Extinction models of robustness for weighted ecological networks

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Abstract

1. Analysis of ecological networks is a valuable approach to understanding the vulnerability of systems to environmental change. The tolerance of ecological networks to co-extinctions, resulting from sequences of primary extinctions, is a widely-used tool for modelling network 'robustness'. Previously, these 'extinction models' have been developed for and applied to binary networks (sometimes weighted by species abundance) and recently used to predict cascades of co-extinctions in plant-pollinator networks. There is a need for robustness models that can be applied to weighted networks (i.e. where interaction frequencies are recorded) and there is a need to understand how the structure of a network affects its robustness.

2. Here, we developed a framework of extinction models that examine the robustness of networks to random and structurally correlated extinctions (i.e. where avalanches of co-extinctions can occur due to feedback between the two trophic levels). Our models can be applied to networks with binary and weighted interactions. We tested how the average and the range of robustness values is impacted by network structure and the impact of structurally-correlated extinctions sampling non-uniformly from the distribution of random extinction sequences. Our findings are illustrated with plant-pollinator networks.

3. We found that the range of robustness values is driven by the structural heterogeneity of species (plants) in the sequence of primary extinctions. With structurally-correlated extinctions, the networks tested were more
robust to ‘avalanche’ extinctions than expected due to the robustness of highly-connected species, but are less robust when preferentially selecting plant species with shared pollinators using a ‘random walk’ model. We found that weighted interactions tend to amplify observed effects and increase variation in robustness.

4. Our framework of models is a new approach to assessing network robustness, permitting it to be calculated with weighted, as well as binary, interaction networks. Models which permit cascades of extinctions vary in their impact on robustness, thus highlighting the vital importance of understanding the model when interpreting robustness.

**Key-words (max 10)**

co-extinction, ecological interactions, extinction models, mutualistic network, network, plant-pollinator communities, pollinators, robustness,

1. **INTRODUCTION**

Network analysis has become an important tool for ecologists seeking to understand the vulnerability of ecosystems to environmental change. Recent research has centred on network approaches for improving our understanding of plant-pollinator communities and extinctions, especially in the light of the widely documented recent declines in key insect pollinators such as honeybees, bumblebees and butterflies (Biesmeijer *et al.*, 2006; Senapathi *et al.*, 2015; Goulson, Lye & Darvill, 2008; Benton, 2006). These trends are concerning for biodiversity, ecosystem function and food security (Potts *et al.*, 2010) as insect pollinators are known to play a vital role in providing ecosystem services (Bailes
et al., 2015). They feed on nectar and pollen provided by plant species, and whilst doing this facilitate the fertilisation of plants via cross pollination (Free, 1993; Lubbock, 1875). In many ecological systems, including plant-pollinator systems, the community can be regarded as a bipartite network comprising two distinct guilds of organisms in which each node represents a species, and species are connected by edges indicating interactions, which may be directly observed, indirectly observed (e.g. pollen analysis) or inferred (Morales-Castilla et al., 2015).

Models of community robustness based on observed plant-pollinator networks (available, for example, from http://www.web-of-life.es and https://www.nceas.ucsb.edu/interactionweb/resources.html) usually fall into one of two types. In the first (see for example Bastolla et al., 2009, James, Pitchford and Plank, 2012), the community is modelled as a dynamical system, in which the population of each species is affected by the interactions that species has with others. The dynamics are typically run to fixation, and the populations at fixation used to determine community robustness, and how it relates to overall network structure.

The second approach, which we adopt here, is to model the tolerance of the network to extinctions. This was pioneered by Albert, Jeong & Barabasi (2000) and rapidly applied to multitrophic food webs (Dunne, Williams and Martinez, 2002) and then mutualistic bipartite networks, especially plant-pollinator networks (Memmott, Waser and Price, 2004; Kaiser-Bunbury et al., 2010).

Extinction models estimate the robustness of a plant-pollinator network by sequentially removing species of the primary type (e.g. plants) and recording the number of surviving species of the secondary type (e.g. pollinators), by applying
some pre-determined rule for species survival. Most models, thus far, have used
simple rules for secondary extinctions e.g. species become extinct when all their
existing links are lost. Network robustness can then be determined from the area
under the curve of the proportion of the secondary type that survive against the
proportion of the primary type removed (Burgos et al., 2007; see Fig. 1a).
These models have been adapted in various different ways. One approach has
been to use specific sequences of primary extinctions, such as ordered by traits
of the nodes (Dunne, Williams & Martinez, 2002; Memmott, Waser and Price,
2004; Pocock, Evans & Memmott, 2012; and Santamaria et al., 2016). Another
development was to allow rewiring of edges (pollinators switching from one
plant to another) based on interactions known from other data (Kaiser-Bunbury
et al. 2010), with the aim of increasing the biological realism of extinction
simulations - an issue that many of these papers acknowledge is lacking. Most
recently, Vieira and Almeida-Neto (2015) allowed feedback in co-extinction
between guilds, which they implemented stochastically, so permitting the
possibility of cascades of extinctions.
Early extinction models showed that the robustness of communities to random
primary extinctions increased with network connectance, the fraction of the
possible interactions that were actually observed (Dunne, Williams & Martinez,
2002) and the resulting robustness was often interpreted in terms of network
also found that cascades were more likely in highly connected networks.
However, more detailed investigation of the impact of network structure on
robustness has been lacking. These studies form the foundation of extinction
models for plant-pollinator communities on which we base our work.
Initially, many of the observed plant-pollinator networks had binary edge weights; interactions between pairs of species were either observed or not. Increasingly researchers are measuring the frequency or importance of interactions to create weighted networks, yielding a better description of the interactions observed (Ings et al., 2009; Tylianakis et al., 2010) and accounting better for under-sampling biases (Bersier, Banašek-Richter & Cattin, 2002). But the lack of weighted robustness models means that the information on the interaction weights, when known, either has to be discarded (e.g. Pocock et al. 2012) or is used simply to weight the binary outcomes by node abundance (Kaiser-Bunbury et al. 2010). Therefore, there is a need for models that are suitable for measuring robustness on weighted networks. Here, we aim to address this need.

One of the features of these extinction models is that when using random sequences of primary extinctions on a single network, there is a broad distribution in the resulting robustness values (see Fig. 1b). Robustness is therefore a product both of structural heterogeneity of the network (eg Pastor et al., 2012) and of the method of producing extinction sequences; we will explore each of these contributions to robustness in this paper.

Here, our aim was firstly to develop a suite of extinction models which can be applied to weighted, as well as binary, bipartite interaction networks. Secondly, we sought to determine how robustness (assessed using our different models) is affected by network structure and how our models influence measured robustness. Throughout this paper we apply our models to mutualistic bipartite networks (specifically plant-pollinator networks) but discuss how they can be applied to any bipartite network (e.g. with trophic, uni-directional interactions).
2. MATERIALS AND METHODS

In this study, we examine the robustness of observed plant-pollinator networks that describe observed interactions between species in a community. A network has $P$ plant nodes and $A$ animal nodes, and contains $E$ interactions between species, encoded in the $A \times P$ matrix $M$. Interactions may be binary (b) or weighted (w).

We illustrate our models and findings using a plant-pollinator network, with data collected by Memmott (1999), from Ashton Court, a site in Bristol, UK. We will refer to this as the Ashton Court (AC) network. This is a well-sampled network (Bluthgen, Menzel & Bluthgen 2006) with interactions recorded over a short period of time (1 month). The AC network is highly resolved: all plants were identified to species ($P=25$) and many pollinators were identified to species level (and the remaining pollinators identified to morphotype: $A=79$). $M^{AC}$ contains 104 species, $E=299$, with connectance (proportion of realised interactions) of 0.151. Interactions in the AC network are weighted by the number of observations. The degree distribution of plant species is highly skewed as is often the case in plant-pollinator networks.

We also present results for two other networks to illustrate the generality of our model outcomes. These networks were collected in Shelfhanger (Sh), Norfolk, UK (Dicks, Corbet & Pywell, 2002); $M^{Sh}$ has $P=16$, $A=36$, $E=89$ and connectance 0.148; and in Ottawa, Canada (Small, 1976); $M^{Ot}$ has $P=13$, $A=34$, $E=141$ and connectance 0.319. Both networks have interactions weighted by the number of observations. We selected these networks because, like the AC network, they
describe northern hemisphere, temperate ecosystems, and have a similar size to the AC network, but differ in having lower and higher connectance respectively.

2.1 Model Development

We took as our starting point the extinction model of Memmott, Waser and Price (2004), who analysed the robustness of binary networks by making species of one type (in their case, pollinators) extinct in a random order, i.e. they used a random primary extinction sequence. From this, we developed two new extinction models that each include sub-sequences of plant extinctions that are determined by network structure. All three models (summarized in Fig. 2) can be applied both to weighted and to binary interaction data by introducing a threshold rule.

In this section, we first describe the features that are common to all our extinction models and then outline the distinctive features of each, highlighting the relationships between ours and previous extinction models.

2.2 Universal Model Features

Starting from the observed matrix $M$, a node of one guild (plants) is removed as a primary extinction. Extinctions result in the loss of interactions from $M$, monitored in the ‘reduced’ matrix $C$. The loss of interactions may, according to the rules of the particular model, result in the secondary loss of nodes of the other guild (pollinators). In our new models (see below) the rules admit the possibility of each secondary pollinator extinction giving rise to further knock-on plant extinction(s). These plant extinctions cannot be considered ‘primary’, but
will take their place in what we shall continue to refer to as a ‘primary extinction sequence’ of the $P$ plant species.

All models proceed until all plant nodes are removed and all species - plants and pollinators - are extinct. The robustness ($R$) of the network is calculated as

$$R = \frac{1}{AP} \sum_{p=0}^{P} a(p),$$

where $p$ is the number of plant species that have gone extinct (from 0 to $P$) and $a(p)$ is the number of pollinator species remaining in the network (from $A$ to 0). $R$ is the normalized ($0 < R < 1$) area under the curve of a graph of the proportion of plant nodes that have gone extinct against the proportion of surviving pollinator nodes (see Fig. 1a). Values of $R$ closer to 1 indicate higher ‘robustness’ of the network to primary extinctions (Burgos et al., 2007 and Albert, Jeong, & Barabási, 2000). We use $a(p)$ as our response variable for all models in order to facilitate comparisons, although we note other options are possible: e.g. Kaiser-Bunbury et al. (2010) used the sum of interaction weights $w(p)$.

The value of $R$ is dependent on the specific sequence of primary extinctions, so running many random extinction sequences will, for all our models, produce a frequency distribution of values of $R$ (Fig. 1b) which we denote $f(R)$.

A key model feature we adopted is a threshold rule for secondary extinctions, so that a node becomes extinction once it has lost a fraction $T$ or more of its observed interactions (binary $M$), or of its observed total interaction weight (weighted $M$). Clearly the value of $T$ that we choose is arbitrary. It must lie in the range $0 < T \leq 1$. [$T=0$ is a pathological case; all pollinator species become
extinct after the first primary plant extinction; $T = 1$ generates the extinction rule for most previous models (Dunne, Williams & Martinez, 2002; Memmott, Waser & Price, 2004; Kaiser-Bunbury et al., 2010), although Vieira introduced a node-specific threshold $\leq 1$. [1] We generated distributions of robustness $f(R)$ for a range of threshold values (0.1 to 1 in 0.1 intervals) for three observed plant-pollinator networks (Ashton Court, Shelfanger and Ottawa) to determine the effect of $T$ on the results of our extinction models. We then chose a threshold of $T = 0.5$ for the remainder of the paper: i.e. a secondary extinction occurs when a node has lost at least half of its interactions (binary $M$) or weights (weighted $M$). It should be noted that the ‘effective $T$’ ($T_{\text{eff}}$) could be greater than $T$; for example, with a binary network and $T = 0.5$, a node linked to 5 others would go extinct after losing 3 edges, giving an effective $T$ of $3/5 = 0.6$. Since most pollinators are observed visiting a relatively small number of plants, the difference between the specified and the ‘effective’ threshold can be noticeable and so we calculate the node-averaged $T_{\text{eff}}$ in all cases.

### 2.3 New extinction model features

We present three distinct models, which we denote: 1. Secondary Only (SO), 2. Deterministic Avalanche (DA) and 3. Random Walk (RW). Each model can be used with binary or weighted interaction data and is prefixed with ‘b’ or ‘w’ to indicate which.

**Model 1. Secondary Only model (bSO and wSO)**

In the Secondary Only model the order of primary plant extinctions is random. All pollinator extinctions are secondary and determined by the threshold rule.
There is no spread of extinctions beyond the secondary extinction of pollinators.

The method is as follows:

1. Select a random plant species (e) from those left in the network (matrix $M$) the first time, then subsequently matrix $C$ for primary extinction
2. Make pollinator species connected to $e$ extinct according to the threshold rule: if they have lost a proportion $\geq T$ of their original edges (bSO) or edge weights (wSO)
3. Count the number of pollinator species remaining, $a(p)$, in the updated network (matrix $C$)

Repeat steps 1 to 3 until there are no species remaining in the network. Then calculate $R$ according to equation 1.

In the special case $T=1$, the bSO and wSO models are identical to each other, and to the model described by Memmott, Waser and Price (2004). Kaiser-Bunbury et al. (2010) employed an adaptation to this special case ($T = 1$) but used the weight of remaining edges $w(p)$ as their response variable, which gives an approximation of the impact on pollinator abundance if pollinator abundance does not change until the extinction of the pollinator.

**Model 2. Deterministic Avalanche Model (bDA and wDA)**

In this model a randomly chosen primary (plant) extinction - a 'trigger' - may produce secondary extinctions (of pollinators) that themselves leave plant species with fewer than a fraction $T$ of their observed interactions. If this happens, there is an 'avalanche' of plant extinctions. During the avalanche the sequence of plant extinctions is not random, but is determined by network
structure. At the end of an avalanche a new, random, trigger is chosen. The method is as follows:

1. Select a random plant species \( e \) from those left in the network \( M \) the first time, subsequently \( C \) for primary extinction – this is a trigger

2.
   a. Make pollinator species connected to \( e \) extinct according to the threshold rule: if they have lost a proportion \( \geq T \) of their original edges (bDA) or edge weights (wDA)
   b. Count the number of pollinator species remaining, \( a(p) \), in \( C \)
   c. Make plant species (there may be more than 1) extinct according to the threshold rule as above
   d. Repeat steps 2a to 2c until there is no further spread of extinctions, then repeat from step 1 with a new trigger

Repeat steps 1 and 2 until there are no species remaining in the network.

Then calculate \( R \) according to equation 1.

Were \( T = 1 \) used here, step 2c would never result in tertiary plant extinctions and no avalanches would occur, so the DA and SO models would be identical. The 'stochastic co-extinction model' developed by Vieira and Almeida-Neto (2015) is a special case of our bDA model where the threshold is applied stochastically and is node specific; specifically, extinctions of nodes at our step 2c occur with probability \( = 1 - \frac{(remaining\ interactions)}{(interactions\ at\ start)} \). We adopt the term 'avalanche' for our spreading deterministic extinctions to differentiate them from the stochastic 'cascades' of Vieira and Almeida-Neto (2015), which occur once only, triggered by the first primary extinction.
Model 3. Random Walk model (bRW and wRW)

The RW model is similar to DA, in that a trigger can cause an avalanche of non-random plant extinctions. This time, the order of plant extinctions within an avalanche is determined by the structure of the plant-plant projection network (described by the $P \times P$ matrix $F$ which quantifies the number of pollinator species shared by each pair of plant species). The full method is as follows:

1. Select a random plant species ($e$) from those left in the network (the first time, subsequently $M$) for primary extinction
2. Create the plant-plant projection network $F$
3. Select the next plant extinction ($f$) from the neighbours of $e$ in $F$ with a probability proportional to edge weights.
4. Make pollinator species connected to $e$ extinct according to the threshold rule: if they have lost a proportion $\geq T$ of their original edges (bRW) or edge weights (wRW)
5. Count the number of pollinator species remaining, $a(p)$, in the updated network (matrix $C$)
6. Identify plant $f$ as the new $e$ and make it extinct
7. Loop through steps 2 to 6. If no neighbours exist in step 3, revert to step 1.

Repeat steps 1 to 7 until there are no species remaining in the network. Then calculate $R$ according to equation 1.
2.4 Natural extensions of our models

We have developed these three models for application to binary and weighted mutualistic bipartite networks and with a random order of primary plant extinctions (i.e. the selection of the next extinction in step 1 of Models 1, 2 and 3 is random). However, we note that these models can easily be modified to use ordered primary extinctions, where the choice of plant in step 1 of Models 1, 2 and 3 is according to a pre-determined rule (based on node degree, biological plant trait etc). The models can also be applied to bipartite networks with unidirectional dependencies (no feedback between the trophic levels, e.g. trophic or host-parasitoid interactions), though in that case avalanches cannot occur.

2.5 Comparison of robustness distributions from the three extinction models

The distribution \( f(R) \) generated from a single network \( M \) will depend on the model used and whether the data are weighted or binary. If there are \( P \) plant species in the network, there are \( P! \) distinct plant sequences. The SO models sample uniformly from these possibilities (i.e. all sequences are equally likely). The DA and RW models do not sample uniformly, because avalanches produce non-random sub-sequences determined by the structure of the network. Using binary and weighted versions of the Ashton Court (AC) network we generated 25,000 extinction sequences using each of the 3 models, in order to assess the effect of \( R \) on model choice. To create values of \( R \) that lie close to the theoretical maximum and minimum we ran bSO with plant extinctions in order of increasing and decreasing degree.
2.6 Assessing node and network-level determinants of variation in robustness

Having described the variation in robustness, we finally sought to assess the attributes which determine this variation under each model. We did this in two ways: by creating null versions of the AC network with defined characteristics, and by examining extinction rank (when a plant goes extinct in an extinction sequence). These calculations are designed to test the role of plant degree in determining the central tendency and spread of $f(R)$ in our three extinction models.

Generating null networks to test effect of degree distribution

To explore the effect of degree distribution $g(k)$ on robustness we created three exemplar null networks from the AC network (in binary form) with different degree distributions. Firstly, we calculated the expected (null) plant and pollinator degree distributions $g_E(k)$ from 10,000 random networks with the same number of plant species ($P = 25$), pollinator species ($A = 79$) and edges ($E = 299$) as the AC network. (Throughout we excluded networks with any disconnected nodes from further consideration.) We then generated 10,000 random networks from the AC matrix ($M_{AC}$) according to 3 randomisation protocols; 1) randomise all values in $M_{AC}$ to generate random plant and pollinator degree distributions; 2) randomise $M_{AC}$ by row only so that the pollinator degree distribution is preserved; 3) randomise $M_{AC}$ by column only so that the plant degree distribution is preserved. We selected, from each set of random networks, the exemplar network which had the best match to $g_E(k)$ for 1) both plants and pollinators, 2) only plants and 3) only pollinators. The best
match was assessed using the sum of the absolute difference between \( g_E(k) \) and the random network degree distribution for either plants, pollinators or both. The bSO, bDA and bRW models were run on the observed AC network and on these three exemplar random networks to produce 10,000 extinction sequences, and a corresponding distribution of robustness \( f(R) \) for each.

**Examining plant extinction rank in relation to \( R \)**

To explore whether (for example) high-degree plants tend to go extinct toward the beginning of a primary extinction sequence, we defined the position in a sequence when a plant became extinct as its extinction rank \( (r) \), \( 1 \leq r \leq P \). We ran each extinction model 25000 times, using binary and weighted versions of the AC dataset, and compared \( h(r) \), the distribution of extinction rank for each species generated by the simulations. From \( h(r) \) for each species we calculated the median extinction rank \( (r_m) \) and degree \( (k) \) and tested for correlation using the Spearman coefficient. We expected that, by definition, \( r \) would be equal across all species for the SO model but not for the DA or RW models, since avalanches and random walks will tend to select (or avoid) high-degree nodes preferentially.

**2.7 Testing on other networks**

We tested our models on the Shelfanger and Ottawa networks. For each network we generated 25,000 extinction sequences, using each of the 3 models, in binary and weighted form. We used a fixed threshold of \( T = 0.5 \) for all cases as we are
not directly comparing the networks, only seeking to confirm the generalities of the resulting $f(R)$ distributions.

3 RESULTS

3.1 Varying the value of the threshold for secondary extinctions

Median robustness $R_m$ increases non-linearly with $T$, and the least robust of our three networks at low $T$ becomes the most robust at high $T$ (Fig. 3a). However, this appears to be an artefact of the relationship between $T$ and $T_{eff}$, the node-averaged effective threshold (Fig. 3b), because $R_m$ increases linearly with $T_{eff}$ and the three networks are increasingly robust in order of increased connectance, as found by Dunne, Williams and Martinez (2002), at all values of $T_{eff}$ (Fig. 3c). The remainder of our results are presented for the AC network only (where $T_{eff} =$ 0.694 for our chosen $T =$ 0.5).

3.2 Robustness Distributions

The distributions $f(R)$ produced by each of the 3 models for binary and weighted data (Fig. 4) are all rather broad, suggesting a strong dependence of $R$ on the order in which plants are made extinct; the computed values span the range generated by primary extinction sequences in bSO with plants removed in increasing and decreasing order of degree ($R =$ 0.178 and $R =$ 0.812 respectively). The bSO model produces a relatively symmetrical $f(R)$ with a median $R_m =$ 0.473 and an inter-quartile range (IQR) 0.411 – 0.534. Using the bSO model as a baseline, the bDA model shifts $f(R)$ to the right (Figure 4b: $R_m =$ 0.512, IQR 0.439 – 0.588), inferring greater robustness, and bRW strongly shifts $f(R)$ to the left
(Figure 4c: $R_m = 0.369$, IQR 0.330 – 0.411) inferring lower robustness. Using weighted, not binary, interaction data increases the IQR of $f(R)$ in all cases.

### 3.3 Network randomisation test

Compared to the results of the binary extinction models for the AC network (Fig. 5a), we found that randomizing the degree distributions caused the distribution of robustness $f(R)$ to be narrower (Fig. 5b-d), and this was especially so when the plant degree distribution is randomised (5c and 5d). This confirms that the observed, highly skewed, plant degree distribution of the AC network drives the broad robustness distributions we generate for this network. Note though that the $R_m$ remain in the same order (RW<SO<DA) in every case, showing the consistency of effect from these models.

### 3.4 Extinction rank of plant species, and the effect on $R$

Plant degree is a predictor of the plant’s extinction rank in the DA and RW models. In the SO models, the expected rank is constant for all plant species, irrespective of degree, because the extinction sequence is entirely random. In contrast, the observed extinction ranks of two example plant species from the DA and RW models are clearly skewed (shown in the insets of Fig. 6). In the DA models median extinction rank is positively correlated with plant degree (bDA: $\rho = +0.803$, $P < 0.0001$; wDA: $\rho = +0.464$, $P = 0.02$). For the RW models, $r_m$ is negatively correlated with $k$ (bRW: $\rho = −0.654$, $P = 0.0004$; wRW: $\rho = −0.723$, $P < 0.0001$). In other words, for the DA models, well-connected plants are resistant to extinction; the model preferentially prunes the low degree plants and network
robustness is high compared to the SO models (Fig. 4b cf. Fig. 4a). In contrast, in the RW models plants with high degree are more vulnerable to extinction (the model preferentially ‘homes in’ on well-connected plants) which results in overall lower robustness of the network (Fig. 4c cf Fig. 4a). Clearly the apparent robustness of a network is dependent on the model used to incorporate feedback and structurally correlated co-extinctions.

4. DISCUSSION

The robustness of an ecological network is a valuable metric, because from it we can learn several things and it provides a quantitative metric for describing the vulnerability of an ecological community to hypothetical extinction scenarios, i.e. sequences of primary extinctions. Based on this we can compare the vulnerability of different networks to these scenarios and determine properties of the environment or of the network that correlate to robustness. However, robustness can also be used to say something about the network itself, in which case the breadth of a robustness distribution is important.

Extinction models that calculate robustness have been around for a long time but until recently ecologists have used simple versions of these models (see Fig. 1). Here, we have created a framework of extinction models, building on those of Memmott, Waser and Price (2004), Kaiser-Bunbury et al. (2010) and Vieira & Almedia-Neto (2015), and applied it here to plant-pollinator networks.

Importantly we have used an extinction threshold of less than one, i.e. pollinators can go extinct before all their plants go extinct. This addition has an ecological motivation - pollinators may decline to extinction due to declining food resources - and adds greatly to the flexibility of the model. Having $T<1$
allows us to create weighted versions of our models and permits the potential for feedback and, hence, avalanches of extinctions cascading between the trophic levels (as first shown by Vieira and Almedia-Neto, 2015).

All of our extinction models, in binary and weighted form produce a broad distribution of robustness values $f(R)$ for every network we analysed, indicating that there is some aspect of the structure of the network which causes this variation. We found this to be the degree distribution of the nodes, specifically the plant degree distribution. Plant pollinator networks tend to have fewer plant species than pollinator species ($P < A$), so the potential for a skewed plant degree distribution in increased, making it more influential on robustness, as indeed was the case in our test network (Memmott 1999).

Though ‘robustness’ has in the past been used to suggest priorities for conservation or management (Pocock, Memmott and Evans. 2012, Devoto et al. 2012), extinction models are not an attempt to predict precisely how an ecosystem would collapse. They do, nonetheless, offer a means to quantify and compare the structure of ecological networks, although to do this we need to ensure we are comparing like-for-like. We found that the median robustness $R_m$ is a linear function of (effective) threshold $T_{eff}$ and the effect is consistent across networks (Fig. 3). However, $T_{eff}$ has a non-linear relationship with the selected threshold value $T$, so if our models are to be used in the future to compare the robustness of ecological networks, it is important to note that $T_{eff}$, not $T$, is the appropriate parameter to ensure comparability.

Plant-pollinator communities are increasingly described with weighted interactions and so our weighted extinction models for robustness are a valuable advance. We found (Fig. 4) that introducing weighted interactions has the effect
of amplifying the outcomes observed for binary data; the inter-quartile range of
the robustness distribution $f(R)$ increases in all models for weighted networks,
and the shifts in median robustness for DA and RW compared to SO are larger.
These effects occur because weights tend to increase the skew of the plant
degree distribution because high degree species accumulate high weights and
low degree species only gain a small fraction of the overall weight in the
network. The double-peaked distribution seen for the weighted DA model can be
explained by competing effects: the model preferentially samples extinction
sequences that have long avalanches and higher $R$ values as well as those with
shorter avalanches and lower robustness, so the paucity of sequences with
intermediate $R$ values creates the two peaks. The exaggeration of effects in $f(R)$
highlights the importance of including interaction weights in robustness
analysis, and in exploring all of the distribution $f(R)$, not just its central tendency.
There are different ways in which feedback between trophic levels can be
applied and we developed two illustrative models: the Deterministic Avalanche
(DA) and the Random Walk (RW) models. These models (and others like the
cascade model developed by Vieira and Almedia-Neto, 2005) may appear to be
generating new outcomes but in reality, they simply produce a non-random
sample of robustness values from those generated by a simple SO model. The
Ashton Court dataset generated a huge range of $R$ values, all of which can be
realised in the Secondary Only models. The DA and RW models preferentially
sample extinction sequences to produce skewed subsets of the SO outcomes (the
$P!$ extinction sequences are not all equally likely, and some will be impossible).
The Deterministic Avalanche Model preferentially samples nodes that are 1 step
away from each other in the network and extinctions can ‘fan out’ from each
trigger (a randomly selected plant extinction). This mechanism can be likened to a ‘breadth first’ search on the network (see for example, Kolaczyk, 2009). On the other hand, the Random Walk model follows a path though the plant projection network $E$ away from trigger in the manner of a “depth first” search (Kolaczyk, 2009).

In ecological terms we can imagine the breadth first approach (DA models) as modelling the gradual collapse of the community along mutualistic dependencies. This is based on the assumption that the mutualistic relationship between plants and pollinators creates the potential for extinctions to travel in both directions between trophic levels (e.g. the presence and/or persistence of pollinators depends on flowers, and the presence and/or persistence of plants depends on pollinators). Avalanches in the DA model are only possible where dependencies are bi-directional, which can be perfect mutualism (each interaction is fully bi-directional) or it can represent a mixture of directed interactions (e.g. mutualism, nectar robbers, pollinator detection). The depth first approach (RW models) can be exemplified as a scenario where a plant disease is spread through the community by visiting pollinators, or a pollinator disease is spread through shared floral resources; a phenomenon observed by McMahon et al. (2015). The Secondary Only models are ecologically applicable when there is a uni-directional dependence in the interactions (e.g. presence of parasites depend upon hosts, but not vice versa) although, of course, they can be applied to mutualistic networks if dependence is assumed to be uni-directional (e.g. persistence of pollinators depends on flowers, but plants do not depend on pollinators).
Overall, the SO, DA and RW models can be used to examine the effect of a range of ecological extinction scenarios. There are many possible applications of the SO model. The DA and RW models are more specific in their application and notably produce opposite effects on the distribution of robustness for a given network. Therefore, this highlights the importance of selecting an appropriate model for the ecological circumstances. We expect that there will be future developments to refine these models to usefully predict the outcomes of possible ecological scenarios.

All of these extinction models are designed to be applied to real network data. Therefore, it is vital to consider the quality and reliability of the data being used. Pollination networks vary hugely in sampling method, period of collection and taxonomic resolution. We caution against comparing the outcomes of extinction models across multiple networks that have come from different sources without consideration of the data, methods and specific question. For example, CaraDonna et al. (2017) highlight the potential pitfalls of assuming that a static network, comprising all accumulated interaction data, is an appropriate representation of a community. Further work in understanding temporal variation and the description of fully-resolved plant-pollinator networks are key to improving the utility of extinction models.

We hope that by improving our understanding of extinction models at a fundamental network level, and by setting out different areas of model extension, our work will guide future developments in the analysis of the vulnerability of ecosystems to environmental change.

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**AUTHORS’ CONTRIBUTIONS**

All authors designed the methodology, discussed the results and commented on the manuscript at all stages. MSB coded the models and analysed the data with technical advice and support from RJ and MJOP. MSB drafted the manuscript. All authors contributed critically to the drafts and gave final approval for publication.

**DATA ACCESSIBILITY**

This work has used the Web of Life database to access ecological network data: [www.web-of-life.es](http://www.web-of-life.es). Model code will be deposited on the University of Bath Research Data Archive.

**REFERENCES**


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Figure 1: The output of an extinction model. (a) For a single extinction sequence the number of surviving pollinator nodes $a$ reduces as the number of plant nodes made extinct, $p$, increases. There are no surviving nodes at the end of the sequence. Robustness ($R$) = 0.5504 computed as the area under $a(p)$, divided by the area of the rectangle, $AP$. (b) In all our extinction models the value of $R$ depends on the order in which plants are made extinct, so many simulations are run to produce a distribution of robustness values $f(R)$. 
Figure 2: Framework of extinction models, with those used in this paper highlighted in colour. All models start from an observed mutualistic bipartite network $M$ that can be binary (prefix b) or weighted (w). For binary data the threshold $T$ is applied to the number of edges; for weighted data it is applied to the weights. The models are split into those that produce entirely random primary extinction sequences: Secondary Only (SO), and those that introduce other methods for determining extinction sequence: Deterministic Avalanche (DA) and Random Walk (RW). (i-iv) indicate previous studies that represent special cases of the models in the framework where $T = 1$: i) Dunne, William and Martinez, 2002; ii) Memmott, Waser and Price, 2004; iii) Kaiser-Bunbury et al. 2010, and where $T$ is applied stochastically and extinctions can ‘stochastically cascade’: iv) Vieira and Almeida-Neto 2015.
Figure 3: The relationship between extinction threshold ($T$), effective threshold ($T_{\text{eff}}$) and median robustness ($R_m$) for the three plant-pollinator networks Ashton Court (triangles), Shelfhanger (squares) and Ottawa (circles) using the bSO model. Variation of (a) $R_m$ with $T$, (b) $T_{\text{eff}}$ with $T$, and (c) $R_m$ with $T_{\text{eff}}$. 
Figure 4: The distribution of robustness $f(R)$ for the Ashton Court network, in binary (left column) and weighted (right column) form, generated by the three extinction models: (a) Secondary only (SO), (b) Deterministic Avalanche (DA) and (c) Random walk (RW). Median robustness $R_m$ for each distribution is indicated by the solid vertical line. Dashed grey lines indicate $R$ values for the bSO model generated by removing plant species in increasing ($R=0.178$) and decreasing ($R=0.812$) degree order.
Figure 5: The effect of node degree distribution on robustness distribution $f(R)$ for (a) the binary Ashton Court network and (b-d) randomised networks generated by the protocols described in section 2.6. Left column (A): pollinator degree distribution (grey - observed; purple – randomised); central column (P): plant degree distributions; right column: summaries of $f(R)$ from the bSO, bDA and bRW extinction models. [Box-plots, with central lines showing median, boxes showing inter-quartile range, and whiskers showing the 95% (2.5–97.5%) interval].
Figure 6: Variation of median extinction rank $r_m$ with degree $(k)$ for all 25 plant species in the Ashton court network for the three extinction models (SO: red, DA: blue and RW: green) and for (a) binary data and (b) weighted data. Smoothed lines are plotted as guides; Spearman's rank correlation show that all these associations are significant: positive for DA (blue) and negative for RW (green). Insets illustrate the extinction rank distribution $h(r)$ for two plant species (*Lathyrus pratensis* $(k=2)$ on the left and *Daucus carota* $(k=46)$ on the right) produced by the DA (blue) and RW (green) models. The corresponding median rank points are indicated with an arrow.