Predicting species abundances in a grassland biodiversity experiment: Trade-offs between model complexity and generality

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Abstract
1. Models of natural processes necessarily sacrifice some realism for the sake of tractability. Detailed, parameter-rich models often provide accurate estimates of system behaviour but can be data-hungry and difficult to operationalize. Moreover, complexity increases the danger of ‘over-fitting’, which leads to poor performance when models are applied to novel conditions. This challenge is typically described in terms of a trade-off between bias and variance (i.e. low accuracy vs. low precision).

2. In studies of ecological communities, this trade-off often leads to an argument about the level of detail needed to describe interactions among species. Here, we used data from a grassland biodiversity experiment containing nine locally abundant plant species (the Jena ‘dominance experiment’) to parameterize models representing six increasingly complex hypotheses about interactions. For each model, we calculated goodness-of-fit across different subsets of the data based on sown species richness levels, and tested how performance changed depending on whether or not the same data were used to parameterize and test the model (i.e. within vs. out-of-sample), and whether the range of diversity treatments being predicted fell inside or outside of the range used for parameterization.
INTRODUCTION

“What a useful thing a pocket-map is!” I remarked.

“That’s another thing we’ve learned from your Nation,” said Mein Herr, “map-making. But we’ve carried it much further than you. What do you consider the largest map that would be really useful?”

“About six inches to the mile.”

“Only six inches!” exclaimed Mein Herr. “We very soon got to six yards to the mile. Then we tried a hundred yards to the mile. And then came the grandest idea of all! We actually made a map of the country, on the scale of a mile to the mile!”

“Have you used it much?” I enquired.

“It has never been spread out, yet,” said Mein Herr: “the farmers objected: they said it would cover the whole country, and shut out the sunlight! So we now use the country itself, as its own map, and I assure you it does nearly as well.”

—from Lewis Carroll, Sylvie and Bruno Concluded, Chapter XI, London, 1895.
A more comprehensive model might unpack the generalized interspecific term into pairwise interspecific interactions between all species (Carrara, Giometto, Seymour, Rinaldo, & Altermatt, 2015; Fort, 2018; Haity, Valdés, Tejera, Picasso, & Fort, 2017; Vandermeeren, 1969). Finally, more complex models might assume that species engage in 'higher-order interactions', for example, where the strength or direction of interactions among a subset of species are modified as a function of the abundance of other species in the community (Letten & Stouffer, 2019; Mayfield & Stouffer, 2017).

Each of these increasingly complex models has some empirical support. For example, neutral models with a single interaction coefficient have adequately explained some patterns in ecological communities (Hubbell, 2001). In other cases, observations are better matched by models with separate intra- versus interspecific terms, reflecting that intraspecific effects often far outweigh interspecific effects (Adler et al., 2018; Broekman et al., in press). Alternatively, in some microbial and plant systems, models that include pairwise interactions among species produce more accurate estimates of species abundances (Carrara et al., 2015; Clark, Lehman, & Tilman, 2018; Vandermeeren, 1969), potentially indicating species-specific impacts on resource availability (Tilman, 1982). Lastly, there is evidence from several systems that interaction strengths differ depending on community composition, suggesting the existence of 'higher-order' interactions (Bairey, Kelsic, & Kishony, 2016; Grilli, Barabás, Michalska-Smith, & Allesina, 2017; Levine, Bascompte, Adler, & Allesina, 2017; Mayfield & Stouffer, 2017; Wilbur, 1972). These differences can be mediated either directly, e.g. by changes in species densities or foraging strategies (Letten & Stouffer, 2019; Tilman, 1982; Tuck et al., 2018), or indirectly, e.g. via pathogens or herbivores (Kulmatiski, Beard, Grenzer, Forero, & Heavilin, 2016; Michalet et al., 2015; Weigelt et al., 2007). Nevertheless, the relative predictive abilities of these different models are rarely compared within a single system, let alone between systems, and the degree to which results can be extrapolated beyond the data used to fit them remains unclear.

The decision about whether to include complex species interactions in an ecological model is a particular example of the 'bias-variance trade-off' (Hastie, Tibshirani, & Friedman, 2017). Any given dataset will include both general phenomena that we might expect to see elsewhere, and particularities that occur in that dataset alone (Levis, 1968). When model performance is tested using the same data that were used to parameterize it, increased complexity can always reduce uncertainty (i.e., variance) by tuning the model to match observations. However, when different subsets of data are used to parameterize the model versus to test model performance, increased complexity can also lead to poorer performance for the testing subset. This phenomenon is known as 'over-fitting', and occurs when parameter tuning during the fitting process draws predictions towards these peculiarities (i.e., bias) (Wenger & Olden, 2012).

The degree to which a model is able to capture general versus particular phenomena can be assessed by dividing data into two or more (ideally independent) subsets, and testing whether models parameterized with one subset of data can make good predictions in the others (i.e. 'cross-validation') (Brewer, Butler, & Cooksley, 2016; Roberts et al., 2017; Wenger & Olden, 2012). Due to data scarcity, cross-validation is typically applied to randomly chosen subsets of data (e.g. 'k-fold cross validation') (Roberts et al., 2017). Overfitting is indicated when predictions for the subset of data that was not used for parameterization are substantially worse than those for the subset that was (i.e. when 'out-of-sample' goodness-of-fit is much lower than 'within-sample' goodness-of-fit). Given sufficient data availability, cross-validation can also be used to test predictive power under specific novel conditions (i.e. 'model transferability') (Wenger & Olden, 2012). By carefully choosing different subsets of data, it is possible to rigorously assess the circumstances under which a particular model is likely to provide accurate extrapolations. For example, to test how general a model is across space and environmental conditions, we might fit the model using data from one site, and test how well it works at another.

A more common approach for deciding whether to accept a more complex model is to compare within-sample goodness-of-fit of different models, with a term to penalize increases in model complexity. In ecology, such tests are often applied using AIC, or other similar information criteria (Aho, Derryberry, & Peterson, 2014; Brewer et al., 2016; Burnham, Anderson, & Huyvaert, 2011; Stone, 1977). Note that AIC is conceptually and theoretically related to cross-validation at the limit where a single observation is retained for testing, and the remainder of the data are used to parameterize a model (‘leave-one-out cross-validation’) (Stone, 1977). Thus, these methods are very efficient from the perspective of data requirements, but are often less useful for assessing model transferability (Brewer et al., 2016; Wenger & Olden, 2012).

Here, we test relationships between model precision and general- ality using a dataset collected as part of the Jena Experiment (‘dominance experiment’; Roscher et al., 2004; Weisser et al., 2017). In this experiment, a diversity gradient was established using nine locally abundant plant species, including replicates of all monocultures and two-species mixtures, and a selection of higher-diversity treatments. In the present analysis, our goal is to characterize the trade-off between good within-sample estimates (low variance) and good out-of-sample predictions (low bias), and to identify models that provide a reasonable compromise between these properties. Because we are interested in community dynamics, we focus on predictions of species-level above-ground biomass (in oven-dried g/m², hereafter ‘abundance’) rather than total biomass summed across species, which is the subject of several other studies of species interactions in biodi- versity experiments (Connolly et al., 2011; Kirwan et al., 2009).

We fitted a series of increasingly complex models, and compared their performance using three types of tests. First, to test for general effects of over-fitting, we fit models using data from monocultures, two-species mixtures, and nine-species mixtures. Thus, all communities within the dominance experiment fall within the range of the observed diversity treatments used to fit these models. Second, to identify models that might be suitable for extrapolation to higher-diversity communities, we fit the same series of models using only data from the minimum number of diversity treatments needed for parameterization (i.e. monocultures or two-species mixtures). Thus, higher diversity communities fall outside of the range of conditions...
used for parameterization. Finally, we compare these results with the ranking of models that would have been obtained had we followed a more typical model selection procedure based on the AIC of models fitted to the full dataset (i.e. all diversity levels). In accordance with the bias-variance trade-off, we expected that the most complex models would yield the best within-sample estimates, especially within the range of diversity levels used to parameterize them, whereas models of lower complexity would provide more generalisable out-of-sample predictions. Furthermore, we expected that AIC would not necessarily be an effective indicator of out-of-sample model performance, due to its strong correspondence to within-sample estimates.

2 | MATERIALS AND METHODS

2.1 | Overview

We designed six models which predicted species abundances in each year as a function of increasingly complex combinations of intra- and interspecific interactions. Details about models and corresponding biological hypotheses are discussed in Section 2.2. We then fit these models to field data, first using plots sown with one, two or nine species and second using plots containing one or two species. Details about the empirical data used to fit and test models are described in Section 2.3. Next, we used these models to predict species abundance in all plots, and compared within-sample and out-of-sample errors among models. We then tested model performance across sown diversity treatments, species, and years. Based on these tests, we identified models that provided the best out-of-sample predictions across these subsets of data. Details about model comparison methods are described in Section 2.4.

2.2 | Model structure

The six models represent increasingly complex hypotheses about intra- and interspecific interactions. All six were adapted from the same underlying model in Equations (1a)–(1b), and estimate the abundance of each species, in each year, as a function of its own abundance in the previous year (i.e. an autoregressive model), and the abundance of other species in the community in the previous year.

First, the intra-only model, Equation (2a), hypothesises that a species is only influenced by its own abundance in the previous year. This model largely serves as a ‘baseline’ and characterizes the predictive power of intraspecific density dependence and autoregressive processes. Second, the intra = inter model, Equation (2b), hypothesises that species are equally influenced by their own abundance and by the abundance of any other species, such that interspecific and intraspecific interactions are equivalent. Third, the intra + inter model, Equation (2c), hypothesizes that species are differentially influenced by interspecific and intraspecific interactions, but assumes that the per-capita interspecific effects are identical. Fourth, the intra*inter model, Equation (2d), is identical to the intra + inter model, except that it hypothesizes that interspecific effects vary depending on the abundance of the focal species (i.e. when a resident species is abundant vs. rare, it experiences differential per-capita effects of interspecific interactions). Fifth, the pairwise model, Equation (2e), hypothesizes that each species has distinct interspecific effects on other members of the community. Note that this model is similar to the classical form of Lotka–Volterra competition. Lastly, the intra*pairwise model, Equation (2f), is identical to the pairwise model, but hypothesizes that the pairwise interspecific effects vary depending on the density of the focal species.

Interspecific and intraspecific interactions can be either negative (net competition) or positive (net facilitation) in all of our models. Interactions between interspecific parameters and the abundance of the focal species (as included in the intra*inter and intra*pairwise models) represent rough approximations of potential higher-order interactions among species (Tuck et al., 2018). We structured the higher-order interactions in this way because the models can then be parameterized using only data from communities sown with one and two species, making them comparable to the other models in extrapolation tests.

The specific functional forms of these six models are adapted from a discrete-time autoregressive Gompertz model. This model is commonly used in population ecology to characterize species dynamics, and can be fit using time-series data and standard linear regression methods (Ives, Dennis, Cottingham, & Carpenter, 2003; Tredennick, Hooten, & Adler, 2017). In its basic form, this model expresses \( A_i(t) \), abundance of species \( i \) in plot \( q \) at time \( t \) as follows:

\[
A_i(t) = A_i(t-1) \times \exp \left( \beta_0 + (\beta_1 - 1) \times \log(A_i(t-1)) \right). \tag{1a}
\]

where \( \beta_0 \) is the intrinsic growth rate and \( \beta_1 \) is the dependence of this year’s abundance on last year’s abundance (i.e., density dependence). Log-transforming Equation (1a) yields a simple linear model:

\[
\log(A_i(t)) = \beta_0 + \beta_1 \times \log(A_i(t-1)) \tag{1b}
\]

to which additive effects of additional covariates (in log space), such as abundances of other species, can be incorporated and fitted using standard regression techniques.

Our six regression models characterize the log abundance of species \( i \) in plot \( q \) at time \( t \), as a function of up to three types of covariates:

- Intra only

\[
\log(A_i(t) + 1) = \beta_0 + \beta_1 + \beta_2 \log(A_i(t-1) + 1) \tag{2a}
\]

- intra = inter

\[
\log(A_i(t) + 1) = \beta_0 + \beta_1 + \log(A_i(t-1) + 1)
+ \beta_2 \log(\Sigma_k |A_{k,q}(t-1) + 1|) \tag{2b}
\]

- intra + inter

\[
\log(A_i(t) + 1) = \beta_0 + \beta_1 + \log(A_i(t-1) + 1)
+ \beta_2 \log(\Sigma_k |A_{k,q}(t-1) + 1|) \tag{2c}
\]

\[
\log(A_i(t) + 1) = \beta_0 + (\beta_1 - 1) \times \log(A_i(t-1)) \tag{2d}
\]

\[
\log(A_i(t) + 1) = \beta_0 + (\beta_1 - 1) \times \log(A_i(t-1))
+ \beta_2 \log(\Sigma_k |A_{k,q}(t-1) + 1|) \tag{2e}
\]

\[
\log(A_i(t) + 1) = \beta_0 + (\beta_1 - 1) \times \log(A_i(t-1))
+ \beta_2 \log(\Sigma_k |A_{k,q}(t-1) + 1|) \tag{2f}
\]
In all of these models, $\beta_0$ is the intercept, $\beta_1$ is a random (categorical) year effect, and $\beta_2$ describes intraspecific density dependence. The number of fitted fixed effect parameters per species for each model is therefore: 2 for intra only and intra = inter; 3 for intra + inter; 4 for intra*inter; 10 for pairwise; and 18 for intra*pairwise.

Because we considered abundances for every species in every plot in which they were sown, even if their abundance was zero, we added $1/gm^2$ to all abundances before log-transforming. We did not include spatial random effects (i.e. plot, block, or spatial coordinates) for two reasons. First, because treatments were randomly assigned to plots, there was no a priori reason to assume spatial autocorrelation that was related to community composition (when included in models, nested plot and block random effects explained, on average, about 2% of variance). Second, because our subsequent statistical tests are based on cross-validation of total model goodness-of-fit, rather than on significance tests of individual model parameters, adjustments in the degrees of freedom related to the random effects structure did not influence our results.

In Equation (2b), $\beta_2$ describes the generalized effect of all intra-specific and interspecific interactions, and $\log(A_i(t+1) = \beta_0 + \beta_1 + \beta_2 \log(A_i(t) + 1) + \Sigma_{k \neq i} \beta_{2,k} \log(A_{k,i}(t) + 1))$ is also included as an offset (i.e. without a fitted covariate) so that the function can still be interpreted as a growth rate. Consequently, for models fit using only data from monocultures, fitted values for $\beta_2$ are identical to those in Equation (2a) (after adjusting for the offset). In Equation (2c) (intra + inter) and Equation (2d) (intra*inter), $\beta_2$ is the generalized effect of interspecific interactions from all non-focal species combined, whereas $\beta_3$ describes how this effect varies as a function of the abundance of the focal species. In Equation (2e) (pairwise) and Equation (2f) (intra*pairwise), $\beta_{2,k}$ is a vector of species-specific interspecific interaction strengths (e.g. the effect of species $k$ on the focal species $i$), $A_{i,k}(t)$ is the observed abundance of species $k$ in plot $q$, and $\beta_{3,k}$ describes how $\beta_{2,k}$ changes as a function of the focal species’ abundance. Note that the fitted $\beta$ values differ among species and models.

We fitted all models using the lmer function from the lme4 package (Bates, Mächler, Bolker, & Walker, 2015) in R version 3.4.2 (R Development Core Team, 2017). We fitted separate regressions for each species and model type. As discussed above, we used two subsets of data to parameterize each model. First, all models were fit using data from all the one, two, and nine species plots (we chose these because they span the full range of diversity treatments, and because all possible species combinations of these diversity levels were represented in the plots). Second, we fit all models using data from communities sown with one and two species, except for the intra only and intra = inter models, for which we fit regressions using only data from monocultures (i.e. we chose the fewest number of diversity levels required to fit the model, based on the types of species interactions that they hypothesized). We then used these fitted models, following Equations (2a)–(2f), to predict biomass of each species in each community and year (i.e. across all sown richness levels). The full, annotated code used for our analyses is available in the file ‘Clark_et.al_JE_Dominance.R’ in the Supporting Information.

2.3 Study site

The ‘dominance experiment’ is part of the Jena Experiment (Roscher et al., 2004; Weisser et al., 2017), which was established on a former agricultural field in the floodplain of the river Saale close to the city of Jena (Germany, 50°55’N, 11°35’E, 130 m) in spring 2002. The region has a mean annual temperature of 9.9°C, and mean annual precipitation is 610 mm (1980–2010; Hoffmann, Bivour, Früh, Kößmann, & Voß, 2014). The dominance experiment is based on a pool of nine grassland species, which often reach high abundances and relative dominance in Central European mesophilic grasslands of the Arrhenatherion type (Ellenberg, 1988). These include five grass species (Alopecurus pratensis L., Arrhenatherum elatius (L.) J. Presl et C. Presl, Dactylis glomerata L., Phleum pratense L., Poa trivialis L.), two legume species (Anthriscus sylvestris (L.) Hoffm., Geranium pratense L.) and two legume species (Trifolium pratense L., T. repens L.). See Table S1 in the Supporting Information for more information about species, and Table S2 for specific data on treatments. Data are available in Weigelt et al. (2016), and in the Jena Experiment Information System (www.the-jena-experiment.de/data). In our study, we used data from the full 13-year duration of the experiment (i.e. 2002–2015).

Sown species richness levels were 1, 2, 3, 4, 6, or 9 species. Each species was equally represented at each richness level, and all possible two-species combinations were present with the same frequency at each richness level of the multi-species mixtures (i.e. 2–9 species). Each combination of species was replicated twice. The design included 2 × 9 monoculture plots, 2 × 36 two-species mixtures (all possible combinations), 2 × 24 three-species mixtures, 2 × 18 four-species mixtures, 2 × 12 six-species mixtures, and 8 nine-species mixtures, resulting in a total of 100 distinct communities (different species compositions) and 206 plots (Roscher et al., 2004).

The experiment was set up in four blocks perpendicular to the Saale River following a gradient in soil texture. Mixtures were randomly assigned to the blocks, ensuring that each block contained the same number of plots per species-richness level. Plots were established by sowing with a constant density of 1,000 germinable seeds per m² (adjusted for germination rates from standard laboratory tests),
which were equally distributed among species in the mixtures. Plots were initially sown across 3.5 × 3.5 m areas, but only maintained in the central 1 × 1 m portion of the plot from 2010 onwards because of the high expense and difficulty of maintaining the larger areas.

The sown species combinations were maintained by weeding all other species not sown into a particular plot (i.e. weeds) two to three times per year. Plots were mowed two times per year (June, September), and mown biomass was removed, as is typical for extensively used meadows in the study region. Plots were not fertilized. Annual above-ground plant biomass production was derived from the sum of two harvests per year, taken at estimated peak biomass (May and August) shortly before mowing. Plant biomass was harvested in two randomly allocated 0.5 × 0.2 m quadrats from 2003 to 2009, and one quadrat (of the same size) in the plot centre after the reduction in plot size in 2010. Harvested biomass was sorted to species after removal of detached dead plant material. Samples were weighed after drying at 70°C for 48 hr. For full species-level dynamics across treatments, see Figure S1 in the Supporting Information.

2.4 Testing model performance

To quantify the goodness-of-fit of each model, we calculated the second-order ‘coefficient of efficiency’ (Legates & McCabe, 1999; Li, 2017; Willmott et al., 1985):

\[
E_j = 1 - \sum (|O_i - P_i|/|O_i - \text{mean}(O)|),
\]

where \(j\) is the order of the coefficient, \(O_i\) is observation \(i\), \(P_i\) is prediction \(i\), and mean\((O)\) is the average taken across observations. Note that \(E_2\) is similar to the classical \(R^2\) metric, except that it measures scatter around the 1-1 line rather than a fitted regression line. Because \(E_2\) can overweight outliers in some cases (Legates & McCabe, 1999), we also repeated all analyses using \(E_1\), which corresponds to absolute error rather than squared error. These results are almost identical to those for \(E_2\), but are presented in Figures S2–S5 for reference.

After fitting models, we tested for differences in goodness-of-fit by aggregating predictions by three different sets of grouping variables—sown richness level, species identity, and year. We chose these groupings following preliminary analyses, which showed significant between-group variability in model goodness-of-fit (see Tables S3a–b for details). These groupings are also consistent with well-supported hypotheses that species interactions vary as a function of species richness (i.e. higher-order interactions), species identity (i.e. non-neutral interactions), and time (i.e. successional status or temporal feedbacks) (Adler et al., 2018; Deyele, May, Munch, & Sugihara, 2016; Mayfield & Stouffer, 2017; Tuck et al., 2018).

To quantify variation in goodness-of-fit, we applied a simple bootstrapping algorithm. For each grouping described above, we resampled from the full pool of observations and model predictions 20,000 times with replacement, and calculated goodness-of-fit for each iteration. Because of differences in sample size among the grouping levels described above, we performed bootstrapping separately for each of the groupings (i.e. we conducted separate stratified sampling within each level of sown richness, species, or year). We then used the resulting distribution of \(E_2\) and \(E_1\) values observed across iterations to calculate a mean, standard error of the mean, and \(p\)-values comparing differences in goodness-of-fit among factors.

Finally, to compare our findings to those that would arise from more classic methods of model comparison, we also calculated AIC for each of the fitted models. For these comparisons, we fitted models to the full set of data from all plots (i.e. plots sown with 1–9 species) and included a random intercept for plot nested within block. Following ‘best practices’ for AIC comparison of models that include different fixed effects, all models were fitted using maximum likelihood (i.e. rather than REML) (Bates et al., 2015). In cases where the random effects structure led to convergence issues, we removed either the plot, or the plot and block random effects (11/54 and 24/54 cases, respectively). Jointly, these approaches were intended to mirror a more typical statistical analysis of ecological data.

3 RESULTS

Considered across all diversity levels, species identities, and years, all six model forms did a similar job of estimating average within-sample abundances, albeit with a slightly increased goodness-of-fit for more complex models, especially for predictions that fell within the range of diversity conditions used for parameterization (Figure 1a). For out-of-sample extrapolations outside of the range of diversity conditions used for parameterization (i.e. those parameterized with richness levels of 1–2 species, hereafter ‘1–2 species models’), models of intermediate complexity (intra + inter and intra*inter) performed significantly better than the other models (Figure 1b, vertical axis). For out-of-sample predictions made within the range of diversity conditions used for parameterization (i.e. those parameterized with richness levels of 1, 2, and 9 species, hereafter ‘1–9 species models’), most models performed similarly, except for intra*pairwise and intra = inter, which performed significantly worse (Figure 1b, horizontal axis). See Tables S4a–k for all regression coefficients, and Tables S5a–h for \(p\)-values corresponding to differences in goodness-of-fit among models. For plots of observed versus predicted values for each model and species, see Figs. S6a–b.

Results were similar within individual diversity levels. For 1–2 species models, out-of-sample extrapolations for intra + inter and intra*inter again had the highest goodness-of-fit, followed closely by intra only and pairwise (Figure 2c–f, vertical axis). For all models, goodness-of-fit declined somewhat as a function of diversity, but the reductions were steeper for intra = inter and pairwise, and were especially steep for intra*pairwise. Thus, despite the increasing number of potential competitors, models of increasing complexity did not improve out-of-sample in more diverse communities, even for 1–9 species models.

When considering each species individually, goodness-of-fit was also consistently highest for the intra only, intra + inter, intra*inter, and pairwise models, whereas intra = inter and intra*pairwise typically
performed poorly, especially for the 1–2 species models (Figure 3). There were, however, some exceptions to this pattern. All models performed well for *A. pratensis*, and the 1–9 species models provided relatively good predictions for *A. sylvestris*, *D. glomerata*, and *G. pratense*. For both types of parameterization, *intra*+*pairwise* had especially high goodness-of-fit for *A. elatius*, and for 1–9 species models it also provided good predictions for *P. trivialis*, *T. repens*, and *T. pratense*. Goodness-of-fit for *intra* = *inter* was especially high for *P. pratense*, and for 1–2 species models of *T. repens*.

Across years, there was a strong decline in out-of-sample predictive ability around 2007 (Figure 4), corresponding to a major decline in legume abundance (especially for *T. pratense*—see Figure S1). However, predictive power recovered rapidly, and by 2009 was roughly equal to that before 2007. All 1–9 species models performed similarly, except *intra* = *inter* for which goodness-of-fit was slightly lower across all years. For 1–2 species models, all followed the same general trend, but *intra* = *inter*, *pairwise*, and *intra*+*pairwise* had particularly low goodness-of-fit, especially around 2006–2008. To account for the potential influence of these outlier years on our analyses, we repeated all of our model fitting and comparisons with years 2006–2008 omitted from the dataset, but found no major changes in our results other than a slight increase in goodness-of-fit for models of the two legumes (not shown).

For our more typical analysis based on the AIC of models fitted to the full dataset, we found that the index typically identified models of higher complexity as the ‘best fitting’ (Table 1). When compared across all species, the *intra*+*pairwise* model had the lowest AIC, which differed from the next best model (*pairwise*) by more than 150 units. Likewise, at the level of individual species, AIC never indicated that models of low complexity provided the best fit, and selected *intra* = *inter* as the best model for one species (*T. pratense*), *pairwise* for four species (*A. pratensis*, *A. elatius*, *G. pratense*, and *P. trivialis*), and *intra*+*pairwise* for four species (*A. sylvestris*, *D. glomerata*, *P. pratense*, and *T. repens*).

4 | DISCUSSION

Our primary result indicates that models of intermediate complexity—which only separate the effects of intra- and interspecific competition—can provide the best extrapolative predictions of species-level abundances in both simple and higher-diversity communities. In particular, our findings support the hypothesis that models of intermediate complexity should usually produce the best out-of-sample predictions, especially for extrapolations (Allen & Starr, 2017; Evans et al., 2013; Wenger & Olden, 2012). In accordance with other studies, we also find that classical methods of AIC comparison, based on the full dataset, selected more complex models than those selected using out-of-sample extrapolations from the models that were fitted using only data from plots sown with 1–2 species. Taken together, these results demonstrate that simple models can yield broadly generalizable predictions in complex systems, but also that commonly used methods of model selection may not always be effective at identifying these models.

In general, we can divide our models into three groups. First, those treating intraspecific and interspecific interactions as equivalent (*intra* = *inter*) almost always yielded poor out-of-sample predictions. In accordance with theory and a recent meta-analysis, this result suggests that differences between intraspecific and...
interspecific interactions are important for explaining species-level abundances (Adler et al., 2018; Broekman et al., in press). Second, models that allowed for differential effects of interspecific and intraspecific interactions (\textit{intra} + \textit{inter} and \textit{intra}*\textit{inter}) provided accurate predictions across most combinations of richness treatments, species, and years. Their strong out-of-sample goodness-of-fit suggests that these models contain sufficient complexity to explain broad trends, but not enough to cause over-fitting (Wenger & Olden, 2012). Finally, models with pairwise interactions among species (\textit{pairwise} and \textit{intra}*\textit{pairwise}) appear, at least in some cases, to be over-fit. In particular, when considered across out-of-sample extrapolations from the 1–2 species models, \textit{pairwise} and \textit{intra}*\textit{pairwise} performed significantly worse than all other tested models except \textit{intra} = \textit{inter}, and performance was especially poor for extrapolations early in the experiment, and in diverse mixtures. Jointly, these results suggest that the added complexity in the pairwise models may be more reflective of peculiarities in the training dataset than of general phenomena (Hastie et al., 2017).

Critically, our findings should not be taken to suggest that some models are more correct than others, but rather that they are more practical from the perspective of prediction given our system and available data. An important caveat for all of our models is that they do not consider any specific mechanisms of competition (e.g. allelopathy, shared resources). Thus, the poorer out-of-sample and extrapolative performance that we find for complex models may be indicative of important omitted mechanisms, rather than of the unimportance of species-specific interactions. For example, in mechanistic resource competition models, pairwise interaction strengths can change dramatically as a result of small shifts in community composition (Letten & Stouffer, 2019; Tilman, 1982). These changes would not be captured by any of the models that we test here, and it may be that simple phenomenological models are better able to average across these different conditions than are more parameter-rich phenomenological models.

4.1 | Biological interpretation of models

The baseline model, which included only intraspecific interactions (\textit{intra} only), often performed similarly to the models with both intraspecific and interspecific interactions (\textit{intra} + \textit{inter} and \textit{intra}*\textit{inter}), however, predictions were usually significantly worse for \textit{intra} only, especially for extrapolations from the 1 to 2 species models. The strong performance of the \textit{intra}-only model indicates that species

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**FIGURE 2** Goodness-of-fit for regression models at each sown richness level, calculated across all species and years. Dashed intervals show estimates for data that were used to fit the regression (i.e. ‘within-sample’), whereas solid lines show fits for data that were not used to fit the regression (i.e. ‘out-of-sample’). Intervals, axis labels, and indices are as described in the legend to Figure 1. See Table S5b in the Supporting Information for \textit{p}-values summarizing differences among model fits. Note that the axes are expanded for 6 and 9 species mixtures [Colour figure can be viewed at wileyonlinelibrary.com]
abundances were strongly regulated by density dependence and temporal autocorrelation, such that observed species abundances were largely explained by their abundance in the previous time step (Petchey et al., 2015). Note, however, that intraspecific density dependence and autocorrelation alone were not sufficient to guarantee accurate out-of-sample predictions. For example, the \textit{intra=inter} also included these components, but performed poorly, indicating that differences between intraspecific and interspecific interactions play an important role in determining abundance in our system.

For the two intermediately complex models (\textit{intra + inter} and \textit{intra*inter}), interspecific interaction coefficients were typically negative, which accords with expectations of strong net competition among communities of locally dominant species in mesic grasslands (see Tables S4a–k) (Weigelt et al., 2007). Similarly, the effects of intraspecific interactions were predicted to be about 5–10 times stronger than interspecific effects, which is consistent with theoretical and empirical results from many ecological systems (Adler et al., 2018; Barabás, Michalska-Smith, & Allesina, 2017; Tuck et al., 2018). The more complex model (\textit{intra*inter}) generally indicated significant reductions in the strength of interspecific competition with increasing focal species abundance, which suggests that competition is asymmetric and size-mediated, as might be expected, for example, light competition (Schwinning & Weiner, 1998). However, given that \textit{intra*inter} rarely performed significantly better than \textit{intra + inter}, it appears that any effects of these size-mediated biological mechanisms were either not very influential, or could be abstracted into the generic interspecific terms in the simpler model.

For the two pairwise models (\textit{pairwise} and \textit{intra*pairwise}), the decline in out-of-sample extrapolative performance early in the experiment and at higher diversity levels could indicate that species interactions vary in ways that we do not account for in Equation (2e–2f)—for example, changes in pairwise competitive interactions over the course of community assembly, or as a function of diversity (Mayfield & Stouffer, 2017; Tilman, 1982). If complex higher-order interactions among species are important, the more complex model (\textit{intra*inter}) generally indicated more significant reductions in the strength of interspecific competition with increasing focal species abundance, which suggests that competition is asymmetric and size-mediated, as might be expected, for example, light competition (Schwinning & Weiner, 1998). However, given that \textit{intra*inter} rarely performed significantly better than \textit{intra + inter}, it appears that any effects of these size-mediated biological mechanisms were either not very influential, or could be abstracted into the generic interspecific terms in the simpler model.

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processes are an important determinant of community dynamics in our system, then accurate extrapolations from complex models may require that we consider a more diverse—and potentially more mechanistic—set of models (Letten & Stouffer, 2019). However, such a result may make generalizing findings to other systems difficult, especially if they differ in the kinds of higher-order processes that are locally important. Alternatively, the relatively good performance of the 1–2 species pairwise models could suggest that the extrapolative models were simply not fitted to enough data, in which case any apparent ‘over-fitting’ might be alleviated by increasing the number of experimental replicates. It would be interesting to test this possibility with larger datasets.

Although many models performed well under some circumstances, none provided significantly better out-of-sample predictions than intra + inter and intra*inter, neither for 1–2 nor for 1–9 species models (with one exception—see Figure 3c). This result accords with a recent meta-analysis of competition experiments, which found that average intraspecific interaction strengths observed across species can be a good proxy for unknown interactions (Fort, 2018). These findings imply that, at least on average, the relative strength of intraspecific versus interspecific interactions is a stronger determinant of species abundances than are individual pairwise effects. Mechanisms that act directly on density dependence, such as species-specific pathogens, may therefore be more important for maintaining diversity in our system than are mechanisms related to pairwise competitive abilities, such as competition for a small number of limiting resources (Hubbell, 2001, 2006), which accords with existing hypotheses about coexistence in the Jena experiment (Curtois et al. 2016; Weisser et al., 2017).

TABLE 1  ΔAIC values for the six models presented in the main text. Columns show different, models, and rows show whether AIC was calculated across all species, sown diversity treatments, and years (‘Total’), or separated by species. Recall that lower AIC values indicate better performance. Bold numbers mark cases where ΔAIC < 2, which is commonly used as a rule of thumb to identify meaningful differences in model performance. Regressions are fit to all observations across all diversity treatments (i.e. 1–9 sown species richness treatments). See main text for details about model fitting and random effects structure.

<table>
<thead>
<tr>
<th>Plant Name</th>
<th>intra only</th>
<th>intra = inter</th>
<th>intra + inter</th>
<th>intra*inter</th>
<th>pairwise</th>
<th>intra*pairwise</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>748.46</td>
<td>2,354.15</td>
<td>484.01</td>
<td>406.83</td>
<td>154.9</td>
<td>0</td>
</tr>
<tr>
<td>Alopecurus pratensis</td>
<td>58.58</td>
<td>96.22</td>
<td>13.15</td>
<td>3.01</td>
<td>0</td>
<td>7.75</td>
</tr>
<tr>
<td>Anthriscus sylvestris</td>
<td>66.13</td>
<td>204.25</td>
<td>52.18</td>
<td>34.75</td>
<td>2.81</td>
<td>0</td>
</tr>
<tr>
<td>Arrhenatherum elatius</td>
<td>47.49</td>
<td>344.6</td>
<td>23.34</td>
<td>25.09</td>
<td>0</td>
<td>1.5</td>
</tr>
<tr>
<td>Dactyliis glomerata</td>
<td>102.65</td>
<td>153.5</td>
<td>63.55</td>
<td>48.13</td>
<td>1.01</td>
<td>0</td>
</tr>
<tr>
<td>Geranium pratense</td>
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<td>209.77</td>
<td>50.56</td>
<td>39.93</td>
<td>0</td>
<td>7.39</td>
</tr>
<tr>
<td>Phleum pratense</td>
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<td>367.42</td>
<td>143.05</td>
<td>120.95</td>
<td>107.22</td>
<td>0</td>
</tr>
<tr>
<td>Poa trivialis</td>
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<td>351.98</td>
<td>66.27</td>
<td>75.66</td>
<td>0</td>
<td>5.97</td>
</tr>
<tr>
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<td>270.47</td>
<td>0</td>
<td>1.89</td>
<td>12.76</td>
<td>17.4</td>
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<tr>
<td>T. repens</td>
<td>161.46</td>
<td>395.96</td>
<td>111.93</td>
<td>97.45</td>
<td>71.12</td>
<td>0</td>
</tr>
</tbody>
</table>

FIGURE 4 Goodness-of-fit of fitted regression models for each year, calculated across all species and sown richness levels. Results are shown only for out-of-sample goodness-of-fit. Intervals show mean ± one standard error of the mean. Axis labels and indices are as described in the legend to Figure 1. Note that the intra + inter and intra*inter trajectories are overlapping. See Table S5d in the Supporting Information for p-values summarising differences among model fits (Colour figure can be viewed at wileyonlinelibrary.com)
Species-level results

Although average predictive accuracy varied greatly across species, the relative performance of the six models usually matched the general order observed for the total dataset. One exception to this pattern was Arrhenatherum elatius, for which goodness-of-fit was low for all models, but with significantly higher goodness-of-fit for the pairwise models. Because its abundance was consistently high, and varied little across plots, treatments, or years, the ‘null expectation’ that we used to predict goodness-of-fit in Equation (3)—that is, mean observed biomass—performed particularly well, which left relatively little room for improvement. A potential explanation for the consistently high abundance of A. elatius is that it is an especially effective competitor for light and nitrogen: it is the tallest species in the dominance experiment, invests a larger fraction of biomass in supporting tissues (Lorentzen et al., 2008; Roscher et al., 2008), and produces the most biomass per unit nitrogen (Roscher, Thein, Schmid, & Scherer-Lorenzen, 2008). Thus, A. elatius may be buffered from the effects of interspecific competition, making models of generic interspecific interactions less effective.

Three other species also have tall stature and produce high biomass per unit nitrogen—Alopecurus pratensis, Dactylis glomerata, and Phleum pratense (Lorentzen et al., 2008; Roscher et al., 2008). Goodness-of-fit for these species was high for all models, except for some poor extrapolations from the 1–2 species intra*pairwise model. As with A. elatius, these species generally had high abundance, and dynamics were consistent across plots and diversity treatments (Figure S1) potentially because their traits shielded them from competitive interactions with most other species. However, unlike A. elatius, their abundances varied over time, such that mean observed biomass alone was not a good predictor. Thus, high goodness-of-fit may indicate that their dynamics were sufficiently complex to differ from the null expectation, but not so complex as to confound our models. That said, high abundance might actually contribute to the strong predictive power of these models by reducing the impact of observation error on predictive outcomes, or increasing temporal autocorrelation (Hastie et al., 2017).

Goodness-of-fit was also relatively high across models for two species that did not have traits associated with strong light or nitrogen competitive ability—Anthiricus sylvestris and Geranium pratense (Lorentzen et al., 2008; Roscher et al., 2008). Potentially because of poorer competitive abilities, both species were slow to establish, and abundances varied greatly among plots and treatments (Figure S1) (Lorentzen et al., 2008). Although strong effects of interspecific competition may explain the relatively poor performance of intra = inter, the high goodness-of-fit for intra only suggests that within-plot autocorrelation alone was sufficient for predicting species dynamics.

Lastly, for three species—Poa trivialis, Trifolium pratense, and T. repens—goodness-of-fit was relatively low for all models. A partial explanation is that over a short period of about 2006–2008 there were especially large declines in abundance for T. pratense and T. repens, and increases in abundance for P. trivialis (see Figure S1). When these years were removed from the dataset, goodness-of-fit increased for the two legumes, but not for P. trivialis. Curiously, this time period did not correspond to any major weather events or changes in experiment management. Instead, it is possible that the initially sown cohort of shorter-lived perennial species (e.g. T. pratense and T. repens) began to die back around this time and failed to establish successful reproductive populations, leading to rapid changes in species relative abundance (Roeder, Schweingruber, Fischer, & Roscher, 2017; Roscher et al., 2011).

Methodological implications of results

There have been many rallying calls for better, and more general, predictive models in ecology (Coelho et al., 2018; Dietze et al., 2018; Evans et al., 2013; Houlahan, McKinney, Anderson, & McGill, 2017; Lawton, 1999; Wenger & Olden, 2012). Not all of these favours reductions in model complexity. In particular, the advent of individual-based models, which can be formulated around rules and behaviours, has been suggested as a potential way to simultaneously satisfy the needs for nuance and generality (Allen & Starr, 2017; Evans et al., 2013; Grimm et al., 2016; Judson, 1994). Nevertheless, to our knowledge, the majority of studies that have empirically tested model performance across a gradient of complexity accord with our findings—that is, that intermittently complex models provide the best extrapolations—including models of bacterial growth (Buchanan, Whiting, & Damert, 1997), fish populations (Wenger & Olden, 2012), and plant communities (Petitpierre, Broennimann, Kueffer, Daehler, BIC, etc.; Burnham et al., 2011; Aho et al., 2014; Brewer et al., 2016; Houlahan et al., 2017; Coelho et al., 2018). While there is nothing inherently wrong with this approach, it is important to remember that these tests are primarily designed to identify models that perform well within the general range of conditions used for parameterization, rather than to estimate how models are likely to perform under novel conditions (Brewer et al., 2016; Stone, 1977). Nevertheless, assuming that such models will also yield good extrapolations can be problematic. Because most ecological systems are inherently complex and interconnected, it is likely that given enough data, significant interactions can be detected among any ecological variables, at least via indirect routes (Levin, 1998; Sugihara et al., 2012). But, much of this complexity is likely to be non-transferable across systems (Lawton, 1999). Hence, if our goal is to obtain generality, we should not necessarily emphasize the particular, but rather should focus on identifying models that perform well across a wide range of contexts (Wenger & Olden, 2012). For example, simpler models associated with AIC values that are > 2 units higher than the best models are unlikely to be explored or considered further, yet they may well provide similar...
within-sample goodness of fit, and better out-of-sample predictions and extrapolations.

Similarly, it is rare to see a thorough analysis of the real impact of higher-order parameters. Figure 1a illustrates this point: including additional terms does indeed improve the within-sample estimates from the 1–9 species models, but not by much. A more detailed analysis of how well simpler models perform—and where, and by how much, they actually get things wrong—is therefore probably more informative than simply accepting some subset of models based on their goodness-of-fit within a single set of conditions (Brewer et al., 2016; Mayfield & Stouffer, 2017; Wenger & Olden, 2012). We do not mean to imply that complexity will never be needed to understand particular aspects of ecological systems. Rather, it may be wise to address complexity judiciously, only after we have convinced ourselves that simpler models fail to provide sufficient precision or generality.

5 | CONCLUSIONS
Because of their complexity, ecological systems will always pose a particular challenge for modellers. Though complexity and nuance may well be important for explaining the abundance and distribution of species in many sites and systems, our results demonstrate that in some cases, simpler models can provide more general predictions. More broadly, our analysis serves as a reminder that the best way to test the generalizability of a fitted model is to use it to make predictions in a new context—and that commonly used model selection tools are not designed to test this property.

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AUTHORS’ CONTRIBUTIONS

DATA AVAILABILITY STATEMENT
All data are publicly available through the PANGEA database (https://doi.org/10.1594/PANGAEA.866358), and can also be accessed through the Jenaexperiment Information System (www.the-jena-experiment.de/data).

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REFERENCES


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