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RESEARCH

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Tree diametric-species diversity is affected by human impact in old Scots pine dominated forest in boreal Fennoscandia



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Gonzalo de Quesada^{*} and Timo Kuuluvainen

Abstract

Background: Forest structural and compositional variability is of fundamental importance for forest ecosystem functioning and species diversity. The purpose of this research was to examine how human impact has affected the compositional-structural diversity of mature pine-dominated boreal forest in boreal Fennoscandia. For this a new approach was used, based on the classification of tree sizes by the diameter at breast height (dbh) and tree species, resulting in a new variable, the diametric-species, the variation of which describes the compositional-structural diversity of the forest. This variable was used to compare the structural-compositional diversity among three forest classes with different degree of human influence, using rarefaction as the main tool of analysis, complemented by analyses based on common diversity indices.

Results: The results showed that the near-natural forest was the most diverse and the managed forest the least diverse. On the other hand, the diversity of near-natural and selectively logged forests were similar, suggesting that selectively logged forests are equal to the natural forest in their compositional-structural diversity. The analysis solely on tree species showed no significant differences among the forest classes of different human impact. The Shannon diversity index showed no significant difference between the forest classes for the diametric-species and tree species classifications only, but the Simpson index signaled a slight difference between the selectively logged and managed forest classes for the diametric-species classification. Furthermore, the Sorensen index detected a difference among forest classes in the diametric-species classification.

Conclusions: Forest utilization had an adverse impact on forest compositional-structural diversity of mature Scots pine forests. The analysis also shows that the novel approach based on diametric-species classification could be a useful tool for forest diversity analysis and comparison, especially in species-poor forests such as the boreal forest.

Keywords: Diametric-species, Boreal forest diversity, Rarefaction analysis, Human intervention, Compositional-structural diversity

Background

Forest structure and composition are essential ecosystem characteristics, because they regulate tree growth and mortality patterns, availability of resources and, habitat conditions for all forest organisms (Lindenmayer et al. 2012, 2014; Pommerening 2002, 2006; von Gadow et al. 2012). Forest structure and composition can affect the probability and spread of forest disturbances (Kuuluvainen 2002). Variation in tree sizes and tree species composition

* Correspondence: gonzalo.dequesadaalzamora@helsinki.fi

strongly contribute to the overall species diversity, because diverse micro- and macrofauna depend on different canopy layers of the forest and large old trees for shelter, nesting and food (Spies 1998; Perry et al. 2008).

Forest structure also includes deadwood, which is a significant component of forest structural diversity (Sturtevant et al. 1997; Siitonen et al. 2000). Deadwood is important for ecosystem functioning as it regulates carbon and nutrient cycling, and water availability (Graham and Cromack 1982; Karjalainen and Kuuluvainen 2002), it is also essential for forest biodiversity because it provides habitat and resources for various kinds of animals, plants and decomposers that are



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Department of Forest Ecology, University of Helsinki, P.O. Box 24, FIN-00014 Helsinki, Finland

important for forest ecosystem functioning (Linder 1998; Brassard and Chen 2006).

Tree species composition is a key characteristic of forest ecosystems (Lindenmayer and Franklin 2002). Tree species diversity contributes positively to productivity, provides habitats and resources to the forest species, and increases forest resilience to disturbances (Hooper and Vitousek 1997; Peterson et al. 1998; Lindenmayer and Franklin 2002). The decline in tree species richness can have an adverse impact on the levels of ecosystem functions (Naeem et al. 1999; Perry et al. 2008). Although boreal forests are characterized by lower tree species richness than temperate and tropical forests, boreal forest structure and dynamics can exhibit great diversity and complexity (Lähde et al. 1999; Ylläsjärvi and Kuuluvainen 2009; Kuuluvainen and Aakala 2011). Moreover, the low number of tree species with diverging ecological characteristics suggests that the presence or absence of even a single tree species may have a considerable impact on total species diversity of the forest.

In boreal forests, human forest utilization has generally had a negative impact on forest diversity (Paillet et al. 2009). In northern Europe, forest management has been more intensive and affected the ecosystem perhaps more than anywhere in the boreal zone. This is because forest management, which is based on clear-cutting, tends to simplify forest composition and structure by removing undesirable species, suppressing fires, and by maintaining even-aged single species stands to enhance wood production (Christensen and Emborg 1996). These practices have led to a continuous decline in the number of old trees and dead wood (Siitonen 2001; Nordlind and Östlund 2003), and the practical disappearance of uneven-aged, structurally complex forests from managed forest landscapes (Kuuluvainen and Gauthier 2018).

The purpose of this research was to examine whether human utilization has impacted compositional-structural diversity of mature pine-dominated boreal forests in Fennoscandia. For this, a new approach was used, based on the classification of trees by their diameter class and species, resulting in a variable called diametric-species. Using this variable, a comparative analysis was carried out using rarefaction curve analysis and complemented by analyses based on diversity indices.

Methods

Study regions

The forests in this research are located in three different regions of Fennoscandia. Häme (Pirkanmaa region) in southwestern Finland, Kuhmo (Kainuu region) in northeastern Finland, and Vienansalo (Russian Karelia) (Fig. 1).

The forests share similar characteristics in their soil, forest site type, species composition, and climatic conditions. Table 1 shows that Kuhmo and Vienansalo regions have almost identical climatic conditions. Although Häme has a slightly higher mean temperature, mean effective temperature sum and longer growing seasons, all three areas are located within the mid-Boreal vegetation zone and present similar growing conditions (Rouvinen and Kuuluvainen 2005).

The forest in all the study regions is mainly pinedominated, with spruce and birch as part of their tree species composition, and with aspen and willow as minor species. Overall, the species composition was representative of natural forests in Fennoscandia (Axelsson and Östlund 2000; Kuuluvainen 2002). The most common and predominant forest site types were Vaccinium-Myrtillus Type (VMT) and Empetrum-Vaccinium Type (EVT). These forest site types are considered medium fertile and mesic (VMT) and poor and sub-dry (EVT) (Cajander 1926; Pyykkö 1996; Korhonen et al. 2000; Rouvinen et al. 2002a) and are usually dominated by pine trees.

The bedrock in Häme is formed mostly by granodiorite and quartz, granite, metabasalt, greenstone and amphibolite gneisses, while in the Kuhmo region a granite vein gneiss is the main component of its bedrock (National board of survey geographical society of Finland 1990), and the bedrock in Vienansalo is formed by an amphibole and biotite gneisses (Gorkovets et al. 2000). Moreover, in all the areas the topsoil covering the bedrock consists of peat deposits and ground moraines (Gorkovets et al. 2000; National board of survey geographical society of Finland 1990).

History of forest utilization of the study regions

To understand the state of these forests it is important to know the history of activities that took place in these regions. Although these regions were settled permanently at different periods of time: Häme in the midsixteenth century, Kuhmo in the seventeenth century, and Vienansalo in the mid-eighteenth century, they went through similar utilization stages until the early twentieth century.

Since the establishment of early settlements in the mid-sixteenth century, Häme was subject to slash and burn cultivation practices until the twentieth century. Furthermore, during the eighteenth century intensive tar extraction and burning practices were carried out in the region, and the selective logging of the forest started, which intensified during the 19th and mid-twentieth century as a result of the increase in population density of the area and the growing forestry industry in southern Finland (Lilja and Kuuluvainen 2005). Currently, many of these forests have become managed production forests, by the use of different silvicultural treatments like thinning and the removal of understory spruce trees (Rouvinen and Kuuluvainen 2005).

As in Häme, Kuhmo was the subject of slash and burn cultivation, tar production, and selective logging became

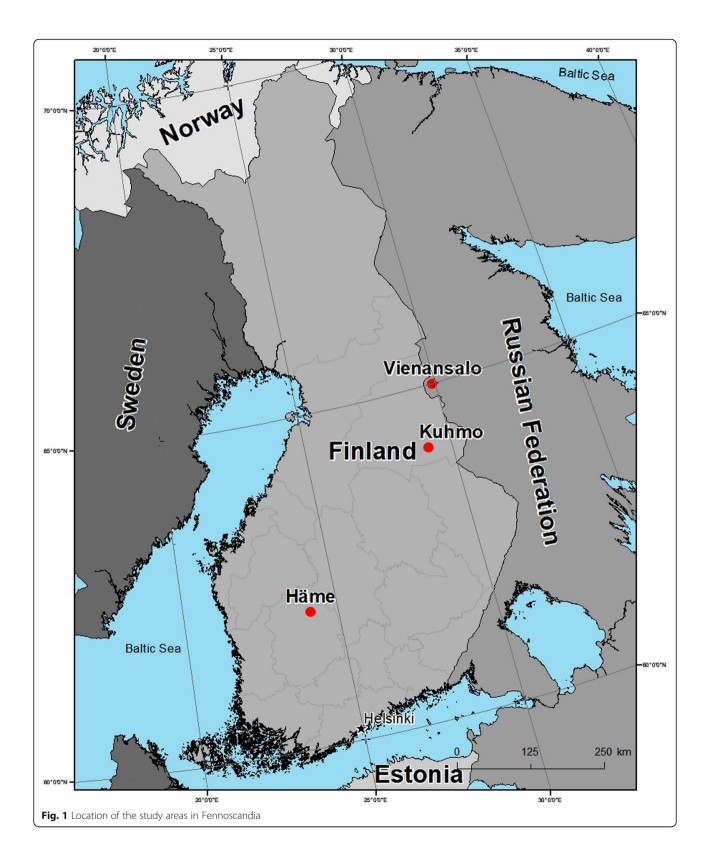


Table 1 Location and climatic characteristics of the three study regions (National board of survey geographical society of Finland 1987; Rouvinen et al. 2002b; Nazarova 2011)

| | Häme | Kuhmo | Vienansalo | | |
|---|--------------|--------------|--------------|--|--|
| Location | 62° N, 24° E | 64° N, 29° E | 65° N, 30° E | | |
| Altitude (m.a.s.l.) | 180 | 200 | 155 | | |
| Mean Temperature (°C) | 3 | 1.5 | 1 | | |
| Precipitation (mm) | 650 | 650 | 650 | | |
| Growing season (days) | 160-165 | 145-150 | 134–159 | | |
| Mean effective temperature sum (threshold, 5 °C) | 1000 | 900 | 900 | | |

an important source of livelihood for the region in the mid-nineteenth century. These practices continued for a long period until the twentieth century (Wallenius et al. 2002).

In Vienansalo, slash and burn cultivation was also carried out, as well as tar burning for domestic purposes (Rouvinen et al. 2002b). There is evidence of selective logging in the area between the 19th and 20th centuries, but the intensity of the logging was low and exclusive to Pine trees (Karjalainen and Kuuluvainen 2002). The fire regime of Vienansalo forests was the same as the ones found in eastern Finland because of the human activities that took place in the area (Kuuluvainen et al. 2002; Wallenius et al. 2004). These forests are considered to be in a more natural conservation state than the forests in Finnish Karelia (Jantunen et al. 2010).

Although Russian Karelia forests show the least amount of forest harvesting activities compared to Finnish forests (Saarinen et al. 2001), all the study areas present similar historical forest utilization backgrounds, and they only differ in the duration and intensity of human intervention.

Sampling and measurements

The sampling of these regions was carried out in three different fieldwork seasons. The first took place in Kuhmo in 1997, the second in 1998 in Vienansalo, and finally in 1999 in Häme. The aim was to collect data from three forest classes with different degrees of human impact (Lilja and Kuuluvainen 2005). These impacts were described by Lommi et al. (2009) and were classified as 1) near-natural stands; 2) selectively logged stands (selective logging was carried out in the early twentieth century, but no other treatments had been performed); 3) managed stands (stands silviculturally treated).

The stands in Kuhmo and Häme were selected from the stand data files of The Finnish Forest and Park Service (Metsahallitus) and The Finnish Forest Research Institute (LUKE), taking into consideration the following criteria: a) Pine dominated forests by volume, b) dominant Pines of at least 90 years old, and c) stand area of at least 3 ha (Rouvinen and Kuuluvainen 2005). The stands in Vienansalo were selected based on the following criteria: a) the area is remote to minimize potential human influence, b) the landscape is typical of the region, and c) there is water access to the area from the local village of Venehjärvi (Kuuluvainen et al. 2002).

A total of 116 sample plots were established in the study areas: 57 in Häme, 32 in Kuhmo, and 27 in Vienansalo. All the sampling plots were rectangular of 20 m × 100 m (0.2 ha) in size. In Häme and Kuhmo the sampling plots were randomized while making sure their location was at least 30 m from the edge of the forest area to avoid the edge effect (Lilja and Kuuluvainen 2005). In Vienansalo the sampling plots were selected randomly from 6 lines running in an east to a west direction within the study area. These lines were 1000 m apart from each other, and the plots were considered if they landed on firm land and within a relatively homogeneous forest patch (Karjalainen and Kuuluvainen 2002).

Out of the 116 sampling plots, 22 were classified as near-natural, 40 as selectively logged, and 54 as managed forests. Furthermore, 32,875 trees were measured: 7527 from the near-natural, 13,169 from the selectively logged, and 12,179 from the Managed forests.

Diametric distributions

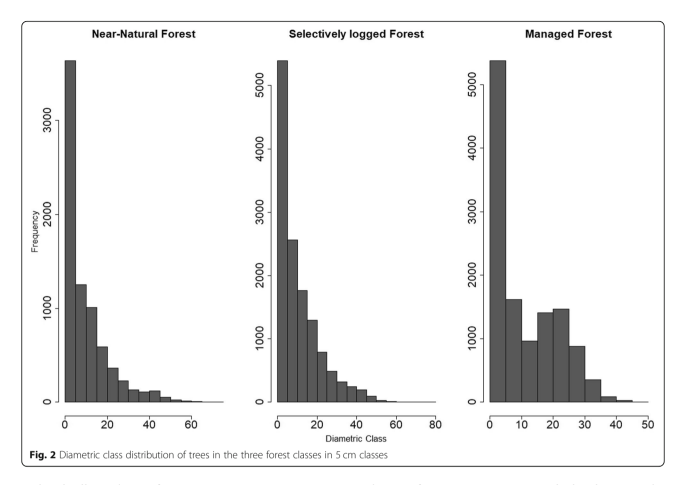
The diametric class distributions were similar in the near-natural and selectively logged forests (Kuuluvainen et al. 2002; Rouvinen and Kuuluvainen 2005). They showed a descending or J-shaped behavior, agreeing with the behavior of an uneven-aged self-regenerating forest, in which the recruitment and mortality are in balanced, and small trees are most abundant (Felfili 1997). On the other hand, the managed forest exhibited a bimodal or rotated-sigmoid form. This behavior agrees with forests that have presented a moderate to severe disturbance or intervention (Lianjun Zhang et al. 2001) (Fig. 2).

Data analysis

The data collected contains information about the different characteristic of the forest, but for this research purpose, only tree species and the diameter at breast height (dbh) were used. The tree species found and their coding is shown in Table 2.

Each tree in the three forest classes was classified by species and diametric class, using 5 cm interval groups. By doing this, a new variable unit was created, called the *diametric-species*. The full classification with the 160 classes is shown in Table 3. Henceforth this new category will be referred to as diametric-species.

The diversity of diametric-species were analyzed using rarefaction curves because they account for sampling biases and adjust for the differences in the number of individuals and samples collected (Gotelli 2001; Gotelli



and Colwell 2011). Rarefaction curves can compare two separate communities of different sizes and number of individuals or species by the interpolation or extrapolation of their curves (Simberloff 1972; Tipper 1979). Because of the difference in the number of diametricspecies in the forest classes, the extrapolation was used to estimate the richness and completeness of the samples, by visualizing the asymptote of the cumulative curves (Chao and Jost 2012; Gotelli and Chao 2013).

 Table 2
 List of the tree species and the code used for the diametric-species classification for the three forest classes

| Code | Tree species |
|------|--------------------------------|
| A | Scots Pine (Pinus sylvestris) |
| В | Norway Spruce (Picea abies) |
| С | Silver Birch (Betula pendula) |
| D | Downy Birch (Betula pubescens) |
| E | Common Alder (Alnus glutinosa) |
| F | Aspen (Populus tremula) |
| G | Willow (Salix sp.) |
| Н | Ash (<i>Sorbus</i> sp.) |
| I | Other hardwood |
| J | Other coniferous |

The rarefaction curves were calculated using the sample-based Bernoulli product model described by Colwell et al. (2012) and the 95% low and high bound confidence intervals were also calculated for each curve. The formula used to calculate the rarefaction curve was:

$$S_{sample}(t) = \sum_{i=1}^{S} \left[1 - (1 - \theta_i)^t \right] = S - \sum_{i=1}^{S} \left(1 - \theta_i \right)^t$$
(1)

And the extrapolation was calculated using the formula:

$$S_{sample}(T + t') = S_{obs} + \sum_{i=1}^{S} \left[1 - (1 - \theta_i)^{(t')} \right] (1 - \theta_i)^T$$
(2)

where $S_{sample}(t)$ is the expected number of species (diametric-species in our case), $S_{sample}(T + t')$ is the estimated number of species extrapolated, t is the random set of sampling units, T is the total number of sampling units, t' is the additional extrapolated sampling units, Sis the total number of species in the assemblage, S_{obs} is the total number of species observed in the sample, i is the species, and θ_i is the detection probability of each species in the assemblage. To avoid any biases due to differences in sample density, the results of the sampleQuesada and Kuuluvainen Forest Ecosystems (2020) 7:8

| Diametric class (cm) | Tree species | | | | | | | | | |
|-------------------------|--------------|--------|--------------|-------------|-------|-------|--------|-----|----------------|------------------|
| | Pine | Spruce | Silver Birch | Downy Birch | Alder | Aspen | Willow | Ash | Other Hardwood | Other Coniferous |
| 1–5 | A1 | B1 | C1 | D1 | E1 | F1 | G1 | H1 | 11 | J1 |
| 6–10 | A2 | B2 | C2 | D2 | E2 | F2 | G2 | H2 | 12 | J2 |
| 11-15 | A3 | B3 | C3 | D3 | E3 | F3 | G3 | H3 | 13 | J3 |
| 16-20 | A4 | B4 | C4 | D4 | E4 | F4 | G4 | H4 | 4 | J4 |
| 21–25 | A5 | B5 | C5 | D5 | E5 | F5 | G5 | H5 | 15 | J5 |
| 26–30 | A6 | B6 | C6 | D6 | E6 | F6 | G6 | H6 | 16 | J6 |
| 31–35 | A7 | B7 | C7 | D7 | E7 | F7 | G7 | H7 | 17 | J7 |
| 36–40 | A8 | B8 | C8 | D8 | E8 | F8 | G8 | H8 | 18 | J8 |
| 41–45 | A9 | B9 | C9 | D9 | E9 | F9 | G9 | H9 | 19 | J9 |
| 46-50 | A10 | B10 | C10 | D10 | E10 | F10 | G10 | H10 | 110 | J10 |
| 51-55 | A11 | B11 | C11 | D11 | E11 | F11 | G11 | H11 | 111 | J11 |
| 56–60 | A12 | B12 | C12 | D12 | E12 | F12 | G12 | H12 | 112 | J12 |
| 61–65 | A13 | B13 | C13 | D13 | E13 | F13 | G13 | H13 | 113 | J13 |
| 66–70 | A14 | B14 | C14 | D14 | E14 | F14 | G14 | H14 | 114 | J14 |
| 71–75 | A15 | B15 | C15 | D15 | E15 | F15 | G15 | H15 | 115 | J15 |
| 76–80 | A16 | B16 | C16 | D16 | E16 | F16 | G16 | H16 | 116 | J16 |

Table 3 Diametric-species classification use in the species area analysis

based rarefaction analysis were plotted against individual abundance as suggested by Gotelli and Colwell (2001).

The non-parametric Chao index was used to determine the lower bound of species richness by estimating the minimum number of species expected from an incidence data (Chao and Chiu 2016). The index takes into consideration the number of uniques and duplicates to determine the minimum number of total species in a community (Gotelli and Chao 2013).

The sample-based Chao2 index was used to determine the minimum species richness and it was only calculated for the diametric-species data because the number of species in the original data is known. The Formula as described by Gotelli and Ellison (2013):

$$Chao2 = S_{(obs)} + \frac{(R-1)}{R} \left(\frac{q_1^2}{2q_2}\right)$$
(3)

Where $S_{(obs)}$ is the observed number of species, R is the number of samples in the data, q_1 is the number of uniques (species that only appear in one sample) and q_2 the number of duplicates (species that appear exactly in two samples).

The Shannon-Wiener, reciprocal Simpson, and Sorensen index were used in this research to complement the rarefaction diversity analyze. They were selected based on their ability to estimate alpha and beta diversity, and their use as common indices in biology, ecology, and diversity studies. The Shannon-Wiener index (H) and the reciprocal Simpson index (D') were calculated using the following formula:

$$H = -\sum_{i=1}^{S} p_i \ln(p_i) \tag{4}$$

$$D' = \left(\sum_{i=1}^{S} p_i^2\right)^{-1}$$
(5)

where *S* is the number of species in the assemblage, and p_i is the proportion of individuals that belong to the species *i*.

The Sorensen index was calculated using the formula:

$$S_S = \frac{2a}{2a+b+c} \tag{6}$$

where a is the number of species common to both samples, b the number of species unique to the first sample and c the number of species unique to the second sample.

The data was analyzed using a free to use software called EstimateS 9.1.0 (Colwell 2003). The software was used to calculate the rarefaction curves and their confidence intervals, as well as the Chao2 index. As a result of the small sample size of the near-natural and selectively logged forest, their curves were extrapolated to be matched with the managed forest larger sample size.

The software randomized the data samples 100 times and calculated the average values to create the curves.

The rarefaction analysis, as well as all the diversity indices, were calculated for the diametric-species dataset and the original data containing only the tree species for comparison (Table 3).

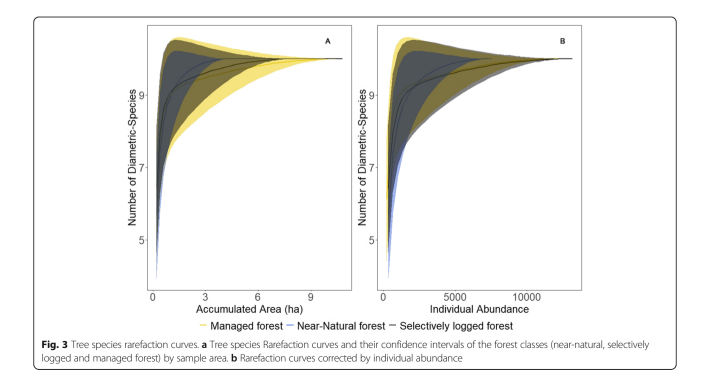
Results

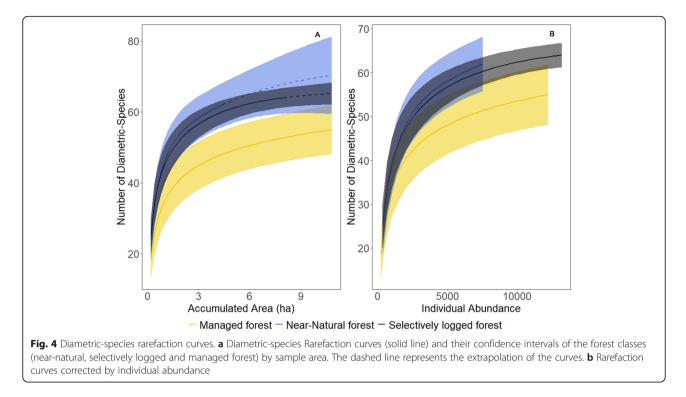
The results of the rarefaction analysis of only tree species show similar behaviors among all three forest classes (Fig. 3). The curves start with a first rapid rise, followed by a slow and steady increase until they approach the asymptote. The near-natural forest reached the asymptote at a sampling area of 4 ha, the selectively logged forest at 7.4 ha and managed forest at 10 ha. Although the forest classes reach the asymptote at different sampling area values, there are no significant differences in their curves' behavior. Furthermore, the near-natural and selectively logged forest confidence intervals were contained within the confidence intervals of the managed forest through the entirety of their trajectory.

The rarefaction curves based on the tree diametricspecies, in contrast to the curves of the sole tree species data, present different behaviors for each forest class (Fig. 4). For the near-natural forest, the rarefaction curve shows the steepest initial rise and it reaches the highest level of diametric-species (70.33) among all the three forest classes. The near-natural forest curve seems to keep rising with the increase of sampling area and it appears that the curve is far from reaching an asymptotic state. The selectively logged forest curve presents also a first steep rise but does not reach as high as the near-natural forest curve (65.15). Additionally, the selectively logged forest curve seems like it is approaching the asymptote. The managed forest, when compared to the near-natural and selectively logged forest, presents a more regular and less steep curve increase and it contains the lowest number of diametric-species (55). To compare the diametric-species in the three forest classes, the near-natural and selectively logged forests curves were extrapolated at 4.4 and 8 ha respectively to match the managed forest larger sample size.

The near-natural forest's confidence interval superimposes that of selectively logged forest throughout its entire trajectory (Fig. 4). Moreover, the near-natural and managed forests confidence intervals slightly overlap through the curves trajectory. The overlap of the near-natural and managed forest confidence intervals could be attributed to the similarities in tree species composition.

The minimum number of diametric-species estimated were: 70.59 for the near-natural forest, 65.39 for the selectively logged Forest, and 64.16 for the Managed forest, but the estimation curve of the Chao2 index (Fig. 5) shows that among the three forest classes, only the selectively logged forest reached a stable point at around 4 ha. The managed and near-natural forest curves do not stabilize and keep rising with the increase in the sampling area, suggesting the estimated minimum number





of species for the near-natural and managed forest could be higher because some rare diametric-species combinations have not been registered.

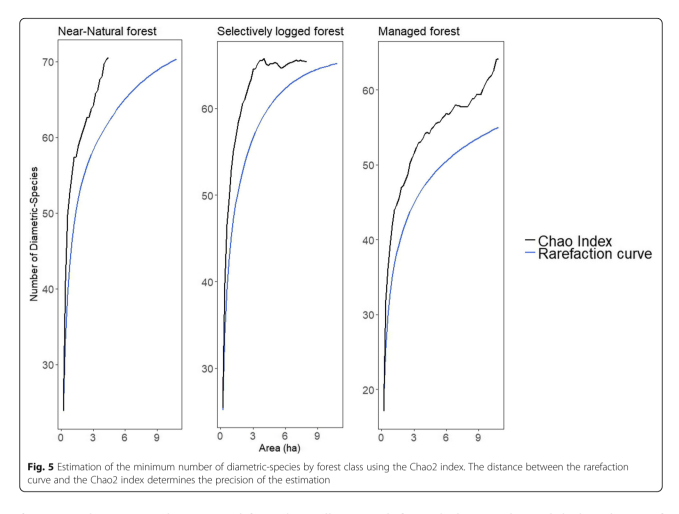
The results of the Shannon index show similar values among the three forest classes for the tree species and the diametric-species classifications. The results show values between 2.9 and 2.91 for the diametric species and a value of 3.97 for all the forest classes of the tree species classification. The reciprocal Simpson index shows values between 10.4 and 12.96 for the diametricspecies classification and 53.70 to 53.79 for just the tree species. The values show a difference between the managed and selectively logged forest for the diametricspecies classification, but because of the near-natural forest small sample size, it was not possible to determine a significant statistical difference with the other forest classes. On the other hand, the tree species classification values for all forest classes show no significant difference (Table 4).

The Sorensen similarity index shows a difference between the forest classes of the diametric-species classification. The results ranged from 0.85 to 0.91 (Table 5). While, the values for the tree species classification were 1 for all the forest classes, meaning a total (100%) similarity between the forest classes.

Discussion

The rarefaction diversity analysis based solely on tree species did not find a diversity difference among forest age classes with different degree of human influence. This may be due to the low number of tree species in the studied boreal forests. On the other hand, when both tree size structure and species composition are considered as diametric-species, the rarefaction analysis shows a significant diversity difference among forest classes. This can be understood in the light of the findings showing that natural boreal forests are characterized by considerable and complex structural and compositional diversity (e.g. Ylläsjärvi and Kuuluvainen 2009).

The diametric-species based rarefaction analysis indicates that near-natural forests have a higher compositional-structural diversity than any other forest class. The results show a noticeable difference between diametric-species diversity of the near-natural and managed forest, suggesting that forest management had an adverse effect on forest compositionalstructural diversity. These differences are the result of forest management practices that reduce the range of the tree diameter distribution by removing large trees for commercial purposes. This can be observed in the managed forest diameter distribution, where no tree individuals over 48 cm in diameter were found. In contrast, the largest tree in the near-natural forest had a diameter of 71 cm. The absence of large trees in the managed forest resulted in a lower number of diametric-species, when compared to the selectively logged and semi-natural forests. This lack of large trees in the managed forest contributes to more homogeneous forest structure, which can result in lower overall diversity of species and ecosystem



functions, when compared to a natural forest (Bunnell and Huggard 1999; Lindenmayer et al. 2012, 2014; Gauthier et al. 2015). These impacts are also attributed to the suppression of small-scale disturbances that control the boreal forest dynamics, particularly important in pine-dominated forests (Kuuluvainen and Aakala 2011).

The rarefaction analysis shows a similar diametricspecies diversity in the near-natural and selectively logged forest. This is supported by Lilja and Kuuluvainen (2005) who concluded that the selectively logged forest has recovered close to a natural state because the activities in the forest had occurred a long time ago and the low number of individuals removed. Alternatively, it is also possible that the nearnatural forest had an undetected higher degree of human intervention (Rouvinen et al. 2002b). The similarities between the near-natural and the selectively logged forest could be relevant from a nature conservation point of view. The selectively logged forest could be considered as a potential future protected area or as a buffer zone if it is located in the vicinity of an already established protected forest.

The Chao2 index values of the near-natural forest show a considerably higher number of estimated diametric-species compared to the other forest classes, which supports the rarefaction curve findings that the near-natural forest is the most compositional-structural diverse forest class. The index curve also shows that more samples are needed to accurately determine the

Table 4 Diversity indices calculated for diametric-species and tree species for each forest class

| Forest class | Shannon (<i>H</i>) | | Reciprocal Simpson (D') | | |
|--------------------|----------------------|--------------|-------------------------|--------------|--|
| | Diametric-species | Tree species | Diametric-species | Tree species | |
| Near-Natural | 2.9 | 3.97 | 11.37 | 53.7 | |
| Selectively logged | 2.91 | 3.97 | 10.4 | 53.79 | |
| Managed | 2.91 | 3.97 | 12.96 | 53.77 | |

 Table 5
 Sorensen index calculated between each diametricspecies forest class

| Sorensen (S _s) | | |
|-----------------------------------|------|--|
| Near-Natural / Managed | 0.85 | |
| Managed / Selectively logged | 0.89 | |
| Near-Natural / Selectively logged | | |

minimum expected number of diametric-species of the near-natural forest. The Chao2 index curve of the selectively logged forest show an accurate measurement of the minimum expected number of diametric-species and suggests the curve has reached the asymptote, this is determined by the distance between the observed richness and the Chao index estimation. (Gotelli and Ellison 2013). The Chao2 index curves also show that rare diametric-species combinations are still present in the managed forest and more samples would be needed to register them all, suggesting the managed forest could have a higher number of minimum diametric-species.

The Shannon index shows no significant difference in diversity among the three forest classes, whether the diametric-species or the original tree species data was used. The reciprocal Simpson index shows no significant difference among forest classes in the tree species classification, but found a slight difference between the managed and selectively logged forest in the diametricspecies classification. It was not possible to determine if there was a statistical difference between the nearnatural forest and the other forest classes because of its small sample size. These results contrast the rarefaction findings and might be the result of the indices susceptibility to evenness (Pielou 1966; Pitkänen 1998; Gotelli 2001; Colwell 2009).

The Sorensen index failed to detect a difference within forest classes in the tree species classification, but found a considerable difference between the near-natural and managed forest classes in the diametric-species classification. Furthermore, this index also shows a high similarity between the near-natural and selectively logged forest supporting the rarefaction analysis findings.

The sample size of the near-natural forest was a limiting factor in the analysis which prevented a further extrapolation of the rarefaction curve because of the method used only reliably extrapolated values up to two times the sample size (Chao and Chiu 2016). The sample size also influences the confidence intervals width because the distance between the lower and upper bounds increase the further the extrapolation moves from the reference sample (Jiroutek et al. 2003; Colwell et al. 2012). Furthermore, a near-natural forest's larger sample size would help to visualize the curve's asymptote and determine the completeness of the data, which in turn would give us a better understanding of the real diversity of this forest class (Chao and Jost 2012).

Conclusions

We carried out forest compositional-structural diversity analysis based on a novel diametric-species classification approach and using rarefaction analysis. Our results from mature pine forest with varying degree of human impact showed that the near-natural forest was the most diverse and the managed forest the least diverse. This indicates that forest utilization has had an adverse impact on forest compositional-structural diversity in mature Scots pine forests. On the other hand, the diversity patterns of nearnatural and selectively logged forests are similar, suggesting that selectively logged forests are similar to natural forest in their compositional-structural diversity.

We conclude that the diametric-species classification, combined with the rarefaction analysis, appears to be a useful approach for quantifying and comparing forest structural and compositional diversity, especially in species-poor forests such as the boreal forest. Thus, this classification could be useful in nature conservation applications such as in assessing the level of forest naturalness and conservation potential.

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Authors' contributions

GQ analyzed and interpreted the data on diametric-species diversity and prepared the manuscript. TK provided the data, verified the methods used were correct to achieve the main goals of the research and reviewed the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

The data that support the findings of this study are available from the University of Helsinki boreal forest dynamics and biodiversity research group but restrictions apply to the availability of these data, which were used under license for the current study, and so are not publicly available. Data are however available from the authors upon reasonable request and with permission of the University of Helsinki boreal forest dynamics and biodiversity research group.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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