1	Protected areas act as a	buffer against	detrimental	effects of	climate change -	- evidence
-	I Totette al cab act ab a	build against	ucu miciitui	cifecto of	chinate change	crucifice

- 2 from large-scale, long-term abundance data
- 3 Running head: Protected areas buffer against climate change
- 4
- 5 Petteri Lehikoinen<sup>1,2</sup>\*, Andrea Santangeli<sup>1</sup>, Kim Jaatinen<sup>3</sup>, Ari Rajasärkkä<sup>4</sup>, Aleksi
- 6 Lehikoinen<sup>1</sup>
- <sup>7</sup> <sup>1</sup> The Helsinki Lab of Ornithology, Finnish Museum of Natural History, P.O. Box 17, 00014
- 8 Helsinki University, Finland; <sup>2</sup> Department of Biology, University of Lund, Sölvegatan 37,
- 9 223 62 Lund, Sweden; <sup>3</sup> Nature and Game Management Trust Finland, Degerbyvägen 176,
- 10 10160 Degerby, Finland; <sup>4</sup> Metsähallitus, National Parks Finland, P.O. Box 81, 90101 Oulu,
- 11 *Finland*.
- 12
- 13 \*Corresponding author: petteri.lehikoinen@helsinki.fi, tel. +358407233383
- 14 Keywords: conservation biology, distribution area, global warming, habitat management, land
- 15 use changes, monitoring
- 16 Paper type: Primary Research Article
- 17
- 18

19 Abstract

Climate change is driving species to shift their distributions towards high altitudes and 20 latitudes, while habitat loss and fragmentation may hamper species ability to follow their 21 22 climatic envelope. These two drivers of change may act in synergy, with particularly 23 disastrous impacts on biodiversity. Protected areas, PAs, may thus represent crucial buffers against the compounded effects of climate change and habitat loss. However, large-scale 24 25 studies assessing the performance of PAs as such buffers remain scarce and are largely based on species occurrence data. Conversely, abundance data have proven to be more reliable for 26 addressing changes in wildlife populations under climate change. We evaluated changes in 27 28 bird abundance from the 1970s–80s to the 2000s inside and outside PAs at the trailing range edge of 30 northern bird species and at the leading range edge of 70 southern species. 29 Abundances of retracting northern species were higher and declined less inside PAs at their 30 trailing range edge. The positive effect of PAs on bird abundances was particularly marked in 31 northern species that rely strongly on PAs, i.e. their density distribution is largely confined 32 33 within PAs. These species were nearly absent outside protected areas in the 2000s. The abundances of southern species were in general lower inside PAs and increased less from the 34 70s–80s to 2000s. Nonetheless, species with high reliance on PAs had much higher 35 36 abundances inside than outside PAs in the 2000s. These results show that PAs are essential in mitigating the retraction of northern species, but also facilitate northward expansions of 37 southern species highly reliant on PAs. Our study provides empirical evidence documenting 38 the role of PAs in facilitating species to adjust to rapidly changing climatic conditions, 39 40 thereby contributing to the mitigation of impending biodiversity loss. PAs may thus allow 41 time for initiating wider conservation programs on currently unprotected land.

### 43 Introduction

44

Anthropogenic climate change has influenced all the ecosystems on Earth (Scheffers et al., 45 2016) and is projected to reduce global biodiversity remarkably (Bellard *et al.*, 2012). Climate 46 change is shifting climatic conditions of species and ecosystems towards higher latitudes and 47 altitudes and species are tracking these changing conditions by shifting their distributions 48 49 (Chen et al., 2011). However, habitat loss and fragmentation driven by anthropogenic land use may impede the ability of species to follow their climatic envelopes (Robillard et al., 50 2015). Indeed, the joint effects of climate change and habitat loss are expected to be 51 disastrous for biodiversity (Oliver et al., 2017, Travis, 2003). Species inhabiting high latitudes 52 and high altitudes may be particularly exposed to the consequences of climate change because 53 they may run out of space into which they can retreat, leading to increased extinction risk 54 (Pacifici et al., 2017). This effect may further be compounded if the arctic and boreal biomes 55 56 at high latitudes warm up at a higher rate than regions at lower latitudes (IPCC, 2013). 57 Evidence is mounting that this may indeed be the case, since boreal and montane species in northern Europe have shifted their distributions faster than southern species (Virkkala & 58 Lehikoinen, 2014), and exhibit rapidly decreasing populations (Lehikoinen et al., 2014, Post 59 et al., 2009). 60

Protected areas (hereafter PAs) are among the most important means for preserving habitats and ensuring species persistence (Watson *et al.*, 2014). Yet their role in mitigating the impacts of climate change has been questioned (Monzón *et al.*, 2011). The main limitations of the current PA network for ensuring nature protection under dynamic and asymmetrical climate change are the static nature of the network, the spatial bias and low coverage (Monzón *et al.*, 2011). These shortcomings relate to species moving outside the network of PAs under shifting

67	climatic conditions (Hannah et al., 2007). There is however evidence that PAs are important
68	for conserving rare species and for the overall maintenance of biodiversity under climate
69	change (Thomas & Gillingham, 2015). PAs have also been found to facilitate species range
70	expansions (Thomas et al., 2012), and are suggested to help species to persist on trailing
71	edges of contracting or moving distributions (Gillingham et al., 2015b). However, the
72	evidence for this is restricted to narrow geographical areas and very few habitats.
73	Understanding the ecological, physiological, genetic and biogeographical mechanisms
74	underpinning species range shifts is fundamental for designing effective conservation
75	strategies and adaptations to climate change (Bonebrake et al., 2018)
76	Most findings concerning changes in species distributions under climate change are based on
77	occurrence data, and although abundance data are limited for many taxa globally, recent
78	evidence shows the increased reliability in generating outcomes for conservation prioritization
79	when abundance data are used (Howard et al., 2014, Johnston et al., 2015). Moreover, many
80	studies have concentrated on projecting future impacts, however attention is also required on
81	studying the current impacts of climate change on species, and for validating and improving
82	projections of future impacts (Pacifici et al., 2017). There is thus a need for studies using
83	large-scale long-term abundance data of wide taxonomic breadth for studying past and present
84	impacts of climate change on species distributions.
85	To heed this call, we studied the impact of PAs in aiding the expansion of populations of

southern species and in maintaining the densities of retracting populations of northern species
under climate change. To achieve this, we used nation-wide Finnish land bird density data
from two time periods, namely 1970–1989 and 2000–2014, available from within and outside
PAs. We evaluated factors affecting bird abundances in the two periods separately for the
leading (i.e. northern) range edge of 70 southern species and for the trailing (i.e. southern)
range edge of 30 northern species (Fig. 1; Table S1).

Based on previous studies conducted using occurrence data (Gillingham et al., 2015b, Hannah 92 93 et al., 2007, Monzón et al., 2011, Thomas et al., 2012, Watson et al., 2014), we hypothesize that abundance changes between the two time periods at both trailing and leading range edges 94 95 are more positive inside PAs than outside. This means that (i) abundances of northern species, the ranges of which are retracting toward the north, will have declined less inside protected 96 areas than outside on the trailing range edge. In addition, we expect that (ii) on the leading 97 98 range edge of southern species the abundance increases would be greater inside protected areas than outside. However, since species differ in their reliance on PAs we hypothesize the 99 presence of an interaction between PA reliance and protection status of a given site (protected 100 101 or unprotected), especially since species less reliant on protection may thrive also outside PAs (Fraixedas et al., 2015). The effect of protection is therefore predicted to be stronger for 102 species highly reliant on site protection, whereas species less reliant to PAs, e.g. farmland 103 104 birds in Finland, exhibit a weaker response to site protection.

105

106 Materials & methods

## 107 Line transect data

108 We used Finnish land bird line transect data spanning a period from the early 1970s to the present day (Virkkala & Lehikoinen, 2014). Transects were 3-6 kilometers long and their 109 locations were pre-set. Transects were surveyed once a year by foot, walking at an average 110 111 speed of 1 km / 45 min. The survey period was between the 21st of May and the 20th of June in South-Finland and between the 10th and the 30th of June in North-Finland. In montane 112 areas, the surveying continued until the 5th of July. Each transect was surveyed in the early 113 morning under favorable conditions for detection of birds, and all observed bird individuals 114 were counted. For each observation, the state and behavior of the bird was recorded (singing, 115

calling, flying, sex, age, nest, brood and flock-size) as detailed as possible. Based on this 116 117 information, all observations were transformed into pairs, which is the census unit (Virkkala & Lehikoinen, 2014). A single individual was always counted as one pair as was a nest and a 118 brood. For flocks of adult birds, the number of individuals was divided by two for 119 transformation into pairs. If the sexes were distinguishable in a flock of adult birds the 120 number of individuals of the more abundant sex was used as the number of pairs. For flocks 121 122 of early breeding passerines, where age and sex could not be identified, the number of individuals were divided by five for transformation into pairs, because flocks may include 123 broods. Thus, the unit of abundance for each species was the number of pairs for censused 124 125 amount of kilometers. This methodology is appropriate to obtain robust relative abundances of species over large areas (Järvinen & Väisänen, 1975, Lehikoinen & Virkkala, 2016). 126 Each transect was classified according to its i) protection status; falling either inside or 127 outside PAs, and ii) time period; either 70s-80s or 2000s according to the year of survey of 128 either 1970–1989 or 2000–2014, respectively (Santangeli et al., 2017). All monitored PAs 129 130 belonged to IUCN protected area categories I-IV (Dudley, 2008). The time of PA establishment is of low importance for the purpose of this study because PAs are typically 131 established in pristine or relatively good quality habitats, such as old-growth forests, 132 133 unditched forest mires or large and wet open mires that have been in a similar state for several decades. Therefore, even the most recently established PAs are comparable in terms of habitat 134 quality with the PAs established decades ago. Transects running across PA boundaries were 135 excluded to avoid edge effects (Santangeli et al., 2017). To account for spatial changes in 136 137 densities between the two time periods (see above), we divided the country into 100 km x 100 138 km squares (hereafter squares) following the Finnish national uniform coordinate system. This resolution represents an optimal tradeoff between the need to retain variation in the data 139 at high enough resolution to depict spatial patterns in abundance, while at the same time 140

allowing large enough sample sizes within each square in the different periods (the same
methodology in Santangeli *et al.*, 2017). Only squares that had line transect data from both
periods and both protection statuses were included in the study. This resulted in 40 squares
with sufficient line transect data and broad spatial coverage throughout the country
(Santangeli et al., 2017; Fig. 1).

146

# 147 Definition of range edges and species-specific measures

148 We studied 128 species of land birds having at least 20 observations during both periods 149 (Lehikoinen & Virkkala, 2016). Each of the species studied were classified as either southern or northern species based on their density distribution in the 70s-80s (Lehikoinen & 150 Virkkala, 2016). This period was chosen for edge definition as it is the starting point for 151 152 studying the population density changes between the time periods of the study. Since the observed population changes in North Europe have been in line with predictions based on 153 climatic suitability (Jiguet et al., 2013, Virkkala et al., 2014a), we contend that the witnessed 154 abundance changes in the defined range edges could be driven mainly by climate change. This 155 contention may be especially true inside PAs, where habitat quality has remained unaltered 156 157 (Virkkala & Rajasärkkä, 2011).

A species was classified as southern if its central gravity of density (introduced in
(Lehikoinen & Virkkala, 2016)) was situated in the southern half of the country and if the
species distribution exhibited a northern edge within the country (i.e. zero density at least in
the northernmost latitudinal row of squares). Conversely, a species was classified as northern
if its central gravity of density was situated in the northern half of the country and if the
species distribution exhibited a southern edge within the country. This yielded a total of 70
southern and 30 northern species (Table S1). 28 of the 128 species studied did not fulfill

either of these criteria, because they occurred throughout the country, and were therebyexcluded from the study.

The range edges of southern and northern species were defined using a percentage of the 167 168 cumulative density sums in the first period, using data from both inside and outside PAs (Fig. 1). To do this, we summed the densities of all squares in latitudinal rows of squares 169 (hereafter rows) and used the cumulative density sums of these rows, starting from the 170 171 southernmost row on southern species and northernmost row on northern species. For northern species, the trailing (i.e. southern) range edge of the distribution was defined as 172 starting from the row where the cumulative density sum of the species in question reached at 173 174 least 95% of the total density of the species and ending at the southernmost row (Fig. 1). For southern species, the leading (i.e. northern) range edge was defined as starting from the row 175 where the cumulative density sum of the species in question reached at least 95% of the total 176 density of the species. The range edge was defined as ending at the northernmost row (Fig. 1). 177 178 We calculated a PA reliance index, hereafter PA reliance, for each species by dividing the 179 mean density within PAs by the overall mean density of that species in Finland. In this calculation we included both time periods but excluded the range edges to avoid circularity. 180 The PA reliance ranged from 1 to 0 where the extreme values corresponded to all and none of 181 the densities of the species being observed inside PAs, respectively. The PA reliance gives a 182 rough estimate of the importance of PAs for the specific bird species. Thus, PA reliance was 183 used to explore the relevance of PAs for the distribution changes of each species in the study. 184 Because PA reliance is most certainly affected by habitat preferences, species were also 185 divided into four categories based on the dominant habitat in which they are found: 1. 186 farmland-urban, 2. forest, 3. wetlands (including rocky outcrops and peatlands) and 4. 187 montane (Lehikoinen & Virkkala 2016). 188

#### 189 Protected areas in Finland

Approximately 78% of the Finnish land area is forested (EEA 2015) and only 12% of the 190 forested land is protected, whereas forestry is operated in the rest of the area (Natural 191 192 Resource Institute Finland, 2016). Logging is prohibited in protected areas and they typically support older and more layered forest, as well as higher volumes of dead wood as compared 193 to unprotected forest lands (Metsähallitus, 2018). Peatlands cover 28% of the Finnish land 194 195 area, out of which 19% are open mires and 81% forest mires. Of the original peatland area c. 60 % has been drained by ditching for forestry purposes (Fraixedas et al., 2017), resulting in 196 pristine forest mires being rare outside PAs. Only 13% of the Finnish peatlands are protected. 197 198 These peatland PAs consist of pristine or relatively good quality habitats including large open mires and the unditched forest mires surrounding them. PAs have also been established to 199 protect threatened habitats including traditional rural habitats, coastal habitats and inland 200 shore meadows (Raunio et al., 2008). In general, major land use actions that could damage 201 nature are strictly prohibited inside PAs. In most PAs recreational activities, e.g. berry and 202 203 mushroom picking and hiking are allowed, while limitations on camping, hunting, use of motor vehicles as well as access to especially sensitive bird nesting areas are in place in many 204 PAs (Metsähallitus, 2018). 205

206

### 207 Statistical analyses

To study density changes at the distribution edges we used generalized linear mixed models (GLMM) separately for southern and northern bird species. The response variable in the models was the observed number of pairs in a given square, period (70s–80s vs 2000s) and protection status (within vs outside PAs). The sample sizes for each species per period and protection status are shown in Table S1. The total sample size for the analysis on northern species was 964, i.e. 241 unique species by square sample units for each combination of
period and status. The total sample size for southern species was 3212, consisting of 803
unique species by square sample units.

216 Squares where a single species was not observed on either of the protection statuses and either of the periods were removed from the dataset. This was done to reduce the number of zeros in 217 the dataset. The species-specific number of squares with data on both periods and protection 218 219 statuses are presented in Table S1. Despite removing the squares with no observations, the 220 datasets on both northern and southern species contained marked amounts of cases with zero pairs observed. 46.1% and 41.1% of all cases represented zero pairs observed for datasets on 221 222 northern and southern species, respectively. The residuals of GLMMs with a logarithmic linkfunction and Poisson distribution showed considerable overdispersion, as depicted by the ratio 223 of Pearson residual deviance to degrees of freedom: 5.91 and 6.91 for models describing 224 northern and southern species, respectively. Since Pearson residuals did not show any 225 systematic patterns against all covariates (Zuur et al., 2009), GLMMs with negative binomial 226 227 distribution (function glmmadmb in R (Fournier et al., 2012)) were used to account for the observed overdispersion. 228

The period was used as an explanatory factor to assess the changes in abundances between 229 70s-80s and 2000s. To assess the effect of protection status and species reliance on PAs, 230 protection status and standardized and centered PA reliance were used as explanatory 231 variables. To account for decreasing abundances towards higher latitudes and longitudes, 232 standardized and centered mean latitude and mean longitude of the squares were set as 233 explanatory variables. Square and protection status specific log-transformed total transect line 234 length was used as an offset in the model, in order to account for sampling effort (see 235 Santangeli et al., 2017). This is important because neglecting sampling effort can cause biases 236 in the analyses especially when presence-absence data are used (Kujala et al., 2013). 237

Transects in PAs covered a total of 3590 kilometers in the first and 10330 kilometers in the second period. Transects outside PAs totaled 5119 kilometers in the first and 9521 kilometers in the second period. In the models, log-transformed total length of the transect lines, latitude, longitude and protected area reliance were continuous variables, the response variable was an integer and protection status and period categorical variables. No strong correlations were detected among the explanatory variables; all Pearson correlation coefficients were below 0.5 (Booth *et al.*, 1994).

245 We built competing models to investigate which explanatory variables had the largest impact on abundances. We included mean latitude and longitude of the squares and period to all the 246 247 models except the simplest intercept-only model. In addition to these fixed explanatory variables the competing models consisted of all the possible subsets of explanatory variables 248 and interactions between period, protection status and PA reliance. Thus, the most complex 249 model contained the main effect of latitude, longitude, protected area reliance, protection 250 251 status and period as well as all two-way interactions and the three-way interaction between 252 protected area reliance, protection status and period.

In all models, we included square and species identity as crossed random factors to account 253 for the lack of independence and potential pseudoreplication arising from repeated 254 255 observations within squares and species. In order to account for the fact that species vary in 256 their densities between PAs and non-PAs, the PA status of the transect was included as random slope in the models that included the main effect of PA status. Species with close 257 phylogenetic relations are more likely similar in their responses to climate change than distant 258 species. We accounted for these phylogenetic relationships in the analysis by running three 259 260 models identical to the most complex competing model (described above) and adding three different levels of phylogenetic grouping. These three models were ranked using AICc values 261 only (Akaike, 1974, Sugiura, 1978) and the level of phylogeny in the model with lowest AICc 262

value was chosen for all the competing models. The phylogenetic grouping levels were:
species alone, species nested within family and species nested within order. The taxonomy
used followed that of IOC World Bird List (Gill & Donsker, 2018). For both northern and
southern species the model with species alone had the lowest AICc values (Tables S2 & S3).
Therefore, these phylogenetic levels were used in all of the competing models. Square,
species and family were categorical variables in the models.

269 Because PA reliance may be driven by habitat use, we also included competing models where PA reliance was replaced with a habitat class variable. Finally, the 26 competing models were 270 evaluated according to their AICc values for both northern and southern species 271 272 (Tables S4 & S5). The model with the lowest AICc value was chosen as the best to explain bird densities at the distribution edges. However, if several models were ranked with  $\Delta AICc \leq$ 273 2, we first investigated whether the top ranked model was more parsimonious than the other 274 models within  $\triangle AICc \le 2$  (i.e. the other models included more uninformative parameters 275 where the 85% confidence interval included zero; (Arnold, 2010)). If the top ranked model 276 277 was less parsimonious, we proceeded with multimodel inference and model averaging (using the R package MuMin (Bartón, 2018)) over the top-ranked models showing no uninformative 278 parameters. 279

The residuals of both models used for explaining abundance of northern and southern bird species on range edges were inspected for the presence of potential spatial autocorrelation, and that of unexplained patterns (Zuur *et al.*, 2009), but none were detected. All statistical analyses were performed with R software version 3.4.4 (R Core Development Team, 2017).

284

285

287 Results

For northern species three top ranked models exhibited differences in AICc-values of less 288 than two units (Table S4). The third ranked model contained the uninformative parameter for 289 290 the three-way interaction between period, PA reliance and PA status. The 85% confidence interval for this parameter ranged from -0.11 to 0.47 (NBGLMM:  $0.18 \pm 0.20$ , z=-0.89, 291 p=0.37). Model averaging was therefore conducted using the two AICc top-ranked models 292 293 that did not contain uninformative parameters. This revealed that the parameter for the interaction between period and protected area reliance, present in the top-ranked model, was 294 uninformative (85% CI: -0.29–0.05; NBGLMM: -0.12  $\pm$  0.12, z=-0.98, p=0.33). For inference 295 296 regarding northern species we therefore chose the simplest, second-ranked model. This model contained the following variables: latitude, longitude, period, protection status, protected area 297 reliance, the interaction between period and protection status and that between protection 298 status and protected area reliance (Tables 1 & S6). 299

Abundances of northern species on their trailing range edge were higher inside PAs (Table 1). Bird abundances decreased with increasing protected area reliance, but an interaction between protection status and PA reliance indicated that the decrease was much less steep inside PAs than outside (Fig. 2a). Moreover abundances of northern species on their trailing range edge declined from the 1970s–80s to the 2000s, and an interaction between protection status and period showed that the change was less negative inside PAs than outside (Fig. 2b).

For southern species, two top AICc-value ranked models exhibited an AICc difference of less
than two units (Table S5). The second-ranked model contained one uninformative parameter
more than the top-ranked model. This parameter, the three-way interaction between period,

protections status and protected area reliance, was uninformative (85% CI: -0.15–0.17;

NBGLMM:  $0.01 \pm 0.11$ , z=0.08, p=0.94). Thus the simpler, top-ranked model was used for

inference for southern species. This model included the following variables: latitude, 311 312 longitude, period, protection status, protected area reliance and the interactions between period and protection status, between protection status and protected area reliance and 313 between period and protected area reliance (Tables 2 & S7). 314 315 Southern species on their leading range edge were less abundant inside than outside PAs (Table 2). Protection status however exhibited interactions with PA reliance and period. 316 317 Inside PAs bird abundances increased markedly with increasing PA reliance (Fig. 2c), whereas they were not affected by PA reliance outside of PAs (Table 2). However, PA 318 reliance interacted positively with time period, showing that bird abundances increased with 319 320 increasing PA reliance from the 1970s–80s to the 2000s (Fig. 2d). Abundances of southern species on their leading range edge showed an overall increase from the 1970s–80s to the 321 2000s (Fig. 2d). The interaction between protection status and period showed that this 322 increase was not as marked inside PAs as outside (Table 2, Fig. 2d). 323

324

325 Discussion

The results show that the abundances of northern species at their trailing range edges have 326 327 declined less inside than outside protected areas, with the abundances of especially species with high PA reliance remaining higher inside than outside PAs. While the latter finding may 328 was partly expected due to generally higher reliance on PAs among northern species (mean 329 PA reliance  $0.63 \pm 0.14$  SD), it underlines the fact that species with the highest reliance on 330 protection were nearly absent outside PAs in the 2000s. This may be explained by the high 331 proportion of old-growth boreal forests in the Finnish protected areas which has been shown 332 to have positive effects on the specialization of the avian community (Häkkilä et al., 2017). 333 334 Our findings therefore emphasize the importance of PAs in mitigating declines of boreal bird

species under climate change. This result is particularly relevant in light of climate change in 335 336 the boreal biome being predicted to be much more rapid than that in other regions, carrying serious consequences for ecosystems and associated wildlife (Loarie et al., 2009). Despite the 337 fact that northern PAs are experiencing among the most rapid rates of climate change (Loarie 338 et al., 2009) they are able to slow down the retraction of species ranges, albeit not fully 339 prevent it. Our results add to the accumulating evidence underscoring the importance of PAs 340 341 which have previously been shown to aid persistence of northern species at temperate region by studying occurrences (Gillingham et al., 2015a) and to facilitate the adaptation of northern 342 species to temperature changes (Gaüzère et al., 2016), as well as support more cold-dwelling 343 344 bird communities (Santangeli et al., 2017) by investigating changes in the thermal signature of bird communities (Devictor et al., 2008). The findings suggest that PAs serve as refuge for 345 northern species of which many are boreal biome specialist facing high pressure from a 346 347 rapidly warming climate (Loarie et al., 2009, Pacifici et al., 2017). The findings are also in line with earlier results showing that the probability of occurrence among northern species 348 349 increased with increasing cover of protected area in a 10 km x 10 km atlas square (Virkkala et al., 2014b). However, that study did not separate surveys conducted inside and outside PAs, 350 351 and used presence-absence data without correcting for observation effort. In contrast, our 352 study explicitly accounted for varying survey effort and has thus been able to reveal significant changes in bird abundances. 353

Our findings regarding southern species show that abundances at the leading range edge were higher and increased more outside than inside PAs from the 1970s–80s to the 2000s, which contradicts our initial hypotheses. The average reliance on protected areas was low in southern species (mean protected area reliance  $(0.39 \pm 0.19 \text{ SD})$ , and the under-representation of species strongly reliant on PAs in the south may be an explanation for the lack of relationship between PA reliance and abundance outside PAs in the 70s–80s. However,

species that are highly reliant on PAs exhibited markedly higher densities inside PAs than 360 361 outside (Fig. 2d). This suggests that PAs are important for expanding populations of southern species with high PA reliance; a conclusion also reached by studies based on occurrence data 362 on birds and butterflies (Gillingham et al., 2015b) and abundance categories on odonates and 363 butterflies (Gillingham et al., 2015a). Species with a high reliance on PAs may use PAs as 364 stepping stones when moving into new areas (Hiley et al., 2013). However increasing PA 365 366 reliance affected abundances positively also outside PAs in the 2000s compared to the 70-80s. This may be the result of PA designation being biased towards boreal habitats. Thus, 367 species with high PA reliance are boreal biome specialists, which may find more suitable 368 369 habitat outside PAs when expanding their distributions from the temperate zone into the boreal zone. This result underscores a striking challenge for national conservation policy 370 posed by climate change, where the managing and expanding of the current PA network in 371 372 order to maintain habitat for northern species needs to be balanced with maintaining habitat important for the northward expansion of southern species. 373

374 Our large-scaled long-term study provides quantitative evidence on the performance of PAs under climate change and as the previous studies concentrate to the temperate region (e.g. 375 Gillingham et al., 2015a, Thomas et al., 2012, Watson et al., 2014), widens the current 376 377 knowledge to cover also boreal region. Besides increasing the geographical coverage of such studies to include one of the regions with highest velocities of climate change (Loarie et al., 378 379 2009), the large-scaled abundance data increases the reliability of our current view of the positive effects of PAs on range shifts under climate change (Howard et al., 2014, Johnston et 380 381 al., 2015). We must stress that the exact transect locations in this study have not necessarily 382 been the same between the study periods. However, we argue that while this may represent a 383 source of random noise across all the data, it is unlikely that it could bias the results given the

large spatial scale of the sampling units and the wide scale of the whole study where rangeedges also vary between species.

This long-term documentation of changes in bird abundances shows that PAs play a key role 386 387 in mitigating the loss of biodiversity under climate change (Johnston et al., 2013), PAs not only facilitate range expansions of species establishing into new areas (Gillingham *et al.*, 388 2015a, Thomas & Gillingham, 2015, Thomas et al., 2012), but also mitigate the retraction of 389 390 species shifting under climate change. Moreover, the existing PAs not only mitigate local extinctions of a large number of northern species, but also abundance losses are lower inside 391 PAs compared to outside. Besides facilitating range expansion, PAs contribute to the increase 392 393 of abundances in already occupied areas of southern species with high reliance on PAs. Climate change may however cause species to be excluded from current PAs due to shifts in 394 climate envelopes (Araújo et al., 2004). Therefore, some PAs may perform poorly in 395 conserving biodiversity under climate change (Araújo et al., 2011), especially given their 396 397 biased distribution, limited area coverage and variable management effectiveness (Watson et 398 al., 2014). As a result, the future of biodiversity conservation in the long term also strongly 399 relies on efforts put in place on currently unprotected land (Santangeli *et al.*, 2016). While 400 there are international commitments to expand the current PA network (Aichi target 11 in 401 (CBD, 2018, Montesino Pouzols et al., 2014), and implement effective biodiversity conservation actions on unprotected land (Santangeli et al., 2016), they require time and 402 403 resources, while biodiversity continues to decline at alarming rates. Under this scenario, our results represent an encouraging message for biodiversity conservation. We show that PAs 404 405 play an important role in mitigating impacts of climate change on biodiversity, providing 406 strongholds for species persistence in the short term, thereby allowing some time for wider protection efforts on unprotected land to start yielding positive effects. Moreover, by 407 buffering against the detrimental effects of climate change, protected areas can buy time 408

- 409 during which the causes of anthropogenic climate change may or may not be remedied.
- 410 Multidisciplinary work in the field of ecology, conservation and social science should
- 411 facilitate the implementation of solutions to mitigate the negative impacts of climate change
- 412 (Bonebrake et al. 2018).
- 413

414 Acknowledgements

- 415 We thank all people who have participated in bird transect surveys inside and outside
- 416 protected areas jointly administered by the Finnish Museum of Natural History and
- 417 Metsähallitus National Parks Finland. AL and AS received financial support from the
- 418 Academy of Finland (grants 275606 & 283664; and 307909, respectively) and PL from Maj
- and Tor Nessling foundation (grants 201600187, 201700195).
- 420
- 421 References
- 422 Akaike H (1974) A new look at the statistical model identification. IEEE Transactions on Automatic
   423 Control, 19, 716-723.
- 424 Araújo MB, Alagador D, Cabeza M, Nogués-Bravo D, Thuiller W (2011) Climate change threatens
   425 European conservation areas. Ecology Letters, 14, 484-492.
- Araújo MB, Cabeza M, Thuiller W, Hannah L, Williams PH (2004) Would climate change drive
   species out of reserves? An assessment of existing reserve-selection methods. Global Change
   Biology, 10, 1618-1626.
- Arnold TW (2010) Uninformative Parameters and Model Selection Using Akaike's Information
   Criterion. Journal of Wildlife Management, 74, 1175-1178.
- Bartón K (2018) MuMIn: Multi-Model Inference. R package version 1.40.4. https://CRAN.R project.org/package=MuMIn.
- Bellard C, Bertelsmeier C, Leadley P, Thuiller W, Courchamp F (2012) Impacts of climate change on
  the future of biodiversity. Ecology Letters, 15, 365-377.
- Bonebrake TC, Brown CJ, Bell JD *et al.* (2018) Managing consequences of climate-driven species
  redistribution requires integration of ecology, conservation and social science. Biological
  Reviews, 93, 284-305.
- Booth GD, Niccolucci MJ, Schuster EG (1994) Identifying proxy sets in multiple linear-regression an aid to better coefficient interpretation. Usda Forest Service Intermountain Research Station
  Research Paper, 1-13.
- 441 CBD (2018) Aichi Biodiversity Targets. https://www.cbd.int/sp/targets/.
- 442 Chen IC, Hill JK, Ohlemüller R, Roy DB, Thomas CD (2011) Rapid Range Shifts of Species
  443 Associated with High Levels of Climate Warming. Science, 333, 1024-1026.

- 444 Devictor V, Julliard R, Couvet D, Jiguet F (2008) Birds are tracking climate warming, but not fast
   445 enough. Proceedings. Biological sciences, 275, 2743-2748.
- 446 Dudley N (ed) (2008) *Guidelines for Applying Protected Area Management Categories. Gland*,
  447 Switzerland: IUCN. WITH Stolton, S., P. Shadie and N. Dudley (2013). IUCN WCPA Best
  448 Practice Guidance on Recognising Protected Areas and Assigning Management Categories
  449 and Governance Types., Gland, Switzerland, IUCN.
- 450 EEA European Environmental Agency (2015) Land use State and impacts (Finland). Retrieved
   451 from <a href="https://www.eea.europa.eu/soer/countries/fi/land-use-state-and-impacts-finland">https://www.eea.europa.eu/soer/countries/fi/land-use-state-and-impacts-finland</a>
- Fournier DA, Skaug HJ, Ancheta J *et al.* (2012) AD Model Builder: using automatic differentiation for
   statistical inference of highly parameterized complex nonlinear models. Optimization
   Methods & Software, 27, 233-249.
- 455 Fraixedas S, Lehikoinen A, Lindén A (2015) Impacts of climate and land-use change on wintering
  456 bird populations in Finland. Journal of Avian Biology, 46, 63-72.
- Fraixedas S, Lindén A, Meller K *et al.* (2017) Substantial decline of Northern European peatland bird
   populations: Consequences of drainage. Biological Conservation, 214, 223-232.
- Gaüzère P, Jiguet F, Devictor V (2016) Can protected areas mitigate the impacts of climate change on
   bird's species and communities? Diversity and Distributions, 22, 625-637.
- 461 Gill F, Donsker DE (2018) IOC World Bird List (v 8.2). DOI: 10.14344/IOC.ML.8.2.
- Gillingham PK, Alison J, Roy DB, Fox R, Thomas CD (2015a) High Abundances of Species in
   Protected Areas in Parts of their Geographic Distributions Colonized during a Recent Period
   of Climatic Change. Conservation Letters, 8, 97-106.
- Gillingham PK, Bradbury RB, Roy DB *et al.* (2015b) The effectiveness of protected areas in the
   conservation of species with changing geographical ranges. Biological Journal of the Linnean
   Society, 115, 707-717.
- Hannah L, Midgley G, Andelman S *et al.* (2007) Protected area needs in a changing climate. Frontiers
   in Ecology and the Environment, 5, 131-138.
- Hiley JR, Bradbury RB, Holling M, Thomas CD (2013) Protected areas act as establishment centres
  for species colonizing the UK. Proceedings of the Royal Society B-Biological Sciences, 280,
  7.
- Howard C, Stephens PA, Pearce-Higgins JW, Gregory RD, Willis SG (2014) Improving species
  distribution models: the value of data on abundance. Methods in Ecology and Evolution, 5,
  506-513.
- Häkkilä M, Le Tortorec E, Brotons L, Rajasärkkä A, Tornberg R, Mönkkönen M (2017) Degradation
  in landscape matrix has diverse impacts on diversity in protected areas. Plos One, 12, 17.
- 478 IPCC (2013) Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to
  479 the Fifth Assessment Report of the Intergovernmental Panel on Climate Change [Stocker,
  480 T.F., D. Qin, G.-K. Plattner, M. Tignor, S.K. Allen, J. Boschung, A. Nauels, Y. Xia, V. Bex and
  481 P.M. Midgley (eds.)], Cambridge, United Kingdom and New York, NY, USA, Cambridge
  482 University Press.
- Jiguet F, Barbet-Massin M, Devictor V, Jonzén N, Lindström Å (2013) Current population trends
   mirror forecasted changes in climatic suitability for Swedish breeding birds. Bird Study, 60,
   60-66.
- Johnston A, Ausden M, Dodd AM *et al.* (2013) Observed and predicted effects of climate change on
   species abundance in protected areas. Nature Climate Change, 3, 1055-1061.
- Johnston A, Fink D, Reynolds MD *et al.* (2015) Abundance models improve spatial and temporal
   prioritization of conservation resources. Ecological Applications, 25, 1749-1756.
- Järvinen O, Väisänen RA (1975) Estimating relative densities of breeding birds by the line transect
   method. Oikos, 26, 316-322.
- Kujala H, Vepsäläinen V, Zuckerberg B, Brommer JE (2013) Range margin shifts of birds revisited –
   the role of spatiotemporally varying survey effort. Global Change Biology, 19, 420-430.
- Lehikoinen A, Green M, Husby M, Kålås JA, Lindström Å (2014) Common montane birds are
   declining in northern Europe. Journal of Avian Biology, 45, 3-14.
- 496 Lehikoinen A, Virkkala R (2016) North by north-west: climate change and directions of density shifts
  497 in birds. Global Change Biology, 22, 1121-1129.

- Loarie SR, Duffy PB, Hamilton H, Asner GP, Field CB, Ackerly DD (2009) The velocity of climate
   change. Nature, 462, 1052-1055.
- 500 Metsähallitus (2018) Hiking in Finland. Retrieved from http://www.nationalparks.fi/en/hikinginfinland
- Montesino Pouzols F, Toivonen T, Di Minin E *et al.* (2014) Global protected area expansion is
   compromised by projected land-use and parochialism. Nature, **516**, 383-386.
- Monzón J, Moyer-Horner L, Palamar MB (2011) Climate Change and Species Range Dynamics in
   Protected Areas. Bioscience, 61, 752-761.
- Natural Resource Institute Finland (2016) Metsien suojelu 1.1.2016. Retrieved from
   <a href="http://stat.luke.fi/metsien-suojelu-112016\_fi">http://stat.luke.fi/metsien-suojelu-112016\_fi</a>
- Oliver TH, Gillings S, Pearce-Higgins JW *et al.* (2017) Large extents of intensive land use limit
   community reorganization during climate warming. Global Change Biology, 23, 2272-2283.
- Pacifici M, Visconti P, Butchart SHM, Watson JEM, Cassola FM, Rondinini C (2017) Species' traits
   influenced their response to recent climate change. Nature Climate Change, 7, 205-208.
- 511 Post E, Forchhammer MC, Bret-Harte MS *et al.* (2009) Ecological Dynamics Across the Arctic
  512 Associated with Recent Climate Change. Science, **325**, 1355-1358.
- R Core Development Team (2017) R: A language and environment for statistical computing, version
   3.4.1. R Foundation for Statistical Computing, Vienna.
- Raunio A, Schulman A, Kontula T (eds.) (2008) Suomen luontotyyppien uhanalaisuus. Suomen ympäristö 8/2008. Osat 1 ja 2. Helsinki, Suomen ympäristökeskus.
- Robillard CM, Coristine LE, Soares RN, Kerr JT (2015) Facilitating climate-change-induced range
   shifts across continental land-use barriers. Conservation Biology, 29, 1586-1595.
- Santangeli A, Arroyo B, Dicks LV, Herzon I, Kukkala AS, Sutherland WJ, Moilanen A (2016)
   Voluntary non-monetary approaches for implementing conservation. Biological Conservation,
   197, 209-214.
- Santangeli A, Rajasärkkä A, Lehikoinen A (2017) Effects of high latitude protected areas on bird
   communities under rapid climate change. Global Change Biology, 23, 2241-2249.
- Scheffers BR, De Meester L, Bridge TCL *et al.* (2016) The broad footprint of climate change from
   genes to biomes to people. Science, **354**.
- Sugiura N (1978) Further analysis of the data by Akaike's information criterion and the finite
   corrections. Communications in Statistics Theory and Methods, 7, 13–26.
- Thomas CD, Gillingham PK (2015) The performance of protected areas for biodiversity under climate
   change. Biological Journal of the Linnean Society, 115, 718-730.
- Thomas CD, Gillingham PK, Bradbury RB *et al.* (2012) Protected areas facilitate species' range
  expansions. Proceedings of the National Academy of Sciences of the United States of
  America, 109, 14063-14068.
- Travis JMJ (2003) Climate change and habitat destruction: a deadly anthropogenic cocktail.
  Proceedings of the Royal Society B-Biological Sciences, 270, 467-473.
- Watson JEM, Dudley N, Segan DB, Hockings M (2014) The performance and potential of protected
  areas. Nature, 515, 67-73.
- 537 Virkkala R, Heikkinen RK, Lehikoinen A, Valkama J (2014a) Matching trends between recent
   538 distributional changes of northern-boreal birds and species-climate model predictions.
   539 Biological Conservation, 172, 124-127.
- Virkkala R, Lehikoinen A (2014) Patterns of climate-induced density shifts of species: poleward shifts
   faster in northern boreal birds than in southern birds. Global Change Biology, 20, 2995-3003.
- 542 Virkkala R, Pöyry J, Heikkinen RK, Lehikoinen A, Valkama J (2014b) Protected areas alleviate
  543 climate change effects on northern bird species of conservation concern. Ecology and
  544 Evolution, 4, 2991-3003.
- 545 Virkkala R, Rajasärkkä A (2011) Climate change affects populations of northern birds in boreal
  546 protected areas. Biology Letters, 7, 395-398.
- Zuur A, Ieno E, Walker N, Saveliev A, Smith G (2009) Mixed Effects Models and Extensions in
   Ecology with R, New York, NY, Springer.
- 549

Table 1. Summary table of the model explaining abundances of northern bird species on their trailing

552 range edges. Significant variables are bolded.

	Estimate	Std. error	z-value	Pr(> z )
(Intercept)	-5.27	0.36	-14.46	<0.001
Period 2000s	-0.94	0.15	-6.31	<0.001
Protected area	1.43	0.21	6.81	<0.001
PA reliance	-0.80	0.34	-2.34	0.02
Latitude	1.67	0.20	8.24	<0.001
Longitude	-0.10	0.10	-1.02	0.31
Period 2000s:Protected area	0.43	0.19	2.22	0.03
Protected area:PA reliance	0.64	0.18	3.58	<0.001

553

- Table 2. Summary table of the model explaining abundances of southern bird species on their leading
- range edges. Statistically significant variables are bolded.

	Estimate	Std. error	z-value	Pr(> z )
(Intercept)	-4.54	0.27	-16.62	<0.001
Period 2000s	0.77	0.07	11.13	<0.001
Protected area	-0.57	0.11	-5.39	<0.001
PA reliance	0.12	0.25	-0.53	0.60
Latitude	-1.53	0.09	-16.46	<0.001
Longitude	-0.14	0.09	-1.66	0.10
Period 2000s:Protected area	-0.33	0.10	-3.29	0.001
Period 2000s:PA reliance	0.20	0.06	3.56	<0.001
Protected area:PA reliance	1.07	0.09	11.84	<0.001

557



Figure 1. Definition of range edges of southern (red) and northern species (blue). Grey 560 squares represent the 100 km x 100 km squares from which abundance data was available for 561 both time periods (1970-89 & 2000-14) and for both protection statuses (protected & 562 unprotected), illustrating the spatial coverage of the data used. Densities of species in 1970-563 1989 are shown as coloured dots inside each square with a resolution of 50 km x 50 km where 564 the intensity of the colour represents the density as illustrated on the left side of the map. 565 Where the ranges between the northern and southern example species overlap, the density of 566 567 northern species is represented by the colour in the upper half of the dot and the density of southern species in the bottom half. Asterisks represent the central point of gravity of the 568 densities and black crosses missing data. The histograms on the side of the map represent the 569 sums of abundances in each 100 km wide latitudinal zones of squares. The latitude 720 570 divides the country in half and is, together with the central point of gravity, used to identify 571 species as either southern or northern. The range edges represent the northernmost (on 572 southern species; the blackbird in this example) or southernmost (on northern species; the 573 574 brambling) 5% of the total density of each species. The range edges for the example species are shown as barred areas. 575

576



577

Figure 2. Model estimates of bird abundance in response to protected area reliance and time period. Figures 2a,c show the estimated effects of protected area reliance on bird abundances inside protected areas (solid line) and outside protected areas (dashed line) during the second time period (2000s) of northern species and southern species, respectively; circles represent data points from protected areas and triangles those from unprotected areas. Shaded bands represent the 95% confidence intervals of the effect of protected area reliance on bird

- abundance. Figure **2b** shows model estimates of the effect of time period on densities of
- northern species (inside PAs: dark blue, outside PAs: light blue) for minimum (0.25; point-
- down triangles), mean (0.63; dots) and maximum (0.96; point-up triangles) PA reliances.
- 587 Figure 2d shows model estimates of the effect of time period on densities of southern species
- 588 (inside PAs: red, outside PAs: dark pink) for minimum (0.01; point-down triangles), mean
- 589 (0.39; dots) and maximum (0.74; point-up triangles) PA reliance indices. The whiskerers in
- 590 figures **2b**,**d** represent standard errors of the parameter estimates.

592 Data is deposited in Zenodo (DOI: 10.5281/zenodo.1422373).

593

- 594 Additional information
- 595 Supplementary information is available in a separate file.

596

- 597 Competing financial interests
- 598 The authors declare no competing financial interests.

599