Temperature-Driven Biodiversity Change: Disentangling Space and Time

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Temperature regimes have multiple spatial and temporal dimensions that have different impacts on biodiversity. Signatures of warming across these dimensions may contribute uniquely to the large-scale species redistributions and abundance changes that underpin community dynamics. A comprehensive review of the literature reveals that 86% of studies were focused on community responses to temperature aggregated over spatial or temporal dimensions (e.g., mean, median, or extremes). Therefore, the effects of temperature variation in space and time on biodiversity remain generally unquantified. In the present article, we argue that this focus on aggregated temperature measures may limit advancing our understanding of how communities are being altered by climate change. In light of this, we map the cause-and-effect pathways between the different dimensions of temperature change and communities in space and time. A broadened focus, shifted toward a multidimensional perspective of temperature, will allow better interpretation and prediction of biodiversity change and more robust management and conservation strategies.

Keywords: climate change, climate velocity, community dynamics, species redistribution, range shifts

Environmental temperature is a primary variable important for biological function at all organizational scales. Even slight temperature changes can dramatically affect biological processes from cells to populations, with strong ecological consequences. At the smallest scale, temperature drives cellular reaction rates through kinetic processes. Individuals respond directly to environmental temperature—for example, by modulating their activity rates (Payne et al. 2016, Payne 2018). Population demographic rates are also temperature sensitive (Dell et al. 2011), with consequences for abundance and occupancy patterns. Demographic changes in combination with individual effects (e.g., foraging velocity, ingestion rates; Dell et al. 2011) lead to shifts in species interaction strengths (disease prevalence, Kock et al. 2018; parasitism rate, Runjie et al. 1996), ultimately translating to change in community dynamics and structure (Kordas et al. 2011, Bellard et al. 2012).

Environmental temperature is highly variable in both space and time. Some aspects of environmental temperature are predictable (e.g., seasonal changes), others are not (e.g., extreme events). Therefore, a temperature regime has multiple dimensions that can be described in both space and time, with the potential to shape biological patterns in different ways (Garcia et al. 2014). For example, as we move from the tropics to the poles, it becomes colder, daily variability decreases, but seasonal variability increases (Wang and Dillon 2014). Distinct signatures of spatial variability also exist; for example, temperature is much more spatially heterogeneous in intertidal systems than in subtidal systems. Along these different axes of temperature dimensions in space and time, there is an additional layer of gradual long-term warming due to anthropogenic climate forcing (IPCC 2013).

Climate change is altering environmental temperature regimes. The spatial arrangement (i.e., spatial heterogeneity) and timing of temperature change is typically abstracted to a statistical distribution, defined presently and by others as temperature magnitude (see box 1 and Garcia et al. 2014). However, by examining only temperature magnitude (e.g., its mean), we are deprived of the detailed dynamics of spatial and temporal temperature change. The position and availability dimensions of temperature change can vary, even without shifts in the overall mean temperature of a region or through time (see box 1 and figure 1 for further definitions; Garcia et al. 2014). For example, the movement of thermal isolines with warming is an example of a shift in the position of temperature in space (Loarie et al. 2009, Burrows et al. 2011, Hamann et al. 2015). The earlier onset of spring represents a change in position of temperature in...
Temperature regimes—the characteristic pattern of temperature variation in space and time for a given scale—vary across the Earth. For example, temperate regions since the Late Quaternary have been characterized by cool and warm periods (i.e., seasonality) through the year and relative cool climates. Four different dimensions of a temperature regime are generally recognized (see Garcia et al. 2014), and how they vary with climate change is described in the present article (see figure 1). Temperature magnitude describes the change in the statistical distribution of temperature for a given locality. The statistical distribution is defined by both its central tendency (e.g., changing mean—i.e., warming or cooling) and its dispersion (e.g., increasing variation, skew or kurtosis—i.e., extreme events). The rate of change in temperature magnitude is defined by the change in the statistical parameter (e.g., mean, standard deviation) per unit time.

Underlying this change in temperature magnitude are changes to temperature position and temperature availability in space and time (figure 1). Temperature temporal position describes the change in the timing (i.e., date) of a specific temperature event within a defined spatial unit. This contrast with temperature temporal availability, which describes a change in the total duration of a specific temperature event within a defined spatial unit. Temperature spatial position is defined by the relocation of temperature to a new area (i.e., isotherm shift change in linear distance) for a given temporal unit (Loarie et al. 2009, Hamann et al. 2015, figure 3). In contrast, temperature spatial availability measures the change in area or size of a temperature available within a geographic location (i.e., the change in geographic space of a temperature regime measured in square kilometers, km²) for a given temporal unit (figure 3). Novel climates are an important component of temperature spatial availability and describe the availability of new climatic space, increasing from an initial area of 0 km² (Williams and Jackson 2007).

Likewise, community metrics also fall into the following broad categories to measure structural and compositional differences in species assembled at local scales: species richness, total abundance, species relative abundance, compositional and trait based (Smith et al. 2009, Magurran and McGill 2011, Hill et al. 2016, Santini et al. 2016). These are important to recognize when matching the measured community responses to the processes driving change (i.e., “Linking community change processes and temperature dimensions”). The total number of different species in a community is measured using species richness metrics. The net loss and gain of species translates to a change in richness. The total abundance of a community is simply the sum of all individuals in a community and is often related to species richness as a result of sampling effects (i.e., more individuals increases the probability of a new species being present). The distribution of individuals between species represents the structure of a community and is often summarized by the shape of species relative abundance distributions. A change in structure occurs with shifts in species relative abundances (e.g., few rare species versus many common species), but these changes are agnostic to species identity (i.e., the same structure, but the assemblage comprises all new species). Therefore, structural change can represent richness and total abundance changes simultaneously. Compositional metrics describe how both species’ relative abundances and identities shift and therefore measure the reorganization of species abundances in a community. The losses or gains of species measures the turnover component of compositional change (Baselga and Leprieur 2015). Finally, trait-based metrics quantify the diversity, range and values of the traits and niche properties of species within a community; these are often relevant to a particular driver of interest (e.g., species thermal limits and warming).
leading to difficulty in identifying with confidence which factors are mechanistic drivers of emergent patterns (Lawton 1999). As with considering changes in the temperature regime, the challenge becomes even greater when the additional complexity of community dynamics through time is of interest (McGill et al. 2015). Certain community dimensions are responsive to environmental changes, such as the composition and relative abundance of assemblages (Hill et al. 2016), whereas others, such as species richness, are less directly responsive to environmental change (Santini et al. 2016), with increases or decreases being highly dependent on measurement scale (Vellend et al. 2017). The multidimensional nature of both temperature and biodiversity variables justifies the aim of the present article: to map predictions of cause-and-effect among different dimensions.

**Literature review**

To determine what dimensions of temperature regime and community responses are most commonly studied, we reviewed articles published from 2005 until 2015 (see the supplemental materials for methods and discussion). We found that, in spite of the complexity in changing temperature regime, most research has been focused on identifying responses to mean warming trends. Of the 156 papers returned from our literature review, 86% were focused on temperature magnitude, and only a small proportion of research has investigated spatial position (3%) and availability (1%) or temporal position (4%) and availability (6%, figure 2). Of the metrics used to describe changes in temperature magnitude—the changing statistical distribution of temperature—mean changes were investigated in 48% of the studies, and 41% of the studies were focused on minimum or maximum temperatures. We also found a strong bias toward species richness (36%) rather than toward species identity (13%) or relative abundance (6%) as the predominant dimension of communities measured in responses to temperature change. We therefore reveal that the spatial and temporal complexity

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**Figure 1. Temperature patterns in defined spatial and temporal units and their change through time.** Temperature is variable in both space and time, but to aggregate to a mean value, a researcher must select scales of space and time to describe a particular statistical temperature distribution. Changes in a statistical distribution (e.g., central tendency and variation) can occur with no changes to the underlying spatial or temporal organization of temperature within those defined units (e.g., 50 square kilometers, 1 year). Likewise, even if no change is observed in the statistical distribution, underlying shifts in space and time may be masked, in both availability and position of temperature. We outline a few, of many, possible scenarios by which temperature magnitude, position, and availability can change.
underlying temperature change is rarely considered as a driver of community dynamics, instead temperature is generally abstracted over spatial or temporal scales to an aggregate mean value.

Objectives and purpose

Trends in the responses of biodiversity that have been detected and attributed to particular dimensions of temperature change in space and time are emerging. No-analogue communities have formed as species reshuffle in response to the development of novel temperature regimes (Urban et al. 2012). Increases and decreases in species richness have been attributed to changing species distributions following locational shifts in thermal isoclines (Devictor et al. 2012, Batt et al. 2017). Changing temperature regimes have also been implicated in driving the increasing relative abundance of widespread and common species, or homogenization, of communities (Davey et al. 2012, Magurran et al. 2015). There is widespread evidence of shifts toward species with thermal preferences for warmer environments (Bates et al. 2014a, Horta e Costa et al. 2014, Guizere et al. 2015, Tayleur et al. 2015, Stephens et al. 2016), a process known as tropicalization or thermophilization.

Notwithstanding these few examples, how different dimensions of temperature change will affect the dynamics of multidimensional communities is poorly established at present. To address this gap, we developed a conceptual framework to guide predictions and explicit quantitative tests of biodiversity change in response to temperature change, measured in the appropriate dimensions of space and time (O’Connor et al. 2015, Houlanah et al. 2017). We further illustrate below why neglecting the dimensionality inherent to both temperature regime and biodiversity change may prevent accurate predictions. We focused on changes in the richness of local communities and homogenization over space as illustrative examples of biodiversity change that are driven by multiple temperature change dimensions.

Linking community change processes and temperature dimensions

Two community processes are relevant to understand community dynamics at anthropogenic time scales—selection and movement (Vellend 2010, 2016). The selection and movement of individuals lead to change in richness, structure, composition of communities, and traits of the species present. These processes underpin local community dynamics that form an important component of biodiversity change (e.g., figure 4; Vellend 2010, 2016).

Abundance-related metrics (composition, total and relative abundance; see box 1) are expected to be most responsive to changes in selection processes acting on communities. For example, under changing environmental conditions, deterministic fitness differences between individuals within a population alter birth and death rates (i.e., demographic effects). This has consequences for population dynamics, and populations within a community increase or decrease in abundance. When species differ in the effect of these selection-based changes, variation in population dynamics between different species within a local community occurs. These population changes manifest themselves as changes in composition and relative abundance.

In contrast to selection processes relating to abundance metrics, changes in species richness and identity are expected to be sensitive to environmental changes that alter movement community processes: The immigration or emigration of individuals (either active or passive—e.g., migration or dispersal) into or out of a local community adds or removes species from a community. Most examples of movement mediated richness change come from colonization of novel species on islands (i.e., MacArthur and Wilson 1967), many of which are driven by human actions in recent times (Sax and Gaines 2008, Vellend et al. 2017). Furthermore, the capacity for species to disperse into communities as they assemble affects climax or equilibrium community richness (Lichter 2000, Makoto and Wilson 2016). Batt and colleagues (2017) reported a novel example from marine benthic fish assemblages in which increasing range size of rare and transient species, through movements to new localities, increased the species richness of any given location within a region.

Movement can also cause additional selection processes by creating interactions between arriving species and those present in the local assemblage and therefore drive additional community changes (Gilman et al. 2010, Urban et al. 2012, Alexander et al. 2015, Vellend 2016). For instance, Alexander and colleagues (2015) found that transplanting competitors into Swiss Alpine plant communities had large effects on the survival, biomass, and flower probability in species of the local assemblage.
How do community processes respond to specific dimensions of temperature change? The effects of a change in temperature magnitude (central tendency and variability) on local communities influence the selection process through population birth and death rates, leading to changes in the relative abundance of species found in a community (figure 4). Temperature-related magnitude changes occur through environmental filtering: selection of individuals with higher relative fitness and selection against individuals with low fitness. Changes in temperature magnitude predict biodiversity change, and this, in part, explains why this approach is so commonly adopted. A proximate cause of these community responses is that species often evolve to optimize temperatures frequently experienced, leading to a peak in performance (Angilletta 2009). In addition, temperature variability can exceed species’ limits (e.g., McKechnie and Wolf 2010, Dowd et al. 2015). For example, modeled population trends of water and sea birds across the United Kingdom, based partly on summer and winter temperature extremes, predict 56% of variation in average the population dynamics of birds in this region (Johnston et al. 2013; see the supplemental materials for further discussion).

However, complexities of temperature change in space and time are missed when considering temperature exclusively from this perspective. How the selection and movement of individuals respond to changes in temperature dimensions depends on the spatial and temporal nature of temperature changes (figures 1, 3). Considering the dimensionality of temperature leads to different predictions for how biodiversity will change with warming (Garcia et al. 2014, Ordonez et al. 2013, described in figure 4).

**Figure 3. Schematic diagram of spatial dimensions of temperature change.** Spatial availability is represented in the present article as differences in the total geographic area of thermal niche at a regional scale. Novel climates may emerge, representing newly available temperature regimes. Spatial position can be measured as forward or backward temperature velocity, which are proxies for different ecological processes of emigration and immigration respectively (Carroll et al. 2015, Hamann et al. 2015). Analogue velocity is the minimum distance necessary to travel to maintain constant temperature conditions (Hamann et al. 2015). Local climate velocity is the rate of temporal change in temperature over the spatial gradient of temperature (Loarie et al. 2009). Adapted from Garcia and colleagues (2014) and Carroll and colleagues (2015).
change in temporal position (e.g., seasonality and the earlier onset of spring, figure 1) and temporal availability (e.g., duration of temperatures above a physiological threshold, figure 1) drive demographic change (Jones and Wiman 2012, Gaillard et al. 2013, Matechou et al. 2014, figure 4). For instance, variation in spring timing (temporal position) between years reduced roe deer (Capreolus capreolus) population growth rates by limiting successful spring recruitment (Gaillard et al. 2013). In another example, broods in common blue butterfly (Polyommatus carus) populations emerged later at higher latitudes, because of differences in the availability of spring and summer temperatures. This change in temporal availability constrained the total broods within the year at northern sites, leading to a smaller overall population size (Matechou et al. 2014).

In ecosystems with seasonal cycles in temperature, a key indirect driver of community change is the mismatch in timing of life history events for species with strong dependencies. This effect is exacerbated if entire groups of species that interact have different capacities to respond to temporal position or, if changes to temporal availability alter how life-history stages transition (i.e., development times). Compelling examples are known of mismatches among resources, consumers, and predators. In a now classic example, Both and colleagues (2009) found that, for passerine birds, climate change led to advances in temporal position of caterpillar prey peak abundance that were unmatched by changes in peak food demand. The predators of these passerine birds did not shift the date of their energy requirements to keep pace with changing prey fledgling availability, and therefore, mismatches occurred at multiple levels across an ecological assemblage (Both et al. 2009). Similar mismatches in key seasonal timings across trophic levels were found with climate change for 726 plant, vertebrate and invertebrate taxa in the United Kingdom (Thackeray et al. 2010). Large-scale compositional changes are expected to occur in communities undergoing mismatches in the timings of species present, because mismatches lead to performance (and abundance) declines of species lacking the resources required within in a specific time window.

**Change in spatial dimensions of temperature change through time**

The distance between habitat patches and habitat area are two key components of spatially explicit ecological theories, such as metapopulation patch dynamics (Hanski 1998) and island biogeography (MacArthur and Wilson 1967). We suggest that parallels can be drawn for spatial temperature dimensions. For example, temperature availability relates to patch or island size, and temperature position relates to interpatch distances or island distance from a mainland. We use analogous ideas to explore the changing position and availability of temperature in generating community dynamics.

**Spatial availability.** Temperature spatial availability measures the geographic area of temperature (i.e., spatial extent) within species’ niche limits (figure 3). The effects of changing temperature spatial availability are dependent on scale. At a local scale, selection processes in communities determine change because temperature availability at a local scale can be viewed as an ecological resource (Magnunson et al. 1979, Roughgarden et al. 1981) for which individuals compete (Melville 2002). Therefore, individuals’ performance and population abundance can be affected by changing geographic area of thermal resources and habitat patches (Matthiopoulos et al. 2015). For example, fragmentation of primary forests leads to patches of matrix that can be many degrees warmer than contiguous forest (Senior et al. 2017), and the size of these warm patches is expected to influence the space use, behavior, and survival of populations of species dependent on forest habitats (Tuff et al. 2016).

At regional to global spatial scales, the available area of thermal niche limits species geographic range sizes and, therefore, a species’ global abundance as the two are strongly linked (Borregaard and Rahbek 2010). Limited availability of areas within the limits of the thermal niche leads to an increased probability of extinctions if populations shrink in geographic area and abundance (Purvis et al. 2000). As such, for a regional community, selection processes are important because the geographic extent of a preferred climate directly constrains species range extents, which deterministically affects species abundances. In the Pleistocene, a period of rapid temperature changes, species’ extinctions occurred at higher rates in regions in which climate refugia were not sufficiently large to maintain viable populations (e.g., Hofreiter and Stewart 2009, Nogues-Bravo et al. 2010). In this period, species with large body sizes were particularly sensitive to temperature availability change because of low density and large ranges (Lyons et al. 2004, Barnosky 2008). The polar bear (Ursus maritimus) provides a modern analogue of a species with increased risk of population extinctions due to spatial availability of temperature-dependent habitat. For this species, there is a predicted 68% reduction in summer habitat availability by the end of the century (Durner et al. 2009).

The spatial context of changes in temperature availability, rather than aggregated temperature data alone, provides additional insights to community responses to temperature change. As one example, if there is greater geographic area of temperature available, at either local or regional scales, we expect populations in environments of more optimal temperatures to increase in size, potentially increasing the total abundance of a community too (Cline et al. 2013). This prediction requires testing in model systems that disentangle the area of temperature availability from habitat size more generally. Figure 5 provides a visual representation of multiple dimensions of community responses, using species- and rank-abundance distributions and community temperature index. These predictions are in contrast with mean temperature change, which predicts that different species may...
Figure 4. Framework linking changing temperature dimensions to processes that drive community level responses. Applying a multidimensional perspective explicitly accounts for temperature changes in space and time that affect biodiversity that occurs through selection and movement pathways. Population-level effects arise when selection influences birth and growth rates, and movement influences immigration and emigration rates. Changes in population and demographic rates drive changes in the relative abundances and richness of regional and local communities. If we aggregate temperature over space and time, we miss the opportunity to attribute community responses to changes in specific temperature dimensions. We also lack the resolution to build a priori hypotheses with mechanistic links between changes in the physical environment and individuals’ responses.

A special case of spatial availability change is the emergence of a novel climate (Williams et al. 2007, Ordonez and Williams 2013). Novel climates can be considered new temperature regimes that were globally unavailable during the evolution of species in the regional fauna, as have occurred frequently throughout Earth’s geological history. Given that the size of available climates is limited by the size of the planet, an area of novel climate space must exclude or replace an area of present day climate space. Within novel climate space, no-analogue species assemblages are expected to form with corresponding shifts in species composition through time because of interspecific differences in climate tolerances (Williams and Jackson 2007).

The ecological implications of emerging novel climates are extremely difficult to anticipate, and ecological surprises are expected to unfold with new species interacting for the first time (Radeloff et al. 2015).

Spatial position. A change in temperature spatial position measures the geographic distance a specific temperature (thermal isocline) shifts after a climatic change (i.e., spatial distance in kilometers). Temperature velocity, the rate of spatial shift in thermal isoclines (figure 3; measured as the rate per km per decade), is a frequently used metric to measure changes in the spatial position (Loarie et al. 2009, Hamann et al. 2015). Therefore, the movement of individuals is an important mechanism underpinning community responses to this temperature dimension. For example, individual leatherback turtle (Dermochelys coriacea) movements tracked 15-degree-Celsius isotherms (McMahon and Hays 2006), and shifts in species ranges are also well documented and are increasingly predicted to track the position of preferred temperatures with warming (Devictor et al. 2008, 2012, Burrows et al. 2011, Pinsky et al. 2013, Hiddink et al. 2015, Sunday et al. 2015). Pinsky and colleagues (2013) demonstrated that range centroids have tracked the position of thermal isoclines with climate variability in the last 50 years in 360 marine taxa.

Identifying whether individuals move within or outside of their geographic range is also important when interpreting biodiversity change (Lenoir and Svenning 2015). Movement outside of a geographic range—that is, colonization of a new habitat—will result in species richness and species identity change for a receiving community (see the tail of rank-abundance distributions in figure 5g). These initially rare species may become more common over longer temporal scales because of increases in population variation.
Species traits cause variation in individuals’ response to the position dimension of temperature change, implicating the importance of trait-based metrics (e.g., Sunday et al. 2015). For example, at the community level the average species’ thermal affinity in a community, often summarized as a community temperature index, is expected to be sensitive to the spatial position dimension of temperature change (figure 4). With the establishment of warmer tolerant colonists, we expect the community temperature index to increase and the distribution of species’ thermal affinities to become increasingly right skewed (figure 5h; ter Hofstede et al. 2010, Bates et al. 2014b). This is in contrast to predictions from changes in temperature availability, according to which only increased variation but no mean change in species thermal affinities is expected (figure 5d cf. 5h). Furthermore, species with high mobility have better capacity to keep pace with spatial shifts in isotherms (Sunday et al. 2015).

Can a multidimensional perspective help disentangle pathways of community change?
Here, we illustrate how two important biodiversity responses to environmental change—community homogenization and local richness change (Dornelas et al. 2014, McGill et al. 2015)—are driven by fundamentally different pathways, rarely disentangled in the literature. We further discuss

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Figure 5. Schematic diagram linking changes in spatial availability and position of temperature with community responses using different biodiversity metrics. In panels (a) and (e), the polygons represent individuals, and the purple (dark) regions represent areas of temperature optimality. In panels (b) and (f), species abundance distributions show a right shift in central tendency with increased abundance, an increase in height with increasing richness. In panels (c) and (g), rank abundance distributions show an increasing tail with higher richness and a shift right with increasing total abundance. Note the long tail for temperature position change. The underlying bars represent species abundance change between time points but maintained at the rank on the x-axis in time point 1. The colors refer to thermal traits (blue (dark) is cool affinity; yellow (light) is warm affinity). In panels (d) and (h), changes in the distribution of species' thermal affinities between time points is the mean of this distribution. Note the same mean for changes in spatial availability and the long tail and changing mean for spatial position.
the potential for disaggregation of communities through time—a community response to climate change that can only be detected by studying the effects of temporal rather than spatial dimensions of temperature change.

**Disentangling community homogenization.** Communities are generally becoming more similar in composition over time or space—a process called homogenization—often quantified as reduced beta diversity (Juraisinski and Kreiling 2007, Baiser et al. 2012, Davey et al. 2012, Savage and Vellend 2014, Maguran et al. 2015, but see Avolio et al. 2015). Identifying the specific dimension of temperature change leading to community homogenization has potential to help estimate the distinct effects of community abundance shifts (selection processes) versus species range shifts or expansions (movement processes, figure 4).

Selection processes cause community homogenization through time when a subset of species in a local assemblage systematically increase or decrease in abundance. For example, across many local stream-fish assemblages in France, temporal changes in community composition were related to losses of individuals—and, therefore, population declines and relative abundance changes—which has favored an increasingly similar set of species since the 1980s (Kuczyński et al. 2017). In this case, community homogenization was linked to selection processes and the timing dimension of temperature change emerged as an important predictor.

Through space, homogenization will occur when the same subset of species increasingly occupy many local communities across a region. For this to occur, species distributions must expand or contract by movement within a region. If range-shifting species display coordinated expansions or contractions across communities spatial homogenization is expected. This form of homogenization is often driven by movement of generalist species undergoing a range expansion (Davey et al. 2012). In cases in which movement processes drive homogenization, species richness will also increase (La Sorte 2006, Davey et al. 2012, Batt et al. 2017). However, the role of temperature position driving homogenization and richness is often unexplored (Davey et al. 2012, Savage and Vellend 2014). A pressing debate is the simultaneous stability of richness with ongoing biotic homogenization of communities (i.e., Maguran et al. 2015 cf. Savage and Vellend 2014) and reordering of communities (e.g., Jones et al. 2017). This debate will benefit if the multiple dimensions of temperature change are identified in studies testing theory.

**Drivers of local richness change.** Local species richness change is commonly attributed to the magnitude dimension of temperature change (Menéndez et al. 2006, Britton et al. 2009, Davey et al. 2012, Tayleur et al. 2015). However, increases in richness must occur because of species movements (i.e., local colonizations), whereas decreases can be due to selection (i.e., decline in situ) or movement (i.e., movement away from site). The rate that species move into a warmer (increased temperature magnitude) environment is determined in part by the position dimension of temperature change, but this could trade-off with greater species losses as warmer temperatures exceed species tolerance limits. Few studies, if any, have acknowledged the interaction between these two processes in driving species richness change.

**Community temporal disaggregation from changes in temporal position and availability of temperature.** Assessing community level responses to changes in timing dimension of temperature change has revealed a distinct fingerprint on community composition, independent of changes in mean temperature. Specifically, Thomsen and colleagues (2015) found that with changing temporal position and availability of temperature, growing seasons are longer causing earlier springs and longer summer periods (these two dimensions were correlated in this study). These temperature changes affect the seasonal timing of peak abundances in the warm and cool affinity species differently. Warm species had later peak abundances, whereas cool species had earlier peak abundances (figure 6a). If communities have varying degrees of thermal niche complementarity—that is, high variation in thermal performance optima (figure 6b)—the temporal
synchrony of species may break down. The extent of asynchrony will depend on species thermal trait distributions across the community (figure 6c). Long-term and high-resolution community time series are necessary to estimate the effects of temporal dimensions of temperature change, which may explain our gaps in understanding of community responses to this temperature dimension (Magurran et al. 2010, Thomsen et al. 2015).

Conclusions

Similar ecological patterns can arise from different combinations of processes (Lawton 1999). We suggest this is also true when measuring community responses to environmental change. To understand biodiversity change on a warming Earth, we must link spatial and temporal structure of temperature and community change—short of this, misattribution of the climatic processes responsible for biodiversity change may occur. Similar community changes can occur through both selection and movement pathways; biodiversity forecasts and management decisions may depend on the relative importance of each. Characterizing the dimensionality of how temperature is changing at scales relevant for biodiversity processes will require closer collaboration between physical scientists and ecologists. This will hopefully lead to an attribution of temperature’s effects beyond average temperature change. We demonstrate a need to build these mechanistic connections into how physical regimes affect biodiversity change, being explicit in space and time.

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Supplemental material

Supplementary data are available at BIOSCI online.

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