Abrupt transitions and its indicators in mutualistic meta-networks: effects of network topology, size of metacommunities and species dispersal

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6 Abstract

Gradual changes in the environment could cause dynamical ecological networks to abruptly shift from one state to an alternative state. When this happens ecosystem functions and services provided by ecological networks gets disrupted. We, however, know very little of how the topology of such interaction networks can play a role in transitions of ecological networks at a spatial scale. In the event of such unwanted transitions, little is known how statistical metrics used to inform such impending transitions, measured at the species-level or at the community-level could relate to network architecture and the scale of spatial interactions. Here, using fifty-six empirical plant-pollinator networks in a spatial setting, I evaluated the impact of network topology and spatial scale of species interactions on abrupt transitions, and on statistical metrics used as predictors to forecast such abrupt transitions. Using generalized Lotka-Volttera equations in a meta-network framework, I show that species dispersal rate and the size of the metacommunity can impact when an abrupt transition can occur. In addition, forecasting such unwanted abrupt transitions of meta-networks using statistical metrics

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of instability was also consequently dependent on the topology of the network, species dispersal rate, and the size of the metacommunity. The results indicated that the plant-pollinator meta-networks that could exhibit stronger statistical signals before collapse than others was dependent on their network architecture and on the spatial scale of species interactions.

keywords: meta-networks, mutualistic networks, abrupt transitions, dispersal, early warning
 signals

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#### 29 Introduction

Ecological systems could abruptly shift from one state to another in response to gradual changes in environmental conditions. Such abrupt transition occurs when environmental 31 conditions cross a specific threshold, the threshold being commonly known as tipping points 32 (Dakos et al., 2018; Scheffer et al., 1993). Tipping points and abrupt transitions are generally observed in ecological systems governed by positive feedback loops. Abrupt transitions can occur in systems ranging from acquatic systems such as algae and macrophyptes (Dakos et al., 2018), mutualistic ecological networks (Baruah, 2022; Dakos and Bascompte, 2014; Lever et al., 2014), and populations exhibiting allee thresholds (Hilker, 2010). When such abrupt transitions occurs, the equilibrium stable state of the system jumps to another state where loss of beneficial ecosytem functions and services could occur (Dunne and Williams, 2009; Hutchings and Reynolds, 2004; Scheffer et al., 2012). Abrupt transitions, thus, can cause long-term radical changes to ecosystems. However, at the scale of multiple communities connected by species dispersal, very few studies have explored how local network architecture and spatial scale of interactions could impact the timing of abrupt transitions 44 Mutualistic communities are examples of communities that exhibit positive feedback loops in networks of interactions between two groups of species (Bascompte et al., 2003; Dakos and Bascompte, 2014; Kéfi et al., 2016). While negative interactions such as intraspecific competition could have stabilising effects, positive interactions such as those observed in plant-pollinator or mutualistic networks could be destabilising and could result in the 49 presence of alternative stable states (Baruah, 2022). As such due to the presence of such alternative stable states, mutualistic communities can transition from a stable state to an alternative state as environmental change causes a change in the system's feedback strength (Lever et al., 2020). The spatial context of such transition has somewhat remain unexplored, particularly on the impact of topological and architectural aspects of such systems such as

network size, nestedness, or connectance. Previous study on spatial mutualistic systems

has suggested that the presence of transitions can be modulated by shift in strength in species phenology or habitat destruction (Fortuna and Bascompte, 2006). Habitat destruction increases the chances of abrupt transition which rewires local networks (Revilla et al., 2015). In addition, with strong mutualistic interactions the amount of habitats required for 59 persistence of all species then depends on the range of species dispersal (Prakash and Roos, 2004). However, it is unknown whether topological features of mutualistic networks such as network size or connectance could influence the timing of occurrence of abrupt transition in 62 response to gradual changes in environment. For instance, could larger networks collapse earlier in comparison to smaller networks? Rarely, ecological communities occur in isolation. More often, ecological communities oc-65 cur in habitat patches across a larger network of patches that is connected by dispersal, which is commonly known as a metacommunity. Metacommunity concept has gained much attention over the last decade and empirical and theoretical studies have provided with an understanding of how local and regional processes work to maintain diversity both at local and regional scale (Limberger et al., 2019; Loreau et al., 2003). Species diversity can stabilize the dynamics of local communities. This stabilization occurs due to differ-71 ent asynchronous responses of species over time to temporal changes in the environment. 72 Such stabilization due to increases in number of species is generally known as species insurance effects (Heino et al., 2015; Loreau et al., 2003; Shoemaker and Melbourne, 2016). Species insurance can occur due to interspecific competition (Lehman and Tilman, 2000), or 75 when species response to biotic environment differs from other species (Loreau et al., 2003). Spatial insurance in metacommunities arises when local communities exhibit asynchronous dynamics. This happens when species dispersal rate is limited or when species composition 78 across local communities vary considerably thereby leading to spatial heterogeniety (Walter et al., 2017; Wang and Loreau, 2014). Species dispersal from habitat patches could potentially rescue local communities from collapses as environment changes. However, the role of 81 species dispersal on timing of transition of meta-networks remains unknown. For instance, does spatial scale of species interactions and the spread of such ecological networks (number 84 of habitat patches) determine whether network collapses occur earlier or later?

Local and regional scale extinctions could occur not only due to local processes such as predation or competition but also due to large scale external disturbances (Cunillera-Montcusí et al., 2021). Large scale disturbances akin to changes in climate could impact species interactions not only locally but also across communities connected in space (Revilla et al., 2015; Thompson and Gonzalez, 2017). In mutualistic communities, changes in the environment could lead to weakening of interactions between species such as phenological mismatches thereby altering local structure of communities (Forrest, 2015; Kudo and Cooper, 2019; Morton and Rafferty, 2017; Renner and Zohner, 2018; Visser and Gienapp, 2019). However, such species mismatches could be rescued when similar communities are 93 accessible to species in a mosaic of communities connected by species dispersal. It is, however, unknown whether such global changes in the environment interacts with local network topological properties to mitigate drastic change. Increases in phenological mismatch could be further compounded by habitat destruction and could lead to substantial changes in 97 network architecture (Revilla et al., 2015). In addition, whether the spatial scale (size of the metacommunity) of such changes matters in the occurrence of transition is somewhat gg not known. 100

There has been statistical tools developed to inform impending transitions that could occur 101 as environment gradually changes (Scheffer et al., 2012). Abrupt transitions could occur 102 when changes in the environment crosses a certain threshold that pushes the ecological sys-103 tem towards another alternative state where ecosystem functions could be lost permanently. 104 However, there are statistical tools that been developed to forecast such impending tran-105 sitions which are commonly known as "early warning signals." Commonly used signals are 106 temporal autocorrelation and variance that could derived using a sliding window approach 107 (see Dakos et al. (2012a) for details) from state based temporal data such as abundance 108 or biomass. However, the utility of such signals are dependent on a host of factors that 109 includes sampling requirements (Arkilanian et al., 2020), data quality (Clements et al.,

2015), eco-evolutionary factors (Baruah et al., 2021, 2020) and type of species interactions 111 (Baruah et al., 2022; Dakos, 2017; Patterson et al., 2021). One important challenge is to 112 test the utility of such signals in a multispecies context embedded within a spatial scale of 113 species interactions. This is especially relevant as multispecies communities rarely occur in 114 isolation and are generally connected by dispersal among habitat patches. The detection of 115 such signals becomes even more challenging as dynamics of such communities are inherently 116 linked to their topological features which also directly impacts the occurrence of transitions 117 (Baruah, 2022; Dakos and Bascompte, 2014). 118

Here, using fifty-six empirical plant-pollinator networks in a spatial context collated from 119 web-of-life.es database, I explore the effects of topological network features such as network 120 size, connectance, or nestedness on timing of transitions and on indicators of temporal 121 and spatial resilience. Using generalized Lotka-Voltterra equations, I model the ecologi-122 cal dynamics of spatially-explicit mutualistic meta-networks to global changes in strength 123 in mutualistic interactions. Specifically, using different metacommunity sizes, I show that 124 timing of abrupt transition depends not only how species disperse across habitat patches, 125 but also on local network topological predictors such as network size and connectance of 126 the network. Furthermore, the threshold at which a species could abruptly transition is 127 influenced by the degree of the species, rate of dispersal and also by the size of the meta-128 community. In addition, how large the metacommunity was also significantly played a role 129 on how early an abrupt transition might occur. Furthermore, when such global transition 130 of mutualistic metacommunities occur, predictability with temporal and spatial resilience 131 indicators also depends on the topological network features and on the rate of dispersal of 132 species. These results argue the importance of understanding the dynamics of communi-133 ties from a spatial perspective and highlights the importance of network architecture on 134 biodiversity maintenance.

## 136 Methods

Using www.web-of-life.es database, I collated 56 empirical plant-pollinator networks (see table S1 for details and the references). These empirical networks were set up in a spatially-138 explicit landscape of different sizes of two, five, ten and twenty habitat patches that deter-139 mined the spatial scale of mutualistic interactions. This spatial scale determined to what extent plant-pollinator interactions were impacted when global changes occur at the scale 141 of the metacommunity. For instance, when changes in climate that drives changes in phe-142 nological interactions occur, does spatial scale of mutualistic interactions matter in delaying an abrupt transition to collapse? 144 These spatially explicit landscape were set up in a two dimensional landscape (Grilli et al., 145 2015). All habitat patches were connected. Following this, I model the ecological dynamics 146

$$\frac{dA_i^k}{dt} = A_i (r_i^k - \sum_j a_{ij}^k A_j + \sum_l \frac{m_{li} \gamma_0 P_l}{1 + h \sum_x m_{xi} \gamma_0 P_x}) + M_i^k - a A_i^k$$
 (1)

Similarly, the dynamics of plants on a spatially explicit metacommunity can be written as:

of mutualistic interactions using generalized Lotka-Volterra equations:

$$\frac{dP_i^k}{dt} = P_i(r_i^k - \sum_j a_{ij}^k P_j + \sum_l \frac{m_{li} \gamma_0 A_l}{1 + h \sum_x m_{xi} \gamma_0 A_x}) + M_i^k - a P_i^k$$
 (2)

where  $A_i^k$ ,  $P_i^k$  represents pollinator and plant abundance for habitat patch k,  $r_i^k$  is the species specific growth rate independent of mutualistic interactions at patch k,  $a_{ij}^k$  represents interspecific competition within each guild of species at patch k,  $\gamma_0$  represents the average mutualistic interaction strength when a plant and a pollinator interacts, with  $m_{li}$  determining the network structure and is either 0 or 1 depending on whether an interaction exist between a plant and a pollinator, h is the handling time which we fixed at 0.15,  $M_i^k$  is the density of species that arrives from all the patches in the metacommunity, and finally a fraction of individuals emigrates from a habitat patch k at the rate a.

In our model simulations, we ensured that mutualism was obligate without any loss of gener-157 ality. This meant that growth rates of species  $r_i^k$  was negative for both the guilds of species 158 and species persistance totally dependent on mutualistic interactions between species. To 159 do that we randomly sampled growth rates  $r_i^k$  from a random uniform distribution within 160 the range of U[-0.05, 0.1] for both plants and the pollinators. In addition, we ensured that 161 intraspecific competition within each guild of species was strictly stronger than interspe-162 cific competition. For that we ensured that  $a_{ii} = 1$  and we sampled  $a_{ij}, i \neq j$  from random 163 uniform distribution ranging from U[0.01, 0.05]. When ensuring that intraspecific compe-164 tition to be stronger than interspecific competition, feasibility of mutualistic community becomes easier provided strength in mutualistic interactions  $\gamma_0$  was high (Barabás et al., 166 2017; Baruah, 2022). Our goal was to finally assess how network properties come into play 167 at a spatial scale in mitigating the timing of occurrence of abrupt collapses as we change 168 the strength in mutualistic interactions. 169

Finally, dispersal among patches was constrained by the spatial scale and as well as distance among patches. Species dispersal among patches decreased exponentially as the distance among patches increased. Specifically, dispersal of species i from patch k can be written as (Thompson and Gonzalez, 2017),

$$M_{i}^{k} = a \sum_{j \neq k}^{n} \frac{e^{-yd_{jk}}}{\sum_{f \neq k}^{n} e^{-yd_{jk}}} N_{i}^{j}$$

Following Thompson and Gonzalez (2017), more than one dispersal route can be taken during a particular time step,  $d_{jk}$  is the distance between patch j and k and y controls the rate at which this distance decreases which we fix at 0.5. Fixing it at 0.5 gives global dispersal, which means that as metacommunity sizes become larger species dispersal does not necessarily gets constrained locally and species could in principle disperse long distance. However, the rate at which they disperse would still depend on distance between patches. Finally, a gives the average rate at which species disperse. Here, when we vary rate of dispersal we ensure that species dispersal remains same across guilds of species although

in nature this might be species-specific. Here, three rates of species dispersal were used, a: 0, 0.05, 0.15.

### Collapse of spatial mutualistic networks

Nestedness measured as NODF ranged from as low as 0 to as high as 0.7, while connectance ranged from . Network size also ranged from as low as 8 to as high as 68.

By gradually decreasing the average mutualistic strength globally, mutualistic networks were 187 forced to collapse. As mutualism among guilds of species was obligate, decreasing average 188 mutualistic strength,  $\gamma_0$ , among species would lead to collapses of species. At a specific 189 mutualistic strength (commonly known as threshold strength or tipping point) collapse of 190 the entire mutualistic network would occur. Collapse of mutualistic networks thus was done 191 by gradually decreasing  $\gamma_0$  from 7 to 0 in steps of 0.25. Globally, across habitat patches 192 this scenario could be linked to changes in the climate such that phenological interaction 193 among plants and pollinators would decrease gradually. For each value of  $\gamma_0$  I simulated the dynamics of the whole metacommunity till it reached ecological equilibrium. Usually, 195 fluctuations in species density stabilize at around 1000 time points. I discarded the initial 196 transient dynamics and estimated equilibrium plant and animal biomass from the last 1000 197 time points. The extinction threshold of species in such mutualistic networks were fixed at 198  $10^{-4}$ . As the strength of mutualistic interactions decreased, loss of species occurred until 199 the entire meta-mutualistic network collapsed. Next, I estimated the slope of the decline 200 of total community abundance as change in total community abundance per change in co-evolutionary mutualistic strength in a habitat patch as 202

$$N_{\gamma_0(t+1)}-N_{\gamma_0(t)}.$$

Next, I determined the point of transition or the mututalistic strength at which a mutualistic network in a habitat patch k transitioned to a collapse state when the total abundance of the

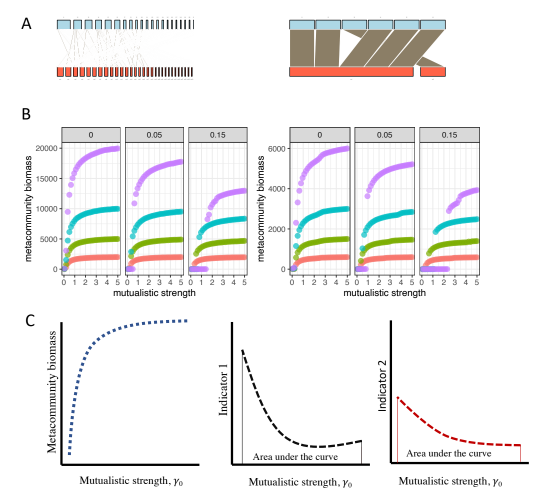
network fell below 80 percent of its equilibrium abundance. Once the mutualistic strength at which a network collapsed in a habitat patch k was determined, I evaluated the relationship of the point of transition with network topology such as connectence and network size, and how such a relationship was influenced by spatial scale of interactions.

# Predictors of temporal and spatial mutualistic network collapses:

I estimated a host of spatial and temporal indicators of collapses at the community level and 211 at the species level. For each mutualistic strength, I estimated spatial variability, regional 212 metacommunity variability, local temporal patch variability. We quantified metacommunity 213 biomass as the sum of all species density across all patches in the metacommunity. I 214 also estimated metacommunity variability defined as the variability in abundance at the 215 metacommunity level (see Wang and Loreau (2014) and Wang et al. (2019) for details) -216  $\frac{\sqrt{(\sum_{ij} w_{ij}}}{N}$ , where  $w_{ij}$  is the covariance matrix of community biomass  $N_i(t)$  at patch i and 217  $\bar{N}$  is the temporal mean of the total metacommunity biomass. Temporal variation or alpha 218 variability was estimated as  $\frac{\sqrt(\sum_i w_{ii}}{\tilde{N}}$  and spatial beta variability was quantified as as the 219 variability at spatial level  $\frac{\sum_{ij} w_{ij}}{(\sum_i \sqrt{w_{ii}})^2}$  (Wang and Loreau, 2014) for each levels of average 220 dispersal rate, metacommunity size, and for each mutualistic threshold strength. 221 At the species level, I estimated standard deviation and temporal autocorrelation coefficient 222 of equilibrium species abundance for each level of mutualistic strength, for three different 223 rates of species dispersal and metacommunity sizes. Temporal autocorrelation coefficient 224 at first-lag and standard deviation are the classic phenomenological early warning signals 225 that are suggested to be useful in forecasting critical transitions. Temporal autocorrelation 226 coefficient at first-lag is measured as,  $y(N_{t+1}) = \alpha y(N_t) + \sigma \epsilon_n$ , where  $\alpha$  is the autocorrelation 227 coefficient of the first-order autoregressive (AR(1)) process.  $\alpha$  is close to 1 for a red-noise and close to 0 for white noise process.  $\alpha$  close to 1 would indicate that the temporal 229

abundance dynamics of a species is highly correlated and 0 would mean uncorrelated. High
correlation would indicate that species is closer to a tipping point.

I wanted to evaluate whether statistical metrics measured at the species level or at the level 232 of the community could increase as strength in mutualistic interaction  $\gamma_0$  decrease, indicat-233 ing an impending critical transition. So for each level of changes in the mutualistic strength of interaction  $\gamma_0$  , temporal autocorrelation and standard deviation was measured at the 235 species level, metacommunity variability was measured at the level of the metacommunity, 236 patch variability was measured at the level of a patch, and spatial variability across habitat 237 patches were estimated. To compare whether these metrics performed well, I estimated 238 the area under the curve (AUC). To elaborate on this, as mutualistic strength decreases, 239 one expectation from these statistical metrics is that that the values of these statistical 240 metrics will increase. For instance, as shown in figure 1C, indicator 1 increases steeply as  $\gamma_0$  decreases, and whether an indicator performs better could be evaluated by calculating 242 the area under the curve of the metric. Stronger the increase, larger would be the AUC of 243 the statistical metric in question.



Indicator 1 has a larger <u>area under the curve</u> and increases steeply as mutualistic strength decreases and metacommunity biomass collapses than indicator 2.

Figure 1: (B) Example equilibrium metacommunity abundance as mutualistic strength is decreased gradually for (A) two different networks which vary in the total number of species for different sizes of metacommunity and rate of species dispersal. (Note that in the depiction of mutualistic networks the line thickness describing interactions between plants and animals decreases as network size increases in order to accommodate the increasing number of interactions). C) As metacommunity biomass collapses due to decrease in mutualistic strength, strength in statistical indicator 1 and statistical indicator 2 increases. However, indicator 1 steeply increases in comparison to indicator 2 and this can be compared by calcuating the area under the curve of indicator 1 and indicator 2. Clearly in this case, indicator 1 is better as its area under its curve is larger than indicator 2.

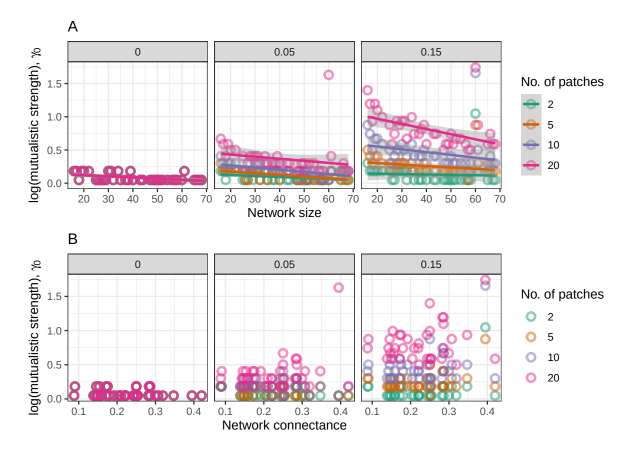


Figure 2: Relationship between the threshold mutualistic strength at which networks collapsed and structural properties of the network namely network size and connectance Note that the y-axis is normalised to have maximum value of 1. A) Network size had a significant impact on the strength at which netowks collapsed modulated mainly by the number of habitat patches. Larger metacommunities collapsed at a higher threshold mutualistic strength than smaller metacommunities. This indicates that smaller metacommunities were able to persist at lower mutualistic threshold strength whereas larger metacommunities collapsed much earlier even when mutualistic strength was high. In addition, size of the network further aggravated at which networks collapsed. Smaller networks collapsed at lower mutualistic strength than larger networks. B) Nestedness didnt have a significant impact on the mutualistic strength at which networks collapsed across a range of dispersal rates and metacommunity sizes. Mutualistic strength was normalised to have values between. Lines and ninety-five percent CI are fit from generalized linear model with quasibinomial error distribution with average threshold strength as response variable, network size/nestedness, number of habitat patches and rate of dispersal as predictor variables.

#### 245 Results

#### Point of transition and network properties

Results indicated that the mutualistic strength at which networks collapsed were determined 247 by network size, species dispersal rate, and the size of the metacommunity. Particularly, the relationship between network size and the strength at which networks collapsed becomes 249 negative at the highest dispersal rate modulated by the size of the metacommunity (Fig. 250 2A). When dispersal rate was zero, the size of metacommunity on the mutualistic strength 251 at which network collapsed remained unaffected. However, at the highest dispersal rate, 252 smaller networks collapsed much earlier than larger networks which became more evident at 253 larger metacommunity sizes. Connectance or nestedness didn't have any significant impacts 254 on the threshold at which mutualistic networks collapsed. This indicated that higher species dispersal might be detrimental as networks could collapse much earlier, particularly when 256 metacommunity sizes were larger. 257

## Indicators of transition and network properties

In an example figure of a mutualistic network (see fig. S1), I show that as metacommunity 259 biomass collapses, indicators such as patch variability, spatial variability, metacommunity 260 variability increases whereas indicators at the species level did not exhibit strong increases as 261 strength in mutualistic interaction decreased. To quantify the strength of such an increased, 262 I measure (as detailed in the methods section) the area under the curve of each of these 263 indicators. Such a measure of AUC would indicate which statistical metric has the strongest increase as the metacommunity collapsed. 265 Results indicated that when species dispersal was zero, size of mutualistic network had no impact on metacommunity variability. But at higher levels of species dispersal, the relation-267 ship between AUC of metacommunity variability and network size becomes negative and 268

was modulated by the metacommunity size (Fig 3A). This indicated that smaller networks

at larger metacommunities exhibited stronger metacommunity variability before an abrupt collapse. As metacommunity size decreases, smaller networks exhibited as strong a signal before collapse as larger networks particularly when there were only two habitat patches. Thus, metacommunity variability will increase and could forecast an abrupt transition provided networks are smaller and metacommunity size is larger. Since, network connectance was negatively correlated with network size, the above result remains similar except that the relationship between AUC of metacommunity variability and connectance becomes positive (Fig. S2).

In case of spatial variability, when dispersal rate was zero, smaller networks exhibited stronger spatial variability before collapse than larger networks. This result was consistent across metacommunity sizes. But as dispersal increased, which networks exhibited stronger spatial variability before collapse was dependent on the size of the metacommunity. Generally, the relationship between AUC of spatial variability and network size was negative and dependent on the size of the metacommunity. Smaller mutualistic networks embedded in larger metacommunities exhibited stronger spatial variability before collapse than smaller networks in smaller metacommunity sizes (Fig. 3B).

Patch variability also known as alpha variability was impacted by size of mutualistic networks and less impacted by species dispersal and number of habitat patches i.e. metacommunity size (Fig. 3C). Particularly, larger networks had larger AUC which indicates that
such networks exhibited stronger patch variability before an abrupt collapse than smaller
networks. At the highest species dispersal, metacommunity size did impact had a slight
impact on AUC of patch variability.

The strength of mutualistic interaction at which species collapsed as the meta-network collapsed was dependent on their degree of interactions. Species which had higher number of interactions i.e., higher degree collapsed later (at a much lower strength of mutualistic strength). This was similar across rate of species dispersal and size of the metacommunity. Strength in warning signals such as standard deviation measured at the species-level

was related to species degree, rate of species dispersal and less related to the size of the 297 metacommunity. Strength of SD, measured as the AUC of species SD, increased as species 298 degree increased indicating that species which have a larger number of unique interactions 299 will display stronger increases in SD as the network collapses due to changes in the envi-300 ronment that weakens mutualistic interactions. Strength in autocorrelation, however, did 301 not related to species degree or species dispersal. In addition, autocorrelation at first-lag, 302 or standard deviation measured at the species level was also not impacted by topological 303 properties or species dispersal or size of the metacommunity (Fig. 4A). However, AUC of 304 standard deviation measured at the species level was impacted by topological properties or species dispersal or the size of the metacommunity (Fig. S5). 306

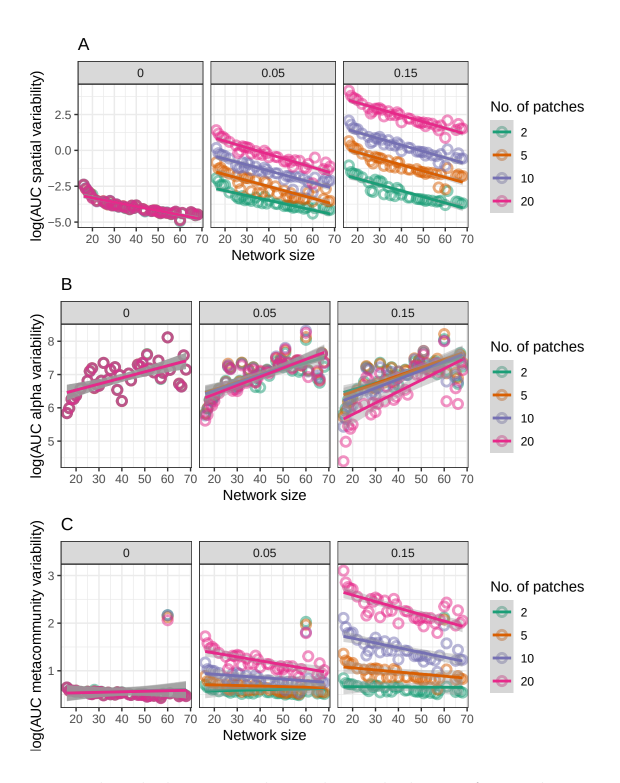


Figure 3: Relationship between network size and area under the curve of statistical metrics such as A) spatial variability, B) Alpha variability, and C) metacommunity variability for three levels of species dispersal (0,0.05,0.15) and four metacommunity sizes (2, 5, 10, 20). Note that the y-axis in all these subplots indicate the strength of increases in these indicators (measured as the AUC of the metric, see figure 1C for description) as a mutualistic meta-network collapsed. Larger AUC would indicated stronger increases as a mutualistic meta-network collapsed.

- Alpha variability or patch variability was impacted by network size and less by species dispersal or metacommunity size. Before an abrupt transition due to decreases in mutualistic strength, larger networks will exhibit stronger patch variability before collapse than smaller
- networks.

  Species-level metrics such as species-level autocorrelation and standard deviation were av-
- eraged over all species in a network before they collapsed as mutualistic strength decreased.
- 313 Species-level indicators did indicate as networks collapsed but its relationship with network
- topology such as connectance and network size were not significant (fig. 4A-B).

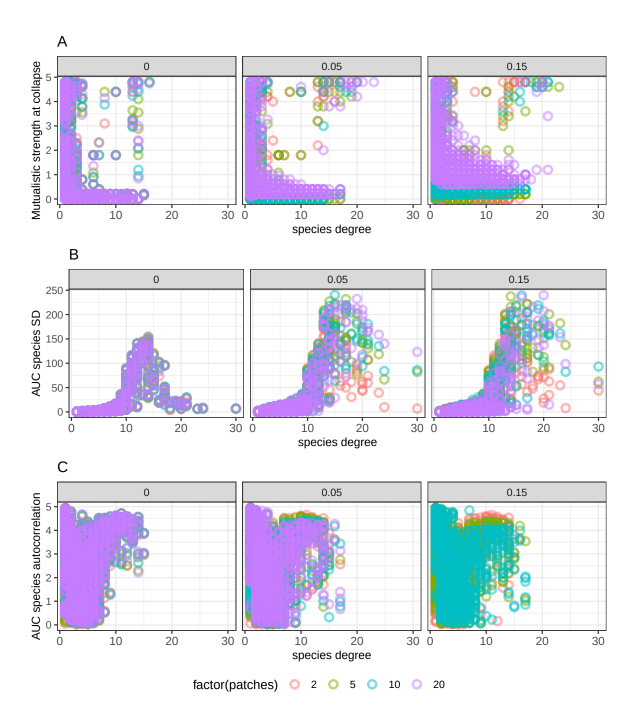


Figure 4: A) Strength of mutualistic interaction at which species in a network collapsed in comparison to its degree. B) Relationship between species degree and area under the curve of SD (standard deviation) measured at the species-level and C) autocorrelation at first-lag for three levels of dispersal (0,0.05,0.15) and four metacommunity sizes (2, 5, 10, 20). Note that the y-axis in all these subplots indicate the strength of increases in leading indicators (measured as the AUC of the metric see figure 1C for description) as a mutualistic meta-network collapsed. Larger AUC would indicated stronger increases as a mutualistic meta-network collapsed.

#### 15 Discussion

Climate change can cause shifts in interaction strengths that could lead to mismatches in species interactions thereby causing negative impacts on species biomass and diversity 317 (Revilla et al., 2015; Thompson and Gonzalez, 2017). In mutualistic networks, positive in-318 teractions between groups of species leads to a positive-feedback mechanism that promotes 319 the occurrence of sudden collapses once environmental change causes interaction strengths to 320 fall below a certain threshold (Baruah, 2022; Kéfi et al., 2016; Lever et al., 2014). Although, 321 studies have tried to understand how such networks are robust to habitat destruction and changes in phenology, little is known about how the architecture of such mutualistic net-323 works at the spatial scale could influence the timing of such collapses. Here, I show that 324 species disperal rate, size of the metacommunity, and network architecture has significant impacts on the threshold at which a mutualistic meta-network collapses. Furthermore, per-326 formance of statistical metrics that could be used to forecast such unwanted collapses, such 327 as metacommunitiy variability, alpha variability or spatial variability, was not only depen-328 dent on the architecture of the mutualistic meta-network but also on species dispersal rate and the size of the metacommunity. 330 When dispersal rate was zero, all mutualistic networks distributed across a spatial scale 331 could be coalesed to independent local communities. Effectively, metacommunity size i.e., 332 number of habitat patches in the metacommunity, thus, has no impact when a network 333 collapsed. This is obvious as species dipsersal is zero and spatial insurance provided by species dispersing across the landscape is effectively negligible (Loreau et al., 2003). As 335 a result, when strength of mutualism decreased, networks collapsed and metacommunity 336 size didnt have any impact when such networks collapsed. There was, however, a slight 337 negative relationship between size of the network and the mutualistic threshold at which 338 such networks collapsed. Particularly, larger networks collapsed later than smaller networks. 339 Earlier studies have indicated that feasiblity of mutualistic networks increases as network 340 size or nestedness increases (Baruah, 2022; Lever et al., 2014). This is intuitive as larger networks have larger number of species which effectively increases network biomass and persistence of the network as a whole.

As species dispersal increases, the effect of metacommunity size becomes more evident. At 344 high species dispersal, the relationship between network size and the mutualistic strength 345 threshold at which networks collapsed became more negative. Smaller networks collapsed much earlier at higher dispersal rate than larger networks. This particular result was evi-347 dent at larger metacommunity sizes (Fig. 2A). At smaller metacommunities, for instance 348 a metacommunity with two habitat patches, species were able to easily disperse to the 349 only other habitat and therefore were able to rescue local communities from collapsing at a 350 much earlier mutualistic strength. As such, in smaller metacommunities, smaller networks 351 collapsed at a similar mutualistic strength as larger networks. The relationship between 352 mutualistic strength at which networks collapsed and network size becomes more negative 353 at higher dispersal rates and larger metacommunity sizes. This can be explained as follows. 354 In such mutualistic metacommunities species were modelled to be obligate mutualists, which 355 indicated that species depended solely on mutualistic interactions for maintaining positive growth rate. As species dispersal increased, indicating that a larger chunk of individu-357 als of a species dispersed from local patches, a stronger average mutualistic interaction 358 was required for a species to maintain an overall positive growth rate locally. As a consequence, at very high dispersal rates, species growth rate decreases locally, and hence, 360 a much stronger mutualistic strength on average would be required to maintain positive 361 growth rate. This particular phenomena did not become an issue in smaller metacom-362 munities, for instance a metacommunity with two habitat patches, as species could easily 363 disperse and compensate for negative growth (Loreau et al., 2003). However, this becomes 364 a problem at larger metacommunities as species now have more than just a few habitats 365 to disperse to. In addition to this, smaller networks, on average, have smaller number of mutualistic interactions when compared with larger networks. As a consequence, smaller 367 networks in larger metacommunities collapsed at a higher mutualistic strength than larger 368 networks in larger metacommunities. Larger networks collapsed later in larger metacommunities solely because they were able to retain higher biomass due to higher number of mutualistic interactions on average, as well as due to larger number of species. Results, therefore, indicate that in order to preserve biodiversity, it is thus imperative to take into account both the sizes of communities (toplogical features) as well as the spatial scale of species interactions (Thompson and Gonzalez, 2017).

Much work has been done in identifying statistical signals that could forecast unwanted 375 critical transitions. These statistical signals are phenomenological in nature and could be 376 easily identified from state-based timeseries data such as abundance or biomass (Baruah 377 et al., 2019; Clements and Ozgul, 2016; Dakos et al., 2012a, 2012b; Scheffer et al., 2012). 378 However, such signals and their efficacy have been questioned recently both in numerical 379 simulations studies (Baruah et al., 2020; Clements et al., 2015; Hastings and Wysham, 2010) 380 and experimental studies (Baruah et al., 2021; Wilkinson et al., 2018). Their application on multi-species communities (Baruah et al., 2022; Dakos, 2017; Dakos and Bascompte, 2014; 382 Patterson et al., 2021), particularly in the context of a spatial structure has been lack-383 ing. Here, I evaluated how the efficacy of commonly used statistical signals measured at the species level (autocorrelation at first-lag, standard deviation), community level (patch 385 variability, spatial variability) and at the metacommunity level (temporal regional variabil-386 ity) in relation to network topology and spatial scale of interactions. Interestingly, species level metrics i.e., standard deviation and autocorrelation at first-lag exhibited increases as 388 strength in mutualistic interaction decreased (AUC greater than 1), however, there was no 389 relationship with network architecture (Fig. S5). Community and metacommunity-based 390 signals such as alpha variability, spatial variability also increased as mutualistic strength 391 decreased. Surprisingly, although smaller networks in larger metacommunities collapsed 392 earlier, however, such networks exhibited stronger increases in spatial variability and meta-393 community variability particularly at higher species dispersal rates (Fig. 3A-C). This indicated smaller networks could exhibit strong statistical signals before collapses which was 395 conditionally dependent on high species dispersal and whether such communities were a 396 part of a large metacommunity. Spatial variability also exhibited a similar pattern. Thus, when global changes weaken mutualistic interactions thereby leading to loss of biodiversity, indicators measured at the community and metacommunity level could potentially inform instability. Our analysis indicated that these indicators and their performance when estimated at the community or metacommunity level will be dependent on species dispersal, network architecture and how large the metacommunity is.

Changes in the environment can weaken species interactions to the point of biodiversity 403 collapse. Recovering lost ecosystem functions and processes is not easy as demographic in-404 formation of species are also lost as biodiversity collapses [(Link and Watson, 2019; Scheffer 405 et al., 2009; schefferAlternativeEquilibriaShallow1993?). In addition, most ecologi-406 cal systems can exhibit a phenomenon called hysteresis whereby even if the original stable 407 environmental conditions are reverted, the ecological system might still not recover. There 408 are tools that have been thus develop to forecast such unwanted transitions Dakos et al. 409 (2012b). When an ecological network collapses both at the local as well as at the spatial 410 scale, signals of global meta-network instability could also be manifested in species in an 411 ecological community. It is relatively unknown, however, which species could exhibit signals of instability, but see Dakos (2017), Baruah et al. (2022) or Patterson et al. (2021). Indeed, 413 the species that exhibited strong increases in standard deviation as networks collapsed, were 414 the ones which had on average a larger number of interactions (Fig. 4B). The relationship between species degree and strength of species SD was non-linear, indicating that species 416 with moderate to high number of interactions would show the strongest increase as net-417 works collapsed. Species that had a relatively larger number of mutualistic interactions, 418 benefited from higher positive growth rate which resulted in stronger increases in standard 419 deviation as networks collapsed than those species that had fewer interactions. However, as 420 shown in Dakos and Bascompte (2014), and also here, species that have fewer interactions 421 would collapse earlier and might not exhibit as strong an increase in standard deviation as networks collapsed. This indicates that, although, one could potentially label species that 423 have higher number of interactions as "indicator species," that, however, would not help 424 in informing instability of "specialist species" as they generally collapsed much earlier (Fig. 426 4A). Nevertheless, these results indicate that standard deviation measured at the species 427 level could potentially increase as mutualistic interactions weaken and more so for species 428 with a moderate to high number of interactions. Autocorrelation at first-lag also increased 429 as interactions weakened, but its relation with species degree was unclear.

Our ability to detect an impending transition is dependent on monitoring dynamics at the vicinity of the transition. However, monitoring population dynamics required intense 431 temporal as well as spatial sampling and could potentially impact forecasting of abrupt tran-432 sitions (Arkilanian et al., 2020; Bruel and White, 2021; Clements et al., 2015). Many other 433 factors could also impact prediction of such transitions including but not limited to long 434 transients and stochasticity (Hastings et al., 2018; Hastings and Wysham, 2010). There has 435 been guite a debate on the appropriate set of methods or tools that could be used to predict 436 future biodiversity states. Studies have indicated that in addition to monitoring dynamics of species abundance, phenotypic traits should be monitored as well. Including information 438 from phenotypic dynamics such as body size could improve forecasts of biodiversity collapse 439 (Clements and Ozgul, 2016), although such an accurate forecast of biodiversity collapse is dependent on the type of environmental perturbation and the type of interactions prevalent 441 in the community (Baruah et al., 2022, 2019). When changes in the environment impact 442 communities both locally and spatially alike, ecological networks could abruptly collapse. However, when such networks collapses depends on the average rate of species dispersal, with higher species dispersal and larger metacommunities causing meta-networks to collapse 445 at a much higher mutualistic strength. In addition, collapses of mutualistic meta-networks 446 was also dependent largely on the topology of the network, with smaller networks collapsing at a higher mutualistic strength than larger networks. These results are pertinent for 448 conservation efforts particularly because it points to the fact that ecological networks at 449 larger metacommunities are as vulnerable or if not more than meta-networks at a much smaller spatial scale. When assessing whether communities are vulnerable to changes in 451 the environment, the results from this research indicates that it is very pertinent to look 452 at factors beyond species demographic rates, that includes the scale of spatial interactions, and topology of local ecological communities.

#### 455 Author contributions

456 GB conceptualized the study, did the analysis and wrote the manuscript.

## 457 Data availability

- The entire paper was written in rmarkdown format and files, data, and R scripts to repro-
- duce the paper is located at.

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## Supplementary document:

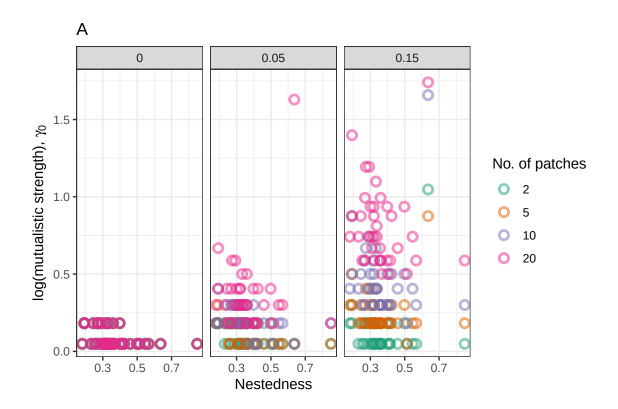


Figure S1: Relationship between the threshold mutualistic strength at which networks collapsed and structural properties of the network namely nestedness (NODF).

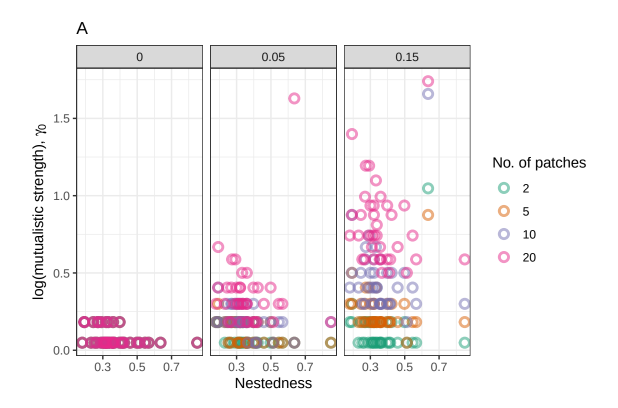


Figure S2: Relationship between the threshold mutualistic strength at which networks collapsed and structural properties of the network namely nestedness (NODF).

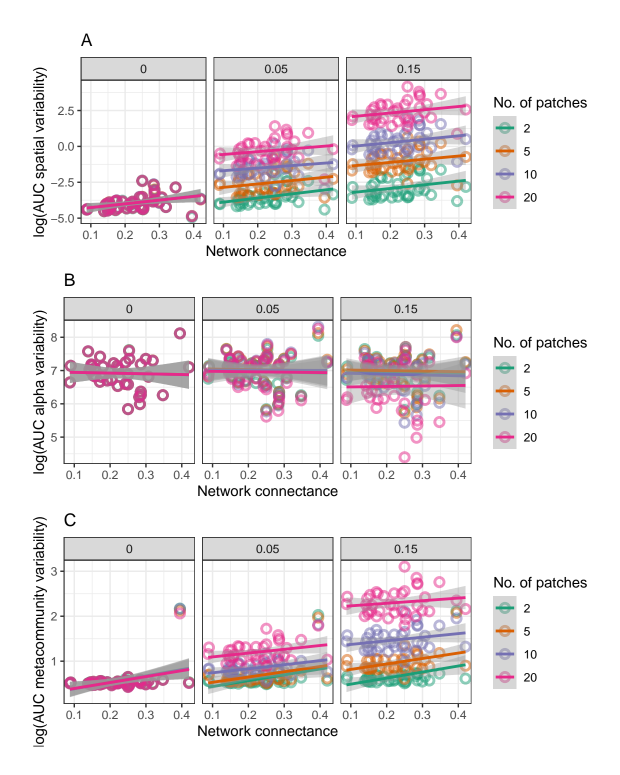


Figure S3: Relationship between network connectance and area under the curve of statistical metrics such as A) spatial variability, B) Alpha variability, and C) metacommunity variability for three levels of species dispersal (0,0.05,0.15) and four metacommunity sizes (2, 5, 10, 20). Note that the y-axis in all these subplots indicate the strength of increases in these indicators (measured as the AUC of the metric, see figure 1C for description) as a mutualistic metacommunity collapsed. Larger AUC would indicated stronger increases as a mutualistic meta-network collapsed.

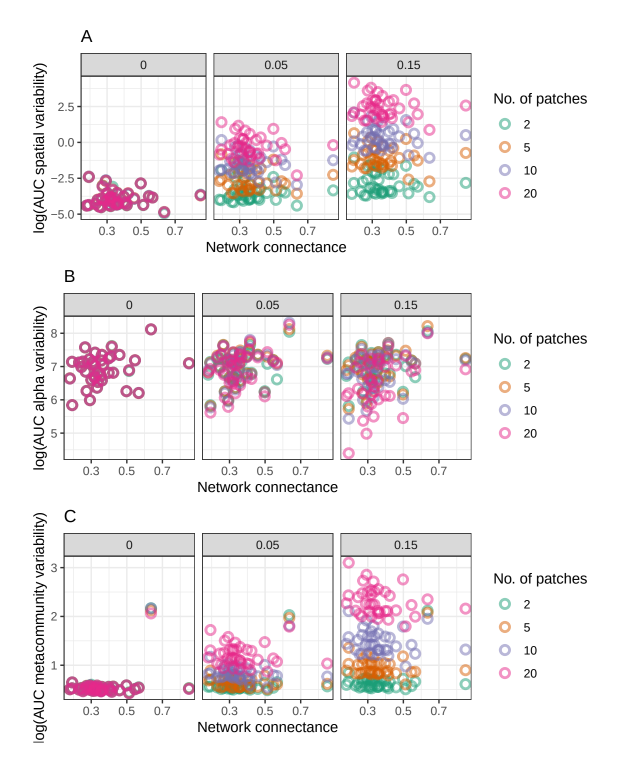


Figure S4: Relationship between network nestedness (NODF) and area under the curve of statistical metrics such as A) spatial variability, B) Alpha variability, and C) metacommunity variability for three levels of species dispersal (0,0.05,0.15) and four metacommunity sizes (2, 5, 10, 20). Note that the y-axis in all these subplots indicate the strength of increases in these indicators (measured as the AUC of the metric, see figure 1C for description) as a mutualistic meta-network collapsed. Larger AUC would indicated stronger increases as a mutualistic meta-network collapsed.

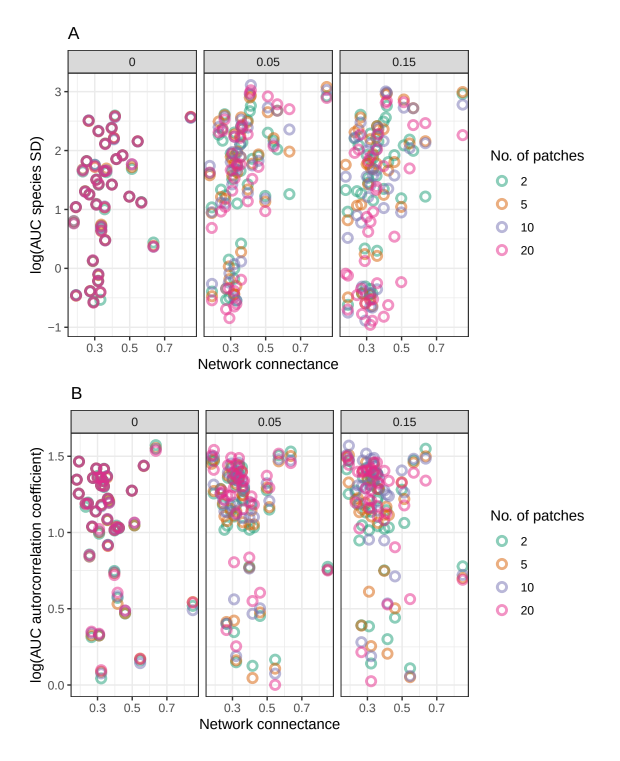


Figure S5: Relationship between network nestedness (NODF) and area under the curve of statistical metrics such as A) species SD, B) Autocorrelation coefficient at first-lag, for three levels of species dispersal (0,0.05,0.15) and four metacommunity sizes (2, 5, 10, 20). Note that the y-axis in all these subplots indicate the strength of increases in these indicators (measured as the AUC of the metric, <sup>35</sup>/<sub>see</sub> figure 1C for description) as a mutualistic metacommunity collapsed. Larger AUC would indicated stronger increases as a mutualistic meta-network collapsed.

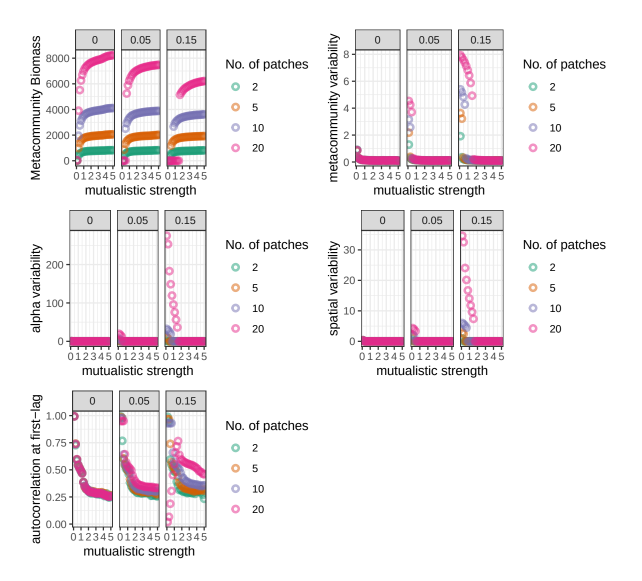


Figure S6: Meta-network collapse and statistical metrics that increased as mutualistic strength decreased for three rates of dispersal, four metacommunity size. A) Example meta-network collapse as mutualistic strength decreased. (B) metacommunity variability as mutualistic strength decreased. (C) alpha variability as mutualistic strength decreased. (D) Spatial variability as mutualistic strength decreased. (E) species autocorrelation as mutualistic strength decreased.