ to state $S_{t+1,c,i+l}$ and $qS_{t,c,i}$ individuals going to $S_{t+1,c,i+l+1}$. If $i + l > n_S$ (respectively $i + l + 1 > n_S$), then concerned individuals go to the next stage. Stage $S \in \{P, T, F_x, M\}$, parity $x \in \{1, 2, 3, 4+\}$. (b) Map of Senegal identifying areas providing field data and localizing the 30x30 simulated area, highlighting the spatial heterogeneity in local carrying capacities k_c (inbox, scale in number of individuals).

154 A mechanistic spatio-temporal model of tsetse fly population dynamics

155 A mechanistic and deterministic compartmental model was developed to predict the spatio-temporal
 156 tsetse fly population dynamics accounting for environmental heterogeneity and including density-
 157 dependence. Individuals were categorized into pupae (P), without differentiating males and females,

158 teneral (T, immature nulliparous females), and parous females with four stages (F₁, F₂, F₃, F₄₊; Fig.
 159 1a) in agreement with ovarian dissection which provides accurate information about the first four
 160 parities (Hargrove & Ackley 2015). Adult males (M) were not considered limiting for breeding. They
 161 could mate from the age of 6 days, regardless of temperature, after which they were only subject to
 162 mortality. They played a role in density-dependent processes. The environment was modelled using
 163 a grid (cell resolution: 250m x 250m; study area: 30 x 30 cells; Fig. 1b). The model was developed
 164 in Python as a discrete-time model with a one-day time step (Supporting Information 6).

165

166 **Within-cell dynamics** - The population size of life stage S at time t in cell c decreased with mortality,
 167 following a negative exponential model of instantaneous rate $\mu_{S,t,c}$ (Eq. 2). Considering the lack of
 168 data on pupa mortality, we used a constant rate (Eq. 3) of 0.01 day⁻¹ (Childs 2011). For adults, the
 169 log of mortality rates increased linearly with temperature after 24°C (Hargrove 2004). Below this
 170 threshold, and for the range of temperatures observed in the field, the literature and the lack of data
 171 suggested a constant mortality rate (Eq. 4). Age-dependence was featured by setting teneral mortality
 172 to twice that of mature females (Eq. 5). Density-dependence occurred when the adult population
 173 exceeded the cell carrying capacity (Eq. 6-7, Hargrove 2004).

174 $S_{t+\Delta t,c} = S_{t,c}e^{-\mu_{S,t,c}\Delta t}$, with stage $S \in \{P, T, F_x, M\}$, parity $x \in \{1, 2, 3, 4+\}$, $\Delta t = 1$ (Eq. 2)

175 $\mu_P = m_P$ (Eq. 3)

176 $\mu_{X,t,c} = \begin{cases} \mu_{X,t,c}(\theta_{t,c} = 24^\circ\text{C}), & \text{if } \theta_{t,c} \leq 24^\circ\text{C} \\ \mu_{X,t,c}(\theta_{t,c}), & \text{if } \theta_{t,c} > 24^\circ\text{C} \end{cases}, X \in \{T, F, M\}$ (Eq. 4)

177 $\mu_{T,t,c} = 2\mu_{F,t,c}$ (Eq. 5)

178 $\mu_{X,t,c} = \beta_{t,c}e^{m_{1,X}\theta_{t,c}+m_{2,X}}$, $X \in \{F, M\}$ (Eq. 6)

179 $\beta_{t,c} = \begin{cases} 1, & \text{if } \frac{A_{t,c}}{k_c} \leq 1 \\ \frac{A_{t,c}}{k_c}, & \text{if } \frac{A_{t,c}}{k_c} > 1 \end{cases}$, with $A_{t,c} = T_{t,c} + \sum_{i=1}^4 F_{i,t,c} + M_{t,c}$ (Eq. 7)

180 In addition, individuals evolved within and between stages as a function of temperature. Pupa
 181 development function $\delta_{P,t,c}$ was fitted on data. For adults and teneral, consistency of experimental
 182 data on the target species was checked against published equations (Hargrove 2004; Eq. 8; Fig. 2):

$$183 \quad \delta_{X,t,c} = d_{1,X}(\theta_{t,c} - 24) + d_{2,X}, X \in \{T, F\} \quad (\text{Eq. 8})$$

184 Each stage was discretized into n_S states, n_S being the longest duration in stage S obtained with its
 185 development rate δ_S calculated at the minimum temperature of the year $\min(\theta_{t,c})$ (Fig. 1a). For
 186 higher temperatures, individuals made a leap forward in the development vector, the interval being
 187 determined by the integer part l of Δ (Eq. 9, Fig. 1a).

$$188 \quad \Delta_{S,t,c} = \delta_{S,t,c}(\theta_{t,c})n_S \quad (\text{Eq. 9})$$

189 To avoid discretization artefacts, individuals were proportionally divided into two successive states
 190 according to the decimal part q of Δ (Fig. 1a). Individuals who reached state n_S evolved to the next
 191 stage, a pupa being produced if teneral or adult females were concerned. After the fourth parity,
 192 females looped back to the start of F_{4+} .

193

194 **Between-cell dynamics** - An original dispersal pattern was designed favouring suitable over hostile
 195 habitats to align with species behaviour. The proportion $p_{t,c}$ of flies leaving cell c at time t was
 196 controlled by a sigmoidal density-dependent dispersal rate (Lloyds-Smith, 2010), adapted for
 197 individuals competing to access resources (Rogers & Randolph 1984) (Eq. 10):

$$198 \quad p_{t,c} = \frac{1}{1 + e^{-g\left(\frac{A_{t,c}}{k_c} - 1\right)}} \quad (\text{Eq. 10})$$

199 with k_c the carrying capacity in cell c , $A_{t,c}$ the number of adults in cell c at time t , and g a shape

$$200 \quad \text{parameter set to 10 meaning that } p_{t,c} \begin{cases} \approx 0, \text{ if } A_{t,c} < 0.5k_c \\ \approx 1, \text{ if } A_{t,c} > 1.5k_c \text{ (Fig. S1).} \\ 0.5, \text{ if } A_{t,c} = k_c \end{cases}$$

201 The spatial distribution of dispersing flies from cell c to neighbouring cells $Prob_{c \rightarrow i \in \{v\}}$ was set by
 202 the relative attractiveness of neighbouring cells $a_{t,i \in \{v\}}$ (Eq. 11-12). This attractiveness was designed
 203 to favour the emptiest cells ($A_{t,i} \ll k_i$) and cells of greatest k_i if equal $A_{t,i}$. An extended Moore

204 neighbourhood of range r was used: flies dispersed from a cell to its $(2r + 1)^2$ neighbours (v),
205 including the cell itself and diagonals. Parameter r is the maximum distance reached daily, in number
206 of cells, rather than the effective distance covered per fly per day, as the trajectory is not linear. It was
207 calibrated on data by looking at the average $\frac{distance(m)}{time(days)}$ between release and capture of marked flies
208 (Fig. S2).

$$209 \quad a_{t,i \in \{v\}} = \frac{\left(1 - e^{-\frac{-k_i}{A_{t,i}}}\right) k_i}{\max(k_{i \in \{v\}})} \quad (\text{Eq. 11})$$

$$210 \quad Prob_{c \rightarrow i \in \{v\}} = \frac{a_i}{\sum_{j \in v} a_j} \quad (\text{Eq. 12})$$

211 **Model analysis**

212 The reference scenario was examined (parameter values provided in Table S2). The individual and
213 joint effects of input variations on aggregated output variance (Table S3) were evaluated through a
214 global sensitivity analysis. Population size and age structure were outputs of interest. As traps do not
215 capture tenerals and old females as efficiently as females of intermediate parities (Sanders 1962),
216 predicted age structure was compared with field data for $\frac{F_{i=1,2,3}}{F_1 + F_2 + F_3}$.

217 A 3-year burn-in period was simulated starting with $T_{0,c} = M_{0,c} = 0.5k_c$ ($A_{0,c} = k_c$), using reference
218 parameter values. Then, parameter values of each of the tested scenarios were applied for three more
219 years. Carrying capacities were spatially heterogeneous (Fig. 1b) but assumed constant over time.
220 Perceived temperatures, estimated daily per cell for a year, were repeated between years.

221 A variance-based global sensitivity analysis was performed using the Fourier Amplitude Sensitivity
222 Testing (FAST) method (Saltelli et al. 2008). Mortality and development functions of each life stage
223 were tuned with weighting coefficients. A common weight was applied to all adult mortalities (T , M ,
224 $F_{1:4+}$) to preserve model hypotheses. A weighting coefficient also was applied to carrying capacities,
225 thus regulating density-dependence magnitude. As the dispersal rate should remain in the range [0-
226 1], the shape parameter g was varied (Fig. S1). Parameters varied by $\pm 5\%$ of their reference value.

227 The same range, when applied to temperature, changed the annual mean by more than 2°C, which
 228 was far greater than what was observed. Therefore, a variation of ±0.3°C was used, corresponding to
 229 the average deviation from the daily mean in the area (Fig. S4). First order and interaction sensitivity
 230 indices were calculated per parameter (Saltelli et al. 2008).

231 Evaluation of control strategies

232 A control strategy was mimicked by increasing adult mortality (from +2.5% to +100%)
 233 homogeneously in space, and assessed with respect to the female population (T+F_{1:4+}) over time
 234 (every year for 5 years) and space. At the end of simulations, two ratios were computed :

235 $\frac{T_{t_{max,c}+F_{1:4+,t_{max,c}}}}{\sum_c T_{t_{max,c}+F_{1:4+,t_{max,c}}}}$ highlighted cells contributing the most to the female population in the area, while

236 $\frac{(T_{t_{max,c}+F_{1:4+,t_{max,c}}})_{scenario}}{(T_{t_{max,c}+F_{1:4+,t_{max,c}}})_{reference}}$ quantified the local impact of increased mortality compared to natural

237 levels. The correlation between environmental variables and the spatial structure of the remaining
 238 population was assessed.

239 3. Results

240 New insights from biological data

241 New equations were calibrated for temperature-dependent processes of the life cycle of tsetse flies
 242 combining published and new observed data (Fig. 2). The log-linear function for adult mortality
 243 (Table S2) differed from published ones for other species (Fig. 2a). Up to 24°C, female mortality rate
 244 was 0.013 day⁻¹, then it grew exponentially to reach 0.023 day⁻¹ at 32°C. Male mortality was higher
 245 than female one (Table S2, Fig. S3).

246 Pupa emergence clearly followed a logistic equation when fitted on observed data, providing a new
 247 pattern compared to Hargrove's equation (2004) (Fig 2b, Eq. 15, Table S4).

248
$$\delta_{P,t,c} = \left(d_{1,P} + \frac{d_{2,P} - d_{1,P}}{1 + e^{\frac{d_{3,P} - \theta_{t,c}}{d_{4,P}}}} \right)^{-1} \quad (\text{Eq. 15})$$

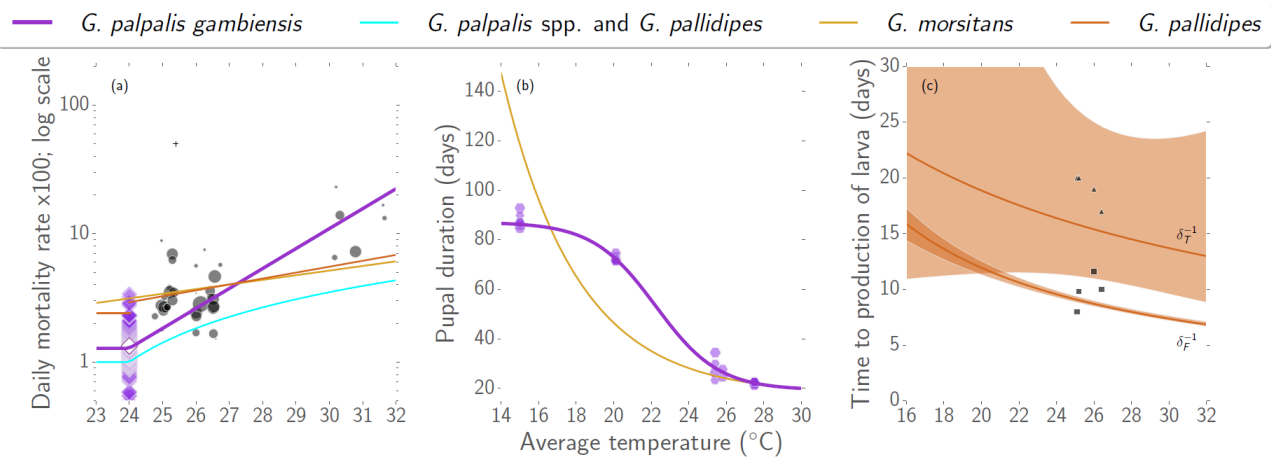


Figure 2. Data (as dots) and predictions (as lines) fitted on new data (if relevant) and from literature for temperature-dependent processes of the model: (a) adult female daily mortality rate (in log-scale); (b) pupal duration (in days); (c) time to larviposition for teneral (T, upper curve, triangles) and parous females (F, lower curve, squares). Data from Pagabeleguem et al. (2016) is shown in grey (the cross in (a) was considered an outlier). New data on *G. p. gambiensis* (from FAO/IPCL and CIRDES) is shown in purple, with the barycentre of mortality rate at 24°C highlighted as a white-filled diamond. Purple thick lines are the newly calibrated equations used in the population dynamics model. Predictions from Barclay's equation (2011) is in cyan. Orange lines correspond to predictions from Hargrove's equations (2004), with filled areas in (c) corresponding to prediction intervals. Equations for time to larviposition were not modified as only few new data was available, which is consistent with Hargrove's equation.

249 Mark-release-recapture data indicated a dispersal range r of one cell, the daily average distance
 250 proved to be less than 250m (Fig. S2).

251 Finally, the spatial heterogeneity of carrying capacities was high, ranging from 7 to 6548 individuals
 252 (median: 145) per cell. On the contrary, spatial variations of local temperatures were small, the
 253 standard deviation over the grid never exceeding 0.67°C at any time step.

254 Reference scenario analysis

255 The reference scenario was closely in line with field observations made before the start of the Niayes'
 256 control program (Fig. 3). Population dynamics was seasonal (Fig. 3b), and driven by temperature as
 257 expected (Fig. 3a). Female population (T+F_{1:4+}) was stable across years with a growth rate of -0.75%
 258 the last simulation year. As observed (Fig. 3d), females between first and third larviposition
 259 (F₁+F₂+F₃) were distributed on average for 40% in F₁, 33% in F₂, and the rest in F₃ (Fig. 3c). The
 260 spatial variability of age structure was 3 to 4 times lower than its temporal variability.

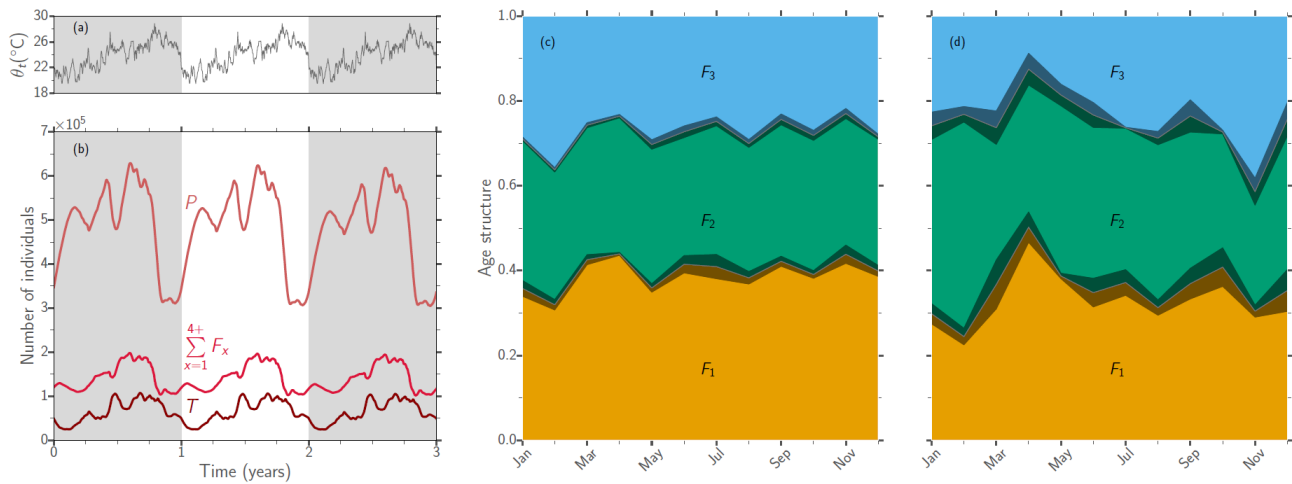


Figure 3. Model predictions for the reference scenario: (a) average daily temperatures over three years (in °C); (b) total number of individuals per stage (P: pupae, T: teneral, F: parous females) in the grid (56.25 km²) over three years of simulation; (c) female age structure ($\frac{F_{i=1,2,3}}{F_1+F_2+F_3}$) during the last year of simulation; (d) observed female age structure (captures and dissection occurred from 2008 to 2011 in the Niayes; results were averaged by month, all years and locations aggregated; grey filled areas are confidence intervals around the mean: $\frac{\pm 1.96 \times sd_{month}}{\sqrt{n_{month}}}$, with sd_{month} the standard deviation and n_{month} the number of measures, i.e. the number of days in the month for simulations, the number of captures for data).

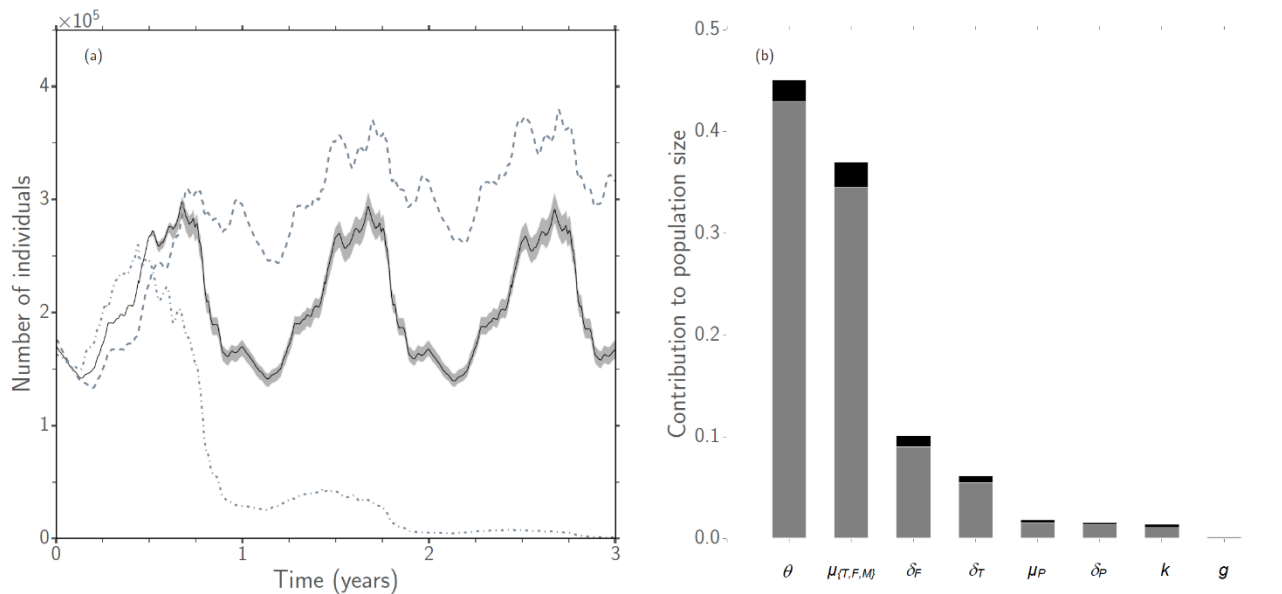


Figure 4. Sensitivity analysis of the model: (a) effect on population size (teneral and adult females) of temperature variations (+5% from reference: dash dot, -5%: dashed) compared to $\pm 5\%$ variations in carrying capacities (grey filling); (b) results of the FAST sensitivity analysis with contribution to population size variance of model parameters (θ : temperature, $\mu_{(T,F,M)}$: adult mortality, δ_X : development of stage X (with X in {F: adult females, T: teneral, P: pupae}), k : carrying capacities, g : the shape parameter in the diffusion process; sensitivity indices for principal effect in grey and for first order interactions in black). All parameters were varied by $\pm 5\%$ from their reference value except temperature varying by $\pm 0.3^\circ\text{C}$.

261 **Temperature and mortality as key factors driving population size**

262 Model predictions other than age structure (Fig. S5) were highly sensitive to temperature (T) and
263 adult mortality ($\mu_{T,F,M}$) variations, and moderately to teneral (δ_T) and parous (δ_F) female
264 development variations (Table S4), while parameters related to pupae (μ_P , δ_P), carrying capacities
265 (k), and dispersal (g) did not contribute to output variance (Fig. 4, Fig. S6). A 5% variation in
266 temperature lead to demographic explosion or extinction, substantially outweighing the effect of a
267 similar variation in carrying capacities (Fig. 4a), reinforcing the need for considering reasonable
268 temperature variations. Temperature and adult mortality explained 78% of population size variance
269 (Fig. 4b). Development of teneral and parous females added up to another 14.5% of explained
270 variance. Unexpectedly, interactions between parameters were not important.

271 **Efficiency of control measures driven by environmental heterogeneity**

272 Increasing adult mortality at levels comparable to what can be obtained during control programs
273 (Hargrove 2003) induced a quick population decline (Fig. 5). A 50% augmentation (i.e. a parous
274 female daily mortality rate of 2.94 day^{-1} and a life expectancy of 51.5 days at 24°C) resulted in a
275 90% decrease in the female population ($T+F_{1;4+}$) in one year (Fig. 5a).
276 Once reaching low local densities, new patterns emerged related to cell-specific properties. On the
277 one hand, as expected, the spatial distribution of individuals was clearly linked to carrying
278 capacities (Fig. 1 vs. Fig. 5b1-3). The greater the adult mortality, the more uneven was the spatial
279 distribution with a progressive concentration of individuals in cells of highest carrying capacities.
280 On the other hand, much more surprisingly, the increase in adult mortality had a heterogeneous
281 impact at the cell level: the local population decrease varied spatially (Fig. 5c2-3) despite a spatially
282 homogeneous increase in mortality, spatial heterogeneity increasing with the level of induced
283 mortality (Fig. 5c2 vs. 5c3).

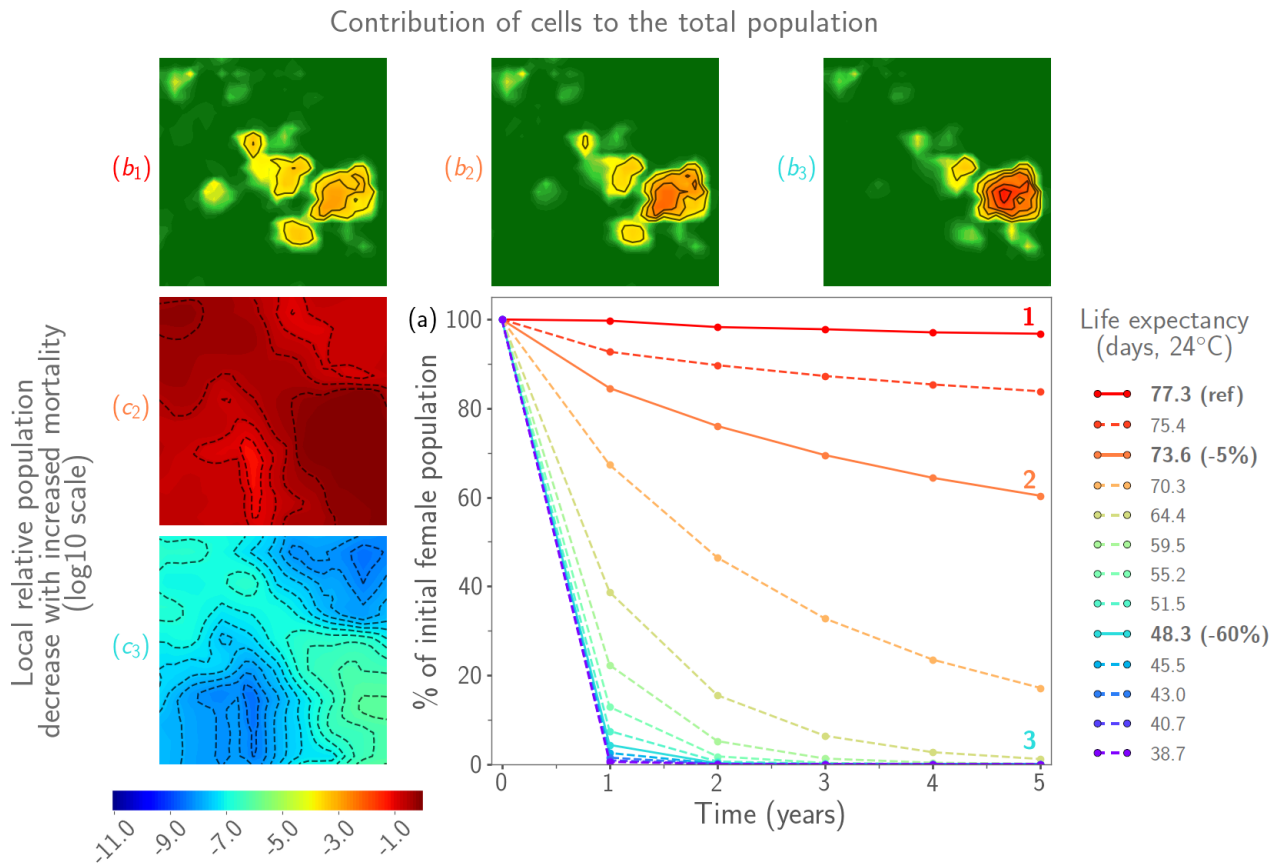


Figure 5. Impact of increasing adult mortality on population size in time and space. (a) relative decrease in female population size compared to the reference scenario while decreasing life expectancy at 24°C (corresponds to increases in adult mortality). Thick lines correspond to (1): no change, (2): +5% of adult mortality, (3): +60%. Spatial patterns during the last time step was assessed for these three scenarios (X from 1 to 3): (b_x) cells contributing the most to female population over the study area in scenario X $\left(\frac{(T_{t_{max},c+F_{1:4},t_{max},c})_{scenarioX}}{(\sum_c T_{t_{max},c+F_{1:4},t_{max},c})_{scenarioX}}\right)$ are in red; (c_x) cells with the highest local impact of increased mortality in scenario x compared to the reference scenario $\left(\frac{(T_{t_{max},c+F_{1:4},t_{max},c})_{scenarioX}}{(T_{t_{max},c+F_{1:4},t_{max},c})_{reference}}\right)$ are in blue (log₁₀ scale).

284 To better understand this latter pattern, three local cell factors related to environmental
 285 heterogeneity were examined: carrying capacity, mean annual temperature, and standard deviation
 286 of annual temperature (Fig. 6). While the carrying capacity had no influence here (Fig. 6a), the local
 287 temperature largely contributed to explain the pattern (Fig. 6b-d). Both a decrease in the mean and
 288 standard deviation of the local annual temperature were associated with a decrease in the local
 289 impact of increasing adult mortality, despite the narrow ranges of variation in the mean (23.7°C to
 290 24.3°C) and standard deviation (1.98°C to 2.37°C). There was no correlation between these two
 291 temperature statistics (Fig. 6c-d).

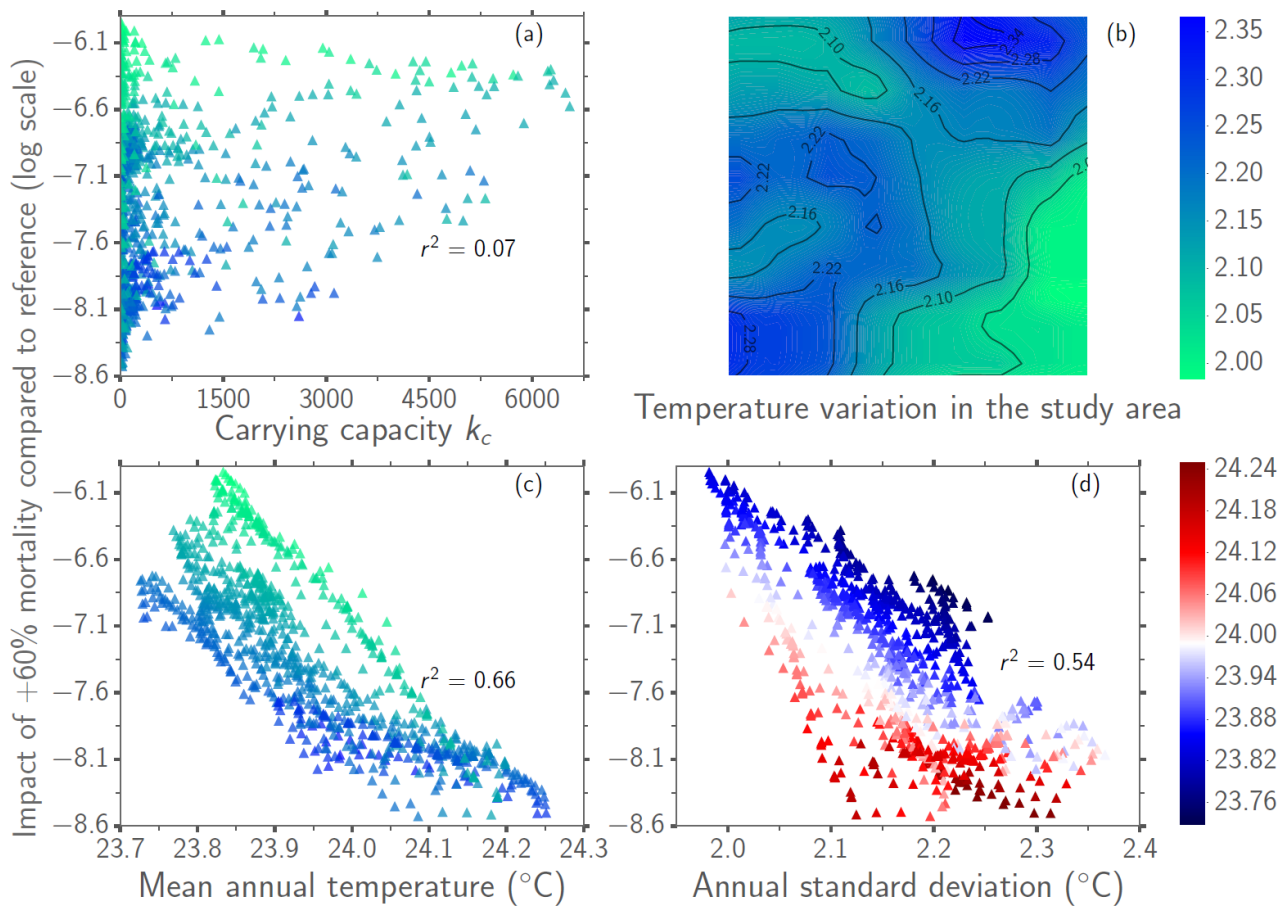


Figure 6. Correlation between environmental variables and the local impact of increasing mortality by 60% (measured as $\frac{(T_{t_{max,c}+F_{1:4+,t_{max,c}})_{scenario3}}}{(T_{t_{max,c}+F_{1:4+,t_{max,c}})_{reference}}$): (a) local carrying capacity k_c ; (b) spatial representation of the annual standard deviation of local temperatures; (c) annual mean temperature; (d) annual standard deviation of temperature. In (a, c, d), the local impact of mortality is on the y-axis (log10 scale), the higher being the value, the smaller the impact. Each point corresponds to a cell of the simulated grid, point colour denoting in (a-b) for the annual standard deviation of local temperature (same colour bar as in (b)), while in (d) to the mean annual temperature of cells (specific colour bar). Correlation coefficient r^2 between axes is shown for (a, c, d).

292 4. Discussion

293 Environmental heterogeneity with respect to carrying capacity and temperature not only drives the
 294 temporal population dynamics of *G. p. gambiensis* at large scale, but also the spatial distribution of
 295 individuals and unexpectedly renders heterogeneous the impact of a homogeneous increase in adult
 296 mortality. Such a heterogeneous impact can be compensated during eradication campaigns by
 297 homogeneous induced sterility when sterile males are released by air and aggregate in the same sites
 298 than wild males (Vreysen et al. 2011), thus warranting a homogeneous sterile to wild ratio, as was
 299 observed in the eradication campaign against *Glossina austeni* on Unguja Island of Zanzibar. We

300 argue that control strategies should account for environmental heterogeneity to increase the chances
301 of success, with emphasis on local areas of high suitability characterized by a high carrying capacity
302 and on local refuges characterized by a cold local temperature within the relevant range for tsetse
303 (23.7-24.0°C here) and a low local variability of temperature over the year (irrespective of carrying
304 capacity). Refuges, highlighted in our study area despite a small surface suitable for tsetse, could
305 jeopardize control efforts by providing areas from which recolonization may occur after control has
306 stopped.

307 In addition, temperature effect on population dynamics both at large and small local scales reinforces
308 the need for investigating further the impact climate change could have on tsetse populations
309 (Terblanche et al. 2008; Moore et al. 2012). It is unlikely that tsetse flies will cross the Sahara, but
310 they could migrate to higher altitudes and invade trypanosoma-free zones, particularly in Eastern and
311 Southern Africa where tsetse distribution is mainly governed by altitude (Solano et al. 2010a). Such
312 population shifts will impact the density of cattle in either direction, which may in turn impact the
313 distribution of wild fauna including lions (Carter et al. 2018). Populations previously isolated from
314 one another could also merge, making developed and adopted control strategies challenging, and
315 conversely, new isolated populations could appear, all the more as temperature is the first driver of
316 landscape friction in tsetse (Bouyer et al. 2015).

317 The mechanistic spatio-temporal model developed to predict *G. p. gambiensis* population dynamics
318 and how these evolve when adult mortality is increased is original compared to already published
319 models. First, it incorporated environmental heterogeneity through a data-driven approach, both
320 accounting for variable temperatures and carrying capacities in space and time. Using realistic
321 patterns instead of theoretical ones (Childs 2011), knowledge-driven ones (Barclay & Vreysen 2013),
322 or aggregated ones assuming a binary occupancy (Lin et al. 2015) evidenced unexpected refuges. The
323 proposed model can be applied to other areas with available data and a known metapopulation
324 structure. Second, new field and laboratory data on mortality, development, and dispersal have been
325 incorporated into the model. Predicted age structure was in very good agreement with field data, and

326 was robust in our simulations, barely impacted by parameter variations. Amplitude and duration of
327 seasons are expected to be major drivers of parity distribution, which could not be assessed here as
328 temperature data were available for only a year. Our results highlight the need for more biological
329 studies to better infer mortality variations with temperature, as well as the crucial need for new
330 methods to thoroughly estimate temperatures as perceived by individuals. Such a complementarity
331 interplay between models, field observations, and laboratory experiments is fundamental to achieve
332 trustworthy predictions.

333 The fact that mortality has a stronger influence on population dynamics than reproduction is
334 consistent with tsetse flies being specialists with a narrow niche. They are willing to avoid mortality
335 at all costs (Pagabeleguem et al. 2016), where other species compensate for losses by boosting birth
336 rates (Southwood et al. 1974). *Glossina* spp. have evolved towards an optimal utilization of energy
337 and resources (Cody 1966), which makes them highly adapted to their ecological niche. Therefore,
338 they are less likely to leave their habitat and expose themselves to other environments, which keeps
339 the population at or near carrying capacity (Southwood et al. 1974).

340 Efficient control methods can only be designed by considering a species ecological strategy
341 (Southwood et al. 1974; Conway 1977). Fast action methods such as chemicals are better suited for
342 species showing high reproductive rates, short generation times, along with broad food preferences
343 and good dispersing abilities (Altieri et al. 1983). In contrast, pests reproducing at lower rates and
344 having longer generation time but good competitive abilities would be more efficiently restrained
345 with cultural control (e.g. insect pests), host resistance, and sterilization (Altieri et al. 1983).
346 Nonetheless, such quite extreme characteristics should be considered in conjunction with species
347 relationships within communities (Ehler & Miller 1978; Altieri et al. 1983).

348 Traps, targets, and insecticide-treated livestock are control tactics that increase adult mortality, which
349 can drastically reduce tsetse populations (Kagbadouno et al. 2011; Dicko et al. 2014; Percoma et al.
350 2018). However, our results indicate also generation time as a contributing factor to population size
351 variations. Such a factor can be indirectly modified using the sterile insect technique, which impair

352 reproduction (Dyck et al. 2005). Obtaining very low tsetse densities is not enough to reach eradication
353 as was demonstrated recently by the failures of three eradication programs against *G. p. gambiensis*
354 in north-western Ghana (Adam et al. 2013), Loos islands in Guinea (Kagbadouno et al. 2011), and
355 the Mouhoun river in Burkina Faso (Percoma et al. 2018). In addition, in view of unexpected local
356 refuges where increasing adult mortality is not as effective as in other areas, it becomes necessary to
357 further assess the effect of combined and spatially targeted control measures to achieve eradication.
358 Our model provides a relevant tool to evaluate such complex control strategies as it originally
359 accounts simultaneously for density-dependent processes, spatial and temporal environmental
360 heterogeneity, and all stages of tsetse lifecycle possibly targeted by control measures. Our framework
361 could also be useful to identify where to focus stakeholders' efforts to minimize impact of other
362 specialist pests, such as the codling moth (*Cydia pomonella*) affecting apple and pear trees, and the
363 sheep ked (*Melophagus ovinus*). Nevertheless, the importance of stochastic events when populations
364 become very small must not be overlooked and these effects should be included in future
365 developments. Our approach gives clues on how to trigger a drastic decline of the population.
366 However, to predict the subsequent population dynamics at low densities and assess final steps of
367 eradication strategies, a deterministic framework becomes irrelevant as it does not enable quantifying
368 the probability of population extinction at local and large scales.

369 Accounting for spatial heterogeneity is essential to better understand and predict tsetse population
370 dynamics, as habitat fragmentation holds the key to population survival when conditions are globally
371 hostile. However, parameters driving tsetse fly dispersal abilities did not structure their final
372 distribution. Landscape ecology must be studied to reveal preferential target zones and identify
373 patches that will need longitudinal surveillance. Optimal management strategies are therefore valid
374 for a given species in a given habitat and should not be generalized without baseline data collection
375 to characterize the ecosystem.

376 To conclude, environmental carrying capacity largely explains the contribution of local source spots
377 to tsetse population dynamics at a large scale, but unfavourable conditions progressively lead such

378 spots to disappear, refuges then being localized in zones with colder and less variable temperature
379 where population decrease due to increasing adult mortality is reduced. Targeted areas for control
380 should be chosen with caution when facing such a heterogeneous habitat.

381 **Authors' contribution**

382 JB and PE designed the study and advised biological details. HC, SA, SPi and PE developed the
383 model. HC conducted the analyses and prepared the figures. HC, SA, SPi, JB and PE discussed the
384 results. HC and PE wrote the manuscript. AD provided model external input data readily usable by
385 the mechanistic model. JB, MTS, BS, MB, MV, SPa, AB collected the data. All authors edited the
386 manuscript.

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395 REVOLINC).

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