North Fennoscandian mountain forests: History, composition, disturbance dynamics and the unpredictable future

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北芬诺斯坎迪亚山地森林分布于斯堪的纳维亚山脉之间，以及挪威北部、瑞典和芬兰，以及紧邻的西北俄罗斯。这些森林在历史上和现有的干扰模式下，包括慢性和急性干扰，以及其强度和频率，以及与植被和地貌的差异。由于其偏远性，生长缓慢和使用的限制，这些山地森林一般不受人类利用的影响，比更易利用的森林在较低海拔和/或纬度区。因此，这些北欧的森林常被称为“欧洲的最后荒野”，即使在人类影响的强度上也已普遍存在。由于其自然性，北芬诺斯坎迪亚山地森林对生物多样性保护、生态系统变化的监测以及社会文化价值观至关重要。

因此，它们也为基本和应用研究提供了独特的参考区域，以及为整个芬诺斯坎地发展森林保护、恢复和基于生态系统的管理方法。然而，当前气候变化的快速变化预计会严重影响这些森林的生态学和动态。
1. Introduction

North Fennoscandian mountain forests are distributed along the remains of ancient mountain ranges, most notably the Scandines mountain range along the border between Sweden and Norway, and the low-mountain regions of northern and northeastern Finland, and the adjacent northwestern Russia (Fig. 1). These regions harbor both northern boreal forests and subarctic/subalpine birch forests, which are collectively henceforward referred to as northern mountain forests. Because of remoteness, low productivity and restrictions of use, the overall human impact on these forests has generally remained significantly lower compared with more southern boreal forests at lower latitudes and/or elevations (Veijola, 1998). However, over the entire region, some low-intensity human use of the forest has occurred for millennia, such as the influence of the Sápmi people and their reindeer herding culture (Josefsson et al., 2009). Reindeer herding continues to be a culturally important land use, which has recently also generated conflicts with other land uses like forestry (Helle and Jaakkola, 2008; Berg et al., 2008).

The exact degree of human impact, i.e. the extent to which the current forests deviate from a natural state, varies strongly across forest types and landscapes, from negligible to local deforestation (Östlund et al., 2015). Nevertheless, these mountain forests still harbor some of the largest areas of relatively natural boreal forest in Northwestern Europe (Aksenov et al., 2014). By natural forest we mean a forest which structure, species composition and processes have not been significantly altered by human activities (Brumelis et al. 2011). Much of our current understanding of natural forest disturbance and successional dynamics and their impact on biodiversity in Fennoscandian conditions is derived from these forests (Kuuluvainen and Aakala, 2011). The ongoing climate change,
with a predicted mean annual increase of mean annual temperature by 4°C and precipitation
by 20% by the end of the 21st century (IPCC, 2013), is likely to profoundly affect the ecology,
dynamics and productivity of these forests in the future (Kellomäki et al., 2008; IPCC, 2013;
Gauthier et al., 2015).

The ecological character and biological diversity of northern mountain forests reflect
to a large extent their Holocene climate and vegetation history. Because of their northern
position, continental ice retreated from these areas as late as around 15-10000 years before
present (Parducci et al., 2012). Tree species colonized the region from different directions,
following the course of retreat of the ice sheet. The developing early Holocene tree
communities were accordingly characterized by the forests surviving the glaciation period
and surrounding the retreating ice to the west, south and east (Kullman, 2008; Paus et al.,
2011; Parducci et al., 2012). Current dominant boreal tree species, pine (Pinus sylvestris L.),
spruce (Picea abies (L.)Karst.) and birch (Betula pubescens Ehrh.), all arrived in early
Holocene 14-11 kyr B.P in the first de-glaciated areas of western Fennoscandia (Kullman,
2008). During the mid-Holocene climatic optimum, thermophilous deciduous tree species
like elm and oak occurred in the region.

The historic timing of formation of boreal tree species dominance at stand or regional
scales differs between species, with birch and pine predating spruce (Kullman, 2008). The
spread and development of regional dominance of spruce has occurred in the last ca. 3000
years, in parallel with long-term climate changes favoring spruce but disfavoring
thermophilous tree species (Kullman, 2001; Giesecke and Bennett, 2004; Bradshaw and
Lindblad, 2005). Concurrently with increasing spruce dominance, high elevation and high
latitude forests retreated downhill and southwards (Payette and Lavoie, 1994; Kullman,
1995). This process was reversed after the termination of the Little Ice Age (Grove, 1988) in
The rate of this recent regain of forest area differs somewhat between geographical regions and between tree species, but has generally accounted for ca. 0.6 vertical meters per year in the northeastern part of the mountain forest region (Aakala et al., 2014; Mathisen et al., 2014) and 0.9 vertical meters per year in the southern part (Kullman and Öberg, 2009). Latitudinal advance has occurred with ca. 150 m per year for birch forest and ca. 70 m per year for pine forests (Hofgaard et al., 2013).

The impact of climate change on forests close to their bioclimatic range limits entail a complex web of both gradual and episodic abiotic and biotic processes (Hofgaard, 1997; Scheffer et al., 2012). The warming climate changes ecosystem structure and composition through changing competitive relationships among the constituent tree species (Kellomäki et al., 2008). Climatic conditions are changing too fast for long-lived plant species such as trees to respond through natural migration, and this is predicted to increase the probability and severity of disturbance events. In addition other change rate-related developments, such as extreme weather events and insect outbreaks are likely to become more common (IPCC, 2013; Gauthier et al., 2015). These perturbations could counteract or reinforce climate-driven changes of the northern mountain forest ecosystems (Hofgaard, 1997; Koven, 2013; Bright et al., 2014). There are historical and modern examples where perturbations, such as cuttings, forest fires or insect outbreaks have pushed northern mountain forest ecosystems to an alternative, non-forested stable state (e.g. Neuvonen et al., 2005; Karlsson et al., 2009; Östlund et al., 2015).

A large share of the remaining natural Fennoscandian mountain forests are protected. These forests together with other forests with long history of low human interference and restrictions of utilization, have provided unique possibilities for fundamental research on the ecology of primeval forests at various scales (Kuuluvainen and
Aakala, 2011). Thus, much of our current understanding of natural forest disturbance and
successional dynamics, and biodiversity concerning Fennoscandian boreal forests as a whole
is derived from these northern mountain forests.

The main purpose of this paper is to review the current understanding of the
dynamics and disturbance ecology of North Fennoscandian mountain forests and to discuss
the implications of this knowledge on forest conservation, restoration and ecosystem-based
management in the context of ongoing climate change. Specifically, we asked: 1) What is the
value of understanding of past ecosystem dynamics considering future conditions? 2) How resilient
are the mountain forests in a changing environment? 3) What are the main shortcomings of
knowledge concerning the ecology of mountain forests? 4) What are the main lessons learned from
ecological research concerning conservation and ecosystem based management?

2. Definition, geographic distribution, environmental conditions and forest communities

Due to the location of the region at high latitudes, forests considered as mountain forests
are growing at lower elevations than elsewhere in Europe. Hence, we loosely define
northern mountain forests as those prevailing in the mountainous region and growing close
to the elevational and latitudinal limit of forest distribution. However, tree line areas are
excluded from this review. Forest is defined as a tree community with minimum tree height
of 5 m and at least 10% of canopy cover (FAO, 2010).

In this paper North Fennoscandian mountain forests are defined to stretch
latitudinally from southern Swedish Lapland at 64°N to 69°N in northern Norway, and
longitudinally from 14°E in Norway to 34°E in the Kola Peninsula, Russia (Fig. 1). In the west,
the Scandes Mountains rise to height of 1000-2100 m above sea level (a.s.l.). The area east
of the northern Scandes Mountains through northern Finland and to Kola Peninsula is
characterized by gently rolling hills (fells), which rarely reach heights above 1000 m a.s.l. The bedrock in the area is mostly composed of ancient Precambrian granites and gneisses, covered by Quaternary and younger Holocene sediments, mainly podzolized moraines (Lidmar-Bergström and Näslund, 2005).

North Fennoscandian mountain forests are unique in that, due to the warming effect of the golf-stream, they grow further north than any other mountain forests. Compared to more southern mountain forests, the light climate is characterized by stronger seasonal changes, with long days in the summer and several months of short days in the winter. The extreme annual variation in solar radiation is reflected in below-zero temperatures and snow cover which prevail for 7-8 months. Also, low sun angles are prevalent for part of the growing season, which emphasizes the effect of topography on the distribution of solar energy, e.g. on south vs. north facing slopes.

The climatic differences within the region are further shaped by the location between the Atlantic and Arctic oceans, and the Eurasian continent. The climate ranges from markedly or slightly oceanic close to the Norwegian coast (Moen, 1999), to variably continental in different areas of central and eastern parts of northern Fennoscandia. This gives rise to large differences in precipitation, which varies from 1000 mm in the western part to 450-500 mm in northern Finnish Lapland and 300-400 mm in more continental parts of northern Norway and Sweden (Moen, 1999; Tikkanen, 2005). A considerable share of the precipitation falls as snow during the winter months.

Although mixed stands also occur, forests are regionally differentiated into spruce, pine or birch dominance due to topography and edaphic conditions and climate regimes. In general, xeric sites are dominated by pine, while mesic sites follow a post-disturbance successional trajectory from birch as a pioneer species, to the dominance of spruce. At
smaller-scales and on sites capable of supporting any of the three main species, tree species
dominance is influenced by within-region characteristics and variation in the disturbance
regimes.

In general, North Fennoscandian mountain forests can be considered as among the
last remnants of natural forests in Europe. Here we adopt the view of naturalness described
by Brumelis et al., (2011) which puts focus on structures, species and processes as the three
core dimensions of forest naturalness. For a large part of the Fennoscandian mountain
forests, trees of all species and wide age ranges occur, volumes of dead wood, species
richness among plants, fungi and animals are high and natural disturbance processes are still
active. As a large fraction of boreal forest species in general are dependent on dead wood
(Stokland et al., 2012), the mountains forests represent important core area for many
species declining elsewhere in Fennoscandian boreal forests.

2.1. Scots pine dominated forests

Scots pine dominated forests in North Fennoscandia often occur on rocky or dry
sandy and moraine soils, on glaciofluvial soils, on eskers and on delta formations (Rauniö et
al., 2008). These forests are generally open and exhibit significant variability in structure and
successional stages as well as small-scale spatial heterogeneity (Aaltonen, 1919). In spite of
their openness, belowground competition plays a major role in tree regeneration and stand
dynamics (Kuuluvainen and Ylläsjärvi, 2011). Mixed-severity fires have historically been one
driver of forest dynamics (Aaltonen, 1917; Hesselman, 1917), although fire cycles have been
longer than previously assumed, up to hundreds of years (Wallenius et al., 2010). Even non-
pyrogenic pine dominated forests exist, but at a low frequency (Zackrisson et al., 1995).
Species dominating the forest floor include dwarf shrubs *Calluna vulgaris*, *Vaccinium vitis-idaea* and *Empetrum hermaphroditum*, and in the bottom layer mosses and lichens such as *Pleurozium schreberi*, *Cladonia arbuscula* and *C. rangiferina*.

The degree of naturalness has been declining during recent decades particularly in the pine-dominated lichen-rich forests in many regions of the mountain forests. For example, in northern Finland, this is associated with factors such as reduction of old forests due to forestry, and increasing trampling and grazing by reindeer which accelerate nutrient cycling (Raunio et al., 2008). In addition, although fire cycles tend to be long in these northern pine forests (see below), the century-long fire suppression represents a loss of naturalness (Drobyshev et al., 2014). These factors are also likely future threats to this habitat type, in addition to effects of climate change (Gauthier et al., 2015).

A prominent feature in natural pine forests is the presence of large standing dead and bark-less snags, so called kelo trees (*sensu* Niemelä et al., 2002, Fig. 2a). These trees have commonly survived a history of repeated surface fires, and hence their wood has a characteristic high resin content, and are thus highly resistant to decay (Niemelä et al., 2002). As such they represent in some cases a forest structure developing over millennia—as living trees for more than 500 years, and as snags for up to several centuries, before finally falling to the ground and slowly decomposing.

2.2. Norway spruce dominated forests

Norway spruce dominated forests cover extensive areas in North Fennoscandian Mountains. These forests are found on mesic to moist sites with varying fertility from productive herb-rich sites to poorer sites where mosses and dwarf shrubs (predominantly
Vaccinium myrtillus L.) dominate the forest floor, and accumulation of poorly decomposed raw humus is prominent (Fig. 2b, Arnborg, 1943; Havas and Kubin, 1981). As fire is rare in Norway spruce dominated forests (Hörnberg et al. 2011; Ohlson et al. 2011), late successional forest dynamics is characterized by gap or patch scale disturbances and regeneration (Hofgaard 1993a, b; Edman et al., 2007; Aakala and Keto-Tokoi, 2011). Disturbance agents often include wind in combination with fungal infections, insects and snow (Edman et al., 2007; Lännepää et al., 2008). A prominent feature among dying spruce is a general senescence prior to mortality, with either persistent slow growth or steadily declining growth in the decades preceding their deaths (Fraver et al. 2008; Lännepää et al., 2008).

In the spruce dominated forests the establishment of tree seedlings is often physically impeded by the ubiquitous mat of dwarf shrubs, grasses, mosses and a layer of poorly decomposed humus, which decreases nutrient availability (Arnborg, 1943; Sirén, 1955). Hence regeneration is strongly dependent on the availability of suitable substrates created by local gap or patch disturbances, such as mineral soil exposed by tree falls and uprootings, and decaying logs on the forest floor (Hofgaard, 1993a, b; Hörnberg et al. 1997; Caron et al., 2009; Grenfell et al., 2011). For spruce, individual decaying logs may remain important regeneration substrate for as long as 150 years (Hofgaard, 1993a). This long-term suitability of logs as regeneration substrate counter balance the low frequency production of viable seeds, with good seed years occurring typically many years apart (Hofgaard, 1993c).

Spruce may also reproduce vegetatively through layering and form long-lasting clones (Öberg and Kullman, 2011). This is in particular a characteristic in abiotic harsh environments such as the forest-tundra transition and in coastal areas. Overall, the diversity of microsites
created by tree falls provide opportunities for recruitment of other tree species in addition to spruce.

Accordingly, late successional spruce forest generally maintain some proportion of birch admixture (Steijlen and Zackrisson, 1987).

Forest fires are rare in spruce-dominated forests (Ohlson et al. 2011), due to the low flammability of the thick humus layer, which retains moisture for extended periods of time (Tanskanen et al., 2005). However, when conditions are beneficial for forest fires to occur, their impact on forest is usually drastic (Wallenius et al., 2005). This is due to the combination of thin poorly insulating bark and low-hanging branches that create ladder-fuels and promote crown scorching and even crown fires. As a result, spruce trees are easily killed by fire creating a huge amount of dead wood. The post-fire succession usually starts with dominance of deciduous species, mainly birch but also aspen (Populus tremuloides) and goat willow (Salix caprea L.) (Sirén, 1955). Hence, similar to wind-throws, forest fire support the presence of deciduous species as an important tree component in otherwise spruce dominated landscape.

2.3. Mountain birch dominated forests

At sites edaphically and climatically suitable for the two conifers, abundance of birch is a characteristic of a transient early-successional stage. However, elsewhere mountain birch dominated forests cover large areas of Northern Fennoscandia (Fig. 2c). The main part of this ecoregion is mainly located in Norway, followed by Sweden and to a lesser extent in Finland and the Kola Peninsula in Russia. These forests characterize the landscape both in coastal and inland areas at elevations and latitudes above and beyond the coniferous boreal
forest. In birch-dominated subalpine and subarctic areas, mountain birch is usually referred
to at subspecies-level (B.p. ssp tortuosa Ledeb. or B.p. ssp czerepanovii (Orlova) Hämët Ahti
are most common in the literature). When it occurs in other areas in mixture with the
coniferous boreal trees, it is generally referred to at species-level, i.e. B. pubescens.

The mountain birch has a wide bioclimatic range, and species composition of the
forest floor varies accordingly. Dwarf shrubs (Vaccinium spp., Empetrum sp.) are generally
most abundant, but herb-rich communities prevail in humid nutrient-rich areas and lichen
dominated communities in dry areas. In Norway and Sweden, where reindeer herding is
associated with rotation of grazing areas, lichen dominated inland areas are commonly used
as winter grazing lands for semi-domesticated reindeer, while herb-rich birch forests are
used as summer grazing lands. During the last decades the lichen biomass has decreased due
to intense grazing practice, and is being replaced by moss-rich communities (Tømmervik et
al., 2009).

Natural birch forest dynamics is driven by cyclic and abrupt population increases of
defoliating insects, such as the autumnal moth (Epírrita autumnata) and winter moth
(Operophera brumata) (Tenow, 1972; Tenow et al., 2007). The frequency and intensity of
these insect outbreaks are linked to climate at local to regional scales, and can cause large
scale stand mortality (Fig. 2d, Tenow and Nilssen; 1990; Neuvonen et al., 2005; Tenow et al.,
2007). Following the defoliation and subsequent stem mortality, the birch forest will slowly
recover its density through mainly growth of basal sprouts (Tenow, 1996; Tenow and Bylund,
2000). Mountain birch has both frequent and high quality seed production throughout its
range, as opposed to spruce and pine, and recruitment is not seed limited. However, grazing
and browsing animals such as sheep in Norway, and reindeer throughout the region might
hinder tree recruitment and maintain open woodland areas (Aune et al., 2011).
3. Disturbance and successional dynamics

The accumulated body of information challenges the conventional view that North Fennoscandian mountain forests mostly represent homogeneous and static communities of the ‘climax type’. The development of forests is slow indeed, because they grow in harsh climatic conditions in the proximity of the range limits of the constituent tree species. However, these forest ecosystems are in a state of constant change even in the absence of major disturbances (Lilja et al., 2006; Edman et al., 2007; Caron et al., 2009; Aakala et al., 2009).

The disturbance agents in mountain forest landscapes vary greatly in their type, severity and spatio-temporal characteristics. Some disturbance factors, such as fires (Kauhanen, 2002; Vajda et al., 2006), storms (Aakala et al., 2009), snow load (Hesselman, 1912; Hofgaard et al., 1991) and insect outbreaks (Tenow et al., 2013), are occasionally capable of causing widespread mortality, whereas some operate at smaller scale but more frequently such as pathogenic fungi and browsing/grazing mammals (e.g. moose and reindeer) (Edman et al., 2007; Lännenpää et al., 2008), and sometimes snow and slush avalanches, which occasionally reach into forested areas (Fig. 2e). Taken together these multi-cause disturbances act as diversity drivers at a wide range of spatial and temporal scales.

At stand scale three basic types of forest dynamics have been suggested to occur (Kuuluvainen, 2009). Two of them are classical modes of forest dynamics: even-aged stand dynamics driven by stand-replacing disturbance, and gap or patch phase dynamics driven by small scale tree mortality mostly in old forests. However, as the review of Kuuluvainen and
Aakala (2011) highlights, intermediate severity disturbances, causing partial tree mortality and leading to complex stands structures consisting of multiple tree age cohorts, are also common (Lassila, 1921; Hofgaard, 1993a). In reality, these types of forest dynamics can be mixed in space and time and are thus not easily separable even at stand scale.

At landscape scale, these different disturbance factors and the associated dynamics often operate in a mixed or nested manner in space and time. More or less continuously occurring deaths of individual or small groups of trees (gap dynamics) (Kuuluvainen, 1994) often appear superimposed on the more infrequent but larger scale partial disturbance events due to insects, wind or fire (Fraver et al., 2008). This kind of disturbance regime that operates at multiple spatial and temporal scales typically creates variable and complex forest structures (Caron et al., 2009).

In the boreal forest, fire is traditionally considered as the dominating disturbance agent (Sirén, 1955; Zackrisson, 1977). In Fennoscandian mountain forests, however, fire has played a smaller role compared to boreal forest at lower elevations and latitudes. This is due to both natural and anthropogenic reasons: these areas show a short fire season due to high precipitation and late snow melt, and the decreasing lightning frequency towards north translate into reduced natural ignitions (Granström, 1993; Larjavaara et al., 2005). In addition, the formerly important anthropogenic driver of ignitions has generally become weaker, especially during the 20th century (Wallenius, 2011). Accordingly, fire history studies using tree rings or cores from peatlands and lake sediments, have shown that prior to significant human influence the fire cycles were in the range of several hundred years (Carcailliet et al., 2007; Wallenius et al., 2010). However, despite fires being rarer than in the southern parts of the boreal forests (Granström, 1993; Larjavaara et al., 2005; Drobyshév et al., 2014), their influence is readily apparent in the age structure and species composition at
xeric sites (Lassila, 1921), and in many places these sites still display structures typical for
surface-fire driven dynamics (Kuuluvainen and Aakala, 2011).

Compared with xeric pine dominated forests, the fire cycles have been longer in
moister Picea abies dominated forests, and it is possible that parts of northern mountain
forests have not burned since the last glaciation (Hyvärinen and Sepponen 1988; Steijlen and
Zackrisson, 1987; Fraver et al., 2008). The intrinsically long fire cycles in the spruce-
dominated Fennoscandian mountain forests indicate that other disturbance agents than fire
play an important role in forest dynamics (Norokorpi, 1979; Hofgaard, 1993; Kuuluvainen
and Aakala, 2011). In the absence of fire, tree mortality is caused by wind, competition,
fungi, insects and stem breakage due snow loads (Hofgaard et al., 1991; Lännepää et al.,
2008), disturbances which are often related to old age and senescence of trees (Fraver et al.,
2008). This type of tree mortality agents drive gap phase dynamics resulting in small-scale
heterogeneity of forest structure, characterizing both age- and physiognomic structure, in
these old growth spruce forests (Hofgaard 1993a,b, Caron et al. 2009). These small-scale gap
dynamics result in a fairly constant background mortality that is then occasionally
punctuated with episodic larger scale disturbances, especially storms (Jonsson and Dynesius,
1993; Kuuluvainen, 1994; Fraver et al., 2008; Aakala et al., 2009). Disturbances such as wind
throws expose mineral soil promoting regeneration, and allowing deciduous trees to
maintain their presence in otherwise conifer-dominated stands over the longer term in
particular (Kuuluvainen, 1994; Grenfell et al., 2011), as well as supporting diversity in the
ground vegetation (Jonsson and Esseen, 1990).

Forest fires are relatively rare in mesic sites in northern mountain forests, but under
favourable conditions, they can be severe and affect large areas (Kauhanen, 2002; Aakala et
al., 2009). An example is provided by the northern spruce forest studied by Aakala et al.
which was still dominated by the post-fire tree cohort regenerated following a fire 317 years earlier. Such stand-replacing fire events have a significant and long lasting impacts on forest landscape structure, composition and biodiversity (Yli-Sirniö et al., 2012). Stand-replacing fires create open sunny habitats with large amounts of charred dead wood, which harbour a high number of saproxylic species, many of which were earlier thought to be confined to shady and moist old-growth forests (Martikainen, 2000). On fertile soils and mesic conditions the successions following fire are first dominated by deciduous species, such as Betula spp., Populus tremula and Salix caprea (Sirén, 1955; Lilja et al., 2006). Later in succession deciduous tree species give way to Picea abies which gains increasing dominance. However, Betula, Populus and Salix trees may remain as a component of the forest for long time periods due to their ability to produce asexual sprouts from the base (Betula and Salix) and the roots (Populus). In particular, this asexual reproduction ability is essential to birch forest recovery after both stand-level and landscape-level insect outbreaks (Tenow, 1996; Tenow and Bylund, 2000). Seed-based recruitment is prohibited or delayed by grazing and field layer competition (Tenow, 1996). The presence of Populus and Salix have important biodiversity consequences since they host a large number of specialized species (e.g. Martikainen, 2000; Kuusinen, 1996).

Although the effect of competition on forest dynamics may be less pronounced than in more productive forests at lower latitudes (Hytteborn et al., 1987) due to the open structure of mountain forests (see Fig. 2), competition may still be a factor regulating tree growth and mortality (Kuuluvainen and Ylläsjärvi 2011). For instance, Aakala et al. (2016) showed that competition clearly influenced the asymmetry of tree crowns, and Fraver et al. (2013) showed that the spatial arrangement of trees within stands regulated inter-tree competition and tree growth in old-growth mountain spruce forest. This latter is further
exemplified by distinct growth releases of neighboring trees when single trees die (Dynesius and Jonsson, 1991; Caron et al., 2009).

In the big picture of spatio-temporal forest dynamics, different variations of uneven-aged (or cohort-aged) dynamics appear to play a dominant role in Fennoscandian mountain forest dynamics. In *Picea* dominated stands occurring on moist sites this is due to absence of fire and thus inherently long fire rotations (Wallenius et al., 2010) and the predominance of gap or patch dynamics. In *dry Pinus* dominated sites, where fires may be more common, tree mortality is often partial or patchy as large *Pinus* trees with their thick heat-insulating bark often survive surface fires. This, and the post-fire regeneration, result over time in forest with several age cohorts of trees (Lassila 1921; Engelmark et al., 1998). In *Betula* dominated forests insect outbreaks are, in the absence of fire, the main driver of stand dynamics.

4. Discussion

Research carried out during the past decades has greatly advanced our understanding of the history, ecology, biodiversity and dynamics of North Fennoscandian mountain forest ecosystems, and the varying but ubiquitous long-term influence of humans on these seemingly natural forest ecosystems (Fraver et al., 2008; Josefsson et al., 2009; Kuuluvainen and Aakala, 2011). It is clear that the predicted rapid changes in climate will greatly influence many key biological processes controlling the dynamics of these forests growing close to their northern distribution range. Some of the changes are gradual but also episodic more drastic disturbances will become more likely (Gauthier et al. 2015). As a consequence, the ongoing climate change will exert unprecedented pressures to the “health” of these forests, which are of paramount importance as biodiversity hotspots and
as the core of the forest protection area network in the whole of Fennoscandia. Critical components of ecosystem health can be defined as the biodiversity, resilience and adaptive capacity of the constituent ecosystems (Gauthier et al., 2015). These ecosystem properties are the crucial components of forest health and should form the basis of conservation and management actions.

Our understanding of the ecology and dynamics of these forests is based on past conditions. A relevant question then is how relevant and useful this knowledge is in the future climatic conditions that will significantly differ from past conditions (Sprugel, 1991; Keane et al., 2009; IPCC, 2013). However, we can identify several important roles that research on past disturbance dynamics can play in forest protection and sustainable ecosystem-based management. First, an often-invoked role is the establishment of baseline or reference information on ecosystem properties and dynamics (e.g. natural range of variability, Willis and Birks, 2006; Keane et al. 2009). Detecting changes is possible only if we have a sufficient understanding of the baseline conditions and their variability. Second, we need knowledge on past forest dynamics to understand how the baseline conditions have historically formed and, for example, what has been the role of human activities in this process (Hofgaard, 1999; Josefsson et al. 2009).

Given that humans have lived in the Fennoscandian mountain forests for millennia we can assume that they to some extent have influenced forest structure and dynamics (e.g. Östlund et al. 2004). Prior to colonization by mainly farmers from the south during the 17th and 18th centuries, only indigenous Sami groups lived in these forests. Sami people have traditionally been viewed as finding subsistence from hunting and gathering and relatively late in history becoming dependent on reindeer herding (see Sommerseth 2010). However, the traditional view is being challenged and no consensus on the onset of reindeer herding
seems to exist. Also recent studies suggest that Sami land-use was more complex than
assumed and may have included small scale shifting cultivation (Östlund et al. 2015). The
land use pattern is evident in pollen profiles from local settlements and provide together
with archeological evidence an insight in to Sami subsistence patterns (Sommerseth 2010). It
is suggested that in some place forest resources may have been overexploited near the tree
line, resulting in ecosystem degradation and subsequent shift in land use (Bergman et al.
2013). However, in general it is difficult to translate the impact of Sami land use into a
broader impact on the forest landscape.

Understanding the historical processes is a pre-requisite for appropriate use of the
past baseline information. For example, in research on forest change over larger spatial
scales, using satellite images and forest inventories, past forest dynamics need to be
understood to attribute detected changes to effective drivers. This is necessary to discern
the influences of past disturbances from the direct influences of climate change due to
warming temperatures, increasing CO₂ concentration or nitrogen deposition (Myneni et al.
2001). Another prerequisite for effective monitoring is looking at deviations from baseline
conditions at multiple spatial scales.

North Fennoscandian mountain forests are occasionally considered to be resilient
against adverse changes, particularly because a large part of them is protected (Aksenov et
al. 2014). This view may also gain support from the slow growth and sometimes quasi-
equilibrium dynamics of late-successional forests (e.g. Caron et al., 2009; Aakala et al., 2009).
However, we have learned that these forests are not as stable as was perhaps once assumed
and ecological theory predicts that ecosystems close to their distribution range are most
vulnerable to sudden state shifts (Atkinson, 1992; Scheffer et al., 2012). For instance, the
Holocene history tells us that major changes in altitudinal positions in the treeline ecotone may occur as a result of the predicted changes in future climate (Aakala et al. 2014). It is generally acknowledged that climate change will strongly affect the future development of forests and the predicted temperature increase (IPCC 2013) suggest that the climate conditions in mountain forests will be displaced towards conditions resembling forests at lower altitudes. It is also clear that disturbance is central in vegetation dynamics and in translating climate change into vegetation response (Overpeck et al., 1990). There is a risk that with accumulating environmental stressors in the rapidly warming climate, critical thresholds are surpassed and abrupt changes in ecosystem conditions can become a reality (Gauthier et al. 2015). Old spruce forests are perhaps most vulnerable to the effects of warming climatic conditions and associated disturbances such as droughts, insect outbreaks, and perhaps also fire (Kellomäki et al., 2008; Kuuluvainen et al. 2014).

However, we lack understanding of the changing role of disturbances, and especially that of fire in mountain forest ecosystems. In the past centuries, the occurrence of fire has to a great extent been determined by human influences in interaction with those of climatic fluctuations (Drobyshhev et al., 2014). First, settlement activities until late 20th century increased fires due to human-caused ignitions. Since forests became a valuable resource, fire was handled more carefully and fire suppression was developed, which resulted in an almost-complete elimination of forest fires over the entire Fennoscandia (Wallenius, 2011; Drobyshhev et al., 2014). This was true also in xeric sites where fires have historically been most prevalent. As trees are long-lived, the lack of fire manifests itself gradually, but eventually leads to changes in structure and composition in forests previously shaped by fires. For example, pine regeneration may be impeded while fire-sensitive spruce may gain dominance especially on more moist and fertile sites capable of supporting spruce (Linder et
The absence of fire is also reducing the occurrence of early successional deciduous forest in the landscape and the regeneration possibilities for rare biodiversity-maintaining pioneer tree species, such as *Salix caprea*. A climate-induced broadening of the distribution of defoliating insects, such as *Epirrita autumnata*, may potentially hasten this development through reduction of deciduous mixture in conifer-dominated mountain forests (Jepsen et al., 2008).

However, predicting the future development of complex ecological systems and their disturbance dynamics is difficult (Messier et al., 2014). In particular, we lack knowledge of possible ecological thresholds of different kinds of mountain forest ecosystems to changing environmental factors, stressors and disturbances. An important question is the interaction of different disturbance factors in a warming climate. We can qualitatively predict, for example, that increased tree mortality due to abiotic factors such as wind or droughts can increase biotic damages (e.g., due to bark beetles) (Kuuluvainen et al., 2014). However, quantitative prediction is difficult because of the unknown ecosystem thresholds and stochastic nature of climate-driven disturbance phenomena, increasing the possibility of unexpected events and surprises (Scheffer et al. 2014). These uncertainties, and the notion that climate change will manifest itself earlier in ecosystems at climatic margins, calls for increasing monitoring of the mountain forest ecosystems.

The North Fennoscandian mountain forests are of paramount importance as biodiversity hotspots, for monitoring of environmental change, and as a reference for forest restoration and ecosystem based management (Kuuluvainen and Aakala, 2011; Halme et al., 2013). They also have a rich cultural heritage. Northern mountain forests are important for the indigenous Sámi people and their reindeer herding culture. However, there is a history of conflicts between reindeer herding and logging activities especially in Finland (Helle and
Jaakkola, 2008) and Sweden (Berg et al., 2008), and there will possibly be increasing pressure for timber harvesting in the so far unprotected mountain forests. New economically important and rapidly increasing land uses affecting mountain forests are nature-based tourism and mining. This complex development and potential conflicts between different competing land uses, highlight the need for conservation and ecologically sustainable management based on adequate understanding and prediction of future trajectories of mountain forest ecosystems.

Then what could be the main lessons learned from ecological research in North Fennoscandian mountain forests concerning conservation and ecosystem-based management? A major conclusion to be drawn from the ecological research literature is that fine scale processes of disturbance and regeneration, and long-term continuity of structural characteristics are essential features of mountain forests and their dynamics. These are also the ecosystem characteristics and their historical range or variability, which need to be respected if biomass harvesting is considered. However, it could be argued that the main value of the remote and slow-growing Fennoscandian mountain forests is not as a biomass resource. Instead they harbor significant values related to cultural heritage, tourism and recreation, biodiversity, and serving as reference monitor of the effects of climate change in marginal forest ecosystems.

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Figure legends

Fig. 1. The geographic region (dashed area) indicating the mountain forests that are in focus in this paper. Grey shows the distribution of forest.

Fig. 2 Pine forest in different site types and developmental stages. (a) Old pine forest with standing and fallen Kelo trees (Maltio Strict Nature Reserve, Finnish Lapland). (b) Paludified pine forest on mountain slope (central Norway). (c) Burned pine forest with deciduous regeneration (Murmansk region, Russia). (d) Abundant pine regeneration after forest fire (Murmansk region, Russia). Photos: Timo Kuuluvaalinen.

Fig. 3 Forest dynamics stages in mostly spruce dominated moist forests. (a) Old spruce dominated forest with thick raw humus layer (Värriö Strict Nature Reserve, eastern Finnish Lapland. (b) Burned spruce forest with abundant deciduous regeneration (Murmansk region, Russia). (c) Canopy gap in old spruce forest with deciduous regeneration (Murmansk region, Russia). (d) young deciduous forest regenerated after fire in a spruce forest (Pallas-Yllästunturi National Park, Finnish Lapland. Photos: Timo Kuuluvaalinen.

Fig. 4. Mountain birch forest is characterizing large parts of the North Fennoscandian mountain region. (a) Birch trees and forest in autumn color (Abisko, northwestern Sweden). (b) Grass-dominated (Descampsia flexuosa) forest floor one year after tree canopy defoliation by autummal moth (Epirrita autumnata) and subsequent nutrient release (Abisko, northwestern Sweden). (c) Slush avalanche track in birch forest, (Tärna Mountains, Sweden). (d) Former mountain birch forest turned into tundra after autumnal moth outbreaks in the 1960s (Kevo area, northern Finland). Photos: Annika Hofgaard (a)-(c) and Timo Kuuluvaalinen (d).
Graphical Abstract