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- 1 Microclimate variability and long-term persistence of fragmented woodland
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- 8
- 9 Abstract

10 Favourable microclimates are predicted to buffer fragmented populations against the effects of 11 environmental change, but ecological timeseries are often too short to establish the extent to which 12 such microsites facilitate population persistence through multiple climate shifts. We investigate the 13 effects of microclimatic heterogeneity on woodland resilience through millennial climate and 14 disturbance shifts near northwest European woodland range limits. We use palaeoecological data 15 from northern Scotland to study the effects of fragmentation on community composition and 16 diversity in a potentially favourable microclimate, and compare palynological timeseries of tree 17 abundance from five sites to assess the effects of favourable (low-lying sheltered) versus more 18 marginal (higher altitude) settings on population persistence and stability. The sheltered site shows 19 persistence of tree cover through Holocene climatic and anthropogenic shifts, including climatically-20 driven regional woodland contraction around 4400 cal BP (calendar years before present), when 21 surviving woods became compositionally differentiated into upland pine and low-lying deciduous 22 communities. A favourable microclimate can thus buffer woodlands against environmental shifts 23 and increase continuity of canopy cover, but it does not generate stable communities. Compositional 24 reorganisation is an essential stress response mechanism and should be accommodated by 25 conservation managers. The replacement of deciduous taxa by Pinus sylvestris after 1060 cal BP 26 represents the decoupling of pine distribution from climate drivers by management intervention. As 27 a result, current microrefugial woodland composition reflects late Holocene human intervention. 28 Alternative models of community composition and behaviour from palaeoecology provide a stronger 29 foundation for managing microsite communities than relict woods in contrasting environmental 30 settings.

- 31
- 32 Keywords
- 33 Paleoecology; climate change; conservation; woodland; fragmentation; Scotland
- 34
- 35 1 Introduction
- 36 Global reductions in woodland size mean that fragmented populations play an increasingly
- 37 significant role in conservation (Haddad et al., 2015). Favourable microclimates in otherwise
- 38 inhospitable landscapes allowed the survival of climate relict tree populations and associated
- 39 biodiversity in the past, and are predicted to buffer populations against ongoing environmental
- 40 change (Hampe and Jump, 2011; Maclean et al., 2015). However, the timescales of modern

- 41 ecological studies are too brief to establish the extent to which microclimates can mitigate the
- 42 negative impacts of fragmentation through multiple climate shifts, including extinction debt and
- 43 local extinction-recolonization dynamics (Saunders et al., 1991; Vellend et al., 2006). Long timeseries
- 44 provide a powerful tool for understanding to what extent locally favourable conditions allow
- 45 populations to persist through multiple environmental changes. They offer insights into the origins
- 46 of modern conservation values in long-fragmented communities and their potential sensitivity to
- 47 future climatic fluctuations (Bhagwat et al., 2012).

48 Northern Scotland is an appropriate location to study interactions between microclimate and 49 woodland resilience because it lies on the range edge for temperate woodland and extant woods are 50 highly fragmented, thus exposing them to recruitment and dispersal stresses. Woods have been 51 repeatedly exposed to climate stresses in the past (Tipping, 1994). This is particularly the case for 52 Pinus sylvestris L. (Scots pine), which underwent multiple phases of population contraction and 53 expansion in response to Holocene climate shifts (Willis et al., 1998). Favourable microclimates are 54 also important for the adaptive capacity of species with a northerly biogeographical distribution: 55 Scots pine is thought to have survived the last glaciation in northerly refugia, including the 56 continental shelf off northwest Scotland, and its range is predicted to shift northeast in response to 57 ongoing climate change (Bhagwat and Willis, 2008; Matias and Jump, 2012). Understanding the 58 extent to which woods in this region were buffered against smaller amplitude climate shifts in the 59 past can help evaluate current and future site potential to retain these populations in the event of 60 more extreme future shifts by indicating whether they served as persistent or transient microclimate 61 refugia (Keppel et al., 2012). Our investigation examines the effects of microclimatic heterogeneity 62 on arboreal resilience in this region. We present new stand-scale pollen evidence for the effects of 63 long-term fragmentation on community composition and diversity in a potentially favourable 64 microclimate, and assess the role of microclimatic buffering on population persistence through a

- 65 comparison of time series of tree abundance from five sites in contrasting settings across this region.
- 66 1.1 Regional context and site description

67 In Scotland, woodland currently constitutes 18% of land cover, 22.5% of which is considered native 68 (Forestry Commission, 2014). This contrasts with the maximum extent of woodland cover around 69 5700 cal BP (calendar years before AD 1950), which has been estimated at 50-60% of the land area 70 (Tipping, 1994; Smout et al., 2005). Abrupt and widespread woodland contraction occurred across 71 northwest Scotland around 4400 cal BP, notably of pine. This is attributed primarily to climate 72 deterioration and resulted in the contraction of pine to near its current range (Fig. 1) (Bennett, 73 1995). For four millennia these woodland fragments have existed within a matrix of blanket peat and 74 heath, with small and dispersed areas of agriculture. The surviving climate relicts are highly valued 75 and form the basis for national and site-based woodland conservation and expansion goals.

76 These high conservation value fragments include the present study site, Ledmore and Migdale 77 National Nature Reserve (NNR) (Fig. 1). It comprises a range of habitats including Quercus (oak) and 78 Betula (birch) woodland, semi-natural P. sylvestris woods, open dwarf shrub heath and mire 79 communities (see Supplementary material: Table A1). These include 95 ha 'old' sessile oakwoods at 80 their northerly limits in Britain, characterised by an acidophilous heath understorey more commonly 81 associated with pinewoods. The 144 ha pinewood on which our study focuses includes 'ancient 82 pinewood indicator' species of orchids, lichens and invertebrates which suggest long-established 83 pine communities (Woodland Trust Scotland, 2015). The 6.9 km² site is topographically diverse, 84 rising from sea level to 228 m OD. Management goals include conserving the distinctive biodiversity 85 mosaic and the ancient woodlands, improving natural regeneration and expanding native woodland 86 cover to form a regional network that increases resilience to climate change impacts on species

87 ranges (Woodland Trust Scotland, 2015). A limited range of management interventions is advocated,

- 88 focused on reducing threats (e.g. thinning forestry plantation to remove exotic species and stimulate
- 89 native tree regeneration).

90 The biogeographical position and composition of the site raise numerous uncertainties about 91 community resilience and appropriate models for management. Maclean et al. (2014) suggest that 92 landscapes with high refugial potential, notably biophysical heterogeneity, support more stable and 93 qualitatively different plant assemblages from those in surrounding regions with lower topographic 94 and climatic heterogeneity. We hypothesise that the comparatively sheltered, east-facing aspect of 95 the study site, and local edaphic and topoclimatic heterogeneity enabled woodland persistence 96 through the Holocene, in contrast with more homogeneous and exposed conditions in adjacent river 97 valleys and upland plateaux. Within this proposed microclimate refugium, the continuity of 98 particular species and origins of current communities are unclear. The NNR lies on the northern edge 99 of current native woodland distribution and equidistant between westerly regions which underwent 100 extensive woodland contraction and easterly woods which show greater continuity of cover. 101 Furthermore, local pine communities suggest affinities with upland woods, while the presence of 102 oak suggests affinities with woods on the Highland fringe (Tipping, 1994), and it is unclear which 103 context provides an appropriate model for assessing and predicting community behaviour.

104 To examine these questions and understand the significance of microclimatic conditions for

105 woodland resilience, the vegetation history from Migdale pinewood is compared with

106 palaeoecological data from two contrasting sites in neighbouring catchments (upland Reidh-lochan

- and low-lying Reidchalmai), and two sites selected to represent the dominant regional upland
- 108 trends: pinewood continuity (Loch an Amair) and mid-Holocene woodland contraction (Torran
- 109 Beithe). The comparatively small diameter of these five sites (Table 1) means that they are sensitive
- to pollen input and thus vegetation dynamics within 50 to a few hundred metres around each site
- 111 (Jacobson and Bradshaw, 1981). When discussing individual sites, we thus use the term
- 112 'microclimate' to refer to vegetation and environmental variability on a sub-landscape scale, at
- 113 which topographic factors can create suitable conditions for localised tree populations and
- 114 woodland communities to survive potentially unfavourable regional climatic regimes (sensu
- Dobrowski, 2011). Modern climate data for the sites is limited since the weather station network is
- sparse in the Highlands, but interpolated data allow us to identify broad rainfall and oceanicity contrasts and temperature similarities between the sites (Table 1) (Averis et al. 2004).
- 118 2 Methods

119 Field sampling, laboratory procedures and statistical methods are described for the Migdale analysis 120 site. Table 1 provides published references detailing the methods used at the four comparative sites. A peat core was extracted from the edge of a valley mire at Migdale, adjacent to mature pinewoods. 121 122 Trees currently grow on the peat surface and woody material preserved in the stratigraphy indicates 123 that they have done so in the past. The full depth of peat was sampled using a closed-chamber 124 Russian peat corer to avoid contamination (Jowsey, 1966). To compare the palynological diversity of 125 modern and past assemblages, pollen was extracted from moss foliage that forms the current 126 ground cover in seventeen vegetation communities around the NNR (Table A1, Fig. A1). Sediment 127 stratigraphic description (Table A2) and pollen analysis follow standard techniques (Moore et al., 128 1991). Pollen and spore nomenclature follow Bennett (1994), with the exception of Sorbus-type (e.g. 129 rowan) (Boyd and Dickson, 1987) and Corylus avellana/Myrica gale (hazel/bog myrtle) (Moore et al., 130 1991). P. sylvestris stomata were identified on pollen slides following Sweeney (2004). The pollen 131 sum consists of a minimum of 500 land pollen grains, excluding aquatics and spores (total land 132 pollen: TLP), although total counts for 18 of the 79 samples were lower due to low pollen

- 133 concentrations (minimum 322 TLP). Values are expressed as a percentage of TLP (for land pollen) or
- 134 TLP + taxon/group (for spores). Microscopic charcoal fragments >10 μ m were tallied on pollen slides.
- 135 Selected percentage data for pollen and spores, and influx data for pollen, pine stomata and
- 136 charcoal are presented (Fig. 2-3). Local pollen assemblage zones that group assemblages of similar
- 137 composition were defined using CONISS (Grimm, 1987). To compare Migdale with the four other
- sites, percentage pollen data are shown for three main arboreal taxa (*Betula, Pinus, Quercus*) to
- examine changes in their relative abundance, while *Pinus* stomata and pollen influx data provide
- proxies for local growth and vegetative population biomass dynamics, respectively (Fig. 3) (Parshall,
 1999; Seppä et al., 2009).
- 141 1999, Seppa et al., 2009).
- 142 To improve chronological comparability, age models were produced for all five sites using the
- 143 Intcal13 calibration curve and classical age-depth modelling techniques (CLAM) (Blaauw, 2010;
- 144 Reimer et al., 2013). The Migdale chronology was constructed from twelve AMS radiocarbon dates
- 145 (Table A3), with time-depth curves at the other four sites constructed from 6-10 radiocarbon dates
- 146 (Table 1). Calendar ages (cal BP) are used throughout, where 0 cal BP = AD 1950.
- 147 Migdale data were analysed using principal components analysis (PCA) and rarefaction. The
- ordination displays changes in assemblage composition and stability through time. Surface pollen
- samples were included as passive samples in the fossil PCA to compare present and past assemblage
- 150 composition (Fig. 4a). Gradient length in an initial detrended correspondence analysis was <2 SD,
- indicating that linear response models are appropriate. Only taxa with a value of \geq 2 % in at least one
- sample were included to avoid rare types biasing the analysis. Data were recalculated to a sum of
- 153 100% and square-root transformed prior to analysis to stabilise variance. Ordinations were carried
- 154 out in Canoco 4.5 (Ter Braak and Smilauer, 2002). Rarefaction analysis provides a robust measure of
- 155 palynological richness (Birks et al., 2016). This was applied to all TLP pollen taxa and implemented in
- psimpoll 3.0 (Bennett, 1998) based on a rarefied sum of 300 TLP (Fig. 4b).
- 157 In the absence of a comprehensive Holocene palaeoclimate synthesis for Scotland, major shifts in
- 158 temperature and/or moisture identified in northern Britain and north-west Europe are summarised
- in Fig. 3 to assess the effect of climate change on woodland dynamics (Anderson et al., 1998; Barber
- and Langdon 2007; Barber et al., 2013; Charman et al., 2006; Charman 2010; Seppä et al., 2009;
- 161 Tipping et al., 2012). This draws on a range of palaeoclimate proxies, primarily humification, testate
- amoebae, chironomids, plant macrofossils and tree-ring widths.
- 163
- 164 3 Results and interpretation
- 165 3.1 Chronology
- 166 The age-depth models underpinning the chronology for each site are presented in Fig. A2. At all
- sites, the radiocarbon dates produced a conformable sequence, with no indications of sediment
- reworking or prolonged hiatuses in sediment accumulation. The age-depth plots indicate that
- sedimentation rates changed through time, likely as a result of a range of bathymetric, catchment
- and climatic factors that influence sediment accumulation. These changes are not discussed in any
- 171 detail since our focus is on vegetation dynamics.
- 172 3.2 Migdale stand-scale succession and dynamics

- 173 Over the last 7790 years cal BP five phases of vegetation compositional stability and transition are
- identified from the pollen zonation (Fig. 2) and PCA analyses (Fig. 4a), as summarised in Table 2.
- 175 Limited overlap between PCA phases indicates significant shifts in woodland composition,
- punctuated by periods of community stability. In brief, the local community was dominated by
- 177 Betula and Pinus (7790-6000 cal BP), Alnus (alder) (6000-4400 cal BP), Betula (4400-600 cal BP), and
- then *Pinus* and *Betula* (600-0 cal BP, AD 2001). Palynological richness fluctuates largely below mean
- Holocene values until c. 4360 cal BP (Fig. 4b). Sustained higher palynological richness from c. 2290-
- 670 cal BP coincides with higher pollen abundance for ruderal taxa. Richness values decline strongly
 to the present, as *Pinus* becomes the dominant pollen producer. Rarefaction values for surface
- 182 samples overlap with subfossil values prior to c. 4360 cal BP (zone MIG1) and since 610 cal BP (zone
- 183 MIG3), but, with one exception, are consistently below intervening values.

184 3.3 Regional range dynamics

185 Inter-sample variability in pollen diagrams, particularly at small sampling sites, is a product of 186 taphonomy and vegetation dynamics immediately around the sampling site (e.g. stand-scale shifts in species distribution or abundance) with smaller contributions from regional pollen production 187 188 (Bradshaw, 2013). To understand climatic influences on tree regeneration across a heterogeneous 189 landscape, we focus on sustained trends as the basis for comparison, rather than finer-resolution 190 variability likely to relate to gap-phase dynamics. Following rapid post-glacial climate amelioration 191 after c. 11 700 cal BP, similar early Holocene woodland succession patterns are evident at 192 Reidchalmai, Loch an Amair and Torran Beithe, where Betula expansion was followed by an increase in Pinus (Fig. 3). As pine pollen is widely dispersed, macrofossil and stomatal evidence is needed to 193 194 securely differentiate local growth from regional pollen influx (Froyd, 2005). Pinus stomata are 195 recorded at Loch an Amair from c. 9900 cal BP, with corresponding pine pollen abundance of only 196 1%, indicating small local populations which are difficult to identify from pollen data alone. There are 197 insufficient sites with stomatal analyses to assess whether small populations were common before 198 observed regional increases in pine pollen. Using 20-25% pollen as a conservative limit for inferring 199 local growth (Bennett, 1984, 1995), pine populations were established by c. 8500 cal BP at Reidh-200 lochan and c. 7700 cal BP at Reidchalmai. After c. 7500 cal BP, pine percentages at Migdale are 201 higher than neighbouring catchments and comparable with values at the regional sites until c. 6000 202 cal BP. Although stomata are absent from the sedimentary sequence, this could suggest pine growth 203 at Migdale.

204

205 With the exception of Loch an Amair, Pinus abundance falls below ~20% by c. 6300-6000 cal BP, 206 coinciding with percentage and influx increases in deciduous taxa, particularly Alnus and Quercus. 207 Alnus values are highest at Migdale, indicating localised or dispersed growth at the other sites 208 (Bennett and Birks, 1990; Froyd & Bennett, 2006; Tipping & McCulloch, 2003). Quercus values reach 209 the 2% TLP level thought to indicate local growth by c. 8000 cal BP (Huntley and Birks, 1983). 210 Although similar values are not recorded at Migdale until c. 5700 cal BP, only at this site do values 211 exceed 10%, which suggests that oak was a significant vegetation component at c. 4900-4770 cal BP 212 (Huntley and Birks, 1983). This overlaps with increased pine representation at Migdale, Reidchalmai 213 and Torran Beithe from c. 5100-4100 cal BP. Sustained reductions in Pinus are recorded at all sites except Loch an Amair from c. 4600-4100 cal BP, although pine stomata persist at some sites after c. 214 215 4100 cal BP, suggesting that small populations remained around sites with (Torran Beithe) and 216 without (Reidh-lochan) a pronounced pine decline until c. 3200-2600 cal BP. 217

Two mid-late Holocene features differentiate Migdale from the other sites: (1) the marked rise in Betula values after c. 4400 cal BP contrasts with relative continuity of pine at Loch an Amair and

- birch at Reidchalmai, and (2) the strong rise in influx and percentage *Pinus* values from c. 1180 cal BP
- is absent from the other sites. The increase in pine influx at Reidchalmai from c. 1340 cal BP likely
- reflects complex fluvial inputs and is not a species-specific response (Tipping and McCulloch, 2003).
- 223
- 4 Discussion

4.1 Microclimate effects on woodland biogeography, turnover and resilience

226 We identify three inter-related factors that influenced woodland persistence: regional climate 227 gradients, landscape-scale topographic and altitudinal factors, and microclimate heterogeneity (that 228 is, variability within each pollen catchment). While woodland dynamics were shaped by time-229 transgressive changes associated with postglacial population colonisation and succession, and 230 regional synchronisation due to climate change, the outcomes were spatially variable as a result of 231 finer-scale topoclimatic heterogeneity. Pinus was an early canopy dominant or co-dominant across 232 the Highlands, but variations in pollen abundance over time and among sites indicate climatic and 233 local constraints on population distribution. The pine population appears to have been 234 discontinuous around Migdale, with low representation in upland and valley settings (Reidh-lochan, 235 Reidchalmai), a persistent early decline at Reidh-lochan from c. 7200 cal BP, and higher values at 236 Migdale and in nearby upland sites at Loch Farlary and Achany Glen (Fig. 1) (Smith, 1996; Tipping et 237 al., 2008b). On a national scale, pine became increasingly restricted to upland habitats from c. 8200 238 cal BP (Bennett, 1984), but even here it is likely to have faced constraints. Independent peat 239 stratigraphic and radiocarbon data indicate the spread of blanket peat before 6000 cal BP and pollen 240 data show the spread of birch, both of which are likely to have constrained pine growth, particularly 241 near northern range edges (Carlisle and Brown, 1968; Gallego-Sala et al., 2016; Tipping, 2008). 242 Although the rate of spread and pollen abundance of Quercus declined as it reached its northern 243 climatic and altitudinal limits, higher pollen frequencies around the Highland fringes suggest that 244 populations were established in sheltered, lower-lying locations (Tipping, 1994). This restricted the 245 realised niche of pine in sheltered valleys at Migdale, Reidchalmai and Achany Glen from c. 6000 cal 246 BP (Smith, 1996).

247 Numerous studies note a correspondence between reductions in pine representation and shifts to 248 wetter climatic conditions, based on independent reconstructions of lake levels and peatland 249 watertables (Anderson et al., 1998; Bridge et al., 1990). Regeneration in marginalised pine 250 populations thus appears to have been synchronised at a regional scale by climate change, but the 251 mechanism of population regulation varied. In the uplands, wetter conditions may have reduced 252 pine regeneration, indicated by declining pollen and macrofossil abundances c. 6500-6000 cal BP (Bridge et al., 1990). At lower altitudes, wetter climate may have contributed to a rise in water-253 254 tables which allowed Alnus to outcompete Pinus in valley mires like Migdale (Bennett and Birks, 255 1990). This combination of climatic, recruitment and competition effects led to extinction-256 recolonization dynamics in pine. This is particularly evident during the mid-Holocene, when 257 macrofossil evidence indicates that pine expanded its range northward around 5400-4200 cal BP in 258 response to lower peatland water-tables (Gear and Huntley, 1991). Stomatal evidence for renewed 259 growth at Reidh-lochan and Loch Farlary (Tipping et al., 2008b) contrasts with a weak pollen influx 260 response and absence of stomata at Migdale (Fig. 3). This suggests that pine colonised drier upland 261 peat surfaces, but gained little advantage in valleys where peat was limited and pine remained 262 subject to competitive exclusion by deciduous taxa and possibly by human impacts.

Anthropogenic disturbance may have selectively advantaged deciduous taxa in sheltered settings.
 Migdale is differentiated from the other sites after c. 5700 cal BP by late expansion and unusually
 high representation of *Quercus* compared with adjacent valleys and regional trends (Fig. 3). Neolithic

farming and selective management is considered causal, inferred from a temporary rise in Poaceae and the occurrence of cereal type and *Rumex* (dock) pollen, with similar disturbance recorded in

268 nearby valleys at Reidchalmai and Achany Glen from c. 5600 cal BP (Smith, 1996). Increased light

- 269 penetration and managed browsing may have allowed oak to replace shorter-lived deciduous trees
- and shrubs, although disturbance was probably low intensity since woodland cover was maintained.
- 271 Drier/warmer climatic conditions during this period could have increased the rate of oak growth in
- 272 this favourable microclimate setting.

273 Stronger inter-site contrasts emerge during the mid-Holocene, with extensive loss of upland 274 pinewoods and fragmentary woodland survival in upland and low-lying catchments (Fig. 3). Climate 275 deterioration, particularly increased wetness, is considered to be a key driver of pine dieback at a 276 regional scale, but the spatial differentiation of relict tree cover into upland pine (Loch an Amair) and 277 broadleaved valley woods (Migdale, Reidchalmai) indicates the need for more complex explanatory 278 mechanisms than rising water tables alone, particularly in low-lying areas with limited peat cover 279 (Bennett, 1995). Steeper slopes, unsuited to blanket peat expansion, and a less oceanic climate may 280 explain Pinus survival around Loch an Amair and in the northeast, respectively (Froyd and Bennett, 281 2006; Tipping, 1994). At Migdale, potential drivers of the transition from alder to birch-dominated 282 woods around 4430 cal BP include poorly understood aspects of climate change like seasonality, 283 which may have created conditions suited to birch growth, rather than persistently humid conditions 284 that previously favoured alder (McVean, 1956). Climate deterioration may also have altered competition outcomes, contributing to reduced recruitment and competitive ability in oak and 285 286 allowing birch to replace it, as occurs now near oak range limits (Atkinson, 1992; Jeffers et al., 2015). 287 Locally increased anthropogenic disturbance from c. 4800 cal BP could have favoured birch over 288 alder (Barthelmes et al., 2010). At all sites in this study where woodland survived the mid-Holocene 289 'collapse', regeneration persisted through subsequent anthropogenic activity. This suggests modest 290 or managed disturbance, and that woodland regeneration was not near a critical threshold (cf. 291 Scheffer et al., 2012).

292 The current mosaic of pine- and oakwoods and open communities that differentiates Ledmore and 293 Migdale NNR from other 'ancient' woodland fragments emerged during the last c. 1000 years. Pine 294 re-expansion takes place in the context of regionally low pine abundance across the northwest. It is 295 not possible to disprove the survival of some individuals at Migdale throughout the Holocene, but 296 low pine pollen values (<7%) at Reidchalmai and Reidh-lochan suggest that there were no sizeable 297 populations at or around Migdale from c. 4170-1180 cal BP, particularly after c. 3330-2600 cal BP 298 when stomata disappear from upland pine decline sites. The replacement of deciduous taxa, apart 299 from birch, by pine after 1060 cal BP and a decline in herbaceous diversity at c. 620 cal BP are 300 interpreted as indicators of silvicultural management, with deliberate selection for pine, probably by 301 planting (Mills and Crone, 2012). Pine abundance had increased further by the nineteenth century, 302 indicated by the presence of fossil fuel-derived spheroidal carbonaceous particles (SCPs) (Rose and 303 Appleby, 2005). This corresponds with local and regional evidence of intensive timber management 304 (Bangor-Jones, 2002; Rydval et al., 2015). This shift represents the decoupling of pine dynamics from 305 climate fluctuations that previously governed local and regional population fluxes and stand 306 composition.

307 4.2 Microclimate buffering and management implications for *Pinus sylvestris*

308 Migdale represents a rare example of continuous deciduous-coniferous woodland cover from the

- and early Holocene through to the present, possibly owing to comparative shelter from westerly climate
- 310 systems. This provided a favourable microclimate which allowed woods to withstand multiple
- climate and disturbance shifts. Uneven topography and varied drainage also may have made the site

less suited to farming, which remains a feature of the wider valley floor at Reidchalmai. This is good

- news for conservation. Although canopy cover was maintained as a result of favourable growing
- conditions and limited farming, community composition was far from stable. In terms of
- 315 management, woodland resilience therefore depends on allowing composition to adapt to changing
- conditions and on appropriate ecological models for anticipating change. Community replacement
- and reassembly in the last c. 1000 years has created floristic affinities between Migdale and long established pinewoods, overriding earlier similarity with deciduous fragments in similar low-lying
- 319 valleys. Pinewood communities thus provide an inappropriate model for anticipating change at this
- 320 site. Challenging existing ecological models and allowing adaptive ecological responses introduces
- 321 uncertainties that may conflict with conservation targets, like the desire to conserve the distinctive
- 322 biodiversity of these woodlands (Hiers et al., 2016; Woodland Trust Scotland, 2015). In view of the
- 323 uncertainties surrounding climate change, long-term evidence emphasises the importance of shifting
- 324 conservation and management focus from compositional stability to functional viability.

While the microclimate at Migdale buffered woodlands against environmental change and mature 325 326 trees around the site demonstrate that conditions are suitable for pine growth, the ecological history 327 of this site suggests that continued community change is highly likely and, over the longer term, local 328 pine populations are probably transient. The transition to pine dominance from c. 1060 cal BP has 329 resulted in a prolonged decline in diversity and the existing pine-dominated stand may still be in a 330 state of flux, characterised by high levels of compositional change more typical of the early Holocene (Fig. 4) (Froyd and Bennett, 2006; Seddon et al., 2015; Tipping et al., 2006). Predicted future milder 331 332 climatic conditions may allow broadleaved species like birch, oak and rowan to expand, thus 333 replaying the successional replacement of pine seen at all except marginal, peat-dominated sites 334 over the course of the Holocene. Birch is also likely to increase due to more wind disturbance (Ray, 335 2008); this is a potential outcome of recent storm damage to pines around Loch Migdale. Scattered 336 pines occur on blanket peat in higher areas of the NNR and sheltered valleys like Migdale may 337 continue to act as seed sources for tree colonisation in adjacent upland areas if environmental 338 conditions, deer numbers and cultural preferences allow.

339 In terms of diversity baselines, current palynological richness across the NNR is low relative to values 340 during Iron Age and Dark Age settlement periods (c. 2290-670 cal BP) and comparable with the early 341 Holocene range of variability (Fig. 4b). High diversity during the woodland grazing period indicates 342 the potential benefits of small-scale, low intensity intervention. It also suggests that the current 343 strategy of thinning planted woods to stimulate natural regeneration could benefit diversity in 344 longer-established stands. Both the early and late Holocene periods of lower diversity correspond 345 with unstable, possibly transitional, assemblages. This emphasises the need for managers to 346 anticipate and manage for change. Surface sample ordination scores indicate high spatial diversity 347 across the NNR (Fig. 4a) and, using space-for-time substitution, this suggests that maintaining spatial 348 heterogeneity across the site, which is one of the current management goals, can help support a 349 dynamic mosaic.

350 The relatively recent origins of the pinewood raise questions about the diagnostic value of the 351 'ancient pinewood indicators' present at the site (Whittet and Ellis, 2013). It appears that continuous 352 canopy cover, rather than the persistence of pine per se, helped maintain distinctive understorey 353 diversity by ensuring the availability of humid, shaded microclimates within the NNR (Bradshaw et 354 al., 2015). While debate continues over the biodiversity and ecosystem service benefits of planted 355 versus native coniferous woodland (Quine and Humphrey, 2010), these findings support existing 356 evidence that semi-native woods of uncertain origin and planted ancient woodland sites are useful 357 in conservation (Roche et al., 2015). Therefore, maintaining distinctive biodiversity seems

- 358 compatible with accommodating change in canopy dominants, as long as woodland cover is
- 359 maintained and community reorganisation is expected and accepted as an essential attribute of 360 resilience.
- 361 5.0 Conclusions

362 Palaeoecological evidence from a currently diverse woodland in a sheltered valley on the northern 363 range limits for pine and oak in Scotland demonstrates that communities within favourable 364 microclimate locations show greater continuity of canopy cover and resilience to climate change 365 than upland catchments, but have undergone significant compositional turnover. The suggestion 366 that microclimate variability arising from biophysical heterogeneity promotes more stable plant 367 communities (Keppel et al., 2012; Maclean et al., 2014) is, therefore, only supported if stability is 368 measured in terms of the continuity of woodland cover; it is not true for composition. Continuity of 369 cover allowed 'ancient' woodland indicator taxa to persist despite changes in canopy composition. 370 Favourable microclimatic conditions and topographic variability conferred low suitability for 371 agriculture and buffered tree populations against climatic shifts. However, prior to late Holocene 372 silvicultural intervention, sheltered conditions mitigated against the survival of pine, which may have 373 been out-competed by deciduous taxa. This highlights the need for more data and alternative 374 models of community composition and behaviour to inform ecological understanding and 375 management of microrefugia communities. Further work is also needed to characterise and map the 376 distribution of favourable long-term microclimates at a landscape-scale in order to understand how 377 they influence ecological responses to changing climate and land-use mosaics over long timescales 378 (Valencia et al., 2016). This will enable palaeoecology to contribute more directly to predictive 379 ecology and climate change conservation strategies by helping to evaluate the likely effectiveness of 380 protected areas under changing climate regimes (Hannah et al., 2002; Lindbladh et al., 2013).

381

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- 385
- 386 7.0 References

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Table 1. Location, current vegetation and comparative characteristics for all study sites

Site	Description		
Migdale	Main study site		
pinewood,	Location: 4°15'22" W 57°53'14" N, 40 m OD		
Ledmore &	Sampling site: valley mire edge		
Migdale NNR	Current vegetation : Betula with mire understorey and P. sylvestris and Quercus		
-	within 100 m		
	Current climate: <14 °C July mean temperature but immediately N of 14-15 °C		
	limit, on boundary between <750 mm and 750-1000 mm annual rainfall,		
	comparable index of oceanicity (mean wet days/monthly mean temperature		
	range) to Reidchalmai, Reidh-lochan and Loch an Amair (Averis et al., 2004)		
	Chronology: 12 AMS ¹⁴ C dates		
Reidchalmai,	Comparison: neighbouring low altitude, valley floor catchment with deciduous		
east Sutherland	woodland		
	Location: 4°9'1" W 58°0'12" N, 90 m OD		
	Sampling site: small infilled lake basin, 80-90 m diameter		
	Current vegetation: improved pastoral grassland within heather moorland with		
	Betula-dominated woods to south		
	Current climate: <14 °C July mean temperature, 750-1000 mm annual rainfall,		
	comparable index of oceanicity to Migdale and Reidh-lochan		
	Chronology: 10 AMS ¹⁴ C dates		
	Sources: Tipping & McCulloch 2003, Tipping et al. 2008b		
Reidh-lochan,	Comparison: neighbouring upland catchment, treeless		
east Sutherland	Location: 4°07'26" W 58°02'13" N, 160 m OD		
	Sampling site: small lake, c.100 m diameter		
	Current vegetation: extensive blanket mire, agriculture to the east		
	Current climate: as Reidchalmai		
	Chronology: 6 bulk ¹⁴ C dates		
	Sources: Froyd 2001, Froyd & Bennett 2006		
Loch an Amair,	Comparison: example of upland pinewood continuity		
East Glen Affric	Location: 4°53'25" W 57°17'20" N, 315 m OD		
	Sampling site: small lake, c.100 m diameter		
	Current vegetation: non-native Pinus contorta plantation with P. sylvestris		
	woodland to north		
	Current climate : <14 °C July mean temperature but immediately N of 14-15 °C		
	limit, >1500 mm annual rainfall, comparable index of oceanicity to Migdale,		
	Reidchalmai and Reidh-lochan		
	Chronology: / bulk ¹⁴ C dates		
T D H	Sources: Froya 2001, Froya & Bennett 2006		
Torran Beithe,	Comparison: example of upland pinewood contraction		
West Glen	Location: 5°6'2" W 57° 14'29" N, 265 m OD		
Attric	Sampling site: peat-filled bedrock basin, c.56 m surface diameter		
	Current vegetation: blanket mire		
	Current climate : <14 °C July mean temperature, >1500 mm annual rainfall,		
	nigner index of oceanicity than the other four sites		
	Chronology: 9 AMIS ¹ C dates		
	Sources: Davies 1999, Tipping et al. 2006		

Table 2. Summary of Migdale stand dynamics based on pollen assemblage zones and ordination (PCA) phases. See Fig. 2 for selected pollen data and Fig. 4a
 for ordination plot

Pollen assemblage	Palynological characteristics	Corresponding PCA phase
zone and age		
MIG3: 610-0 cal BP	Renewed expansion of <i>Betula</i> , marked rise in <i>Pinus</i> , very low values for <i>Quercus</i> , <i>Alnus</i>	Phase 5 (430-0 cal BP): shift towards
(AD 2001)	and <i>Corylus, Myrica, Calluna</i> and fern values decline, and herbaceous pollen abundance	Pinus with Betula and away from
	and diversity is reduced. Low charcoal values. Spheroidal carbonaceous particles	species scores for other deciduous
	indicative of fossil fuel burning post-c. AD 1850 occur from 12 cm; extrapolated date of	trees, heath and herb taxa
	310 cal BP (AD 1640) using ¹⁴ C-derived chronology appears too old, likely due to lower	
	decomposition and compaction in upper sediments above youngest radiocarbon date	
MIG2b: 2230-610 cal	Differentiated from zone MIG2a by lower arboreal pollen sums and increased	Weaker correspondence between
BP	abundance of mire taxa (Calluna, Myrica, Sphagnum and Erica). Higher Pinus	zones and phases: zone MIG2b
	percentage and influx values, especially from 1090 cal BP. More abundant disturbance	includes part of PCA phase 3 and all of
	indicators and cereal type pollen (P. lanceolata, Potentilla-type, Asteraceae and	phase 4 (1090-550 cal BP)
	Rumex). Charcoal values rise. More minerogenic peat with fine sand and silt at c. 2120-	
	1790 cal BP	
MIG2a: 4430-2230	Shift to Betula dominance with reductions in the other arboreal taxa, particularly Pinus	Phase 3: shift in sample scores to new
cal BP	and Alnus. Sorbus and Salix are the main exceptions. Myrica gale-type is more	quasi-equilibrium (3990-1180 cal BP)
	abundant, along with Hordeum group, Plantago lanceolata, Potentilla-type and other	
	herbaceous pollen types. Reduced frequencies for <i>Pteridium</i> and Pteropsida spores	
MIG1b: 5940-4430	High Alnus percentage and influx values, peaking around 5130-5050 cal BP, with	Overlaps with phase 2 quasi-stable
cal BP	secondary peaks at c. 5580 cal BP and 4750-4650 cal BP. High total pollen influx and	state (5830-4360 cal BP)
	more rapid peat accumulation. Subsequent alder reduction corresponds with increases	
	in Betula, Pinus and Corylus (5580-5160 cal BP), then Quercus and Poaceae (5020-4770	
	cal BP). Maximum <i>Quercus</i> abundance from 5020-4700 cal BP. Short-lived percentage	
	and influx increased in <i>Pinus</i> values from c. 4840-4600 cal BP. <i>Hordeum</i> group pollen is	
	recorded more frequently from 4800 cal BP	
MIG1a: 7790-5940	High but erratic values for <i>P. sylvestris</i> and <i>Betula</i> , rising <i>Alnus</i> frequencies, increased in	Phase 1 (7790-6050 cal BP):
cal BP	Quercus relative and influx values late in zone, Salix and Calluna values decline. High	characterised by variable sample
	representation for Pteropsida and Pteridium aquilinum spores. Peak charcoal values	scores



Figure 1. Locations of study sites in northern Scotland, with other pollen studies mentioned in text

and current range limits of (a) Caledonian pine forest and (b) old sessile oak woods with *llex* and

Blechnum in UK (not mapped in Republic of Ireland) (source: JNCC)



574 Figure 2. Selected percentage pollen and spore data from Migdale, with influx data for charcoal and spheroidal carbonaceous particles (SCPs). Clear

575 exaggeration curve x10

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Figure 3. Comparison of data from Migdale, Reidchalmai, Reidh-lochan, Loch an Amair and Torran Beithe, showing (a) percentage data (%TLP) for *Betula* and *Quercus* and (b) percentage (%TLP) and influx (pollen grains or stomata cm⁻² yr⁻¹) data for *Pinus* pollen and stomata (stomata unavailable for Reidchalmai), with a qualitative summary of the main climate shifts (see text for references)



Figure 4. (a) Principal components analysis and (b) palynological richness data for Migdale with 95% confidence intervals, both showing five main communities (phases). Surface samples (1-17) passively ordinated on PCA and plotted against fossil rarefaction values. Taxon abbreviations: Aln – *Alnus*, Bet – *Betula*, Call – *Calluna vulgaris*, Cyp – Cyperaceae, Myr – *Myrica gale*-type, Pin – *Pinus sylvestyris*, Poac – Poaceae, Que – *Quercus*, Rub – Rubiaceae. The first and second principal components (eigenvalues) account for 51.1% and 26.6% of the variance, respectively