Bioclimatic context of species' populations determines community stability

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Bioclimatic context of species' populations determines community stability

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Abstract

Aim: It is important to understand the factors affecting community stability because ecosystem function is increasingly at risk from biodiversity loss. Here, we evaluate how a key factor, the position of local environmental conditions within the thermal range of the species, influences the stability of butterfly communities at a continental scale.

Location: Spain, UK and Finland.

Time period: 1999–2017.

Major taxa studied: Butterflies.

Methods: We tested the following hypotheses about how species responses to temperature anomalies aggregate to influence stability: Hypothesis 1, species have contrasting responses to local temperature anomalies at opposing edges of their thermal range; hypothesis 2, communities with central thermal range positions have higher community stability; and the impacts of thermal range position on community stability are driven by hypothesis 3, population asynchrony, or hypothesis 4, additive population stability. Data were analysed at 876 sites for 157 species.

Results: We found some support for hypothesis 1, because there were interactions between thermal range and response to temperature anomalies such that species at different range edges could provide weak compensatory dynamics. However, responses were nonlinear, suggesting strong declines with extreme anomalies, particularly at the hot range edge. Hypothesis 2 was supported in part, because community stability increased with central thermal range positions and declined at the edges, after accounting for species richness and community abundance. Thermal range position was weakly correlated with asynchrony (hypothesis 3) and population stability (hypothesis 4), although species richness and population abundance had larger impacts.
1 | INTRODUCTION

The relationships between diversity, stability and ecosystem functioning have long been debated (Elton, 1958; May, 1972; Tilman & Downing, 1994), and they remain in focus owing to recently observed declines (Díaz et al., 2019) and projected (Ceballos et al., 2020) declines in biodiversity. Understanding the responses of species and communities to environmental disturbance is also a priority for conservation, given the projected increases in the magnitude and frequency of extreme climatic events (Donohue et al., 2016; Ummenhofer & Meehl, 2017).

Currently, there are large concerns around insect declines (Cardoso et al., 2020), although debate about their severity and extent is ongoing (Simmons et al., 2019; Thomas et al., 2019). Butterflies, one of the most monitored and studied insect groups, show evidence of declines across Europe (Warren et al., 2021) and in North America (Forister et al., 2021). Consequently, understanding the drivers influencing the dynamics of populations and communities of this insect group is important, because synchronous fluctuations in abundance can lead to short-term losses of ecosystem function (Greenwell et al., 2019) and long-term declines to reduction in mean function provision.

Community and population stability can vary spatially. Butterflies, along with declines in abundance, exhibit recent distributional changes (Warren et al., 2021). Consistent with the Grinnellian niche concept (Grinnell, 1917a, Grinnell, 1917b), distributions of butterfly species are driven largely by abiotic factors, such as climate (Settele et al., 2008), and their population dynamics are driven by weather (Palmer et al., 2017; WallisDeVries et al., 2011). In combination, we can expect that populations are most abundant and stable near the centre of their niche range and are most sensitive to environmental variation at range edges (Brown, 1984; Mills et al., 2017; Osorio-Olvera et al., 2020). Consequently, the local dynamics of communities might be impacted by the larger-scale biogeographical context of the constituent species, in addition to local factors, such as weather anomalies.

The overall stability of a community is contingent on several factors. Community stability often increases with biodiversity (Cardinale et al., 2012; Jiang & Pu, 2009), although the strength of this relationship can vary between systems (Campbell et al., 2011). Early mechanistic explanations for diversity–stability relationships included the insurance hypothesis (Yachi & Loreau, 1999) and the portfolio effect (Doak et al., 1998; Tilman et al., 1998). These are now combined in modelling frameworks demonstrating that higher species richness increases asynchrony in population dynamics and, along with mean population stability, influences the overall community stability (Thibaut & Connolly, 2013; Wang & Loreau, 2014).

Therefore, when considering butterflies and other insect groups, it is necessary to consider factors that generate asynchrony or that lead to higher overall population stability. Asynchrony is often associated with competition, but this might be less important in mobile animal communities where competition is diffuse (Loreau & De Mazancourt, 2008). Many animal communities at similar trophic levels are characterized by low competition owing to limiting similarity (Macarthur & Levins, 1967) or specialization (Futuyma & Moreno, 1988). Butterflies are resource generalists as adults (Dennis, 1992; Sharp et al., 1974) but more often specialists as larvae, owing to the adaptations required to circumvent structural or chemical plant defences (Ehrlich & Raven, 1964), which might limit interspecific competition. Although interspecific competition is present in phytophagous insects (Kaplan & Denno, 2007) and can be present in some butterfly species with shared host plants (Millan et al., 2013), or through apparent competition (Audusseau et al., 2021), asynchrony might be driven primarily by differential responses to weather anomalies, given the known effects of climate and weather on distributions and population dynamics.

Likewise, weather can have large impacts on population stability, particularly in association with range position. The links between distribution, abundance and geographical range have long been of interest in macroecological theory (e.g., abundant centre hypothesis; Brown, 1984), often supported by mixed evidence (Sagarin & Gaines, 2002). But substitution of the geographical position with niche position (abundant niche centre hypothesis; Martínez-Meyer et al., 2013; Yañez-Arenas et al., 2014) shows support for increased abundance (Osorio-Olvera et al., 2020), positive population trends (Manthey et al., 2015) and genetic variation (de Mazancourt et al., 2014) towards niche centres. Consistent with this, butterflies are more impacted by weather anomalies towards geographical...
range edges (Mills et al., 2017), although the effect of niche position is less explored. Consequently, the mean population stability of a community might be impacted not only by the absolute size of a weather anomaly, but also by the niche position of the population. Therefore, for both the main factors influencing community stability (i.e., asynchrony and population stability), the bioclimatic context of species and their responses to key weather variables might be key factors to understand and predict differences in community stability at large scales.

A challenge with selecting a key environmental driver for community stability is that species vary in their sensitivity to different environmental variables (Lawson et al., 2015). Functional responses to many variables, although identified (McDermott Long et al., 2017; Roy et al., 2001), are not typically known in detail, limiting our understanding of how responses combine to influence the whole community. Therefore, the approach applied here is to select a single influential type of anomaly and evaluate how species responses contribute to community stability, asynchrony, mean population stability or any other stabilizing mechanism. Likewise, to understand the impact of bioclimatic context, we apply the abiotic niche (Grinnell, 1917b; Hutchinson, 1957) and the abundant niche centre (Martínez-Meyer et al., 2013; Yañez-Arenas et al., 2014) concepts along a single major axis. This allows us to understand how the range position of species combines with local climatic anomalies to influence community stability at large scales. We select here temperature, owing to its influence on key biological rates that impact fitness (Kingsolver, 2009) and because it gives insight into the potential ecological impacts of climate change (Altwegg et al., 2017; Palmer et al., 2017; Pandori & Sorte, 2019).

Temperature responses are relatively well characterized in ectotherms (Angilletta et al., 2002, 2010), with biological rates typically increasing up to an optimum temperature before subsequently declining rapidly (Briere et al., 1999; Huey & Kingsolver, 1989; Shi & Ge, 2010). Therefore, an expectation is that populations of species at the cold edge of their temperature range (position in the temperature range is termed here “thermal range position”) will have contrasting responses to local temperature anomalies compared with populations of species that are at the hot edge (i.e., position in the thermal range is a potential mechanism driving asynchrony in population dynamics of different species; Jiguet et al., 2010). Position in the thermal range can also impact the average stability of a population because those near their optimum might have milder responses to anomalies than those at range edges where populations are prone to crashes (Mills et al., 2017; Oliver et al., 2014). However, temperature

![Figure 1](https://example.com/fig1.png)

**Figure 1** Hypothesized responses to temperature anomalies at five sites across a thermal range for species showing different combinations of thermal specialism and local adaptation. The mean temperature at the five sites is indicated by the vertical dashed lines, with colour representing the mean temperature (e.g., blue to red, cold to hot mean temperatures). In ectotherms, thermal performance (e.g., rates of growth, development and survival) are characterized by a shape that increases to a maximum before decreasing more sharply. We define local adaptation as the matching of the thermal performance curve for a species to the mean local conditions. (a,c) Species with low local adaptation have a single curve across their range. (e,g) In contrast, adapted species have their maximum thermal performance at the mean conditions at a site. To generate the potential impacts of temperature anomalies on population growth rate for these adaptation–specialism combinations, we simulated temperature observations for the five sites: one site at the optimum temperature for the performance curve shown in (a,c), then at 80, 90, 110 and 120% of this value. Linear models predicting logarithmic growth rate were then fitted to these data with fixed effects matching the structure of a model described below in Equation (1). (b,d,f,h) The expected mean responses. Further details about the simulation are presented in the Supporting Information (Appendix S2).
responses might also be influenced by local adaptation or acclimation (Huey & Kingsolver, 1989), impacting the expected mean population stability and asynchrony in the community.

Here, we test how the position in the thermal range and temperature anomalies combine to influence community stability at a European scale. We test the following four hypotheses (H1–H4): H1: populations at the cold and hot range edges of their thermal range will have contrasting responses to local temperature anomalies (see Figure 1 for a simulation articulating H1); H2: communities with a more mixed composition of species’ thermal range positions will have more stable dynamics overall. The two further hypotheses test possible mechanisms driving the effects of thermal range position: H3, communities composed of species from a mix of range positions will have contrasting population responses to temperature anomalies, leading to asynchrony in abundance and higher overall community stability; and H4, populations near the centre of species’ thermal ranges will be more stable, meaning that communities with more populations at the thermal range centres will, overall, be more stable (for a diagram articulating H2, H3 and H4, see Supporting Information Figure S1). We used the inverse of temporal variation in total community abundance as a proxy for community stability (Tilman, 1999) because it is tightly related to ecosystem function owing to mass ratio effects (Dangles & Malmqvist, 2004; Grime, 1998; Smith & Knapp, 2003).

2 | MATERIALS AND METHODS

2.1 | Data

Data were collected with butterfly monitoring schemes carried out in Finland, Spain (Catalonia) and the UK. The schemes consist of a network of sites where volunteers count butterflies along transects following a standardized framework called the “Pollard walk” (Pollard & Yates, 1993). We then processed counts using generalized additive models (GAMs) to provide an index of abundance per site and year (Dennis et al., 2013; Schmucki et al., 2016). In the case of missing counts, we interpolated from the GAM fitted to counts made at other sites in the same bioclimatic zone (Metzger et al., 2013; Schmucki et al., 2016). This approach generates unbiased estimates of abundance and performs better than interpolations from simple linear regressions (Dennis et al., 2013). However, to assure robust estimates, we removed indices of abundance with >50% of missing observations (Schmucki et al., 2016).

The schemes started in different years, and the number of sites per region varied: Finland (1999, number of sites = 107), Spain (1994, number of sites = 130) and the UK (1976, number of sites = 2128). Therefore, Finland, the country with the shortest scheme, set the study period (1999–2017). From each of the schemes, only those sites with >10 years of data were retained, leaving 59, 55 and 762 sites from Spain, Finland and the UK, respectively. The data included 157 species, and Spain had the highest species richness and the highest number of species (S) unique to the country (S = 131, unique = 74), followed by Finland (S = 58, unique = 12) and the UK (S = 58, unique = 8). We converted counts into densities by dividing the indices of abundance by the transect length, thus standardizing the measure of abundance across transects of different lengths.

Temperature data were obtained from the European Climatic and Assessment Dataset (ECAD) project (Haylock et al., 2008; Klok & Klein Tank, 2009). ECAD provides gridded daily temperatures at a 0.1° scale through interpolations from observations collected by a network of meteorological stations. The temperature was collected to the nearest degree at each site from the period of 1999–2017.

2.2 | Thermal range construction

The locations of the transect sites within the thermal range of each species were measured in environmental (temperature) space. We took this approach because our focus was the position of a species in its temperature niche, rather than geographical position, because changes in landscape features and seasonal variation in climate at a continental scale can decouple the relationship between geographical space and weather (Loarie et al., 2009).

It was also necessary to select the temperature for a given time period because, given our broad biogeographical scale, different locations will have different positions in the temperature niche over time (e.g., sites in Finland are warmer in summer than sites in the UK but colder in winter). Furthermore, given that our analysis included many species with varying phenologies, it required a broad period. We selected spring (March–May) because it coincides with the developmental period for many species and should be a broad enough period to capture differences across our spatial scale without averaging out anomalies. To standardize the position in niche space, we first took the daily temperatures at each site for the spring period and calculated the mean for the preceding 30 years (1988–2017) to obtain an average site temperature (T). We then transformed these mean values (Equation 1) such that the maximum mean temperature for each species across all sites was given a score of one and the minimum mean temperature minus one. Thus, every site-by-species combination had a position within the thermal range between minus one and one:

\[
\text{Thermal range position}_{ij} = 2\left[\frac{T_{ij} - \min(T)}{\max(T) - \min(T)}\right],
\]

where \(T_{ij}\) is the mean temperature of the site for species \(i\) at site \(j\), and \(T\) is the set of mean temperatures from all sites for that species.

The thermal range preserved the relative difference between the temperatures at the sites where a species was observed but standardized thermal range differences between species, hence a score of +0.75 covers 87.5% of the total projected temperature range for a species independent of the absolute size of the temperature range. Note that the temperature range across all our monitoring sites from Catalonia (Spain) to Finland will not always capture the full thermal range for every species (e.g., some extend beyond the
2.3 | Hypothesis 1: Negative covariance in response to temperature anomalies at thermal range edges

We predicted that the population responses of a species, in terms of change in the logarithm of abundance between one year ($N_{t-1}$) and the next ($N_t$), $\ln(N_t/N_{t-1})$, to a local temperature anomaly will be dependent on the position of the site within the thermal range for the species and that populations at the cold and hot edges will have contrasting responses to a given direction of temperature anomaly (see articulation of the hypothesis in Figure 1). To test this hypothesis, we fitted the relationship between the change in population size in response to a local temperature anomaly (the difference each year between the observed seasonal mean temperature and the average seasonal temperature conditions at a site over the last 30 years) and the thermal range position for all species in a hierarchical Bayesian model.

We predicted, on average, a nonlinear response, with large temperature anomalies either above or below the mean reducing population size, but also that the shape/direction of this response might be influenced by thermal range position and would vary between species. Given that thermal performance might be nonlinear (Figure 1a), we included fixed and random quadratic and cubic terms for temperature anomaly and interactions of each of these terms with thermal range position (Equation 2). The cubic model allowed the shape of the response to vary between species and to vary with range position, accounting for varying amounts of factors such as local adaptation (Figure 1b,f). It also allowed us to account for sharp drops in performance with large anomalies. We also included a fixed and random density dependence term of the logarithm of the population size in the previous year and random effects of site and year to account for repeated measures (further discussion regarding model alternatives is provided in Supporting Information Appendix S3).

To start, we fitted the a priori plausible maximal model (Equation 2). Fitting this model produced singular fits in higher-order random effect terms. Consequently, we simplified the model by progressively removing the singular highest-order terms one at a time until we obtained a model with sensible precision overall random effects (Equation 2) (Barr et al., 2013). The removal of these higher-order terms reduced the deviance information criterion (DIC; Spiegelhalter et al., 2002) and the Watanabe–Akaike information criterion (WAIC; Watanabe & Opper, 2010) scores of the model fit.

$$\ln(N_t/N_{t-1})_{\text{ijkl}} \sim N(\gamma_{ijkl} \sigma^2)$$

$$\gamma_{ijkl} = \ln(N_{t-1}) + \text{temp anomaly}_i + \text{temp anomaly}_j^2 + \text{temp anomaly}_k^2 + \text{trange}_l + \text{temp anomaly}_i \cdot \text{trange}_l + \text{temp anomaly}_j \cdot \text{trange}_l + \text{temp anomaly}_k \cdot \text{trange}_l + \text{temp anomaly}_i^2 \cdot \text{trange}_l + \text{temp anomaly}_j^2 \cdot \text{trange}_l + \text{temp anomaly}_k^2 \cdot \text{trange}_l + \text{Site}_i + \text{Year}_j + \text{Species}_k + f(\theta_y)$$

Here, subscripts refer to site $i$, species $j$ and year $k$; $\sigma^2$ refers to the variance and $f(\theta_y)$ the value of the Gaussian spatial field at location $x'y'$; trange refers to the thermal range position and temp anomaly the local temperature anomaly.

We used the integrated Laplace approximation (INLA) method for approximate Bayesian inference (Rue et al., 2009), which provides a method for fitting a Matérn covariance spatial autocorrelation function, through a weak solution to a stochastic partial differential equation (SPDE) (Gómez-Rubio, 2020; Lindgren et al., 2011; Lindgren & Rue, 2015). Further information about prior selection is provided in the Supporting Information (Appendix S3).

2.4 | Hypothesis 2: Community stability from thermal range position

We predicted that communities with central thermal range positions on average will have more stable community dynamics. To measure community stability, we used the inverse coefficient of variation (CV) for yearly abundance counts summed across all species at a site across the sampling period (maximum of 30 years), hence it was a measure of variation in total community abundance. This approach is similar to stability measures used for studies of ecosystem functioning (e.g., Donohue et al., 2016; Hautier et al., 2015), although here we do not extrapolate our results to function. Given that time series were not identical in length, we used a bias correction for smaller sample sizes, because the CV can be underestimated from shorter time series, and we also limited the data to sites with ≥10 years of data (Equations 3.1 and 3.2):

$$\text{CV}_j \dagger = \frac{\sigma_j}{\mu_j} \left(1 + \frac{1}{4n_j}\right), \quad (3.1)$$

$$\text{Community stability}_j = \frac{1}{\text{CV}_j \dagger}. \quad (3.2)$$

where $\dagger$ refers to the bias-corrected estimate of the coefficient of variation at site $j$ and $n$ to the sample size, $\sigma$ to the standard deviation of abundance, and $\mu$ to the mean abundance.

Community stability is also known to be affected by species richness (Tilman & Downing, 1994) and mean abundance (Taylor, 1961). Therefore, at each site, we calculated the mean thermal range position of the species (using range position from average yearly temperatures), the mean community abundance (measured as density to compare across transects of different lengths) and the species richness.

For the model, we predicted a quadratic effect of thermal range position on community stability, because stability should increase...
with a mean range score of zero and decrease when species are, on average, at either side of the range edges, and we also included linear effects of species richness and mean community abundance to control for their effects on stability:

\[
\text{Community stability}_i \sim N(\mu, \sigma^2). \quad (4)
\]

\[
y_j = \text{Mean trange}_j + \text{Mean trange}_j^2 + \text{Species richness}_j \text{ Mean abundance}_j + f(\theta_y).
\]

Symbols used here are the same as in Equation 2.

2.5 Hypothesis 3: Asynchrony

We hypothesized two mechanisms that might explain the effect of thermal range position on community stability: \(H_3\), communities composed of species from a mix of range positions will have contrasting population responses to temperature anomalies, leading to asynchrony in abundance and higher overall community stability; and \(H_4\), populations near the centre of species’ thermal ranges will be more stable, meaning that communities with more populations at the thermal range centres will, overall, be more stable.

We predicted that if asynchrony was influenced by opposing thermal range positions, the mean synchrony in the community (i.e., averaged across all pairs of species) should be lowest in communities with central average range positions (i.e., concave up relationship). To test this, we calculated the average population synchrony (measured as the mean of all pairwise correlations across species’ time series at a site, including only species pairs with \(\geq 10\) concurrent observations). We then constructed a linear model (Equation 5) for synchrony against mean range position and mean range position². Given that synchrony is expected to increase with species richness (Ives et al., 1999; Thibaut & Connolly, 2013), this was calculated for each site and included in the model.

\[
\mu \text{ Synchrony}_j \sim N(\mu_{\text{syn}}, \sigma^2), \quad (5)
\]

\[
y_{\text{syn}} = \text{Mean trange}_j + \text{Mean trange}_j^2 + \text{Species richness}_j.
\]

Here, \(\mu\) indicates the mean, and other symbols are as in Equation 2.

Given that we were modelling an estimate of mean synchrony, the regression was weighted by the inverse of the variance to control for variation in the precision of the estimate of the mean owing to differences in sample size. INLA does not offer a straightforward method for implementing weighted regression; consequently, we used the “brms” package (Bürkner, 2017) to fit a weighted regression using Hamiltonian Monte Carlo through Stan (Carpenter et al., 2017).

2.6 Hypothesis 4: Population stability

We predicted that average population stability might increase at the centre of the thermal range, leading to increased community stability. We calculated the stability of the population dynamics of each species at each site, as in Equations 3.1 and 3.2, along with the mean population abundance. Observations included only species–site combinations with \(\geq 10\) observations in total, as in Equation 4. To test our hypothesis, we constructed a model with fixed and random effects of range position, range position², and mean population abundance, along with random intercepts of year and site. After fitting, the year intercept was found to be near singular and was removed from the model. Population abundance was also Ln-transformed before the fit because there were a few large populations that had an outsized influence on coefficient estimates. Information on priors is provided in the Supporting Information (Appendix S1).

\[
\text{Population stability}_{ik} \sim N(y_{ik} | \sigma^2). \quad (6)
\]

\[
y_{ik} = \text{range} + \text{range}^2 + \text{Mean abundance} + \text{range} + \text{range}^2 + \text{Mean abundance}_i + f(\theta_y) + \text{Site}_i + \text{Species}_i.
\]

\[
\text{Site}_i \sim N(0 | \sigma^2_j),
\]

\[
\text{Species}_i \sim N(0 | \sigma^2_i).
\]

All models were constructed in R v.4.1.1 (R Core Team, 2021). The hierarchical Bayesian models were fitted using the “INLA” package (Lindgren & Rue, 2015), and the package “brinla” was used to aid in diagnostics (Faraway et al., 2021). Polygons for barrier models were taken from the “maturaleza” package (South, 2017), and spatial fields were plotted using functions from the paper by Krainski et al. (2018). Code supporting the results is archived at DOI: 10.5281/zenodo.6350070.

3 RESULTS

3.1 Hypothesis 1: Responses to temperature anomalies across the thermal range

The model fit did not conform precisely to a thermal performance curve, although responses of population growth to anomalies were nonlinear. Species in the warmer half of their thermal range were most impacted by large anomalies, showing decreased growth rates with hot anomalies and increased growth with cold anomalies (Figure 2a). The response to temperature anomalies was weaker overall at the cold edge of the range, and there was evidence of contrasting responses, with species at the hot edge performing worse with high-temperature anomalies and better with low-temperature anomalies compared with species at the cold edge (Table 1; Figure 2a). However, at extreme hot temperatures the populations from all range positions declined, and towards the warm edge the declines were substantial. The variance of the random effects (Table 1) suggested that there was interspecies variety in both the shape of the response and the importance of the interaction terms.
The spatial field demonstrated relatively broad-scale regional correlations (range: 95 km) in average population changes in the areas such as the south-east of England, showing lower growth rates relative to much of the interior, and the north-eastern tip of Spain, showing higher growth rates than the interior (Figure 2b).

### 3.2 Hypothesis 2: Community stability

Community stability was influenced by thermal range position, with stability increasing from low mean range positions to central range positions, although the fit suggested that stability declined only at the very upper end of mean range positions (Figure 3a,c). Species richness increased stability; however, the 95% credible interval for mean community abundance overlapped with zero, although it was still positively associated with community stability. The spatial field showed much smaller-scale spatial correlations compared with those found for growth rates with a geographical range of c. 25 km (Figure 3b; Supporting Information Table S1).

### 3.3 Hypothesis 3: Asynchrony

We found that synchrony tended towards being slightly concave up with thermal range position (Figure 4a). However, species richness had a clearer effect, with species richness decreasing synchrony (Figure 4b,c; Supporting Information Table S2).

### 3.4 Hypothesis 4: Population stability

For mean population stability, rather than an increase in the centre of the range, stability was lowest at the hottest range edge (Figure 5a; Supporting Information Table S2), although the overall effect was weak. However, mean population abundance increased stability (Figure 5b).

### 4 DISCUSSION

We tested four hypotheses about how the position of species within their thermal range and anomalies interact to influence population responses and community stability: H₁, species will have contrasting responses to temperature anomalies at different ends of their thermal ranges; H₂, communities consisting of populations with central thermal range positions on average will be more stable; H₃, communities composed of species from a mix of range positions will have contrasting population responses to temperature anomalies, leading to asynchrony in abundance and higher overall community stability; and H₄, populations near the centre of species’ thermal ranges will...
Information Table S1). The mechanisms (H; ) showing smaller reductions in performance at both edges of the thermal range are more responsive to temperature anomalies, with the cold edge showing larger declines (H; ). However, responses were highly nonlinear relative to expectations (Figure 2a), and populations at the hot edge performed worse with high-temperature anomalies and vice versa. Our analysis of responses to temperature anomalies (Figure 2a) suggests that the composition of thermal range positions at a site is unlikely to aggregate in a simple, consistent way to impact community stability. This is because the degree of synchrony between species responses might be contingent upon the relative size of the temperature anomaly. For example, two species at either side of the centre of their thermal ranges might have small asynchronous responses to moderate weather variation, but in an extreme high-temperature event, both species might be pushed beyond thermal limits (Sunday et al., 2014) and crash synchronously. Our test of the asynchrony mechanism supports this, because synchrony decreased only slightly in the centre of the range, with species richness having a much larger effect.

A few reasons might explain why we found only a weak impact of asynchrony with thermal range position. First, although temperature is an important driver, butterfly species are impacted by other weather variables, such as precipitation (Herrando et al., 2019; Roy et al., 2001) and aridity (Oliver et al., 2015). This might disrupt the expected opposing responses of species at different thermal range edges. Second, temperature can also affect species in unpredictable ways owing to its impact on natural enemies or host plants, and phenology might also vary across the study region, affecting the life stage impacted by the anomaly (McDermott Long et al., 2017). Consequently, asynchrony might be a larger driver of stability in butterfly communities in general, but the contrasting responses are not strongly connected to the thermal range constructed here. Instead, large temperature anomalies might lead to synchrony rather than asynchrony.

Our results suggest that populations at the hot edge of the thermal range are, in general, more responsive to weather anomalies than those at the cold edge. Previous work has shown that butterfly populations are more variable at range edges (Mills et al., 2017; Oliver et al., 2012), but greater declines in population growth rates towards the hottest edge have also been noted in US butterflies (Breed et al., 2013) and birds (Jiguet et al., 2010). We provide further support for population instability at the hot edge with our analysis on the mechanisms (H; and H; ), finding that populations at the hot edge are the least stable. These results are consistent with abundant niche centre hypotheses (Martínez-Meyer et al., 2013; Yañez-Arenas et al., 2014), and we add here new evidence for niche position impacting both population and community stability, in addition to abundance (Osorio-Olvera et al., 2020). But the increased impact of temperature variation and the decline in populations at the warmer edge are also suggestive of impacts of climate warming on butterfly populations. Negative impacts of climate change on population trends have been noted in many taxa, including butterflies (Martay et al., 2017); therefore, management might be required to enhance the stability of populations at the warm edge with future climate change (Oliver et al., 2015). Furthermore, the results suggest that relative niche position might be a simple but important indicator

### TABLE 1 Estimated regression parameters for relationships between population change, temperature anomalies and thermal range position using the hierarchical Bayesian model in Equation 2

| Fixed effects          | Mean     | Lower credible interval | Upper credible interval |
|------------------------|----------|-------------------------|-------------------------|
| Intercept              | 0.636    | 0.556                   | 0.717                   |
| lnNt−1                 | −0.360   | −0.388                  | −0.322                  |
| Temperature anomaly    | 0.026    | 0.007                   | 0.045                   |
| Temperature anomaly²   | −0.003   | −0.015                  | 0.010                   |
| Temperature anomaly³   | −0.008   | −0.012                  | −0.005                  |
| Range position         | −0.018   | −0.051                  | 0.015                   |
| Temperature anomaly:Range position | 0.004 | −0.020                  | 0.015                   |
| Temperature anomaly²:Range position | 0.014 | −0.002                  | 0.030                   |
| Temperature anomaly³:Range position | −0.007 | −0.011                  | −0.002                  |

| Variances random effects | σ | 0.550 | 0.546 | 0.553 |
|--------------------------|---|-------|-------|-------|
| Intercept site           | 0.014 | 0.012 | 0.016 |
| Intercept year           | 0.018 | 0.012 | 0.028 |
| Intercept species        | 0.062 | 0.050 | 0.077 |
| lnNt−1                   | 0.021 | 0.013 | 0.031 |
| Temperature anomaly      | 0.0064 | 0.0052 | 0.0081 |
| Temperature anomaly²     | 0.0032 | 0.0024 | 0.0040 |
| Temperature anomaly³:Range position | 0.0073 | 0.0057 | 0.096 |
| Temperature anomaly²:Range position | 0.0041 | 0.0031 | 0.0056 |
| Range spatial field (km) | 94.85 | 55.27 | 156.37 |
| Standard deviation spatial field | 0.054 | 0.047 | 0.062 |

Note: Credible intervals are set to contain 95% of the posterior. lnNt−1 refers to the log of the abundance of the previous year.
of the populations and communities most at risk from climate change (e.g., Settele et al., 2008).

Although populations towards the hot end of the thermal range were more unstable, the communities at these locations were only marginally less stable than in the centre, and communities in Finland, at the cold edge, were overall the least stable. This is likely to be attributable to a combination of these communities having lower abundance and species richness than the communities in the UK and Spain, respectively; Spain, in particular, has much higher species richness than the other two countries. We do not include asynchrony directly in Equation 4, which ultimately drives the species richness effect (Thibaut & Connolly, 2013), and we only include species richness, which could lead to an underestimation of the negative impacts of communities dominated by populations at the hot range edge. Therefore, a more structural account of the initial impact of species richness on asynchrony, then of asynchrony on community stability (e.g., Olivier et al., 2020), could be informative for understanding community stability at this broad spatial scale.

Furthermore, the compositional change in species between the countries might also account for some differences in community stability. The UK contains the northern range limits of many species owing to its cool climate (Warren et al., 2001), and the species assemblage that survives in the UK might be an adaptable and resilient subset of the wider European assemblage (Thomas, 1993). Average trait differences between the assemblages might also explain differences in community stability beyond range position, because mobility has been shown to impact recent population trends in butterflies (Eskildsen et al., 2015), and numerous traits impact the
sensitivity to climate change for many species (Pacifici et al., 2017). Finally, Spain also contains the largest number of unique species, and although there are large temperature differentials between sites in Spain, some species might have thermal range limits beyond those encountered. Consequently, extending the study to include a larger area and to include data from other European butterfly monitoring schemes might provide an increased resolution for understanding population and community responses to temperature anomalies at the European scale.

After accounting for the effects of thermal range position and temperature anomalies, we found differing patterns in the spatial fields that represent average rates of population change and community stability across the study area (Figures 2b and 3b). There is a general expectation that sites closer to one another should have similar dynamics because they share various environmental conditions (Tobler, 1970), which we found. The spatial pattern for rates of change of interannual population was in the range of 100 km, suggesting similar patches of average population change at regional scales for our countries. The spatial field for community stability, however, showed a smaller-scale patterning (range c. 25 km). The different spatial patterns are likely to be influenced by different drivers. Population dynamics might be driven by regional weather conditions (Breed et al., 2013), possibly interacting with broad land-cover types (Stefanescu et al., 2011) or factors such as elevation. The Spanish sites offer a possible example of this, with the north-eastern tip of Catalonia, which is associated with higher elevation and greater amounts of woodland cover (García Viñas et al., 2006), showing more positive trends than the adjacent interior. Alternatively, the smaller-scale pattern of community stability might be related to landscape heterogeneity (Oliver et al., 2010) or small-scale variations in site quality owing to differences in local-scale habitat management. These processes require further investigation, but our results demonstrate the potential of INLA for deriving new hypotheses around the drivers of community and population variability at varying scales.

We note some additional limitations with the research. First, the environmental data are ultimately coarse grained (c. 11 km × 11 km), and species are not solely responsive to changes in mean temperature. Insects are influenced by the microclimate (Duffy et al., 2015), fine-scale habitat features (Aguirre-Gutiérrez et al., 2017) and multiple landscape factors. Second, ectotherm development is influenced by the cumulative effects of temperature (e.g., growing degree days), and in these cases the average temperatures or anomalies can be misleading (Denny, 2017; Sunday et al., 2014). Linking community patterns with detailed mechanisms, such as context-dependent growth rates, is a major challenge and currently has been attempted only at small scales (e.g., White et al., 2020). In future, modelling efforts might be better able to connect species physiology to large-scale patterns observed for butterflies by using standardized schemes (Johnston et al., 2019), but this would require tighter integration of the cycle of data collection, theory, modelling and testing than is typically practised (Boul et al., 2021; Dietze et al., 2018). Finally, our data might contain observational errors, which have been shown to explain variation in community stability (de Mazancourt et al., 2014). There has been exploratory work on the possible impacts of species detectability in butterfly counts (Isaac et al., 2011), but it remains unclear whether adjusting for detectability would improve estimates of real abundance. Given that we measure variation in relative abundance across years at fixed locations, where detectability should be reasonably constant, it is unlikely to have a large impact on our results.

![Figure 5](image-url)

**FIGURE 5** Marginal relationships between population stability and (a) thermal range position, or (b) natural logarithm of mean abundance. Spanish sites are shown in red, Finnish sites in blue and UK sites in green. Lines show marginal fits with 95% credible intervals. Note that (a) does not show the full extent of the residuals, in order to show the model fit better.
5 | CONCLUSIONS

In summary, we have demonstrated that the interaction of thermal range position and local anomalies influences both interannual population change and community stability in butterflies at a broad biogeographical scale. We found that range edge communities were less stable and that populations near the hot edge of the thermal range were most strongly influenced by temperature anomalies. Responses were also nonlinear, meaning that many species might be impacted strongly by extreme heat events. Aggregating population responses across a single-niche dimension, although informative, does not simply predict community stability, because we found that small temperature anomalies can produce weak compensatory dynamics, but large extreme events might synchronize dynamics. Our results suggest that niche position is an important determinant of community and population stability, but the larger sensitivity of populations at the hot edge suggests that community stability at hot locations might be most impacted by climate change.

AUTHOR CONTRIBUTIONS

L.C.E. designed the study and performed the analysis. R.S. processed the butterfly indices before analysis. L.C.E., T.H.O. and Y.M. wrote the first draft. All other authors contributed substantially to revisions.

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DATA AVAILABILITY STATEMENT

Standardized monitoring data for Catalonia and Finland are available upon request to the Catalan butterfly monitoring scheme and Finnish butterfly monitoring scheme, respectively. UK monitoring data are freely available from the Environmental Information Data Centre (EIDC). Community-level data and code in support of the results are available at: 10.5281/zenodo.6350070.

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BIOSKETCH

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SUPPORTING INFORMATION

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