

1 **Above-ground biomass accumulation patterns in moorlands after prescribed burning**
2 **and low-intensity grazing**

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1 **Abstract**

2 Shrub-dominated ecosystems such as moorlands are recognized internationally as cultural
3 landscapes with high biodiversity conservation value. These ecosystems are commonly
4 managed using prescribed burning to reduce the impact of wildfires, increase biodiversity and
5 ecosystem productivity for grazing. Given that ecosystem responses are sensitive to the
6 above-ground balance within the vegetation, knowledge of the above-ground biomass
7 accumulation patterns on moorlands is an important issue for planning management action.
8 Here, we used the replicated long-term manipulative grazing and burning experiment at Moor
9 House (UK) to explore the cumulative effects of multiple fires and low-grazing. The study
10 comprised a comparison between no-burn reference plots (no-burn since ca. 1923) and an
11 experiment where all plots were burned in 1954/55. Within the experiment, the effects of low
12 sheep grazing vs. no grazing and three burning rotations were tested (no-burn since 1954/55,
13 repeat-burning at 10- and 20-year intervals). We hypothesized that prescribed burning and
14 grazing will interact, affecting both the above-ground biomass and vegetation height. The
15 results reveal that although the main above-ground biomass was constrained in three fractions
16 (litter, *Calluna* and bryophytes) there was no significant effect of sheep-grazing or its
17 interaction with prescribed burning (graze×burn) on any biomass variables or vegetation
18 height. Significant reductions in above-ground biomass and vegetation height were only
19 produced by repeated burning. There were no significant differences in biomass or vegetation
20 height between the no-burn since 1954/55 treatment and reference plots. Moreover, *Calluna*
21 biomass and vegetation height showed a positive significant asymptotic association with time
22 since the last burn with an asymptote at 20 and 15 years after fire, respectively. This work
23 demonstrates that burning rotations lower than 20 years reduced the above-ground biomass
24 and vegetation height on this moorland compared to stands unburned for more than 50 years.
25 In order to maximize the C fixation, fire return intervals should be around the *Calluna*
26 biomass accumulation asymptote 20 years since last fire. Furthermore, the vegetation height
27 asymptote of 36 cm, indicating when the vegetation is at its maximum stage, could be a
28 useful tool for guiding when to implement prescribed burning for carbon accumulation
29 purposes.

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1 **Highlights:**

- 2 • The effect of prescribed burning and low-grazing pressure in moorlands was assessed
- 3 • We examined the vegetation height and above-ground biomass accumulation patterns
- 4 • Repeated burning produce reductions in vegetation height and above-ground biomass
- 5 • Low sheep-grazing has no effect on any biomass variables
- 6 • Fire-return interval must be around 20 years since burning to maximize C fixation

7

8 **Keywords:** Fire, return interval, *Calluna vulgaris*, litter, sheep grazing, shrub-dominated

9 ecosystems.

10

1 **Introduction**

2 For many shrub-dominated ecosystems around the world, prescribed burning, the deliberate
3 application of fire under specific conditions (Fernandes and Botelho, 2003), is a widely-used
4 management tool to reduce the impact of wildfires, prevent succession to woodlands,
5 improve wildlife habitats, or increase biodiversity and ecosystem productivity for grazing
6 livestock (Pakeman et al., 2003; Calvo et al., 2005; Borghesio, 2009; Keeley et al., 2012; Lee
7 et al., 2013a). As these benefits usually last for only a few years after fire, repeated prescribed
8 burning is needed for effective management (Ascoli et al., 2009). Therefore, prescribed
9 burning planning requires an understanding of the vegetation response to repeated burning,
10 which involves the implementation of long-term monitoring programs (Fernandes et al.,
11 2013). These long-term monitoring programs help to unravel the potential consequences of
12 the use of prescribed burning on ecosystems, their recovery and their carbon balance (Ascoli
13 et al., 2009; Velle et al., 2012), as well as to define the most effective rotation interval
14 required under present and future global climatic change scenarios (Keeley, 2005).
15 Unfortunately, despite the importance of having well-designed management plans, there is a
16 lack of long-term empirical studies testing the effects of repeated prescribed burning on
17 shrub-dominated ecosystems (e.g. Boer et al., 2009).

18 One of the main difficulties in prescribed burning practices is to quantify the patterns in
19 above-ground carbon balances. Many complex interactions may exist between fire and other
20 management practices where biomass is consumed, such as with grazing. It is well known
21 that grazing is an important biomass consumer in many terrestrial ecosystems and it can
22 interact with fire in many shrub-dominated ecosystems (Rigolot et al., 2002; Ascoli et al.,
23 2013; Johansson and Granström, 2014). Grazing can also influence the above-ground
24 biomass accumulation patterns after fire, which are primarily controlled by a balance between
25 plant growth, litter production (both negatively affected by grazing; Evlagon et al., 2012) and
26 decomposition rates (positively affected; Riggan et al., 1988). This is especially true in the
27 early post-fire years when pasture quality is greatest (Fuhlendorf et al., 2009). As a
28 consequence, prescribed burning might be expected to be implemented in grazed systems at
29 longer time intervals as grazing slows down the biomass accumulation (Johansson and
30 Granström, 2014). However, despite the great interest in prescribed burning as a management
31 tool in shrub-dominated ecosystems, there are few studies that investigate the post-fire
32 biomass accumulation patterns under the influence of grazing (but see Rigolot et al., 2002,
33 Ascoli et al., 2013). Studies of the interactions between plant growth rates and management
34 treatments such as grazing and repeated burning are essential for developing appropriate

1 management strategies within shrub-dominated ecosystems. Indeed, repeated prescribed
2 burning along-side grazing are the predominant tools for the management of north-west
3 European moorlands (Harris et al., 2013a; Lee et al., 2013b; Velle et al., 2014).

4 In Great Britain upland moors, many of them growing on blanket bog (ombrotrophic mire)
5 have a very high conservation value of international significance (Bain et al., 2011; Lee et al.
6 2013a). These moors are currently cultural landscapes that have been created and maintained
7 by anthropogenic activity, mainly sheep grazing and prescribed burning (Rosenburgh et al.,
8 2013). Whilst fire has been used for hundreds, perhaps thousands of years (Simmons, 2003),
9 its use has increased in the last 200 years to enhance the productivity of the moors for sheep
10 grazing and especially for red grouse (*Lagopus lagopus scoticus* Latham) (Harris et al.,
11 2011a; Lee et al., 2013a). Nowadays, approximately, 65% of British upland moors are
12 managed using prescribed burning for the benefit of red grouse (Sotherton et al., 2009).
13 Hence, their sustainable management is important in terms of both the local economy and
14 biodiversity (Harris et al., 2011a).

15 Prescribed burning will inevitably affect the vegetation carbon balance from the moorland
16 systems as the fire moves through it. Carbon balance will depend on the biomass consumed
17 by the fire (initial instantaneous loss) and the ecosystem resilience (Mitchell et al., 2000), i.e.
18 the time it takes for the ecosystem to recover via plant growth and biomass accumulation
19 during the inter-fire interval. Where prescribed burning is done carefully within the approved
20 burning season (winter months in Great Britain; Anon, 2007), using “cool burn” or
21 “pressurized fuel-assisted” burning (Harris et al., 2011a), these losses should be minimized as
22 some vegetation remains after the fire and the peat should be left relatively unaffected. After
23 burning there is often relatively rapid vegetation recovery and hence carbon accumulation
24 during the post-fire succession, and the overall aim should be to produce a balanced budget
25 over a specified time period. In this sense, there is still little information available in
26 moorlands subjected to prescribed fires, being often conflicting. In a moorland area in central
27 England, Allen et al. (2013) used a modelling approach to predict that over a 50 year period
28 the longer the fire-rotation interval, greater the accumulation of above-ground biomass
29 (vegetation and litter). However, Clay et al. (2010) observed, in terms of carbon budgets, that
30 prescribed burning can reduce global C releases in comparison to long-term unburned areas;
31 i.e., taking into account fluvial and gaseous fluxes such as dissolved organic carbon,
32 particulate organic carbon, excess dissolved CO₂, release of CH₄, net ecosystem respiration
33 of CO₂, and uptake of CO₂ through primary productivity. Therefore, that knowledge of the
34 above-ground biomass accumulation patterns on moorlands is fundamental for global change

1 research, and for planning management action; therefore, further research is needed to
2 disentangle the effect of different fire rotation intervals in defining above-ground biomass as
3 C sink and source. Undoubtedly, such knowledge will assist in determining the fire rotation
4 interval that optimizes C fixation by means of vegetation growth.

5 As far as moorland conservation management in Great Britain is concerned, a major issue
6 that needs to be addressed is a quantification of the effects of low-intensity grazing, current
7 on many moorland ecosystems, and repeated prescribed burning on the above-ground
8 biomass during the prescribed burning/post-fire recovery cycle. To address this, we measured
9 above-ground biomass within the replicated long-term manipulative grazing and burning
10 experiment at Moor House National Nature Reserve (Rawes and Hobbs, 1979; Lee et al.,
11 2013a). This experiment has a history of approximately 90 years of known low-grazing
12 pressure and fire rotations at different intervals (10, 20 and 56/57 years). This experiment,
13 therefore, represents a unique opportunity to quantify the cumulative effects of multiple fires
14 and low-grazing on biomass accumulation and its related parameters such as the dry weight
15 of component fractions (e.g. *Calluna vulgaris* (L.) Hull, litter, bryophytes, graminoids and
16 other vascular plants) and vegetation height. For all of these measures we assessed the effect
17 of (a) grazing (grazing vs. no grazing), (b) the different rotation intervals (short-, long-, and
18 unburned for 50+ years) and their interaction. In addition, we modelled plant growth through
19 time since the last burning; as both the total accumulated, and as the absolute growth rate
20 (AGR).

21 Essentially, our aim is to determine what is the optimal fire return time based on the
22 biomass patterns found. For that, we made a comparison of biomass accumulation with
23 literature sources and we tested two main hypotheses: First, that the shorter fire return-
24 intervals will produce greater reductions in above-ground biomass and vegetation height.
25 Second, it has been shown that grazing slows down the above-ground biomass accumulation
26 by consumption, especially in early post-fire years when pasture quality is greater (Velle et
27 al., 2012; Johansson and Granström, 2014); therefore, we expect that biomass reduction by
28 grazing will be greater in shorter fire return-intervals.

29

1 **Material and methods**

2 *Study area*

3 The study site is within the Moor House National Nature Reserve (hereafter referred as Moor
4 House, Table 1), which is located in the northern Pennines, a range of hills that form the
5 backbone of England (54°41'34.4"N, 2°24'28.1"W). The experimental site is on the eastern
6 side of Hard Hill, a gently-sloping, high-level plateau (600-650 m; Heal and Smith, 1978);
7 and it is situated on blanket bog (>50 cm peat, the widely-accepted definition for blanket peat
8 in the UK; Costigan et al., 2005). The vegetation at Moor House can be described as *Calluna*
9 *vulgaris*-*Eriophorum vaginatum* blanket mire (M19) and *Eriophorum vaginatum* blanket and
10 raised mire (M20) communities within the British National Vegetation Classification (NVC;
11 Rodwell, 1991). Thus, the most common species are *Calluna vulgaris*, *Eriophorum*
12 *vaginatum* L., *Pleurozium schreberi* (Brid.) Mitt. and *Sphagnum capillifolium* (Ehrh.) Hedw.
13 Here, *Calluna* was the dominant species in the vegetation sampled (i.e. biomass and height);
14 with no presence of other woody shrub species. The climate is oceanic/sub-arctic rather than
15 temperate, cool, wet and windy (abridged for Heal and Smith's (1978) description of the
16 Moor House climate). The January and July mean temperatures at Moor House are 2.7°C and
17 14.4°C, respectively. The Moor House annual precipitation is 1314 mm (data derived from
18 UK Meteorological Office 5-km monthly gridded climatic data averaged between 1961-2005;
19 Perry and Hollis, 2005), and precipitation was much greater in winter (January mean = 130
20 mm) compared to summer (July mean = 82 mm).

21

22 *Experimental design*

23 The experimental design is detailed in Marrs et al. (1986). Briefly, the experiment was set up
24 on Hard Hill at Moor House in 1954/5 and was designed to test the effects of low-intensity
25 grazing vs. no sheep grazing in combination with three prescribed burning rotations (a 10-
26 and 20-year rotation plus a no-further-burn treatment). In 1954/5 four replicate moorland
27 blocks (A-D), each 90 m × 60 m, were burned along an elevational gradient (A = NY743330;
28 B = NY740330; C = NY736330; D = NY738331). Blocks A, B and D were burned in 1954
29 and block C in 1955. At the start of the experiment the vegetation was considered to have
30 remained unburned for at least 30 years (Rawes and Hobbs, 1979). Within each block, two
31 main-plot treatments (60 m × 30 m) were allocated randomly, these treatments were: sheep
32 grazing and no sheep grazing, referred to here as Grazed (G) and Enclosed (E), respectively
33 (Table 1). The sheep grazing pressures have varied during the experimental period, but have

1 always been low. The densities on this moorland type vegetation were estimated at ca. 0.1-0.3
2 sheep ha⁻¹ in the 1960s when the overall sheep grazing density over the entire reserve was
3 15,400 sheep in the summer months, an average of 4.4 sheep ha⁻¹ across all vegetation types
4 (Rawes and Welch, 1969). The formalization of grazing rights under the Commons
5 Registration Act (1965) was completed for Moor House in 1972 and grazing was then
6 restricted to 7,000 sheep, a halving of sheep numbers (2 sheep ha⁻¹). There was a further
7 reduction in sheep numbers following the foot and mouth outbreak in 2001 when some
8 grazing rights were extinguished and a new stocking density of 0.5 sheep ha⁻¹ established
9 over the whole moor. Hence the already light stocking density present when the experiment
10 was set up has been reduced on two occasions during the study to an approximate 15-fold
11 reduction.

12 Within each main-plot, three burning-rotation treatments were allocated randomly to sub-
13 plots (30 m × 30 m), these were: (i) Short-rotation burning approximately every 10 years (S),
14 (ii) Long-rotation burning approximately every 20 years (L), and (iii) No burning since
15 1954/5 (N). Prescribed burning in the weather conditions prevailing at Moor House is very
16 difficult and in some years burning is impossible, accordingly burning timings could not be
17 applied fully in accordance with the planned schedule but were applied as follows: 1954/55
18 (All), 1965 (S), 1975 (S & L), 1984 (S), 1995 (S & L) and 2006 (S). Thus, the experimental
19 data here represents six burns in S, three burns in L and one burn with subsequent recovery
20 after 56/7 years (N) since the start. However, since monitoring was done in 2011 the years
21 between sampling and last burn for each treatment were: S=5 years, L=16 years and N=56/57
22 years since the last burn (Table 1). In addition to the formal experiment, each block had an
23 associated unburned reference plot (denoted R, 10 m × 10 m) set up outside the burn area
24 delimited in 1954/5, but sheep grazed. The exact positions of these unburned reference
25 controls have been lost over time but in 2011 the approximate positions of these plots were
26 relocated using a combination of map locations and aerial photography to ensure they were
27 placed outside the original burn areas. We accept they may not be in identical positions but
28 they are extremely close. These reference plots were deemed to have remained unburned for
29 at least ca. 87 years, although the exact date of the last burn remains unknown; for calculation
30 purposes we have assumed the elapsed time since last burn was 90 years.

31

32 *Biomass monitoring after prescribed burning experiments*

33 In July 2011, above-ground biomass was harvested from each of the 28 sub-plots (Table 1).
34 In each sub-plot, three 0.25 m² quadrats were sampled in the buffer zone surrounding the

1 vegetation monitoring plot to minimise disturbance to ongoing research. In each quadrat, first
2 the vegetation height was measured with a ruler (cm) in five points (i.e. centre and the four
3 quadrat corners), and then plot average was taken. Here, vegetation height was predominantly
4 dictated by *Calluna* height. Afterwards vegetation was harvested to 2 cm from the solid peat
5 surface and separated crudely in the field into two fractions (*Calluna* and the remainder). In
6 the laboratory, the two fractions from each sample were re-sorted to produce a sample of
7 *Calluna* and remaining vegetation per plot; the *Calluna* sample was then oven-dried and
8 weighed. The remaining vegetation was then quartered; one quarter selected randomly was
9 retained for further separation and the other three-quarters oven-dried and weighed. The
10 retained quarter was separated into four fractions: litter, bryophytes, graminoids and a
11 combined fraction containing all other vascular plants. The litter was distinguished from live
12 material by absence of chlorophyll. In these vegetation samples *Calluna* was the only shrub
13 species detected. All fractions were then oven-dried and weighed. The proportions of each
14 fraction in the sub-sample were applied to the entire sample to derive component weights.

15

16 *Data analyses*

17 Biomass data from all burns were converted to g m^{-2} dry weight. Unfortunately, the statistical
18 design at Moor House was unbalanced (no enclosed reference plot) so the data were,
19 therefore, analyzed as follows: First, a comparison of burning treatment effects on vegetation
20 height and biomass variables (total, *Calluna*, litter and bryophytes biomass) in the grazed
21 plots. Here, the data for all four grazed treatments (S, L, N, R) were compared using Linear
22 Mixed-effects models to account for the spatial structure of the data (random effects);
23 Block/Plot/Sampling quadrat. We accept that the R treatment was not positioned randomly
24 within the experimental design. Second, a comparison of the formal experiment was done.
25 Here, the R treatment was excluded and the effects of grazing (G, E) and burning rotation (S,
26 L, N) and their interactions on vegetation height and biomass variables (total, *Calluna*, litter
27 and bryophytes biomass) were assessed using the same methodology. In both analyses the
28 biomass of graminoids and the combined fraction containing other vascular plants were not
29 analyzed statistically, because these fractions were detected in less than 20% of the plots.

30 Afterwards, we analyzed the plant growth through elapsed time since the last burning
31 using two variables: *Calluna* biomass and vegetation height. An understanding of growth is
32 essential to understand ecological processes including the interactions between plants and
33 factors affecting communities such as grazing and burning. Here, these relationships were
34 assessed using non-linear mixed models based on the idea that plant size may approach an

1 asymptote because of limiting below-ground resources or ontogenetic changes (Paine et al.,
2 2012). In the case of *Calluna* non-linear mixed Gompertz growth curves were fitted; the
3 choice of Gompertz curves was to maintain consistency with previous published works on
4 British heath/moorland areas (Chapman et al., 1975; Miller, 1979). The Gompertz regression
5 model (function “SSgompertz”) full equation is $y = \text{Asym} \times \exp(-b2 \times b3^{\text{Age}})$, where y is
6 biomass variables, Asym is the asymptote, $b2$ is the y-intercept and $b3$ determines the rate at
7 which the asymptote is reached. Vegetation height was modelled using a non-linear logistic
8 function (function “SSlogis”) where the full equation is $y = \text{Asym} / (1 + \exp((x_{\text{mid}} - \text{Age}) / \text{scal}))$,
9 where y is the vegetation height, Asym is the asymptote, x_{mid} is the inflection point
10 of the curve and scal determines the scale parameter of the age since the last burning
11 parameter. In all non-linear analyses, the spatial structure of the data and treatments applied
12 (grazing and burning) were included as random factors; Block/Graze/Burning/Sampling.
13 Next, the absolute growth rate (AGR, biomass = $\text{g m}^{-2} \text{ year}^{-1}$; height = cm year^{-1}) were
14 derived from the non-linear models fits using the methodologies and scripts described in
15 Paine et al. (2012). These growth rates were expressed as functions of time since the last
16 burning and their confidence intervals were derived from population prediction intervals
17 (Paine et al., 2012). Because the litter, bryophytes, graminoids and other vascular plants
18 group biomass showed no change since the time of last burning (continuous values along the
19 sequence, $P > 0.050$); the total biomass pattern was controlled mainly by the *Calluna* biomass
20 pattern. As a consequence only a detailed description of *Calluna* biomass change is reported.

21 All statistical analyses were performed in the R statistical environment (version 3.1.0 R
22 Development Core Team, 2014); Linear and non-linear Mixed-effects modelling was
23 performed using the “nlme” package (Pinheiro et al., 2014).

24

25

1 **Results**

2 *Biomass distribution within the vegetation*

3 The biomass distribution within the vegetation at Moor House showed that there were three
4 main fractions; litter, *Calluna* and bryophytes; these fractions represented almost 97% of the
5 total biomass (Fig. 1). Litter was the most abundant biomass fraction (36% - 67%) followed
6 by *Calluna* (5% - 43%) and bryophytes (6% - 27%). The other two fractions, either the
7 graminoids or other vascular plants biomass, made up a trivial contribution to the total
8 biomass. Graminoid biomass was more abundant in short-rotation treatments (S) independent
9 of grazed treatment (ES = 1.1% and GS = 0.7%), whereas the biomass of other vascular
10 plants was only abundant in short rotation enclosed plots (ES = 0.9%). At the same time,
11 there were few differences in the distribution of the three main biomass fractions (litter,
12 *Calluna* and bryophytes) between grazing treatments (E vs. G), although there were
13 consistent differences between burning treatments (Fig. 1). *Calluna* biomass was greater at
14 the longer rotation interval treatment (N, followed by L and S), whereas bryophytes biomass
15 was greater in the shortest rotation interval (S).

16

17 *Burning rotations effects on biomass and height in the grazed plots*

18 As expected, there were no significant differences between the no-burn since 1954/55
19 treatment (N) and reference plots (R) on any of the five biomass and height measures
20 considered ($P > 0.050$, Table 2). In contrast, the short- and long- burning rotation treatments
21 (S and L) produced significant reductions of total biomass compared to the reference plots (P
22 < 0.050 , Table 2). These reductions were greatest in the short-term rotation treatment (S)
23 which had the lowest biomass value ($S = 1198 \pm 165 \text{ g m}^{-2}$) followed by long-term rotation
24 ($L = 1593 \pm 119 \text{ g m}^{-2}$). Indeed, the short-rotation treatment was the only burning treatment
25 that reduced *Calluna* biomass and vegetation height compared to the reference plots (R, $P <$
26 0.001 , Table 2). In contrast, bryophytes biomass was only reduced by the long-term rotation
27 treatment (L; $P = 0.043$), whereas, the litter biomass was not affected by any burning rotation
28 ($P > 0.050$).

29

30 *Grazing and burning rotations effects on biomass and height in the formal experiment*

31 Within the formal experiment, the grazing treatments showed no effect on their own nor in
32 interaction with prescribed burning (graze \times burn) on any biomass variables or height ($P >$
33 0.050). Significant differences were only found in four biomass variables (total, *Calluna* and
34 bryophytes biomass and height) with respect to the prescribed burning rotations ($P < 0.050$).

1 The results were similar to the previous analysis; prescribed burning rotation treatments S
2 and L produced reductions on biomass in comparison with the no-burn since 1954/55
3 treatment (N) ($P < 0.050$). Total biomass was lower in short rotations than in no burn
4 treatments ($N = 2037 \pm 25$ vs. $S = 1342 \pm 194$, t -value = -3.57 , $P = 0.002$), *Calluna* biomass
5 was reduced by burning rotation ($N = 833.70 \pm 70.25$ vs. $L = 656.98 \pm 81.39$ vs. $S = 83.50 \pm$
6 81.39 , F -value = 46.44 , $P < 0.001$), bryophytes biomass was reduced by long-rotation burning
7 ($N = 415.52 \pm 106.80$ vs. $L = 166.10 \pm 113.25$, t -value = -2.26 , $P = 0.041$) and vegetation
8 height by short-rotation burning ($N = 35.83 \pm 1.28$ vs. $S = 22.29 \pm 1.81$, t -value = -7.48 , $P <$
9 0.001). Litter biomass showed no response to burning rotation ($P > 0.050$).

10

11 *Calluna biomass and height accumulation patterns through time*

12 The non-linear mixed-effects analysis showed that *Calluna* and vegetation height had a
13 positive asymptotic association with elapsed time since the last burning (Table 3, Fig. 2).
14 Both variables increased with elapsed time since the last burning until they stabilized.
15 *Calluna* biomass reaches an asymptote of $795.87 \pm 80 \text{ g m}^{-2}$ approximately at 20 years after
16 fire and vegetation height reaches an asymptote of $36.31 \pm 0.84 \text{ cm}$ approximately at 15 years
17 after fire (Table 3). The AGR peak for *Calluna* was at 8 years after the last fire with a value
18 of $76.08 \text{ g m}^{-2} \text{ year}^{-1}$, whereas the AGR peak for vegetation height was earlier at 4 years, with
19 a value of $4.61 \text{ cm year}^{-1}$ (Fig. 2b, d).

20

21 **Discussion**

22 *Biomass distribution within the vegetation and comparison with literature sources*

23 The data collected at Moor House provides comprehensive information on above-
24 ground biomass budgets during the prescribed burning/post-fire recovery cycle on high-
25 elevated moorlands. The above-ground material was accumulated almost entirely in
26 three biomass component fractions: litter, *Calluna* and bryophytes, with negligible
27 amounts of graminoids and other vascular plant species. This almost certainly reflects the
28 impact of severe climate and the lack of exposure to pollutants as other moorlands (Tallis,
29 1988; Caporn and Emmett, 2009; Lee et al., 2013a). Moor House being colder, wetter and
30 less polluted have a lower productivity (e.g. *Calluna* biomass), more litter and peat
31 accumulation than low-land moorlands where productivity should be enhanced by the drier
32 (less water-logged), warmer conditions and a greater nitrogen loading (Lee et al., 2013b;
33 Table 4). At the same time, Moor House has retained a substantive bryophyte component
34 including the peat-forming *Sphagnum* species (Rawes and Hobbs, 1979; Lee et al., 2013b)

1 probably because of a lack of exposure to present and past pollutants (nitrogen deposition;
2 Tallis, 1988; Caporn and Emmett, 2009; Lee et al., 2013a) which has favoured the biomass
3 increases of bryophytes in this ecosystem and especially under the shorter-rotation intervals
4 (S). On the other hand, the deep litter layer acts as an inhibiting recolonization *sensu* Connell
5 and Slatyer (1977), reducing the development of graminoids and other vascular plants groups
6 (Lee et al., 2013a), as a result their above-ground biomass in all blocks was trivial (<1.2%).

7 The lower productivity is supported by evidence from other studies, it is clear that the total
8 biomass data for older stands at Moor House that ranged from 2076-2223 g m⁻² were
9 considerably lower than comparable data from the literature (5,240-10,000 g m⁻²; Table 4).
10 Literature values for younger moorland stands at Teesdale, Kincardineshire or Dartmoor were
11 within the same orders of magnitude as the Moor House total biomass values for greater age
12 ranges (Moor House=2076-2223 g m⁻² at age range 56/57-90 years vs. younger Moors=1820-
13 2930 g m⁻² at age range 6-25 years; Table 4). The vegetation at Moor House does not appear
14 to follow the traditional *Calluna* four-phase, life-cycle model of Watt (1947, 1955), where the
15 transition from mature to degenerate phases involves the older branches falling over with
16 gaps being created in the middle (Barclay-Estrup & Gimingham, 1969). At Moor House, the
17 degenerate phase does not appear to be pronounced rather the mature phase continues
18 vertically and with horizontal growing along the ground and through other vegetation; the
19 stems adventitiously rooting in the bryophyte/litter layer. This was first noted by Forrest's
20 (1971) observations that at Moor House the moorland vegetation was in a presumed steady-
21 state but the degenerate phase of *Calluna* was seldom seen. It will be interesting to monitor
22 these patterns in the future especially if either conditions become warmer and drier as a result
23 of climate change, or there is a reduction in burning management.

24

25 *Burning rotations effects on biomass and height*

26 This study has provided a detailed long-term insight into the above-ground biomass dynamics
27 of high-land moorland communities in Great Britain managed using repeated prescribed
28 burning rotation intervals. Our results indicate that the shorter fire return-intervals (5 years-S
29 and 16 years-L after last fire) produced greater *Calluna* biomass and height reductions in
30 comparison with unburned controls (56/57 years-N and 90 years old-R), and this was equal
31 independently of the type of analysis done (grazed plots or formal experiment); hence,
32 Hypothesis 1 is accepted with respect to burning rotations. As expected, when the fire
33 frequency increased there is a reduction of the dominance of *Calluna* (Littlewood et al., 2010;
34 Allen et al., 2013). In contrast, short-burning rotations do not optimize the C fixation based

1 on vegetation growth as the *Calluna* biomass asymptote is held at 20 years, equal to long-
2 rotation burning treatment in this moor. However, short rotations appears to be associated
3 with an increase in peat-forming species (*Eriophorum* spp. and *Sphagnum* spp.; Lee et al.,
4 2013a), being fundamental for moorland diversity and conservation, and also for carbon
5 fixation in the peat (Lee et al., 2013a). It seems that no burning for more than 50 years (N and
6 R treatments) will produce a stand dominated mainly by *Calluna* and litter with low biomass
7 of bryophytes and other vascular plant species, and with a community composition changing
8 towards *Calluna vulgaris-Hypnum jutlandicum* with a reduction of peat forming species (Lee
9 et al., 2013a). However, it is important to consider in the interpretation of results that
10 differences in biomass accumulated or height can also be attributed to shorter regeneration
11 times between treatments (i.e. vegetation at different successional stage). Irrespective, the
12 results observed here provide an overview of biomass dynamics and accumulation patterns
13 for similar moorlands.

14

15 *Grazing effects on above-ground biomass and height*

16 In contrast to our expectation (Hypothesis 2), sheep grazing had no effect on any above-
17 ground biomass variables (Total, *Calluna*, litter and bryophytes) or vegetation height.
18 Therefore, grazing does not reduce ecosystem production neither interacts with fire, thus
19 Hypothesis 2 is rejected. These results were not at all unexpected because the summer-only
20 sheep grazing intensity was very low in the 1960s (0.1-0.3 sheep ha⁻¹, Rawes and Welch,
21 1969), and this has been reduced as part of a conservation management plan twice over the
22 course of this study. The actual grazing pressure on a given spot on the site is a function of
23 both the absolute numbers on the reserve and the movements of sheep between vegetation
24 types to find more productive grassland communities (Rawes and Welch, 1969). In any case,
25 this study adds weight to the evidence that grazing removal in slow growing high-altitude
26 moorlands produce a dwarf shrub species dominance, especially *Calluna* (Hartley and
27 Mitchell, 2005).

28

29 *Calluna height and biomass accumulation patterns through time*

30 After prescribed burning the ecosystem recovers from a combination of resprouting by
31 *Calluna* and bryophytes (e.g. biomass and height increases) and the colonization of new
32 species including *Calluna* from seedlings and bryophytes from propagules (Lee et al., 2013b).
33 This is effectively known as the “reorganization and aggrading phases” for moorlands as
34 described by Bormann and Likens (1979) for forest. In this study, our results for *Calluna*

1 biomass and vegetation height showed these patterns since the absolute growth rates were
2 increasing over the first 8 and 4 years, respectively (maximum rates of 76.08 g m⁻² year⁻¹ for
3 biomass and 4.61 cm year⁻¹ for height). This is transformed into a rapid post-fire
4 accumulation of *Calluna* biomass and vegetation height. It is interesting to mention that the
5 AGR values might help to determine what kind of management is optimal in view of the
6 pasture availability based on growth. In this case, the greater biomass production will be
7 produced by burning rotations lower than 10-12 years, i.e. when AGR for biomass is
8 maintained at maximum values.

9 The main increase in total biomass resulted only from *Calluna* growth, with all other
10 components (biomass of litter, bryophytes, graminoids and other vascular plants group)
11 contributing in a very variable manner. Graminoids and other vascular plants constituted a
12 trivial component of the total biomass, whereas litter and bryophyte biomass stayed constant
13 over time since last burning. This result may be due to the practice of “cool burns” (discussed
14 at Harris et al., 2011a), as a consequence of high rainfall and the relatively moist vegetation
15 at Moor House, this produced relatively little damage to the underlying moss and litter layers,
16 or even some *Calluna* stems. However, further research is needed to confirm this hypothesis
17 through estimates of pre- and post-fire biomass.

18 The vegetation at Moor House achieved a modelled asymptote at 20 years for *Calluna*
19 biomass and 15 years for vegetation height, suggesting that after 20 years more or less
20 equilibrium conditions are reached. Essentially this is evidence to support the view that the
21 system was approaching to “steady-state” (Bormann and Likens, 1979). In this “steady-state”
22 any growth in the above-ground vegetation must be compensated either by respiration or
23 transfer into the peat. This is an interesting result since the time needed to reach the biomass
24 asymptote at Moor House (20 years) is much lower than more productive lowland heaths
25 where asymptotes are reached at more than 36-40 years (Chapman and Webb, 1978; Miller,
26 1979), or even in similar low-productive moorlands at Kerloch where an asymptote was
27 estimated at 25 years (Miller, 1979). It seems that if the management of this moorland is
28 focused on optimizing C fixation by means of biomass accumulation the fire rotational
29 interval should be around 20 years. Irrespective, the differences in biomass asymptotes
30 between these contrasting moorlands indicate the difficulties of managing sites using simple
31 prescriptions, instead of develop site-specific management plans.

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1 *Management implications*

2 The present work demonstrates that burning rotations lower than 20 years reduced the above-
3 ground biomass and vegetation height on this moorland compared to stands unburned for
4 more than 50 years. The modelled outputs suggest that an asymptote in *Calluna* growth and
5 vegetation height occurs at 20 and 15 years after fire, respectively. In the case of *Calluna*
6 biomass the asymptote is produced simultaneously as the longest rotation interval for this
7 moorland (long-rotation of 20 years). In contrast, no grazing effect was detected on biomass
8 or vegetation height in this plant community. Therefore, in order to maximize the C fixation
9 in similar moorlands, fire-return intervals should be around the *Calluna* accumulation
10 asymptote, i.e. 20 years since last fire. However, this return-interval could reduce the
11 component of some important peat-forming species such as *Sphagnum* and *Eryophorum* (Lee
12 et al., 2013a), which are favoured in 10-years rotation intervals.

13 Finally, the analogous accumulation curves for *Calluna* biomass and vegetation height
14 (same response to burning treatments) are an interesting result to assist in heathlands
15 management. Land managers usually use moorlands age or *Calluna* biomass to identify the
16 optimal time to burn (Harris et al., 2011a). However, since here *Calluna* biomass and
17 vegetation height were similar, the use of vegetation height could be a new faster way to
18 assess the optimal time to burn since it is more reliable and faster to measure by land
19 managers in the field. This approach was suggested by Harris et al. (2012) for Peak District
20 moorlands and they suggested using 25 cm height as a yardstick to determine when
21 prescribed burning should be applied to maintain diversity and an upper value of 40 cm as the
22 maximum that should be allowed otherwise there would be a predicted loss of species
23 diversity. The asymptote at Moor House was within this range (36 cm) suggesting that
24 vegetation height might be a useful tool for guiding when to implement prescribed burning.

25

26

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1 **Table 1.** Details of experimental treatments within each of the four replicate blocks for the
 2 Moor House study site, where post-fires biomass accumulation patterns were studied over
 3 each sub-plot (n=28). The treatment history of experimental plots description is included.
 4 Key to burning treatments: S=short-rotation; L=long-rotation; N=no-rotation post 1954/5,
 5 R=reference plots, unburned for at least 87 years.
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Main-plot treatments	Sub-plot treatments			
	Burning rotation	No of burns since 1954	Year of last burn	Years between sampling (2011) and last burn
Enclosed	N	1	1954/55	56/57
Enclosed	S	6	2006	5
Enclosed	L	3	1995	16
Grazed	N	1	1954/55	56/57
Grazed	S	6	2006	5
Grazed	L	3	1995	16
Grazed	R	0	1924?	Minimum 87

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1 **Table 2.** Effects of time since last burn on above-ground vegetation biomass (g m^{-2}) and
 2 vegetation height (cm) in the Hard Hill grazing and burning experiment at Moor House; the
 3 data presented are derived from a Mixed-effects modelling: The treatments were three
 4 burning rotations within the formal experiment, 5 years after last burn-short (S), 16 years
 5 since last burn-long (L) rotations plus no-burn after 1954/5 (N) compared to the reference
 6 plots (R) as intercept. Arithmetic means ($\pm\text{SE}$, $n=4$ plots, each with three sub-samples/plot)
 7 are presented along with the statistical estimates from the analyses.
 8

Variables	Mean \pm SE	Estimate \pm SE	t-value	p-value
Total Biomass (g m^{-2})				
R(\sim 90)	2223 \pm 201	2223.17 \pm 218.06	10.20	<0.001
N(56/57)	2079 \pm 144	-143.92 \pm 02.80	-0.71	0.496
L(16)	1593 \pm 119	-629.79 \pm 02.80	-3.11	0.013
S(5)	1198 \pm 165	-1024.83 \pm 02.80	-5.05	<0.001
<i>Calluna vulgaris</i> (g m^{-2})				
R(\sim 90)	705 \pm 73	704.95 \pm 82.61	8.53	<0.001
N(56/57)	808 \pm 80	103.13 \pm 104.02	0.99	0.347
L(16)	672 \pm 39	-33.30 \pm 104.02	-0.32	0.756
S(5)	60 \pm 16	-645.33 \pm 104.02	-6.23	<0.001
Litter (g m^{-2})				
R(\sim 90)	993 \pm 107	993.11 \pm 131.55	7.55	<0.001
N(56/57)	758 \pm 91	-235.59 \pm 156.64	-1.50	0.167
L(16)	801 \pm 105	-191.86 \pm 156.64	-1.22	0.252
S(5)	811 \pm 110	-181.68 \pm 156.64	-1.16	0.276
Bryophytes (g m^{-2})				
R(\sim 90)	522 \pm 98	522.54 \pm 137.05	3.81	<0.001
N(56/57)	513 \pm 183	-9.98 \pm 171.65	-0.06	0.955
L(16)	120 \pm 56	-402.78 \pm 171.65	-2.35	0.043
S(5)	319 \pm 73	-203.47 \pm 171.65	-1.19	0.266
Height (cm)				
R(\sim 90)	37.3 \pm 1.2	37.25 \pm 2.04	18.30	<0.001
N(56/57)	37.4 \pm 1.2	0.17 \pm 2.88	0.06	0.955
L(16)	35.0 \pm 1.3	-2.25 \pm 2.88	-0.78	0.455
S(5)	23.0 \pm 2.2	-14.25 \pm 2.88	-4.95	<0.001

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1 **Table 3.** Parameters for the non-linear mixed models trough time since last burning for
 2 *Calluna* biomass (g m⁻²) and vegetation height (cm) accumulation curves in the Hard Hill
 3 grazing and burning experiment at Moor House.
 4

Variable	Model selected		Estimate±SE	t-value	p-value
		Asym	795.87±80	10.08	<0.001
<i>Calluna</i> biomass (g m ⁻²)	Gompertz	b ₂	8.07±3.55	2.28	0.035
		b ₃	0.78±0.04	20.12	<0.001
		Asym	36.31±0.84	43.11	<0.001
Height (cm)	Logistic	xmid	4.09±1.64	2.49	0.015
		scal	1.97±3.65	0.54	0.591

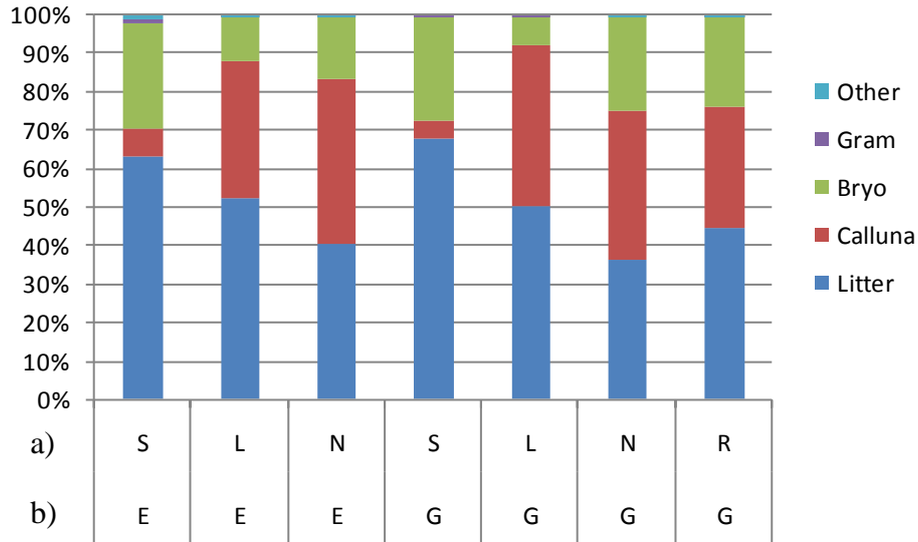
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1 **Table 4.** Comparison of total above-ground biomass data (g m^{-2}) from Moor House with
 2 literature values; np = information not presented.
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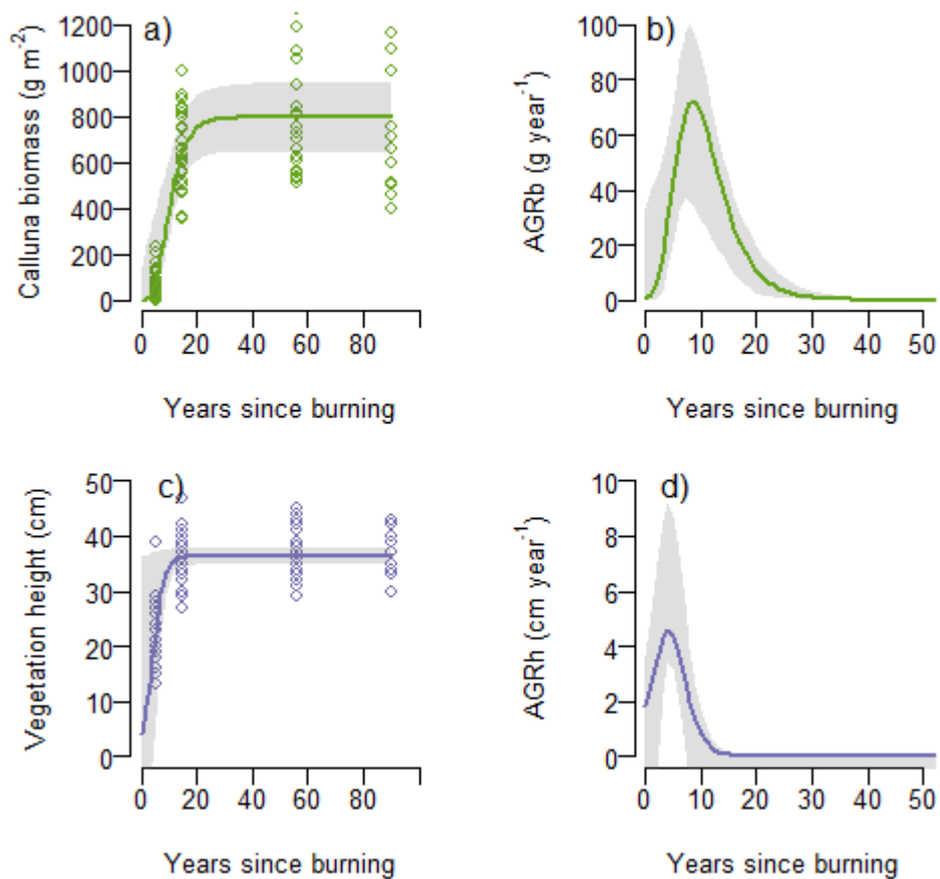
Location	Site	Elevation (m)	Stand Age (years)	Biomass (g m^{-2})	Reference
Peak District	Bamford	300-420	3-14	163±37-2138±72	Harris et al. (2011a)
			38	5364±569-8019±270	
Peak District	Broomhead	300-460	2-15	132±17-2279±190	
			40-50	6540±844-7114±701	
Peak District	Howden	272-540	2-15	267±9-2262±60	
			50	5577±470-6401±596	
Peak District	Midhope	270-480	3-15	334±11-2673±90	
			40	5241±660- 10024±337	
Peak District	Snailsden	350-470	3-16	517±39-2375±185	
			40	5250±662-6507±549	
Moor House	Hard Hill Burning Experiment	600-632	90	2223±201	This study
Moor House	Hard Hill Burning Experiment	600-632	56/57	2079±144	This study
Hexham	Blanchard moor	305	10	1920	Robertson and Davies (1965)
Kincardineshire	North Cairn o' Mount	274	15	2930	Robertson and Davies (1965)
Kincardineshire	Kerloch Moor	140-280	25	1840	Kayll (1966)
Teesdale	np	290-850	6	600	Bellamy and Holland (1966)
Teesdale	np	290-850	14	2000	Bellamy and Holland (1966)
Dorset	Poole Basin	90	?	1820	Chapman (1967)
Dartmoor		320-340	?	2000	Chapman (1967)

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Fig. 1. Distribution of the five main fractions of biomass (%) in the Hard Hill grazing and burning experiment at Moor House. The treatments were: a) burning treatments: short-rotation (S), long-rotation (L), no-burn after 1954/5 (N) and reference plots (R); b) grazing treatments: enclosed (E) and grazed (G). The colour codes were: blue=Litter, red=*Calluna*, green=Bryophytes, purple=Graminoids and sky-blue=Other vascular plants.



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Fig. 2. Observed and predicted values from non-linear mixed effects models through years since last burning in the Hard Hill Grazing and Burning experiment at Moor House. Relationship between (a) *Calluna* biomass and years since burning modelled by a Gompertz model, (b) absolute growth rate (AGRb) on a time basis for *Calluna* biomass, (c) vegetation height and years since the last burning modelled by a logistic model, and (d) absolute growth rate (AGRh) on a time basis for vegetation height. Grey bands indicate 95% confidence intervals.