## ORIGINAL PAPER

# Disturbances, elevation, topography and spatial proximity drive vegetation patterns along an altitudinal gradient of a top biodiversity hotspot

Pedro V. Eisenlohr · Luciana F. Alves · Luís Carlos Bernacci · Maíra C. G. Padgurschi · Roseli B. Torres · Eduardo M. B. Prata · Flavio Antonio M. dos Santos · Marco Antônio Assis · Eliana Ramos · André Luís C. Rochelle · Fernando R. Martins · Mariana C. R. Campos · Fernando Pedroni · Maryland Sanchez · Larissa S. Pereira · Simone A. Vieira · José Ataliba M. A. Gomes · Jorge Y. Tamashiro · Marcos A. S. Scaranello · Cora J. Caron · Carlos Alfredo Joly

Received: 8 May 2013/Accepted: 21 August 2013/Published online: 31 August 2013 © Springer Science+Business Media Dordrecht 2013

**Abstract** The correlation between vegetation patterns (species distribution and richness) and altitudinal variation has been widely reported for tropical forests, thereby providing theoretical basis for biodiversity conservation. However, this relationship may have been oversimplified, as many other factors may influence vegetation patterns, such as disturbances, topography and geographic distance. Considering these other factors, our primary question was: is there a vegetation pattern associated with substantial altitudinal variation (10-1,093 m a.s.l.) in the Atlantic Rainforest—a top hotspot for biodiversity conservation—and, if so, what are the main factors driving this pattern? We addressed this question by sampling 11 1-ha plots, applying multivariate methods, correlations and variance partitioning. The *Restinga* (forest on sandbanks along the coastal plains of Brazil) and a lowland area that was selectively logged 40 years ago were floristically isolated from the other plots. The maximum species richness (>200 spp. per hectare) occurred at

A. L. C. Rochelle · F. R. Martins · L. S. Pereira · J. Y. Tamashiro ·

M. A. S. Scaranello · C. J. Caron · C. A. Joly

Present Address: P. V. Eisenlohr (⊠) Departamento de Botânica, Universidade Federal de Minas Gerais, Belo Horizonte, MG 31270-901, Brazil e-mail: pedrov.eisenlohr@gmail.com

L. F. Alves · L. C. Bernacci · R. B. Torres · J. A. M. A. Gomes Centro de Pesquisa e Desenvolvimento de Recursos Genéticos Vegetais, Instituto Agronômico de Campinas (IAC) CP 28, Campinas, SP 13012-970, Brazil

E. M. B. Prata

P. V. Eisenlohr · L. F. Alves · M. C. G. Padgurschi · F. A. M. dos Santos ·

Departamento de Biologia Vegetal, Instituto de Biologia, CP 6109, Universidade Estadual de Campinas (UNICAMP), Campinas, SP 13083-970, Brazil

Programa de Pós-Graduação em Botânica, CP 478, Instituto Nacional de Pesquisas da Amazônia (INPA), Manaus, AM 69060-001, Brazil

approximately 350 m a.s.l. (submontane forest). Gaps, multiple stemmed trees, average elevation and the standard deviation of the slope significantly affected the vegetation pattern. Spatial proximity also influenced the vegetation pattern as a structuring environmental variable or via dispersal constraints. Our results clarify, for the first time, the key variables that drive species distribution and richness across a large altitudinal range within the Atlantic Rainforest.

**Keywords** Atlantic Rainforest · Conservation hotspot · Multivariate analysis · Species distribution · Species richness

# Introduction

The investigation of vegetation patterns has been recognized as a crucial step in providing theoretical basis for biodiversity conservation (e.g., Ivanauskas et al. 2006; Marques et al. 2011). One of the most widely reported type of vegetation patterns for tropical forests is the variation in species distribution and richness along elevational gradients (Austin and Greig-Smith 1968; Beals 1969; Gentry 1988; Rodrigues and Shepherd 1992; Auerbach and Shmida 1993; Rahbek 1995; Lieberman et al. 1996; Vázquez and Givnish 1998; Oliveira-Filho and Fontes 2000; Zhao et al. 2005; Sanchez et al. 2013). However, it is unlikely that studies on this subject have found a realistic model because species distributions and richness can be influenced by other factors, such as the geographic distance between locations (e.g., Hubbell 2001; Diniz-Filho et al. 2012), disturbances of human or natural origin (e.g., Connell 1978; Vázquez and Givnish 1998; Laurance 2001; Pereira et al. 2007) and topography (e.g., Bourgeron 1983; O'Brien et al. 2000; Bohlman et al. 2008). When investigating ecological patterns, controlling or accounting for a set of factors that are potentially important for determining these patterns is essential to obtaining reliable results (Simberloff 1983; Rey Benayas and Scheiner 2002).

The Brazilian Atlantic Rainforest, a top hotspot for biological conservation (Myers et al. 2000; Laurance 2009; Ribeiro et al. 2011), lacks realistic, comprehensive models for understanding local floristic patterns. Despite this problem, several studies have addressed important questions concerning vegetation patterns along elevation gradients. Oliveira-Filho and Fontes (2000) reported that altitude is one of the principal factors influencing floristic differentiation in the Atlantic Forest sensu lato of southeastern Brazil. In an

M. A. Assis · E. Ramos

M. C. R. Campos School of Plant Biology, University of Western Australia, M089 35 Stirling Highway, Crawley, WA 6009, Australia

F. Pedroni · M. Sanchez Departamento de Ciências Biológicas e da Saúde, Universidade Federal de Mato Grosso, Pontal do Araguaia, MT 78698-000, Brazil

## S. A. Vieira

Departamento de Botânica, Instituto de Biociências de Rio Claro, CP 199, Universidade Estadual Paulista Júlio de Mesquita Filho, Rio Claro, SP 13506-900, Brazil

Núcleo de Estudos e Pesquisas Ambientais (NEPAM), Universidade Estadual de Campinas (UNICAMP), Campinas, SP 13083-867, Brazil

analysis of the floristic similarity between rainforest and semi-deciduous forest sites in São Paulo State, Ivanauskas et al. (2000) detected the establishment of certain groups relative to the elevation. Scudeller et al. (2001) found that the distinctions between the provincial and coastal Atlantic plateaus are linked to altitude, precipitation and temperature. Sanchez et al. (2013) found that the Serra do Mar rainforests along the seashore, mountain slopes and highlands differed dramatically in their floras, physiognomies and origins. At the local scale, Rodrigues and Shepherd (1992) and Moreno et al. (2003) also found consistent floristic links relative to altitude. Thus, well-supported relationships between altitude and floristic variations have been reported to the Atlantic Forest.

Our objective was to enhance the existing theoretical support for conservation initiatives for protecting lands in moist tropical forests, considering the existing need for more comprehensive approaches that include a set of potentially relevant predictors for vegetation patterns along altitudinal gradients. Strong particularities and novelties of this study are the thorough documentation of: (1) a broad and standardised sampling effort (11 ha of vegetation in equally sampled plots); (2) sampling conducted along a single conservation unit (Serra do Mar State Park) while minimising environmental factors that might interfere with the investigation; (3) the measurement of variable sets for the few factors that might interfere with the investigation, such as the topography and disturbance history, allowing for a consistent analysis of the vegetation patterns throughout the elevational range.

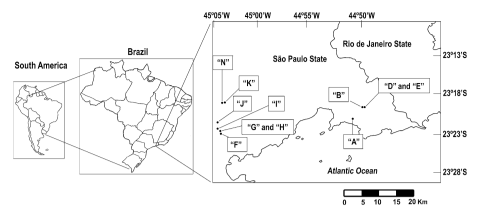
We addressed the following questions: (1) is there a pattern in the floristic composition associated with the altitudinal variation that we analysed (10–1,093 m) for trees in the Atlantic Rainforest; i.e., do areas having a similar altitude exhibit higher species similarity than those areas that are more distant with respect to altitude? (2) Does the species richness differ from areas of lower altitude to areas of higher altitude? (3) What are the principal factors driving the variation in tree species composition and richness?

## Methods

#### Study area

We studied the Atlantic Rainforest in the Picinguaba and Santa Virginia Research Station of the Serra do Mar State Park, near Ubatuba and São Luis do Paraitinga municipalities in São Paulo State, southeastern (SE) Brazil (Fig. 1), which covers an elevation range of 1,083 m (10–1,093 m a.s.l.). The coordinates and other data on the areas studied are presented in Table 1; additional data can be found in Alves et al. (2010) and Joly et al. (2012). The climate is tropical (Af according to the Köeppen system), with higher rainfall in summer (Morellato et al. 2000) and without significant rainfall variations along the slope. The topography of the region includes the coastal plain, isolated hills and mountains of the elongated "Morraria" Coast and the inner slopes of the "Serrania" Coast (Ponçano et al. 1981).

The soils are generally low in nutrients, with low fertility and high levels of aluminium. In area "A" (*Restinga* forest), the soil type is classified as Quartzipsamment, whereas in the other areas the soils are Inceptisols (Table 1). Shallow and well-drained soils predominate along the slope, with the exception of the *Restinga* forest. Soils become more aged with elevation, according to "Kr" and "Ki" weathering indices—at the top, there is predominance of oxides (high levels of  $Al_2O_3$ ). Soil carbon and nitrogen contents increase significantly with altitude. On the other hand, pH does not strongly vary along the elevational gradient (in general <4). The litter production has been higher at lower elevations (8–10 Mg ha<sup>-1</sup> at plots lower than 100 m a.s.l. and 7 Mg ha<sup>-1</sup> above 400 m a.s.l.). Those



**Fig. 1** Locations of the 11 study areas along an altitudinal gradient in the Atlantic Rainforest (SE Brazil). The codes for the localities are presented in Table 1

data were provided by SC Martins (unpublished data), Sousa Neto et al. (2011), Vieira et al. (2011) and Sanchez et al. (2013).

The codes for the localities where the sampling was completed are listed in Table 1 in alphabetical order according to their positions in the landscape. These codes follow those of the "BIOTA Functional Gradient" Project (Joly et al. 2012). It should be noted that the altitudinal belts (Table 1) do not follow the Brazilian classification system for vegetation (IBGE 2012), but rather used an alternative classification based on the field observations of the research team. The relative locations of the 11 1-ha areas are shown in Fig. 1. Area "A" (*Restinga* or Sandy Forest, at approximately sea level) is located where the major local rivers flow into the region. In this area, the soil is exposed to the seasonal water table (César and Monteiro 1995). Areas "B", "D", "E", "F", "G", "H", "I" and "J" are located at increasing altitudes along the slope of the Serra do Mar, whereas areas "K" and "N" are located at the summit (Table 1). According to long-time residents, area "F" was selectively logged ~40 years ago for the tree *Hieronyma alchorneoides* (Phyllanthaceae). A more detailed description of the study area was provided by Joly et al. (2012).

Data collection and preparation of matrices

We measured all individual trees with a diameter equal to or greater than 4.8 cm at 1.30 m above the ground (DBH) in each 1-ha area (100 m× 100 m, a widely used sampling design in tropical forests, thus allowing comparisons with other studies) (Table 1). The sampling protocol was described in detail by Alves et al. (2010) and Joly et al. (2012).

To investigate the patterns of species distribution and variation in species richness, we first constructed three matrices, each of which included the 11 study areas. The first matrix contained data on the numbers of individuals of each species. The second matrix was composed of the following environmental variables (Table 1): topographic variables—aspect, slope and standard deviation of the slope (SD slope is reported in the tables and figures), 3D surface area (SurfArea), average elevation (Elevation) and elevational range (ElevRange); disturbance variables—gaps.area<sup>-1</sup> (Gaps), basal area of standing dead stems (BAdead) and number of trees with two or more stems (MultStem); and two synthetic climatic variables—annual precipitation (AnnuPrec) and annual mean temperature (AnnMTemp). The units used for each variable are provided in Table 1.

| Table | 1 Codes and other en                        | Table 1 Codes and other environmental information for the 11 sampled areas along an altitudinal gradient in the Atlantic Rainforest, SE Brazil  | the 11 samp                        | oled areas alon                    | g an altitu               | dinal gradient                 | in the Atla                  | ntic Rain                                  | forest, SE Bra                            | zil                        |                              |
|-------|---|---|------------------------------------|------------------------------------|---------------------------|--------------------------------|------------------------------|--|---|----------------------------|------------------------------|
| Areas | Areas Forest type                           | Geographic<br>coordinates<br>(lat, long)  | Average<br>elevation<br>(m a.s.l.) | Elevational<br>range<br>(m a.s.l.) | Aspect                    | Slope<br>(%) ±SD               | Multiple<br>stemmed<br>trees | Total gaps. area $^{-1}$ (m <sup>2</sup> ) | Basal area of standing dead stems $(m^2)$ | Annual<br>rainfall<br>(mm) | Annual mean temperature (°C) |
| "A"   | Restinga                                    | 23°21′21′′S, 44°51′04′′W  | 10                                 | 1                                  | -0.956                    | $1.10 \pm 0.7$                 | 117                          | 2,662                                      | 0.40                                      | 2,406                      | 22.60                        |
| "В"   | Lowland Forest                              | 23°20'14''S, 44°50'06''W  | 46                                 | 23                                 | -1.000                    | $14.7\pm5.8$                   | 69                           | 2,700                                      | 0.44                                      | 2,406                      | 22.60                        |
| "D"   | Lowland Forest                              | 23°20'08"S, 44°50'00"W  | 57                                 | 26                                 | -0.807                    | $12.6\pm5.5$                   | 65                           | 3,526                                      | 0.89                                      | 2,324                      | 22.25                        |
| "E"   | Lowland Forest                              | 23°20'04"S, 44°49'57"W  | 73                                 | 25                                 | -0.876                    | $11.1 \pm 4.8$                 | 75                           | 2,774                                      | 1.03                                      | 2,324                      | 22.25                        |
| "Ғ"   | Lowland Forest                              | 23°22′53″S, 45°04′46″W  | 105                                | 28                                 | -0.956                    | $11.9\pm6.8$                   | 184                          | 4,770                                      | 1.12                                      | 2,582                      | 21.90                        |
| "Є    | Submontane Forest                           | 23°22'26''S, 45°04'51''W  | 188                                | 22                                 | -0.987                    | $12.2 \pm 4.6$                 | 46                           | 1,970                                      | 1.95                                      | 1,975                      | 18.31                        |
| "Н"   | Submontane Forest                           | 23°22'25''S, 45°04'55''W  | 208                                | 16                                 | -0.988                    | $9.20\pm3.5$                   | 56                           | 3,245                                      | 1.73                                      | 1,975                      | 18.31                        |
| "I"   | Submontane Forest                           | 23°22'01"S, 45°05'01"W  | 350                                | 48                                 | -0.852                    | $27.3\pm6.3$                   | 48                           | 1,627                                      | 0.83                                      | 1,865                      | 17.46                        |
| "΄    | Submontane Forest                           | 23°21′58″S, 45°05′03″W  | 375                                | 46                                 | -1.000                    | $25.5\pm8.0$                   | 31                           | 1,749                                      | 2.11                                      | 1,865                      | 17.46                        |
| "К"   | Montane Forest                              | 23°19′32″S, 45°04′05″W  | 1,066                              | 47                                 | 0.850                     | $25.5\pm8.3$                   | 4                            | 2,314                                      | 3.16                                      | 1,724                      | 16.28                        |
| "N"   | Montane Forest                              | 23°19′36′′S, 45°04′32″W   | 1,025                              | 34                                 | 0.787                     | $15.8\pm5.4$                   | 64                           | 3,860                                      | 3.14                                      | 1,724                      | 16.28                        |
| Other | data are reported by A ion (IBGE 2012), but | Other data are reported by Alves et al. (2010) and Joly et al. (2012). Altitudinal belts (see :"Forest type" column) do not follow the Brazilian classification system for vegetation (IBGE 2012), but rather used an alternative classification based on the field observations of the BIOTA research team | et al. (2012)<br>assification h    | . Altitudinal b<br>based on the fi | elts (see :<br>eld observ | "Forest type"<br>ations of the | column) d<br>BIOTA rese      | o not foll<br>earch tean                   | ow the Brazili<br>1                       | ian classif                | cation system for            |

Biodivers Conserv (2013) 22:2767–2783

SD standard deviation

To acquire data on the topographic variables, we first used the elevation information from each  $10 \times 10$  m corner point determined in the field using high-precision equipment (a total station) to create a digital elevation model (DEM) at a 1-ha scale. The DEM was created using the "topo-to-raster" tool, an interpolation method implemented in ArcGIS to generate DEMs (Hutchinson 1989). This procedure allowed us to store topographic information in a raster format with a high spatial resolution. The new elevation data from the DEM was fitted to probability density functions to extract the expected value (e.g., the mean for a Gaussian distribution) and deviation for each plot. The 3D surface variable was also estimated from the DEM. We estimated gaps area<sup>-1</sup> as the surface area directly under a canopy opening, extending to the base of the forest edge, according to the extended gap definition of Runkle (1982). Gaps are relatively small-sized in our Atlantic forest site. The mean gap area varies between 60.4 and 115.2  $m^2$ , and the maximum total gap area per plot was 2,598.9 m<sup>2</sup>, corresponding to less than 30 % of the plot area. The climatic variables were extracted from WORLDCLIM (Hijmans et al. 2005). Although some authors do not recommend using WORLDCLIM for climatic interpolation in mountainous areas (Sklenář et al. 2008), to our knowledge, there is no disadvantage in the Atlantic Forests.

Finally, we constructed a coordinate matrix, containing the latitude and longitude of each area, for the purpose of assessing and controlling spatial autocorrelation and performing variance partitioning (below).

Variations in floristic patterns and correlations with predictor variables

To ordinate the 11 areas without interference from any environmental variable, we performed a DCA (Detrended Correspondence Analysis) using PC-ORD 6.0 (McCune and Mefford 2011). We discarded the NMS (Non-Metric Multidimensional Scaling), a widely recommended method (McCune and Grace 2002), following the same reasons that Santos et al. (2012). We removed the 243 singletons because the Chi squared distance used in DCA is sensitive to rare species (McCune and Grace 2002). The DCA was performed using the options "down-weight rare species" and "rescale axes".

We used OLS multiple linear regressions (Quinn and Keough 2002) to analyse the predictability of the environmental variables on the floristic variation summarised by the two principal axes of the DCA. We initially discarded variables with very weak correlations (r < 0.3) with the DCA axis and removed colinearities, keeping only the most important variable (>r with the respective DCA axis) in each group of collinear variables. After this procedure, no variables remaining with a variance inflation factor >10 (Quinn and Keough 2002), and we then chose the best model (lowest AICc—Corrected Akaike Information Criterion; Burnham and Anderson 2002). However, we detected a strong outlier in the linear trend of the OLS models, the *Restinga* plot (area "A"). We then opted to present both DCAs (DCA1, with *Restinga*, and DCA2, without *Restinga*), illustrating the overall floristic patterns, while discarding the linear model that included *Restinga*. Further information on the OLS procedure can be found in the Common procedures for all OLS models section below.

Variations in species richness and correlations with predictor variables

To avoid a dependent relationship between the numbers of individuals and species, we estimated the species richness by running an individual-based rarefaction analysis (Gotelli and Colwell 2001) using ECOSIM, version 7.71 (Gotelli and Entsminger 2004). We compared the estimated species richness among sites using the same sampling effort

(numbers of individuals) with 95 % confidence intervals. This level of sampling corresponds to the number of individuals in area "B" (1,082 trees) because this site had the lowest value among the studied areas. The obtained species richness was hereafter treated as the estimated species richness.

Finally, we performed a multiple linear regression analysis (OLS) to investigate whether changes in species richness could be explained by our environmental predictors. The colinearities were removed, as described above, with one exception: although altitude was less correlated with species richness than was basal area of standing dead stems, the effect of removing the multicolinearity was more powerful when we discarded the last variable; therefore, we decided on the altitude. We used the same methods described in the floristic variation section above to select the best model (see also the Common procedures for all OLS models section below).

Common procedures for all OLS models

We followed the general format of the procedures proposed by Eisenlohr (2013) for the aforementioned OLS multiple linear regression models and heeded the caveats identified by this same author. We used correlograms to apply Moran's I coefficient as an indicator of spatial autocorrelation (SAC) using SAM 4.0 (Rangel et al. 2010) to check for possible violations of the assumption of spatial independence in the residuals of both the full and selected OLS models (Diniz-Filho et al. 2003; Diniz-Filho et al. 2008). The number and size of distance classes used the defaults for SAM. The significance of SAC was detected by sequential Bonferroni criteria (Fortin and Dale 2005). When significant SAC in the residuals of the OLS models was detected, we used spatial filters (Moran's Eigenvector Maps—MEMs; Dray et al. 2006), obtained using the "spacemakeR" package in the R environment (The R Foundation for Statistical Computing 2012), as additional predictors. We selected MEMs until we found randomness (non-significant SAC) in the residuals. When we found SAC in the full model (i.e. prior to AICc selection), we used MEMs as fixed variables in the selection (Diniz-Filho et al. 2008).

We confirmed the assumptions of homoscedasticity and linearity by creating graphical displays of the residuals analysis (X: predicted values; Y: residuals), demonstrating an even distribution of the points above and below the zero-residual horizontal line (Quinn and Keough 2002). The normality of the residuals was confirmed by using the Shapiro–Wilk test.

The significance of the linear model was assessed using the F test. The relative influence of each environmental variable on the floristic data was assessed based on partial regression coefficients and t tests. In all of the tests, we used 5 % as the significance level.

## Variance partitioning

We also performed variance partitioning using a partial regression analysis to obtain the components that explain the variation in each response variable (ordination axes and species richness) (Legendre and Legendre 2012). The environmental variables were the same as in the respective OLS models, and the spatial variables (MEMs) were forward-selected using the "packfor" package for the R environment. The forward procedure is justified by our requirement to obtain only MEMs that provide significant contributions to the response variables (Bellier et al. 2007). Note that the MEMs selected for variance partitioning were not necessarily the same MEMs chosen for the removal of spatial autocorrelation from the OLS residuals because the objectives were distinctly different.

Axis 1 of the DCA2 did not have any significant spatial variables, and thus, we did not perform a variance partitioning in this case.

# Results

Considering the overall sampling from 11 ha of the Atlantic Rainforest in southeastern Brazil, we registered a total of 16,796 trees, 563 species, 195 genera and 68 families (including 'singletons'). A summary of the vegetation data from each area is presented in Table 2, and other details are provided by Joly et al. (2012).

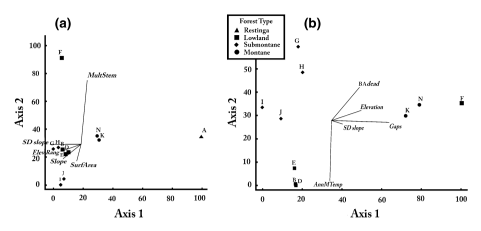
Variations in floristic patterns and correlations with predictor variables

The first axis of DCA1 showed contrast between areas "A" (*Restinga*), "K"+"N" (montane forests) and the other areas (Fig. 2a). Topography (slope and SD slope) and elevational range were the most highly correlated with this axis (Table 3). Considering the strong differences between soils of *Restinga* and other areas, we might infer that the edaphic component was also important in such floristic differentiation. For the second axis, areas "F" (lowland forest, selectively logged) and "I"+"J" (submontane forest) were dissociated from each other and from other areas (Fig. 2a). The disturbance variables (mainly multiple stems) appeared to be the most relevant in this case (Table 3).

The first axis of DCA2 showed dissociation between areas "F", "K"+"N" and the other areas (Fig. 2b). Disturbance (total gap area and basal area of standing dead stems) and elevation variables once again assumed great importance (Fig. 2b; Table 3). The linear model indicated gaps and average elevation as being significant variables (Table 4). The second axis suggested a significant role for the basal area of standing dead stems and the annual mean temperature (Fig. 2b); however, only the spatial filter (MEM) added to account for the spatial autocorrelation was significant (Table 4). Thereafter, climatic variables did not present a primary role in the floristic patterns. Since there is predominance of oxides at the top and soil carbon and nitrogen contents increase along the slope, soil could also be important to explain the floristic dissociation between "K"+"N" and the

| Site<br>(area) | Number<br>of trees | Species richness | Genus<br>richness | Family richness |
|----------------|--------------------|------------------|-------------------|-----------------|
| "A"            | 1,671              | 85               | 61                | 32              |
| "В"            | 1,082              | 136              | 87                | 39              |
| "D"            | 1,286              | 156              | 94                | 42              |
| "Е"            | 1,240              | 141              | 93                | 41              |
| "F"            | 1,381              | 106              | 72                | 38              |
| "G"            | 1,496              | 151              | 101               | 41              |
| "H"            | 1,528              | 152              | 98                | 41              |
| "I"            | 1,993              | 203              | 111               | 50              |
| "J"            | 1,832              | 211              | 117               | 49              |
| "K"            | 1,851              | 178              | 83                | 44              |
| "N"            | 1,436              | 148              | 73                | 39              |
| All            | 16,796             | 563              | 195               | 68              |

**Table 2**Number of trees and<br/>the species, genus and family<br/>richness for 11 sites sampled<br/>along an altitudinal gradient in<br/>the Atlantic Rainforest (SE<br/>Brazil)



**Fig. 2** DCA ordinations from species abundance data along an altitudinal gradient in the Atlantic Rainforest (SE Brazil). *Arrows* indicate a posteriori Pearson correlations with environmental variables. **a** Ordination with *Restinga* and **b** without *Restinga*. The abbreviations are defined in the Methods section. Weak (r < 0.3) and redundant variables were not represented

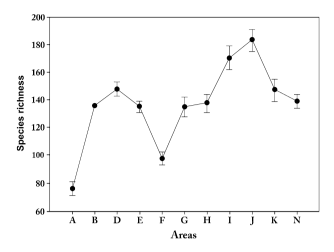
| <b>Table 3</b> DCA ordination results           for species distribution patterns | -               | DCA1 (with <i>Restinga</i> ) |        | DCA2 (without Restinga) |        |
|---|-----------------|------------------------------|--------|-------------------------|--------|
| along an altitudinal gradient in<br>the Atlantic Rainforest (SE Bra-              |                 | Axis 1                       | Axis 2 | Axis 1                  | Axis 2 |
| zil) and a posteriori Pearson cor-<br>relations for environmental                 | Eigenvalue      | 0.610                        | 0.4193 | 0.4200                  | 0.1864 |
| variables   | Gradient length | 3.419                        | 3.111  | 2.539                   | 1.513  |
|   | Gaps            | 0.003                        | 0.823  | 0.701                   | -0.113 |
|   | MultStem        | 0.271                        | 0.881  | 0.639                   | -0.052 |
|   | BAdead          | -0.116                       | -0.007 | 0.485                   | 0.492  |
|   | Slope           | -0.450                       | -0.420 | -0.102                  | 0.026  |
|   | SD slope        | -0.596                       | -0.024 | 0.308                   | -0.156 |
|   | SurfArea        | -0.225                       | -0.458 | -0.159                  | -0.039 |
|   | Elevation       | 0.054                        | -0.064 | 0.502                   | 0.269  |
|   | ElevRang        | -0.476                       | -0.293 | 0.076                   | 0.015  |
|   | AnnMTemp        | 0.182                        | 0.318  | -0.131                  | -0.665 |
| The abbreviations are defined in the "Methods" section                            | AnnuPrec        | 0.124                        | 0.503  | 0.037                   | -0.524 |

other areas. Both models exhibited high coefficients of determination (Axis 1:  $R_{Adi}^2 = 0.7380$ ; Axis 2:  $R_{Adi}^2 = 0.7175$ ).

The floristic gradients were at least moderate for all the axes (except for Axis 2 from DCA2), which was evidenced by the eigenvalues and the gradient length (Table 3). These results indicated an intermediate to high level of species turnover for the main trends of floristic variation along this altitudinal variation.

Variations in species richness and correlations with predictor variables

Areas "I" and "J" (submontane forest) were the richest in species among the 11 areas investigated (Fig. 3). The poorest locality in species richness was area "A" (*Restinga* forest), followed by area "F" (lowland forest, selectively logged) (Fig. 3).



**Fig. 3** Comparison of species richness among the 11 areas along an altitudinal gradient in the Atlantic Rainforest (SE Brazil) at the point of equal numbers of individuals (individual-based rarefaction algorithm). Narrow bars indicate 95 % confidence intervals

Multiple stems negatively predicted the variation in estimated species richness and the standard deviation of the slope positively predicted that variation (Table 4). This model also demonstrates a high predictive power ( $R_{Adi}^2 = 0.8362$ ).

Variance partitioning

We detected the primary role of the spatially structured environmental factors (fraction [b]) on the floristic patterns emerged from Axis 2 of the DCA2 (Fig. 4a); 22 % of the species distribution remained undetermined (fraction [d]; Fig. 4a). The variations in species richness also indicated a high level of explanation for [b] and demonstrated a high level of explanation for the sum of components [a], [b] and [c], whereas component [d] represented only 3 % of the explanation (Fig. 4b). The isolated spatial fraction ([c]) was more relevant than the isolated environmental fraction ([a]) in both variance partitioning calculations.

# Discussion

The ordination analysis indicated several floristic patterns: (i) the strong dissociation of *Restinga* ("A") relative to the other areas, most likely due to topographic and edaphic differences (the *Restinga* soil is classified as a Quartzipsamment and is subject to seasonal exposure to the water table); (ii) the strong dissociation of the lowland area ("F"), which appears to be influenced by past disturbance, reflected by an increase in the number of the multiple stemmed trees and total gaps.area<sup>-1</sup> variables; (iii) the equivalence of blocks "I"+"J", most likely due to topographic factors; and (iv) "K"+"N", which might be influenced by gaps.area<sup>-1</sup> and the elevation average. Explanations for these patterns are possible when considering the geomorphological processes, disturbances and/or elevation zonation.

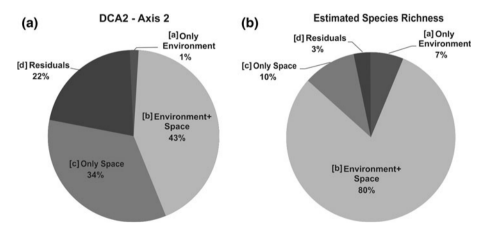
The geomorphological processes occurring in the *Restinga* forest area (Araújo and Lacerda 1987; Assis et al. 2011) are markedly different from those in the other studied areas, as described by Sanchez et al. (2013). The processes operating in *Restinga* lead to a

| U                |       |                   | ,       |                |         |
|------------------|-------|-------------------|---------|----------------|---------|
|                  | Beta  | Std. Err. of Beta | В       | Std. Err. of B | p level |
| Axis 1           |       |                   |         |                |         |
| Constant         | -     | -                 | -141.85 | 48.16          | 0.02    |
| Gaps             | 0.74  | 0.17              | 0.07    | 0.02           | <0.01   |
| Elevation        | 0.55  | 0.17              | 0.13    | 0.04           | 0.01    |
| Axis 2           |       |                   |         |                |         |
| Constant         | -     | -                 | 3.44    | 113.70         | 0.98    |
| AnnMTemp         | 0.18  | 0.31              | 3.47    | 5.86           | 0.57    |
| MEM 1            | 1.03  | 0.31              | 155.34  | 47.00          | 0.01    |
| Species richness |       |                   |         |                |         |
| Constant         | -     | -                 | 132.712 | 13.705         | <0.01   |
| SD slope         | 0.499 | 0.14              | 6.96    | 1.84           | <0.01   |
| MultStem         | -0.68 | 0.14              | -0.46   | 0.09           | <0.01   |
|                  |       |                   |         |                |         |

 
 Table 4
 Summary of the linear regression models prepared for DCA2 and estimated species richness along an altitudinal gradient in the Atlantic Rainforest (SE Brazil)

The abbreviations are defined in the "Methods" section. Beta: standardised coefficients. B: unstandardised coefficients. Significance p level ( $\leq 0.05$ ) in bold

Axis 1:  $R_{Adj}^2 = 0.7380$ ; p < 0.01. Axis 2:  $R_{Adj}^2 = 0.7175$ ; p < 0.01; Species richness:  $R_{Adj}^2 = 0.8362$ ; p < 0.01



**Fig. 4** Variance partitioning among the components that explain the variations in **a** DCA2 Axis 2 scores and **b** estimated species richness along an altitudinal gradient in the Atlantic Rainforest (SE Brazil). Axis 1 of the DCA2 did not have any significant spatial variables, and thus, we did not perform a variance partitioning in this case

restrictive environment (Assis et al. 2011) and, consequently, to species selectivity due to environmental filters (Ackerly 2003; Assis et al. 2011). Disturbances, in turn, can affect water conditions and temperature (Zhao et al. 2005), which are principal factors affecting the distribution of vegetation (Holdridge 1947; Gentry 1988). The logging of *Hyeronima alchorneoides* in area "F" appears to have been largely responsible for the local loss of certain species due to changes in the dynamics and succession (Villela et al. 2006), as the logged species was a large tree reaching a large volume of wood. Human disturbances have also appeared to influence species distributions in other studies (e.g., Vázquez and Givnish

1998), and it has been highlighted as a factor influencing biodiversity in the Atlantic Forest (Pereira et al. 2007; Tabarelli et al. 2010).

We also found evidence of the influence of gaps on floristic variation. The opening of gaps influences the germination and growth of forest species (Paz and Martínez-Ramos 2003). Gap area seems to be a key variable influencing species composition, since it determines the amount of light that reaches the area of the gap (Barton et al. 1989). Therefore, different species colonise gaps of different sizes (Whitmore 1996).

The dissociation between "I"+"J" and the other areas appears to be related to topographic variables, as described below. Regarding the floristic differences between higher ("K"+"N") and lower elevation areas, regional studies performed in the Brazilian Atlantic forest have indicated that altitude is important for detecting floristic affinities (e.g., Oliveira-Filho and Fontes 2000; Meireles et al. 2008; Sanchez et al. 2013). The relationship between vegetation patterns and altitude is well established (Rahbek 1995; Givnish 1999; Körner 2007), and several different patterns have been observed (He and Chen 1997). In the same conservation unit used for this study, Sanchez et al. (2013) addressed altitudinal limits for certain tree species, where most cases of restricted distribution (51 %) occurred at higher elevations. Thus, our results confirm that segregation patterns are driven by a set of factors related to altitude.

At each level of a slope, the vegetation is a result of a confluence of the constituent species, climate, soils and disturbances (e.g., Zhao et al. 2005). The confluence of these factors, which have been occurring for thousands of years, might result in elevation zonation, reflecting differences in species diversity levels for different regions of the slope (Zhao et al. 2005). However, two observations did not support the pattern of decreases in diversity with increases in altitude: (i) there was no significant correlation between species richness and altitude, and (ii) the species richness in higher altitude areas ("K" and "N", >500 m, in montane rainforest) decreased relative to "I" and "J". This result would suggest a "midaltitude bulge" pattern for this gradient (Lomolino 2001), thereby confirming the results provided by Zhao et al. (2005), among others. However, because we did not sample the vegetation along the complete altitudinal range due to logistic limitations (Joly et al. 2012), this potential pattern could not be confirmed. Further research on this question is needed.

It has been proposed that depending on the substrate, species diversity might display an increase or decrease along an elevation gradient (Wilson et al. 1990). Throughout the area investigated in this study, the soil carbon, nitrogen and total alkalinity levels were highest in "I" and "J" (the richest species areas), which can be explained by topographic factors influencing the microclimate (SC Martins, unpublished data). The slope and surface area could help us to understand the floristic segregation of areas "I" and "J", as we found a significant interference from variations in slope (SD Slope) on species richness. Accordingly, topography was important for some of the trends of variation in species distribution, in agreement with Guerra et al. (2013), for example, and topography mainly affected species richness is expected where local-scale variations in topography result in higher soil water availability. This result is expected because higher species richness is usually found in forests with low seasonal variation in climate (Gentry 1988; Pyke et al. 2001).

The number of multi-stemmed trees per hectare exhibited a significant negative correlation with species richness. Probably many adult trees were dying as a consequence of slope, thus opening new gaps in the forest. These gaps allowed the increased branching of the remaining trees, resulting in a greater number of multiple stems; thus, multiple stems would be associated with gap opening. This phenomenon is supported by the fact that the death of many adults could result in a decline in species richness (personal observations). For example, a peak in multiple stems and gaps.area<sup>-1</sup> was observed at one of the areas ("F") having the lowest species richness among all studied sites. Thus, we propose that despite the expected high species richness in areas with wider gaps (e.g., Terborgh 1992), in the investigated forests, a more complex pattern has occurred, perhaps due to the significant effects of logging in area "F".

The summit might experience more fog, with associated decline in species richness, and increase in floristic differentiation (e.g., Bertoncello et al. 2011). However, our field observations indicate that the presence of low-level cloud cover and fog formation is not frequent in the Montane Forest plots. Despite a clear decline in air and soil temperatures (Vieira et al. 2011), probably such climate condition is not a main driver on vegetation patterns (Alves et al. 2010). We believe that our Montane forest site is located in a transitional area exposed to varying degrees of cloud impaction, thus it might be considered as a lower montane forest under some influence of clouds. Additional studies on micrometeorology and forest hydrology are necessary to understand this topic.

The studied areas "I" and "J" had a unique level of richness, since it is unusual to observe an alpha diversity of more than 200 species for Atlantic Rainforest sites (Siqueira 1994; Tabarelli and Mantovani 1999; Scudeller et al. 2001). Our results surpass the species richness found in other studies developed in Neotropical forests (see Tabarelli and Mantovani 1999) and contrast with the argument given by these authors that Atlantic Rainforests has lower species richness when compared to other Neotropical forests.

The variance partitioning indicated a clear influence from the spatially structured environment and a secondary influence of the "pure" spatial component on variations in species composition and richness. The former comprises a well-known pattern in the Atlantic Forest (e.g., Oliveira-Filho and Fontes 2000; Scudeller et al. 2001) despite the lack of specific approaches to assessing this component of variation. The influence of the "pure" spatial component suggests limitations to biotic dispersal and indicates the importance of stochastic mechanisms (Hubbell 2001). Both components (the spatially structured environment and "pure" space) could be related to the well-known (Martins 1991; Scudeller et al. 2001; Caiafa and Martins 2010) geographically restricted distribution of tree species in the Atlantic Forest (see Diniz-Filho et al. 2012). Thus, spatial proximity is undoubtedly a crucial factor driving the vegetation patterns along the altitudinal range investigated in this study.

We found evidence for a floristic variation influenced by certain environmental variables, such as disturbances, topography and elevation related variables as well as spatial proximity, and a high regional richness (a total of 563 species in the investigated area) and extraordinarily high species richness (>200 spp.) in two montane areas. These results indicate that the hillside forest studied here is an important location for additional detailed studies to investigate the processes that generate and maintain its floristic composition and diversity and is worthy of increasing efforts to preserve its biological heritage.

**Acknowledgments** We thank the BIOTA/FAPESP Program for supporting the BIOTA Functional Gradient Project (Atlantic Ombrophyllous Dense Forest: Floristic Composition, Structure and Functioning within the "Serra do Mar" State Park, Brazil FAPESP Grants); the Graduate Programs in Plant Biology and Ecology at UNICAMP; CNPq for the PhD scholarship granted to the first author; and CNPq, CAPES and FAPESP for Grants awarded to the other authors and researchers directly or indirectly involved with this paper.

## References

Ackerly DD (2003) Community assembly, niche conservatism, and adaptive evolution in changing environments. Int J Plant Sci 164(3 Suppl):165–184

- Alves LF, Vieira SA, Scaranello MA, Camargo PB, Santos FAM, Joly CA, Martinelli LA (2010) Forest structure and live aboveground biomass variation along an elevational gradient of tropical Atlantic moist forest (Brazil). For Ecol Manag 260:679–691. doi:10.1016/j.foreco.2010.05.023
- Araújo DSD, Lacerda LD (1987) A natureza das restingas. Ciência Hoje 6:42-48
- Assis MA, Prata EMB, Pedroni F, Sanchez M, Eisenlohr PV, Martins FR, Santos FAM, Tamashiro JY, Alves LF, Vieira SA, Piccolo MC, Martins SC, Camargo PB, Carmo JB, Simões E, Joly CA (2011) Florestas de restinga e de terras baixas na planície costeira do sudeste do Brasil: vegetação e heterogeneidade ambiental. Biota Neotrop 11:103–121. doi:10.1590/S1676-06032011000200012
- Auerbach M, Shmida A (1993) Vegetation change along an altitudinal gradient on Mt Hermon, Israel—no evidence for discrete communities. J Ecol 81:25–33
- Austin MP, Greig-Smith P (1968) The application of quantitative methods to vegetation survey. II. Some methodological problems of data from rain forest. J Ecol 56:827–844
- Barton AM, Fetcher N, Redhead S (1989) The relationship between treefall gap size and light flux in a Neotropical rain forest in Costa Rica. J Trop Ecol 5:437–439. doi:10.1017/S0266467400003898
- Beals EW (1969) Vegetation change along altitudinal gradients. Science 165:981–985
- Bellier E, Monestiez P, Durbec JP, Candau JN (2007) Identifying spatial relationships at multiple scales: principal coordinates of neighbour matrices (PCNM) and geostatistical approaches. Ecography 30:385–399. doi:10.1111/j.0906-7590.2007.04911.x
- Bertoncello R, Yamamoto K, Meireles LD, Shepherd GJ (2011) A phytogeographic analysis of cloud forests and other forest subtypes amidst the Atlantic forests in south and southeast Brazil. Biodivers Conserv 20:3413–3433. doi:10.1007/s10531-011-0129-6
- Bohlman SA, Laurance WF, Laurance SG, Nascimento HEM, Fearnside PM, Andrade A (2008) Importance of soils, topography and geographic distance in structuring central Amazonian tree communities. J Veg Sci 19:863–874. doi:10.3170/2008-8-18463
- Bourgeron PS (1983) Spatial aspects of vegetation structure. In: Golley FB (ed) Ecosystems of the world: tropical rain forest ecosystems, structure and function. Elsevier, Amsterdam, pp 29–47
- Burnham KP, Anderson DR (2002) Model selection and multimodel inference: a practical informationtheoretic approach, 2nd edn. Springer, New York
- Caiafa AN, Martins FR (2010) Forms of rarity of tree species in the southern Brazilian Atlantic rainforest. Biodivers Conserv 19:2597–2618. doi:10.1007/s10531-010-9861-6
- César O, Monteiro R (1995) Florística e fitossociologia de uma floresta de restinga em Picinguaba (Parque Estadual da Serra do Mar), Município de Ubatuba-SP. Naturalia 20:89–105
- Connell JH (1978) Diversity in tropical rain forests and coral reefs. Science 199:1302–1309
- Diniz-Filho JAF, Bini LM, Hawkins BA (2003) Spatial autocorrelation and red herrings in geographical ecology. Glob Ecol Biogeogr 12:53–64. doi:10.1046/j.1466-822X.2003.00322.x
- Diniz-Filho JA, Rangel TFLVB, Bini LM (2008) Model selection and information theory in geographical ecology. Glob Ecol Biogeogr 17:479–488. doi:10.1111/j.1466-8238.2008.00395.x
- Diniz-Filho JAF, Siqueira T, Padial AA, Rangel TF, Landeiro VL, Bini LM (2012) Spatial autocorrelation analysis allows disentangling the balance between neutral and niche processes in metacommunities. Oikos 121:201–210. doi:10.1111/j.1600-0706.2011.19563.x
- Dray S, Legendre P, Peres-Neto PR (2006) Spatial modelling: a comprehensive framework for principal coordinate analysis of neighbour matrices (PCNM). Ecol Model 196:483–493. doi:10.1016/j. ecolmodel.2006.02.015
- Eisenlohr PV (2013) Challenges in data analysis: pitfalls and suggestions for a statistical routine in vegetation ecology. Braz J Bot. doi:10.1007/s40415-013-0002-9
- Fortin MJ, Dale MRT (2005) Spatial analysis: a guide for ecologists. Cambridge University Press, Cambridge
- Gentry AH (1988) Changes in plant community diversity and floristic composition on environmental and geographical gradients. Ann Miss Bot Gard 75:1–34
- Givnish TJ (1999) On the causes of gradients in tropical tree diversity. J Ecol 87:193–210. doi:10.1046/j. 1365-2745.1999.00333.x
- Gotelli NJ, Colwell RK (2001) Quantifying biodiversity: procedures and pitfalls in the measurement and comparison of species richness. Ecol Lett 4:379–391. doi:10.1046/j.1461-0248.2001.00230.x
- Gotelli NJ, Entsminger GL (2004) EcoSim: null models software for ecology. Version 7.71. http://together. net/~gentsmin/ecosim.htm. Accessed 12 Nov 2012
- Guerra TNF, Rodal MJN, Lins e Silva ACB, Alves M, Silva MAM, Mendes PGA (2013) Influence of edge and topography on the vegetation in an Atlantic Forest remnant in northeastern Brazil. J For Res 18:200–208. doi:10.1007/s10310-012-0344-3
- He J-S, Chen W-L (1997) A review of gradient changes in species diversity of land plant communities (In Chinese with an English summary). Act Ecol Sin 17:91–99

- Hijmans RJ, Cameron SE, Parra JL, Jones PG, Jarvis A (2005) Very high resolution interpolated climate surfaces for global land areas. Int J Climat 25:1965–1978. doi:10.1002/joc.1276
- Holdridge LR (1947) Determination of world plant formations from simple climatic data. Science 105:367-368
- Hubbell SP (2001) The unified neutral theory of biodiversity and biogeography. Princeton University Press, Princeton
- Hutchinson MF (1989) A new procedure for gridding elevation and stream line data with automatic removal of spurious pits. J Hydrol 106:211–232. doi:10.1016/0022-1694(89)90073-5
- IBGE (2012) Manual Técnico da Vegetação Brasileira. IBGE, Rio de Janeiro
- Ivanauskas NM, Monteiro R, Rodrigues RR (2000) Similaridade florística entre áreas de Floresta Atlântica no estado de São Paulo. Braz J Ecol 1(2):71–81
- Ivanauskas NM, Rodrigues RR, Souza VC (2006) Restoration Methodology: the importance of the regional floristic diversity for the forest restoration successfulness. In: Rodrigues RR, Martins SV, Gandolfi S (eds) High diversity forest restoration in degraded areas. Nova Science Publishers, New York, pp 63–76
- Joly CA, Assis MA, Bernacci LC, Tamashiro JY, Campos MCR, Gomes JAMA, Lacerda MS, Santos FAM, Pedroni F, Pereira LS, Padgurschi MC, Prata EMB, Ramos E, Torres RB, Rochelle A, Martins FR, Alves LF, Vieira AS, Martinelli LA, Camargo PB, Aidar MPM, Eisenlohr PV, Simões E, Villani JP, Belinello R (2012) Florística e fitossociologia em parcelas permanentes da Mata Atlântica do sudeste do Brasil ao longo de um gradiente altitudinal. Biota Neotrop 12:125–145. doi:10.1590/S1676-06032012000100012
- Körner C (2007) The use of 'altitude' in ecological research. Trop Ecol Evol 22:569–574. doi:10.1016/j.tree. 2007.09.006
- Laurance WF (2001) Fragmentation and plant communities: synthesis and implications for landscape management. In: Bierregaard RO Jr, Gascon C, Lovejoy TE, Mesquita RCG (eds) Lessons from Amazonia: the ecology and conservation of a fragmented forest. Yale University Press, New Haven, pp 158–168
- Laurance WF (2009) Conserving the hottest of the hotspots. Biol Conserv 142:1137. doi:10.1016/j.biocon. 2008.10.011
- Legendre P, Legendre L (2012) Numerical ecology, 3rd edn. Elsevier Science BV, Amsterdam
- Lieberman D, Lieberman M, Peralta R, Hartshorn GS (1996) Tropical forest structure and composition on a large-scale altitudinal gradient in Costa Rica. J Ecol 84:137–152
- Lomolino MV (2001) Elevation gradients of species density: historical and prospective views. Glob Ecol Biogeogr 10:3–13. doi:10.1046/j.1466-822x.2001.00229.x
- Marques MCM, Swaine MD, Liebsch D (2011) Diversity distribution and floristic differentiation of the coastal lowland vegetation: implications for the conservation of the Brazilian Atlantic Forest. Biodivers Conserv 20:153–168. doi:10.1007/s10531-010-9952-4
- Martins FR (1991) Estrutura de uma floresta mesófila. Editora da UNICAMP, Campinas
- McCune B, Grace JB (2002) Analysis of Ecological Communities. MjM Software Design, Gleneden Beach McCune B, Mefford MJ (2011) PC-ORD—multivariate analysis of ecological data, version 6.0. MjM Software Design, Gleneden Beach
- Meireles LD, Shepherd GJ, Kinoshita LS (2008) Variações na composição florística e na estrutura fitossociológica de uma floresta ombrófila densa alto-montana na Serra da Mantiqueira, Monte Verde, MG. Rev Bras Bot 31:559–574. doi:10.1590/S0100-84042008000400003
- Morellato LPC, Talora DC, Takahashi A, Bencke CC, Romera EC, Ziparro VP (2000) Phenology of Atlantic Rain Forest trees: a comparative study. Biotropica 32:811–823. doi:10.1111/j.1744-7429.2000. tb00620.x
- Moreno MR, Nascimento MT, Kurtz BC (2003) Estrutura e composição florística do estrato arbóreo em duas zonas altitudinais na Mata Atlântica de encosta da região de Imbé, RJ. Acta Bot Bras 17:371–386. doi:10.1590/S0102-33062003000300005
- Myers N, Mittermeier RA, Mittermeier CG, Fonseca GAB, Kent J (2000) Biodiversity hotspots for conservation priorities. Nature 403:853–858. doi:10.1038/35002501
- O'Brien EM, Field R, Whittaker RJ (2000) Climatic gradients in woody plant (tree and shrub) diversity: water–energy dynamics, residual variation, and topography. Oikos 89:588–600. doi:10.1034/j.1600-0706.2000.890319.x
- Oliveira-Filho AT, Fontes MAL (2000) Patterns of floristic differentiation among Atlantic forests in southeastern Brazil, and the influence of climate. Biotropica 32:793–810. doi:10.1111/j.1744-7429. 2000.tb00619.x
- Paz H, Martínez-Ramos M (2003) Seed mass and seedling performance within eight species of *Psychotria* (Rubiaceae). Ecology 84:439–450. doi:10.1890/0012-9658(2003)

- Pereira JAA, Oliveira-Filho AT, Lemos-Filho JP (2007) Environmental heterogeneity and disturbance by humans control much of the tree species diversity of Atlantic montane forest fragments in SE Brazil. Biodivers Conserv 16:1761–1784. doi:10.1007/978-1-4020-6444-9\_13
- Ponçano WL, Carneiro CDR, Bistrichi CA, Almeida FFM, Prandini FL (1981) Mapa Geomorfológico do estado de São Paulo, vol 1. Instituto de Pesquisas Tecnológicas do Estado de São Paulo, São Paulo
- Pyke CR, Condit R, Aguilar S, Lao S (2001) Floristic composition across a climatic gradient in a neotropical lowland forest. J Veg Sci 12:553–566. doi:10.2307/3237007
- Quinn GP, Keough MJ (2002) Experimental design and data analysis for biologists. Cambridge University Press, Cambridge
- Rahbek C (1995) The elevational gradient of species richness: a uniform pattern? Ecography 18:200–205. doi:10.1111/j.1600-0587.1995.tb00341.x
- Rangel TFLVB, Diniz-Filho JAF, Bini LM (2010) SAM: a comprehensive application for spatial analysis in macroecology. Ecography 33:46–50. doi:10.1111/j.1600-0587.2009.06299.x
- Rey Benayas JM, Scheiner SM (2002) Plant diversity, biogeography and environment in Iberia: patterns and possible causal factors. J Veg Sci 13:245–258. doi:10.1111/j.1654-1103.2002.tb02045.x
- Ribeiro MC, Martensen AC, Metzer JP, Tabarelli M, Scarano F, Fortin MJ (2011) The Brazilian Atlantic Forest: a shrinking biodiversity hotspot. In: Zachos FE, Habel JC (eds) Biodiversity hotspots: distribution and protection of conservation priority areas. Springer, Heidelberg, pp 405–434
- Rodrigues RR, Shepherd GJ. 1992. Análise de variação estrutural e fisionômica da vegetação e características edáficas, num gradiente altitudinal na Serra do Japi. In: Morellato LPC (org). História natural da Serra do Japi. Editora da UNICAMP, Campinas, pp 64–93
- Runkle JR (1982) Patterns of disturbance in some old-growth mesic forests of North America. Ecology 63:1533–1556. doi:10.2307/1938878
- Sanchez M, Pedroni F, Eisenlohr PV, Oliveira-Filho AT (2013) Changes in tree community composition and structure of Atlantic rain forest on a slope of the Serra do Mar range, Southeastern Brazil, from near sea level to 1000 m of altitude. Flora 208:184–196. doi:10.1016/j.flora.2013.03.002
- Santos RM, Oliveira-Filho AT, Eisenlohr PV, Queiroz LP, Cardoso DBOS, Rodal MJN (2012) Identity and relationships of the Arboreal Caatinga among other floristic units of seasonally dry tropical forests (SDTFs) of north-eastern and Central Brazil. Ecol Evol 2:409–428. doi:10.1002/ece3.91
- Scudeller VV, Martins FR, Shepherd GJ (2001) Distribution and abundance of arboreal species in the Atlantic ombrophilous dense forest in southeastern Brazil. Plant Ecol 80:705–717. doi:10.1023/A: 1011494228661
- Simberloff D (1983) Competition theory, hypothesis-testing, and other community ecology buzzwords. Am Nat 122:626–635
- Siqueira MF (1994) Análise florística e ordenação de espécies arbóreas da Mata Atlântica através de dados binários. Dissertation, Universidade Estadual de Campinas
- Sklenář P, Bendix J, Balslev H (2008) Cloud frequency correlates to plant species composition in the high Andes of Ecuador. Basic Appl Ecol 9:504–513. doi:10.1016/j.baae.2007.09.007
- Sousa Neto E, Carmo JB, Keller M, Martins SC, Alves LF, Vieira SA, Piccolo MC, Camargo P, Couto HTZ, Joly CA, Martinelli LA (2011) Soil-atmosphere exchange of nitrous oxide, methane and carbon dioxide in a gradient of elevation in the coastal Brazilian Atlantic forest. Biogeosciences 8:733–742. doi:10.5194/bg-8-733-2011
- Tabarelli M, Mantovani W (1999) A riqueza de espécies arbóreas na floresta atlântica de encosta no estado de São Paulo (Brasil). Rev Bras Bot 22:217–223. doi:10.1590/S0100-84041999000200012
- Tabarelli M, Aguiar AV, Ribeiro MC, Metzger JP, Peres CA (2010) Prospects for biodiversity conservation in the Atlantic Forest: lessons from aging human-modified landscapes. Biol Conserv 143:2328–2340. doi:10.1016/j.biocon.2010.02.005
- Terborgh J (1992) Diversity and the tropical rain forests. Scientific American Library, New York
- The R Foundation for Statistical Computing (2012) R: a language and environment for statistical computing. Vienna (Austria). http://www.R-project.org/. Accessed 15 Dec 2012
- Vázquez JAG, Givnish TJ (1998) Altitudinal gradients in tropical forest composition, structure, and diversity in the Sierra de Manantlán, Jalisco, México. J Ecol 86:999–1020. doi:10.1046/j.1365-2745. 1998.00325.x
- Vieira SA, Alves LF, Duarte-Neto PJ, Martins SC, Veiga LG, Scaranello MA, Picollo MC, Camargo PB, Carmo JB, Neto ES, Santos FAM, Joly CA, Martinelli LA (2011) Stocks of carbon and nitrogen and partitioning between above- and below-ground pools in the Brazilian coastal Atlantic Forest elevation range. Ecol Evol 1:421–434. doi:10.1002/ece3.41
- Villela DM, Nascimento MT, De Aragão LEOC, Da Gama DM (2006) Effect of selective logging on forest structure and nutrient cycling in a seasonally dry Brazilian Atlantic forest. J Biogeogr 33:506–516. doi:10.1111/j.1365-2699.2005.01453.x

- Whitmore TC (1996) A review of some aspects of tropical rain forest seedling ecology with suggestion for further enquiry. In: Swaine MD (ed) The ecology of tropical forest tree seedlings. UNESCO, Paris, pp 3–39
- Wilson JB, Lee WG, Mark AF (1990) Species diversity in relation to ultramafic substrate and to altitude in southwestern New Zealand. Vegetatio 86:15–20. doi:10.1007/BF00045131
- Wolf JA, Fricker GA, Meyer V, Hubbell SP, Gillespie TW, Saatchi SS (2012) Plant species richness is associated with canopy height and topography in a neotropical. Remote Sens 4:4010–4021. doi:10. 3390/rs4124010
- Zhao C-M, Chen W-L, Tian Z-Q, Xie Z-Q (2005) Altitudinal pattern of plant species diversity in Shennongjia mountains, Central China. J Integr Plant Biol 47:1431–1449. doi:10.1111/j.1744-7909.2005. 00164.x