Eco-evolutionary diversification of trait convergence and complementarity in

mutualistic networks

Francisco Encinas-Viso<sup>1,\*</sup>, Rampal S. Etienne<sup>2,\*\*</sup> and Carlos J. Melián<sup>3,\*\*</sup>

1) NRCA & Centre for Australian National Biodiversity Research, CSIRO, GPO Box 1600

Canberra, Australia

2) Groningen Institute for Evolutionary Life Sciences, University of Groningen, The Netherlands

3) Department of Fish Ecology and Evolution, Center for Ecology, Evolution and Biogeochemistry,

EAWAG, Swiss Federal Institute of Aquatic Science and Technology, Switzerland

Keywords: Diffuse coevolution, specialization, sexual reproduction, assortative mating, dispersal

limitation, phylogenetic relatedness, obligate mutualism, morphological constraints, individual

based model.

Type of Article: Letters

Number of figures: 6 in color; Number of tables: 1

\* Corresponding author: franencinas@gmail.com

\*\* Joint last authorship

Mutualistic networks show high levels of convergence, complementarity and nestedness. Convergence and complementarity have been attributed to coevolutionary interaction selection under a phenotype matching scenario. It is not clear, however, whether phenotype matching fits empirical patterns better than a phenotype difference scenario. Here, we present a spatially and genetically explicit plant-animal trait diversification model combining the phenotype matching and phenotype difference model. Our model predicts high levels of convergence, complementarity, and nestedness. Comparing the predictions with observed levels of complementarity and convergence in an empirical plant-hummingbird mutualistic network, we find good predictions except for convergence patterns in plants. Our results indicate that combining the spatial structure of plant-animal trait diversification with population dynamics in a phenotypic difference model may alter predictions in a direction that is more in line with observations, and hence are important to be included in future studies of coevolutionary dynamics in mutualistic networks.

### Introduction

Since Darwin's book "On The Origin of Species" (Darwin, 1859), the idea of coevolution, defined as reciprocal evolutionary trait change between species, has sparked interest from biologists trying to understand how species interactions generate trait changes. The first clear indication of coevolution was Darwin's moth example (Darwin, 1862) showing that the long corolla from the orchid Angraecum sesquispedale could only be reached by a pollinator species with a similar or larger proboscis length. Following the moth and orchid mutualism model system, several studies have modeled coevolutionary dynamics of a few species (Ferriere et al., 2007; Law et al., 2001; Ferdy et al., 2002; Gomulkiewicz et al., 2003; Jones et al., 2009), particularly highly specialized (i.e. obligatory mutualists) systems of plant-animal interactions, such as the fig-fig wasp mutualism (Bronstein et al., 2006). These studies have determined the ecological conditions for coevolutionary stable systems in highly specialized plant-animal interactions (Law et al., 2001; Jones et al., 2009) Janzen (1980) argued that high specialization between plants and animals was not the only example of coevolution, but coevolution can also be the product of multiple-species interactions, a term that he coined "diffuse coevolution". Diffuse coevolution means that selection on traits is determined by the interaction of more than two species and not only based on pairwise interactions. This concept is based on the idea of "syndromes", where, for example, plants have a set of traits that attract a specific group of pollinator species with traits complementary to those of the plants (Jousselin et al., 2003; Guimarães et al., 2011). Evolutionary convergence, i.e. independent evolution of similar features among different lineages of the same plant or animal community may partly explain the formation of "syndromes" produced by the presence of specific mutualist partner species (Howe and Smallwood (1982); Waser et al. (1996); Bascompte and Jordano (2007); Stayton (2008); Losos (2011)). Theoretical studies suggest that evolutionary convergence and complementarity are generated by coevolutionary interaction selection driven by a "phenotype matching model" (Nuismer et al. 2005; Kopp and Gavrilets 2006). For example, in highly specialized two-species interactions such as the fig-fig wasp mutualism (Bronstein et al., 2006), plants coevolve with their most efficient pollinator to strengthen the complementarity of their matching traits. The phenotype matching model assumes that the probability of a successful interaction decreases with increasing distance between the phenotypes of the interacting individuals (Nuismer et al., 2012). Examples are plant and animal phenology

(Jordano et al. 2003; Olesen et al. 2010; Encinas-Viso et al, 2012) or proboscis and corolla lengths
(Agosta and Janzen 2005; Santamaría and Rodríguez-Gironés 2007; Stang et al. 2007).

Only two theoretical studies have explored the importance of coevolutionary selection and pheno-

type matching in the emergence of convergence, complementarity and network structure in speciesrich mutualistic networks (Nuismer et al., 2012; Guimarães et al., 2011). Guimarães et al. (2011)
studied a coevolutonary model with a fixed network structure and showed that convergence in a
one-dimensional trait may emerge as a consequence of coevolutionary selection for a complementarity trait between trophic levels. Nuismer et al. (2012) explored the importance of coevolutionary
selection in a multispecifies context to study the emergence of convergence, complementarity and
network structure in mutualistic networks. They found that weak coevolutionary selection produces
little to no convergence but generates high complementary. As coevolutionary selection intensifies,
variation in the trait values of animal and plant species is reduced and high convergence emerges but
complementarity decreases. Furthermore they found antinested networks regardless of the strength
of coevolutionary selection.

Nuismer et al. (2012) also used an alternative model called "phenotypic difference" in which the success probability of an interaction increases with the degree to which the animal trait exceeds the plant trait as might be the case for fruit and beak size in a dispersal mutualism (Lambert 1989; da Silva and Tabarelli 2000) or proboscis and corolla length in some pollination mutualisms (Inouye 1980; Borrell 2007; Anderson and Johnson 2009; Anderson et al. 2010). They showed that under the pheotype difference model networks were more nested when coevolutionary selection was weak. Furthermore, similar to the phenotypic matching model, convergence increases with the strength of coevolution in the phenotypic difference model (but only when interactions depend strongly of phenotypes); however complementarity is weaker for interactions mediated by phenotype difference than those mediated by phenotype matchings. Taken together, these results suggest that it is not trivial to simultaneously explain high degrees of convergence, complementarity and nestedness in species-rich mutualistic networks as observed in empirical data (Bascompte and Jordano (2007).

The failure of successfully explaining three patterns simultaneously may also be a consequence of fac-

 $_{\circ}$  tors currently lacking in coevolutionary models of mutualistic networks, such as the role of genetics,

gene flow, and assortative mating in plant-animal trait diversification (Gavrilets et al. (2000); de Aguiar

et al. (2009); Schluter (2009); Doebeli (2011); Seehausen et al. (2014)), and spatial structure and population dynamics. For example, although gene flow can be seen as an homogenizing factor, there is increasing evidence showing the importance of gene flow in the speciation process (Nosil, 2008) as well as the importance of spatial structure de Aguiar et al. (2009). However, more importantly, there is no doubt that considering genetic processes (e.g. drift, selection) are essential to understand biodiversification patterns. In this study we present an integrative model to study the patterns of convergence, complementarity and nestedness in a model that includes genetic processes, sexual reproduction, spatial structure, population and trait diversification dynamics. Our model lies between a phenotype matching and phenotype difference model, as an interaction can only occur when the animal trait exceeds the plant trait, but the probability of success does not increase with this difference. Furthermore, we use a definition of convergence that accounts for phylogenetic relatedness. In contrast to previous studies, our model shows high levels of convergence, complementarity and nestedness. Comparing these predictions to convergence and complementarity patterns observed in a plant-humming bird mutualistic network, we find that the model is in agreement with observed level of plant-animal complementarity and convergence for the humming bird community, but conflicts with convergence in the plant community. Our results indicate that combining the spatial structure of plant-animal trait diversification with population dynamics in a phenotype difference model alter

# Eco-evolutionary trait diversification in mutualistic networks

included in future studies of evolutionary patterns in mutualistic networks.

To study the effect of spatial structure and plant-animal trait diversification on convergence, complementarity and nestedness in mutualistic networks, we develop a spatial quantitative genetics model for plant-animal interactions with overlapping generations. Individuals have fixed genome size and we start with one plant and animal species with all individuals having the same genetic composition at the outset (see Figure 1 for the steps of the model). Following population demography males and females individuals in both communities must be close enough in spatial, genetic, and morphological space for an obligate mutualistic interactions to occur (Figure 1 step B) and to produce offpsring

predictions in a direction that is more in line with observations, and hence are important to be

with a new genotype-phenotype after mutation, recombination and phenotypic plasticity (Figure 1 step C, D and E). We follow the species definition of Nei et al. (1983), where a species is a group of interbreeding and successfully reproducing individuals that are reproductively isolated from other 101 groups. In our model two individuals can mate successfully if their genetic similarity value is larger 102 than or equal to the minimum genetic similarity value. Thus, speciation is defined as a group of 103 genetically related individuals, where two individuals in a sexual population can be incompatible, but 104 still conspecific as long as they can exchange genes indirectly through other conspecifics (de Aguiar 105 et al., 2009; Melián et al., 2010). This is the definition of 'ring species' (Moritz and Schneider, 1992). 106 Please refer to Figure 1, Table 1 and SI for a detailed explanation of the terms included in the model 107 and how we model population dynamics, diversification dynamics, quantitative trait dynamics and 108 neutral locus evolution, respectively. 109

### Convergence, complementarity, nestedness and connectance

### Evolutionary convergence and complementarity

125

A focal species is considered to have convergent traits with another species if it is phenotypically more similar to this other species than to the genetically most similar species (the sister species) 113 (see figure 2 and SI). The number of convergences potentially increases with the number of species 114 present. For example, if we have ten species and we exclude one of them as the sister species of the 115 focal species, we have nine species to calculate convergence. If we find that two out of nine species are phenotypically similar enough to the focal species, we count two (out of nine,  $\sim 22\%$ ) convergences. 117 We repeat this by changing the focal species and calculate the mean number of convergence events 118 over all species. In contrast to previous approaches that used the mean pair-wise difference between 119 traits of species (Guimarães et al., 2011) or the variance of species traits in a guild as a proxy to 120 predict convergence (i.e., large values weak convergence whereas small values of the variance may 121 indicate strong convergence, (Nuismer et al., 2012)), our method considering phylogenetic relatedness is that it excludes cases of development of very similar trait values in sister species due to shared evolutionary history and therefore it does not overestimate convergence events.

The condition for complementarity between two mutualistic partners is simply that the phenotypic

similarity between them, is larger than some predefined phenotypic threshold value

We refer to the SI for a detailed explanation of phenotypic similarity and mean genetic and phenotypic species similarity, and a precise definition of complementarity and convergence in terms of these similarities.

#### Nestedness and connectance

To study the connection between convergence and complementarity with network properties, we mea-131 sured two topological properties of plant-animal mutualistic networks: nestedness and connectance. 132 We estimated nestedness using the NODF algorithm developed by (Almeida-Neto et al., 2008) be-133 cause of its statistical robustness. NODF is based on standardized differences in row and column fills 134 and paired matching of occurrences. Connectance measures the proportion of realized interactions 135 among all possible interactions in a network. It is defined as  $C = \frac{k}{P*A}$ , where k represents the number 136 of realized interactions between plant and animal species and P and A represent the number of plant 137 and animal species in the network, respectively (Jordano et al., 2003). 138

### Simulations

151

We simulated equal population sizes for plants and animals with  $J^P = J^A = 1,000$  individuals. 140 Genome size, L, of each individual was 150 loci. Initial trait distributions for the plants,  $Z^P = [z_i^P]$ 141 and animals,  $Z^A = [z_i^A]$ , were generated following equation 2 plus a normally distributed environ-142 mental effect,  $\epsilon$ ,  $\mathcal{N}(\mu_{\epsilon}=0,\sigma_{\epsilon}^2=1)$ . To ensure plant mating conditions are met at the beginning of 143 the simulation all animal individuals have a higher phenotypic trait value that the plant individuals. 144 Geographic distances between each pair of individuals i and j for the plants,  $d_{ij}^{P}$ , and animals, 145  $d_{ij}^A$ , were calculated as follows: 1) Euclidean coordinates of a two-dimensional space  $(x_i, y_i)$  were 146 sampled from a uniform distribution  $(x_i = [0, 1], y_i = [0, 1])$  for each individual for the plants and 147 animals; 2) Using these coordinates we calculated a matrix of relative Euclidean distances between 148 the individuals for the plants,  $d_{ij}^P$ , and animals,  $d_{ij}^A$ . This procedure was repeated for each of the 149 geographic distance matrices  $(D^{PA}, D^P, D^A)$  (See Table 1 and SI). 150

We ran 2,000 generations for each replicate for a total of 500 replicates, where a generation is

the update of the effective population size  $(J^P = J^A = 1,000)$ , i.e. the number of steps equal to the effective population size. Steady-state was verified by checking the constancy of speciation 153 events during the last 1000 generations. We calculated convergence, complementarity, nestedness and 154 connectance at steady-state. Convergence and complementarity events were calculated for a whole 155 range ([0.0, 1.0]) of their respective thresholds,  $t_{conv}$  and  $t_{comp}$ . We explored parameter combinations 156 with mutation rate,  $\mu \in \{10^{-4}, 10^{-2}\}$ , minimum genetic similarity,  $q_{min} = 0.97$ , maximum distance 157 for finding a mate and disperse,  $d_{max} \in \{0.1, 0.3\}$ , and a maximum geographic distance to find a 158 mutualistic partner,  $d_{max}^{PA}=0.3$ . We implemented the model in Python (and tested in IPython (Pérez 159 and Granger, 2007)). 160

## Confrontation to empirical data

We tested our model's predictions of convergence and complementarity on a plant-humming bird net-162 work with 38 hummingbird species and 133 plant species (Maglianesi et al. (2014)). To compute 163 phenotypic similarity we used empirical values of corolla length and bill length from plants and hum-164 mingbirds, respectively (REF). To compute phylogenetic similarity for the hummingbirds we used 165 a well resolved recently published comprehensive phylogeny (McGuire et al, 2014) to obtain phyloge-166 netic relationships for 24 out of 38 humming bird species; the remaining 14 species were not present in 167 the phylogenetic tree. For the plant species we constructed a phylogenetic tree using sequence data 168 of 69 species from GenBank (data on rbcL, matK and ndhF genes collected by PhyloGenerator using 169 the option THOROUGH to allow also data from close relatives if these relatives were not present 170 in the community), aligned them using MUSCLE (REF) and coonstructed the maximum likelihood 171 (ML) tree using RAXML (REF). We excluded 64 plant species from the analysis (see Suppl. Infor-172 mation) because there was no sequence data available or the phylogenetic relationships were not well 173 resolved. 174

We used the phylogenetic trees with their respective branch lengths to calculate a genetic distance matrix among species. Using both phylogenetic trees (hummingbirds and plants) we simulated nucleotide sequences of 100 bp with the program SeqGen (Rambaut and Grassly, 1997) following the Jukes-Cantor model of molecular evolution. These simulated sequences were then used to calculate

the genetic distance matrix using the R package seqinr in R (R Core Team, 2013). To compare the convergence values obtained from the empirical data with our model predictions, we generated 1000 replicates from the simulations (bootstrapping) with each replicate containing the same number of plant and animal species as the empirical data. Mean values as well 0.05 and 0.95 CI were generated from these 1000 replicates. Complementarity and convergence were calculated for each of the replicates across the whole range of convergence,  $t_{conv}$ , and complementarity,  $t_{comp}$ , thresholds. We used a conservative definition for convergence in which not only the sister species, but the 30% most genetically similar species were excluded.

#### Results

#### 188 Trait evolution

Population dynamics and diversification dynamics changed plant and animal community trait distributions (i.e. corolla and proboscis lengths) with bimodal distributions being the most commonly 190 produced patterns across replicates (figure 3). At the species level, a gradient of species phenotypes 191 with common species presenting lower mean and higher variance than rare species emerged. Mean and variance of the trait values were correlated for most replicates (Spearman- $\rho > 0.41$ , p < 0.05) 193 and the distributions of abundance for plant or animal species were highly skewed and significantly 194 different from a normal distribution (Lilliefors's test, all p < 0.001). Abundance predicted plant or 195 animal mean species traits in approximately 70% of the replicates (Spearman- $\rho > 0.32$ , p < 0.05) 196 and trait variance for all replicates (0.39 < Spearman- $\rho$  < 0.79, all p < 0.05). Mean and variance of 197 species trait values significantly differed between common and rare plant or animal species (inset in 198 figure 3) suggesting a strong impact of diversification by producing a gradient of species phenotypes 199 in mutualistic networks. 200

# Convergence and complementarity

Evolutionary convergence events occurred in all replicate simulations (see equations SI-4 and SI-5 with an example of evolutionary convergence events in animals and plants represented in figure

4). Convergence events were heterogeneously distributed across species with most events occurring

occurred on average in  $17.3 \pm 6\%$  of all possible convergence events with more than 95% of these 206 events involving some of the three most common species. These results show that evolutionary 207 convergence is not randomly distributed across pairs of species but highly aggregated during the 208 diversification process. Evolutionary convergence can also be visualized using a scatter plot of the 209 genotype-phenotype map for all pairs of individuals within the plant and animal communities (see 210 Figure SI-1 Supplementary Information). As expected from equation ??, there is a positive genotype-211 phenotype relationship. The scatter plot contains three main clouds of points that consistently 212 occur in our simulations for the plants, P, and animals, A: 1) pairs of individuals of the same 213 species with high genetic  $(q_{ij} > q_{min})$  and phenotypic  $(p_{ij} > 0.9)$  similarity, 2) pairs of individuals 214 of the same species with genetic similarity below  $q_{min}$   $(q_{ij} < q_{min} = 0.97)$  and high phenotypic 215 similarity  $(p_{ij} > 0.9)$ . These are incompatible individuals of the same species for mating, yet with 216 high phenotypic similarity,  $p_{ij} > 0.9$ , and 3) highly genetically dissimilar individuals from different 217 species,  $q_{ij} \ll q_{min}$ , but with the presence of highly phenotypically similar individuals  $(p_{ij} > 0.9)$ . 218 This last category shows evidence of evolutionary convergence between species in plants and animals. 219 An increase in mutation rate increases the genetic divergence between species, as expected, but it 220 does not change the genotype-phenotype relationship qualitatively (see Figure SI-1 Supplementary 221 Information). 222 Evolutionary complementarity occurred with a similar frequency as evolutionary convergence in 223 each replicate (see equation SI-5 and compare the initial with the final trait distributions in figure 224 3), but with a larger variation ( $20 \pm 18\%$ ). Connectance values were consistently medium or high 225  $(\overline{C} = 0.5 \pm 0.07, \text{ figure 5}), \text{ mostly larger than reported in empirical data where it ranges between 0.05}$ 226 and 0.25. Nestedness values were always high  $(\overline{N} = 69.97 \pm 13.4 \text{ (figure 5)})$ , as observed in the empir-227 ical plant-pollinator networks. Convergence, complementarity and nestedness did not show signs of 228 trade-offs and were uncorrelated across all replicates (0.08 < Spearman- $\rho$  < 0.27, all p > 0.1) with the exception of a positive correlation between trait complementarity and evolutionary convergence 230 in the plant community (Spearman- $\rho = 0.61$ , all p < 0.05). Our results, using phylogenetic related-

ness and phenotypic similarity for the estimation of evolutionary convergence and complementarity

under weak coevolutionary selection, show evolutionary trait convergence and complementarity in all

232

between common species (0.42 < Spearman- $\rho$  < 0.89, all p < 0.05). Evolutionary convergence

our replicate simulations but with little and large variation, respectively. These results suggest that in our model, trait convergence in plant and animal communities is largely independent or positively correlated with trait complementarity for the animal and plant community, respectively.

## Comparison of model predictions with empirical data

Our model predicts well plant-animal complementarity and convergence for animals but not for 238 plants in a empirical plant-humming bird mutualistic network (Figure 6). The observed proportion of 239 complementarity events for the empirical plant-humming bird data is within the CI for a broad range 240 of values of the complementarity threshold,  $t_{comp}$  (Figure 6a). Our model consistently predicts higher proportion of convergence events than the observed proportion in the plant community (Figure 6b). 242 Predictions in the proportion of convergence events quickly increase for a high convergence threshold 243 value both in the empirical data and in our model predictions (red lines Figure 6b) and saturates 244 around the same observed values for medium and low convergent threshold values. Predictions of the proportion of convergence events for the humming bird community are within the estimated CI 246 for all the range of convergence threshold values (Figure 6c). These results show that predictions for plant-animal complementarity and convergence in the humming bird community are robust against 248 a broad range of threshold values.

#### 250 Discussion

Multispecific coevolution has been poorly studied so far due to its complexity, which involves numerous processes and mechanisms in the ecology and evolution of species interaction networks. Nevertheless, there is accumulating empirical data (REFS) for and increasing interest in understanding the
drivers of multispecific coevolution in ecological communities (REFS). Our model comes closer to
observed patterns of complementarity and convergence in the plant-hummingbird community than
previous models.

# The evolution of convergence and complementarity

258

tionary processes ('diffuse coevolution')(Janzen, 1980; Thompson and Cunningham, 2002; Jordano 259 et al., 2003; Bascompte and Jordano, 2007) and therefore convergence events are molded by similar 260 ecological (or niche) selective pressures. The model of (Nuismer et al., 2012), which studied a model 261 of 'phenotype differences' has shown that for weak coevolutionary selection trait values in animal and 262 plant species can be highly variable and non-convergent, but trait values of animal and plants species 263 show high complementarity (i.e. they are positively correlated). However, strong coevolutionary selection decreases variation in the trait values of animal and plant species increasing convergence and 265 simultaneously complementarity decreases (i.e. correlations between traits of interacting species are 266 weakened). Interestingly, the results of Guimarães et al. (2011) show that trait convergence may 267 in part emerge as a consequence of selection for a complementarity trait between the plants and animals. This model, contrary to Nuismer et al (2012), did consider background evolution (i.e. evo-269 lution driven by other forces not related to mutualistic interactions Guimarães et al. (2011), which provides support to our results. 271 Interestingly, the model of 'phenotype differences' considered here has shown to make unlikely the 272 evolution of convergence and complementarity by coevolutionary selection (Nuismer et al., 2012). 273 However, our model shows that by considering a more realistic model of phenotype differences in a 274 more realistic setting gives results that are more like the observations than the results of previous 275 models. In other words, it is possible to observe the evolution of both convergence and complementarity. It remains to be seen whether the action of both (selective an non-selective forces) and other 277 models of plant-animal interaction (e.g. 'phenotypic matching') will be able to generate the observed patterns of high convergence, complementarity and nestedness in species-rich mutualistic networks. 279 Non-selective forces underlying trait dynamics can produce convergence. For example, Stayton 280 (2008) simulated evolution along phylogenies according to a Brownian motion model of trait change 281 and demonstrated that rates of convergence can be quite high when clades are diversifying under only 282 the influence of genetic drift. Furthermore, other types of constraints in the production of variation 283 can also lead to convergence. If the variation produced is limited, then unrelated species are likely to 284 produce the same variation, which may then become fixed in the population by genetic drift (Stayton,

Previous studies have argued that evolutionary convergence is the product of multispecific coevolu-

2008; Losos, 2011). Developmental constraints or the evolution of genetic networks by non-adaptive processes may also be explanations for the convergence of traits (Solé et al., 2002; Lynch, 2007; Losos, 2011), but the role of developmental constraints or genetic networks in determining convergence in 288 species-rich mutualistic networks has yet not been explored. For example, the tinkering of traits 289 by evolutionary forces largely affects developmental pathways (e.g. gene regulatory networks) (Solé 290 et al., 2002). Developmental pathways are not static but can diverge through time randomly without substantially affecting the phenotype (Wagner, 2008). This concept, also called developmental system 292 drift (DSD) (True and Haag, 2001), might play an important role in the evolution of convergence 293 in morphological traits and it should be considered as another process where drift can act (Ohta, 294 2002), for example, by random wiring in gene regulatory networks. Our results based on a method 295 that excludes cases of the development of a similar trait in related but distinct species descending 296 from the same ancestor show that additional constraints such as dispersal limitation and assortative 297 mating limit the production of variation and lead consistently to convergence in distinct lineages in 298 species-rich mutualistic networks. 299

Evolutionary complementarity is also consistently observed in our results but with a larger variation 300 than convergence. Complementarity is argued to be the main result of tight coevolution between 301 mutualistic species by mechanisms, such as trait-matching (e.g. corolla length-proboscis length) 302 (Jordano et al., 2003). There is empirical (Anderson and Johnson, 2008) and theoretical evidence 303 (Gomulkiewicz et al., 2000) for coevolutionary hot spots (Thompson, 1999), which suggests that local 304 selective regimes can promote the coevolution of traits (Gomulkiewicz et al., 2000; Ferdy et al., 2002; 305 Gomulkiewicz et al., 2003; Jordano et al., 2003; Bronstein et al., 2006; Thompson and Cunningham, 306 2002; Thompson, 2009; Jones et al., 2009). In contrast, our results show that medium levels of 307 complementarity can emerge from relatively non-selective forces and constraints occurring at several 308 levels, from geographic limits to encounter partners and disperse to the genetic and morphological 309 constraints on producing viable offspring. In addition, our model fitting show our predictions fit 310 well to the observed plant-humming bird complementarity across a broad range of complementarity 311 threshold values (Figure 6).

Our predictions are able to reproduce some observed patterns of convergence and complementarity in a plant-hummingbird community. Observed convergence values were close to those predicted by the model for the hummingbird community, but we predicted more convergence values than those observed in the plant community. According to previous models, fewer convergence events could be indicative of weak coevolutionary selection processes acting on the plant species. Although this is a possible explanation for our overestimation of convergence, we think that our convergence estimates may have been biased by the fact that we could not consider all the plant species of the community ( $\approx 50\%$ ) only around 60% of the hummingbird species. Our study is the first to quantify and compare convergence and complementarity values from an empirical dataset with model predictions and we believe this a major leap towards the understanding of multispecific coevolution and community assembly.

### The problem of calculating convergence

One prominent problem in the literature is how to quantify convergence. This problem is probably related to the lack of a rigorous or precise definition of convergence (REFS). Previous studies (Nuismer et al, Guimaraes et al) proposed different ways of doing this, but their measurements of 327 convergence do not consider phylogenetic relatedness. For example, Guimarães et al. (2011) measured convergence as "the mean pair-wise difference between traits of species at same trophic level" 329 considering all the species of the guild. (Nuismer et al., 2012) calculated the variance as a proxy to 330 predict convergence (i.e., large values weak convergence whereas small values of the variance may 331 indicate strong convergence). The main problem with these measurements that ignore phylogenetic relatedness is that they will tend to overestimate convergence events. Most definitions of conver-333 gence agree that it is an evolutionary pattern in which "similar phenotypes evolve independently in 334 multiple lineages" (Stayton, 2015). Therefore, it is important to consider phylogenetic distance to 335 correct for shared and therefore dependent evolutionary history. Our model with our new measure 336 of convergence, which considers phylogenetic relatedness and phenotypic similarity, shows that in 337 the absence of abiotic selection trait convergence and complementarity always evolve but with little 338 and large variation, respectively. Therefore, even with a more conservative estimate of the number 339 of convergence events, the results show that the modelled processes (or background evolution sensu 340 Guimarães et al. (2011)) are crucial for the emergence of these evolutionary patterns.

### The emergence of mutualistic network structure

Nuismer et al. (2012) explored the connection between convergence and complementarity to nest-343 edness patterns in mutualistic networks. They show that coevolutionary selection tend to decrease 344 nestedness and it generates even more strongly antinested networks when coevolutionary selection 345 increases by favoring the emergence of reciprocal specialization. In contrast, nestedness values were 346 very high in our model, as in real mutualistic networks. Previous neutral models taking into account 347 ecological drift (Krishna et al., 2008; Canard et al., 2012), produced high values of nestedness which 348 suggests that random interactions and species abundance distribution ('neutral forbidden links' (Ca-349 nard et al., 2012)), are determinants of the structure of mutualistic networks. Connectance values 350 obtained from our simulations are close to the predictions of other neutral network models (Canard 351 et al., 2012). However, compared to real mutualistic networks with similar diversity as ours (24 plant 352 and animal species on average), our connectance values ( $\overline{C} = 0.5$ ) are higher than the reported webs (C = 0.28) (Olesen and Jordano, 2002). Interestingly, Nuismer et al. (2012) found that only assuming 354 coevolutionary selection forces also leads to an increase in connectance. This means that both basic genetic and ecological processes and coevolutionary selection can increase connectance in mutualistic 356 networks. The question is why observed mutualistic webs have a lower connectance than those pre-357 dicted by our model and previous models. We conjecture that this difference in connectance values 358 might be due to different types of forbidden links (i.e. biological constraints impeding plant-animal interactions), such as phenology (Encinas-Viso et al., 2012; Olesen et al., 2008) or environmental 360 fluctuations that were not explicitly included in our approach. 361

In summary, our results show the emergence of convergence, complementarity and nestedness 362 following basic genetic and ecological processes. Our predictions fit well to the observed plant-363 hummingbird complementarity and hummingbird convergence. In contrast to previous studies show-364 ing antinested networks, we found highly nested values in agreement with empirical mutualistic 365 networks (Bascompte et al., 2003). Our results suggest that diversification dynamics combining 366 ecological (demography and dispersal limitation), population genetics (mutation, recombination, as-367 sortative mating and drift) and morphological constraints may form the basic processes producing 368 the key patterns of mutualistic networks, from trait convergence and complementarity to connectance 369 and nestedness.

# Acknowledgments

- $_{\it 372}$   $\,$  We thank Martina Stang and Ole Seehausen for useful discussions. FEV and RSE were supported by
- VIDI and VICI grants from the Netherlands Organization for Scientific Research (NWO), awarded to
- RSE. CJM was supported by the Swiss National Science Foundation (SNSF-project 31003A-144162).

# 75 References

- de Aguiar, M., M. Baranger, E. Baptestini, L. Kaufman, and Y. Bar-Yam, 2009. Global patterns of speciation and diversity. Nature 460:384–387.
- Almeida-Neto, M., P. G. aes, P. G. aes, R. Loyola, and W. Ulrich, 2008. A consistent metric for
- nestedness analysis in ecological systems: reconciling concept and measurement. Oikos 117:1227–
- <sub>380</sub> 1239.
- Anderson, B. and S. Johnson, 2008. The geographical mosaic of coevolution in a plant-pollinator mutualism. Evolution 62:220–225.
- Bascompte, J. and P. Jordano, 2007. Plant-animal mutualistic networks: The architecture of biodiversity. Annu. Rev. Ecol. Evol. Syst. 38:567–593.
- Bascompte, J., P. Jordano, C. J. Melian, and J. M. Olesen, 2003. The nested assembly of plant-animal mutualistic networks. Proc. Natl. Acad. Sci. USA 100:9383–9387.
- Bronstein, J. L., R. Alarcon, and M. Geber, 2006. The evolution of plant-insect mutualisms. New Phytol. 172:412–428.
- Canard, E., N. Mouquet, L. Marescot, K. Gaston, D. Gravel, and D. Mouillot, 2012. Emergence of structural patterns in neutral trophic networks. Plos One 7:e38295.
- Cavalli-Sforza, L. L. and A. W. Edwards, 1967. Phylogenetic analysis. models and estimation procedures. Am J Hum Genet. 19:233–257.
- Darwin, C., 1859. On the origin of the species. Murray.
- 394 , 1862. Fertilisation of Orchids. Murray.
- Doebeli, M., 2011. Adaptive diversification. Princeton University Press, Princeton, NJ.
- Encinas-Viso, F., T. Revilla, and R. Etienne, 2012. Phenology drives mutualisic network structure and diversity. Ecol Lett 15:198–208.

- Ferdy, J., L. Despres, and B. Godelle, 2002. Evolution of mutualism between globeflowers and their pollinating flies. Journal of Theoretical Biology 217:219–234.
- Ferriere, R., M. Gauduchon, and J. L. Bronstein, 2007. Evolution and persistence of obligate mu-
- tualists and exploiters: competition for partners and evolutionary immunization. Ecology Letters
- 402 10:115-126.
- Gavrilets, S., R. Acton, and J. Gravner, 2000. Dynamics of speciation and diversification in a metapopulation. Evolution 54:1493–1501.
- Gomulkiewicz, R., S. L. Nuismer, and J. N. Thompson, 2003. Coevolution in variable mutualisms.
- The American Naturalist 162:S80–S93.
- 407 Gomulkiewicz, R., J. N. Thompson, R. D. Holt, S. L. Nuismer, and M. E. Hochberg, 2000. Hot
- spots, cold spots, and the geographic mosaic theory of coevolution. The American Naturalist
- 156:156-174.
- Guimarães, P., P. Jordano, and J. Thompson, 2011. Evolution and coevolution in mutualistic net-
- works. Ecol Lett 14:877–885.
- Hoban, S., O. Gaggiotti, C. Consortium, and G. Bertorelle, 2013. Sample planning optimization tool
- for conservation and population genetics (spotg): a software for choosing the appropriate number
- of markers and samples. Methods in Ecology and Evolution 4:209–303.
- Howe, H. and J. Smallwood, 1982. Ecology of seed dispersal. Ann Rev. Ecol. Syst. 13:201–228.
- Janzen, D., 1980. When is it coevolution? Evolution 34:611–612.
- Jones, E., R. Ferrière, and J. Bronstein, 2009. Eco-evolutionary dynamics of mutualists and ex-
- ploiters. Am Nat 174:780–794.
- Jordano, P., J. Bascompte, and J. Olesen, 2003. Invariant properties in coevolutionary networks of
- plant-animal interactions. Ecol. Lett. 6:69–81.
- Jousselin, E., J.-Y. Rasplus, and F. Kjellberg, 2003. Convergence and coevolution in a mutualism:
- evidence from a molecular phylogeny of ficus. Evolution 57:1255–1269.

- Kondrashov, A. and M. Shpak, 1998. On the origin of species by means of assortative mating. Proc.
- R. Soc. Lond. B 265:2273–2278.
- Krishna, A., P. Guimarães, P. Jordano, and J. Bascompte, 2008. A neutral-niche theory of nestedness
- in mutualistic networks. Oikos 117:1609–1618.
- Law, R., J. L. Bronstein, and R. Ferrière, 2001. On mutualists and exploiters: plant-insect coevolution
- in pollinating seed-parasite systems. Journal of Theoretical Biology 212:373–389.
- Losos, J., 2011. Convergence, adaptation, and constraint. Evolution 65:1827–1840.
- Lynch, M., 2007. The evolution of genetic networks by non-adaptive processes. Nature Reviews
- 431 Genetics 8:803–813.
- Maglianesi, M., N. Bluthgen, K. Böhning-Gaese, and M. Schleuning, 2014. Morphological traits de-
- termine specialization and resource use in plant-humming bird networks in the neotropics. Ecology
- 95:3325-3334.
- Melián, C. J., D. Alonso, D. Vázquez, J. Regetz, and S. Allesina, 2010. Frequency-dependent selection
- predicts patterns of radiations and biodiversity. PLoS Comput Biol 6:e1000892.
- 437 Moritz, C. and C. J. Schneider, 1992. Evolutionary relationships within the Ensatina eschscholtzii
- complex confirm the ring species interpretation. Syst Biol 41:273–291.
- Nei, M., T. Maruyama, and C.-I. Wu, 1983. Models of evolution of reproductive isolation. Genetics
- 103:557-579.
- Nuismer, S. L., P. Jordano, and J. Bascompte, 2012. Coevolution and the architecture of mutualistic
- networks. Evolution 67:338–354.
- Ohta, T., 2002. Near-neutrality in evolution of genes and gene regulation. Proc Natl Acad Sci USA
- 99:16134-16137.
- Olesen, J. and P. Jordano, 2002. Geographic patterns in plant-pollinator mutualistic networks.
- Ecology 89:2416-2424.

- Olesen, J. M., J. Bascompte, H. Elberling, and P. Jordano, 2008. Temporal dynamics in a pollination network. Ecology 89:1573–1582.
- <sup>449</sup> Pérez, F. and B. Granger, 2007. IPython: a System for Interactive Scientific Computing. Comput.
- sci. Eng. 9:21–29.
- R Core Team, 2013. R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria. URL http://www.R-project.org/.
- Rambaut, A. and N. Grassly, 1997. Seq-gen: An application for the monte carlo simulation of dna
- sequence evolution along phylogenetic trees. Computer Applications In The Biosciences 13:235–
- 455 238.
- Schluter, D., 2009. Evidence for ecological speciation and its alternative. Science 323:737–741.
- Seehausen, O., R. K. Butlin, I. Keller, C. E. Wagner, J. W. Boughman, P. A. Hohenlohe, C. L.
- Peichel, G. P. Saetre, et al., 2014. Genomics and the origin of species. Nature Reviews Genetics
- 459 15:176–192.
- Solé, R., R. Ferrer-Cancho, J. Montoya, and S. Valverde, 2002. Selection, tinkering, and emergence
- in complex networks. Complexity 8:20–33.
- Stang, M., P. Klinkhamer, and E. van der Meijden, 2006. Size constraints and flower abundance
- determine the number of interactions in a plant-flower visitor web. Oikos 112:111–121.
- Stang, M., P. Klinkhamer, N. Waser, I. Stang, and E. van der Meijden, 2009. Size-specific interaction
- patterns and size matching in a plant-pollinator interaction web. Ann Bot 103:1459–1469.
- Stayton, C. T., 2008. Is convergence surprising? An examination of the frequency of convergence in
- simulated datasets. J. Theo. Biol. 252:1–14.
- Thompson, J. N., 1999. Specific hypotheses on the geographic mosaic of coevolution. The American
- 469 Naturalist 153:S1–S14.
- 470 ———, 2009. The coevolving web of life. The American Naturalist 173:125–140.

- Thompson, J. N. and B. M. Cunningham, 2002. Geographic structure and dynamics of coevolutionary selection. Nature 417:735–738.
- True, J. R. and E. S. Haag, 2001. Developmental system drift and flexibility in evolutionary trajectories. Evolution & Development 3:109–119.
- Wagner, A., 2008. Neutralism and selectionism: a network-based reconciliation. Nature Reviews

  Genetics 9:965–974.
- Waser, N., L. Chittka, M. Price, N. Williams, and J. Ollerton, 1996. Generalization in pollination systems, and why it matters. Ecology 77:1043–1060.

# Supplementary Information

# The model: evo-evolutionary trait diversification in mutualistic

# 481 networks

To study the role of spatial structure and plant-animal trait diversification on convergence, complementarity and nestedness, we start by considering a landscape consisting of several individual plants (P) and animal pollinators (A). We model obligate mutualism for both partners and the number of individuals at each trophic level is fixed and equal to the environmental carrying capacity for the given community. Genetic, phenotypic and species composition change in time and space due to population dynamics. At each time step replacement of dead individuals are followed by the offspring of the same or another species (the key terms and model steps are summarized in figure 1 and table 1, respectively). In this section we explain how we model populations, traits, and diversification.

# Population dynamics

503

- Our model is a stochastic individual-based model with overlapping generations. The population consists of  $J_P$  and  $J_A$  haploid gonochoric (i.e. separated sexes) individuals with an explicit genome of size L each and equal sex ratios at the outset. The genome of each individual is composed of L-1 loci determining assortative mating and one neutral locus. Both plant and animal population reproduce sexually and are spatially structured. First an individual plant k and an animal k' are randomly removed (death). They are replaced by reproduction. There are four conditions for producing viable offspring for the plants and animals. These four conditions are 1) geography, 2) genetics, 3) obligate mutualism and 4) morphology:
- 1. Geography: a female and a male individual within the plant and animal populations are randomly chosen among all females and males within a maximum distance,  $d_{max}$ , of the dead plant k and dead animal k'. This requires two geographic distance matrices, one for plants,  $D^P = [d_{ij}^P]$ , and one for animals,  $D^A = [d_{ij}^A]$ , containing all the pairwise distances.
  - 2. Genetics: to produce viable offspring the female and the male in the plant and animal popula-

tions must have a genetic similarity value of the assortative mating loci,  $q_{QO}$ , higher than the minimum genetic similarity to have viable offspring,  $q_{min}$ ,  $(q_{QO} > q_{min})$ . This process reflects assortative mating and it requires two genetic similarity matrices, one for plants,  $Q^P = [q_{ij}^P]$ , and one for animals,  $Q^A = [q_{ij}^A]$ , containing all the pairwise similarity values.

- 3. Obligate mutualism: in order to have a mutualistic interaction the geographic distance between the female, Q (animal or plant), and one of the two male animal or plant individuals, represented here as j, must be lower than the maximum distance,  $d_{max}^{PA}$ ,  $(d_{jQ}^{PA} < d_{max}^{PA})$ . This requires one geographic plant-animal distance matrix,  $D^{PA} = [d_{ij}^{PA}]$ , containing all the pairwise distances.
- 4. Morphology: the female plant need the presence of an animal pollinator with a larger or equallysized proboscis than the corolla of the plant, thus the phenotype of the selected pollinator,
  represented here as j, must satisfy  $z_{\mathbb{Q}}^{P} \leq z_{j}^{A}$ . This requires two phenotype distributions, one
  for the plants,  $Z^{P} = [z_{i}^{P}]$  and one for the animals,  $Z^{A} = [z_{i}^{A}]$ . This trait-mediated interaction
  relationship is a variation on the "phenotype differences" model Nuismer et al. (2012).

The offspring arising from this mating event will occupy the geographic position of the just deceased individuals.

## Diversification dynamics

504

505

506

507

508

509

511

To quantify speciation events we calculate the genetic distance between each pair of individuals based on the assortative mating loci. We represent the genome of each individual by a sequence of L-1 loci, where each locus can be in two allelic states, +1 or -1. The assortative mating loci of each plant individual i in a population of size  $J_P$  is represented as a vector:  $S^i = (S^i_1, S^i_2, ..., S^i_L)$ , where  $S^i_u$  is the  $u^{th}$  locus of individual i. The genetic similarity based on assortative mating loci between individuals i and j is calculated as the sum of identical loci across the genome

$$q_{ij}^{P} = \frac{1}{L} \sum_{u=1}^{L} S_{u}^{i} S_{u}^{j} \tag{1}$$

where  $q_{ij}^P \in \{-1, 1\}$  with the genetic similarity matrix,  $Q^P = [q_{ij}^P]$ , containing all pairwise genetic similarity values for plants (the same for animals,  $Q^A = [q_{ij}^A]$ ). The genome of the offspring is obtained by a block cross-over recombination of a female genome,  $S^{\mathbb{Q}}$ , and a male genome,  $S^{\mathbb{Q}}$ , where a locus

l in the genome of the parents is randomly chosen partitioning the genome of each individual in two blocks. All genes beyond that locus l in either genome are swapped between the two parents and eventually form two new genomes. One of the two new genomes is randomly chosen for the offspring. The offspring's genome undergoes mutations at mutation rate  $\mu$ . Figure 1 describes the recombination-mutation process.

At the beginning of the simulations all individuals are genetically identical (all  $q_{ij}^P$  and  $q_{ij}^A = 1$ ); 534 hence they are all able to mate and produce viable offspring regardless of their spatial location. 535 The genetic similarity between individuals of a guild can be visualized as an evolutionary spatial 536 graph (Melián et al., 2010), where nodes correspond to individuals and the edges correspond to 537 the geographic distances between a pair of individuals satisfying the genetic similarity condition 538 for mating,  $q_{ij}^P(q_{ij}^A) > q_{min}$ . The connectance of the graph will decrease when generations move 539 forward because of the processes described in the previous section: 1) spatial constraints for mating 540 driving assortative mating and dispersal limitation; 2) genetic divergence driven by the threshold 541 for mating (incompatibilities), mutation and recombination forming the genome of the offspring; 3) 542 obligate mutualistic interactions driven by spatial proximity of individuals of the other guild, and 4) 543 morphological constraints following the phenotype differences scenario. 544

These four set of processes drive genetic divergence and speciation. We followed the species definition of Nei et al. (1983), which states that species are groups of individuals that are reproductively isolated and can interbreed to produce fertile offspring. In our model this is realized through allowing two individuals to mate successfully if their genetic similarity value is larger or equal to the minimum value,  $q_{min}$ . Thus, speciation is defined as a group of genetically related individuals, where two individuals in a sexual population can be conspecific while also being incompatible, as long as they can exchange genes indirectly through other conspecifics (de Aguiar et al., 2009; Melián et al., 2010). This is the definition of 'ring species' (Moritz and Schneider, 1992).

Genetic divergence will eventually produce the formation of two genetically incompatible clusters of individuals, i.e. two species. This speciation process, also called 'fission-induced' speciation, continues to form more clusters and genetic divergence between individuals of different species increases. However, the diversification dynamics will fluctuate due to random extinctions (death of last individual of a species). A stochastic balance between speciation and extinction is eventually reached giving the final steady-state of the metacommunity.

### Quantitative trait dynamics

We model each individual plant and animal with a quantitative trait,  $z^P$  and  $z^A$ , respectively. The processes described in figure 1 govern two quantitative traits, one for each guild: proboscis or bill length  $(z_i^A)$  in pollinators and corolla length  $(z_i^P)$  in plants. The quantitative trait of offspring is determined by the additive genetic effects of the genome (i.e. no epistasis) after the process of randomly choosing one of the two new genomes and mutation (figure 1) plus a normally distributed environmental effect,  $\epsilon$ ,  $\mathcal{N}(\mu_{\epsilon} = 0, \sigma_{\epsilon}^2 = 1)$  (Guimarães et al., 2011). The phenotype of the plant offspring i is  $z_i^P = g_i^P + \epsilon$  and the genetic component  $(g_i^P)$  of the phenotype of offspring i is

$$g_i^P = L + S_o^i \tag{2}$$

with  $S_o^i = \sum_{u=1}^L S_u^i$ . Hence  $g_i^P$  is calculated as the sum of alleles across the genome (Kondrashov and Shpak, 1998) plus the number of loci to avoid negative trait values ( $g_i^A$  is calculated similarly for animals). We assumed that the magnitude of the influence (i.e., effect sizes) of any given locus on this quantitative trait is equal across all the loci (Seehausen et al., 2014). This means that two individuals with a different combination of alleles in the genome can express the same quantitative trait (Losos, 2011).

#### Neutral locus evolution

We considered a neutral locus to estimate genetic divergence among species for the calculation of 574 convergent events (see section "Evolutionary convergence"). The neutral locus is located at the end 575 of the genome at the position L and it has k possible allelic states. The locus is completely unlinked 576 from the rest of the genome that contains the assortative mating loci. We used low mutation rates for 577 this neutral locus,  $\mu_{neutral} = 10^{-7}$ , and the k allele mutation model (i.e. model in which each allele 578 can mutate to any of the other k-1 possible alleles with equal probability (Hoban et al. (2013)). We 579 used the Cavalli-Sforza distance to calculate the matrix of genetic distances among species (Cavalli-580 Sforza and Edwards (1967)). 581

# Convergence, complementarity and nestedness

583 Evolutionary convergence

#### 584 Phenotypic similarity

The phenotypic similarity for plants  $(p_{ij}^P)$  between individual i and j is defined as

$$p_{ij}^{P} = 1 - \frac{|z_{i}^{P} - z_{j}^{P}|}{z_{max}^{P}}$$
 (3)

where  $z_i^P$  and  $z_j^P$  are the phenotypic similarity values of i and j, respectively, and  $z_{max}^P$  is the maximum value of the phenotype distribution,  $Z^P$ . Thus, the elements  $p_{ij}^P \in \{0,1\}$  of the phenotypic similarity matrix,  $\mathcal{P}^P = [p_{ij}^P]$  represent all pairwise values for plants (the same for animals,  $\mathcal{P}^A = [p_{ij}^A]$ ).

#### 589 Mean genetic and phenotypic species similarity

We define evolutionary convergence as the similarity between average species phenotypes from distantly related species. We assume that two species are distantly related, in phylogenetic terms, if they do not come from a direct common ancestor, i.e. they are not sister species. To exclude sister species from the analysis we need to calculate the mean genetic similarity among species of the same guild. The mean genetic similarity between a plant species k and a plant species l is

$$\hat{q}_{kl}^{P} = \frac{1}{n_k n_l} \sum_{i=1}^{n_k} \sum_{j=i}^{n_l} q_{ij}^{P} \tag{4}$$

where  $q_{ij}^P$  is the genetic similarity between an individual i of plant species k and an individual j of plant species l, and  $n_k$  and  $n_l$  are the absolute abundances of plant species k and l, respectively. The elements  $\hat{q}_{kl}^P$  form the matrix  $Q_s^P = [\hat{q}_{kl}^P]$  from which the sister species of each species in the guild can be identified (The elements for animals,  $Q_s^A = [\hat{q}_{kl}^A]$ , are calculated in the same way as we did for the plants). To calculate evolutionary convergence we need to know the average phenotypic similarity between two species. We define phenotypic similarity between species k and l as

$$\hat{p}_{kl}^P = \frac{1}{n_k n_l} \sum_{i=1}^{n_k} \sum_{j=i}^{n_l} p_{ij}^P \tag{5}$$

which is analogous to the definition of eq. 4, but now considering phenotypes instead of genotypes.

This will build a species phenotypic similarity matrix  $P_s^P = [\hat{p}_{kl}^P]$  (the species phenotypic similarity matrix,  $P_s^A = [\hat{p}_{kl}^A]$ , is calculated analogously for the animals). We then focus on each species in turn and exclude its sister species to avoid cases of parallel evolution to calculate the number of convergences related to the focal species. We define a focal plant species k and a non-sister plant species l to be convergent if phenotypic similarity between them is higher than between focal and sister species  $(\hat{p}_{k,sister}^P < \hat{p}_{kl}^P)$  and higher than a certain phenotypic threshold value  $t_{conv}$  ( $\hat{p}_{kl}^P > t_{conv}$ ); convergent species is calculated analogously for the animals).

#### Evolutionary complementarity

#### 610 Plant-animal interactions

609

In addition to the genetic and geographic constraints for mating, we consider two other conditions for plants and animals: obligate mutualism and morphological constraints. Obligate mutualism applies to the plants and animals to reproduce but the morphological constraints only apply to plants. We therefore need a geographic distance matrix,  $D^{PA}$ , to describe the geographic distance between plant and animal individuals. Plant-animal mutualistic interactions are here described as follows: plants benefit from the presence of specific pollinators that are able to pollinate them and animals benefit from the presence of plants that provide resources for them. Thus, we have two extra conditions for mating:

- 1. Female plants need the presence of an animal pollinator (i.e., male and female represented as j) within a close distance,  $d_{jQ}^{PA} < d_{max}^{PA}$ . The pollinator must have a larger or equally-sized proboscis than the corolla of a plant,  $z_{Q}^{P} \leq z_{j}^{A}$ . This corresponds to a morphological constraint for individual interactions observed between plant and pollinator species (Stang et al., 2009, 2006).
- 2. Animals need the presence of a plant (male or female represented as j) within a close geographic distance,  $d_{jk}^{PA} < d_{max}^{PA}$ .

# **Tables**

Table 1: Glossary of mathematical notation and parameter values  $\,$ 

Notation	Definition	Values
$J^P,J^A$	Effective population size of plants (P) and animals (A)	1,000
$d_{ij}^P, d_{ij}^A$	Geographical pairwise distance plants (P) and animals (A)	variable
$d_{max}$	Maximum geographical distance to mate and disperse	0.3
$D^P, D^A$	Geographic distance matrix with all $d_{ij}^P$ and $d_{ij}^A$ values	variable
$d_{ik}^{PA} \ d_{max}^{PA} \ D^{PA}$	Geographical distance between plant $i$ and animal $k$	variable
$d_{max}^{PA}$	Maximum geographical distance to find a mutualistic partner	0.3
	Geographic distance matrix with all the $d_{ik}^{PA}$ values	variable
$\begin{array}{c} q_{ij}^P, q_{ij}^A \\ Q^P, Q^A \end{array}$	Genetic similarity between ind. $i$ and $j$ in (P) and (A)	variable
$Q^{P}, Q^{A}$	Genetic similarity matrix with all the $q_{ij}^P$ and $q_{ij}^P$ values	variable
$q_{min}$	Minimum genetic similarity to have viable offspring	0.97
$z_i^P, z_i^A \ Z^P, Z^A$	Quantitative trait of ind. $i$ in (P) and (A)	variable
$Z^P, Z^A$	Quantitative trait distribution in (P) and (A)	variable
$p_{ij}^P, p_{ij}^A \ \mathcal{P}^P, \mathcal{P}^A$	Phenotypic similarity between ind. $i$ and $j$ in (P) and (A)	variable
$\mathcal{P}^{P},\mathcal{P}^{A}$	Phenotypic similarity matrix with all the $p_{ij}^P$ and $p_{ij}^A$ values	variable
L	Size of the genome	150
$g_i^P,g_i^A$	Genetic component of phenotype of offspring in (P) and (A)	variable
$\epsilon$	Environmental component of phenotype of offspring	$\mathcal{N}(0,1)$
$\mu$	Mutation rate per locus	$10^{-4} - 10^{-2}$
$\hat{q}_{kh}^P, \hat{q}_{kh}^A  Q_s^P, Q_s^A$	Mean genetic simil. between species $k$ and $h$ in (P) and (A)	variable
$Q_s^P, Q_s^A$	Species genetic simil. matrix with all $\hat{q}_{kl}^P$ and $\hat{q}_{kl}^A$ values	variable
$\hat{p}_{kh}^P, \hat{p}_{kh}^A$	Mean phen. simil. between species $k$ and $h$ in (P) and (A)	variable
$P_s^P, P_s^A$	Species phen. simil. matrix with all $\hat{p}_{kl}^P$ and $\hat{p}_{kl}^A$ values	variable
$\hat{p}_{kh}^{PA}$	Mean trait similarity plant species $k$ and animal species $h$	variable
$P_s^P, P_s^A \\ \hat{p}_{kh}^{PA} \\ P_s^{PA}$	Phenotypic simil. matrix with all $\hat{p}_{kh}$ values	variable
$^{h}conv$	Phenotypic threshold to calculate convergence events	variable
$^{h}comp$	Phenotypic threshold to calculate complementarity events	variable

# **Figures**

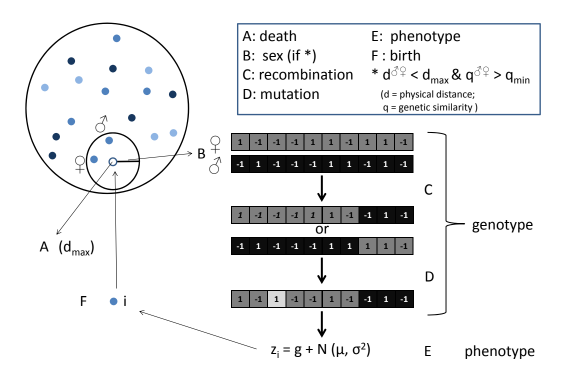


Figure 1: The death-birth cycle per time step. Individuals are represented as filled circles and blue tones represent variation of phenotypes. (A) an individual k is randomly selected to die and leaves an empty location in the landscape. (B) a female individual,  $\mathcal{P}$ , is randomly selected among all females satisfying the condition  $d_{kQ} < d_{max}$ . We then choose randomly a male,  $\sigma$ , among all males satisfying  $d_{k\sigma} < d_{max}$  and  $q_{Q\sigma} > q_{min}$  with  $q_{min}$ , the minimum genetic similarity required for mating. In addition to these two constraints, two more are required to complete mating. For the condition of obligate mutualism, the geographic distance between the female (animal or plant), and an animal (or plant) individual j, must satisfy  $d_{iQ}^{PA} < d_{max}^{PA}$ . Finally, female plants need the presence of an animal pollinator with a larger or equally-sized proboscis than the corolla of the female plant, thus individual pollinators represented as j, must satisfy  $z_{Q_P} \leq z_{j_A}$ . In (C) and (D) we calculate the genome of the new offspring once these constraints are satisfied. (C) Genomes are composed of L loci where each locus can be in two allelic states (-1, 1) and undergo block crossover recombination between female (dark gray) and male (black). A position l in the genome of the parents is randomly chosen partitioning the genome in two blocks. All genes beyond the l locus in either organism's genome is swapped between two parents and two new genomes are formed. (D) One of the two new genomes is randomly chosen for the offspring i,  $S_o^i$ , and it might undergo mutation (light gray). (E) The phenotype expression of offspring i is  $z_i = g_i + \epsilon$  with  $g_i = L + S_o^i$  and  $\epsilon$  are the genetic and environmental component of the phenotype, respectively. (F) The offspring i occupies the site of the dead individual k.

# $\mathbf{Q}_{\mathrm{s}}$ matrix

	а	b	С
a		0.85	0.97
b			0.89
С			

# $P_s$ matrix

	а	b	С
а		0.98	0.90
b			0.92
С			

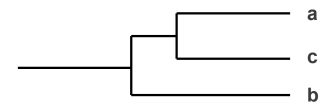


Figure 2: A simple example of evolutionary convergence using species a, b and c. The upper matrix  $(Q_S = [\hat{q}_{kl}])$  shows species a and c are genetically closely related,  $\hat{q}_{ac} = 0.97$ , while genetically distant from species b ( $\hat{q}_{ab} = 0.85$ ,  $\hat{q}_{cb} = 0.89$ ). A clear description of these genetic relationships can be represented with a cluster tree or dendrogram, as shown in the lower part of the figure. Thus, we establish that species a and c are sister species. The species phenotypic similarity matrix,  $P_S = [\hat{p}_{kh}]$  shows that species a and b are phenotypically highly similar ( $\hat{p}_{ab} = 0.98$ ) and highly genetically dissimilar ( $\hat{q}_{ab} = 0.85$ ) (i.e. more than the average intraspecific genetic similarity or sister species 0.97), indicating an event of evolutionary convergence.

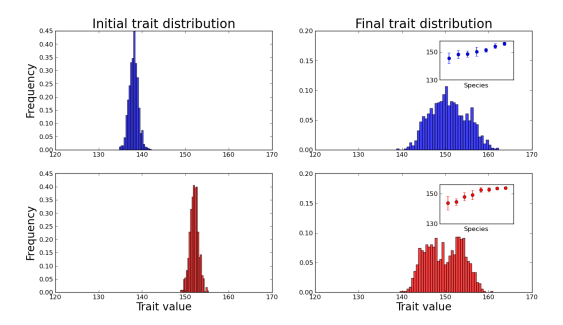


Figure 3: Changes in trait distribution of plants (top, blue) and animals (bottom, red). Left and right panels show the initial and final trait distribution, respectively. The insets in the right panels show the mean trait and standard error for each species sorted from the most common to the most rare. Initial trait distributions changed towards higher variance, and in most replicates, towards bimodal distribution in both guilds. Shown is the outputs from one replicate with parameters values  $q_{min} = 0.97$ ,  $d_{max} = d_{max}^{PA} = 0.3$ ,  $\mu = 5 \times 10^{-3}$  and  $J_P = J_A = 1,000$ .

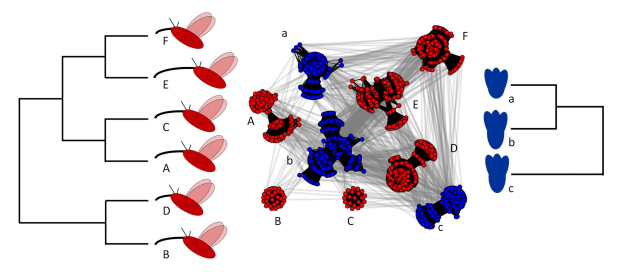


Figure 4: Evolutionary convergence and complementarity in plant-pollinator networks. Trees at the left and right side show genetic similarities between animal (red) and plant (blue) species, respectively. Mean species trait, proboscis and corolla length, is sketched with cartoons next to their respective position in the trees. Animals, composed by six species, have two evolutionary convergence events (A-B and F-D). Plants, composed by three species, have one convergent event (b-c). The central part of the figure shows the network of plantanimal interactions, where each node (colored filled circles) represents an individual. The network is composed of two types of links: genetic relatedness links (black solid) forming clusters that represent species and plant-animal individual-based interaction links (gray). The network shows variability in terms of genetic relatedness and plant-animal interactions within a species (i.e. high intraspecific variability). This figure is an example from one simulation. Parameters used are as in figure 3,  $q_{min} = 0.97$ ,  $d_{max} = d_{max}^{PA} = 0.3$ ,  $\mu = 5 \times 10^{-3}$  and  $J^P = J^A = 1,000$ .

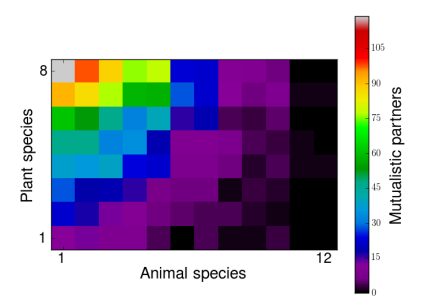


Figure 5: Plant-animal species interaction network. Plant species are represented in rows and animal species in columns. The color gradient indicates the number of mutualistic partners (i.e. individuals interacting) shared between plant and animal species. This matrix comes from one replicate with nine plant and thirteen animal species. The network shows high level of nestedness (N=0.72) and intermediate level of connectance (C=0.5). Parameters used are  $q_{min}=0.97$ ,  $d_{max}=d_{max}^{PA}=0.3$ ,  $\mu=5\times10^{-3}$  and  $J_P=J_A=1,000$ .

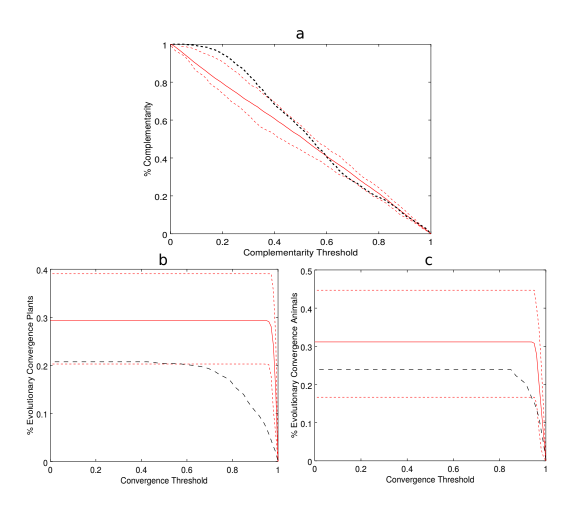


Figure 6: Comparison of model's predictions with estimations of convergence and complementarity from an empirical data of a plant-pollinator community. a) The proportion of complementarity events (y-axis) as a function of the complementarity threshold (x-axis) for the empirical data (dotted black line) and for the model (continuous and dotted red lines represent mean, 0.05 and 0.95 CI values, respectively). Predictions are within the CI for most complementarity threshold values. Empirical data deviates from model predictions for complementarity threshold values around 0.4 and lower. b) The proportion of convergence events in the plant community (y-axis, 69 species) as a function of the convergence threshold (x-axis) for the empirical data (dotted black line) and for the model (continuous and dotted red lines represent mean, 0.05 and 0.95 CI values, respectively). Convergence events in the empirical data strongly deviates from model predictions for convergence threshold values ranging between 1 and approx. 0.82. In that range, model predicts much higher proportion of convergence events than the empirical observations. c) The proportion of convergence events in the animal community (y-axis, 24 species) as a function of the convergence threshold (x-axis) for the empirical data (dotted black line) and for the model (continuous and dotted red lines represent mean, 0.05 and 0.95 CI values, respectively). Convergence events in the empirical data are within the CI of model predictions for most convergence threshold values.