# Analysis of critical transitions at the Global Forest

# 2 Abstract

Global forests provide critical habitat for many species, essential ecosystem services, and are coupled to atmospheric dynamics through exchanges of energy, water and gases. One of the most important changes produced in the biosphere is the replacement of forest areas with human dominated landscapes. These areas contain many fewer species than the natural ecosystem they replaced, and this is one of the main drivers of extinctions in the Anthropocene. Another effect is that habitat replacement leads to fragmentation, altering the sizes of patches, the structure and function of the forest. In different kinds of vegetation, including forests, the patch size distributions follow power laws. This have been used as a signal of critical phase transitions, where the system state changes suddenly at a critical point; in the case of forest patches, this implies an 10 abrupt change in connectivity that causes a increased fragmentation level and this poses an additional risk to forest function as an habitat and ecosystem. All these patterns have been observed at relatively small scales, however. Here we studied the distribution of forest patch sizes at a global level, its changes over the last fourteen years, and different signals of a critical transition. We defined wide regions of connected forest at continental level and also studied big islands. The conditions that indicate that a region is near a critical 15 fragmentation threshold are: a patch size distributions following a power law; temporal fluctuations of the 16 largest patch following a power law; variance of largest patch fluctuations increasing in time; and a negative 17 skewness of the largest patch fluctuations. Most regions, except the Eurasian mainland, followed a power-18 law distribution. We found that only the tropical forest of Africa and South America met the five criteria 19 and thus seemed to be near a critical fragmentation threshold. This implies that the combined influence of human pressures and climate forcings might trigger undesired effects of fragmentation, such as species loss and degradation of ecosystems services, in these regions. The simple criteria proposed here could be used to estimate the distance to a fragmentation threshold in forest around the globe, and provide a guide to direct conservation efforts at a continental level.

# 25 Introduction

Forests are some the most important ecosystems on earth, providing habitat for a large proportion of species and contributing extensively to global biodiversity (Crowther *et al.*, 2015). In the previous century human activities have influenced global bio-geochemical cycles (Bonan, 2008; Canfield *et al.*, 2010), with one of the most dramatic changes being the replacement of 40% of Earth's formerly biodiverse land areas with

- landscapes that contain only a few species of crop plants, domestic animals and humans (Foley et al., 2011).
- These local changes have accumulated over time and now constitute a global forcing (Barnosky et al., 2012).
- Another global scale forcing that is tied to habitat destruction is fragmentation. Fragmentation is defined as
- 4 the division of a continuous habitat into separated portions that are smaller and more isolated, this produces
- 5 multiple interwoven effects: reductions of biodiversity between 13% and 75%, decreasing forest biomass and
- 6 changes in nutrient cycling (Haddad et al., 2015). The effects of fragmentation are not only important from
- 7 an ecological point of view but also human activities and ecosystem services supply and flow are deeply
- 8 influenced by the level of landscape fragmentation (???).
- 9 Complex interactions between species and feedbacks at different levels of organization (Gilman et al., 2010)
- can produce abrupt changes called critical transitions (Scheffer et al., 2009). These abrupt state shifts cannot
- be linearly forecasted from past changes, and are thus difficult to predict and manage (Scheffer et al., 2009).
- 12 Critical transitions had been detected mostly at local scales (Drake & Griffen, 2010; Carpenter et al., 2011),
- but the accumulation of changes in local communities that overlap geographically can propagate and cause
- an abrupt change of the entire system at larger scales (Barnosky et al., 2012). Coupled with the existence
- 15 of global scale forcings, this implies the possibility that a critical transition could occur at a global scale
- 16 (Rockstrom et al., 2009; Folke et al., 2011).
- There are two general classes of critical transitions. In the first, a sudden shift between states is mostly
- 18 irreversible because of the presence of two stable states and no coexistence between these states is allowed.
- This behavior is called hysteresis an represent a catastrophic regime shift (Scheffer et al., 2001), also called
- 20 first order transitions. This could be the result of a positive feedback mechanisms (Martín et al., 2015), for
- example the growth of forest in a site is more probable if it is surrounded by forest than if it is surrounded
- by cultivated or degraded land (Xu et al., 2015). This has been observed in semiarid regions where the
- 23 transition from a vegetated to a desertic state occurs as a result of human pressures and climatic change
- <sup>24</sup> (Kéfi et al., 2007a), in biological invasions (Taylor & Hastings, 2005), and biology and social sciences in
- 25 general (Solé, 2011).
- The second class are continuous critical transitions (Solé & Bascompte, 2006), or second order transitions. In
- 27 this case there is a narrow region where the system changes from one domain to another, pushed by external
- 28 conditions. The change in this case is sudden, but continuous and theoretically reversible. This region of
- 29 change can be characterized by a critical point where the change begins. At the critical point we can observe
- 30 distinctive spatial patterns: scale invariant fractal structures characterized by power law patch distributions
- 31 (Stauffer & Aharony, 1994). The simplest model where we can observe this kind of transition is a random
- p map, in which each site has an independent probability p of being occupied. Even in this simple setting,

- starting with p = 0, when p reaches a certain value the landscape changes from fragmented to a connected
- <sup>2</sup> state. Another more relevant model concerns forest fires and ecological memory: biomass is consumed in a
- 3 fire, and afterwards some time must pass to build up fuel before a subsequent fire can occur (Zinck & Grimm,
- 4 2009). The behavior of this last model is called a self-organized criticality where the system drives itself to
- 5 the critical region, so the critical behavior is observed for a wide range of parameter with the condition that
- a double separation of scales exist (???). The rate of producing new fires (lightning) should be much slower
- 7 than the rate of growth of trees and in turn the growth of trees is slow compared with the velocity of fire
- 8 spread.
- 9 The spatial phenomena observed in continuous critical transitions deal with connectivity, a fundamental
- property of general systems and ecosystems from forests (Ochoa-Quintero et al., 2015) to marine ecosystems
- (Leibold & Norberg, 2004) and the whole biosphere (Lenton & Williams, 2013). When a system goes from
- 12 a fragmented to a connected state we say that it percolates (Solé, 2011). Percolation implies that there is a
- path of connections that involves the whole system. Thus we can characterize two domains or phases: one
- dominated by short-range interactions where information cannot spread, and another in which long range
- 15 interactions are possible and information can spread over the whole area. (The term "information" is used
- in a broad sense and can represent species dispersal or movement.)
- 17 Thus, there is a critical "percolation threshold" between the two phases, and the system could be driven
- 18 close to or beyond this point by an external force; climate change and deforestation are the main forces that
- could be the drivers of such a phase change in contemporary forests (Bonan, 2008; Haddad et al., 2015).
- 20 Species' dispersal strategies are influenced by percolation thresholds in three-dimensional forest structure
- 21 (Solé et al., 2005), and it has been shown that species distributions also have percolation thresholds (He
- 22 & Hubbell, 2003). This implies that pushing the system below the percolation threshold could produce a
- biodiversity collapse (Pardini2010; Bascompte et al., 1996; Solé et al., 2004); conversely, being in a connected
- 24 state (above the threshold) could accelerate the invasion of forest into prairie (Loehle et al., 1996; Naito &
- 25 Cairns, 2015).
- <sup>26</sup> One of the problems with critical transitions is that the value of the critical threshold is not known in
- 27 advance; in addition, because near that point a small change can precipitate a state shift of the system,
- 28 they are difficult to predict. Several methods have been developed to detect the closeness of the critical
- point, e.g. a deceleration in recovery from perturbations, or an increase in variance in the spatial or temporal
- pattern (Hastings & Wysham, 2010; Carpenter et al., 2011; Boettiger & Hastings, 2012; Dai et al., 2012).
- In forests, we can observe processes such as demographic fluctuations, spatial heterogeneities, and/or dis-
- persal limitation, all processes that introduce stochasticity that can convert a catastrophic transition to a

- second order transitions (Martín et al., 2015). Thus a continuous transitions should be more probable than
- 2 catastrophic transitions, and some evidence of a continuous critical transition related to drought and fire
- has been found in tropical forest (Pueyo et al., 2010), and grasslands (???, ???).
- 4 The necessary condition we need to be near a continuous critical threshold is that the patch size distribution
- 5 follows a power-law. This distributions are present in a wide range of conditions, for example in arid
- 6 ecosystem vegetation (Kéfi et al., 2007b; Scanlon et al., 2007), ant colonies (Vandermeer et al., 2008), and
- 7 attached microalgae (Saravia et al., 2012; Dal Bello et al., 2014). The presence of power-laws is not a
- 8 sufficient condition because they can be produced by different mechanisms. One hypothesis is that these
- 9 power-laws were produced by facilitation mechanisms (Manor & Shnerb, 2008; Irvine et al., 2016), that are
- likely to be present in forest (Bulleri et al., 2008).
- 11 In this study, our objectives were to evaluate what models best describe forest patch distribution at a
- 12 continental scale; examine how patch size and connectivity change between regions and across time; and
- 13 look for evidence that forests are near critical fragmentation thresholds. The advantage of using data at a
- continental scale is that for very large systems the transitions are very sharp (Solé, 2011) and thus much
- 15 easier to detect than at smaller scales, where noise can mask the signals of the transition.

# 16 Methods

### 17 Area definition

- We analyzed mainland zones at a continental scale, specifying that close patches of continuous forest must
- connect the region [TODO: what does this mean?], big islands of more than XXXX km<sup>2</sup>-like Madagascar—
- 20 were treated as separate areas, unless they were < 1 km from the mainland, while smaller islands were
- excluded. With these criteria we defined three regions in America: South America temperate forest (SAT),
- subtropical and tropical forest up to Mexico (SAST), and USA and Canada forest (NA). Europe and north
- 23 Asia were all connected and treated as one region (EUAS). Other regions were south Asia (SEAS), Africa
- <sup>24</sup> (AF), and Australia and islands (OC) (Supplementary figure S1-S6). Each region have subregions that
- <sup>25</sup> correspond to separate areas inside a continent or to nearby islands (Table 1).
- 26 To define patches we use the MODerate-resolution Imaging Spectroradiometer (MODIS) Vegetation Con-
- 27 tinuous Fields (VCF) Tree Cover dataset, version 051 (DiMiceli et al., 2015). This dataset is produced at
- <sup>28</sup> 250-m resolution, globally from 2000 to 2014. There are several definition of forest (Sexton et al., 2015), but
- $_{29}$  we choose a 30% threshold to convert the percentage tree cover to a binary image of forest and non-forest

- pixels. This was the definition used by the United Nations' International Geosphere-Biosphere Programme
- <sup>2</sup> (Belward, 1996), and studies of global fragmentation (Haddad et al., 2015). This definition avoids the errors
- 3 produced by low discrimination of MODIS VCF between forest and dense herbaceous vegetation at low
- 4 forest cover (Sexton et al., 2015). Patches of contiguous forest were determined in the binary image by
- 5 grouping connected pixels using a neighborhood of 8 forest units (Moore neighborhood). To fit the patch
- size distributions we set a minimal patch size  $(X_{min})$  at nine pixels to avoid artifacts at patch edges due to
- <sup>7</sup> discretization (Weerman et al., 2012).

### 8 Percolation theory

<sup>9</sup> A more in-depth introduction to percolation theory can be found elsewhere (Stauffer & Aharony, 1994) and a

review from an ecological point of view is available (Oborny et al., 2007). Here to explain the basic elements

of percolation theory we formulate a simple model: we represent our area of interest by a square lattice

and each site of the lattice can be occupied—e.g. by forest—with a probability p. The lattice will be more

p occupied when p is greater, but the sites are randomly distributed. We are interested in the connection

between sites so define a neighborhood, the eight adjacent sites surrounding any particular site. The sites

that are neighbors define a patch. When there is a patch that connects the lattice from opposite sides it

is said that the system percolates. The percolating patch suddenly appears at some value of p called the

17 critical point  $p_c$ .

Thus percolation is characterized by two well defined phases: the unconnected phase (called subcritical in

physics) when  $p < p_c$ , in which species cannot travel far inside the forest, as it is fragmented; in a general

sense, information cannot spread. The second is the connected phase (supercritical) when  $p>p_c$ : species

21 can move inside a forest patch from side to side of the area (lattice), i.e. information can spread over the

whole area. Near the critical point several scaling laws arise: the structure of the patch that spans the area

is fractal, the size distribution of the patches is power-law, and other quantities also follow power-law scaling

<sup>24</sup> (Stauffer & Aharony, 1994).

The value of the critical point  $p_c$  depends on the geometry of the lattice and on the definition of the

26 neighborhood, but other power-law exponents only depend on the lattice dimension. Close to criticality the

distribution of patch sizes is:

(1) 
$$n_s(p_c) \propto s^{-\alpha}$$

where  $n_s(p)$  is the number of patches of size s. The exponent  $\alpha$  does not depend on the details of the

model and it is called universal (Stauffer & Aharony, 1994). These scaling laws can be applied for landscape

- structures that are approximately random, with correlations if the forest sites are correlated over short
- <sup>2</sup> distances (Gastner et al., 2009). In physics this is called "isotropic percolation universality class", and
- $\alpha$  corresponds to an exponent  $\alpha = 2.05495$ ; if we observe that the patch size distribution has another exponent
- 4 it will not belong to this universality class and some other mechanism should be invoked to explain it.
- <sup>5</sup> Percolation thresholds can also be generated by models that have some kind of memory (Hinrichsen, 2000;
- 6 Odor, 2004): for example, a patch that has been exploited for many years will recover different than a
- recently deforested forest patch. In this case the system can belong to a different universality class, or in
- some cases there is no universality, in which case the value of  $\alpha$  will depend on the parameters and details
- 9 of the model (Corrado et al., 2014).

#### 10 Patch size distributions

- We fitted the empirical distribution of forest patch areas, based on the MODIS-derived data described above,
- to four distributions using maximum likelihood estimation (Goldstein et al., 2004; Clauset et al., 2009).
- 13 The distributions were: power-law, power-law with exponential cut-off, log-normal, and exponential. We
- 14 assumed that the patch size distribution is a continuous variable that was discretized by remote sensing data
- 15 acquisition procedure.
- Besides the hard  $X_{min}$  limit we set due to discretization, the power-law distribution needs a lower bound
- for its scaling behaviour. This lower bound is also estimated from the data by maximizing the Kolmogorov-
- Smirnov (KS) statistic comparing the empirical to fitted cumulative distribution functions (Clauset et al.,
- <sup>19</sup> 2009). We also calculated the uncertainty of the parameters using a non-parametric bootstrap method (Efron
- <sup>20</sup> & Tibshirani, 1994), and computed corrected Akaike Information Criteria (AIC<sub>c</sub>) and Akaike weights for
- each model (Burnham & Anderson, 2002). Akaike weights  $(w_i)$  are the weight of evidence in favor of model
- i being the actual best model given that one of the N models must be the best model for that set of N
- 23 models.
- <sup>24</sup> Additionally, we computed the goodness of fit of the power-law model following the bootstrap approach
- described by Clauset et. al (2009), where simulated data sets following the fitted model are generated, and a
- <sub>26</sub> p-value calculated as the proportion of simulated data sets that has a KS statistic less extreme than empirical
- data. The criterion to reject the power law model was  $p \leq 0.1$  but as we have a very large n, meaning that
- negligible small deviations could produce a rejection (???), we chose a  $p \leq 0.05$  to reject the power law
- 29 model.
- 30 To test for differences between the fitted power law exponent for regions and subregions we used a weighted

- linear regression with a residual auto-correlation structure. For this we use a generalized least squares pro-
- <sup>2</sup> cedures: function gls from package nlme (???). The weights were the bootstraped 95% confidence intervals
- 3 and we added an auto-regressive model of order 1 to the residuals to account for temporal autocorrelation.

### 4 Largest patch dynamics

5 The largest patch is the one that connects the highest number of sites in the area. This has been used

extensively to indicate fragmentation (Gardner & Urban, 2007; Ochoa-Quintero et al., 2015). The relation

of the size of the largest patch  $S_{max}$  to critical transitions has been extensively studied in relation to

e percolation phenomena (Bazant2000;Botet2004; Stauffer & Aharony, 1994), but seldom used in ecological

studies (but see (2009)). When in a connected state  $(p > p_c)$  the landscape is almost insensitive to the loss

of a small fraction of forest, but close to the critical point the same loss can have important effects (Solé

<sup>11</sup> & Bascompte, 2006; Oborny et al., 2007), because at this point the largest patch will have a filamentary

12 structure, i.e. extended forest areas will be connected by thin threads. Small losses can thus produce large

13 fluctuations.

One way to evaluate the fragmentation of the forest is to calculate the proportion of the largest patch against

15 the total area (???). The total area of the regions we are considering (Supplementary figure S1-S6) may not

be the same than the total area that the forest could potentially occupy, thus a more accurate way to evaluate

the weight of  $S_{max}$  is to we use the total forest area. We calculate the proportion of the largest patch for each

year, dividing  $S_{max}$  by the total forest area of the same year:  $RS_{max} = S_{max} / \sum_i S_i$ . This has the effect of

reducing the  $S_{max}$  fluctuations produced due to environmental or climatic changes influences in total forest

area. When the proportion  $RS_{max}$  is large (more than 60%) the largest patch contains most of the forest so

there are fewer small forest patches and the system is probably in a connected phase. Conversely, when it is

low (less than 20%), the system is probably in a fragmented phase.

It is also possible that a system with high  $RS_{max}$  is near a percolation threshold, and this has to be evaluated

24 using the temporal fluctuations. We calculate the fluctuations around the mean with the absolute values

 $\Delta S_{max} = S_{max}(t) - \langle S_{max} \rangle$ , using the proportions of  $RS_{max}$ . To characterize the fluctuations we fitted

three empirical distributions: power-law, log-normal, and exponential, using the same methods described

previously. We expect that large fluctuation near a critical point have heavy tails (log-normal or power-law)

<sup>28</sup> and that fluctuations far from a critical point have exponential tails, corresponding to Gaussian processes.

29 As the data set spans 15 years we have do not have enough power to reliably detect which distribution is

better (Clauset et al., 2009). To improve this we performed the goodness of fit test described above for all

- the distributions. We generated animated maps showing the fluctuations of the 2 largest patches to aid in
- 2 the interpretations of the results.
- 3 A robust way to detect if the system is near a critical transition is to analyze the increase in variance of the
- 4 density (Benedetti-Cecchi et al., 2015), in our case density means the area of forest divided by total area.
- 5 One of the problems is that density is difficult to determine because we do not know the total area that the
- 6 forest can potentially colonize, this has to be estimated introducing more uncertainty in the density. The
- 7 other problem is that the variance increase in density appears when the system is very close to the transition
- 8 (Corrado et al., 2014), this practically does not constitute an early warning indicator. An alternative is
- <sub>9</sub> to analyze the variance of the fluctuations of the largest patch  $\Delta S_{max}$ : a maximum is attained at the
- critical point but a significant increase occurs well before the system reaches the threshold (Corrado et al.,
- 11 2014). In addition, before the critical fragmentation the skewness of the distribution of  $\Delta S_{max}$  should be
- 12 negative, implying that fluctuations below the average are more frequent. We characterized the increase in
- the variance using quantile regression: if variance is increasing the slopes of upper or/and lower quartiles
- should be positive or negative.
- 15 All statistical analyses were performed using the GNU R version 3.3.0 (R Core Team, 2015), using code
- 16 provided by Cosma R. Shalizi for power law with exponential cutoff model and the poweRlaw package
- (Gillespie, 2015) for fitting the other distributions. Image processing was done in MATLAB (Version??).
- 18 The complete source code for the processing of images and analysis, the patch size data and the figures with
- each fit (year by region by subregion) is available at figshare http://dx.doi.org/10.6084/m9.figshare.XXXXXX.

# 20 Results

- The power law distribution was selected as the best model in most of the cases (Supplementary Figure S7).
- 22 In a small number of cases (4 of 300) the power law with exponential cutoff was selected, but the value of the
- parameter  $\alpha$  was similar by  $\pm 0.02$  to the pure power law. Additionally the patch size where the exponential
- tail begins is very large, thus we used the power law parameters for this cases (See Supplementary data,
- 25 region EUAS3). In finite-size systems the favored model should be the power law with exponential cutoff,
- because the power-law tails are truncated to the size of the system (Stauffer & Aharony, 1994). Here the
- 27 regions are so large that the cutoff is practically not observed.
- 28 There is only one region that does not follow a power law: Eurasia mainland, which follows a log-normal
- 29 distribution. The log-normal and power law are both heavy tailed distributions and difficult to distinguish,
- <sub>30</sub> but in this case Akaike weights have very high values for log-normal (near 1), meaning that this is the

- only possible model. In addition, the goodness of fit tests clearly rejected the power law model in all cases
- <sup>2</sup> for this region (Supplementary table S1, region EUAS1). In general the goodness of fit test rejected the
- power law model in fewer than 10% of cases. In large forest areas like Africa mainland (AF1) or South
- 4 America tropical-subtropical (SAST1), larger deviations are expected and the rejections rates are higher so
- 5 the proportion is 30% or less (Supplementary Table S1).
- Taking into account the bootstrapped confidence intervals of each power law exponent  $(\alpha)$  and the temporal
- autocorrelation, there is no significant differences between  $\alpha$  for the regions with the biggest (greater than
- <sub>8</sub> 10<sup>7</sup> km<sup>2</sup>) forest areas (Figure 1 and Supplementary figure S8). There is also no difference between these
- 9 regions and smaller ones (Supplementary Table S2 & S3), and all the slopes of  $\alpha$  are not different from 0
- <sub>10</sub> (Supplementary Table S3). This implies a global average  $\alpha = 1.908$  with a bootstrapped 95% confidence
- interval between 1.898 and 1.920.
- The proportion of the largest patch relative to total forest area  $RS_{max}$  for regions with more than  $10^7 \text{ km}^2$
- 13 of forest is shown in figure 2. South America tropical and subtropical (SAST1) and North America (NA1)
- have a higher  $RS_{max}$  of more than 60%, and other big regions 40% or less. For regions with less total
- 15 forest area (Supplementary figure S9 & Table 1), the United Kingdom (EUAS3) has a very low proportion
- 16 near 1%, while other regions such as New Guinea (OC2) and Malaysia/Kalimantan (OC3) have a very high
- proportion. SEAS2 (Philippines) is very interesting because it seems to be under 30% until the year 2005,
- fluctuates in the range 30-60%, and then stays over 60% (supplementary figure S9). This seems to be an
- example of a transition from an unconnected to connected state, and deserves further in-depth study.
- We analyzed the distributions of fluctuations of the largest patch relative to total forest area  $\Delta RS_{max}$  and
- fluctuations of the largest patch  $\Delta S_{max}$ . The model selection for  $\Delta S_{max}$  results in power law distributions for
- 22 all regions (Supplementary table S6). The goodness of fit test (GOF) did not reject power laws in any case,
- but nor could it reject the other models except in a few cases, because of the small number of observations.
- We found some differences in  $\Delta RS_{max}$ : Eurasia mainland (EUAS1), New Guinea (OC2), Malaysia (OC3),
- 25 New Zealand (OC6, OC8) and Java (OC7) all follow an exponential distribution (Supplementary Table S7).
- <sup>26</sup> We only considered fluctuations to follow a power law when this distribution was selected for both absolute
- 27 and relative fluctuations.
- 28 The animations of the two largest patches (Supplementary data) qualitatively shows the nature of fluctuations
- 29 and if the state of the forest is connected or not. If the largest patch is always the same patch over time,
- the forest is probably not fragmented; this happens for regions with  $RS_{max}$  of more than 40% such as AF2
- Madagascar), NA1 (North America), and OC3 (Malaysia). Moreover these regions maintain always the same
- largest patch, which could represent a core area for conservation. [TODO: I'd move previous sentence to

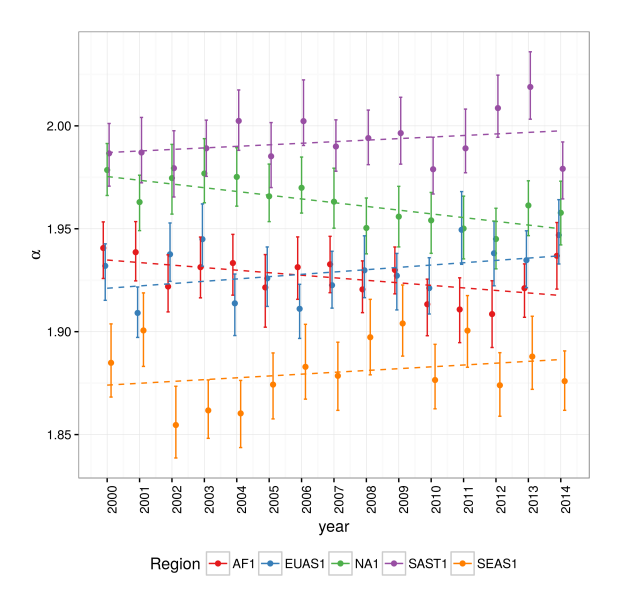


Figure 1: Power law exponents ( $\alpha$ ) of forest patch distributions for regions with total forest area >  $10^7$  km². Dashed horizontal lines are the fitted generalized least squares linear model, with 95% confidence interval error bars estimated by bootstrap resampling. The regions are AF1: Africa mainland, EUAS1: Eurasia mainland, NA1: North America mainland, SAST1: South America subtropical and tropical, SEAS1: Southeast Asia mainland. For EUAS1 the best model is log-normal but the exponents are included here for comparison.

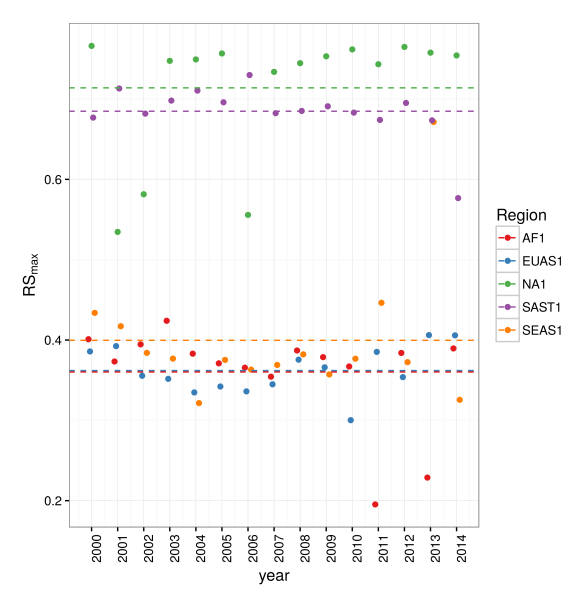


Figure 2: Largest patch proportion relative to total forest area  $RS_{max}$ , for regions with total forest area  $> 10^7~\rm km^2$ . Dashed lines are averages across time. The regions are AF1: Africa mainland, EUAS1: Eurasia mainland, NA1: North America mainland, SAST1: South America tropical and subtropical, SEAS1: Southeast Asia mainland.

- discussion In the regions with  $RS_{max}$  between 40% and 30% the identity of the largest patch changes could
- 2 change or stay the same in time. For OC7 (Java) the largest patch changes and for AF1 (Africa mainland) it
- stays the same, as this is the zone where the critical transition occurs. Only for EUAS1 (Eurasia mainland)
- 4 did we observe that the two patches largest patches are always the same, implying that this region is probably
- 5 composed of two independent domains and should be divided in further studies. For regions with  $RS_{max}$  less
- than 30%, the largest patch always changes, e.g. in SAST2 (Cuba) and EUAS3 (United Kingdom), reflecting
- their fragmented state. In the case of SEAS2 (Philippines) a transition is observed, with the identity of the
- 8 largest patch first variable, and then constant after 2010.
- <sub>9</sub> The results of quantile regressions are identical for  $\Delta RS_{max}$  and  $\Delta S_{max}$  (supplementary table S4). Among
- the biggest regions, Africa (AF1) has upper and lower quantiles with significantly negative slopes, but the
- 11 lower quantile slope is lower, implying that negative fluctuations and variance are increasing (Figure 3).
- Eurasia mainland (EUAS1) has only the upper quantile significant with a positive slope, suggesting an
- 13 increase in the variance. North America mainland (NA1) exhibits a significant lower quantile with positive
- 14 slope, implying decreasing variance. Finally, for mainland Australia, all quantiles are significant, and the
- 15 slope of the lower quantiles is greater than the upper ones, as variance is decreasing (Supplementary figure
- 16 S10). These results are summarized in Table 1.
- 17 The conditions that indicate that a region is near a critical fragmentation threshold are that patch size
- distributions follow a power law; temporal  $\Delta RS_{max}$  fluctuations follow a power law; variance of  $\Delta RS_{max}$  is
- increasing in time; and skewness is negative. All these conditions were true only for Africa mainland (AF1)
- 20 and South America tropical & subtropical (SAST1).

Table 1: Regions, subregions and indicators of closeness to a critical fragmentation threshold. Where,  $RS_{max}$  is the largest patch divided by the total forest area,  $\Delta RS_{max}$  are the fluctuations of  $RS_{max}$  around the mean, skewness was calculated for  $RS_{max}$  and the increase or decrease in the variance was estimated using quantile regressions, NS means the results were non-significant.

Region			Average	Patch Size	$\Delta RS_{max}$	Skewness	Variance
		Description	$RS_{max}$	Distrib	Distrib.		
AF	1	Africa mainland	0.36	Power	Power	-1.8630	Increase
	2	Madagascar	0.65	Power	Power	-0.2478	NS
EUAS	1	Eurasia, Mainland	0.36	LogNormal	$\operatorname{Exp}$	0.4016	Increase

Region			Average	Patch Size	$\Delta RS_{max}$	Skewness	Variance
		Description	$RS_{max}$	Distrib	Distrib.		
	2	Japan	0.94	Power	Power	0.0255	NS
	3	United Kingdom	0.07	Power	Power	2.1330	NS
NA	1	North America	0.71	Power	Power	-1.5690	Decrease
	5	Newfoundland	0.87	Power	Power	-0.7411	NS
OC	1	Australia, Mainland	0.28	Power	Power	0.0685	Decrease
	2	New Guinea	0.97	Power	Exp	0.1321	Decrease
	3	Malaysia/Kalimantan	0.97	Power	Exp	-0.9633	NS
	4	Sumatra	0.92	Power	Power	1.3150	Increase
	5	Sulawesi	0.87	Power	Power	-0.3863	NS
	6	New Zealand south island	0.76	Power	Exp	-0.6683	NS
	7	Java	0.38	Power	Exp	-0.1948	NS
	8	New Zealand north island	0.75	Power	Exp	0.2940	NS
SAST	1	South America, Tropical and subtropical forest up	0.68	Power	Power	-2.7760	Increase
	2	to Mexico Cuba	0.21	Power	Power	0.2751	NS
SAT	2 1	South America,	0.21 0.60	Power	Power	-1.5070	NS Decrease
NAI	1	Temperate forest	0.00	1 Owel	1 OMGI	-1.5070	Decrease
SEAS	1	Southeast Asia, Mainland	0.40	Power	Power	3.0030	NS

Region		Average	Patch Size	$\Delta RS_{max}$	Skewnes	s Variance
	Description	$RS_{max}$	Distrib	Distrib.		
2	Philippines	0.54	Power	Power	0.3113	Increase

# Discussion

- <sup>2</sup> We found that the forest patch distribution of most regions of the world followed power laws spanning
- seven orders of magnitude. These include tropical rainforest, boreal and temperate forest. Power laws have
- 4 previously been found for several kinds of vegetation, but never at global scales as in this study. Interestingly,
- 5 Eurasia does not follow a power law, but it is a geographically extended region, consisting of different domains
- 6 (as we observed in the Supplementary Information animations); the union of two independent power law
- <sup>7</sup> distributions produces a lognormal distribution (Rooij et al., 2013). Future studies should split this region
- 8 into two or more new regions, and test if the underlying distributions are power laws.
- 9 Several mechanisms have been proposed for power laws: the first is related self organized criticality (SOC),
- when the system is driven by its internal dynamics to the critical state; this has been suggested mainly for
- fire-driven forests (Zinck & Grimm, 2009; ???). Real ecosystems do not seem to meet the requirements
- of SOC dynamics: their dynamics are influenced by external forces, and interactions are non-homogeneous
- (i.e. vary from place to place) (???). Moreover, SOC for forest-fire dynamics requires the memory effect: fire
- scars in a site should accumulate and interfere with the propagation of a new fire. Puevo et al. (2010) do
- not find any such effect, and suggested that other mechanisms produce the power laws patterns observed.
- 16 Other studies also found that SOC models did not reproduce the patterns of observed fires. The SOC
- models produce power law exponents  $\alpha = 1.16$  that is much lower than the confidence interval we found for
- $\alpha$  (1.898–1.920). Thus a mechanism which resembled SOC i.e. with a double separation of scales, is not a
- 19 plausible explanation for the global forest dynamics.
- The mechanism suggested by Pueyo et al. (2010) was isotropic percolation, when a system is near the critical
- 21 point power law structures arise. This is equivalent to the random forest model that we explained previously,
- 22 and requires the tuning of an external environmental condition to carry the system to this point. We did
- 23 not expect forest growth to be a random process at local scales, but it is possible that combinations of
- <sup>24</sup> factors cancel out to produce seemingly random forest dynamics at large scales. If this is the case the power
- law exponent should be theoretically near  $\alpha = 2.05495$ , but this value is outside the confidence interval we

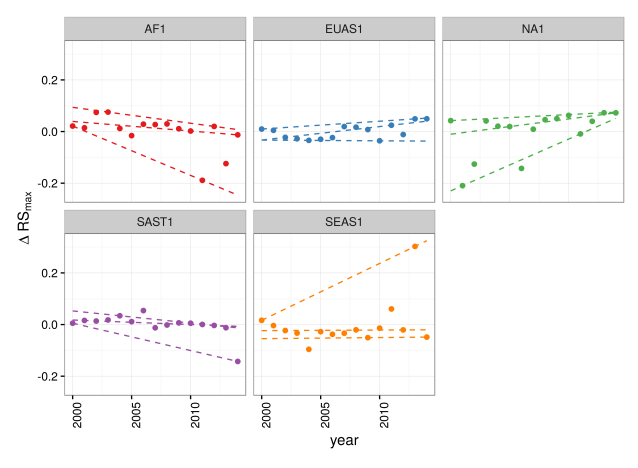


Figure 3: Largest patch fluctuations for regions with total forest area  $> 10^7 \mathrm{km}^2$ . The patch sizes are relative to the total forest area of the same year. Dashed lines are 90%, 50% and 10% quantile regressions, to show if fluctuations were increasing. The regions are AF1: Africa mainland, EUAS1: Eurasia mainland, NA1: North America mainland, SAST1: South America tropical and subtropical, SEAS1: Southeast Asia mainland.

observed, and thus other explanations are needed.

The third mechanism suggested as the cause of pervasive power laws in patch size distribution was facilitation (Manor & Shnerb, 2008; Irvine et al., 2016): a patch surrounded by forest will have a smaller probability of been deforested or degraded than an isolated patch. We hypothesize that models that include facilitation could explain the patterns observed here. The model of Scanlon et al. (2007) represented the dynamics of savanna trees includes a global constraint (mean rainfall), a local facilitation mechanism, and two states (tree/non-tree), producing power laws across a rainfall gradient without the need for tuning an external parameter. The results for this model showed an  $\alpha = 1.34$  which is also different that our results. Another model but with three states (tree/non-tree/degraded), including local facilitation and grazing, was also used to obtain power laws patch distributions without external tuning, and exhibited deviations from power laws 10 at high grazing pressures (Kéfi et al., 2007b). The values of the power law exponent  $\alpha$  obtained for this 11 model are dependent on the intensity of facilitation, when facilitation is more intense the exponent is higher 12 but the maximal values they obtained were still lower than the ones we observed. The interesting point is 13 that the value of the exponent is dependent on the parameters, thus the observed  $\alpha$  might be obtained with 14 some parameter combination. Kefi et al. (2007b) proposed that a deviation in power law behavior with 15 the form of an exponential decay or cut-off could be a signal of a critical transition. At the continental scales studied here, we did not observe exponential cut-offs, but we do observed other signals of a transition, 17 this confirms that different mechanisms could produce different spatial patterns near the transition and that early warnings based only on spatial patterns are not universal for all the systems (Weerman et al., 2012; Kéfi et al., 2014).

It has been suggested that a combination of spatial and temporal indicators could more reliably detect critical transitions (Kéfi et al., 2014). In this study, we combined five criteria to detect the closeness to a fragmentation threshold. Two of them were spatial: the forest patch size distribution, and the proportion of the largest patch relative to total forest area  $(RS_{max})$ . The other three were the distribution of temporal fluctuations in the largest patch size, the trend in the variance, and the skewness of the fluctuations. Each one of these is not a strong individual predictor, but their combination gives us an increased degree of confidence about the closeness of a critical transition.

We found that only the tropical forest of Africa and South America met the five criteria and thus seem to
be near a critical fragmentation threshold. This means that the combined influence of human pressures and
climate forcings might trigger all the undesired effects of fragmentation in these extended areas. A small but
continuous increase in forest loss could produce a biodiversity collapse (Solé et al., 2004), and this threshold
effect has been observed in different kind of models, experimental microcosms (???), field studies (Pardini et

al., 2010; Martensen et al., 2012) and food webs (Martinson et al., 2012). Of these two areas, Africa is more affected, because the proportion of the largest patch relative to total forest area  $(RS_{max})$  is near 30%, this could indicate that the transition is already started. Moreover, this region was estimated to be potentially bistable, with the possibility to completely transform into a savanna (Staver et al., 2011). The region of South America tropical forest has a  $RS_{max}$  of more than 60% suggesting that the fragmentation transition is approaching but not yet started.

When habitat is reduced species population will decline proportionally; this can happens even when the habitat fragments retain connectivity. As habitat reduction continues, the critical threshold is approached and connectivity will have large fluctuations (Brook et al., 2013). This could trigger several synergistic effects: populations fluctuations and the possibility of extinctions will rise, increasing patch isolation and 10 decreasing connectivity (Brook et al., 2013). This positive feedback mechanism will be enhanced when the 11 fragmentation threshold is reached, resulting in the loss of most habitat specialist species at a landscape scale 12 (Pardini et al., 2010). Some authors argue that as species have heterogeneous responses to habitat loss and 13 fragmentation, and that as biotic dispersal is limited, the importance of thresholds is restricted to local scales 14 or even that its existence is questionable (Brook et al., 2013). Fragmentation is by definition a local process 15 that at some point produces emergent phenomena over the entire landscape, even if the area considered is infinite (Oborny et al., 2005). In addition, after a region's fragmentation threshold connectivity decreases, 17 there is still a large and internally well connected patch that can maintain sensitive species (Martensen etal., 2012). What is the time needed for these large patches to become fragmented, and pose a real danger of 19 extinction to a myriad of sensitive species? If the forest is already in a fragmented state, a second critical transition could happen, this is called the desertification transition (Corrado et al., 2014). Considering the actual trends of habitat loss, and studying the dynamics of non-forest patches—instead of the forest patches as we did here—the risk of this kind of transition could be estimated. To improve the estimation of non-forest patches other data set as the MODIS cropland probability should be incorporated (Sexton et al., 2015). The simple models proposed previously could also be used to estimate if these thresholds are likely to be continuous and reversible or discontinuous and irreversible, and the degree of protection (e.g. using the set-asides strategy (Banks-Leite et al., 2014)) than would be necessary to stop this trend.

The effectiveness of landscape management is related to the degree of fragmentation, and the criteria to direct the efforts could be focusing on the regions that are near a transition (Oborny et al., 2007). Regions that are in an unconnected state require large efforts to recover a connected state, but regions that are near a transition could be easily pushed to a connected state; feedbacks due to facilitation mechanisms might help to maintain this state.

- 1 It will be possible that these thresholds will sum up and produce a cascading effect to reach a planetary
- 2 tipping point, crossing the fragmentation critical point could negative effects on biodiversity and ecosystem
- services (Haddad et al., 2015), but it also could produce feedback loops at different levels of the biological
- 4 hierarchy. This means that a critical transition produced at a continental scale could have effects at the
- <sup>5</sup> level of communities, food webs, populations, phenotypes and genotypes (Barnosky et al., 2012). All these
- 6 effects interact with climate change, thus there is a potential production of cascading effects that will lead
- 7 to a global collapse. The risk of such event will be higher if the dynamics of separate continental regions
- are coupled (Lenton & Williams, 2013). Using the time series obtained in this work the coupling of the
- 9 continental could be further investigated. It was proposed that to predict the possibility of a global scale
- shift the different small scale ecosystems should be studied in parallel, as forest comprises a major proportion
- of such ecosystem we think that the transitions of forest could drive all these smaller scale ones and thus
- 12 it could be used as a proxy for all the underling changes and a successful predictor of a planetary tipping
- 13 point.

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