# **International Journal of Climatology**



# Mire microclimate: groundwater buffers temperature in waterlogged versus dry soils

Journal:	International Journal of Climatology
Manuscript ID	JOC-20-0599.R2
Wiley - Manuscript type:	Research Article
Date Submitted by the Author:	n/a
Complete List of Authors:	Fernández-Pascual, Eduardo; Universidad de Oviedo, Departamento de Biología de Organismos y Sistemas Correia-Álvarez, Eva; Independent researcher, NA
Keywords:	bog, CHELSA, climatic model, datalogger, fen, peatland, soil temperature, wetland
Country Keywords:	Spain

SCHOLARONE™ Manuscripts

## 1 Mire microclimate: groundwater buffers temperature in

## 2 waterlogged versus dry soils

- 3 Short title: Testing the thermal buffer of mire groundwater
- 4 Eduardo Fernández-Pascual<sup>1+</sup>, Eva Correia-Álvarez<sup>2</sup>
- <sup>1</sup> Universidad de Oviedo, <sup>2</sup> Independent researcher
- 6 + Correspondence: Departamento de Biología de Organismos y Sistemas,
- 7 Universidad de Oviedo, C/ Catedrático Rodrigo Uría, 33006 Oviedo/Uviéu, Spain.
- 8 Email: efernandezpascual@gmail.com. Telephone: +34985104787.

#### 9 Financial support

- 10 E.F.P. received financial support from the Government of Asturias and the FP7 -
- 11 Marie Curie COFUND programme of the European Commission (Grant 'Clarín'
- 12 ACB17-19).

#### 13 **Abstract**

- 14 Ecosystems adapt differently to global warming through microclimate factors. Mires
- are sensitive wetland habitats that strongly rely on local soil properties, which makes
- 16 them a good model to understand how local climate parameters counteract the
- 17 effects of climate change. We quantified the temperature buffering effect in
- waterlogged mire soils as compared with adjacent dry soils.
- 19 We buried dataloggers at 5 cm depth in waterlogged and dry points in 8 mires of the
- 20 Cantabrian Mountains (Spain, Southwestern Europe) and recorded soil
- 21 temperatures for c. 5 years. We also compared our local measures with air
- temperatures predicted by the CHELSA model.

- 23 Waterlogged soils had less diurnal thermal amplitude (-2.3 °C), less annual thermal
- 24 amplitude (-5.1 °C), cooler summer maximums (-4.3 °C) and warmer winter
- 25 minimums (+0.8 °C). CHELSA air temperatures only correlated significantly (p <
- 26 0.05) with the winter minimum temperatures (Pearson's r > 0.83), and CHELSA
- 27 predictions were less accurate (higher RMSE) for waterlogged soils, except for the
- 28 summer maximums.
- 29 We conclude that mire soils show a thermal buffer effect that insulates them from
- 30 the surrounding landscape. This effect is stronger at the warm end of the climatic
- 31 spectrum, i.e. during summer and at lower elevations. These results highlight the
- 32 potential refugial character of mires under global warming, and the need to integrate
- 33 microclimate measurements into climate change models.

#### Keywords

34

36

bog, CHELSA, climatic model, datalogger, fen, peatland, soil temperature, wetland

#### Introduction

- 37 Climate change (IPCC 2014) affects global biodiversity, from drylands (Huang et al.
- 38 2016) to forests (Seidl et al. 2017) and oceans (Hoegh-Guldberg et al. 2017). An
- 39 accurate prediction of species responses requires to focus on physiologically
- 40 relevant climatic variables related to critical plant growth periods (Gardner, Maclean,
- 41 and Gaston 2019). Traditional models of species responses to global warming are
- based on macroclimatic data from weather stations. Recently, several authors have
- 43 pointed out the need of complementing these models with *in situ* microclimatic
- 44 measures (Lembrechts, Nijs, and Lenoir 2019; Lembrechts et al. 2019; Maclean,
- 45 Mosedale, and Bennie 2019; Philippov and Yurchenko 2019). As has been shown
- 46 for European forests (Zellweger et al. 2020), ecosystems usually respond to broad
- 47 climatic changes through local processes. Increasingly there are downscaling efforts
- 48 based on atmospheric microclimate networks distributed in complex terrain that

49 focus on topographic effects on air temperature (Scherrer and Körner 2010), but 50 hydrologic factors under edaphic control require more attention (Ashcroft and Gollan 51 2013). 52 Mires are permanent semi-terrestrial peatlands whose soils remain waterlogged but 53 not inundated during most of the year (Wheeler and Proctor 2000). The term mire 54 encompasses peatlands that are classified as either ombrotrophic (rain-fed bogs) or 55 minerotrophic (groundwater-fed fens). These are azonal habitats whose existence 56 depends on local soil properties rather than macroclimatic zonation (Breckle 2002). 57 Mires meet certain criteria that make them especially vulnerable to climate change 58 (Horsák et al. 2018): (a) preponderance of species that evolved under a cold climate; 59 (b) low productivity due to nutrient limitation, making them sensitive to increased 60 nutrient cycling caused by warming (Cornelissen et al. 2007); and (c) scattered 61 distribution pattern, which limits species dispersal and migration (Pearson and 62 Dawson 2005). Therefore, mires are priority habitats for biodiversity conservation, 63 harbouring high numbers of endangered species (Bergamini et al. 2009), and 64 supporting highly adapted floras in spatially reduced areas (Grootjans et al. 2006). 65 Worryingly, mires retain high levels of methane and carbon which can be released 66 due to global warming (Koffi et al. 2020). Habitat distribution models have predicted 67 a loss of mire area as a consequence of ongoing climate change (Essl et al. 2012). 68 It has been known for a relatively long time that groundwater can produce a buffer 69 effect on soil temperature, keeping soils warmer than air during cold periods, and 70 vice versa (Frederick 1974; Ellenberg 1988; Geiger, Aron, and Todhunter 2009). 71 Clara M. Frederick (1974) showed the existence of a buffer effect comparing soil 72 temperatures taken at Cedar Bog (Ohio, United States) with temperatures from a 73 neighbouring agricultural station. Recently, more soil temperature measurements 74 have become available for mires of North America (Raney, Fridley, and Leopold 75 2014), Western Europe (Fernández-Pascual et al. 2015) and Central Europe 76 (Horsák et al. 2018). Their comparison with air temperatures derived from models 77 has shown that mire soils are indeed warmer in winter and cooler in summer, thus

- 78 giving support to the existence of the groundwater thermal buffer (Fernández-
- 79 Pascual et al. 2015; Horsák et al. 2018). Furthermore, the effect has been linked to
- the composition of mire flora and fauna (Horsák et al. 2018; Schenková et al. 2020),
- the growth rings of mire trees (Raney et al. 2016) and the role of mires as glacial
- 82 refugia (Jiménez-Alfaro et al. 2016; Dítě et al. 2017).
- 83 Since root-zone temperature is a major determinant of plant ecophysiology (Körner
- and Paulsen 2004), the groundwater buffer effect is expected to allow mire plants to
- live in a wider range of air temperatures than they could otherwise. Indeed, mires
- 86 have a relatively homogeneous flora across their range despite their broad
- 87 distribution. In Europe, fens are distributed from the Iberian Peninsula to boreal
- 88 Fennoscandia and from low valleys to the alpine belt (Peterka et al. 2017). In the
- 89 arctic, temperate-continental and suboceanic regions of boreal Russia mires make
- 90 up an important part of the landscape, from 20 to 80 % of the surface of different
- 91 regions stretching from Europe to the Pacific (Yurkovskaya 2012). In the United
- 92 States, fens are distributed across the glaciated Midwest and Northeast, as well as
- 93 portions of the Appalachian Mountains and mountainous West (Bedford and Godwin
- 94 2003). However, continued aridification is expected to significantly reduce the overall
- 95 extent of wetlands as it has been shown in the Midwestern United States (Johnson
- 96 and Leopold 1994).
- 97 It is evident that the groundwater buffer effect will play a determinant role in the
- 98 response of mire habitats to climate change (Raney 2014). Both bogs and fens are
- actively peat forming and are dependent on precipitation, whereas fens rely also on
- 100 sources of telluric water from mineral ground. Moreover, they show a mosaic of
- 101 microreliefs and a specific plant cover zonation which makes them especially
- sensitive to small changes in wetness. Climate change affects the quantity, timing
- and spatial distribution of precipitation, leading to changes in surface wetness which
- alter the intensity or organic decomposition by disturbing the conditions for plant
- grow and the depth of air penetration (Bragg 2002). Global warming would also result
- in warmer groundwater delivered to fens, but there is a gap of knowledge about the

110

111

112

113

114

115

116

117

118

119

120

121

122

123

124

125

126

127

128

129

130

131

132

133

134

rate of such changes and their ecological consequences within the mires (Raney 2014).

As is the case for all azonal habitats, locally measured temperatures are essential to understand these processes. Available references recorded temperatures only on waterlogged soils, using model-derived air temperatures for comparison (Fernández-Pascual et al. 2015; Horsák et al. 2018; Schenková et al. 2020); or recorded temperatures in both wet and dry spots but for less than a year, lacking representation throughout the growth cycle of mire vegetation (Raney, Fridley, and Leopold 2014; Raney et al. 2016). This article provides the first measurement of the thermal buffer against surrounding non-mire areas, based on soil temperatures recorded during a period of five years. These measures can be used to support previous evidence about the general effects of soil moisture on thermal buffering (Raney, Fridley, and Leopold 2014; Fernández-Pascual et al. 2015; Horsák et al. 2018; Raney 2014) and to determine the magnitude of this buffering within microrefugia habitats. Specifically, we tested the hypotheses that, when compared to adjacent dry soils, waterlogged mire soils are (i) warmer in winter and (ii) colder in summer; and have less thermal amplitude in (iii) daily and (iv) annual scales. In addition, we compared in situ measurements with data derived from the CHELSA climatic models (Karger et al. 2017).

#### **Materials and methods**

This study took place in the temperate oceanic region of north-western Spain (43° N, 5° W) (Fig. 1a). Local geography and climate are dominated by the Cantabrian Mountains (> 1500 m above sea level), which run parallel to the coast and trap the prevailing NW Atlantic winds. In the study area, average annual precipitation ranges from 800 mm at the low elevations to 1800 mm at the high mountains, while average annual temperature ranges from 12.5 °C to 5 °C (source = Agencia Estatal de Meteorología, http://www.aemet.es/es/serviciosclimaticos/). Under the most extreme emissions scenario, the latest projections expect temperatures to increase

by 4 °C and precipitation to decrease by c. 10 % by the end of the century, although the precipitation projections are subjected to great uncertainty (Amblar-Francés et al. 2017). The present humid and mild climate of this region harbours the southwestern limit of mire communities in Europe (Jiménez-Alfaro, Díaz González, and Fernández-Pascual 2011; Fernández Prieto, Fernández Ordóñez, and Collado Prieto 1985). Rain-fed raised bogs (https://eunis.eea.europa.eu/habitats/260) are very rare and appear locally on raised hummocks of acid valley mires (https://eunis.eea.europa.eu/habitats/526) which can be found from the coast to just below the treeline, in poorly drained valleys and former glacial lakes. Glacial lakes undergoing silting develop transition mires and quaking bogs communities (https://eunis.eea.europa.eu/habitats/1637) in the water-to-land transition. Spring fens appear in the mountains above 1000 m; they range from soft-water poor fens (https://eunis.eea.europa.eu/habitats/279) on acid bedrocks, to alkaline calcareous fens (https://eunis.eea.europa.eu/habitats/277) on limestone.

We selected 8 mire sites representing the regional elevation gradient of mire vegetation and the different mire types (Table 1). Although we classify two of these sites as rain-fed bogs based on their vegetation (*Oxycocco-Sphagnetea* Br.-Bl. et Tx. ex Westhoff et al. 1946), we must note that these bogs are very poorly developed on hummocks over valley mires and are in tight connection to the underlying water table coming from streams. Therefore, soil waterlogging in all our sites is fed by streams or springs. In each site, we buried two dataloggers (M-Log5W, GeoPrecision, Ettlingen, Germany; accuracy: +/- 0.1 °C (at 0 °C), resolution: 0.01 °C) at a depth of 5 cm below the upper layer of the soil. In this ecosystem, it is at this depth that true soil begins to develop, under the porous upper layers made up of live mosses. We installed one datalogger in a flat waterlogged spot within the mire; the other one in the close vicinity, but in a flat and dry upland area outside the mire (Fig. 1b). The vegetation at the sampling points was always either mire or pasture, with no shrubs, trees or any other landscape features shading the measuring points. Dataloggers recorded temperature once every hour and stayed on site for

165

166

167

168

169

170

171

172

173

174

175

176

177

178

179

180

181

182

183

184

185

186

187

188

189

190

191

192

approximately five years, after which we retrieved them and downloaded their records. At the moment of retrieval, the internal clock of all dataloggers had not deviated for more than four hours.

All data processing and analysis was performed in R (R Core Team 2019). To clean the logs we took the following steps: (i) removing records from the first week after installation, to account for the installation process and the settling of the soils; and (ii) keeping only time series with records for both the dry and waterlogged points, because some of the dataloggers had failed at different points in time. Afterwards, we calculated four bioclimatic variables for each datalogger: (1) the mean diurnal range; i.e. the average for the whole period of the daily differences between the maximum and the minimum temperatures recorded in the day; (2) the maximum temperature of the warmest month; i.e. the average of the daily maximum records, for the warmest month; (3) the minimum temperature of the coldest month; i.e. the average of the daily minimum records, for the coldest month; and (4) the annual range; i.e. the difference between the maximum temperature of the warmest month and the minimum temperature of the coldest month. These are classical bioclimatic variables that indicate the limiting factors (extreme temperatures) and seasonality (diurnal and annual range) that organisms must tolerate (O'Donnell and Ignizio 2012). To compare our measurements with model-based predictions of air temperature, we downloaded from CHELSA the same bioclimatic variables for our measuring coordinates (https://chelsa-climate.org/bioclim/).

To test if the differences between dry and waterlogged points were significant, we used paired t-tests (n = 8 paired sites per each of the four bioclimatic variables). Tests were one-tailed, according to the original hypotheses: the dry point would have a higher diurnal range, a higher maximum temperature, a lower minimum temperature, and a higher annual range. To compare our soil measurements with the CHELSA air temperatures we calculated Pearson's correlation between both sets of values, and we also calculated the root-mean-square error (RMSE) of the CHELSA values as a measure of their accuracy (n = 8 sites per combination of

bioclimatic variable and dry/waterlogged condition). We calculated RMSE using its implementation in the package *Metrics* (Hamner and Frasco 2018). We checked whether our data complied with the normality assumption of the t-test and Pearson's correlation using the Shapiro-Wilk normality test; the assumption was met in all cases (p > 0.05). To check whether the analyses were influenced by the fact that recording periods did not match across all eight sites, we repeated the analyses using only full-year, contemporaneous data series (01 Jan 2015 - 31 Dec 2017, excluding La Vega Comeya); the results of these analyses were the same as the ones presented in this article (Supplementary Tables S1 and S2).

#### Results

193

194

195

196

197

198

199

200

201

202

- 203 Dataloggers recorded temperatures for five years in five of the sites, four years in
- 204 two, and two years in one. Time series showed considerably less variation at
- waterlogged points, for all eight site comparisons (Fig. 2).
- 206 Bioclimatic variables (Table 2 and Fig. 3) supported this notion. The mean annual
- 207 range was wider at dry points, as was the mean diurnal range. Maximum
- 208 temperatures of the warmest months were usually higher at dry measuring points.
- 209 The opposite was true for the minimums of the coldest months, in which case
- 210 temperature was generally colder at dry points, although the difference was less
- 211 pronounced than for the maximums (Fig. 3).
- 212 Patterns regarding the maximum temperatures and annual range were especially
- 213 noticeable at both low (El Molinucu, La Malva) and high (La Recoleta) elevations.
- 214 The pattern for the diurnal range, however, was more prominent just at the low sites
- 215 (El Molinucu, La Malva), whereas the minimums of the coldest months showed no
- 216 specific pattern (Table 2).
- 217 T-tests supported the original hypotheses; namely waterlogged measuring points
- 218 had (a) smaller diurnal fluctuations (t = -3.05, p = 0.009, effect size = -2.29 °C, 95 %
- 219 CI =  $-\infty$ , -0.87); (b) lower maximums (t = -3.04, p = 0.009, effect size = -4.28 °C, 95

- % CI =  $-\infty$ , -1.61); (c) higher minimums (t = 2.86, p = 0.012, effect size = 0.77 °C, 95
- % CI = 0.26,  $\infty$ ), and (d) smaller annual fluctuations (t = -3.95, p = 0.003, effect size
- 222 = -5.05 °C, 95 % CI = -∞, -2.63).
- 223 The CHELSA climatic model (Table 3 and Fig. 4) provided air temperatures that
- 224 significantly correlated (p < 0.05) with soil temperatures only in the case of the
- 225 minimums of the coldest month, both at dry points (Pearson's r = 0.90) and
- 226 waterlogged points (Pearson's r = 0.84). The correlation between CHELSA and soil
- values was marginally significant (p = 0.05) in the case of the maximums of the
- warmest month, but only at dry points (Pearson's r = 0.71). The values of CHELSA
- 229 did not correlate with soil temperatures in the rest of the cases (p > 0.05). RMSE
- 230 indicated that the predictions of CHELSA were more accurate at dry points for the
- cases of the annual range, the diurnal range and the winter min; the CHELSA values
- were more accurate at waterlogged points in the case of the summer max.

#### Discussion

233

247

234 The results presented here quantify the size of the thermal buffer effect that takes 235 place in waterlogged mire soils (Frederick 1974), when compared with adjacent dry 236 soils. The mire thermal buffer had been compared previously with air temperatures 237 derived from models, with generally similar results (Fernández-Pascual et al. 2015; 238 Horsák et al. 2018). The mire buffer had also been compared to dry soils at 10 cm 239 depth (Raney, Fridley, and Leopold 2014; Raney et al. 2016); our results confirm 240 those findings at 5 cm depth and extend them to the full year. Also, in concordance 241 with our results, the air temperature measured at 0.5 m from the surface of boreal 242 Russian bogs is lower at the wetter zones, at least during the warmest months 243 (Philippov and Yurchenko 2019). Thus, the pattern is reproducible among years (out 244 to five years), seems to take place both at southern and northern latitudes, and may 245 affect not only the temperature in the soil but also air temperatures within a certain 246 distance from the soil. The thermal buffer effect of mire soils that makes their

temperatures less extreme than the surrounding landscape is therefore a general

phenomenon. The existence of this buffer effect highlights the importance of using fine-scale microclimatic data to assess vegetation responses to climate change (Storlie et al. 2014; Lembrechts, Nijs, and Lenoir 2019; Zellweger et al. 2020).

One important difference with previous studies is the magnitude of the buffer during winter. The articles that had used model air temperatures as a control concluded that the buffer effect was stronger at the cold end of the thermal spectrum, i.e. in winter and at night (Fernández-Pascual et al. 2015; Horsák et al. 2018). In the case of this investigation, the situation is the reverse: the effect is weaker when considering the minimum temperatures of the cold period. This indicates the importance of identifying root temperatures when working with plant communities. At high elevations of the study region, the soil can remain covered by snow for periods of winter, and this has its own insulating effect on soil temperatures (Körner 2003). Indeed, snow cover has being described as one of the vertical features that affects vegetation distribution in a local manner (Maclean, Mosedale, and Bennie 2019). In any case, since this study was conducted at the southern limit of mire distribution, more research on the winter buffer is needed at high latitudes, since it might have a deeper biological significance there.

In these southern mires, the buffering effect is much stronger during the summer (Fig. 5). This is most noticeable in the two lowest elevations, El Molinucu and La Malva. La Malva shows especially high summer temperatures within the dry soils (> 40 °C). It must be noted that this difference of > 15 °C between the dry and waterlogged points of La Malva, which are separated by a few meters, is almost four times the warming expected in the study region at the end of this century (+ 4 °C) under the more extreme emissions scenario (Amblar-Francés et al. 2017). Whereas the rest of the study sites are flat, La Malva is a calcareous spring in a southeast-facing slope on limestone, a place experiencing sub-Mediterranean conditions at the micro-scale (Sánchez de Dios, Benito-Garzón, and Sainz-Ollero 2009), and surrounded by a forest of evergreen oaks (*Quercus rotundifolia* Lam., *Quercus faginea* Lam.). This suggests the importance of groundwater in the existence of mire

278

279

280

281

282

283

284

285

286

287

288

289

290

291

292

293

294

295

296

297

298

299

300

301

302

303

304

305

vegetation in Mediterranean areas (De Hoyos et al. 1996), not only from the water-availability aspect, but also providing cooling regulation during summer (Ellenberg 1988). The buffering is also prominent at one of the highest elevations, La Recoleta, which may be explained by the high saturation of water in this kind of habitat, a quaking bog, i.e. a former glacial lake undergoing silting. However, the relatively small number of sites in our study does not allow to explore further the moderating factors that may drive differences in the strength of the buffer. More studies focusing on each kind of habitat are needed to understand the relation between hydrology conditions and microclimatic effects.

Most models currently employed to predict vegetation and species distribution use macroclimatic parameters, like the ones obtained from CHELSA (Karger et al. 2017). This study revealed that CHELSA values correlate relatively well with soil temperatures in the case of the minimum temperatures of the coldest month, but not with the rest of the bioclimatic variables under examination. Moreover, CHELSA predictions are less accurate for waterlogged soils, except for the summer max, which might be the consequence of the noise introduced by sun-heated soil. In addition to the water buffering effect, other features may explain the differences between our results and CHELSEA values, like the height of temperature measurement (5 cm below the upper layer of the soil in our study versus 2 m above the ground in CHELSA); the temporal scale of observation (hourly based for a 5 years period in our study versus a 34 years period); and the spatial resolution (c. 1 m in our study versus c. 1 km). Be that as it may, our results bring out the importance of considering local factors as drivers of microclimatic changes (Zellweger et al. 2019). These factors contribute to landscape heterogeneity, producing safe sites that can act as micro-refuges and buffer species from regional climatic warming.

In summary, this article shows that waterlogged mire soils have a thermal buffer when compared to adjacent soils, contributing to their behaviour as mild island habitats in a landscape that can be more thermally variable (Raney 2014; Horsák et al. 2018). The effect occurs during cold and warm periods, but it is stronger during

the summer, at least in the study area. However, this climatic buffering within microrefugia relies strongly on the water regime, a key factor for its conservation. Long-term decreases in precipitation could reduce the groundwater discharge into the mires, with potentially fatal consequences if the mires dry out during warmer summers. Rain-fed bogs will be affected more immediately than spring fens since the latter are somewhat decoupled from short-term rainfall variation. However, longterm regional drying would affect all types of mires. For the study area, the latest projections envision a decrease of c. 10 % in annual precipitation at the end of the century, but these projections are still subjected to a high level of uncertainty (Amblar-Francés et al. 2017). Future recording schemes are needed to monitor the evolution of the soil thermal buffer in drying regions, to obtain local soil temperatures from other latitudes, and from more microtopographies within the same mire. We must also consider that the microclimatic effect is not limited to mires as it can also be found in other habitats where a gradient of moisture exists (Ashcroft and Gollan 2013), such as the mosaic of microtopographies (snow patches, fellfields) that is characteristic of alpine environments (Scherrer and Körner 2010; Ohler, Lechleitner, and Junker 2020). This study provides useful microclimate parameters to improve the current models that predict the impact of global warming on moisture-driven ecosystems.

#### Data availability

306

307

308

309

310

311

312

313

314

315

316

317

318

319

320

321

322

323

324

325

330

- The original data, as well as the R code for the analysis and creation of the manuscript, can be accessed at the GitHub repository https://github.com/efernandezpascual/mires. Upon publication, a version of record
- 329 of the repository will be deposited in Zenodo.

#### Supporting information

- 331 Tables S1-S2 Results of the statistical analyses, repeated using only the
- 332 contemporaneous full-year data series.

#### References

- 334 Amblar-Francés, P, MJ Casado-Calle, MA Pastor-Saavedra, P Ramos-Calzado, and
- E Rodríguez-Camino. 2017. Guía de escenarios regionalizados de cambio
- 336 climático sobre España a partir de los resultados del IPCC-AR5. Madrid:
- 337 Ministerio de Agricultura y Pesca, Alimentación y Medio Ambiente. Agencia
- 338 Estatal de Meteorología.
- 339 Ashcroft, Michael B., and John R. Gollan. 2013. "Moisture, thermal inertia, and the
- spatial distributions of near-surface soil and air temperatures: understanding
- factors that promote microrefugia." Agricultural and Forest Meteorology 176: 77–
- 342 89. https://doi.org/https://doi.org/10.1016/j.agrformet.2013.03.008.
- 343 Bedford, Barbara L., and Kevin S. Godwin. 2003. "Fens of the United States:
- distribution, characteristics, and scientific connection versus legal isolation."
- 345 *Wetlands* 23 (3): 608–29. https://doi.org/10.1672/0277-
- 346 5212(2003)023[0608:FOTUSD]2.0.CO;2.
- 347 Bergamini, A., M. Peintinger, S. Fakheran, H. Moradi, B. Schmid, and J. Joshi. 2009.
- 348 "Loss of habitat specialists despite conservation management in fen remnants
- 349 1995-2006." Perspectives in Plant Ecology, Evolution and Systematics 11 (1): 65–
- 350 79. https://doi.org/10.1016/j.ppees.2008.10.001.
- 351 Bragg, O. M. 2002. "Hydrology of peat-forming wetlands in Scotland." Science of the
- 352 Total Environment 294 (1-3): 111–29. https://doi.org/10.1016/s0048-
- 353 9697(02)00059-1.
- 354 Breckle, Siegmar-Walter. 2002. Walter's Vegetation of the Earth: the Ecological
- 355 Systems of the Geo-biosphere. Berlin Heidelberg New York: Springer.
- 356 Cornelissen, Johannes H. C., Peter M. Van Bodegom, Rien Aerts, Terry V.
- Callaghan, Richard S. P. Van Logtestijn, Juha Alatalo, F. Stuart Chapin, et al.
- 358 2007. "Global negative vegetation feedback to climate warming responses of leaf

- 359 litter decomposition rates in cold biomes." *Ecology Letters* 10 (7): 619–27.
- 360 https://doi.org/10.1111/j.1461-0248.2007.01051.x.
- 361 De Hoyos, D, Jesús Muñoz Fuente, A Negro, Juan José Aldasoro, JC Vega, and
- Gonzalo Moreno Moral. 1996. "A survey on Cantabrian mires (Spain)." *Anales Del*
- 363 *Jardín Botánico de Madrid* 54 (1): 472–89.
- 364 Dítě, Daniel, Tomáš Peterka, Zuzana Dítětová, Petra Hájková, and Michal Hájek.
- 365 2017. "Arcto-Alpine species at their niche margin: the Western Carpathian refugia
- of Juneus castaneus and J. triglumis in Slovakia." Annales Botanici Fennici 54 (1–
- 367 3): 67–82, 16. https://doi.org/10.5735/085.054.0311.
- 368 Ellenberg, Heinz. 1988. "Spring areas and adjacent swamps." In Vegetation Ecology
- of Central Europe, 313–13. Cambridge: Cambridge University Press.
- 370 Essl, Franz, Stefan Dullinger, Dietmar Moser, Wolfgang Rabitsch, and Ingrid
- 371 Kleinbauer. 2012. "Vulnerability of mires under climate change: implications for
- 372 nature conservation and climate change adaptation." Biodiversity and
- 373 *Conservation* 21: 655–69.
- 374 Fernández-Pascual, Eduardo, Borja Jiménez-Alfaro, Michal Hájek, Tomás E. Díaz,
- and Hugh W. Pritchard. 2015. "Soil thermal buffer and regeneration niche may
- favour calcareous fen resilience to climate change." Folia Geobotanica 50 (4):
- 377 293–301. https://doi.org/10.1007/s12224-015-9223-y.
- 378 Fernández Prieto, José Antonio, María del Carmen Fernández Ordóñez, and Miguel
- Angel Collado Prieto. 1985. "Datos sobre la vegetación de las turberas de
- esfagnos galaico-asturianas y orocantábricas." *Lazaroa* 7: 443–71.
- 381 Frederick, Clara May. 1974. "A natural history study of the vascular flora of Cedar
- Bog, Champaign County, Ohio." The Ohio Journal of Science 74 (2): 65–116.
- 383 Gardner, Alexandra S., Ilya M. D. Maclean, and Kevin J. Gaston. 2019. "Climatic
- predictors of species distributions neglect biophysiologically meaningful

- 385 variables." *Diversity and Distributions* 25 (8): 1318–33.
- 386 https://doi.org/10.1111/ddi.12939.
- 387 Geiger, Rudolf, Robert H. Aron, and Paul Todhunter. 2009. The Climate Near the
- 388 *Ground*. Lanham: Rowman & Littlefield.
- 389 Grootjans, A. P., E. B. Adema, W. Bleuten, H. Joosten, M. Madaras, and M.
- Janáková. 2006. "Hydrological landscape settings of base-rich fen mires and fen
- 391 meadows: an overview." Applied Vegetation Science 9: 175–84.
- http://search.ebscohost.com/login.aspx?direct=true&db=aph&AN=25691039&la
- 393 ng=es&site=ehost-live.
- Hamner, B, and M Frasco. 2018. "Metrics: Evaluation Metrics for Machine Learning."
- 395 R package version 0.1.4. https://CRAN.R-project.org/package=Metrics."
- 396 Hoegh-Guldberg, Ove, Elvira S. Poloczanska, William Skirving, and Sophie Dove.
- 397 2017. "Coral reef ecosystems under climate change and ocean acidification."
- 398 Frontiers in Marine Science 4 (158). https://doi.org/10.3389/fmars.2017.00158.
- 399 Horsák, Michal, Vendula Polášková, Marie Zhai, Jindřiška Bojková, Vít Syrovátka,
- 400 Vanda Šorfová, Jana Schenková, Marek Polášek, Tomáš Peterka, and Michal
- 401 Hájek. 2018. "Spring-fen habitat islands in a warming climate: partitioning the
- 402 effects of mesoclimate air and water temperature on aquatic and terrestrial biota."
- 403 Science of the Total Environment 634: 355–65.
- 404 https://doi.org/https://doi.org/10.1016/j.scitotenv.2018.03.319.
- 405 Huang, Jianping, Haipeng Yu, Xiaodan Guan, Guoyin Wang, and Ruixia Guo. 2016.
- 406 "Accelerated dryland expansion under climate change." *Nature Climate Change*
- 407 6 (2): 166–71. https://doi.org/10.1038/nclimate2837.
- 408 IPCC. 2014. Climate Change 2014: Synthesis Report. Contribution of Working
- 409 Groups I, II and III to the Fifth Assessment Report of the Intergovernmental Panel
- on Climate Change [Core Writing Team, R.K. Pachauri and L.A. Meyer (eds.)].
- 411 Geneva: IPCC.

- 412 Jiménez-Alfaro, Borja, Tomás Díaz González, and Eduardo Fernández-Pascual.
- 413 2011. "Grupos de vegetación y hábitats de tremedales neutro-basófilos en las
- 414 montañas pirenaico-cantábricas." *Acta Botanica Barcinonensia* 53: 47–60.
- 415 Jiménez-Alfaro, Borja, Laura García-Calvo, Pedro García, and José Luis Acebes.
- 416 2016. "Anticipating extinctions of glacial relict populations in mountain refugia."
- 417 Biological Conservation 201: 243–51.
- 418 Johnson, Anne M., and Donald J. Leopold. 1994. "Vascular plant species richness
- and rarity across a minerotrophic gradient in wetlands of St. Lawrence County,
- 420 New York, USA." Biodiversity and Conservation 3 (7): 606–27.
- 421 https://doi.org/10.1007/BF00114204.
- 422 Karger, Dirk Nikolaus, Olaf Conrad, Jürgen Böhner, Tobias Kawohl, Holger Kreft,
- 423 Rodrigo Wilber Soria-Auza, Niklaus E. Zimmermann, H. Peter Linder, and Michael
- 424 Kessler. 2017. "Climatologies at high resolution for the earth's land surface
- 425 areas." Scientific Data 4 (1): 170122. https://doi.org/10.1038/sdata.2017.122.
- 426 Koffi, Ernest N., Peter Bergamaschi, Romain Alkama, and Alessandro Cescatti.
- 427 2020. "An observation-constrained assessment of the climate sensitivity and
- future trajectories of wetland methane emissions." Science Advances 6 (15):
- 429 eaay4444. https://doi.org/10.1126/sciadv.aay4444.
- 430 Körner, Christian. 2003. Alpine Plant Life: Functional Plant Ecology of High Mountain
- 431 *Ecosystems*. 2nd ed. Berlin Heidelberg New York: Springer.
- 432 Körner, Christian, and Jens Paulsen. 2004. "A world-wide study of high altitude
- 433 treeline temperatures." Journal of Biogeography 31: 713–32.
- 434 http://10.0.4.87/j.1365-2699.2003.01043.x
- http://search.ebscohost.com/login.aspx?direct=true&db=aph&AN=12823821&la
- 436 ng=es&site=ehost-live.
- 437 Lembrechts, Jonas J., Jonathan Lenoir, Nina Roth, Tarek Hattab, Ann Milbau, Sylvia
- Haider, Loïc Pellissier, et al. 2019. "Comparing temperature data sources for use

- in species distribution models: from in-situ logging to remote sensing." Global
- 440 Ecology and Biogeography 28 (11): 1578–96. https://doi.org/10.1111/geb.12974.
- 441 Lembrechts, Jonas J., Ivan Nijs, and Jonathan Lenoir. 2019. "Incorporating
- 442 microclimate into species distribution models." *Ecography* 42 (7): 1267–79.
- 443 https://doi.org/10.1111/ecog.03947.
- 444 Maclean, Ilya M. D., Jonathan R. Mosedale, and Jonathan J. Bennie. 2019.
- 445 "Microclima: an R package for modelling meso- and microclimate." Methods in
- 446 Ecology and Evolution 10 (2): 280–90. https://doi.org/10.1111/2041-210x.13093.
- 447 O'Donnell, Michael S, and Drew A Ignizio. 2012. "Bioclimatic predictors for
- supporting ecological applications in the conterminous United States." US
- 449 Geological Survey Data Series 691 (10).
- 450 Ohler, Lisa-Maria, Martin Lechleitner, and Robert R. Junker. 2020. "Microclimatic
- 451 effects on alpine plant communities and flower-visitor interactions." Scientific
- 452 Reports 10 (1): 1366. https://doi.org/10.1038/s41598-020-58388-7.
- 453 Pearson, Richard G., and Terence P. Dawson. 2005. "Long-distance plant dispersal
- and habitat fragmentation: identifying conservation targets for spatial landscape
- 455 planning under climate change." Biological Conservation 123 (3): 389–401.
- 456 https://doi.org/https://doi.org/10.1016/j.biocon.2004.12.006.
- 457 Peterka, Tomáš, Michal Hájek, Martin Jiroušek, Borja Jiménez-Alfaro, Liene Aunina,
- 458 Ariel Bergamini, Daniel Dítě, et al. 2017. "Formalized classification of European
- fen vegetation at the alliance level." *Applied Vegetation Science* 20 (1): 124–42.
- 460 https://doi.org/10.1111/avsc.12271.
- 461 Philippov, Dmitriy A., and Victoria V. Yurchenko. 2019. "Data on air temperature,
- relative humidity and dew point in a boreal *Sphagnum* bog and an upland site
- 463 (Shichengskoe mire system, North-Western Russia)." Data in Brief 25: 104156.
- 464 https://doi.org/https://doi.org/10.1016/j.dib.2019.104156.

- 465 Raney, Patrick A. 2014. *Identifying potential refugia from climate change in wetlands*.
- Syracuse, New York: Doctoral thesis, State University of New York.
- 467 Raney, Patrick A, Jason D Fridley, and Donald J Leopold. 2014. "Characterizing
- 468 microclimate and plant community variation in wetlands." *Wetlands* 34 (1): 43–53.
- 469 https://doi.org/10.1007/s13157-013-0481-2.
- 470 Raney, Patrick A, Donald J Leopold, Martin Dovčiak, and Colin M Beier. 2016.
- 471 "Hydrologic position mediates sensitivity of tree growth to climate: groundwater
- subsidies provide a thermal buffer effect in wetlands." Forest Ecology and
- 473 *Management* 379: 70–80.
- 474 R Core Team. 2019. "R: a language and environment for statistical computing."
- 475 https://www.r-project.org/.
- 476 Sánchez de Dios, Rut, Marta Benito-Garzón, and Helios Sainz-Ollero. 2009.
- 477 "Present and future extension of the Iberian submediterranean territories as
- determined from the distribution of marcescent oaks." *Plant Ecology* 204 (2): 189–
- 479 205. https://doi.org/10.1007/s11258-009-9584-5.
- 480 Schenková, Jana, Vendula Polášková, Martina Bílková, Jindřiška Bojková, Vít
- 481 Syrovátka, Marek Polášek, and Michal Horsák. 2020. "Climatically induced
- temperature instability of groundwater-dependent habitats will suppress cold-
- 483 adapted Clitellata species." *International Review of Hydrobiology* 105: 85–93.
- 484 https://doi.org/10.1002/iroh.201902006.
- 485 Scherrer, Daniel, and Christian Körner. 2010. "Infra-red thermometry of alpine
- 486 landscapes challenges climatic warming projections." Global Change Biology 16
- 487 (9): 2602–13. https://doi.org/10.1111/j.1365-2486.2009.02122.x.
- 488 Seidl, Rupert, Dominik Thom, Markus Kautz, Dario Martin-Benito, Mikko Peltoniemi,
- 489 Giorgio Vacchiano, Jan Wild, et al. 2017. "Forest disturbances under climate
- 490 change." *Nature Climate Change* 7 (6): 395–402.
- 491 https://doi.org/10.1038/nclimate3303.

510

492 Storlie, Collin, Andres Merino-Viteri, Ben Phillips, Jeremy VanDerWal, Justin 493 Welbergen, and Stephen Williams. 2014. "Stepping inside the niche: microclimate 494 data are critical for accurate assessment of species' vulnerability to climate 495 change." Biology Letters 10 (9): 20140576. 496 https://doi.org/doi:10.1098/rsbl.2014.0576. 497 Wheeler, B. D., and M. C. F. Proctor. 2000. "Ecological gradients, subdivisions and 498 terminology of north-west European mires." Journal of Ecology 88 (2): 187–203. 499 https://doi.org/10.1046/j.1365-2745.2000.00455.x. 500 Yurkovskaya, Tatiana. 2012. "Mires on the map of Russia." In Mires from Pole to 501 Pole, edited by Tapio Lindholm and Raimo Heikkilä, 31–38. Helsinki: Finnish 502 Environment Institute. 503 Zellweger, Florian, Pieter De Frenne, Jonathan Lenoir, Duccio Rocchini, and David 504 Coomes. 2019. "Advances in microclimate ecology arising from remote sensing." 505 Trends in Ecology & Evolution 34 (4): 327-41. 506 https://doi.org/https://doi.org/10.1016/j.tree.2018.12.012. 507 Zellweger, Florian, Pieter De Frenne, Jonathan Lenoir, Pieter Vangansbeke, Kris 508 Verheyen, Markus Bernhardt-Römermann, Lander Baeten, et al. 2020. "Forest

microclimate dynamics drive plant responses to warming." Science 368 (6492):

772–75. https://doi.org/10.1126/science.aba6880.

511 **Tables** 

513

514

515

516

Table 1: Mire sites included in this study, indicating the type of fen, pH, conductivity, elevation, coordinates, and length of the temperature recording period. Groundwater pH and conductivity values are the average of 5–10 measures per site taken with a MM40 Portable Multimeter (Crison Instruments S.A., Alella, Spain). Coordinates are in decimal degrees WGS84.

Site	Habitat	рН	Conductivity (µS/cm)	Elevation (m)	Latitude	Longitude	Records (days)
El Molinucu	Raised bog	4.2	47.2	284	43.3924	-5.5392	1421
La Malva	Alkaline fen	8.2	427.7	700	43.1176	-6.2543	1347
La Vega Comeya	Raised bog	5.9	102.4	822	43.2856	-4.9885	664
La Bruxa	Alkaline fen	8.0	271.8	1528	43.0232	-6.2113	1850
La Veiga Cimera	Acid fen	4.6	13.9	1552	43.0252	-6.2539	1850
La Recoleta	Quaking bog	7.2	245.9	1768	43.0167	-6.1112	1854
El Riotuertu	Alkaline fen	7.3	400.1	1820	43.0096	-5.9479	1852
La Vega Lliordes	Alkaline fen	7.3	293.3	1878	43.1504	-4.8464	1809

Table 2: Groundwater buffer effect per mire and bioclimatic variable. The buffer effect was calculated as the difference between the value in the waterlogged and the dry points.

Site	Annual	Diurnal	Summer	Winter min	
	range	range			
El Molinucu	-7.80	-4.90	-6.59	1.21	
La Malva	-10.81	-5.14	-10.25	0.56	

Site	Annual	Diurnal	Summer	Winter min
Oile	range	range	max	vviiitei iiiiii
La Vega Comeya	-3.63	-2.99	-3.51	0.12
La Bruxa	-1.09	-0.88	0.99	2.08
La Veiga Cimera	-3.41	-2.21	-3.47	-0.06
La Recoleta	-8.81	-2.62	-8.85	-0.04
El Riotuertu	-3.62	1.12	-2.28	1.34
La Vega Lliordes	-1.26	-0.71	-0.28	0.98

Table 3: Comparison of soil bioclimatic variables with CHELSA air temperatures, per bioclimatic variable and groundwater situation. RMSE is the root-mean-square error, a measure of the accuracy of the predictions (lower values indicate higher accuracy).

Variable	Groundwate r	t	df	р	Pearson's r	RMSE (°C)
Annual range	Dry	0.012	6	0.991	0.00	3.62
Annual range	Waterlogge d	-0.334	6	0.750	-0.14	4.71
Diurnal range	Dry	0.016	6	0.988	0.01	3.09
Diurnal range	Waterlogge d	1.354	6	0.225	0.48	4.62
Summer max	Dry	2.450	6	0.050	0.71	5.86
Summer max	Waterlogge d	1.527	6	0.178	0.53	2.95

Variable	Groundwate r	t	df	р	Pearson's r	RMSE (°C)
Winter min	Dry	5.115	6	0.002	0.90	4.04
Winter min	Waterlogge d	3.809	6	0.009	0.84	4.82

#### **Figures**

Figure 1: (a) Location of the study sites in northern Spain. (b) Representation of the sampling design at El Riotuertu site. The position of the dry and waterlogged temperature loggers is indicated, the blue area represents the extent of the waterlogged mire habitat.

Figure 2: Hourly soil temperature records at the mire sites. The blue series was recorded within the mire, in a waterlogged area. The red series was recorded in a neighbouring dry area. Dataloggers were buried at 5 cm depth. The amplitude of diurnal temperature range in dry sites was reduced with increasing elevation, where the thermal buffer effect was also less apparent.

Figure 3: Average bioclimatic variables in the dry and waterlogged points. The bars represent the mean value, and the brackets the standard error of 8 records.

Figure 4: Scatter plots of the bioclimatic variables predicted by CHELSA air temperatures vs. soil temperatures measured in situ, in dry and waterlogged points.

542

543

544

545

546

547

548

549

Figure 5: Complementary temporal analysis of the groundwater buffer effect during peak hot events. Variation of soil temperatures at paired sites (dry vs. waterlogged points) during exceptionally warm days of the warm season. Exceptionally warm days were considered those with a daily maximum (at the dry point) belonging to the 90th percentile of daily maximums for the site. The 1:1 line (corresponding to no its it.
ie waterios buffer effect) is shown, points falling to the right of this line show a buffer effect (i.e. soil temperatures at the waterlogged point are cooler than the neighbouring dry site).

# 1 Mire microclimate: groundwater buffers temperature in

# 2 waterlogged versus dry soils

- 3 Short title: Testing the thermal buffer of mire groundwater
- 4 Eduardo Fernández-Pascual<sup>1+</sup>, Eva Correia-Álvarez<sup>2</sup>
- <sup>1</sup> Universidad de Oviedo, <sup>2</sup> Independent researcher
- 6 + Correspondence: Departamento de Biología de Organismos y Sistemas,
- 7 Universidad de Oviedo, C/ Catedrático Rodrigo Uría, 33006 Oviedo/Uviéu, Spain.
- 8 Email: efernandezpascual@gmail.com. Telephone: +34985104787.

### 9 Financial support

- 10 E.F.P. received financial support from the Government of Asturias and the FP7 -
- 11 Marie Curie COFUND programme of the European Commission (Grant 'Clarín'
- 12 ACB17-19).

13

#### Abstract

- 15 Ecosystems adapt differently to global warming through microclimate factors. Mires
- are sensitive wetland habitats that strongly rely on local soil properties, which makes
- 17 them a good model to understand how local climate parameters counteract the
- 18 effects of climate change. We quantified the temperature buffering effect in
- 19 waterlogged mire soils as compared with adjacent dry soils.
- 20 We buried dataloggers at 5 cm depth in waterlogged and dry points in 8 mires of the
- 21 Cantabrian Mountains (Spain, Southwestern Europe) and recorded soil
- 22 temperatures for c. 5 years. We also compared our local measures with air
- 23 temperatures predicted by the CHELSA model.
- 24 Waterlogged soils had less diurnal thermal amplitude (-2.3 °C), less annual thermal
- 25 amplitude (-5.1 °C), cooler summer maximums (-4.3 °C) and warmer winter
- 26 minimums (+0.8 °C). CHELSA air temperatures only correlated significantly (p <
- 27 0.05) with the winter minimum temperatures (Pearson's r > 0.83), and CHELSA
- 28 predictions were less accurate (higher RMSE) for waterlogged soils, except for the
- 29 summer maximums.
- We conclude that mire soils show a thermal buffer effect that insulates them from
- 31 the surrounding landscape. This effect is stronger at the warm end of the climatic
- 32 spectrum, i.e. during summer and at lower elevations. These results highlight the
- potential refugial character of mires under global warming, and the need to integrate
- 34 microclimate measurements into climate change models.

#### Keywords

35

bog, CHELSA, climatic model, datalogger, fen, peatland, soil temperature, wetland

#### Introduction

37

65

38 Climate change (IPCC 2014) affects global biodiversity, from drylands (Huang et al. 39 2016) to forests (Seidl et al. 2017) and oceans (Hoegh-Guldberg et al. 2017). An 40 accurate prediction of species responses requires to focus on physiologically 41 relevant climatic variables related to critical plant growth periods (Gardner, Maclean, 42 and Gaston 2019). Traditional models of species responses to global warming are 43 based on macroclimatic data from weather stations. Recently, several authors have 44 pointed out the need of complementing these models with in situ microclimatic 45 measures (Lembrechts, Nijs, and Lenoir 2019; Lembrechts et al. 2019; Maclean, 46 Mosedale, and Bennie 2019; Philippov and Yurchenko 2019). As has been shown 47 for European forests (Zellweger et al. 2020), ecosystems usually respond to broad 48 climatic changes through local processes. Increasingly there are downscaling efforts 49 based on atmospheric microclimate networks distributed in complex terrain that 50 focus on topographic effects on air temperature (Scherrer and Körner 2010), but 51 hydrologic factors under edaphic control require more attention (Ashcroft and Gollan 52 2013). 53 Mires are permanent semi-terrestrial peatlands whose soils remain waterlogged but 54 not inundated during most of the year (Wheeler and Proctor 2000). The term mire 55 encompasses peatlands that are classified as either ombrotrophic (rain-fed bogs) or 56 minerotrophic (groundwater-fed fens). These are azonal habitats whose existence 57 depends on local soil properties rather than macroclimatic zonation (Breckle 2002). 58 Mires meet certain criteria that make them especially vulnerable to climate change 59 (Horsák et al. 2018): (a) preponderance of species that evolved under a cold climate; 60 (b) low productivity due to nutrient limitation, making them sensitive to increased 61 nutrient cycling caused by warming (Cornelissen et al. 2007); and (c) scattered 62 distribution pattern, which limits species dispersal and migration (Pearson and 63 Dawson 2005). Therefore, mires are priority habitats for biodiversity conservation, 64 harbouring high numbers of endangered species (Bergamini et al. 2009), and

supporting highly adapted floras in spatially reduced areas (Grootjans et al. 2006).

66 Worryingly, mires retain high levels of methane and carbon which can be released 67 due to global warming (Koffi et al. 2020). Habitat distribution models have predicted 68 a loss of mire area as a consequence of ongoing climate change (Essl et al. 2012). 69 It has been known for a relatively long time that groundwater can produce a buffer 70 effect on soil temperature, keeping soils warmer than air during cold periods, and 71 vice versa (Frederick 1974; Ellenberg 1988; Geiger, Aron, and Todhunter 2009). 72 Clara M. Frederick (1974) showed the existence of a buffer effect comparing soil 73 temperatures taken at Cedar Bog (Ohio, United States) with temperatures from a 74 neighbouring agricultural station. Recently, more soil temperature measurements 75 have become available for mires of North America (Patrick A Raney, Fridley, and 76 Leopold 2014), Western Europe (Fernández-Pascual et al. 2015) and Central 77 Europe (Horsák et al. 2018). Their comparison with air temperatures derived from 78 models has shown that mire soils are indeed warmer in winter and cooler in summer, 79 thus giving support to the existence of the groundwater thermal buffer (Fernández-80 Pascual et al. 2015; Horsák et al. 2018). Furthermore, the effect has been linked to 81 the composition of mire flora and fauna (Horsák et al. 2018; Schenková et al. 2020), 82 the growth rings of mire trees (Raney et al. 2016) and the role of mires as glacial 83 refugia (Jiménez-Alfaro et al. 2016; Dítě et al. 2017). 84 Since root-zone temperature is a major determinant of plant ecophysiology (Körner 85 and Paulsen 2004), the groundwater buffer effect is expected to allow mire plants to 86 live in a wider range of air temperatures than they could otherwise. Indeed, mires 87 have a relatively homogeneous flora across their range despite their broad 88 distribution. In Europe, fens are distributed from the Iberian Peninsula to boreal 89 Fennoscandia and from low valleys to the alpine belt (Peterka et al. 2017). In the 90 arctic, temperate-continental and suboceanic regions of boreal Russia mires make 91 up an important part of the landscape, from 20 to 80 % of the surface of different 92 regions stretching from Europe to the Pacific (Yurkovskaya 2012). In the United 93 States, fens are distributed across the glaciated Midwest and Northeast, as well as 94 portions of the Appalachian Mountains and mountainous West (Bedford and Godwin

2003). However, continued aridification is expected to significantly reduce the overall
 extent of wetlands as it has been shown in the Midwestern United States (Johnson
 and Leopold 1994).

It is evident that the groundwater buffer effect will play a determinant role in the response of mire habitats to climate change (Patrick A. Raney 2014). Both bogs and fens are actively peat forming and are dependent on precipitation, whereas fens rely also on sources of telluric water from mineral ground. Moreover, they show a mosaic of microreliefs and a specific plant cover zonation which makes them especially sensitive to small changes in wetness. Climate change affects the quantity, timing and spatial distribution of precipitation, leading to changes in surface wetness which alter the intensity or organic decomposition by disturbing the conditions for plant grow and the depth of air penetration (Bragg 2002). Global warming would also result in warmer groundwater delivered to fens—and in drier conditions, but there is a gap of knowledge about the rate of such changes and their ecological consequences within the mires (Patrick A. Raney 2014).

As is the case for all azonal habitats, locally measured temperatures are essential to understand these processes. Available references recorded temperatures only on waterlogged soils, using model-derived air temperatures for comparison (Fernández-Pascual et al. 2015; Horsák et al. 2018; Schenková et al. 2020); or recorded temperatures in both wet and dry spots but for less than a year, lacking representation throughout the growth cycle of mire vegetation (Patrick A-Raney, Fridley, and Leopold 2014; Raney et al. 2016). This article provides the first measurement of the thermal buffer against surrounding non-mire areas, based on soil temperatures recorded during a period of five years. These measures can be used to support previous evidence about the general effects of soil moisture on thermal buffering (Patrick A-Raney, Fridley, and Leopold 2014; Fernández-Pascual et al. 2015; Horsák et al. 2018; Patrick A. Raney 2014) and to determine the magnitude of this buffering within microrefugia habitats. Specifically, we tested the hypotheses that, when compared to adjacent dry soils, waterlogged mire soilsoils

150

151

- are (i) warmer in winter and (ii) colder in summer; and have less thermal amplitude in (iii) daily and (iv) annual scales. In addition, we compared *in situ* measurements
- with data derived from the CHELSA climatic models (Karger et al. 2017).

#### **Materials and methods**

- 128 This study took place in the temperate oceanic region of north-western Spain (43° 129 N, 5° W) (Fig. 1a). Local geography and climate are dominated by the Cantabrian 130 Mountains (> 1500 m above sea level), which run parallel to the coast and trap the 131 prevailing NW Atlantic winds. In the study area, average annual precipitation ranges 132 from 800 mm at the low elevations to 1800 mm at the high mountains, while average 133 annual temperature ranges from 12.5 °C to 5 °C (source = Agencia Estatal de 134 http://www.aemet.es/es/serviciosclimaticos/). Under the 135 extreme emissions scenario, the latest projections expect temperatures to increase 136 by 4 °C and precipitation to decrease by c. 10 % by the end of the century, although 137 the precipitation projections are subjected to great uncertainty (Amblar-Francés et 138 al. 2017). The present humid and mild climate of this region harbours the south-139 western limit of mire communities in Europe (Jiménez-Alfaro, Díaz González, and 140 Fernández-Pascual 2011; Fernández Prieto, Fernández Ordóñez, and Collado 141 Prieto 1985). Rain-fed raised bogs (https://eunis.eea.europa.eu/habitats/260) are 142 very rare and appear locally on raised hummocks of acid valley mires 143 (https://eunis.eea.europa.eu/habitats/526) which can be found from the coast to just 144 below the treeline, in poorly drained valleys and former glacial lakes. Glacial lakes 145 undergoing silting develop transition mires and quaking bogs communities 146 (https://eunis.eea.europa.eu/habitats/1637) in the water-to-land transition. Spring 147 fens appear in the mountains above 1000 m; they range from soft-water poor fens 148 (https://eunis.eea.europa.eu/habitats/279) on acid bedrocks, to alkaline calcareous 149 fens (https://eunis.eea.europa.eu/habitats/277) on limestone.
  - We selected 8 mire sites representing the regional elevation gradient of mire vegetation and the different mire types (Table 1). Although we classify two of these

153

154

155

156

157

158

159

160

161

162

163

164

165

166

167

168

169

170

171

172

173

174

175

176

177

178

179

180

sites as rain-fed bogs based on their vegetation (Oxycocco-Sphagnetea Br.-Bl. et Tx. ex Westhoff et al. 1946), we must note that these bogs are very poorly developed on hummocks over valley mires and are in tight connection to the underlying water table coming from streams. Therefore, soil waterlogging in all our sites is fed by streams or springs. In each site, we buried two dataloggers (M-Log5W, GeoPrecision, Ettlingen, Germany; accuracy: +/- 0.1 °C (at 0 °C), resolution: 0.01 °C) at a depth of 5 cm below the upper layer of the soil. In this ecosystem, it is at this depth that true soil begins to develop, under the porous upper layers made up of live mosses. We installed one datalogger in a flat waterlogged spot within the mire; the other one in the close vicinity, but in a flat and dry upland area outside the mire (Fig. 1b). The vegetation at the sampling points was always either mire or pasture, with no shrubs, trees or any other landscape features shading the measuring points. Dataloggers recorded temperature once every hour and stayed on site for approximately five years, after which we retrieved them and downloaded their records. At the moment of retrieval, the internal clock of all dataloggers had not deviated for more than four hours.

All data processing and analysis was performed in R (R Core Team 2019). The original data, R code for the analysis and creation of the manuscript can be accessed at the GitHub repository https://github.com/efernandezpascual/mires. (The GitHub repository is private until publication, for peer review a copy can be downloaded at https://drive.google.com/file/d/1wnrNuQXvXcCruhKXdoFaRcm6tEZPaZkN/view?us p=sharing). To clean the logs we took the following steps: (i) removing records from the first week after installation, to account for the installation process and the settling of the soils; and (ii) keeping only time series with records for both the dry and waterlogged points, because some of the dataloggers had failed at different points in time. Afterwards, we calculated four bioclimatic variables for each datalogger: (1) the mean diurnal range; i.e. the average for the whole period of the daily differences between the maximum and the minimum temperatures recorded in the day; (2) the maximum temperature of the warmest month; i.e. the average of the daily maximum

182

183

184

185

186

187

188

189

190

191

192

193

194

195

196

197

198

199

200

201

202

203

204

205

206

records, for the warmest month; (3) the minimum temperature of the coldest month; i.e. the average of the daily minimum records, for the coldest month; and (4) the annual range; i.e. the difference between the maximum temperature of the warmest month and the minimum temperature of the coldest month. These are classical bioclimatic variables that indicate the limiting factors (extreme temperatures) and seasonality (diurnal and annual range) that organisms must tolerate (O'Donnell and Ignizio 2012). To compare our measurements with model-based predictions of air temperature, we downloaded from CHELSA the same bioclimatic variables for our measuring coordinates (https://chelsa-climate.org/bioclim/).

To test if the differences between dry and waterlogged points were significant, we used paired t-tests (n = 8 paired sites per each of the four bioclimatic variables). Tests were one-tailed, according to the original hypotheses: the dry point would have a higher diurnal range, a higher maximum temperature, a lower minimum temperature, and a higher annual range. To compare our soil measurements with the CHELSA air temperatures we calculated Pearson's correlation between both sets of values, and we also calculated the root-mean-square error (RMSE) of the CHELSA values as a measure of their accuracy (n = 8 sites per combination of bioclimatic variable and dry/waterlogged condition). We calculated RMSE using its implementation in the package Metrics (Hamner and Frasco 2018). We checked whether our data complied with the normality assumption of the t-test and Pearson's correlation using the Shapiro-Wilk normality test; the assumption was met in all cases (p > 0.05). To check whether the analyses were influenced by the fact that recording periods did not match across all eight sites, we repeated the analyses using only full-year, contemporaneous data series (01 Jan 2015 - 31 Dec 2017, excluding La Vega Comeya); the results of these analyses were the same as the ones presented in this article (Supplementary Tables S1 and S2).

#### Results

207

- 208 Dataloggers recorded temperatures for five years in five of the sites, four years in
- 209 two, and two years in one. Time series showed considerably less variation at
- 210 waterlogged points, for all eight site comparisons (Fig. 2).
- 211 Bioclimatic variables (Table 2 and Fig. 3) supported this notion. The mean annual
- 212 range was wider at dry points, as was the mean diurnal range. Maximum
- 213 temperatures of the warmest months were usually higher at dry measuring points.
- 214 The opposite was true for the minimums of the coldest months, in which case
- 215 temperature was generally colder at dry points, although the difference was less
- 216 pronounced than for the maximums (Fig. 3).
- 217 Patterns regarding the maximum temperatures and annual range were especially
- 218 noticeable at both low (El Molinucu, La Malva) and high (La Recoleta) elevations.
- 219 The pattern for the diurnal range, however, was more prominent just at the low sites
- 220 (El Molinucu, La Malva), whereas the minimums of the coldest months showed no
- 221 specific pattern (Table 2).
- 222 T-tests supported the original hypotheses; namely waterlogged measuring points
- had (a) smaller diurnal fluctuations (t = -3.05, p = 0.009, effect size = -2.29  $^{\circ}$ C, 95  $^{\circ}$ C
- 224 CI =  $-\infty$ , -0.87); (b) lower maximums (t = -3.04, p = 0.009, effect size = -4.28 °C, 95
- 225 % CI =  $-\infty$ , -1.61); (c) higher minimums (t = 2.86, p = 0.012, effect size = 0.77 °C, 95
- % CI = 0.26, ∞), and (d) smaller annual fluctuations (t = -3.95, p = 0.003, effect size
- 227 = -5.05 °C, 95 % CI =  $-\infty$ , -2.63).
- 228 The CHELSA climatic model (Table 3 and Fig. 4) provided air temperatures that
- significantly correlated (p < 0.05) with soil temperatures only in the case of the
- 230 minimums of the coldest month, both at dry points (Pearson's r = 0.90) and
- waterlogged points (Pearson's r = 0.84). The correlation between CHELSA and soil
- values was marginally significant (p = 0.05) in the case of the maximums of the
- warmest month, but only at dry points (Pearson's r = 0.71). The values of CHELSA
- 234 did not correlate with soil temperatures in the rest of the cases (p > 0.05). RMSE

236

237

238

239

240

241

242

243

244

245

246

247

248

249

250

251

252

253

254

255

256

257

258

259

260

261

262

indicated that the predictions of CHELSA were more accurate at dry points for the cases of the annual range, the diurnal range and the winter min; the CHELSA values were more accurate at waterlogged points in the case of the summer max.

#### **Discussion**

The results presented here quantify the size of the thermal buffer effect that takes place in waterlogged mire soils (Frederick 1974), when compared with adjacent dry soils. The mire thermal buffer had been compared previously with air temperatures derived from models, with generally similar results (Fernández-Pascual et al. 2015; Horsák et al. 2018). The mire buffer had also been compared to dry soils at 10 cm depth (Patrick A Raney, Fridley, and Leopold 2014; Raney et al. 2016); our results confirm those findings at 5 cm depth and extend them to the full year. Also, in concordance with our results, the air temperature measured at 0.5 m from the surface of boreal Russian bogs is lower at the wetter zones, at least during the warmest months (Philippov and Yurchenko 2019). Thus, the pattern is reproducible among years (out to five years), seems to take place both at southern and northern latitudes, and may affect not only the temperature in the soil but also air temperatures within a certain distance from the soil. The thermal buffer effect of mire soils that makes their temperatures less extreme than the surrounding landscape is therefore a general phenomenon. The existence of this buffer effect highlights the importance of using fine-scale microclimatic data to assess vegetation responses to climate change (Storlie et al. 2014; Lembrechts, Nijs, and Lenoir 2019; Zellweger et al. 2020).

One important difference with previous studies is the magnitude of the buffer during winter. The articles that had used model air temperatures as a control concluded that the buffer effect was stronger at the cold end of the thermal gradientspectrum, i.e. in winter and at night (Fernández-Pascual et al. 2015; Horsák et al. 2018). In the case of this investigation, the situation is the reverse: the effect is weaker when considering the minimum temperatures of the cold period. This indicates the

importance of identifying root temperatures when working with plant communities. At high elevations of the study region, the soil can remain covered by snow for periods of winter, and this has its own insulating effect on soil temperatures (Körner 2003). Indeed, snow cover has being described as one of the vertical features that affects vegetation distribution in a local manner (Maclean, Mosedale, and Bennie 2019). In any case, since this study was conducted at the southern limit of mire distribution, more research on the winter buffer is needed at high latitudes, since it might have a deeper biological significance there.

263

264

265

266

267

268

269

270

271

272

273

274

275

276

277

278

279

280

281

282

283

284

285

286

287

288

289

290

291

In these southern mires, the buffering effect is much stronger during the summer (Fig. 5). This is most noticeable in the two lowest elevations, El Molinucu and La Malva. La Malva shows especially high summer temperatures within the dry soils (> 40 °C). It must be noted that this difference of > 15°C15 °C between the dry and waterlogged points of La Malva, which are separated by a few meters, is almost four times the warming expected in the study region at the end of this century (+ 4 °C) under the more extreme emissions scenario (Amblar-Francés et al. 2017). Whereas the rest of the study sites are flat, La Malva is a calcareous spring in a southeastfacing slope on limestone, a place experiencing sub-Mediterranean conditions at the micro-scale (Sánchez de Dios, Benito-Garzón, and Sainz-Ollero 2009), and surrounded by a forest of evergreen oaks (Quercus rotundifolia Lam., Quercus faginea Lam.). This suggests the importance of groundwater in the existence of mire vegetation in Mediterranean areas (De Hoyos et al. 1996), not only from the wateravailability aspect, but also providing cooling regulation during summer (Ellenberg 1988). The buffering is also prominent at one of the highest elevations, La Recoleta. which may be explained by the high saturation of water in this kind of habitat, a quaking bog, i.e. a former glacial lake undergoing silting. However, the relatively small number of sites in our study does not allow to explore further the moderating factors that may drive differences in the strength of the buffer. More studies focusing on each kind of habitat are needed to understand the relation between hydrology conditions and microclimatic effects.

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

310

311

312

313

314

315

316

317

318

319

320

Most models currently employed to predict vegetation and species distribution use macroclimatic parameters, like the ones obtained from CHELSA (Karger et al. 2017). This study revealed that CHELSA values correlate relatively well with soil temperatures in the case of the minimum temperatures of the coldest month, but not with the rest of the bioclimatic variables under examination. Moreover, CHELSA predictions are less accurate for waterlogged soils, except for the summer max, which might be the consequence of the noise introduced by sun-heated soil. In addition to the water buffering effect, other features may explain the differences between our results and CHELSEA values, like the height of temperature measurement (5 cm below the upper layer of the soil in our study versus 2 m above the ground in CHELSA); the temporal scale of observation (hourly based for a 5 years period in our study versus a 34 years period); and the spatial resolution (c. 1 m in our study versus c. 1 km). Be that as it may, our results bring out the importance of considering local factors as drivers of microclimatic changes (Zellweger et al. 2019). These factors contribute to landscape heterogeneity, producing safe sites that can act as micro-refugees refuges and buffer species from regional climatic warming.

In summary, this article shows that waterlogged mire soils have a thermal buffer when compared to adjacent soils, contributing to their behaviour as mild island habitats in a landscape that can be more thermally variable (Patrick A. Raney 2014; Horsák et al. 2018). The effect occurs during cold and warm periods, but it is stronger during the summer, at least in the study area. However, this climatic buffering within microrefugia relies strongly on the water regime, a key factor for its conservation. Long-term decreases in precipitation could reduce the groundwater discharge into the mires, with potentially fatal consequences if the mires dry out during warmer summers. Rain-fed bogs will be affected more immediately than spring fens since the latter are somewhat decoupled from short-term rainfall variation. However, long-term regional drying would affect all types of mires. For the study area, the latest projections envision a decrease of c. 10 % in annual precipitation at the end of the

century, but these projections are still subjected to a high level of uncertainty (Amblar-Francés et al. 2017). Future recording schemes are needed to monitor the evolution of the soil thermal buffer in drying regions, to obtain local soil temperatures from other latitudes, and from more microtopographies within the same mire. We must also consider that the microclimatic effect is not limited to mires as it can also be found in other habitats where a gradient of moisture exists (Ashcroft and Gollan 2013), such as the mosaic of microtopographies (snow patches, fellfields) that is characteristic of alpine environments (Scherrer and Körner 2010; Ohler, Lechleitner, and Junker 2020). This study provides useful microclimate parameters to improve the current models that predict the impact of global warming on moisture-driven ecosystems.

#### Data availability

The original data, as well as the R code for the analysis and creation of the manuscript, can be accessed at the GitHub repository https://github.com/efernandezpascual/mires. Upon publication, a version of record of the repository will be deposited in Zenodo. (The GitHub repository is private until publication, for peer-review a copy can be downloaded at https://drive.google.com/file/d/1wnrNuQXvXcCruhKXdoFaRcm6tEZPaZkN/view?us p=sharing)

## **Supporting information**

<u>Tables S1-S2</u> Results of the statistical analyses, repeated using only the contemporaneous full-year data series.

#### References

Amblar-Francés, P, MJ Casado-Calle, MA Pastor-Saavedra, P Ramos-Calzado, and E Rodríguez-Camino. 2017. *Guía de escenarios regionalizados de cambio* 

- 346 climático sobre España a partir de los resultados del IPCC-AR5. Madrid:
- 347 Ministerio de Agricultura y Pesca, Alimentación y Medio Ambiente. Agencia
- 348 Estatal de Meteorología.
- 349 Ashcroft, Michael B., and John R. Gollan. 2013. "Moisture, thermal inertia, and the
- spatial distributions of near-surface soil and air temperatures: understanding
- factors that promote microrefugia." Agricultural and Forest Meteorology 176: 77–
- 352 89. https://doi.org/https://doi.org/10.1016/j.agrformet.2013.03.008.
- 353 Bedford, Barbara L., and Kevin S. Godwin. 2003. "Fens of the United States:
- distribution, characteristics, and scientific connection versus legal isolation."
- 355 *Wetlands* 23 (3): 608–29. https://doi.org/10.1672/0277-
- 356 5212(2003)023[0608:FOTUSD]2.0.CO;2.
- 357 Bergamini, A., M. Peintinger, S. Fakheran, H. Moradi, B. Schmid, and J. Joshi. 2009.
- 358 "Loss of habitat specialists despite conservation management in fen remnants
- 359 1995-2006." Perspectives in Plant Ecology, Evolution and Systematics 11 (1): 65–
- 360 79. https://doi.org/10.1016/j.ppees.2008.10.001.
- 361 Bragg, O. M. 2002. "Hydrology of peat-forming wetlands in Scotland." *Science of the*
- 362 Total Environment 294 (1-3): 111–29. https://doi.org/10.1016/s0048-
- 363 9697(02)00059-1.
- 364 Breckle, Siegmar-Walter. 2002. Walter's Vegetation of the Earth: the Ecological
- 365 Systems of the Geo-biosphere. Berlin Heidelberg New York: Springer.
- 366 Cornelissen, Johannes H. C., Peter M. Van Bodegom, Rien Aerts, Terry V.
- Callaghan, Richard S. P. Van Logtestijn, Juha Alatalo, F. Stuart Chapin, et al.
- 368 2007. "Global negative vegetation feedback to climate warming responses of leaf
- litter decomposition rates in cold biomes." *Ecology Letters* 10 (7): 619–27.
- 370 https://doi.org/10.1111/j.1461-0248.2007.01051.x.

- 371 De Hoyos, D, Jesús Muñoz Fuente, A Negro, Juan José Aldasoro, JC Vega, and
- Gonzalo Moreno Moral. 1996. "A survey on Cantabrian mires (Spain)." *Anales Del*
- 373 *Jardín Botánico de Madrid* 54 (1): 472–89.
- 374 Dítě, Daniel, Tomáš Peterka, Zuzana Dítětová, Petra Hájková, and Michal Hájek.
- 375 2017. "Arcto-Alpine species at their niche margin: the Western Carpathian refugia
- of Juncus castaneus and J. triglumis in Slovakia." Annales Botanici Fennici 54 (1-
- 377 3): 67–82, 16. https://doi.org/10.5735/085.054.0311.
- 378 Ellenberg, Heinz. 1988. "Spring areas and adjacent swamps." In *Vegetation Ecology*
- of Central Europe, 313–13. Cambridge: Cambridge University Press.
- 380 Essl, Franz, Stefan Dullinger, Dietmar Moser, Wolfgang Rabitsch, and Ingrid
- 381 Kleinbauer. 2012. "Vulnerability of mires under climate change: implications for
- 382 nature conservation and climate change adaptation." Biodiversity and
- 383 *Conservation* 21: 655–69.
- 384 Fernández-Pascual, Eduardo, Borja Jiménez-Alfaro, Michal Hájek, Tomás E. Díaz,
- and Hugh W. Pritchard. 2015. "Soil thermal buffer and regeneration niche may
- favour calcareous fen resilience to climate change." Folia Geobotanica 50 (4):
- 387 293–301. https://doi.org/10.1007/s12224-015-9223-y.
- 388 Fernández Prieto, José Antonio, María del Carmen Fernández Ordóñez, and Miguel
- Angel Collado Prieto. 1985. "Datos sobre la vegetación de las turberas de
- esfagnos galaico-asturianas y orocantábricas." *Lazaroa* 7: 443–71.
- 391 Frederick, Clara May. 1974. "A natural history study of the vascular flora of Cedar
- Bog, Champaign County, Ohio." The Ohio Journal of Science 74 (2): 65–116.
- 393 Gardner, Alexandra S., Ilya M. D. Maclean, and Kevin J. Gaston. 2019. "Climatic
- 394 predictors of species distributions neglect biophysiologically meaningful
- 395 variables." Diversity and Distributions 25 (8): 1318–33.
- 396 https://doi.org/10.1111/ddi.12939.

- 397 Geiger, Rudolf, Robert H. Aron, and Paul Todhunter. 2009. *The Climate Near the*
- 398 *Ground*. Lanham: Rowman & Littlefield.
- 399 Grootjans, A. P., E. B. Adema, W. Bleuten, H. Joosten, M. Madaras, and M.
- Janáková. 2006. "Hydrological landscape settings of base-rich fen mires and fen
- 401 meadows: an overview." Applied Vegetation Science 9: 175-84
- http://search.ebscohost.com/login.aspx?direct=true&db=aph&AN=25691039&la
- 403 ng=es&site=ehost-live.
- Hamner, B, and M Frasco. 2018. "Metrics: Evaluation Metrics for Machine Learning.
- 405 R package version 0.1.4. https://CRAN.R-project.org/package=Metrics."
- 406 Hoegh-Guldberg, Ove, Elvira S. Poloczanska, William Skirving, and Sophie Dove.
- 407 2017. "Coral reef ecosystems under climate change and ocean acidification."
- 408 Frontiers in Marine Science 4 (158). https://doi.org/10.3389/fmars.2017.00158.
- 409 Horsák, Michal, Vendula Polášková, Marie Zhai, Jindřiška Bojková, Vít Syrovátka,
- Vanda Šorfová, Jana Schenková, Marek Polášek, Tomáš Peterka, and Michal
- Hájek. 2018. "Spring-fen habitat islands in a warming climate: partitioning the
- 412 effects of mesoclimate air and water temperature on aquatic and terrestrial biota."
- 413 Science of the Total Environment 634: 355–65.
- 414 https://doi.org/https://doi.org/10.1016/j.scitotenv.2018.03.319.
- Huang, Jianping, Haipeng Yu, Xiaodan Guan, Guoyin Wang, and Ruixia Guo. 2016.
- 416 "Accelerated dryland expansion under climate change." *Nature Climate Change*
- 417 6 (2): 166–71. https://doi.org/10.1038/nclimate2837.
- 418 IPCC. 2014. Climate Change 2014: Synthesis Report. Contribution of Working
- 419 Groups I, II and III to the Fifth Assessment Report of the Intergovernmental Panel
- on Climate Change [Core Writing Team, R.K. Pachauri and L.A. Meyer (eds.)].
- 421 Geneva: IPCC.

- 422 Jiménez-Alfaro, Borja, Tomás Díaz González, and Eduardo Fernández-Pascual.
- 423 2011. "Grupos de vegetación y hábitats de tremedales neutro-basófilos en las
- 424 montañas pirenaico-cantábricas." Acta Botanica Barcinonensia 53: 47–60.
- 425 Jiménez-Alfaro, Borja, Laura García-Calvo, Pedro García, and José Luis Acebes.
- 426 2016. "Anticipating extinctions of glacial relict populations in mountain refugia."
- 427 Biological Conservation 201: 243–51.
- 428 Johnson, Anne M., and Donald J. Leopold. 1994. "Vascular plant species richness
- and rarity across a minerotrophic gradient in wetlands of St. Lawrence County,
- 430 New York, USA." Biodiversity and Conservation 3 (7): 606–27.
- 431 https://doi.org/10.1007/BF00114204.
- 432 Karger, Dirk Nikolaus, Olaf Conrad, Jürgen Böhner, Tobias Kawohl, Holger Kreft,
- 433 Rodrigo Wilber Soria-Auza, Niklaus E. Zimmermann, H. Peter Linder, and Michael
- Kessler. 2017. "Climatologies at high resolution for the earth's land surface
- 435 areas." Scientific Data 4 (1): 170122. https://doi.org/10.1038/sdata.2017.122.
- 436 Koffi, Ernest N., Peter Bergamaschi, Romain Alkama, and Alessandro Cescatti.
- 437 2020. "An observation-constrained assessment of the climate sensitivity and
- future trajectories of wetland methane emissions." Science Advances 6 (15):
- 439 eaay4444. https://doi.org/10.1126/sciadv.aay4444.
- 440 Körner, Christian. 2003. Alpine Plant Life: Functional Plant Ecology of High Mountain
- 441 *Ecosystems*. 2nd ed. Berlin Heidelberg New York: Springer.
- 442 Körner, Christian, and Jens Paulsen. 2004. "A world-wide study of high altitude
- treeline temperatures." Journal of Biogeography 31: 713–32.
- 444 http://10.0.4.87/j.1365-2699.2003.01043.x
- http://search.ebscohost.com/login.aspx?direct=true&db=aph&AN=12823821&la
- 446 ng=es&site=ehost-live.
- Lembrechts, Jonas J., Jonathan Lenoir, Nina Roth, Tarek Hattab, Ann Milbau, Sylvia
- Haider, Loïc Pellissier, et al. 2019. "Comparing temperature data sources for use

- in species distribution models: from in-situ logging to remote sensing." Global
- 450 Ecology and Biogeography 28 (11): 1578–96. https://doi.org/10.1111/geb.12974.
- 451 Lembrechts, Jonas J., Ivan Nijs, and Jonathan Lenoir. 2019. "Incorporating
- 452 microclimate into species distribution models." *Ecography* 42 (7): 1267–79.
- 453 https://doi.org/10.1111/ecog.03947.
- 454 Maclean, Ilya M. D., Jonathan R. Mosedale, and Jonathan J. Bennie. 2019.
- 455 "Microclima: an R package for modelling meso- and microclimate." *Methods in*
- 456 Ecology and Evolution 10 (2): 280–90. https://doi.org/10.1111/2041-210x.13093.
- 457 O'Donnell, Michael S, and Drew A Ignizio. 2012. "Bioclimatic predictors for
- 458 supporting ecological applications in the conterminous United States." US
- 459 Geological Survey Data Series 691 (10).
- 460 Ohler, Lisa-Maria, Martin Lechleitner, and Robert R. Junker. 2020. "Microclimatic
- 461 effects on alpine plant communities and flower-visitor interactions." Scientific
- 462 Reports 10 (1): 1366. https://doi.org/10.1038/s41598-020-58388-7.
- Pearson, Richard G., and Terence P. Dawson. 2005. "Long-distance plant dispersal
- and habitat fragmentation: identifying conservation targets for spatial landscape
- planning under climate change." Biological Conservation 123 (3): 389–401.
- 466 https://doi.org/https://doi.org/10.1016/j.biocon.2004.12.006.
- 467 Peterka, Tomáš, Michal Hájek, Martin Jiroušek, Borja Jiménez-Alfaro, Liene Aunina,
- Ariel Bergamini, Daniel Dítě, et al. 2017. "Formalized classification of European
- fen vegetation at the alliance level." *Applied Vegetation Science* 20 (1): 124–42.
- 470 https://doi.org/10.1111/avsc.12271.
- 471 Philippov, Dmitriy A., and Victoria V. Yurchenko. 2019. "Data on air temperature,
- relative humidity and dew point in a boreal *Sphagnum* bog and an upland site
- 473 (Shichengskoe mire system, North-Western Russia)." Data in Brief 25: 104156.
- 474 https://doi.org/https://doi.org/10.1016/j.dib.2019.104156.

- 475 Raney, Patrick A. 2014. *Identifying potential refugia from climate change in wetlands*.
- 476 Syracuse, New York: Doctoral thesis, State University of New York.
- 477 Raney, Patrick A, Jason D Fridley, and Donald J Leopold. 2014. "Characterizing
- 478 microclimate and plant community variation in wetlands." *Wetlands* 34 (1): 43–53.
- 479 https://doi.org/10.1007/s13157-013-0481-2.
- 480 Raney, Patrick A., Donald J. Leopold, Martin Dovčiak, and Colin M. Beier. 2016.
- 481 "Hydrologic position mediates sensitivity of tree growth to climate: groundwater
- subsidies provide a thermal buffer effect in wetlands." Forest Ecology and
- 483 *Management* 379: 70–80.
- 484 R Core Team. 2019. "R: a language and environment for statistical computing."
- 485 https://www.r-project.org/.
- 486 Sánchez de Dios, Rut, Marta Benito-Garzón, and Helios Sainz-Ollero. 2009.
- 487 "Present and future extension of the Iberian submediterranean territories as
- determined from the distribution of marcescent oaks." *Plant Ecology* 204 (2): 189–
- 489 205. https://doi.org/10.1007/s11258-009-9584-5.
- 490 Schenková, Jana, Vendula Polášková, Martina Bílková, Jindřiška Bojková, Vít
- 491 Syrovátka, Marek Polášek, and Michal Horsák. 2020. "Climatically induced
- temperature instability of groundwater-dependent habitats will suppress cold-
- 493 adapted Clitellata species." *International Review of Hydrobiology* 105: 85–93.
- 494 https://doi.org/10.1002/iroh.201902006.
- 495 Scherrer, Daniel, and Christian Körner. 2010. "Infra-red thermometry of alpine
- 496 landscapes challenges climatic warming projections." Global Change Biology 16
- 497 (9): 2602–13. https://doi.org/10.1111/j.1365-2486.2009.02122.x.
- 498 Seidl, Rupert, Dominik Thom, Markus Kautz, Dario Martin-Benito, Mikko Peltoniemi,
- 499 Giorgio Vacchiano, Jan Wild, et al. 2017. "Forest disturbances under climate
- 500 change." *Nature Climate Change* 7 (6): 395–402.
- 501 https://doi.org/10.1038/nclimate3303.

502	Storlie, Collin, Andres Merino-Viteri, Ben Phillips, Jeremy VanDerWal, Justin
503	Welbergen, and Stephen Williams. 2014. "Stepping inside the niche: microclimate
504	data are critical for accurate assessment of species' vulnerability to climate
505	change." <i>Biology Letters</i> 10 (9): 20140576.
506	https://doi.org/doi:10.1098/rsbl.2014.0576.
507	Wheeler, B. D., and M. C. F. Proctor. 2000. "Ecological gradients, subdivisions and
508	terminology of north-west European mires." Journal of Ecology 88 (2): 187–203.
509	https://doi.org/10.1046/j.1365-2745.2000.00455.x.
510	Yurkovskaya, Tatiana. 2012. "Mires on the map of Russia." In Mires from Pole to
511	Pole, edited by Tapio Lindholm and Raimo Heikkilä, 31–38. Helsinki: Finnish
512	Environment Institute.
513	Zellweger, Florian, Pieter De Frenne, Jonathan Lenoir, Duccio Rocchini, and David
514	Coomes. 2019. "Advances in microclimate ecology arising from remote sensing."
515	Trends in Ecology & Evolution 34 (4): 327–41.
516	https://doi.org/https://doi.org/10.1016/j.tree.2018.12.012.
517	Zellweger, Florian, Pieter De Frenne, Jonathan Lenoir, Pieter Vangansbeke, Kris
518	Verheyen, Markus Bernhardt-Römermann, Lander Baeten, et al. 2020. "Forest
519	microclimate dynamics drive plant responses to warming." Science 368 (6492):
520	772–75. https://doi.org/10.1126/science.aba6880.
521	

**Tables** 

Table 1: Mire sites included in this study, indicating the type of fen, pH, conductivity, elevation, coordinates, and length of the temperature recording period. Groundwater pH and conductivity values are the average of 5–10 measures per site taken with a MM40 Portable Multimeter (Crison Instruments S.A., Alella, Spain). Coordinates are in decimal degrees WGS84.

Site	Habitat	рН	Conductivity (µS/cm)	Elevation (m)	Latitude	Longitude	Records (days)
El Molinucu	Raised bog	4.2	47.2	284	43.3924	-5.5392	1421
La Malva	Alkaline fen	8.2	427.7	700	43.1176	-6.2543	1347
La Vega Comeya	Raised bog	5.9	102.4	822	43.2856	-4.9885	664
La Bruxa	Alkaline fen	8.0	271.8	1528	43.0232	-6.2113	1850
La Veiga Cimera	Acid fen	4.6	13.9	1552	43.0252	-6.2539	1850
La Recoleta	Quaking bog	7.2	245.9	1768	43.0167	-6.1112	1854
El Riotuertu	Alkaline fen	7.3	400.1	1820	43.0096	-5.9479	1852
La Vega Lliordes	Alkaline fen	7.3	293.3	1878	43.1504	-4.8464	1809

531

532

Table 2: Groundwater buffer effect per mire and bioclimatic variable. The buffer effect was calculated as the difference between the value in the waterlogged and the dry points.

Site	Annual	Diurnal	Summer	Winter min
Onc	range	range	max	vviii(Ci iiiiii
El Molinucu	-7.80	-4.90	-6.59	1.21
La Malva	-10.81	-5.14	-10.25	0.56
La Vega Comeya	-3.63	-2.99	-3.51	0.12
La Bruxa	-1.09	-0.88	0.99	2.08
La Veiga Cimera	-3.41	-2.21	-3.47	-0.06
La Recoleta	-8.81	-2.62	-8.85	-0.04
El Riotuertu	-3.62	1.12	-2.28	1.34
La Vega Lliordes	-1.26	-0.71	-0.28	0.98

**5**33

Table 3: ResultsComparison of the linear models predicting the soil bioclimatic variables from the with CHELSA air temperatures, per bioclimatic variable and groundwater situation. RMSE is the root-mean-square error, a measure of the accuracy of the predictions (lower values indicate higher accuracy).

536

537

Variable	Groundwate r	t	df	р	Pearson's r	RMSE (°C)
Annual range	Dry	0.012	6	0.991	0.00	3.62
Annual range	Waterlogge d	-0.334	6	0.750	-0.14	4.71
Diurnal range	Dry	0.016	6	0.988	0.01	3.09
Diurnal range	Waterlogge d	1.354	6	0.225	0.48	4.62
Summer max	Dry	2.450	6	0.050	0.71	5.86
Summer max	Waterlogge d	1.527	6	0.178	0.53	2.95
Winter min	Dry	5.115	6	0.002	0.90	4.04
Winter min	Waterlogge d	3.809	6	0.009	0.84	4.82

### 539 Figures

540

541

542

543

544

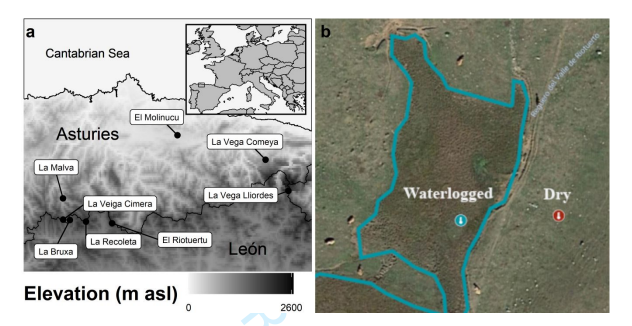


Figure 1: (a) Location of the study sites in northern Spain. (b) Representation of the sampling design at El Riotuertu site. The position of the dry and waterlogged temperature loggers is indicated, the blue area represents the extent of the waterlogged mire habitat.

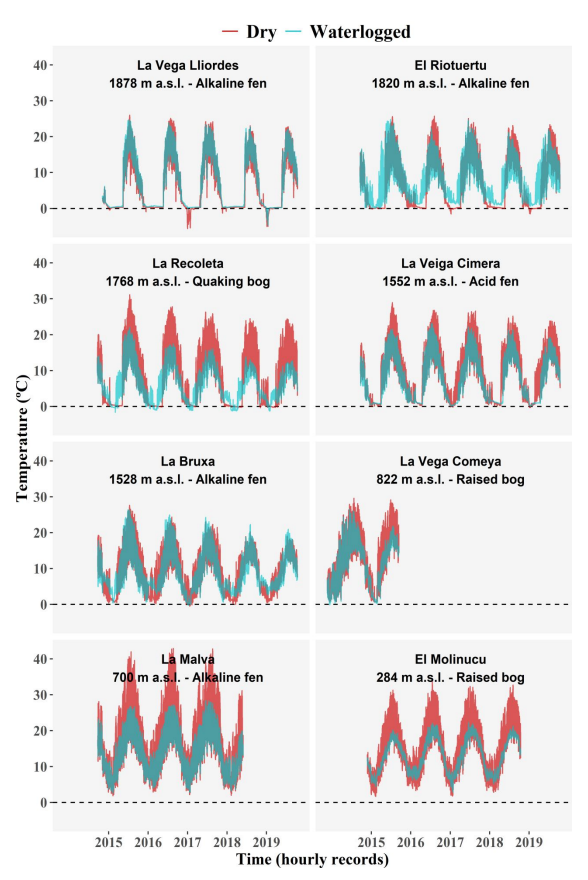


Figure 2: Hourly soil temperature records at the mire sites. The blue series was recorded within the mire, in a waterlogged area. The red series was recorded in a neighbouring dry area. Dataloggers were buried at 5 cm depth. The amplitude of diurnal temperature range in dry sites was reduced with increasing elevation, where the thermal buffer effect was also less apparent.

51

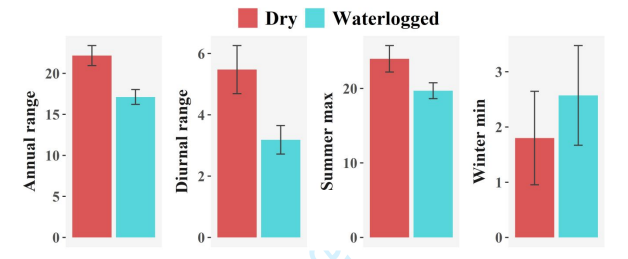


Figure 3: Average bioclimatic variables in the dry and waterlogged points. The bars represent the mean value, and the brackets the standard error of 8 records.

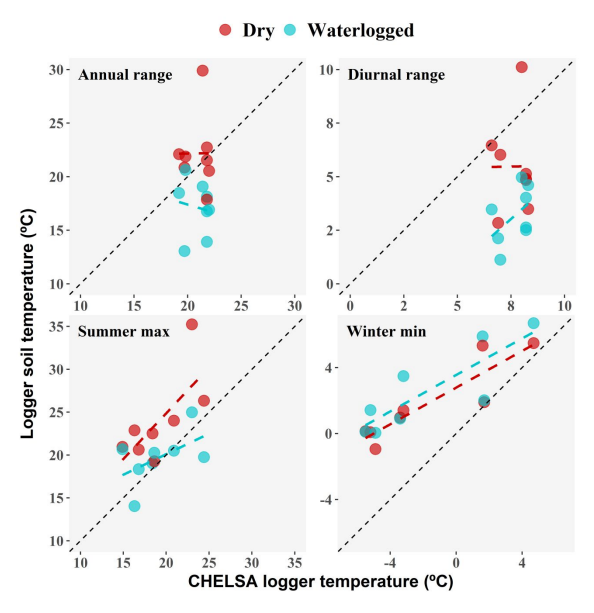


Figure 4: Scatter plots of the bioclimatic variables predicted by CHELSA air temperatures vs. soil temperatures measured in situ, in dry and waterlogged points.

556

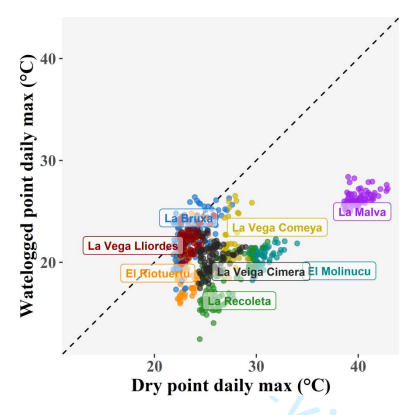


Figure 5: Complementary temporal analysis of the groundwater buffer effect during peak hot events. Variation of soil temperatures at paired sites (dry vs. waterlogged points) during exceptionally warm days of the warm season. Exceptionally warm days were considered those with a daily maximum (at the dry point) belonging to the 90th percentile of daily maximums for the site. The 1:1 line (corresponding to no buffer effect) is shown, points falling to the right of this line show a buffer effect (i.e. soil temperatures at the waterlogged point are cooler than the neighbouring dry site).

# **Supporting information for Mire microclimate:**

# 2 groundwater buffers temperature in waterlogged versus

## 3 dry soils

- 4 Table S1: Results of the t-tests comparing the paired waterlogged-dry soil
- 5 bioclimatic variables. Analysis performed with full-year, contemporaneous data
- 6 series (2015-2017).

Trait	t.statistic	t.p.value	t.estimate	t.conf.int.1.	t.conf.int.2.	
Annual	-3.609308	0.00562013	-5.1431459	-Inf	-2.374177	
range	0.000000	7	0.1101100		2.017111	
Diurnal	-2.643586	0.01917802	-2.2947974	-Inf	-0.607996	
range	2.0.0000	4		•••	2.237.000	
Summer	-2.737935	0.01691408	-4.3721276	-Inf	-1.269120	
max		8		•••		
Winter min	2.730278	0.01708671	0.7710183	0.2222728	Inf	

- 7 Table S2: Results of the linear models predicting the soil bioclimatic variables from
- 8 the CHELSA air temperatures, per bioclimatic variable and groundwater situation.
- 9 RMSE is the root-mean-square error, a measure of the accuracy of the predictions
- 10 (lower values indicate higher accuracy). Analysis performed with full-year,
- 11 contemporaneous data series (2015-2017).

Variable	Groundwate r	t	df	р	Pearson's r	RMSE (°C)
Annual	Dry	0.180	5	0.865	0.08	3.68
range	ыу	0.100	3	0.003	0.00	3.00

Variable	Groundwate r	t	df	р	Pearson's r	RMSE (°C)
Annual range	Waterlogge d	0.424	5	0.689	0.19	3.90
Diurnal range	Dry	0.489	5	0.645	0.21	3.03
Diurnal range	Waterlogge d	2.359	5	0.065	0.73	4.47
Summer max	Dry	2.213	5	0.078	0.70	6.92
Summer max	Waterlogge d	0.970	5	0.376	0.40	3.68
Winter min	Dry	6.774	5	0.001	0.95	4.35
Winter min	Waterlogge d	6.812	5	0.001	0.95	5.06
12			4			

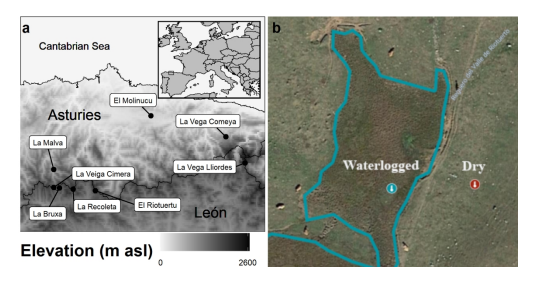


Figure 1: (a) Location of the study sites in northern Spain. (b) Representation of the sampling design at El Riotuertu site. The position of the dry and waterlogged temperature loggers is indicated, the blue area represents the extent of the waterlogged mire habitat.

169x84mm (300 x 300 DPI)

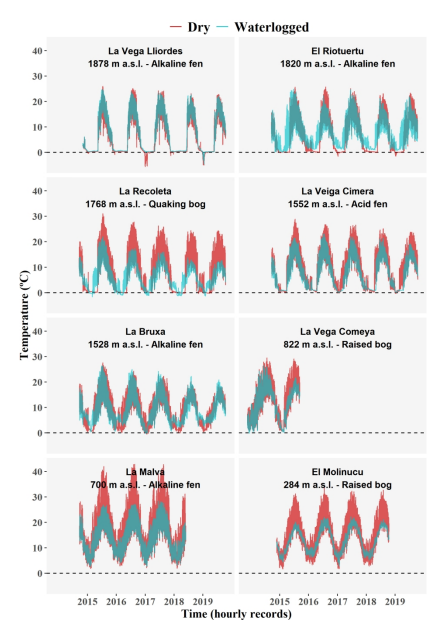


Figure 2: Hourly soil temperature records at the mire sites. The blue series was recorded within the mire, in a waterlogged area. The red series was recorded in a neighbouring dry area. Dataloggers were buried at 5 cm depth. The amplitude of diurnal temperature range in dry sites was reduced with increasing elevation, where the thermal buffer effect was also less apparent.

169x249mm (300 x 300 DPI)

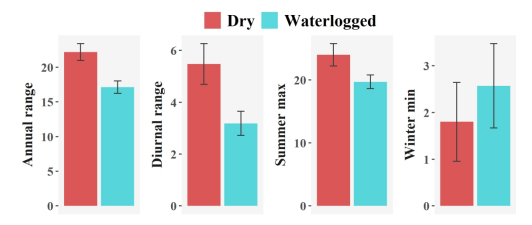


Figure 3: Average bioclimatic variables in the dry and waterlogged points. The bars represent the mean value, and the brackets the standard error of 8 records.

169x69mm (300 x 300 DPI)

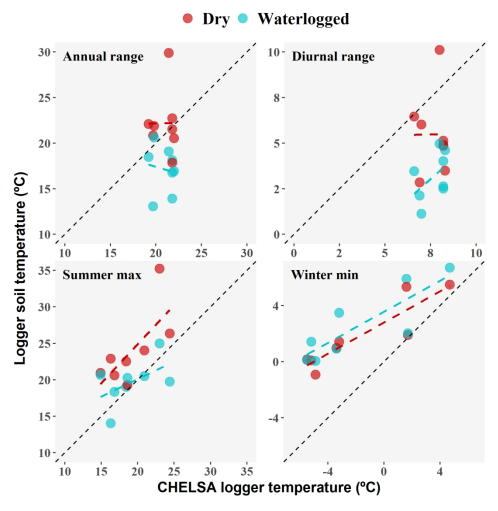


Figure 4: Scatter plots of the bioclimatic variables predicted by CHELSA air temperatures vs. soil temperatures measured in situ, in dry and waterlogged points.

169x169mm (300 x 300 DPI)

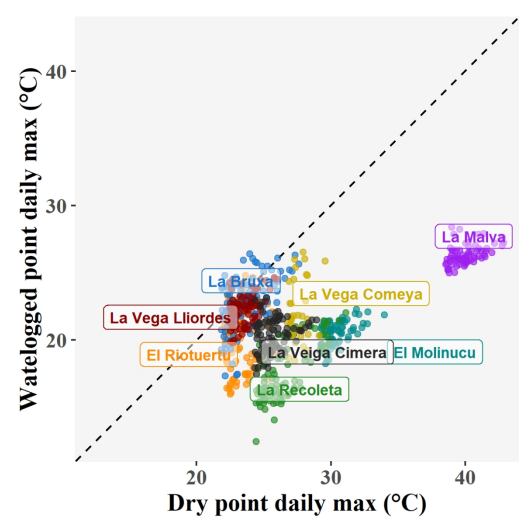


Figure 5: Complementary temporal analysis of the groundwater buffer effect during peak hot events. Variation of soil temperatures at paired sites (dry vs. waterlogged points) during exceptionally warm days of the warm season. Exceptionally warm days were considered those with a daily maximum (at the dry point) belonging to the 90th percentile of daily maximums for the site. The 1:1 line (corresponding to no buffer effect) is shown, points falling to the right of this line show a buffer effect (i.e. soil temperatures at the waterlogged point are cooler than the neighbouring dry site).

99x99mm (300 x 300 DPI)

# Mire microclimate: groundwater buffers temperature in waterlogged versus dry soils

Eduardo Fernández-Pascual<sup>1</sup>\*, Eva Correia-Álvarez<sup>2</sup>

- <sup>1</sup> Universidad de Oviedo, <sup>2</sup> Independent researcher
- \* Correspondence: Departamento de Biología de Organismos y Sistemas, Universidad de Oviedo, C/ Catedrático Rodrigo Uría, 33006 Oviedo/Uviéu, Spain. Email: efernandezpascual@gmail.com. Telephone: +34985104787.

### **Graphical abstract**



Mires are semi-terrestrial wetlands that remain waterlogged for most of the year. Mire groundwater produces a thermal buffer effect that insulates these habitats from the surrounding landscape, especially at the warm end of the climatic spectrum, i.e. during summer and at lower elevations. This highlights the potential refugial character of mires from global warming, and the need to integrate in situ microclimate measurements into climate change models.