Stochastic frontiers or regression quantiles for estimating the self-thinning surface in higher dimensions?

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Received: 5 November 2019 / Accepted: 12 January 2020 / Published online: 13 August 2020
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Abstract Stochastic frontier analysis and quantile regression are the two econometric approaches that have been commonly adopted in the determination of the self-thinning boundary line or surface in two and higher dimensions since their introduction to the field some 20 years ago. However, the rational for using one method over the other has, in most cases, not been clearly explained perhaps due to a lack of adequate appreciation of differences between the two approaches for delineating the self-thinning surface. Without an adequate understanding of such differences, the most informative analysis may become a missed opportunity, leading to an inefficient use of data, weak statistical inferences and a failure to gain greater insight into the dynamics of plant populations and forest stands that would otherwise be obtained. Using data from 170 plot measurements in even-aged Larix olgensis (A. Henry) plantations across a wide range of site qualities and with different abundances of woody weeds, i.e. naturally regenerated non-crop species, in northeast China, this study compared the two methods in determining the self-thinning surface across eight sample sizes from 30 to 170 with an even interval of 20 observations and also over a range of quantiles through repeated random sampling and estimation. Across all sample sizes and over the quantile range of 0.90 ≤ τ ≤ 0.99, the normal-half normal stochastic frontier estimation proved to be superior to quantile regression in statistical efficiency. Its parameter estimates had lower degrees of variability and correspondingly narrower confidence intervals. This greater efficiency would naturally be conducive to making statistical inferences. The estimated self-thinning surface using all 170 observations enveloped about 96.5% of the data points, a degree of envelopment equivalent to a regression quantile estimation with a τ of 0.965. The stochastic frontier estimation was also more objective because it did not involve the subjective selection of a particular value of τ for the favoured self-thinning surface from several mutually intersecting surfaces as in quantile regression. However, quantile regression could still provide a valuable complement to stochastic frontier analysis in the estimation of the self-thinning surface as it allows the examination of the impact of variables other than stand density on different quantiles of stand biomass.

Keywords Larix olgensis · Normal-half normal distribution · Site productivity · Woody weeds · Sample size · Quantile selection
Introduction

In plant population and community ecology, self-thinning refers to the progressive decline in stand density (i.e., number of plants per unit area) through competition-induced mortality as individual plants grow larger and collectively accumulate more biomass while competition intensity increases and site resources become limiting for all to survive (Harper 1977; Begon et al. 2006). A series of Japanese studies on density effects, intraspecific competition and self-thinning among plants over a decade soon after the Second World War gave rise to the self-thinning rule for even-aged plant populations (Kira et al. 1953; Koyama and Kira 1956; Shinozaki and Kira 1956; Yoda et al. 1957, 1963). The rule delineates a density-dependent upper frontier of mean plant mass or total stand biomass for even-aged pure stands at full site occupancy in a given environment with a power function:

$$B = KN^{\beta_i},$$  

where $B$ represents either mean plant mass or the total stand biomass per unit area, $N$ is stand density, $K$ is a species-specific and environment-specific constant, and $\beta_i$ is the self-thinning exponent, postulated to be $-1.5$ for mean plant mass or $-0.5$ for total stand biomass (Yoda et al. 1963; White and Harper 1970; White 1980, 1981, 1985; Westoby 1984; Whittington 1984). When plotted on a log–log graph, this power function becomes the self-thinning boundary line that regulates the growth trajectories of even-aged plant populations from herbs to trees in the two-dimensional space of log transformed $B$ and $N$ (Gorham 1979; Norberg 1988; Pretzsch 2002). For trees in forest stands managed for timber production in particular, the quadratic mean diameter ($D_m$), an easily obtainable stand attribute, is often used instead of average tree mass or stem volume as a measure of average tree size in the determination of size-density relationships for stand density management. In this case, the maximum size-density relationship of Reineke (1933), which defines a species-specific upper limit of stand density for a given quadratic mean diameter on log scales, has proven to be a special case of the self-thinning boundary line (Pretzsch 2002, 2009; Pretzsch et al. 2012).

For quite some time in much of the earlier literature, it was generally understood that the self-thinning boundary line was species-specific and site-independent as reviewed by Westoby (1984). This presumed invariability with soil fertility or site productivity was largely due to the limitation of the original simplistic graphical analysis of the classic self-thinning experiment by Yoda et al. (1963), where a single boundary line was drawn visually on top of the self-thinning trajectories of horseweed (Erigeron canadensis (L.) Cronquist) populations grown at different levels of soil fertility. A statistically more rigorous analysis of the same experimental data by Bi (2004) some forty years later revealed that the intercept of the self-thinning boundary line increased with soil fertility. This revelation effectively extended the originally presumed species-specific two-dimensional boundary line into a three-dimensional self-thinning surface over a gradient of soil fertility. Such a surface was also observed in even-aged forest stands of both coniferous and broadleaf tree species (Bi 2001; Weiskittel et al. 2009; Zhang et al. 2013). In addition to soil fertility or site productivity, other environmental and climatic variables as well as stand type and history have been found to influence the self-thinning boundary line (Weiskittel et al. 2009; Zhang et al. 2013; Brunet-Navarro et al. 2016; Condés et al. 2017; Kweon and Comeau 2017; Andrews et al. 2018). Furthermore, attempts have been made to delineate the boundary line for uneven-aged or mixed-species stands (Puetmann et al. 1992; Woodall et al. 2003, 2005; Ducey and Knapp 2010; Long and Shaw 2012; Rivoire and Le Moguèdec 2012; Reyes-Hernandez et al. 2013; Brunet-Navarro et al. 2016; Andrews et al. 2018; Bravo-Oviedo et al. 2018; Quiñonez-Barraza et al. 2018; Salas-Eljatib and Weiskittel 2018; Weiskittel and Kuehne 2019; Kimsey et al. 2019; Herberich et al. 2020). These developments have further extended the three-dimensional self-thinning surface exemplified by Bi (2001, 2004) into higher dimensions:

$$B = KN^{\beta_i}X_2^{\beta_i} \ldots X_l^{\beta_i},$$  

where $X$’s are other independent variables indexed by $i = 2, ..., l$ in addition to stand density.

The delineation of the two-dimensional self-thinning boundary line has mostly in the past relied heavily on either visual or more systematic selection of data points that lie close to a perceived or arbitrarily determined upper boundary (Bi and Turvey 1997; Solomon and Zhang 2002; Sun et al. 2010; Deng and Li 2014; Brunet-Navarro et al. 2016; Ge et al. 2017).

Only the selected data are then used to formally delineate the perceived boundary line through traditional methods such as least squares regression or principle component analysis (e.g., Bi and Turvey 1997; Zhang et al. 2005; Marchi 2019). However, no matter how data selection is done, some degree of subjectivity cannot be avoided, resulting in a certain lack of objectivity in the estimated self-thinning boundary line. To overcome this problem, two econometric methods, namely stochastic frontier analysis (SFA) and quantile regression (QR), were introduced to the estimation of the self-thinning boundary line by Bi et al. (2000), Bi (2001, 2004), Cade and Guo (2000) and Zhang et al. (2005). These two methods have since become the most commonly used techniques for estimating the self-thinning boundary line or surface in two and higher dimensions. Stochastic frontier
analysis has been performed in some studies (e.g., Montigny and Nigh 2007; Weiskittel et al. 2009; Charru et al. 2012; Reyes-Hernandez et al. 2013; Kweon and Comeau 2017; Quínonez-Barraza et al. 2018; Kimsey et al. 2019), while quantile regression has been carried out in others (e.g., Vospernik and Sterba 2015; Xue et al. 2015; Riofrío et al. 2016; Condés et al. 2017; Andrews et al. 2018). In far fewer studies, both methods have been used (Sun et al. 2010; Zhang et al. 2013; Salas-Eljatib and Weiskittel 2018).

Among these studies, the number of independent variables other than stand density that were incorporated into the model specification generally in the form of Eq. 2 has varied between one and four (e.g., Weiskittel et al. 2009; Reyes-Hernandez et al. 2013; Zhang et al. 2013; Kweon and Comeau 2017). Concomitantly, the number of data points ranged from tens to hundreds and thousands (e.g., Comeau et al. 2010; Sun et al. 2010; Zhang et al. 2013; Condés et al. 2017; Kweon and Comeau 2017), and even to more than ten thousand in the extreme case (Vospernik and Sterba 2015). However, the rational for using stochastic frontier analysis or quantile regression has in most cases not been clearly explained and there appears to be a general lack of adequate appreciation of the differences between the two methods as statistical estimators for delineating the self-thinning surface. Without a thorough understanding of such differences, the best and most informative method for analysing the valuable empirical data at hand may become a missed opportunity, leading to an inefficient use of data, a failure to gain a greater insight into stand dynamics that would otherwise be obtained from the analysis, and more practically, leading to a suboptimal stand density management scheme for even-aged forest plantations. Even more seriously, the validity of results may be called into question if statistical inferences are not made on a valid or reliable basis.

The two methods were briefly compared by Zhang et al. (2005) in a short research note for estimating the self-thinning boundary line using 262 data points from even-aged pure stands of eastern white pine (Pinus strobus L.) as an example. Following this initial work, two brief comparisons of the methods were reported by Sun et al. (2010) and Zhang et al. (2013). The former used 160 observations from 10 experimental plots that were repeatedly measured in a spacing trial of Chinese fir (Cunninghamia lanceolata (Lamb) Hook.), and the latter analysed data of ponderosa pine (Pinus ponderosa Lawson & C. Lawson) from 109 long-term research plots and 59 temporary inventory plots across California. In addition, the study by Socha and Zasada (2014) on the allometric relationships between density and tree dimensions in birch stands on post-agricultural lands in Poland also contained some comparative results. A further and more considered comparison of these two methods was made in a recent study by Salas-Eljatib and Weiskittel (2018), particularly for determining the maximum stand density, i.e. the maximum number of trees at a given reference average diameter, using 178 observations from 130 plots in mixed Nothofagus forests.

All these comparisons have been made in a two-dimensional space, except for Zhang et al. (2013) which included site index in the analysis and were limited to the comparative positions of the estimated self-thinning boundary line. The methodological attributes and statistical properties of the two approaches that researchers need to carefully consider before choosing one or the other for their particular research objectives and for the nature and size of data at hand remain to be further evaluated, particularly when the analysis goes beyond two dimensions. This paper aimed to further evaluate and highlight the pros and cons of these two methods particularly for estimating the self-thinning surface in higher dimensions through a more detailed comparison across a range of data sizes, (i.e. number of data points), to help researchers make the best choice for their analyses. In doing so, the two econometric methods were briefly reviewed first to provide the minimum but most essential details on their statistical properties as parametric boundary estimators. They were then compared in the estimation of the self-thinning surface in higher dimensions using cross-sectional data from even-aged L. olgensis plantations across a range of sample sizes through repeated random sampling with replacement and model estimation. Finally, some practical guidance for choosing the best method is provided for researchers to address their particular research questions.

Materials and methods

Stochastic frontier analysis and quantile regression

Stochastic frontier analysis originated from the economic concept of a production function which relates an upper boundary of maximum attainable output to any given quantities of a set of inputs in a production process (see Farrell 1957; Forsund et al. 1980; Schmidt 1985). This econometric approach was first proposed some 40 years ago by Aigner et al. (1977), Battese and Corra (1977) and Meusen and van den Broeck (1977). Over the past four decades, it has been intensively studied and applied to the estimation and measurements of productivity and technical efficiency in economics and a very wide range of other fields (see Kumbhakar and Lovell 2000; Fried et al. 2008; Kumbhakar et al. 2018 for major reviews). The model specification commonly used for a stochastic frontier function takes the form of what is known in economics as the Cobb–Douglas production function with a composed error structure:

\[
Y = AX_1^{\theta_1}X_2^{\theta_2} \ldots X_k^{\theta_k} e^\varepsilon e^{-\eta}
\] (3)
where \( Y \) is the dependent variable which is called output in econometrics, \( X \)'s are inputs, i.e., independent variables, and \( \beta \) and \( \alpha \)'s are parameters, \( e^v \) and \( e^u \) are two error components. Taking logarithms, the model becomes

\[
y = \alpha + X\beta + \varepsilon
\]

(4)

where \( y = \ln Y, \alpha = \ln A, \ln \) denotes natural logarithm, \( X \) is a vector of log-transformed independent variables, and \( \beta \) is a vector of parameters. The composed error term, \( \varepsilon = v - u \), is a compound random variable with two components and each is assumed to be independently and identically distributed across observations. The first error component \( v \) is assumed to be normal with zero mean and constant variance \( \sigma_v^2 \) and is intended to capture the effects of random factors external to producers on the frontier. The second random variable \( u \) is non-negative and is specified to capture the effects of technical efficiency of producers. Because \( u \geq 0 \), it was assigned a half-normal distribution and an exponential distribution in the three seminal papers published in 1977, effectively giving the stochastic frontier function a normal-half normal and a normal-exponential specification. By allowing \( u \) to follow a truncated normal distribution \( N^+(\mu, \sigma_u^2) \) obtained by truncating at zero of the normal distribution with mean \( \mu \) and variance \( \sigma_u^2 \), Stevenson (1980) generalised the normal-half normal model into a normal-truncated normal specification for a more flexible representation of the patterns of technical efficiency \( u \) in the data. The flexibility is realised because the mode of \( u \) in the truncated normal distribution is no longer arbitrarily set at zero, but is allowed to be estimated together with other parameters in the stochastic frontier model. Unlike the half-normal and exponential densities which always have a mode at 0, the truncated-normal density has a mode at 0 only when \( \mu \leq 0 \), but a mode at \( \mu \) otherwise.

Because \( u \geq 0 \) by specification, the composed error term, \( \varepsilon = v - u \), is asymmetric and its expectation, \( E(\varepsilon) = -E(u) \leq 0 \), as \( v \) and \( u \) are distributed independently of each other. As shown in Kumbhakar and Lovell (2000):

\[
E(\varepsilon) = -\sigma_u \sqrt{2/\pi}
\]

and

\[
V(\varepsilon) = \sigma_v^2 + \sigma_u^2(1 - 2/\pi)
\]

(6)

for the normal-half normal model, and

\[
E(\varepsilon) = -\sigma_u
\]

and

\[
V(\varepsilon) = \sigma_v^2 + \sigma_u^2
\]

(7)

(8)

for the normal-exponential model. For the normal-truncated normal model:

\[
E(\varepsilon) = -\frac{\mu a}{2} - \frac{\sigma_u a}{\sqrt{2\pi}} \cdot \exp\left\{-\frac{1}{2} \left( \frac{\mu}{\sigma_u} \right)^2 \right\}
\]

(9)

and

\[
V(\varepsilon) = \sigma_v^2 + \frac{\mu^2 a^2}{2}(1 - \frac{a}{2}) + \frac{a}{2} \left( \frac{\pi - a}{\pi} \right) \sigma_u^2
\]

(10)

where the pre-truncation mean parameter \( \mu \) becomes the mode of the truncated normal distribution, \( \sigma_v^2 \) is the pre-truncation variance of normal distribution \( N(\mu, \sigma_u^2) \), \( a = \Phi(\mu/\sigma_u)^{-1} \) and \( \Phi(\cdot) \) is the standard normal cumulative distribution function.

Similar to the half normal distribution, the exponential distribution is also a special case of the truncated normal distribution, and as such the three most commonly used distributions for \( u \) are all captured by the truncated normal distribution (Meesters 2014). Although other alternative distributions have been proposed for \( u \) in econometrics research over the past four decades, stochastic frontier models based on the three distributions, particularly the normal-half normal model, have remained the choice of applied researchers in the vast majority of empirical work (Kumbhakar and Lovell 2000; Parmeter and Kumbhakar 2014; Kumbhakar et al. 2018). The model parameters can be estimated by the maximum likelihood method, of which comprehensive descriptions can be found in Aigner et al. (1977), Stevenson (1980), Greene (1997), Kumbhakar and Lovell (2000) and Kumbhakar et al. (2018).

Another econometric approach that lends itself to boundary estimation is quantile regression introduced by Koenker and Basset (1978) at almost the same time when stochastic frontier analysis was first proposed. As reviewed by Kumbhakar et al. (2018) and shown by references therein, it has recently been embraced for the estimation of production frontiers and measurements of efficiency. Quantile regression extends the least squares regression framework, traditionally focused on the conditional mean function onto non-central parts of the conditional distribution so that the \( \tau \) th quantile \( Q_{\tau}(X) \) of the conditional probability distribution of a response variable \( Y \) given \( X \), a vector of predictor variables, can be estimated for any chosen value of \( 0 < \tau < 1 \) (Koenker and Hallock 2001; Koenker 2005, 2017). The estimator of \( Q_{\tau}(X) \) proposed by Koenker and Basset (1978) minimises the following asymmetric loss function:

\[
\arg\min_{\beta} \left\{ \sum_{Y \geq X\beta} \tau |Y - X\beta| + \sum_{Y < X\beta} (1 - \tau) |Y - X\beta| \right\}
\]

(11)

which gives positive and negative residuals the weight of \( \tau \) and \( 1 - \tau \), respectively. The minimization is achieved through linear programming as detailed by Koenker (2005)
The quantile regression estimator is consistent and asymptotically normally distributed under some regularity conditions (Koenker and Basset 1978). Under the assumption of independent and identically distributed (i.i.d.) errors, its asymptotic variance-covariance matrix can be obtained analytically through the estimation of the so-called sparsity function by differentiating a theoretical quantile function or by using a difference quotient of empirical quantiles. In the case of non i.i.d. errors, the derivation is more complicated as it uses a local estimate of the sparsity function to compute a Huber sandwich estimate (Koenker 2005; Hao and Naiman 2007). To avoid making the i.i.d. assumption, which is often not the case in quantile regression applications, and still obtain robust results, resampling methods such as bootstrapping are commonly implemented in quantile regression applications (Kocherginsky et al. 2005; Koenker 2005; Hao and Naiman 2007; Tarr 2012). As the estimator uses a well-known M-function, the absolute value function, it is robust or more resistant than the traditional least squares estimator to the influence of outliers, particularly for inner quantiles. However, the variance of quantile regression estimator is U-shaped with \( \tau \) changing from zero to one. The implication of such a pattern of estimator variance for extreme quantiles close to the data boundary is that the estimated frontiers or boundary lines can be quite variable even with a very small change in \( \tau \), particularly when data are sparse around the edges, and may even cross over each other, posing difficulties for statistical inferences and interpretation of results.

**Data**

The data set for this study came from 146 temporary square or rectangular plots established between 1991 and 2016 in even-aged larch (L. olgensis) plantations under the management of four forestry agencies across two provinces in northeast China: Mengjiagang (MJG) Forest Farm, Linkou (LK) and Dongjingcheng (DJC) Forestry Bureaux in Heilongjiang Province, and Songjianghe (SJH) Forestry Bureau in Jilin Province. These plots covered much of the natural distribution of the species in northeast China (see Peng et al. 2018), and a wide range of site quality and climatic conditions (Table 1). The exact initial planting densities of the individual plots are unknown but they were most likely to be 3300, 4400 and 6600 ind. ha\(^{-1}\), the most common planting densities for larch plantations in northeast China (Yu 2008). For planting densities at such high levels, the plot size of 0.04–0.09 ha would contain enough trees for summarizing key stand attributes and describing stand structure and dynamics, although larger plots would be preferable (see García 1998; García and Batho 2005; Wagner et al. 2010; Zhou et al. 2019). Some plots were in plantations that had been thinned many years prior to measurement as judged by stumps remaining on the forest floor, but their silvicultural histories were not recorded. Although in even-aged plantations, there were only 38 plots where larch was the

| Table 1  | Number of plots, geographical locations, plot size, age ranges, key stand attributes and climatic conditions across four forest management areas: Mengjiagang (MJG), Linkou (LK), Dongjingcheng (DJC) and Songjianghe (SJH) |
|----------|--------------------------------------------------------------------------------------------------|
| Number of plots | MJG  | LK  | DJC  | SJH  |
| Longitude (E) | 130°32′–130°52′ | 129°40′–130°34′ | 128°07′–130°02′ | 127°12′–127°50′ |
| Latitude (N) | 46°20′–46°30′ | 45°51′–45°59′ | 43°30′–44°18′ | 41°44′–42°21′ |
| Elevation (m) | 250 | 400–600 | 450–900 | 700–1000 |
| Mean max. temp. (°C) | 35.6 | 37.0 | 36.5 | 34.8 |
| Mean ann. temp (°C) | 1.3 | 2.5 | 3.0 | 2.7 |
| Mean min. temp. (°C) | −34.7 | −39.0 | −40.1 | −44.1 |
| MAR (mm) | 550 | 580 | 550–600 | 830 |
| Plot size (ha) | 0.06, 0.09 | 0.04, 0.06 | 0.06 | 0.04, 0.06 |
| Age (years) | 14–42 | 17–28 | 15–29 | 8–35 |
| MDH (m) | 12.1–24.2 | 9.7–23.3 | 14.0–34.4 | 2.2–38.5 |
| Site Index (m) | 15.0–25.1 | 15.6–22.1 | 19.6–23.2 | 10.3–20.8 |
| N (trees·ha\(^{-1}\)) | 450–3078 | 550–2767 | 1367–3078 | 1367–2450 |
| N\(_{w}\) (trees·ha\(^{-1}\)) | 0–717 | 0–450 | 0–333 | 0–1275 |
| G (m\(^2\)·ha\(^{-1}\)) | 10.4–33.9 | 6.8–21.3 | 13.4–31.9 | 1.6–37.2 |
| G\(_{w}\) (m\(^2\)·ha\(^{-1}\)) | 0–5.3 | 0–2.9 | 0–5.6 | 0–13.0 |
| Mortality (ind·ha\(^{-1}\)) | 0–667 | 0–125 | 0–33 | 0–896 |

MDH stands for mean dominant height, i.e., average height of 100 tallest trees per hectare. N and N\(_{w}\) are the stand densities of larch and woody weeds; G and G\(_{w}\) are stand basal areas of the two groups. MAR is mean annual rainfall.
only species. In the remaining plots, larch was the dominant species in both number and size, as naturally regenerated non-crop trees had emerged and mostly grown under the dominant larch, possibly reflecting the differences in previous land use and in site preparations across the plantations. Such non-crop trees in commercial plantations are usually regarded and termed as woody weeds in forest research and management (e.g., Bi and Turvey 1994; Rose et al. 2006). The woody weeds included Korean pine (Pinus koraiensis Siebold & Zucc.) and three broadleaf species: Betula platyphylla Sukaczew, Quercus mongolica Fisch. ex Ledeb. and Populus davidiana Dode. As mostly small trees, they represented less than 30% of the total number of trees and less than 30% of the total stand basal area in most plots (Table 1).

Except for 24 plots on the MJG Forest Farm which were measured twice, all plots had a single measurement for a total of 170 plot measurements from the 146 plots. For the purpose of this paper, the potential correlation between the two sets of equations. The biomass estimates of individual larch trees were summed up for each plot and converted to total aboveground stand biomass on a pro rata basis. Standing dead trees were not included in the biomass calculation.

### Deterministic model specification and estimation

The deterministic model specification extended that of Bi (2001) for estimating the self-thinning surface by incorporating stand density and basal area of non-larch species in the equation:

\[
B = KN_1^{0.12} S^{0.18} e^{0.3} W
\]

where \(B\) represents stand biomass of larch trees in kg/ha, \(K\) is a constant, \(N\) stands for the stand density of larch in trees/ha, \(\beta_1\) is the self-thinning exponent, \(S\) is a relative measure of site productivity and takes any value between 0 and 1 because it was calculated as a ratio between site index and the maximum site index of 26 m for larch plantations in northeast China, \(W\) is a relative measure of the abundance of woody weeds in the larch stands, \(\beta_2\) and \(\beta_3\) are parameters. The formulation of \(W\) was based on a still-to-be published study by the second author on self-thinning in relatively even-aged eucalypt regrowth forests in Australia. It took both the density and basal area of woody weeds in the larch stands into account:

\[
W = \sqrt{\left(\frac{N_w}{N + N_w}\right)^2 + \left(\frac{G_w}{G + G_w}\right)^2}
\]

where \(N_w\) is the density of woody weeds in trees/ha, \(G\) and \(G_w\) indicate, respectively, the basal area of larch and that of woody weeds in m²/ha.

This formulation of \(W\) was used following an exploratory analysis with the aim to overcome the collinearity that arose from the correlation between \(N_w\) and \(G_w\) if the two variables were included separately in the model specification. The ill effects of collinearity on parameter estimation and interpretation in linear regression was well recognised and described by Belsley (1991). In pure larch stands, \(W = 0\); when woody weeds are present, \(0 < W < 1\). When \(W = 0\) in Eq. (6), woody weeds have no influence whatsoever on the self-thinning surface; when \(W > 0\), however, they are expected to have a negative impact on the self-thinning surface because of their competition with larch for site resources. Varying between 0 and 0.82, the calculated \(W\) had a positively skewed distribution with a median of 0.12, a mean of 0.18, a third quartile of 0.30 and a 90th percentile of 0.46 (Fig. 1).
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Estimating the self-thinning surface as stochastic frontiers

To estimate the self-thinning surface as stochastic frontiers in the four-dimensional space, Eq. (12) was linearized by taking the natural logarithm of both sides and the linearized equation was then given a composed error term \( \varepsilon = v - u \):

\[
\ln B = k + \beta_1 \ln N + \beta_2 \ln S + \beta_3 W + \varepsilon
\]  

(14)

where \( k = \ln K \), and the other variables are as previously defined. While the first error component, \( v \), was assumed to follow \( N(0, \sigma_v^2) \) as reviewed previously, the second error component, \( u \), was considered to follow either a half normal, or an exponential or a truncated normal distribution. The combination of distributional assumptions of the two error components led to three stochastic frontier models, namely, normal-half normal (NH), normal-exponential (NE) and normal-truncated normal (NT). The three models were estimated through the maximum likelihood estimation using the Quasi–Newton method in the QLIM procedure of SAS/ETS (SAS Institute Inc. 2014). Convergence was easily reached for the first two simpler models with no more than 20 iterations. Their parameter estimates and asymptotic standard errors were compared, together with their model fit summaries. However, convergence was not attained for the more complex NT model even after 200 iterations. To overcome the problem of non-convergence and obtain valid parameter estimates, the model was repeatedly estimated 2000 times, each time using 170 observations randomly sampled with replacement from the data set. Using the parameter estimates from the converged cases out of the 2000 rounds of estimation, the mean and standard error were calculated for all model parameters and taken as their parameter estimates and standard errors. Finally, values of \( \hat{\varepsilon} \), i.e., estimated residuals of the composed error term \( \varepsilon \) in Eq. (14), were obtained for the three stochastic frontier models. Their residual frequency distributions were then plotted and examined with the aid of simple descriptive statistics including the mean, variance, skewness and kurtosis.

Fig. 1 Scatter plot matrix of stand biomass (B), stand density (N), relative site productivity index (S) and abundance of woody weeds (W) for 170 even-aged larch stands; frequency distributions of the four variables are displayed as kernel density plots in the diagonal graphic cells
Estimating the self-thinning surface as regression quantiles

Like the stochastic frontier models, Eq. 12 was similarly linearized for estimating the self-thinning surface as regression quantiles:

\[ \ln B_r = k_r + \beta_1 \ln N + \beta_2 \ln S + \beta_3 W + \varepsilon_r \]  \hspace{1cm} (15)

where \( B_r, k, \beta_1, \beta_2, \) and \( \beta_3 \) are previously defined variable and parameters corresponding to the chosen quantile \( r \). But unlike the composed error term in the stochastic frontier models, \( \varepsilon_r \) here represents the \( r \) th quantile of the error term, which is set to zero as required by the model specification of quantile regression (Hao and Naiman 2007). As such, \( r \) also indicates the proportion of nonpositive residuals, or in other words, the proportion of data points on or under the self-thinning surface as defined by the \( r \)th conditional quantile of \( \ln B_r \) for any given values of \( N, S \) and \( W \) in Eq. (15). As there were 170 data points available for estimating the self-thinning surface in this study, 10 values of \( r \) from 0.90 and 0.99 with an even increment of 0.01 were used in the QUANTREG procedure of SAS/STAT (SAS Institute Inc. 2013). For the parameter estimates associated with each value of \( r \), their approximate standard errors and 95% confidence limits were computed using the resampling method through 2000 repetitions.

Comparative performances of the two methods across sample sizes

The estimations described above used all 170 observations and so the performances of stochastic frontiers and regression quantiles could only be compared on the basis of one sample size. However, as evident from the literature reviewed previously, the sample size, i.e., the number of observations or data points, used in the estimation of the self-thinning frontiers varied from as small as 30 observations to as large as thousands, even tens of thousands of plot measurements in the extreme case (e.g. Bi 2004; Condés et al. 2017; Kimsey et al. 2019). As statistical estimators, the two methods would naturally differ in statistical efficiency and so their comparative performances can be expected to change with sample size from small to large. For a more complete and informative comparison of the two methods, the estimations described above were carried out over a range of sample sizes through repeated random sampling with replacement from the 170 observations. In doing so, the sample size varied from 30 to 170 with an even interval of 20 observations and for each sample size the sampling was repeated 2000 times. Each time, the self-thinning surface was estimated as stochastic frontiers as well as quantile regression. As there were little differences in the estimated self-thinning surface among the three stochastic frontier models, only the normal-half normal model was used in the comparison for the sake of parsimony. Considering the range of sample size, 20 values of \( r \) from 0.80 to 0.99 with an increment of 0.01 were used in the quantile regression for estimating the self-thinning surface. For each sample size, the distribution of the 2000 estimates of each parameter and its descriptive statistics were obtained for the stochastic frontier model and quantile regression over the 20 selected quantiles. These descriptive statistics included the mean, variance, coefficient of variation, skewness and kurtosis calculated as the excess kurtosis, which is three less than the standardized fourth central moment. They were compared between the stochastic frontier model and quantile regression across the range of sample sizes to evaluate the comparative performances of the two methods.

Results

The self-thinning surface estimated as stochastic frontiers

There were little differences in the estimated self-thinning intercept \( \kappa \) and slope \( \beta_1 \) in Eq. 14 among the three stochastic frontier models with different distributional specifications for the composed error term \( \varepsilon \) (Table 2). The estimated value of \( \kappa \) was 15.90 for the normal-half normal (NH) model, 15.72 for the normal-exponential (NE) model and 15.95 for

| \( \nu \) | \( k \) | \( \beta_1 \) | \( \beta_2 \) | \( \beta_3 \) | \( \mu \) | \( \sigma_r \) | \( \sigma_u \) |
|---|---|---|---|---|---|---|---|
| NH | 15.90 | -0.48 | 1.54 | -1.52 | 0.14 | 0.88 |
| | (0.62) | (0.08) | (0.25) | (0.23) | (0.04) | (0.06) |
| NE | 15.72 | -0.47 | 1.29 | -1.73 | 0.18 | 0.56 |
| | (0.55) | (0.07) | (0.23) | (0.21) | (0.03) | (0.06) |
| NT | 15.95 | -0.50 | 1.32 | -1.69 | -177.52 | 0.11 | 2.86 |
| | (0.76) | (0.10) | (0.44) | (0.35) | (5269.24) | (0.08) | (10.49) |

Unlike the two simpler models, tabulated values for the NT model were based on results from repeated sampling and estimation (see text)
the normal-truncated normal (NT) model. The estimated value of $\beta_1$ was $-0.48$ for the NH model, $-0.47$ for the NE model and $-0.50$ for the NT model. All three estimates of $\beta_1$ were significantly different from zero but not significantly different from $-0.50$, the hypothesized self-thinning slope at $a = 0.05$. The estimated value of $\beta_2$ and $\beta_3$, the two parameters associated with the relative site productivity $S$ and the abundance of woody weeds $W$, were, respectively, $1.54$ and $-1.52$ for the NH model, $1.29$ and $-1.73$ for NE model, $1.32$ and $-1.69$ for the NT model. The differences in the three estimates of the same parameter were not significant at $a = 0.05$. The standard errors of parameter estimates were comparable among the two simpler models, but they were noticeably larger for the NT model (Table 2). When the three estimated self-thinning surfaces were sliced at particular values of $S$ and $W$ and the resulting self-thinning frontiers plotted together, they were hardly visually distinguishable (Fig. 2).

The estimated value of $\sigma_v$ was smaller for the NH model than for the NE one (Table 2). Consistent with the stochastic frontier specifications, the distribution of $\hat{\epsilon}$ was negatively skewed and leptokurtic for all three models (Fig. 3). The NH model had only six positive residuals (i.e., $\hat{\epsilon} > 0$ observations above the estimated self-thinning surface) and a mean $\hat{\epsilon}$ of $-0.67$, while the NE model had 18 such residuals and a mean $\hat{\epsilon}$ of $-0.56$. With 12 positive residuals and a mean $\hat{\epsilon}$ of $-0.59$, the NT model was in the middle of the two simpler models. The distribution of $\hat{\epsilon}$ for the NH model had a smaller variance and was also slightly less skewed and less leptokurtic than the other two stochastic frontier models for which the variance, skewness and kurtosis of $\hat{\epsilon}$ were almost the same.

During parameter estimation, convergence was easily reached for the two simpler NH and NE models, but the convergence rate of the NT model was less than $58\%$, with only 1156 cases attaining convergence out of the 2000 repeated estimations. Among these converged cases, about $7.4\%$ of the estimated mean of the pre-truncation normal distribution $\hat{\mu}$ were positive, but only within a narrow open interval of $(0, 0.81)$. For the remaining cases, $\hat{\mu}$ was either a negative or an unreasonably large negative number. The lower $25\%$ values of $\hat{\mu}$ lay below $-12$ and even dropped to a 6-digit negative number in the extreme case. The estimates of the pre-truncation standard deviation $\hat{\sigma}_\epsilon$ were correspondingly large or extremely large, ranging from a single digit to a three-digit number. Pearson’s correlation
coefficient between $\hat{\mu}$ and $\hat{\sigma}_u^2$ was almost $-1$, indicating a close negative linear relationship. Among the unconverged cases, the values of $\hat{\mu}$ and $\hat{\sigma}_u$ reached when optimization was terminated after a large number of iterations were even larger negative and positive numbers than those in the lower 25% of the estimates of the converged cases as described above.

The self-thinning surface estimated as regression quantiles

Parameter estimates for the self-thinning surface obtained through quantile regression changed systematically as $\tau$ increased from 0.80 to 0.99 (Table 3). Among the 20 values of $\tau$, the parameter estimates and their standard errors were

| $\tau$ | $k$ | $\beta_1$ | $\beta_2$ | $\beta_3$ | $\hat{\mu}$ | $\hat{\sigma}_u$ |
|-------|-----|-----------|-----------|-----------|-------------|----------------|
| 0.80  | 15.70 | -0.48 | 1.37 | -1.75 | 0.90 | 16.22 | -0.56 | 0.83 | -1.99 |
|       | (0.73) | (0.10) | (0.35) | (0.31) | (1.01) | (0.15) | (0.33) | (0.35) |
| 0.81  | 16.20 | -0.55 | 1.22 | -1.92 | 0.91 | 16.11 | -0.54 | 0.85 | -2.01 |
|       | (0.75) | (0.10) | (0.34) | (0.31) | (1.13) | (0.16) | (0.35) | (0.38) |
| 0.82  | 16.12 | -0.54 | 1.26 | -1.82 | 0.92 | 15.84 | -0.49 | 0.87 | -2.00 |
|       | (0.78) | (0.11) | (0.33) | (0.32) | (1.18) | (0.16) | (0.40) | (0.42) |
| 0.83  | 16.12 | -0.54 | 1.26 | -1.82 | 0.93 | 15.50 | -0.45 | 0.91 | -1.99 |
|       | (0.80) | (0.11) | (0.33) | (0.31) | (1.27) | (0.18) | (0.38) | (0.44) |
| 0.84  | 16.31 | -0.57 | 1.18 | -1.88 | 0.94 | 14.88 | -0.36 | 0.84 | -1.92 |
|       | (0.84) | (0.12) | (0.32) | (0.32) | (1.39) | (0.19) | (0.42) | (0.51) |
| 0.85  | 16.36 | -0.57 | 1.17 | -1.83 | 0.95 | 15.06 | -0.39 | 0.75 | -1.98 |
|       | (0.87) | (0.12) | (0.31) | (0.32) | (1.57) | (0.22) | (0.45) | (0.57) |
| 0.86  | 16.39 | -0.58 | 1.13 | -1.85 | 0.96 | 15.07 | -0.38 | 0.90 | -1.82 |
|       | (0.89) | (0.12) | (0.30) | (0.31) | (1.78) | (0.24) | (0.52) | (0.64) |
| 0.87  | 16.31 | -0.56 | 1.12 | -1.87 | 0.97 | 15.95 | -0.51 | 0.57 | -2.17 |
|       | (0.92) | (0.13) | (0.31) | (0.31) | (2.31) | (0.31) | (0.68) | (0.76) |
| 0.88  | 16.27 | -0.56 | 1.03 | -1.91 | 0.98 | 15.14 | -0.39 | 0.94 | -1.48 |
|       | (0.97) | (0.14) | (0.30) | (0.32) | (3.19) | (0.44) | (1.68) | (1.08) |
| 0.89  | 16.19 | -0.55 | 0.93 | -1.94 | 0.99 | 13.28 | -0.13 | 1.11 | -0.89 |
|       | (1.01) | (0.14) | (0.32) | (0.33) | (8.14) | (1.11) | (3.11) | (2.68) |
closest to the stochastic frontier estimates when $\tau = 0.80$. As $\tau$ increased from 0.81 to 0.91, the estimated self-thinning intercept $k_2$ in Eq. (9) varied within a narrow range between 16.11 and 16.36, while the estimated self-thinning slope $\beta_2$, also varied narrowly between $-0.54$ and $-0.58$. At the same time, the estimates of $\beta_3$, generally decreased from values greater than 1.20 to less than 0.90, while the estimates of $\beta_3$, varied between $-1.82$ and $-2.01$. As $\tau$ further increased beyond 0.91 to 0.99, the parameter estimates for the self-thinning surface exhibited much larger variations. The estimated value of $k_3$ varied mostly between 15.00 and 16.00 until it dropped down to 13.28 when $\tau$ reached 0.99. Correspondingly, the estimated value of $\beta_{1r}$ varied between $-0.36$ and $-0.51$ before it became a much flatter slope of $-0.13$ when $\tau = 0.99$. The estimated value of $\beta_{2r}$ varied mostly within 0.57 and 0.94 before reaching a much greater value of 1.11 when $\tau = 0.99$, while that of $\beta_{3r}$ varied within the range of $-2.17$ to $-1.48$ before a large increase to $-0.89$ at the most extreme quantile. Over the more extreme quantiles, a slight change in $\tau$, even as small as 0.01, could lead to relatively large changes in parameter estimates. As expected, the standard errors of parameter estimates became increasingly larger as $\tau$ increased beyond 0.90 to 0.99 (Table 3). Because of the large standard errors, the estimated self-thinning slope $\beta_{1r}$ was not significantly different from zero at $\alpha = 0.05$ when $\tau$ ranged from 0.95 and 0.99, and also $\beta_{2r}$, the parameter associated the relative measure of site productivity $S$. Further detailed numerical and graphical examinations of the self-thinning surfaces estimated at different values of $\tau$ revealed that they intersected each other within the empirical data space delineated in Fig. 1, particularly for high and extreme quantiles with $\tau \geq 0.90$.

**Comparative performances of the two methods across sample sizes**

The repeated sampling and estimation of the self-thinning surface through the normal-half normal stochastic frontier model revealed systematic changes in the variability, skewness and kurtosis, but not the mean of parameter estimates, across sample sizes. The means of 2000 values of $\hat{k}$, $\hat{\beta}_1$, $\hat{\beta}_2$ and $\hat{\beta}_3$ each changed little with sample size and were almost identical to the estimated parameters of the normal-half normal model presented in Table 2 when sample size reached 170 (Fig. 4). The absolute values of coefficient of variation (CV) of the four parameters were expectedly the largest for the smallest sample size of 30, being at least or more than twice the values for the largest sample size of 170 (Fig. 5). As sample size increased, the absolute value of CV decreased rapidly until sample size reached 90 and from there onwards the decline became more gradual for all four parameters. Concomitantly, the 90% confidence interval of parameter estimates (i.e. interval containing the middle 90% of values of parameter estimates) was increasingly narrower (Fig. 4). The skewness of $\hat{k}$ and $\hat{\beta}_1$ were both close to zero for the smallest sample size of 30 and then diverged, with the former showing a slight increase and the latter a slight decrease as sample size became larger (Fig. 6). The skewness of $\hat{\beta}_2$ decreased linearly with sample size, from 0.33 and 0.50 at sample size 30 and 50 to $-0.26$ when sample size reached 170. Contrastingly, the skewness of $\hat{\beta}_3$ was consistently positive across sample sizes and did not exhibit a systematic pattern of change, falling between 0.31 and 0.42 for all sample sizes except for the value of 0.21 for the largest sample size. In comparison to the skewness, the pattern of change in the kurtosis with sample size resembled a flattened and elongated tick mark for all four estimated parameters, with the largest sample sizes being the most leptokurtic (Fig. 7). For $\hat{k}$, $\hat{\beta}_1$ and $\hat{\beta}_3$, the kurtosis was mostly positive, falling between 0 and 1, but for $\hat{\beta}_2$, it was within $\pm 0.24$ among the eight sample sizes. As expected, $\hat{k}$ and $\hat{\beta}_1$ were closely and negatively correlated and their Pearson’s correlation coefficient ($r$) was almost $-1$. In contrast to this negative correlation, $\hat{k}$ was positively correlated with $\hat{\beta}_2$ ($r = 0.81$) and, to a much lesser extent, with $\hat{\beta}_3$ ($r = 0.46$).

In comparison to stochastic frontiers, the self-thinning surface estimated as regression quantiles through repeated random sampling with replacement across the eight sample sizes exhibited both similarities and differences in the distributional characteristics of its four parameters. The degree of differences varied with sample size and also depended upon the selected quantile $\tau$. The estimated values of the self-thinning intercept and slope, $\hat{k}$, $\hat{\beta}_1$, and $\hat{\beta}_{1r}$, were very similar to the values of $\hat{k}$ and $\hat{\beta}_1$ estimated by the stochastic frontier model for sample sizes smaller than 100 and also for larger samples but only when $\tau < 0.90$. As $\tau$ and sample size increased, $\hat{k}_r$ became increasingly smaller than $\hat{k}$ and at the same time $\hat{\beta}_{1r}$ became increasingly greater than $\hat{\beta}_1$. When $\tau$ reached 0.99 at the largest sample size of 170, the differences were the largest and became marginally significant statistically as $\hat{k}_r$ and $\hat{\beta}_{1r}$ just touched the 90% confidence limits of $\hat{k}$ and $\hat{\beta}_1$ (Fig. 4). Unlike the similarities that $\hat{k}_r$ and $\hat{\beta}_{1r}$ exhibited to $\hat{k}$ and $\hat{\beta}_1$, the estimated values of $\hat{\beta}_{2r}$ and $\hat{\beta}_{3r}$ were consistently smaller than those of $\hat{\beta}_2$ and $\hat{\beta}_3$ over the range of $\tau$ across all sample sizes except for the extreme quantiles with $\tau \geq 0.98$ at large sample sizes in the case of $\hat{\beta}_3$.

Although $\hat{k}$ and $\hat{\beta}_{1r}$ were similar in value to $\hat{k}$ and $\hat{\beta}_1$, their variabilities were much greater and therefore the 90% confidence intervals were much wider than that of $\hat{k}$ and $\hat{\beta}_1$ particularly for high and extreme quantiles with $\tau \geq 0.90$. For any given quantile, the variability of $\hat{k}$ and $\hat{\beta}_{1r}$ decreased with sample size, but the decrease was much less steeper for high and extreme quantiles (Fig. 4). The other two parameters, $\hat{\beta}_{2r}$ and $\hat{\beta}_{3r}$, also exhibited a similar pattern of change.
in variability, but only less prominently. The skewness and kurtosis of the quantile regression parameter estimates were quantile-dependent and showed a much greater magnitude of variation than that of the stochastic frontier parameter estimates across the eight sample sizes (Figs. 6, 7). As sample size increased, the quantile-dependent variation was increasingly amplified, with skewness swinging between positive and negative and kurtosis oscillating from leptokurtic to platykurtic. Similar to $\xi$, $\tilde{\beta}_1$, $\tilde{\beta}_z$, and $\tilde{\beta}_t$, were also closely and negatively correlated across sample sizes with correlation coefficient ($r$) varying from $-0.85$ to a value approaching $-1.00$, except for the smallest sample size where the correlation became weak and turned positive with an $r$ of 0.19.

**Fig. 4** A multi-panel display of mean and 90% confidence limits of 2000 estimates for each of the four parameters of the self-thinning surface estimated through the normal-half normal stochastic frontier model (blue lines) and through quantile regression over 20 evenly spaced $\tau$ values from 0.80 to 0.99 (red curves) across the eight sample sizes as denoted in the top stripe of each column.

**Discussion**

Since their introduction to the estimation of the self-thinning boundary line some 20 years ago by Bi et al. (2000), Bi (2001, 2004), Cade and Guo (2000) and Zhang et al. (2005), stochastic frontier analysis and quantile regression have become the two most commonly adopted econometric approaches by ecologists and forest biometricians in the determination of the self-thinning frontier or surface in two and higher dimensions. The two approaches were both motivated in their early development by the desire to determine an unobservable upper boundary of maximum attainable output for a given vector of inputs in a production process.
and subsequently to evaluate and benchmark the efficiencies of individual production units against the production frontier (Kumbhakar and Lovell 2000; Parmeter and Kumbhakar 2014; He 2017; Kumbhakar et al. 2018). While stochastic frontier analysis has so far been applied predominantly in economics and related fields (Kumbhakar et al. 2018), quantile regression, as a natural extension of the traditional least squares regression, has enjoyed a much wider audience and has rapidly become a more general method in applied statistics (Koenker 2005, 2017; Hao and Naiman 2007; Davino et al. 2013). Although both approaches can serve the same purpose of boundary delineation in applied statistics far beyond econometrics (e.g., Brandt 2016; Rosenberg et al. 2017), the statistical thinking and reasoning behind them are quite different. An adequate appreciation of the differences between the two approaches would be essential for researchers to select the best method to estimate the self-thinning surface, particularly in high dimensions, for their data at hand and to gain a greater insight into how variables other than stand density would impact upon the self-thinning surface.

The three stochastic frontier models, each with a different distributional specification for \( u \), the non-negative one-sided error component in the composed error term of Eq. (14), resulted in little and practically immaterial differences in the estimated self-thinning surface (Fig. 2). In this case, a natural question would arise: do distributional assumptions matter in the estimation of self-thinning surface? Despite the vast number of stochastic frontier analysis undertaken in theoretical as well as applied research with empirical data, surprisingly few studies have been devoted to discerning the impact that alternative shapes of the distribution of \( u \) can have on the frontier and efficiency estimation and few attempts have been made to explicitly address specification testing in stochastic frontier models (Guo et al. 2018; Kumbhakar et al. 2018). Under the assumption of a normally distributed \( v \) in the composed error term, Wang et al. (2011) proposed the Pearson \( \chi^2 \) and Komolgorov-Smirnov type statistics to test the goodness of fit of the specific distributional assumption on \( u \) through simulating the quantiles of the composed error distribution. However, the power of these tests against plausible alternative distributions is somewhat low, making it hard to distinguish the sum of a normal and an exponential from the sum of a normal and a half-normal distribution, unless the variance of the normal component is very small or the sample size is very large (Wang et al. 2011). Furthermore, a rejection from the test does not necessarily imply that the distributional assumption on \( u \) is incorrect as it could be that the normality distributional assumption on \( v \) or the parametric form of the stochastic frontier model is mis-specified (Kumbhakar et al. 2018). An alternative test of the distributional assumptions on both \( v \) and \( u \) was proposed by Chen and Wang (2012) through a centred residuals-based method of moments estimator for stochastic frontier models. Unlike the commonly used maximum likelihood estimator for stochastic frontier models, this moment estimator has yet to be implemented in general purpose statistical software.
Philosophically, it does not matter that the stochastic frontier models cannot be distinguished statistically if they give more or less the same results as it only becomes a problem when the models give substantively different results (Wang et al. 2011). The choice of a distribution for \( u \) out of an ever-increasing number of theoretical candidates such as half normal, exponential, truncated normal, gamma, Pearson, uniform, Beta, Weibull, doubly truncated normal, and even a mixture distribution, as reviewed by Kumbhakar and Lovell (2000), Kumbhakar et al. (2018) and Horrace and Parmeter (2018), is often practically driven through available statistical software to implement the method rather than an underlying theoretical link between a model of productive inefficiency in econometrics and the exact shape of the corresponding distribution. Although conceptually pleasing, some alternative or more complex frontier models may lead to difficult estimation problems as found by Ritter and Simar (1997) with the normal-gamma model and also shown by the difficulty of the NT model in attaining convergence in our case. Possibly as a consequence, the half-normal assumption for the one-sided inefficiency term is almost without question the most common distribution for \( u \) in practical applications (Kumbhakar et al. 2018). In the case of this study, the estimated variance of the normally distributed error component \( \sigma^2 \) for the NH model was about 60% of that for the NE model based on the estimates of \( \sigma \), for the two models (Table 2). Correspondingly there were only six positive residuals for the former against 18 such residuals for the latter. Therefore, the relatively simple half-normal

![Fig. 6 Multi-panel display of the skewness of the 2000 estimates for each of the four parameters of the self-thinning surface estimated through the normal-half normal stochastic frontier model (blue lines) and through quantile regression over 20 evenly spaced \( r \) values from 0.80 to 0.99 (red curves) across the eight sample sizes as denoted in the top stripe of each column](image-url)
distribution became the obvious choice for $\mu$ out of the three distributions for estimating the self-thinning surface.

Another reason for choosing the half normal distribution was the difficulty of the more flexible NT model in attaining convergence during parameter estimation and the unreasonably wide range and high degree of variation of $\hat{\mu}$ and $\hat{\sigma}$, the estimate values of the mean and standard deviation of the pre-truncation normal distribution. With a convergence rate of less than 58%, the normal-truncated normal (NT) model was the slowest and most difficult to converge during parameter estimation. Among the 1156 estimations that reached convergence, a quarter ended up with $\hat{\mu}$ being a large negative or an unreasonably large negative number and $\hat{\sigma}$ being a correspondingly large or extremely large positive number. Among the unconverged cases, the values of $\hat{\mu}$ and $\hat{\sigma}$ when optimization was terminated after a large number of iterations were even larger negative and positive numbers than that in the lower 25% of the converged cases. In the cases where $\hat{\mu}$ and $\hat{\sigma}$ tended to negative and positive infinity, Meesters (2014) proved mathematically that the truncated distribution was equal to an exponential distribution. As the relationship between the truncated normal and exponential distribution is at the boundary of the parameter space of the truncated normal distribution, he deduced that a maximum of the likelihood based on the truncated normal distribution may not exist if the underlying data is exponentially distributed. During parameter estimation, this non-existence of a maximum may result in a nonconverging optimization routine or result in numerical problems before finding the maximum as observed in this study. Another cause of the
slow- and non-convergence of the normal-truncated normal model might be the high degree of collinearity between \( \mu \) and \( \sigma^2_n \) as their Pearson’s correlation coefficient (\( r \)) was almost -1 among the 1156 converged cases. Both variables appear in the log-likelihood function for parameter estimation, although \( \sigma^2_n \) in a reparameterized form (see Stevenson 1980; Kumbhakar and Lovell 2000). On the other hand, because of the wide range and high degree of variation of \( \hat{\mu} \), about 7.4% of its estimates were small positive values within an open interval of (0, 0.81) and the remaining 92.6% were negative. Therefore, it is almost certain that the truncated-normal density had a mode at 0. Furthermore, as its 95% confidence interval contained 0, \( \hat{\mu} \) might not be judged to be significantly different from zero in a statistical sense, making it hard to distinguish the truncated normal distribution from its special case, the half-normal when \( \hat{\mu} = 0 \) in this case. These reasons further strengthen our case for choosing the half-normal distribution for \( \mu \) in the stochastic frontier estimation of the self-thinning surface.

Unlike the maximum likelihood-based estimation of parameters in stochastic frontier analysis, quantile regression does not require any distributional assumption to be made for the quantile-specific error term \( \varepsilon_q \) as in Eq. (15) because its parameter estimates are obtained by minimising the asymmetric loss function as shown in Eq. (11) through linear programming. However, as shown in Figs. 4, 6 and 7, the parameter estimates, their variance, skewness and kurtosis were all quantile-dependent and these distributional characteristics also changed with sample size. Consequently, which value of \( \tau \) to choose for the estimation of the self-thinning surface becomes a question that has to be addressed with sound statistical as well as biological reasoning for the size and nature of the empirical data under analysis. The value of \( \tau \) chosen for estimating the self-thinning frontier, mostly in the two-dimensional space of stand biomass or tree size versus stand density, has ranged largely from 0.90 to 0.99 in the literature, with 0.95 \( \leq \tau \leq 0.99 \) being the most common choices (Sun et al. 2010; Zhang et al. 2013; Vospernik and Sterba 2015; Xue et al. 2015; Riofrío et al. 2016; Condés et al. 2017; Andrews et al. 2018). Only in the most extreme cases where the research interest was for the estimated boundary line to completely envelope all data points, an extremal quantile with \( \tau \) greater than 0.99 was used (e.g. Zhang et al. 2005; Salas-Eljatib and Weiskittel 2018). The number of observations in these studies ranged from tens to hundreds to thousands. Often the objectives of the analyses were twofold: (1) delineating the self-thinning boundary line to envelope most, if not all, the data points; and, (2) making statistical inferences about the boundary line across species, forest types or other environmental gradients. When estimating the self-thinning surface in higher dimensions, a balance between the two objectives may have to be found in choosing the value of \( \tau \) for a given data set.

As demonstrated by the results of this study, for sample sizes less than 100 data points, a value of \( \tau < 0.90 \) may have to be chosen, otherwise the confidence interval of parameter estimates, such as the self-thinning slope, may become far too wide for making any statistically significant and biologically meaningful inferences. Even for sample size greater than 100 data points, high and extreme quantiles with \( 0.95 \leq \tau \leq 0.99 \) should be selected after a careful comparison of multiple \( \tau \) values. The need for such a comparison stems from the following reasons: (1) the confidence intervals of parameter estimates may be too wide as shown in Fig. 6; (2) the self-thinning surfaces estimated at different quantiles could intersect each other; and, (3) parameter estimates at the extremal quantile of \( \tau = 0.99 \) may become illogical, leading to a rather flattened self-thinning slope (Fig. 4). These problems can potentially further complicate the interpretation of results. Without such a careful comparison to strike a balance between the two objectives, quantile regression would likely carry a certain degree of subjectivity, defying the original purpose of its use i.e. to avoid the long-standing problem of subjectivity in the estimation of the self-thinning boundary line as described in the introduction.

Across all eight sample sizes and over the quantile range of \( 0.90 \leq \tau \leq 0.99 \) that is most applicable for estimating the self-thinning surface in higher dimensions, the normal-half normal stochastic frontier model proved to be superior to quantile regression in the sense of statistical efficiency. Its parameter estimates had much lower degrees of variability and correspondingly narrower confidence intervals (Figs. 4, 5). This greater efficiency would naturally be conducive to detecting differences among species, forest types, site classes and other environmental and climatic conditions through statistical inferences. When the entire data set of all 170 observations were used in the estimation, the model had only six positive residuals and therefore the estimated self-thinning surface enveloped about 96.5% of the data points. This degree of envelopment would be equivalent to a regression quantile estimation with a \( \tau \) of 0.965, but it was achieved with a much greater degree of precision through the stochastic frontier estimation. Such comparative performances of the two methods could also be seen from the results reported but not noted or elaborated by Zhang et al. (2013). Furthermore, the stochastic frontier estimation could lead to a more objective self-thinning surface because it did not involve the subjective selection of a particular value of \( \tau \) to peel through the data space in high dimensions as in quantile regression.

However, quantile regression could still provide a valuable complement to stochastic frontier analysis in the estimation of the self-thinning surface as it allowed the examination of the impact of variables other than stand density, namely \( S \) and \( W \) in this case, on different quantiles.
of stand biomass. The quantile regression estimates of $\beta_2$ and $\beta_3$, the two parameters associated with relative site productivity $S$ and the abundance of woody weeds, exhibited somewhat different patterns of quantile-dependence (Table 3, Fig. 4). When $\tau$ increased from 0.80 to 0.95, $\beta_2$ generally decreased but $\beta_3$ showed no obvious change trend. As $\tau$ further increased, both $\beta_2$ and $\beta_3$ increased, but with the former less so than the latter. The effect of site productivity and the impact of the abundance of woody weeds on the accumulation of stand biomass were not the same across the conditional quantiles. Such differences, when carefully examined, would provide a greater insight into the growth dynamics of the even-aged larch stands growing under the self-thinning surface. This complementarity between stochastic frontier analysis and quantile regression argues for the application of both methods in the estimation of the self-thinning surface in high dimensions, especially when dealing with data that do not conform to the distributional assumptions of stochastic frontier models.

Conclusions

For estimating the self-thinning surface in higher dimensions using cross-sectional data, the normal-half normal (NH) stochastic frontier model proved to be superior to quantile regression over the quantile range of $0.90 \leq \tau \leq 0.99$ across small as well as large sample sizes. The superiority lies not only in the statistical efficiency of parameter estimation but also in the greater objectivity of the stochastic frontier model as it does not involve the subjective selection of a particular value of $\tau$ as in quantile regression. The higher statistical efficiency and greater degree of objectivity are conducive to making statistical inferences about the parameters of the self-thinning surface. However, quantile regression can still provide a valuable complement to stochastic frontier analysis in the estimation of the self-thinning surface because it can deal with data that do not conform to the distributional assumptions of stochastic frontier models and help reveal the impact of variables other than stand density on stand biomass at levels of $\tau$ below the estimated surface.

Acknowledgements We are indebted to the academic staff, and past and present postgraduate students of the Department of Forest Management, School of Forestry, Northeast Forestry University, who collected the field data.

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