Individual tree mortality of silver birch (*Betula pendula* Roth) in Estonia

Kobra Maleki, Andres Kiviste

The functioning of complex forest ecosystems is intimately related to their structural properties. Tree mortality is a major driver of forest stand dynamics and therefore plays an important role in the formation of forest structure. Data from the Estonian Network of Forest Research Plots (ENFRP) was used to estimate the mortality probability of silver birch trees (*Betula pendula* Roth) by using logistic models. In this study several spatial and non-spatial variables were tested to determine the most important mortality explanatory factors. Additionally, thinning variables were defined and implemented into the mortality models, to examine whether thinning practices could modify the stand structure and density, then leading to a lower mortality rate. The results of this study showed that tree mortality models that included either a five-year diameter growth rate (Id5), as a measure of tree vitality, or the tree relative diameter (drel) as a measure of competition, or both these two variables, were substantially better than any models not including these variables. In addition, any measures of spatial aggregation (agg) and species proportion (sp) within the zone of influence markedly improved the model predictions, though the mortality probability of trees declined where there was higher aggregation and species mixture. Our results also suggested that if thinning were conducted around the birch trees, depending on the thinning intensity, the number of neighbors is effectively reduced, and consequently the competition load within the influence zone decreases, leading to healthier growth and lower mortality rates of the shade-intolerant birch trees. We thus recommend to adopt thinning regimes in mixed forest stands to foster tree species diversity, and at the same time provide adequate growing space for birch trees within the stands. This will improve the forest structure and increase the adaptive capacity of forests, which is increasingly important under changing climatic conditions.

Keywords: *Betula pendula* Roth, Diameter Growth, Species Proportion, Aggregation, Competition

**Introduction**

Tree mortality is an important driver of forest stand dynamics as it determines the formation of forest structure (Pederson 1998, Van Mantgem & Stephenson 2007). Structural properties of forests are of great importance since they are related to ecosystem functioning and productivity, in its growth and productivity (Bobiec 2002, Froliking et al. 2009).

Tree mortality occurs when tree vigor declines due to intolerance of the tree to the negative influence of stress factors, such as drought and competition. These stress factors can lead to a wide variety of causes of tree mortality. To achieve a better understanding of mortality, numerous attempts to categorize its causes have been reported (Vanclay 1994, Pederson 1998, Yang et al. 2003). As a simplified approach, mortality has been generally considered either as regular, due to competition or ageing, or as irregular, caused by ecological and catastrophic events, such as storm or fire (Monserud 1976). The probability of regular mortality is believed to be U-shaped, indicating relatively high mortality rates for small and large trees due to light competition and senescence, respectively (Monserud & Sterba 1999, Loimer et al. 2001, Fraver et al. 2008). The probability of irregular mortality is rather fluctuating, and thus quite difficult to predict, since it may be triggered by various biotic and abiotic factors, which change temporally and spatially, e.g., the attack of pests or diseases may cause a comparatively small mortality in one year, or eliminate the host species from the region (Vanclay 1994).

The ability of a tree to withstand stress factors is affected by a multitude of interacting factors such as tree size, tree viability, competition among trees and stand density (Hamilton 1986). However, diagnosing these factors as contributors to tree mortality and evaluating their relative importance can be very difficult (Das et al. 2008); especially in times of global changes it is important to broaden our understanding of tree mortality as a primary driver of changes in the composition and structure of forest communities. Maintaining a forest is essential to provide important services, such as carbon sequestration, water purification, and timber production, as well as future habitats for a large diverse number of species (Dietze & Moorcroft 2011). Measurements of tree size can be easily
collected (tree diameter and tree height) or computed (tree growth and tree basal area) and to some extent they contain valuable, but not sufficient, information about the probability of tree mortality (Monserud & Sterba 1999, Yang et al. 2003). Diameter growth rate (Buchman 1983), or as a substitute, crown size (Avila & Burkart 1992, Monserud & Sterba 1999) have proven to be suitable measures of tree vitality in mortality studies.

Competition is the next well-documented contributor to tree mortality. Competition for limited resources (such as water, light and nutrients) is a fundamental ecological process, which has strongly modulated the mortality of suppressed trees (Keddy 1989, Szwagrzyk & Szewczyk 2001). Competition among individuals can be either asymmetric or symmetric. Under asymmetric competition, larger trees are not affected by smaller trees and can therefore benefit of a greater share of resources, whereas in symmetric competition all trees have an equal share of resources, and therefore impose some competitive influence on their neighbors, regardless of their sizes (Cannell et al. 1984, Freckleton & Watkinson 2001). Occasionally, some trees may withstand the competition pressure, but will eventually end up in a “death spiral” (Franklin et al. 1987), since trees suffering competition over long periods will weaken and eventually die due to other reasons (Wyckoff & Clark 2000, Antos et al. 2008, Lännepää et al. 2008).

Theoretically, inclusion of the spatial effects in modeling the competitive environment that surrounds an individual tree is expected to improve the prediction ability of mortality (Biging & Dobbertin 1995). An increment in local density is assumed to increase the density-dependent mortality, or self-thinning, since many species are unable to endure deep shading in their dense neighborhood (He & Duncan 2000, Gómez-Aparicio et al. 2009). In young dense stands where trees are subjected to self-thinning, such assumption was proven to be valid (Oliver & Larson 1996). Several studies have been conducted to define the limit to the number of trees with a given average size growing in a stand (the so-called “limiting density”), depending on the forest type and the environmental conditions (Reineke 1933, Zeide 1987, Hynnen 1993, Nilson 2006, Von Gadow et al. 2015). On the other hand, when stands grow to maturity, generally the size of surviving trees and their spacing increase, and stands become less clumped. In such stands the density-dependent mortality decreases, and trees may die because of other factors, such as pathogen and insect attacks (Kenkel 1988, Franklin & Van Pelt 2004).

Regarding the fact that any desired tree species may be absent due to self-thinning in a final regular stand, thinning during the early self-thinning years is an acceptable management tool to control stand quality and supports favorable tree species (Siimard et al. 2004). Early thinning improves the tree growth by reducing the competition stress for the remaining trees (Oliver & Larson 1996), while the thinning of suppressed trees at intermediate development stages reduces economical loss from tree deaths (Perry 1985). Thinning to different reduced densities influences the tree mortality rate, and individual trees may react differently to the same treatment. For example, small trees survive better in a less competitive neighborhood (Uriarte et al. 2004), while the mortality rate of large trees increases due to exposing their crowns to strong winds when neighboring trees are removed (Thorpe et al. 2008).

We focused our study on the mortality of silver birch trees (Betula pendula Roth), an ecologically essential broadleaf tree species (Hynnen et al. 2010) and the second most important tree species in Estonia in terms of forest cover (31.2% and this coverage is expanding – Yearbook of Forest 2013). As a pioneer tree species (Fischer et al. 2002), silver birch grows naturally in boreal and northern temperate biomes. This species is intolerant to shade and remains vital and vigorous where it occurs as a dominant species in a relatively wide spacing, and under less favorable environmental conditions for other tree species (Hynnen et al. 2010).

The main objectives of this study were to: (i) filter the variables that are meaningful for the mortality of silver birch trees; (ii) test these variables in an empirical model for their ability to predict mortality of silver birch trees; and (iii) examine whether thinning practices effectively minimize the rate of tree mortality in managed forests.

Materials and methods

Study data

Data from the Estonian Network of Forest Research Plots (ENFRP) was used for this study. This network, consisting of 730 permanent plots, was established during the period 1995-2004, and contains the data for all the main forest types in Estonia (Kiviste et al. 2015). The Republic of Estonia lies on the eastern shores of the Baltic Sea and along the southern shore of the Gulf of Finland (57°3’09.5” N, 21°5’28.4” E). The climate varies from maritime to continental. Average temperature ranges from 16.0°C to 18.1°C in July (the warmest month) and from -3.5°C to -7.6°C in February (the coldest month). Mean annual precipitation is within the range 600-700 mm, and decreases from east to west (Tarand et al. 2013).

Clear cutting is the predominant management system in Estonia; consequently overstory trees are mostly even-aged. Most stands have been managed to maintain pure stands, though near-natural situations can occur under good forest growth conditions in any stand after some decades without management. The ENFRP permanent plots were circular with a radius ranging from 10 to 30 meters. Depending on the stand age and density, they were delineated so that every plot contained at least 100 trees in the overstory. Within each plot the azimuth, the distance from plot center, the diameter at breast height (d), the tree species identity, and any defects of trees were recorded. For every fifth tree, and also for dominant and rare tree species, the tree height and the height to the live crown base were also measured. The measurements were repeated at intervals of five years.

To carry out this study, we selected 422 research plots with three consecutive measurements where silver birch was present. Other plots of the ENFRP network were excluded because either the plots did not include silver birch or at least three measurements taken at five-year intervals were not available. We used the first five-year interval, i.e., the period between the first and second measurements, for the evaluation of growth and thinning rates, and the second five-year interval, i.e., the period between the second and third measurements, for the mortality study. During the first period, trees were harvested within 93 plots with different thinning intensity ranging from 0.1 to 80.0% of the total number of trees (0.3-66% of the basal area). Seventy-four plots were discarded from the analysis because thinning was practiced within the

**Fig. 1** – (a) The distribution of study plots throughout Estonia, and (b) the proportion of three major tree species within the study plots.
plots in the second interval, or before the first interval. Overall, 61,685 recorded tree measurements were considered in 348 selected plots. The species composition of the plots was 25.6% Scots pine (Pinus sylvestris L.), 33.9% silver birch, 26.8% Norway spruce (Picea abies L.) and 13.7% of several other species (e.g., Alnus glutinosa L., Alnus incana L. and Populus tremula L.). Fig. 1 shows the distribution of the selected study plots throughout Estonia (a) and the proportion of the three major tree species within those plots (b).

Predictors of tree mortality

Based on the concept of the influence-zone (Staebler 1951) we assumed an imaginary circle in which the center is defined by a tree, and its radius is 40% of the average height of trees in the first storey on each plot (Sims et al. 2009, Maleki et al. 2015). In order to avoid biased estimations due to the interference from immediate non-measured neighboring trees outside the plot boundary, we established a boundary strip (buffer zone) inside the monitoring plot with a width equal to the radius of the influence zone. The mortality study was performed only for trees inside the reduced plot (reference trees) for which the neighbors’ information was available. The distribution of plots by number of reference trees, and the plots with an average number of neighbors within the reference trees’ influence zones by the radius of the influence zone are presented in Fig. 2a and Fig. 2b, respectively.

A list of selected variables studied for the contribution in tree death is provided in Tab. S1 of the Supplementary Material. As we here aimed to investigate the suitability of neighborhood spatial information to predict tree mortality, the variables were classified into spatial (where the neighboring trees’ co-ordinates within the influence zones were required) and non-spatial (where the neighboring trees’ co-ordinates were not used).

Non-spatial variables are simple functions of stand or tree level measurements. In Tab. S1 of the Supplementary Material the most common stand variables like density (N&lt;sub&gt;0&lt;/sub&gt;), basal area (G, m²/ha), site index (SI&lt;sub&gt;100&lt;/sub&gt; m) and stand age (age, year), were calculated from plot data. Variables N&lt;sub&gt;0&lt;/sub&gt; and G can be treated as measures of stand density (Eid & Tuhus 2001, Burgman et al. 1994), while G can be considered as a measurement of symmetric competition (Bravo et al. 2001). The site index SI&lt;sub&gt;100&lt;/sub&gt; was used to measure the site productivity, calculated as the average height of a stand at the reference age of 100 years (Nilson 1999).

For non-spatial tree level measurements the following variables were used: tree diameter at breast height (dth, cm), relative tree diameter (d&lt;sub&gt;th&lt;/sub&gt;) calculated as the ratio of tree and stand diameters, tree five-year diameter growth (id&lt;sub&gt;th&lt;/sub&gt; cm), tree basal area (g, m²), and the sum of the trees’ basal areas (per plot) larger than the reference tree (BAL, m² ha⁻¹·Wykoff et al. 1983). The tree diameter integrates the past competitive interactions of the reference trees (Soares & Tomé 1999), while d&lt;sub&gt;th&lt;/sub&gt; shows the dominance of a reference tree related to other trees in the stand. The tree diameter growth rate (id&lt;sub&gt;th&lt;/sub&gt;) is a measure of tree vitality often used in mortality models (Hamilton 1986, Das et al. 2007). Fig. 2c shows the relationship between the diameter growth and the relative diameter in the reference tree data set. Trees with negative diameter increment were not excluded from the data set in order to avoid bias toward positive growth, because the occurrence of undetectable overestimation of growth is unavoidable. The value of BAL can be used as a measure of asymmetric competition (Monserud & Sterba 1999, Yang et al. 2003). When considering the influence of the neighborhood, the alternative estimators of the relative diameter and the sum of the basal area of larger trees (d&lt;sub&gt;th&lt;/sub&gt; and BAL) were calculated only for trees within the influence zone around each reference tree.

Spatial variables in Tab. S1 of the Supplementary Material consisted of the well-known Hegyi (1974) competition index (CI) as a measure of neighbouring trees’ competition, an aggregation index (agg - Clark & Evans 1954) to measure clumping and the different measures of species proportion (sp). Aggregation illustrates the spatial pattern of neighboring trees within the influence zone, where values < 1.0 indicate an aggregated distribution of trees, and values > 1.0 indicate a tendency toward a regular distribution. The species proportion defines the degree of spatial segregation of tree species in the stand. In this study the different values of sp were calculated as follows: (i) the proportion of silver birch trees (spb); (ii) the proportion of other species that are not silver birch (sp); (iii) the proportion of Norway spruce trees (spns); and (iv) the proportion of Scots pine and other deciduous trees (spn), for all neighboring trees within the influence zone.

An important issue in stand development is the self-thinning, when an increased density-dependent mortality rate is expected due to high competition among individuals. Traditionally, the allometric relationship between stand density (N&lt;sub&gt;0&lt;/sub&gt;) and diameter (D) has been applied for approximating the self-thinning line (Burkhart & Tomé 2012). However, Nilson (2005, 2006) showed the advantage of using stand sparsity (L, m) by a transformation of the stand density (N&lt;sub&gt;0&lt;/sub&gt; - eqn. 1).
Model fitting and statistical analyses

Mortality is a discrete event that can only take two values (alive or dead), hence logistic functions are widely applied to model probability of tree mortality (Monserud 1976, Yao et al. 2001, Bravo-Oviedo et al. 2006). Since data from many plots were used and the number of reference trees were not equal for studied plots, we devised the mortality probability in a generalised linear mixed effects logistic regression model for that part of predictor variables, del intercept, 3): $p = \frac{1}{1 + e^{-(a + bX + yZ + \epsilon)}}$

Assuming that $v$ is the number of the predictor variables, $N$ is the number of observations, and $n$ is the number of plots, then $\alpha$ is the model intercept, $X$ is a $N \times v$ matrix of the $v$ predictor variables, $b$ is a $v$ column vector of the fixed effects model coefficients, $Z$ is the $N \times n$ design matrix for the random effect (the random complement to the fixed $X$), $y$ is a vector of the random effect (the random complement to the fixed $b$) and $\epsilon$ is a $N$ column vector of the residuals for that part of $p$ that is not explained by the model $Xb + Zy$.

Since the coefficient of determination is not appropriate when discrete variables are modeled (Bravo-Oviedo et al. 2006), the goodness-of-fit of the models was assessed using a log-likelihood statistic (log-likelihood, Akaikes Information Criteria (AIC), and Akaikes weights (AICc). The probability that the model was the best with the lowest expected information loss was determined by the smallest value of AIC and the biggest AICc (Wagenmakers & Farrell 2004). $\Delta$AIC indicates the differences in AIC values between a given model and the model with the lowest AIC value. According to Burnham & Anderson (2002), we assumed that the models with $\Delta$AIC < 2.0 had a “substantial empirical support” and that these models were almost equal, whereas the models with $\Delta$AIC > 4.0 were considerably different. The accuracy of the models in correctly classifying the trees in dead/alive groups was also tested by the area under the receiver operating characteristic (ROC) curve (Bravo-Oviedo et al. 2006). The area under a ROC curve (AUC) quantifies the accuracy of the model, where values > 0.7 indicate good accuracy (Fawcett 2006). Also, a classification of alive and dead trees for each model was performed using the ideal cut-point, where the sensitivity (true mortality rate) and specificity (true survival rate) curves crossed (Hosmer & Lemeshow 2000). The val.prob function of the rms package in the R statistical software (R Development Core Team 2014) was used to assess the accuracy of the models, and to validate the predicted probabilities of the logistic fits against the observed mortality.

Models were developed and selected as follows. Initially, pairwise models were calculated between mortality and each variable presented in Tab. S1 (Supplementary Material). The importance of each variable in the pairwise fits was evaluated using the analysis of variances (ANOVA) and the 10 most effective variables were selected based on their AIC values. The significance of thinning variables, $l_{int}$, and $C_{int}$ on mortality probability of silver birch trees was also tested. About 400 models were fitted with all possible combinations of these selected variables with the restriction that no model could simultaneously include variables representing a similar factor (e.g., one model could only have either CI or BAL as a measure of competition). Models were then compared using AIC values, and the best models, as well as the models with an AICc value greater than zero, were selected. Finally, the contribution of thinning to the mortality of individual trees was assessed. To this purpose, variables $l_{int}$ and $C_{int}$ were separately added to the selected model as new predictors, and the changes to the quality measure of the full statistical model were investigated. For all models, the glm function from the lme4 package in the R statistical software was applied. Furthermore, for each combination of variables the variance inflation factor (VIF) was determined to ensure that the models were not strongly biased by multicollinear-ity.

Results

Over the 348 selected research plots the five-year mortality rate of silver birch trees was 9.67%, i.e., out of 6009 silver birch reference trees 581 died. The mortality rate in pure stands was higher than in mixed stands (Tab. 1). There was a stronger survival probability for a silver birch tree when growing in a neighborhood that consisted of species other than birch, as the mortality rate decreased from 16.67% in a pure silver birch stand to 6.09% for single silver birch trees surrounded by other species, mainly Norway spruce and Scots pine.

The results from the pairwise analyses indicated that tree growth (dbh), tree size (dbh), as well as competition (sensu CL and sBu.b), structure (agg and sp), and stand size (CI), were the most important predictors of individual tree mortality (Fig. 3). The mortality probability of individual trees was negatively correlated with dbh, id3, sp, and agg, but the correlation was positive for competition and stand size. The other studied variables (see Tab. S1 in the Supplementary Material) showed non-significant effects in the current mortality study. In an attempt to take the site quality into consideration, the site index at reference age of 50, and the site type as a nominal variable, as alternatives to $S_{site}$ were also tested, but they did not provide any improvement to the model performance.

The highest ranked logistic models with different combinations of variables are presented in Tab. 2 in two categories: some models included only non-spatial mortality predictors, and both spatial and non-spatial variables were fitted into other models. The models consisting of spatial variables far exceeded those with non-spatial measurements in terms of AIC and AICc. The VIF values for all combinations were less than 3.0, indicating a low multicollinearity. Additionally, we tested different transformations of dbh, $d_{agg}$ and $i_{sad}$, and found then

<table>
<thead>
<tr>
<th>Tab. 1 - Mortality rate of silver birch trees depending on silver birch proportion.</th>
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<tbody>
<tr>
<td>Proportion of birth in influence zone (%)</td>
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<tr>
<td>---------------------------------</td>
</tr>
<tr>
<td>81-100</td>
</tr>
<tr>
<td>61-80</td>
</tr>
<tr>
<td>41-60</td>
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<tr>
<td>21-40</td>
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<td>0-20</td>
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inferior to the untransformed variables (data not shown).

Fig. 4 shows a comparison of the predicted mortality probability of reference trees with their observed mortality probability for the best logistic model. The predictive performance of the best model was good; however, a closer analysis of the estimated mortality probability revealed an evidence of negligible overestimation of the mortality prospects of very small silver birch trees (dbh < 40 cm) with low survival probability, and also a slight underestimation of the mortality probability of the small proportion of large silver birch trees (dbh > 40 cm), see sensitivity and specificity values in
Tab. 3 - The best combinations of variables predicting silver birch mortality when thinning is included as predictor in the model used. (id): five-year diameter growth of tree (cm); (dbh): relative dbh of reference trees for each plot; (Clrel): Hegyi’s competition measure of thinned trees inside the zone of influence; (influ): thinning intensity of the plot; (agg): aggregation of trees inside the zone of influence; (sp): proportion of other species than silver birch within the influence zone. ΔAIC and AICc are the statistical measures of models. AUC, TPR and FPR are area under curve, true positive rate and false positive rate of dead trees, respectively. Coeff are coefficients of the models. TPR and TNR are the sensitivity and specificity of the models, respectively.

<table>
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<tr>
<th>Variables</th>
<th>ΔAIC</th>
<th>AICc</th>
<th>AUC</th>
<th>TPR</th>
<th>TNR</th>
<th>Intercept</th>
<th>Coeff</th>
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<th>Coeff</th>
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<tbody>
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<td>0.00</td>
<td>0.655</td>
<td>0.819</td>
<td>0.833</td>
<td>0.856</td>
<td>2.324</td>
<td>-1.977</td>
<td>-3.263</td>
<td>-1.105</td>
<td>-1.008</td>
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<tr>
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<td>0.819</td>
<td>0.828</td>
<td>0.856</td>
<td>2.327</td>
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<td>-3.232</td>
<td>-1.103</td>
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</tr>
<tr>
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<td>0.819</td>
<td>0.811</td>
<td>0.863</td>
<td>2.303</td>
<td>-1.981</td>
<td>-3.262</td>
<td>-1.098</td>
<td>-0.993</td>
</tr>
</tbody>
</table>

Tab. 3. More detailed examinations also revealed that some models had overestimated the likelihood of smaller tree survival. Moreover, a tendency to underestimate the survival likelihood of silver birch trees with small relative diameters in the unthinned plots was detected.

As shown in Tab. 3, the inclusion of the non-spatial variable Iinflu in the best model did not seem to improve its performance. However, a significant improvement was observed when the spatial thinning variable Clrel was included. Thus, thinning practices in silver birch stands demonstrated that there was a negative influence on the probability of tree mortality (see Coeff, in Tab. 3) by reducing the competition. Therefore, an excessive decrease in the number of trees due to high thinning intensity, and consequently a reduction in competition, could also decrease the tree mortality rate (from 10.1% in unthinned plots to 6.6% in thinned plots). Additionally, the accuracy estimated by AUC was excellent (AUC>0.8) for all the models in Tab. 3 (Hosmer & Lemeshow 2000). These high values indicated that many of the dead trees (true positives) were correctly classified.

Discussion

Drivers of silver birch mortality

In this study, the silver birch trees with an increasing diameter growth rate clearly indicated an increment in survival. Recent radial growth has been frequently used as an indicator of tree health and vigor (Kobe 1996, Yao et al. 2001). Accordingly, it is biologically meaningful to assume an improved survival for trees showing higher growth rates, while the opposite is expected for trees showing feeble growth. As a consequence, the probability of any tree to survive can be inferred from its diameter increment (Waring 1987). On the other hand, since radial growth rate varies systematically by tree size after an initial increase, the radial growth rate often decreases as the tree diameter increases, and therefore, diameter growth rate tends to overestimate the mortality probability of bigger trees as compared with smaller trees. It has been proposed that basal area growth rate is a better measure of tree survival than diameter growth rate (Pederson 1998, Ireland et al. 2014). Basal area growth rate represents the tree mass growth more closely, as it increases with tree size (Clark & Clark 1999, Di Filippo et al. 2012). Similar to Das & Stephenson (2015), our results did not indicate any better prediction of tree mortality when basal area growth is included in the model (data not shown). On the other hand, we observed a systematic overestimation of the mortality probability of small birch trees.

While the mortality of small birch trees in young and dense stands increased due to self-thinning (Barnes et al. 1998), in older stands the high mortality of small trees can be attributed to the suppression by dominating trees—where the non-spatial explanatory variable dbh appeared superior to capture this trend. By some means, the hierarchical position of trees within the stand is measured by dbh and indirectly indicates their competitive status (Burkhart & Tomé 2012). We found that competition is a significant mortality driver, in accordance with the extensive literature corroborating such relationships in the Estonian forests (Laarmann et al. 2009, Sims et al. 2009) and in other forests (Monserud et al. 2004, Temesgen & Mitchell 2005, Bravo-Oviedo et al. 2006, Das et al. 2011). Moreover, the reversed J-shaped (L-shaped) size-dependent mortality pattern of our studied trees (Fig. 3) can be explained by the insufficient number of large trees, and the high mortality rates of young trees within the stands where the competition for resources is high (Coomes & Allen 2007, Olano et al. 2009, Dietze & Moorcroft 2011).

The competition-induced mortality is presumed to decline as local tree density decreases, and the overall plot density showed to be related to the strength of competitive effects (Gonzalez et al. 2004, Temesgen & Mitchell 2005). The variable dbh could have served as a simple index of competition, but it did not take the variation of stand density into account. Therefore, equipping the model with spatial measures that represented the neighborhood properties of reference trees did appear unavoidable. Surprisingly, contrary to our assumption, adding the new calculations of BAL (BALdbh) and dbh(BALdbh) only for immediate neighbors inside the influence zone diminished the predictive power of the model. This can be explained by the expected increasing uniformity in tree spacing with tree age or size, due to mortality from competition (Jose et al. 1991, Moer 1997, Druczenbrod et al. 2005). Therefore, for some small birch trees in old and regular stands, and also for some big birch trees in young and clumped stands, the real values of competition and relative diameter may not be measured by considering the influence zone (here as a ratio of stand height). Finally, the inclusion into the model of the aggregation measure (agg) could take into consideration the horizontal spatial arrangement of tree positions inside the influence zone. The aggregation index has been successfully used to assess the regularity of tree positions (Pommerning 2002, Aguirre et al. 2003, Brumels et al. 2005). In our study, agg efficiently proved the negative influence of clumping (agg<0) on the survival prospects of birch trees (see Tab. 3, negative coefficients for agg in the mortality models).

Several studies reported that the competition and growth of silver birch trees, and consequently their survival prospects, are strongly related to the identity of neighboring species (Kattami & Lintunen 2010, Jõgiste et al. 2010, Hynynen et al. 2011). Their findings are consistent with our results, in that including sp to the best non-spatial mortality model definitely improved its fit. The mortality likelihood of a birch tree was negatively related to the species mixture of its immediate neighbors (decreasing from pure stands to mixed stands - Tab. 2, Tab. 3). According to the Janzen-Connell’s type effect, this may be caused either by the increased risk of attack by species-specific herbivores or diseases, when a silver birch tree grows in close proximity to many other con-specifics (Janzenn 1970, Connell 1971). Also, the inter-specific competition among hetero-specific individuals (mainly Norway spruce and Scots pine) appeared lower than the competition among con-specific individuals (i.e., neighboring silver birch trees). Due to a similar demand for light and resources, the competition among neighboring silver birch individuals increases, and therefore the competition-induced mortality is relatively high. Norway spruce and birch trees have similar site requirements, but when occurring together the formers seem more shade-tolerant than the light-demanding birches, and also show different temporal growth patterns (Tahvanainen & Fors 2008, Hynynen et al. 2011). Moreover, birch trees easily overwhelm and suppress the
shade-intolerant Scots pine trees due to their vigorous early growth (Hyunen et al. 2017); however, their co-occurrence is relatively infrequent due to different site requirements.

The effect of thinning on silver birch mortality

Thinning to different residual densities produces varying effects on tree mortality, depending on the tree species with different characteristics and site conditions (He & Duncan 2000, Canham et al. 2001, Uriarte et al. 2004). Some studies have found higher mortality risk during the first years after thinning, either due to mechanical damage to the remaining trees (Nyland 1994, Caspersen 2006), or by shifting the population towards smaller trees more prone to environmental stresses, or even to windfall for larger residual trees on the margin of cutting areas (Jónsson et al. 2007, Fortin et al. 2008). In contrast, some studies have reported that thinning operations may rescue trees from density-dependent mortality (Powers et al. 2010). In our case, the mortality of silver birch trees was affected negatively by thinning operations within the plots (up to 3.5% decline in mortality rate of birch trees in thinned stands). The slight improvement of performance of the mortality model obtained by including CI indicated that the competition imposed by neighboring trees, which were cut during the thinning operation, partially limited the survival of birch trees. As explained before, density-dependent mortality or self-thinning (Drew & Flewelling 1977) occurs when the stand sparsity is smaller than the limiting sparsity. Thinning operations have successfully increased the stand sparsity, and released the shade-intolerant birch trees that showed a vigorous growth within a less competitive environment.

Data limitation

The areas included in the Estonian Network of Forest Research Plots (ENFRP) were mostly located in managed forests, i.e., only a few old dense stands were measured in this study. The record of the age of individual trees was not available and the age was only determined for tree cohorts. Since regular mortality of a tree could be dependent on its age (Monserrud 1976), including the exact age of trees may have improved the model’s strength. However, determining the age of individual trees in practical forest management is too laborious. Another factor of uncertainty is the inclusion of small plots when applying spatial variables. As mentioned above, a buffer zone inside the plot (near the plot boundary) was applied to avoid the biased estimations due to edge effects. Consequently, a relatively large number of trees were excluded from mortality analyses, and this loss was more relevant for small plots. However, for the simulation of the neighborhood, a border method (Ripley 1981) was used as a means to eliminate this effect, as described by Lilleleht et al. (2014). This method is simple to implement in comparison with other methods, and guarantees the removal of all edge effects (Kint et al. 2004).

Additionally, one might expect that building the mortality model to include soil descriptors and climate variables will also improve its performances. In this study, mortality was assessed over periods of five years as the exact year of tree death was unknown. Thus, it was difficult to connect extreme events, such as very cold temperatures or hot summers, which can usually cause high tree mortality. Regarding the costly assessment of annual tree mortality and soil analysis around individual trees for large data sets, a dendrochronological analysis seems to be a more effective method for climate studies (Linares et al. 2009). Further studies are needed to investigate the efficiency of the mentioned variables and other possible mortality predictors in defining the live/death status of individuals.

Conclusions

Natural mortality of individual trees is a stochastic and irregular phenomenon for which we attempted to find the most relevant explanatory variables. Although a number of known and unknown factors affecting tree mortality makes its modeling complicated, the fitted models used in this study did produce results that satisfactorily explained mortality of Estonian silver birch trees. Five-year diameter growth, relative diameter, species proportion, and aggregation were the most appropriate explanatory variables in our mortality models. In order to maintain high survival probabilities, forest management plans and practices should pay a special attention to growth performances of trees, species compositions, tree density and forest stand structure. Furthermore, the reduction of stand density would provide more growing space for light-demanding birch trees, as demonstrated by the reduced mortality predicted by our model when thinning was included as predictor. However, root and stem damage caused by heavy machinery used for thinning operations, and the higher wind exposure of trees after thinning should also be considered. Finally, based on our results, silver birch trees should preferably be managed in mixed stands where they occur along with other tree species, and attempts should be made to minimize the clumping (agy>1) within their neighborhoods.

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