Needle longevity, shoot growth and branching frequency in relation to site fertility and within-canopy light conditions in *Pinus sylvestris*

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Abstract – Changes in needle morphology, average needle age, shoot length growth, and branching frequency in response to seasonal average integrated quantum flux density ($Q_{\text{int}}$) were investigated in *Pinus sylvestris* L. in a fertile site (old-field) and an infertile site (raised bog). In the fertile site, the trees were 30 years old with a dominant height of 17–21 m, and with average ± SD nitrogen content (% of dry mass) of 1.53 ± 0.11 in the current-year needles. In the infertile site, 50 to 100-yr-old trees were 1–2 m tall, and needle N content was 0.86 ± 0.12%. Relationships between the variables were studied using linear correlation and regression analyses. With increasing irradiance, shoot length ($L_s$) and shoot bifurcation ratio ($R_b$, the number of current-year shoots per number of shoots formed in the previous year) increased in the fertile site. The shoot length distributions became more peaked (positive kurtosis) and biased towards lower values of $L_s$ (positive skewness) with increasing $Q_{\text{int}}$ in this stand. The shoot distributions were essentially normal in the infertile site. Large values of $R_b$ combined with the skewed distributions of shoot length resulted in conical crowns in the fertile site. In contrast, lower bifurcation ratio, normal shoot length distributions and low rates of shoot length growth led to flat-topped crowns in the bog. Average needle age was independent of $Q_{\text{int}}$, but was larger in the infertile site. Thus, reduced rates of foliage production in the infertile site were somewhat compensated for by increased foliage longevity, and we suggest that shoot growth rates may have directly controlled the needle life span via reduced requirements for nutrients for the growth and via reduced self-shading within the canopy. Needle age and $Q_{\text{int}}$ independently affected needle structure. Needle age only moderately altered needle nutrient contents, but the primary age-related modification was the scaling of needle density with age. The density was similarly modified by age in both sites, but the needles were denser in the infertile site. Given that denser needles are more resistant to mechanical injury, larger density may provide an additional explanation for enhanced longevity in the infertile site. Our study demonstrates that site fertility is an important determinant of the plastic modifications in crown geometry, and needle life span in *P. sylvestris*.

bifurcation ratio / branching / irradiance / leaf life span / leaf density / site fertility

Résumé – Longévité des aiguilles, croissance des pousses et fréquence de ramification en relation avec la fertilité du site et les conditions de lumière dans la canopée de *Pinus sylvestris*. Les changements dans la morphologie des aiguilles, l’âge moyen des aiguilles, la croissance en longueur des pousses, la fréquence de la ramification ont été étudiés en réponse à la densité du flux quantique intégré ($Q_{\text{int}}$) moyen saisonnier chez *Pinus sylvestris* L. dans un site fertile (anciennement cultivé) et dans un site pauvre (tourbière). Dans le site fertile, les arbres étaient âgés de 30 ans, avec une hauteur dominante de 17–21 m, et une teneur en azote (g kg$^{-1}$ de matière sèche) moyenne de 15,3 ± 1,1 dans les aiguilles de l’année. Dans le site pauvre, les arbres, âgés de 50 à 100 ans, avaient une taille de 1 à 2 m, la teneur en azote des aiguilles était de 8,6 ± 1,2 g kg$^{-1}$. Les relations entre les variables ont été étudiées en utilisant les analyses de corrélation linéaire et de régression. Lorsque l’irradiation est croissante, la longueur de la pousse ($L_s$) et le rapport de ramification ($R_b$, nombre de pousses de l’année par nombre de pousse formées au cours de l’année précédente) augmentent dans le site fertile, mais pas dans le site pauvre. Malgré une fréquence plus élevée de ramification, le contrôle apical est exerçée par une irradiation plus élevée dans le site fertile. Les distributions des longueurs de pousses deviennent plus pointues (kurtosis positive) et biasées vers les valeurs les plus faibles de $L_s$ (skewness positive) avec un $Q_{\text{int}}$ en augmentation dans ce site. Les fortes valeurs de $R_b$, combinées avec des distributions skewness des longueurs de pousses conduisent à des canopies coniques dans le site fertile. Par opposition, un rapport plus faible de la ramification, distributions normales des longueurs de pousses, et une faible croissance en longueur des pousses conduisent à la formation de canopées aplatis dans la tourbière. L’âge moyen des aiguilles était indépendant du $Q_{\text{int}}$, mais il était plus élevé dans le site le plus pauvre. Cependant, les taux réduits de production foliaire dans la station pauvre étaient, en quelque sorte, compensés par l’accroissement de longévité du feuillage, et nous suggérons que les taux de croissance des pousses peuvent avoir contrôlé directement la durée de vie des aiguilles par une réduction des besoins en nutriments pour la croissance et par une réduction de l’ombre dans la canopée. L’âge des aiguilles et $Q_{\text{int}}$ affectent indépendamment la structure des aiguilles. L’âge des aiguilles modifie seulement modérément la teneur en nutriments des aiguilles, mais la modification primaire liée à l’âge, était l’échelle de densité d’aiguilles. La densité était pareillement modifiée par l’âge dans les deux stations, mais les aiguilles étaient plus denses dans le site pauvre. Étant donné que des aiguilles plus denses sont plus résistantes aux blessures mécaniques, une plus grande densité peut fournir une explication additionnelle à la longévité renforcée dans les stations pauvres. Notre étude montre que la fertilité de la station est un important déterminant des modifications plastiques de la géométrie de la couronne et la durée de vie des aiguilles chez *P. sylvestris*.

rapport de bifurcation / ramification / irradiance / durée de vie de la feuille / densité de feuille / fertilité de la station

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1. INTRODUCTION

Crown architectural characteristics control the light harvesting efficiency of the canopy and species competitive potential [40, 64, 78, 84]. Differences in branching angle, branch length, and frequency of branching modify the aggregation of the foliage on the branches [19, 20, 40, 78], and thereby change the degree of self-shading within the canopy. Because the requirements for efficient light usage and acquisition vary with incident quantum flux density [64], a specific canopy constitution is not appropriate for all natural light levels. As the result of evolutionary adaptations in crown architecture to incident irradiance, there exists an array of various crown morphologies, and genetic heterogeneity in crown geometry provides a major explanation for species separation along gap-understory gradients [40, 84].

The species also possess considerable phenotypic plasticity for modification of canopy architecture, and thus, the foliar exposition characteristics [64]. Understory individuals of many plant species have flat crowns with the foliage arranged in a few planar layers to minimize self-shading within the canopy [11, 38, 39, 74, 77]. In contrast, plants in open habitats have conical crowns with a large number of leaf layers [3, 6, 39, 60, 74, 88] that have a greater within canopy shading, but larger photosynthesizing leaf area. Such important alterations in crown shape are the consequence of light-related adaptability in branching frequency, branch length and branching angles [11, 15, 41, 65, 76, 77, 81]. Thus, understanding the environmental modifications in these characteristics is of paramount significance to characterize tree crown growth and light interception capacity [21, 36].

Apart from light, all environmental and soil variables that modify growth and development may potentially have important influences on canopy geometry, but much less is known of canopy morphological responses to these external factors [84]. There is evidence that, in conifers, branchiness may increase with decreasing site water availability [5]. In addition, increases in soil nutrient availability generally lead to enhanced branch extension growth [47, 67], as well as higher fractional biomass investment in foliage [59], and greater total plant foliar area [47, 70, 73]. The branching responses to nutrient availability have not been investigated extensively in trees, and it is not clear whether the nutrient-related increase in branch extension is sufficient to support the extra foliar area, or whether the improved nutrition also leads to greater shoot production and more frequent branching. However, enhanced branching in higher nutrient availability is likely, because increases in branch length only, lead to larger biomass costs for mechanical support of branches [20, 46]. In herbaceous species, there is evidence of more frequent branching at higher nutrient availabilities [73], but the potential effects of nutrient limitations on plastic changes of crown architecture to light availability have not been characterized.

Adjustments in needle longevity also influence the total foliar area on the tree, and thereby the self-shading within the canopy. There is phenomenological evidence that decreases in light [37, 39, 45, 72] or nutrient availability [66] may result in increases in average needle life span, but the mechanisms responsible for extended needle longevity are still not entirely understood. Despite the lack of knowledge at the mechanistic level, such increases in needle longevity are relevant, and may largely compensate for the limited new foliage production in plants growing in shortage of light and/or nutrients. Moreover, limited shoot growth may directly lead to greater needle life span because of reduced self-shading within the canopy [1]. Thus, changes in crown architecture and in needle longevity may be closely interrelated.

We studied relationships of shoot growth, branching frequency and average needle age versus long-term integrated average quantum flux density in infertile and fertile sites in temperate conifer species *Pinus sylvestris* L. This species colonizes a wide range of early-successional habitats with strongly varying soil water and nutrient availabilities [42, 58], and is apparently a very plastic species that may readily change the crown architectural variables [36] and biomass allocation [33, 34] in response to changes in light availability. The primary objective of our study was to determine whether both the light and nutrient availabilities alter canopy architecture and needle life span, and whether the effects are interactive or independent. Although *P. sylvestris* is a plastic species, we have previously demonstrated that its ability for needle physiological and morphological [55] and shoot architectural [54] acclimation to light availability is considerably lower in the low than in the high fertility site. Thus, we expected similar differences in the plasticity also in canopy architecture. The conifers strongly reduce foliar area in response to decreases in soil nutrient availability [2, 42, 86], and it is logical to assume that the investments in woody support framework also parallel the major changes in needle area. As the characteristics of canopy architecture, we study average shoot lengths, shoot length distributions and branching frequency, which collectively allow quantitative estimation of conifer crown development [36].

To gain mechanistic insight into the variability in needle longevity between and within the sites, we also studied foliage structure, and needle nitrogen and phosphorus contents in needles of various age. Given that light and nutrient availabilities may independently modify needle morphological variables in *P. sylvestris* [55], and that these characteristics may directly alter leaf life span by altering the sensitivity of the foliage to mechanical damage [51], we hypothesized that light availability and site fertility have independent effects on needle longevity as well, and that these effects are related to site-to-site differences in needle morphological characteristics.

2. MATERIALS AND METHODS

2.1. Study sites

A monospecific even-aged homogeneous *Pinus sylvestris* plantation (1400 trees ha\(^{-1}\), 29–31 years old, dominant height 17–21 m) on an old field at Ahumadalu, Estonia (58°19’ N, 27°17’ E, elevation ca. 60 m above sea level) was chosen as a representative nutrient-rich habitat. The soil was a pseudogley with moderately acidic (pH in 1 M KCl of 4.3) humus horizon ([55] for specific details). In the understory, the dominants were the shrub *Rubus idaeus* L. and the herbaceous species *Epilobium angustifolium* L., *Impatiens parviflora* DC. and *Urtica dioica* L., which are indicators of nitrogen-rich early-successional habitats [16].
Although needle morphological characteristics and nutrient contents transported to the laboratory within an hour from collection, the branches were enclosed in plastic bags, and tree. In the infertile stand, 2–4 branches per tree were sampled. After collection, the branches were enclosed in plastic bags, and transported to the laboratory within an hour from collection. Although needle morphological characteristics and nutrient contents transported to the laboratory within an hour from collection, the branches were enclosed in plastic bags, and tree. In the infertile stand, 2–4 branches per tree were sampled. After collection, the branches were enclosed in plastic bags, and transported to the laboratory within an hour from collection. Although needle morphological characteristics and nutrient contents may potentially vary during the season [30, 43], such effects were not evident in our data [55].

The nutrient-limited site was a scattered woodland (200 trees ha−1) dominated by P. sylvestris and Betula pubescens Ehrh. at Männikjärve raised bog, Endla State Nature Reserve, Estonia (58° 52' N, 26° 13' E) on thick – up to 8 m in the centre of the bog – Sphagnum peat [85]. The average height of ca. 50–100 year-old trees was only 1–2 m. The organic soil was strongly acidic throughout the entire profile (pHKCl = 2.59). Eriophorum vaginatum L., Rhynchospora alba (L.) Vahl and Scheuchzeria palustris L. dominated the herb layer, and Calluna vulgaris (L.) Hull, Chamaedaphne calyculata (L.) Moench, Empetrum nigrum L. and Ledum palustre L. the dwarf-shrub layer. A thorough description of this site is given in Niinemets et al. [55]. According to the previous study, the plants were limited both by low P and N availabilities in this site [55].

2.2. Foliage sampling and long-term light availability estimations

Because the fertile site was very homogeneous, three 19–20 m tall trees in the centre of the forest were selected for detailed sampling. In the infertile site, 22 trees with heights ranging from 0.8 to 2 m were selected in the central areas of the bog. In addition, seven larger trees (height 2.9–8.7 m) with apparently better nutrition were chosen at the edge of the bog and on the adjacent dried peatlands to attain a larger gradient in nutrient availability [29]. The trees sampled in this site were 20–150 years old according to the increment cores taken at the ground level (average ± SE = 43 ± 8 yr.). Only mature, reproductive phase trees were considered, and we did not observe any significant effect of tree age on studied crown and foliage characteristics (P > 0.05). Insignificant effects of tree age on foliage structure and branching are in agreement with previous observations in mature trees [49, 52]. In fact, tree-to-tree differences in height were primarily associated with differences in tree nutrient status (figure 1). Both the N (figure 1A) and P (figure 1B) contents of the uppermost unshaded needles were positively correlated with tree height for both sites pooled, and also for the infertile site considered separately. This suggests that although there were site differences in average tree age, comparisons of foliage and crown characteristics between the sites are valid.

The sampling was conducted in Sept. 1998 in both sites, and in Oct. 1999 in the fertile site, and late Aug. 1999 in the infertile site. Entire branches (n = 68) were harvested along the light gradient in tree canopies. In the fertile site, 4–5 branches were taken from each tree. In the infertile stand, 2–4 branches per tree were sampled. After collection, the branches were enclosed in plastic bags, and transported to the laboratory within an hour from collection.

Figure 1. Correlation of the sampled tree height with the nitrogen (A) and phosphorus (B) contents of uppermost unshaded foliage (integrated quantum flux density Qint > 30 mol m−2 d−1). The linear regressions were fitted to the entire set of data (dashed lines, filled symbols correspond to the fertile, and open symbols to the infertile site), and separately to the infertile habitat (solid lines).

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where statistical comparison of these sample shoots demonstrated that \[8, 41, 81, 83\], the frequency of branching is often described by the bifurcation ratio \(M\). Given that needle dry mass per unit area, \(A_t\), was computed as the product of needle circumference (\(C\)) and \(L_n\) approximating the needle cross-section geometry by half-ellipse [55]. The projected needle area, \(A_p\), was computed as \(W_n L_n\). The sample needles were weighted after oven-drying at 70 °C for at least 48 h, and needle dry mass per unit total (\(M_A\), g m\(^{-2}\)) and projected area (\(M_P\)) were calculated. The assumption of half-elliptical needle cross-section geometry was also employed to find needle volume (\(V\), [55]) and the \(V/A_t\) ratio (mm). Given that needle dry mass per unit area, \(M_A\), is a product of \(V/A_t\) and needle density [50], needle density (\(D\), g cm\(^{-3}\)) was computed as \(M_A/(V/A_t)\). All shoots in each age-class were dried at 70 °C, separated between needle and woody biomass, and weighted. Shoot dry matter content (\(\delta_n\)) was further calculated as the weighted average of needle and shoot axis dry to fresh mass ratios. For 26 shoots, needle and stem fresh masses were determined separately, allowing to compute needle (\(\delta_n\)) and shoot axis (\(\delta_s\)) dry matter contents. The statistical comparison of these sample shoots demonstrated that \(\delta_n\) was significantly larger than \(\delta_s\) (\(P < 0.05\) according to a t-test), but also that the differences were minor (average \(\pm SE = 0.612 \pm 0.020\) g g\(^{-1}\) for \(\delta_n\) and 0.600 \(\pm\) 0.032 g g\(^{-1}\) for \(\delta_s\)).

2.4. Calculation of shoot bifurcation ratio

Assuming that branching in plants follows a geometric sequence, the frequency of branching is often described by the bifurcation ratio [8, 41, 81, 83], \(R_b\):

\[
R_b = \frac{N_a}{N_{a+1}},
\]

where \(N_a\) is the number of branches of age \(a\) and \(N_{a+1}\) is the number of branches in the next older age-class [61, 87]. In a more general form:

\[
N_a = N_0 R_b^{a-1},
\]

where \(N_0\) is the number of shoots in the youngest age class (\(a = 1\)). Logarithming equation (3) allows to linearize the relationship, and thus, we calculated the average bifurcation ratio from the slope of \(\log N_a\) vs. \(a\):

\[
\log N_a = \log(N_0 R_b) - a \log R_b.
\]

Only branches with a minimum of four shoot age classes present were used for the analysis, and the maximum number of shoot age classes available was 15. Equation (3) gave good fits to the data (figure 2) with the fractions of explained variance (\(r^2\)) generally exceeding 0.90. This indicates that the concept of bifurcation ratio is valid for Pinus sylvestris, and also that the value of \(R_b\) was almost constant throughout the life span of the branches. Thus, \(R_b\) may be used as an estimate of long-term trends in crown architectural development in this species.

2.5. Determination of average needle age

Dry mass-averaged needle age (\(\Lambda\)) was computed for each branch as:

\[
\Lambda = \frac{\sum_{i=1}^{n} \Lambda_i M_i}{M_T},
\]

where \(i\) is the number of specific needle age class of age \(\Lambda_i\), \(M_i\) is the dry mass of all needles in this age-class, \(n\) is the number of needle age-classes present and \(M_T\) is the total needle dry mass on the branch.

Current-year needles were assigned an age of 1.0 yr. in these calculations. It is important that the average needle age for a specific branch depends not only on needle longevity, but also on shoot bifurcation ratio. For a common needle life-span, more frequent branching leads to a greater fraction of needles present in younger needle age classes than in the case of less frequent branching.

2.6. Measurement of needle carbon, nitrogen and phosphorus contents

Total needle nitrogen and carbon contents were estimated by an elemental analyser (CHN-O-Rapid, Foss Heraeus GmbH, Hanau, Germany), and phosphorus contents by inductively coupled plasma emission spectroscopy (Integra XMP, GBC Scientific Instruments, Melbourne, Australia). In some cases, standard Kjeldahl digestion was applied, and N content was estimated by indophenol method and P content by molybdenum blue method [28]. All methods gave essentially identical estimates of the contents of chemical elements [55].

2.7. Statistical analysis of data

To analyse the relationships among foliage nutrient content, shoot irradiance, needle age, shoot branching and needle architecture, linear correlation and regression techniques were employed [71]. All statistical effects were considered significant at \(P < 0.05\). Given that the characteristics of shoot length distribution, shoot length as well as the bifurcation ratios of the uppermost shoots in the tree crown differed considerably from the rest of the data, we also examined the leverage statistics (\(h\)) and studentized residuals to determine whether these cases influenced the regression models more than others [4]. The values of leverage statistic, which vary from 0.0 (no effect on the model) to 1.0 (completely determining the model), were always less than 0.25, suggesting that these data did not bias the regressions considerably. This conclusion was further corroborated by the finding that removal of the uppermost data points did not change the conclusions with respect to the statistical significance of the relations (figures 3–5).
If $Q_{\text{int}}$ was a significant determinant of a specific dependent variable, $Y_i$, site differences (Site, fixed effect) were separated by analyses of covariance:

$$Y_i = \mu + Q_{\text{int}} + \text{Site} + Q_{\text{int}} \times \text{Site} + \varepsilon,$$

where $\mu$ is the overall mean of the dependent variable and $\varepsilon$ is the error variance. If the interaction term, $Q_{\text{int}} \times \text{Site}$, was not significant ($P > 0.05$) the separate slope model (Eq. (6)) was followed by the common slope ANCOVA model to test for the intercept differences. One-way analysis of variance was employed if $Q_{\text{int}}$ was not a significant determinant of the dependent variable. The comparisons were conducted with and without the potentially influential upper canopy values of the fertile site. However, the observed differences were not sensitive to these data, indicating that the relationships were robust.

Tree crowns are composed of modular units [68], and there is a growing consensus that these moduli – branches – function essentially autonomously [32, 69, 74, 75]. Therefore, branch rather than tree was the experimental unit in the current study. However, branches on the same tree share a common pathway for nutrient, water and assimilate transport, and the repeated measurements conducted within a tree may confound the true statistical effect of irradiance and needle nutrient contents on shoot growth and branching morphology. We tested the possible tree effect ($T$) within each site by the following model:

$$Y_i = \mu + X_i + T + \varepsilon,$$

where $X_i$ is the independent variable ($Q_{\text{int}}$ or leaf N or P content). The statistical significance of the effect of the independent variable on $Y_i$ was always the same whether or whether not $T$ was included. Thus, these analyses demonstrated that the reported statistical effects were not attributable to the repeated measurements within the trees, further supporting the autonomy of branches within the tree.

As a second way to test for the possible effect of repeated measurements, we also computed the average values of all variables for each tree. Again, the statistical significance of all relationships was qualitatively the same for this and for the entire dataset as reported in the Results.

Overall, all the information was available for 14 branches from the fertile site and for 54 branches from the infertile site. The bias towards the infertile site reflects the circumstance that previous investigations have primarily studied $P. \text{sylvestris}$ characteristics in relation to light environment in nutrient rich sites (e.g., [33, 35, 36]). Due to the constraints applied for shoot length distributions and for bifurcation ratio calculations, the number of data points was reduced for these characteristics.

3. RESULTS

3.1. Shoot length and shoot length distributions in relation to light and site fertility

Average length of current-year shoots ($L_s$) increased with increasing needle nitrogen content per mass in both sites (figure 3A, $r^2 = 0.33$, $P < 0.02$ for the correlation with the average values per tree in the infertile site). However, $L_s$ was positively correlated with needle phosphorus content per mass ($r^2 = 0.40$, $P < 0.02$) and integrated quantum flux density ($Q_{\text{int}}$, figure 3B) only in the fertile site, but not in the infertile site. Because the average lengths of different age-classes were strongly ($r^2 > 0.80$) correlated, the relationships were similar with shoot lengths of other shoot age classes.

Shoot lengths were similar in low irradiance at the fertile and infertile sites (figure 3B), but the values of $L_s$ were lower in high light at the infertile habitat, indicating a lower plasticity with respect to growth adjustment to light in this site. According to one-way ANCOVA (site as the categorical variable, $Q_{\text{int}}$ as the covariate), both the site, and site $X Q_{\text{int}}$ interaction were significant determinants of $L_s$ ($P < 0.001$).

Nitrogen and phosphorus contents per unit dry mass were independent of $Q_{\text{int}}$ at the infertile site ($r^2 = 0.05$, $P > 0.2$ for N, and $r^2 = 0.00$, $P > 0.8$ for P), but strong positive dependencies were observed at the fertile site ($r^2 = 0.66$, $P < 0.001$ for N and $r^2 = 0.36$, $P < 0.05$ for P, see also [55]) complicating the correlations between light, nutrients and shoot characteristics. Nevertheless, when the interrelations between N, P and light availability were accounted for by a multiple linear regression analysis, only $Q_{\text{int}}$ was a significant determinant of most of foliar characteristics at the fertile site.

Kurtosis and skewness of the $L_s$ distributions were positively correlated ($r^2 = 0.71$, $P < 0.001$ for the fertile, and $r^2 = 0.44$, $P < 0.001$ for the infertile site). Kurtosis increased with increasing irradiance (figure 4A) at the fertile site, indicating
that shoot distributions became more peaked at higher irradiance. Similarly, the skewness scaled positively with irradiance in the fertile site (figure 4B), suggesting that there were less long shoots at high irradiance than expected on the basis of normal distribution. Thus, the apical dominance increased with increasing irradiance in this site. Skewness and kurtosis were independent of irradiance (figure 4A, B) and N content at the infertile site (for the average values per tree, \( r^2 = 0.11, P > 0.3 \) for the skewness, and \( r^2 = 0.17, P > 0.2 \) for the kurtosis). Analyses of covariance demonstrated that the slopes of the kurtosis vs. \( Q_{\text{int}} \) and skewness, vs. \( Q_{\text{int}} \) relationships were significantly lower at the infertile site (\( P < 0.01 \)). Thus, the shoot length distributions were essentially normal at the infertile site, and became increasingly asymmetric and peaked with increasing irradiance at the fertile site.

### 3.2. Effects of irradiance and nutrient availability on branching frequency and biomass partitioning within the shoot

The finding that the lengths of shoots of all age classes were strongly correlated, indicates that the growth conditions were similar throughout the branch life time, and supports the calculation of the bifurcation ratio as the slope of the shoot number vs. shoot age relationship (figure 2, Eq. (4)).

The bifurcation ratio (figure 2, Eq. (4)) at low to moderate light (\( Q_{\text{int}} < 20 \text{ mol m}^{-2} \text{ d}^{-1} \)) was not different between the infertile (average ± SE = 1.35 ± 0.21) and fertile (1.42 ± 0.12) site (figure 4C, means were not significantly different at \( P > 0.7 \) according to ANOVA). The bifurcation ratio scaled positively with irradiance in the fertile site (figure 4C), indicating that increased irradiance led to more frequent branching. In contrast, the bifurcation ratio did not respond to increases in irradiance in the infertile site, and the general mean of 1.314 ± 0.025 for all data from this site was similar to the value observed in low light in the fertile site.

The bifurcation ratio was positively related to average shoot length in both sites, but the explained variance was larger in the fertile than in the infertile site (figure 5, \( r^2 = 0.38, P < 0.005 \) for the correlation with the average values per tree in the infertile site). The slope of the \( R_b \) vs. \( L_s \) relationship of 0.26 cm\(^{-1}\) was larger (\( P < 0.001 \) according to ANCOVA) in the fertile than in the infertile site (0.07 cm\(^{-1}\)), demonstrating that the length of mother shoots controlled the branching less in the infertile site.

The ratio of current needle to shoot axis dry mass (\( \gamma \)) was positively related to irradiance in the fertile site (figure 4D), but not in the infertile site. However, \( \gamma \) was significantly lower (\( P < 0.001 \), analysis of covariance) at the infertile than at the fertile site, indicating that biomass requirement for needle support was larger in the nutrient-poor site.

Shoot dry matter content (\( \delta_s \), weighted average of needle and shoot axis dry matter contents) was significantly larger (\( P < 0.001 \)) with average ± SE = 0.551 ± 0.010 g g\(^{-1}\) in the

**Figure 4.** Effects of \( Q_{\text{int}} \) on the distribution characteristics of the length of current year shoots (A, B), on the bifurcation ratio (C, Eqs. (2–4), figure 2), and the partitioning of dry mass between needles and shoot axes (D). The inset in A shows frequency distributions of normalised shoot length for representative branches (denoted by arrows in A and B) from the fertile (filled bars) and infertile sites (open bars). Data presentation as in figure 3.
nutrient and light effects on canopy architecture in *Pinus* infertile than in the fertile stand (0.476 ± 0.005 g g⁻¹). The ratio of needle to shoot axis dry mass was positively related to \( s \) in the infertile site (\( r^2 = 0.18, P < 0.001 \)), but not in the fertile site (\( r^2 = 0.00, P > 0.9 \)).

3.3. Dependence of needle average age on light and nutrient availability

The maximum needle age observed was six years at the infertile and four years at the fertile site, suggesting that the site fertility significantly altered needle longevity. When the sites were considered separately, mass-weighted average needle age (\( \Lambda \), Eq. (5)) was independent of needle nitrogen content in both the infertile (\( r^2 = 0.04, P > 0.3 \)) and fertile stand (\( r^2 = 0.09, P > 0.5 \)). However, the needles were considerably older in the infertile site with an average \( \Lambda \pm SE \) for all shoots of 2.27 ± 0.05 yr. than the needles in the fertile site (1.70 ± 0.05 yr., the means are significantly different at \( P < 0.001 \) according to one way ANOVA). When the data for both sites were pooled, there was a strong negative correlation between needle nitrogen content and average needle age (figure 6A). A similar relationship was also observed for foliar \( P \) contents (\( r^2 = 0.48, P < 0.001 \)). The average needle age (\( \Lambda \)) was not significantly influenced by irradiance (figure 6B).

In both sites, \( \Lambda \) was negatively related to average shoot length (\( r^2 = 0.16, P < 0.02 \) for the infertile and \( r^2 = 0.49, P < 0.001 \) for the fertile site). For all data pooled, the explained variance (\( r^2 \)) was 0.37 (\( P < 0.001 \)), indicating strong interrelatedness of growth and needle longevity. \( \Lambda \) was positively related to needle density (\( r^2 = 0.18, P < 0.01 \)) and to shoot dry matter content (\( r^2 = 0.14, P < 0.005 \)). Thus, apart from scaling with growth, life span of more resistant needles tends to be larger.

3.4. Age effects on foliage morphological and chemical characteristics

Needle to axis mass ratio decreased with increasing shoot age (figure 7), and this decrease was stronger in the fertile site (\( P < 0.001 \) for the interaction term – age X site – according to a covariance analysis). Like for the current year shoots, the average ratio of needle to woody biomass of all needled shoot age classes pooled was significantly (\( P < 0.05 \) according to one-way ANOVA) lower in the infertile (1.84 ± 0.10 g g⁻¹) than in the fertile site (2.29 ± 0.22 g g⁻¹).

Needle dimensions – length, width, and thickness – as well as needle total area (\( A_T \)), and \( A_T \) to projected needle area ratio were mainly affected by \( Q_{int} \) in needles of all age classes, but were independent of needle age in both sites (table I). Needle dry mass per unit area (\( M_A \)) also increased with increasing irradiance (figure 8A, B), and was strongly affected by needle age (figure 8A, C, table I).

Given that \( M_A \) is the product of needle density (\( D \)) and volume to \( A_T \) ratio (\( V/A_T \)), the effects of \( Q_{int} \) and needle age on \( D \) and \( V/A_T \) were also studied to unravel the age effects on \( M_A \). Needle age did not significantly influence \( V/A_T \), but

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**Figure 5.** Correlations between average shoot length (\( L_s \)) and shoot bifurcation ratio, \( R_b \), in the fertile site (filled symbols) and in the infertile site (open symbols). Data presentation as in figure 3. The inset displays the relationship between \( R_b \) and \( L_s \) without the two uppermost data points in a better resolution (\( r^2 = 0.68, P < 0.005 \) for the fertile site).

**Figure 6.** Average needle age (\( \Lambda \), Eq. (5)) as a function of needle nitrogen content (\( N_M \), A) and irradiance (B). Within each site, \( \Lambda \) and \( N_M \) were not significantly related (\( r^2 = 0.11, P > 0.2 \) for the infertile and \( r^2 = 0.24, P > 0.1 \) for the fertile site). Given that neither the site effect at the common \( N_M \) nor the site X \( N_M \) interaction were significant according to ANCOVA (\( P > 0.2 \)), data were fitted by a single regression line in A. Symbols and regression lines as in figure 3.
Needle density strongly increased with increasing age ([table I](#)), providing an explanation for the age-related increases in \( M_A \). At the fertile site, irradiance was positively correlated with both \( D \) and \( V/A_T \), but more strongly with \( V/A_T \) ([figure 8B](#)) than with \( D \) \((r^2 = 0.16, P > 0.06 \text{ for } 1\text{-yr}, r^2 = 0.35, P < 0.05 \text{ for } 2\text{-yr} \text{ and } r^2 = 0.03, P > 0.8 \text{ for } 3\text{-yr needle})\). At the infertile site, similar fractions of explained variance were observed for both \( V/A_T \) ([figure 8D](#)) and \( D \) \((r^2 = 0.13, P < 0.02 \text{ for } 1\text{-yr}, r^2 = 0.14, P < 0.02 \text{ for } 2\text{-yr} \text{ and } r^2 = 0.10, P > 0.06 \text{ for } 3\text{-yr needle})\).

Needle nitrogen contents, \( N_M \), were independent of needle age in the fertile site ([table IA](#)), but \( N_M \) increased in the second-year needles relative to the first-year needles in the infertile site ([table IB](#)), suggesting that older needles remained physiologically competent. Foliage carbon contents increased with increasing needle age in both sites ([table I](#)), possibly because of age-related accumulation of certain carbon-rich compounds such as lignin or terpenoids. Increases in foliar carbon content were paralleled by modifications in needle density ([figure 9](#)).

The explained variance of all leaf structure and chemistry vs. irradiance relationships generally decreased with increasing needle age, possibly indicating that needles became less plastic with advancing age. Despite this, the interaction term, age \( \times Q_{\text{int}} \), was insignificant in all relationships \((P > 0.2)\). Accordingly, age and light independently altered needle morphology and chemistry.

4. DISCUSSION

4.1. Shoot growth characteristics

Monotonic increases in height growth and length of individual shoots in response to irradiance are frequently

![Figure 7](#). Needle to shoot axis dry mass ratio in relation to shoot age in the fertile (filled symbols) and infertile site (open symbols). According to a co-variation analysis (age as the covariate, site as the factor), both the site, and shoot age \( \times \) site interaction were significant determinants of the mass ratio \((P < 0.001 \text{ for both})\).

![Figure 8](#). Correlations of ([A, B]) needle dry mass per unit area \( (M_A) \) and ([C, D]) needle volume to total area ratio \( (V/A_T) \) with \( Q_{\text{int}} \) in needles of various age in the fertile ([A, C]) and the infertile site ([B, D]). \( M_A \) is the product of \( V/A_T \) and needle density. Current-year needles were attributed an age of 1-yr. Data for each needle age-class were fitted by separate linear regressions as depicted in A.
Table 1. Needle morphological characteristics, and nitrogen and carbon contents (average ± SE) in relation to needle age in the fertile (A) and the infertile (B) site, and the statistical significance of the effects of age and integrated quantum flux density ($Q_{int}$) on needle variables$^1$.

### A. Fertile site

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<th>Variable</th>
<th>Needle age</th>
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<tr>
<td>Total needle area ($A_T$, mm$^2$)</td>
<td>16.4 ± 8.9a</td>
<td>16.2 ± 10a</td>
<td>16.0 ± 10a</td>
<td>ns. 2</td>
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<td>$A_T$ to projected area ratio ($A_T$/AP)</td>
<td>2.588 ± 0.016a</td>
<td>2.532 ± 0.020a</td>
<td>2.562 ± 0.022a</td>
<td>ns. 0.005</td>
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<td>Dry mass per $A_T$ (g m$^{-2}$)</td>
<td>89.8 ± 3.2a</td>
<td>95.1 ± 3.7b</td>
<td>99.1 ± 3.0c</td>
<td>0.001 0.001</td>
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<tr>
<td>Density (g cm$^{-3}$)</td>
<td>0.488 ± 0.010a</td>
<td>0.530 ± 0.010b</td>
<td>0.561 ± 0.010c</td>
<td>0.001 0.001</td>
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<td>Volume to $A_T$ ratio ($V/A_T$, mm)</td>
<td>0.184 ± 0.006a</td>
<td>0.176 ± 0.050a</td>
<td>0.177 ± 0.005a</td>
<td>ns. 0.001</td>
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<td>Length (mm)</td>
<td>52.3 ± 1.5a</td>
<td>51.8 ± 1.5a</td>
<td>54.0 ± 1.8a</td>
<td>ns. 0.001</td>
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<td>Width (mm)</td>
<td>1.203 ± 0.044a</td>
<td>1.22 ± 0.05a</td>
<td>1.136 ± 0.040a</td>
<td>ns. 0.001</td>
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<td>Thickness (mm)</td>
<td>0.603 ± 0.016a</td>
<td>0.614 ± 0.026a</td>
<td>0.589 ± 0.015a</td>
<td>ns. 0.001</td>
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<tr>
<td>Nitrogen content (%)</td>
<td>1.531 ± 0.023a</td>
<td>1.479 ± 0.032a</td>
<td>1.518 ± 0.046a</td>
<td>ns. 0.01</td>
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<tr>
<td>Carbon content (%)</td>
<td>48.17 ± 0.27a</td>
<td>48.75 ± 0.21b</td>
<td>48.83 ± 0.23b</td>
<td>0.01 0.02</td>
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### B. Infertile site

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<tr>
<th>Variable</th>
<th>Needle age</th>
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<tr>
<td>Total needle area ($A_T$, mm$^2$)</td>
<td>71.5 ± 4.1a</td>
<td>68.8 ± 4.1a</td>
<td>72.3 ± 3.4a</td>
<td>58.5 ± 6.4a</td>
<td>83.4 ± 9.6a</td>
<td>ns. ns.</td>
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<tr>
<td>$A_T$ to projected area ratio ($A_T$/AP)</td>
<td>2.541 ± 0.008a</td>
<td>2.532 ± 0.009a</td>
<td>2.575 ± 0.047a</td>
<td>2.490 ± 0.022a</td>
<td>2.497 ± 0.023a</td>
<td>ns. ns.</td>
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<tr>
<td>Dry mass per $A_T$ (g m$^{-2}$)</td>
<td>93.7 ± 1.5a</td>
<td>103.3 ± 1.6b</td>
<td>114.5 ± 1.7c</td>
<td>110.7 ± 5.1bc</td>
<td>126.6 ± 2.3c</td>
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<tr>
<td>Density (g cm$^{-3}$)</td>
<td>0.611 ± 0.009a</td>
<td>0.688 ± 0.015b</td>
<td>0.730 ± 0.014b</td>
<td>0.727 ± 0.031b</td>
<td>0.778 ± 0.023b</td>
<td>0.001 0.001</td>
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<tr>
<td>Volume to $A_T$ ratio ($V/A_T$, mm)</td>
<td>0.1539 ± 0.0021a</td>
<td>0.1514 ± 0.0024a</td>
<td>0.1579 ± 0.0023a</td>
<td>0.1543 ± 0.0043a</td>
<td>0.163 ± 0.008a</td>
<td>0.005 ns.</td>
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<tr>
<td>Length (mm)</td>
<td>25.5 ± 1.1a</td>
<td>24.6 ± 1.1a</td>
<td>25.8 ± 1.0a</td>
<td>21.2 ± 2.1a</td>
<td>28.8 ± 2.3a</td>
<td>ns. ns.</td>
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<tr>
<td>Width (mm)</td>
<td>1.073 ± 0.021a</td>
<td>1.074 ± 0.024a</td>
<td>1.089 ± 0.016a</td>
<td>1.102 ± 0.021a</td>
<td>1.153 ± 0.039a</td>
<td>0.01 ns.</td>
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<tr>
<td>Thickness (mm)</td>
<td>0.510 ± 0.009a</td>
<td>0.503 ± 0.011a</td>
<td>0.510 ± 0.008a</td>
<td>0.491 ± 0.017a</td>
<td>0.521 ± 0.027a</td>
<td>0.005 ns.</td>
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<tr>
<td>Nitrogen content (%)</td>
<td>0.866 ± 0.022a</td>
<td>0.993 ± 0.032b</td>
<td>0.896 ± 0.037ab</td>
<td>nd. 3</td>
<td>nd. 0.02 ns.</td>
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<tr>
<td>Carbon content (%)</td>
<td>49.25 ± 0.14a</td>
<td>50.13 ± 0.10b</td>
<td>50.39 ± 0.10c</td>
<td>nd. nd.</td>
<td>0.001 ns.</td>
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$^1$ Means with the same letter are not significantly different ($P > 0.05$). The means were compared either by co-variation analyses when $Q_{int}$ significantly correlated with the specific foliar characteristic or by one way analyses of variance when $Q_{int}$ was insignificant in the former analysis. The interaction term, age x $Q_{int}$, was insignificant in all cases ($P > 0.2$). Thus, the co-variation analyses only included the factor and the covariate (common slope model). After the analysis of variance, Bonferroni test was employed to separate the significantly different means; 2 ns.: not significant; 3 nd.: not determined.

observed in conifers [12, 23, 38, 44, 79, 88]. As our study indicates, this relationship is strongly affected by site fertility (figure 3). Average shoot length responded to irradiance in the fertile site, but did not depend on irradiance in the infertile site (figure 3B). The fact that shoot length was positively correlated with needle nitrogen (figure 3A) and phosphorus contents in the infertile site provides conclusive evidence that the growth was chiefly limited by nutrients rather than by light in this site.

In conifers, absolute rates of lateral canopy expansion respond to irradiance similarly to height growth [12, 88]. Yet, the height growth increment generally exceeds the lateral growth such that the ratio of lateral to vertical growth may be negatively related to irradiance [12, 15, 88]. A relatively larger increase of vertical relative to horizontal growth with increasing irradiance is a major factor leading to various crown geometries – flat in low irradiance vs. conical in high irradiance. Thus, the arrested height growth may provide an explanation for the flat crown shape in the open environments in the infertile site.

The distributions of shoot length in forest trees are generally peaked and asymmetric with a greater number of short than long shoots [77] as was also observed in *P. sylvestris* in the fertile site (figure 4A, B). Similarly to previous observations in conifers [80, 90], the number of short shoots relative to long shoots increased progressively with increasing light availability in the fertile site (figure 4A, B) indicating a stronger apical control at higher irradiance. Although shoots branched more frequently at higher irradiance in the fertile site (figure 4C), stronger apical control permitted preferential resource investment in height growth. In contrast, apical control was released in the infertile site, where the shoot
possibly because of the positive scaling of increases with increasing light availability in the canopy [74], photosynthesizing leaf area. Leaf area density generally greater shoot number per unit crown volume and for greater 77]. More frequent branching at higher irradiance results in apical control of shoot growth in the infertile site. Growth was limited by nutrients in the infertile site, low sink are required for effective apical control [89]. Given that assimilate, either in the leader shoot or in the stem and roots, the mechanisms of hormone action are still unknown [13, 90]. Yet, there are conclusive data indicating that strong sinks for evidence that hormones are involved in the apical control, but resulted in primarily horizontal canopy extension. There is competition for resources by many independent growth points distributions were essentially normal (figure 4A, B), and the competition for resources by many independent growth points resulted in primarily horizontal canopy extension. There is evidence that hormones are involved in the apical control, but the mechanisms of hormone action are still unknown [13, 90]. Yet, there are conclusive data indicating that strong sinks for assimilate, either in the leader shoot or in the stem and roots, are required for effective apical control [89]. Given that growth was limited by nutrients in the infertile site, low sink activities may provide a mechanistic explanation for lower apical control of shoot growth in the infertile site.

4.2. Branching morphology

Bifurcation ratio ($R_b$, Eq. (2)) is an important branch parameter [22, 40] that may strongly affect the shoot density in the canopy [11], and thereby the aggregation of the leaf area. Although there exist non-plastic species with bifurcation ratios independent of long-term light availability [11, 61, 65, 87], $R_b$ is generally positively related to $Q_{\text{int}}$ [7, 11, 41, 65, 76, 77]. More frequent branching at higher irradiance results in greater shoot number per unit crown volume and for greater photosynthesizing leaf area. Leaf area density generally increases with increasing light availability in the canopy [74], possibly because of the positive scaling of $R_b$ with irradiance.

The dependence of $R_b$ on $Q_{\text{int}}$ in P. sylvestris in the fertile site indicates that it is a plastic species, but also that it requires high nutrient availabilities for maximum branching intensity and foliar area development. Although the high light environment favours conical crowns with multiple leaf layers (Introduction), P. sylvestris formed such crowns only in the high nutrient availability site. In the infertile site, branching morphology was not plastically modified in response to irradiance, and reduced shoot length growth, low rate of branching (figures 3 and 4C) and more horizontal branch inclination angles (personal observations) led to flat crowns with a few needle layers at all irradiances in this site. Because the flat crowns allow maximization of exposed needle area, such a foliar arrangement is particularly apt to low understory irradiances. Yet, the minimization of self-shading is not necessarily advantageous in high irradiance, because it increases the risk of photoinductive damage [64]. Given that the photosynthetic capacities were strongly reduced in the infertile relative to the fertile site [55], the probability for photoinduction at a common incident quantum flux density (62) for a review) was greater in the infertile than in the fertile site. Thus, we conclude that nutrient availability strongly curbed the morphological adjustment of crown shape and that the resulting crown architectures were not optimal for the specific environmental conditions.

Previously, the correlation between shoot length and bifurcation ratio has been used to model the canopy architecture in P. sylvestris [34, 36]. However, as our study demonstrates (figure 5), this relationship is considerably weaker in nutrient-limited environments where the shoots of the same length branch more frequently than the branches in the fertile site.

4.3. Dry matter partitioning between stems and foliage within the branch

Partitioning of shoot biomass between needles and shoot axes may be an additional determinant of foliar area in the tree. Conifers may decrease needle to shoot axis mass ratio with increasing irradiance [14, 33, 39, 44], thereby allowing more extensive needle area development at a common biomass investment in branches in low light. However, in our study, there was an increase in the fractional investment in needles with increasing $Q_{\text{int}}$ in the fertile site, and no effect of $Q_{\text{int}}$ in the other site (figure 4D). In other works, it has been observed that the fractional investment in needles was independent of irradiance [38, 56]. We cannot currently explain these contrasting patterns between the studies. However, given that conifers’ branches must sustain extensive snow loads in the winter, the requirements for mechanical stability may provide a possible explanation for the larger biomass investment in support in low irradiance. The branches are more horizontal in the lower canopy of P. sylvestris [35, 79], and thus, have effectively longer lever arms with greater biomass requirements for mechanical support [26, 46].

By the same token, the circumstance that the branches were essentially horizontal in the bog, and vertical in the forest (personal observations) may be a reason for lower needle to shoot axis mass ratio in the infertile site (figure 4D). In addition, stand density was less in the infertile (200 trees ha$^{-1}$) than in the fertile (1400 trees ha$^{-1}$) site. According to the simulation studies, the risk of snow damage is larger in stands with lower density [63], because average wind speeds are higher in less dense stands. Thus, the evidence collectively suggests that the lower biomass investment in the needles in the infertile site may reflect greater snow loads and mechanical stress in the winter.
4.4. Modification of average needle age by site nutrient availability

Decreases in shoot growth and branching in the infertile site can somewhat be compensated for by increased needle life span (figure 6). We hypothesized that the bifurcation ratio and needle longevity may be interrelated, because in species possessing leaves with a longer life span, an extensive foliar area can be formed with a lower frequency of branching. Although we did not observe such a relationship within the sites, the assumption was fulfilled for the patterns across the sites (cf. figures 4C and 6).

We studied average needle age, and therefore, it is relevant to consider that the relationship between the average needle age calculated as the mass-weighted average (Equation (5)) and needle longevity depends directly on shoot bifurcation ratio. This is because in a branching canopy, the mass of younger needle age classes is always progressively larger than the mass of older needle age classes, and this leads necessarily to a lower average needle age. For example, if the individual mother and daughter shoots have similar average needle mass, the total needle mass of daughter shoots is equal to \( R_t \) times the mass of needles in mother shoots. Nevertheless, the maximum needle age observed in our study was six years in the bog and four years in the forest. In addition, the needle to shoot axis dry mass ratio was significantly larger for older needle age classes in the bog as well (figure 7). Thus, we argue that modifications in average needle age (figure 6) truly mirror the changes in needle longevity.

According to the leaf life span model of Ackerly [1], the foliar life span reflects the shading patterns within the shoot. The model predicts that leaf life span increases with decreasing the rate of leaf production, because the older leaves intercept higher irradiances longer in branches with less rapid new leaf production, and accordingly, their carbon balance turns zero later than in branches with more rapid foliar production. Given that low needle growth rates were accompanied by increased average needle age in \( P. \text{sylvestris} \), our results also agree with the model.

Although nitrogen and other limiting elements are retranslocated from senescing conifer foliage [17, 27, 31, 52], large quantities of nitrogen are necessary to increase the efficiency of light harvesting in shaded needles [9, 10]. Given that needles in most shaded shoot positions are the first to abscise [48], keeping high N contents in older, but functionally active needles may be an important acclimation response to increase the light interception efficiency in the low light environments [48]. Maintenance of high N contents may also be the prerequisite for high needle longevity [48]. We found that the needle nutrient contents were independent of needle age in the fertile site, and were even larger in the second- than in the current-year needles in the infertile site (table I). Larger N contents in the second-year needles hint at lower rate of nutrient loss in the bog than in the forest. Reduced rate of age-related declines in needle N contents and thus, in the functional activity of needles in the infertile site, may be directly associated with decreased growth rates and less extensive shading in this site. Thus, the foliar nitrogen vs. age dependencies also indirectly support the argument that the shoot growth rates may exert an effective control over needle longevity.

4.5. Morphological and chemical characteristics of needles of various age

Apart from carbon balance arguments, needle longevity may directly depend on foliage structural characteristics that improve needle resistance to mechanical damage as well as on the speed of age-related modifications in foliage structural characteristics. There is conclusive evidence of secondary needle growth in conifers [18, 25]. As the needles age, the number of xylem and phloem layers increases in the vascular bundles of conifer needles [18, 25]. Possibly because of the secondary needle growth, needle thickness and width increased with increasing needle age in \( Picea \text{abies} \) [48]. However, we did not find any significant age effect on needle width and thickness (table I), and needle volume to total area ratio was also independent of age in \( P. \text{sylvestris} \) (figure 8C, D). Given that with increasing the number of xylem and phloem layers in the needle vascular bundles, phloem and xylem become increasingly compressed [18, 25], restricted expansion of vascular bundles because of lignified cell walls of the bundle sheath cells may explain away the missing age effect on needle thickness and width in the present study. Increases in needle carbon concentrations with needle age (table I) indirectly support the idea of advancing lignification with needle age. Lignin is a carbon-rich chemical (65.4% C, calculated according to [24]), and increases in foliar lignin contents are generally accompanied by increases in C contents [57].

As in other conifers [10, 48], the explained variance in leaf structure vs. \( Q_{\text{int}} \) relationships consistently decreased with increasing needle age (figure 8). However, there were no irradiance x age interactions, indicating that irradiance did not influence the age-related modifications in needle structure. An increase in needle dry mass per unit area (table I, figure 8A, B) was the primary age-caused change in needle morphology. Similar changes in \( M_A \) with age have also been observed previously [27, 48, 82]. As the former [48] and current work demonstrate, changes in \( M_A \) primarily result from modifications in needle density (table I). Such adjustments in density are compatible with smaller and more tightly packed cells, greater fraction of cell walls in the leaves and greater lignification [50], and accordingly with mechanically more resistant leaves [51]. Given that the needle density was consistently larger in the infertile relative to the fertile site (table I), it is probable that changes in density may be an important driver of needle longevity.

In conclusion, we demonstrated extensive light-related plastic adjustments in crown architectural characteristics in \( P. \text{sylvestris} \), but also that the plasticity strongly depended on site nutrient availability. There was evidence that the crowns developing in the infertile site were not optimal for light interception and plant performance. Although the growth was limited by nitrogen in this site, and the effective light capture was not of paramount importance for growth, the crowns with low self-shading likely led to high risk of photoinhibitory damage in high irradiance. We also found an extension in needle life span with decreasing nutrient availability. This nutrient availability related modification in needle longevity is possibly the direct consequence of altered shoot growth and within canopy shading patterns. Yet, changes in needle morphological characteristics that alter needle resistance to
mechanical lesions may also play a role in needle longevity. From a practical perspective, the interrelationships between crown structural characteristics may be used to model the canopy architecture in *P. sylvestris* [36]. However, our study indicates that such relationships strongly depend on nutrient availability, and we conclude that site fertility controls on branching and shoot length growth must be included in the future canopy architecture models.

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