



## Cronfa - Swansea University Open Access Repository

This is an author produced version of a paper published in : *Theoretical Population Biology* 

Cronfa URL for this paper: http://cronfa.swan.ac.uk/Record/cronfa25962

## Paper:

Irvine, M., Bull, J. & Keeling, M. (2016). Aggregation dynamics explain vegetation patch-size distributions. *Theoretical Population Biology*, *108*, 70-74.

http://dx.doi.org/10.1016/j.tpb.2015.12.001

This article is brought to you by Swansea University. Any person downloading material is agreeing to abide by the terms of the repository licence. Authors are personally responsible for adhering to publisher restrictions or conditions. When uploading content they are required to comply with their publisher agreement and the SHERPA RoMEO database to judge whether or not it is copyright safe to add this version of the paper to this repository. http://www.swansea.ac.uk/iss/researchsupport/cronfa-support/

# Aggregation dynamics explain vegetation patch-size distributions

MA. Irvine<sup>a,\*</sup>, JC. Bull<sup>b</sup>, MJ. Keeling<sup>a</sup>

<sup>a</sup>Centre for Complexity Science, University of Warwick, Coventry, CV4 7AL, UK. <sup>b</sup>Department of Biosciences, Wallace Building, Swansea University, Swansea, SA2 8PP, UK.

## Abstract

Vegetation patch-size distributions have been an intense area of study for theoreticians and applied ecologists alike in recent years. Of particular interest is the seemingly ubiquitous nature of power-law patch-size distributions emerging in a number of diverse ecosystems. The leading explanation of the emergence of these power-laws is due to local facilitative mechanisms. There is also a common transition from power law to exponential distribution when a system is under global pressure, such as grazing or lack of rainfall. These phenomena require a simple mechanistic explanation. Here, we study vegetation patches from a spatially implicit, patch dynamic viewpoint. We show that under minimal assumptions a power-law patch-size distribution appears as a natural consequence of aggregation. A linear death term also leads to an exponential term in the distribution for any non-zero death rate. This work shows the origin of the breakdown of the power-law under increasing pressure and shows that in general, we expect to observe a power law with an exponential cutoff (rather than pure power laws). The estimated parameters of this distribution also provide insight into the underlying ecological mechanisms of aggregation and death.

*Keywords:* patch-size distribution, pattern formation, spatial ecology, aggregation

Preprint submitted to Theoretical Population Biology

<sup>\*</sup>Corresponding author. Email address: M.A.Irvine@Warwick.ac.uk (MA. Irvine )

## 1 1. Introduction

Vegetation patch-size distributions have been under intense study in re-2 cent years [1, 2, 3, 4, 5]. It has been shown that a power-law provides a 3 good fit to the patch-size distribution under a robust range conditions, how-4 ever there are marginal cases to this. Kéfi et al. [6] analysed patch-size 5 distributions in semi-arid vegetation in the Mediterranean and found that 6 there was not only a power-law distribution evident in the patch-size distri-7 bution, but also a truncated exponential term, when the system was under 8 increased grazing pressure. Similar power-law distribution phenomena have 9 also been detected in a number of other ecosystems including mussel beds 10 [7] and marine benthic diatoms [8]. This phenomena of a power-law distri-11 bution transitioning to an exponential distribution under increasing stress 12 has recently shown to be robust, where diverse ecological models are able to 13 reproduce these results [2]. 14

The leading explanation of this power-law pattern formation in ecology 15 is due to local interactions driving the large-scale behaviour [9, 10]. Scanlon 16 et al. [11] supported through the use of numerical simulation of spatially-17 explicit models of vegetation growth combined with a global effect on the 18 population density interpreted as the amount of rainfall or other global pro-19 cesses. The local positive feedback process driving the patch formation is 20 through facilitation of neighbourhood sites that increase the birth rate and 21 decrease the death rate [5]. This explanation does not answer how a power-22 law forms at the patch level, whether it is due to a competition effect between 23 larger clusters dominating the landscape or an aggregation of smaller clus-24 ters. There is also an open question of how patches aggregating together 25 drives these observed patterns. 26

Models of aggregation and fragmentation have been considered in other 27 areas in ecology such as the size of fish schools [12] and marine diatoms [13]. 28 Aggregation phenomena has been more generally studied in the Physical sci-29 ences [14], including processes such as polymerisation [15], coagulation of 30 aerosols [16] and flocculation [17]. Although these examples include clusters 31 that may diffuse, aggregation phenomena may also be considered in the case 32 where clusters are immobile [18]. Aggregation of vegetation clusters, how-33 ever, has not been previously considered as an explicit driving force of the 34 evolution of the patch-size distribution. Our novel contribution here is to 35 apply established theory of aggregation dynamics to the system of vegeta-36 tion clusters and derive a new model of aggregation with global death that 37

<sup>38</sup> is applicable to vegetation dynamics.

In this article, spatially implicit models of vegetation clusters are investi-39 gated by considering how patches form and aggregate. The general conditions 40 under which a power-law distribution is expected to emerge are explored as 41 well as when there is a breakdown of the power law distribution due to an ex-42 ponential truncation. By adopting a patch-centric viewpoint, the impact of 43 aggregation on the resulting distribution along with other processes may be 44 studied directly. This represents a powerful new approach to understanding 45 the origin of these distributions, by explicitly modelling the patch-size dy-46 namics without the need to infer the patch-size distribution from a spatially 47 explicit model [5]. 48

Further, the connection between the power-law exponent and the persis-49 tence of the distribution in this model is explored. We begin by defining a 50 novel model of aggregation with linear death and then deriving an asymp-51 totic solution when the death rate is small. This analytic result is compared 52 to a simulation study of vegetation with local and global growth properties 53 subjected to a global disturbance. For small disturbance, the power law ex-54 ponent closely matches the exponent expected from the model. The conclu-55 sion is that the power-law clustering observed in many vegetation ecosystems 56 may simply be an aggregation effect and the exponential truncation observed 57 when there is increased stress is due to an increase in the linear death rate 58 of clusters. 59

#### 60 2. Theory

The idea developed here is to model the patches themselves as opposed 61 to an individual spatial site as is done in probabilistic cellular automata 62 [19, 20]. We denote  $c_k(t)$  as the density of patches of size k at time t, where 63 time is taken to be continuous. A continuous model of patch-sizes can be 64 studied, however for the present k shall take positive integer values only, 65  $k \in \{1, 2, \ldots\}$ . A kernel of aggregation gives the rate at which patches of 66 size i and j aggregate together to form a patch of size i + j, this kernel 67 is denoted K(i, j). Finally it is assumed there is a constant rate at which 68 patches of size 1 or monomers enter the system. These assumptions are 60 general and can include many different phenomena, including static clusters 70 and diffusing monomers [18]. The governing master equation, also known as 71

<sup>72</sup> the Smoluchowski equation [21] is then

$$\frac{\mathrm{d}}{\mathrm{d}t}c_k = \frac{1}{2}\sum_{i+j=k} K(i,j)c_ic_j - \sum_{j\geq 1} K(j,k)c_jc_k + \delta_{k,1},\tag{1}$$

where  $\delta_{k,1}$  is the Kronecker-delta function that is 1 when k = 1 and 0 other-73 wise. For convenience, time has been re-scaled such that the rate at which 74 aggregation occurs is 1. It is instructive to imagine a single unit or monomer 75 coming into contact with a cluster and calculating the rate at which this oc-76 curs for larger as opposed to smaller clusters. If a > 0 then, assuming the size 77 of the monomer is negligible, the monomer rate equation is  $K(i) = i^{-a}$ . This 78 means smaller clusters are favoured and the growth rate reduces as clusters 79 grow larger in size. An ecological explanation of this could be due to the 80 self-limitation through competition a larger cluster experiences with itself, 81 thus reducing its potential for growth. Smaller clusters have more space and 82 thus can grow at a quicker rate. 83

When a < 0, larger clusters are favoured for growth compared with 84 smaller clusters, this can be seen as a form of the Allee effect [22]. In the 85 regime when a < 0, small clusters are more susceptible to environmental 86 perturbation and as such, have a lower propensity for growth. At the other 87 length scale, larger clusters of vegetation are able to regulate their environ-88 ment more and thus have greater resources for growth (An example species 89 where this holds is ribbed mussels [23], where larger clusters provide protec-90 tion and shelter for new mussels). This example of an Allee effect can be 91 demonstrated by again considering the rate at which single units of vegetation 92 aggregate to a cluster. If a i > j, then  $K(1, i) = 1 + i^{-a} > 1 + j^{-a} = K(1, j)$ 93 i.e. the rate at which a larger patch recruits new growth is greater than for a 94 smaller patch. A value for a then can give an indication of whether there is 95 strong small cluster growth at the expense of large clusters forming or if the 96 converse holds. 97

An alternative explanation of the aggregation exponent a is due to the edge effects of a cluster. A single individual vegetation unit aggregates to a cluster proportional to the edge of that cluster. If all clusters are non-fractal then it would be expected that a vegetation unit aggregates at rate  $i^{1/2}$ , since the length of a non-fractal object scales as a square root with its area. For a general fractal cluster with boundary dimension d, it would be expected that an individual unit scales as  $i^{1/d}$ .

Various properties are desirable for the kernel. Firstly symmetry, where the rate at which patches of size i and j aggregate does not depend on the <sup>107</sup> ordering of the patches i.e. K(i, j) = K(j, i). Secondly, scaling homogeneity, <sup>108</sup> where the rate at which patches of a certain size aggregate scales by some <sup>109</sup> factor  $K(mi, mj) = m^{\lambda}K(i, j)$ . The simplest kernel that satisfies these con-<sup>110</sup> ditions is the constant kernel K(i, j) = 1, corresponding to the case where <sup>111</sup>  $\lambda = 0$ . When this form of kernel is assumed, the tail-solution (for large k) <sup>112</sup> has the simple form [24]

$$c_k \sim \frac{1}{\sqrt{4\pi}} \frac{1}{k^{3/2}}.$$
 (2)

The tail of the patch-size distribution is a power law with exponent 3/2, where the power law nature of the solution is a consequence of the injection term (where births of patch size one enter the system) and the non-linear aggregation term in the equation. The equation can be solved analytically for more general kernels of the type

$$K(i,j) = i^{-a} + j^{-a}.$$
 (3)

<sup>119</sup> This type of kernel also admits an analytic solution in the large patch-size <sup>120</sup> limit [25, 26] with a steady state distribution of the form where

118

$$c_k \sim C k^{-\tau} \tag{4a}$$

$$\tau = \frac{3-a}{2}, \quad C = \sqrt{\frac{1-a^2}{4\pi}} \cos\left(\frac{\pi a}{2}\right).$$
 (4b)

For a steady state to exist we require -1 < a < 1 and hence the scaling 121 exponent can be found on the interval  $\tau \in (1,2)$ . The dynamics of the 122 equation can be assessed by defining the cross-over time, which is the time 123 taken for a density of patches of a certain size to reach its asymptotic value. 124 The cross-over time for a patch of size  $k_*$  to the steady state solution  $c_{k_*}$ 125 is found to take the form  $t = (k_*)^z$  where z = (1+a)/2. The scaling of 126 the cross-over time and the patch-size exponent can be related by the simple 127 linear equation  $\tau = 2 - z$ . This gives a linear relationship between the static 128 exponent at stationarity and its dynamic exponent. 129

A real vegetation system is not purely defined by an aggregation process however. In particular in the previous example there is no death either of single vegetation units or patch clusters. Death may lead to changes in the exponent of the stationary distribution and so it is important to include in any model of vegetation clustering. It is also assumed that a death event does not lead to fragmentation of the cluster. A modified Smoluchowski equation <sup>136</sup> with a linear death term can then be produced as

$$\frac{\mathrm{d}}{\mathrm{d}t}c_k = \frac{1}{2}\sum_{i+j=k}K(i,j)c_ic_j - \sum_{j\geq 1}K(j,k)c_jc_k + \mu(k+1)c_{k+1} - \mu(k)c_k, \quad (5a)$$

$$\frac{\mathrm{d}}{\mathrm{d}t}c_1 = -\sum_{j\ge 1} K(j,1)c_jc_1 + 1 + \mu(2)c_2 - \mu(1)c_1.$$
(5b)

The general additive aggregation kernel is again taken to be of the form 137  $K(i, j) = i^{-a} + j^{-a}$ , where a represents the scaling parameter of the rate at 138 which aggregates of a certain size join. If it is equally likely for a cluster 139 of a certain size to aggregate with a cluster of any other size then the scale 140 parameter a = 0. For a pure aggregation system with no fragmentation, this 141 leads to a cluster scaling of 3/2.  $\mu(k)$  defines the death rate, which is the 142 rate at which individual units are lost from a patch, where a patch of size k143 transitions to a patch of size k-1 due to exogenous or endogenous factors. 144 A number of different forms of this death rate may be considered dependent 145 on the biological details of the system. For example if each individual has a 146 constant rate of death regardless of the size of patch its contained, such as due 147 to lack of rainfall or grazing, is then  $\mu(k) = \mu k$ . If death occurs at the edge 148 of a patch then the death rate is  $\mu k^{1/d}$ , where d is the boundary dimension 149 of the patches. The simplest form of the death rate is where  $\mu(k) = \mu$  for 150 all k. In order to gain insight into the effect of a death rate on the resulting 151 patch-size distribution, we assume the final form of the death rate. 152

In order to gain analytic tractability on the model a constant aggregation kernel is assumed (a = 0, K = 2) together with a constant death rate for each individual within a patch. The strategy for deriving a solution is similar to the strategy in Krapivsky et al. [26]. A constant kernel K(i, j) = 2 is used. Eq. (5) is rewritten as

$$\frac{\mathrm{d}}{\mathrm{d}t}c_k = \sum_{i+j=k} c_i c_j - 2c_k \sum_{j\geq 1} c_j + \mu c_{k+1} - \mu c_k,$$
(6a)

$$\frac{\mathrm{d}}{\mathrm{d}t}c_1 = -2c_1 \sum_{j\ge 1} c_j + 1 + \mu c_2 - \mu c_1.$$
(6b)

The asymptotic tail of the resulting patch-size distribution is then sought in order to gain an understanding of how the linear death rate affects the stationary distribution. By using the asymptotic approximation and assuming k is large, the k-th coefficient in this expansion and hence the density of 162 patches of size k is

$$c_k = k^{-3/2} \exp(-\Lambda k),\tag{7}$$

where  $\Lambda = \log(1 + \mu N)$  and N is the total population size (See Appendix A 163 for a derivation). The solution is therefore a power law with an exponential 164 truncation of factor  $\Lambda$ . When the death rate is 0,  $\Lambda = 0$  and hence the patch-165 size distribution is a pure power law as is expected. A large death rate will 166 lead to a solution that is dominated by an exponential decay term, hence the 167 patch-size distribution is expected to have a smooth transition from a pure 168 power law to an exponential distribution. A dimensionality argument of Eq. 6 169 [27] also leads to a power law exponent of the form 3/2. This solution can 170 be compared to the general power-size distribution with exponential cut-off 171 N(k) given by 172

$$N(k) = Ck^{-\alpha} \exp(-k/k_x), \tag{8}$$

where  $k_x$  is the patch-size above which N(s) decreases faster than power-law [2]. Matching terms and assuming  $\mu N$  is small gives the following simple relationship between the cross-over patch size  $k_x$  and death rate  $\mu$  as

$$k_x = \frac{1}{N\mu}.$$
(9)

The model therefore predicts an inverse relationship between patch-size crossover and death rate. This also predicts that when the death rate is small enough, the cross-over patch-size  $k_x$  may be larger than the system size and as such the exponential tail may not be observed in empirical distributions.

## 182 3. Results

173

In order to compare the model predictions of patch formation in an aggre-183 gation system with a constant death rate to the prediction of the patch-size 184 distribution obtained in Eq. 7 is compared to a simple probabilistic cellu-185 lar automata model of vegetation growth. The cellular model is similar to 186 the one discussed in [11], the model is defined on a toroidal lattice where 187 each site can exist in one of two states: occupied (1) and empty (0). The 188 occupied state propagates through nearest neighbour growth at rate  $\beta$ , as 189 well as through a background constant birth probability  $\gamma$ . The alive sites 190 transition to a dead site with a constant death probability  $\mu$ . Hence if  $n_x$  is 191 the number of alive sites in the neighbourhood of site x, the transitions can 192

<sup>193</sup> be summarised as

$$P_x(0 \to 1) = \epsilon \min\{1, \gamma + \beta n_x/4\},\tag{10a}$$

$$P_x(1 \to 0) = \epsilon \mu, \tag{10b}$$

where  $\epsilon$  sets the total reaction rate of the system and was implemented to reduce the probability of multiple events occurring within the same neighbourhood. The minimum function is used here to guarantee the probability of transitioning to an alive state is one in the rare case when the sum of the two probabilities increases above one.

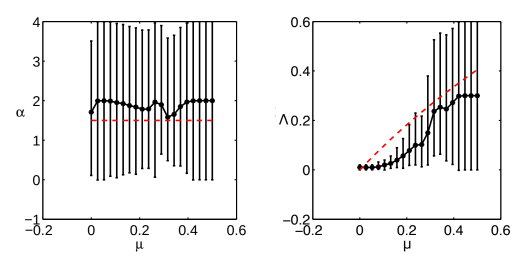
Simulations were conducted for constant local growth, birth rate and reaction rate  $\beta = 0.2, \gamma = 0.01, \epsilon = 0.1$  and over a range of death rates. Simulations were ran for lattice length L = 500 and for 1000 replicates of each parameter set. The patch-size distribution was recorded for each simulation run after 600 time-steps. This was chosen so that when  $\mu = 0$ , the lattice is approximately 50% occupied. The following power-law with exponential truncation was fitted to the distribution using a maximum likelihood method

$$f(K = k) = Ck^{-\alpha} \exp(-\Lambda k), \tag{11}$$

for some normalising factor C. The resulting maximum likelihood estima-206 tors were found using a downhill simplex method implemented in Matlab 207 R2014a [28]. The approximate solution to the aggregation equation predicts 208 a constant power-law exponent  $\alpha$  of 3/2. This is close to the inferred value 209 from simulation for the range of  $\mu$  values studied (Fig. 1a). The exponential 210 factor  $\Lambda$  is zero when the death rate is zero (Fig. 1b), as predicted. For 211 increasing death rate,  $\Lambda$  increases again as predicted. Overall there is an 212 increase in the exponential factor for increasing death rate as is predicted, 213 however the functional form of the increase is not fully captured by the mean 214 field approximation. 215

#### 216 4. Discussion

<sup>217</sup> Changing the focus away from explicit spatial modelling of vegetation <sup>218</sup> patch formation and instead focusing on the dynamics of patch-sizes gives a <sup>219</sup> unique insight into the underlying aggregation-fragmentation processes. Here <sup>220</sup> we have primarily focused on solutions to equations where the aggregation <sup>221</sup> kernel that governs the rate at which patches of two sizes will aggregate by



(a) comparison of power-law exponent from simulation to theoretical prediction

(b) comparison of exponential factor from simulation to theoretical prediction

Figure 1: Exponents of patch-size distribution compared to simulations. Theoretical values shown as red dashed lines, while simulation calculated values are given as black dots with 95% confidence intervals. The theoretical values for the power-law exponent  $\alpha$  and the exponential factor  $\Lambda$  are derived in Eq. 7. As predicted for small values of the death rate the power-law component of the patch-size distribution is constant whilst there is an increase in the exponential component for increasing death rate.

either a constant or power law kernel. For a system where there is aggrega-222 tion only the resulting patch-size distribution is that of a pure power law, 223 with exponent that is dependent on the exponent of the power law aggrega-224 tion kernel. The introduction of a linear death term, where an individual is 225 lost from a patch at rate  $\mu/k$  gives rise to a power law with exponential tail 226 distribution of the form  $c_k \sim k^{-\alpha} \exp(-\Lambda k)$ . This solution holds generally 227 when there is a linear death term and power-law aggregation kernel, even 228 when the kernel is composed of a sum of two power-laws. Further,  $\alpha$  is de-220 pendent on the specifics of the aggregation term alone and  $\Lambda$  is dependent on 230 the death rate alone. This separation of the aggregation and fragmentation 231 term implies, in principle, the ability to infer aggregation and death processes 232 through observing the converged patch-size distribution alone, hence this is 233 applicable to inferring process from a single spatial snapshot. 234

Kéfi et al. [6, 2] predicts that a power-law distribution in the patch-size 235 distribution occurs when a global environmental death rate is small. This 236 transitions to an exponential distribution when there is greater stress on the 237 system through this global death rate term. The model used is a spatially-238 explicit one with a local growth term and a background death rate. The 239 model proposed in this article can be seen as a deterministic equivalent of 240 this spatially-explicit model. Through the derived solution in this article it is 241 observed that there should always be an exponential tail to the distribution 242 if the death rate is non-zero. Similar arguments have also been made recently 243 [29], but notably none have explained the origin of a power law with exponen-244 tial tail observed in vegetation systems. The derived model then, provides a 245 theoretical origin to the observed spatial patterns in vegetation ecosystems 246 that are under a pressure that can be considered constant throughout space 247 (rainfall, grazing etc.). As an example, if all sites had the same death rate 248 regardless of its neighbourhood, such as for a grazer, the death rate would 249 be  $\mu k$  for a patch of size k. The model also suggests that a power-law with 250 exponential tail is a more accurate description of the patch-size distribution, 251 although when the death rate is small, the exponential tail may not be ob-252 served directly. This approach would be able to provide further insights into 253 the nature of the patch-size distribution for other systems where disturbance 254 may be spatially distributed. 255

The model also gives insight into how there can be a continuous array 256 of power-law exponent observed in nature. The aggregation with no death 257 model predicts that power-laws exponents in the range (1,2) are physically 258 possible, which is what has been observed in a number of ecosystems [7, 6, 8]. 259 The model predicts that a change in the exponent of a patch-size distribu-260 tion is related to a change in the power-law exponent of the aggregation 261 kernel. A simple dimensionality argument can be used to show that in the 262 aggregation and death model with a kernel that has a general power law 263 scaling as described in Eq. 3, the resulting stationary distribution will have 264 the same exponent as that in the model with no death [27]. The conclusion 265 of how to relate the patch-size distribution to the system dynamics is that 266 both the power-law exponent and the presence of an exponential cut-off does 267 give an indication of the underlying dynamics. More complex fragmentation 268 processes than the one discussed would alter these conclusions however, as a 269 non-linear fragmentation process will also lead to self-similar solutions and 270 thus the two processes are confounded when only the stationary state is ob-271 served [30], such processes include storms and other strong weather events 272

that could split a single cluster of vegetation into multiple clusters. The size of the system where the dynamics occur, such as in the lattice model, may also have an impact on the exponents of the patch-size distribution due to finite-size effects [31].

Other possible extensions to the model could include multi-species sys-277 tems, where patches are formed of multiple species each with their own in-278 trinsic death rates. Multi-species systems have already been considered in 279 the physical sciences and as such this would make for an interesting avenue of 280 future research [32]. Where the aggregation process is indistinguishable be-281 tween two different species, this leads to similar results laid out in this article 282 [33]. However, more complex interactions such as inter-specific competition 283 would inevitably lead to a more complex relationship between the exponent 284 death term and the underlying death rates. The model equations were scaled 285 such that the rate of aggregation and rate at which single vegetation units 286 are created is one. This was done for convenience since we were interested in 287 studying the scaling alone, whereas these parameters change the constant of 288 the patch-size distribution only. Another extension then would be to explic-289 itly calculate the constant for the patch-size distribution and study how this 290 changes as the other system rates change. 291

## 292 Acknowledgement

The work conducted in this article was funded as part of an EPSRC studentship.

### <sup>295</sup> Appendix A. Derivation of asymptotic solution

A moment-generating function is used to find the steady state solution to this equation in a similar fashion to the one described in Krapivsky et al. [26]. Firstly define the total number of all patches as  $N = \sum_{k\geq 1} c_k$  and then sum Eq. (6a-b) in order to obtain

$$\frac{dN}{dt} = \sum_{k\geq 1} \sum_{i+j=k} c_i c_j - 2 \sum_{k\geq 1} c_k \sum_{j\geq 1} c_j + 1 + \sum_{k\geq 1} \mu c_{k+1} - \sum_{k\geq 1} \mu c_k, \quad (A.1)$$

$$\frac{dN}{dt} = N^2 - 2N^2 + 1 - \mu c_1, \tag{A.2}$$

$$\frac{dN}{dt} = -N^2 + 1 - \mu c_1. \tag{A.3}$$

<sup>300</sup> Dynamically, consider when N is at equilibrium. If  $\mu = 0$  then the stationary <sup>301</sup> solution is N = 1. If  $\mu > 0$  then the equilibrium solution is necessarily <sup>302</sup> bounded between one and zero as N and  $c_1$  are always positive.

The moment-generating function  $C(z,t) = \sum_{k=1}^{\infty} c_k z^k$  is now considered. Multiplying Eq. (6) by  $z^k$  and summing over all k gives the following

$$\frac{\mathrm{d}}{\mathrm{d}t}C = C^2 - 2NC + z + \mu \sum_{k \ge 1} z^k c_{k+1} - \mu \sum_{k \ge 1} z^k c_k$$
$$= C^2 - 2NC + z + \frac{\mu}{z}C - \mu C - \mu c_1.$$
(A.4)

The  $C^2$  term is derived using the relationship

$$\left(\sum_{k\geq 1} a_k\right)^2 = \left(\sum_{i\geq 1} a_i\right) \left(\sum_{j\geq 1} a_j\right) = \sum_{k\geq 1} \sum_{i+j=k} a_i a_j.$$
(A.5)

The new moment generating function defined as A(z,t) = C(z,t) - N(z,t) is considered in order to derive the final stationary solution. The time derivative is calculated by combining Eq. A.4 with Eq. A.3

$$\frac{d}{dt}A(z,t) = \frac{d}{dt}C(z,t) - \frac{d}{dt}N(t) 
= C^2 - 2NC + \frac{\mu}{z}C + z - \mu C - \mu c_1 - 1 + N^2 + \mu c_1 
= A^2 + \frac{\mu}{z}C - \mu C + z - 1 
= A^2 + \mu \frac{1-z}{z}A + \mu \frac{1-z}{z}N + z - 1.$$
(A.6)

Note that the right-hand side is quadratic in terms of *A*. Setting the timederivative to zero gives the steady-state solution of the moment-generating function as

$$A = \mu \frac{z-1}{z} + \sqrt{\mu^2 \frac{(1-z)^2}{z^2} - 4\left(\mu \frac{1-z}{z}N + z - 1\right)}.$$
 (A.7)

In order to proceed it is assumed that the death rate  $\mu$  is small and only the leading order term is kept. Hence

$$A \approx 2\sqrt{1 - z - \mu \frac{1 - z}{z}N}.$$
(A.8)

The strategy is to find A in terms of the power series  $\sum_{k=1}^{\infty} c_k z^k$ , where  $c_k$  is a function of  $\mu$ . Assuming z is sufficiently close to one such that  $z + \mu(1 - z)N/z < 1$ , the expansion of  $\sqrt{1-x}$  is used to obtain

$$A_{\text{approx}} = 2\sum_{k=0}^{\infty} \frac{\Gamma(3/2)}{\Gamma(3/2-k)\Gamma(k+1)} (1+\mu N)^{1/2-k} \left(-z-\mu N/z\right)^k.$$
 (A.9)

<sup>318</sup> Using the relationship  $\Gamma(z)\Gamma(1-z) = \frac{\pi}{\sin(\pi z)}$ , cancelling the  $(-1)^k$  terms and <sup>319</sup> absorbing all constants into a constant c term

$$A_{\text{approx}} = c \sum_{k=0}^{\infty} \frac{\Gamma(k-1/2)}{\Gamma(k+1)} (1+\mu N)^{1/2-k} \left(z+\mu N/z\right)^k.$$
(A.10)

<sup>320</sup> Using the binomial expansion, this becomes

$$A_{\text{approx}} = c \sum_{k=0}^{\infty} \sum_{i=0}^{k} \frac{\Gamma(k-1/2)}{\Gamma(k+1)} \frac{\Gamma(k+1)}{\Gamma(i+1)\Gamma(k-i+1)} (1+\mu N)^{1/2-k} (\mu N)^{k-i} z^{2i-k}.$$
(A.11)

In order to find the k-th coefficient as  $k \gg 1$  the leading order of the *i* term in the binomial is considered. Given that  $\mu N \ll 1$ , the *i* dependent terms are dominated by i = k. Hence the kth term of the expansion where k is large is

$$c\frac{\Gamma(k-1/2)}{\Gamma(k+1)}(1+\mu N)^{1/2-k}z^k.$$
(A.12)

<sup>327</sup> By using the asymptotic approximation  $\Gamma(n+a)/\Gamma(n) \sim n^a$  and assuming <sup>328</sup> k is large, the k-th coefficient in this expansion and hence the density of <sup>329</sup> patches of size k is

$$c_k = ck^{-3/2} \exp(-\Lambda k), \qquad (A.13)$$

331 where  $\Lambda = \log(1 + \mu N)$ .

#### 332 **References**

321

326

330

[1] M. Scheffer, J. Bascompte, W. A. Brock, V. Brovkin, S. R. Carpenter,
V. Dakos, H. Held, E. H. van Nes, M. Rietkerk, G. Sugihara, Nature
461 (2009) 53–59.

[2] S. Kéfi, M. Rietkerk, M. Roy, A. Franc, P. C. De Ruiter, M. Pascual,
Ecology letters 14 (2011) 29–35.

- [3] M. Rietkerk, S. C. Dekker, P. C. de Ruiter, J. van de Koppel, Science
   305 (2004) 1926–1929.
- <sup>340</sup> [4] G. M. Oborny, B. György Szabó, Oikos 109 (2005) 291–296.
- <sup>341</sup> [5] A. Manor, N. M. Shnerb, Physical review letters 101 (2008) 268104.
- [6] S. Kéfi, M. Rietkerk, C. L. Alados, Y. Pueyo, V. P. Papanastasis,
  A. ElAich, P. C. De Ruiter, Nature 449 (2007) 213–217.
- [7] F. Guichard, P. M. Halpin, G. W. Allison, J. Lubchenco, B. A. Menge,
   The American Naturalist 161 (2003) 889–904.
- [8] E. Weerman, J. Van Belzen, M. Rietkerk, S. Temmerman, S. Kéfi,
  P. Herman, J. V. de Koppel, Ecology 93 (2012) 608–618.
- [9] M. Pascual, M. Roy, F. Guichard, G. Flierl, Philosophical Transactions
  of the Royal Society of London B: Biological Sciences 357 (2002) 657–
  666.
- <sup>351</sup> [10] M. Roy, M. Pascual, A. Franc (2003).
- [11] T. M. Scanlon, K. K. Caylor, S. A. Levin, I. Rodriguez-Iturbe, Nature
   449 (2007) 209–212.
- <sup>354</sup> [12] H.-S. Niwa, Journal of theoretical biology 195 (1998) 351–361.
- [13] G. A. Jackson, Deep Sea Research Part A. Oceanographic Research
   Papers 37 (1990) 1197–1211.
- <sup>357</sup> [14] D. J. Aldous, Bernoulli (1999) 3–48.
- <sup>358</sup> [15] R. M. Ziff, Journal of Statistical Physics 23 (1980) 241–263.
- <sup>359</sup> [16] W. Koch, S. Friedlander, Journal of Colloid and Interface Science 140
   (1990) 419–427.
- [17] K. D. Danov, I. B. Ivanov, T. D. Gurkov, R. P. Borwankar, Journal of
   colloid and interface science 167 (1994) 8–17.
- [18] P. Krapivsky, J. Mendes, S. Redner, The European Physical Journal
   B-Condensed Matter and Complex Systems 4 (1998) 401–404.

- <sup>365</sup> [19] P. Hogeweg, Applied mathematics and computation 27 (1988) 81–100.
- <sup>366</sup> [20] H. Balzter, P. W. Braun, W. Köhler, Ecological modelling 107 (1998)
   <sup>367</sup> 113–125.
- <sup>368</sup> [21] M. Von Smoluchowski, Z. Phys. 17 (1916) 557–585.
- <sup>369</sup> [22] P. A. Stephens, W. J. Sutherland, Trends in Ecology & Evolution 14
   (1999) 401–405.
- <sup>371</sup> [23] M. D. Bertness, E. Grosholz, Oecologia 67 (1985) 192–204.
- <sup>372</sup> [24] H. Hayakawa, Journal of Physics A: Mathematical and General 20 (1987)
   <sup>373</sup> L801.
- <sup>374</sup> [25] P. L. Krapivsky, J. F. F. Mendes, S. Redner, physical review B 59 (1999)
   <sup>375</sup> 15950–15958.
- P. L. Krapivsky, S. Redner, E. Ben-Naim, A kinetic view of statistical
   physics, Cambridge University Press, 2010.
- [27] C. Connaughton, R. Rajesh, O. Zaboronski, Physical Review E 69 (2004)
  061114.
- [28] J. C. Lagarias, J. A. Reeds, M. H. Wright, P. E. Wright, SIAM Journal
  on optimization 9 (1998) 112–147.
- <sup>382</sup> [29] S. Pueyo, Landscape ecology 26 (2011) 305–309.
- <sup>383</sup> [30] M. Ernst, P. Van Dongen, Physical Review A 36 (1987) 435.
- [31] C. Connaughton, R. Rajesh, O. Zaboronski, Physical Review E 78 (2008)
  041403.
- [32] C. Pilinis, Atmospheric Environment. Part A. General Topics 24 (1990)
   1923–1928.
- [33] R. D. Vigil, R. M. Ziff, Chemical engineering science 53 (1998) 1725–
   1729.