Running Title: Genotype-species networks

Title: **Genotypic variation in foundation species generates network structure that may drive community dynamics and evolution**

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**ABSTRACT**

Although genetics in a single species is known to impact whole communities, little is known about how genetic variation influences species interaction networks in complex ecosystems. Here, we examine the interactions in a community of arthropod species on replicated genotypes (clones) of a foundation tree species, *Populus angustifolia* James (narrowleaf cottonwood), in a long-term, common garden experiment using a bipartite “genotype-species” network perspective. We combine this empirical work with a simulation experiment designed to further investigate how variation among individual tree genotypes can impact network structure. Three findings emerged: 1) the empirical “genotype-species network” exhibited significant network structure with modularity being greater than the highly conservative null model; 2) as would be expected given a modular network structure, the empirical network displayed significant positive arthropod co-occurrence patterns; and 3) furthermore, the simulations of “genotype-species” networks displayed variation in network structure, with modularity in particular clearly increasing, as genotypic variation increased. These results support the conclusion that genetic variation in a single species contributes to the structure of ecological interaction networks, which could influence ecological dynamics (e.g., assembly and stability) and evolution in a community context.

**Key words**: species interactions,foundation species, network ecology, genotype-species networks**,** modularity, nestedness, centralization, co-occurrence, common garden experiment

**INTRODUCTION**

The evolution of species interactions in complex communities is a central question at the interface of ecology and evolutionary biology and in understanding diverse theories of community structure ranging from random and neutral, to highly co-evolved (Clements 1916; Gleason 1926; Hubbell 2006; Shuster *et al.* 2006; Ricklefs 2008). Indirect effects in interaction networks have been theoretically (MacArthur 1955; May 1972; Allesina *et al.* 2011) and empirically (Bondavalli & Ulanowicz 1999; Bastolla *et al.* 2009; Bascompte *et al*. 2010) shown to influence the structure and dynamics of communities and ecosystems. For example, Bastolla *et al.* (2009) found support for decreased interspecific competition and increased co-existence of species in mutualistic networks. Network ecology has provided a means to approach the complexities of multi-species interactions by incorporating indirect effects, with some studies suggesting that ecological dynamics that are linked to evolutionary processes may lead to structural differences in mutualistic and trophic networks (Thébault & Fontaine 2011). In addition, phylogenetic analyses have revealed that shared evolutionary history contributes to ecological network structure (Rezende et al. 2007; Rafferty and Ives 2013). Collectively, these studies point to the importance of evolutionary processes in determining network structure and the potential for network structure to impact ecological dynamics.

The field of community genetics studies the genetic basis for interactions in complex communities primarily focusing on foundation species (i.e. species that define much of the structure of a community by creating locally stable conditions; Ellison *et al.* 2005). Community genetics research has demonstrated how genetic variation in a single species can determine the structure of whole communities and ecosystem processes (Wimp et al. 2005; Whitham *et al.* 2006; Crutsinger et al. 2006; Rowntree *et al.* 2011; Gugerli *et al.* 2013). This approach of focusing on species that are “central” (*sensu* Borrett 2013) in ecosystems (i.e., interacting with a large proportion of the species in a community) has provided a framework to address genetically based evolutionary questions in complex ecosystems (Whitham et al. 2006).

In this study, we merge network ecology and community genetics to examine how genetic variation in a foundation species influences the structure of ecological networks in complex communities. Specifically, we examine networks of foundation species genotypes interacting with associated species, which we refer to as “genotype-species” networks. Based on the findings of the many studies of the community level impacts of genotypic variation, we hypothesize that genotypic variation in foundation species produces patterns in the structure of interactions among genotypes of the foundation species and associated species. Furthermore, the genotype-species networks should exhibit structure similar to plant-herbivore networks.

To test these hypotheses, we first examine the structure of an empirical network of interactions between narrowleaf cottonwood (*P. angustifolia* James) and its associated canopy arthropod community using data from a long-term (20 year), common garden experiment with trees of known genetic identity. We use a conservative, null model approach to test for significant structural patterns in this “real” genotype-species network. Second, to further support and explore the effect of genotypic variation on network structure we conducted a simulation experiment using a community genetics model where we vary the intensity of the genotypic effect and explore three whole network metrics. Previous community genetics studies have focused on either community composition or trophic chains, and network ecology has incorporated an evolutionary perspective primarily via phylogenetic or model based simulation approaches. Our study seeks to empirically evaluate interaction network structure within an evolutionary framework by including intraspecific, heritable variation via the foundation species’ genotypes. Investigating the impact of genotypically based, and therefore heritable, variation on network structure provides a window into the dynamics of the architecture of the species interactions comprising complex communities within an evolutionary context.

**METHODS**

**Empirical network analysis**

A network of interactions between canopy arthropod species and genotypes of *Populus angustifolia* James (narrowleaf cottonwood) were collected from a common garden following the methods of Keith *et al.* (2010). Narrowleaf cottonwood is a foundation species (Ellison *et al.* 2005) of western riparian forests supporting a rich community of associated species (Whitham *et al.* 2006). In 1991 replicate clones of genotypes collected at random to represent the genetic variability of the Weber River Watershed (Utah, U.S.A.) and randomly planted in a common environment (Ogden Utah, U.S.A., latitude = 41.248146, longitude = −111.999830, elevation = 1302 m) to both minimize and randomize the effect of local environmental variation with respect to genotype (Martinsen *et al.* 2001). Surveys of arthropods of four replicate trees for each of 10 genotypes (n = 40 trees) were conducted in August of 2008 using timed sampling of similarly sized branches. Arthropods, including mainly herbivorous and sap feeding insects but also leaf modifying arthropods and mites (see Appendix Table A1), were identified to morphological species in the field and selected specimens were identified to their highest taxonomic resolution possible and vouchered in the Museum of Arthropod Biodiversity at Northern Arizona University, Flagstaff, AZ USA. Previous studies in this system have demonstrated community compositional effects of tree genotype on arthropods using an ordination-based approach (Keith *et al.* 2010); and in this study, we further explore the impacts of genotypic variation on community level interactions using a bipartite network approach, which examines networks where interactions typically occur between two main groups in a network (e.g. plants and pollinators). Using bipartite networks provides a set of analytical tools and hypotheses developed by the studies of plant-mutualist (Bascompte 2010), plant-herbivore (Thébault & Fontaine 2010) and other networks that we apply to a foundation tree, *Populus angustifolia*, and its arthropod community. As the genotypes were planted for the experiment, rather than through natural processes of establishment and selection by the local communities, we interpret these connections as directed effects of the genotype on the associated community. In addition, although these species comprise multiple trophic levels from herbivores to predators, all of these species were observed in close contact with the tree, typically using it as a resource. Hereafter, we use network terminology to refer to genotypes of narrowleaf cottonwood, as well as species in the network, as “nodes” and their connections, in this case the frequency of interactions between an individual tree of known genotype and an arthropod species, as “edges” comprising “genotype-species” networks.

We then analyzed the structure of this empirical genotype-species network. As no previous analyses have been conducted on genotype-species networks, we chose three network metrics (modularity, nestedness and centralization) and a measure of species co-occurrence (C-Score; Stone & Roberts 1990). As there are many characteristics of networks that could be examined that each have a multitude of metrics associated with them, we describe our chosen set of metrics and our use of them in more detail here. Modularity is a measurement of the degree to which the network is comprised of groups of nodes that are relatively more connected to each other, and for our analyses we used the method developed by Newman (2004) to find the maximally likely partitioning of the network graph using the “greedy optimization” algorithm. Nestedness was calculated for the observed network using the nestedness temperature metric developed by Atmar and Patterson (1993, see also Santamaría & Rodríguez-Gironés 2006), which measures the degree to which species tend to interact with subsets of the community across “parts” of the network. We used Freeman’s (1979) generalized degree centralization of a graph (*G*), for all nodes (*v*) in *G*, to compare how each network varied in tendency of one or a few nodes to dominate the edges in the network, and, thus, is a useful indicator of the distribution of interactions among all species in the network. We measured the co-occurrence patterns among the associated community using Stone and Robert’s (1990) C-score, which is the number of “checkerboard units” averaged across all pairs of species, which provides an average measure of species associations. The checkerboard unit is calculated as (*r*i-*S*ij)(*r*j-*S*ij), where *r* is the total number of occurrences for a given species and *S*ij is the total number of times a species pair occurs, and the C-score has been shown to have robust statistical properties (Gotelli 2001).

We tested for the significance of all network statistics and the C-score using a conservative null-model based randomization method. The null model consisted of a non-sequential sampling algorithm that constrains the matrices to maintain the marginal totals (i.e. the total interactions across all genotypes for a given species or all species for a given genotype), which has been recommended for bipartite networks (Blüthgen 2010). After an initial series of 100 “burn-in” permutations of the original matrix, a set of 5000 randomizations was generated and used to test the significance of the metrics. For each metric we calculated a non-directional p-value as the proportion of null model metrics that were equal to or more extreme than the original (i.e. “observed”) metric, and because this approach produces an empirical distribution it avoids the need to make assumptions of the distribution for our statistics of interest. We also calculated a standardized score as *z* = (*x* - μ)/σ, which in the co-occurrence literature is typically referred to as a standardized effect size (SES), where *x* is the observed metric and μ and σ are the mean and standard deviation for the null simulation metrics, respectively. It is important to note here that due to the limitation of the currently accepted the null modeling approaches, in particular the absence of well studied and implemented methods for weighted matrices (Gotelli & Ulrich 2012), it was necessary to reduce each network, including the empirical network, to an un-weighted (i.e. presence-absence) matrix.

**Simulating genotypic effects on network structure and co-occurrence patterns**

To generalize the patterns found with the empirical network, we conducted a simulation experiment testing the effect of genotypic variation on genotype-species network structure. We used the community genetics simulation method previously developed in Shuster *et al.* (2006). Briefly, this method uses a combination of both individual-based and mass-action modeling approaches. It starts by creating a population of individuals assigned a genotypic value (i.e. genotypes). Each genotype is comprised of multiple replicate individuals (e.g. clones randomly planted in a common garden), which are assigned phenotypes as a single numeric value for each individual sampled from a uniform distribution, which represents the traits of individual “trees.” Next, a set of species representing the associated community was assigned an average genotypic value that was then mapped to a numerical phenotypic value using an additive, bi-allelic genetic system, determining the abundance of each arthropod species on each tree. “Environmental” variance was introduced through random draws from a uniform distribution, simulating the introduction variance arising from non-genetically based sources. Last, mass-action methods were used to determine the equilibrium population value for each associated species given the phenotypic values for each foundation species individual and the carrying capacity of each associated community member (Appendix Eq. A.1).

We used this method to conduct simulations of communities with different levels of foundation species’ genotypic effect on the community. To do this, we adjusted the co-efficient determining the foundation species’ genotypic effect for each species in the associated community, which modifies the fidelity of the mapping of a tree genotype to phenotype to an arthropod species by varying the power of the co-efficient per the method of Shuster *et al.* (2006; Appendix Table A1). In addition, to explore the impact of variation in associated species abundances, we produced two sets of simulations, one (n = 80) with a carrying capacity (K) of 100 individuals (i.e. even species abundances), and a second set (n = 80) with carrying capacities for the dependent species selected from a Poisson distribution (λ = 5), ranging from K = 100 to K = 5 (i.e. un-even species abundances). This distribution was chosen for its similarity to the highly skewed, “un-even” distribution of species total abundances commonly observed in ecological communities (Blüthgen et al. 2008). As intended the diversities of the even communities (Shannon’s H = 3.25±0.05 SD) were greater than the un-even communities (Shannon’s H = 2.25±0.15 SD), which were very similar to the diversity of the empirical canopy arthropod community (Shannon’s H = 2.15). The simulated communities were generated using the same initial phenotypes for both the “foundation species” individuals (n = 100; 10 individuals for each of 10 genotypes) and the “associated species” (25 total). In line with previous simulations of community level genetic effects, our simulated communities displayed genetic impacts that were similar to the results of Shuster *et al.* (2006) for both the even and un-even simulations. The simulated trees differed in their individual phenotypic trait values (Fig. 2A) and composition of interactions among genotypes (Fig. 2B). To test and compare the effect of genotype on the four metrics, we used correlation analysis employing Kendall’s τ as our correlation coefficient to capture non-linear monotonic relationships.

**Simulation and analytical software**

All simulations and analyses were conducted in R version 3.0.2 (R Core Team 2014). Simulations were conducted using the *ComGenR* package (Lau 2015), and simulation scripts are publically available (<https://github.com/MKLau/cg_simulations>). Non-metric multidimensional scaling (NMDS) ordination and nestedness tests were conducted with *vegan*, and networks were plotted using *bipartite*.

**RESULTS**

Supporting our hypothesis that a genotype-species network would display non-random structure, in the field we found significant structure and co-occurrence patterns among associated species in the empirical network. The cottonwood genotype and canopy arthropod network (Fig. 1) had levels of modularity (observed = 0.20, *z* =3.82, *P =* 0.038), nestedness (observed = 38.17, *z* = -3.00, *P* = 0.003), centralization (observed = 0.31, *z* = 3.90, *P* = 0.003) and C-Score (observed = 0.47, *z* = 1.95, *P* = 0.025) that all differed from the null model based expectation. Modularity, centralization and C-Score all showed greater structure than the null expectation (i.e. more modules, greater centralization and more checkerboard units); however, the observed nestedness was lower than the null model.

As with the empirical network patterns, we observed genotypic effects in our simulations that supported the hypothesis of a genetic basis to network structure. All four metrics of simulated network structure and co-occurrence patterns increased with increasing genotypic effect, but for some metrics the effect depended on the evenness of the associated community. Even community network structure (i.e. even species abundances) and co-occurrence patterns showed increasing structure with increasing genotypic effect with modularity (Fig. 3a; τ = 0.82, *P* < 0.001), nestedness (Fig. 3b; τ = 0.83, *P* < 0.001), centralization (Fig. 3c; τ = 0.89, *P* < 0.001) and C-score (Fig. 3d; τ = 0.70, *P* < 0.001) all showing significant, positive correlations with genotypic effect.

The evenness of the associated communities affected how network structure changed with increasing genotypic effect. Three of the four metrics for the un-even community structure displayed clear significant correlations with genotypic effect: modularity (Fig. 3a; τ =0.69, *P* < 0.001), nestedness (Fig. 3b; τ = -0.28, *P* = 0.001) and centralization (Fig. 3c; τ = -0.87, *P* < 0.001). The C-score for the un-even community simulations across all levels of genotypic effect was not significant (Fig. 3d; τ = -0.01, *P* = 0.95); however, the relationship was clearly not monotonic and a Bayesian information theoretic breakpoint analysis of means using a normal distribution (Priyadarshana and Sofronov 2014) detected a significant break in C-scores corresponding to a genotypic effect of 0.008. Analyzing the un-even C-scores above and below this threshold separately, we found a significant correlation between C-score and genotypic effect of τ = 0.83 (*P* < 0.001) above the threshold. Below the threshold the correlation was not significant (τ = 0.06, *P* = 0.58).

**DISCUSSION**

Our results support the hypothesis that genotypic variation in a foundation species contributes to ecological network structure. In the field, we observed significant empirical network structure for all network metrics, including modularity, nestedness and centralization, which were also corroborated by significant co-occurrence patterns of species among trees. These findings emerged even though they were tested against a conservative null model that restricted tree genotype and associated species totals that greatly reduced the probability of detecting significant structure. Our simulation experiment, which manipulated both genotypic effect and community evenness, displayed significant effects of foundation species genotypic variation with all metrics in the “even” community simulations, also supporting the empirical findings. The “un-even”, simulated networks were also impacted by genotypic variation, but the presence of more frequently interacting species lead to a different response to the modularizing affect of genotype. Last, although the goal of the simulation was not to quantitatively replicate the empirical patterns, all of the metrics observed for the empirical network fell within the range of the simulated values (Fig 3, dashed lines). Taken together, these two lines of evidence (empirical and simulation based) support the conclusion that genotypic differences among individuals of a foundation species can generate interaction network structure in ecological communities.

**Genotypic variation contributes to network structure**

Significant network and co-occurrence patterns in the empirical genotype-species network indicate a strong genetic basis to ecological interaction network structure. In terms of the network structure metrics, modularity in plant-herbivore networks is often attributed to the specialization of a few species on a set of host species (Fontaine *et al.* 2011), while centralization generally can arise through the presence of a few generalists interacting with a larger number of specialist species (Sazima *et al.* 2010). Applying this to the empirical network, modularity could be arising from the specialization of some highly interactive arthropods on particular genotypes (e.g. the gall aphid, *Pemphigus betae*; Keith *et al*. 2010). It is likely that the same mechanism is contributing to species co-occurrence patterns, as the C-Score was much less than the null expectation, indicating that species tended to aggregate (i.e. co-occur), as would be the expected pattern if groups of species track similar genotypes and co-occur more frequently on preferred genotypes.

The results of the simulation experiment revealed the underlying impacts of genotypic variation in structuring interaction networks, further supporting the generality of the empirical findings. We found that when species total abundances are even, all metrics of network structure and co-occurrence patterns increase with the influence of variation among foundation species genotypes. A more complicated set of patterns was observed in the un-even network simulations. Modularity increased dramatically with increasing effect of genotype, as it did in the even simulations, however, both nestedness and centralization decreased, while the C-score increased only after a threshold of the influence of genotypic variation. The decreasing levels of nestedness and centralization, as well as the threshold before the increase in the C-score, are likely the result of the counterbalancing effects of genotype versus variation among species in the frequency of interactions. At lower levels of genotypic effects numerically dominant species abundances are determining the structure of the network. Then, as the influence of genotype increases, genotypic variation begins to form modules, and thereby both increasing co-occurrences of species within modules and breaking nestedness and decentralizing the network by compartmentalizing the influence of the dominant species. Although the simulations such as these are an obvious oversimplification of real ecological dynamics (Gotelli 2001; Blüthgen et al. 2008), as they do not incorporate the interactions among genotypes or among species (e.g. competition and facilitation), the behavior of these models is consistent with an underlying genetic basis for ecological network structure and can serve as a falsifiable framework for future studies.

**Genetic variation and the evolution of ecological networks**

Why should foundation species’ genotypic variation have effects on ecological network structure? Multiple empirical approaches support our findings. First, the multivariate plant resource/defense phenotype demonstrates extensive genetic variation in n-dimensional traits including chemical defenses, ontogeny, induction, phenology, sink-source relationships, productivity, leaf litter, and other genetics based traits (Larson & Whitham1997; Schweitzer *et al*. 2008; Grady *et al.* 2011; Holeski *et al.* 2012 and others). In combination, these factors result in each tree genotype differing from other tree genotypes in important traits that affect communities of organisms ranging from microbes to vertebrates. Thus, the greater the variation in the multivariate plant resource/defense phenotype, the greater the variation in the associated communities a species will support. Second, genotypes that are more similar in their genetics and the traits they express support more similar communities than those that differ in their genetics and the traits they express (genetic similarity rule of Bangert *et al.* 2008; Barbour *et al.* 2009; Zytynska *et al.* 2011). Third, genetically based interactions between strongly interacting species can create greater habitat variation, which in turn supports different communities on different genotypes (e.g., Bailey *et al.* 2006; Shuster *et al.* 2006; Keith *et al.* 2010; Lamit *et al.* 2011, 2014; Busby *et al.* 2015). One example of this, Busby *et al.* (2015), showed that the genetics-based interaction of the tree and a common pathogen played a major role in defining the arthropod community, which differed among genotypes based on the plant’s resistance to the pathogen. Thus, species trait values could be averaging over ecologically relevant variation in traits of individuals within species. In combination, these findings provide a mechanistic explanation for why trait differences among individual tree genotypes should direct community network structure.

A genetic basis to network structure has implications for the consequences of selection and evolution in a community context. Directional, stabilizing and divergent selection on a foundation species (e.g. pathogens and climate change) will alter the structure of these interaction networks in ways that can affect the outcome of selection, which occurs in a community context (Shuster *et al.* 2006; Turcotte et al. 2012; Fortuna et al. 2013; Thompson 2013). For example, in line with the predictions of selection imposed by “super-generalist” species in co-evolutionary dynamics (Guimarães *et al.* 2011), if directional selection on a foundation species selects for a “generalist” genotype, i.e. a genotype that interacts with a large number of species, we would predict a reduction in modularity and an increase in species co-occurrences, which would likely increase the frequency of species interactions. This would also increase the centralization of the network, potentially making the community less resistant to disturbance if it were to impose a different selection pressure that selects against the generalist genotype.

Our finding of a genetic basis to ecological interaction network structure has two consequences for community dynamics. First, these network level effects are only manifested in a community context, and because the structure of the network has pervasive effects on important community members (e.g. herbivores, mutualists, pathogens), changes in the community network are likely to feed back to influence the fitness of individual trees (e.g., Schweitzer et al. 2012). This conclusion is further supported by the rich experimental documentation that different genotypes support different community and ecosystem phenotypes (review by Whitham et al. 2012), and that major species’ interactions are genotype specific (e.g. interacting foundation species hypothesis; Keith *et al.* 2010; Busby *et al.* 2014). Furthermore, the contrasting pattern of lower nestedness with higher modularity observed in the empirical network and the simulations suggests the possibility that the tree genotype modules break up the nested structure of the arthropod community. If this is the case, the network level impact of tree genotype could simultaneously reduce the impact of herbivores and increase competition among species that affect tree vigor potentially leading to increased fitness, as the individual trees within each module would be comprised of a subset of the population thus reducing the resource base that the herbivores within that module would be utilizing. Second, the primary theories of community dynamics don't consider intraspecific variation at the community level and are thus missing a potentially important mechanism directing how communities and whole ecosystems change through time. Specifically, our results do not support neutral theory and its hypothesized species equivalence (Hubbell 2006) and the individualistic theory of Gleason in which species associations are the result of “coincidence” and that every species is a “law unto itself” (1926). As our local scale analyses of common garden studies demonstrate extensive community structure, and now interaction network modularity, they also are in opposition to Ricklefs’ (2008) concept of the “disintegration” of local communities in which “local coexistence can be understood only in terms of the distributions of species within entire regions…”. The existence of a genetic basis to network structure does support the community dynamic concepts of Clements (1916). However, the modular structure of interactions, which in the case of our genotype-species network are primarily the result of direct interactions rather than modification of environmental conditions, is not in line with the Clements’ concept of tightly bound communities, and are more “loosely” coupled, as one would expect in a dynamic system where species interactions can shift due to stochasticity and in response to a multiple dimensions of gradients both abiotic and biotic of which at least part is genetically based.

**Conclusion**

Understanding and predicting the responses of ecosystems is critical given the current pace of change ecosystems are currently experiencing (Scheffer et al. 2012). Species interactions play an integral role in ecosystems (Agrawal *et al.* 2007; Winfree *et al.* 2011) and how species interactions lead to evolutionary changes in participant species has been a topic of biology since Darwin’s writings on orchid pollinators (Darwin 1862). Studies of co-evolution between pairs and small groups of species have expanded the theory and support for evolutionarily dynamic species interactions (Ehrlich & Raven 1964; Jones *et al.* 2009; Thompson 2013). Studies of plant-mutualist and plant-herbivore networks have more recently provided a broader perspective, addressing more complex relationships among the many species in a community (Bascompte *et al.* 2010; Thébault & Fontaine 2010) and revealing the importance of the community context of both ecological and evolutionary dynamics (Thompson 2013). As a genetic basis of traits is a key requirement for evolution by natural selection, and given that multiple studies have demonstrated rapid evolution of species in response to climate change (reviewed in Parmesan 2006), as well as the importance of co-evolutionary dynamics in ecological networks (Guimarães et al. 2011; Moya-Laraño 2011), further exploration of the structure of genotype-species networks is key to understanding evolutionary dynamics in real ecosystems.

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**Appendix A**

**Equation A.1.** Simulation model mathematical description.

**Table A1.** Table of arthropod identifications from the empirical genotype-species network.

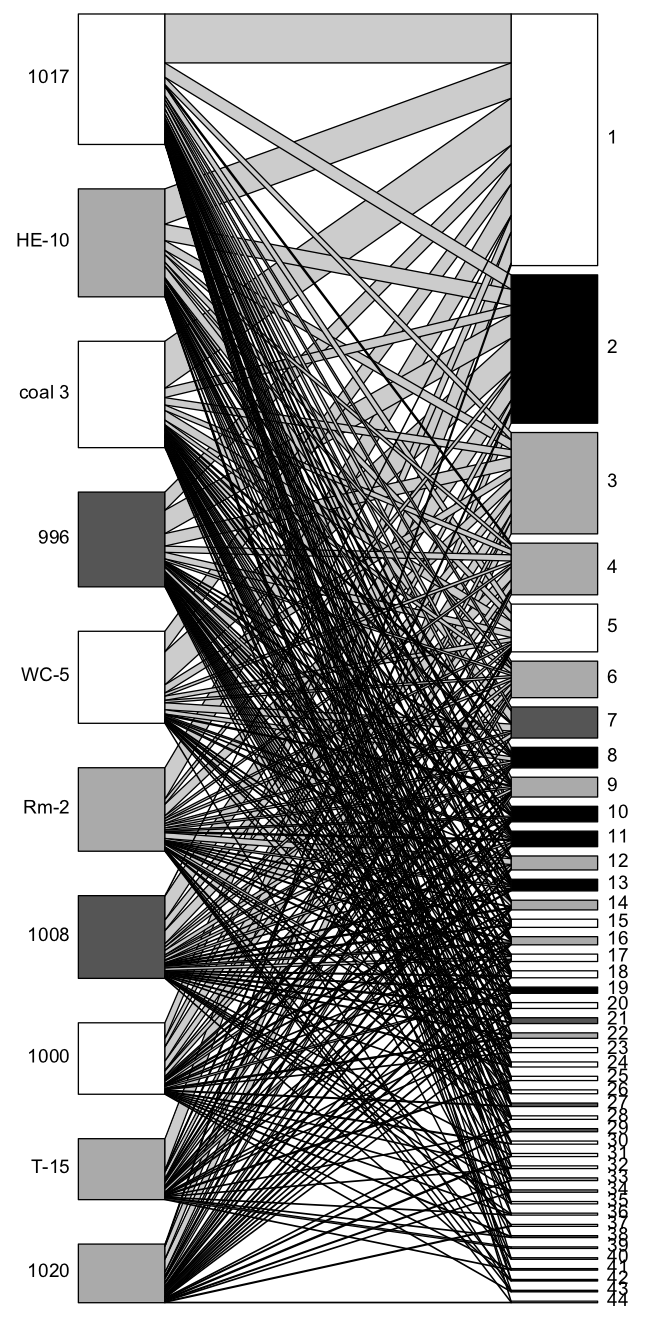
**Figure Legends**

**Figure 1**. The structure of the empirical bipartite network of 10 individual plant genotypes (left nodes) and 44 arthropod species (right nodes) living in the canopies of narrowleaf cottonwood (*Populus angustifolia*) in a common garden. Nodes are scaled by the marginal totals for genotypes (i.e. total arthropod abundance) and species (i.e. species total abundance). The nodes are arranged in order of increasing interactions, which corresponds to increasing centralization bottom to top, and are colored by module membership. Lines show interactions between genotypes and species scaled by the average abundance of each arthropod species across all individuals of a given genotype. Plant genotype names follow the naming presented in Keith *et al.* (2010) and taxonomic identification of the arthropod community is provided in the Appendix (Table A1).

**Figure 2.** Genotypic effect in simulated communities produced differences in individual phenotypic traits that contributed to variation in the associated community. The density plot (A) shows the variation in the simulated foundation “tree” phenotypic values among individuals with each line showing the distribution for a different genotype, while the NMDS ordination (B) plots the community composition of each genotype shown as the multivariate centroid for each genotype (±1 SD).

**Figure 3**. Bivariate plots showing how network and co-occurrence patterns of the simulated networks vary with increasing genotypic effect. When species’ total abundances were even (black), modularity (A), nestedness (B), centralization (C) and the C-Score (D) increased as the effect of foundation species genotype increased. When species total abundances were un-even (grey), only modularity showed a similar pattern of increasing structure (A), both nestedness (B) and centralization (C) decreased, while the C-Score (D) increased structure at higher levels of genotypic effect. Dashed lines show the values of these metrics for the empirical network, which all fall within the ranges of the simulations.

**Figure 1**



**Figure 2**



**Figure 3**

