Enhanced Structural Complexity Index: An Improved Index for Describing Forest Structural Complexity

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The horizontal distribution of stems, stand density and the differentiation of tree dimensions are among the most important aspects of stand structure. An increasing complexity of stand structure is often linked to a higher number of species and to greater ecological stability. For quantification, the Structural Complexity Index (*SCI*) describes structural complexity by means of an area ratio of the surface that is generated by connecting the tree tops of neighbouring trees to form triangles to the surface that is covered by all triangles if projected on a flat plane. Here, we propose two ecologically relevant modifications of the *SCI*. The degree of mingling of tree attributes, quantified by a vector ruggedness measure, and a stem density term. We investigate how these two modifications influence index values. Data come from forest inventory field plots sampled along a disturbance gradient from heavily disturbed shrub land, through secondary regrowth to mature montane rainforest stands in Mengsong, Xishuangbanna, Yunnan, China. An application is described linking structural complexity, as described by the *SCI* and its modified versions, to changes in species composition of insect communities. The results of this study show that the Enhanced Structural Complexity Index (*ESCI*) can serve as a valuable tool for forest managers and ecologists for describing the structural complexity of forest stands and is particularly valuable for natural forest swith a high degree of structural complexity.

Keywords: Forest Structure Index; Structural Complexity; Stem Map; Species Composition; NMDS

Introduction

The importance of ecosystem structure to species richness has been established through many studies. Already in the early 1960s MacArthur and MacArthur (1961) showed that the physical structure of a plant community was of greater importance than the composition of plant species in determining bird diversity. A meta-analysis by Tews et al. (2004) found that 85% of 85 reviewed studies on habitat heterogeneity and species richness conducted between 1960 and 2003 found a positive correlation between richness and structural variables. As plants play an important role in shaping the physical structure of many environments (Lawton, 1983; McCoy & Bell, 1991) the structural complexity of plant communities has frequently been used as an indicator of the diversity in other taxa (Whittaker, 1972; Franzreb, 1978; Temple et al., 1979; Aber, 1979; Recher et al., 1996; Moen & Gutierrez, 1997). Moreover, the habitat heterogeneity hypothesis (Simpson, 1949; MacArthur & Wilson, 1967) relates this positive association between species diversity and structural complexity by suggesting that more complex environments provide increased niche space and thus facilitate specialization and avoidance of competition through spatial segregation (Cramer & Willig, 2005). This further implies that the structural complexity of a forest is of great importance for the number and composition of species inhabiting it (Willson,

1974; Ambuel & Temple, 1983; Freemark & Merriam, 1986; Spanos & Feest, 2007).

The multidimensional character of forest stands makes it hard to describe structural complexity holistically. Canopy layering, the presence of particular understory species, trees with different bark types, and decaying logs and hollow trees (Doherty et al., 2000) have all been considered to be important components of structural complexity. However, three-dimensional stand structures are probably the most important of all characteristics (Pretzsch, 1997). According to Pretzsch (2009), the horizontal distribution pattern of trees, stand density, the differentiation of dimensions, and species intermingling constitute the most important aspects of stand structure that influence growth processes, habitats, species richness, and stability of forest ecosystems. Kimmins (2005) suggests the spatial arrangement of plants, both horizontally and vertically, the structure of tree canopies and the presence of canopy gaps, and snags, and coarse woody debris are the principal characteristics that influence the diversity of animals. While some of these attributes are hard to define and difficult to measure in the field. tree stem diameter and position are standard in measurement protocols of forest inventories in countries like, for example, the USA (United States Department of Agriculture Forest Service, 2011) and Germany (Polley, 2007). Hence, this study focuses on these variables and defines structural complexity as the spatial arrangement of plant dimensions, both horizontally and vertically (Zenner & Hibbs, 2000).

Several indices have been developed in the past decades to provide interpretable metrics of structural complexity and thus facilitate comparisons among stands (Pommerening, 2002; LeMay & Staudhammer, 2005; McElhinny, 2005). LeMay & Staudhammer (2005) identify three groups of indices: 1) indices based on tree attributes, 2) indices of spatial heterogeneity and 3) indices combining tree attributes and spatial heterogeneity. While indices in groups 1) and 2) focus on only one aspect of forest structural complexity, indices in group 3) intend to retain more information and thus may provide more comprehensive measures of structural complexity.

In this study we propose an index that integrates the horizontal distribution of trees, stand density, and the differentiation and intermingling of tree dimensions into one measure, while requiring only data on tree position and diameter-at-breastheight (*DBH*) for its calculation. Our index is a modification of the Structural Complexity Index (*SCI*) developed by Zenner & Hibbs (2000). In the first part of this study, we describe limitations of the *SCI* and propose two modifications to create an Enhanced Structural Complexity Index (*ESCI*). In the second part, the *SCI* and *ESCI* are calculated for forest inventory data from an upland landscape in Xishuangbanna, South China, and results are compared. In the last part, we compare how the *SCI* and *ESCI* correlate with turnover in insect species composition.

The Structural Complexity Index (SCI)

Zenner & Hibbs (2000) introduced the SCI, a formula that mathematically integrates both vertical (size differentiation) and horizontal (spatial position) components of forest structure. It is based on the position of trees whose xy-coordinates are complemented with a tree attribute, such as DBH or height, as a z-coordinate. By a spatial tessellation approach (Delaunay, 1934) each tree is connected to its neighbours such that traingles are defined that form a continuous faceted surface, i.e. a triangulated irregular network (TIN) (Figure 1). If tree height is selected as the z-coordinate, this TIN can be visualized as connecting the tops of neighbouring trees. Instead of tree height, any measured continuously or ordinally scaled tree attribute can be chosen as the z-coordinate. The SCI is defined as the surface area of the TIN in three dimensional space divided by the area covered by its projection on a plane surface (Equation (1)). If all trees have the same z-value (e.g. all trees have the same height or basal area as in an even aged plantation) the SCI equals 1, the lower limit of the SCI. For structurally more complex forest stands the SCI is >1.

$$SCI = \frac{\text{surface area of TIN}}{\text{projected area of TIN}}$$
(1)

Limitations of the SCI

We illustrate basic characteristics of the *SCI* with a set of four simple forest stands which differ in their structural complexity but have the same *SCI* value. All stands cover the same area but vary in number of trees, range of tree height, and spatial mingling of trees with different heights (**Table 1**, **Figure 2**). Observe that a stand composed of 36 regularly planted trees with a range of tree heights between 14 and 18 meters (**Figure 2(a)**) has the same *SCI* value as a stand with 8 trees and a range

of tree heights between 1 and 12 meters (**Figure 2(c)**, **Table 2**). Intuitively, these two stands have very different structures, and it may be considered an undesirable property of the *SCI* that it is not able to differentiate these stands.



Figure 1.

Spatial distribution of stems (upper panel). Triangulated irregular network calculated for stems in the upper panel (lower panel).

Table 1.

Stand characteristics of example forest stands from Figure 2.

Stand	Area [m ²]	No. of trees	Range of tree heights [m]		Standard deviation of tree heights	No. of individual heights
a)	1600	36	14 - 18	16.0	2.03	2
b)	1600	36	1 - 21	11.0	6.92	6
c)	1600	8	1 - 12	7.03	5.38	3
d)	1600	14	1 - 21	10.0	8.55	5



Figure 2.

Four different forest stands (a, b, c, d) having the same Structural Complexity Index (*SCI*) value. For stand characteristics see **Tables 1** and **2**.

Table 2.

Structural Complexity Index (*SCI*) and Enhanced Structural Complexity Index (*ESCI*' and *ESCI*) values of example stands from **Figure 2** and **Table 1**.

Stand	SCI	ESCI	ESCI	TIN area [m ²]	projected area [m ²]	VRM	Trees/10 m ²
a)	1.17	1.33	1.63	1295.77	1111.11	1.14	.23
b)	1.17	1.17	1.43	1295.77	1111.11	1.00	.23
c)	1.17	1.33	1.40	1295.77	1111.11	1.15	.05
d)	1.17	1.17	1.27	1295.77	1111.11	1.00	.09

The Enhanced Structural Complexity Index (ESCI)

We propose two modifications to the *SCI* to avoid the ambiguity described above and to better distinguish specific properties of structural complexity.

The two proposed modifications of the SCI towards the Enhanced SCI (ESCI) are:

1) Incorporation of triangle orientations;

2) Incorporation of triangle orientations + stem density.

The first modification enables the index to distinguish forest type a from type b, as in **Figure 2**. The two types do not differ in stem density, and trees in both stands are located on a regular grid with a spacing of 4 m. However, they differ in the range of tree heights and in the mingling of tree dimensions. In forest type b, there is a clear trend from small trees in the first row towards larger trees in the last row. This can be imagined as a forest edge in which tree height gradually increases towards the forest interior or as adjacent strips of clear cuts, each at a different age. In contrast, forest type a is composed of only two distinct tree heights; rows of 14 m high trees alternate with rows of 18 m high. The resulting TINs have the same surface area for both stands, resulting in the same *SCI* value.

To distinguish between these stand types, the orientation of triangles in the TINs is quantified by a vector ruggedness measure (*VRM*) (Equation (3)), adapted from a method proposed by Hobson (1972) and Sappington et al. (2007). Here, unit vectors are used to represent the orientation of a triangle (**Figure 3**). To centre a unit vector at triangle *i*, the cross product of triangle sides a_i and b_i is calculated. This results in a vector that is divided by its own length to standardize length to one. To ensure that all unit vectors are oriented upwards, unit vectors with a negative *z*-coordinate are mirrored. All unit vectors are summed, resulting in a new vector whose strength (*VSTR*, Equation (2)) is divided by the number of triangles in the TIN (*n*) and subtracted from 2 (Equation (3)). The resulting *VRM* is a dimensionless measure that ranges from 1 (non-rugged) to 2 (most rugged).

The *ESCI*' is calculated by multiplying the *SCI* with the *VRM* (Equation (4)). Based on the *ESCI*' forest types a (*ESCI*' = 1.6) and b (*ESCI*' = 1.3) can clearly be distinguished (see **Table 2**).

$$VSTR = \left| \sum_{i=1}^{n} \frac{a_i \times b_i}{|a_i \times b_i|} \right|$$
(2)

$$VRM = 2 - \frac{VSTR}{n}$$
(3)

$$ESCI' = SCI * VRM \tag{1}$$

The second modification enables one to discriminate among forest types a and c and types b and d (**Figure 2**). Compared to forest types a and b, types c and d contain fewer trees (8 and 12 trees respectively) and cover different height ranges (**Table 1**).

Since stem density is considered an important structural characteristic (e.g. Pretzsch, 2009), the number of stems per unit area is included in the *ESCI* by multiplying the number of stems per 10 m² with the *ESCI* (Equation (5)). Hence, the index calculated for stands with low stem densities is lower than for stands with high stem densities. To avoid too much weight being assigned to the stem density, a value of 1 is added to the number of stems per 10 m². The *ESCI* is a dimensionless measure >1 that increases with stem density, the intermingling of trees with different attributes, and differences between tree attributes.

$$ESCI = ESCI'^* (1 + No. \text{ of stems per } 10 \text{ m}^2)$$
(5)

Case Study from Mengsong, Xishuangbanna, Yunnan, China

Data Collection and Analysis

To investigate the *ESCI* and the *ESCI*' compared to the *SCI*, these indices were calculated for 28 plots of a forest inventory carried out in Mengsong township, Xishuangbanna, Yunnan, China (UTM/WGS84: 47 N 656355 E, 2377646 N, alt = 1600 m) (**Figure 4**). Plots cover the study site along a disturbance gradient from heavily disturbed shrub land, through secondary regrowth to mature montane rainforest stands. Each plot consists of nine circular 10 m-radius sub-plots (314.16 m²) arranged on



Figure 3.

Unit vectors centred at each triangle of a TIN. The vector ruggedness measure (*VRM*) quantifies the dispersion of these vectors: (a) Low *VRM* (low vector dispersion); (b) High *VRM* (high vector dispersion). *VRM* is used as a measure of the diversity of triangle orientations, hence a measure to describe the mingling of tree dimensions.



Figure 4.

Location of the study site Mengsong in Xishuangbanna, China. Black squares show plot locations within the site.

a square grid with 50 m spacing (**Figure 5**). For sub-plots larger 250 m², mean and median *SCI* values have been shown to be scale-invariant (Zenner, 2005).

In each sub-plot, *DBH* and position (azimuth and distance to sub-plot centre) were recorded for all trees with a $DBH \ge 10$ cm. *SCI*, *ESCI*', and *ESCI* were calculated for each sub-plot using basal area of each tree as the z-coordinate. Sub-plots without at least three trees were assigned *SCI*, *ESCI*' and *ESCI* values of zero. The *R* statistical software (R Core Team, 2012) and the *geometry* package (Grasman et al., 2011) were used for the Delaunay triangulation. Per plot values were derived by averaging all 9 sub-plot values per plot.

In five sub-plots (No. 1, 3, 5, 7, 9) per plot, insects were collected using Malaise traps in April-May 2011. Malaise traps were left in place for one week per plot and for each plot, species composition was determined using high-throughput DNA metabarcoding of the cytochrome oxidase subunit I (COI) barcode gene (Yu et al., 2012). Insect species were approximated with 97% threshold-similarity Operational Taxonomic Units (OTUs), each representing a cluster of similar COI sequences.

Insect community composition was examined by non-metric multidimensional scaling (NMDS) (Legendre & Legendre, 1998; Quinn & Keough, 2002). NMDS maps the position of plots in species space, in our case represented by the Jaccard dissimilarities of Hellinger transformed per plot OTU counts (Yu et al., 2012), onto a predefined number of axes in an iterative search for an optimal solution. NMDS is commonly regarded as the most robust unconstrained ordination method in community ecology (Minchin, 1987). The R package vegan (Oksanen et al., 2007) was used for the ordination analysis. To investigate whether a relationship exists between structural complexity and turnover in insect species composition we used the function *envfit* in the package *vegan* to calculate the fit of environmental variables to the ordination scores. envfit maximises the correlation of environmental variables to the matrix of ordination scores and uses a permutation test to evaluate the probability of obtaining the resulting or a higher r^2 value. Pvalues stated here are based on 1000 permutations. To test whether differences in correlation coefficients between SCI, ESCI'

and *ESCI* are significant we took 1000 subsamples (without replacement) of 26 observations from our field data. NMDS ordinations were calculated for each subsample and the corresponding r^2 values with *SCI*, *ESCI*² and *ESCI* were calculated. Subsequently, we applied Mann-Whitney U tests to investigate whether differences of mean r^2 values of the replicated samples were significant.

Results

From a total of 2890 trees that were recorded, each sub-plot contained on average 11.47 trees, with numbers ranging from 0 to 34 trees per sub-plot. The basal area of individual trees varied from 78.61 cm² to 11,750 cm².

Across all sub-plots, *ESCI* values are consistently higher and cover a larger range than do *ESCI*' values. The same is observed if *ESCI*' and *SCI* are compared (**Figure 6**).

In addition to the observed differences in index value range, the indices treat special cases of tree arrangements differently. For example, sub-plots 49_2 and 372_9 (Figure 7, upper panels) have similar *SCI* values (1% difference), but their *ESCI*' values differ substantially (22.75% difference) (Table 3). However, in Figure 7, sub-plots 49_2 and 90_1 still have similar structural complexities, as measured by the *ESCI*'.



Figure 5.

Plot design: Cluster plot consisting of 9 sub-plots arranged on a square grid with 50 meters spacing between subplot centres.



Figure 6.

Box and ladder plot comparing paired *SCI* and *ESCI*' observations (left) and paired *SCI* and *ESCI* observations (right).



Figure 7.

Depiction of tree positions (*xy*-coordinates; in meters) complemented with basal area (*BA*) as the *z*-coordinates of sub-plots listed in **Table 3**. Upper panels: Sub-plot 49_2 has a *SCI* value similar to sub-plot 372_9. Based on the *ESCI*' value 49_2 is similar to 90_1. Lower panels: Sub-plots with similar *ESCI*' values which would be distinguished by their *ESCI* values. Note: *ESCI* ranks the sub-plots in a different order than *SCI*.

Table 3.

Index values calculated for sub-plots shown in Figure 7.

sub-plot	projected area [m ²]	TIN area [m ²]	SCI	ESCI'	ESCI	VRM	No. of trees
49_2	47.53	24911.01	524.15	528.26	595.51	1.01	4
372_9	222.35	119133.61	535.79	1014.81	1790.07	1.89	24
90_1	113.11	33290.73	294.31	532.38	667.95	1.81	8
189_9	36.94	40896.36	1107.08	1445.78	1629.86	1.31	4
398_3	43.86	40417.48	921.44	1452.41	1729.81	1.58	6
192_8	166.83	133784.66	801.91	1473.12	2551.62	1.84	23

The effect of the incorporation of the stem density term becomes clear when the sub-plots in **Figure 7** (lower panels) are examined. According to their *SCI* values, sub-plot 189_9 has a higher structural complexity than does 398_3 or 192_8. *ESCI* ranks sub-plot 189_9 as the least structurally complex and 192_8 as the most structurally complex (**Table 3**).

The number of NMDS dimensions was fixed to three after visual examination of the scree plot of stress vs dimensions. Three dimensions provided a satisfactory stress value of 9.24%. There were highly significant (p < .001) correlations between all three structural complexity indices and the NMDS ordination. The *SCI* shows the lowest r^2 value of .51 followed by the *ESCI*' with $r^2 = .54$ and *ESCI* with $r^2 = .59$. Mean r^2 values based on 1000 sample replicates were significantly different among all combinations of indices (p < .0001).

Discussion

The number of stems per 10 m^2 is a critical factor in the *ESCI* formula. In the data set used in this study, we found that $(1 + \text{No. of stems per m}^2)$ results in a range of values from 1 to

2.02, for 0 and 32 trees per sub-plot respectively. This range does not assign a weight to the number of stems that outweighs the other terms of the equation. SCI, VRM and stem density are hence of approximately equivalent importance in determining the ESCI value. For other data sets with a higher number of stems per unit area, the relation to the base of 10 m^2 probably leads to density values that dominate the resulting ESCI value. A higher stem density might occur, for example, in different forest types, succession stages with high stem numbers, or if the DBH threshold is lower than 10 cm. In these cases it might be more informative to calculate ESCI' and stem density separately to describe structural complexity and to make inferences. The addition of 1 to the number of stems per 10 m^2 assigns a greater weight to low stem densities but has little influence if stem densities are high. This weighting takes into account that with increasing density, the potential for a spatial differentiation decreases.

Sub-plots in which stems were mapped for this analysis cover an area of 314.16 m², which is sufficiently large for the calculation of the *SCI* (Zenner, 2005). Nevertheless, due to the high scale dependency of forest structure (Franklin et al., 2002; Zenner, 2005), this plot size is probably at the lower bound of an adequate size. A further investigation of the variability of *ESCI* and *ESCI* values at varying scales is recommended to achieve deeper insights regarding the minimum plot size for calculations of these indices.

In the structural assessment of forests, the inclusion of small diameter trees was found to enhance the detection of structure types (Zenner et al., 2011). In future studies it might be interesting to analyse whether *ESCI*' and *ESCI* show a similar response if trees with a DBH < 10 cm are considered.

Comparing SCI, ESCI' and ESCI, we find that the stronger emphasis on the intermingling of tree dimensions through the incorporation of the VRM as a measure of surface ruggedness improves the ability of the index to discriminate among stand conditions. This might result in a better utility of the index. For example, it could be used for a separation of forest edges with increasing tree height from other spatial arrangements of tree dimensions typical for forest interiors. Such discrimination makes sense from an ecological point of view since forest edges are characterized by a distinct microclimate and resource spectrum, with corresponding effects on species composition and abundance (McDonald & Urban, 2006). If in addition to the VRM, the stem density of a forest is taken into account, the ability to unambiguously characterize forest structural complexity is increased again. Ecologists and forest managers likewise consider stem density as an important aspect of structural complexity because it determines the mean growing space per plant and hence is an indicator of competition for resources within the stand (Pretzsch, 2009). Changing stem density is the principal way forest managers manipulate forests (Davis et al., 2001), and these changes alter forest habitats with consequences for forest organisms.

A possible ecological relevance of the modifications of the *SCI* is indicated by the correlation of the index values with the NMDS ordination, which describes turnover in insect species composition. Compared to the *SCI*, significantly stronger correlations with the NMDS ordination were observed for the *ESCI* and the *ESCI*. The observed correlations support the habitat heterogeneity hypothesis by suggesting that the structural complexity of a habitat influences the insect species community composition. The higher r^2 values for *ESCI* and *ESCI* suggest that

these indices perform at least as well and possibly better in detecting this relationship. Nevertheless, the increase in r^2 values was only moderate, and hence it would be valuable to test the indices for structural complexity against other data sets from different geographic regions and taxa and to assess associations at multiple spatial scales.

Conclusion

The results of this study show that the suggested modifications to the *SCI* are valuable improvements that increase the ability to characterize the structural complexity of forests. *ESCI* and *ESCI* allow for a more complete view of a forest structure than the *SCI*. This makes these indices relevant to ecologists, forest scientists, and forest managers who are interested in the relationship between ecosystem structure and biodiversity.

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