Going against the flow: potential mechanisms for unexpected downslope range shifts in a warming climate

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However, most of the studies that reported expected range shifts towards higher elevations have detected species moving towards lower elevations as well. Table 1 provides an illustrative, non-comprehensive survey of such studies from the recent years: in summary they demonstrate that ca 65% of the species have shifted their mid-range positions upslope, 10% have not changed their mid-range positions, and 25% have shifted their mid-range positions downslope (Table 1). In addition, according to a global review of the literature published until the beginning of the 21st century (Parmesan and Yohe 2003) ca 20% of the species have adjusted their ranges towards lower elevations and/or southern latitudes. Hence, a considerable fraction of the investigated species has shown range shifts that are inconsistent with the forecasted effects of climate warming. These downslope movements seem very unlikely to occur as a direct consequence of rising temperatures, but the potential mechanisms involved have received little attention.

Stochastic fluctuations in the positions of individuals, or populations, together with measurement errors, represent one such potential “mechanism”. However, many, though not all of the studies reporting downslope shifts have explicitly tested the observed changes in single species’ ranges for significant deviation from random fluctuations. For plants, significant downslope shifts have been reported for 5 of 46 species displaying significant mid-range shifts between the periods 1905–1985 and 1986–2005 (Table S2...
Table 1. Studies that have documented changes of the mid-range positions (i.e. mean, median or optimum) of species' elevational distributions in relation to current climate change. Only studies comparing at least two inventories are listed here. This list is informal and not intended to be exhaustive.

<table>
<thead>
<tr>
<th>Species groups</th>
<th>Sampling periods</th>
<th>Geographic locations</th>
<th>Number of species</th>
<th>Percentage of species</th>
<th>Mean shift of the central position</th>
<th>Author references</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moth</td>
<td>1965 and 2007</td>
<td>Mount Kinabalu, Borneo</td>
<td>102</td>
<td>59%</td>
<td>+68 m between surveys</td>
<td>Chen et al. (2009)</td>
</tr>
<tr>
<td>Animal</td>
<td>1993–2003</td>
<td>Tsaratanana mountains, Madagascar</td>
<td>30</td>
<td>73%</td>
<td>+6.5 m per decade</td>
<td>Raxworthy et al. (2008)</td>
</tr>
<tr>
<td>Bird</td>
<td>1970s and 2000s</td>
<td>Alps, France</td>
<td>41</td>
<td>10%</td>
<td>–2 m between periods</td>
<td>Archaux (2004)</td>
</tr>
</tbody>
</table>

Climate warming may cause temporary downslope range shifts by facilitating or releasing species from biotic limits, leading to competitive release or facilitation, which can cause temporary downslope range shifts. One potential mechanism is habitat-mediated changes in species distributions and competition, which have already been shown to cause downslope range shifts (Harrington and Kettler 1995, Archaux 2004). In addition, however, the observed downslope shifts may also result from changes in species interactions. Indeed, the relevant question is: which proportion of species would we expect to shift their mid-range position downslope by chance alone? (i.e. due to random population fluctuations, data idiosyncrasies, and observer errors) despite an average upslope trend of climate warming? Data for testing the occurrence of significant downslope movement are scarce, and the relevant question is: which proportion of species would we expect to shift their mid-range position downslope by chance alone? (i.e. due to random population fluctuations, data idiosyncrasies, and observer errors) despite an average upslope trend of climate warming?
null expectation of the proportion of downslope mid-range positions in species mid-range elevation within a single statistical population. The gray filling illustrates the expected proportion of random downslope shifts in mid-range positions despite a warming-driven upslope trend of z m (<50%). The vertical broken line displays the average upslope trend of z m.

The stress-gradient hypothesis gives the key to understanding potential downslope range shifts despite climate warming. We start outlining our conceptual model by restricting the temperature-elevation-stress relationship to elevational ranges where the macroclimate does not present drought-induced stress gradients that complicate the effect of the elevational gradient (Callaway 1998, Callaway et al. 2002). Thus, this concept will apply to many mountainous regions, notably in moist temperate and tropical areas. As a corollary, many species in such systems will likely be characterized by realized climatic niches being smaller than fundamental ones (Vetaas 2002) in particular towards lower elevations (due to biotic interactions). For example, Vetaas (2002) has suggested that competition plays an important role in constraining the climatic ranges of four Himalayan Rhododendron species in their native habitat as compared to their climatic ranges in ornamental gardens and arboreta, three of which being able to grow there under a far wider range of climatic conditions (generally warmer). Therefore, any mechanism that alleviates competitive exclusion is likely to induce changes in species realised distributions.

Secondly, the importance of competition in structuring communities is likely reduced by increased levels of disturbance (Dayton 1971, Connell 1978, Huston 1979, McAuliffe 1984, Brooker and Callaghan 1998, Brooker 2006). Indeed, the increasing frequency of climatic extremes, indicated by changes in the interannual variability around mean values of climate parameters, have already been reported to influence species range margins (Zimmermann et al. 2009). The interplay between the idea of disturbance-related release from competition and the above-mentioned stress-gradient hypothesis gives the key to understanding potential downslope range shifts despite climate warming.
from their realised niche), but differing in the way they fill their potential distribution area (as defined from their fundamental ecophysiological niche) (Fig. 2c, d). Hereafter, we will consistently refer to the “potential distribution” as the range corresponding to the species’ fundamental ecophysiological niche, and to the “realised distribution”
as the range corresponding to the species’ realised niche. The first illustrative species is strongly limited at the lower margin of its elevational distribution by competitors from below (low “realisation” of the potential distribution), whereas the second one is much less limited by biotic interactions (high “realisation” of the potential distribution). It is difficult to infer the physiological climatic requirements of these two species from their actual range since their elevational distributions are identical. Nevertheless, they might respond differently to climate warming because of this distinct “realisation” of their ecophysiological range.

Let us now introduce the effect of climate warming in our conceptual model (red colours in Fig. 2). As argued above, warming-induced disturbances are likely to transiently reduce the importance of competition along the elevation gradient (Fig. 2b). Simultaneously, climate warming is likely to shift species’ potential distribution along the elevation gradient towards higher elevations (Fig. 2c, d). However, the transient reduction in the importance of competition at the lower margin of a species’ elevational distribution is likely to particularly benefit species that currently have a greater part of their potential distribution unfilled because of competition. For such species, climate warming allows range expansion towards lower elevations from which they had hitherto been competitively excluded (Fig. 2c). In contrast, species which are currently little limited by competition and largely fill their potential distribution areas along the elevation gradient will not be able to move downwards due to the upward shift of habitats climatically suitable to them (Fig. 2d). Figure 2e gives a full account of the climatically driven range shifts along the elevation gradient conceivable under this model.

To sum up, species with a low “realisation” of their potential distribution areas along the elevation gradient are especially good candidates for downslope shifts, if they additionally have good dispersal abilities and a wide fundamental climatic niche. For example, Vetaas (2002) has suggested that *Rhododendron* species with their lower margins hitherto set by competition can grow under warmer climatic conditions, i.e. expand towards lower elevations. Additionally, species mid-range shifts might result not only from expansions at range margins alone, but also from changes in species local abundance (Fig. 2e). For example, species with low competitive abilities, but a wide potential distribution along the elevation gradient associated with a high degree of plasticity (Fig. 2c) might be already present downwards as remnant populations living at environmental extreme sites where the competitive species are excluded by abiotic factors (Eriksson 2000). In this latter case, downslope mid-range shifts might simply result from an increasing abundance of the species with these outlier populations acting as “expansion foci”. Finally, we note that downslope range shifts under this model would be likely to be temporary, as the importance of competition may become reasserted if climate change slows down or come to a halt, while species will eventually be forced upwards in elevation if climate change continues and conditions at lower elevations shift beyond the fundamental climatic niche of the species.

**Habitat modification as an alternative mechanism causing downslope range shifts**

Species may also shift downslope as a direct consequence of habitat modification, with or without involving competitive release, either following natural disturbances (windthrows, fires, and avalanches), human-induced disturbances or permanent habitat changes (recreational activities, land use changes, and management practices), or due to other local changes in habitat suitability. For example, in the Swiss Central Alps, Hättenschwiler and Körner (1995) have suggested that the cessation of forest cattle grazing and the high level of nitrogen deposition may have led to denser and more exuberant ground vegetation, thereby enhancing the replacement of *Pinus sylvestris* seedling populations by those of *P. cembra* below the present lower margin of *P. cembra* adult trees. Similarly, Archaux (2004) has suggested that the increase of conifer areas at the expense of broad-leaved trees due to changes in forest management might cause both coniferous- and deciduous-forest bird species to shift their mean elevation downwards. We note that habitat modification in conjunction with climate warming may explain upslope range shifts as well. As an illustration, in the Swiss Alps, Gehrig-Fasel et al. (2007) have reported that >90% of upslope shifts in the local tree line are due to ingrowth and the filling of gaps indicating that land use is the primary driver over climate warming in many instances, although the two drivers may also act in combination. In a conceptual model involving climate change and herbivory pressure, Cairns and Moen (2004) have highlighted a potential pathway for the interaction of both climate change and herbivore pressure on tree line fluctuations leading to upslope migration, a stationary state, or retrogression of tree lines. Hence, habitat modification has often been claimed to be an important driver of elevational range shifts, acting in concert with climate warming or even outweighing it (Hättenschwiler and Körner 1995, Archaux 2004, Cairns and Moen 2004, Gehrig-Fasel et al. 2007, Vittoz et al. 2009).

Nevertheless, habitat modification may well involve increased disturbance levels and thereby cause release from competition independently of climate warming, but with similar effects on species’ elevational distributions. Human-induced disturbances are likely to be more frequent in lowland areas, however, given a generally increasing degree of anthropogenic habitat modification towards lower elevations of most mountainous settings (Nogués-Bravo et al. 2008). Consequently, the reduction in the importance of competition due to disturbances should be more important there, allowing range expansions of the realised distribution of species towards lower elevations as long as climatically suitable sites are available down below their lower margins. Therefore, species that fill only part of their potential distribution areas along the elevation gradient (Fig. 2c) are more likely to shift downwards in response to habitat modification alone, especially if their potential distribution areas do not shift upwards. For example, it has been suggested that unplanned vegetation destruction (burning and grazing), removing the pressure from competitive dominants from below, has enabled alpine and subalpine species in New Zealand to increase their elevational distributions downwards (Halloy and Mark 2003).
Towards a unified view of climate warming and habitat modification effects on downslope range shifts

Let us now consider a more unified point of view integrating the effects of climate warming and habitat modification. On the one hand, habitat modification may act in concert with climate warming to cause a reduction in the importance of competition at the lower margin of a species’ elevational distribution and to allow a potential downslope shift in its mid-range position due to potential expansions at the competitively-released margin (Fig. 2c). This is especially true for species that fill only part of their potential distribution areas along the elevation gradient (Fig. 2c). On the other hand, habitat modification may restrict upslope range expansions due to climate warming or even cause regression patterns at the higher margin of a species’ elevational distribution (Fig. 2e). For example, species may partially fail to colonise new climatically suitable areas at the higher margins of their elevational distributions if they are constrained by habitat fragmentation (Honnay et al. 2002). Additionally, other non-climatic factors such as restricted space towards mountain tops (Colwell et al. 2008), limited dispersal ability (Bossuyt et al. 1999, Hermó et al. 1999, Svenning and Skov 2006), and/or edaphic constraints may restrict, or at least delay warming-induced upslope shifts. Finally, natural habitat modification may further delay warming-induced upslope shifts through unexpected patterns of regression at the higher margins of species’ elevational distributions (Fig. 2e). For example, Cannone et al. (2007) have suggested that warming-induced permafrost degradations at high elevations may trigger habitat disturbances, in the form of debris flow and landslides, causing unexpected patterns of regression in vegetation coverage above 2500 m. This constitutes another migration barrier that restricts upslope migrations to disturbance-adapted species (Cannone et al. 2007).

Such limitations to upslope migrations are coherent with observations in different species groups: extinctions at the lower margins of species distributions have been reported to be more common than colonisations at the higher margins (Wilson et al. 2005, 2007, Kelly and Goulden 2008, Moritz et al. 2008, Lenoir et al. 2009). This may sometimes result in no upslope migration, but rather in local changes in species abundance over time (Wilson and Nilsson 2009). In such cases, mortality-induced shifts may take place more rapidly than do recolonisation-induced shifts associated with both migration and establishment processes (Davis 1989). The resulting pattern is a transient “lean” upslope (Breshears et al. 2008). The few establishments of a given species towards higher elevations may fail to compensate for the losses at lower elevations leading to transient declines in species richness or biotic attrition not only at lower elevations (Colwell et al. 2008), but across the whole elevation gradient (Fig. 2 in Wilson et al. 2007)). This configuration is transient and, again, is likely to open a “window of opportunity” for highly vagile and plastic species that might shift either upslope or downslope to fill the gaps initiated either by climate warming or habitat modification. Such a process leaves biological communities with reduced numbers of species, and dominated by more mobile and opportunistic species (Warren et al. 2001). Increased frequency of windthrows across central European forests during the last few decades (Usbeck et al. in press) is one example of disturbances that is likely to produce “windows of opportunities” for vagile species with a high degree of plasticity. Thus, habitat modifications strongly interact with climate warming and contribute to bias competitive release even further towards lower elevations, making downslope range shifts of some species more likely than with climate warming alone.

Other plausible causes of downslope range shifts

Of course, downslope range shifts could be driven by changes in other aspects of climate than mean temperature, e.g. precipitation regime, snow cover duration, water balance, or seasonality in climate parameters. These complex aspects of climate variability may heavily influence species range margins (Zimmermann et al. 2009), and thus more complex environment-competition interactions are likely to cause unexpected range shifts in response to climate warming. At high elevations, for example, warmer temperatures may decrease the winter snow cover duration (Beniston 2005), and thus may cause frost damage at the higher margin of a species’ elevational distribution, which in turn may alleviate the competitive effect of this species on other ones potentially migrating towards lower elevations (Fig. 2c). Consequently, the competitive control that this species exerts on the distributions of the species above will likely become less tight. Additionally, the influx towards areas vacated by upslope shifting competitive species is likely to occur both from above and below. This should result in some species shifting upslope and others shifting downslope, and others even expanding towards both sides without changing in their mid-range position (Fig. 2c). Therefore, complex environment-competition interactions may also cause downslope range shifts, but temporarily before other stronger competitors invade.

A case study from French mountain forests: proportion of random downslope movements

According to our null expectation, the proportion of random downslope mid-range shifts despite an average upslope trend of z m is dependent on the range of random variation among shifts (Fig. 1). We used data from a previous study focusing on the shifts in the elevational position of plant species’ maximum probability of presence (optimum) (Lenoir et al. 2008) to estimate this range of random variation among shifts and then assess the proportion of random downslope mid-range shifts despite an average upslope trend of z m. That study found an average upslope trend of 65 m, among 171 plant species, between a 1905–1985 dataset and a carefully matched 1986–2005 dataset for French mountain forests. Each dataset comprised 3991 surveys. To estimate the range of random variation among shifts, we constructed two random
datasets, each built by randomly drawing (with replacement) 3991 forest surveys from the 1905–1985 dataset described by Lenoir et al. (2008). Because the increasing trend in mean annual temperature and the warmest records have mostly occurred since the late 1980s (Jones et al. 2001), we chose to draw our two random datasets from the first period to avoid potential strong differences in climatic conditions between the two randomly-drawn datasets. Indeed, it was important that the two bootstrap samples represented an identical range of environmental conditions. We then used the same analytical method as described in Lenoir et al. (2008) to compute species elevation optimum for each of the 171 studied species in each of the two bootstrap samples. The estimated elevational optimum position was bounded between the lowest and the highest elevations in the 1905–1985 dataset. However, instead of computing the difference in each species optimum elevation between two different periods, we here computed the difference in each species optimum elevation between two bootstrap samples from the same period to get the distribution of differences in optima expected from stochasticity alone. We repeated this procedure 1000 times using the “boot” library (Canty and Ripley 2008) in R (R Development Core Team 2009). Averaging across the 1000 iterations, we found a mean difference in species optimum elevation of $-0.4\ m$ (standard deviation: $10.5\ m$) and a confidence interval for mean at 95% ranging from $-14.2\ m$ (standard deviation: $10.7\ m$) to $13.3\ m$ (standard deviation: $10.6\ m$). Finally, to assess the proportion of downslope mid-range shifts expected at random despite an average upslope trend of 65 m in French mountain forest plants, we simply repositioned, for each of the 1000 iterations, the distribution of random differences in optima expected from stochasticity alone. We repeated this procedure 1000 times using the “boot” library (Canty and Ripley 2008) in R (R Development Core Team 2009). Averaging across the 1000 iterations, we found a mean difference in species optimum elevation of $-0.4\ m$ (standard deviation: $10.5\ m$) and a confidence interval for mean at 95% ranging from $-14.2\ m$ (standard deviation: $10.7\ m$) to $13.3\ m$ (standard deviation: $10.6\ m$). Finally, to assess the proportion of downslope mid-range shifts expected at random despite an average upslope trend of 65 m in French mountain forest plants, we simply repositioned, for each of the 1000 iterations, the distribution of random differences in optima $65\ m$ upwards, i.e. to the right, by adding $65\ m$ to the location of each shift (Fig. 3a). We then estimated the expected proportion of random downslope shifts, despite an average upslope trend of 65 m, as the proportion of shifts in the left tail of the distribution, truncated at $0\ m$ shift (see solid dark gray bars in Fig. 3a). Across the 1000 iterations (Fig. 3b), we found a proportion of 16% (standard deviation: 4%) of species expected to have optima below the 1905–1985 elevational mean, by chance alone, which is about half the proportion of downslope shifts we originally found (30%) between 1905–1985 and 1986–2005 (Lenoir et al. 2008). Thus, the number of species found to move downwards along the elevation gradient is approximately twice as high as expected by chance under the observed general upward trend. We note that this estimated proportion of species “going against the flow” (14%) is higher than the number of species with significant downslope shifts (5/171) found when individual species optimum elevation were tested for significant difference from a constant species optimum elevation expectation (Table S2 in Lenoir et al. (2008)). However, this probability reflects the conservative nature of the test we used for testing differences in individual species optima along the elevation gradient (Lenoir et al. 2008).

The five forest plant species displaying significant individual downslope shifts despite an average upslope trend of 65 m are *Clinopodium vulgare*, *Dryopteris dilatata*, *Quercus pubescens*, *Rubus fruticosus* and *Saxifraga cuneifolia*. Two of these species have seeds dispersed by birds (*Quercus pubescens* and *Rubus fruticosus*), the spores of *Dryopteris dilatata* are tiny and hence easily going with the wind, while *Clinopodium vulgare* is dispersed by epizoochory (Rameau et al. 1993). Hence, four of these five species have efficient dispersal mechanisms, most likely an important trait for allowing downslope range shifts despite climate warming.

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**Figure 3.** Distribution of (a) observed and random differences in optima along the elevation gradient for 171 plant species using published data from French mountain forests (Lenoir et al. 2008), the distribution of random differences in optima representing a single illustrative case of the 1000 bootstrap iterations, and (b) distribution of the proportion of random downslope mid-range shifts despite an average upslope trend of 65 m for the 1000 bootstrap iterations. Solid light gray bars represent the observed shifts in species optimum elevation between 1905–1985 and 1986–2005. Unfilled bars represent the distribution of random shifts in species optimum, computed by comparing two bootstrap samples drawn from the 1905–1985 dataset, after repositioning this distribution 65 m upslope, i.e. to the right (see text for details). Solid dark gray bars represent the proportion of downslope mid-range shifts expected at random, despite an average upslope trend of 65 m. The vertical dark broken line displays the average upslope trend of 65 m. The vertical white broken line displays the proportion of random downslope mid-range shifts despite an average upslope trend of 65 m for the single illustrative case of the 1000 bootstrap iterations.
Although *Saxifraga cuneifolia* has low dispersal abilities (barochory) and a distribution mainly restricted between 1500 and 2000 m, it also occurs sporadically in the lowlands, reaching down to 300 m in French mountain forests (Rameau et al. 1993). Thus, *Saxifraga cuneifolia* might represent a species strongly limited by competitors from below (Fig. 2c). Additionally, *Rubus fruticosus*, and to a lesser extent *Clinopodium vulgare*, and *Quercus pubescens* seedlings, are highly reactive to canopy opening, i.e. positively influenced by disturbance. While we do not intend to validate our conceptual model with these examples, we note that these five downslope shifting species have a series of traits making them likely to respond to climate change in a way outlined by our model. However, a much more thorough empirical testing of this model is clearly needed.

**Conclusion**

We suggest that downslope range shifts of species may constitute an indirect biotic response to both climate warming and habitat modification rather than representing just random effects due to stochastic fluctuations in population distributions or observer errors. The concept presented here should become part of a general framework for future studies of changes in species distributions in response to climate warming. In our conceptual model, we assume, on a timescale too short for adaptive change, that downslope shifts primarily occur for species that are strongly limited by competition at their lower elevation range margin, and therefore have a realised distribution that do not fill their potential distribution areas almost completely along the elevation gradient. To test this hypothesis, one could select two sets of species: one set of species that have significantly shifted downslope and another set of species that have significantly shifted upslope. One could then compare their realised distribution in their natural habitats with their potential distribution areas additionally assessed by experiments (common garden or ecotone). In such an experiment, we would expect larger differences between the realised and the potential distributions along the elevation gradient for the set of species that have significantly shifted downslope. Although downslope range shifts, particularly where solely driven by warming-induced competitive release, should be only transient, we underpin the necessity to take the hitherto neglected downslope range shifts of species more explicitly into consideration when making predictions of the effects of future climate change scenarios on species distributions.

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