

**Effects of long-term removal of sheep-grazing in a series of British upland plant communities:
insights from plant species composition and traits**

Josu Alday^{a,b,c}, John O'Reilly^d, Rob J. Rose^e, Rob H. Marris^a

^aSchool of Environmental Sciences, University of Liverpool, Liverpool L69 3GP, UK,

^bJoint Research Unit CTFC – AGROTECNIO, Av. Alcalde Rovira Roure 191 E25198 Lleida, Spain,

*^cDepartment of Crop and Forest Sciences, University of Lleida, Av. Alcalde Rovira Roure 191
E25198 Lleida, Spain,*

^dPtyxis Ecology, Railway Cottages, Lambley, Northumberland CA8 7LL, UK,

^eCEH Lancaster, Centre for Ecology & Hydrology, Bailrigg, Lancaster LA1 4AP, UK.

Marris calluna@liverpool.ac.uk

Alday josucham@gmail.com

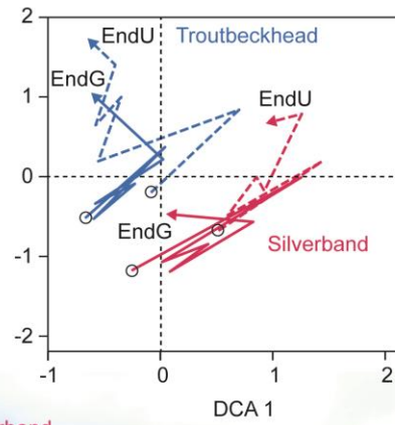
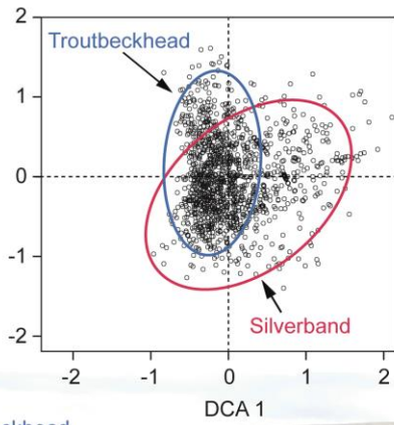
O'Reilly john@ptyxis.com

Rose robrose73@icloud.com

Corresponding author Marris: calluna@liverpool.ac.uk

Running title:

Long-term effects of removing sheep-grazing



Troutbeckhead

DCA 1



Silverband

DCA 1



Abstract

Environmental and management pressures are considered a threat for preserving plant communities worldwide. Identification of long-term impacts of changing management practices on plant community composition must, therefore, be a major priority to ensure improvement in conservation value. Land abandonment/wilding is one topical areas where there is little available information on long-term impacts. To address this, here, we describe plant species compositional change over relatively long-time scales in a range of four British upland communities (high-level grasslands, intermediate grasslands, blanket bogs and high-level bogs) and its relationships with a series of life-history and plant trait variables. Our aims were to determine whether the business-as-usual sheep grazing practice was maintaining these communities, and if grazing was to be removed as part of abandonment/wilding strategies, would there be a conservation benefit. We used a series of long-term experimental grazing-exclosure studies at twelve sites (>40 years) on the Moor House National Nature Reserve in northern England. Each site consisted of paired plots; i.e. sheep-grazed versus ungrazed. Our results showed that there was relatively little change in dominant plant species with most change occurring within sub-dominant species. Similarly, different temporal responses in plant species richness were detected between the experiments (richness was reduced in three experiments, only one increased). In any case, the vegetation temporal trajectories were moving in similar direction in grazed and ungrazed plots at most sites. Interestingly, blanket bog experiments showed a clear compositional convergence for both grazed and ungrazed plots, although, the between-site differences related to a combination of elevation and past burning testament. Finally, the bryophytes (especially liverworts) and lichens were the groups that contributed most to the reductions in species richness. Moreover, implementation of a no-stock grazing strategy under a land abandonment/wilding approach will not bring about much plant change in the short-term in the plant communities studied here.

Key-words:

Moorland

Blanket bog

Upland grassland

Ordination

Exclosures

1. Introduction

Land abandonment is a topical and contentious issue in northern Europe with a forecast that large areas of agricultural land will be affected in the next 20-30 years (Renwick et al., 2013). Most of this land is predicted to be in upland areas and in more northerly latitudes (Renwick et al., 2013; Kapfer and Popova, 2020). Much of the land likely to be affected is grazing land which plays an important role in global ecosystem functioning and human well-being (O'Mara, 2012; Steinfeld et al., 2006). However, many grassland and shrublands maintained by grazing are over-grazed, which is a key driver reducing productivity and sustainability (Conant, 2010; O'Mara, 2012) and altering plant communities, soil water and nutrient availability (Conant 2010; McSherry and Ritchie 2013). It is surprising, therefore, given this importance that a recent review suggested that few studies linked livestock pressures to biodiversity impacts (Kok et al, 2020). As most of the land is likely to be abandoned in northern Europe has a High Nature Value (Renwick et al., 2013), there must be scope to devise ecological restoration strategies to improve their conservation value (Xie et al., 2020).

One positive approach to land abandonment that is currently gaining momentum is re-wilding/wilding (henceforth referred to as wilding), which in its simplest form is the removal/reduction in livestock grazing with the intention that this will improve the conservation benefit of grassland and shrublands (Sandom et al., 2013; Marrs et al., 2018, 2020). This belief is inherent in the “wet-desert” description of the British uplands by Frank Fraser Darling (1955), who argued that sheep-grazing, in particular, was very detrimental to British upland plant communities, reducing the species diversity. Indeed, these uplands have recently been described as “sheep-wrecked” by Monbiot (2013). However, since Darling’s time evidence from wide-scale surveys have shown that factors other than over-grazing can damage upland vegetation including acid deposition, nitrogen deposition and climate change (Mitchell et al. 2017). The situation is even more complex as recent evidence suggests that acid deposition has declined from the latter parts of the twentieth century and plant species richness has responded positively, even though there have also been an increase in dominant plant species (Mitchell et al., 2018), and biotic homogenization (Ross et al., 2012; Pakeman et al. 2019). Part of the difficulty in managing upland

habitats is that the slow dynamics of these communities make it extremely difficult to predict the effects of management decisions taken. Clearly, if the reduction of grazing policy (i.e. wilding) is to be implemented wholesale, conservation managers need to know the likely outcomes of grazing removal on the long-term ecosystem dynamics.

In essence, before implementing new policies land managers need to know how ecosystems will respond to the changed management, given that these changes will interact with the other changing environmental drivers (Lindenmayer and Likens, 2018). It is also useful to understand the ecological requirements of the plant component species in order to describe the impacts of management through time (Alday et al. 2013a). Therefore, ideally, any conservation management should be validated using evidence derived from long-term manipulative experiments (Pullin and Knight, 2009; Sutherland and Wordley, 2018). Unfortunately, to set such experiments up in advance of their need obviously requires considerable foresight.

One series of such long-term, experimental grazing enclosure studies were set up between 1953 and 1967 on a range of common plant community types of upland habitats at the Moor House National Nature Reserve in the North Pennines of England (Milligan et al., 2016, 2018). At each study area, two paired-plots were established, one fenced to exclude sheep grazing, the other where sheep grazing was allowed. These study areas collectively encompass a large proportion of British upland plant communities, including those dominated by dwarf-shrubs, grasses and sedges, and soil types ranging from deep blanket peat through to brown-earth soils (Rawes and Welch, 1969). Moreover, these different plant communities experience different grazing pressures (Rawes and Welch, 1969), and these pressures have themselves changed through time, reducing from ca. 4.4 to ca. 2 sheep ha⁻¹ across the entire grazing unit in 1972, and a further reduction to ca. 1 sheep ha⁻¹ in 2001. The studies, therefore, compare no sheep grazing to the “business-as-usual” grazing pressure across the North Pennines reserve over a considerable timescale (Milligan et al., 2016, 2018; Marrs et al., 2018). As these upland plant communities have been traditionally considered degraded by overgrazing (McGovern et al., 2011), conservation managers expected that grazing reductions would lead to an improvement in the conservation value of the

vegetation, by increasing diversity including bryophytes and lichens, and by reducing the abundance of unpalatable species (Marrs et al., 2020).

From these long-term studies, some preliminary results have been published on species change in the individual study areas up to the early 2000s, covering experimental periods of between 24 and 45 years (Rawes, 1981, 1983; Milligan et al., 2016). However, as vegetation in upland areas tends to respond slowly because of adverse climatic conditions there is a need to obtain a much longer-term view of plant community change in response to grazing to inform conservation policies such as wilding. For example, Chen et al. (2020) have recently shown that information derived for short-term studies can be misleading when viewed in the light of longer-term data. Here, therefore, we report additional data (up to 2016) for vegetation change in eight of the nine experiments used by Milligan et al. (2016) plus data from four additional exclosures reported by Milligan et al. (2018) on the Moor House grazing unit, representing experimental periods of 48-63 years for twelve experiments composing four different plant community types (Milligan et al. 2016, 2018).

Our aims were first to describe the long-term changes in plant species composition in the four plant community types comparing grazed versus ungrazed plots and then to test two hypotheses: (a) grazing would maintain the plant community composition at a quasi-equilibrium state through time, and, (b) rewilding by grazing removal would improve the conservation value of the vegetation relative to the grazed situation. To test both hypothesis we analyzed the change in both (1) species composition to provide relative measures of the differential effects on sheep grazing *versus* no sheep grazing on the different types of upland communities across time, and (2) plant traits to provide additional information on potential indicator groups that would inform conservation managers about likely drivers of vegetation change (Pakeman, 2004; de Bello et al., 2010). Thus, we have assessed the long-term effectiveness of sheep grazing for preserving the original communities and the removal of livestock in rewilding schemes as part of an ecological restoration strategy. These results will help define practices that prevents future negative impacts on upland habitats. Based on these findings, we put forward recommendations for

effective conservation and management of British uplands plant communities and other areas of northern Europe where stock removal may be implemented (Renwick et al., 2013).

2. Methods

2.1. Study sites

Moor House NNR is situated in northern England, straddling the Pennines (Appendix A: Fig. S1). The site was established in 1952 as an experimental reserve where long-term manipulative studies could be implemented. Between 1953 and 1967 a series of grazing exclosures were set up at 12 sites across the entire reserve. The sites covered a range of the most common vegetation types that occur in the North Pennines and across the UK; i.e. from relatively productive *Agrostis-Festuca* grassland on brown-earth soils at the near-neutral end of the soil spectrum through grasslands dominated by *Festuca ovina* or *Nardus stricta*, to *Juncus squarrosus*, *Eriophorum* spp. and dwarf shrub *Calluna vulgaris*, *Erica tetralix* or *Empetrum nigrum*-dominated vegetation on acidic blanket bog (least productive). A detailed description of Description of the twelve-paired monitored sheep-grazing and exclosure plots that are used in this work is included in Appendix A: Table S1.

2.2. Experimental design

Here, we used the complete dataset containing information from the 12 sites, each with paired plots; one being fenced to exclude sheep and the other left open to allow free-range, sheep-grazing (Rawes, 1981, 1983, Marrs et al., 1988). However, the 12 sites can be viewed as four long-term, structured-experiments based on the vegetation type that dominate each site (see Milligan et al., 2016 for further details). These are outlined in Appendix A: Table S1, summarized as: (1) High-level (*Agrostis/Festuca*) grasslands, on the more-base-rich soils with three replicate sites (Hard Hill, Little Dun Fell, Knock Fell), starting in 1955/6 (Rawes, 1981); (2) Intermediate (*Nardus stricta/Juncus squarrosus*) grasslands, with two replicate sites (Cottage Hill, River Tees), starting in 1967; (3) Blanket bog (*Calluna-Eriophorum*) with five replicate sites (4 Hard Hill Burning Blocks (HHA, HHB, HHC, HHD) and Bog Hill), starting between 1953 and 1955, and (4) High-level bog, *Eriophorum*-dominated, with two replicate sites (Silverband, Troutbeck Head) starting in 1966 (Rawes, 1983).

Throughout, point-quadrats have been used to measure vascular plant, bryophyte and lichen species abundance. To keep long-term recording structure, in all experiments the point-quadrat frame was positioned using a permanently-marked reference system within the plot with the sampling positions selected randomly at the outset. On many occasions, height-stratified pins (0-10 cm, 10-20 cm, 20-30 cm and >30 cm) were used to record vascular plants to provide information on canopy composition. However, the exact way in which the pin frame was used has varied between experiments and on different sampling occasions. For example, not all pins were sampled on every occasion, or only a selection of pins was sampled on a height-stratified basis. Full details of the pin frame technique are given in Marrs et al. (1986) and Milligan et al. (2016) and all data are available in Rose et al. (2018, 2020). As a consequence of these slight monitoring differences, a detailed pre-analysis data preparation was needed which is described in detail in Appendix A.

2.3. Data Preparation and analyses

All data preparation and analyses were performed in the R Statistical Environment v.3.5.2 (R Core Team, 2019) using the ‘vegan package’ (Oksanen et al., 2019).

First, from the clean abundance-score database a series of variables were derived, including three diversity variables, species richness, Shannon-Wiener and Simpson’s diversity indices and 41 variables based on taxonomy, life-form and traits (details in Appendix A: Table S2). For taxonomic and life-form variables, the total number of hits per class was calculated; for traits, abundance-weighted mean values were calculated (Milligan et al., 2016). The three diversity variables were calculated using the ‘*specnumber*’ and ‘*diversity*’ functions of the ‘vegan’ package (Oksanen et al., 2019).

The species datasets for each of the four experiments were analyzed independently. Three datasets were Hellinger-transformed (‘*decostand*’ function) followed by Detrended Correspondence Analysis (DCA) using the ‘*decorana*’ function. No transformation was used for the high-level bog dataset because transformation produced conflicting eigenvalue ranks. Species that were present only once were removed before the analysis. The amount of variation explained by each axis was expressed as a percentage of the

total variation explained in the data, estimated using Økland's (1999) method. The correlations between the DCA ordination axes and the three diversity measures were then calculated using the '*envfit*' function with 9999 permutations; as the three variables showed similar significant results ($P < 0.001$) and arrow directions only species richness data are discussed further. The distribution of sites within each experiment was visualized in ordination space as standard-deviational bivariate ellipses (SD-ellipses, 95% confidence limits) using the '*ordiellipse*' function; the trajectories of the grazed and ungrazed treatments were then plotted through time for each site. The area of each of the ellipse within the ordination space was considered as a measure of the size of the overall species pool.

The species abundance data was thereafter converted to a distance matrix using the '*vegdist*' function and the Bray and Curtis index, and the significance of each of the 41 trait variables were then assessed using the '*adonis*' function with 9999 permutations stratified by site and grazing treatment (i.e. restricting permutations to maintain special dependence). Those significant variables were then analyzed using redundancy analysis (function '*rda*') after standardization ('*decostand*' function) with the interaction of grazing treatment x time as the constraining variables (EtG = grazing through elapsed time; EtU = ungrazed treatments through time). Summary statistics from all DCA and RDA analyses are presented in Appendix A: Table S3.

3. Results

3.1. Plant community response to removal of sheep grazing

3.1.1. High-level grasslands

The three high-level grasslands sites were separated along DCA axis 1 (40% of variance), with Little Dun Fell and Hard Hill towards the negative end and Knock Fell at the positive end with Hard Hill being more or less subsumed within the Little Dun Fell community space (Fig. 1b). Hard Hill had the smallest species pool, Little Dun Fell was intermediate and Knock Fell had the largest with the deviational ellipses being 1.58, 4.62, 6.96 units respectively. However, most of the within-site variation is described across axis 2 (major ellipse axes for all sites; 25% of variance). The site separation was linked with a species gradation

along DCA axis 1 from an *Avenella flexuosa*, *Carex bigelowii*, and *Nardus stricta* community at the negative end (Fig. 1a) to *Agrostis canina*, *A. capillaris*, *Anthoxanthum odoratum*, *Carex caryophylla*, *Deschampsia cespitosa*, *Festuca rubra*, *Luzula campestris/multiflorum*, *Achillea millefolium*, *Rumex acetosa* and *Thymus polytrichus* at the positive end. The bryophytes showed a similar gradation (Fig. 1a) with *Dicranum fuscescens*, *Polytrichum strictum* and the lichens *Cladonia uncialis* and *C. portentosa* at the negative end through to *Hylocomium splendens*, *Lophocolea bidentata* and *Thuidium tamariscinum* at the positive end. In contrast, Axis 2 (25% of the variance) reflected a bryophyte/lichen rich community at the negative end with vascular plants at the positive end (Fig. 1a), these effects were more marked in Little Dun Fell and Knock Fell. Interestingly, species richness was correlated positively with axis 1 and negatively with axis 2 indicating that Knock Fell quadrats with vascular, lichen and bryophytes were the more diverse ones (Fig. 1a).

The three high-level grassland sites also showed marked differences in successional trajectories between grazed and ungrazed plots (Fig. 1c-e). Both grazed and ungrazed plots at Hard Hill were compositionally stable, fluctuating slightly around the same ordination area with temporal vectors of similar magnitude (Fig. 1d). In contrast, both the Little Dun Fell and the Knock Fell sites showed different compositional trajectories between the grazed and ungrazed plots. At Little Dun Fell the successional trajectory of the ungrazed plots moved in the positive direction of axis 2 towards increased vascular plants and reduced bryophytes/lichens; whilst the grazed plots showed a more erratic trajectory ending also on positive side of axis 2, although some distance from the ungrazed plots (Fig. 1c). At Knock Fell the compositional trajectories of ungrazed plots moved towards the positive end of axis 2 with increased graminoid cover and reduced bryophytes/lichens, while the grazed plots were moving in a positive direction on axis 1 with increasing graminoids (Fig. 1e). At all sites the trajectories were moving in the opposite direction to species richness.

3.1.2. Intermediate grasslands

The two intermediate grassland sites were significantly separated along DCA axis 1 ($p < 0.05$; 55% of variance); River Tees site was located at the negative side and Cottage Hill on the positive side of DCA

axis 1 with compositional differences between both (Fig. 2b). The Cottage Hill site deviational ellipse covered almost double the ordination space of the River Tees site, 3.13 and 1.49 units respectively, indicating a greater species pool (species heterogeneity), although the main within-sites variation was represented along DCA axis 2 (major ellipses axes; 20% of variance). The site compositional differences were also translated into species gradients (Fig. 2a) from *Agrostis canina*, *A. capillaris*, *Deschampsia cespitosa*, *Nardus stricta* and *Hypnum jutlandicum* at the negative end of DCA axis 1 (correlated with the river River Tees site) through to *Avenella flexuosa*, *Eriophorum angustifolium* and *E. vaginatum* at the positive end (correlated with river Cottage Hill site), Bryophytes moved from *Hypnum jutlandicum* through to *Aulacomnium palustre*, *Barbilophozia floerkei*, *Calypogeia muelleriana*, *Ptilidium ciliare* and *Sphagnum capillifolium* (Fig. 2a). DCA Axis 2 showed a gradation from a *Juncus squarrosus*-*Luzula campestris/multiflora* and bryophyte-dominated community with *Barbilophozia floerkei*, *Calypogeia muelleriana*, *Lophocolea bidentata*, *Ptilidium ciliare*, *Plagiothecium undulatum* at the negative end and *Deschampsia cespitosa*, *Eriophorum vaginatum*, *Pleurozium schreberi*, *Rhytidiadelphus squarrosus*, *Sphagnum capillifolium* at the positive end. Species richness was mainly correlated negatively with axis 2 (Fig. 1a).

Both sites showed similar successional trajectories for both grazed and ungrazed plots converging towards the upper central ordination space where *Pleurozium schreberi*, *Rhytidiadelphus squarrosus*, *Sphagnum capillifolium* was located. At the River Tees site, both plots moved in a positive direction on DCA axis 2, with the grazed plots moving less far than the ungrazed plot (Fig. 2c). At Cottage Hill, both plots also moved in a positive direction on DCA axis 2, but the ungrazed plot trajectory was displaced further to the right in a positive direction (Fig. 2c). At both sites the trajectories were moving in the opposite direction to species richness.

3.1.3. Blanket bogs

The five blanket bogs sites were superimposed around the center of the ordination with the deviational ellipses located along the entire gradient of DCA axis 1 (major ellipse axes, 39% of variance). This indicates that although there is a lack of compositional differences between sites (main species are shared

at the ordination center, Fig. 3a), the DCA axis 1 showed similar within-site differences for all experiments in response to changes in bryophytes/lichens (i.e. main successional trends). In contrast, there was a slight separation of experiments up DCA axis 2 (24% of variance; in order from lowest to highest Bog Hill through Plots HHC > HHB > HHA to HHD; Fig. 3b) essentially reflecting a reduction in species richness, at least partly linked to increasing elevation. The analysis of deviational ellipses areas showed that the Bog Hill ellipse was at least half that of the Hard Hill sites at 1.64 units compared to 3.76, 3.72, 3.33 and 5.01 units for HHA, HHB, HHC and HHD respectively, which may reflect differences in sampling intensity. Interestingly, the trajectories of all sites moved in a reverse direction along DCA axis 1, which similar movements and direction for grazed and ungrazed plots towards communities with more bryophytes (Fig. 3c-g).

The main species in these blanked bog sites, such as *Calluna vulgaris*, *Dicranum scoparium*, *Eriophorum vaginatum*, *Rubus chamaemorus*, were centered around the origin and the main first gradient reflected to a large extent successional changes in distribution of bryophytes and lichens (Fig. 3a). The bryophytes at the negative end of DCA axis 1 were *Hypnum jutlandicum*, *Plagiothecium undulatum*, *Pleurozium schreberi* through to *Campylopus flexuosus*, *Calypogeia muelleriana*, *Cephalozia bicuspidata*, *Lophozia ventricosa* and *Pohlia nutans* at the positive end. DCA axis 2 reflected a gradient from *Cephalozia connivens*, *Cladonia portentosa*, *Empetrum nigrum* and *Sphagnum capillifolium* at the negative end to Green algae, *Calluna vulgaris*, *Campylopus flexuosus*, *Eriophorum angustifolium* and *Kurzia pauciflora* at the positive end (Fig. 3a). Species richness was mainly correlated negatively with axis 2 (Fig. 3a).

3.1.4. High-level bogs

The two high-level bogs occupied relatively similar positions in the DCA ordination center with considerable overlap, although DCA axis 1 (28% of variance) showed that compositional differences between plots of both sites were caused by species richness (negatively correlated, Fig. 4a). The deviational ellipse for Silverband occupied more than double the ordination space of Troutbeckhead, 4.46 compared to 1.61 units (Fig. 4b), with the major ellipse axes diagonal to both axes. The DCA axis 1

gradient shows *Avenella flexuosa*, *Calluna vulgaris*, *Empetrum nigrum* and *Vaccinium myrtillus* among the vascular plants at the negative end and *Carex nigra*, *Narthecium ossifragum*, *Rubus chamaemorus* and *Trichophorum cespitosum* at the positive end (Fig. 4a). The bryophytes were mainly placed at the negative end of axis 1, and placed in the same general direction as increasing species richness (Fig. 4a). DCA Axis 2 (27% of the variance) showed a gradient from a community dominated by *Festuca ovina*, *Carex nigra*, *Vaccinium myrtillus* and *Rubus chamaemorus* through to one dominated by *Avenella flexuosa*, *Calluna vulgaris*, *Empetrum nigrum*.

The successional trajectories of grazed and ungrazed plots for both sites started in similar areas of the ordination. At Troutbeckhead both treatments moved in a negative manner on axis 1 and a positive one on axis 2, moving towards *Avenella flexuosa*, *Calluna vulgaris* and in the same direction as increasing species richness (Fig. 4c); the ungrazed plot moved slightly further than the grazed one. (Fig. 4c). At Silverband, the successional trajectories grazed trajectory moved positively on axis 1 and was fairly close to the grazed Troutbeck Head plot at the end of the study period. The ungrazed plot moved marginally in a positive direction on axis 2 (same as the other sites) but also moved marginally positively on axis 1, and in an opposite direction to species richness (Fig. 4c).

3.2. Trait and life-history responses to removal of sheep grazing

In many respects the RDA analyses showed similarities between the four experimental groups in that the general effect of elapsed time in species composition was in a positive direction on axis 1 (Fig. 5a). For three experiments this was in the opposite direction to species richness, the exception was the high-level bogs where species richness was correlated with successional trajectories of grazed and ungrazed plots (Fig. 5a). These results agree with the previous DCA results reported above. All three climatic variables were associated with the vegetation temporal trajectories to some extent; on both grasslands and high-level bog they were associated with the ungrazed plot whereas on the blanket bog they were linked to the grazed plot. The Ellenberg values were also correlated with the vegetation temporal trajectories, however, in both grasslands and the high-level bog Ellenberg values were correlated with ungrazed trajectory,

while, in the blanket bogs they were with the grazed one (Fig. 5b). The grazing index showed mixed effects being placed towards the negative end of the vegetation temporal trajectories in the intermediate grassland and the blanket bog, but with the grazed treatment in the high-level grasslands and high-level bog (Fig. 5b).

The taxonomic groups showed that liverworts and lichens were at the negative end of the four successional trajectories and vascular plants and mosses at the positive end (more recent samplings); except the intermediate grasslands where vascular plants, grasses and herbaceous species were found earlier in the succession (Fig. 6a). In the high-level grasslands the vascular plants were mainly graminoids/Poaceae and herbaceous species (Fig. 7a). In terms of clonality, there was a tendency for an increase in species with little vegetative spread in all experiments, plus species with shortly-creeping rhizomes in the high-level grasslands, extensively-creeping species in the intermediate grasslands, blanket bog and high-level bog and turf-forming species in both blanket bog and high-level bog.

The vascular plant life-histories showed some similarities in three experiments (intermediate grasslands, blanket bog, and high-level bog) with hemi-cryptophytes placed at the negative end of the successional trajectories (older monitoring) and chamaephytes nearer the positive end (most recent samplings, Fig. 7a). No other life-history forms were significantly explaining species successional trends in these experiments. In the high-level grasslands, therophytes and chamaephytes were placed at the early part of the succession and hemi-cryptophytes in the later stages, geophytes being intermediate. The bryophyte life-histories showed a transition along the successional gradients with smooth and rough mats (high-level grasslands only), tufts and turfs at the negative end (older monitoring) and cushions, rough mats, turfs (blanket bog only) and species with little vegetative spread at the positive end (most recent monitoring, Fig. 7b).

4. Discussion

In this study, we compared the long-term (48-63 years) change in plant species composition of vegetation in the British uplands under experimental conditions comparing the effects of two grazing pressures, free-

range sheep grazing versus no grazing pressure. We also investigated how the plant traits and life-history information might be responsible for detected differences, and hence could be used as potential indicators to inform conservation managers. Surprisingly, our results showed that the first hypothesis, related to quasi-equilibrium state of plant community composition maintained by grazing, was partially accepted because only two sites (Hard Hill high-level grassland, Silverband high-level bog) showed either cyclic or little change through time in vegetation composition of the grazed sites; all others experiments showed some directional change through time. Similarly, the second hypothesis, related to the positive effect of rewilding by grazing removal over vegetation composition, was also partially accepted because most sites showed a greater directional change of ungrazed plots than their grazed counterparts, nevertheless, this greater directional change was usually accompanied by a reduction in species diversity.

Our results did, however, highlight five important points. First, there has been no wholesale change in the composition of the dominant plant species (Grime 1998) even though they have fluctuated in abundance (*sensu* Miles 1979) during the last 48-63 years. However, sub-dominant species (Grime, 1998) have fluctuated and/or declined in abundance. Thus, part of the compositional differences were caused mainly by change in these sub-dominant species, some of them being more susceptible to management practices and environmental changes than the dominant ones, and this will obviously hamper their conservation. These results are in accordance with recent studies on synchrony and stability which highlighted that the dominant plant species was the main driver of community stability rather than species richness (Valencia et al., 2020). Second, different temporal responses in species richness were detected between the experiments which were dependent on species pools of the surrounding community types. For example, in three of the four experiments (high-level and intermediate grasslands and blanket bog) there has been a reduction in species richness through time, which is in agreement with successional trends described previously (Milligan et al., 2016). However, in the blanket bogs species richness was orthogonal to the trajectories and this may, in part, reflect differences in past site histories (legacy effects; Muirinen et al. 2019), Troutbeckhead being intact and Silverband recovering from frequent erosion damage (Rawes, 1983).

Third, the temporal trajectories for both grazed and ungrazed plots were moving in similar directions in most experiments (similar successional trends *sensu* Alday et al. 2011). The similarity of successional trends in each site between both plots was expected as local species pools are identical for both of them, and grazing was reduced across the experiment. The exception is the high-level grasslands, where site-dependent different responses were found between the grazed and ungrazed plots. For example, the species composition at Little Dun Fell changed the most through time with both grazed and ungrazed plots moving in similar directions comparable to previous sites, in contrast, Knock Fell showed an intermediate response with the ungrazed plot showing a similar trajectories to Little Dun Fell, but the grazed plot moving erratically less, whereas Hard Hill experiment showed remarkable stability with a small movement of both plots through time. It seems that these grasslands are more heterogeneous ecosystems than supposed, in view of these different successional responses (Seabloom et al. 2005).

Fourth, all of the blanket bog sites showed more or less similar trajectories for both grazed and ungrazed plots (compositional convergence *sensu* Alday et al. 2011), with site differences related to a combination of (a) elevation with the lowest Bog Hill being more species-rich than the higher-elevation Hard Hill plots, or (b) a reduction in species diversity in the Hard Hill plots associated with the burning treatment in 1954/55 (Milligan et al., 2018). Unfortunately, it is impossible to disentangle the real cause. However, the similarity in successional trends between both grazing treatments suggests that the initial effects of grazing exclosure has disappeared through time. Finally, where a reduction in species richness was detected, this was mainly due to a reduction in bryophytes, especially liverworts, and lichens. This is a common feature in the British uplands over the last century, being detected in a range of large-scale, repeat surveys (Britton et al., 2009a,b, 2018), almost certainly as a result of elevated pollutant impacts and/or increase vegetation growth (see below).

It is also important to consider those species that are producing the compositional changes through time. In the intermediate grasslands and two of the high-level grasslands, although the community composition in both grazing treatments were moving in the same direction with a general trend towards domination by graminoids, different dominants were detected in each experiment; *Avenella flexuosa* and

Carex bigelowii at Little Dun Fell, *Agrostis capillaris*, *Deschampsia cespitosa* and *Festuca rubra* at Knock Fell, *Deschampsia cespitosa* at River Tees and *Eriophorum vaginatum* at Cottage Hill. The intermediate grasslands were considered transitional between grassland and blanket bog communities by Rawes (1981), however, these new data suggest that Cottage Hill plant composition is moving towards blanket bog, while River Tees plant composition is moving towards a grassland. Similarly, on the blanket bogs and high-level bogs, *Calluna vulgaris* and *Empetrum nigrum* increased through time along with the pleurocarpus mosses *Hypnum jutlandicum* and *Pleurozium schreberi*. These results are consistent with those from wider-scale, repeat surveys (Ross et al., 2012; Pakeman et al., 2019; Mitchell et al., 2017, 2018), indicating the importance of species pool heterogeneity in shaping grasslands composition (Questad and Foster 2008).

In relation to successional speed, in both grassland experiments and the high-level bogs, the ungrazed plots moved a greater distance than the grazed ones during the same time frame, indicating greater compositional responses. This suggests that grazed impacts in these communities may be associated with increasing biomass and the reduction in light impinging on the bryophytes (van der Wal et al., 2005). Again, there were two exceptions, the first was at the high-level Hard Hill grassland, where this site showed a more or less-stable pattern through time with no evidence of the temporal trajectories at the other two sites. The other exceptions were the blanket bog; all sites showed similar trajectories through time but no consistency between the distance travelled between the grazed and ungrazed treatments. Part of the reasons for these inconsistencies might be because of the differential grazing patterns across the reserve, in part controlled by interactions between the underlying geology/soil types interacting with the way the sheep are managed (by allowing them to freely range over a large area of differing vegetation types). The high-level grasslands occur on acid brown-earths and the other communities are on either peats or soils with a high surface organic content. The grazing unit is a "Common" under English law and the sheep are put onto the moor predominantly from the south-western edge of the reserve, where they gravitate to the best grazing (high-level grasslands) and when forage on these grasslands is depleted, they move to less-nutritious vegetation (Rawes and Welch, 1966; 1969). Hard Hill, is some distance (2.5 km)

from the other high-level grasslands, it is separated by blanket bog, and had a much reduced sheep density in the early 1960s, averaging 2.1 sheep ha⁻¹ over the season with an estimated annual herbage removal of 5.5 kg DM ha⁻¹ compared to 3.3 and 4.2 sheep ha⁻¹ and 7.4 and 7.6 kg DM ha⁻¹ at Little Dun Fell and Knock Fell respectively (Welch and Rawes, 1964). In a similar vein, the Hard Hill blanket bog sites have a low sheep grazing density (Table 1), but they too are much closer to sheep entry point/good grazing than Bog Hill. However, these differential grazing pressures were measured in the 1960s before the two reductions in overall sheep numbers (Welch and Rawes, 1965, 1966; Rawes and Welch, 1966 1969), but we would predict that the grazing pressure on the high-level grasslands would be reduced less than that of the blanket bogs because the sheep would tend to concentrate on them as the grazing there is most nutritious (Marrs et al., 2020). The importance of size and spatial distribution of species-rich grasslands within a grass-heath matrix has recently been shown to influence grazing impacts where red deer (*Cervus elaphus* L.) were the main herbivores (Moore et al., 2015). Here, the correlation of the grazing index with the grazed treatment in the high-level grasslands and high-level bogs confirms these hypotheses as these communities are closest to the sheep entry points and should have a greater grazing pressure. However, further studies on grazing patterns on these experiments are needed to confirm these predictions.

The analysis of the changes in plant traits provided to us some insight into the potential mechanisms involved in the successional trends and in the responses of grazed and ungrazed plots. For example, the positive association of all climate variables with vegetation temporal trajectories suggest that the species are responding to a warmer and wetter climate of the last years, which has been suggested as a major driver of vegetation change in the Scottish Highlands (Britton et al., 2017; Mitchell et al., 2018). At the same time, all the Ellenberg indices were also significantly correlated with the vegetation temporal trajectories. This suggest that through time there has been an increase in species that grow best in less acidic, more fertile, wetter and with more light habitats (Fig. 5b). This provides further tentative evidence for the suggestion damage to plant communities caused by interactions with atmospheric pollution are beginning to be reversed (Rose et al., 2016; Mitchell et al., 2018). Atmospheric SO₂ concentrations, a major cause of acidification since its peak in the 1970s had been reduced by 90% in 2008 and N

emissions have also reduced over this period, but only by 50% (ROTAP, 2012; Monteith et al., 2016).

The continuing increase in weighted N values suggests that elevated N may still be having an impact over these habitats. We accept that these conclusions are speculative and require further work is needed to unravel the exact mechanisms involved.

4.1. Limitations of this investigation

Like any long-term study, the experiment could have been improved with hindsight. Few of the experiments had appropriate baseline monitoring although all treatments were set up on “uniform vegetation” (Rawes and Hobbs, 1979). In some cases, one plot had a baseline sample taken but the other did not, hence, the variability of vegetation between some of the paired plots at the starting positions (Fig. 1). For the blanket bog sites, the current pin-quadrat monitoring started in 1972/3, ca. twenty years after the exclosures were erected, and there has been substantive changes in both atmospheric sulphur and nitrogen pollutant loads as well as overall sheep numbers which will almost certainly have impacted on the flora, and specially the bryophytes and lichens. Lastly, the monitoring methods were designed to assess species occurrence at a standardized detection level, therefore, species that are “new” or “disappear” may still be present at low frequency. In this regard, Milligan et al. (2018) argued that the recent appearance of *Sphagnum subnitens* might result from recent elevated nitrogen loads through a combination of colonization or increased abundance from an existing low population that was below the detections level in previous surveys. In any case, the analysis of long-term experiments simultaneously is a powerful tool to describe long-term vegetation trends to inform management choices (Alday et al. 2013b), as well as, to define some ecological indicators such as in our case the use of liverworts and lichens as indicators of grazing removal for these Upland habitats.

4.2. Conclusions and implications for conservation

Overall, the dominant plant species within these communities were rather stable, although lichens and bryophytes and some rare species were not. This suggests that the main vegetation is unlikely to change much over time if sheep numbers were to be reduced, or even removed, through wilding schemes (Monbiot, 2013; Sandom et al., 2013). One clear result is that plant community change takes a long time in these upland areas and wilding may take a very long time to produce compositional changes. The likely winners could be some of the sub-dominant species which might recover if grazing were to be removed, for example the seven focal species reported by Marrs et al. (2020), i.e. *Chamaenerion angustifolium*, *Dryopteris dilatata*, *Geum rivale*, *Narthecium ossifragum*, *Potentilla erecta*, *Rumex acetosa* and *Rubus chamaemorus* in a small-scale study of plant nutrition in some of these experiments. However, it took between 10 and 20 years in high-level bog and 60 years in the high-level grasslands for these focal species to increase (Marrs et al., 2020). As the seedbanks in these plant communities have been shown to be very depauperate (Lee et al. 2013, 2020), it is unlikely that additional species will colonize unless they are introduced by management, probably with some form of disturbance (Marrs et al., 2020). Moreover, in these experiments, and indeed across the entire grazing unit, there was no evidence of substantive tree invasion up to 2001 (24-45 years, Milligan et al., 2016) and this is confirmed here over a longer period (48-63 years). Restriction/control of tree invasion should be fundamental if these upland communities are to be conserved as they are or encouraged if succession to scrub woodland was deemed a desirable outcome. Irrespective, we predict that if sheep were removed at least in the short- to medium-term there would be a period of relative stability in plant composition. It can be argued that vegetation change might differ if grazing removal was implemented at larger scales than considered here (Ross et al., 2012); this is possible, but it remains to be tested.

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References

- Alday J.G., Marrs R.H., Martínez-Ruiz, C., 2011. Vegetation convergence during early succession on coal wastes: a six-year permanent plot study. *J. Veg. Sci.* 22, 1072-1083.**
- Alday J.G., Cox E.S., Pakeman R.J., Harris M.P.K. Le Duc M.G., Marrs, R.H., 2013a. Effectiveness of *Calluna*-heathland restoration methods after invasive plant control. *Ecol. Eng.* 54, 218-226. <https://doi.org/10.1016/j.ecoleng.2013.01.038>.
<https://doi.org/10.1111/j.1654-1103.2011.01308.x>.**
- Alday, J.G., Cox, E.S., Pakeman, R.J., Harris, M.P., Le Duc, M.G., & Marrs, R.H., 2013b. Overcoming resistance and resilience of an invaded community is necessary for effective restoration: a multi-site bracken control study. *J. Appld Ecol.* 50, 156-167. <https://doi.org/10.1111/1365-2664.12015>.
- Britton, A.J., Beale, C.M., Towers, W., Hewison, R.L., 2009a. Biodiversity gains and losses: Evidence for homogenisation of Scottish alpine vegetation. *Biol. Conserv.*, 142, 1728-1739.
DOI:10.1016/j.biocon.2009.03.010.
- Britton, A.J., Fisher, J.M., 2009b. Terricolous alpine lichens are sensitive to both load and concentration of applied nitrogen and have potential as bioindicators of nitrogen deposition. *Env. Poll.*, 158, 1296-1302. <https://doi.org/10.1016/j.envpol.2010.01.015>.
- Britton, A.J., Hester, A.J., Hewison, R.L., Potts, J.M., Ross, L.C, 2017. Climate, pollution and grazing drive long-term change in moorland habitats. *Appl. Veg. Sci.* 20, 194-203. DOI:10.1111/avsc.12260.
- Britton, A.J., Mitchell, R.J., Fisher, J.M., Riach, D.J., Taylor, A.F.S., 2018. Nitrogen deposition drives loss of moss cover in alpine moss-sedge heath via lowered C:N ratio and accelerated decomposition. *New Phyt.* 218, 470-478. DOI: 10.1111/nph.15006.

- Chen, Q.Q., Bakker, J.P., Alberti, J., Smit, C., 2020. Long-term management is needed for conserving plant diversity in a Wadden Sea salt marsh. *Biod. & Cons.* 29, 2329-2341. DOI: 10.1007/s10531-020-01976-w.
- Conant R.T., 2010., Challenges and Opportunities for Carbon Sequestration in Grassland Systems: A Technical Report on Grassland Management and Climate Change Mitigation Integrated Crop Management. FAO, Rome.
- Darling, F.F., 1955. West Highland Survey: an essay in human ecology. OUP, London, UK.
- de Bello, F., Lavorel, S., Gerhold, P., Reier, U., Partel, M., 2010. A biodiversity monitoring framework for practical conservation of grasslands and shrublands. *Biol. Conserv.* 143, 9-17. DOI: 10.1016/j.biocon.2009.04.022.
- Grime, J.P., 1998. Benefits of plant diversity to ecosystems: immediate, filter and founder effects. *J. Ecol.* 86, 902-910. DOI: 10.1046/j.1365-2745.1998.00306.x.
- Kapfer, J., & Popova, K. (2020). Changes in subarctic vegetation after one century of land use and climate change. *J. Veg. Sci.* DOI: 10.1111/jvs.12854.
- Kok, A., de Olde, E.M., de Boer, I.J.M., Ripoll-Bosch, R., 2020. European biodiversity assessments in livestock science: A review of research characteristics and indicators. *Ecol. Ind.* 112105902. <https://doi.org/10.1016/j.ecolind.2019.105902>.
- Lee, H., Alday, J.G., Rosenburgh, A., Harris M., McAllister, H. Marris R.H., 2013. Change in propagule banks during prescribed burning: a tale of two contrasting moorlands. *Biol. Cons.* 165, 187-197. DOI: <http://dx.doi.org/10.1016/j.biocon.2013.05.023>.
- Lee, H., McAllister, H.A., Cho, K., Rose, R.J., O'Reilly, J., Furness M., Marris R.H., 2020. Effects of long-term removal of sheep grazing on the seedbanks of high-level grasslands and blanket bogs. *Proceedings of the National Institute of Ecology*, in press.
- Lindenmayer, D.B., Likens, G.E., 2018. Effective ecological monitoring. 2nd ed. CSIRO Publishing, Clayton, Victoria, Australia.

- McGovern, S., Evans, C.D., Dennis, P., Walmsley C., McDonald, M.A., 2011. Identifying drivers of species compositional change in a semi-natural upland grassland over a 40-year period. *J. Veg. Sci.* 22, 346–356. Doi: 10.1111/j.1654-1103.2011.01256.x.
- McSherry M.E., Ritchie M.E., 2013. Effects of grazing on grassland soil carbon: a global review *Glob. Change Biol.* 19, 1347-1357 doi:10.1111/gcb.12144.
- Marrs, R.H., Rawes, M. Robinson, J.S., Poppitt, S.D., 1986. Long- term studies of vegetation change at Moor House NNR: guide to recording methods and database. (Merlewood Research and Development Paper 109). Institute of Terrestrial Ecology, Grange-over-Sands, UK.
- Marrs, R.H., Bravington, M.J., Rawes, M., 1988. Long term vegetation change in the *Juncus squarrosus* grassland at Moor House NNR in northern England. *Vegetatio* 76, 179-187. <https://www.jstor.org/stable/20038332>.
- Marrs, R.H., Sánchez, R., Connor, L., Blackbird, S., Rasa, J., Rose, R.J., 2018. Effects of removing sheep grazing on soil chemistry, plant nutrition and forage digestibility: lessons for rewilding the British uplands. *Ann. Appld Biol.* 173, 294–301. DOI:10.1111/aab.1246.
- Marrs, R.H., Lee, H., Blackbird, S., Connor, L., Girdwood, S.E., O'Connor, M., Smart, S.M., Rose, R.J., O'Reilly, J., Chiverrell, R., 2020. Release from sheep-grazing appears to put some heart back into upland vegetation: a comparison of nutritional properties of plant species in long term experiments. *Ann Appld Biol.* 171, 152-162. DOI: 10.1111/aa b.12591.
- Miles, J., 1979. *Vegetation dynamics*. Chapman and Hall, London.
- Milligan, G., Rose, R.J., Marrs, R.H., 2016. Winners and losers in a long-term study of vegetation change at Moor House NNR: effects of sheep-grazing and its removal on British upland vegetation. *Ecol. Ind.* 68, 89–101. <http://dx.doi.org/10.1016/j.ecolind.2015.10.053>.
- Milligan, G., Rose, R.J., O'Reilly, J., Marrs, R.H., 2018. Effects of rotational prescribed burning and sheep grazing on moorland plant communities: Results from a 60-year intervention experiment. *Land Deg. Dev.* 29, 1397–1412. <https://doi.org/10.1002/ldr.2953>.

- Mitchell, R.J., Hewison, R.L., Britton, A.J., Brooker, R.W., Cummins, R.P., Fielding, D.A., Fisher, J.M., Gilbert, D.J., Hester, A.J., Hurskainen, S., Pakeman, R.J., Potts, J.M., Riach, D., 2017. Forty years of change in Scottish grassland vegetation: Increased richness, decreased diversity and increased dominance. *Biol. Cons.* 212, 327-336. DOI: 10.1016/j.biocon.2017.06.027.
- Mitchell, R.J., Hewison, R.L., Fielding, D.A., Fisher, J.M., Gilbert, D.J., Hurskainen, S., Pakeman, R.J., Potts, J.M., Riach, D., 2018. Decline in atmospheric sulphur deposition and changes in climate are the major drivers of long-term change in grassland plant communities in Scotland. *Env. Poll.* 235, 956-964. DOI: 10.1016/j.envpol.2017.12.086.
- Moore, E. K. Britton, A.J. Iason, G., Pemberton, J., Pakeman R.J., 2015. Landscape-scale vegetation patterns influence small-scale grazing impacts. *Bio. Cons.* 192, 218–225.
<http://dx.doi.org/10.1016/j.biocon.2015.09.019>.
- Monbiot, G., 2013. *Feral: Searching for enchantment on the frontiers of Rewilding*. Allen Lane, London.
- Monteith, D., Henrys, P., Banin, L., Smith, R., Morecroft, M., Scott, T., Andrews, C., Beaumont, D., Benham, S., Bowmaker, V., Corbett, S., Dick, J., ; Dodd, B., ; Dodd, N., ; McKenna, C., ; McMillan, S., Pallett, D., Pereira, M.G., Poskitt, J., Rennie, S., Rose, R., Schafer, S., Sherrin, L., Tang, S., Turner, A., Watson, H., 2016. Trends and variability in weather and atmospheric deposition at UK Environmental Change Network sites (1993-2012). *Ecol.Ind.* 68, 21-35.DOI: 10.1016/j.ecolind.2016.01.061.
- Muurinen, L., Oksanen, J., Vanha-Majamaa, I., Virtanen, R., 2019. Legacy effects of logging on boreal forest understorey vegetation communities in decadal time scales in northern Finland. *For. Ecol. Man* 436, 11-20. <https://doi.org/10.1016/j.foreco.2018.12.048>.
- Økland, R.H., 1999. On the variation explained by ordination and constrained ordination axes. *J. Veg. Sci.* 10, 131-136. <https://doi.org/10.2307/3237168>.
- Oksanen, J., Blanchet, F.G., Friendly, M., Kindt, R., Legendre, P., McGlinn, D., Minchin, P.R. O'Hara, R.B., Simpson, G.L., Solymos, P., Stevens, M.H.H., Szoecs, E., Wagne H., 2019. Package 'vegan' Community Ecology Package Version 2.5-5, 2019-05-08. (accessed 1/8/2019).

- Pakeman R.J., 2004. Consistency of plant species and trait responses to grazing along a productivity gradient: a multi-site analysis. *J. Ecol.* 92, 893–905. <https://www.jstor.org/stable/359938.7>.
- Pakeman, R.J., Fielding, D.A., Everts, L., Littlewood N.A., 2019. Long-term impacts of changed grazing regimes on the vegetation of heterogeneous upland grasslands. *J. Appld Ecol.* 56, 1794–1805. DOI: 10.1111/1365-2664.13420.
- Pullin, A.S., Knight, T.M., 2009. Doing more good than harm - Building an evidence-base for conservation and environmental management. *Biol. Cons.*, 142, 931-934. DOI: 10.1016/j.biocon.2009.01.010.
- O'Mara F.P., 2012. The role of grasslands in food security and climate change *Ann. Bot.* 110, 1263-1270 doi:10.1093/aob/mcs209.
- Questad, E.J., Foster, B.L., 2008. Coexistence through spatio-temporal heterogeneity and species sorting in grassland plant communities. *Ecol. Lett* 11. 717-726. <https://doi.org/10.1111/j.1461-0248.2008.01186.x>
- R Core Team, 2019. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. URL <https://www.R-project.org> [accessed 1/1/2019].
- Rawes, M., 1981. Further results of excluding sheep from high-level grasslands in the north Pennines. *J. Ecol.* 69, 651-669. <http://www.jstor.org/stable/2259690>.
- Rawes, M., 1983. Changes in two high altitude blanket bogs after the cessation of sheep grazing. *J. Ecol.*, 71 219-235. <http://www.jstor.org/stable/2259974>.
- Rawes, M, Hobbs, R., 1979. Management of semi-natural blanket bog in the northern Pennines. *J. Ecol.* 67, 789-807. doi:10.2307/2259215.
- Rawes, M., Welch, D., 1966. Further studies in sheep grazing in the northern Pennines. *J. Br. Grassld Soc.* 21, 56-61. DOI: 10.1111/j.1365-2494.1966.tb00447.x.
- Rawes, M., Welch, D., 1969. Upland productivity of vegetation and sheep at Moor House National Nature Reserves, Westmorland, England. *Oikos Suppl.* 11, 1-69.

- Renwick, A.J., Jansson, T., Verburg P.H., Revoredo-Giha, C., Britz, W., Gocht, A. McCracken, D. 2013., Policy reform and agricultural land abandonment in the EU. *Land Use Pol.* 30, 446– 457. <http://dx.doi.org/10.1016/j.landusepol.2012.04.005>.
- Rose R., Monteith, D.T., Henrys, P., Smart, S., Wood, C., Morecroft, M., Andrews, C., Beaumont, D., Benhan S., Bowmaker, V., Corbett, S., Dick, J., Dodd, B., Dodd, N., Flexen, M. McKenna, C., McMillan, S., Pallett. D., Rennie, S., Schafer, S., Scott, T., Sherrin, L., Turner, A., Watson, H., 2016. Evidence for increases in vegetation species richness across the UK Environmental Change Network sites resulting from changes in air pollution and weather patterns. *Ecol. Ind.* 68, 21-35. DOI: 10.1016/j.ecolind.2016.01.061.
- Rose, R.J., Marrs, R.H., O'Reilly, J., Furness, M., 2018. Long-term vegetation monitoring data (1961-2013) from moorland burning plots established at Hard Hill, Moor House in 1954. NERC Environmental Information Data Centre. <https://doi.org/10.5285/0b931b16-796e-4ce4-8c64-d112f09293f7>.
- Rose, R.J., Marrs, R.H., O'Reilly, J., Furness, M., Wood, C.M., 2020. Long-term monitoring of vegetation in enclosure and grazing plots at Moor House National Nature Reserve, 1953-2016 <https://doi.org/10.5285/c72ab043-1b02-42c9-94e8-c1cae42b3dc8>.
- Ross, L.C., Woodin, S.J., Hester, A.J., Thompson D.B.A., Birks H.J.B., 2012. Biotic homogenization of upland vegetation: patterns and drivers at multiple spatial scales over five decades. *J. Veg. Sci.* 23, 755–770. Doi: 10.1111/j.1654-1103.2012.01390.x.
- RoTAP, 2012. Review of Transboundary Air Pollution in the UK. Acidification, Eutrophication, Ground Level Ozone and Heavy Metals in the UK. Report Prepared for DEFRA. CEH, Edinburgh, UK, pp. 292.
- Sandom, C.J. Hughes, J., Macdonald, D.W., 2013. Rooting for Rewilding: Quantifying Wild Boar's *Sus scrofa* Rooting Rate in the Scottish Highlands. *Rest. Ecol.* 21, 329-335. <https://doi-org.liverpool.idm.oclc.org/10.1111/j.1526-100X.2012.00904.x>

- Stewart, G.B., Coles, C.F., Pullin, A.S., 2005. Applying evidence-based practice in conservation management: Lessons from the first systematic review and dissemination projects. *Biol. Cons.* 126, 270-278. DOI: 10.1016/j.biocon.2005.06.003.
- Seabloom, E W., Bjørnstad, O.N., Bolker, B.M., Reichman, O.J., 2005. Spatial signature of environmental heterogeneity, dispersal, and competition in successional grasslands. *Ecol. Mono* 75, 199-214. <https://doi.org/10.1890/03-0841>
- Steinfeld, H, Gerber, P, Wassenaar, T, Castel, V, Rosales, M, De Haan, C., 2006. Livestock's long shadow. FAO, Rome .
- Sutherland, W.J., Wordley, C.F.R., 2018. A fresh approach to evidence synthesis. *Nature*, 558, 364-366. DOI: 10.1038/d41586-018-05472-8.
- Valencia E., de Bello F., Galland T., Adler P.B. Lepš J., E-Vojtkó A., van Klink R., Carmona C.P., Danihelka J., Dengler J., Eldridge D.J., Estiarte M., García-González R., Garnier E., Gómez-García D., Harrison S.P., Herben T., Ibáñez R., Jentsch A., Juergens N., Kertész M., Klumpp K., Louault F., Marrs R.H., Ogaya R., Ónodi G., Pakeman R.J., Pardo I., Pärtel M., Peco B., Peñuelas J., Pywell R.F., Rueda M., Schmidt W., Schmiechel U., Schuetz M., Skálová H., Šmilauer P., Šmilauerová M., Smit C., Song M., Stock M., Val J., Vandvik V., Ward D., Wesche K., K. Wiser S.K., A. Woodcock B.A., P. Young T.P., Yu F., Zobel M., Götzenberger, L. 2020. Synchrony matters more than species richness in plant community stability at a global scale. *PNAS* 117 (39), 24345-24351. <https://doi.org/10.1073/pnas.1920405117>.
- Van der Wal, R., Pearce, I.S.K., Brooker, R.W., 2005. Mosses and the struggle for light in a nitrogen-polluted world. *Oecologia* 142, 159-168. DOI:10.1007/s00442-004-1706-0.
- Welch D., Rawes M., 1964. The early effects of excluding sheep from high-level grasslands in the north Pennines. *J. Appld Ecol.* 1, 281-300. <https://www.jstor.org/stable/2401313>.
- Welch D., Rawes M., 1965. The Herbage Production of Some Pennine Grasslands. *Oikos* 16, 39-47. <https://www.jstor.org/stable/3564863>.

- Welch D., Rawes M., 1966. The intensity of sheep grazing on high-level blanket bog in upper Teesdale. *Irish J. Agric. Res.* 5, 185-196. <https://www.jstor.org/stable/25555409>.
- Xie, Z., Game, E.T., Hobbs, R.J., Pannell, D.J., Phinn, S.R., McDonald-Madden, E., 2020. Conservation opportunities on uncontested lands. *Nat. Sustain.* 3, 9–15. <https://doi.org/10.1038/s41893-019-0433-9>.

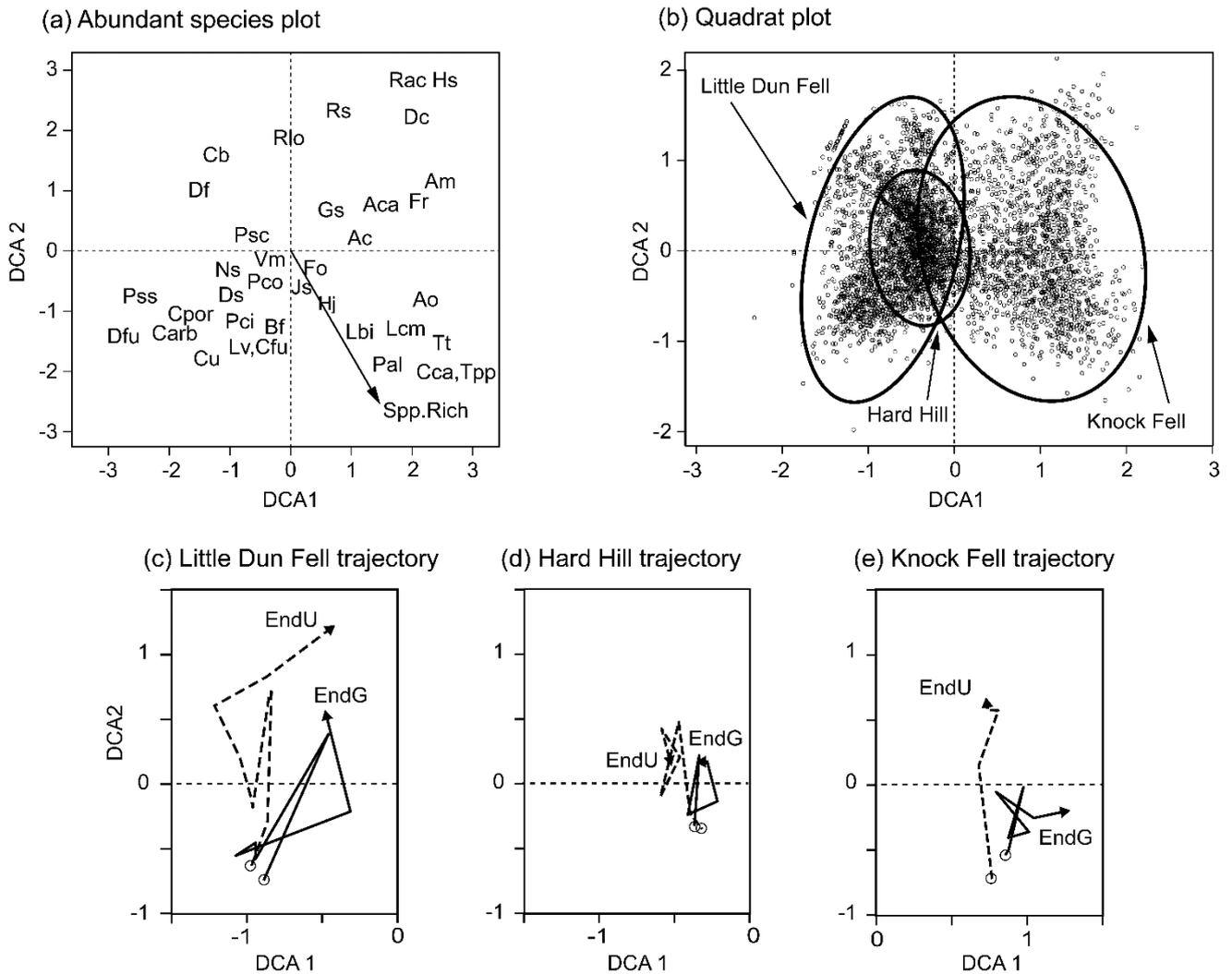


Fig. 1. Decorana plots from the grazed and ungrazed treatments from three high-level grassland sites (Hard Hill, Knock Fell and Little Dun Fell) at Moor House NNR: (a) the most abundant species, (b) quadrat plot with sites displayed by standard deviation ellipses (95%CL) and (c-e) temporal trajectories through time. Species Key: Ac=*Agrostis canina* s.l.; Aca=*Agrostis capillaris*; Af=*Avenella flexuosa*; Am=*Achillea millefolium*; Ao=*Anthoxanthum odoratum*; Bf=*Barbilophozia floerkei*; Carb=*Cladonia arbuscula*; Cb=*Carex bigelowii*; Cca=*Carex caryophylla*; Cfu=*Cladonia furcata*; Cpor=*Cladonia portentosa*; Cu=*Cladonia uncialis*; Dc=*Deschampsia cespitosa*; Dfu=*Dicranum fuscescens*; Ds=*Dicranum scoparium*; Fo=*Festuca ovina*; Fr=*Festuca rubra*; Gs=*Galium saxatile*; Hj=*Hypnum jutlandicum*; Hs=*Hylocomium splendens*; Js=*Juncus squarrosus*; Lbi=*Lophocolea bidentata*; Lcm=*Luzula campestris/multiflora*; Lv=*Lophozia ventricosa*; Ns=*Nardus stricta*; Pal=*Polytrichastrum alpinum*; Pci=*Ptilidium ciliare*; Pco=*Polytrichum commune*; Psc=*Pleurozium schreberi*; Pss=*Polytrichum strictum*; Rac=*Rumex acetosa*; Rlo=*Rhytidiadelphus loreus*; Rs=*Rhytidiadelphus squarrosus*; Tpp=*Thymus polytrichus*; Tt=*Thuidium tamariscinum*; Vm=*Vaccinium myrtillus*. Start and end points of trajectories are illustrated with and open circles and diamonds respectively; dotted lines = ungrazed; solid = grazed.

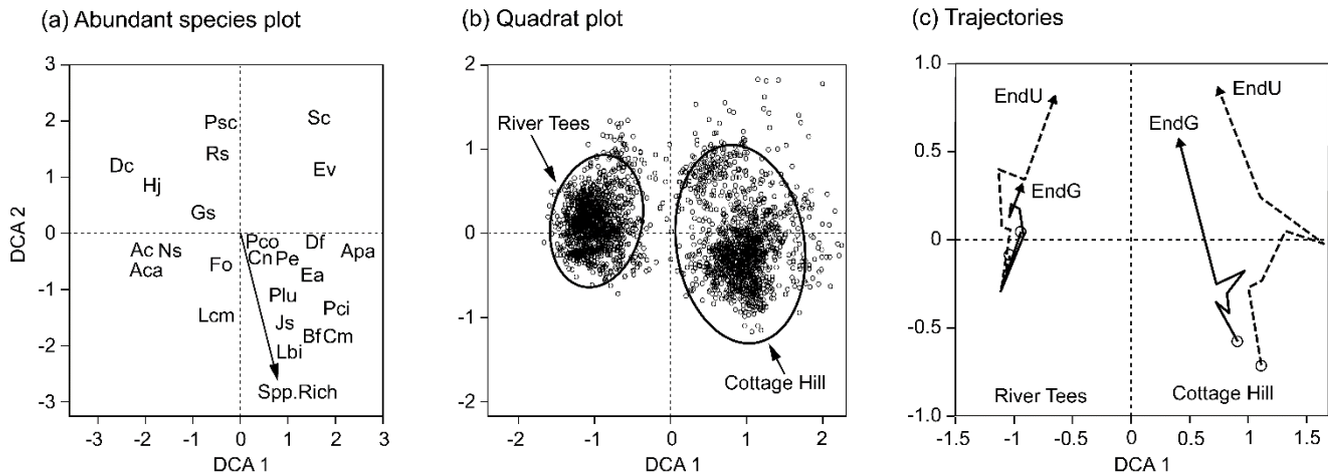


Fig. 2. Decorana plots from the grazed and ungrazed treatments from two Intermediate grassland sites (Cottage Hill and River Tees) at Moor House NNR: (a) the most abundant species, (b) quadrat plot with sites displayed by standard deviational ellipses (95%CL) and (c) temporal trajectories through time. Species Key: Ac=*Agrostis canina* sl; Aca=*Agrostis capillaris*; Af=*Avenella flexuosa*; Apa=*Aulacomnium palustre*; Bf=*Barbilophozia floerkei*; Cm=*Calypogeia muelleriana*; Cn=*Carex nigra*; Dc=*Deschampsia cespitosa*; Ea=*Eriophorum angustifolium*; Ev=*Eriophorum vaginatum*; Fo=*Festuca ovina*; Gs=*Galium saxatile*; Hj=*Hypnum jutlandicum*; Js=*Juncus squarrosus*; Lbi=*Lophocolea bidentata*; Lcm=*Luzula campestris/multiflora*; Ns=*Nardus stricta*; Pci=*Ptilidium ciliare*; Pco=*Polytrichum commune*; Pe=*Potentilla erecta*; Plu=*Plagiothecium undulatum*; Psc=*Pleurozium schreberi*; Rs=*Rhytidiadelphus squarrosus*; Sc=*Sphagnum capillifolium*. Start and end points of trajectories are illustrated with and open circles and diamonds respectively; dotted lines = ungrazed; solid = grazed.

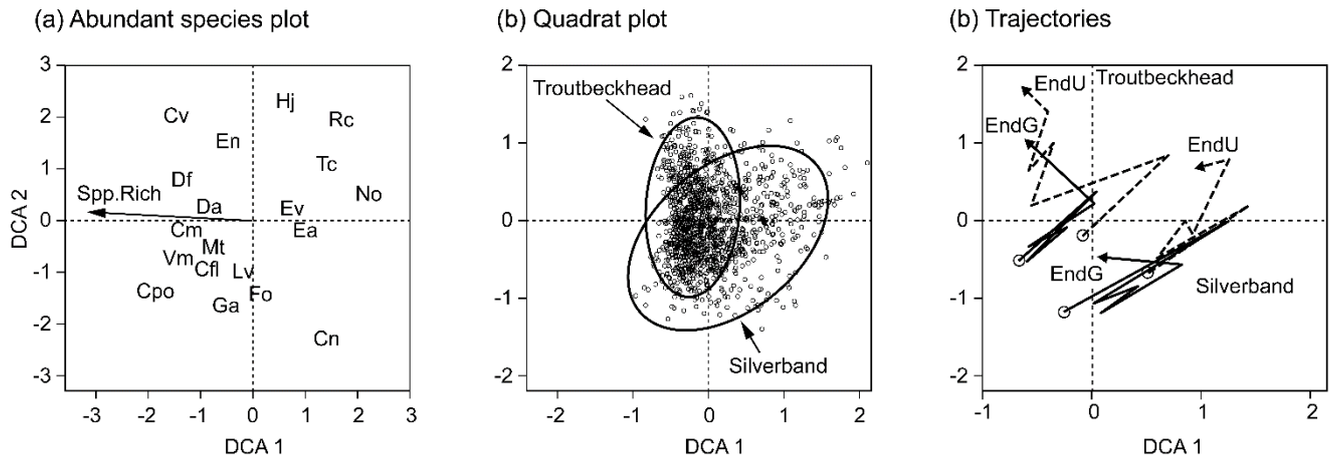


Fig. 3. Decorana plots from the grazed and ungrazed treatments from five blanket bog sites (Bog Hill and Hard Hill with Blocks denoted A, B, C, D) at Moor House NNR: (a) the most abundant species, (b) quadrat plot with sites displayed by standard deviational ellipses (95%CL) and (c-g) temporal trajectories through time. Species Key: Cbi=*Cephalozia bicuspidata*; Cco=*Cephalozia connivens*; Cfl=*Campylopus flexuosus*; Cm=*Calypogeia muelleriana*; Cpor=*Cladonia portentosa*; Cv=*Calluna vulgaris*; Ds=*Dicranum scoparium*; Ea=*Eriophorum angustifolium*; En=*Empetrum nigrum*; Ev=*Eriophorum vaginatum*; Ga=Green alga terrestrial ; Hj=*Hypnum jutlandicum*; Kpa=*Kurzia pauciflora*; Lv=*Lophozia ventricosa*; Plu=*Plagiothecium undulatum*; Pn=*Pohlia nutans*; Psc=*Pleurozium schreberi*; Rc=*Rubus chamaemorus*; Sc=*Sphagnum capillifolium*. Start and end points of trajectories are illustrated with and open circles and diamonds respectively; dotted lines = ungrazed; solid = grazed.

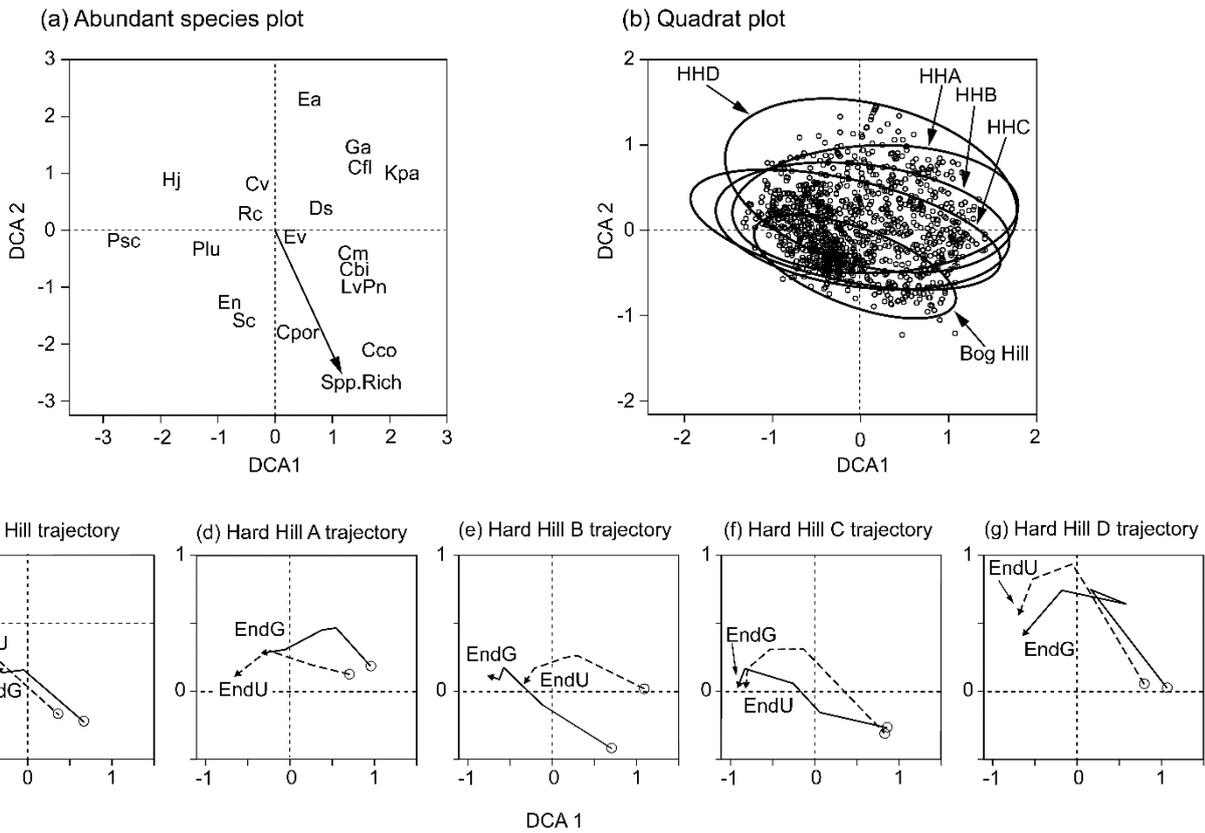


Fig. 4. Decorana plots from the grazed and ungrazed treatments from two high-level blanket bog sites (Silverband and Troutbeckhead) at Moor House NNR: (a) the most abundant species, (b) quadrat plot with sites displayed by standard deviational ellipses (95%CL) and (c) temporal trajectories through time. Species Key: Af=*Avenella flexuosa*; Cfl=*Campylopus flexuosus*; Cm=*Calypogeia muelleriana*; Cn=*Carex nigra*; Cpor=*Cladonia portentosa*; Cv=*Calluna vulgaris*; Da=*Diplophyllum albicans*; Ea=*Eriophorum angustifolium*; En=*Empetrum nigrum*; Ev=*Eriophorum vaginatum*; Fo=*Festuca ovina*; Ga=Green alga terrestrial ; Hj=*Hypnum jutlandicum*; Lv=*Lophozia ventricosa*; Mt=*Mylia taylorii*; No=*Narthecium ossifragum*; Rc=*Rubus chamaemorus*; Tc=*Trichophorum cespitosum*; Vm=*Vaccinium myrtillus*. Start and end points of trajectories are illustrated with and open circles and diamonds respectively; dotted lines = ungrazed; solid = grazed.

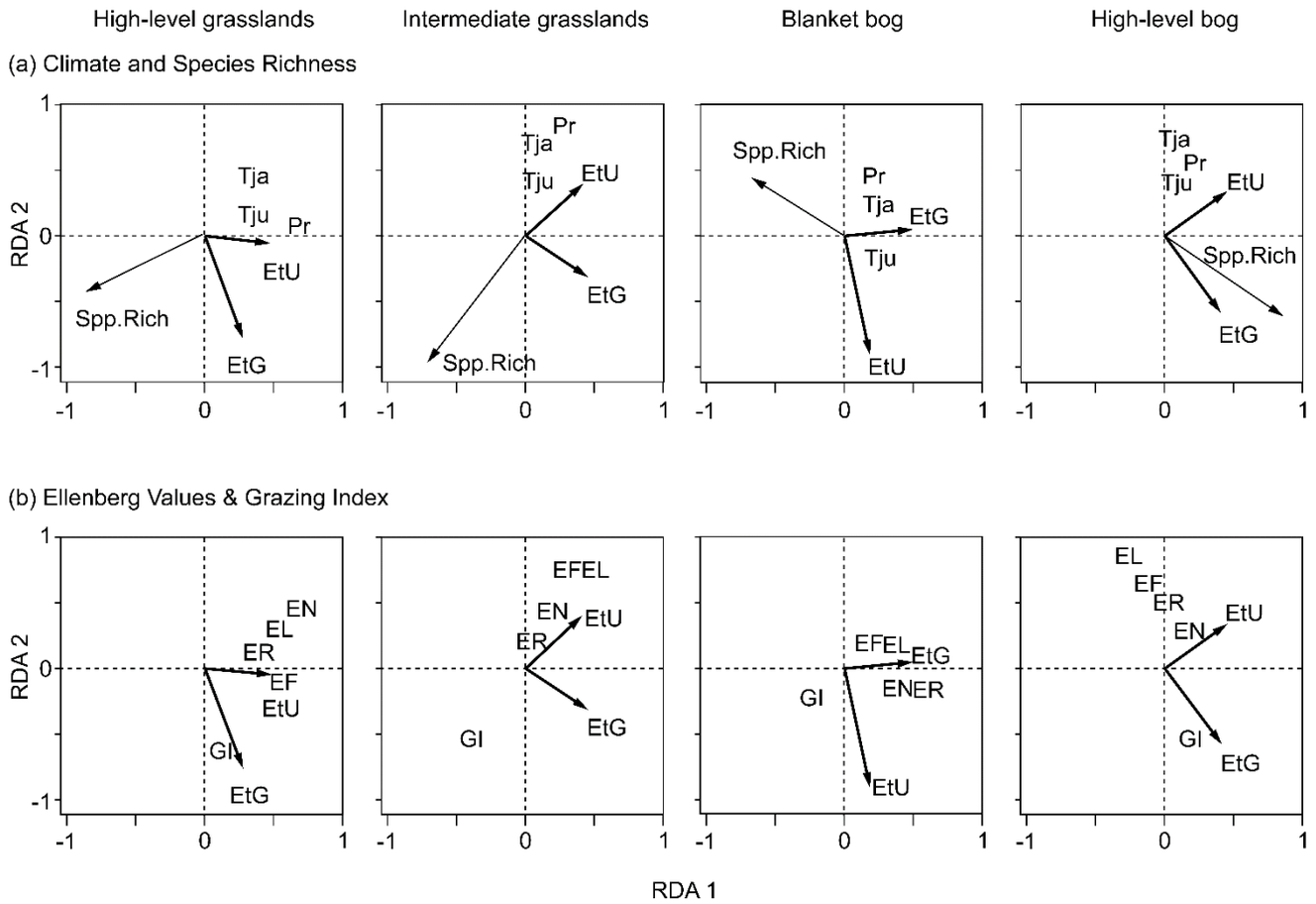


Fig. 5. Comparison of the changes in abundance-weighted traits derived from the redundancy analysis for (a) climate variables (mean annual January temperature, mean annual July temperature and precipitation), and (b) Ellenberg values and grazing index with respect to with respect to the temporal trajectories in the grazed, (EtG) and ungrazed plots, (EtU) for the four experimental groups at Moor House NNR. The species richness correlation was fitted over the climate variables plot (a): High-level grasslands $r^2 = 17.9\%$; Intermediate grasslands $r^2 = 54.9\%$; Blanket bog $r^2 = 12.1\%$; High-level bog $r^2 = 48.7\%$ (all $p < 0.001$). Variable codes are presented in Appendix A:Table S2.

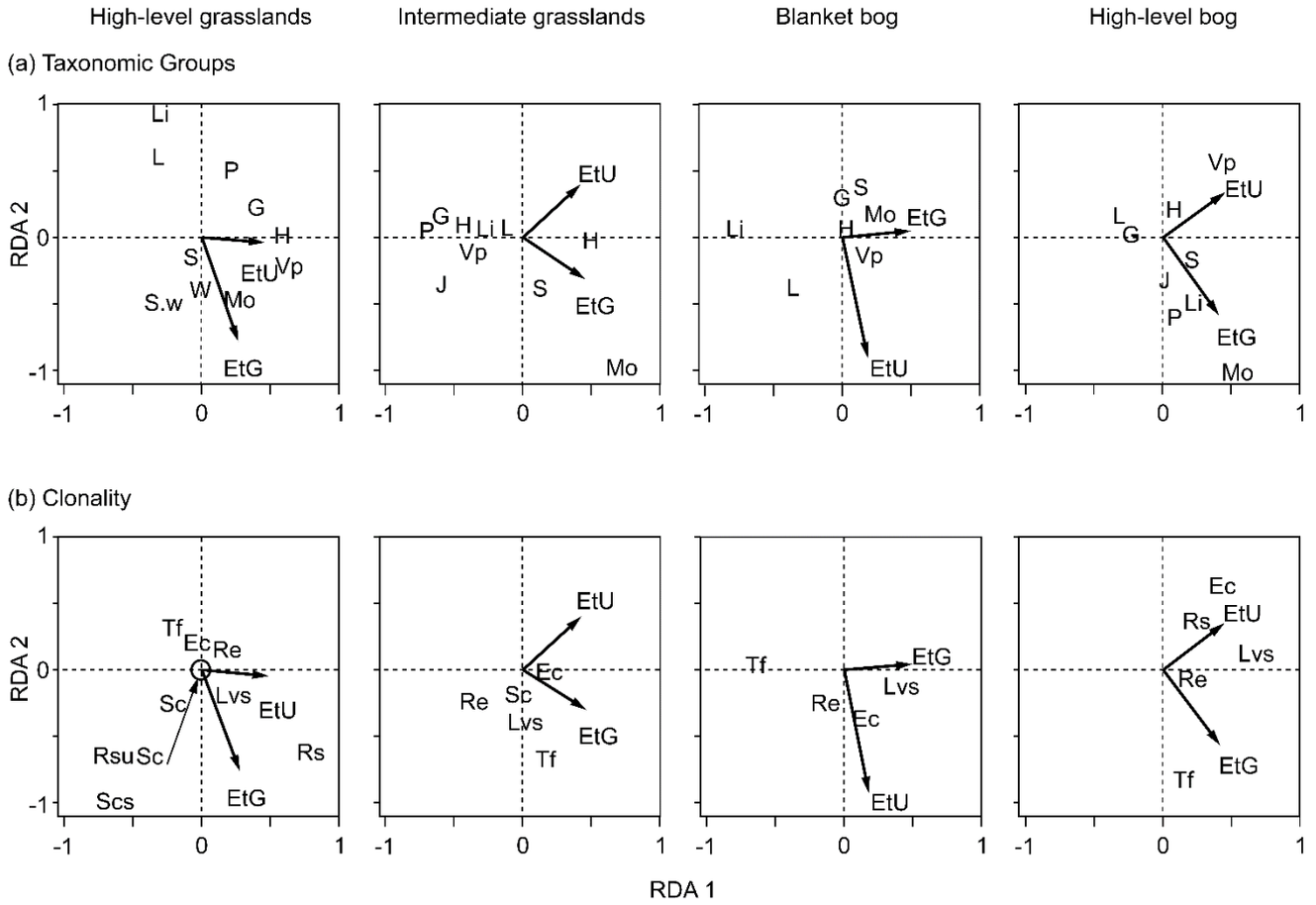


Fig. 6. Comparison of the changes in abundance-weighted derived from the redundancy analysis for (a) taxonomic and (b) clonality variables with respect the temporal trajectories in the grazed, (EtG) and ungrazed plots, (EtU) for the four experimental groups at Moor House NNR. Variable codes are presented in Appendix A:Table S2.

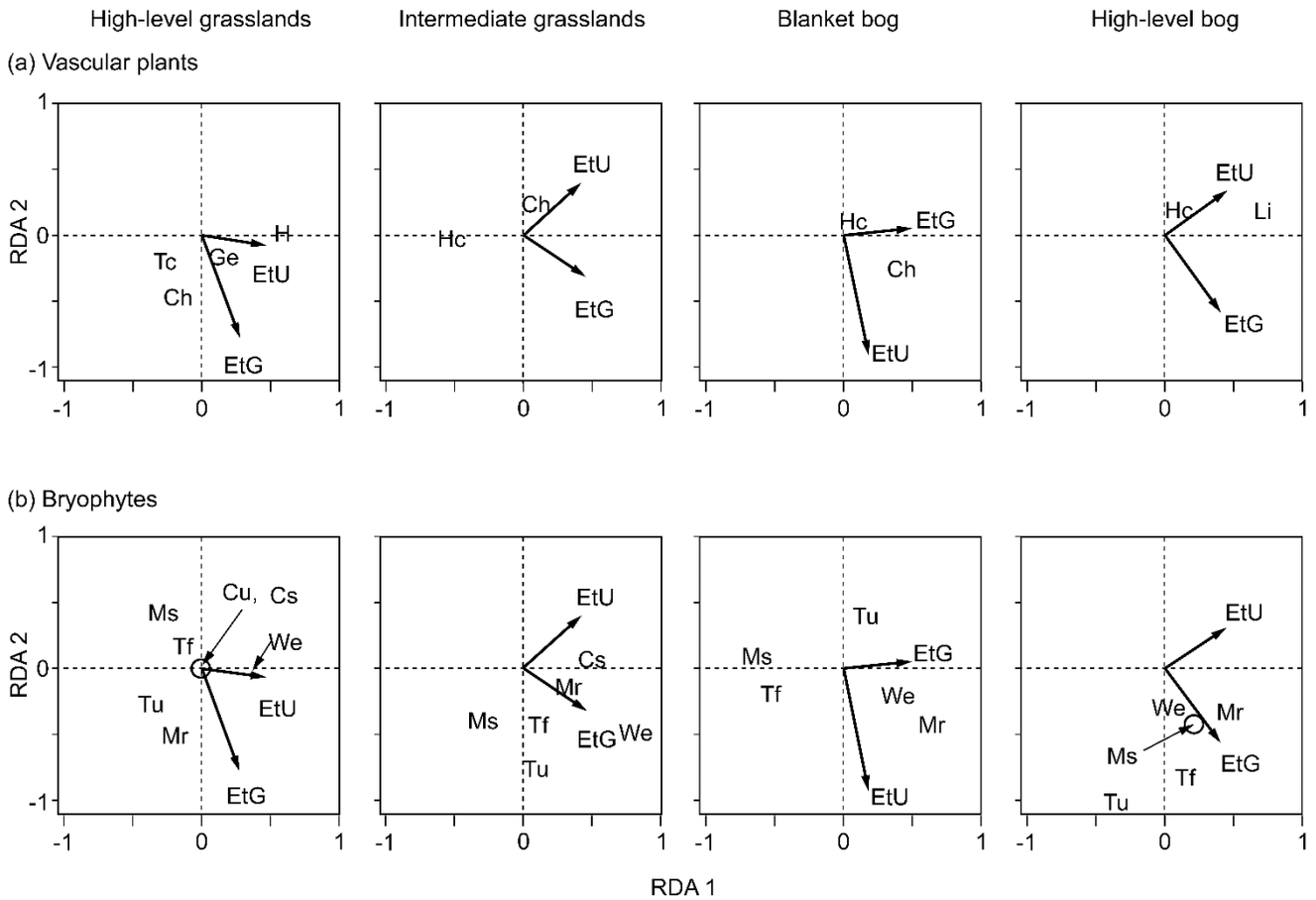


Fig. 7. Comparison of the changes in abundance–weighted derived from the redundancy analysis for life-history variables values with respect the temporal trajectories in the grazed, (EtG) and ungrazed plots, (EtU) for the four experimental groups at Moor House NNR: (a) vascular plants, and (b) bryophytes. Variable codes are presented in Appendix A:Table S2.

Appendix A. Data preparation, Supplementary Tables and Figures

Effects of long-term removal of sheep-grazing in a series of British upland plant communities: insights from plant species composition and traits

Alday, O'Reilly, Rose, Marrs

Data preparation

The dataset was voluminous and complex and required a substantive pre-processing, first to bring species nomenclature to the same standard: Stace (2019) for vascular plants, Atherton et al. (2010) for bryophytes and Dobson (2000) for lichens, and second to combine some taxa that were recorded inconsistently. These two pre-processing steps and changes were outlined in Milligan et al. (2016).

Whilst all data collection within each experiment was internally consistent, there were differences in methods of stratified random sampling between experiments; for example, vascular plants were recorded as number of hits per

pin strata, whereas bryophytes and lichens were counted as either present/absent. Accordingly, the following procedure was adopted to achieve a common recording methodology and sampling intensity across all experiments:

1. All species hits per pin from all height strata were summed to provide pin totals.
2. These summed values were converted to presence/absence data using the ‘decostand’ function. Taken together, these two steps reduced all data collected at a single pin to either 1 or 0, ensuring the vascular plants, bryophytes and lichens were quantified on the same scale.

The sum of all presences was calculated at each sampling position; depending on the experiment, this was either a pin-frame.

Appendix references

- Atherton, I., Bosanquet, S., Lawley, M., 2010. Mosses and liverworts of Britain and Ireland: a field guide. British Bryological Society, Plymouth, UK.
- Dobson, F., 2000. Lichens: an illustrated guide to the British and Irish species. Richmond Publishing Company, Richmond, UK.
- Critchley, C.N.R., 2000. Ecological assessment of plant communities by reference to species traits and habitat preferences. *Biod.and Cons.* 9, 87-105. DOI: 10.1023/A:1008976014424.
- Critchley, C.N.R., Smart, S.M., Poulton, S.M.C. & Myers, G.M., 1996. Monitoring the consequences of vegetation management in Environmentally Sensitive Areas. *Aspects Appld Biol.* 44, 193-201.
- Eddy, A., Welch, D., Rawes, M., 1969. The vegetation of the Moor House National Nature Reserve in the northern Pennines England. *Vegetatio* 16, 239-284. <https://www.jstor.org/stable/20035417>.
- Hill, M.O., Preston, C.D., Roy, D.B., 2004. PLANTT: Attributes of British and Irish plants, status, size, life history, geography and habitats. CEH, Huntingdon, UK.
- Hill, M.O., Preston, C.D., Bosanquet, S.D.S., Roy, D.B., 2007. BRYOATT: Attributes of British mosses, liverworts and hornworts. CEH, Huntingdon, UK.
- Lee, H., Alday, J.G., Rose, R.J., O’Reilly, J., Marrs, R.H., 2013. Long-term effects of rotational prescribed-burning and low-intensity sheep-grazing on blanket-bog plant communities. *J. Appld Ecol.* 50, 625-635. <https://doi.org/10.1111/1365-2664.12078>.
- Milligan, G., Rose, R.J., O’Reilly, J., Marrs, R.H., 2018. Effects of rotational prescribed burning and sheep grazing on moorland plant communities: Results from a 60-year intervention experiment. *Land Deg. Dev.* 29, 1397–1412. <https://doi.org/10.1002/ldr.2953>.
- Rawes, M., Welch, D., 1966. Further studies in sheep grazing in the northern Pennines. *J. Br. Grassld Soc.* 21, 56-61. DOI: 10.1111/j.1365-2494.1966.tb00447.x.
- Stace, C.A., 2019. New flora of the British Isles, 4th edn. C and M Floristics, Middlewood Green, Suffolk. UK.

Table S1

Description of the twelve-paired monitored sheep-grazing and exclosures plots at Moor House NNR in north-west England.

Experiment	Site Name (Code)	British National Grid reference	Elevation (m)	Year fence erected	¹ First year sampled Ungr/Graz	Last year sampled (no. of years ungrazed)	² Number of samplings Ungr/Graz	Vegetation type according to (Eddy et al., 1969)	³ Total area of pure stands of the vegetation types on the Moor House reserve (ha)	⁴ Sheep Grazing Density (sheep ha ⁻¹)
High-level Grassland	Hard Hill (HH)	NY 725 330	690	1954	1955/62	2015 (61)	7/6	<i>Festucetum</i>	180	2.6
	Little Dun Fell (LDF)	NY 704 331	830	1954	1955/62	2016 (62)	7/8	<i>Festucetum</i>	-	5.8
	Knock Fell (KF)	NY 717 312	750	1955	1956/62	2014 (59)	8/8	Limestone Agrostofestucetum	125	5.8
Intermediate Grassland	Cottage Hill (CH)	NY 758 336	550	1967	1967	2016 (49)	7/8	<i>Juncus squarrosus</i> grassland	373	1.4
	River Tees (RT)	NY 747 344	550	1967	1967	2015 (48)	7/7	<i>Nardus stricta</i> grassland	416	2.8
Blanket Bog ³	Hard Hill Burning AN (HHB.AN)	NY 743 330	600	1954	1972	2013 (59)	5/5	<i>Calluna-Eriophorum</i>	1169	0.01-0.30
	Hard Hill Burning BN (HHB.BN)	NY 740 330	610	1954	1972	2013 (59)	5/5	<i>Calluna-Eriophorum</i>	1169	0.01-0.30
	Hard Hill Burning CN (HHB.CN)	NY 736 330	617	1955	1972	2013 (58)	5/5	<i>Calluna-Eriophorum</i>	1169	0.01-0.30
	Hard Hill Burning DN (HHB.DN)	NY 738 331	632	1954	1972	2013 (59)	5/5	<i>Calluna-Eriophorum</i>	1169	0.01-0.30
	Bog Hill (BH)	NY 767 328	550	1953	1971	2016 (63)	4/4	<i>Calluna-Eriophorum</i>	1169	na
High-level Bog	Silverband (SB)	NY 710 309	690	1966	1966	2015 (49)	7/7	<i>Eriophoretum</i>	323	0.25
	Troutbeck Head (TB)	NY 722 317	690	1966	1966	2014 (48)	7/7	<i>Eriophoretum</i>	419	0.5

¹In some experiments only the enclosed plot was sampled at the start.²Some experiments had additional samplings of one treatment for unspecified reasons.³The total area of these communities makes up 3019 ha, i.e. 79% of the reserve area of 3842 ha, the remaining vegetation comprised predominantly re-colonising peatland, sandstone scree and mosaics of the above vegetation classes (Eddy et al. 1969).⁴Sheep grazing density was determined by dropping volume measurement (Rawes and Welch 1969); data were not available (na) for one site.⁵Hard Hill Grazing and Burning experiment: data have been abstracted for each of the grazed and ungrazed no-burn treatment plots (coded N) in each of four replicate blocks (A-D) (Lee et al. 2013; Milligan et al., 2018).**Table S2**

Diversity and trait variables derived for each of the four experiments at Moor House National Nature Reserve. Taxonomic groupings were derived from Stace (2019) and Atherton et al. (2010); trait data were derived from Hill et al. (2004) for vascular plant data, Hill et al. (2007) for bryophyte data and the suited species database for the grazing index for vascular plants (Critchley et al., 1996; Critchley, 2000). The importance of these variable in explaining the species composition assessed using the Adonis analysis is also presented as R² and F values; were significant (P<0.001) unless denoted as * (P<0.05). Species traits that showed no significant relationships included thalloid mats (Bryophyte life-forms) and vascular plant species with root suckering, shortly-creeping stolons and extensively-creeping stolons (Clonality).

Grouping	Classes or Variables	Code	Grasslands				Blanket Bog			
			High-level		Intermediate		Blanket Bog		High-level	
			R ²	F	R ²	F	R ²	F	R ²	F
Diversity	Species richness	SR	8.34	402.0	6.64	209.7	13.18	195.7	9.45	152.3
	Shannon-Weiner Index	SW	2.65	127.8	6.79	214.2	5.00	74.3	2.67	43.1
	Simpson's Index	SI	0.66	31.7	1.49	47	6.65	98.8	1.43	23.1
Taxonomic classes	Vascular plants	Vp	7.34	546.5	5.51	415.4	26.84	751.2	11.84	305.3
	Mosses	Mo	5.69	423.7	6.53	492.4	6.66	186.5	3.05	78.6
	Liverworts	Li	3.16	235.2	11.45	864.1	11.25	315	5.82	150
	Lichens	L	1.94	144.8	0.04	2.9	2.0	55.9	2.77	71.4
	Poacea	P	5.3	394.8	21.4	1614.7	-	-	6.71	173.1
	Juncaceae	J	3.2	238.5	7.08	534.1	-	-	0.05	1.2
	Graminoids	G	10.28	766.2	11.08	836.1	4.16	116.4	8.00	206.3
	Sphagnaceae	S	0.29	21.7	0.68	51.2	5.25	147.1	0.89	23
	Herbaceous	H	0.29	21.7	0.73	54.9	4.19	114.6	7.11	183.2
	Semi-woody	S.w	2.83	210.6	-	-	-	-	-	-
	Woody	W	2.93	217.9	-	-	-	-	-	-
Vascular plant – Life-form classes	Chamaephyte	Ch	3.41	163.6	0.72	20.5	23.47	385.8	10.44	182.5
	Geophyte	Ge	0.38	18.2	-	-	-	-	-	-
	Hemi-cryptophyte	Hc	7.08	339.7	5.51	157.7	8.58	141.1	9.8	171.4
	Therophyte	Th	0.44	21.3	-	-	-	-	-	-
Bryophytes Life-form classes	Creeping solitary	Cs	0.04	1.8*	-	-	-	-	-	-
	Cushions	Cu	0.04	2.0*	-	-	-	-	-	-
	Mats rough	Mr	6.54	337.2	13.00	548.3	10.75	220.5	2.75	44.7
	Mats smooth	Ms	1.04	53.8	8.31	350.8	10.98	225.2	7.09	115.3
	Turfs	Tu	2.1	108.1	0.92	38.8	18.1	371.1	0.97	15.8
	Tufts	Tf	4.55	234.6	6.50	274.2	3.25	66.6	3.31	53.8
	Wefts	We	6.54	337.2	0.32	13.7	2.59	53.2	0.34	5.5

Table S2 continued										
Grouping	Classes or Variables	Code	Grasslands				Blanket Bog			
			High-level		Intermediate		Blanket Bog		High-level	
			R ²	F	R ²	F	R ²	F	R ²	F
Clonality	Little vegetative spread	Lvs	4.02	261.4	1.36	51.0	22.33	404.8	7.35	152.9
	Tussock-forming	Tf	3.42	222.4	0.58	21.7	7.71	139.7	4.3	89.5
	Shortly-creeping	Sc	0.91	59.4	0.16	6.0	-	-	-	-
	Extensively-creeping	Ec	8.17	531.5	22.89	859.3	4.39	79.5	8.12	169
	Rhizome shortly-creeping	Rs	10.9	709.4	-	-	-	-	7.27	151.3
	Rhizome Extensively-creeping	Re	7.22	469.7	3.52	132.2	4.05	73.3	6.14	127.8
Ellenberg Values	Light	EL	8.05	525.4	4.49	209.4	20.79	414.9	5.05	91.3
	Moisture	EF	4.55	296.5	23.08	1075.8	13.91	277.6	5.8	104.9
	Reaction	ER	14.06	917.1	12.13	565.1	4.75	94.8	5.8	104.8
	Fertility	EN	8.12	529.5	2.69	125.1	4.68	93.4	6.42	116.1
Climate Values	Mean annual Jan temp (T.Jan °C)	TJa	6.54	357.4	4.48	206.3	21.18	374.9	4.97	81.6
	Mean annual Jul temp (T.Jul °C)	TJu	11.06	604.4	33.47	1541.9	7.49	132.6	4.06	66.6
	Annual precipitation (mm)	Pr	4.53	247.5	3.75	172.7	8.30	146.9	6.08	99.8
Grazing	Grazing Index	GI	6.66	347.7	22.29	890.7	21.21	348.3	6.45	106.6

Table S3

Summary statistics from (a) the DCA analyses of species community data, and (b) the RDA analysis of species traits in four grazing enclosure experiments at Moor House NNR, northern England. In (a) eigenvalues (λ) and gradient lengths (GL) are presented for the first four DCA axes; (b) eigenvalues are presented for the first four RDA axes, along with the partitioning of inertia.

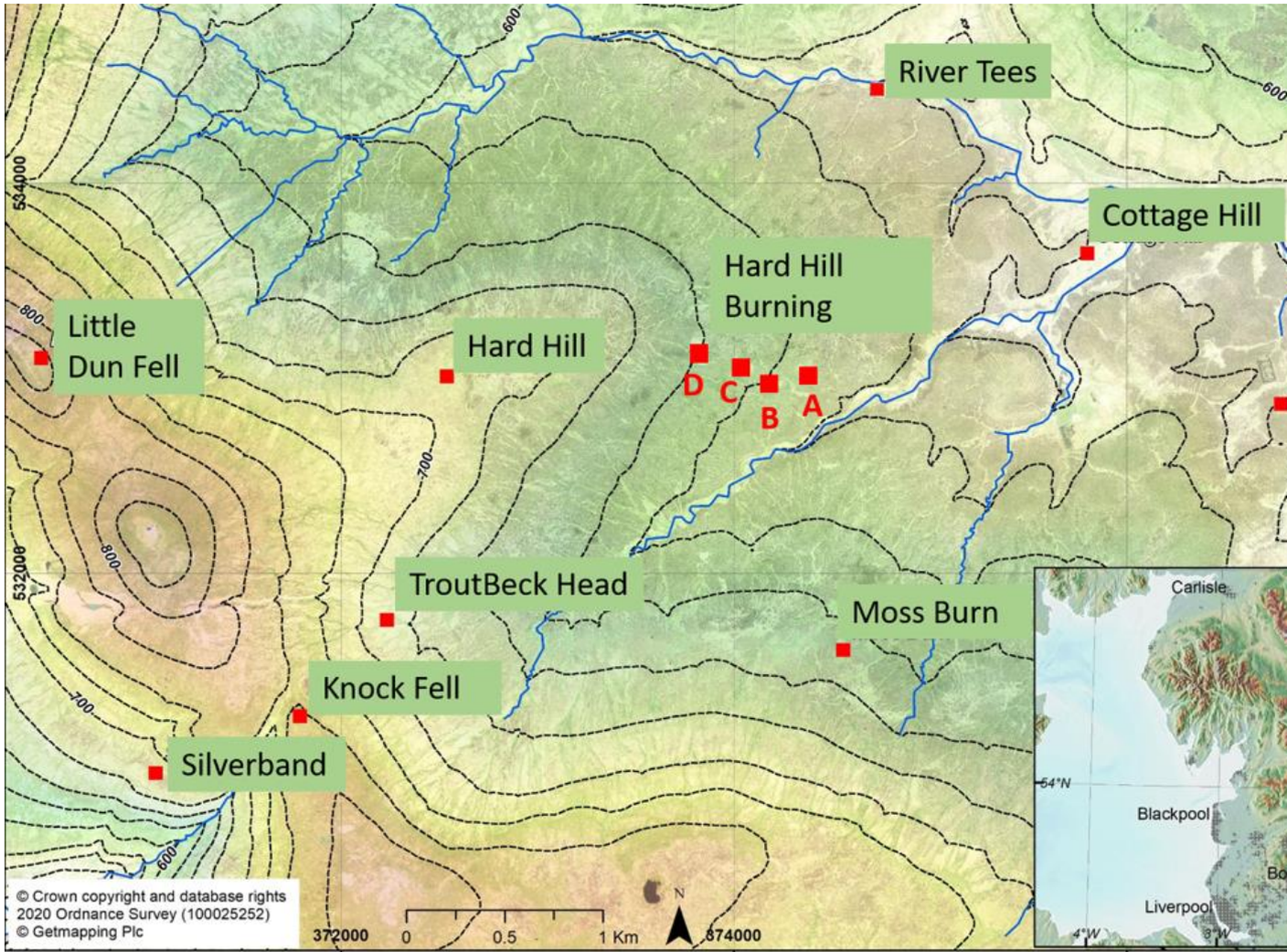
(a)

Experiment	λ_1	λ_{12}	λ_3	λ_4	GL ₁	GL ₂	GL ₃	GL ₄
High-level grasslands	0.471	0.297	0.223	0.193	4.51	4.11	3.93	3.72
Intermediate grasslands	0.548	0.196	0.127	0.120	4.04	3.38	1.65	2.62
Blanket bog	0.337	0.210	0.170	0.142	2.74	2.67	2.58	2.29
High-level Blanket Bog	0.303	0.299	0.286	0.201	3.48	3.76	3.27	2.56

(b)

Experiment	λ_1	λ_{12}	λ_3	λ_4	Total Inertia	Constrained Inertia	Unconstrained Inertia	% Explained
High-level grasslands	7.555	4.883	2.550	2.080	40.00	2.80	37.20	6.99
Intermediate grasslands	8.870	4.423	3.196	2.411	32.00	2.55	29.45	7.98
Blanket bog	13.906	1.945	1.555	1.282	27.00	2.61	24.39	9.66
High-level Blanket Bog	8.423	3.991	3.222	1.954	30.00	2.23	37.77	7.43

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Fig. S1. Location of the Moor House Natural National Reserve and the distribution of the individual study sites (R.C. Chiverrell).