The patterns of nearest neighbor trees in a temperate forest

Xiongwen Chen,
Kimberly A Bowman

Introduction

Forest structure, meaning the arrangement of trees, species, sizes, or age distributions in a vertical and horizontal space (Goff & Zedler 1968), is an emergent property of a forest community (Ponge 2005). Individuals of forest plant species are often aggregated (Greig-Smith 1982), so it is essential to study the nearest neighbor trees in any forest community. The most immediate neighborhood (i.e., a community of trees) is one of the crucial characteristics reflecting forest structure and revealing individual trees’ contribution to the heterogeneity at a forest stand scale (McRoberts 2012, Zenner & Peck 2018). Information on nearest neighborhood distribution in different forest types is needed because these nearest individuals (or species) can play critical roles in forest communities, such as those related to economic impact (e.g., timber production and windbreak) and ecosystem health (e.g., species invasion – Iverson & Prasad 1998, Poland & McCullough 2006).

In ecology, neighborhood patterns have been crucial in explaining species coexistence and community structure. For example, niche theories propose that environmental heterogeneity and biological interactions may lead to spatial clustering because different species have varied niches (Grubb 1977); dispersal-assembly theories indicate that dispersal limitation may contribute to the emergence of spatial clustering (Wong & Whitmore 1970, Hubbell 1997). The Janzen-Connell hypothesis (Janzen 1970, Connell 1971) predicts that widely dispersed seeds far away from parent plants are essential in avoiding the detrimental influence of pathogens, herbivores, seed predators, and seedling competition. These mechanisms have been proposed to explain the diversity of forests as they maintain the survival of many different plant species within one localized region.

The nearest neighbor trees (NNTs) are essential for reflecting forest structure and spatial heterogeneity in a forest stand. It is not clear whether different tree species have varied patterns of NNTs in a small area due to biological interactions, whether big trees affect the nearest neighbors for diversity and recruitment, or whether a universal linear relationship between the distance of NNTs and their average DBH exists. In this study, the information of NNTs at two plots (each 30 × 100 m) in a temperate mixed broadleaf forest in Southern USA was collected by field survey. Our results indicated that approximately 80% of NNTs were within a distance of 1.5-4.0 m. Tulip poplar, oaks, and hickory trees did not have the same species as NNTs or were very limited. Carolina buckthorn had itself as an NNT but with fewer other species. Sugar maple could serve as the NNT for oaks, hickory and others. The relationships between the distance of an NNT and its cumulative percentage were different among various species or groups. Overall, for trees and their NNTs, there existed complicated relationships between their sizes (e.g., height and DBH). Big trees might affect NNTs in diversity and recruitment. The suggested linear relationship between tree size and distance was not observed. The results could be helpful to manage forest structure (tree species and NNT) and provide evidence to improve the scaling theory on NNTs.

Keywords: Forest Structure, Heterogeneity, Scaling, Species Interaction, Tree Size

Here, diversity means that the nearest neighbor tree in a natural forest is a different species. However, it is unclear whether different tree species may vary in response to interacting forces in a small area.

Mapping multiple individual tree species over an area is required to study the nearest neighbors. Plot data, such as from forest inventory, are always helpful. In the USA, forest inventory data were used to create maps of tree species abundance and distribution for the entire eastern part of the country (Wilson et al. 2012). However, the forest inventory data are not spatially continuous because the plots are small and discrete; also, seedlings were not included in the forest inventory. Their results may be helpful to explain species patterns at large scales. But for large plots (e.g., 10 ha), topography and forest types may affect the outcome (Wiegand et al. 2007a, Bianchi et al. 2021). For example, a steep slope may disperse seeds far away. Also, it needs a high cost to survey a large plot. For a local scale with relatively homogenous environmental conditions, it would be interesting to have spatially continuous data (e.g., a relatively big plot or transect) including seedlings to study the patterns of nearest neighbor trees.

Studying plant species occurrences, spatial patterns, and nearest neighbor associations in communities at small scales may show their biological interactions (Wiegand et al. 2007b, Bianchi et al. 2021). There are many different methods to study spatial patterns based on the research pur-
pose, such as pair correlation function, Ripley function, O-ring function, nearest neighbor distance index, Voronoi polygon analysis, or uniform angle index (Pommerening & Grabarnik 2019). Direct tree-tree interactions can operate at a small local scale, such as 2-4 m in a tropical forest (Wiegand et al. 2007a). After studying the plant diversity change in temperate forest communities at different spatial scales, it was found that biotic interactions explained the most variation at fine scales (Yuan et al. 2011). Species pair-correlation or association with the nearest neighbors may evaluate their interactions. Different interactions (e.g., positive and negative) could form distinct patterns in nearest neighbor trees (Lieberman & Lieberman 2007, Liu et al. 2020). Some species or big trees can have many different species as neighbors, these are called accumulator species; while repeller species have limited neighbors, these are called accumulator trees, as they can have many different species as neighbors. This study investigates the pattern of nearest neighbor trees in a temperate forest on a small scale based on the absolute distance. The hypothesis is that to be complementary, the nearest neighbor to a tree would have a high probability of being a different species and of entirely different morphology (e.g., one is big and the other is small). The specific objectives include (i) determining whether there is a general pattern for nearest neighbor trees, whether different tree species have similar patterns for nearest neighbor trees, such as in species composition and tree size, and whether big trees affect their nearest neighbors, such as in size and recruitment; (ii) whether there exists a universal scaling relationship ($d_i = c \cdot r_i$) for nearest neighbor trees in this temperate forest, whether different tree species have similar patterns in this scaling relationship; and (iii) implications for forest management at a small scale.

**Materials and methods**

**Study site**

The study site is located at Guntersville State Park ($34°23′24″$ N, $86°12′57″$ W) in Marshall County of Alabama, USA. The forest community is composed of broadleaved (mainly hickories and oaks) forest in the southern region of the USA, and the oldest trees are about 100 years old. There were limited human disturbances (e.g., no timber harvesting) in this forest community during the last five decades. The entire area of plots is within a relatively flat section at the southeastern aspect with a slope of around 10 degrees. This could minimize the contribution of topographic differences. The elevation of the site is about 200 m above sea level, and the soil is a red clay soil type with a silt loam texture (https://www.nrcs.usda.gov). The average air temperature in this region is around 18°C, and the mean annual precipitation is about 1350 mm (based on recent 20 years’ data from the nearby meteorological station at Huntsville International Airport).

**Plot information and methods**

Two plots were established. Each plot size is 0.3 ha (100 m in length and 30 m in breadth). These two plots are parallel but separated at 40 m to avoid spatial autocorrelation. All living trees, including woody seedlings in the plots, were surveyed. Tree seedlings are plants with a woody stem but less than 1.3 m in stem height and thus no diameter at breast height (DBH). Based on abundance, the major tree species of the two plots included Carolina buckthorn (Frangula caroliniana [Walter] A. Gray), tulip poplar (Liriodendron tulipifera L.), sugar maple (Acer saccharum Marshall), shagbark hickory (Carya ovata [Mill.] K. Koch), red oak (Quercus rubra L.), white oak (Quercus alba L.), American elm (Ulmus americana L), black cherry (Prunus serotina Ehrh.), black locust (Robinia pseudoacacia L) and others. Some 1368 tree individuals (including seedlings) on plot 1 and 591 tree individuals on plot 2 were recorded.

For each tree in the plots, its location ($x$, $y$), DBH, canopy diameter, and tree total height were recorded. Only one tree (or seedling) with the shortest distance was selected based on its distance to nearest trees. Tree height was measured by a Pulse Rangerfinder® Hypsometer (Laser Technology Inc., Centennial, CO, USA). The mean canopy diameter of each tree was estimated from measurements in two opposite directions (e.g., north-south and east-west) by a tape with an accuracy of 1 cm. The circumference at breast height (1.3 m)
was measured by a tape with an accuracy of 1 mm. DBH was calculated by transforming circumference. For each seedling, the circumference at the stem bottom (interface at the soil surface) was used to estimate its stem diameter. Each tree was considered as a standard cylinder, and the tree occupying volume \(V\) was calculated from canopy diameter \((CD)\) and tree height \((h)\) as the following (eqn. 1):

\[
V = \pi \cdot h \cdot (CD/2)^2
\]  

\((1)\)

As for the big tree (height ≥ 5 m) size, height was classified as follows: [5-10 m], i.e., height ≥ 5 m but < 10 m; [10-15 m]; [15-20 m]; and ≥ 20 m. For each big tree, the nearest neighbor tree species and whether it was a seedling were recorded. The ratio of species for big trees was calculated as the species number of the nearest neighbors divided by the species number of big trees in each height class. If the ratio of species is more than 1.0, the big trees are considered accumulator species that can increase species diversity; otherwise, the big trees are considered repellers. The same idea was used for seedlings; the ratio of seedlings was calculated as the number of seedlings within the nearest neighbors divided by the number of big trees in each class. If the ratio of seedlings is high (such as 1.0), the big trees may be considered beneficial to seedlings.

The distance \((d)\) between each tree and its nearest neighbor tree was calculated from their locations. The average DBH of the nearest tree pair was estimated as \((DBH1+DBH2)/2\). The ratios of height, DBH, and volume for the nearest tree pair were calculated by \(h_1/h_2, DBH_1/DBH_2, \) and \(V_1/V_2\), respectively. To make the result easy to read, here these ratios \((h_1/h_2, DBH_1/DBH_2, \) and \(V_1/V_2\) were calculated with the greater value as the numerator and were thus always 21.

Spearman’s correlation was conducted between the distance of nearest neighbors and morphological characters of trees (e.g., ratio in height, DBH, and \(V\)) by SAS software v. 9.3, (SAS Institute Inc., Cary, NC, USA) with the statistical significance at \(p < 0.05\).

**Results**

There were 1368 trees, including 703 seedlings at plot 1, and 591 trees, including 110 seedlings at plot 2. Tree height varied from 0.1 m to 43.4 m in the plots, and the average tree height was 4.3 m in plot 1 and 12.7 m in plot 2. The average DBH was 4.7 cm in plot 1 with the range from 0.3 cm to 84.4 cm, and the average DBH was 14.3 cm in plot 2 with the range from 0.15 cm to 95.6 cm.

The distance of nearest neighbor trees ranged from 0.1 m to 7.19 m for overall individuals in plot 1 and 0.1 m to 11.42 m in plot 2. There were approximately 76% of nearest neighbor trees within a 1.5 m distance in plot 1 and 82% of nearest neighbor trees within a distance of 4 m in plot 2 (Fig. 1).

Some tree species did not have the same species as the nearest neighbor trees or were very limited (Tab. 1), such as tulip poplar, oaks, and hickory trees. Other species only had the same species as nearest neighbor trees and limited other species, such as Carolina buckthorn. Others could live with the same species or other species, such as American elm and black locust. Oaks and hickory trees both lived with sugar maple as nearest neighbor trees. The relationships between the distance
of nearest neighbor trees and the cumulative percentage were different among varied species or groups (Fig. 2). The correlation between the distance of nearest neighbor trees and the accumulated percentage was linearly significant for tulip poplar, American elm, and black locust in plot 1. For other species (or groups), there existed nonlinear relationships (e.g., logarithmic form) with a saturation point at a distance of 4 m.

There were no clear relationships in size (e.g., height, DBH, and V) between trees and their nearest neighbors (Fig. 3). High variations in size ratios usually appeared within the distance of 4 m. Around a big tree (i.e., a high ratio), there were usually many small trees (i.e., small ratios) at multiple vertical layers. The patterns in ratio distribution were slightly different in the two plots (e.g., the nearest distance and multiple layers). Some negative relationships seemed to exist for these outlier points, such as \( h_1/h_2 \) and \( V_1/V_2 \), but these relationships were not statistically significant. Similarly, patterns existed among different tree species or groups, such as tree height vs. tree height of nearest neighbors. Big trees may affect the diversity of nearest neighbors (Fig. 4a). The ratios of species numbers were higher in plot 2 than in plot 1. Big trees may also affect the nearest seedling distribution (Fig. 4b). The ratio of seedlings in the big trees was relatively high in the height class of \( \geq 20 \) m in plot 1, but it was relatively high in the height class of \( 10-15 \) m in plot 2.

A positive linear relationship between the distance of nearest neighbor trees and their average DBH was not observed in both plots (Fig. 5). Similar patterns were also observed at the species or group level.

**Discussion**

Different tree species may have varied characteristics to influence the nearest neighbor trees since the plots are small and relatively homogeneous in environmental factors. Differences between plots in this study (as in Fig. 1 and Fig. 4) may have been due to or augmented by trees and stand characteristics. For example, plot 1 had over twice as many trees (1368 vs. 591) and more than double the percentage of saplings (51% vs. 19%) as plot 2. Bias may exist for the trees around plot boundaries, but it should be limited due to the sample size of hundreds of trees within each plot. Based on whether the nearest neighbor tree species was the same species/group or not, three groups could be classified: those that mainly had homospecific neighbors, those that had mostly heterospecific neighbors, and those that had both. The emergent interactions with the nearest neighbor trees in the community networks may be varied for species at different locations (e.g., plot 1 and plot 2). There were many Carolina buckthorn and sugar maple in plot 1 and plot 2, respectively. But most Carolina buckthorn lived with themselves as nearest neighbor trees, while sugar maple lived with nearest oaks, hickories, and other trees. It appears that sugar maple is a prevalent tree species and can live with oaks, hickory, and others as the nearest neighbors. However, if the space between any pairs of trees is too close (such as 10 cm in this study), natural resources, such as light and soil water, may become limited, causing one or multiple trees to suffer dieback from poor nutrition or moisture. Both the dispersal-assembly theories and the Janzen-Connell hypothesis may explain the pattern of nearest neighbor trees.

The result of nearest tree species is consistent with the previous finding that the heterogeneous information from a fine scale can provide different outcomes and model-based estimates than those from a large scale (Wilson et al. 2012, Zemmer & Peck 2018). Accumulator or repeller species were used to describe this kind of variation in a tropical forest (Punchi-Manage et al. 2015). In this case, some patterns existed in the association of trees and their nearest neighbor species. The detailed description of the patterns may provide a deep insight.
into the biology of the species interactions in community networks and generate hypotheses. The regime in the distance distribution for nearest neighbor trees could be different among various species or groups. For tulip poplar, American elm, and black locust, there was a significant correlation between the distance of nearest neighbor trees and accumulated percentage in plot 1. The relationship was different for the other species (oaks, hickory, and Carolina buckthorn) but these relationships follow logarithmic forms. Our results support that multiple scales may exist in the distribution of distances with nearest neighbors within the smaller-scaled level itself (Wiegand et al. 2007a,b). The distance threshold (90% of accumulated frequency) may be related to the spatial autocorrelation in the distribution of the nearest neighbor trees. These thresholds varied slightly with species. The distance threshold for different species was about 2 m for Carolina buckthorn, 2.5 m for black locust, 2.5 m for tulip poplar, 3 m for hickories, 3 m for oaks, and 3 m for American elm, respectively. These trees have approximately 90% of nearest neighbor trees within 2-4 m, which could mean relatively high connectivity to the nearest neighbor trees compared to those in more than 10 m. It was proposed that the connectivity to the nearest neighbor trees could be influenced by (i) population density and local community dynamics; (ii) habitat quality and spatial configuration; and (iii) species’ migration and behavior (Moilanen & Nieminen 2002). Further multiscale research on community structure could be conducted from this data set. The related processes might include disturbance, mortality, colonization, reproduction, recruitment, growth, competition, and senescence (Coff & West 1975, Bonan 1988). The net balance of negative or positive species-species interactions could result in distinct spatial patterns in the placement of neighboring species and have the potential to generate spatial structure (Lieberman & Lieberman 2007). The smaller distances to nearest heterospecific neighbors were more likely to be accumulators; otherwise, the long distance might mean repellers (Punchi-Manage et al. 2015). The result indicates that the distance frequency to nearest neighbor trees might be used to characterize the structural heterogeneity in a temperate forest community.

To co-exist, trees and their nearest neighbors are hypothesized to be of either small or differing sizes (e.g., height, diameter, and occupying volume). However, this relationship was not evident in this study, where the ratios of tree size were varied. Tree size is an essential structural element that affects resource use, growth, reproduction, and allometry at a small scale (West et al. 2009). A more or less inverse relationship between the sizes was often found in multi-layered temperate forests because of species’ shade tolerance (Peterken 1996); such as fast-growing, short-lived tree species can reach over-story canopy from gaps while shade-tolerant trees grow beneath them (Lieberman & Lieberman 2007). This structure is considered to have high vertical heterogeneity with “balanced” or “equilibrium” population structures (Goodburn & Lorimer 1999, Rubin et al. 2006). Since only one tree was selected for the nearest neighbor in this study, analysis of this relationship may be incomplete. For example, the size ratios might be massive for a forest structure with many low-ratio values. The big trees (tall trees) in this study could affect the nearest neighbors in species diversity and seedlings because there were many low-ratio values. The big trees (tall trees) in this study could affect the nearest neighbors in species diversity and seedlings because there were many low-ratio values. The big trees (tall trees) in this study could affect the nearest neighbors in species diversity and seedlings because there were many low-ratio values.

The scaling relationship between the nearest neighbor trees and their average DBH. Some previous studies indicated that the asymmetric competition for light among trees in forest stands could affect the scaling relationships (King 1994, Purves & Pacala 2008, Ishihara et al. 2016). Trees could have diverse scaling relationships in their average diameter and the nearest distance for the tree pairs because these varied scaling relationships exist in tree allometric scaling (Chen 2018). This result may provide a case for the limitation of the Metabolic Scaling Theory.

The results from this study could provide implications for forest management.Traditional forest management emphasizes species composition at stand level or releasing suppressed trees, but does not quantitatively include the distance to the nearest neighbor trees. Management practices could be made more efficient if new tree configurations in a stand were considered before conducting alterations. In this region, logging, thinning, and prescribed burning are mainly related to the nearest neighbor trees. First, species-species interactions from the nearest neighbor trees need to be considered. We need to realize that some species may not have the same species as their nearest neighbor trees, such as tulip poplar trees. But in this study, sugar maple trees lived with all other species as nearest neighbors. This emergent pattern could be caused by species interactions within community networks, possibly related to pathogens or allelopathy. After logging or thinning, it may be better if some tree species are not the nearest
neighbors to themselves. For example, in the absence of disturbance or human activities (e.g., prescribed burning), the Carolina buckthorn may form a patch because it may like to live as its own nearest neighbor so that other species cannot invade. This characteristic is suitable for even-aged plantations. Second, maintaining a distance with the nearest neighbor trees, such as 3 m or more, may result in fewer suppressed trees. Based on given tree sizes, it is possible to estimate the naturally-occurring optimum distance for the nearest neighbors from the algorithms in this study. Third, the dimensions of nearest neighbor trees need to be managed. Under a big tree, only small trees with shade tolerance can survive. Taking out a big tree or trimming its big canopy (e.g., branch) may increase the growth of neighbor trees. Big trees might affect the nearest neighbors in species diversity and seedling recruitment, but this pattern may be inconsistent across sites. Forth, the scaling relationships derived from Metabolic Scaling Theory, such as the distance of nearest neighbor trees and their average DBH, may not be applicable in a southern USA temperate forest area. These scaling relationships should be tested before applications.

Conclusions
After studying the patterns of nearest neighbor trees at a small scale in a temperate forest through intensive tree mapping, there existed some patterns in suitable species and distances to nearest neighbors, which differed by species or group. However, detailed biological mechanisms need to be investigated further. Realizing the patterns of suitable species and distances to nearest neighbors may be helpful to the management of local natural forests and plantations, especially when considering the management practices of logging, thinning, and prescribed burning. Some results, which conflict with the scaling theory on large scales, may complement the development of the Metabolic Scaling Theory. Current advances in quality and reductions in the cost of remotely sensed tools are providing new opportunities for spatially explicit monitoring.

Acknowledgments
The authors are thankful to Matthew Shaw, Lawson Quick, and Michael Brown for their help in fieldwork and also to the Guntersville State Park Administration for their help in fieldwork and also to the authors. This research was supported by the USDA National Institute of Food and Agriculture McIntire Stennis project (1008643) and the Capacity Building Program (2021-38821-34596).

References
Bianchi E, Bugmann H, Hobi ML, Bigler C (2021). Spatial patterns of living and dead small trees in subalpine Norway spruce forest reserves in Switzerland. Forest Ecology and Management 494: 119315. - doi: 10.1016/j.foreco.2021.119315

Bonan GB (1988). The size structure of theoretical plant populations: spatial patterns and neighborhood effects. Ecology 69: 1721-1730. - doi: 10.2307/1938152

Chen X, Li B-L (2003). Testing the allometric scaling relationships with seedlings of two species. Acta Oecologica 24: 125-129. - doi: 10.1016/S1436-020X(03)00062-6

Chen X (2006). Tree diversity, carbon storage and soil nutrient in an old-growth forest at Changbai Mountain, Northeast China. Communications in Soil Science and Plant Analysis 37: 363-375. - doi: 10.1080/00103620500440210

Chen X (2009). Trends of forest inventory data in Alabama, USA during the last seven decades. Forestry 83: 517-526. - doi: 10.1093/forestry/cpq034

Chen X (2018). Diverse scaling relationships of tree height and diameter in five tree species. Plant Ecology and Diversity 11: 147-155. - doi: 10.1016/j.pladed.2017.07.004.18.04528

Chen X (2020). Managing the forest landscape: exploring the quantitative interplay between forestland patches, areas and landowner numbers in counties from Alabama, USA. Small-Scale Forestry 19: 505-520. - doi: 10.1080/11467999.2020-09451-8

Chen X, Niu J (2020). Relationships between tree height and tree species richness at small scales. Acta Oecologica 109: 103668. - doi: 10.1016/j.actao.2020.103668

Connell JH (1971). On the roles of natural enemies in preventing competitive exclusion in some marine animals and in rain forest trees. In: “Dynamics of Populations”. Center for Agricultural Publishing and Documentation, Wageningen, The Netherlands, pp. 298-312.

Enquist BJ, West GB, Brown JH (2009). Extensions and evaluations of a general quantitative theory of forest structure and dynamics. Proceedings of the National Academy of Sciences USA 106: 7046-7051. - doi: 10.1073/pnas.0812030106

Goff FG, Zedler PH (1968). Structural gradient analysis of upland forests in the western Great Lakes area. Ecological Monograph 38: 65-86. - doi: 10.2307/1948537

Goff FG, West D (1975). Canopy-understory interaction effects on forest population structure. Forest Science 21: 98-108. - doi: 10.1093/forestscience/21.2.98

Goodburn JM, Lorimer CG (1999). Population structure in old-growth and managed northern hardwoods: an examination of the balanced diameter distribution concept. Forest Ecology and Management 118: 11-29. - doi: 10.1016/S0378-1119(99)00047-2

Greig-Smith P (1982). Quantitative plant ecology (3rd edn.). Blackwell, Oxford, UK, pp. 359. [Online] URL: http://books.google.com/books?id=fSfr9fhP8A0c

Grubb P (1977). The maintenance of species richness in plant communities: the importance of the regeneration niche. Biological Review 52: 107-145. - doi: 10.1111/j.1469-185X.1977.tb01347.x

Hubbell SP (1997). A unified theory of biogeography and relative species abundance and its application to tropical rain forests and coral reefs. Coral Reefs 16: 59-521. - doi: 10.1007/s003380050237

Hubbell SP (2001). The unified neutral theory of biodiversity and biogeography. Princeton University Press, Princeton, NJ, USA, pp. 392.

Ishihara Mi, Konno Y, Umeki K, Ohno Y, Kikuzawa K (2016). A new model for size-dependent tree growth in forests. PLOS One 11: e0155219. - doi: 10.1371/journal.pone.0155219

Iversen LR, Prasad AM (1998). Predicting abundance of 80 tree species following climate change in the eastern United States. Ecological Monograph 68: 465-485. - doi: 10.1890/0012-9615(1998)068[0465:PAOTSF]2.0.CO;2

Janzen DH (1970). Herbivores and the numbers of tree species in tropical forests. American Naturalist 104: 501-518. - doi: 10.1086/S02628

King DA (1994). Influence of light level on the growth and morphology of sapling in a Panamanian forest. American Journal of Botany 81: 948-957. - doi: 10.2307/2445287

Kotliar NB, Wiens JA (1990). Multiple scales of patchiness and patch structure: a hierarchical framework for the study of heterogeneity. Oikos 59: 253-260. - doi: 10.2307/5455472

Liebmer M, Liebmer D (2007). Nearest-neighbor tree species distributions in tropical forest: the role of chance, and some consequences of high diversity. Oikos 116: 377-386. - doi: 10.1111/j.0030-1299.2007.21370.x

Li Y, Hui G, Yu S, Luo Y, Yao X, Ye S (2017). Nearest-neighbor relationships in Pinus yunnanensis var. tenuifolia forests along the Nanpan River, China. iForest 10: 746-753. - doi: 10.3832/ifor2405-010

Liu P, Wang W, Bai Z, Guo Z, Ren W, Huang J, Xu Y, Yao J, Ding Y, Zang R (2020). Competition and facilitation co-regulate the spatial patterns of boreal tree species in Kanas of Xinjiang, northwest China. Forest Ecology and Management 467: 11867. - doi: 10.1016/j.foreco.2020.11.8167

McRoberts RE (2012). Estimating forest attribute parameters for small areas using nearest neighbor techniques. Forest Ecology and Management 272: 3-12. - doi: 10.1016/j.foreco.2011.06.039

Mollanen A, Nieminen M (2002). Simple connectivity measures in spatial ecology. Ecology 83: 1131-1145. - doi: 10.1890/0012-9658(2002)083[1131:SCMISE]2.0.CO;2

Parker KE (1996). Natural woodland: ecology and conservation in northern temperate regions. Cambridge University Press, Cambridge, UK, pp. 522. [Online] URL: http://books.google.com/books?id=p3nyNmXPC6

Poland TM, McCullough DC (2006). Emerald ash borer invasion of the urban forest and the threat to North America’s ash resource. Journal of Forestry 104: 118-124. - doi: 10.1093/jof/104.3.118

Pommerening A, Grabarnik P (2019). Individual-based methods in forest ecology and management. Springer, Cham, Switzerland, pp. 411. [Online] URL: http://link.springer.com/content/pdf/10.1007%2F978-3-311-01522-1.pdf

Ponge JF (2005). Emergent properties from organisms to ecosystems: towards a realistic approach. Biological Review 80: 403-411. - doi: 10.1111/j.1469-185X.2002.00672X

Punch-Maier R, Wiegand T, Wiegand K, Getzin S, Huth A, Gunatilleke CVS, Gunatilleke IAUN (2015). Neighborhood diversity of large trees shows independent species patterns in a mixed dipterocarp forest in Sri Lanka. Ecology 96: 320
Patterns of nearest neighbor in a forest

1823-1834. - doi: 10.1890/14-1477.1
Purves D, Pacala S (2008). Predictive models of forest dynamics. Science 320: 1452-1453. - doi: 10.1126/science.1155359
Rubin BD, Manion PD, Faber-Langendoen D (2006). Diameter distributions and structural sustainability in forests. Forest Ecology and Management 222: 427-438. - doi: 10.1016/j.foreco.2005.10.049
West GB, Enquist BJ, Brown JH (2009). A general quantitative theory of forest structure and dynamics. Proceedings of National Academy of Sciences USA 106: 7040-7045. - doi: 10.1073/pnas.0812294106
Wiegand T, Gunatilleke S, Gunatilleke N, Okuda T (2007a). Analyzing the spatial structure of a Sri Lankan tree species with multiple scales of clustering. Ecology 88: 3088-3102. - doi: 10.1890/06-1350.1
Wiegand T, Gunatilleke S, Gunatilleke N (2007b). Species associations in a heterogeneous Sri Lankan Dipterocarp forest. American Naturalist 170: E77-E95. - doi: 10.1086/521240
Wilson BT, Lister AJ, Riemann RI (2012). A nearest-neighbor imputation approach to mapping tree species over large areas using forest inventory plots and moderate resolution raster data. Forest Ecology and Management 271: 182-198. - doi: 10.1016/j.foreco.2012.02.002
Wong YK, Whitmore TC (1970). On the influence of soil properties on species distribution in a Malayan lowland dipterocarp forest. Malayan Forestry 33: 42-54. [online] URL: http://www.cabdirect.org/cabdirect/abstract/19700605763
Yuan Z, Gazol A, Wang X, Lin F, Ye J, Bai X, Li B, Hao Z (2011). Scale specific determinants of tree diversity in an old growth temperate forest in China. Basic and Applied Ecology 12: 488-495. - doi: 10.1016/j.baae.2011.07.008
Yuan Z, Ali A, Jucker T, Ruiz-Benito P, Wang S, Jiang L, Wang X, Lin F, Ye J, Hao Z, Loreau M (2019). Multiple abiotic and biotic pathways shape biomass demographic processes in temperate forests. Ecology 100: e02650. - doi: 10.1002/ecy.2650
Zenner EK, Peck JE (2018). Floating neighborhoods reveal contribution of individual trees to high substand scale heterogeneity. Forest Ecology and Management 412: 29-40. - doi: 10.1016/j.foreco.2018.01.054