



Wind and gravity in shaping *Picea* trunks

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Abstract

Key message Spruce trunk tapering corresponds closely to tapering required to resist bending forces caused by wind and gravity.

Abstract Understanding why trunks (tree stems) are the size that they are is important. However, this understanding is fragmented into isolated schools of thought and has been far from complete. Realistic calculations on minimum trunk diameters needed to resist bending moments caused by wind and gravity would be a significant step forward. However, advancements using this biomechanical approach have been delayed by difficulties in modelling bending of trunks and wind gusts. We felled and measured five Norway spruces (*Picea abies*) in an unthinned monoculture in southeastern Finland planted 67 years earlier. We then focused on forces working on storm-bent (maximally bent) trees caused by gravity and the strongest gust in a 1-h simulation with a large-eddy simulation model. The weakest points along the trunks of the three largest trees resisted mean above-canopy wind speeds ranging from 10.2 to 12.7 m s⁻¹ (3.3-fold in the strongest gust), but the two smallest were well protected by a dense layer of leaves from the bending tops of larger trees, and could have resisted stronger winds. Gravity caused approximately one quarter of the critical bending moments. The wind that breaks the trunks in their weakest points is close to breaking them in other points, supporting the importance of bending moments caused by wind and gravity in the evolution of trunk taper.

Keywords Gravity · Norway spruce (*Picea abies* [L.] Karst.) · Sail area · Stem · Thigmomorphogenesis · Trunk taper · Wind drag · Wood

Introduction

Understanding why trees and their trunks (stems) are the size that they are is important for evaluating the potential of forests to mitigate climate change and produce timber.

Therefore it is surprising that that the scientific understanding of tree height and diameters along the trunk is fragmented. For example, a question concerning the dimensional determinants of a particular tree trunk may cause surprise and be considered too general by experts in narrow fields, even though understanding trunk dimensions should be considered one of the largest questions in applied ecology.

Research on trunk dimensions can be classified in two ways. First, the classification can be based on the object of the study, i.e. the state of the forest. Some studies focus on those experiencing natural successions (Anderson-Teixeira et al. 2013), others on tallest trees in old-growth forests (Van Pelt et al. 2016), on plantations subject to self-thinning (Yoda et al. 1963) or impacts of silvicultural treatments (Bianchi et al. 2020). Second, the classification can be based on whether the approach is descriptive or theoretical. The majority of research on trunk dimensions in forest sciences and research related to forests' role in climate change mitigation is mainly descriptive (e.g. Chave et al. 2014), while much of the physiological and ecological research attempts

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to explain the causes of the described patterns based on theories. These theories may be grouped based on the function on which the focus is transport, storage or biomechanical support as explained in the following paragraphs.

Trees passively transport sap (water) up in the sapwood, and the resistance caused by the length of the path or need to lift sap against gravity has been used as the basis for modelling maximum tree height (Koch et al. 2004) and growth deceleration in plantations (Ryan and Yoder 1997). However, as the heartwood is not contributing to sap transport, diameters along the trunk cannot be understood based on sap transport only, unless heartwood is considered a waste produced, e.g. due to ageing (Collalti et al. 2019) or difficulties in using the same sapwood when branches die and grow (Chiba et al. 1988). Phloem transport down the trunk may be similarly limiting tree height due to path length (Woodruff 2013), but again, does not explain diameters, unless trunk circumference needs to be increased to increase transport capacity.

Trees store water (Scholz et al. 2011) and energy (Schiestl-Aalto et al. 2015) in their woody tissues and this is likely to influence trunk dimensions in certain conditions. For example, baobab (*Adansonia digitata*) trees probably have unusually fat trunks to store water needed to level out seasonal variation in water availability (Chapotin et al. 2006), and lignotubers located at the trunk base can store energy and nutrients enabling rapid sprouting (Canadell and López-Soria 1998). Trunk dimensions are, therefore, potentially influenced by storage needs, but this is unlikely to be common and may be restricted to the rare trees that do not form metabolically dead heartwood and therefore cannot increase sapwood volume by adjusting the sapwood-to-heartwood ratio such as the above-mentioned baobabs (Patrut et al. 2010).

The third general function of trunks in addition to transport and storage, and the only for which heartwood is useful, is to biomechanically support the leaves, branches and trunk sections above the height at which the focus is. Common sense tells that trees exposed to wind or heavy loads need to have thicker trunks for a given height and crown size. These mechanisms have been studied experimentally for over two centuries (Telewski 2016), and the term “thigmomorphogenesis” has become established in the recent decades to describe the responses of plants to mechanical stimuli (Pruyn et al. 2000). Two very different mechanisms may serve as a basis for modelling trunk dimensions biomechanically. Elastic buckling (Euler buckling) can permanently bend trunks if the tree fresh mass and permanent loads, such as epiphytes and lianas, exceed the limit that the trunk can support. Modelling can be performed easily (McMahon 1973), and normally a “safety factor” is computed describing how far the height of the tree is from a height that leads to buckling. This approach has been used

in well-known modelling approaches (e.g. West et al. 1999). However, most trees, with the exception of certain rainforest understorey trees, are far from elastic buckling. For example, Niklas (1994) reported an average safety factor of four. Because of non-linearities, a safety factor of four implies that the plant’s weight is only 1.6% of the weight that would lead to elastic buckling. Furthermore, the safety factor is a misleading concept and should not be interpreted as an indication of biomechanical safety. A safety margin is needed for engineer-designed structures, as they are built and then need to resist variable forces without subsequent adjustments to the structure. However, thanks to thigmomorphogenesis, trees can tune their structure (Bonnesoeur et al. 2016) and a small safety factor is therefore not dangerous, as supporting tissue can be increased according to demand from the increasing height or weight. The wide usage of the theory on elastic buckling shocked Mattheck (2012) and he wrote: “Much to the surprise of the author, failure by buckling has nevertheless been discussed by McMahon (1973), and comparisons have been made between measured height-diameter relations and relations calculated from Euler’s buckling theory.” The other, more useful, biomechanical approach is based on trunks breaking. Brief buckling e.g. due to a temporary load of snow may not be a problem for the tree if it recovers and is erect most of the time. However, when modelling trunk breakage, even a short period to which the tree has not been able to acclimatize may be fatal. This modelling approach is challenging to follow, as wind speeds are variable in space and time, and trunks, branches and leaves streamline in wind. In both buckling and breaking approaches, diameters needed along the trunk for a given height and other characteristics can be computed based on biomechanics. However, these approaches do not limit height if the diameters are not limited.

All trees need trunks to transport, to provide biomechanical support and probably also to store, and theories and modelling to understand trunk dimensions should ideally incorporate all of these with appropriate weights. However, in practice realistic modelling of even one of these aspects at a time is challenging. Therefore, it is useful to consider their relative importance. One challenge is that scientists are often experts on only one of these three functions and may therefore overestimate its importance, even though some reviews on all them are available (Badel et al. 2015). Second, if building or maintaining a trunk that is superior in any of the three functions causes an energetic cost, all functions would evolve close to the needed level even if the cost improving it relative to the others would be minuscule. Therefore, demonstrating that, e.g. transporting sap higher than the current height of the tallest trees (Koch et al. 2004) does not necessarily mean that sap transport is the main factor determining maximum heights. Instead, in evolutionary timescales for example sap transport capacity could improve

to a height determined mainly by the biomechanics and energetics of maintaining the living biomass.

We did not properly assess the relative importance of how the functions of trunks influence their dimensions, as that would need to be done by incorporating them into one model. In this paragraph, we just note a few pieces of evidence that indicated to us the direction to take. One approach is to consider the marginal construction and maintenance costs of increasing capacity. Tissue suitable for storage or sap transport may be increased by increasing sapwood to heartwood ratio. Furthermore, sap transport efficiency may be boosted without compromising transport safety by increasing the density of conduits in angiosperm wood, probably with little or no additional construction costs (Larjavaara 2021). However, significant strengthening of the trunk is not possible without substantial additional construction costs, either by increasing diameters or wood density (Larjavaara and Muller-Landau 2012).

Another approach to know about the relative importance of factors influencing trunk dimensions is to compare them in variable environmental conditions that demand variable transport, storage and biomechanical support needs. This approach underlines the importance of sap transport if height and diameters along the trunk vary according to water availability. The very tallest trees would then be expected to be found in climates and soils with the most abundant water, which is not the case, even though the driest climates have a low canopy height (Klein et al. 2015). If storage function was critical in determining trunk dimensions, then seasonality should increase trunk volumes relative to leaf area, which may be the case (Chapotin et al. 2006) but probably only in the case of exceptional species. Finally, with biomechanical support being the most significant, tree heights and forest biomasses should vary depending on winds. This is the case for example with variable distances from the edge and therefore variable wind regimes (Brüchert and Gardiner 2006). Another perspective on the importance of biomechanics is provided by comparing trees to lianas, which do not have the same biomechanical support needs. Lianas have similar transport and storage needs as trees, and much higher leaf area for a given stem basal area (Ichihashi and Tateno 2015), which is very likely due to differing biomechanical needs, highlighting their importance to trunk diameters. These considerations led us to explore the role of wind forces and gravity as key determinants of trunk diameters, which is the focus of this article.

The importance of wind and gravity as a cause of trunk breakage is perhaps what common sense would suggest to be the main factor explaining trunk dimensions. This approach was pioneered in the nineteenth century (Metzger 1893) and regularly discussed from perspectives described with keywords such as “uniform stress model” (Mäkelä 2002) and thigmomorphogenesis when the focus is on the formation of

the tissue leading to this uniform stress (Pruyn et al. 2000). However, we argue that it still remains underrepresented and that this is probably due to methodological challenges from variable winds. In addition, the streamlining mentioned above and the rarity of the strongest storms that are critical for tree survival and therefore probably drive evolution cause extra challenges. Interesting studies are available on small trees secured on the roof of a moving car (Butler et al. 2012) and medium-sized trees during the leafless period (Niklas and Spatz 2000), but small (Larjavaara 2015) and leafless (Mattheck 2000) trees have different biomechanical constraints than large foliated trees. Large foliated trees have also been examined in impressive studies representing simple (Morgan and Cannell 1994), more realistic (Spatz and Bruechert 2000) or excellent detail in tree dimensions (Jackson et al. 2019). However, none of these studies focused on maximally bent trees.

The objective of this study was to increase our comprehension of determinants concerning tree size and trunk taper, as modified by selective pressures caused by exposure to storm-strength winds, and to examine whether trees are adapted and acclimatized via thigmomorphogenesis to those. To this end, we modelled wind in a canopy of a mature storm-bent stand and computed gravity- and wind-caused forces on segments along the trunks based on destructive sampling of *Picea abies* [L.] Karst. trees (Norway spruces). We then focused on the winds that break the trunks at their weakest segment and expected diameters at other segments to be only slightly larger than what was needed to resist the bending moments caused by this wind and gravity.

Methods

Picea abies is a common tree species in its natural range of Northern Europe and Central European mountains and is also planted widely in Central European lowlands and North America (Caudullo et al. 2016). In Finland, *Picea abies* trunk volumes make up 30% of all tree trunk volumes and the volume of harvested trees is 38% of total (Peltola 2014). It regenerates in intermediate or fertile soils, is the most shade tolerant of the common tree species in Finland and will, therefore, invade all but the most infertile sandy or peat soils when sufficient time since disturbance has passed (Kuuluvainen and Aakala 2011). *Picea abies* trees have a straight trunk and long conical crown often reaching the ground. In Finland, the lower branches shed normally only from the lower crown layers in the deep shade of conspecifics. New branches develop annually, forming whorls of branches. Its wood is of low density at 374 kg m^{-3} (Kantola and Makela 2006), especially when compared to angiosperms (Chave et al. 2009).

We based our study on data collected in 2001 to investigate crown development in three sites around southern Finland in stands after canopy closure (Kantola and Mäkelä 2004). However, to reduce the complexity of wind gust-related analysis, only a single plot featuring flat terrain is included in this study. The other plots were excluded because of hilly terrain, which alters low-altitude winds in a complex manner (Gardiner et al. 2016). The included plot, described in more detail by Kantola and Mäkelä (2004), was located in Punkaharju municipality at 61° 49' N, 29° 19' E, now part of Savonlinna in southeastern Finland. The local climate is conducive to tree growth, as abundant lakes level out temperature fluctuations during the growing season. The soils in the plot are well above average fertility for the region, classified as *Oxalis*-type (Cajander 1949), leading to a site index, H_{100} of 32 m. The monoculture of *Picea abies* trees was planted 67 years prior to data collection.

Three stands with varying thinning histories were studied in the plot but two were excluded from our study because of thinnings, as explained below. The included unthinned stand had a basal area of 44 m² ha⁻¹ and stand density of 805 ha⁻¹. Five sample trees representing various canopy layers were felled, and their trunks, branches and leaves (i.e. needles) were measured and weighed as described in detail by Kantola and Mäkelä (2004). In summary, trunk diameters were measured below each whorl of branches, and all branches were cut and measured and a subset of them taken to a laboratory for more detailed measurements. The heights and diameters of the five trees at a 1.3-m height ($d_{1.3}$) can be seen in Fig. 8 in “Results”. The percentage of the trunk with living branches of the five sample trees differed between 42 and 63, being greatest for dominant trees and smallest for trees grown in more suppressed positions. And further, the more suppressed trees also had the lightest-weight crowns compared to more dominant ones, which was consistent with the pipe model theory (Kantola and Mäkelä 2004).

For this study, we divided the five tree trunks into “segments” and estimated their angle relative to vertical and location relative to the base based on bending and length of all segments below. From the angle relative to vertical we computed their projected area perpendicular to wind direction (i.e. frontal area) and fresh mass based on volumes. We assumed the centre of each segment to be in the whorl of branches and extremes to be located halfway between neighbouring whorls. We divided the unmeasured lower branchless trunk into four segments, with the lowest centred at a height of 1.3 m, the remaining three at regular intervals between 1.3 m and the lowest whorl and assumed diameter to simply change linearly, as we anticipated this lowest part of the trunk to contribute only little to the bending moments or to the bending of the trunk.

The streamlining of trees is complex, and therefore the common approach is to simulate upright trees but with

reduced wind drag estimated with a coefficient (Gardiner et al. 2016). We instead focused on the strongest gust and “storm-bent” trees, i.e. trees bent along their trunks as much as they can without breaking (see Fig. 6). This focus was based on the reasoning that even though acclimation is likely to be mainly driven via thigmomorphogenesis by signals from normal wind speeds (Bonnesoeur et al. 2016), trunks are probably tuned to resist the strongest gusts based on normal winds. Maximum strain in both compression and tension may be assumed to equal the ratio of modulus of rupture and modulus of elasticity (but see Niez et al. 2019). In a bending segment or cylinder, the maximum tension occurs in the outermost fibres of the convex side and maximum compression on the opposite side. However, to simplify the calculations, we assumed rigidity of the segments (as can be seen in Fig. 3) and bending was realised by assuming a change (α) in the deviation of the axis of the segment relative to the segment below:

$$\alpha = \sin^{-1} \frac{2l\sigma}{dE}, \quad (1)$$

where l is the length of the segment, σ the modulus of rupture obtained from tree-pulling experiment is 36.26 Mpa (Peltola et al. 2000), d the diameter of the segment at its centre and E the modulus of elasticity is 7730 Mpa (Peltola et al. 2000).

We used the projected area of trunks, branches and leaves (we call their sum “sail area”) first for estimating wind speeds and then to compute wind-caused horizontal forces (Online Resource 1).

In addition to the five felled trees, we measured the $d_{1.3}$ of all trees less than 7 m away from the felled ones. We estimated their sail area and its vertical storm-bent distribution by fitting two simple linear regressions to the variables. We then first computed the storm-bent height based on the model in Fig. 1 and then its sail area based on the model in Fig. 2, in which a linear relationship was expected based on biomechanics, as the bending moment is expected to scale roughly with the product of the sail area and the length of the lever (tree height) and the strength of the trunk with the cube of its diameter (Ennos 2012). We plotted these models for all three stands but observed the fit to be tight in the unthinned plot only. We surmised that as the previous thinning occurred only 14 years prior to the measurement, the trunk dimensions relative to the sail area (Online Resource 1, Figs. S1 and S2) were possibly still unbalanced because of too little time since the thinning. We, therefore, excluded these stands from the analysis.

The mean $d_{1.3}$ of the five felled trees was 0.272 m and they ranged from 0.213 to 0.328 m, while the surrounding trees around these five had a mean of 0.260 m and a range from 0.167 to 0.382 m. Because of the tight fit of models in Figs. 1 and 2, we do not think that extrapolating to some

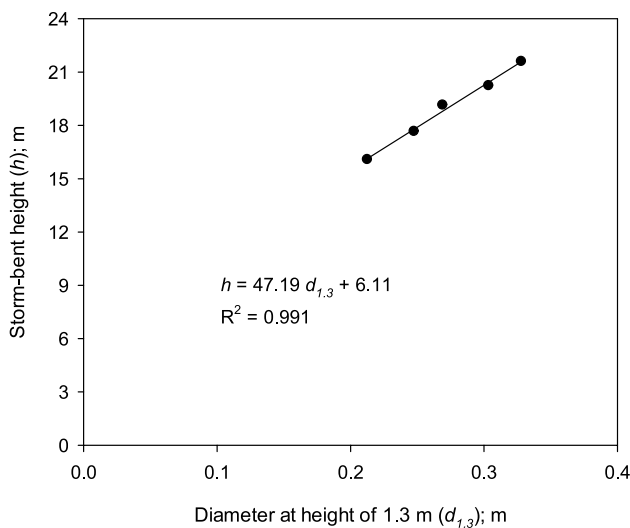


Fig. 1 Storm-bent height of the five felled trees plotted against $d_{1.3}$ and a fitted linear regression model

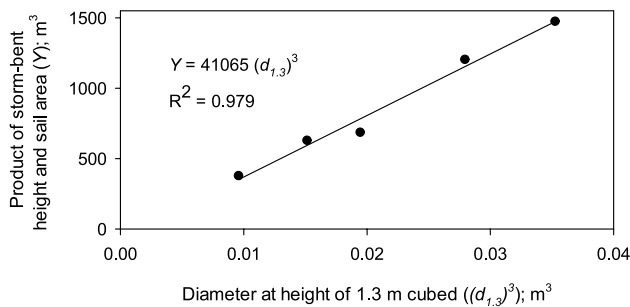


Fig. 2 Storm-bent height of the five felled trees multiplied by their sail area (projected area of trunk, branches and leaves) plotted against the cube of $d_{1.3}$ and a fitted linear regression model

distance out of the range was likely to cause a significant bias.

We wanted to focus on the strongest wind gusts that the trees can stand and, therefore, used turbulence resolving large-eddy simulation (LES) model to describe wind behaviour above and within forest canopies. Because of significant horizontal movement of trees in gusts, we had to assume that the forest canopy had a horizontally homogenous sail area and therefore sail area per unit volume (i.e. plant area density) for each 1.5-m thick layer. The large-eddy simulation model PALM (Maronga et al. 2015) was employed to obtain a time-accurate and spatially resolved description of fully developed boundary layer turbulence over continuous forest canopy. The PALM model is specifically tailored for atmospheric boundary layer turbulence applications and has been optimized for massively parallel supercomputing environments. The model implements the conservation equations governing atmospheric boundary layer turbulence

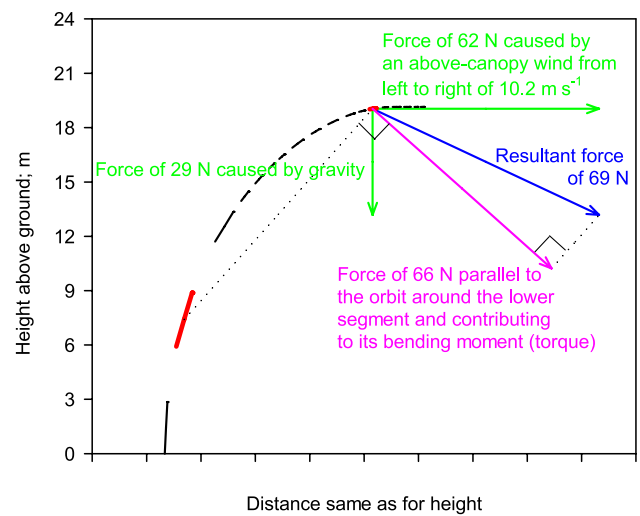


Fig. 3 An example of how we computed the bending moments from the forces caused by gravity and wind blowing from left to right. The “dashed” line represents storm-bent Tree3 with 18 uneven segments visible out of its 35 segments. The vectors show how we computed the moment caused by the 11th topmost segment to the 3rd lowest segment (both of which are highlighted with a thicker red line)

employing finite-difference discretization on a staggered Cartesian grid. The system of equations is solved using a third-order accurate Runge–Kutta time-stepping scheme and fifth-order accurate upwind biased spatial discretization scheme (Wicker and Skamarock 2002). The forest canopy is modelled assuming a porous homogenous medium within each 1.5-m layer, whose porosity varies according to the measured vertical sample-averaged plant area density distribution of the trees.

A vast majority of the drag caused by the forest canopy was assumed to be pressure drag, and, therefore, the drag force (f) is implemented in PALM as

$$\vec{f} = C_d P |\vec{u}| \vec{u}, \tag{2}$$

where C_d is the drag coefficient for forest canopy, P is the vertical plant area density profile of the forest, and \vec{u} is the spatially and temporally resolved wind velocity vector whose magnitude is denoted as $|\vec{u}|$. We set C_d at 0.2 as suggested by Katul (1998). The wind simulations were performed on a rectangular domain with L_x of 3.84 km, L_y of 1.28 km and L_z of 0.52 km as streamwise, lateral and vertical dimensions, respectively. Wind was driven with a prescribed pressure gradient at $z > 250$ m, allowing the lower-altitude flow to attain a constant momentum flux layer, which is characteristic of atmospheric boundary layer flows (Stull 2012). The magnitude of the pressure gradient was set sufficiently high to achieve very high Reynolds number conditions, which ensures that the associated turbulence solution attains a state that is independent of wind speed. That is, if the wind speed

were further increased, the turbulent structures and dynamics would remain statistically identical. This Reynolds number independence allows one representative turbulent wind solution to be freely scaled (especially upward) to represent other wind conditions. The simulation for the (scalable) reference wind was initially run for one hour to allow the flow to reach a statistically stationary state. The simulation was then continued for an additional hour during which detailed wind velocity time series is collected every 3 s (at 1/3 Hz) across the entire depth of the forest canopy from a 0.5-km² monitoring plane with 409 × 205 locations. This time series contains a sample of 105.6 × 10⁶ instantaneous wind events impacting the forest canopy. As the main interest is on gusts whose duration is sufficient to cause further displacements in the tree trunks, two consecutive wind events are averaged to yield a conservative approximation for a 3-s gust. Thus, the time series contained approximately 50 × 10⁶ gust events, which is considered a sufficiently large sample size to capture rare gust events that impose the largest risk for trunk failure. The gust events causing the maximal bending moments were searched by considering the forest canopy to contain trees with uniform horizontal cross-sections (just for the sake of wind gust analysis). The bending moment for each model tree was computed for all 3-s gust events and the maximum events (time and location) were stored. The wind speed profile spanning across the tree height was then obtained from this location and instance. The selected gust event provided the most realistic estimation for the critical velocity distribution during a probable failure event.

In addition to the normal simulation named “Dense”, we performed a second simulation with half of the sail area removed from all heights above ground (i.e. “Thinned”) and a third simulation with trunks and branches remaining but leaves removed (i.e. “Leafless”). However, it is important to note that these two secondary simulations violate the basis of our modelling of trees evolved via thigmomorphogenesis to withstand a given above-canopy wind speed by equal strain along the trunk, as a sudden thinning or defoliation would disturb the balance to which trees have acclimated and trunks would therefore likely break from a severely underbuilt segment before full bending is reached.

We computed the bending moments by adding moments from all segments and associated branches and leaves above the segment in question (Fig. 3). We obtained the weights, i.e. the vertical forces, by adding water contents of 0.79 for the trunk, 1.41 for the branches and 2.24 for leaves (Kantola and Makela 2006; Kärkkäinen 1985) to the dry masses (Kantola and Mäkelä 2004) and multiplying by the gravity constant (9.82 m s⁻²). We did not take physical contact between the trees into account.

The critical bending moment, i.e. the maximum bending moment that a cylindrical segment can resist (m_r) is

$$m_r = \frac{\pi \sigma d^3}{32}, \quad (3)$$

where σ is the modulus of rupture and d is the diameter of the segment (Ennos 2012). The sum of gravity- and wind-caused bending moments that cause this same m_r for the trunk segment is

$$m_r = r^2 \sum m_u + \sum m_g, \quad (4)$$

where $\sum m_g$ is the sum of all gravity-caused bending moments of all the segments and associated branches and leaves above, $\sum m_u$ is the sum of all wind-caused bending moments from segments and associated branches and leaves above in a reference above-canopy mean wind speed and r is the ratio of the maximum and reference (to compute $\sum m_u$) mean above-canopy wind speeds based on the wind profile obtained from the PALM model. These steps are shown as a flowchart in Fig. 4.

We then computed critical wind speeds that break the trunks in their weakest segments and compared diameters of other segments to those needed to resist this wind. We did not “tune” the approach or parameters to obtain a desirable fit. Below, we report the results from the analysis planned before beginning analysing the data with the exception of the exclusion of recently thinned plots.

Results

Most of the sail area of the five felled trees is caused by leaves and is located, once the trees are storm-bent, at a height of 15–21 m (Fig. 5a). When the surrounding trees are added, the layer of dense sail area thickens, mainly upward (Fig. 5b), but is still surprisingly thin for a tree species having an unusually long crown. The lack of thinnings in the studied stand has probably resulted in unusually small crown ratios and thin trunks enabling considerable bending, both of which thin the layer of dense sail area in a storm-bent stand.

The gust wind speeds are weak below 8 m and increase roughly linearly upwards through the mainsail area in Dense and Thinned stands (Fig. 5c). However, gust wind is significant down to the ground in the Leafless stand (Fig. 5c).

The weight of the branchless lower parts of the trunks of all five felled trees is important, but they cause bending moments only to the lower segments of the trunk. These moments are small, as the segments are nearly vertically aligned (Fig. 6). The weights from the upper segments and associated branches and leaves that produce potentially more significant bending moments are roughly evenly divided by those caused by the trunk, branches and leaves (Fig. 6). The comparison between trees illustrates how trees with larger $d_{1,3}$ (Tree4 and Tree5) have correspondingly heavier crowns

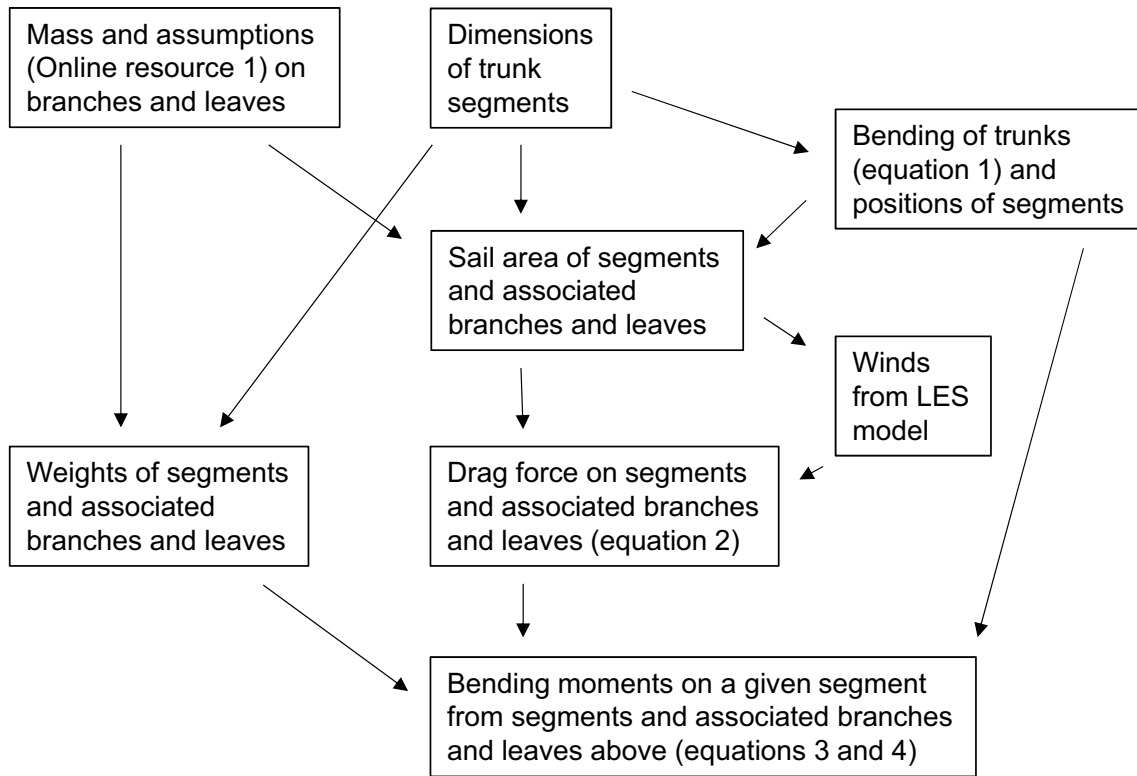


Fig. 4 Calculation of bending moments on segments

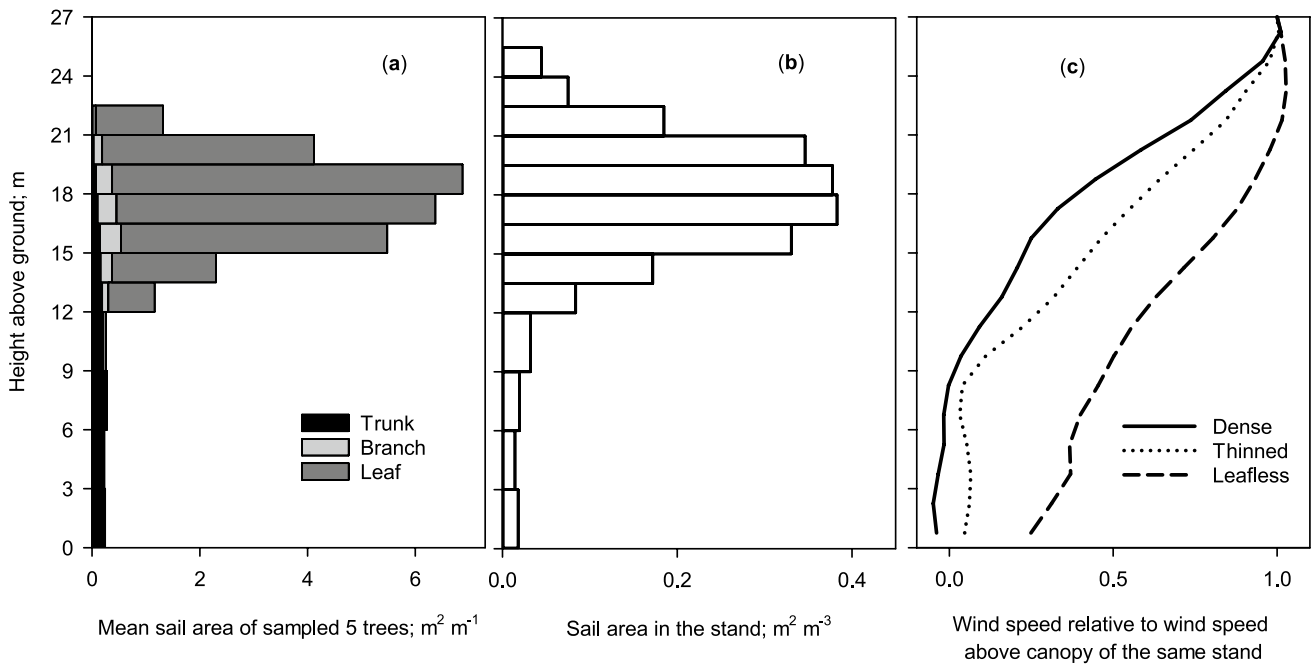


Fig. 5 Sail area and winds in a gust at various heights in the canopy and just above

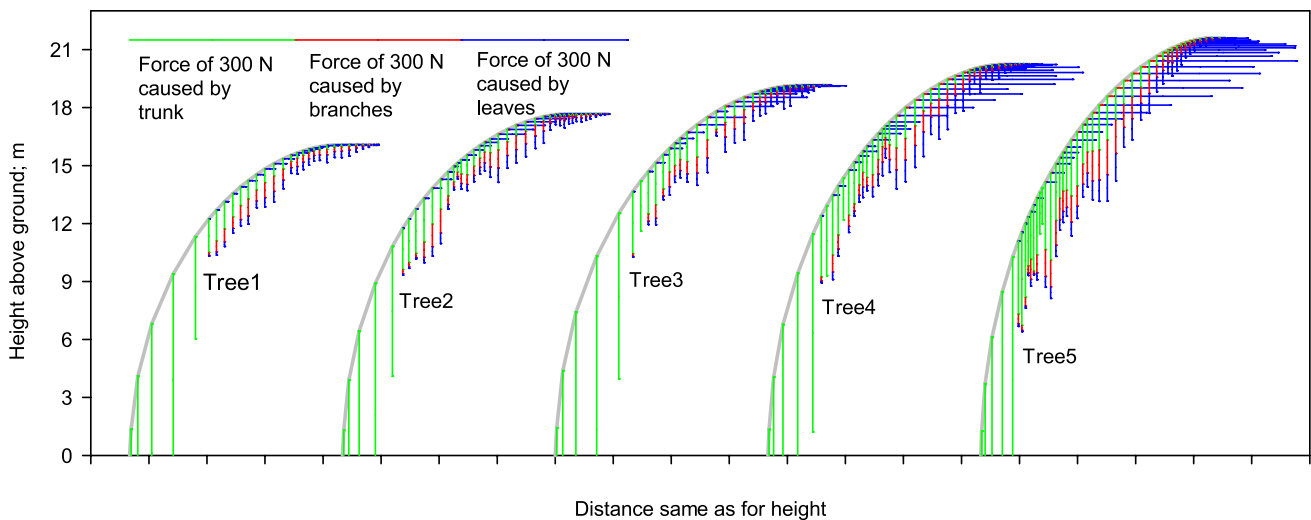


Fig. 6 The five felled trees shown as storm-bent. The number of the poorly visible topmost segments that have bent to horizontal ranges from 4 (Tree5) to 11 (Tree2). The green, red and blue horizontal lines represent force vectors caused by wind in the dense simulation on each segment, with the colour indicating whether the drag is caused

by the trunk, branches or leaves. The vertical lines represent forces caused by gravity. The length of vertical vectors from the lowest segments is not shown. The bottom end of a vector is -5.7 from the lowest segment of Tree5 with the same scale below the 0-level of the Y-axis as above

but the differences are small. The differences between the five trees are much more significant when the horizontal vectors caused by wind are examined (Fig. 6). The smallest trees experience much greater forces caused by gravity than wind, whereas both forces are of the same magnitude in the crowns of the largest trees. However, the wind-caused forces act higher up along the trunk and their direction also causes greater strengthening requirements for the lower trunk. Because the top of storm-bent Tree1 is only at a height of 16.1 m, it is well protected by more rigid taller trees (Fig. 6). Interestingly, because the shorter trees bend more, the horizontal displacement caused by wind is approximately the same for all five trees, ranging from 12.7 m (Tree5) to 14.3 m (Tree3).

Gravity from all segments and associated branches and leaves above the height at which the focus is 18–27% of the bending moment that breaks a tree at a height of 1.3 m (Fig. 7). This proportion increases upwards to a height of 12–15 m with the lowest branches and then decreases down to a rounded 0% for the tops of the trees. However, as the bark is included in the used d and the wood characteristics are unusual for the topmost segments, the estimated proportion is likely to be a severe underestimation. Nevertheless, the proportion of gravity relative to the critical bending moment clearly decreases upwards in the canopy.

Figure 8 demonstrates the dimensions of the five felled trees without wind and in addition to the measured diameters, the diameters needed to resist an above-canopy mean wind of 10.2 m s^{-1} , which is the speed that is at the limit of breaking Tree4. This can be seen from the dotted red line

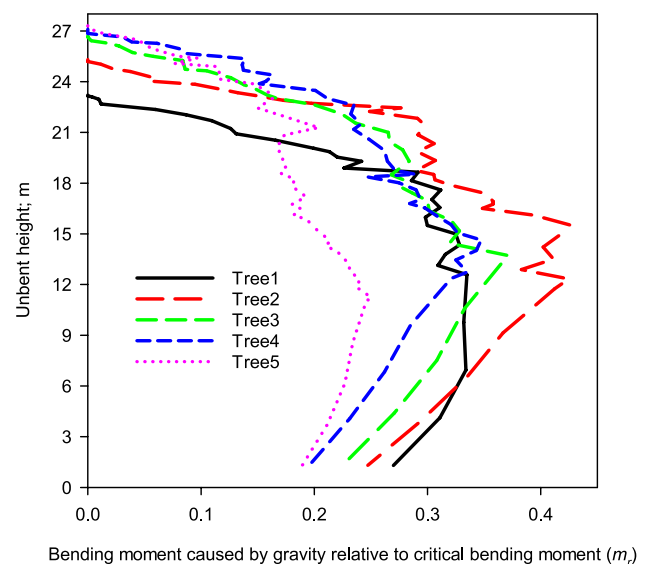


Fig. 7 The relative importance of the bending moment caused by gravity acting on segments and associated branches and leaves above the segment in question

contacting the solid black line at a height of 13.9 m. Tree3 and Tree5 are able to resist similar mean above-canopy wind speeds (12.7 m s^{-1} and 11.3 m s^{-1}), and therefore, the modelled taper is similar to the measured taper (Fig. 8). However, for Tree2 and especially Tree1, a significantly thinner trunk would be sufficient to withstand the simulated gust with an above-canopy mean wind of 10.2 m s^{-1} . The simulated gust increases wind speeds considerably, reaching

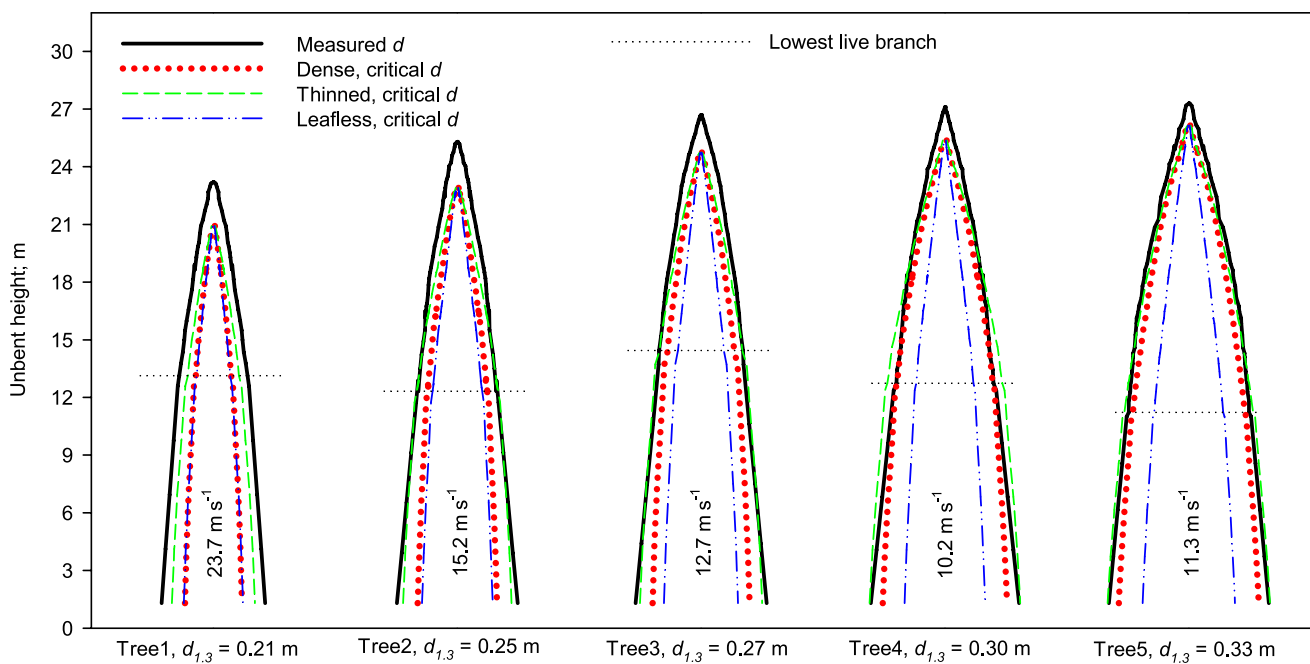


Fig. 8 The dimensions of five felled tree trunks (solid black) and dimensions sufficient to withstand wind and gravity (dotted and dashed lines) in a meteorological situation that causes a mean wind above the canopy of the dense stand (w) of 10.2 m s^{-1} , which is the critical speed that nearly breaks Tree4. The heights on vertical axis

and diameters on the horizontal axis are not proportional. Diameters at a height of 1.3 m are given at the bottom. The critical above-canopy wind speed for the dense stand is indicated inside the trunks. The lowest living branches were at heights of 11.2–14.5 m

34.2 m s^{-1} above-canopy (height of 29.25 m) and decreasing downwards as shown in Fig. 5c, with a speed of 25.9 m s^{-1} in the upper part (height of 21.75 m) of the storm-bent main canopy and 5.6 m s^{-1} in the lower part (height of 12.75 m).

The above-canopy mean wind speed in the thinned stand is surprisingly similar to that above the dense stand, and rounds to the same 10.2 m s^{-1} in the equivalent meteorological situation and is slightly weaker in the strongest gust at 33.2 m s^{-1} . However, the winds are stronger within the canopy, and for all except Tree1, greater diameters would have been needed to resist breaking (Fig. 8), indicating that thinnings increase the risk of stem breakage.

The wind simulation for a leafless stand resulted in an above-canopy mean wind speed of 14.6 m s^{-1} (gust 36.3 m s^{-1}) in the same meteorological situation as discussed above and the wind penetrated the stand much more (Fig. 5c). A significantly smaller diameter for all trees and along all heights would be sufficient in this situation (Fig. 8), as sail areas of the trees decreased.

Discussion

We developed a novel approach to model bending moments of storm-bent trees caused by wind and gravity and applied this to an unthinned middle-aged *Picea* stand that originated

from planted seedlings. We focused on winds that break the weakest segments and observed a close match of modelled and the actual diameters along other segments their trunks for most of the trees (Fig. 8). Therefore, we may conclude that these bending moments are probably important in determining trunk diameter and shape, but we are unable to compare importance of alternative determinants of tree size such as sap transport. The relatively small contribution of a tree's own mass (Fig. 7) indicates that, if to simplify only gravity or wind can be included in the modelling, wind would probably be a better choice, even in a dense unthinned stand (e.g. Larjavaara 2010) with small sail areas relative to fresh masses. The studied trees were probably much closer to elastic buckling than plants in the dataset of Niklas (1994) and may be close to bending even in windless conditions due to the extra weight of snow.

Our simulated winds may be compared to those within (at a height of 9 m) and above (at a height of 23 m) a 16-m tall *Pinus sylvestris* stand during a summer microburst that toppled over trees approximately 300 m from the wind measurements (Järvi et al. 2007). The microburst caused one-minute mean wind speeds of ca. 14 m s^{-1} above and 5 m s^{-1} within the canopy. The above-canopy speed is close to the winds that our five trees can resist, with the exception of Tree1 (Fig. 8). Furthermore, the wind speed within the relatively sparse *Pinus* canopy corresponds to values

that may have been expected based on our wind profiles (Fig. 5c). However, the variation in wind speed measured by Järvi et al. (2007) was much lower, as their “instantaneous” above-canopy wind speeds peaked at only just above 20 m s^{-1} . This may indicate that our biomechanical computations overestimated the resistance of trees to bending forces. However, as the damaged *Pinus* trees were located some distance away from the anemometers, it is likely they experienced much stronger wind speeds than recorded at the specific location of the sensors.

Our objectives were to understand more about trunk taper based on wind and the risks that trees potentially take, whereas the majority of research linking taper, wind and risks inversely attempt to estimate risks from taper and winds (Gardiner et al. 2008). The demand for advice from forest managers is substantial both in plantations (Gardiner et al. 2016) and urban setting (Sæbø et al. 2003), and advances have been impressive (Gardiner et al. 2019). However, a pessimist may argue that scientists will never be “wiser” than an acclimated tree in “understanding” the local wind profile and risks caused by extreme gusts. From an evolutionary perspective, trees balance between having their trunks breaking in a storm and overinvesting in trunk tissue and being overtopped by their neighbours growing faster. A winning strategy optimally balancing between the deadly “ditches” on both sides depends on the position of the other ditch. Hence, in a situation with fierce competition and high likelihood of being overtopped by neighbours, such as in middle-aged dense plantations, the risk on trunk breakage in a storm is increased. Therefore, the most fruitful theoretical (not just statistical and descriptive) way to estimate the risk of trunk breakage may be based on competition for height from an evolutionary perspective. Physical modelling, such as that used in this article but inversely, is more promising for trees in situations that have not acclimated to, e.g. after their neighbours have been harvested (e.g. Peltola et al. 1999).

In our simulation of the strongest gust, it is remarkable how a *Picea abies* monoculture, characterised by long, conical, and slender crowns, forms a relatively thin layer of dense sail area of sail area at approximately 18 m above ground during a gust. To support a larger leaf mass, a tree needs to build a thicker trunk to resist the wind drag and gravity acting on this additional mass. Even without additional height when unbent, the additional diameter reduces bending and the storm-bent height increases. Because trees with thicker trunks are normally also taller, they have greater wind drag caused by bending moments because of greater sail area and this area being located in greater winds because of greater unbent height but also reduced bending. The thicker trees in a stand are responsible for blocking wind and protecting the smaller “biomechanical free-riders”. This mechanism operates as a balancing force, i.e. negative feedback, in stand

development, thanks to which height growth of shorter trees is boosted relative to the tall ones.

Tree1 is much thicker and Tree2 is to some extent thicker than they need to be to resist the modelled gust. Their positions in the canopy may have weakened rapidly, leaving their thicker trunks as a legacy of a time when they needed strength for a larger leaf area, but biomechanically they would not then need new diameter growth. Also, the transport-focused perspective offers an alternative explanation. When trees become suppressed in the canopy, they rapidly lose their lower branches and their crown length grows more slowly than their height, reducing their crown ratio. This change in growth pattern may be regarded as an evolutionary response to competition for light (e.g. Mäkelä 1985). In this process, active wood, i.e. sapwood, related to the receding branches loses its connection to the foliage and gradually turns into inactive heartwood. Empirical evidence and eco-evolutionary balance theories suggest that active wood area and foliage area are in balance with each other (Chiba et al. 1988; Mäkelä and Valentine 2006; Shinozaki et al. 1964). Losing the active wood related to the receding branches, therefore, creates a need for new diameter growth to build new sapwood, as the existing inactive wood can no longer be used for water transport. If we assume that all these selective pressures, related to biomechanics, water transport, and competition for light, are present in the tree population, then our results suggest that biomechanics dominate trunk dimensions of dominant trees (see also Mäkelä and Valentine 2006), while with suppressed trees the balance has possibly shifted from biomechanics towards sap transport. Another reason for our result that smaller trees have larger diameters than apparently necessary may be that our wind model severely overestimates the steepness of the vertical wind profile. It is also possible that suppressed trees occasionally experience unusually strong gusts that penetrate the canopy but which was not our “strongest gust” due to our sampling, and are therefore seemingly overbuilt. Suppressed trees could also be prepared for surviving the gust that breaks their suppressors. These questions could be studied by analysing how tree size influences mortality in storms.

The tops of all five trees appear overbuilt. We can try to understand this by comparing small trees of the same height that may initially seem to have nearly identical biomechanical constraints. Coincidentally, both small *Picea* trees and residue treetops have commonly been used as Christmas trees in Finland and are easy to differentiate even from a distance. Treetops need to resist much stronger winds but can streamline easier, as their bases are tilted thanks to the bending lower trunk. Probably most importantly, treetops cannot rely on the “shrub strategy” of bending all the way to the ground to remain unharmed (Larjavaara 2015). This makes small trees resistant to the strongest winds and heaviest snow loads as they can bounce back after a gust has

passed or the snow has melted. Treetops, however, cannot rely on ground support during gusts, but this is probably not a problem for the well-streamlined tops of *Picea abies* (Fig. 6). Snow weight, which may be significant in the region especially when temperatures are close to freezing or when direct condensation occurs on trees, is a possible reason for the seemingly overbuilt tops in our dataset (Peltola et al. 1999).

We focused on an unthinned boreal monoculture, i.e. nearly the simplest stand imaginable—only treetops could potentially have been easier to understand in an ice-free climate. We nevertheless had to make many simplifying assumptions. The risk of resonating with the wind is a serious concern in designing structures, such as bridges, and the risk of trees swaying with a pulsing wind has often been the focus of trunk breakage literature (Niklas and Spatz 2012). However, air flow modelling does not seem to create such winds (Gardiner et al. 2019) and is rarely seen in dozens of videos found on the Internet that depict uprooting or trunk breakage (ML personal observation), but scientific evidence is still needed (Moore et al. 2018). Similarly, torsional forces have attracted some attention (Skatter and Kucera 2000), but it is likely that strengthening the trunks to resist twisting could be achieved easier by adjusting wood characteristics without increasing trunk diameters. Uprooting possibly being more common than trunk breakage is one argument against the biomechanical modelling of trunks, but this does not rule out the importance of trunk dimensions on trunk failure. In their evolutionary history, trees have probably balanced the risks of uprooting and trunk breakage depending on the level and variability of risks and on the cost of strengthening them. Our assumptions that the same level of streamlining occurs at all heights (Online Resource 1) and invariable, modulus of rupture (σ) and modulus of elasticity (E), may be far from realistic but probably do not interfere significantly with our comparison between trees and along the trunk of one tree, except perhaps in the tops which may in reality be more flexible due to juvenile wood and, therefore, e.g. the relative importance of gravity would be underestimated (Fig. 7). Choosing the value for drag coefficient (C_d) was rather arbitrary as always. Furthermore, we did not attempt to include physical contact with neighbours influencing the bending forces. Such canopy contacts may be harmful, as tree tissue may be damaged, but on the other hand, they may save a tree that is supported by a neighbour in extreme winds.

Our greatest concern relates to dealing with streamlining and the homogeneity of the sail area. We assumed 50% streamlining for branches and none for leaves (Online Resource 1). This is probably an underestimation (Peltola et al. 1999), but perhaps surprisingly it does not strongly influence this kind of analysis related to trunk diameters,

as despite streamlining reducing wind drag caused by a given wind speed, it increases wind speeds within the stand. For example, the Thinned simulation with half of the sail area removed corresponds to the Dense simulation with streamlining reducing the projected area to half its original size. This allows us to estimate the sensitivity of our results to assumptions on streamlining. Interestingly, the wind-caused bending moments were larger for two of our five trees, with 50% stronger streamlining, while they were smaller for three trees. This indicates that our results are not very sensitive to streamlining, as the increasing wind speed due to streamlining compensates for the reduced sail area. Similarly, the spatial grouping of sail area is probably important and drastically influences both winds and the drags that they cause. However, again it is possible that reduced winds for a given wind speed cause greater within-canopy winds thanks to the clustering of sail area, and their impacts may roughly even out as with the cause of streamlining.

Our approach could be utilized in several applications. Evolutionary simulations could optimize trunk dimensions by considering the benefits of being a biomechanical free-rider and relying on larger neighbour trees to withstand wind, but potentially face local extinction if all canopy species or individuals take excessive risks and rely on trunks of others not breaking. Other mechanistic modelling approaches (Kalliokoski et al. 2016), which are potentially especially valuable when optimizing forest management in changed conditions, may also benefit from the incorporation of wind- and gravity-driven trunk diameter modelling, e.g. by increasing detail in the direction pointed by Eloy et al. (2017).

Author contribution statement ML and AM developed the research idea, AK designed and implemented the data collection procedure supervised by AM, MA performed the wind simulations and wrote the first draft of its description, ML performed the other analyses, prepared the figures and wrote the first draft of the other sections, and all authors participated in producing the final version of the manuscript.

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Declarations

Conflict of interest The authors declare that they have no conflict of interest.

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