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# Long-term changes in populations of rainforest birds in the Australia Wet Tropics bioregion: a climate-driven biodiversity emergency — Source link $\square$

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### 16 Abstract

17 Many authors have suggested that the vulnerability of montane biodiversity to climate change worldwide is significantly higher than in most other ecosystems. Despite the extensive variety 18 of studies predicting severe impacts of climate change globally, few studies have empirically 19 20 validated the predicted changes in distribution and population density. Here, we used 17 years of bird monitoring across latitudinal/elevational gradients in the rainforest of the Australian 21 22 Wet Tropics World Heritage Area to assess changes in local abundance and distribution. We used relative abundance in 1977 surveys across 114 sites ranging from 0-1500m above sea 23 24 level and utilised a trend analysis approach (TRIM) to investigate elevational shifts in abundance of 42 species between 2000 - 2016. The local abundance of most mid and high 25 26 elevation species has declined at the lower edges of their distribution by >40% while lowland species increased by up to 190% into higher elevation areas. Upland-specialised species and 27 regional endemics have undergone dramatic population declines of almost 50%. The 28 "Outstanding Universal Value" of the Australian Wet Tropics World Heritage Area, one of the 29 most irreplaceable biodiversity hotspots on Earth, is rapidly degrading. These observed impacts 30 are likely to be similar in many tropical montane ecosystems globally. 31

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Keywords: climate change; rainforest; birds; Australia; population declines; elevational
 distribution; biodiversity; species vulnerability; world heritage; escalator to extinction;
 montane ecosystems.

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#### 40 Introduction

41 There is widespread recognition that climate change is rapidly becoming the most significant threat to global biodiversity and natural ecosystems [1, 2]. At a global scale, estimates of total 42 species extinctions projected over the remainder of the century vary considerably between 43 ecosystems, taxa and methods of analysis [3]. However, in all studies, the level of predicted 44 impacts is disturbing with potential losses of between 15-35% of all species [3, 4]. The 45 projected impact of climate change is expected to be especially severe in mountain ecosystems 46 with up to 84% of mountain species globally facing a high extinction risk [5-7]. Mountain 47 regions provide a host of critical ecosystem services [8], support a disproportionate amount of 48 the world's biological diversity and, harbour many phylogenetically unique species [5, 9-12]. 49 50 Mountain regions contain roughly 87% of the world's vertebrate biodiversity [13], 54% of which is completely restricted to mountain ecosystems [13]. The global significance of 51 montane ecosystems is heightened in the tropics [14, 15]; approximately 50% of the world's 52 species of plants and vertebrates are believed to be endemic to 34 identified global biodiversity 53 hotspots [16], 85% of which include large areas of tropical forest or montane cloud forest [17]. 54 Tropical species are considered to be particularly sensitive to climate change [18-20] and the 55 warming rates are relatively high in tropical mountains [18]. Consequently, tropical montane 56 biodiversity is not only globally important but particularly threatened [7, 21]. Despite the global 57 significance and high vulnerability of tropical ecosystems, there have been few studies 58 demonstrating observed impacts of climate change in the tropics [22, 23]. The paucity of 59 60 tropical studies makes it difficult to measure and predict the impacts of climate change relative 61 to other drivers like habitat loss [24], especially given that most studies are short-term or lacking abundance data [25]. There is a need for increasing monitoring and improving 62 63 understanding of the impacts of climate change in tropical montane ecosystems [12].

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On mountains, biotic communities and abiotic conditions change abruptly over short distances, 65 with greater elevational than lateral turnover in species composition [26]. Across all elevations, 66 67 assemblages on mountains with high rates of past temperature change exhibit more rapid diversification, highlighting the importance of climatic fluctuations in driving the evolutionary 68 69 dynamics of mountain biodiversity [12]. Globally, increasing evidence indicates that species are responding to climate change by shifting their geographical distributions [27]. These shifts 70 often follow warming temperatures poleward and upslope [6, 10]. Montane species are of 71 particular concern in this respect, as they are expected to experience reduced distribution area, 72

increased population fragmentation, and increased risk of extinction with upslope movement 73 into ever-smaller area [28, 29]. The high degree of specialization to narrow temperature ranges 74 that montane species typically exhibit has raised concern over their future under climate change 75 [10, 29, 30]. It is widely expected that montane species will experience further upslope shifts 76 in the future and, in the absence of broad latitudinal shifts due to the geographical features of 77 montane ecosystems, such movements will leave species with less habitable area as they 78 79 approach mountain peaks [29, 31]. Left with nowhere else to go, montane species are predicted to become increasingly susceptible to the stochastic extinctions or declining populations [32]. 80 81 This so-called "escalator to extinction" [33] has been predicted, and now observed, in a number of places and taxa around the world [28, 34-36]. 82

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The rainforests in the Australian Wet Tropics bioregion in north-east Queensland are globally 84 significant for high biodiversity value based on high endemism, evolutionary significance and 85 phylogenetic distinctiveness [37-39]. These biodiversity values resulted in the region being 86 listed as a World Heritage Area in 1988 and being described as the sixth most irreplaceable 87 protected area globally [40] and the second most irreplaceable World Heritage Area [41]. The 88 high endemism and relictual nature of the biodiversity within the region is largely attributed to 89 90 the influence of historical fluctuations in rainforest area over the Quaternary and the restriction of rainforest to cool, moist, upland refugia [37]. This biogeographic history imposed a non-91 92 random extinction filter across the region resulting in most of the regionally-endemic species being cool-adapted upland species [42, 43]. It is this biogeographic history, with the resulting 93 94 concentration of endemic species in the cool uplands, that has made the biodiversity of the region so unique but highly vulnerable to a warming climate. Predictions about the future of 95 this biodiversity under anthropogenic climate change are grim, particularly for the upland 96 97 regionally-endemic species and habitat types [5, 44-46]. In 2003, species distribution 98 modelling of the endemic vertebrates suggested the potential for catastrophic impacts over the coming century with more than 50% of these species predicted to go extinct due to a complete 99 loss of suitable climatic space [5]. These predictions drove a greater research effort in the 100 region in the interim years and there have been extensive region-wide surveys of many 101 vertebrate and invertebrate taxa [47-50]. 102

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Are the declines in species ranges predicted in 2003 concordant with observed spatial trends in species abundance patterns over subsequent years? Unfortunately, the answer is yes. We present quantitative evidence, based on the long-term monitoring of vertebrates across the 107 entire bioregion, for significant declines and shifts in the spatial distribution of populations. Field monitoring clearly demonstrates that the previously projected impacts are clearly 108 concordant with observed shifts in species abundance. Previous analyses of the climate change 109 impact in the region have relied on either modelled distribution changes using various IPCC 110 emission scenarios or coarse comparisons of changes pre-2008 compared to post-2008. Here, 111 we examine in high spatial, temporal and taxonomic detail the observed changes in the 112 rainforest bird assemblages of the region between 2000-2016, based on the regional-scale 113 standardised surveys from the Williams Wet Tropics monitoring program (updated from 114 115 Williams, VanDerWal (47)). We use this long-term dataset to test for bird species that have undergone significant changes in local abundance and/or elevational. We predicted that bird 116 assemblages should systematically shift upwards in elevation and that the local abundance of 117 individual species would decline on the lower (warmer) edge of their distribution and increase 118 at the higher (cooler) edge of their distribution [31]. We tested for trends across time in local 119 abundance (site/elevation specific) and assemblage shifts across elevation and used an area-120 weighted trend to examine trends in total population size. These impacts are likely to be 121 representative of impacts in many mountain ecosystems across the globe [7]. 122

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#### 126 **Results**

127 Overall, across all 42 species over the 17 years, there has been a significant decline in local abundance of rainforest birds of approximately  $12 \pm 1.4\%$  (~ -0.2% per year) (Figure 1a, Table 128 1). However, this overall trend masks complex, and often contrasting trends, within different 129 ecological subsets of the rainforest bird assemblage (Table 1). Habitat generalists increased by 130 more than 50% from 2000 to 2011 and then steadily declined until 2016 (13 species, 3.3%/year, 131 overall trend  $9 \pm 4.1$  %, Figure 1b, Table 1). Local abundances of rainforest specialist species 132 have declined on average by approximately 20% (29 species, -1.7%/year, overall trend  $-20 \pm$ 133 1.3 %, Figure 1C, Table 1). Regionally endemic species, a subset of habitat specialists, showed 134 135 the strongest decline with a loss of  $\sim 34 \pm 1.7\%$  in local abundance (10 species, -2.4%/year, Figure 1d, Table 1). Population trends of habitat generalists and specialists were significantly 136 different (trend difference  $0.05 \pm 0.002$ , p<0.05). However, these average multi-species trends 137 in ecological groupings also mask variable trends for individual species (Figure S1) and 138

assemblages in different elevational bands (Table S3). Species-specific trends in local
abundance and total population size (local abundance trends weighted by area within each
elevational band) are presented in the Appendix (Figure S1) (Temporary link to interactive
Appendix -

- 143
- 144 https://alejandrodelafuente.shinyapps.io/BirdsPopTrendAWT/? ga=2.148260535.10245279
- 145 <u>38.1618546081-1577712465.1581926346</u>.

Table 1. Multi-species indicator trends between 2000 – 2016 (overall) and separate trends pre-2009 and post-2008. Population trends were classified following Pannekoek and van Strien (51) into one of the following categories depending on the overall multiplicative slope and its 95% confidence interval. Strong increase/Steep decline: increase/decline significantly more than 5% per year. Moderate increase/decline: significant increase/decline, but no more than 5% per year. Stable: no significant increase or decline, and it is certain than trends are less than 5% per year. Uncertain: no significant increase or decline, but it is not certain if trends are less than 5% per year.

	Overall			2000 - 2008			2009 - 2016		
Indicator	Slope	s.d.	Trend classification	Slope	s.d.	Trend classification	Slope	s.d.	Trend classification
All species	0.998	0.001	Moderate decline	1.003	0.002	Stable	0.982	0.002	Moderate decline
Endemic species	0.976	0.001	Moderate decline	0.967	0.003	Moderate decline	0.985	0.004	Moderate decline
Lowland species	1.03	0.002	Moderate increase	1.074	0.005	Strong increase	1.008	0.005	Stable
Midland species	0.986	0.001	Moderate decline	0.992	0.003	Moderate decline	0.98	0.003	Moderate decline
Upland species	0.971	0.001	Moderate decline	0.952	0.003	Moderate decline	0.972	0.003	Moderate decline
Habitat generalists	1.033	0.002	Moderate increase	1.054	0.005	Moderate increase	0.974	0.005	Moderate decline
Rainforest specialists	0.983	0.001	Moderate decline	0.981	0.002	Moderate decline	0.986	0.002	Moderate decline

#### 154

Lowland specialist species have undergone a strong and significant increase of  $72 \pm 7\%$  (6) 155 species, 3%/year, Figure 2a). Mid-elevation specialists declined by  $21 \pm 1.4\%$  (16 species, -156 1.9%/year. Figure 2b) and upland specialists have undergone declines of  $44 \pm 4\%$  (13 species, 157 -2.9%/year, Figure 2c). Population trends for lowland specialists were significantly different 158 from the trends in both midland (trend difference  $0.043 \pm 0.002$ , p<0.05) and upland specialists 159 (trend difference  $0.059 \pm 0.002$ , p<0.05). Additionally, upland specialists have declined 160 significantly more than midland specialists (trend difference  $0.015 \pm 0.002$ , p<0.05), 161 suggesting that the pattern of decline increases with increasing elevation. 162

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Shifts in patterns of local abundance are species-specific and highly variable, often involving 164 complex spatio-temporal and non-monotonic trends (for example, Brown Gerygone, Figure 165 S1, Figure S1.1). However, the overall multi-species trends that summarise the observed shifts 166 in bird assemblages along the elevational gradient are what we would predict under a warming 167 climate: there have been significant upslope shifts in bird abundance patterns across the 17 168 years of this study. We demonstrated this by examining the trends in local abundance for each 169 group of elevational specialists in their original preferred elevation (baseline averaged 170 abundance at each elevation across 1996-2003) and in the adjacent elevational bands over time. 171 We predicted, for example, that lowland species should increase in the midlands, and upland 172 species should decline at lower elevations. The observed spatial shifts in elevational 173 distribution and abundance are in accord with the earlier predictions based on distribution 174 changes. Lowland specialist species have moderately increased in local abundance in the 175 176 lowlands (+17%). On the other hand, lowland specialists' local abundance has dramatically increased into the midlands (~190% increase) (Figure 3). Midland specialist species have 177 declined in the lowlands by approximately 42%, by 22% in the midlands and are currently 178 stable in the uplands (Figure 3). Upland specialist species have declined everywhere, with a 179 180 catastrophic 49% decline in the lower-elevation midlands and a 33% decline in the uplands (Figure 3). Detailed trends for individual species at each elevation category are in Appendix 181 182 (Table S3).

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#### 185 **Discussion**

There is a temptation for policy makers and environmental managers to consider that 186 187 biodiversity within a protected area is safe, this is a dangerous, and in this case incorrect, assumption. The montane rainforest birds of the Australia Wet Tropics World Heritage Area 188 189 are in danger of extinction due to the increasingly severe impacts of a changing climate. In 2003, Williams et al. [52] predicted catastrophic extinction levels within the regionally 190 191 endemic species of the Australian Wet Tropics World Heritage Area by the end of the 21st century. These early predictions were based on bioclimatic envelope models of the changes in 192 193 species distributions with increasing temperature. However, the reliability of predictions using this simple approach is debateable, with a likelihood of overestimating the severity of impacts 194 [53]. Therefore, a more intensive monitoring effort was conducted throughout the region, 195 covering ~94% of the environmental space present and providing standardised estimates of 196 local abundance for many species. 197

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An earlier analysis using these local abundance measures of rainforest birds across elevation 199 [45] predicted that 74% of rainforest bird species would become threatened by the end of the 200 21<sup>st</sup> century. demonstrated a tight relationship between elevation and the assemblage structure 201 202 of birds based on empirical, site-based relative abundance across the elevational gradient. Based on this relationship, they predicted that bird assemblages would systematically move 203 upwards in elevation as the climate warmed. However, despite the development of more 204 sophisticated modelling approaches and the availability of more field data, the qualitative 205 outcomes of predictions remained similar to the 2003 predictions. Here, we demonstrate that 206 207 these predicted trends, whether based on species distribution models or empirical abundance, were concordant with expectations. Our results here demonstrate that the rainforest bird 208 assemblage in the Australian Wet Tropics bioregion is clearly, and consistently, moving uphill 209 in a classic demonstration of the "escalator to extinction" [33]. 210

211

Overall, we are on track for severe impacts on rainforest birds, particularly the upland specialists. Many of these species are endemic to the region and/or include species that are evolutionary distinct and/or ecologically specialised [38]. Upland bird populations have

declined since 2000 by 33% within their core range and the more marginal populations in the midlands have crashed by almost 50% (Figure 3). Midland species, although stable in the cooler uplands at this stage, have declined by >20% within their core area and >40% in the hotter lowlands. Lowland species are mostly increasing in abundance thus far with increases in abundance in the lowlands and strong increases in range and abundance into the midlands (Figure 3, Table 1).

The pattern in temporal change in abundance of rainforest birds presented in this study is 221 consistent with observed species elevational shift worldwide [27, 34-36, 54-56]. Globally, 222 evidence suggests a strong uphill elevational shift across different taxa, indicating that this 223 trend could be a generality. However, some studies have pointed out some idiosyncratic results 224 found in species-specific trends within different communities [27, 54]. According to this, our 225 226 results showed that, even though the uphill distribution shift is marked across most species, some birds showed stable trends, while others showed pattern opposite than expected (e.g., 227 228 Acanthiza katherina, figure S1). Those individual exceptions to the overall pattern highlighted potential interspecific differences in dispersal capacity, resilience, and adaptability between 229 bird species in the Wet Tropics, whose impact in the overall community "re-shuffling" at the 230 ecosystem level should be further studied. Overall, this study supports growing evidence tof 231 the rapid impact that climate change is having on tropical ecosystems [28] and provides the 232 first evidence of a climate-induced elevational shift in tropical rainforest of Australia. 233

So, what is driving these shifting abundance patterns? Is it the direct impacts of temperature 234 on the birds or is it an indirect effect via food resources or other biotic interactions? There is 235 little confidence in our ability to predict the potential impacts of the complex changes in biotic 236 237 interactions due to these marked changes in abundance and geographic shifts in distributions. Upland and midland assemblages are likely coming under increased pressure due to changing 238 239 biotic interactions and community structure associated with the increasing abundance of lower elevation species as they shift higher up the mountain. The changes shown in this study suggest 240 241 that, thus far, the impacts on species have been largely direct, or at least directly associated with an important resource, rather than changes in competitive interactions between bird 242 243 species. We argue this on the basis that upland bird species have declined in their core range (>850m) despite there being no significant increase in midland birds in the uplands thus far. 244 245 Midland birds have only declined by about 20% in the midlands despite the influx of lowland species. Lowland species have increased (albeit with more recent trends of decline) and there 246 has been no noticeable change in species composition or new invading species in the lowlands 247

to date. Thus far, there is no evidence supporting "lowland attrition" of species in this system 248 [31]; however, this is not entirely surprising given that we have little knowledge of the upper 249 thermal limits of the lowland species because they already occur in the hottest part of the region 250 Shoo, Anderson (57) suggested that the elevational pattern of abundance for the Lewin's 251 Honeyeater was directly influenced by temperature and not competitive interactions. This 252 conclusion was based on a demonstration that the elevational abundance profile of the species 253 varied as predicted by temperature in the populations on mountains on Cape York, a hotter 254 montane system, within a very different bird assemblage to that in the Wet Tropics region[57]. 255 256 However, this does not account for other biotic interactions that might also be changing such as diseases, or other non-bird competitors such as ectotherms (invertebrates mostly) that might 257 be increasing in uplands. 258

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There are many indirect mechanisms that could potentially exacerbate the impact that climate 260 change will ultimately have on bird species. Increased pressure from parasites with increasing 261 temperature [58], decreased reproductive success during prolonged dry seasons [59, 60], 262 increased habitat and population fragmentation [61-63] and increased environmental 263 marginality [64]. In this region, Williams, Shoo (50) hypothesised that dry season bottlenecks 264 and changes in net primary productivity could have a strong influence on bird populations. Net 265 primary productivity is limited by both temperature and water availability and could possibly 266 increase in the uplands with a warming climate, potentially alleviating some of the more direct 267 impacts of high temperatures. However, the strong declines in the abundance of upland bird 268 species shown here suggests that any positive influence of increased net primary productivity 269 has been swamped by the negative impacts of increasing temperature. There is existing 270 evidence highlighting the importance of changing rainfall patterns, especially harsher dry-271 272 seasons [59, 65] and decreasing water input from cloud interception [66]. It seems most likely that the driving factor behind many of the changes demonstrated in this study are the increasing 273 frequency and intensity of heat waves [67]. We need to increase our understanding of the 274 impacts of extreme climatic events so we can make more robust predictions than those that rely 275 276 only on changes in average conditions.

277

There have been clear impacts on biodiversity in almost every ecosystem and taxa across the globe due to anthropogenic climate change [2]; and now the world urgently needs to reduce

emissions and adapt as much as possible to minimise future impacts. Our efforts need to be 280 firmly focussed on the difficult question – what can we do? Managing habitats at the landscape 281 scale via habitat restoration, threat abatement and enhancing dispersal pathways represents one 282 avenue for local adaptation efforts to increase the resilience of biodiversity (Shoo et al. 2013, 283 [68]). The maintenance and restoration of movement pathways and corridors to facilitate 284 species movement into refugial areas will be vitally important [69], however, our results here 285 suggest that facilitating movement also warrants caution. While it is imperative that many 286 species can move into cool refugia [70], our data demonstrating the observed movement of 287 288 lowland, generalist species into the upland refugia, represents a potential threat to the upland endemics via increased competition with more generalist species. 289

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It is clear that montane systems are of paramount importance due to their high biodiversity value, many specialists and endemics and their role as the best-available cool refugia [70] and that these biodiversity hotspots are particularly threatened by climate. It is particularly disconcerting that, even in a fully protected and well-managed system such as the Australian Wet Tropics World Heritage Area, observed impacts are significant and accelerating. Most other tropical, montane biodiversity hotspots across the globe also face additional threats such as ongoing habitat degradation, poaching and urban encroachment.

#### 298

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#### 300 Conclusions

Upland bird species, of great conservation importance, are declining in abundance and 301 contracting their range to higher elevations in the montane rainforests in the Australian Wet 302 Tropics World Heritage Area in a classic example of the "escalator to extinction" [33]. These 303 species are suffering the combined, and increasing, threat of reduced distribution area, reduced 304 305 local population density and more fragmented and isolated populations, potentially causing a loss of genetic diversity in many species; factors that increase their vulnerability to extinction 306 [69]. Low-elevation rainforest species that are often more generalist, geographically 307 widespread and locally common are increasing in abundance and range size, potentially 308 resulting in yet another negative pressure on the upland specialists and an overall 309 homogenisation of the rainforest avifauna [71]. These changes are likely to be indicative of 310

impacts across all montane ecosystems globally, especially in important biodiversity hotspots associated with tropical mountain ecosystems. The important next step is to determine how we can slow, prevent, or adapt to these impacts to prevent the loss of the unique biodiversity of

- these regions around the world.
- 315

#### 316 Methods

#### 317 Study area

The Australian Wet Tropics bioregion is composed of mixed tropical rainforest in an area of approximately 1.85 million hectares. The terrain is rugged and dominated by mountain ranges, tablelands, foothills and a lowland coastal plain. The elevation varies from sea level to highlands at 1000 meters, with isolated peaks reaching up to 1,620 meters [37]. Annual rainfall varies between 1200-8000mm with rainforest covering most areas with annual rainfall above ~1500mm.

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#### 325 Data collection

Rainforest birds in the Australian Wet Tropics bioregion were monitored between 1996 and 326 2016 across the region at locations ranging from 0 to 1500 meters above sea level. All 327 individuals were recorded either by call or visual observation. Each survey was based on a 30 328 minute-150 meter transect with two observers. Surveys occurred within two hours of sunrise. 329 All long-term monitoring sites were located within large, unfragmented areas of rainforest with 330 continuous forest cover over the available elevational gradient. For further details of methods, 331 species observed, site localities and species traits see Williams, VanDerWal (47), Williams, 332 Shoo (50). 333

334

Data from four mountain ranges were included in the analyses presented here (Atherton Uplands, Carbine Uplands, Spec Uplands and Windsor Uplands), representing a total of 1977 surveys across 124 different sites. Analysis across these sites was possible based on coverage of the elevational gradient and consistency and numbers of surveys over time. Years with an entire elevation category missing were not included in the trend analyses (from 1996 to 1999). See Table S5 in supplementary information for complete breakdown of survey numbers by elevation and year.

342

#### 343 Species included

Initially, all species for which the survey technique was unsuitable (e.g. water birds) or when 344 call identification was considered unreliable due to the presence of multiple species with 345 similar calls were excluded to ensure maximum reliability in the trend analyses. This resulted 346 in a dataset of 54 species. Of these, analysis of population trends across elevation and time was 347 possible for 42 species where there was sufficient data across both elevation and years to 348 reliably analyse temporal changes in species abundance. We grouped species by habitat 349 (rainforest specialist, generalist) and elevation specialisation (lowland 0-450 m, midland 451-350 850m, upland >850). These elevational bands were selected to have the finest scale division of 351 352 elevation possible with a relatively equal band width and sample size in each band. Species were categorised as rainforest specialist if rainforest represented their main or core breeding 353 habitat [47]. Elevation specialisation was defined by the elevational abundance profile of each 354 species using a mean abundance for all surveys conducted between 1996-2004 within each 355 elevational band as the baseline elevation abundance profile (Table S1, Figure S1.1). Species 356 were assigned as a specialist in that elevational band when >70% of their elevational abundance 357 profile occurred within that band. 358

359

#### **360 Population indices and trends**

361 Overall population indices and trends for all 42 species were modelled using the rtrim package [72], an R-package based on the TRIM software (Trends and Indices for Monitoring data. 362 TRIM v. 3.54. [73]). TRIM is designed to analyse time-series of counts and produce unbiased 363 yearly indices and standard errors using log-linear models. The programme also estimates the 364 dispersion factor, correcting for over-dispersion, and takes account of serial correlation 365 between counts at the same site in different years [73]. This method has proved to be robust in 366 367 trend calculation with missing years [74-77]. We used model 2 in rtrim, which assumed all years as possible changing points in the population trend [78]. Overall trends for each species 368 369 were calculated with both unweighted data and data weighted by the geographic area within each of the elevational bands. The weighted trends give an estimate of the changes in the 370 species total population size as it takes into account the area within each elevational band as 371 well as local abundance changes. Given that the difference in results using weighted and 372 unweighted data was very marginal (Figure S1) and our primary focus here was to examine 373 site-specific changes in local abundance and elevational distribution shifts in abundance, we 374 have only presented in the results section the unweighted trends. Area-weighted trends of total 375 population size are included in the Appendix (Figure S1b). 376

Individual species indices produced by TRIM were combined into multi-species indicators for 377 each predefined group. Multi-species indicators were calculated using the MSI-tool in R [79]. 378 This tool uses species-specific annual indices and their standard errors provided by TRIM to 379 calculate annual multispecies indicator with confidence interval, accounting for sampling error, 380 using the Monte Carlo simulation method. This method calculates a mean and a SE from 1000 381 simulated multi-species indicators and back-transforms these to an index scale, then repeats the 382 process 10000 times. Those indicators are considered a measure of biodiversity change, where 383 a reduction in index mean will occur if more species are declining than increasing and vice 384 385 versa [75, 80]. We tested for significant differences between the multi-species indicators using the TREND DIFF-function, based on a Monte Carlo procedure (1000 iterations) and report 386 the average difference with SE in the multiplicative trends and the significance of this 387 difference. 388

389

Additionally, we explored the influence of changing trends over the study period by separately
examining the trends prior to, and after, 2008 (the midpoint of the time-series). This enables
some consideration of how the trends would have been observed over shorter time periods.

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Finally, to examine the population changes of each species within each elevational band, we
estimated the local trends for each species along the elevational gradient. Multi-species trends
in each elevation category were combined for each of the elevation specialist species groups.
Differences within groups were tested using the TREND\_DIFF-function. R (version 3.6.2) was
used in all the analyses [81].

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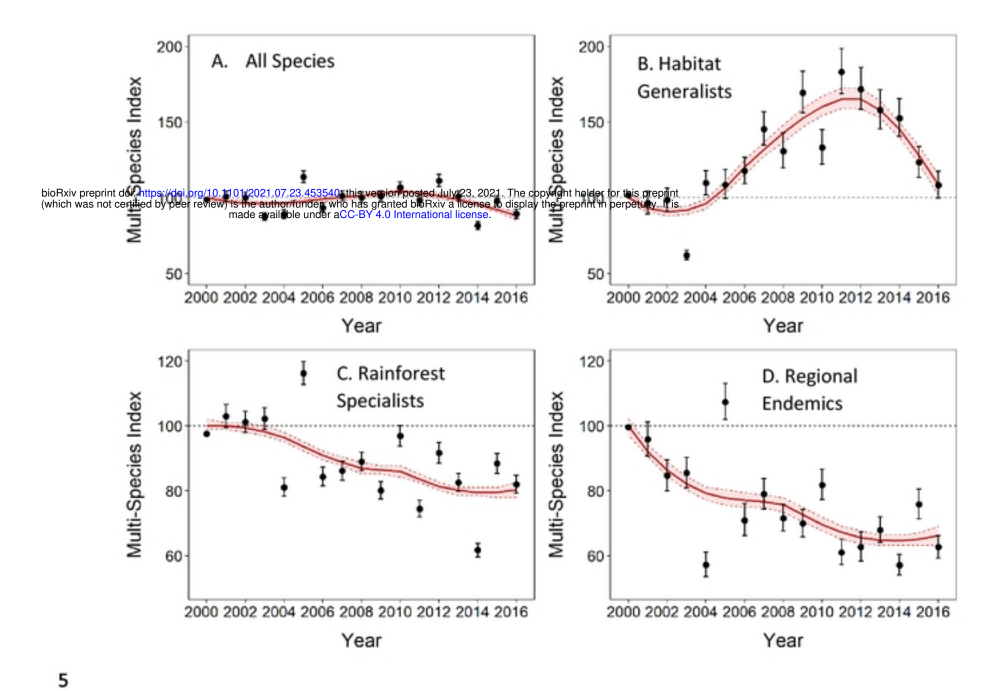
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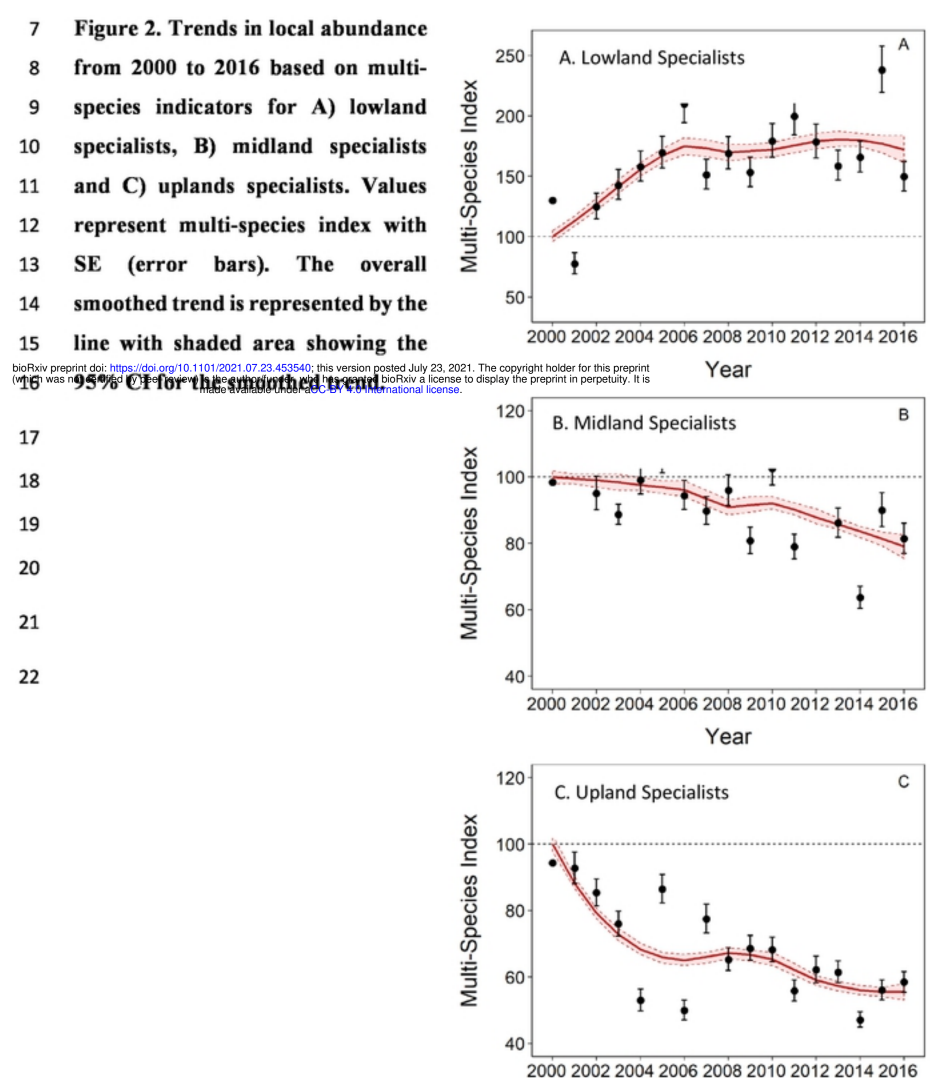
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Figure 1. Trends in local abundance from 2000 to 2016 based on the multi-species indicators for a) all species, b) habitat generalists, c) regional endemic species and d) rainforest specialists. Values are the multi-species index with SE (error bars) with the smoothed trend line with 95% CI (shaded area).

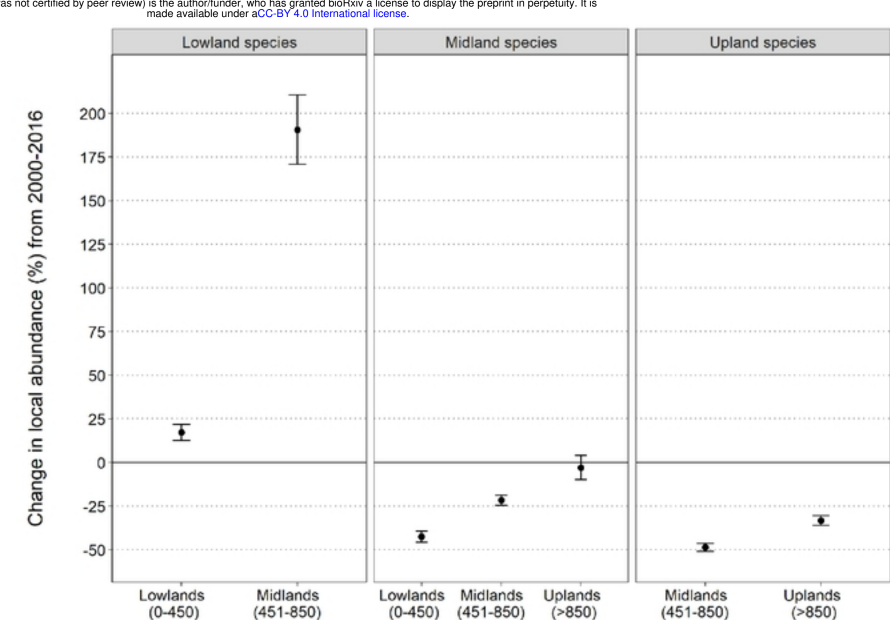




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Year

Figure 3. Elevational shifts in local abundance patterns in bird assemblages between 2000 to 2016 across the elevational gradient. Baseline population set to 0 with values above the baseline representing an increase in local population abundance and values below the baseline representing a decline. Values represent the change in local abundance over the full time period based on the mean multiplicative slope of the multi-species trend with its SE (error bars). See the full abundance trends over time for each combination of elevational band and species group in Appendix (Figure S4, Table S4).



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Elevation gradient (m a.s.l.)

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