Climate Effects on Prey Vulnerability Modify Expectations of Predator Responses to Short- and Long-Term Climate Fluctuations

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Climate changes affect the distribution and abundance of organisms, often via changes in species interactions. Most animals experience predation, and a number of models have investigated how climate fluctuations can influence predator–prey dynamics by affecting prey abundance through changes in resource availability. However, field studies have shown that prey vulnerability is a key feature determining the outcome of predator–prey interactions, which also varies with climatic conditions, via changes in prey body condition or in habitat characteristics (e.g. vegetation cover). In this theoretical work, we explore, with large mammals of African savannas in mind, how the interplay between climate-induced changes in prey abundance and climate-induced changes in prey vulnerability affects the immediate and long-term responses of predator populations. We account for prey body condition and habitat effects on prey vulnerability to predation. We show that predictions on how predator abundance responds to climate fluctuations differ depending on how climate influences prey vulnerability (habitat characteristics vs. prey body condition). We discuss how species traits influence the relative importance of the different sources of vulnerability. For example, our results suggest that populations of cursorial predators (such as spotted hyaenas) are expected to fare better than populations of ambush predators (such as African lions) in African ecosystems that will be characterised by an aridification. This study highlights the importance of understanding, and accounting for, the vulnerability factors associated to a given predator–prey pair, and improves our comprehension of predator–prey relationships in a changing climate.

Keywords: population dynamics, traits, vulnerability, climate changes, predator–prey interactions

INTRODUCTION

The Earth’s climate is changing rapidly, largely because of human activities (IPCC, 2014b). Temperature is rising globally and will continue to do so, precipitation regimes are, or will be, locally altered, and extreme climatic events will become more common (IPCC, 2014b). Climate changes already have ecological impacts and are, or will soon be, a major driver of species dynamics...
and survival (Maclean and Wilson, 2011; Pacifi et al., 2017). So far, most studies on the impacts of climate changes have focused on describing changes in population distribution and abundance (e.g. Parmesan et al., 1999; Martay et al., 2017), phenology and demography (e.g. Inouye et al., 2000; Moyes et al., 2011) or behaviour (Candolin and Wong, 2012). Even though these changes can sometimes emerge from the direct effect of climate on individuals, they often result from cascading effects occurring through biotic interactions, such as trophic interactions (e.g. Visser et al., 2006). Therefore, understanding how climate changes will affect species through their biotic interactions is fundamental to grasp the full picture of the impacts of climate changes.

Climate changes have the potential to modify predator–prey interactions, which are central in the functioning of populations, communities and ultimately ecosystems (Bastille-Rousseau et al., 2018). First, by altering primary production quality and quantity, climatic conditions influence the body condition of large mammalian herbivores (rainfall: Bourgarel et al., 2002; Owen-Smith, 2002; NAO: Mysterud et al., 2001; snow: Saether and Gravel, 1988). Herbivore body condition, in turn, influences demographic parameters, such as survival (Bender et al., 2007) or fecundity (Cook et al., 2004), which ultimately affect herbivore population abundance. As a result, climate indirectly influences prey abundance through density-dependent processes (Saether, 1997; Forchhammer et al., 1998). For example, several studies linked rainfall to herbivore abundance in savanna ecosystems (East, 1984; Ogutu et al., 2008), or showed the influence of climatic variations on the dynamics of several populations of Northern ungulates (Forchhammer et al., 1998; Post and Stenseth, 1999). Extreme climatic conditions also affect herbivore demographic parameters and ultimately population abundances (e.g. drought: Mduma et al., 1999; Foley et al., 2008). These changes in herbivore abundance originating from climate fluctuations will affect predator populations, as the maintenance of predator populations largely depends on prey abundance (East, 1984; Carbone and Gittleman, 2002). Prey body condition is thus a first pathway through which climate changes may alter predator–prey interactions and such climate-driven impact on prey demography has already been taken into account in pioneering predator–prey modelling works (e.g. Wilmers et al., 2007a,b; blue arrows in Figure 1).

Predator–prey relationships are also influenced by prey vulnerability, which may arise from intrinsic (inherent to the prey) or extrinsic (arising from the environment) sources. Prey body condition is typically an intrinsic factor that can determine prey vulnerability to predators. Indeed, several studies pointed out that predators tend to select the weakest individuals among prey and that prey in poorer body condition are more likely to be preyed upon (in savanna ecosystems: FitzGibbon and Fanshawe, 1989; Pole et al., 2004; in temperate ecosystems: Husseman et al., 2003; Atwood et al., 2007). On the other hand, habitat characteristics are extrinsic factors that can modulate prey vulnerability to their predators and that can be influenced by climatic conditions. Several studies stressed out the importance of vegetation structure and cover, which interact with the predator hunting technique and the prey escape technique to ultimately affect the hunting outcome. This was shown in savanna (Funston et al., 2001; Hopcraft et al., 2005) and temperate ecosystems (Husseman et al., 2003; Lone et al., 2014). Other habitat characteristics can interfere with the predator hunting success. For example, prey were either more vulnerable to predators with increasing amount of snow (Mech et al., 1971; Post et al., 1999) or less vulnerable to predators as snow hardness decreased (Stenseth et al., 2004).

Hence, the true prey availability to predators results from the interplay between prey abundance and prey vulnerability, which is characterised by different sources of vulnerability (red arrows in Figure 1). Little is known on the relative importance of climate-induced changes on the different components of prey availability to predators, and particularly on the different sources of prey vulnerability. Here, we integrate both prey abundance and vulnerability effects (Figure 1) in a theoretical study investigating the immediate response of predator populations to a year of a specific climatic condition, as well as the long-term response of predator populations to changes in average climatic conditions. This study aims at providing general insights on the interplay between climate-induced changes in prey abundance and prey vulnerability. The model is, on purpose, not designed for a specific predator–prey system, so that people studying predator–prey interactions worldwide can adapt the model to fit the system they are interested in. However, we designed our study with large mammalian herbivores and carnivores of African savannas in mind so that the parameters and the functional relationships used are easy to relate to a real world. Annual rainfall is expected to decrease in southern Africa and increase in eastern Africa and generally to become more variable (IPCC, 2014a). We therefore specifically focussed on the influence of rainfall conditions on predator–prey interactions in African savanna ecosystems. We modelled the population dynamics of a predator preying on a density-dependent age-structured prey population following the model described in Wilmers et al. (2007a,b), but we further and originally accounted for prey body condition and environmental effects on prey vulnerability to predation (Figure 1).

**MATERIALS AND METHODS**

**The Predator–Prey Model**

The Prey Population Model Without Predation

We considered a female-only prey population (Caswell, 2001) which, in any year \( t \), had a total population size of \( N(t) \) made of \( J(t) \) juveniles and \( A(t) \) adults. Fluctuations from year to year were driven by survival and reproduction of individuals, which are now described.

Body condition is a critical individual characteristic, as it links environmental conditions to demography. For simplicity, we assumed a linear relationship between body condition and survival, and thus only modelled the latter. We modelled the effect of body condition on reproduction using a relationship between the probability to reproduce and the probability of survival. Each year, individual survival probabilities \( p_s(t) \) and \( p_a(t) \), for juveniles and adults, respectively, were drawn from
Beta distributions with shape parameters $\alpha_j$ and $\beta_j$ for juvenile prey, and $\alpha_a$ and $\beta_a$ for adult prey, respectively, and differed from one individual to another, allowing to integrate individual heterogeneity. However, new probabilities were drawn each year and were not preserved through the individual's life, and thus varied randomly within individuals from year to year. The method-of-moments gives:

$$\alpha_j = p_j \left( \frac{p_j (1 - p_j)}{Sd_{pj}^2} - 1 \right)$$  \hspace{1cm} (1a)$$

$$\beta_j = (1 - p_j) \left( \frac{p_j (1 - p_j)}{Sd_{pj}^2} - 1 \right)$$  \hspace{1cm} (1b)$$

for juvenile prey, and

$$\alpha_a = p_a \left( \frac{p_a (1 - p_a)}{Sd_{pa}^2} - 1 \right)$$  \hspace{1cm} (1c)$$

$$\beta_a = (1 - p_a) \left( \frac{p_a (1 - p_a)}{Sd_{pa}^2} - 1 \right)$$  \hspace{1cm} (1d)$$

for adult prey, where $p_j$ and $p_a$ are the mean survival rates and $Sd_{pj}$ and $Sd_{pa}$ are the standard deviations for juvenile and adult prey population, respectively. In any year $t$, mean survival rates $p_j$ and $p_a$ were:

$$p_j (t) = P_j \times g_j (t)$$  \hspace{1cm} (2a)$$

$$p_a (t) = P_a \times g_a (t)$$  \hspace{1cm} (2b)$$

where $P_j$ and $P_a$ are parameters defining the maximum survival rates for juveniles and adults, respectively (see Table 1 for all default values of model parameters), and $g_j$ and $g_a$ are scaling functions allowing to account for density- and resource-dependence in survival. These scaling functions (bounded between 0 and 1) are those used by Wilmers et al. (2007a) and follow:

$$g(N(t)) = \frac{\gamma^\delta}{\gamma^\delta + \frac{N(t)}{R}}$$  \hspace{1cm} (3)$$

where $R$ is the annual rainfall of the site. $R$ varied from 300 to 900 mm by 100 mm increment, which are coherent values for annual rainfall in African savannas (Sankaran et al., 2005; Chamaillé-Jammes and Fritz, 2009). We assumed that rainfall determined directly and linearly prey resource availability (Rutherford, 1980). We thus considered rainfall as a proxy of the resource availability and the ratio $N(t)/R$ as a number of individuals sharing the amount of available resource for year $t$. The half-saturation parameter $\gamma$ determined the per capita resource availability at which the maximum survival rate was reduced by half. Finally, $\delta$ was a shape parameter setting the beginning and the strength of the density-dependence (as in Wilmers et al., 2007a).

Juvenile survival is expected to be the first demographic parameter affected by increasing density, followed by age at first reproduction, reproductive success and finally adult survival (Eberhardt, 1977). This hypothesis has found support in large
TABLE 1 | Definitions and values of the state variables and parameters for prey and predators used in the model.

| Variables | Definition/description | Values |
|-----------|------------------------|--------|
| \( N(t) \) | Prey population size at the beginning of year \( t \) | \( N_1 = 10,000 \) |
| \( J(t) \) | Number of juvenile prey at the beginning of year \( t \) | \( J_1 = 2500 \) |
| \( A(t) \) | Number of adult prey at the beginning of year \( t \) | \( A_1 = 7500 \) |
| \( Y(t) \) | Predator population size at the beginning of year \( t \) | \( Y_1 = 10 \) |
| \( P(t) \) | Pool of prey vulnerable to predation for year \( t \) | Calculated at each time step |
| \( M(t) \) | Pool of prey consumed at the end of year \( t \) | Calculated at each time step |
| \( \alpha_s \) | First shape parameter of the Beta distribution from which are drawn adult prey individual survival rate | Calculated at each time step |
| \( \alpha_j \) | First shape parameter for Beta distribution from which are drawn juvenile prey individual survival rate | Calculated at each time step |
| \( \beta_a \) | Second shape parameter for Beta distribution from which are drawn adult prey individual survival rate | Calculated at each time step |
| \( \beta_j \) | Second shape parameter for Beta distribution from which are drawn juvenile prey individual survival rate | Calculated at each time step |
| \( P_a \) | Maximum adult prey survival probability | 0.95 |
| \( P_j \) | Maximum juvenile prey survival probability | 0.8 |
| \( \hat{S}_{da} \) | Standard deviation, variation of adult prey survival | 0.05 |
| \( \hat{S}_{dj} \) | Standard deviation, variation of juvenile prey survival | 0.05 |
| \( g_a \) | Effect of increased population density on adult survival | Calculated at each time step |
| \( g_j \) | Effect of increased population density on juvenile survival | Calculated at each time step |
| \( \rho_a \) | Mean adult prey survival probability | 0.3 |
| \( \rho_j \) | Mean juvenile prey survival probability | \( \rho_j = P_j \times g_j \) |
| \( \gamma_{ja} \) | Adult half-saturation, point at which adult prey survival is lowered by half | 30 |
| \( \gamma_{ja} \) | Juvenile half-saturation, point at which juvenile prey survival is lowered by half | 20 |
| \( \delta_{da} \) | Shape parameter, defines the onset of density-dependence for adult prey | 2 |
| \( \delta_{dj} \) | Shape parameter, defines the onset of density-dependence for juvenile prey | 2 |
| \( R \) | Annual level of rainfall | \( R \) varies between 300 and 900 mm, by 100 mm increment |
| \( F_{\text{est}} \) | Value determining fecundity value when adult survival is close to 0 | 100 |
| \( F_{\text{est}} \) | Rate at which fecundity value reaches 1 | 15 |
| \( T_h \) | Prey threshold of survival probability below which predators can catch and consume the prey | \( T_h = 1 \) for scenarios 1 and 3; \( T_h = 0.5 \) for scenarios 2 and 4 |
| \( H_{\text{min}} \) | Lowest value that can be taken by the handling time across all scenarios | 5 |
| \( H_{\text{max}} \) | Highest value that can be taken by the handling time across all scenarios | 8 |
| \( B \) | Maximum rate of change of the handling time | 0.015 |
| \( V \) | Rainfall value for which the rate of change of the handling time is maximum (or inflexion point) | 600 |
| \( \mu \) | Encounter rate in the predator functional response | 0.0002 |
| \( h \) | Predator handling time for one prey in the predator functional response | \( h = 5 \), for scenarios 1 and 2; \( h \) defined by \( H = h_{\text{min}} + \frac{H_{\text{max}} - h_{\text{min}}}{R_{\text{max}} - R_{\text{min}}} \), for scenarios 3 and 4 |
| \( \tau \) | Interference between predators in the predator functional response | 0.3 |
| \( \lambda_{\text{max}} \) | Predator maximum growth rate | 1.8 |

For herbivore demographic parameters, we chose values that were consistent with values reported in the literature on African large herbivores (e.g., Owen-Smith, 1990; Grange et al., 2015). For large carnivore parameters, we chose values that were either consistent with the published literature (e.g., Schaller, 1972), or consistent with our field knowledge of lion and hyaena ecology, or, for parameters that have never been estimated, we adjusted them to allow the model to run and give realistic outcomes.

ungulate studies (Bonenfant et al., 2009). As our model was conceptualised for large mammals (as for the one of Wilmers et al., 2007a), we assumed a greater sensitivity of juvenile survival to increasing density by assuming that \( \gamma_j \) was lower than \( \gamma_a \).

In summary, both rainfall \( R \) and population size \( N \) determined \( g \) and ultimately the average survival rates \( p_j \) and \( p_a \) in the population (Figure 2A). Once mean survival probability had been computed, we determined the fate of each individual by randomly drawing a value from a uniform distribution on the [0–1] interval and comparing it to the individual’s survival probability. If the value was lower than the survival probability, the individual survived; if not, it died.

We assumed that juveniles became adults in 1 year (a realistic assumption for some herbivore species; Estes, 1991) and only adults were able to reproduce. Therefore, the number of juveniles at the beginning of a given year depended only on the fecundity...
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FIGURE 2 | Summary of the functional relationships underlying the model. (A) Effect of increased number of individuals per resource unit (approximated by rainfall) on mean juvenile and adult prey survival rates adapted from Wilmers et al. (2007a). Mean adult and juvenile prey survival rates has been represented for two values of δ parameter, which controls the onset and the strength of the density dependence. Juvenile prey survival rate has been set to be the first impacted by increasing density. (B) Probability of reproduction as a function of survival probability \( p_{ai}(t) \). (C) Functional relationship between rainfall and predator handling time \( h \) as defined by Eq. 5 for parameters given in Table 1. The hunting success varies inversely with the handling time.

of adults the year before. We also considered that, for each adult, the probability of reproducing \( (P_f) \) was a function of its survival probability \( p_{ai}(t) \), and the link between \( P_f \) and \( p_{ai}(t) \) was as follows:

\[
P_f \left( p_{ai}(t) \right) = \frac{1}{1 + f_{low} \times e^{-f_{growth} \times p_{ai}(t)}}
\]

(4)

where \( f_{low} \) is a parameter determining the probability of reproduction \( P_f \) of individuals whose survival probability is close to 0 and \( f_{growth} \) a parameter defining the speed at which \( P_f \) reaches the value of 1 with increasing \( p_{ai} \) (Figure 2B). We compared, for each individual, values drawn randomly from a uniform distribution on the \([0,1] \) interval with their reproduction probability. If the value was lower than the reproduction probability, the individual produced one offspring; if not, it did not reproduce.

Integrating Predation Into the Prey Model
We modelled an unstructured predator population (similarly to Wilmers et al., 2007b), which, in any year \( t \), had a total population size of \( Y(t) \). Predator population dynamics resulted from the combination of a functional response, which describes how the intake rate of a predator varies with prey abundance (Holling, 1959; Abrams and Ginzburg, 2000), and a numerical response, which relates predator intake to changes in its abundance (Bayliss and Choquenot, 2002).

We integrated the influence of prey body condition by accounting for the fact that only adult prey in poor condition, i.e. with a low survival probability, were vulnerable to predators, whereas all juvenile prey were vulnerable to predators,
irrespectively of their body condition. To do so, we defined a threshold of survival probability $T_h$. Any adult prey with an individual survival probability $p_{a\varepsilon}(t)$ below this threshold was integrated to the pool of adult prey vulnerable to predators $A_{vuln}$. Rainfall and prey density both affected $g_p(t)$ and thus $p_{a\varepsilon}(t)$, the mean survival probability within the population. Consequently, they played a crucial role in determining the proportion of adult prey in poor body condition. As $p_{a\varepsilon}(t)$ decreased, there were more adult prey with low individual survival probabilities, and therefore more adult prey with a survival probability under the vulnerability threshold and exposed to predation.

We integrated the influence of habitat characteristics that could affect prey vulnerability through the manipulation of the handling time $h$, which is classically defined as the time needed by a predator to subdue and consume a prey and during which the predator cannot capture another prey (Abrams and Ginzburg, 2000). In our model, this time to subdue one prey corresponded to all predation attempts (with one predation attempt being a predation sequence: search, encounter and attack) leading to a successful prey capture. For African savannas, we assumed that an increase in annual rainfall leads to a greater vegetation cover (taller grass, more leaves in bushes and trees) and hence better concealment opportunities for predators to approach their prey undetected (Hopcraft et al., 2005). Hence, we considered that an increase in the climatic variable $R$ would increase prey vulnerability and, consequently, decrease the number of attempts leading to a successful attack, i.e. lead to lower handling time $h$ values (an alternate situation where handling time increases with rainfall is presented in Appendix A of the Supplementary Material). Therefore, $h$ varied with rainfall according to the following equation:

$$H(R) = h_{\text{min}} + \frac{h_{\text{max}} - h_{\text{min}}}{1 + e^{B(R - V)}}$$

(5)

with $H(R)$ the value of handling time for a given year $t$ and the rainfall value $R$, $h_{\text{min}}$ is the lowest value of $h$, $h_{\text{max}}$ is the maximum value of $h$, $B$ is the maximum rate of change and $V$ is the rainfall value at which the rate of change of $h$ is maximum (or inflexion point). This equation allowed the handling time to vary non-linearly and negatively with rainfall (Figure 2C). One value of $h$ was calculated for each rainfall value and was then incorporated into the functional response $I(t)$.

For the functional response, we assumed that the number of prey killed and consumed by one predator during a given year $t$ followed a type II Beddington–DeAngelis functional response. This equation allows the integration of interference between predators when searching for and capturing prey and is formulated as follows:

$$I(P(t), Y(t)) = \frac{\mu P(t)}{1 + \tau Y(t) + \mu hP(t)}$$

(6)

where $P(t)$ is the number of vulnerable prey, $Y(t)$ the number of predators, $\mu$ a parameter defining the encounter rate between predators and their prey, $h$ the parameter defining the handling time and $\tau$ the interference between predators (Abrams and Ginzburg, 2000). Finally, this type of functional response allows both prey and predator densities to impact predators’ intake and therefore the regulation of predator populations.

Predation occurs all year around. Consequently, to account for the depletion of prey that occurred throughout the year and could affect the estimation of the number of prey actually captured, we recursively applied the functional response presented in Eq. 6 over daily time steps, adjusting prey number at each time step. We named $M(t)$ the total number of prey killed by predators and calculated it as follows:

$$M(t) = P(t) - \sum_{d=1}^{365} I(P(d), Y(t))$$

(7)

where $P(t)$ is the number of vulnerable prey to predation, $P(d)$ the pool of prey vulnerable to predation at the beginning of day $d$ and $Y(t)$ the number of predators at the beginning of year $t$.

The numbers of juvenile and adult prey killed by a given predator during a year are given by:

$$M_j(t) = M(t) \times \frac{I(t)}{P(t)}$$

(8)

$$M_a(t) = M(t) - M_j(t)$$

(9)

The $M_j(t)$ juveniles killed in year $t$ were randomly removed from the $J(t)$ juveniles present at the beginning of the year $t$. Similarly, the $M_a(t)$ adults killed in year $t$ were randomly removed from the pool of vulnerable adults $A_{vuln}$, i.e. individuals whose survival probabilities were below the threshold of survival probability $T_h$. For simplification, we assumed that adult and juvenile prey were equally profitable to predators. Similar assumption can be found in other modelling studies (e.g. Wilmers et al., 2007b).

Finally, the predator numerical response was given by:

$$Y(t + 1) = \lambda_{\text{max}} \times P_{\text{rel}}(t) \times Y(t)$$

(10)

where $Y(t)$ is the number of predators present in year $t$, $\lambda_{\text{max}}$ is the maximum growth rate of the predator population and $P_{\text{rel}}(t)$ was calculated as follows:

$$P_{\text{rel}}(M(t), Y(t)) = \frac{M(t)}{\frac{1}{h_{\text{min}}} \times 365 \times Y(t)}$$

(11)

with $h_{\text{min}}$ the smallest value possible for the handling time. $P_{\text{rel}}$ was therefore bounded between 0 and 1 and quantified predators hunting efficiency over a year by expressing the amount of prey actually eaten in year $t$ as a proportion of the maximum amount of prey that predators could have eaten.

Simulations

To study how climatic conditions may impact predator populations through their influence on prey body condition or habitat characteristics, we designed four scenarios:

Scenario 1: Climatic conditions influence prey availability through their effect on prey body condition and ultimately on prey abundance. Prey vulnerability to predators did not vary with prey body condition (all prey were vulnerable to predators) nor with habitat characteristics. Climatic conditions therefore
influenced predator–prey interactions only through their impacts on prey survival and fecundity, i.e. through prey abundance only (blue arrows in Figure 1). This demographic effect of climate is present in all scenarios.

Scenario 2: Prey vulnerability was affected by climatic conditions through their influence on prey body condition (red arrows in Figure 1). The poorer a prey body condition, the more vulnerable to predators the prey was. Therefore, climatic conditions influence prey availability through their effect on prey body condition affecting both abundance and prey vulnerability to predation.

Scenario 3: Prey vulnerability was affected by climatic conditions through climate-driven changes in habitat characteristics (e.g. vegetation) (red arrows in Figure 1). Therefore, climatic conditions influence prey availability through their effect on prey body condition affecting prey abundance and on habitat characteristics affecting prey vulnerability to predation.

Scenario 4: Prey vulnerability was affected by both prey body condition and climate-driven changes in habitat characteristics. In this scenario combining scenarios 2 and 3, climatic conditions influence prey availability through their effect on prey body condition and habitat characteristic and, ultimately, prey abundance and vulnerability to predation.

While in scenarios 1 and 3, predators could capture any prey individual, in scenarios 2 and 4, predators could capture any juvenile, but could only subdue adult prey that were in poor body condition. These differences between scenarios were modelled by setting the value of the threshold of survival probability $Th$ at 1 in scenarios 1 and 3, and at 0.5 in scenarios 2 and 4 (Table 1). As vulnerability to predators was independent from habitat characteristics in scenarios 1 and 2, the handling time was set constant and equal to $h = 5$ in these scenarios. In scenarios 3 and 4, prey vulnerability to predators was dependent on habitat characteristics and handling time $h$ varied with climatic conditions following Eq. 5 (for a summary of the model functioning, see Figure 3).

We focussed on these four scenarios only, as the goal of this study was to assess how the addition of prey vulnerability to traditional predator–prey models that considered prey abundance only would change the expectations in terms of responses of predator populations to climatic changes.

For simulations, parameters were given default values considered realistic for African large mammals. These default values and initial values for state variables are presented in Table 1. The sensitivity of the main results to some of these parameter values was investigated by sensitivity analyses (see section below). As we were mostly interested in how the link between climatic conditions and prey vulnerability could affect predator dynamics, predators were characterised by identical parameter values in the different scenarios, but for the threshold $Th$ and handling time $h$ values. Therefore, in our work, all differences in predator dynamics between scenarios emerged from differences in these two parameters.

**Immediate Response of Predator Populations to Specific Rainfall Conditions**

We assessed, for each scenario, the immediate response of the predator population to specific rainfall conditions. We simulated over 198 years the dynamics of the predator population under the long-term annual rainfall 600 mm, then imposed a predefined rainfall in year 199 (different values were tested: from...
changes in parameter \( h \) of parameter values, we conducted sensitivity analyses for demographic rates. In addition, we also tested for the influence of changes in \( \lambda_{\text{max}} \) values, which is a central parameter of the predator numerical response. Finally, we carried out a sensitivity analysis for the \( \delta \) parameter, which describes the shape of density dependence and determines the magnitude of the prey population response, the number of individuals per resource units and ultimately prey dynamics. We always varied this parameter simultaneously for adult and juvenile, ensuring that juvenile survival never exceeded adult survival (see Figure 2A). Sensitivity analyses were conducted by increasing or decreasing each of these parameters, one at a time, by 20% around its default value. For each parameter value, simulations were replicated 500 times. Results of the sensitivity analyses are presented in Appendix B of the Supplementary Material.

An example of the behaviour of the model for one run showing fluctuations of rainfall, prey and predator population size over 200 years is presented in Supplementary Figure S7 (Appendix C of the Supplementary Material), and a summary of the key steps of the modelling process and of the different scenarios can be found in Figure 3. All simulations were conducted in R 4.0.2 (R Core Team, 2020).

RESULTS

Immediate Response of Predator Populations to Specific Rainfall Conditions

In scenario 1, i.e. when climatic conditions influenced prey abundance only (through survival and fecundity) and not prey vulnerability, the population growth rate of predators was not influenced by the specific climatic condition of the year (Figure 4A). This result was not surprising as according to the model design, the demographic consequences of climate are expected to be visible in prey population the following year \((t + 1)\) and, therefore, would only impact predator growth rate calculated over the year \(t + 1\) and \(t + 2\). In contrast, in the three other scenarios, i.e. when climatic conditions influenced prey vulnerability, either through changes in prey body condition (scenario 2) or through changes in habitat characteristics (scenario 3) or both (scenario 4), the population growth rate of predators 1 year was influenced by the specific climatic condition of the year. Population growth rates in scenarios 2 and 3 showed opposite patterns (Figure 4A). Predators from scenario 2 benefited from below-average rainfall (600 mm) and their populations increased, as shown by the growth rate being greater than 1, but there was a negative effect of above-average rainfall (Figure 4A). In scenario 3, there was a negative effect of below-average rainfall and a positive effect of above-average rainfall on predator growth rate (Figure 4A). Because scenario 4 is a combination of scenarios 2 and 3, the population growth rate of predators from scenario 4 was intermediate between those of scenarios 2 and 3 (Figure 4A).
rainfall generally followed a similar pattern than those observed from scenario 3. However, the difference in the size of predator populations between scenarios 4 and 3 increased with increasing rainfall (Figure 4B). Conversely, the size of predator populations was virtually identical between scenarios 4 and 2 at the wettest end of the rainfall gradient (Figure 4B). In all cases, the size of predator populations in scenario 4 was always equal or lower than in the other scenarios (Figure 4B).
DISCUSSION

Our work shows how predator populations could respond differently to changing climatic conditions depending on how climate influences prey vulnerability (habitat characteristics vs. body condition). For example, in a context of aridification, a predator–prey system where prey vulnerability depends mainly on prey body condition (scenario 2) would be characterised by larger predator populations than a system where prey vulnerability is mainly affected by the habitat (scenario 3) (see section “How Species Traits Influence the Relative Importance of the Different Sources of Vulnerability” for the role of species traits). It thus highlights the importance of having a good understanding of the processes determining vulnerability at play within the predator–prey pair studied, as integrating climate influence on prey vulnerability, in addition to resource-mediated climate effects on prey abundance, leads to different responses of predator populations (on the short- and the long-term) to changes in climatic conditions.

How Species Traits Influence the Relative Importance of the Different Sources of Vulnerability

Ultimately, whether the predator–prey pair studied will respond according to one scenario or another will depend upon the traits of the species involved. Here, we will illustrate the importance of the predator hunting mode and the prey escape tactics, as well as the importance of the relative size of the predator and the prey. Predator hunting mode (cursorial or ambush) and prey escape tactics (fleeing, fighting or freezing) are key behavioural traits involved in predator–prey interactions that can affect how climatic conditions contribute, indirectly via forage resources or vegetation cover, to determine the vulnerability of prey to predation. For example, cursorial predators, such as spotted hyenas Crocuta crocuta (Cooper, 1990) and wild dogs Lycaon pictus (Pole et al., 2004) in African savannas, or wolves in temperate ecosystems (Mech et al., 2015), are active predators, i.e. they can chase down their prey over long distances and are more likely to kill individuals in poor body condition. They therefore exemplify predators from scenario 2 (climatic conditions influence prey vulnerability through changes in prey body condition only), as prey in poorer condition because of a shortage of food will be particularly more vulnerable to predation from these predators (Mills, 1995; Pole et al., 2004; Atwood et al., 2007). Conversely, ambush predators, such as African lions Panthera leo and leopards Panthera pardus in African savannas, or cougars Puma concolor and lynx Lynx lynx in temperate ecosystems, stalk their prey before attacking them by surprise when they come within short distance (Elliott et al., 1977; Hussman et al., 2003). As such, they rely less on prey body condition. However, to approach their prey undetected, ambush predators need concealment. Therefore, their hunting success is prone to be influenced by vegetation (Funston et al., 2001; Lone et al., 2014), and negatively affected by an increased aridity and a subsequent decrease in vegetation density (scenario 3, climatic conditions influence prey vulnerability through changes in habitat characteristics only). Whereas prey escape tactics might not be that crucial for the outcome of an encounter with an ambush predator, a climate-driven decrease in prey body condition will negatively influence prey fleeing capacity leading to an overall disadvantage against cursorial predators (scenario 2).

Predator body size influences the prey body size targetted by the predator (Radloff and Du Toit, 2004; Owen-Smith and Mills, 2008a) and climate-driven changes in prey body condition can interfere with the chance of successfully subduing an adult prey depending on the relative size of the predator and the prey considered (Mech et al., 2001; Metz et al., 2012). For example, for lion–impala interactions, it is very likely that climatic conditions will not affect the likelihood of lions to capture an adult impala Aepyceros melampus, while in the case of lion–buffalo Syncerus caffer interactions, lions are expected to successfully hunt adult buffaloes, which are dangerous prey for them, in dry conditions only (scenario 2, Owen-Smith, 2008). This is also true at the interspecific level: prey species that are usually inaccessible because they are too difficult to be captured under normal climatic conditions may become vulnerable due to unfavourable conditions, allowing predators to kill young individuals of larger species (e.g. with elephants: Salnicki et al., 2001; Loveridge et al., 2006).

Importance of Integrating Climate-Driven Changes in Prey Vulnerability

Our results show how predator population responses to changes in climatic conditions differ when we take into account climate-driven changes in prey vulnerability in addition to climate-driven changes in prey abundance (existing work takes into account climate-driven changes in prey abundance only – Figure 1). For example, while the overall trend depicted in all four scenarios on the long term is that there are more predators when rainfall increases, because of an overall increase in prey population abundance, the originality of our study is to illustrate how prey vulnerability and its drivers influence the link between predator populations and climate.

When changes in climatic conditions influence prey vulnerability through changes in prey body condition (scenario 2), a below average rainfall event leads to poorer herbivore body condition and hence results in an increase in the pool of adult prey vulnerable to predators (Supplementary Figure S8A). Unsurprisingly, this, in turn, favours the immediate growth rate of the predator population. This result is in accordance with field observations, which reported a positive effect of drought conditions over the short term on large carnivore populations (Mills, 1995). These dry-condition increases in prey vulnerability can be assimilated to “resource pulses” for the predators and to be beneficial to them (Holt, 2008; Yang et al., 2008). Contrarily, above average rainfall events negatively affect the population growth rate of predators, as very few prey adults are vulnerable in years characterised by high levels of resources (Supplementary Figure S8A). Over the long term, predator populations benefit from the overall increase in prey populations associated with increasing rainfall (Supplementary Figure S9). However, predators in scenario 2 can only subdue a portion of adult prey and have a smaller pool of available prey than...
predators in scenario 1. They are therefore maintained at smaller population size. In addition, the proportion of vulnerable adult prey decreases with increasing rainfall (Supplementary Figure S8B), which leads to the observed increase in the difference between the size of predator populations in scenarios 1 and 2.

When changes in climatic conditions influence prey vulnerability through changes in habitat characteristics and habitat-driven hunting success increases with rainfall (scenario 3), predator growth rates are negatively impacted by below average rainfall and positively influenced by above average rainfall. This is typically the case in African savannas where grass height is related to rainfall (Rutherford, 1980) and lions are hunters that are more successful in tall grass (Funston et al., 2001). Over the long term, predator populations benefit from the combined linear increase of prey abundance (Supplementary Figure S9) and non-linear increase of hunting success (i.e. decrease of handling time, Figure 2C), leading to an overall non-linear increase as rainfall increases.

Finally, when changes in climatic conditions influence prey vulnerability through changes in both prey body condition and habitat characteristics (scenario 4), the growth rate of predator populations combines the effects of both sources of vulnerability and predators are affected by opposite effects of different sources of vulnerability. For below average rainfall, predators benefit from the positive effect of lower prey body conditions, which buffers the negative effect of a lower hunting success due to habitat characteristics, and inversely for above rainfall level, leading to the observed intermediate pattern of predator growth rate (Figures 2C, 4A and Supplementary Figure S8A). Over the long term, predator populations of scenario 4 follow the same patterns as the populations in scenario 3 with the exception that they can only subdue a proportion of adult prey. They subsequently remain at a smaller population size.

Contrasting Predator Responses to Short- and Long-Term Climate Fluctuations

Our study highlights the importance of the temporal scale considered as the patterns observed differed whether we considered the short- or long-term temporal scale. While a sudden arid event can be beneficial to predator population, increasing aridity over time leads to an overall decrease in predator population size. The explanation of such differences lies in the fact that a predator response over the long term integrates the predator responses over the short term, along with the long-term response of prey, which declines with increasing aridity (Supplementary Figure S9). Therefore, our results illustrate that the long-term response of predator populations to changes in mean annual rainfall cannot be predicted only from the immediate response of predator populations to specific rainfall conditions.

Considerations on the Model

Modelling studies are difficult exercises where one walk on a thin line between complexity/realism and simplification that is required for conceptualisation, i.e. building some theoretical understanding. We have tried to do this successfully but we want to underline here a few simplifications made in this study (and already made in classical models of predator–prey interactions – e.g. Wilmers et al., 2007a,b). It is noteworthy that the code of the model is available1 so that one can parameterise it for a specific system and add the complexity that would be considered critical. First, in our model, juvenile prey contribute as much energy as adult prey. Second, the predator population is not age-structured. Third, the predator–prey system modelled here involves one prey and one predator only, yet multi-species assemblages often characterise both prey and predator communities. Predators often prey on several prey species, with primary and secondary prey, and display prey switching depending on prey availability and catchability (Owen-Smith and Mills, 2008b; Elbroch et al., 2013). The presence of an alternative prey could influence model outcomes for scenarios 2–4: when the vulnerability of the primary prey decreases, if the predator can switch to an alternate prey species available and more vulnerable, the decrease in predator population growth rate or over the long term may be weaker or not observed. In contrast, the presence of competitors preying on the same prey could intensify the decrease in the number of vulnerable prey available to each predator owing to additive effects (Melis et al., 2009) and negatively influence predator population size.

CONCLUSION

Despite its general simplicity, this theoretical study highlights the importance of understanding, and accounting for, the vulnerability factors in predator–prey relationships, and sheds light on the complexity of predicting predator–prey relationships in a changing climate. We believe it is a prospective study that suggests interesting future research directions. Based on our work, we advise that future theoretical and empirical works should integrate climate-driven changes in prey vulnerability if we want to grasp the full picture of the impacts of climate changes on predator–prey interactions.

DATA AVAILABILITY STATEMENT

The code used for the model and the simulations is available from a GitHub repository: https://github.com/aissamorin/climate-prey-vulnerability-and-predators.

AUTHOR CONTRIBUTIONS

MV and SC-J conceived the ideas. AM and SC-J designed the model. All authors analysed the results, wrote the manuscript, and gave approval for publication. All authors contributed to the article and approved the submitted version.

1https://github.com/aissamorin/climate-prey-vulnerability-and-predators
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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fevo.2020.601202/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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