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Taylor, Emily S.; Levy, Peter E.; Gray, Alan. 2017. **The recovery of Sphagnum capillifolium following exposure to temperatures of simulated moorland fires: a glasshouse experiment.** *Plant Ecology & Diversity*, 10 (1). 77-88.<https://doi.org/10.1080/17550874.2017.1302017>

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This is an Accepted Manuscript of an article published by Taylor & Francis Group in Plant Ecology & Diversity on 04/04/2017, available online: <https://doi.org/10.1080/20423489.2017.1408190>

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9 **Abstract**

10 *Background*. In the UK, government legislation allows the use of prescribed fire in peatlands for 11 land management purposes. The use of fire, however, remains controversial, partly because of a 12 distinct lack of data on the response of key peatland species to fire. *Sphagnum* species are key 13 components of peatland ecosystems, yet a fundamental knowledge gap in the debate is the 14 response of *Sphagnum* species to fire. *Aims*. To determine if a widespread species (Sphagnum 15 capillifolium) has the ability to recover from exposure to high temperatures, analogous to those 16 recorded in managed peatland fires. *Methods*. Samples of *S. capillifolium* were exposed to a range 17 of temperature treatments. Recovery was monitored using chlorophyll fluorescence, $CO₂$ exchange 18 and physical damage and new growth assessed. *Results*. We found that the degree of recovery of *S.* 19 *capillifolium* was related to the temperature treatment, post-treatment environmental conditions 20 and pre-treatment stem moisture content. The slowest recovery was found when samples were heated to 400 °C for 30 seconds. *Conclusions*. Our results demonstrate that *S. capillifolium* has the 22 ability to recover following exposure to the temperatures experienced in prescribed fire, provided 23 that at least some living material remains. Our results suggest that prescribed burning in the spring 24 may allow for a quicker recovery than autumnal fires.

25

26 **Introduction**

27 Prescribed fire is a key management tool used on peatlands in Britain, to promote the 28 regrowth of *Calluna vulgaris* (L.) Hull (*Calluna* hereafter) and grasses for grazing by game 29 birds and livestock. There is considerable debate over the impact of fire on peatlands, and in 30 particular, on the impacts on species of ecological and conservation importance, such as 31 *Sphagnum* species. The debate surrounding prescribed fire remains contentious, partly due to 32 the polarised views of the protagonists, but also because of the lack of evidence for the 33 effects of fire on taxa including *Sphagnum* (Davies et al. 2016). Much of the data comes

34 from studies on wildfires, which may be much more severe (sensu Keeley 2009) than 35 prescribed fires, and result in greater depth of burn and exposure of bare peat (Benscoter 36 2006; Maltby et al. 1990). The effects of prescribed fires may be qualitatively quite 37 different, and it was this that we aimed to investigate here. *Sphagnum* mosses are key peat-38 forming species and store large quantities of carbon (Rydin and Jeglum 2006). Their 39 capacity for holding water and locking up nutrients, together with their recalcitrant litter, 40 allows them to survive in, and maintain, the nutrient-poor and acidic peatland environment 41 (Clymo and Hayward 1982; Jones et al. 1994; Kuhry et al. 1993; Rydin and Jeglum 2006). 42 As key components of peatlands, understanding the response of the *Sphagnum* species to 43 land management and the environment is of fundamental importance to peatland 44 conservation.

45 In England, burning on blanket bog is only allowed as part of a pre-approved plan for 46 conservation and restoration in a defined season (Anon 2007), and Wales has a similar set of 47 regulations (Anon 2008). In Scotland, burning can only be legally carried out during a 48 defined season, and only where *Calluna* constitutes more than 75% of the vegetation cover 49 (Anon 2011); these guidelines are currently under review. Understanding the response of 50 *Sphagnum* is a crucial aspect of these guidelines and needs to be based on evidence if the 51 debate on the use of fire is to progress (see Davies et al. 2016). To date, little research has 52 looked at the direct effect of fire on the *Sphagnum*. Observations suggest that the impact on 53 the *Sphagnum* may depend on vegetation and environmental characteristics that influence 54 temperature at the moss surface and the penetration depth and duration of high temperatures.

The typical adiabatic flame temperature of wood burning in air is 1980 \degree C (Griffiths and

56 Barnard 1995), so very high temperatures can be reached in the vegetation canopy during the

57 passage of a fire. At the moss surface, temperatures can reach up to 600° C for relatively

58 short periods (<30 seconds) (Davies 2005; Hamilton 2000), but typically, the moss layer is

59 not exposed to such high temperatures (see Hobbs and Gimingham 1984). The limited 60 available data suggest that the temperature at 2 cm below the moss surface rarely exceeds 50 $^{\circ}$ C (Davies 2005), although, Harris et al. (2011) recorded maximum temperatures of 62 approximately 600 °C 1 cm above the ground level in *Calluna* moorland. High surface 63 temperatures can potentially affect *Sphagnum* growth through cell damage in the uppermost 64 capitulum, the site of the majority of photosynthetic activity (Rydin and Jeglum 2006). 65 However, fire may damage only the upper sections of stems, allowing re-growth from side 66 shoots (Rydin and Jeglum 2006). This has been observed in the field in at least some 67 circumstances (c.f. Clymo and Duckett 1986; Hamilton 2000).

68 The depth and duration of high temperatures will depend on the amount, composition and 69 distribution of fuel above the moss layer, the moisture content and bulk density of vegetation 70 and moss layer, and meteorological conditions (e.g. see Harris et al. 2011; Santana and Marrs 71 2014, 2016). A high density of above-ground fuel will prolong the residence time of the 72 fire, causing greater heating and evaporation, and may allow the fire to penetrate the peat 73 (Ashton et al. 2007; Davies et al. 2013). 'Hot spots' have been observed in *Calluna* fires in 74 the moss layer immediately around the woody stems of *Calluna* (Davies 2005; Hamilton 75 2000). Conversely, high *Sphagnum* moisture contents could result in reduced temperatures 76 and depth of penetration, as thermal energy would be dissipated by evaporation. A wetter 77 *Sphagnum* layer may also have quicker recovery following the drying effect of the fire. 78 However, given that fuel loads and moisture contents differ considerably among (and within) 79 fires (Legg et al. 2010), the thermal impact of fire on *Sphagnum* will also vary. Here, we 80 used a representative range of maximum temperatures, fire residence times and moisture 81 contents to assess the effects of high temperature on photosynthesis and recovery in 82 *Sphagnum capillifolium* (Ehrh.) Hedw *sensu lato*.

89 **Materials and methods**

90 *Experimental design*

91 *Sphagnum capillifolium* was exposed experimentally to a high temperature over a short 92 duration, reproducing the temperature dynamics in the range recorded in *Calluna* fires 93 (Davies 2005). *S. capillifolium* was collected from Whim Moss, Penicuik, south-east 94 Scotland (NT203532), an ombrotrophic blanket bog classified as M19 *Calluna vulgaris*-95 *Eriophorum vaginatum* National Vegetation Classification (NVC, Rodwell 1991) blanket 96 mire. The site lies at 280 m a.s.l., with mean temperature of 8.6 \degree C (Sheppard et al. 2013). 97 Clumps of *S. capillifolium* (6 cm deep, 5 cm diameter) were collected a maximum of two 98 days prior to the start of each run of the experiment from four separate hummocks (so enough 99 material could be gathered), and placed into 5-cm diameter fibre pots (Grow It, Spalding, 100 UK). Clumps were kept as intact as possible, ensuring the number of stems in each pot was 101 representative of natural stem densities; the number of stems per pot varied between 42 and 102 83. For each run, 96 pots of *S. capillifolium* were placed within a tray containing a bed of *S.* 103 *capillifolium* cuttings, to help maintain near natural moisture conditions with regularly 104 watering in a glasshouse that was programmed to track external air temperatures $(+/- 2 \degree C)$. 105 The tray was divided into four blocks so any variation in conditions across the tray could be 106 reflected in the statistical models. Each pot was individually watered with distilled water, 107 using a syringe, to maintain *S. capillifolium* moisture content to around 90% (moisture 108 content is expressed throughout as mass of water / initial fresh mass of moss x 100, as this 109 could be measured gravimetrically non-destructively over the whole course of the 110 experiment; initially, the moss was near to saturation). A pilot study ensured that *S.* 111 *capillifolium* samples could remain healthy under these conditions as indicated by 112 chlorophyll florescence (see Taylor 2015).

113 The experiment was run three times in spring, autumn and winter, with one of three different 114 temperature treatments (see below) randomly assigned to each of the 96 pots per run (Table 115 1). Pots were randomly assigned to one of four measurement procedures: chlorophyll 116 fluorescence, CO2 exchange, growth measurements or moisture content analysis, as both the 117 chlorophyll fluorescence and moisture content analysis were destructive. Although the winter 118 experiment in February 2013 was initiated only a month earlier in the seasonal cycle 119 compared to the spring experiment (in March 2012), the prevailing weather conditions were 120 quite different. In the two weeks prior to the winter experiment, the locally-measured air 121 temperature averaged 1.4 \degree C, and was below freezing for much of the time. In the two weeks 122 prior to the spring experiment, the air temperature averaged 7.4 °C , and the plants were 123 physiologically active. Hence, we think these experiments approximate the typically 124 contrasting conditions in these seasons, even though the timing in terms of the seasonal cycle 125 was not large.

126 The temperature treatments were carried out by placing each pot in a perforated steel 127 chamber, heated from above by a butane-propane gas burner (Parasene Weed Wand 550, 128 Parasene, UK). The flame was held in place for the desired length of time once the surface of 129 the pot reached the desired maximum temperature. The perforated steel chamber diffused the 130 direct heat from the flame, so that temperatures could be better controlled at the moss 131 surface. Temperature was logged (CR21X, Campbell Scientific, Utah, USA) every 2 s using 132 k-type twisted thermocouples at, 2-cm and 5-cm depth. The temperature treatments were: 100 (100 °C for 3 s); 400 (400 °C for 3 s); 400+ (400 °C for 30 s); and 400+D (400 °C for 30s 134 and air drying the *Sphagnum* for three days prior to treatment). Control pots were treated the 135 same except that no heating was applied. For post-treatment recovery, the pots were 136 maintained at stem moisture content of around 90%. The mean, minimum and maximum 137 values of air temperature and photosynthetic photon flux density (PPFD) were also measured.

138 *Chlorophyll fluorescence*

139 Chlorophyll fluorescence was used to measure plant stress (Krause and Weis 1991; Maxwell 140 and Johnson 2000), based on previous evaluations on *Sphagnum* (e.g. Hájek and Beckett 141 2008; Manninen et al. 2011; van Gaalen et al. 2007). The technique works on the principle 142 that the ratio between variable florescence (F_v) and maximal florescence (F_m) approximates 143 the maximum quantum yield of PSII, ranging between 0.75 and 0.84 in healthy mosses (e.g. 144 Bates et al. 2013; Green et al. 1998; Hájek and Beckett 2008; Manninen et al. 2011; Proctor 145 2003; van Gaalen et al. 2007), with lower values indicating stress (Maxwell and Johnson 146 2000).

147 Chlorophyll fluorescence measurements were made using a Continuous Excitation

148 Chlorophyll Fluorimeter (HandyPEA, Hansatech Instruments Ltd, UK) on the capitulum of

149 one stem from each pot on 8 days beginning on the first day of temperature exposure and up

150 to 100 days after exposure. Each capitulum was dark-adapted for 20 min, prior to

151 measurements at a PPFD of 1500 μ mol m⁻² s⁻¹. During the autumn run chlorophyll

152 fluorescence was assessed 100 days after treatment at 5-mm intervals down the stem.

153 *CO2 exchange*

154 Gas exchange measurements were made five times on eight pots per treatment from day 3 to 155 99, using an infra-red gas analyser (LI-6400XT, Li-Cor, Lincoln, NE, USA) in an open gas 156 exchange system, with a sample chamber designed to measure whole pots of *Sphagnum*. 157 Each pot of *Sphagnum* was carefully transferred into an inert plastic pot of the same size as 158 the gas exchange measurements. Air from the sample chamber was circulated through a 159 column of silica gel to remove excess water vapour, such that the humidity of incoming and 160 outgoing air from the sample chamber was similar. The Licor LI-6400XT was set to control 161 the system flow rate (500 µmol air s⁻¹), chamber air temperature (20 °C), incoming CO₂

162 concentration (400 μmol mol⁻¹), and PPFD (0 or 2000 μmol photons m² s⁻¹ using a 6400-18 163 RGB light source, Licor, Lincoln, NE, USA). CO2 concentrations were logged at 10 164 sintervals and averaged once stable, typically over 5 to 10 min. Because of the difficulties in 165 quantifying leaf area, photosynthesis and respiration were expressed as μ mol CO₂ (g dry 166 mass¹ s⁻¹, correcting all CO₂ mixing ratios to a dry air basis. The dry mass of *Sphagnum* 167 was calculated by oven-drying samples at the end of each run of the experiment (day 100) at 168 -70 °C for 5 days before weighing. No respiration measurements were made during the spring 169 run and measurements were made on only three occasions during the first half of the autumn 170 run, as pots were infected with mould which covered some or all of the surface of the 171 *Sphagnum*. No other runs of the experiment were affected by mould.

172

173 *Physical damage and new growth*

174 The depth of physical damage and bleaching (loss of pigment) was measured down the stem 175 from the capitulum. Reduced structural integrity was estimated by gently running a finger 176 across the surface of the pot showing breakage in brittle stems from pots assigned to the 177 whole pot gas exchange and new growth.

178 The number and dry mass of new auxiliary stems was measured at the end of each run of the 179 experiment and the length of the new stems was measured in a subset of samples. New stems 180 oven-dried at 70 °C for 5 days, weighed and new growth calculated as the ratio of dry 181 biomass (new growth plus original sample) to the original dry biomass to take into account 182 the difference in the number of stems between pots.

183 *Statistical analyses*

184 Linear mixed-effects models were used for analysing chlorophyll fluorescence,

185 photosynthesis and respiration data, accounting for the repeated measurement design. Initial

- 186 models were composed of all fixed and random effect terms (Table 2). In subsequent models,
- 187 non-significant fixed effect terms were dropped one by one (using AIC) to derive a model
- 188 with the smallest AIC that consisted only significant fixed effect terms, as indicated by Wald
- 189 tests. Statistics were carried out using R (v R i386 3.0.1) (R Core Team 2013) with mixed
- 190 effects modelling computed using the package lme4 (Bates et al. 2009).

191 --

192 **Results**

193 *Chlorophyll florescence*

194 Control plants had an *Fv/Fm* ratio closest to 0.7 but had distinct low periods during spring 195 and winter (Figure 1a). In spring and autumn, the control plants had higher *Fv/Fm* ratios than 196 plants from temperature treatments. The linear mixed effects model showed that both the 197 fixed terms of Day and Treatment were significant as well as the interaction between Day 198 and Treatment. Between-pot variability was found to be the largest random effect (Table S1, 199 Figure S2). Physiological damage was confined to the upper portion of stems, where the 200 *Fv/Fm* ratio was reduced in the top 20 mm of stems in the 400 and 400+ treatments compared 201 to the control (Figure 2).

202 *CO2 exchange*

203 Net CO₂ exchange under full light (2000 μ mol m² s⁻¹), *A*_{max}, varied considerably between 204 runs and treatments, ranging from 84 ± 13 to 252 ± 26 µmol g⁻¹ (dry weight) day⁻¹ with highest 205 values in control pots during autumn (Figure 1b). *A*max in temperature-treated pots was only 206 noticeably lower during the first half of the spring and autumn runs. In general, the 207 respiration rate was less variable between treatments than *A*max (Figure 1c) during both 208 autumn and winter. Respiration rate also varied less between runs, ranging from -105±33 to - 209 13 \pm 22 µmol g⁻¹ (dry weight) day⁻¹ in autumn and -103 \pm 11 to 17 \pm 12 µmol g⁻¹ (dry weight) $210 \, \text{day}^{-1}$ in winter.

211 The linear mixed effects models for *A*max, respiration and fluorescence showed that the Day 212 and Treatment fixed-effect terms were significant (Table S2). In contrast, Day was not found 213 to be significant in models of respiration rate (Table S3) with Hummock and Block the best 214 random effects terms to explain the variance beyond the Treatment effect (Figure S3).

215 Common to both the models of *A*max and respiration was that the random effects explained 216 little of the within-treatment variance

217 *Physical damage and new growth*

218 In control pots, bleaching was largely absent, only occurring for short periods on one or two 219 stems per pot after particularly warm and dry conditions. Most stems in high-temperature 220 treatments showed some bleaching (Figure 3) of the upper parts and capitulum, with the 100 221 \degree C treatment showing the least amount of bleaching (Figure 4), and bleaching being more 222 pronounced a few days after heat treatment.

223 Depth of physiological damage was confined to the upper portion of stems; the *Fv/Fm* ratio 224 was reduced in the 400 and 400+ treatments in the top 20 mm of stems (Figure 2). In the 400 225 ^oC treatment, the extent of damage increased with residence time. The greatest depth of 226 damage occurred in the 400+D treatment (Figure 4). No damage was found down stems in 227 control pots (Figures 2 and 4). A loss of structural integrity of the capitula was found in all 228 400+ treatments in autumn and winter, but was not seen at all in the control treatment (Figure 229 5).

230 New growth during the duration of the experiment arose in new, smaller and more elongated 231 auxiliary stems in all treatments after 100 days in spring and winter (Figure 6). No new 232 growth was found in any of the pots in autumn. Two distinct zones of growth were apparent 233 in both spring and winter runs with new stems growing from upper side innovations and base 234 innovations which grew from the lowest portion of the original stem. In both spring and 235 winter, most new growth occurred in pots which had been subjected to the higher 236 temperature treatments as opposed to control pots. Significantly more new growth was seen 237 in spring in both control and temperature treatment pots than in winter.

238 The distance down the stem where new side innovations grew was correlated to the depth of 239 bleaching, as the new side innovations grew from unbleached areas below the bleached stem 240 (Figure 7).

241

242 *Discussion*

243 Our results show that the photosynthetic capacity of *S. capillifolium* was reduced following 244 exposure to high temperatures, and that higher temperatures and longer residency times 245 caused more physical damage. However, we found that *S. capillifolium* has the capacity to 246 recover its photosynthetic capacity by producing new auxiliary growth.

247 *Photosynthetic capacity and CO2 exchange*

248 The photosynthetic capacity, (chlorophyll fluorescence and CO2 assimilation) of the upper 249 sections of *S. capillifolium* was found to vary considerably between temperature treatments, 250 and between each run of the experiment carried out in the different seasons. The highest 251 *Fv/Fm* ratio closest to healthy plants ratio (around 0.75 Demmig and Bjorkman 1987), was 252 found in control pots. Treated plants showed a general increase in *Fv/Fm* after an initial drop 253 following high temperature treatment, but with quicker recovery to healthy *Fv/Fm* ratios in 254 treatments with lower maximum surface temperatures and residency times. The ascending 255 order of severity, indicated by the reduction in photosynthetic capacity and damage 256 (bleaching) sustained, of the treatments can be summarised as control <100 <400<400+ (with 257 the increased temperature residency time) <400+D (greatest damage caused when *S.* 258 *capillifolium* was dried prior to temperature treatment.).

259 Other than the controls, the least reduction in photosynthetic capacity was seen in pots

260 exposed to 100 \degree C, suggesting that this treatment did not cause severe damage. A similar

261 effect was seen for bleaching (results not shown). Pots treated with a maximum surface

262 temperature of 400 $^{\circ}$ C showed the greatest reduction in Fv/Fm . Little difference was detected 263 in Fv/Fm among 400 °C treatments where residence time and pre-treatment moisture content 264 were varied. This suggests that the maximum temperature reached at the surface of the 265 *Sphagnum* layer may be a sufficient indicator of the short-term impact on photosynthetic 266 capacity. As shown by here, damage to plant cells brought about by fire, such as protein 267 denaturation or lipid mobility (Levitt 1972) can be brought about by exposure to surface 268 temperatures of around 400 °C for just 3 s in *S. capillifolium* at a pre-treatment moisture 269 content of around 90%.

270 Another important observation was that the *Fv/Fm* varied both within a run and between 271 runs, suggesting that both short-term changes in environment and seasonality are important. 272 This was demonstrated by the control pots, which did not show the steady increase in *Fv/Fm* 273 over time as seen in temperature-treated pots, but considerable variation between sample 274 days. Stem moisture content was found to account for the most within treatment and sample 275 day variation with lowest *Fv/Fm* in control pots corresponding to lower stem moisture 276 content and a particularity warm period during spring. During the winter run of the 277 experiment, it was also found that the lowest *Fv/Fm* found in control pots occurred after a 278 period of a few days when the Sphagnum had frozen.

279 An optimum stem moisture content for photosynthesis has been shown in *Sphagnum,* with 280 declining rates of CO2 assimilation coupled with a reduction in stem moisture content 281 (Clymo 1973; Johansson and Linder 1980; Strack et al. 2009; Titus et al. 1983; Williams and 282 Flanagan 1996). The moisture content needed for maximum photosynthesis varies between 283 species (Clymo 1973; Strack et al. 2009; Williams and Flanagan 1996) and seasonally 284 (Johansson and Linder 1980; Titus et al. 1983). Specifically, the *Fv/Fm* measured using 285 chlorophyll fluorescence has been shown to decline with reduced stem moisture content in 286 *Sphagnum* (van Gaalen et al. 2007). *Sphagnum* has been shown to tolerate desiccation to a

287 critical moisture threshold (Schouwenaars and Gosen 2007) when reached net photosynthesis 288 ceases (Schipperges and Rydin 1998). This suggests that the drying experienced in control 289 pots during this study was survivable and did not drop below this threshold. In the 400+D 290 treatment, *Sphagnum* were dried to a moisture content of 80% prior to treatments, and they 291 remained consistently drier, up to 88 days than other treatments. This could be caused by the 292 water transport and holding capacity of the *Sphagnum* being compromised by exposure to 293 high temperatures. This suggests that high temperatures caused by fire may make *Sphagnum* 294 vulnerable to long-term damage brought about by drought, by increasing the likelihood of 295 drying below the critical threshold. Therefore, post-fire conditions may be important and 296 short-term environmental changes may have long-term influences on productivity in 297 *Sphagnum* (Backéus 1988; McNeil and Waddington 2003).

298 *A*max largely reflected the treatment effects on *Fv/Fm* with the exception of the lack of 299 treatment effect on *A*max in the winter run. During spring and autumn, the control pots showed 1300 higher A_{max} than the 400 °C temperature treatments with some degree of recovery shown in 301 the temperature pots in spring. Recovery of the temperature treatments was not seen in either 302 winter or autumn, but this could be because measurements were not continued for the 303 duration of the autumn run because of mould contamination. There was no significant 304 difference in A_{max} in the 100 °C treatment, suggesting that higher temperatures had a more 305 detrimental effect on photosynthesis, which was further supported by the *Fv/Fm* 306 observations. There was no clear treatment effect on respiration, despite it still being a 307 significant term in the mixed effects model. This is likely to be due to differences in 308 respiration rates observed on day 1 between the control pots and temperature treated pots in 309 autumn.

310 There was considerable variation between sampling days particularly in winter *A*max but no 311 detectable difference between treatments. During both autumn and winter, *A*max in temperature-312 treated pots followed the same temporal pattern of *A*max of control pots. This suggests that *A*max 313 in all pots was determined by other factors beyond the temperature treatments.

314 *Sphagnum* has been found to exhibit strong seasonal variation in productivity, with short-day 315 photoperiods (Gerdol 1995; Li and Glime 1991) and low temperatures associated with up to a 316 five-fold reduction in growth (Gerdol et al. 1998). The findings here support this seasonality 317 with the lack of CO2 assimilation during the winter run of the experiment. However, the 318 lower stem moisture content experienced throughout the winter run could also account for 319 low *A*max in the control pots. This suggest that the implications for prescribed burning may be 320 that if photosynthesis and growth rates are lower during the colder and shorter days of winter, 321 then rates of recovery could be much slower following fires which have taken place from 322 October to February that burns which happen at from March to April. Seasonality and timing 323 of fire is therefore an important consideration when reducing the impact on *Sphagnum* is a 324 goal.

325 In real fires in the field, the deposition of ash on to the moss layer may have detrimental 326 effects on photosynthesis, but very little is known about this. Future work using laboratory-327 based simulated fires could usefully separate the effects of ash deposition from the effects of 328 high temperature, and examine any interaction effects.

329 *New growth*

330 New growth in side and base innovations were only found in spring and autumn. Low light 331 levels in the winter run of the experiment, could account for the lack of new growth 332 observed. An additional control on growth is night-time temperature and *S. capillifolium* has 333 been demonstrated to have a five-fold increase in growth at a night-time temperature of 15 \degree C 334 compared to 5 $\rm{^{\circ}C}$ (Gerdol et al. 1998). The temperatures recorded during autumn declined 335 from around day 50, making them lower than those in spring, so these low temperatures 336 could also contribute to the lack of growth observed.

337 It is important to highlight the need to take into account post-burn conditions when assessing 338 *Sphagnum* recovery especially when assessing fire severity. For example, management burns 339 occurring in the spring may show faster rates of *Sphagnum* recovery due to the more 340 favourable growing conditions than those found in winter. Thus, post-burn recovery may 341 have as much to do with season than with the fire itself as post-fire conditions, most notably 342 the moisture status of the *Sphagnum* layer and height of the water table may retard or 343 promote growth (Robroek et al. 2007; Rochefort et al. 2002). This would make it necessary 344 to include post-burn environmental variables in methods that assess fire severity.

345 Our observation regarding regeneration was that side innovations appeared to be very similar 346 to those described by Clymo and Duckett (1986) who suspected that the ability of *Sphagnum* 347 to produce new shoots was a widespread and important mechanism to overcome disturbance 348 (see also Hamilton 2000; Rochefort et al. 2002). Regeneration and production of new 349 innovations has been observed in the field (Burch 2009; Hamilton 2000),where patches of 350 *Sphagnum* produced new green capitula on the surface and capitula regained colouration 351 after bleaching, following a fire. This suggests that the side innovations observed here were 352 not just a product of experimental conditions.

353 The most severe temperature treatment (400+D) used here was intended to be fatal to *S.* 354 *capillifolium* but was still found to result in new growth. This suggests even higher surface 355 temperatures and longer temperature residency times are needed to kill S*. capillifolium*. 356 Clymo and Duckett (1986) demonstrated *Sphagnum* growth 30 cm below the surface, 357 suggesting that high temperatures would have to penetrate very deep within the *Sphagnum*

358 layer to prevent any regeneration. However, at this depth, light may limit regeneration after 359 fire, and complete consumption would prevent any regeneration. Nevertheless, partial 360 consumption may allow sufficient light for side or base innovations to proliferate and hence 361 allow recovery.

362

363 *Conclusions*

364 The aim of this research was to determine the short-term responses of *Sphagnum* 365 *capillifolium* to fire. We found that the rate of photosynthesis was reduced by exposure to 366 high surface temperatures. High temperature also increased the extent of bleaching and 367 capitulum loss. Importantly though, within the range of surface temperatures and residence 368 times used here, no critical threshold was found to cause widespread death of *S. capillifolium*. 369 Even in the treatment specifically designed to be lethal, new auxiliary stem growth was 370 found. Our results provide evidence that *S. capillifolium* has the ability to recover from the 371 high temperatures experienced in typical prescribed fires, provided that at least some living 372 material remains. The experiment also suggests seasonal effects are important to *S.* 373 *capillifolium* recovery, and that recovery may be conditional on the fire timing. Although 374 these results demonstrate that *S. capillifolium* has the ability to survive a fire event, it is 375 important to consider these results within the context of management burning regimes. Lee et 376 al. (2013) for instance have demonstrated that, although *Sphagnum* may survive a fire, long-377 term frequent burning (every 10 years) can reduce the propagule bank within the peat. This 378 could reduce the capacity for recovery from fire events which wholly consume the *Sphagnum* 379 layer. Future research into the impact of the types of fires simulated here, which are 380 supported by the current best practice guidance and legislation, should include other 381 *Sphagnum* species, particularly those from differing micro-habitats, to establish if the

- 382 findings are generalizable to other *Sphagnum* spp. and identify micro-habitats or species
- 383 which may be most vulnerable.

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Table 1 Temperature treatments, designed to simulate conditions recorded in *Calluna vulgaris* fires, used on samples of *Sphagnum capillifolium* for three runs of an experiment to determine its capacity for recovery. Burn Season refers to the time of year the pots were exposed to each temperature treatment. All pots were observed and recovery measurements made for a total of 100 days after being exposed to each temperature treatment which is termed Observation Period in the table. $n = 32$ per treatment per run of the experiment. *400+D indicates where the treatment was carried out on pots of *Scapillifolium* subjected to three days of drying prior to the treatment. As some treatments were repeated in different runs total pots varied by treatment; control (n=96), 100 (n=32), 400 $(n=64)$, 400+ $(n=64)$, 400+D $(n=32)$.

Table 2 Fixed and random effects terms used in mixed effects modelling of the repeated measures of chlorophyll fluorescence and CO2 exchange of *Sphagnum capillifolium* samples exposed to temperature treatments designed to simulate conditions recorded in *Calluna vulgaris* fires.*Moisture content term only used in chlorophyll fluorescence model as stems were harvested for moisture content analysis only on days fluorescence measurements were made.

Figure Captions

Figure 1: (a) *Fv/Fm* ratio of *Sphagnum capillifolium* stems subjected to temperature treatments designed to simulate conditions recorded in *Calluna vulgaris* fires. The experiment was repeated in three seasons; spring, autumn and winter (n=8 per treatment per sampling time). Treatments were: control, no temperature treatment; 400 surface exposed to 400oC for 3 seconds; 400+, surface exposed to temperatures between 350 and 450oC for 30 seconds; and 400+D, where the moss sample was dried prior to exposure to surface temperatures between 350 and 450oC for 30 seconds. Points show mean $Fv/Fm \pm SEM$ bars. **(b)** A_{max} in each treatment group (described above) during each run. Points show mean ±SEM bars. Positive values show CO2 uptake (indicating photosynthesis). **(c)** Respiration of pots in each treatment group in autumn and winter. Points show mean ±SEM bars. Respiration is expressed as a negative quantity in our sign convention. No data were available from the spring experiment.

Figure 2: *Fv/Fm* ratio of *Sphagnum capillifolium* stems subjected to temperature treatments designed to simulate conditions recorded in *Calluna vulgaris* fires. The *Fv/Fm* ratio is shown in relation to distance down the stem, starting from the capitulum. Data are from a sub-sample of pots in the autumn experiment (n=6 per treatment). Treatments were either a control without temperature treatment (C), exposed to 400 $^{\circ}$ C for 3 seconds (400), or exposed between 350 and 450 °C for 30 seconds $(400+)$.

Figure 3: Examples of bleached *Sphagnum capillifolium* subjected to temperature treatments designed to simulate conditions recorded in *Calluna vulgaris* fires. (a) Pot showing the characteristic pale areas of bleaching of the capitula (red arrow). Bright green capitula are growth innovations from the stem below. (b) Several stems with individual branches bleached (red arrow). The lack of colouration in the lower stem is a normal response to low light levels. (c) *S. capillifolium* plant showing bleached capitulum (red arrow) and new growth innovation near the top of the plant (green side stem). (d) *S. capillifolium* plant showing bleached stem (red arrow) and new growth innovation near the bottom of the plant (green side stem) which were characteristically smaller and thinner than those arising further up the stem. When bleaching occurred, capitula became brittle to touch and easily broke away from the stem.

Figure 4 Depth of bleaching down stems of *S. capillifolium* (mean ± SEM) exposed to temperature treatments designed to simulate conditions recorded in *Calluna vulgaris* fires in a sub-sample of pots from experiments carried out in autumn and winter (n= 6 stems per pot, 16 pots per treatment per run of the experiment). Treatments were either a control without temperature treatment (C), exposed to 400 $\rm{^{\circ}C}$ for 3 seconds (400), exposed to between 350 $\rm{^{\circ}C}$ and 450 °C for 30 seconds (400+), or dried prior to exposure to between 350 °C and 450 °C for 30 seconds (400+D). No permanent bleaching was recorded in any control pots. All pots were harvested on day 100. Means with different letters are significantly different (Welch Two Sample t test: t=-4.6, df=39.6,*P*=<0.05 and t=-5.1, df=25.1, *P*=<0.05 respectively).

Figure 5: Capitulum decay in *S. capillifolium* plants subjected to temperature treatments designed to simulate conditions recorded in *Calluna vulgaris* fires. Bars show the number of stems showing capitulum decay (bleaching and/or reduced structural integrity) at each sampling time for each treatment during autumn and winter (n=8 stems per treatment per sampling time per run).). Treatments were either a control without temperature treatment (C), exposed to 400 °C for 3 seconds (400), exposed to between 350 °C and 450 °C for 30 seconds (400+), or dried prior to exposure to between 350 °C and 450 °C for 30 seconds (400+D). No capitulum decay occurred in control pots.

Figure 6: Bars show the number and location of regenerating stems of *S. capillifolium* following temperature treatments designed to simulate conditions recorded in *Calluna vulgaris* fires. Plots show a subsample of 16 pots per treatment per run showing the total number of new side and base innovations in spring and winter experiments. The total number of new innovations (base+side) was significantly higher in the 400 treatment compared to the 100 and control treatments in spring (Welch Two-sample t test: $t=3.3$, $df=38$, $P=<0.05$ and $t=.3.2$, df=37, *P*=<0.05 respectively). There were significantly more new innovations in the 400+ and 400+D treatments compared to the control treatment in autumn (Welch Two Sample t test: t=2.6 df=39, p=<0.05 and t=3.4, df=36, *P*=<0.05 respectively). No significant difference was found between the 400+ and 400+D treatments.

Figure 7: The location of regenerative growth in relation to the mean depth of bleaching in *S. capillifolium,* following temperature treatments designed to simulate conditions recorded in *Calluna vulgaris* fires. Points show the mean \pm SEM from a subsample of 16 pots per treatment per run from the winter experiment. Treatments were either a control without temperature treatment (C), exposed to between 350 °C and 450 °C for 30 seconds (400+), or dried prior to exposure to between 350 $\rm{^{\circ}C}$ and 450 $\rm{^{\circ}C}$ for 30 seconds (400+D).

