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Microclimate affects the distribution of grassland birds, but not forest birds, in an Alpine environment

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1	Microclimate Affects the Distribution of Grassland Birds, but not Forest Birds,
2	in an Alpine Environment
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29 Abstract

30 Predictions derived from species distribution models (SDMs) are strongly influenced by the 31 spatial scale at which species and environmental data (e.g. climate) are gathered. SDMs of 32 mountain birds usually build on large-scale temperature estimates. However, the topographic 33 complexity of mountain areas could create microclimatic refuges which may alter species 34 distributions at small spatial scales. To assess whether fine-scale data (temperature and/or 35 topography) improve model performance when predicting species occurrence, we collected 36 data on presence-absence of bird species, habitat and fine-scale temperature at survey points 37 along an elevational gradient in the Alps (NW Italy). Large-scale temperature data, and both 38 large- and fine-scale topography data, were extracted from online databases for each point. 39 We compared species models (fine-scale vs large-scale) using an information-theoretic 40 approach. Models including fine-scale temperature estimates performed better than 41 corresponding large-scale models for all open habitat species, whereas most forest/ecotone 42 species showed no difference between the two scales. Grassland birds such as Northern 43 Wheatear Oenanthe oenanthe and Water Pipit Anthus spinoletta were positively associated 44 with warmer microclimates. These results suggest that alpine grassland species are potentially 45 more resistant to the impact of climate change than previously predicted, but that indirect 46 effects of climate change such as habitat shifts (forest- and shrub encroachment at high 47 elevations) pose a major threat. Therefore, active management of alpine grassland is needed 48 to maintain open areas and to prevent potential habitat loss and fragmentation. SDMs based 49 solely on large-scale temperatures for open habitat species in the Alps should be re-assessed.

50 51

52 Key words: information-theoretic approach, mountains, species distribution models,
53 temperature, topography.

55 Introduction

56 Species distribution models (henceforth SDMs) are a widely used tool in conservation (Guisan 57 and Thuiller 2005, Rodríguez et al. 2007, Franklin 2013) for a range of taxa (Ongaro et al. 58 2018, Lewthwaite et al. 2018, Hof and Allen 2019). In the face of climate change, SDMs have 59 become particularly important in predicting current and/or future distributions of species under 60 different climate change scenarios (Avalos and Herández 2015, Jackson et al. 2015, 61 Lehikoinen and Virkkala 2016). These studies usually rely on macroclimate data, which 62 describe climatic conditions at a relatively large scale (approximately one square kilometre or 63 more; Zellweger et al. 2019) derived from national networks, weather stations or online 64 databases (e.g Worldclim; Hijmans et al. 2005).

65 However, mountain environments are often poorly represented by conventional climate 66 station data, and uncertainty for interpolated climatic values is high (Hijmans et al. 2005). 67 Furthermore, local temperature can vary substantially due to the topographic complexity in 68 mountain areas (Scherrer and Körner 2010, Gunton et al. 2015), thus creating a mosaic of 69 microclimatic conditions over small spatial scales. Depending on discipline, microclimates 70 have been defined in various ways. In this study, we adopt the definition by Bramer et al. (2018) 71 who defined microclimate as fine-scale climate variations at spatial resolutions of < 100m, 72 which are influenced by fine-resolution biotic and abiotic variations (topography, soil type and 73 vegetation). Topographic variables like aspect and slope can markedly alter microclimate by 74 influencing the amount of incoming solar radiation between different exposed slopes. Between 75 north and south exposed slopes, temperature can differ by approximately 1°C if slopes are 76 gentle (<5°) but can increase up to 5°C if slopes are steep (40°; Gubler et al. 2011). Moreover, 77 these differences could subsequently influence snow accumulation processes and thus the 78 rate of snow melt in spring (Gubler et al. 2011).

There is mounting evidence of the importance of microclimate in influencing habitat selection. For example, Bramblings *Fringilla montifringilla* tend to rest in higher densities in areas with warm microclimatic conditions (Zabala et al. 2012). In Mountain Chickadees *Poecile gambeli*, microclimates influence the selection of foraging sites (Wachob 1996). Microclimates

can also act as thermal refuges, which enable individuals to persist despite unfavourable
ambient conditions (Wilson et al. 2015). This has been shown in Northern Bobwhites *Colinus virginianus*, which mitigated thermal stress by seeking thermally-buffered microclimatic sites
during hot days (Carroll et al. 2015). Furthermore, Northern Bobwhite nest site selection was
proven to be influenced by microclimate: Individuals nested in cooler and moister microclimatic
conditions compared to surrounding non-nesting locations (Tomecek et al. 2017, Carroll et al.
2018).

90 Only a few studies have investigated the role of microclimate within a mountain context. 91 Frey et al. (2016) showed that fine-scale temperature metrics were strong predictors of bird 92 distributions, with temperature effects being larger than vegetation effects on occupancy 93 dynamics in mountain forests (but see Viterbi et al. 2013). In the Alps, the habitat of the alpine 94 Rock Ptarmigan Lagopus muta helvetica is characterised by a wide variety of microclimates 95 over small spatial scales with individuals choosing colder sites in summer (Visinoni et al. 2015). 96 Beside the direct impact on birds, microclimate also plays a crucial role in habitat selection 97 in insects. It has been demonstrated that in *Parnassius apollo*, a mountain specialist butterfly, 98 larval habitat selection is related to ambient temperature. Larvae selected warm microclimates 99 when ambient temperatures fell below a threshold of 27°C, whereas cold microclimates were 100 selected when this threshold was exceeded (Ashton et al. 2009). Microclimate can further 101 influence oviposition (Stuhldreher et al. 2012), and the precise microclimatic conditions for 102 thermoregulation are actively sought by montane species of the genus *Erebia* (Kleckova et al. 103 2014). In this respect, microclimate won't only shape the distributions of these butterfly species, 104 but it will also indirectly influence bird species which rely on caterpillars as a food source for 105 chick rearing.

Microclimate thus has the potential to influence many aspects of an organism's life cycle. It could help to buffer or to compound the effects of climate change (Spasojevic et al. 2013). To assess the impact of climate change on current or future distributions of species it is crucial to gather climate data at the most appropriate scale in order to increase model accuracy (Barton et al. 2018, Randin et al. 2009). However, predictions for future geographic

111 distributions of mountain birds under a range of climate change scenarios have thus far been 112 based on models which have considered climate variables measured at large scales, usually 113 ca. 1 km² (Chamberlain et al. 2013, 2016, Brambilla et al. 2016, 2017a). Given the potential 114 for bird responses to microclimatic conditions in mountains (Frey et al. 2016, Visioni et al. 115 2014), it may be more appropriate to consider the role of climate measured at finer spatial 116 resolutions in determining mountain bird distributions. This is particularly important given that 117 environmental conditions in mountains typically change over very small spatial scales thanks 118 to steep elevation gradients (Scherrer and Körner 2010, Gunton et al. 2015).

119 In this study, we investigated the role of microclimate for a range of Alpine ecotone and 120 open habitat species. There were two specific aims. First, to evaluate if models including a 121 microclimatic variable (in this case temperature) show better performance than models using 122 large-scale climate estimates. This will inform future modelling studies, and should help to 123 improve predictions of future impacts of climate change on Alpine birds where microclimatic 124 effects are evident. Second, to assess if models including topographic variables (slope and 125 aspect) in combination with climatic variables (fine and large scale) increase model 126 performance. This will assess the extent to which topographic variables should be included in 127 SDMs of alpine bird species. Based on previous studies, which showed that microclimate can 128 influence bird distributions within mountain habitats (Frey et al. 2016, Visinoni et al. 2015), we 129 hypothesise that models using fine-scale temperature estimates will show better model 130 performance than models using large-scale temperature estimates.

131

132 Methods

133 Study area and point selection

The study was carried out in Val Troncea Natural Park (44°57'28" N; 6°56'28" E) in the western Italian Alps. At lower elevations, the area is dominated by larch *Larix decidua*. The natural treeline is typically found at around 2200 m asl, but varies depending on local conditions. Typical shrub species are *Juniperus nana* (henceforth Juniper) and *Rhododendron ferrugineum* (henceforth Rhododendron) which rapidly encroached wide areas of grasslands after the decline of agro-pastoral activities. Grasslands are mainly dominated by *Festuca curvula*, *Carex sempervirens*, and *Trifolium alpinum*. Scree and rocky areas occur
 predominantly at higher elevations, above approximately 2700 m asl.

Point counts were carried out along an elevational gradient ranging from 1750 m to 2820 m encompassing forest, ecotone and open habitats. Point count locations coincided with the centroids of a pre-existing grid at a scale of approximately 150 x 150 m (there was some variation, due to access constraints for example; Probo et al. 2014) along the western facing slope of the valley. All points were spaced a minimum of 200 m apart.

147

148 Bird surveys

Point counts (n = 221) were carried out from mid-May to mid-July 2017 following the methods of Bibby et al. (2000), using a 10 minute count period. At each point count location, all individual birds seen or heard were recorded within a 100 m radius (estimated with the aid of a laser range finder). Point counts commenced 1-1.5 h after sunrise and continued until 1200 h. Surveys did not take place in excessively wet or windy conditions. Each point count location was visited once.

155

156 Habitat data collection

157 At each point count location, habitat data were collected through the visual estimation of the 158 percentage cover of canopy (i.e. vegetation above head height), the dominant shrub species, 159 open grassland and bare rock (including scree and unvegetated areas) within a 100 m radius 160 of the point's centre. The dominant shrub species were defined into four groups: 161 Rhododendron, Juniper, bilberry (Vaccinium myrtillus and V. gaultherioides) and other (e.g. 162 Green Alder Alnus viridis, Willow Salix spp, and also including young trees less than two 163 meters in height, mostly European Larch Larix decidua). Furthermore, the number of mature 164 trees (greater than c. 20 cm in diameter at breast height) within a 50 m radius of a point count 165 location was counted. These estimates have been shown to correlate well with estimates of

166 land cover derived from remote sensing and have been used as the basis of predictive models

167 for several species considered here (Chamberlain et al. 2013, 2016, Jähnig et al. 2018).

168

169 Temperature measurements

170 At each point count location, temperature was measured with hygro buttons (Plug & Track[™]), 171 using methods based on Frey et al. (2016). Each button was stuck on the bottom of a small 172 plastic cup, which was attached upside down to a bamboo stick to protect the button against 173 wind, direct sunlight and water. Mean button height was 40.89 cm (min = 28 cm, max = 47174 cm). Hygro buttons were programmed to record temperature every 5 minutes. They were 175 placed 24 hours before a point count commenced and were collected 24 hours after the point 176 count ended, which resulted in a total recording time of 48 hours. At every hygro button 177 location, button height, distance to slope, substrate and canopy presence/absence was 178 recorded.

179

180

181 Statistical analysis

182 Temperature modelling

183 For each point count location, minimum, maximum and mean temperatures were derived over 184 the 48 hour recording period. All temperature measurements were checked for collinearity by 185 calculating Pearson's correlation coefficient. Mean temperature was strongly correlated with 186 both minimum (r = 0.80) and maximum temperature (r = 0.73) over the recording period. 187 Therefore, temperature modelling was undertaken with mean temperature values. The same 188 procedure was repeated for night-time temperatures. Minimum, maximum and mean night-189 time temperatures were obtained for the time period between 23.00 pm and 03.00 am over the 190 same recording period at each point. There was a strong positive correlation of mean night-191 time temperature between minimum (r = 0.97) and maximum night-time temperature (r = 0.89). 192 The objective of the first analysis was to model temperature in relation to date and 193 elevation. This model was then used to predict a standardised temperature at each point count

194 location, set at a fixed date, which was representative of the fine-scale temperature at that 195 point controlling for seasonal effects. This procedure provided data which was analogous to 196 the larger scale temperature data (see below). This standardised temperature was then used 197 subsequently as a variable in species distribution models. Note that all subsequent modelling 198 steps were performed separately for mean temperature and mean night-time temperature. 199 However, models with night-time temperature were very similar to those using mean 200 temperature, so we focus on the latter. Further details on night-time temperature models are 201 given in the Electronic Supplementary Material (ESM) Table S1.

First, to investigate if temperature recording was influenced by characteristics of the hygro button's position, it was analysed using a generalised linear model in relation to button height, distance to slope, substrate underneath the button and canopy presence/absence, specifying a normal error distribution. None of the variables showed a significant effect on mean temperature (p > 0.05), therefore they were not considered further in the analysis.

207 In the next modelling step, standardised temperature estimates were derived separately 208 for open Alpine grassland and forest/ecotone habitat, i.e. models were used to estimate 209 temperature for a given elevation whilst accounting for seasonal variation. Points were 210 classified as Alpine grassland if there was no canopy within 100m radius of the point count 211 centre (following Chamberlain et al. 2013). For open habitat points (n = 93), temperature was 212 modelled in relation to date and elevation. Date was described as the number of days passed 213 since the start of the field season, where day 1 = 27-May-2017. Canopy cover was added to 214 the model structure for points located in forest and ecotone habitat (n = 128). In both cases, a 215 normal distribution was specified. Prior to modelling, all variables were scaled and centred 216 using the scale function in R. Collinearity was assessed using Variance inflation factors (VIFs), 217 calculated using the 'corvif' function (package 'AED', Zuur et al. 2009), and by considering 218 Spearman correlations between continuous variables. All variables had VIF < 3, and no pair of 219 variables showed a correlation > 0.7, indicating low levels of inter-correlation. These models 220 were used to derive a standardised temperature for each point, based on the elevation at that 221 point, the canopy cover (for forest/ecotone habitat) and for a date fixed at 15th June.

222

223 Species distribution models

Birds detected within a 100-m radius of a point count location were used to analyse species distribution (presence/absence of individual species). Bird species were considered in the modelling process if they were present on at least 15 % of the points; below this threshold model performance is consistently poor (Chamberlain et al. 2013).

228 The commonest species were modelled in relation to four different variable sets: (i) habitat 229 (HABITAT), (ii) habitat + temperature (TEMP), (iii) habitat + topography (TOPO), (iv) habitat + 230 temperature + topography (COMB; Table 2). Temperature and topographic variables were 231 used at two different scales (large-scale/ fine-scale). Fine-scale temperature estimates were 232 derived from the temperature modelling approach described above, whereas large-scale 233 temperature data for each point were extracted from the Worldclim database (Hijmans et al. 234 2005) by calculating the average temperature within a 1000 m radius of the point count centre. 235 Topographic variables (aspect and slope) were derived from a Digital Elevation Model (DEM) 236 at a spatial resolution of 10 metres. Aspect was transformed as $x = -1^* \cos[\mathcal{O}(\pi/180)]$, where \mathcal{O} 237 is measured in degrees. Values ranged from 1 where solar insolation was higher (south-facing 238 slopes) to -1 (north-facing slopes) where it was lower.

The mean aspect (transformed values) and slope was calculated within a 100 m (finescale) and a 1000 m (large-scale) radius of the point count centre for the analysis. Habitat variables were kept at a constant scale in the models (as the objective was to test scale effects in temperature and topography).

Habitat models of Lesser Whitethroat *Sylvia curruca* and Dunnock *Prunella modularis* were tested for non-linear relationships with Rhododendron and Juniper cover as suggested by previous work(Jähnig et al. 2018). Habitat models with and without quadratic terms for shrub species cover were compared using AIC. Lesser Whitethroat models showed lower AIC values for the habitat model without quadratic terms. Therefore these were omitted in further modelling steps. The addition of the quadratic term for Rhododendron cover reduced the AIC of the habitat model for Dunnock by $\Delta AIC > 2$, hence it was included in the next modelling steps.

The occurrence probability of each species was modelled in relation to the different variable sets using a binomial logistic regression, after controlling for potential collinearity (as above). In the case of open habitat species, we found high VIFs for the variables rock and grass cover. After the removal of rock cover, all VIFs were below the threshold of three. As a result, rock cover was removed from all models for open habitat species.

255 Data were analysed using an information theoretic approach with the MuMIn package in 256 R version 3.5.2; (R Development Core Team 2018, Bartón 2013). This entailed deriving full 257 models for each variable set at each scale (except habitat which was kept constant in all 258 models) using generalised linear models (R package lme4; Bates et al. 2015). This approach 259 served two goals. First, model-averaged parameter estimates were derived for all 260 combinations of variables in each full model set in order to identify variables that were most 261 closely associated with bird distribution. p-values derived from the model-averaged parameter 262 estimates and their SEs were considered to represent significant effects when p < 0.05. 263 Second, the Akaike information criterion corrected for small sample size (AICc) was 264 determined for each individual model and was used to assess model performance for different 265 variable combinations at different scales in the full model. In this way it was possible to assess 266 which combination of the four different variable sets produced the best models, and at which 267 scale.

At each scale, the residuals for all full models were extracted and tested for spatial autocorrelation using Moran's *I* (Moran 1950). Significant spatial autocorrelation was found for models of Eurasian Skylark *Alauda arvensis*, Tree Pipit *Anthus trivialis* and Water Pipit. For these species, spatial effects were incorporated by modelling their distributions using Generalized Additive Models (GAMs) from the mgcv package (Wood 2011) by fitting smoothed terms for latitude and longitude in the model, following Wood (2017).

274

275 **Results**

In total, 862 individuals of 40 species were recorded in 221 point counts over an elevational
range of 1750 – 2800 m a.s.l. There were seven species that were recorderd on at least 15%

of the points within forest and ecotone habitat: Dunnock, Lesser Whitethroat, Chaffinch *Fringilla coelebs*, Mistle Thrush *Turdus viscivorus*, Coal Tit *Parus ater*, rock bunting *Emberiza cia*, Tree Pipit and three species within open habitat: Eurasian Skylark, Water Pipit and
Northern Wheatear.

The best model to predict rock bunting occurrence was always the null model for each model set at each scale, with no model-averaged parameter estimates being significant. Therefore, this species was not considered further in the analysis.

285

286 Forest and ecotone species

287 Habitat variables such as trees and shrubs were the variables most commonly associated with 288 species occurrence within the HABITAT model for forest and ecotone species. In general, the 289 results of the HABITAT models were in line with previous findings by Jähnig et al. (2018). 290 Juniper showed a positive relationship with Coal Tit, Dunnock and Lesser Whitethroat, but was 291 negatively related to Tree Pipit presence. Rhododendron was positively associated with Mistle 292 Thrush and Lesser Whitethroat presence, whereas it showed a non-linear relationship with 293 Dunnock presence. The number of mature trees showed a positive relationship with forest 294 species (Chaffinch, Mistle Thrush and Coal Tit). Habitat associations among the species 295 remained mostly constant in TEMP, TOPO and COMB models (for full details see ESM Table 296 S2, S4).

297 Each variable set at each scale performed equally well for Lesser Whitethroat, Mistle 298 Thrush and Coal Tit (Table 3). (Note that full details of all models are given in ESM Table S3). 299 Large-scale temperature and topographic variables were included in the best performing 300 model for Dunnock, temperature being negatively associated with Dunnock presence (Table 301 4, Fig. 1). In contrast, large-scale temperature showed a positive relationship with Chaffinch 302 presence in models including only large-scale temperature (Table 4, Fig. 1), or in models 303 including a combination of large-scale temperature and topographic variables. In both species, 304 large-scale model sets performed better than their fine-scale equivalents. Large-scale models 305 for TOPO and COMB were the best performing models for Tree Pipit, whose presence was

306 more closely associated with large-scale topographic variables such as aspect, for which it 307 showed a strong negative relationship indicating a preference for westerly over southerly 308 slopes (Fig. 2). Beside Tree Pipit, only Mistle Thrush showed a negative association with 309 aspect. No other species showed any association with slope or aspect. Furthermore, Tree Pipit 310 was the only species that showed better model performance ($\Delta AICc \leq 2$) for the large-scale 311 TOPO model compared to all fine-scale models and the large-scale TEMP model. All other 312 species showed better (Chaffinch) or equal model performance of TEMP models compared to 313 TOPO models at both scales.

314

315 Open habitat species

The HABITAT model for each open species did not show any habitat associations among the recorded variables. However, all fine-scale models (TEMP, TOPO and COMB) showed a positive association between grass cover and Skylark presence while Juniper cover was only positively associated in the TEMP and COMP models.

320 Models including fine-scale temperature and topography performed best (Δ AlCc \leq 2) for 321 Northern Wheatear. The best performing models of Skylark and Water Pipit included both fine-322 scale TEMP and COMB models. Fine-scale temperature was positively associated with Water 323 Pipit and Northern Wheatear presence, whereas Eurasian Skylark presence was negatively 324 associated (Table 4, Fig. 3).

325 At a fine scale, TEMP models showed better model performance than TOPO models for 326 Northern Wheatear and Water Pipit, whereas on a large scale, model sets for TEMP and TOPO 327 were overlapping (Northern Wheatear, Water Pipit). The large-scale TOPO model showed 328 equal model performance compared to the large-scale TEMP model for Skylark, but AIC_c was 329 still higher compared to fine-scale COMB. In addition, aspect showed a positive relationship 330 with Northern Wheatear (Fig.2, fine-scale COMB model) and Skylark presence (large-scale 331 TOPO model) while slope was positively related to Skylark presence in the fine-scale TOPO 332 model.

333

334 Discussion

335 Models including fine-scale temperature estimates (TEMP, COMB) showed better model 336 performance ($\Delta AIC_c < 2$) than corresponding large-scale models for all three open habitat 337 species. Northern Wheatear and Water Pipit were both positively associated with warm 338 microclimates while Skylark presence was negatively associated with fine-scale temperature. 339 These results contrast with previous findings from the same region of the Alps (Chamberlain 340 et al. 2013, 2016), where model predictions were based on large-scale climatic variables. In 341 these studies, SDMs (based on temperature change and assuming no change in habitat) 342 suggested that under warmer conditions, Skylark and Northern Wheatear would show an 343 increase in their distribution whereas Water Pipit distribution would decrease. Therefore, for 344 Water Pipit and Skylark distributions, our findings suggest opposite associations between fine-345 scale and large-scale temperature.

346 Differences in model predictions at different spatial scales have been reported for a range 347 of studies, and thus identifying the appropriate scale represents a major problem when 348 forecasting suitable habitat in order to inform conservation planning (Elith and Leathwick 2009, 349 Randin et al. 2009, Franklin et al. 2013, Logan et al. 2013, Scridel et al. 2018). To improve 350 SDMs, it is therefore necessary to carefully select predictors (e.g. temperature variables) and 351 their spatial resolution. In the case of microclimate, local topography could create areas with 352 suitable climatic conditions under which it would still be possible for a species to persist under 353 the impact of climate change. Through the use of large-scale climate data, these areas might 354 not be recognised by SDMs (Austin et al. 2011). Besides affecting the future distribution of a 355 species, microclimate can also influence many other aspects of a species' life cycle.

There is evidence that microclimate can be important in influencing habitat selection in mountain birds which may explain our findings. For example, it has been shown that Horned Larks *Eremophila alpestris* adjusted the amount of incubation time in response to microclimatic conditions (Camfield and Martin 2009) by spending less time on the nest as temperatures in the nest surrounding increased, which may imply energy savings in warmer microclimates. Furthermore, microclimate and aspect strongly influenced nestling survival in Water Pipits

362 (Rauter et al. 2002). Nests which were located at ENE-facing slopes (temperature maximum 363 in the morning) had more fledglings than those on WSW-facing slopes (temperature maximum 364 in the afternoon). In contrast, foraging habitat selection by alpine White-winged Snowfinches 365 Montifringilla nivalis, a high altitude specialist, was influenced by solar radiation (Brambilla et 366 al. 2017b). Snowfinches preferred to forage at colder sites (low solar radiation) throughout the 367 season. These studies illustrate that behaviour, foraging habitat selection and choice of nest 368 sites could be driven by microclimatic conditions thereby affecting bird species distributions. 369 Therefore, we would strongly recommend considering microclimate as a predictor in future 370 SDMs for open habitat Alpine species.

In contrast to the open habitat species considered, forest and ecotone species showed no association with fine-scale temperature. One possible reason might be the buffering effect of vegetation. Körner et al. (2007) showed that temperature can vary strongly between forest and open alpine grassland along the elevation gradient with intermediate values at the treeline ecotone. Furthermore, canopies can buffer the diurnal amplitude of air temperature in the forest (Chen et al. 1999).

377 For two species (Dunnock and Chaffinch) large-scale models including temperature (TEMP, 378 COMB) performed better than fine-scale models. The probability of occurrence of Chaffinch 379 was positively associated with large-scale temperature, whereas the probability of Dunnock 380 presence was negatively affected. A future increase in temperature could therefore affect the 381 distribution of Chaffinches by expanding its range towards higher elevations. In contrast, the 382 distribution of Dunnocks might be severely limited. Bani et al. (2019) demonstrated that 383 Dunnock distribution experienced a lower range contraction along the elevational gradient 384 during the last 35 years, but a simple dispersal into higher elevations as a response to 385 environmental change might not be possible because it's preferred nesting habitat in our study 386 area, Rhododendron, has a slow rate of colonisation to the extent that treeline shifts towards 387 higher elevations are likely to be more rapid than upwards shifts in this species (Komac et al. 388 2016).

The mismatch between temperature and available future habitat can also affect open habitat species considered in this study. Due to increasing temperatures, shifts in major habitat types (i.e. forest and shrub encroachment; Harsch et al. 2009) may lead to habitat fragmentation and/or loss of open alpine grassland at higher elevations. This process might even be exacerbated by the abandonment of pastoral activities which formerly have maintained the forest limit at lower elevations than would be possible under climatic constraints only (Gehrig- Fasel et al. 2007).

397 Topography

398 For the majority of species, COMB models performed equally well in comparison with TEMP 399 models at both spatial scales. Combining temperature with topographic variables increased 400 model performance only for Northern Wheatear and Skylark at a fine scale. For the former 401 species, occurrence was more closely related with south-facing slopes. At a large scale, the 402 probability of Tree Pipit presence was higher on westerly slopes. However, in general 403 topographic variables were rarely associated with species occurrence. The influence of aspect 404 on the occurrence of some species could be explained by its effect on snow melt patterns 405 during spring. Thermal differences among slopes with different exposition, which are caused 406 by the amount of received solar radiation, could lead to an early snow melt on south-exposed 407 slopes whereas north-exposed slopes might stay snow covered for a longer period (Keller et 408 al. 2005). These early snow free areas could potentially benefit Northern Wheatears by making 409 suitable nesting sites available earlier. Furthermore, it has been shown that differences in 410 temperature among slopes can influence plant species diversity in temperate mountains 411 (Winkler et al. 2016) with south-exposed slopes favouring a higher degree of species richness 412 and diversity which may in turn influence insect availability.

413

414 Conservation implications

415 Previous studies from the Italian Alps have indicated that increasing temperatures could have
416 detrimental effects for certain Alpine species in the future (Chamberlain et al. 2013), with some

³⁹⁶

species being potentially impacted by both temperature and habitat shifts (Water Pipit), while
for others, loss of habitat due to forest and shrub encroachment will likely be more important
(Northern Wheatear, Skylark).

420 However, our results have shown that species such as Water Pipit and Northern 421 Wheatear are positively associated with warm microclimates which could indicate that both 422 species are potentially more resistant to the impact of a warming climate than previously 423 emphasised by large-scale temperature modelling (e.g. Chamberlain et al. 2013). As a 424 consequence, our results imply that changes in habitat in the form of advancing treelines and 425 the encroachment of formerly open areas by shrubs and trees (Gehrig-Fasel et al. 2007, 426 Leonelli et al. 2011) are currently the major threat to those Alpine species, rather than direct 427 effects of temperature. Therefore, it becomes particularly important to actively manage open 428 areas within mountain environments. This could be achieved by targeted grazing techniques 429 such as mineral mix supplements (Pittarello et al. 2016) or temporary night camp areas (Tocco 430 et al.2013). Both techniques lead to the mechanical damage of shrubs (including saplings) and 431 eventually result in a reduction of shrub cover (Probo et al. 2013, 2014).

432

433

434

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438

439 **Figure legends**

440

Fig. 1 Relationship between large-scale temperature and the probability of occurrence of
Dunnock and Chaffinch based on the large-scale COMB model. Shading indicates the 95%
confidence interval.

444

Fig. 2 Relationship between aspect and the probability of occurrence for Tree Pipit and Northern Wheatear for the large-scale TOPO and the fine-scale COMB model, respectively. Note that aspect was modelled as an index from 1 (south-facing) to -1 (north facing), but here we present the axis as the equivalent cardinal direction for ease of interpretation. Shading indicates the 95% confidence interval.

451 Fig. 3 Relationship between fine-scale temperature and probability of occurrence for open
452 habitat species for the fine-scale COMB model. Shading indicates the 95% confidence interval.

Parameter	Scale	Description					
Canopy	fine	Percentage cover of canopy (above head height) within a radius					
		of 100 m of the point count centre					
Rod fine		Percentage cover of Rhododendron within a radius of 100 m of					
		the point count centre					
Jun	fine	Percentage cover of Juniper within a radius of 100 m of the point					
		count centre					
Vac	fine	Percentage cover of bilberry within a radius of 100 m of the point					
		count centre					
Oth	fine	Percentage cover of shrubs different from Juniper,					
		Rhododendron and bilberry within a radius of 100 m of the point					
		count centre					
Grass	fine	Percentage cover of grass within a radius of 100 m of the point					
		count centre					
Rock	fine	Percentage cover of rock within a radius of 100 m of the point					
		count centre					
Trees	fine	Number of mature (greater than ca. 20 cm in diameter) trees					
		within a radius of 50 m of the point count centre					
Temp	fine	Modelled fine-scale standardised average temperature of the					
		point count centre					
Temp	large	Large-scale average temperature within 1000 m of the point					
		count centre extracted from WorldClim.					
Aspect	fine	The average direction a slope is facing within a 100 m radius of					
		the point count centre transformed as $x = -1*\cos[O(\pi/180)]$,					
		where \emptyset is measured in degrees.					
Aspect	large	The average direction a slope is facing within a 1000 m radius of					
		the point count centre transformed as $x = -1*\cos[O(\pi/180)]$,					
		where \emptyset is measured in degrees.					
Slope	fine	The average inclination of the surface within a 100 m radius of					
		the point counts centre measured in degrees.					
Slope	large	The average inclination of the surface within a 1000 m radius of					
		the point counts centre measured in degrees.					

 Table 1
 Variables considered in the analysis, and the scale at which they were measured.

Table 2 Variable combinations for each model set. Model sets for TEMP, TOPO and COMB were considered at two different scales (fine and large) and included temperature, slope and

Full	Parameter					
models	Falanelei					
HABITAT	Rod + Jun + Vac + Oth + Grass + (Rock) + (Trees)					
TEMP	Rod + Jun + Vac + Oth * Grass + (Rock) + (Trees) + Temp					
ΤΟΡΟ	Rod + Jun + Vac + Oth * Grass + (Rock) + (Trees) + Slope + Aspect					
COMB	Rod + Jun + Vac + Oth * Grass + (Rock) + (Trees) + Temp + Slope +					
	Aspect					

aspect at their matching scale. The variables Rock and Trees were omitted from the habitat model for open habitat species (Northern Wheatear, Water Pipit and Skylark).

Table 3 ΔAIC_c value for each model set at each scale for all species. A ΔAIC_c value of zero indicates the best performing model. Note that in some cases, the best performing models were identical in different model sets, hence a value of zero can appear more than once for a given species. Original AIC_c values are listed in ESM Table S3.

	HABITAT	TEMP		Т	ОРО	COMB		
Species	fine-scale	fine-scale	large-scale	fine-scale	large-scale	fine-scale	large-scale	
Prunella modularis	3.9	2.7	0.8	3.9	3.9	2.7	0	
Sylvia curruca	0	0	0	0	0	0	0	
Parus ater	1.4	1.4	0	1.4	1.4	1.4	0	
Turdus viscivorus	1.2	1.2	1.2	0	1.2	0	1.2	
Fringilla coelebs	13.9	6.3	0	13.1	12.9	6.3	0	
Anthus trivialis	10	8.5	10	9.8	0	8.4	0	
Oenanthe oenanthe	7.7	3.5	7.7	5.8	6.5	0	6.5	
Anthus spinoletta	4.6	0.9	4.6	3.2	4.6	0	4.6	
Alauda arvensis	6.0	0.9	2.7	2.3	4.8	0	2.7	

Species	Mode	Scal	Paramet		SE	Ζ	р
	Ι	е	er	Estimate			
Prunella modularis	COM	large	Rod	1.285	0.618	2.068	
	В						0.038
			Jun	0.818	0.308	2.632	
							0.008
			Oth	0.580	0.270	2.126	
							0.033
			Temp	- 0.886	0.406	2.163	
							0.030
Sylvia curruca*	TEM	fine	Rod	1.029	0.265	3.835	≤
	Р			0.004	0.005	0.000	0.001
			Jun	0.624	0.265	2.332	0.010
			Dook	1 171	0 467	0 407	0.019
			Rock	- 1.174	0.467	2.487	0.012
Parus ater*	COM	large	Jun	0.677	0.333	2.017	0.012
	B	large	Jun	0.077	0.555	2.017	0.043
	D		Oth	0.657	0.277	2.351	0.040
			0	01001	0.277	2.001	0.018
			Grass	0.938	0.434	2.145	
							0.031
			Trees	1.126	0.303	3.678	≤
							0.001
Turdus viscivorus*	TOP	fine	Vac	- 1.700	0.823	2.049	
	0						0.040
			Aspect	- 0.644	0.316	2.017	
							0.043

Fringilla coelebs	TEM	large	Trees	2.453	0.809	3.004	
	Р						0.002
			Temp	1.500	0.462	3.218	
					0	0.2.0	0.001
							0.001
Anthus trivialis	TOP	large	Jun	- 1.187	0.399	2.939	
	0						0.003
			Aspect	- 2.614	0.644	4.022	≤
							0.001
Oenanthe	COM	fine	Temp	0.736	0.305	2.386	
oenanthe	В						0.017
			Aspect	0.622	0.288	2.133	
							0.032
Anthus spinoletta	COM	fine	Temp	1.336	0.577	2.281	
	В						0.022
Alauda arvensis	COM	fine	Jun	0.616	0.282	2.158	
	В						0.030
			Grass	1.010	0.436	2.289	
							0.022
			Temp	-	0.447	1.980	
				0.896			0.047

Table 4 Significant model averaged parameters of the best model for each species. The model type, scale (large or fine), estimate, standard error (SE), test value (*z*) and *p*-value are given for each parameter. Note that species where there was more than one model in the best model set (i.e. $\Delta AIC_c < 2$) are marked with *. In these cases, the model with the lowest AIC_c is presented, but competing models are shown in ESM, Table S3, along with full details for all species.



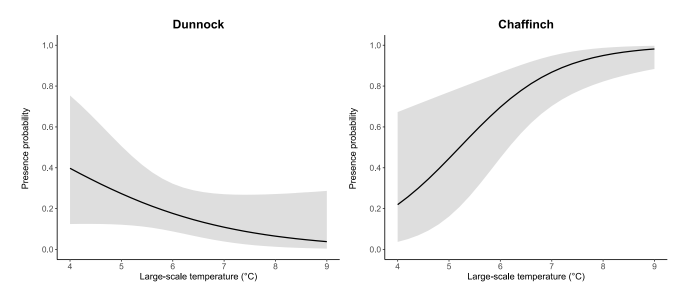


Figure 2

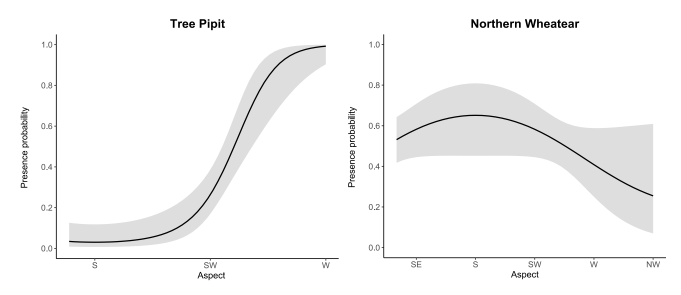


Figure 3

