When Shape Matters: Using a Simple Mathematical Model to Estimate Critical Area Sizes in Conservation

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When shape matters: using a simple mathematical model to estimate critical area sizes in conservation

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Abstract
In the analysis of anthropogenic impact on the environment arises the question of whether the shapes of preserved habitat fragments play an important role in the conservation of wild species. In this work we use a very simple mathematical model based on a reaction-diffusion equation to analyze the effects of geometric shape and area on the permanence of populations in habitat fragments. Our results indicate that a dimensionless quantity calculated from a combination of biological variables is the main component that determines if the species survives in the preserved fragment and whether its geometric shape is important. We provide a methodology to calculate critical area sizes for which population size is most affected by fragment shape. The calculation is based on four quantitative variables: maximum per capita reproduction rate, per capita mortality rate outside the fragment, carrying capacity in the conserved environment and mobility in the disturbed environment. The methodology is illustrated by a preliminary study, in which the model is used to estimate threshold area sizes for habitat fragments for the threatened species *Sapajus xanthosternos.*

KEYWORDS:
area, conservation, fragment, geometric shape, mathematical model, threshold

1 | INTRODUCTION

Landscape ecology studies traditionally use landscape pattern indices, a.k.a landscape metrics, to predict ecological responses, they are mostly associated to patch size, shape and habitat amount and aggregation. Detailed ecological research on patch-scale alterations on biodiversity were never as relevant, considering that half of the forests of the planet are less than 500 m from the forest edge, and mostly are patches smaller than 10 ha. Moreover, habitat fragmentation, isolation and creation of edge environments initiate long-term responses of organisms and ecosystems processes that percolate through the landscape.
The edge of forests may be a barrier to animal movements\textsuperscript{3,4}, thus, the area and shape of forest patches are appropriate metrics to assess the effects of spatiotemporal changes in the landscape configuration over biodiversity measures, such as, species richness, community structure and organisms’ abundance\textsuperscript{5,6}. These alterations in landscape patterns are also used as predictors of ecological processes, namely the probability of population extinctions and migrations\textsuperscript{7}.

Recently, a vivid debate on the importance of considering several spatial scales to assess the changes in biodiversity has arisen and most studies indicate that scale selection is species-sensitive\textsuperscript{8}. It is well-known that anthropogenic and natural landscape alterations disproportionately impact forest-core species, which are almost four times more prone to extinction than edge-tolerant and habitat generalist species\textsuperscript{9}. Accordingly, it has been shown that forest-core species are vulnerable to hunting and predation when moving through non-forest habitats, and that abundance of forest-core animals is consistently larger at about 400 m away from the edge, confirming that edge effects operate at a small spatial scale\textsuperscript{9}.

The impacts of edge effects on populations of forest-dependent species are influenced by the ratio of forest core in relation to forest edge. Thus, edge effects are strongly related to the shape of forest fragments, but they are less intensive in the core of larger forests fragments\textsuperscript{10,11}. Forest-core species are more strongly affected in small forest fragments with convoluted shapes\textsuperscript{12,13}. The population abundance of these forest species tend to decline under edge effects and ultimately increase the risk of local extinction\textsuperscript{9}.

Traditional approaches in landscape ecology use diversity measures, such as species richness and equitability, to explore how changes in habitat structure and landscape configuration may lead to species loss, alterations in the animal community composition, and biodiversity erosion\textsuperscript{13}. Fewer landscape ecology studies address the influence of those landscape modifications on population dynamics. Here we use population dynamics models (partial differential equations) to understand the population vulnerabilities associated to changes in the spatial configuration of forest patches. This approach allows us to study the link between the spatial configuration of the habitat and the population dynamics\textsuperscript{14}.

The paper is subdivided in three other essential sections besides this introduction. In section 2 we present the model, the biological hypotheses and parameters, and quickly review some metrics of fragment shape. In section 3 we deduce general ecological results from the model and illustrate its application in a particular preliminary study directed to a particular species (Sapajus xanthosternos). Finally, in section 4 we discuss some of the general and particular results, delineating also directions for future research.
2 | MATERIALS AND METHODS

2.1 | The model

The main focus of this paper is to deduce conditions for which the shape of a habitat fragment does have a significant impact on the chances of conservation of a certain species that occupies it. Since this question is quite general, we are immediately caught in a tension, trying to avoid two opposing dangers: oversimplification of biological traits and narrowness of results due to an excessive number of hypotheses.

For the particular model used in this work, each species may be represented by a combination of five basic biological parameters: the time it would take a population to be doubled under the most favorable conditions ($T_r$ in time units), the carrying capacity in an undisturbed/preserved environment ($K$ in number individuals per area unit), the time that it would be necessary for the population to be halved in a disturbed/hostile environment ($T_m$ in time units), a dispersion coefficient ($D$ units of area$^{-1}$ per unit of time) and a non-dimensional coefficient of mobility ($\eta$) that identifies if individuals tend to disperse faster or slower in the disturbed environment when compared to its movement in the preserved areas.

Of course, under many aspects, this is an oversimplification of any biological species, but here we argue for this approach, underlying the following points:

1. If our objective is to gain insight, from a broad perspective, on the effects of shape and area size on the survival of species, we cannot rely on detailed dynamics which are species-dependent.

2. This generality allows for an easier application in specific cases, requiring just a few biological parameters to adequately apply the methodology outlined by the model. Detailed models often require the estimation of a much higher number of parameters, many of which may demand intricate experiments to be successfully determined.

3. As long as the model and methodology is not considered as an absolute tool and is used only as a guideline to provide rough estimates of threshold area sizes, there should be no harm in approximating the complex population dynamics of the real populations for the simplified one presented here.

It is worth to mention that the model is effective in illustrating the importance of the relations between certain scales that are connected with mobility, reproduction rates and area of the fragment. In this sense, the model should provide some insight from the perspective of ecological theory and conservation, going beyond the simple applications of statistical methods of analysis, which, of course, have their own importance and scope.
2.1.1 | The biological hypotheses

In the process of mathematical modeling it is important to clarify the biological hypotheses assumed so that we know exactly what is included and what is not in the dynamics presented by the model. The main assumptions considered in our model are enumerated below:

1. The model describes the dynamics of one particular species at a time. No interspecific relations are modeled.

2. The region to be analyzed can be divided into two clearly different landscapes. One represents a preserved and favorable environment to the species while the other corresponds to a hostile ambient, where the population could be sustained only for an definite amount of time (extinction is unavoidable in the hostile ambient).

3. Within the preserved environment the species grows logistically up to a carrying capacity.

4. In the hostile ambient the population decays exponentially.

5. Individuals are forced to move around in the environment in both types of landscapes, either due to overpopulation forces, foraging or other species-dependent factors. The particular details are not taken into account in the model, instead, those forces are represented by two parameters, one for movement in the preserved region and other for the hostile one.

The first hypothesis simply means that we are not explicitly describing complex ecological relations between species. These relations, of course, are very important in the real biological system and may be the decisive factor between the survival or the extinction of a particular population. In our approach, those relations are “embedded” in the form of the division of the region in a preserved or hostile environment. In the preserved region, all the ecological relations necessary for the survival of the population are present while in the hostile region those conditions are absent. In this way, the analysis can be focused in just one species.

The second hypotheses is related to a very frequent occurrence in the study of anthropogenic impact on wild populations. In many instances is is possible to observe a “natural” landscape, where wild populations reproduce and a “disturbed” one, that has been modified to serve another purpose, for instance, raising of cattle or agriculture\textsuperscript{15}. Although the biological hypotheses do not explicitly account for a continuous transition between those types of environment, in the sense that each location is either classified as “preserved” or “disturbed”, the mathematical formulation does provide a certain smoothness in the transition from one scenario to the other. In this type of model, populations that live closer to the transition frontier tend to reproduce, on average, slower than those in more “interior” areas.

Referring to the third hypothesis, although some species tend to conform to the specific form of Logistic growth\textsuperscript{16}, in our model, this particular form of dynamics is used just to represent the factual observation that every population has its growth limited to the natural resources available. In this case, those are supposed to be directly proportional to the size of the preserved area. Similar models with analogous limited-growth functions (a generalized Logistic model was used\textsuperscript{17}) were tested and very
close results were obtained so we have some evidence that our analysis in independent of the particular choice of a Logistic growth, but rather that the fundamental fact is the limited carrying capacity of the environment.

The fourth hypothesis accounts for mortality in the hostile environment. It can be proved\textsuperscript{19} that if each individual has a fixed probability of dying per time unit then the dynamics of the whole population can be approximated by an exponential decay. The important factor here is that the species is tending to extinction in the hostile environment.

The last hypothesis involves the dispersion of the individuals. Animal movement is an extremely complex topic and many factors do contribute to the observed movement pattern at the population level. When enough data is available, precise mathematical models can be fitted to describe complex inter and intra-specific interactions. In particular, Moorcroft et al.\textsuperscript{19} have successfully used a reaction-diffusion equation to model wolf pack territory formation in Yellowstone Park, being able to closely fit the model to empirical data on population distributions. Due to its generality, the model we propose is much less detailed and we do not expect such model to precisely describe the microscopic individual movement dynamics. Instead, we rely on a macroscopic parameter, the diffusion (or dispersion) coefficient, to represent a general tendency of dispersion, whatever is the particular biological mechanics behind it. Particular population estimates in well-known regions may then be used to fit the dispersion coefficient, avoiding the need of intricate empiric research on the movement pattern of the modeled species.(see Section for further details).

Finally, there is the observation that individuals may move at different speeds depending if they are in a preserved or hostile area. While the precise estimation of such difference may be subject to technical difficulties, we find this assumption necessary, given the generality of the model, which could be applied to many different species and terrains. In Section 3.1.2 we show that parameter \( \eta \), which is related to this difference in movement, is a parameter that does not have a very strong impact on the equilibrium population. Since the results are fairly robust to this factor, the model may still be applicable even if it is not possible to estimate precisely the difference in mobility between hostile and preserved areas.

2.1.2 The variables and the simulation space

The dynamics proposed occurs in a three-dimensional space \((\mathbb{R}^3)\), having two Cartesian coordinates, \((x, y) \in \mathbb{R}^2\), for the spatial distribution and another one for time, \( t \in \mathbb{R} \). The population is then described by a density function \( u(x, y, t) \), so if \( \Omega \subset \mathbb{R}^2 \) represents a fragment, \( P(\Omega) = \int \int _\Omega u(x, y, t) \, dx \, dy \) is the total population in it the fragment at time instant \( t \). The dimensions of \( x \) and \( y \) are compatible with length (kilometer, mile, meter), while \( u \) may be expressed in individuals per area using convenient units of population and area compatible with \( x \) and \( y \) (thousands, hundreds for population; square meters, miles or kilometers for area). The time variable may be measured in years, months or other useful scale.

The domain of simulation was chosen as a rectangular set \( L = [a, b] \times [c, d] \). Each point inside the domain was classified as being either favorable to the species (preserved habitat) or hostile to it (impacted environment). We defined \( \Omega \subset L \) as the set that
FIGURE 1 The image used in this example is merely illustrative. a) Rectangular satellite image of a preserved environment and surrounded by a hostile environment. The geographical coordinates (12°0′49.60″ S 55°24′29.35″ W) belong to the highlighted region. Source: Google Earth. b) The domain of the model is a rectangular box \( L \) that includes a region of preserved environment represented by \( \Omega \) and the complementary region \( L - \Omega \) that stands for the hostile environment. The population is described by a density function \( u \equiv u(x, y, t) \) that depends on the position in a two-dimensional space, \((x, y)\) and a time variable \( t \). The boundary of the domain is denoted by \( \partial L \) and is composed only of the four sides of the rectangle \( L \).

represents the fragment of preserved habitat while \( L - \Omega \) stands for the hostile environment. Figure 1 illustrates an example of this type of separation.

### 2.1.3 The reaction-diffusion equation

Reaction–diffusion models are applied to the study of a wide variety of natural systems. Pattern formation in animal coats, spatial distribution of slime molds and formation of galaxies, ecological invasion by alien species, chemical signalization in insects and even evolutionary phenomena are some examples of applications. For an excellent review of its use in Mathematical Ecology, we indicate the book by Okubo.

To model the population dynamics in \( L \), we propose the following equations:

\[
\frac{\partial u}{\partial t} = \begin{cases} 
\text{div}(Du) + ru(1 - u/K) & \text{if } (x, y) \in \Omega \\
\text{div}(DHu) - \mu u & \text{if } (x, y) \in L - \Omega 
\end{cases}
\] (1)

where \( D \) and \( DH \) are the dispersion coefficients in the preserved habitat and the hostile environment, respectively, \( r \) is the per capita reproduction rate in the habitat, \( \mu \) is the per capita mortality rate in the hostile environment and \( K \) is the carrying capacity in the preserved habitat.
The dynamics is separated in two regions, \( \Omega \) and \( L - \Omega \). The first term in both equations represents the mobility of the population in each environment, so the ratio \( \eta = D_H / D \) indicates where the movement is faster, if \( \eta > 1 \) individuals move faster in the hostile environment while \( \eta < 1 \) means the opposite.

The use of a diffusion equation to incorporate the characteristics of complex movement patterns may be criticized as an oversimplification. While it is true that this type of equation may be deduced in many different ways\(^2^6\), a common misconception is to think that it can only represent Brownian motion, that is, the movement displayed by particles randomly colliding and moving in every direction. One way to interpret differently this equation is to suppose that, in microscopic scale, individuals carefully avoid overcrowded regions, moving in the direction of less populated areas in a velocity that is proportional to the stimulus given by the Weber-Fechner law\(^2^7\). Under such assumptions, the resulting macroscopic dynamics is exactly a diffusion equation\(^1\).

This can be seen by simply writing the term corresponding to the flux of individuals, \( \vec{z} = D \nabla u \) as \( \vec{\eta} = Du \vec{\nabla} \), where

\[
\vec{\nu} = \frac{\nabla u}{u}; \quad u \neq 0.
\]

The biological interpretation is clear now: individuals move on the opposite direction of the gradient, but with a speed/probability that is proportional to the ratio of the concentration difference perceived and the concentration of individuals. This ratio incorporates the effect described by the Weber-Fechner law, that the perception of changes in the intensity of stimuli is relative to the total present stimulation. Just to give a simple concrete example of this psychophysical law: a match lighted in total darkness provides a much higher stimulus than one lighted in plain daylight. So the model can be interpreted as implying that the population is trying to avoid overcrowded areas, moving in the opposite direction of increasing population density. Although the Weber-Fechner is not an “absolute” natural law and has been subject to criticism and also that alternatives formulations have been proposed for modeling stimuli perception\(^2^8,2^9\), recent results seem to find new evidence supporting it\(^3^0\). In any case, here it serves the purpose of illustrating how the diffusion equation may be re-interpreted as portraying a more complex behavior than just Brownian motion.

As stated in section 2.1.2 the domain of the equation is \( L = [a, b] \times [c, d] \). This choice of domain was made for the following reasons:

1. A rectangular domain facilitates the implementation of numerical methods to solve the differential equation.

2. The domain is chosen in a way so that it included that fragment of interest (\( \Omega \)) and that the boundary conditions do not have a significant impact on the results.

---

\(^1\)This interpretation came to our knowledge through personal communication with Prof. Wilson Castro Ferreira Jr. Up to date we are not aware of any publication that treats this topic in detail.
The boundary conditions for the mathematical problems may be homogeneous Dirichlet conditions:

\[ u|_{\partial L} = 0 \]  

(2)

where \( \partial L \) stands for the boundary of \( L \). Another possibility is to use the Robin homogeneous condition is given by:

\[ \frac{\partial u}{\partial n}|_{\partial L} = k_r u \]  

(3)

where \( \partial u/\partial n \) is the normal derivative (pointing outwards region \( L \)) and \( k_r > 0 \) is a constant that defines the intensity of the flux.

Homogeneous Dirichlet conditions imply that individuals die at the boundaries while Robin homogeneous conditions represents a migration that is proportional to the population density at boundaries. We will not stress too much the role of the boundary conditions because the focus of the model is to analyze the effect of the geometric shape and area of \( \Omega \). The domain \( L \), in this case is just a convenient choice that allows for an easier numerical simulation. We tested our results using both conditions and a variety of values for \( k \) and the results were very similar, independently of the boundary conditions used.

One last technical observation on the model is that there is a discontinuity in the diffusion coefficient. This is not a relevant modeling problem for our particular case since, in this work, the solution of the differential equation is approximated using a discretized version of the domain, where discontinuities are not an issue.

### 2.2 Numerical methods

To simulate the model we first divided the domain \( L \) into the two regions \( \Omega \) and \( L - \Omega \) and, observing that \( D \) and \( D_H \) are constant, obtained the two partial differential equations:

\[ \frac{\partial u}{\partial t} = D \Delta u + ru(1 - u/K), \quad x \in \Omega \]  

(4)

and

\[ \frac{\partial u}{\partial t} = D_H \Delta u - \mu u, \quad x \in L - \Omega \]  

(5)

where \( \Delta \) stands for the two-dimensional Laplacian operator.

To approximate the solution of those equations, we adopted a finite-difference scheme (centered, second order) for the spatial Laplacian operator (\( \Delta \)) and Runge-Kutta methods of combined orders 2 and 3 to solve the resulting system of ordinary differential equations. The application of these methods is considered routine in numerical analysis and can be consulted, for example, in Burden and Faires[31].

To implement such numerical schemes we used two softwares: Matlab R2016b (registered for academic use under license number 1115837) and the open source Scilab (version 6.1.0). Two independent teams worked on the codes, each using a different...
software for comparison of results and avoidance of coding errors. For the Runge-Kutta methods we used the built-in functions provided by both softwares: `ode23` for Matlab and `ode` (with method option "rk") for Scilab.

2.3 Estimates for $r$, $\mu$ and $\eta$

The model uses five mathematical parameters to simulate the dynamics: $r$, $\mu$, $K$, $D$ and $D_H$. To estimate those parameters, we need some biological estimates, as mentioned in Section 2.1:

1. $T_r$: supposing the conditions are very favorable to the population (resource abundance, low competition), $T_r$ is the time the population takes to double its numbers.

2. $T_m$: supposing the whole region was transformed into a hostile environment, $T_m$ is how long would it take for the population to be halved.

3. $\eta$: if we denote by $d$ the average daily displacement of individuals in the preserved habitat and $d_H$ in the hostile environment, then $\eta$ may be interpreted as an estimate of $d_H/d$. Here the time scale of one day was mentioned, but it can be adapted according to the biological convenience.

The exact method for providing those estimates are left to the expert biologists for each case. From these three biological parameters and some population estimates in known areas and a least squares fit it is possible to estimate the remaining parameters ($K$ and $D$). In Section 3.2 we provide a case study as a guideline.

Parameter $r$: From the estimate of $T_r$, and taking into account that, for populations that are small relative to the carrying capacity, the Logistic growth can be approximated by a Malthusian one, we obtain $r$ as:

$$r = \frac{\ln(2)}{T_r}$$

Parameter $\mu$: Since the population decays exponentially on the hostile environment, the instantaneous rate of decay $\mu$ can be directly obtained from the estimate $T_m$:

$$\mu = \frac{\ln(2)}{T_m}$$
2.4 Measures of compactness

One of the central questions approached in this paper is to evaluate how important fragment shape is to the conservation of populations. To conduct a mathematical analysis we need, then, some measure of “compactness” that can be used to represent the characteristics of a certain shape, so that its impact on population can be quantified. A discussion of the roles of such measures and a review of some particular formulas can be consulted in Li et al.[13]. Other authors[13],[34] present also some shape measures commonly used in landscape ecology analysis. Below, we briefly review some of the measures and choose which suits best the scope of this work.

If $\Omega$ is the two-dimensional set that represents the fragment we denote by $\rho(\Omega)$ as the perimeter (we suppose that the boundary of $\Omega$ can be described as a smooth function) and $a(\Omega)$ its area.

Mcgarigal et. al.[34][p.104] present two of the most common measures of compactness, PARA:

$$\text{PARA} = \frac{\rho}{a}$$

and a Shape Index (SHAPE):

$$\text{SHAPE} = \frac{0.25\rho}{\sqrt{a}}.$$  \hspace{1cm} (9)

Another possible measure is the IPQ[35], given by:

$$\text{GE}_{IPQ} = \frac{4\pi a}{\rho^2}.$$  \hspace{1cm} (10)

The measure given by PARA is not convenient because it is scale-dependent, so it was not adopted in this work. SHAPE is not dependent on scales and it is directly related to the IPQ.

In the IPQ measure a circle has the maximum measure of compactness value of 1. The intuition is simple, as the perimeter increases for the same area, the measure decreases, resulting in a value between 0 and 1. The use of the square of the perimeter avoids the impact of scaling factors (units of length), resulting in a reliable measure. To make the results more intuitive, we use the acronym GE to stand for “geometric efficiency” as the measure of compactness.

Here we propose a slight modification of the IPQ, using for the measure of compactness the following formula:

$$\text{GE} = \frac{16a}{\rho^2}.$$  \hspace{1cm} (11)

The change is only a re-scaling of IPQ but, in our simulations, the discretization of the domain causes the regions to be transformed into a union of small squares, which also affects the length of its perimeter and, to a minor degree, its area. Due to this slight deformation of regions, we find it useful to use the square as the reference figure instead of the circle. In this measure, the square has a compactness measure of value 1.
SHAPE is directly related to GE:

\[
SHAPE = \frac{1}{\sqrt{GE}}.
\]  

(12)

so any results obtained in terms of GE can be easily converted to SHAPE and vice-versa.

3 | RESULTS

This section divided in two parts. The first is dedicated to general results and the second to a case study of the species *Sapajus xanthosternos*.

3.1 | General results

For a clearer presentation, we separated the general results into three further sections. First, the combination of parameters that are important to the simulations are identified, then a sensitivity analysis is performed and finally the impact of geometric shape on species population is discussed.

3.1.1 | Identifying the key parameters

Before we addressed the question relative to the impact of geometric shape of a fragment on populations, it was important to analyze the role of the parameters when shape was fixed. We chose a fragment as a circle and used six parameters to simulate the model: \(A_i\) (total area of the fragment), \(r\) (per capita reproduction rate in the preserved habitat), \(K\) (carrying capacity), \(\mu\) (per capita mortality rate in the hostile environment), \(\eta\) (mobility coefficient) and \(D\) (dispersion coefficient).

Each simulation was conducted for a time limit of \(100T_r\), that is, if the population is doubled in a year under the most favorable conditions, then the model was simulated for 100 years. For each simulation, we evaluated the total final population \(P_\Omega(A_i, r, \mu, K, D)\) and then re-scaled it according to the maximum possible population \(P_{\text{MAX}} = A_i \cdot K\), obtaining:

\[
P^* = \frac{P_\Omega}{P_{\text{MAX}}}. \tag{13}
\]

This re-scaling provides an estimate of the border effects on the fragment. For instance, if \(P^* = 0.8\), this implies that the influence of the outside hostile environment causes a decrease of 20% of the maximum potential population in the fragment.

By simulating the model it was possible to obtain the re-scaled population as a function: \(P^*(A_i, r, \mu, K, D)\) and we used this relation to study how the parameters affected the survival of the species. Allowing all parameters to vary in each simulation, the trend was not clear \(P^*\) when we looked individually at parameters \(A_i, r, \mu, \eta\) or \(D\), while parameter \(K\) had no impact at all in
Instead, we found that $P^*$ is dependent on the non-dimensional groupings:

$$\alpha = \sqrt{\frac{A_i r}{D}}$$

$$\lambda = \frac{\mu}{r}$$

(14)

and the non-dimensional parameter $\eta$.

To illustrate such dependency, we chose three different values of $\lambda$ (1/2, 1 and 2) and for each value of $\lambda$, the following distributions for the other parameters was chosen:

$$A_i \sim U(0.25, 3), \ D \sim U(0.001, 0.25)$$

$$r \sim U(0.25, 1.4), \ \eta \sim U(0.5, 2)$$

(15)

where $U(a, b), a, b \in \mathbb{R}$ stand for a uniform distribution between $a$ and $b$. Note that once $r$ and $D$ are established, $\mu$ and $D_H$ are automatically determined by the relations $\mu = \lambda r$ and $D_H = \eta D$.

These intervals were conveniently chosen to illustrate the relations and similar results are obtained with other intervals that comprise a similar spectrum for $\alpha$. It is worth to stress that since $\alpha$, $\lambda$ and $\eta$ are non-dimensional, they are not affected by the choice of units used to describe population density. A total of 250 random combination of parameters was chosen for each fixed value of $\lambda$. In Figure 2 we present illustrations of how $P^*$ behaves in relation to some of the parameters, this illustrates the important role of $\lambda$ and $\alpha$. The results were robust to the choice of both initial and boundary conditions.

Another way to analyze the impact of the parameter is to measure the correlation between the parameters and $P^*$. In Table 1 we present the Spearman coefficient of correlation for the parameters and the variable $P^*$. The results confirm that $\alpha$ is the most important factor in the determination of $P^*$.

**TABLE 1** Spearman correlation coefficient for the simuations of this section. $\alpha$, has the largest value, illustrating its importance to the survival of the species. Since parameters $A_i$, $r$ and $D$ are directly related to $\alpha$, there are also a significant correlation.

| Parameter | $\lambda = 0.1$ | $\lambda = 1$ | $\lambda = 10$ |
|-----------|-----------------|---------------|---------------|
| $\alpha$ (1) | 0.9622          | 0.9748        | 0.8486        |
| $A_i$ (1)  | 0.5264          | 0.5417        | 0.4037        |
| $D$ (1)    | -0.7330         | -0.6660       | -0.6196       |
| $r$ (1)    | 0.3219          | 0.3308        | 0.4039        |
| $\eta$ (2) | 0.2020 (0.1)    | -0.0113 (0.85)| 0.0723 (0.26) |

1 All coefficients have a $p$-value smaller than $10^{-8}$.
2 $p$-value inside parenthesis.
FIGURE 2 Seven hundred and fifty simulations of $P^* = P_{10}/P_{\text{MAX}}$, the re-scaled population inside $\Omega$. Parameters $r$ (per capita reproduction rate), $\eta$ (mobility outside the fragment), $A_t$ (fragment area) and $D$ (dispersion coefficient) were allowed to vary randomly according to the distributions given in Equation 15. a) Results of $P^*$ in relation with $A_t$, the behavior is strongly dependent on the values of the other parameters. b) Results of $P^*$ in relation with $r$. c) Results of $P^*$ in relation with $D$. d) Results of $P^*$ in relation with $\alpha$, for each value of $\lambda$, a clear trend can be observed, indicating that $\alpha$ and $\lambda$ are key parameters to the survival of the species in the fragment.

Another way of deriving the importance of $\alpha$, $\lambda$ and $\eta$ is to write a non-dimensional version of the model. By defining new non-dimensional variables as $w = x/\sqrt{D/r}$, $v = y/\sqrt{D/r}$, $\tau = rt$ and $h = u/K$, model 1 is transformed to:

$$\frac{\partial h}{\partial \tau} = \begin{cases} \frac{\partial^2 h}{\partial w^2} + \frac{\partial^2 h}{\partial v^2} + h(1 - h) & \text{if } (w, v) \in \tilde{\Omega} \\ \eta \left( \frac{\partial^2 h}{\partial w^2} + \frac{\partial^2 h}{\partial v^2} \right) - \lambda h & \text{if } (w, v) \in \tilde{\Omega} \in \tilde{L} - \tilde{\Omega} \end{cases} \quad (16)$$

where $\tilde{L}$ and $\tilde{\Omega}$ are transformations of the domain $L$ and the fragment region $\Omega$ to $(w, v)$ mapping. For the particular case of the simulations in this section, $\Omega$ was taken as a circle of area $A_t$, having a radius of $R = \sqrt{A_t/\pi}$, so the transformed region $\tilde{\Omega}$ is a circle of radius $\tilde{R} = R/\sqrt{D/r} = \sqrt{A_t r/(D \pi)}$. Re-arranging the terms, we obtain:

$$R = \frac{1}{\sqrt{\pi}} \sqrt{\frac{A_t r}{D}} = \frac{\alpha}{\sqrt{\pi}}. \quad (17)$$

Equations 16 and 17 show that the behavior of the non-dimensional model (which is independent of scaling) depends only on the non-dimensional groupings $\eta$, $\lambda$ and $\alpha$ (which shall be called parameters henceforth).
The results of both the simulations and the dimensional analysis show clearly that parameters $\alpha$ and $\lambda$ are essential to the survival of the species. These results are biologically intuitive: $\alpha^2$ is proportional to both $A$, and $r$, meaning larger areas and fast reproduction rates favor the survival of the species (hence the positive correlations between $\alpha$ and $P^*$). Also, $\alpha^2$ inversely proportional to $D$, meaning that large mobility associated with a hostile surrounding environment impacts negatively the population. Parameter $\lambda$ indicates how hostile is the environment outside the environment, being proportional to $\mu$ and inversely proportional to $r$, with more hostile environments leading to lower populations. Parameter $\eta$ had a smaller influence on the final results, as can be seen in the curves in Figure 2-d), for each fixed curve with fixed $\lambda$, $\eta$ was allowed to vary between 0.5 and 2, with little impact on the final re-scaled population $P^*$ which was also confirmed by the analysis of the correlation coefficients in Table 1.

To analyze the relative impact of parameters $\alpha$, $\lambda$ and $\eta$ we performed a sensitivity analysis of model 16 presented in the section below.

### 3.1.2 Sensitivity analysis

In this section we consider $P^*$ as a function of $\alpha$, $\lambda$ and $\eta$. The simulations are analogous to the ones in Section 3.1.1. We chose the shape of the fragment as a circle (with radius given by Equation 17), taking all parameters with the same distribution:

$$\alpha \sim \lambda \sim \eta \sim U(0.05, 20).$$

To evaluate the sensitivity we used the slopes of the direct one-dimensional regressions ($P^* \times \alpha$, $P^* \times \lambda$ and $P^* \times \eta$, from the multilinear regression ($P^* = a\alpha + b\lambda + c\eta + d$), the standardized regression coefficients and the Spearman coefficient. In Table 2 we display the results of the analysis. They clearly confirm the strong influence of parameter $\alpha$, with $\lambda$ and $\eta$ playing a secondary role.

| Parameter | Linear 1 | Multilinear 2 | Standardized 3 | Spearman 4 |
|-----------|----------|---------------|----------------|------------|
| $\alpha$  | 0.0482   | 0.0479        | 0.8414         | 0.8864 (< 10^{-4}) |
| $\lambda$ | -0.0095  | -0.0081       | -0.1365        | -0.2889 (< 10^{-4}) |
| $\eta$    | 0.0091   | 0.0081        | 0.1319         | 0.2195 (< 10^{-3}) |

1 Individual linear regression in each parameter.
2 Simultaneous linear regression in all parameters.
3 Multilinear regression on the standardized variables \( x_i^* = (x_i - \bar{x}) / \sqrt{\text{var}(x)} \).
4 Upper bound for $p$-values inside the parenthesis.

TABLE 2 Results of sensitivity analysis of model 16 for parameters $\alpha$, $\lambda$ and $\eta$. $P^* = P_\Omega / P_{\text{MAX}}$ is the dependent variable. The Spearman coefficient and the slopes of the linear, multilinear and standardized regressions indicate that $\alpha$ plays the leading role in the survival of the species, followed by $\lambda$ and $\eta$ that have a much weaker impact.
3.1.3 | The impact of geometric shape

In the previous sections it was clearly established that $\alpha$ and $\lambda$ are key parameters that define species survival in the model. The simulations indicated that, for each fixed $\lambda$, it is possible to divide the region of variation of $\alpha$ in three quite distinct regions. For very low values of $\alpha$, the population is unable to survive, with $P^* \to 0$. For very high values, boundary effects have little impact on the dynamics, so $P^* \approx 1$ with the population approaching the maximum value possible. Finally, there is an intermediate region where $P^*$ makes a transition between those extreme values, and this is strongly related to the effect of geometric shape.

For very large or very small values of $\alpha$, geometric shape in unimportant, because either the area size is too big or too small in relation to the scales of reproduction and mobility (which is the measure that parameter $\alpha$ incorporates). In the intermediate region the geometric shape plays crucial role in determining how strong is the boundary effect, which may lead either to extinction or preservation.

The central problem analyzed in this paper can now be addressed as a very precise question: for each point in the two-dimensional parameter space $(\alpha, \lambda)$, is it possible to determine if geometric shape affects significantly the population estimates by the model? To answer this question we designed procedures to create random shapes and also to deform a preexisting shape, affecting its measure of compactness (as defined in Section 2.4).

For each point in the $(\alpha, \lambda)$ map we took an initial geometric shape for the region $\Omega$, which we defined as $S_0$. This shape had a total area equal to $\alpha^2$ to correspond to a dimensional simulation of a figure of area $A$, and an initial GE, $G_0 = GE(S_0)$ larger than 0.65, to allow for variation. By applying deformations on this initial shape we generated a set of 30 other shapes, $S_1, \ldots, S_{30}$, with the same area but with compactness measures $G_1, G_2, \ldots, G_{30}$ in the set $(0.05 \cdot G_0, G_0)$. In Appendix A we discuss in more detail the methods used to create deformations of the initial figure as to obtain shapes with smaller compactness values GE. This process is then repeated 10 times, so we obtain a collection of 300 shapes for each $(\alpha, \lambda)$ point.

For this set of shapes we simulated the non-dimensional model for a time horizon of $\tau = 100$ (which corresponds to the time horizon of $T = 100/r$ in the dimensional model) calculating the final re-scaled population $P_i^* = P(S_i)/\alpha^2$ for each shape, where $P(S_i)$ denotes the total population inside $S_i$ after the simulation of the model. We obtained thus numbers between between 0 and 1 that correspond to the fraction of the maximum possible population achieved by that particular combination of parameters and geometric shape.

Given the series of points $(G_i, P_i^*)$ relating figures with different measures of compactness and the final re-scaled population, it is possible then to calculate regressions of $P^*$ by $G$ that estimate the impact of the measure of compactness in the final re-scaled population. Since both variables are non-dimensional and are scaled between 0 and 1, we adopted the slope of the linear regression to estimate the correlation, obtaining the impact of geometric shape as a function of $\alpha$ and $\lambda$, $S \equiv F(\alpha, \lambda)$.

In Figure 3 we present a scheme that illustrates the whole process described in this section.
\[ \alpha = \frac{\Delta r}{D}, \lambda = \frac{\mu}{r}, GE = \frac{1}{\mu r^2}, D_p = \eta D, P^* = P(S_i)/\alpha^2 \]

For each point \((\alpha, \lambda)\) create 10 initial shapes of area \(\alpha^2\). Calculate \(G_0\) the measure of compactness for each initial shape (\(G_0 > 0.65\)). Calculate the slope of linear regression of \((G_j, P_j^*)\) for each point \((\alpha, \lambda)\), obtaining the distribution \(F(\alpha, \lambda)\). In regions where \(F(\alpha, \lambda) \approx 0\), shapes have low impact on population.

For each initial shape, applying deformation techniques, generate 30 other shapes with \(GE\) in \((0.05 G_0, G_0)\) and same area. With the 300 figures obtained, simulate the model for each shape calculating the total re-scaled population \(P^*\), obtaining a collection of points \((G_j, P_j^*)\).

**FIGURE 3** Scheme for the simulations of geometric shape impact on population. Each initial shape is deformed, reducing its compactness, and the impact on population recorded. The distribution \(S(\alpha, \lambda)\), of linear regression slopes, measures the relation between compactness and population levels.

**FIGURE 4** a) Five examples of collections of points \((G_j^*, P_j^*)\), for different values of \(\alpha\) and fixed \(\lambda = 1\). For each collection we calculate the slope of the linear regression, \(F(\alpha, 1)\). b) \(F(\alpha, 1)\) denotes the values of the slope of linear regression of \((G_j^*, P_j^*)\), for \(\lambda = 1\) and \(\alpha \in (2, 30)\).

In Figure 4 we present examples of distributions of \(G_j, P_j^*\) and a curve \(F(\alpha, \lambda)\) for \(\lambda = 1\). The distributions clearly illustrate that, when \(\lambda = 1\), we have the maximum impact of the geometric shape for values of \(\alpha \in (5, 10)\).

In Figure 5 we present the map of slopes obtained from the simulations, \(F(\alpha, \lambda)\). In the map is drawn also two curves \(f(\alpha) = 0.1348e^{0.0074906\alpha}^3\) and \(g(\alpha) = 0.0096e^{0.2494\alpha}\). \(C^*\) is used to estimate area sizes that are greatly impacted by fragment shape. To estimate this curve, for each fixed value of \(\lambda\) we estimated the corresponding \(\alpha\) for which \(F(\lambda, \alpha)\) was at its maximum value, next using a least squares fit, we adjusted a curve to fit the points. In this way \(C^*\) represents critical points where fragment shape has maximum impact and can be used to estimate areas sizes \((A^*)\) for which fragment shape is very important. Finally,
FIGURE 5 Two-dimensional mapping of the impact of geometric shape on the population. $F(\alpha, \lambda)$ denotes the values of the slope of linear regression of $(G_j^*, P_j^*)$. Two curves $\lambda(\alpha)$ are drawn in the mapping: $C^*$ given by the equation: $f(\alpha) = 0.1348 e^{0.007490604 \alpha^3}$ and $C_{0.4}$, with equation $g(\alpha) = 0.0096 e^{0.2494 \alpha}$. Curve $C^*$ is constructed by estimating, for each fixed value of $\lambda$ the corresponding value of $\alpha$ that leads to the highest value for $F(\lambda, \alpha)$ while $C_{0.4}$ is a part of the level set $F(\alpha, \lambda) = 0.4$. $C^*$ represent critical values where fragment shape has maximum impact while $C_{0.4}$ represent a point where the impact of shape is already low.

$C_{0.4}$ represents a part of the level set $F(\alpha, \lambda) = 0.4$. This line allows us to estimate, area sizes ($A_{0.4}$) for which fragment shape is relatively weak ($F = 0.4$ and smaller for larger areas).

If we recall that $\alpha = \sqrt{A/r/D}$ we can establish approximate formulas for the critical area size ($A^*$) and relatively safe area size ($A_{0.4}$):

$$A^* \approx \frac{D}{r} \left( \frac{\ln(\lambda/0.1348648)}{0.007490604} \right)^{2/3}$$

$$A_{0.4} \approx \frac{D}{r} \left( \frac{\ln(\lambda/0.0096)}{0.2494} \right)^2.$$  

(19)  

(20)

Please note that the relatively small effects produced by variations in $\eta$ are not included in these equations.
3.2 | A preliminary case study

In this section we present a case study for the species *Sapajus xanthosternos*. We must stress that this study has the main purpose of illustrating the application of the model and how to interpret its results. For more robust biological conclusions concerning the species, further empirical work in estimating the parameters should be conducted.

It is possible to estimate $r$ the maximum per capita growth rate if the maximum annual growth rate of the species is known. For this particular species, it was estimated that, given the most favorable conditions, the population would have a maximum growth rate of 13% per year, which leads to a correspondent logistic growth rate of

$$ r = \ln 1.13 \approx 0.1222 \text{/year}. \quad (21) $$

Mortality rates in the hostile environment are harder to estimate, and we had to rely on an educated guess by the expert to arrive at a reference value. The information provided is that a group of 64 individuals could not last longer than a year, if exposed continuously to the hostile environment in this time interval. With such information and, considering an exponential decay in the hostile environment it is possible to obtain:

$$ \mu = \ln 64 \approx 4.1588 \text{/year}. \quad (22) $$

Since *Sapajus xanthosternos* is an arboreal species and the hostile environment, in this particular case, is considered to be mostly deforested areas dedicated monoculture, individuals tend to move and disperse faster in the impacted than in the preserved areas. The meaning here is that even though the individuals might move faster in arboreal environment, they tend to move more often in deforested areas, always looking for forest cover, resulting in a higher average speed in the open areas. Considering this information and also by expert advice, we adopted then a value of $\eta = 2$, meaning that, on average, individuals move two times faster outside the preserved areas.

To estimate the remaining parameters, $D$ and $K$, we collected information on population estimates of populations of *Sapajus xanthosternos* living in ecological sanctuaries near hostile environments. Since these population estimates are based on extrapolation from number of individuals sighted, we selected only those studies that were conducted in smaller regions (total area below 1000 ha) so that these estimates can be considered more precise. It is also worth to mention that 1000 ha is the largest forest area size in which *Sapajus xanthosternos* was recorded. After this process we obtained 5 fragments with different area sizes, shapes and populations. Such data can be consulted in Table 3.

To obtain images of the fragment shapes we used the software QGIS, using the CBERS4A downloader plugin, which provides access to the online database of CBERS4A satellite images. The images used correspond to the years in which the population estimates studies were conducted each individual fragment. These are displayed in Figure 6. For each region, the model was
TABLE 3 Data on *Sapajus xanthosternos* populations in preserved fragments, available in Culot *et al.* (2019) and comparison with simulations of the model with optimal values $D^* \approx 9.1746 \times 10^{-4}$ year$^{-1}$·km$^{-2}$, $K^* \approx 26.8033 \times 10^{-4}$ individuals·km$^{-2}$.

| Region name/number | Total Area (ha) | Population estimate | Simulation | Geographical Coordinates |
|--------------------|-----------------|---------------------|------------|--------------------------|
| Água Sumida - 1    | 240             | 52.80               | 51.22      | $(-48.29082778, -22.63317778)$ |
| Monal - 2          | 374             | 164.48              | 85.76      | $(-48.08360278, -22.69596944)$ |
| Pouso Alegre - 3   | 350             | 26.98               | 70.02      | $(-45.96666667, -22.21666667)$ |
| Mata São José - 4  | 230             | 56.35               | 44.68      | $(-47.47743300, -22.35881600)$ |
| Sara - 5           | 501             | 76.85               | 114.69     | $(-48.19536111, -22.66642222)$ |

1 Number of individuals. Left: estimates based on number of sightings. Right: estimates based on simulations of the mathematical model.

2 (Longitude, Latitude)

simulated until the population reached an equilibrium. At the end of the simulation we calculated the total population inside the protected region, as in Figure 6, obtaining thus a vector with five population estimates provided by the mathematical model.

We then used the least squares method to determine the optimal values of $D$ and $K$ that led to the best fit.

![Images of preserved fragments for the *Sapajus xanthosternos* case study.](1)

**FIGURE 6** Images of preserved fragments for the *Sapajus xanthosternos* case study. The blue area represents an ecological reserve with preserved environment where we have estimates for the population, the green area stands for preserved environment but not within the reserve while the white area corresponds to the hostile environment. Name of the reserves: (a) Água Sumida (1) (b) Monal (2) (c) Pouso Alegre (3) (d) Mata São José (4) (e) Sara (5). Simulations of the model were performed in these domains and the results were compared with the field estimates for the populations (see Table 3).
In Table 3 we present the results from simulations using the least squares fit for \( D \) and \( K \). The optimal value for \( D \) was \( D^\ast \approx 9.1746 \times 10^{-4} \text{ year}^{-1} \cdot \text{km}^{-2} \) and for \( K \) was \( K^\ast \approx 26.8033 \times 10^{-4} \text{ individuals} \cdot \text{km}^{-2} \). This particular value of \( K^\ast \) was considered biologically plausible by the expert biologist and within his own estimates.

With the estimated value of \( D^\ast \) now it is possible to use equations [19] and [20] to establish to estimate the most sensible \( (A^\ast) \) and a relatively safe \( (A_{0.4}) \) area sizes for the species \textit{Sapajus xanthosternos}. Results are presented in Table 4.

**TABLE 4** Results for the species \textit{Sapajus xanthosternos}. Areas were estimated using parameters estimated through simulations \((D)\), expert advice \((\lambda)\) and bibliographical research \((r)\) combined with equations [19] and [20]. \( A^\ast \) stands for the size estimate in which the population is most sensible to changes in fragment shape. \( A_{0.4} \) is an estimate for a fragment size in which geometric shape does not have much impact on the population.

| Type of area (critical/safe) | km² | ha   |
|-----------------------------|-----|------|
| \( A^\ast \) (1)           | 0.6132 | 61.32 |
| \( A_{0.4} \) (2)          | 8.062 | 806.20 |

1 Calculated using \( A^\ast \approx \frac{D}{r} \left( \frac{\ln(\lambda/0.1348648)}{0.007490604} \right)^{2/3} \).
2 Calculated using \( A_{0.4} \approx \frac{D}{r} \left( \frac{\ln(\lambda/0.0096)}{0.2494} \right)^{2} \).

4  | DISCUSSION

The simple model developed and the simulations allowed us to identify clearly the most important parameters that determine how strongly populations are affected by fragment shape. The results show clearly that these parameters are associated with mobility of the species, reproduction rate, area of the fragment and mortality rate in the hostile environment. All these parameters are combined in the two non-dimensional groupings \( \alpha \) and \( \lambda \).

The two-dimensional mapping of the behavior of the model in relation to parameters \( \alpha \) and \( \lambda \) allowed us to establish functional relationships (i.e. [19] and [20]) for a \( A^\ast \), area size for which the population is very sensible to geometric shape, and \( A_{0.4} \), an estimate of a sufficient large area where geometric shape does not have much impact on population size. These relations shows is that the area threshold is proportional to the mobility (represented by \( D \)) and inversely proportional to \( r \) (rate of reproduction), reflecting the intuitive ideas that more mobile species need larger areas and those who reproduce faster need smaller ones. Also, the fact that the parameter that represents mortality in the hostile environment (\( \lambda \) or \( \mu \)) is present as an argument of a logarithm function reflects the fact, also shown in the sensitivity analysis, that its impact on the area threshold is smaller.

Another, somewhat obvious, biological result that can be derived from the equations is that species with smaller body sizes should also need smaller areas for their preservation. This is because smaller species tend to reproduce faster[10], meaning that
parameter \( r \) should be greater for these species, leading to smaller critical areas. Finally, more lethal hostile environments also lead to increase in the area needed for preservation of the species. This is demonstrated in the results simply by recalling that \( \lambda = \mu / r \), so any increase in \( \mu \) leads to a correspondent increase in \( \lambda \) and, since both \( A^* \) and \( A_{0,4} \) are increasing functions of \( \lambda \), any increase in mortality rates in the hostile environment also leads to larger areas needed for preservation.

Finally, we highlight a few points where further improvement of the model and results can be made:

1. Incorporate the use of finite element methods for the simulations of the model, since those can incorporate better the details of geometric shape of fragments.
2. Apply the model to a number of different species and fragments.
3. Improve the quality of parameter estimation, either by using larger data bases with information on fragment populations or better/more numerous estimates of reproduction/mortality rates and mobility.

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**Consent to participate**

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**Consent for publication**

The article does not use any third-part sensitive information that needs consent to be published. The authors consent to the publishing of the manuscript.
Availability of data and material

All the relevant data used in the manuscript is available in the published scientific references. The software used in the simulations is licensed for academic user and is registered in the name of the first author.

Code availability

The codes used in the simulations are available upon request by electronic mail to the first author. We observe, however, that the authors do not make the commitment to provide an user-friendly code and/or explain the details of the implementations. The methods used in the codes are explicitly described in the manuscript.

Author contributions

Raul Abreu de Assis elaborated the model, directed the analysis, helped create codes for the simulations, and led the writing of the manuscript. Gustavo Cannale proposed the biological problem, provided biological references, estimated part of the parameters for *Sapajus xanthosternos*, provided advice in biological issues and helped writing the manuscript. Mazilio Coronel Malavazi, Rubens Pazim, Odair Fonseca and Moiseis Cecconello created routines in both MATLAB and Scilab, executed numerical simulations and provided mathematical criticism for the model and simulations. All authors contributed critically to the drafts and gave final approval for publication. Not applicable.

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APPENDIX

A GEOMETRICAL METHODS OF SHAPE DEFORMATION

To generate figures with various values of GE, we adopted three methods. Method 1 is based on “stretching” the figure in one direction, method 2 is based on adding irregularities at the boundaries, increasing the perimeter without stretching the figure in any direction, finally, the method 3 is a combination of the first two. In figure [A1] we present some examples of figures generated by the three methods.

Given a target area $a_{r_0}$, a polygon $P$ with $nv$ vertices is generated using polar coordinates, with angle $a_i = \frac{2\pi i}{nv+1}$, and ray $p_i$, $i = 0, 1, \ldots, nv-1$, such that the area of $P$ is $ar_{0}$. For each region $P$, two preserving area geometric transformations are applied
on $P$ in order to modify its measure of geometric efficiency (GE). One of these transformation consists in stretching $P$ alongside one of the axis. The second one consists in stretching $np$ (out of $nv$) vertices for a factor $r_i, i = 1, 2 \ldots, np$.

The data used in this article was create according to the following steps:

1. Initially, for a given $ar_0$, a region $P$ is created with $nv$ vertices, following a uniform distribution $U([a, b])$ and following a uniform distribution $U([c, d])$ such that $GE > 0.65$;

2. Using the first transformation method on $P$, ten regions $R_i$ are created with geometric efficiency $\frac{GE}{10}, i = 1, 2, \ldots, 10$;

3. Using the second transformation method on $P$, ten regions $S_i$ are created with geometric efficiency $\left(0.2 + \frac{i-1}{12}\right)GE, i = 1, 2, \ldots, 10$. In this case, $np$ is a random variable following a uniform distribution $U([e, f])$ and $r_i$ follows a uniform distribution $U([g, h])$;

4. Using a combination of those two methods, ten regions $T_i$ are created like this: first, $\tilde{T}_i$ is created, using the second transformation method, with geometric efficiency $\tilde{E}G = \left(0.2 + \frac{i-1}{12}\right)E\tilde{G}$ and then $T_i$ is created, using the first transformation method, with geometric efficiency $\gamma \tilde{E}G, \gamma < 1$.

For a given $ar_0$, steps 1-4 previously describe are repeated ten times so that, for such $ar_0$, thirty regions are created.
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