Light detection and ranging explains diversity of plants, fungi, lichens, and bryophytes across multiple habitats and large geographic extent

Abstract. Effective planning and nature management require spatially accurate and comprehensive measures of the factors important for biodiversity. Light detection and ranging (LIDAR) can provide exactly this, and is therefore a promising technology to support future nature management and related applications. However, until now studies evaluating the potential of LIDAR for this field have been highly limited in scope. Here, we assess the potential of LIDAR to estimate the local diversity of four species groups in multiple habitat types, from open grasslands and meadows over shrubland to forests and across a large area (~43,000 km²), providing a crucial step toward enabling the application of LIDAR in practice, planning, and policy-making. We assessed the relationships between the species richness of macrofungi, lichens, bryophytes, and plants, respectively, and 25 LIDAR-based measures related to potential abiotic and biotic diversity drivers. We used negative binomial generalized linear modeling to construct 19 different candidate models for each species group, and leave-one-region-out cross validation to select the best models. These best models explained 49%, 31%, 32%, and 28% of the variation in species richness ($R^2$) for macrofungi, lichens, bryophytes, and plants, respectively. Three LIDAR measures, terrain slope, shrub layer height and variation in local heat load, were important and positively related to the richness in three of the four species groups. For at least one of the species groups, four other LIDAR measures, shrub layer density, medium-tree layer density, and variations in point amplitude and in relative biomass, were among the three most important. Generally, LIDAR measures exhibited strong associations to the biotic environment, and to some abiotic factors, but were poor measures of spatial landscape and temporal habitat continuity. In conclusion, we showed how well LIDAR alone can predict the local biodiversity across habitats. We also showed that several LIDAR measures are highly correlated to important biodiversity drivers, which are notoriously hard to measure in the field. This opens up hitherto unseen possibilities for using LIDAR for cost-effective monitoring and management of local biodiversity across species groups and habitat types even over large areas.

Key words: airborne laser scanning; ecospace; generalized linear model; remote sensing; species richness; terrain structure; vegetation structure.

INTRODUCTION

Nature management typically aims to create, restore, or conserve specific landscape or vegetation structures or natural processes that are favorable for high levels of biodiversity (Polasky et al. 2008, Landis 2017). However, explaining variation in biodiversity across different organism groups and habitats remains a major challenge (Pennisi 2005). A number of abiotic environmental factors related to soil and hydrology are known to influence local terrestrial biodiversity (Pharo and Beattie 1997, Ejræs and Bruun 2000, Moeslund et al. 2013a,
Brunbjerg et al. 2017b). On the other hand, despite the fact that for example vegetation structure and temporal continuity has long been recognized as important drivers of local diversity (MacArthur and MacArthur 1961, Hermy et al. 1999), the role of biotic structures and resources as well as spatiotemporal continuity remains hard to quantify and disentangle (Elton 1966, Nordén et al. 2014, Brunbjerg et al. 2017b). Here, we use a comprehensive biodiversity inventory to investigate if light detection and ranging data (LIDAR) acquired by airborne laser scanning can adequately represent both the abiotic environment and the biotic factors shaping local biodiversity, and hence allow for effective prediction of the variation in local richness of plants, fungi, lichens, and bryophytes across a large spatial extent.

Light Detection and Ranging is increasingly used as a tool for exploring, explaining, and predicting biodiversity (Ceballos et al. 2015, Peura et al. 2016, Zellweger et al. 2016, Guo et al. 2017, Vihervaara et al. 2017). An airborne LIDAR scanner records a three-dimensional set of points at sampling densities of typically 0.1–100 points/m² using a multi-sensor system combining laser ranging, systematic scanning, high accuracy positioning, and attitude recording (Wehr and Lohr 1999). Since both terrain- and vegetation surfaces reflect the laser signal, a LIDAR point cloud includes direct information on both topography and vegetation structures.

The potential of LIDAR-based metrics for investigating and predicting species diversity has already been recognized for local-to-regional scale studies of various species groups (Veihmas et al. 2009, Lopatin et al. 2016, Peura et al. 2016, Thers et al. 2017, Mao et al. 2018). Such studies typically used LIDAR-based indicators of general vegetation structure, such as vegetation height, variance of point height in individual height layers (Froidevaux et al. 2016), and the count (or density) of points in various height layers (Veihmas et al. 2009, Mao et al. 2018). Most studies using these vegetation-structure measures were restricted to forests (but see Thers et al. 2017) and have shown that local species richness can be modeled with explanatory power up to 66% for plants and up to 82% for fungi (Lopatin et al. 2016, Peura et al. 2016, Peura et al. 2016, Thers et al. 2017). Instead of vegetation-structure measures, other studies have used terrain measures derived from LIDAR-based digital terrain models (DTM) such as aspect, elevation above sea level, slope, topographic wetness (Moeslund et al. 2013a, Mao et al. 2018), or depth-to-water indices (Bartels et al. 2018). For example, recent work showed that the predictive power of LIDAR-based terrain measures were ~20% and 5–16% for predicting local plant (Moeslund et al. 2013a) and bryophyte diversity (Bartels et al. 2018), respectively. All the studies of local alpha biodiversity that we are aware of, address only one species group or one habitat type and hence none of them are able to generalize their findings across multiple habitat types and species groups. In fact, several of these studies conclude that the next step is to evaluate and validate their modeling results at broader spatial scale and across multiple habitats and species groups (Peura et al. 2016).

Here, we present a nationwide evaluation of how measures of terrain and vegetation structure, represented by LIDAR measures, can be used to study local biodiversity patterns across multiple terrestrial habitat types and several species groups in Northern Europe. More specifically, we addressed the following questions: (1) to what extent can LIDAR-derived measures (termed “LIDAR measures” hereafter) predict local species richness of vascular plants, macrofungi, bryophytes and lichens across national extent and various habitat types? (2) What are the most important LIDAR measures for predicting local biodiversity, and which aspects of the locally measured environment do they represent?

**Methods**

**Study area**

Data for this study were collected in Denmark (excluding the island of Bornholm), which has an area of ~43,000 km². Denmark is a North European country in the temperate climate zone and is characterized by a lowland landscape (maximum elevation ~170 m above sea level).

**Biodiversity data**

Data on biodiversity were collected in the non-winter periods of 2014–2015 as part of a comprehensive biodiversity project covering 130 sites (40 × 40 m) distributed throughout Denmark (Fig. 1, Brunbjerg et al. 2017a). One hundred of the study sites represented natural and seminatural habitats. Ten of these were believed to be biodiversity hotspots, while 90 plots were selected by stratified random sampling to cover 5 replicates of 18 combinations of positions along three major natural gradients: fertility (rich, poor), moisture (dry, moist, wet), and successional stage (early, mid, late). Additionally, 15 intensively (ploughed and harvested every year) and extensively (grazed and set-aside) cultivated fields as well as 15 managed forest sites were included. For logistic reasons, study sites were clustered into 15 clusters in five regions as shown in Fig. 1. Three sites were left out from analyses either because (1) they were completely inundated during the period where LIDAR data were recorded (causing these data to be erroneous) or (2) their shape was altered by construction works during the biodiversity data collection period.

Leading experts carefully identified all plant, bryophyte, lichen, and macrofungal species found in each of the study sites, both those found on soil, stones, and dead wood as well as those on living herbs and trees (Fig. 1). Each site was inventoried once for lichens, twice for plants and bryophytes, and three times for fungi (in the fruiting season, August–November). Each inventory had a duration of up to 1 h. Subsequently, species not readily identifiable in the field were identified in a lab using appropriate equipment.
All details on the collection of biodiversity data can be found in Brunbjerg et al. (2017a).

**LIDAR data**

We used the latest nationally covering LIDAR point cloud data set collected for Denmark (2015) to quantify and identify terrain and vegetation structures of importance for local biodiversity. This data set was recorded using fixed-wing airplanes and Riegl LMS-680i (Horn, Austria) scanners operating in the near-infrared wavelength (1,550 nm) in a parallel line scan pattern. The airplanes’ flying height was 680 m above ground level and their speed 240 km/h. The data were collected during the leaf-off season in the spring of 2014 (East Denmark) and the fall, winter, and spring of 2014–2015 (West Denmark). The data set has a nominal minimum point density of 4.6 points/m², except for water areas, and is freely available as 1 × 1 km tiles composed of points from multiple strips (available online). In the current study, we also

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used the gridded (0.4 × 0.4 m) digital terrain model (DTM) that were based on the point cloud data set described above (for details see Danish Ministry of Environment 2015, also available online [see footnote 6]).

To represent vegetation and terrain structure, we calculated 25 measures based on the LIDAR data described above. With one exception (terrain roughness, 0.5 m resolution, measure 22 in Table 1), all measures were rasterized in 10 m resolution. For each measure, we calculated its average within circles of 20 m radius around the center of each study site based on these raster-layers. LIDAR data processing was carried out using the OPALS software package v 2.2.0.0 (Pfeifer et al. 2014). The full OPALS script, which also holds the exact settings for each calculation, is available in Data S1 in the Supporting Information. All measures and their relevant characteristics are detailed in the following. However, to understand all calculation details please refer to the references given in Table 1, which also provides an overview of the LIDAR measures used in this study.

Vegetation-structure measures.—To represent succession stage and moisture balance in both vegetation and soil we retrieved the point amplitude (measure 1, Table 1) directly from the points in the LIDAR point cloud. A point’s amplitude is high if the target reflecting the laser light is flat and has a high reflectivity. It is low for tall canopies where the light energy is distributed between a number of returns, for complex or opaque surfaces such as leaves, and for surfaces with low reflectivity. At the wavelength used here, vegetation surface reflectivity (and thus point amplitude) relates to leaf water content (Junttila et al. 2018) and soil moisture (Zlinszky et al. 2014).

To reflect canopy complexity and number of canopy layers we retrieved the number of echoes (measure 2) returned by each laser pulse emission. Single echoes are returned from continuous surfaces (e.g., flat arable fields) larger than the sensor footprint (the area illuminated by each pulse of laser light from the sensor), while multiple echoes are generated when the pulse hits several surfaces at different distances from the sensor (e.g., a relatively open forest with shrubs, under-forest and trees having leaves or twigs at different elevations). Note that in dense forests, some LIDAR pulses may not penetrate and reach surfaces below the upper parts of the canopy and therefore the number of echoes may be relatively low here. Since the number of echoes correlates with the number of overlapping vegetation layers (Wagner et al. 2006), it also represents the leaf area index (LAI). In the point cloud data set used here, the upper limit for number of echoes recorded was five.

To represent vegetation height we subtracted the local DTM from the height of all the individual LIDAR points giving the normalized height (measure 3) and subsequently we computed the tree canopy top height (measure 4). The latter differs from the first in the calculation procedure. Canopy top height is based on the 90th percentile of points above 3 m and below 50 m (Mücke et al. 2014), and is consequently undefined when the surface height is outside this span.

To mirror vegetation penetrability (measures 5–6) we calculated the echo ratio and the root mean square (RMS) of the echo return number. Both measures reflect the penetrability of the canopy (Höfle et al. 2012). Echo ratio is high where the surface is impenetrable and relatively flat and lower where the surface is uneven or penetrable. The RMS of the echo return number is high when the vegetation is relatively tall and dense, and low when vegetation is low or impenetrable. The differences between the two measures are the spatial scale at which they originate and the way they are calculated. The echo ratio was originally calculated for circles centered at each point in the point cloud and with a radius of 1.5 m (the search radius), while each echo return number relates only to the LIDAR footprint, which had a radius of ~0.1 m.

To reflect vegetation density in different canopy layers we calculated the layer density, typically referred to as the point count, in six height intervals aboveground (measures 7–12, see also Fig. 2), starting with 1.5–5 m, upward in steps of 5 m until 30 m (similar to Zellweger et al. 2014). To approximate the local leaf area index (LAI), we calculated the pseudowaveform (measure 13, see also Fig. 2) following van Aardt et al. (2012). This measure is low when the local LAI is high meaning that the canopy is dense and the LIDAR pulses hardly penetrate the canopy. If LAI is lower, the LIDAR pulse can penetrate further into the canopy giving a higher pseudowaveform value. This measure may appear to be similar to measures 5 and 6 but is calculated differently and was originally calculated at a different spatial scale; 0.5 m radius circles centered at each point in the point cloud.

To estimate biomass, we developed a new index of relative biomass (measure 14, see also Fig. 2) by calculating a weighted combination of multiple structural attributes based on the recommendations of McElhinny et al. (2005). Biomass correlates with vegetation height, but is also influenced by vegetation density and vegetation layering. Therefore, we combined normalized height, echo ratio, and number of echoes in a weighted sum to create a measure of relative biomass (Eq. 1). The weighting was selected to obtain a value equal to vegetation height in the simplest cases (large trees, no significant understory), and higher if vegetation is denser or has more layers than in these simple cases

\[
\text{biomass} = \frac{N_z}{3} + \frac{N_z \times \text{ER}}{36} + \frac{N_z \times \#\text{echoes}}{6}.
\]  

\(N_z\) is normalized height of the first LIDAR echoes, \(\text{ER}\) is echo ratio, \(\#\text{echoes}\) is the number of echoes generated by each LIDAR pulse. Before using this measure in our analyses, we checked that it correlated highly with measured factors typically thought to mirror the actual biomass such as litter mass, dead wood volume, vegetation height and the total basal area of trees with a diameter

\[
\text{biomass} = \frac{N_z}{3} + \frac{N_z \times \text{ER}}{36} + \frac{N_z \times \#\text{echoes}}{6}.
\]
| No. | Name                        | Alias              | Var | Unit | Represents                                                                 | Hypothesis, biodiversity | Reference                  |
|-----|-----------------------------|--------------------|-----|------|----------------------------------------------------------------------------|----------------------------|---------------------------|
| 1   | point amplitude             | ENT                | ENT | m    | succession, leaf and soil moisture                                         | depends on success        | Junttila et al. (2018), Zlinszky et al. (2014) |
| 2   | number of echoes            | count              | count | m    | leaf area index, canopy complexity and number of canopy layers              | is higher in more complex vegetation communities |                           |
| 3   | normalized height           | normalized Z       | RAN | m    | vegetation height                                                          | is higher when vegetation height varies | Mao et al. (2018)         |
| 4   | tree canopy top height      | canopy top height  | m    | height of tree canopies above 3 m                                          | may be higher in taller forests when other factors apply as well; may signify old growth forests, which often have high biodiversity |                           |
| 5   | vegetation penetrability 1  | echo ratio         | ENT | %   | vegetation penetrability                                                   | is lower when vegetation is very dense and higher at intermediate levels | Höfle et al. (2012)       |
| 6   | vegetation penetrability 2† | echo no. RMS       | count | m    | vegetation penetrability                                                   | is lower when vegetation is very dense and higher at intermediate levels |                           |
| 7–12| layer density               | PCount[height interval] | count | m    | vegetation density of a given layer in the following height intervals: 1.5–5, 5–10, 10–15, 15–20, 20–25, and 25–30 m | is higher in open landscapes and forests with shrub layers | Zellweger et al. (2014) |
| 13  | local LAI                   | pseudowaveform     | VAR  | m    | local leaf area index (LAI)                                                | is lower when vegetation is very dense and higher at intermediate levels | van Aardt et al. (2012)   |
| 14  | relative biomass            | ENT                | ENT | m    | biomass, litter mass, deadwood                                             | may be higher when biomass is high; may also be high in open habitats with low biomass levels |                           |
| 15  | crown base height           | m                  | m    | height of tree crown bases                                                 | is higher when tree crown base is high, as this may indicate old growth forest | Mao et al. (2018)         |
| 16  | crown span                  | m                  | m    | vertical extent of tree crowns                                             | is higher when crown span is high as this may indicate old growth forest |                           |
| 17  | shrub layer height          | m                  | m    | shrub layer height                                                          | is higher in forests with shrub layers |                           |
| 18  | canopy openness             | VAR                | radian | m    | light conditions                                                           | Is higher when the canopy is not too closed (forests with gaps) | Doneus (2013)             |
| 19  | terrain slope               | DTM Slope          | radian | m    | soil moisture, heat balance, bare soil                                     | depends on terrain slope  | Moeslund et al. (2013a)   |
| 20  | topographic wetness index   | TWI                | m    | soil moisture                                                              | depends on moisture balance |                           |
| 21  | heat load index             | DTM Heat           | VAR  | m    | heat balance                                                               | is often lower when the terrain is very dry may be higher when terrain varies more | Hengl and Reuter (2009), McCune and Keon (2002), Zlinszky et al. (2012) |
| 22  | terrain roughness           | DTM SigmaZ 0.5     | m    | microscale (0.5-m resolution) terrain heterogeneity/roughness             |                           |                           |
at breast height above 40 cm. This was indeed the case (Appendix S1: Table S1).

To represent the height of tree crown bases (i.e., the lowest point of a crown) we calculated the crown base height (measure 15). This is based on the fifth percentile of the height distribution of all LIDAR points above 3 m and below 50 m (Mao et al. 2018).

To reflect the sizes of tree crowns we calculated the crown span (measure 16). This is the height difference between the tree canopy top height (measure 4) and the crown base (measure 15).

As an estimation of the shrub layer height (measure 17), we calculated the 90th percentile of the normalized heights between 0.3 and 3 m.

To represent light conditions, we calculated canopy openness (measure 18) for all points categorized as “ground,” but contrary to terrain openness (see description of measures 23–24), we calculated this considering vegetation points as well. Therefore, canopy openness relates to the actual occlusion of sky view of ground points by the canopy around them. Canopy openness is high for ground points inside canopy gaps, and low for ground points beneath a closed canopy.

**Terrain-structure measures.**—To represent key features of the local terrain (e.g., soil moisture or heat balance; Moeslund et al. 2013b), we calculated terrain slope (measure 19) and terrain aspect (used for heat load index calculation, see below) directly from the DTM.

As a proxy for local moisture conditions we used the topographic wetness index (TWI, measure 20, Hengl and Reuter 2009) from Moeslund et al. (2013a). To match the resolution of the rest of the measures we aggregated (average) this TWI layer to 10 × 10 m.

To reflect local heat balance, we calculated the heat load index (measure 21) based on terrain aspect, following the heat load index formula in McCune and Keon (2002). This index reaches maximum values on southwest-facing slopes and zero on northeast-facing slopes.

To estimate local terrain roughness (measure 22), we used the points classified in the point cloud as “ground” to calculate sigma Z at a 0.5 × 0.5 m resolution (i.e., a robust indicator of standard deviation) with a search radius of 0.75 m. Note, that this was the only LIDAR measure not rasterized at 10 × 10 m resolution enabling us to test for micro-scale terrain heterogeneity effects.

To represent local and landscape scale terrain heterogeneity, we calculated the terrain openness (measures 23–24, also known as sky-view factor; Doneus 2013) at 10- and 150-m spatial scales (kernel radius). Terrain openness is defined as the angle of a cone (having the radius of the kernel) turned upside down, with its tip restrained to the point of interest, when it touches the points closest to the surface normal vector. This measure is high in flat (relative to the scale at which it is calculated) areas and low in heterogeneous terrains.

To estimate the terrain linearity (measure 25) we calculated the difference between minimum and maximum terrain openness (see above). Maximum openness is high if at least some part of the terrain is open, whereas minimum openness is high when the terrain is open in all directions surrounding the point of interest. In randomly rough surfaces, minimum and maximum openness are quite similar, but in terrain locations with linear features, maximum openness is high (along a ditch or embankment for example) while minimum openness is low (along the sides of a linear terrain feature). Therefore, the difference in minimum and maximum openness is high where linear features with a clear direction, typically human-made, occur (Zlinszky et al., 2015).

To enable a test of the importance of variability in the LIDAR measures we calculated a number of variability measures: standard deviation, root mean square error, and Shannon entropy and in some cases the range. We did this only for LIDAR measures for which we believed it made ecological sense (the measures marked with a variability measure in Table 1).

**Locally measured environmental data**

To support the ecological interpretation of our LIDAR measures, we used data for a number of biotic and abiotic factors. These factors were measured or estimated at each of the study sites. The protocols for these
measurements and estimates can be found in Brunbjerg et al. (2017a). We obtained data on the following 15 locally measured or estimated factors for this study: mean difference of day and night temperatures for (1) air and (2) ground surface, respectively, (3) median light intensity all year, (4) median soil moisture in May, (5) leaf nitrogen (N), (6) leaf phosphorus (P) and (7) leaf N:P ratio, (8) soil N, (9) soil P, and (10) soil pH, (11) litter mass, (12) total basal area of trees larger than 40 cm diameter at breast height, (13) deadwood volume, (14) mean herb layer height, and (15) temporal continuity (year since the most recent major disturbance of habitat).

**Data preparation**

For a given LIDAR measure, its variability measures (e.g., the root mean square error, Shannon entropy, and...
standard deviation of vegetation height) were always highly correlated (Spearman’s rho >0.7; Appendix S1: Fig. S1). Consequently, for further analysis we retained only the variability measure (for each LIDAR measure) showing the highest mean correlation to the species richness of all four species groups (Table 1 shows which variability measure we retained).

For statistical analysis, we used the species richness of plants, bryophytes, lichens, and macrofungi as response variables, modeling each species group individually. We used both the LIDAR measures and their respective variability measures (as shown in Table 1, 25 LIDAR measures and seven variability-measures, in total 32) as predictors in our models.

Prior to analysis, the nature of each predictor’s relationship to the response variable was checked visually and the predictor in question was either logarithmically or square-root transformed if needed to ensure normality (Table 1). In a few cases (i.e., two to four, depending on species group) this check caused us to suspect quadratic relationships. In these cases, we used Akaike’s information criterion (AIC) to evaluate if including the squared term of the predictor improved the model (see Statistical modeling for further description of the modeling approach). For this, we used a backward stepwise model selection procedure based on Akaike’s information criterion (the function stepAIC in the MASS package for R, version 7.3-49; Venables and Ripley 2002). Since this evaluation did not reveal any quadratic relationships, we used only linear terms in the statistical modeling described below.

**Statistical modeling**

We used generalized linear models (GLMs) to examine the explanatory power of the airborne LIDAR-based measures for local species richness. Species richness (count data) is usually expected to follow a Poisson distribution. However, initial implementation of GLMs with a Poisson error distribution and logarithmic link function were overdispersed. Therefore, we used negative binomial GLMs.

To evaluate the performance of all LIDAR-based measures, avoid issues related to multi-collinearity, select the best model and evaluate each predictor’s cross-model importance for local species richness, we implemented the following procedure, which is described in detail in the following paragraphs. We (1) constructed 19 candidate models for each species group pinpointing the best predictors of local species richness among each of 19 sets of uncorrelated LIDAR measures. We then (2) selected the best model for each species group based on fivefold leave-one-region-out cross validation. Finally, we (3) calculated cross-model importance values for each individual predictor based on Akaike weights. These steps are detailed in the following.

**Candidate models.**—We wished to evaluate the performance of all predictors, as there is no consensus on which LIDAR measures act as the most optimal predictors of local diversity. To avoid issues with multi-collinearity, we therefore (1) calculated the pairwise correlations between all possible combinations of predictors, and subsequently (2) divided the predictors into 19 sets of uncorrelated predictors (i.e., Spearman’s rho ≤0.7 with any other predictor in the set; Appendix S1: Fig. S1 shows all pairwise correlations) making sure that each predictor was present in at least one of these sets. Then, (3) for every combination of species group and predictor set, we constructed negative binomial GLM models having species richness as response variable, and each predictor in a particular predictor set as explanatory variables. Finally, (4) for each of these full models (having all predictors in a set as explanatory variables), we used a backward stepwise model selection procedure based on Akaike’s information criterion (the function stepAIC in the MASS package for R, version 7.3-49; Venables and Ripley 2002), to throw away unimportant predictors. This procedure resulted in 19 candidate models, each having only the LIDAR measures important for local diversity as predictors, for every species group.

All modeling details including all full and candidate models are shown in Appendix S1: Tables S2–S5.

**Explanatory power of LIDAR-based measures for local species richness.**—To evaluate the explanatory power of the LIDAR measures for local biodiversity, we selected the overall best model between the set of 19 candidate models for each species group. This was achieved by conducting a fivefold leave-one-region-out cross validation for every candidate model. Thus, for each region (see Biodiversity data and Fig. 1) we predicted species richness using models calibrated on data from the other four regions. We used nonparametric rank correlation (Spearman’s rho) between predicted and observed values to select the best model for each species group. This procedure was adopted to secure robust model selection with respect to overfitting, potential multi-collinearity and spatial autocorrelation. During model selection, we did not encounter issues with non-normally distributed model residuals.

**Cross-model importance of individual LIDAR-based measures and their relation to locally measured environmental factors.**—To evaluate the cross-model importance of the individual LIDAR measures we constructed an importance measure based on Akaike weights following Johnson and Omland (2004). To account for the fact that some variables were only allowed into a model once (if highly correlated with other predictors), and others were included in many or all models, we had to modify the importance measure for each predictor. Therefore, initially each standardized coefficient was weighted with the model’s Akaike weight following Johnson and Omland (2004), summed and then finally this resulting value was multiplied by a predictor weight. This predictor weight was the number of times the variable was
RESULTS

LIDAR-biodiversity relations

We found that our LIDAR-based measures have considerable predictive power for species richness in all species groups investigated (Table 2). Our best models, which had four to seven LIDAR measures as predictors, yielded explanatory powers ($R^2$) of 0.49, 0.31, 0.32, and 0.28 for fungi, lichens, bryophytes, and plants, respectively. For all model details, see Appendix S1: Tables S2–S5.

### Table 2. Best model (based on highest cross validation score) and variable importance details.

| LIDAR measure               | Measure number | Coefficient | Absolute importance | Coefficient | Absolute importance | Coefficient | Absolute importance | Coefficient | Absolute importance |
|-----------------------------|----------------|-------------|---------------------|-------------|---------------------|-------------|---------------------|-------------|---------------------|
| Vegetation structures       |                |             |                     |             |                     |             |                     |             |                     |
| Point amplitude             | 1              | 0.037       | -0.29               | 0.219       | -0.14               | 0.129       | 0.10                | 0.073       |                     |
| Point amplitude (ENT)       | 1              | 0.017       | 0.18                | 0.120       | 0.21                | **0.209**   | 0.000               |             |                     |
| Relative biomass (ENT)      | 14             | 0.000       |                     | 0.000       |                     | 0.000       | -0.13               | **0.080**   |                     |
| Canopy openness (VAR)       | 18             | 0.000       | 0.25                | 0.224       |                     | 0.003       | 0.001               |             |                     |
| Crown span                  | 16             | 0.26        | 0.003               | 0.000       |                     | 0.000       | 0.000               |             |                     |
| Layer density (1.5–5.0 m)   | 7              | 0.31        | 0.011               | 0.35        | 0.016               | 0.002       | 0.28                | **0.257**   |                     |
| Layer density (10–15 m)     | 9              | 0.000       |                     | 0.000       |                     | 0.001       | -0.22               | **0.222**   |                     |
| Layer density (25–30 m)     | 12             | 0.011       |                     | 0.000       | -0.09               | 0.026       | 0.000               |             |                     |
| Shrub layer height          | 17             | **0.380**   | **0.356**           | 0.24        | **0.218**           |             | 0.021               |             |                     |
| Terrain structures          |                |             |                     |             |                     |             |                     |             |                     |
| Heat load index (VAR)       | 21             | 0.14        | **0.125**           | 0.41        | **0.411**           | 0.22        | **0.218**           |             | 0.000               |
| Terrain roughness           | 22             | 0.002       | -0.28               | 0.220       |                     | 0.000       | 0.000               |             |                     |
| Terrain slope               | 19             | 0.22        | **0.317**           | 0.50        | **0.597**           | 0.14        | 0.130               | 0.001       |                     |

**Notes:** If a standardized coefficient is given, the predictor in question was included in the best model for that particular species group. Since exclusion from the best models does not imply that a predictor is not important for the diversity of a specific species group, absolute importance values of the most important predictors in the study (having absolute importance ≥0.02 for at least one species group) are also shown. The absolute importance values of the three most important predictors are shown with boldface type. The names of all predictors that are among these three most important for at least one species group are also highlighted in boldface type. All details on the modeling results are available in Appendix S1:Tables S2–S5. CVS, cross-validation score.
The relative importance of LIDAR measures for biodiversity

Three LIDAR measures were important for three of the four species groups (fungi, lichens, and bryophytes): terrain slope, shrub layer height, and variation in local heat load (Table 2). These were all positively related to local diversity. In addition to these three, four other LIDAR measures (i.e., seven in total) were ranked among the three most important for at least one of the species groups: point amplitude entropy, shrub layer density (1.5–5 m), medium-tree layer density (10–15 m), and variation in relative biomass (Table 2). Generally, these measures were also included in the best models for each species group (Table 2). While some of the measures were important for multiple species groups, some showed importance for only one or two species groups. These are detailed in Table 2 and illustrated in Fig. 3.

Fig. 3. LIDAR point cloud cross sections and field photographs of characteristic species-rich locations for the four species groups. High species richness for bryophytes (top panel) was related to relatively steep terrain with relatively wet soils and a dense shrub layer. Locations with high vascular plant species richness (second from top) were open areas with low variability in biomass and high density in the shrub (1.5–5.0 m) layer and low density of trees. For macrofungi (third panel from top) species richness were highest in areas with steep terrain with relatively wet soils but variable soil moisture levels, and a high degree of typical features for old-growth forest such as large crown spans, dead wood, high litter mass, and dense understory. High species-richness sites for lichens (bottom panel) were found in steep areas with a tall understory and variable canopy openness and on relative dry soils with variable moisture levels but little micro-topographic variation.
TABLE 3. Spearman’s rho (only statistically significant values are shown, \( P < 0.05 \)) of pairwise correlations between the most important predictors (having absolute importance \( \geq 0.02 \) for at least one species group) and 15 environmental variables measured at each study site.

| LIDAR measure          | Measure number | Night-day temp. diff. (ground) | Night-day temp. diff. (air) | Median light intensity | Median soil moisture | Mean leaf N content | Mean leaf P content | Mean leaf N:P ratio | Mean soil N content | Mean soil P content | Mean soil pH | Litter mass | Tot. bas. area of trees > 40 cm (DBH) | Deadwood volume | Mean herb layer height |
|------------------------|----------------|-------------------------------|----------------------------|------------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|--------------|-------------|----------------------------------------|------------------|----------------------------------------|
| Vegetation structure   |                |                               |                            |                        |                     |                     |                     |                     |                     |                     |              |             |                                        |                  |                                        |                  |
| Point amplitude        | 1              | 0.47                          | 0.51                       | 0.46                   | −0.24               | 0.27                | 0.19                | −0.42               | −0.31               | −0.47               | 0.24         |            |                                        |                  |                                        |                  |
| Point amplitude (ENT)† |                |                               |                            |                        |                     | −0.27               |                     |                     |                     |                     |              |             |                                        |                  |                                        |                  |
| Relative biomass (ENT) † | 14             | −0.65                         | −0.66                      | −0.63                  | 0.28                | −0.19               |                     | 0.67                | 0.61                | 0.58                | −0.19        |            |                                        |                  |                                        |                  |
| Canopy openness (VAR)  | 18             | 0.52                          | 0.51                       | 0.45                   | −0.34               | −0.32               | −0.32               | −0.32               | −0.42               | −0.39               | 0.33         |            |                                        |                  |                                        |                  |
| Crown span             | 16             | −0.76                         | −0.73                      | −0.77                  | 0.39                | −0.27               |                     | 0.79                | 0.77                | 0.73                | −0.34        |            |                                        |                  |                                        |                  |
| Layer density (1.5–5.0 m)† | 7              | −0.55                         | −0.49                      | −0.59                  | 0.25                |                     |                     | 0.62                | 0.41                | 0.50                |              |            |                                        |                  |                                        |                  |
| Layer density (10–15 m) † | 9              | −0.74                         | −0.74                      | −0.74                  | 0.32                | 0.20                | −0.22               | −0.20               | −0.35               | 0.78                | 0.68        | 0.69       | −0.41                                  |                  |                                        |                  |
| Layer density (25–30 m) † | 12             | −0.50                         | −0.51                      | −0.49                  | 0.34                |                     |                     | 0.48                | 0.73                | 0.56                | −0.37        |            |                                        |                  |                                        |                  |
| Shrub layer height†    | 17             | −0.58                         | −0.57                      | −0.59                  | 0.27                | −0.19               |                     | 0.64                | 0.44                | 0.50                |              |            |                                        |                  |                                        |                  |
| Terrain structure      |                |                               |                            |                        |                     |                     |                     |                     |                     |                     |              |             |                                        |                  |                                        |                  |
| Heat load index (VAR) † | 21             | −0.20                         | −0.18                      | −0.24                  | 0.27                | −0.20               | −0.23               | −0.38               | 0.30                |                     |              |            |                                        |                  |                                        |                  |
| Terrain roughness      | 22             | −0.20                         | −0.18                      | −0.24                  | 0.27                | −0.20               | −0.23               | −0.38               | 0.30                |                     |              |            |                                        |                  |                                        |                  |
| Terrain slope†         | 19             | −0.29                         | −0.34                      | −0.29                  | −0.36               |                     |                     |                      |                     |                     |              |            |                                        |                  |                                        |                  |

Notes: The names of predictors and environmental variables in bold denote those involved in at least one strong (\( r > 0.7 \)) relationship. The environmental factors are divided into the ecospace (Brunbjerg et al. 2017b) components position and expansion while continuity factors are not shown since no important (absolute importance \( > 0.02 \)) predictors were significantly correlated with any of these. The measure number relates to Table 1 where descriptions of the LIDAR measures can be found.

ENT, Shannon entropy; VAR, variation; temp., temperature; diff., difference; N, nitrogen; P, phosphorus; tot., total; bas., basal; DBH, diameter at breast height.
† Predictors marked that were ranked among the three most important for at least one of the species groups (see Table 1).
The seven LIDAR measures ranked as most important for local biodiversity were strongly correlated to several of the locally measured abiotic and biotic variables (Table 3). Generally, the most important LIDAR measures representing vegetation structure (i.e., vegetation density, shrub layer height, and relative biomass variation) correlated (all negatively) most strongly with the measured diurnal temperature differences and local light conditions and were less related to the other measured abiotic factors. Additionally, out of all the measured biotic factors, these LIDAR measures correlated most strongly (all positively) with litter mass, the volume of dead wood, and the coverage of old trees in the study sites (Table 3). The terrain LIDAR measures (i.e., local heat balance and terrain slope) were mainly related to locally measured soil moisture (Table 3). Generally, these LIDAR measures showed weaker relationships to the measured environment than those reflecting vegetation structure (Table 3).

The LIDAR measures most strongly correlated to local measurements of both abiotic and biotic factors (Spearman’s rho > 0.7) were the span of tree crowns and the vegetation density in the medium-tree (10–15 m) to upper (up to 30 m) height layers (Table 3). We also note that some of the factors typically thought to be important for local biodiversity such as soil moisture and vegetation height, were actually quite strongly related to a couple of our LIDAR measures (e.g., the topographic wetness index; Appendix S1: Table S1). However, these were not among the most important LIDAR measures for local biodiversity identified in this study.

**DISCUSSION**

*The ability of LIDAR to explain local species richness*

Using predictors from no other sources than LIDAR, our models explained a considerable amount of the variation in local biodiversity. However, for some species groups the explanatory power was substantially higher than for others. Notably, LIDAR measures explained the variation in diversity of macrofungi considerably better than the diversity of the other species groups. Although grasslands can hold quite a number of fungi species (Heilmann-Clausen and Vesterholt 2008), this group of organisms is notoriously known for its strong associations to old-growth structures, with old forests typically holding many species of macrofungi (Heilmann-Clausen and Vesterholt 2008). These forest structures are well represented by LIDAR derived measures and also known to be important for the diversity of plants, lichens, and bryophytes (Camathias et al. 2013, Zellweger et al. 2015, Lopatin et al. 2016, Mao et al. 2018). However, for these non-fungal groups the importance of terrain structures, microclimate, and soil-related factors are generally found to be more important than vegetation structures (Camathias et al. 2013, Zellweger et al. 2015). In particular, the local diversity in these groups strongly depend on soil characteristics (Ejrnæs and Bruun 2000, Ilomets et al. 2010, Ódor et al. 2013) and these characteristics are not well represented by LIDAR. This may explain the differences in predictive power between the species groups we observed here. Also, some aspects of local diversity may be better represented by functional diversity, and therefore future studies could consider analyzing this important part of diversity in addition to species diversity (Villéger et al. 2008). Such an approach might potentially reveal a stronger relationship between local diversity and environmental structure.

To our knowledge, this is the first study demonstrating the suitability of LIDAR-based measures for predicting local (i.e., a few decameters) biodiversity patterns across several species groups and across all major temperate terrestrial ecosystems including fields, grasslands, wetlands, heathlands, dunes, scrubs, and forests. So far, only a few studies have studied the extent to which LIDAR measures can predict diversity across multiple species groups, and these have included only one habitat type (see, for example, Zellweger et al. 2015, 2016). Intuitively, one could expect LIDAR to predict local diversity in forests better than in open landscapes since LIDAR represents the more complex, three-dimensional, vegetation structure in forests particularly well. However, in our study the explanatory power obtained for all species groups corresponds well to, or is even higher than, results from earlier studies relating LIDAR measures to species richness of fungi, lichens, plants, and bryophytes (Camathias et al. 2013, Moeslund et al. 2013a, Thers et al. 2017, Bartels et al. 2018). This suggests that LIDAR is not only suitable for management and planning of diversity in forests, but is probably more broadly applicable and likely to be a valuable support tool for nature management and planning in open landscapes as well.

*Importance of individual LIDAR measures and their relation to locally measured environmental factors*

A similar set of LIDAR measures were important for both bryophyte and lichen diversity but for fungi and plant diversity the set of important LIDAR measures differed notably. Hence, sites with high local species richness were structurally different in several aspects depending on the species group in question. The most important LIDAR measures for local biodiversity represented both vegetation (shrub layer height, point amplitude entropy, variation in relative biomass, shrub and medium-tree layer density) and terrain structures (slope of the terrain and variation in local heat load). The two terrain-structure measures correlated mostly with local soil moisture conditions, while the vegetation-structure measures were mainly associated with local light conditions and diurnal temperature variations, as well as biotic factors such as litter mass, stand age, and the amount of dead wood.
In previous studies, local terrain structure has been shown to affect both the occurrence, abundance, and species richness of macrofungi (Peura et al. 2016, Thers et al. 2017, Chen et al. 2018). Here, we found fungal species richness to be highest in areas with steep terrain with relatively wet soils, while at the same time having many typical features for old-growth forest such as large crown spans, large amounts of dead wood, high litter mass, and a dense shrub layer. This supports that this group is typically strongly associated with old-growth structures (Heilmann-Clausen and Vesterholt 2008) and suggests that steep slopes in this case could reflect refugia from human impact (Odgaard et al. 2014).

The most important LIDAR measures for local diversity of bryophytes and lichens were almost the same. This hints that in nature the same natural factors determine the local diversity patterns of these two species groups, a finding that was also highlighted by Pharo and Beattie (1997). However, for lichens, terrain slope was the most important predictor and had a strong positive relationship to local species richness, whereas for bryophytes, variation in heat load index was an important predictor. Since we found terrain slope negatively, and variation in heat load index positively related to locally measured soil moisture, we believe these results support (Pharo and Beattie 1997); bryophyte richness is higher in relatively moist sites and lichen richness is higher in the drier sites. For bryophytes and lichens, local diversity decreased with mean point amplitude and increased with point amplitude variation. As point amplitude can be interpreted as a measure of successional stage from bare soil (high point amplitude) to closed forest (low point amplitude, see Methods), this indicates that species richness of bryophytes and lichens is often higher in late successional stages (old growth forests and old scrubland). Furthermore, local diversity of both groups was positively related to shrub layer height, which were associated with light availability, microclimatic conditions, litter mass, stand age and the amount of dead wood. These findings correspond to the current knowledge based on single-habitat studies. For example, Mills and Macdonald (2004) and Zellweger et al. (2015) showed that microsite bryophyte diversity in forests was clearly affected by dead wood characteristics, and local levels of soil moisture, temperature and solar radiation among others. Similarly, Leppik et al. (2013) found that forest lichen diversity increased with stand age and soil moisture. Note, that while we assess the importance of LIDAR measures for all lichens and bryophytes, considerable differences in predictor importance can be expected among edaphic and epiphytic species (Camathias et al. 2013).

For vascular plants, we found high species richness at localities with high density in the shrub layer and low density of medium sized trees, and in areas with low variability in relative biomass and a high mean point amplitude (indicating early successional stage, i.e., open landscapes). These results suggest that plant diversity is often high in open landscapes, for example, grasslands, which are known hotspots for plants in Northern Europe (Habel et al. 2013). On the other hand, they also indicate that areas with relatively many shrubs or small trees are rich in plant species. A combination of two processes might explain this. First of all, grasslands are threatened by shrub encroachment (Timmermann et al. 2015) and the diversity–density relationship could therefore reflect an extinction debt to unfavorable habitat conditions following encroachment. Secondly, shrubs create additional microhabitats in open grasslands and could thereby increase richness. The later supports that plant diversity can be promoted by the presence of single-standing trees and bushes in otherwise homogenous grassland swards (Moeslund et al. 2017). Contrary to previous findings, we did not find evidence that terrain-related factors are important for determining the species richness of plants. For example, terrain controlled soil moisture has been found important for local diversity of plants in open habitats and is generally regarded as important for plant species richness (Moeslund et al. 2013a, Silvertown et al. 2015). However, we found no clear indications of a relationship between soil moisture and plant diversity. We included forests in this study and here soil moisture is probably less important for shaping local vegetation patterns (Zellweger et al. 2015) due to the more moist local climate mediated by trees. This may explain the lack of this otherwise important relationship in the present study. However, local environmental variation unaccounted for by LIDAR might also mask the effect of soil moisture. Exploring this in more detail could be the focus of future studies.

Recently, Brunbjerg et al. (2017b) proposed the *ecospace* framework. Within this framework, three components define an *ecospace*: position, expansion, and continuity. *Position* is given by all relevant abiotic factors for the local diversity at a given site, for example soil moisture, pH, nutrient ion availability, and temperature. *Expansion* represents the resources (diversification and build-up of organic matter) for species to live on and from, for instance the amount of dead wood, flowers, insects, carcasses, dung, and leaf litter. *Continuity* refers to the spatial and temporal extension of expansion and position. Our LIDAR measures captured major aspects of the environmental variation related to build up and diversification of organic matter (i.e., the *ecospace expansion sensu* Brunbjerg et al. [2017b]). For example, the measured litter mass, the basal area of old trees (having stem diameter at breast height > 40 cm) and the volume of dead wood were all highly correlated with at least one of our LIDAR measures. Several studies have reached a similar conclusion in forests (Camathias et al. 2013, Zellweger et al. 2015, Lopatin et al. 2016). However, our results demonstrate for the first time that LIDAR can be used to estimate expansion-related factors along the full successional gradient from open wetlands, grasslands, and fields to shrubs and forests. This opens interesting perspectives for applying LIDAR more
broadly in nature management and planning (see section Application perspectives).

The LIDAR measures used in this study were good representatives of soil moisture and local temperature conditions, but failed to act as proxies for other ecosphere position related factors such as soil pH and nutrient status. By nature, LIDAR does not record anything belowground, nor any chemical properties. This renders LIDAR alone unsuitable for recording soil and leaf chemistry, but there could be differences in terrain and vegetation structure across habitat types mirroring these factors of which we are unaware. Denser point clouds and terrestrial laser scanning combined with new machine learning techniques (Liu et al. 2018), spectral information (Lausch et al. 2016), and emerging environmental DNA technologies (Bush et al. 2017) might potentially remedy this situation in the future. Notably, advances within the field of imaging spectroscopy shows promising potential for mitigating the lack of information on for instance nutrient balance and soil pH, characterizing LIDAR measures. For example, Singh et al. (2015) recently showed how imaging spectroscopy can be used to map the percentage of for instance nitrogen, carbon, lignin, and fibers in plant leaves. Including such information may improve the prediction of local biodiversity significantly by complementing the information found in LIDAR measures. On the other hand, some studies have also demonstrated that adding spectral remote-sensing-based measures in the modeling of local biodiversity patterns did not improve modeling performance (Leutner et al. 2012, Ceballos et al. 2015). Therefore, we incite researchers to carefully design studies giving insight into how LIDAR and spectral remote sensing can be amalgamated to effectively complement each other and consequently improve the prediction of local biodiversity.

Spatial scale affects ecological patterns and processes (Levin 1992), and therefore our selection of grain size (10 m) can affect the importance of each of the measures that we included here. For example, our LIDAR measures were generally quite poor proxies for ecospace continuity (Brunbjerg et al. 2017b). Since spatial continuity by definition is a broad-scale factor (Nordén et al. 2014), it is not surprising that this factor was poorly represented by our LIDAR measures. Though, by considering LIDAR point clouds for a larger area than we did in this study, we believe LIDAR could be valuable for estimating spatial continuity and we urge researchers to attempt to do this in future studies. On the other hand, temporal continuity is tricky to estimate from LIDAR, at least in open landscapes where the structures and vegetation patterns characteristic of long continuity are difficult to capture. For example, considerable plant species turnover can occur over time from abandoned fields to heathland and grassland, without noticeable changes in vegetation structure (Ejrnæs et al. 2008). Contrary, in forests, as we have shown, some of the structures characteristic of old-growth can be estimated with high confidence using LIDAR. Clearly, more work and possibly technological advances are needed to find methods to effectively estimate temporal continuity using LIDAR.

Application perspectives

Using LIDAR, researchers and managers have gained the ability to cover large areas (even whole nations) in adequately fine detail for nature planning and management (McElhinny et al. 2005). In fact, LIDAR has the potential to play a crucial role in this applied field by enabling detailed mapping and assessments for decision makers and field biologists, while at the same time covering the desired extent. The best candidates for LIDAR measures with potential for supporting conservation planning and management are those being notoriously difficult to quantify in the field, having high importance across species groups and a plausible ecological interpretation. Generally, a measure such as shrub layer height is indeed complicated and time-consuming to quantify in the field, but had high importance for most species groups in this study and seemed to capture several of the factors of importance for local biodiversity patterns (see previous discussion). Another such measure is the variability of the terrain aspect-based heat load index, which proved to represent soil moisture well and be important for most species groups in our study. While more accurate, terrain-based, wetness measures exist (Hengl and Reuter 2009, Moeslund et al. 2013a), this indicator has the advantage that it is computationally efficient since it can be calculated without the need to delineate watersheds. Hence, these LIDAR measures are two out of potentially several that may be successfully implemented in planning and management, and used to, for example, create quick first impressions of local biodiversity patterns across large areas in the future.

In summary, our results show that LIDAR alone can provide reasonable predictive power for biodiversity, giving insights into local biodiversity patterns and their potential drivers. By refining these methods, for example by (1) including full-waveform LIDAR (Anderson et al. 2016), (2) further investigate LIDAR-based measures for assessing continuity and maybe soil chemistry (see previous discussion), and (3) possibly combining LIDAR with other available data and methods, this technology could open new avenues offering reliable fine-grain overviews of the biodiversity patterns and potentially also dynamics for whole countries. This would allow hitherto unseen possibilities for evidence-based biodiversity management (Brunbjerg et al. 2016, Mao et al. 2018), and help to precisely target field-based biodiversity monitoring nationwide.

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LITERATURE CITED

Anderson, K., S. Hancock, M. Disney, and K. J. Gaston. 2016. Is waveform worth it? A comparison of LiDAR approaches for vegetation and landscape characterization. Remote Sensing in Ecology and Conservation 2:5–15.

Bartels, S. F., R. T. Caners, J. Ogilvie, B. White, and S. E. MacDonald. 2018. Relating bryophyte assemblages to a remotely sensed depth-to-water index in boreal forests. Frontiers in Plant Science 9:858.

Brunbjerg, A. K., J. Bladt, M. Brink, J. Fredshavn, P. Mikkelsen, J. E. Moeslund, B. Nygaard, F. Skov, and R. Eijrnæs. 2016. Development and implementation of a high nature value (HNV) farming indicator for Denmark. Ecological Indicators 61:274–281.

Brunbjerg, A. K., et al. 2017a. A systematic survey of regional multitalent biodiversity: evaluating strategies and coverage. bioRxiv. https://doi.org/10.1101/158030.

Brunbjerg, A. K., H. H. Bruun, J. E. Moeslund, J. P. Sadler, J-C. Svenning, and R. Eijrnæs. 2017b. Ecospace: A unified framework for understanding variation in terrestrial biodiversity. Basic and Applied Ecology 18:86–94.

Bush, A., et al. 2017. Connecting Earth observation to high-throughput biodiversity data. Nature Ecology & Evolution 1:0176.

Camathias, L., A. Bergamini, M. Küchler, S. Stofer, and A. Balsweneiler. 2013. High-resolution remote sensing data improves models of species richness. Applied Vegetation Science 16:539–551.

Ceballos, A., J. Hernández, P. Corvalán, and M. Galleguillos. 2015. Comparison of airborne LiDAR and satellite hyperspectral remote sensing to estimate vascular plant richness in deciduous Mediterranean forests of Central Chile. Remote Sensing 7:2692–2714.

Chen, Y., Z. Yuan, S. Bi, X. Wang, Y. Ye, and J-C. Svenning. 2018. Macrofungal species distributions depend on habitat partitioning of topography, light, and vegetation in a temperate mountain forest. Scientific Reports 8:13359.

Danish Ministry of Environment. 2015. Danmarks Højde-model, DHM/Punktsky. Data version 2.0. Danish Ministry of Environment, Copenhagen, Denmark.

Doneus, M. 2013. Openness as visualization technique for interpretative mapping of airborne lidar derived digital terrain models. Remote Sensing 5:6427–6442.

Eijrnæs, R., and H. H. Bruun. 2000. Gradient analysis of dry grassland vegetation in Denmark. Journal of Vegetation Science 11:573–584.

Eijrnæs, R., J. Liira, R. S. Poulsen, and B. Nygaard. 2008. When has an abandoned field become a semi-natural grassland or heathland? Environmental Management 42:707–716.

Elton, C. S. 1966. The pattern of animal communities. Methuen, London, UK.

Frodéveaux, J. S. P., F. Zellweger, K. Bollmann, G. Jones, and M. K. Obrist. 2016. From field surveys to LiDAR: Shining a light on how bats respond to forest structure. Remote Sensing of Environment 175:242–250.

Guo, X., N. C. Coops, P. Tompalski, S. E. Nielsen, C. W. Bater, and J. John Stadt. 2017. Regional mapping of vegetation structure for biodiversity monitoring using airborne lidar data. Ecological Informatics 38:50–61.

Habel, J. C., J. Dengler, M. Janisiovi, P. Török, C. Wellstein, and M. Wiezik. 2013. European grassland ecosystems: threatened hotspots of biodiversity. Biodiversity and Conservation 22:2131–2138.

Heilmann-Clausen, J., and J. Vesterholt. 2008. Conservation: selection criteria and approaches. Pages 325–347 in L. Boddy, J. C. Frankland, and P van West, editors. Ecology of saprotrophic basidiomycetes. Elsevier, London, UK.

Hengt, T., and H. I. Reuter. 2009. Geomorphometry: concepts, software, applications, First edition. Elsevier, Amsterdam, The Netherlands.

Hermy, M., O. Honnay, L. Fribank, C. Grashof-Bokdam, and J.-E. Lawesson. 1999. An ecological comparison between ancient and other forest plant species of Europe, and the implications for forest conservation. Biological Conservation 91:9–22.

Höfle, B., M. Hollaus, and J. Hagenauer. 2012. Urban vegetation detection using radiometrically calibrated small-footprint full-waveform airborne LiDAR data. ISPRS Journal of Photogrammetry and Remote Sensing 67:134–147.

Houmets, M., L. Truu, R. Pajula, and K. Sepp. 2010. Species composition and structure of vascular plants and bryophytes on the water level gradient within a calcareous fen in North Estonia. Estonian Journal of Ecology 59:19–38.

Johnson, J. B., and K. S. Omland. 2004. Model selection in ecology and evolution. Trends in Ecology & Evolution 19:101–108.

Junttila, S., J. Sugano, M. Vastaranta, R. Linnakoski, H. Kaartinen, A. Kukko, M. Holopainen, H. Hyyppä, and J. Hyyppä. 2018. Can leaf water content be estimated using multispectral terrestrial laser scanning? A case study with Norway Spruce seedlings. Frontiers in Plant Science 9:299.

Landis, D. A. 2017. Designing agricultural landscapes for biodiversity-based ecosystem services. Basic and Applied Ecology 18:1–12.

Lausik, A., et al. 2016. Linking earth observation and taxonomic, structural and functional biodiversity: local to ecosystem perspectives. Ecological Indicators 70:317–339.

Leppik, E., I. Jüriado, A. Suija, and J. Liira. 2013. The conservation of ground layer lichen communities in alvar grasslands and the relevance of substitution habitats. Biodiversity and Conservation 22:591–614.

Leutner, B. F., B. Reineking, J. Müller, M. Bachmann, C. Beierkuhnlein, S. Dech, and M. Wegmann. 2012. Modelling forest x-diversity and floristic composition—on the added value of LiDAR plus hyperspectral remote sensing. Remote Sensing 4:2818–2845.

Levin, S. A. 1992. The problem of pattern and scale in ecology: the Robert H. MacArthur Award lecture. Ecology 73:1943–1967.

Liu, Z., C. Peng, T. Work, J.-N. Candau, A. DesRochers, and D. Kneeshaw. 2018. Application of machine-learning methods in forest ecology: recent progress and future challenges. Environmental Reviews 26:339–350.

Lopatin, J., K. Dolos, H. J. Hernández, M. Galleguillos, and F. E. Fassnacht. 2016. Comparing generalized linear models and random forest to model vascular plant species richness
using LiDAR data in a natural forest in central Chile. Remote Sensing of Environment 173:200–210.

MacArthur, R. H., and J. W. MacArthur. 1961. On bird species diversity. Ecology 42:594–598.

Mao, L., J. Dennett, C. W. Bater, P. Tompalski, N. C. Coops, D. Farz, M. Kohler, B. White, J. J. Stadt, and S. E. Nielsen. 2018. Using airborne laser scanning to predict plant species richness and assess conservation threats in the oil sands region of Alberta’s boreal forest. Forest Ecology and Management 409:29–37.

McCune, B., and D. Keon. 2002. Equations for potential annual direct incident radiation and heat load. Journal of Vegetation Science 13:603–606.

McEllhiney, C., P. Gibbons, C. Brack, and J. Bauhus. 2005. Forest and woodland stand structural complexity: its definition and measurement. Forest Ecology and Management 218:1–24.

Mills, S. E., and S. E. Macdonald. 2004. Predictors of moss and liverwort species diversity of microsites in conifer-dominated boreal forest. Journal of Vegetation Science 15:189–198.

Moeslund, J. E., L. Arge, P. K. Bocher, T. Dalgaard, M. V. Odgaard, B. Nygaard, and J.-C. Svenning. 2013a. Topographically controlled soil moisture is the primary driver of local vegetation patterns across a lowland region. Ecosphere 4:9/1.

Moeslund, J. E., L. Arge, P. K. Bocher, T. Dalgaard, and J.-C. Svenning. 2013b. Topography as a driver of local terrestrial vascular plant diversity patterns. Nordic Journal of Botany 31:129–144.

Moeslund, J. E., A. K. Brunbjerg, K. K. Clausen, L. Dalby, C. Fligaard, A. Juel, and J. Lenoir. 2017. Using dark diversity and plant characteristics to guide conservation and restoration. Journal of Applied Ecology, 54:1730–1741.

Mücke, W., B. Deak, A. Schroiff, N. Pfeifer, and H. Heilmeier. 2014. Estimation of vertical forest layer structure based on small-footprint airborne LiDAR. Pages 63–66 in N. Pfeifer and A. Zilinszky, editors. Proceedings of the international workshop on remote sensing and GIS for monitoring of habitat quality. Department of Geodesy and Geoinformation, Vienna University of Technology, Vienna, Austria.

Nordén, B., A. Dahlberg, T. E. Brandrud, Ø. Fritz, R. Ejrnæs, and O. Ovaskainen. 2014. Effects of ecological continuity on species richness and composition in forests and woodlands: a review. Ecoscience 21:34–45.

Odgaard, M. V., P. K. Bocher, T. Dalgaard, J. E. Moeslund, and J.-C. Svenning. 2014. Human-driven topographic effects on the distribution of forest in a flat, lowland agricultural region. Journal of Geographical Sciences 24:76–92.

Ödor, P., I. Király, F. Tíny, F. Bortignon, and J. Nascimbene. 2013. Patterns and drivers of species composition of epiphytic bryophytes and lichens in managed temperate forests. Forest Ecology and Management 306:256–265.

Pennisi, E. 2005. What determines species diversity? Science 309:90.

Peura, M., R. Silveyya Gonzalez, J. Müller, M. Heurich, L. A. Vierling, M. Mönnkönen, and C. Bässler. 2016. Mapping a ‘cryptic kingdom’: performance of lidar derived environmental variables in modelling the occurrence of forest fungi. Remote Sensing of Environment 186:428–438.

Pfeifer, N., G. Mändlburger, J. Otepka, and W. Karel. 2014. OPALS—A framework for Airborne Laser Scanning data analysis. Computers, Environment and Urban Systems 45:125–136.

Pharo, E. J., and A. J. Beattie. 1997. Bryophyte and lichen diversity: a comparative study. Australian Journal of Ecology 22:151–162.

Polasky, S., et al. 2008. Where to put things? Spatial land management to sustain biodiversity and economic returns. Biological Conservation 141:1505–1524.

R Core Team. 2018. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. www.R-project.org

Silvertown, J., Y. Araya, and D. Gowing. 2015. Hydrological niches in terrestrial plant communities: a review. Journal of Ecology 103:93–108.

Singh, A., S. P. Serbin, B. E. McNeil, C. C. Kingdon, and P. A. Townsend. 2015. Imaging spectroscopy algorithms for mapping canopy foliar chemical and morphological traits and their uncertainties. Ecological Applications 25:2180–2197.

Thers, H., A. K. Brunbjerg, T. Læsøe, R. Ejrnæs, P. K. Bocher, and J.-C. Svenning. 2017. Lidar-derived variables as a proxy for fungal species richness and composition in temperate Northern Europe. Remote Sensing of Environment 200:102–113.

Timmermann, A., C. Damaaard, M. T. Strandberg, and J.-C. Svenning. 2015. Pervasive early 21st-century vegetation changes across Danish semi-natural ecosystems: more losers than winners and a shift towards competitive, tall-growing species. Journal of Applied Ecology 52:21–30.

van Aardt, J., J. Wu, J. McClinchy, K. J. Wessels, R. S. Mathieu, T. K. Bowdoin, D. E. Knapp, and G. P. Asner. 2012. On using discrete return LIDAR distributions as a proxy for waveform LIDAR signals when modeling vegetation structure. In IEEE International Geoscience and Remote Sensing Symposium. IEEE, Munich, Bavaria, Germany.

Vehmas, M., K. Eerikäinen, J. Pehukurinen, P. Packalén, and M. Maltamo. 2009. Identification of boreal forest stands with high herbaceous plant diversity using airborne laser scanning. Forest Ecology and Management 257:46–53.

Venables, W. N., and B. D. Ripley. 2002. Modern applied statistics with S, Fourth edition. Springer, New York, New York, USA.

Vihervaara, P., et al. 2017. How essential biodiversity variables and remote sensing can help national biodiversity monitoring. Global Biodiversity and Conservation 10:43–59.

Villegger, S., N. W. H. Mason, and D. Mouillot. 2008. New multidimensional functional diversity indices for a multifaceted framework in functional ecology. Ecology 89:2290–2301.

Wagner, W., A. Ullrich, V. Ducic, T. Melzer, and N. Studnicka. 2006. Gaussian decomposition and calibration of a novel small-footprint full-waveform digitising airborne laser scanner. ISPRS Journal of Photogrammetry and Remote Sensing 60:100–112.

Wehr, A., and U. Lohr. 1999. Airborne laser scanning—an introduction and overview. ISPRS Journal of Photogrammetry and Remote Sensing 54:68–82.

Zellweger, F., F. Morsdorf, R. S. Purves, V. Braunisch, and K. Bollmann. 2014. Improved methods for measuring forest landscape structure: LiDAR complements field-based habitat assessment. Biodiversity and Conservation 23:289–307.

Zellweger, F., V. Braunisch, F. Morsdorf, A. Baltensweiler, M. Abegg, T. Roth, H. Bugmann, and K. Bollmann. 2015. Disentangling the effects of climate, topography, soil and vegetation on stand-scale species richness in temperate forests. Forest Ecology and Management 349:36–44.

Zellweger, F., A. Baltensweiler, C. Ginzler, T. Roth, V. Braunisch, H. Bugmann, and K. Bollmann. 2016. Environmental predictors of species richness in forest landscapes: abiotic factors versus vegetation structure. Journal of Biogeography 43:1080–1090.

Zilinszky, A., B. Deak, A. Kania, A. Schroiff, and N. Pfeifer. 2015. Mapping Natura 2000 habitat conservation status in a pannonic salt steppe with airborne laser scanning. Remote Sensing 7:3091–3109.

Zilinszky, A., A. Schroiff, A. Kania, B. Deak, W. Mücke, Á. Vári, B. Szekely, and N. Pfeifer. 2014. Categorizing grassland vegetation with full-waveform airborne laser scanning: a feasibility study for detecting Natura 2000 habitat types. Remote Sensing 6:8056–8087.
DATA AVAILABILITY

The biodiversity data used for this study is stored in a permanent university repository under Aarhus University: http://bios.au.dk/om-instituttet/organisation/biodiversitet/projekter/data/. The LIDAR data is publicly available through kortforsyningen.dk, which is the official map service of Denmark. To use this service, one must register as a user, which is free of charge. The LIDAR data comes in $10 \times 10$ km batches holding $1 \times 1$ km tiles. Each tile is named after the coordinates (UTM Zone 32 N) they cover. Use the coordinates found in the biodiversity data set for each study site, to pinpoint the correct LIDAR tiles.