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Water relations in the soil crust lichen *Psora decipiens* are optimized via anatomical variability

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1	Water relations in the soil crust lichen <i>Psora decipiens</i> are optimized via anatomical variability
2	C. COLESIE, L. WILLIAMS, B. BÜDEL

3 Abstract

4 Biological soil crusts are communities composed of cryptogamic organisms such as lichens, mosses, cyanobacteria and green algae that form a skin on soils in areas where vascular plants are excluded or 5 limited by water availability or temperature. The lichen Psora decipiens (Hedw.) Hoffm. is a 6 7 characteristic key organism in these communities in many different biomes. The species has a generalistic ecology and high morphological variation, which contributes to the ability of the species 8 9 to withstand environmental changes. We investigated whether different populations, based on site and 10 associated morpho-anatomical differences, incorporate functional water relations and how/whether 11 this was mediated by changes in abiotic factors. Samples were collected from two climatically distinct 12 sites, one "dry" site in southern Spain, and one "wet" site in the Austrian Alps. Our results showed 13 that samples from the dry site had a significantly thicker epinecral layer, higher specific thallus area, a 14 faster water uptake and contained more water per dry weight, all of which contributed to a much 15 slower drying rate. Both populations showed a highly adjusted water gain that incorporates functional 16 water relations and diffusion properties as a result of local water availability. We show eco-17 physiological and morphological mechanisms that underlie the high variability in P. decipiens and 18 draw conclusions around the ecological benefits for this generalistic lichen species such as optimized 19 water relations and light exploitation.

20

21 Keywords

Biological soil crusts, hydrological characteristics, epinecral layer, lichens, phenotypic plasticity,
 drying rate

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Introduction

27 The concept of ecotypes, defined as distinct genotypes (or populations) within a species, resulting 28 from adaptation to local environmental conditions (Hufford & Mazer 2003), describes the geographic 29 variation within a species and the balance between local adaptation and intra-specific hybridization 30 (Begon et al. 2006). In lichenology, to distinguish between differentially exposed populations, the 31 concept of ecotypes can be used, for example to describe different altitudinal distributions (Nadveina 32 et al. 2014). Another interpretation of structural changes in lichen thalli from different populations is 33 often described as phenotypic plasticity. For example, different rates of photosynthesis occur as a 34 result of differences in thallus hydration, due to structural changes, when comparisons are made 35 between north- and south facing populations of Ramalina capitata (Ach.) Nyl. (Pintado et al. 1997) 36 and vagrant compared with attached thalli of Cetraria aculeata (Schreber) Fr. (Pérez-Ortega et al. 37 2012). Increasing the amount of rhizinae is a modification that can improve thallus hydration and 38 therefore photosynthesis, when comparing epilithic versus epiphytic populations of Parmelia 39 pastillifera (Harm.) Hale (Tretiach & Brown 1995). Finally, Catillaria corymbosa (Hue) I. M. Lamb, 40 an Antarctic endemic species, showed an increased water retention capacity and therefore more 41 photosynthesis in shaded localities (Sojo et al. 1997).

42 In drylands, vascular plants are limited due to low water availability and therefore biological soil 43 crusts (BSC) become a dominant component of vegetation (Pointing & Belnap 2012; Büdel et al. 44 2014). BSCs also occur in high alpine areas above the tree-line, typically when the mean temperature 45 of the warmest month is below 10°C (Körner, 1998), seventy to eighty percent of the precipitation 46 falls as snow, and snow cover lasts 270 to 300 d (Auer et al., 2002). BSCs are small scale 47 poikilohydric lichen-bryophyte and microbial communities that are considered to be ecosystem 48 engineers (Pointing & Belnap, 2012) that stabilize soil (Belnap et al. 2003), and make significant 49 contributions to the carbon and nitrogen fixation budgets (Elbert et al. 2012). Lichens often form the 50 major proportion at the climax stage of these communities (Büdel et al. 2009). Studies on soil lichen 51 populations, including their functional aspects and morphological differences, are need to enable us to

52 predict climate change scenarios in these areas; these habitats are considered to be at high risk in 53 recent climate change projections (IPCC 2012).

54 Here we studied the differences in morpho- anatomical and hydrological traits of two populations of 55 Psora decipiens (Hedw.) Hoffm., a typical lichen occurring in the climax stage of biological soil crusts 56 worldwide (Büdel 2003, Galun & Garty 2003, Rosentreter & Belnap 2003. Two contrasting areas 57 with distinct populations were selected: a high alpine environment ("Site Hochtor", Fig. 1A) and arid 58 badlands ("Site Almeria", Fig. 1B). Preliminary observations indicated (1) strong variance in overall 59 thallus structure and appearance and (2) higher vitality of the alpine population (based on mean thallus 60 size and vivid color). Based on this field observation we hypothesized, that local water availability is a 61 major driver for morpho-anatomical differences in lichens and thallus water relations can be optimized 62 to local necessities.

63

Material and Methods

64 Study sites, organisms and sampling

We chose two sampling sites that support natural BSC occurrence, but differ considerably in their environmental conditions. The first, and humid "Site Hochtor", was a high alpine area at an elevation of 2500 to 2600 m asl. of the Großglockner massif, Austria. The second site was the dry site, "Site Almeria", located in South East Spain. This location was considered as one of the driest and sunniest in Europe, and has been described in previous studies (Lázaro *et al.* 2001, Cantón *et al.* 2004).

Psora decipiens is a pale pink squamulose-crustose lichen with conspicuous white-pruinose, often upturned margins. The lower cortex is white, poorly developed or absent and the attached hyphal net penetrates into the substrate. It is a cosmopolitan lichen and is well known as indicator species for intact BSC climax stages (Fig. 1C and D).

Samples (30) were randomly collected at each site in areas with homogenous BSC cover and no shading from plants. Each sample unit was a 9.2 cm² section of an intact *P. decipiens* dominated BSC, where several thalli were growing. Sampling at site Hochtor took place in August 2013 and at site Almeria in June 2013. Samples were dried at room temperature within 3 d of collection, transported and stored frozen. In the laboratory the surrounding soil crust was removed from the thalli, which were

then washed. The number of replicates used in each experiment is indicated below.

80 Climate measurements

81 Both investigation sites were equipped with similar climate stations, capable of monitoring air 82 temperature and humidity, solar radiation (Photosynthetically Active Radiation, PAR), $UV_{(A+B)}$ -83 radiation and precipitation. Climate data were recorded from both sites over 2 y, from April 2012 -84 March 2014 at site Almeria, and from August 2012 - July 2014 at site Hochtor. Air temperature and 85 relative humidity were measured 1.5 m above ground, and solar radiation at 2 m above ground. The 86 snow cover was estimated from the length of time the various sensors were inactive. Mean (± standard 87 deviation) values were calculated for summer and winter air temperature, humidity, PAR and UV_A and UV_B combined radiation. Additionally maximum and minimum temperature, and maximum UV and 88 89 PAR radiation were extracted from the dataset.

90 Thallus morphology and anatomy

91 *Thickness.* The thicknesses of the lichen thalli, the epinecral layer, the photobiont layer and the 92 medulla were measured on freezing microtome sections, n= 400 for the photobiont layer and medulla 93 measurements and n= 100 for the epinecral layer measurements from each site, using the AxioVision 94 software (Carl Zeiss, Jena, Germany). To visualize thallus internal relationships between photo- and 95 mycobiont, the ratio between the photobiont layer and medulla was calculated.

96 Specific thallus area. To calculate the specific thallus area (STA, mm² mg⁻¹) thallus size was first 97 determined by binocular microscopy using the above mentioned software. A standard procedure was 98 used to delineate the extent of each lichen thallus. The lichen thalli were wetted to ensure maximum 99 surface area, placed on scale paper and photographed. The corresponding dry masses (DM) of these 100 thalli were determined by weighing after 3 d of oven drying at 60°C (n=20).

Specific thallus mass. This parameter was emphasized by Kershaw (1985) as an important lichen
 parameter for water loss and water uptake and is the inverse equivalent of STA (see above).

103 Hydrological traits

Drying rate. Photosynthetic activity was monitored by measuring the efficiency of PSII 104 105 photochemistry using an imaging-PAM chlorophyll fluorometer (Heinz Walz, GmbH, Effeltrich, 106 Germany). Specimens of roughly the same size (\pm 10 mm²), which had been submerged in water 107 overnight, were first weighed, then placed in a sealed plastic chamber on a wire net over a saturated 108 NaCl solution which maintained a level of humidity of 75% rH equivalent to a water potential of -37 109 MPa at room temperature (Pardow & Lakatos 2013). Initially the maximum quantum yield of PSII 110 F_v/F_m of the hydrated and dark adapted (30 min) samples was measured as a reference. Subsequently, 111 short saturation pulses were applied every 2 minutes to determine the fluorescence parameters for 112 calculating PSII yield (Y = Fv/Fm). The time was measured until a threshold of 0.2 PSII yield was 113 reached for n=18 replicates (3 thalli per chamber). To determine the absolute water loss during the 114 measurement, samples were weighed again, directly after the measurement. Drying rate was 115 calculated, and expressed as the time need for 1 μ l of water to evaporate from one mm² of thallus $(\min \mu l^{-1} mm^{-1}).$ 116

117 *Water uptake*. To ensure full water saturation prior to weighing, the samples were submerged in 118 distilled water for 30 min. Excessive water and droplets were carefully shaken off before measurement 119 of maximum wet mass (WM_{max}). The corresponding dry mass (DM) of these thalli was determined by 120 weighing after 3 d at 60°C. The maximum water uptake relative to the thallus specific dry mass of the 121 samples (n=36) was calculated as $WM_{max} - DM/DM$ (Pérez 1997).

Repellency. To measure water repellency/hydrophobicity of individual lichen thalli, the water drop penetration time (WDPT) was measured for n= 30 replicates. The WDPT test consists of placing a drop of water on the surface of the epinecral layer and measuring the time until complete absorption occurs. This is a commonly used test because of its simplicity (Letey *et al.* 2000) and the value of information it provides, as it was considered to be the most indicative and sensitive way for the hydrological consequences of water repellency to be investigated (Doerr 1998, Leelamanie *et al.* 2008).

129 Optimum water content. CO₂ gas exchange measurements were conducted under controlled laboratory 130 conditions using a portable mini cuvette system (GFS 3000, Walz Company, Effeltrich, Germany). 131 The response of net photosynthesis (NP) and dark respiration (DR) to thallus water content (WC) was 132 determined for three replicates (each replicate was composed of about 20 individual squamules) from 133 each of the sites. Complete drying-out cycles (from water saturated to air dry thalli) were measured at 134 750 μ mol photons m⁻² s⁻¹ (saturating light), ambient CO₂, at 17°C (which is within the optimal 135 temperature range for CO₂-gas exchange of this species). Samples were weighed between each 136 measurement and thallus water content (WC) was later calculated as a percentage of dry mass. Dry 137 mass was determined after five days in a desiccator over silica gel. Ninety percent of maximum net 138 photosynthesis was considered to be a reasonable estimate for optimal water content. 139 *Water holding capacity*. Water-holding capacity (WHC) was calculated by: saturated wet mass – dry 140 mass (mg) / thallus area (cm⁻²) after shaking surplus water off the lichen thallus. The corresponding

141 dry mass (DM) of these thalli (n=20) were determined by weighing after drying for 3 d at 60°C.

142 Thallus area was measured as described above.

143 Statistics

To determine differences between the lichens anatomical features and the climate of the two sites student's t-tests were used (Statistica 10, Stat soft). All data was normally distributed. Significance level was defined at P < 0.05.

147

Results

148 Climate measurements

The data (Table 1) demonstrate the pronounced differences in temperature, humidity and precipitation between the sites, and also within the sites based on seasonality. In Almeria it was significantly warmer, drier and brighter than at the alpine site Hochtor. However in site Hochtor the growing season only lasted about 3 months in a year, due to the persistent snow cover.

153 Thallus morphology and anatomy

154 Thalli from Almeria had thicker epineeral layers, but only half the medulla thickness compared to samples from Hochtor (Fig. 1E and F; Table 2). The photobiont layer showed no difference between 155 156 the sites, therefore, the photobiont layer to the medulla-ratio of the thalli from site Almeria was twice 157 as high. This indicates significant differences in the internal structure of the thalli between the 158 populations. STA was also higher for samples from site Almeria (Table 2), indicating that these thalli 159 can support relatively large areas despite their reduced thallus matter. Corresponding to this finding, 160 STM was significantly higher for samples from site Hochtor, indicating that these samples are thicker 161 than those from site Almeria.

162 Hydrological traits

163 Both populations had similar water holding capacities per area (Table 2) and showed some common 164 drying characteristics, independent of sampling site and thallus size (Fig. 2). For example, the initial 165 activity of PS II was similar, with high yields of PSII at ca. 0.6 (blue color, Fig. 2). Additionally, these 166 levels of activity remained relatively stable for certain amounts of time independent of thallus size 167 (around 0.6). As soon as a threshold of desiccation was reached, the actual drying event occurred 168 quickly, and no fluorescence signal could be detected shortly after this. In contrast to these common 169 features, the actual drying rate was different between the two populations. Thalli from site Almeria 170 dried six-times slower than those from site Hochtor (Table 3, Fig. 2). Additionally, P. decipiens 171 specimens from site Almeria took up water faster (1.2 ± 0.27 sec compared to 168.4 ± 15.8 sec for 172 samples from site Hochtor) and contained more water per dry mass than those from site Hochtor 173 (Table 2). The optimum WC for CO_2 exchange extended over a significantly narrower range, (109-156 174 % DM) for samples from site Almeria than for samples from site Hochtor (131-195 % DM).

175

Discussion

In the present study we have demonstrated distinct differences between two populations of the lichen species *P. decipiens,* derived from climatically contrasting habitats. Morpho-anatomical differences appear to be reflected in differences in functional water relations and diffusion properties as a result of local water availability. We have identified eco-physiological and morphological mechanisms that underlie the high variability in *P. decipiens* that determine its ecological fitness in a particular habitat.

181 The most striking difference between the two populations is water gain. Considering the differing STA 182 and the maximum water uptake relative to the thallus specific dry mass, thalli from both sites have the 183 same water holding capacity on an area basis (WHC in Table 2). This indicates that, not only surface 184 area, but also thallus internal structure must be studied to understanding the drying processes in 185 lichens. During the actual drying process we need to consider the amount of water in the thallus, based 186 on thallus area. The resulting calculation shows that drying from optimal water content to completely desiccated lichen thalli takes 2.5 minutes mm⁻² for thalli from Hochtor, but 15 minutes mm⁻² for 187 Almeria thalli. This conclusion is supported by the chlorophyll florescence results, which show a 188 189 much slower rate of decrease in activity in Almeria thalli (Fig. 2).

190 Samples from the dry site Almeria show both improved water uptake and reduced water loss. A 191 reduction of water loss in lichens seems remarkable, because lichens, as poikilohydric organisms are 192 known for their passive water control and unregulated loss of water over the whole thallus surface. 193 Nevertheless, Beckett (1995) showed that lichens from dry habitats seem to make better use of their 194 water, by maintaining turgor down to low relative water contents. The reduced rates of water loss 195 found here, may be the result of increased diffusive resistance caused by the thicker epineeral layer. 196 This layer is often described as amorphous, and is composed of decomposing hyphae with indistinct 197 cell lumina; it forms a very dense layer that can act as a barrier to water loss, more so than the upper 198 cortex. In the revised generic concept of Parmelioid lichens (Crespo et al. 2010) a pored epicortex is 199 an important diagnostic feature beside molecular, morphological and chemical evidence. The 200 epicortex proabably provides an extra layer of protection against evaporation. A thick epineeral layer, 201 as present in the samples from the dry site Almeria may therefore account for such a reduction in 202 water loss. Nevertheless, developing a thick epinecral layer also seems contradictory to improved 203 water gain, because it is also reported to have hydrophobic properties (Lakatos et al. 2006). The 204 occurrence of hydrophobins, proteins unique to mycelial fungi, has been suggested to be important for 205 the survival of lichens (Wessels 2000). According to Honegger (1991), a hydrophobic lining of gas 206 spaces allows efficient apoplastic transport of water and solutes between the symbionts, and permits 207 optimal gas exchange during wet periods. Therefore, very low water repellency of the samples from 208 site Almeria is unsurprising. The suggested explanation for this result derives from the structure of the

209 epinecral laver itself and how this influences the lichen surface (Fig. 1C and D). In dried thalli, the 210 epinecral layer has open cracks (Fig. 1E), thus increasing the surface area and facilitating water uptake 211 by cohesion and adhesion, leading liquid water towards the photobiont layer (Fig. 1E and F). With WHC_{shaking} being close to 50 mg H₂O cm⁻² (Table 2, 10 mg H₂O cm⁻² equates to 0.1 mm dew or rain) 212 213 these lichens use the rare rain evens more often than dewfall (Gauslaa et al. 2014). Both findings 214 support the explanation that a functional role of the epineeral layer is to influence the lichen 215 hydrology. To our knowledge, this has not been previously described. The function of the epineeral 216 layer is usually suggested to be protection against high light stress (Büdel & Lange 1994, Büdel et al. 217 1997, Rikkinen 1995, Kappen et al. 1998, Dietz et al. 2000). For chlorolichens, drying combined with 218 light exposure can be particularly harmful (Gauslaa et al. 2012) and it was suggested that the ability to 219 recover, correlates positively with increasing species-specific water holding capacities (WHC). In 220 Almeria, light intensities and UV-radiation are much higher throughout the year (Table 1), and in this 221 population the thicker epineeral layer is therefore necessary to prevent light damage. In the Hochtor 222 site, where light intensity is less and exposure time to both PAR and UV are shortened due to snow 223 cover, such protection would not be required or beneficial to the lichen.

224 In the wet site of Hochtor, the lichens experience a different set of stresses. Water saturation for many 225 hours a day may result in negative carbon gain for two reasons. First high respiration rates during the 226 night and under the snow cover can influence carbon balance negatively, and second CO₂ diffusion 227 resistance is high in water supra-saturated thalli, thus reducing the substrate for photosynthesis 228 (Cowan et al. 1992). The high WHC_{shaking} values indicate that these lichens are more often exposed to 229 rain events of about 40 mm precipitation (Gauslaa et al. 2014), which underlines their frequent water 230 saturation. For lichens, it is essential to minimize periods of water supra-saturation. Indeed, the 231 functional aspects that we report here for the lichens from the wet site Hochtor are fast desiccation 232 times, high hydrophobicity, low maximum water uptake relative to the thallus specific dry weight and 233 a broad range of thallus water contents for optimal photosynthesis. All of these may be explained by 234 specific thallus anatomy, which includes a very thick medulla layer and is best quantified by the high 235 STM values (Table 2). The medulla is the fungal zone in the lichen thallus, composed of hyphae, with 236 cell walls often incrusted with crystalline secondary metabolites. It was shown that lichen substances

237 did not maintain the water-free diffusion pathways (Lange et al. 1997) and the authors suggested that these pathways are rather maintained by structural changes. Together with numerous hydrophobic air 238 239 spaces in the medulla, supra-saturation with water is minimized or even avoided (Lange et al. 1993). 240 The morpho-anatomical adjustments leading to water repellency of the upper layers involve reduced 241 STA for the samples from site Hochtor. The parameter of STA is analogous to specific leaf area 242 (SLA), in higher plants, which can yield information about life strategies. Species with low SLA 243 conserve acquired resources, due to their large dry matter content, high concentration of cell walls and 244 secondary metabolites, and high leaf and root longevity (Marron et al. 2003). By applying these 245 features to lichens, the lower the STA, the lower the fitness and the more lichen material is needed to 246 support the same surface area. Our results show higher STA values for lichens from the dry site 247 Almeria, which means that these lichens have a higher fitness. One conclusion from this result might 248 be that anatomical and functional adjustments within P. decipiens are more easily made towards 249 dryness and high light stresses than towards water supra-saturation, a common phenomenon in the wet 250 site Hochtor. This conclusion may be of general interest towards regions with climate change 251 predictions that include increased flood and heavy rain risks (IPCC 2012), as the conditions are 252 expected to occur in the future in many areas that sustain natural BSCs covers. Accumulations of such 253 events may influence the natural BSC more severely than increasing drought.

254 This study suggests that two populations show variations in morpho-anatomical traits that result from 255 their native environments climatic differences. These differences could result from ecotypic variation 256 or phenotypic plasticity. Increasingly, recent studies on plant plasticity describe not only growth rates 257 documentation and morphological parameters, but also functional aspects of plasticity. The plasticity 258 of functional traits (both long- and short term) can contribute to the ability of species to occupy 259 diverse and variable habitats in nature (Sultan et al. 1998). Phenotypic plasticity plays an important 260 role in community ecology because it contributes to the ability of species to withstand environmental 261 changes, such as those caused by human disturbance. The timescale of such changes is often too short 262 for an evolutionary response, thus species that lack sufficient plasticity might be at risk of altered 263 reproduction, degradation or extinction (Sultan 2000). On the other hand, ecotypic variation would 264 result in the two populations also being different on a genetic level. This would suggest that the

observed differences are not due to the species plasticity and could therefore be at risk to climate change and habitat loss. This study has focused on purely morphological and physiological characteristics of the lichen *P. decipiens*. In order to reveal a complete picture of the variation of this important soil crust lichen molecular analysis has to be included. The next step is to investigate the genetic diversity and acclimation potential of both the algal and fungal partners of *P. decipiens* from the two populations at the climatically distinct sites.

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- **392** Figure captions:

Figure 1: Characteristics of *Psora decipiens*. A) Study site Almeria overview, B) study site Hochtor overview; C) natural appearance at study site Almeria (photo: Martin Westberg); D) natural appearance at the study site Hochtor; E) cross section from site Almeria with the thick cracked (arrows) epinecral layer; F) cross section from site Hochtor with a continuous epinecral layer and a thick medulla including a thick photobiont layer. White scale bars indicate 1 cm, black bars indicate 20µm.

- Figure 2: Dehydration kinetics of *Psora decipiens*. False color chlorophyll fluorescence images of the
- 400 effective quantum yield (Y) of photosystem II distribution over three thalli from site Almeria (upper

401 row) and site Hochtor (lower row), obtained using an Imaging PAM (Walz GmbH, Effeltrich, 402 Germany). Yield intensity is color coded covering a range from 1-0 with red indicating very high <text> 403 values and violet low values. Red flags indicate exact Yield values at a chosen area of interest in the 404

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Characteristics of Psora decipiens. A) Study site Almeria overview, B) study site Hochtor overview; C) natural appearance at study site Almeria (photo: Martin Westberg); D) natural appearance at the study site Hochtor; E) cross section from site Almeria with the thick cracked (arrows) epinecral layer; F) cross section from site Hochtor with a continuous epinecral layer and a thick medulla including a thick photobiont layer. White scale bars indicate 1 cm, black bars indicate 20µm.

188x211mm (300 x 300 DPI)



Dehydration kinetics of Psora decipiens. False color chlorophyll fluorescence images of the effective quantum yield (Y) of photosystem II distribution over three thalli from site Almeria (upper row) and site Hochtor (lower row), obtained using an Imaging PAM (Walz GmbH, Effeltrich, Germany). Yield intensity is color coded covering a range from 1-0 with red indicating very high values and violet low values. Red flags indicate exact Yield values at a chosen area of interest in the picture. Pictures where taken every 2 minutes.

297x67mm (300 x 300 DPI)

Table 1: Climate data at the study sites. Climate data are differentiated by summer and winter for both investigation sites. Summer: April-September, Winter: October-March, Almeria: April 2012-March 2014, Hochtor: August 2012-July 2014. Shown are mean values with standard deviation where appropriate, maximum and minimum values recorded across the measuring period, average rainfall per season and snow cover duration. PAR and UV are based on daily average and maximum values. a: values are significantly different within the sites (summer vs. winter), b: values are significantly different summer vs. site Hochtor summer).

		Site Almeria		Site Hochtor	
Parameter		Summer	Winter	Summer	Winter
Air temp. (°C)	Average	23.0 ± 6.6^{ab}	13.6 ± 5.8^{ab}	2.0 ± 4.6^{ab}	$\textbf{-3.7}\pm3.0^{ab}$
	Max	43.8	34.8	19.86	14.6
	Min	2.8	0.0	-7.5	-18.6
Humidity. (%)	Average	51.4 ± 20.5^{b}	60.6 ± 19.7^{b}	92.4 ± 12.3^{b}	93.1 ± 9.5^{b}
PAR (µmol m ⁻² s ⁻¹) Average		962.5 ^{ab}	619.8 ^{ab}	441.8 ^{ab}	152.8 ^{ab}
	Max	2650	2406	2680	1862
UV_{A+B} (µmol m ⁻² s ⁻¹) Average		89.69 ^{ab}	60.6 ^{ab}	77.8 ^{ab}	33.8 ^{ab}
	Max	346.9	266.8	384.2	244.6
Rain (mm)	Average	25.5 ^{ab}	91.8 ^{ab}	558.3 ^{ab}	75.45 ^{ab}
Snow cover Year		None	None	3 months	6 months

Table 2: Comparison of anatomical and hydrological characteristics of lichen thalli deriving from the wet exposed site Hochtor and the dry site Almeria. Shown are mean values with standard deviation and significance levels from t-tests.

Thallus morphology			Significance of and we	Significance of difference between dry and wet exposed thalli		
parameter	Site Almeria	Site Hochtor	t	df	р	
Epinecral layer thickness (µm)	92.2 ± 18.8	70.1 ± 9.8	7.3	98	0.000	
Photobiont layer thickness (µm)	91.5 ± 16.2	93.0 ± 20.2	-0.7	398	0.4	
Medulla thickness (µm)	102.4 ± 38.6	224.6 ± 59.4	-24.4	398	0.000	
Photobiont/ Medulla ratio	0.89	0.41				
Specific thallus area (mm ⁻² mg)	4.7 ± 0.9	2.9 ± 0.3	5.6	19	0.005	
Specific thallus mass (mg cm ⁻²)	22.3 ± 3.8	36.7 ±7.6	5.6	19	0.005	
	Thallus hydrolo	ogy				
Maximum water uptake relative to the thallus specific dry weight (mg H ₂ O/mg dry weight)	2.1 ± 0.7	1.4 ± 0.1	-4.1	34	0.000	
WC opt (%)	109.7 -156.1	131.8 - 195.4	-7	2	0.03	
WHC (mg H ₂ 0 cm ⁻²)	46.4 ± 12.8	51.6 ± 9.9	-2.5	19	0.1	
WDPT (sec)	1.2 ± 0.3	168.4 ± 15.8	-4.1	28	0.000	
Drying rate $(\min \mu l^{-1} mm^{-2})$	34.1 ± 28.8	5.3 ± 4.3	-4.2	34	0.000	
	I					