Complex elevational shifts in a tropical lowland moth community following a decade of climate change

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Abstract

Aim: Climate change is driving many species towards higher latitudes and higher elevations. However, empirical studies documenting these changes have largely focused on presence/absence based range shifts in temperate regions. Studies in lowland tropical ecosystems that control for detection probabilities are especially lacking.

Location: Hong Kong SAR, China.

Methods: By analysing a 15-year trapping dataset of geometrid moths along a lowland elevation gradient (0–600 m) in Hong Kong, we used multiple metrics and approaches to estimate occurrence shifts with elevation, changes to range limits, and community-wide shifts in elevation. Our approaches used Bayesian occupancy models to account for false absences, which may bias naïve measures of range shifts over time.

Results: Over the study period, we detected only subtle changes in forest cover but a notable increase in maximum temperatures (~0.5°C per decade) and extreme weather events. Of geometrid moths, one out of the 123 examined species exhibited increased occurrence probabilities across all elevations after accounting for uncertainty in detection; all other species exhibited no significant change in occupancy with elevation. However, at least two species became newly established in Hong Kong over the decade, and 32 species showed significant elevational shifts in lower or upper range limits. At the community level, geometrid moths showed a noticeable upslope shift at all but one of the examined sites.

Main conclusions: The complex patterns observed highlight the difficulties and limitations in detecting climate change impacts on diverse tropical communities. Our conservative results indicate early responses of tropical species over a relatively short timespan to a decade of environmental change, and the necessity of long-term monitoring for providing insights into the management and conservation of vulnerable species.
1 | INTRODUCTION

Climate change is affecting biodiversity across the globe (Araújo & Rahbek, 2006; García, Cabeza, Rahbek, & Araújo, 2014). Distribution shifts towards higher altitudes and latitudes have been observed in many taxa (Hickling, Roy, Hill, Fox, & Thomas, 2006; Pecl et al., 2017). Despite the large number of resurvey and monitoring studies that have compared distributions between historic and current communities (Chen, Hill, Ohlemüller, Roy, & Thomas, 2011; Lenoir & Svenning, 2015), much variation exists in range shift studies exclusively examining shift patterns; variation in methods particularly complicates the attribution and magnitude of range shifts detected across species groups, local climates and habitats (Guo, Lenoir, & Bonebrake, 2018).

Directly calculated distribution limits or range centre shifts often suffer from detectability issues, as imperfect detection and false absences can cause apparent range shifts even when no true shift occurs (Tingley & Beissinger, 2009). Variation in sampling effort across time (documentation of historic sampling effort is usually limited or unavailable) is especially important to consider in comparing distributions in the context of environmental change (Bates et al., 2014; Brown et al., 2016; Tingley & Beissinger, 2009). Furthermore, short-term studies are more likely to track population fluctuation signals, rather than real climate change effects, especially species with high population variability such as invertebrates and annual plants (McCain, Szewczyk, & Bracy Knight, 2016). Under such circumstances, the use of occupancy modelling to correct for imperfect detection when examining range shifts has gained popularity (Moritz et al., 2008; Rowe et al., 2015; Tingley, Koo, Moritz, Rush, & Beissinger, 2012). Occupancy modelling utilizes hierarchical logistic regression models to separate observation processes from occupancy processes to control detection probability variation commonly encountered in ecological sampling (MacKenzie et al., 2002; Tingley & Beissinger, 2009). Partitioning variation caused by observation allows for a more accurate analysis of ecological variation and has been shown to significantly reduce false absence rates compared to traditional direct comparisons (Tingley et al., 2012).

Geographical biases also occur in range shift research in that most studies have focused on species in temperate regions where historical records are often more readily available (Forister et al., 2010; Kelly & Goulden, 2008; La Sorte & Thompson, 2007; Larsen, 2012; Menéndez, González-Megías, Jay-Robert, & Marquéz-Ferrando, 2014; Moritz et al., 2008; Rubidge, Monahan, Parra, Cameron, & Brashares, 2011; Tingley et al., 2012), while studies in the tropics are generally rare (Chen, Hill, Ohlemüller, et al., 2011; Freeman & Freeman, 2014; Lenoir & Svenning, 2015). The few tropical studies that have been conducted have largely focused on highland montane areas (elevation>1,000 m a.s.l.; Peh, 2007; Raxworthy et al., 2008; Chen et al., 2009; Feeley et al., 2011; Forero-Medina, Terborgh, Socolar, & Pimm, 2011; Jump, Huang, & Chou, 2012; Freeman & Freeman, 2014; see also Figure 1), while lowland tropical habitats (elevation <1,000 m) hold the highest levels of alpha biodiversity among all terrestrial habitats (Brooks et al., 2006; Pimm et al., 2014). Climate change is projected to have large impacts on tropical species (Bonebrake, 2013; Deutsch et al., 2008; Khaliq, Hof, Prinzinger, Böhning-Gaese, & Pfenninger, 2014), particularly in lowland areas where topographic variation is limited and more variable climates for buffering warming impacts may be unavailable (Bonebrake & Deutsch, 2012; Bonebrake & Mastrandrea, 2010; Colwell, Brehm, Cardelús, Gilman, & Longino, 2008; Malhi et al., 2008). Moreover, tropical lowland ecosystems are also under great threat of anthropogenic disturbance and habitat loss (Guo et al., 2018; Nogués-Bravo, Araújo, Romdal, & Rahbek, 2008).

Here, using an annual continuous trapping dataset of geometrid moths, we investigated species range shifts along a lowland elevational gradient in Hong Kong, at the northern edge of the tropics. Moths of the family Geometridae are relatively well studied both biologically and taxonomically (Holloway, 1993; 1996; 1997; Kendrick, 2002). They are also ecologically important in natural systems (Schowalter, Hargrove, & Crossley, 1986) as well as sensitive to climate change (Chen et al., 2009) and habitat disturbance (Beck, Schulze, Linsenmair, & Fiedler, 2002), making Geometridae ideal indicator taxa for environmental change studies. By examining climate and forest cover change from 2000 to 2014 and controlling...
for detection probabilities with occupancy models, we tested (a) whether occurrence probabilities across elevation shifted for each species during our study period, (b) whether each species showed shifts of upper or lower elevation boundaries while accounting for false absences, and (c) whether the community of geometrid moths at re-sampled sites showed systematic elevational trends over time.

2 | METHODS

2.1 | Study system

Continuous sampling of geometrid moths in Hong Kong started in 1994 and continued through 2014 by one of the authors using Robinson light traps (Fry & Waring, 1996), often in conjunction with white ground sheets. Moths were sampled in different months throughout the year along an elevational gradient from sea level up to the highest hill (Tai Mo Shan, 957 m) in Hong Kong SAR. (Figure 2). Tai Mo Shan is largely covered by subtropical evergreen forest which has regenerated since the 1940s. The majority of Tai Mo Shan’s forest lies within protected areas established in the 1970s and 1980s (Dudgeon & Corlett, 2004). In total, 556 moth samplings events were opportunistically made between 1994 and 2014, with a maximum of 14 sampling events per year. During each sampling event, light traps were set from dusk to midnight (but sometimes for fewer hours) and species checklists were recorded.

2.2 | Local climate and habitat change

To quantify local climate change during the study period, we downloaded monthly averaged daily mean temperature, monthly averaged daily maximum temperature, monthly averaged daily minimum temperature and accumulated monthly rainfall from two weather stations at the top and bottom of our study site from the Hong Kong Observatory (Figure 2; https://www.hko.gov.hk/cis/climat_e.htm). The elevational distance for our two weather stations is approximately 940 m apart, and the geometric distance between these two weather stations is approximately 4.5 km. We also calculated the annual number of extreme hot days (daily Tmax>33°C), annual number of extreme hot nights (daily Tmin>28°C), annual number of extreme cold days (daily Tmin<12°C), annual number of frost days (daily Tmin<0°C), annual number of extreme cold nights (daily Tmax<12°C), and annual number of extreme cold nights (daily Tmax<12°C).
number of dry days (daily rainfall = 0 mm) and annual number of flood days (daily rainfall > 100 mm) for Hong Kong from the same source. We were only able to find climate data from 1997 to 2014. To quantify the changes of these climatic variables over the study period, we applied simple linear regressions with year as an explanatory variable and the yearly climatic variables as response. Frost days were very rare during the sampling period, so we did not analyse these data statistically, but given the important ecological influence of frost on Lepidoptera (Wang et al., 2016), we qualitatively compared frost days over time.

We used a high-resolution (30 m by 30 m) forest cover change dataset as an indicator of land use change over the last decade (Hansen et al., 2013). We generated forest cover loss and gain profiles across elevational bands at 100 m intervals for our study area by overlaying these forest data onto a fine-scale digital elevation map (SRTM Arc-Second Global, 30 metres in resolution, https://lta.cr.usgs.gov/SRTM1Arc). We also created 90 m by 90 m, 180 m by 180 m and 360 m by 360 m buffer zones around our 24 sampling sites and calculated the adjacent forest cover loss and gain of each site using the same dataset.

2.3 | occupancy models and occurrence probability change

Using occupancy modelling in a Bayesian framework, we estimated and tested for changes in occurrence probability from 2000 to 2014 across elevation. Because sampling was highly uneven at sites across years, we undertook a series of data partitioning steps prior to analysis. We thus subset the moth dataset to two time periods: 2000–2004 (historic) and 2010–2014 (current). We excluded data from before 2004 and from 2005 to 2009, when sampling was sparse, and focused on the two periods of greatest data concentration to maximize inferential power. Additionally, by averaging over 5 years of sampling, our study design minimizes bias from random population fluctuations that can bias range shifts (McCain et al., 2016).

Within each major time period (i.e., 2000–2004 and 2010–2014), detections and non-detections of moths were assumed to represent imperfect samples of the true occurrence state. Sampling effort also varied across sites, where some sites were visited a few times per year, and other sites were visited up to 14 times per year (Supporting information Table S1). To avoid over-fitting of detectability to high-frequency sampling sites, we randomly selected 15 sampling events per site per sampling period to include in the analysis for all sites that had >15 visits per time period.

Additionally, because only a small number of sites (six out of 24) were visited in both time periods, we applied an "unpaired site" model parameterization as described by Tingley and Beissinger (2009), which looks for changes in occupancy across a defining gradient (here, elevation), rather than explicit colonization–extinction at paired sites. We only included sites visited more than once, resulting in 16 historical sites (92 sampling events) and 14 current sites (98 sampling events). Despite the fact that the geographic distances between sampling sites were greater in the historic sampling, the elevational distances are similar (Two-tailed T test, historic 177.4 m, current 167.8 m, $p = 0.6$). Detailed site information is shown in Figure 2 and Supporting information Table S1.

We only analysed species that occupied at least 10% of both historic and current sites, as occupancy models cannot reliably be fit in data-sparse situations. Consequently, we focused the analysis only on species for which we were confident that we could successfully fit an occupancy model, resulting in 123 species in total (Supporting information Table S3).

As all moths were collected by the same person with identical trap design, for each species, we assumed a constant detectability between the two sampling periods. While peak flying time for adult moths in Hong Kong is either at the beginning of the wet season (March to May), or at the end of the wet season (September to November), or both (Kendrick, 2002), sampling was concentrated during these seasons for both historic and current periods. Additionally, there was no seasonal bias in sampling within elevational bins across the two time periods (Supporting information Figure S1).

For occupancy parameterization, we included both sampling period and elevation in our models, resulting in four competing occupancy models: sampling period, sampling period + elevation, elevation and constant. We did not include quadratic terms because the elevation gradient covered is such that it was unlikely that a "bell-shaped" curve could be fitted. We built occupancy models in a Bayesian framework using the package “R2WinBUGS” in R 3.3.1 (Gelman, Sturtz, Ligges, Gorjanc, & Kerman, 2013; R Core Team, 2014). For each model (4 models for each of 123 species = 492 models in total), we used three chains and sampled 10,000 iterations with a thinning rate of 8. For each model, we calculated Bayesian information criterion (BIC) scores (Ntzoufras, 2011). As all BIC scores of competing models for each species were close (differences usually within 10 units), we calculated BIC weights for each occupancy model to produce occupancy curves along elevational bands by model averaging the candidate set (Hooten & Hobbs, 2015). We then calculated 95% confidence intervals based on unconditional variance of averaged models according to Buckland, Burnham, and Augustin (1997), and assumed that non-overlapping confidence intervals within elevational bands identified significant changes in occurrence probability.

2.4 | Elevational limit shifts

In addition to comparing estimated occurrence probabilities for each species, we tested explicitly for shifts in range limits by calculating the cumulative probability of false absence where raw occupancy data suggest a range shift has occurred (Moritz et al., 2008; Tingley & Beissinger, 2009). This approach is robust in minimizing errors of reporting range shifts when no actual range shift has occurred (Tingley et al., 2012). To calculate the probability of false absence ($P_{fa}$) for an apparent range shift, we used the detection probability ($p$) for each species from the best occupancy model (lowest BIC value) and the formula as reported by Tingley...
and Beissinger (2009). We considered a $P_{fa} < 0.05$ as a significant range shift; that is, species were likely to have truly changed their occupancy status rather than an apparent shift arising from false absences.

As all the other metrics we compared so far in this study were drawn from species that occur in both time periods, we made a direct comparison of species checklists to check for potential changes in colonization or local extinction. During the entire study period (1994 to 2014), the moth sampling data were not only limited to our study site (Tai Mo Shan elevational gradient), but also included 5,358 species occurrence records spread across 182 sites in Hong Kong. Using that list, we compared species of different time periods and listed 15 candidate species that only appeared in either the current or historic sampling period. We verified this list with the same $P_{fa}$ approach described above, but this time we ad hoc calculated detection probability for each species by dividing its total number of detections by the total number of visits to occupied sites, as we could not fit occupancy models for these species due to the hap hazard nature of sampling. We calculated the $P_{fa}$ of sites with visits where we failed to record each species and also used visits from the geographically nearest sites for unpaired sites. Additionally, we compared those 15 species to records from iNaturalist, a citizen science program with wide participation in Hong Kong (https://www.inaturalist.org/projects/hong-kong-moths), to see whether any validated records (records with a photo associated that could be verified as to the species identification) exist within our study period but were not recorded in our own surveys.

2.5 Community level elevation changes

We calculated “community elevation scores” for each shared site between time periods ($n = 6$: 70 m, 175 m, 210 m, 291 m, 307 m, 468 m) similar to Chen et al., (2009). This community metric treats the elevational centre of each species in the historic sampling period as a fixed “trait.” By averaging these traits of species occurring at one site together, we produced the “community elevation score” for that site. A lower community elevation score in the current period indicates a higher proportion of species with lower elevational centres occurring in that site, which suggests a community-wide shift to higher elevations. Paired-Wilcoxon tests were performed to determine whether community changes across all sites were significant. To calculate the elevational centre for each moth species, we first calculated the relative occurrence probability for each species by dividing occurrence in each site by the number of sampling times in that site. Then, we standardized these occurrence probabilities to one, and used them as weights to average the elevations of sites for each species.

3 RESULTS

Environmental data suggest that the climate has changed in Hong Kong over the sampling period, but the natural environment has mostly remained constant. For climate data, we detected no significant change in annual mean temperature, mean daily minimum temperature or annual mean precipitation from 1997 to 2014 (Supporting information Figures S2 and S3). However, mean maximum temperature significantly increased at an estimated warming rate of 0.46°C per decade for the low-elevation site (SK, 16 m a.s.l) and 0.51°C per decade for the high-elevation site (TMS, 955 m). For extreme weather, we detected a 10.8 days per decade increase in the annual number of extreme hot days and a 7.5 days per decade increase in the annual number of extreme cold days (Supporting information Figure S4). We found no significant change in annual number of extreme hot nights, dry days or flood days. In terms of extreme cold, 15 frost days within 3 years occurred during the current sampling period, compared to 4 frost days within 2 years during the historic sampling. For forest cover, the amount of change was negligible in that the highest change rate was −4.1% (0.54 ha) forest cover loss below 100 m (a.s.l) and +1.2% (10.26 ha) forest gain at 500–600 m. Otherwise, forest cover changes were less than 1% (0.9 to 13.59 ha) of the forested area. Buffers of different sizes around sampling points also suggested minimal forest cover change. During the study period, for the 90 m by 90 m, 180 m by 180 m and 360 m by 360 m buffer, the average forest cover losses were −0.31%, −0.36% and −0.24%, respectively. Average forest cover gains were +0%, +0.04% and +0.06%, respectively.

The total number of moth detections (counting one per species per sample) recorded in historical sites (1,199) was higher than the number of individuals at current sites (958). However, only one of the 123 modelled species showed significant (but limited) changes of occupancy with elevation over time (less than 10%; Supporting information Figure S5). This species, Callix ghosha, exhibited increased occurrence probability across all elevations. Of all moth distributions we examined, the majority (72 out of the 123 species) could be best explained by a “constant” model. The model with “sampling period” was the best model for 48 species, and the “elevation” model explained the distributions for the remaining three species.

When checked with $P_{fa}$, 32 out of 123 moth species showed lower or upper range limit shifts (Supporting information Table S3; without considering $P_{fa}$, the number would be 96 out of 123). Three species showed both lower and upper range limit shifts (expansions and downward shifting). Three species only shifted their upper range limits while 26 species only shifted lower range limits (including C. ghosha). For lower range limits, 22 species extended downwards, and seven species contracted upwards. For upper range limits, four species extended upwards and two species contracted downwards (Figure 3 and Supporting information Table S3).

Of the 15 candidate species that only occurred in one time period, the $P_{fa}$ approach found that Alex palparia and Herochroma viridaria (Supporting information Table S2) were likely truly absent from historic recordings and were newly established in the current sampling period. Records from iNaturalist of these two species in the current sampling period (2010–2014) outside of the study site.
further confirm the recent colonization of Hong Kong by these two species. Based on iNaturalist records and our own surveys, *Calletaera dentata* may also be newly established—however, we cannot rule out the possibility of false absence due to limited historic sampling effort in our surveys relative to its detectability (Supporting information Table S2). For the other 12 species, due to limited sampling intensity in our dataset, the Pfa method could not rule out the possibility of false absence. However, iNaturalist records indicated that they were actually present elsewhere in Hong Kong across the whole sampling period. In this case, we do not know whether they went locally extinct or are new arrivals to the Tai Mo Shan area, but we know they were and are present in Hong Kong.

Community-wide, we found that the average elevational centres of species present at re-surveyed sites decreased over time at 5 out of 6 sites, indicating widespread upslope shifts (Wilcoxon signed-rank test, $p = 0.06$). For these five sites, current community elevation scores were 3 to 27 m lower than historic scores, representing more lowland distributed species having been recorded in higher elevations (Figure 4). For our lowest elevation site (70 m), however, the community elevation index increased by 12 m, with more species of higher elevations increasing ($n = 28$, mean elevation centre 345.4 m) than species of lower elevations disappearing ($n = 25$, mean elevation centre 314.8 m).

### DISCUSSION

This study represents one of the first examinations of climate change effects along an elevational gradient in the lowland tropics (Figure 1). Although moths are generally nocturnal, day-time temperatures can still affect resting individuals and larvae (Xing et al., 2018). With minimal habitat change but significant warming over the past decade (as reflected by a 0.5°C increase in maximum temperatures), we detected complex changes of geometrid moth species at both species and community levels. Increased averaged daily maximum temperatures, and increased numbers of extreme hot and cold days, result in higher fluctuations in daily temperatures. These fluctuations potentially are more challenging for organisms than pure increases of daily mean temperatures (Paaijmans et al., 2013; Vasseur et al., 2014). This might explain why, despite the relatively short resurvey timespan (15-year study duration), we detected significant changes in the moth communities, particularly in range limits and at community levels, despite using a highly conservative inferential approach. These changes not only indicate high sensitivity of moths to environmental change, but also provide possible early warning signs for larger climate change impacts in the future (e.g., 48 of 123 moth species showed temporal trends in occupancy).

| Range Shift | Species | Elevation Shift (m) | Diagram |
|-------------|---------|--------------------|---------|
| **Expansion** (+80 m, -16 m) | *Antitrygodes agrata vicina* | 650 | ![Expansion Diagram] |
| **Lower and upper limits** (-127 m, -114 m) | *Synegia eumeleata* | 650 | ![Lower and upper limits Diagram] |
| **Lower limits** (-109.6 m) | *Rutellerona pseudocestaria* | 650 | ![Lower limits Diagram] |
| **Lower limits** (+149.7 m) | *Luxiaria mitmorhapes* | 650 | ![Lower limits Diagram] |
| **Upper limits** (-223 m) | *Perixera griseata* | 650 | ![Upper limits Diagram] |
| **Upper limits** (+83 m) | *Thalassodes immisaria* | 650 | ![Upper limits Diagram] |

**Figure 3** Different range shift scenarios, the number of moth species exhibiting each scenario (detection-corrected), and representative moth species occurrences for each scenario in Tai Mo Shan area, Hong Kong SAR (for full list of species results see Supporting information Table S3). Hollow circles represent absence, and filled circles represent presence. Numbers of moths depicted are the total numbers of species belonging to each scenario. Numbers in parentheses are average elevational shifts for all species belonging to each scenario, with positive values indicating upwards shifts and negative values indicating downward shifts.
We found high heterogeneity among and within the different approaches with respect to how different moth species have changed over the past decade. Compared to the historic sampling period, only one species (Collix ghosha) showed a significant change in occurrence probability and was found to be more common in the current sampling period. However, we documented two new species as recently established in Hong Kong. Many species also exhibited distribution changes, especially expansion downwards from their lower range limits (Figure 3). Consistent with this result, we found that the lowest elevation site had more high-elevation species compared to the historic community (Figure 4). However, communities at all other sites recorded more lowland species than high-elevation species, indicating an overall upslope shifting trend across most of the elevational gradient. The different responses detected between community and range margins could result from the limited number of sites with resampling (n = 6) allowing calculation of community elevation scores. In other tropical montane sites, similar heterogeneous shifts have been reported in resurveys of birds (Forero-Medina et al., 2011; Freeman & Freeman, 2014), plants (Feeley et al., 2011; Jump et al., 2012), moths (Chen et al., 2009; Chen, Hill, Shiu, et al., 2011) and reptiles (Raxworthy et al., 2008). In addition, studies across different taxa in other continents and cooler climates have also found similar heterogeneous shift patterns (Forister et al., 2010; Rowe et al., 2015; Tingley et al., 2012). The high heterogeneity in species responses documented here in the Hong Kong geometrids is plausibly a consequence, at least in part, of complex local climatic conditions, especially considering the increase of both extreme hot and cold events in recent years. Species experiencing more variable environments, such as mountain tops, are likely to be more adaptable to such variable environments (Ghalambor, Huey, Martin, Tewksbury, & Wang, 2006), and we suspect that the increase of frost days and extreme weather events are the key drivers of the observed large proportion of lower range margin shifts.

Non-climatic factors may also be responsible for the complex changes observed in Hong Kong moths since 2000. Tropical areas are characterized by strong and complex biotic interactions (Schemske, Mittelbach, Cornell, Sobel, & Roy, 2009), which could produce complicated effects for different taxa under climate change with different traits (Rehm, 2014; Van der Putten, Macel, & Visser, 2010). Our heterogeneous shift patterns in range limits could be a result of important biotic interactions such as interspecific competition (Rubidge et al., 2011), trophic interactions (Brodie et al., 2017; Pounds, Fogden, & Campbell, 1999) or time lags (Forero-Medina et al., 2011). For example, plants (including host plants of moths, which could change without detection by the satellite image forest cover change criteria we used here) are known to lag warming impacts due to relatively low dispersal abilities (Rehm, 2014), while predatory birds might react differently and move faster or more slowly (Forero-Medina et al., 2011; Rehm, 2014). These interactions could indirectly influence moth presence to a greater extent than direct temperature or precipitation changes and produce complex shift patterns.

Additionally, the context of this study system might also affect the interpretation of these results. Community elevation scores results may only represent patterns of changed site occupancy but not “true” elevational shifts, as it only includes six sites. In addition, rare species are common in the tropics (Price, Diniz, Morais, & Marques, 1995) and are potentially more sensitive to climate change impacts (Iverson, Schwartz, & Prasad, 2004). However, such species also have low detection probabilities, which therefore require greater sampling effort to detect changes and shifts. Different species might also react to sampling methods differently and produce underestimated or overestimated changes in occupancy or abundance (e.g., Eumelea ludovicata is not easily attracted to light or light traps but is frequently observed in Hong Kong).

Finally, these results lead to recommendations for long-term monitoring programs in the tropics for the detection and management of climate-driven species redistribution (Basset et al., 2017; Bonebrake et al., 2018). First, sampling effort should be balanced between sites and sampling methods should be diversified to cover broad communities. Second, repeated sampling is crucial to effectively calculate detection probability and record abundance changes. Third, citizen science data (in this case iNaturalist records) can be used as a reliable source to verify or support findings from standardized monitoring efforts. Where possible, more population-level studies are also needed to examine species redistributions and the processes underlying these changes (Brown et al., 2016). And for those species that do show trends of range shifts and abundance changes, we need to prioritize and support the monitoring of their populations and distributions, in case intervention is required if further climate change endangers their persistence. Meanwhile, species with different life history traits will respond differentially to the same
change in climate (Garnas, 2018). Collecting additional natural history information on such species is important and will allow more sophisticated trait-based analyses which can uncover mechanistic processes behind range shifts (Dulle et al., 2016).

CONFLICT OF INTEREST.
Authors declare no conflict of interest.

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DATA ACCESSIBILITY
Data are available in the Supporting information (Figures S1–S5 and Tables S1–S4).

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REFERENCES
Araújo, M. B., & Rahbek, C. (2006). How does climate change affect biodiversity? Science, 313, 1396–1397.
Basset, Y., Lamarre, G. P., Ratz, T., Segar, S. T., Decaëns, T., Rougerie, R., ... Ramirez, J. A. (2017). The Saturniidae of Barro Colorado Island, Panama: A model taxon for studying the long-term effects of climate change? Ecology and Evolution, 7, 9991–10004. https://doi.org/10.1002/ece3.3515
Bates, A. E., Pecl, G. T., Frusher, S., Hobday, A. J., Wernberg, T., Smale, D. A., ... Holbrook, N. J. (2014). Defining and observing stages of climate-mediated range shifts in marine systems. Global Environmental Change, 26, 27–38. https://doi.org/10.1016/j.gloenvcha.2014.03.009
Beck, J., Schulze, C. H., Linsenmair, K. E., & Fiedler, K. (2002). From forest to farmland: Diversity of geometrid moths along two habitat gradients on Borneo. Journal of Tropical Ecology, 18, 33–51. https://doi.org/10.1017/S026646740200202X
Bonebrake, T. C. (2013). Conservation implications of adaptation to tropical climates from a historical perspective. Journal of Biogeography, 40, 409–414. https://doi.org/10.1111/jbi.12011
Bonebrake, T. C., Brown, C. J., Bell, J. D., Blanchard, J. L., Chauvenet, A., Champion, C., ... Dell, A. I. (2018). Managing consequences of climate driven species redistribution requires integration of ecology, conservation and social science. Biological Reviews, 93, 284–305. https://doi.org/10.1111/brv.12344
Bonebrake, T. C., & Deutsch, C. A. (2012). Climate heterogeneity modulates impact of warming on tropical insects. Ecology, 93, 449–455. https://doi.org/10.1890/11-1187.1
Bonebrake, T. C., & Mastrandrea, M. D. (2010). Tolerance adaptation and precipitation changes complicate latitudinal patterns of climate change impacts. Proceedings of the National Academy of Sciences, 107, 12581–12586. https://doi.org/10.1073/pnas.0911841107
Brodie, J. F., Strimbas-Mackey, M., Mohd-Azlan, J., Granados, A., Bernard, H., Giordano, A. J., & Helmy, O. E. (2017). Lowland biotic attrition revisited: Body size and variation among climate change ‘winners’ and ‘losers’. Proceedings of the Royal Society London B: Biological Sciences, 284, 20162335.
Brooks, T. M., Mittermeier, R. A., da Fonseca, G. A., Gerlach, J., Hoffmann, M., Lamoreux, J. F., ... Rodrigues, A. S. (2006). Global biodiversity conservation priorities. Science, 313, 58–61. https://doi.org/10.1126/science.1127609
Brown, C. J., O’Connor, M. I., Poloczanska, E. S., Schoeman, D. S., Buckley, L. B., Burrows, M. T., ... Richardson, A. J. (2016). Ecological and methodological drivers of species’ distribution and phenology responses to climate change. Global Change Biology, 22, 1548–1560. https://doi.org/10.1111/gcb.13184
Buckland, S. T., Burnham, K. P., & Augustin, N. H. (1997). Model selection: An integral part of inference. Biometrics, 53, 603–618. https://doi.org/10.2307/2533961
Chen, I. C., Hill, J. K., Ohlemüller, R., Roy, D. B., & Thomas, C. D. (2011). Rapid range shifts of species associated with high levels of climate warming. Science, 333, 1024–1026. https://doi.org/10.1126/science.1206432
Chen, I. C., Hill, J. K., Shiu, H. J., Holloway, J. D., Benedick, S., Chey, V. K., ... Thomas, C. D. (2011). Asymmetric boundary shifts of tropical montane Lepidoptera over four decades of climate warming. Global Ecology and Biogeography, 20, 34–45. https://doi.org/10.1111/j.1466-8238.2010.00594.x
Chen, I. C., Shiu, H. J., Benedick, S., Holloway, J. D., Chey, V. K., ... Thomas, C. D. (2009). Elevation increases in moth assemblages over 42 years on a tropical mountain. Proceedings of the National Academy of Sciences, 106, 1479–1483. https://doi.org/10.1073/pnas.0809320106
Colwell, R. K., Brehm, G., Cardelús, C. L., Gilman, A. C., & Longino, J. T. (2008). Global warming, elevational range shifts, and lowland biotic attrition in the wet tropics. Science, 322, 258–261.
Deutsch, C. A., Tewksbury, J. J., Huey, R. B., Sheldon, K. S., Ghulamber, C. K., Haak, D. C., & Martin, P. R. (2008). Impacts of climate warming on terrestrial ectotherms across latitude. Proceedings of the National Academy of Sciences, 105, 6668–6672. https://doi.org/10.1073/pnas.0709472105
Dudgeon, D., & Corlett, R. (2004). The ecology and biodiversity of Hong Kong (pp. 336). Hong Kong: Friends of the Country parks.
Dulle, H. I., Forister, M. L., McCall, A. C., Sanders, N. J., Fordyce, J. A., Thorne, J. H., … Holbrook, N. J. (2014). Defining and observing stages of climate mediated evolution in species with mobile life stages. Proceedings of the National Academy of Sciences, 111, 2088–2092. https://doi.org/10.1073/pnas.0909686107
Feeley, K. J., Silman, M. R., Bush, M. B., Farfan, W., Cabrera, K. G., Böhning-Gaese, K., & Hof, C. (2016). Changes in abundances of forest understory birds on Africa’s highest mountain suggest subtle effects of climate change. Diversity and Distributions, 22, 288–299. https://doi.org/10.1111/ddi.12405
Forister, M. L., McCall, A. C., Sanders, N. J., Fordyce, J. A., Thorne, J. H., O’Brien, J., ... Shapiro, A. M. (2010). Compounded effects of climate change and habitat alteration shift patterns of butterfly diversity. Proceedings of the National Academy of Sciences, 107, 6668–6672. https://doi.org/10.1073/pnas.0909472105
Forerero-Medina, G., Terborgh, J., Socolar, S. J., & Pimm, S. L. (2011). Elevational ranges of birds on a tropical montane gradient lag behind warming temperatures. PLoS ONE, 6, e28535. https://doi.org/10.1371/journal.pone.0028535
Fry, R., & Waring, P. (1996). A guide to moth traps and their use. The Amateur Entomologists, 24, 4–60.
Garcia, R. A., Cabeza, M., Rahbek, C., & Araújo, M. B. (2014). Multiple dimensions of climate change and their implications for biodiversity. Science, 344, 1247579. https://doi.org/10.1126/science.1247579
Schemske, D. W., Mittelbach, G. G., Cornell, H. V., Sobel, J. M., & Roy, K. (2009). Is there a latitudinal gradient in the importance of biotic interactions? Annual Review of Ecology, Evolution, and Systematics, 40, 245–269. https://doi.org/10.1146/annurev.ecolsys.39.110707.173430

Schowalter, T. D., Hargrove, W., & Crossley, D. A. Jr (1986). Herbivory in forested ecosystems. Annual Review of Entomology, 31, 177-196. https://doi.org/10.1146/annurev.en.31.010186.001141

Tingley, M. W., & Beissinger, S. R. (2009). Detecting range shifts from historical species occurrences: New perspectives on old data. Trends in Ecology & Evolution, 24, 625–633. https://doi.org/10.1016/j.tree.2009.05.009

Tingley, M. W., Koo, M. S., Moritz, C., Rush, A. C., & Beissinger, S. R. (2012). The push and pull of climate change causes heterogeneous shifts in avian elevational ranges. Global Change Biology, 18, 3279-3290. https://doi.org/10.1111/j.1365-2486.2012.02784.x

Van der Putten, W. H., Macel, M., & Visser, M. E. (2010). Predicting species distribution and abundance responses to climate change: Why it is essential to include biotic interactions across trophic levels. Philosophical Transactions of the Royal Society of London B: Biological Sciences, 365, 2025–2034.

Vasseur, D. A., DeLong, J. P., Gilbert, B., Greig, H. S., Harley, C. D., McCann, K. S., ... O'Connor, M. I. (2014). Increased temperature variation poses a greater risk to species than climate warming. Proceedings of the Royal Society of London B: Biological Sciences, 281, 20132612.

Wang, X., Liu, H., Gu, M. B., Boucek, R., Wu, Z. M., & Zhou, G. Y. (2016). Greater impacts from an extreme cold spell on tropical than temperate butterflies in southern China. Ecosphere. 7(5), e01315. https://doi.org/10.1002/ecs2.1315.

Xing, S., Bonebrake, T. C., Ashton, L. A., Kitching, R. L., Cao, M., Sun, Z., ... Nakamura, A. (2018). Colors of night: Climate-morphology relationships of geometrid moths along spatial gradients in southwestern China. Oecologia, 188, 537-546. https://doi.org/10.1007/s00442-018-4219-y

BIOSKETCH

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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