Do fragment size and edge effects predict carbon stocks in trees and

2

lianas in tropical forests?

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25 Summary

1. Tropical forests are critical for protecting global biodiversity and carbon stores. While forest degradation and fragmentation cause negative impacts on trees, many woody lianas benefit, with associated negative effects on carbon storage. Here we focus on the key question of how abiotic environmental changes resulting from tropical forest fragmentation mediate the allocation of carbon into trees and lianas.

2. We focus on the globally threatened Brazilian Atlantic Forest, in forest fragments spanning 13
to 23 442 ha in area and at fragment edges and interiors. Within each fragment, we established
two transects: one at the edge and one in the interior. Each transect consisted of ten 10 × 10 m
plots spaced at 20 m intervals. Within each plot we sampled living trees with diameter ≥4.8 cm
at 1.3 m above ground, living lianas with diameter ≥1.6 cm at 10 cm above ground, and several
microclimatic and soil variables.

37 3. Fragmentation changed a broad suite of abiotic environmental conditions recognized as being
associated with forest carbon stocks: edges and smaller fragments were hotter, windier, and less
humid, with more fertile and less acid soils at edges. Tree carbon stocks were thus higher in
forest interiors than at edges, and were positively related to fragment size in interiors, but were
not impacted by fragment size at edges.

42 4. Trees and lianas showed different responses to fragmentation: in interiors of small fragments,
43 tree carbon stocks declined whereas liana carbon stocks increased; and at edges, tree carbon
44 stocks were not affected by fragment size, whereas liana carbon stocks were highest in smaller
45 fragments. These patterns were strongly related to changes in abiotic environmental conditions.
46 5. We conclude that the abiotic changes across the fragmentation gradient, rather than liana
47 proliferation, were more likely to reduce tree carbon stocks. Cutting of lianas is frequently

- 48 promoted for restoring forest carbon in human-modified tropical forests. However, this approach
- 49 may not be effective for restoring forest carbon stocks in fragmented forests.
- 50
- 51 **Keywords**: Carbon cycle; Biomass; Microclimate; Air Temperature; Desiccation; Soil fertility;
- 52 REDD+.
- 53

54 Introduction

The loss, fragmentation and disturbance of tropical forests represent a major threat to 55 56 biodiversity (Laurance et al. 2006; Gibbs et al. 2010; Solar et al. 2015) and globally important ecosystem services, including carbon storage and climate regulation (Magnago et al. 2014, 57 58 2015a; Lewis *et al.* 2015). For example, tropical forests contribute to over a third of the net 59 primary productivity in global terrestrial ecosystems (Field 1998), and the largest tropical forest countries harbour 45% of the terrestrial global carbon stocks (Bonan 2008). Tropical forest 60 deforestation and disturbance contributes to 6-17% of the global anthropogenic CO₂ emissions, 61 62 which is second only to the burning of fossil fuels (van der Werf *et al.* 2009). In countries such 63 as Brazil, CO₂ emissions from forest losses and disturbance account for 22% of total emissions 64 (Tollefson 2013). Understanding how land-use change affects the carbon stocks in tropical forest 65 fragments is critical, especially given the potential to implement carbon-based payments for 66 ecosystem services and biodiversity conservation (e.g. the United Nations Reducing Emissions 67 from Deforestation and Forest Degradation [REDD+] program) (Berenguer et al. 2014; Magnago et al. 2015a). 68

69 Outside of tropical peatlands, the vast majority of carbon held within tropical forests is 70 stored in mature trees (Chave *et al.* 2005; Berenguer *et al.* 2014), yet this is also the stock that is 71 most vulnerable to anthropogenic disturbance (Laurance et al. 2000; Berenguer et al. 2014). In 72 fragmented landscapes, alterations to the microclimate at forest edges include increased wind 73 speeds, higher temperatures via canopy openness, and lower humidity (Laurance *et al.* 2002; Magnago et al. 2015b). These impacts alter tree functional attributes (hardwood to softwood 74 species) and drive high rates of tree mortality (Laurance et al. 2000; Briant, Gond & Laurance 75 2010; Magnago et al. 2014, 2015a; b), reducing tree carbon stocks (Laurance et al. 2006; 76 Berenguer et al. 2014). 77

78	While such environmental changes reduce carbon stocks in trees, there is a growing body
79	of evidence suggesting that many woody lianas benefit from forest disturbance and
80	fragmentation (Laurance et al. 2001; Schnitzer & Bongers 2002). Several complementary
81	mechanisms can explain the competitive advantage of lianas over trees in certain environmental
82	conditions, including a greater capability to access deep-water facilitating resistance to drought
83	events, the ability to cause mechanical damage to trees which opens up gaps, and rapid
84	colonization and proliferation with high light intensity or soils with increased fertility due to
85	nutrient deposition (Laurance et al. 2002, 2014; Schnitzer & Bongers 2002; Restom & Nepstad
86	2004; Magrach et al. 2014). Lianas may also benefit from the increase in global atmospheric
87	CO ₂ , which can cause a greater increase in rates of fecundity, recruitment and growth for lianas
88	than tree species (Schnitzer & Bongers 2011; Laurance et al. 2014).
89	Liana proliferation would negatively affect the carbon storage potential of forests
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100 fragmentation mediate the allocation of carbon into trees and lianas. Previous studies in tropical

101	forests have shown how edge effects usually change the abiotic environment of fragments
102	(Laurance et al. 2002; Magnago et al. 2015b) and that these changes can affect forest structure,
103	reducing tree carbon stocks and increasing the dominance of lianas (Laurance et al. 2014;
104	Magrach et al. 2014; Magnago et al. 2015ab). We therefore measured tree carbon stocks, liana
105	abundances and carbon stocks, soil nutrients and acidity, and microclimatic conditions to answer
106	three main questions: (i) are microclimatic and soil conditions controlled by the fragmentation
107	gradient (fragment size and edge-interior location)?; (ii) do environmental changes following
108	fragmentation reduce trees carbon stock and increase lianas abundances and carbon stocks?; and
109	(iii) does liana prevalence affect tree carbon stocks?
110	
111	Material and Methods
112	Study area
113	This study was based in the state of Espírito Santo, in South-East Brazil. Within the
114	region, we focused on the municipalities of Sooretama, Linhares and Jaguaré (18°54'-19°15'S
115	and 39°54–40°15'W, 28–65 m.a.s.l) (Fig. S1; Magnago et al. 2014), which contain a landscape
116	matrix composed mainly of pastures, Eucalyptus spp., coffee and papaya plantations (Rolim et
117	al. 2005), plus many forest fragments, including two larger than 20,000 ha (see Text S1 for more
118	details). The climate is tropical wet (Köppen classification), with mean annual precipitation of
119	1403 mm and a dry season from May to September when precipitation is 33 mm per month
120	(Peixoto & Gentry 1990).
101	

122 Tree and liana sampling

Fieldwork was conducted from January 2011 to January 2012. We created permanent plots along 123 transects within nine fragments ranging from 13 to 1318 ha in area (mean = 334 ha) and within 124 two control fragments larger than 20,000 ha in Reserva Natural Vale (RNV) and Reserva 125 126 Biológica de Sooretama (REBIO) (Table S1; Magnago et al. 2014). We established one edge and 127 one interior transect within each of our nine fragments, with transects consisting of ten 10×10 m plots spaced at 20 m intervals. We also established six transects within control fragments: one 128 edge and one interior transect in RNV, and two edge and two interior transects in REBIO, with a 129 130 mean distance of 17.1 (\pm 10.4) km between transects. Based on mapping of IBGE (1987), all transects were on Yellow Ultisol soils. The spatially auto-correlated plots were aggregated into a 131 132 single 0.1 ha transect for all of our analyses. Although larger sample areas reduce error in 133 diversity and forest structure metrics (Baraloto et al. 2012), 0.1 ha is frequently used for assessing tropical forest carbon stocks (Saatchi et al. 2011; Stegen et al. 2011; Magnago et al. 134 2015a). 135

A major review of 33 abiotic and biotic edge parameters showed that 10 parameters extended less than 25 m from forest edge, while all but one factor (wind speed) extended less than 300 m (Laurance *et al.* 2002). Hence, we established edge transects ~5 m inside the fragment and parallel to the forest edge to capture all edge parameters, and interior transects with distance \geq 300 m from the nearest edge (although the smallest fragments did not permit a longer minimum distance from edge).

Within each plot, we sampled every living tree with a diameter at breast height (DBH)
≥4.8 cm at 1.3 m height. We identified the botanical material using collections at the CVRD
Herbarium of the Reserva Natural Vale and VIES Herbarium of the Federal University of

145	Espírito Santo, and with assistance from experts in the identification of specific plant families
146	(e.g. Myrtaceae and Sapotaceae). Botanical material collected in the fertile stage was deposited
147	at the CVRD Herbarium. Within the plots, we also measured every liana ≥ 1.6 cm diameter at
148	10 cm above ground (Diameter at Soil Height [DSH]). We only sampled individuals with \geq 50%
149	of their roots inside the plot (see Phillips et al. 2009).
150	
151	Trees and lianas carbon estimation
152	Following previous studies in the region (Magnago et al. 2015a), we used Chave et al.'s
153	(2005) equation to estimate the Above Ground Biomass of Trees (AGBt):
154	
155	ABGt = $\rho . \exp(-1.499 + 2.148 \ln(\text{DBH}) + 0.207(\ln(\text{DBH}))^2 - 0.0281(\ln(\text{DBH}))^3)$
156	
157	where $\rho = \text{wood density (g/cm^3)}$.
158	
159	For estimating live Liana Above Ground Biomass (AGBl) we used Schnitzer et al.'s (2006)
160	equation:
161	$AGBI = exp(-1.484 + 2.657 \ln(DSH))$
162	
163	We assume that 50% of AGB of each tree and liana individual is represented by carbon
164	(Malhi et al. 2004; Chave et al. 2005). Tree species data for wood density on dry weight (g/cm ³)
165	were obtained from The Global Wood Density (GWD) database, subsection Tropical South
166	America (http://hdl.handle.net/10255/dryad.235; (Chave et al. 2009; Zanne et al. 2009). We

167 made two following adjustments (following Flores & Coomes 2011; Hawes *et al.* 2012): (i) for

tree morphospecies only identified to the family or genus level, we used the average wood density of the minor taxonomic group; (ii) for species not in the GWD database, we used the average wood density for the species' genus.

- 171
- 172 Sampling microclimatic and soil variables

In each plot, we sampled the microclimatic variables of maximum air temperature (°C), 173 maximum wind speed (km/h) and relative humidity (%) using two Kestrel Weather and 174 Environmental Meters (model 4500, Nielsen-Kellerman Company, USA). These measures are 175 176 known to respond to fragmentation and edge effects (Chen et al. 1999; Didham & Lawton 1999; Laurance et al. 2002). The data for the two control fragments were obtained from Magnago et al. 177 (2015b). To standardize microclimatic data collection among the sample plots, all measures were 178 179 recorded once per plot during a 15-minute period (between 08:00-11:00 and then 14:00-17:00) and at 1.5 m above ground. Because microclimatic parameters show natural variation (e.g., 180 warmer days have higher humidity and stronger winds than other days), we also recorded 181 microclimatic data approximately 10 m from the edge in the matrix adjacent to each fragment. 182 Each fragment's microclimatic data is then expressed as the percentage change from the matrix 183 184 values (Table S1).

For soil variables we collected three replicate samples of the topsoil (0-10 cm depth) in each 10×10 m plot. These were mixed to form one composite sample per plot, which were then analyzed in the Soil Analysis Laboratory, Federal University of Viçosa (UFV). We selected the following variables as measures of soil fertility (Hazelton & Murphy 2007; Ronquim 2010): Sum of Bases (calculated by the sum of Ca+Mg+K+Na), Effective Cation Exchange Capacity, organic matter content, available phosphorus and pH in H₂O (Table S1).

191 Data Analysis

192 *General modelling framework*

All statistical analyses and model selections were constructed by applying Generalized 193 Linear Mixed Models (GLMM) using each fragment as a random factor (Bolker et al. 2009). We 194 195 used a Negative Binomial error distribution with log link function for response variable of count 196 data (abundance), since these data showed overdispersion, preventing the use of Poisson error. A Gaussian error distribution was used for continuous response variables. However, when response 197 variable was a ratio, we applied a log-normal distribution. These analyses were performed using 198 the "glmer.nb" function for Negative Binomial models and "lmer" function for Gaussian and 199 log-normal models in the "lme4" package. All models were validated using the relation between 200 standardized residuals with standardized normal quantiles, using the function "ggmath" in 201 "lme4". 202

To determine the best model, we used an information theoretical approach based on the 203 Akaike Information Criterion of Second Order (AICc), which is indicated for small sample sizes, 204 and the best model was indicated by the AICc lower value (Burnham & Huyvaert 2011). We 205 used the "dredge" function from the "MUMIN" package to test all possible combinations of the 206 207 variables included in the full model, including the null model. The approach of using AICc combined with a low number of parameters (two fixed explanatory variables and the interaction 208 between them) in the GLMM full model avoided overfitted models and reduced the chances of 209 210 type I error (Burnham & Anderson 2002 and Zuur et al. 2009). For all best models, we used residuals maps to assess whether spatial autocorrelation was determining model outcomes 211 212 (Dormann et al. 2007; Zuur et al. 2009). All analyses were performed in the R environment, 213 version 2.15.3 (R Development Core Team. 2012).

Are microclimatic and soil conditions controlled by the fragmentation gradient? 215 For each microclimatic and soil response variable (all continuous) we evaluated the 216 217 GLMM framework considering two explanatory variables: (i) fragment size (continuous) and (ii) 218 edge-interior location (categorical), plus the interaction between them. We then constructed a 219 PCA with all soil and microclimatic variables together. The final PCA1 included only the variables with significant Spearman's correlation values with the PCA1 axes and included Sum 220 of Bases, Effective Cation Exchange Capacity, maximum air temperature, maximum wind speed 221 222 and relative humidity (Figure S2A). Therefore, the final PCA1 did not select organic matter, phosphorus and pH. The significance of each axis of the PCA1 was obtained by 999 Monte 223 Carlo's permutation tests. As only axis 1 was significant (p=0.001) (see Figure S2A), we used it 224 225 as our composite metric of abiotic changes (Abiotic Change Axis 1, hereafter). We analysed the response of Abiotic Change Axis 1 using the GLMM framework to evaluate the effect of 226 fragment size, edge-interior location, and the interaction between fragment size and edge-interior 227 location. To develop a more complete gradient of environmental changes, we repeated the PCA1 228 procedures also including fragment size (Figure S2B). The Spearman's correlation values of all 229 230 variables used in PCA2 are shown in Figure S3. Again, only axis 1 was significant (p=0.001) (Figure S2B), which we then used as our composite metric of all microclimatic and soil changes 231 and fragment size reductions (Fragmentation Axis 1, hereafter; Figure S3). 232

233

234 Does fragmentation alter trees and lianas carbon stocks?

We used GLMMs to examine the impact of fragmentation on trees and lianas carbonstocks and also on liana abundance. First we ran each GLMM using two explanatory variables:

(i) fragment size and (ii) edge-interior location, plus the interaction between them. We thenrepeated this GLMM replacing fragment size with Fragmentation Axis 1.

We evaluated the effect of fragmentation on the relative dominance of lianas (see Phillips 239 et al. 2002) using a GLMM to examine the ratio between lianas and trees (for both abundance 240 and carbon stock metrics). Again, our first GLMM used two explanatory variables: (i) fragment 241 242 size and (ii) edge-interior location, plus the interaction between them. We then repeated this replacing fragment size with Fragmentation Axis 1. 243 244 245 *Does liana prevalence affect trees carbon stock?* We used GLMMs to examine whether there was a negative association between tree 246 carbon stocks (response variable) and liana abundance or liana carbon stock (explanatory 247 248 variables; (Durán & Gianoli 2013). As there was a strong Spearman's correlation between liana abundance and their carbon stock (r=0.8; p<0.0001), we did not include these two variables 249 within the same full model and instead ran full models separately and then compared the best 250

251 models for each variable (see Table S7).

252

253 Results

254 Are microclimatic and soil conditions controlled by the fragmentation gradient?

Overall, our composite Abiotic Axis 1 was strongly predicted by fragment size (GLMM; t=-5.350, p<0.0001, Figure S4A) and edge-interior location (GLMM; t=6.546, p<0.0001, Figure S4B). This was reflected by the changes in the specific microclimatic variables, although there was some variation in their individual responses. Air temperature was significantly higher in small fragments (GLMM; t=-3.06, p=0.01; Figure 1A) and near forest edges (GLMM; t=3.56, p<0.01; Figure 1B). In contrast, air humidity showed a significant interaction between fragment size and edge-interior location (GLMM; t=-3.05, p=0.01; Figure 1C), with a higher accrual of air humidity with increasing fragment size in the interior than at the edge. Wind speed showed a significant interaction with fragment size and edge-interior location (GLMM; t=2.28, p<0.05; Figure 1E), with a significant negative influence of fragments size for plots located in interiors (F=8.79, p=0.01; Figure 1E), but no effect for those located near edges (F=0.2, p=0.66; Figure 1E).

For soil, the best model showed that edges had significantly higher Effective Cation 267 268 Exchange Capacity and were thus more fertile when compared with the interior (GLMM; t=2.971; p<0.01; Figure 2A). Also, there was a slightly non-significant negative effect of 269 fragment size on Effective Cation Exchange Capacity (GLMM; t=-1.983, p=0.06; Figure 270 271 2B). Soils from transects located in forest edges showed higher values of Sum of Bases (GLMM; t=3.64, p<0.01; Figure 3C) and lower acidity (GLMM; t=2.55, p<0.05; Figure 2D) than interiors. 272 273 The best models also suggested a slight (but non-significant) influence of fragment edges on 274 available phosphorous (GLMM; t=2.033, p=0.06; Figure 2E) and organic matter contents 275 (GLMM; t=-2.02, p=0.07; Figure 2F).

276

277 Does fragmentation alter tree's and liana's carbon stocks?

We sampled 4,140 trees and 8,236 lianas. As expected, across all transects, much more carbon was stored by trees than lianas (Table S1). On average, trees stored 97.7% (\pm 1.8%) of the total carbon (i.e., trees + lianas carbon stocks) in interiors and 87.1% (\pm 7.9%) at edges. Total carbon in interior transects (61.8 \pm 25.8 Mg) was nearly three-fold that at edges (22.8 \pm 11.6 Mg). 282 Considering only tree's carbon stock, there was a 69% reduction from 60.7 Mg in interiors to283 18.7 Mg at edges.

Trees and lianas responded very differently to fragmentation. Following the fragment size gradient (interior plus edge), tree carbon stocks ranged from 115.7 Mg/transect in the largest (> 23,000ha) to 46.4 Mg/transect in the smallest (13 ha) fragment, representing a 60% reduction. In contrast, liana carbon stocks increased by 200%, from 1.6 Mg/transect in the largest to 4.79 Mg/transect in the smallest fragment.

Tree carbon stocks - There was a significant interaction effect of fragment size and edge-289 interior location (GLMM; t= -4.435; p=0.001; Figure 3A): decreasing fragment size had a 290 significant negative effect on trees carbon stock in interiors (F = 17.4; P = 0.002; Figure 3A), but 291 no significant effect at edges (F = 1.3; P = 0.3; Figure 3A). The best model also showed 292 significant reductions of tree carbon stocks at edges versus interiors (GLMM; t= -11.6; 293 294 p<0.0001; Figure 3B). Very similar results were obtained when we replaced fragment size with Fragmentation Axis 1. There was a significant interaction effect between Fragmentation Axis 1 295 and edge-interior location (GLMM; t= -4.9; p=0.0003; Figure 3C), and increasing environmental 296 297 impacts had a significant negative effect on tree carbon stocks in interiors (F = 13.82; P = 0.004; Figure 3C), but no significant effect at edges (F = 0.096; P = 0.7; Figure 3C). The best model 298 299 also showed significant reductions of tree carbon stocks at edges versus interiors (GLMM; t= -6.045; p<0.0001; Figure 3D). 300

Liana carbon stocks - There was a significant interaction effect of fragment size and edge-interior location on liana carbon stocks (GLMM; t= -3.14, p=0.01; Figure 3E). However, in contrast with trees, decreasing fragment size had a positive effect on liana carbon stocks in forest interiors (F = 6.47: P = 0.03; Figure 3E), but had a more strongly positive effect at edges 305 (F = 32.1; P = 0.0002; Figure 3E). The best models also revealed significantly higher liana carbon stocks at fragment edges than interiors (GLMM; t= 5.38; p= 0.0002; Figure 3F). Again, 306 similar results were obtained when we replaced fragment size with Fragmentation Axis 1. There 307 was a significant interaction effect of Fragmentation Axis 1 and edge-interior location on liana 308 carbon stocks (GLMM; t= -2.664, p=0.014; Figure 3G), as increasing environmental impacts had 309 a slightly non-significant positive effect in forest interiors (F = 4.1: P = 0.07; Figure 3G), but a 310 strongly positive effect at edges (F = 16.8: P = 0.002; Figure 3G). Also, the best models showed 311 no significant change in liana carbon stocks between edges and interior habitats (GLMM; t= 1.5; 312 313 p=0.15; Figure 3H).

Liana abundance - Liana abundance increased in smaller fragments (GLMM; t= -4.221, p<0.0001; Figure S5A) and at fragment edges (GLMM; t= 18.259, p<0.0001; Figure S5B), but there was no significant interaction between fragment size and edge-interior location. The models using Fragmentation Axis 1 showed similar results, with the best model revealing

significant increases in lianas with harsher environmental conditions (GLMM; t= -3.91,

p<0.0001; Figure S6A) and at fragment edges (GLMM; t= 6.58, p<0.0001; Figure S6B).

Relative prevalence of trees and lianas - For carbon stocks, the best GLMM models of the ratio of lianas to trees showed higher dominance of lianas in smaller fragments (GLMM; t= -

322 6.353, p<0.0001; Figure 3I) and at fragment edges (GLMM; t= 9.982, p<0.0001; Figure 3J).

323 Similar relationships were found when replacing fragment size with Fragmentation Axis 1, with

a higher dominance of lianas in harsher conditions (GLMM; t = -4.232, p < 0.001; Figure 3L) and

at fragment edges (GLMM; t=4.186, p<0.001; Figure 3M).

For abundance, there was a higher dominance of lianas relative to trees in smaller fragments (GLMM; t= -5.189, p<0.001; Figure 3N) and at fragment edges (GLMM; t= 18.091, p<0.0001; Figure 3O). Again, similar relationships emerged when replacing fragment size with
Fragmentation Axis 1, with relatively higher abundance of lianas in harsher environmental
conditions (GLMM; t= -4.739, p<0.001; Figure 3P) and at fragment edges (GLMM; t= 6.259,
p<0.0001; Figure 3Q).

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333 Does liana prevalence affect tree carbon stocks?

Although both measures of liana prevalence were negatively associated with tree carbon 334 stocks (Table S7), liana abundance was a much stronger predictor than liana carbon stocks 335 336 (model comparisons showed Delta AICc of 7). The single best model revealed a significant effect of liana abundance due to the interaction between edge-interior location on tree carbon 337 stocks (GLMM: t=-4.7, p<0.0001; Figure 4): only forest interiors had a strong association 338 339 between higher liana abundance and lower tree carbon stocks (F = 6.4: P = 0.03; Figure 4), whereas there was no significant association at edges (F = 0.23: P = 0.64; Figure 4). We found 340 no clear pattern of spatial autocorrelation in any of our model residuals (Supplementary Material 341 2). 342

343

344 Discussion

Fragmentation and edge effects are reducing the potential of forests to store carbon pantropically (Laurance *et al.* 2002; Durán *et al.* 2013; Magnago *et al.* 2015ab). Here we showed that fragmentation processes change a broad suite of abiotic environmental conditions, which could benefit lianas over trees, and that trees and lianas reveal contrasting responses to fragmentation area and edge effects. We examine these results in detail by first assessing our research questions before investigating their management implications for tropical forests.

352 Abiotic drivers of trees and lianas

Fragment size and edge location capture almost all of the variation in the abiotic variables we recorded. These variables were associated with significant reductions in trees carbon stock, but only within fragment interiors (Laurance *et al.* 2001, 2006). Many studies suggest that forest succession at edges alters forest structure to protect the fragment interiors from the hotter and drier matrix (Laurance *et al.* 2006). Our study adds to this by suggesting that structural changes at edges occur regardless of fragment size.

359 Many of the abiotic variables could explain the loss of tree carbon stocks. We observed increased wind speeds in edges (Figures 2 and 4) and it is known that this can extend far (up to 360 361 400 m) into forest fragments, causing biomass loss and posterior mortality due to the physical 362 damage in tree canopies (Laurance et al. 1998, 2000, 2002; Magnago et al. 2015b). The decrease of air humidity and increase of air temperatures could also have contributed to the 363 overall reduction in tree carbon stocks: droughts and extreme high temperatures are related with 364 carbon reduction due to enhanced tree mortality and changed tree species composition at 365 366 regional scales (Rolim et al. 2005; Laurance et al. 2006; Allen et al. 2010). In addition, these 367 microclimatic changes can promote the replacement of slow growing hardwood species with fast growing softwood species, which have lower carbon storage capacity (Laurance et al. 2006). 368

369

370 Abiotic drivers of liana dominance

Liana's abundance and carbon stock was generally higher within small fragments and at edges, where the soils were more fertile and the microclimate was hotter and drier (Schnitzer & Bongers 2002; Laurance *et al.* 2014; Durán *et al.* 2015). There are four good reasons why lianas are more successful than trees in these conditions. First, their flexible stems mean they are not
affected by elevated wind speeds. Second, their stomatal control and deep roots mean they can
survive high temperatures and low humidity whilst maintaining productivity during dry events
(Chen *et al.* 2015);

Third, lianas could benefit from increased soil fertility stimulated by: (i) the burning of 378 biomass in the matrix, which rapidly supplies nutrients and increases pH at forest edges (Moreira 379 et al. 2009); and (ii) the wind-blown input of soluble nutrients from artificial fertilizers applied in 380 nearby agricultural plantations (Selle 2007; Moreira et al. 2009). Finally, liana growth could be 381 382 further enhanced by plant-soil feedbacks (Kulmatiski et al. 2008; van der Putten et al. 2013), as the high rates of tree mortality and turnover in fragment edges, plus the high levels of leaf 383 384 production by lianas and pioneer trees (Laurance et al. 2002; van der Heijden et al. 2015), can 385 return greater amounts of nutrients to the top soil

386

387 Interactions between trees and lianas

Could some of the patterns we observed be driven by competition between the plant 388 forms and the interaction with disturbance? For example, lianas can be stronger competitors with 389 390 trees in more stressful and disturbed regimes due to their better-developed root system and higher root extension rate (Schnitzer & Bongers 2002, 2011; Restom & Nepstad 2004). These 391 adaptions allow lianas to exploit soil resources, such as nutrients and deep-water (Schnitzer & 392 393 Bongers 2002, 2011). Although our snapshot study cannot demonstrate causation, it does provide some insights into the likelihood of liana-tree competition and highlights some areas for new 394 395 research. In particular, if lianas and trees do compete, our results indicate that this either only happens in forest interiors, or that there is a threshold level at which lianas no longer have an 396

effect (Figure 4), because the composition of trees at edges has changed to one with more
defenses against lianas (e.g., fast growth in pioneer species) (van der Heijden & Phillips 2009;
Magrach *et al.* 2016). Both are plausible, but the fact that trees carbon stock was consistently low
in forest edges strongly suggests that microclimatic variables at edges can affect trees
independently of liana dominance.

Evidence for liana-tree competition was stronger in forest interiors, where there was a 402 very strong negative relationship between lianas dominance and trees carbon stock (Figure 5). 403 Again, it is not clear if this association is driven by the contrasting responses of lianas and trees 404 405 to abiotic variables that change with forest fragmentation or through direct competition for water, soil, and light. In practice, these factors are likely to occur simultaneously: for example, due to 406 wind increases, fragmentation causes mechanical damage to trees (Laurance et al. 2002), 407 408 potentially benefitting lianas, which are better adapted to exploit the canopy gaps (Schnitzer et al. 2014). In turn, this could increase mechanical damage to trees resulting from lianas 409 themselves, as well as increase competition for soil water – increasing the likelihood for further 410 tree mortality. Experiments are needed to assess to what extent lianas are driving changes in tree 411 412 communities in undisturbed forest interiors, or merely responding to them. 413 It is important to note that our snapshot study did not assess the effect of drought (Brando et al. 2014; Rowland et al. 2015), and greater insights into the relevance of competitive 414 interactions between trees and lianas could be gleaned by undertaking experimental 415 416 manipulations across gradients of soil moisture and water availability. In addition, we did not have wood density values to adjust the carbon equation of lianas. Thus, if liana wood density 417

418 changes with habitat disturbance, mainly in forest edges and small fragments where lianas could

419 be softer than in interiors or larger fragments (as expected for trees; Laurance *et al.* 2006), then it

420 is possible that lianas carbon stocks are over-estimated in edges and/or smaller fragments.

421 Implications for forest management and conclusions

Liana cutting is widely used as a management technique to enhance tree productivity and 422 423 restore the functionality of fragments or to reduce collateral damage during selective logging (Edwards, Fisher & Boyd 2010; Kainer, Wadt & Staudhammer 2014). However, the utility of 424 such management will depend on the presence of competition between plant forms. Liana 425 dominance varies among different tropical forests with, for example, reduced liana abundance 426 427 and biomass across altitudinal gradients in the Atlantic Forest (Alves et al. 2012) and, between forest ecosystems, and across soil fertility and climate gradients (Schinitzer & Bongers 2011). 428 Furthermore, lianas are not normally the first cause of edge-related tree mortality, which is 429 430 particularly intense in the few years following edge creation (Laurance et al. 1998), in part because microclimatic changes are especially strong near new edges, which are hotter, drier and 431 windier (Laurance et al. 2002). Since most trees along newly formed edges are not 432 physiologically acclimated to these conditions, they simply die standing or via wind throw, 433 434 especially the big trees (Laurance et al. 2000). In this scenario, lianas only proliferate after the 435 canopy opens due to the initial tree mortality (Schinitzer & Bongers 2011) and appear to be responding to rather than driving change. As such, removing lianas is unlikely to prevent tree 436 mortality. 437

Liana removal could even jeopardise recovery of forest structure and functionality in the likely scenario that lianas themselves help to buffer forest interiors. Here, abiotic changes would continue to depress trees carbon stock after lianas have been cleared by management, and clearance could even worsen conditions. For example, Rolim *et al.* (2005) showed that El Niño

drought negatively affects the above-ground biomass of trees species even in forest interiors of a 442 large Lowland Atlantic Forest fragment. Therefore, we are not advocating that lianas dominance 443 cannot cause impacts on trees, but similar studies should be conducted in more tropical forest 444 regions before making general conclusions about how to manage forest to enhance carbon stocks 445 near edges. Our study also shows that, for the Brazilian Atlantic Forest, fragmentation-induced 446 447 changes of several microclimatic and soil variables can only be reduced by increasing fragment size and reducing edge-interior ratios. This would necessitate a major program of forest 448 restoration (Lamb & Parrotta 2005). 449

450 We conclude that tree carbon stocks are negatively impacted by fragmentation via direct links with altered microclimatic and soil conditions. These include low air humidity and high air 451 452 temperatures, which can increase mortality rate and change species composition, factors likely to 453 alter plant physiology, or increased wind speed, which can cause mechanical damage and mortality (Laurance et al. 2006; Magnago et al. 2015ab). As the vast majority (83%) of the 454 remaining Brazilian Atlantic Forests are found within fragments of less than 50 ha and 46% is 455 located less than 100 m from an edge (Ribeiro et al. 2009), lianas are likely to be an increasingly 456 important feature of this biome. Further research is needed to understand their competitive 457 458 interactions with trees under changing climatic conditions, their carbon storage potential in 459 disturbed forests, and their ability to facilitate edge closure.

460

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473	References
474	Allen, C.D., Macalady, A.K., Chenchouni, H., Bachelet, D., McDowell, N., Vennetier, M.,
475	Kitzberger, T., Rigling, A., Breshears, D.D., Hogg, E.H. (Ted), Gonzalez, P., Fensham, R.,
476	Zhang, Z., Castro, J., Demidova, N., Lim, J.H., Allard, G., Running, S.W., Semerci, A. &
477	Cobb, N. (2010) A global overview of drought and heat-induced tree mortality reveals
478	emerging climate change risks for forests. Forest Ecology and Management, 259, 660–684.
479	Alves, L.F., Assis, M.A., van Melis, J., Barros, A.L.S., Vieira, S.A., Martins, F.R., Martinelli,
480	L.A. & Joly, C.A. (2012) Variation in Liana Abundance and Biomass Along an Elevational
481	Gradient in the Tropical Atlantic Forest (Brazil). Ecological Research, 27, 323–332.
482	Baraloto, C., Molto, Q., Rabaud, S., Hérault, B., Valencia, R., Blanc, L., Fine, P.V.A. &
483	Thompson, J. (2012) Rapid Simultaneous Estimation of Aboveground Biomass and Tree
484	Diversity Across Neotropical Forests: A Comparison of Field Inventory Methods.
485	<i>Biotropica</i> , 45 , n/a–n/a.
486	Berenguer, E., Ferreira, J., Gardner, T.A., Aragão, L.E.O.C., De Camargo, P.B., Cerri, C.E.,
487	Durigan, M., Oliveira, R.C. De, Vieira, I.C.G. & Barlow, J. (2014) A large-scale field

- 488 assessment of carbon stocks in human-modified tropical forests. *Global Change Biology*,
 489 **2005**, 3713–3726.
- 490 Bolker, B.M., Brooks, M.E., Clark, C.J., Geange, S.W., Poulsen, J.R., Stevens, M.H.H. & White,
- J.-S.S. (2009) Generalized linear mixed models: a practical guide for ecology and evolution.
 Trends in Ecology & Evolution, 24, 127–135.
- Bonan, G.B. (2008) Forests and climate change: forcings, feedbacks, and the climate benefits of
 forests. *Science*, **320**, 1444–1449.
- 495 Brando, P.M., Balch, J.K., Nepstad, D.C., Morton, D.C., Putz, F.E., Coe, M.T., Silvério, D.,
- 496 Macedo, M.N., Davidson, E. a, Nóbrega, C.C., Alencar, A. & Soares-Filho, B.S. (2014)
- Abrupt increases in Amazonian tree mortality due to drought-fire interactions. *Proceedings*of the National Academy of Sciences of the United States of America, 111, 6347–52.
- 499 Briant, G., Gond, V. & Laurance, S.G.W. (2010) Habitat fragmentation and the desiccation of
- forest canopies: A case study from eastern Amazonia. *Biological Conservation*, 143, 2763–
 2769.
- 502 Burnham, K.P., Anderson, D.R. & Huyvaert, K.P. (2011) AIC model selection and multimodel
- 503 inference in behavioral ecology: some background, observations, and comparisons.
- 504 *Behavioral Ecology and Sociobiology*, **65**, 23–35.
- 505 Chave, J., Andalo, C., Brown, S., Cairns, M.A., Chambers, J.Q., Eamus, D., Folster, H.,
- 506 Fromard, F., Higuchi, N., Kira, T., Lescure, J.P., Nelson, B.W., Ogawa, H., Puig, H., Riera,
- B. & Yamakura, T. (2005) Tree allometry and improved estimation of carbon stocks and
 balance in tropical forests. *Oecologia*, 145, 87–99.
- 509 Chave, J., Coomes, D., Jansen, S., Lewis, S.L., Swenson, N.G. & Zanne, A.E. (2009) Towards a
- 510 worldwide wood economics spectrum. *Ecology Letters*, **12**, 351–366.

511	Chave, J., Olivier, J., Bongers, F., Chatelet, P., Forget, PM., van der Meer, P., Norden, N.,
512	Riera, B. & Charles-Dominique, P. (2008) Above-ground biomass and productivity in a rain
513	forest of eastern South America. Journal of Tropical Ecology, 24, 355–366.
514	Chen, YJ., Cao, KF., Schnitzer, S.A., Fan, ZX., Zhang, JL. & Bongers, F. (2015) Water-
515	use advantage for lianas over trees in tropical seasonal forests. The New phytologist, 205,
516	128–36.
517	Chen, J.Q., Saunders, S.C., Crow, T.R., Naiman, R.J., Brosofske, K.D., Mroz, G.D., Brookshire,
518	B.L. & Franklin, J.F. (1999) Microclimate in forest ecosystem and landscape ecology -
519	Variations in local climate can be used to monitor and compare the effects of different
520	management regimes. BioScience, 49, 288–297.
521	Didham, R.K. & Lawton, J.H. (1999) Edge Structure Determines the Magnitude of Changes in
522	Microclimate and Vegetation Structure in Tropical Forest Fragments1. Biotropica, 31, 17-
523	30.
524	Dormann, C.F., Mcpherson, J.M., Arau, M.B., Bivand, R., Bolliger, J., Carl, G., Davies, R.G.,
525	Hirzel, A., Jetz, W., Kissling, W.D., Ohlemu, R., Peres-neto, P.R., Schurr, F.M. & Wilson,
526	R. (2007) Methods to account for spatial autocorrelation in the analysis of species
527	distributional data : a review., 609–628.
528	Durán, S.M. & Gianoli, E. (2013) Carbon stocks in tropical forests decrease with liana density
529	Carbon stocks in tropical forests decrease with liana density. <i>Biology letters</i> , 2013–2016.
530	Durán, S.M., Sánchez-Azofeifa, G.A., Rios, R.S. & Gianoli, E. (2015) The relative importance
531	of climate, stand variables and liana abundance for carbon storage in tropical forests. Global
532	Ecology and Biogeography, 24, 939–949.
533	Edwards, D.P., Fisher, B. & Boyd, E. (2010) Protecting degraded rainforests: Enhancement of

- forest carbon stocks under REDD+. *Conservation Letters*, **3**, 313–316.
- Field, C.B. (1998) Primary Production of the Biosphere: Integrating Terrestrial and Oceanic
 Components. *Science*, 281, 237–240.
- 537 Flores, O. & Coomes, D.A. (2011) Estimating the wood density of species for carbon stock

assessments. *Methods in Ecology and Evolution*, **2**, 214–220.

- 539 Gibbs, H.K., Ruesch, A.S., Achard, F., Clayton, M.K., Holmgren, P., Ramankutty, N. & Foley,
- J.A. (2010) Tropical forests were the primary sources of new agricultural land in the 1980s
 and 1990s. *Proceedings of the National Academy of Sciences*, **107**, 16732–16737.
- 542 Hawes, J.E., Peres, C.A., Riley, L.B. & Hess, L.L. (2012) Landscape-scale variation in structure
- and biomass of Amazonian seasonally flooded and unflooded forests. *Forest Ecology and Management*, 281, 163–176.
- Hazelton, P. & Murphy, B. (2007) Interpreting Soil Test Results: What do all the Numbers

546 mean? - by P.; B.Murphy. *NSW Department of Natural Resources*.

- van der Heijden, G.M.F. & Phillips, O.L. (2009) Liana infestation impacts tree growth in a
- 548 lowland tropical moist forest. *Biogeosciences Discussions*, **6**, 3133–3158.
- van der Heijden, G., Powers, J.S. & Schnitzer, S.A. (2015) Lianas reduce forest-level carbon
 accumulation and storage. *Nature*, **112**, 13267–13271.
- 551 Kainer, K.A., Wadt, L.H.O. & Staudhammer, C.L. (2014) Testing a silvicultural
- recommendation: Brazil nut responses 10 years after liana cutting. *Journal of Applied Ecology*, **51**, 655–663.
- Kulmatiski, A., Beard, K.H., Stevens, J.R. & Cobbold, S.M. (2008) Plant-soil feedbacks: a metaanalytical review. *Ecology letters*, 11, 980–92.
- Lamb, D., Erskine, P.D. & Parrotta, J. a. (2005) Restoration of degraded tropical forest

557	landscapes.	Science	(New	York,	N.Y.),	310,	1628-	-32
-----	-------------	---------	------	-------	--------	------	-------	-----

- Laurance, W.F., Andrade, A.S., Magrach, A., Camargo, J.L.C., Valsko, J.J., Campbell, M.,
- 559 Fearnside, P.M., Edwards, W., Lovejoy, T.E. & Laurance, S.G. (2014) Long-term changes
- 560 in liana abundance and forest dynamics in undisturbed Amazonian forests. *Ecology*, **95**,
- 561 1604–1611.
- Laurance, W.F., Delamônica, P., Laurance, S.G., Vasconcelos, H.L. & Lovejoy, T.E. (2000)
 Rainforest fragmentation kills big trees. *Nature*, 404, 836.
- Laurance, W.F., Ferreira, L. V., Rankin-de Merona, J.M. & Laurance, S.G. (1998) Rain forest
- fragmentation and the dynamics of Amazonian tree communities. *Ecology*, **79**, 2032–2040.
- Laurance, W.F., Lovejoy, T.E., Vasconcelos, H.L., Bruna, E.M., Didham, R.K., Stouffer, P.C.,
- 567 Gascon, C., Bierregaard, R.O., Laurance, S.G. & Sampaio, E. (2002) Ecosystem decay of
- 568 Amazonian forest fragments: A 22-year investigation. *Conservation Biology*, **16**, 605–618.
- Laurance, W.F., Nascimento, H.E.M., Laurance, S.G., Andrade, A., Ribeiro, J.E.L.S., Giraldo,
- 570 J.P., Lovejoy, T.E., Condit, R., Chave, J., Harms, K.E. & D'Angelo, S. (2006) Rapid decay
- of tree-community composition in Amazonian forest fragments. *Proceedings of the*
- 572 *National Academy of Sciences of the United States of America*, **103**, 19010–19014.
- 573 Laurance, W.F., Pérez-Salicrup, D., Delamônica, P., Fearnside, P.M., D'Angelo, S., Jerozolinski,
- A., Pohl, L. & Lovejoy, T.E. (2001) Rain forest fragmentation and the structure of
 Amazonian liana communities. *Ecology*, 82, 105–116.
- 576 Lewis, S.L., Edwards, D.P. & Galbraith, D. (2015) Increasing human dominance of Tropical
- 577 Forests. *Science*.
- 578 Lewis, D.B. & Feit, S.J. (2015) Connecting carbon and nitrogen storage in rural wetland soil to
- 579 groundwater abstraction for urban water supply. *Global Change Biology*, **21**, 1704–1714.

580	Magnago, L.F.S., Edwards, D.P., Edwards, F.A., Magrach, A., Martins, S. V. & Laurance, W.F.
581	(2014) Functional attributes change but functional richness is unchanged after
582	fragmentation of Brazilian Atlantic forests. Journal of Ecology, 102, 475–485.
583	Magnago, L.F.S., Magrach, A., Laurance, W.F., Martins, S. V., Meira-Neto, J.A.A., Simonelli,
584	M. & Edwards, D.P. (2015a) Would protecting tropical forest fragments provide carbon and
585	biodiversity cobenefits under REDD+? Global Change Biology, 21, 3455–3468.
586	Magnago, L.F.S., Rocha, M.F., Meyer, L., Martins, S.V. & Meira-Neto, J.A.A. (2015b)
587	Microclimatic conditions at forest edges have significant impacts on vegetation structure in
588	large Atlantic forest fragments. Biodiversity and Conservation, 2305–2318.
589	Magrach, A., Rodríguez-Pérez, J., Campbell, M. & Laurance, W.F. (2014) Edge effects shape
590	the spatial distribution of lianas and epiphytic ferns in Australian tropical rain forest
591	fragments. Applied Vegetation Science, 17, 754–764.
592	Magrach, A., Senior, R.A., Rogers, A., Nurdin, D., Benedick, S., Laurance, W.F., Santamaria, L.
593	& Edwards, D.P. (2016) Selective logging in tropical forests decreases the robustness of
594	liana-tree interaction networks to the loss of host tree species. Proceedings of the Royal
595	Society of London B: Biological Sciences, 283.
596	Malhi, Y., Baker, T.R., Phillips, O.L., Almeida, S., Alvarez, E., Arroyo, L., Chave, J., Czimczik,
597	C.I., Di Fiore, A., Higuchi, N., Killeen, T.J., Laurance, S.G., Laurance, W.F., Lewis, S.L.,
598	Montoya, L.M.M., Monteagudo, A., Neill, D. a., Vargas, P.N., Patino, S., Pitman, N.C. a,
599	Quesada, C.A., Salomao, R., Silva, J.N.M., Lezama, A.T., Martínez, R.V., Terborgh, J.,
600	Vinceti, B. & Lloyd, J. (2004) The above-ground coarse wood productivity of 104
601	Neotropical forest plots. <i>Global Change Biology</i> , 10 , 563–591.
602	Moreira, F.M.D.S., Nóbrega, R.S.A., Jesus, E.D.C., Ferreira, D.F. & Pérez, D.V. (2009)

603	Differentiation in the fertility of Inceptisols as related to land use in the upper Solimões
604	river region, western Amazon. Science of the Total Environment, 408, 349-355.
605	Peixoto, A.L. & Gentry, A. (1990) Diversidade e composição florística da mata de tabuleiro na
606	Reserva Florestal de Linhares (Espírito Santo, Brasil). Revista Brasileira de Botânica, 13,
607	19–25.
608	Phillips, O.L., Higuchi, N., Vieira, S., Baker, T.R., Chao, K.J. & Lewis, S.L. (2009) Changes in
609	Amazonian forest biomass, dynamics, and composition, 1980-2002. Geophysical
610	Monograph Series, 186 , 373–387.
611	Phillips, O.L., Vásquez Martínez, R., Arroyo, L., Baker, T.R., Killeen, T., Lewis, S.L., Malhi,
612	Y., Monteagudo Mendoza, A., Neill, D., Núñez Vargas, P., Alexiades, M., Cerón, C., Di
613	Fiore, A., Erwin, T., Jardim, A., Palacios, W., Saldias, M. & Vinceti, B. (2002) Increasing
614	dominance of large lianas in Amazonian forests. Nature, 418, 770–774.
615	Poorter, L., McDonald, I., Alarcón, A., Fichtler, E., Licona, JC., Peña-Claros, M., Sterck, F.,
616	Villegas, Z. & Sass-Klaassen, U. (2010) The importance of wood traits and hydraulic
617	conductance for the performance and life history strategies of 42 rainforest tree species. The
618	<i>New phytologist</i> , 185 , 481–92.
619	van der Putten, W.H., Bardgett, R.D., Bever, J.D., Bezemer, T.M., Casper, B.B., Fukami, T.,
620	Kardol, P., Klironomos, J.N., Kulmatiski, A., Schweitzer, J.A., Suding, K.N., Van de
621	Voorde, T.F.J. & Wardle, D.A. (2013) Plant-soil feedbacks: the past, the present and future
622	challenges (ed M Hutchings). Journal of Ecology, 101, 265–276.
623	Restom, T.G. & Nepstad, D.C. (2004) Seedling growth dynamics of a deeply rooting liana in a
624	secondary forest in eastern Amazonia. Forest Ecology and Management, 190, 109–118.
625	Ribeiro, M.C., Metzger, J.P., Martensen, A.C., Ponzoni, F.J. & Hirota, M.M. (2009) The

- Brazilian Atlantic Forest: How much is left, and how is the remaining forest distributed?
 Implications for conservation. *Biological Conservation*, 142, 1141–1153.
- 628 Rolim, S.G., Jesus, R.M., Nascimento, H.E.M., do Couto, H.T.Z. & Chambers, J.Q. (2005)
- Biomass change in an Atlantic tropical moist forest: the ENSO effect in permanent sample
- 630 plots over a 22-year period. *Oecologia*, **142**, 238–246.
- Ronquim, C. (2010) Conceitos de fertilidade do solo e manejo adequado para regiões tropicais. *Boletim de Pesquisa e Desenvolvimento*, **8**, 26.
- Rowland, L., da Costa, A.C.L., Galbraith, D.R., Oliveira, R.S., Binks, O.J., Oliveira, A.A.R.,
- Pullen, A.M., Doughty, C.E., Metcalfe, D.B., Vasconcelos, S.S., Ferreira, L. V, Malhi, Y.,
- Grace, J., Mencuccini, M. & Meir, P. (2015) Death from drought in tropical forests is
 triggered by hydraulics not carbon starvation. *Nature*, **528**, 119–122.
- 637 Saatchi, S.S., Harris, N.L., Brown, S., Lefsky, M., Mitchard, E.T.A., Salas, W., Zutta, B.R.,
- Buermann, W., Lewis, S.L., Hagen, S., Petrova, S., White, L., Silman, M. & Morel, A.
- 639 (2011) Benchmark map of forest carbon stocks in tropical regions across three continents.
- 640 *Proceedings of the National Academy of Sciences*, **108**, 9899–9904.
- 641 Schnitzer, S.A. & Bongers, F. (2002) The ecology of lianas and their role in forests. *Trends in*642 *Ecology and Evolution*, **17**, 223–230.
- Schnitzer, S.A. & Bongers, F. (2011) Increasing liana abundance and biomass in tropical forests:
 emerging patterns and putative mechanisms. *Ecology letters*, 14, 397–406.
- 645 Schnitzer, S.A., Dewalt, S.J. & Chave, J. (2006) Censusing and Measuring Lianas: A
- 646 Quantitative Comparison of the Common Methods. *Biotropica*, **38**, 581–591.
- 647 Schnitzer, S. a., van der Heijden, G.M.F., Mascaro, J. & Carson, W.P. (2014) Lianas in Gaps
- 648 Reduce Carbon Accumulation in a Tropical Forest. *Ecology*, in press, 140515081851004.

649 Selle, G.L. (2007) Ciclagem de nutrientes em ecossistemas florestais. *Bioscience Journal*, 23,
650 29–39.

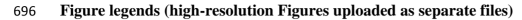
651	Solar, R.R. de C., Barlow, J., Ferreira, J., Berenguer, E., Lees, A.C., Thomson, J.R., Louzada, J.,
652	Maués, M., Moura, N.G., Oliveira, V.H.F., Chaul, J.C.M., Schoereder, J.H., Vieira, I.C.G.,
653	Mac Nally, R. & Gardner, T.A. (2015) How pervasive is biotic homogenization in human-
654	modified tropical forest landscapes? (ed H Cornell). Ecology Letters, 18, 1108-1118.
655	Stegen, J.C., Swenson, N.G., Enquist, B.J., White, E.P., Phillips, O.L., Jørgensen, P.M., Weiser,
656	M.D., Monteagudo Mendoza, A. & Núñez Vargas, P. (2011) Variation in above-ground
657	forest biomass across broad climatic gradients. Global Ecology and Biogeography, 20, 744-
658	754.
659	Tollefson, J. (2013) Brazil reports sharp drop in greenhouse emissions. <i>Nature</i> , 11–13.
660	van der Werf, G.R., Morton, D.C., DeFries, R.S., Olivier, J.G.J., Kasibhatla, P.S., Jackson, R.B.,
661	Collatz, G.J. & Randerson, J.T. (2009) CO2 emissions from forest loss. Nature Geoscience,
662	2 , 737–738.
663	Zanne, A., Lopez-Gonzalez, G., DA, C., Ilic, J., Jansen, S., SL, L., RB, M., NG, S., MC, W. &
664	Chave, J. (2009) Data from: Towards a worldwide wood economics spectrum. Ecology

665 *Letters*.

Zuur, A.F., Ieno, E.N., Walker, N.J., Saveliev, A. a, Smith, G.M. & Ebooks Corporation. (2009)
Mixed Effects Models and Extensions in Ecology with R. *Statistics for Biology and Health*,
579 p.

670 Supplementary Materials include:

- 671 Text S1 Conservation value of the study region.
- Figure S1 Study area and forest fragments sampled in South-eastern Brazil.
- Figure S2 Principal Correlation Analyzes (PCA) of microclimate, soil and fragment size values.
- Figure S3 Correlations between PCA Axis 1 and microclimate, soil and fragment size values.
- Figure S4 Best GLMM model of the PCA Axis 1 with fragment size and edge-interior location.
- Figure S5 Graphs of best model of lianas abundance in relation to fragment size gradients.
- Figure S6 Graphs of best model of lianas abundance in relation to fragmentation abiotic
- 678 environmental gradients.
- Table S1 Identification and variable values of sampled fragments.
- 680 Table S2 Model selection of the Generalized Linear Mixed Models for all microclimate
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- Table S4 Model selection of the Generalized Linear Mixed Models for Fragmentation Axis 1
- 685 (PCA2) in relation to fragment size and edge-interior location.
- Table S5 Model selection of the Generalized Linear Mixed Models for carbon stocks of treesand lianas and for ratio of lianas to trees in relation to fragment size and edge-interior location.
- Table S6 Model selection of the Generalized Linear Mixed Models for carbon stocks of trees
- and lianas and for ratio of lianas to trees in relation to environmental changes (Fragmentation
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- Table S7 Model selection of the Generalized Linear Mixed Models for tree carbon stocks inrelation to lianas abundances, lianas carbon stocks and edge-interior location.
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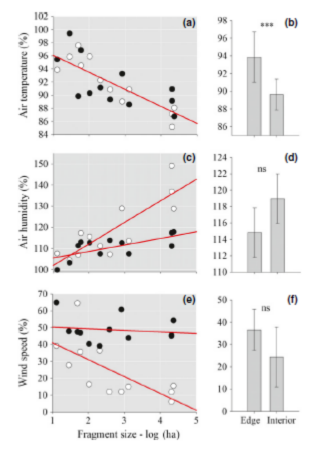


Figure 1. Graphs of best models for the effects of fragments size and edge-interior location on microclimatic variables. (A-B) Effects of fragment size and habitats on air temperature; (C-D) Effects of fragment size and habitat on air humidity; (E-F) Effects of fragment size and habitat on wind speed. Black circles = Edge; White circles = Interior. Circles represent values obtained after the summation of raw residuals to the expected values for each variable, being assumed average values for other covariates. Errors bars represent the 95% of confidence intervals. ***=p<0.001; ns=No significant results.

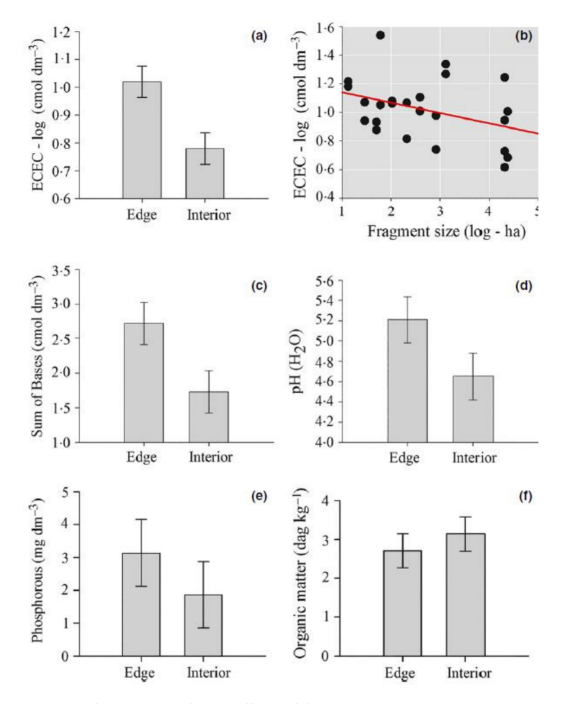


Figure 2. Graphs of best models for the effects of fragments size and edge-interior location on soil attributes. (A) Edge-interior and (B) fragment size effects on soil fertility via Effective Cation Exchange Capacity (ECEC); (C) edge-interior effect on soil fertility via Sum of Bases; (D) edge-interior effect on soil acidity - pH in H₂O; (E) edge-interior effect on phosphorous; (F) edge-interior effect on organic matter. Errors bars represent the 95% of confidence intervals. Circles represent values obtained after the summation of raw residuals with the expected values for each variable, assuming average values for other covariates.

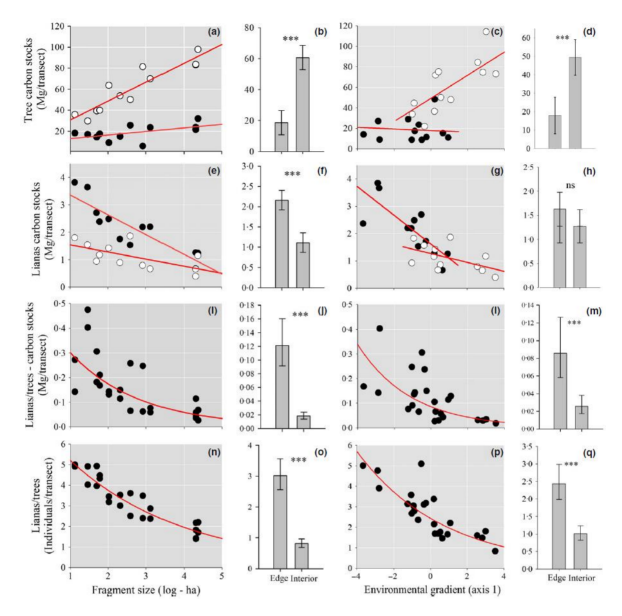
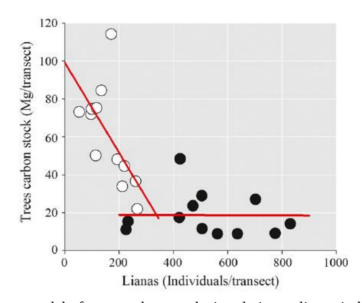


Figure 3. Graphs of best models for the effects of fragments size (A, E, I, N), fragmentation 714 abiotic environmental gradients (Fragmentation Axis 1; C, G, L, P) and edge-interior location (B, 715 D, F, H, J, M, O, Q) on trees carbon stocks (A, B, C, D), lianas carbon stocks (E, F, G, H), ratio 716 of lianas to trees carbon stocks (I, J, L, M), and ratio of lianas to trees individuals (N, O, P, Q). 717 Circles represent values obtained after the summation of raw residuals with the expected values 718 for each variable, assuming average values for other covariates. Black and white circles represent 719 720 edge and interior location, respectively, when there is a significant effect of the interaction term between fragment size and edge-interior location; when only black circles are shown, there is no 721 722 significant interaction effect. Errors bars represent the 95% of confidence intervals.

***=p<0.001; ns=No significant results. Negative values for Fragmentation Axis 1 represent
 more stressful and disturbed conditions, positive values more benign conditions (see Fig. S3).

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Figure 4. Graph of best model of trees carbon stocks in relation to lianas individuals. Circles
represent values obtained after the summation of raw residuals with the expected values for each

variable. Black and white circles represent edge and interior location, respectively, and thus the

- raginificant effect of the interaction term between fragment size and edge-interior location.
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