Strong self-regulation and widespread facilitative interactions in phytoplankton communities

Coralie Picoche^{1,2}, Frédéric Barraquand^{1,2*} January 27, 2020

- 1 University of Bordeaux, Integrative and Theoretical Ecology, LabEx COTE, Bât. B2 Allée Geoffroy St-Hilaire, 33615 Pessac, France; coralie.picoche@u-bordeaux.fr
- **2** CNRS, Institute of Mathematics of Bordeaux, 351 Cours de la Libération, 33405 Talence, France; frederic.barraquand@u-bordeaux.fr
 - $oldsymbol{*}$ corresponding author: Email: frederic.barraquand@u-bordeaux.fr

Abstract

- 1. The persistence of phytoplanktonic diversity in spite of competition for basic resources has long been a source of wonder and inspiration to ecologists. To sort out, among the many coexistence mechanisms suggested by theory and experiments, which ones actually maintain diversity in natural ecosystems, long-term field studies are paramount.
- 2. We analysed a large dataset of phytoplankton abundance time series using dynamic, multivariate autoregressive models. Phytoplankton was counted and identified down to the genus level, every two weeks over twenty years, at ten sites along the French coastline. Multivariate autoregressive models allowed to estimate biotic interaction networks, while also accounting for abiotic variables that may drive part of the phytoplankton fluctuations. We then analysed the ratio of intra- to inter-taxa interactions (measuring self-regulation, itself a measure of niche differentiation), the frequency of negative vs positive interactions, and how stability metrics (both at the network and genus level) relate to network complexity and genus self-regulation or abundance.
- 3. We showed that a strong self-regulation, with competition strength within a taxon (genus) an order of magnitude higher than between taxa, was present in all phytoplanktonic interaction networks. This much stronger intragenus competition suggests that niche differentiation rather than neutrality is commonplace in phytoplankton. Furthermore, interaction networks were dominated by positive net effects between phytoplanktonic taxa (on average, more than 50% of interactions were positive). While network stability (sensu resilience) was unrelated to complexity measures, we unveiled links between self-regulation, intergenus interaction strengths and abundance. The less common taxa tend to be more strongly self-regulated and can therefore maintain in spite of competition with more abundant ones.
- 4. Synthesis: We demonstrate that strong niche differentiation, widespread facilitation between phytoplanktonic taxa and stabilizing covariances between interaction strengths should be common features of coexisting phytoplankton communities in the field. These are structural

properties that we can expect to emerge from plausible mechanistic models of phytoplankton communities. We discuss mechanisms, such as predation or restricted microscale movement, that are consistent with these findings, which paves the way for further research.

Keywords: phytoplankton; coexistence; facilitation; mutualism; niche theory; time series; networks

1 Introduction

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How species or close genera can coexist together in spite of competition is one of the main puzzles of
   community ecology, especially for primary producers that seemingly share the same basic resources
   (Hutchinson, 1961). Many theoretical studies of competition models have shown that competitive
   exclusion is likely in those circumstances, unless mechanisms involving spatial or temporal variation
   are at play (Armstrong & McGehee, 1976, 1980; Chesson & Huntly, 1997; Huisman & Weissing, 2001;
   Li & Chesson, 2016; Chesson, 2018). Neutral theory, that assumes a non-equilibrium coexistence
   maintained by dispersal and equal competitive abilities for all species (Hubbell 2001, though there
   are exceptions, see Volkov et al. 2003, 2007), has been proposed as a solution to explain highly
   diverse communities (Hubbell, 2001; Rosindell et al., 2011).
       However, the evidence gathered from terrestrial plant communities starts to suggest that, in fact,
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   niche rather than neutral processes may be paramount to explain coexistence, with intraspecific
   competition dwarfing interspecific competition in most cases (Adler et al., 2010, 2018b). Whether
   these conclusions drawn mostly from studies of terrestrial plants apply to other ecosystems and taxa
   is currently little known (but see Mutshinda et al. 2009).
       Moreover, competition may not be the rule: the meta-analysis by Adler et al. (2018b) reported a
   large number of facilitative interactions (30%) and several reviews (Brooker et al., 2008; McIntire
   & Fajardo, 2014; Kinlock, 2019) have highlighted that facilitation may be much more widespread
   than ecologists usually tend to think. Although some theoretical studies suggest that facilitative
   interactions can be destabilizing (sensu resilience) and therefore undermine coexistence in Lotka-
   Volterra models (Coyte et al., 2015), multiple other modelling (Gross, 2008) and empirical (Brooker
   et al., 2008; Cavieres & Badano, 2009) studies have suggested that facilitative interactions can
   to a large degree benefit coexistence, especially when multiple interaction types are considered
   simultaneously (Mougi & Kondoh, 2012; García-Callejas et al., 2018).
       Here, we analyse a spatially replicated, long-term community-level dataset, consisting of ten
   multivariate time series of phytoplankton abundance along the French coastline. We do so using
   multivariate autoregressive (MAR) models, which allows to estimate interactions between genera.
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Although many ecological studies focus on interactions between species, competition has been shown experimentally to occur between different genera of phytoplankton (Titman, 1976; Descamps-Julien & Gonzalez, 2005). The genus level is also a rather fine taxonomic scale for phytoplankton interaction studies, as most studies are restricted to interactions between different classes or even phyla (Ives et al., 2003; Hampton et al., 2008; Griffiths et al., 2015). Studying interactions between different genera of phytoplankton therefore both makes empirical sense in light of competition experiments and allows to estimate better-resolved networks. We focus here on genera that belong mostly to diatoms and dinoflagellates. To put our results into a more general context, we then compare our interaction strength estimates to previously published interaction networks produced under the same statistical framework, both in plankton and other empirical systems.

38 Material and methods

39 Sampling methods

All phytoplankton samples were collected by Ifremer coastal laboratories as part of the National
Phytoplankton and Phycotoxin Monitoring Network (REPHY, 2017). Since 1987, this monitoring
program has required the sampling of 26 sites along the French coastline every 2 weeks within 2
hours of high tide to document both biotic (phytoplankton counts) and abiotic (water temperature,
salinity) variables. We focused on sites which had the longest time series. We also excluded time
series which had missing data for over 6 months or an average delay between sampling dates above
20 days. This reduced the number of study sites to 10 sites nested within 4 regions (Brittany, Oléron,
Arcachon and the Mediterranean Sea; Fig. S1 and Table S1).

Abiotic variables (temperature, salinity) were measured directly from the boat during the sampling
process while water samples for biotic analyses were fixed with a Lugol's solution and examined later.
Phytoplankton cells above 20 μm were identified at the lowest possible taxonomic level and counted
with the Utermöhl method using an optical microscope (Utermöhl, 1958). Throughout the years
and sites, more than 600 taxa were identified at different taxonomic levels. We aggregated them at
the genus (or group of genera when not possible) level based on previous work (Table S2; Hernández

Fariñas et al. 2015; Barraquand et al. 2018), except for cryptophytes and euglenophytes in Arcachon, which could not be identified below the family level. Although the taxonomic resolution used here may seem coarse in comparison to land plants, it is in fact more refined than 86% of the MAR(1) studies of phytoplankton listed in Table S4.

For each region, the MAR(1) analysis focused on the most abundant and most frequently observed genera to avoid most of the gaps in the time series. When gaps did not exceed a month, missing values were linearly interpolated; remaining missing values were replaced by a random number between 0 and half of the lowest observed abundance (Hampton et al., 2006). Time series are plotted in Fig. S2. We tested extensively this and other methods to deal with missing data in a previous publication on a subset of this dataset (Barraquand et al., 2018). All time series were scaled and centered before MAR analyses.

$_{65}$ MAR(1) model

Multivariate autoregressive (MAR) models are used to determine the interspecific interactions and abiotic effects shaping a community's dynamics (Ives *et al.*, 2003). MAR(1) models are based on a stochastic, discrete-time Gompertz equation which relates the log-abundance of each of the S taxa at time t+1 to log-abundances of the whole community at time t, with possible interactions between taxa, and effects of V abiotic variables at time t+1. These assumptions are encapsulated in eq. 1:

$$\mathbf{n}_{t+1} = \mathbf{B}\mathbf{n}_t + \mathbf{C}\mathbf{u}_{t+1} + \mathbf{e}_t, \mathbf{e}_t \sim \mathcal{N}_{\mathcal{S}}(0, \mathbf{Q})$$
(1)

where \mathbf{n}_t is the $1 \times S$ vector of log-abundance of phytoplankton taxa, \mathbf{B} is the $S \times S$ community (interaction) matrix, \mathbf{C} is the $S \times V$ environment matrix describing the effects of V variables (stacked in vector \mathbf{u}_{t+1}) on growth rates, with V=2 in our case (temperature and salinity). The noise \mathbf{e}_t is a $1 \times S$ noise vector which covers both process and observation error, following a multivariate normal distribution with a variance-covariance matrix \mathbf{Q} . \mathbf{Q} is diagonal and we have previously showed that this parsimonious choice did not affect qualitatively the results (Barraquand et al., 2018). We used the MARSS package (Holmes et al., 2014) v3.9, in R v3.3.2 (Venables & Smith,

⁷⁸ 2013), to estimate parameters with a maximum likelihood procedure.

Our previous analysis of the Arcachon region, for which more covariables were available (Bar-79 raquand et al., 2018), revealed that hydrodynamics and hydrology had more influence on phyto-80 plankton dynamics than nutrients on the two-week timescale. Because temperature and salinity sum 81 up seasonal changes in light as well as hydrology (salinity is inversely related to freshwater inflow), these represent the two key drivers needed to account for abiotic influences (Scheef et al., 2013). They are therefore used to summarize the abiotic environment in the remainder of the article. The analysis of real data in Barraquand et al. (2018) was complemented by that of simulated data mimicking the study design, which confirmed the ability of MAR(1) models to infer biotic interactions and abiotic forcings. Fitting a more sophisticated model (threshold autoregressive model) did not reveal extra non-linearities or a storage effect in the Arcachon subset of the data (Barraquand et al., 2018). Other aspects of the MAR(1) modelling are likewise quite robust: using two abiotic variables (temperature and salinity) in this study rather than the full set used in Barraquand et al. (2018) led to almost identical covariate effects and interaction estimates for the Arcachon study sites. Even if some departures from the true data-generating model may not always be detectable through MAR(1) diagnostics (e.g., residuals), the analysis of nonlinear simulations has showed that MAR(1) models are in general robust to nonlinearities if the inference focuses on interaction sign and order of magnitude of model coefficients (Certain et al., 2018), which is how these models are used here. For ease of interpretation of MAR(1) interaction coefficients, we also prove the correspondence between the magnitude of intra/inter interaction strength in a MAR(1) model and a multispecies Beverton-Holt model, i.e., a discrete-time Lotka-Volterra model (Cushing et al., 2004), in the Supporting Information. In this study, the number of phytoplankton taxa (S) and the community composition vary 100 slightly between regions but sites share on average 67% of their taxa. In order to have comparable 101 models, we also keep the same 2 covariates, i.e., water temperature and salinity, that were measured 102 at all study sites. Therefore, the dimension of the dynamical system depends on the (square of

the) number of phytoplankton taxa we study, which ranges between 7 (Mediterranean Sea) and

14 (Brittany). The smallest system still requires 63 parameters to be estimated (49 for the 7×7

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interaction matrices and 14 for the 7×2 environment matrices) if we consider all possible interactions between taxa. To reduce this dimensionality and remove unnecessary parameters, we built different 107 'interaction scenarios' based on known phylogenetic information (as suggested in Violle et al., 2011; 108 Narwani et al., 2017). The null interaction scenario assumed no interaction between genera (diagonal 109 interaction matrix) and was compared to four other interaction scenarios. The first interaction 110 scenario assumed that interactions could only occur between phylogenetically close organisms, i.e., 111 within a class (groups were then diatoms, dinoflagellates, and other phytoplanktonic organisms) 112 while the second interaction scenario further differentiated pennate and centric diatoms. The third interaction scenario considered the reverse hypothesis, that only unrelated organisms could interact 114 (i.e., a diatom could only interact with a dinoflagellate or a cryptophyte, but not with another diatom), and the last interaction scenario did not constrain the interactions at all (full interaction 116 matrix). We selected the best scenario by comparing BIC (Fig. S3), which proved to be satisfactory 117 in our previous analyses of both real data and similar simulated datasets (Barraquand et al., 2018, 118 Appendix 2). The second interaction scenario, hereafter called the pennate-centric scenario, had the 119 lowest BIC for all sites (Fig. S3). This parsimonious scenario was therefore chosen as the basis for 120 further investigations of network structure. 121

Analysis of interaction strengths

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The interaction matrix obtained from MAR(1) analyses can be used to determine the stability 123 of a discrete-time dynamical system (Ives et al., 1999, 2003). To investigate stability-complexity relationships, we compared the maximum modulus of the eigenvalues of the pennate/centric matrices 125 for each site to network descriptors. The maximum modulus is analogous to the real part of the leading eigenvalue for continuous time models, and measures resilience while still accounting for 127 some variability properties (Ives et al., 1999). However, because most theory on stability-complexity 128 has been developed in continuous time (e.g. Allesina & Tang, 2015), we numerically checked that the 129 maximum modulus of the eigenvalues in a discrete-time interaction matrix and its continuous-time 130 model counterpart yield similar information in the Supporting Information. We then compared 131 this resilience measure to complexity metrics, such as the interaction strength distribution (sign, mean and variance) and weighted connectance (Bersier *et al.*, 2002). Weighted connectance is a
measure of the proportion of realized links compared to all possible links, taking into account the
shape of the flux distribution. This metric is adapted to weighted interaction matrices but cannot
accommodate for both positive and negative coefficients: we therefore chose to focus on interaction
strength only (absolute values of the coefficients), irrespective of interaction sign. In contrast, mean
and variance of the off-diagonal coefficients, which can affect the stability of a community (Allesina
Tang, 2015), are computed on raw values of the coefficients. Interaction coefficient variance is
multiplied by the number of taxa, according to theory (Allesina & Tang, 2015).

In addition to these network-level metrics, we also computed the average vulnerability (average effect of other taxa on a focal taxon, eq. S5) and impact (average effect of a focal taxon on other taxa, eq. S6) on both raw and absolute values of the coefficients. Vulnerability and impact can be related to in-strength and out-strength in the meta-analysis of Kinlock (2019). We then compared these to the regulation a focal species exerted on itself. Raw values indicate the average effect that can be expected on the growth rate of a taxon from the rest of the community (i.e., is the effect of others mostly positive or negative?), while absolute effects characterise the strength of all types of interactions on a taxon (i.e., is a taxon strongly affected by the others?).

Finally, we compared the observed ratio between mean self-regulation (intrataxon interaction strength) and mean intertaxa interaction strength to other published studies based on a MAR(1) model. A list of references is given in Table S4. Authors usually reported only coefficients that were significant at the 95% significance threshold, thus ignoring potentially many weak effects, which we had to set to 0. There are therefore two ways of computing the mean intertaxa interactions, i.e., taking the mean value of all coefficients outside of the matrix diagonal, including zeroes (which decreases the estimated mean intertaxa interaction strength, Fig. 4), or taking the mean value of statistically significant intertaxa coefficients only (which increases the estimated mean intertaxa interaction strength, Fig. S9). We considered both.

Results

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Interaction estimates

et al., 2013) - i.e., Jacobian community matrices on the logarithmic abundance scale (Ives et al., 161 2003). Best-fitting models corresponded to a phylogenetically-structured interaction scenario, where 162 interactions only occurred between closely related genera (Fig. S3). This led to sparse, modular 163 matrices that have two main features. First, we observed a strong self-regulation for all sites (Fig. 1, diagonal elements of all matrices), a feature that we had previously highlighted in a more detailed 165 analysis on one of the considered study regions (Barraquand et al., 2018). The ratio of mean 166 intragenus to intergenus interaction coefficients varied between 6 and 10, not counting coefficients 167 set to 0 in the estimation process. When we included the zeroes in the interaction matrix in 168 the computation of the intra/inter mean interaction strength, the ratio rose to 21-43. Therefore, 169 intragenus interactions were on average one order of magnitude stronger than intergenus interactions. 170 Second, although the percentage of facilitative interactions varied among sites (between 40% 171 and 71% of interactions in the selected models), facilitation remained predominant in 9 sites out 172 of 10 (only Lazaret, in the Mediterranean Sea, has 60% negative interactions). Our observational 173 setup being nested, with sites within regions, we could examine whether locally positive interactions 174 remain positive in a regional context: the percentage of consistently positive interactions at the 175 regional level varied between 30% and 53%, higher than the percentage of similarly defined negative 176 interactions (between 15% and 40%), except for sites in the Mediterranean Sea. 177 We found that the percentage of true mutualism (+/+) was substantial: averaged over all sites, 178 32% of all interactions were (+/+) while only 12% of them were (-/-), see also Fig. S5. The sign 179 correspondence was not always maintained between regions: the only interaction that was non-zero 180 in the 10 sites (CHA/SKE) was mutualistic in Men er Roue only (Brittany) and mixed (+/-) in 181 all other sites. Within the same region, however, interactions measured in different sites tended to 182 keep the same sign. In the 3 sites of Oléron, for instance, there were 4 interactions which remained 183 positive for both taxa involved (CHA/GUI, DIT/GUI, LEP/THP, SKE/THP), 3 of them being also

Using MAR(1) autoregressive models, we produced interaction matrices (Ives et al., 2003; Hampton

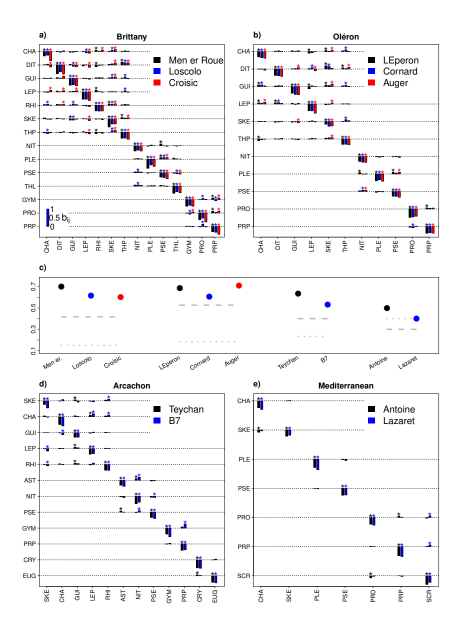


Figure 1: Interaction matrices estimated in 10 sites along the French coastline. The most parsimonious interaction scenario is presented. Taxon j (in columns) has an effect on taxon i's growth rate (in rows) proportional to the bar height, which corresponds to the $\mathbf{B} - \mathbf{I}$ matrix (community composition in Table S2). The scale for the coefficient values is given at the bottom left of panel a). 95% significance of coefficients is marked by asterisks (*). The fraction of positive interactions in each matrix is given by points in c) while the dashed (resp., dotted) line represents the ratio of interactions remaining positive (resp., negative) for all sites of a given region.

mutualistic in some of the Brittany sites. This contradicts previous observations that mutualistic interactions tend to be more context-dependent than competitive interactions (Chamberlain et al., 186 2014). 187

Interaction network analysis 188

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The stability (sensu resilience, Ives & Carpenter 2007) of all interaction matrices was not strongly 189 affected by the percentage of positive interactions or the mean and variance of the intergenus 190 interactions (Fig. 2). There was a slight increase in stability with weighted connectance, with a drop 191 in eigenvalue modulus for weighted connectances between 0.09 and 0.1. The maximum modulus of 192 the interaction matrix eigenvalues remained between 0.65 and 0.80. 193

Given that a direct complexity-stability (sensu resilience) link was not obvious, we investigated whether the matrix coefficients had some particular structure that could help theoretical ecology to make better null models of joint community dynamics and interactions (James et al., 2015). We defined two scores, vulnerability (summed effect of others on the focal taxon growth rate, eq. S5) and impact (summed effect of the focal taxon onto other taxa's growth rates, eq. S6). Relations between inter- and intra-genus interactions emerged (Fig. 3): genera that were more self-regulating 199 also had also a higher vulnerability score and a lower impact score. Those two influences are likely to trade-off: a high degree of self-regulation somehow buffers the effect of outside influences on population dynamics. Taxa that were less self-regulating were also more likely to have a stronger effect onto other taxa. As these genera tended to be more abundant (Fig. S7), this could be mediated by the average density of a genus. It is important to note, however, that these trends are weak and there is therefore a considerable amount of randomness dominating the interaction matrix: many scenarios of self-regulation vs limitation by others are therefore possible.

Aside from the trade-offs of Fig. 3, we found no remarkable patterns of covariation between 207 matrix elements (Fig. S5) other than a mean-variance scaling of interaction coefficients (Fig. S6).

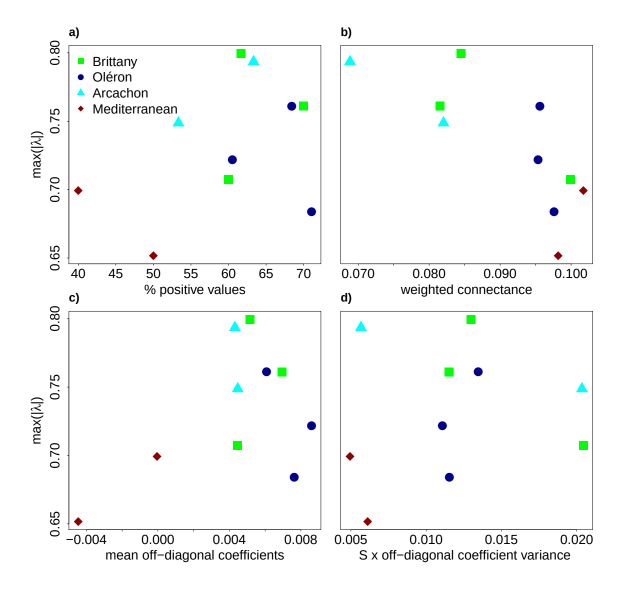


Figure 2: Relation between stability and complexity of the interaction networks. The maximum modulus of the eigenvalues of the interaction matrix **B** indicates stability *sensu* resilience. Off-diagonal coefficient variance is multiplied by the dimension of the network, that is the number of species in the region. Each color or shape corresponds to a given region. The formula for weighted connectance is given in the Supporting Information.

Literature comparison

Finally, we sought to put these results in a broader context by compiling the intra vs inter group estimates of previous MAR(1) studies of long-term observational count data (listed in Table S4).

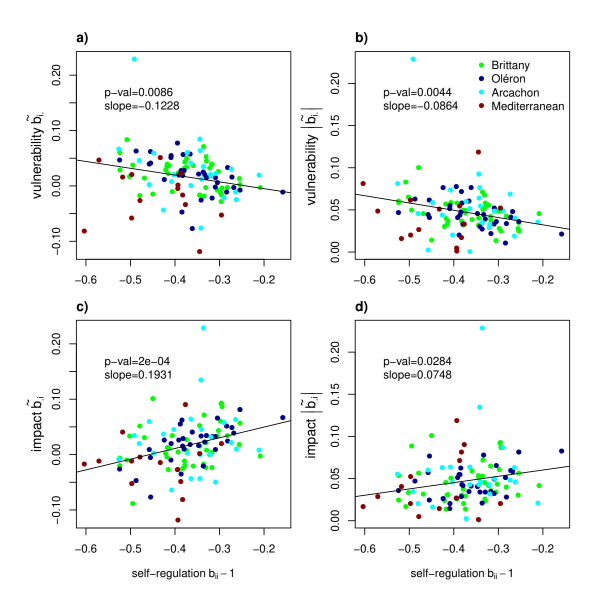


Figure 3: Relation between vulnerability/impact and self-regulation. Average vulnerability (effects of others on the focal taxon growth rate, a-b) and impact (effects of the focal taxon on others' growth rates, c-d), as well as self-regulation, are computed for untransformed (a-c) or absolute (b-d) values of the coefficients of the interaction matrix $(\mathbf{B} - \mathbf{I})$ for the 10 study sites. Each color corresponds to a given region (Fig S1). Linear regressions are shown as black lines.

We found that the order of magnitude of intra/inter interaction strengths considered here is not particularly above those found for most planktonic systems to which MAR(1) models have been

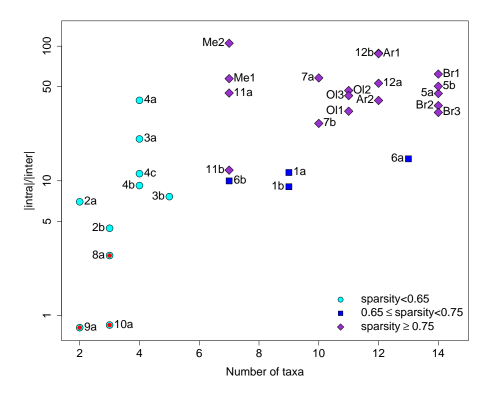


Figure 4: Ratio of intra- to inter-group interaction strength in Multivariate AutoRegressive (MAR) models. The reference for each study is given in Table S4. Codes beginning with letters correspond to the present study (Ar: Arcachon; Ol: Oléron; Br: Brittany; Me: Mediterranean Sea). The symbol color and shape correspond to the sparsity of the interaction matrix (e.g., the proportion of null interactions in the matrix). Red dots correspond to terrestrial and/or low dimension predator-prey systems, giving a lower bound for the intra/inter ratio. Intergroup interactions were set to 0 when they were not specified in the articles (in most cases, authors removed non-significant interactions at the 95% threshold; Fig. S9 is the same figure taking into account only significant interactions)

fitted, considering that our systems are relatively high-dimensional and that the higher the number of taxa, the larger the intraspecific regulation (Barabás et al., 2017). We included in Fig. 4 not only 215 plankton studies but also a couple of vertebrate or insect studies on less diverse communities, where 216 interactions are stronger, in order to provide lower bounds for the intra/inter ratio. The conclusion from this comparison seems to be that, unlike small communities that can be tight-knit, any diverse 218 field system of competitors and facilitators has evolved large niche differences making intragroup 219 competition much larger in magnitude than intergroup interactions.

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Discussion

22 Strong self-regulation and facilitation

We found very large niche differences between genera, translating into much higher intragenus than intergenus effects on growth rates (i.e., strong self-regulation), together with a high degree of facilitative net interactions.

The intra/intertaxa interaction strength ratio (Levine & HilleRisLambers, 2009) that we found, 226 from 6-10 to above 20, depending on whether one includes interactions set to zero in the estimation 227 process, could appear very high in light of previous intra/interspecific competition strength estimates 228 of 4 to 5 by Adler et al. (2018b). Additional estimates using the unconstrained interaction matrix 229 yielded ratios between 8 and 11 depending on the site (Table S3 and Fig. S8 in the Supporting 230 Information), but weak interspecific effects are likely to be inflated in the full model. Therefore, a intra/inter ratio of 10 seems like a conservative estimate, twice that of Adler et al. (2018b). Even 232 though their model is a different one, i.e., Lotka-Volterra competition, we prove in the Supporting Information that the intra/inter ratio should remain commensurate in a MAR(1) model for diffuse 234 competition. Unfortunately, the skew in species abundances is such that the map between Lotka-235 Volterra and MAR(1) intra/inter ratios is limited for individual coefficients, but averaged over 236 species pairs as we compute it, this ratio should be similar for both frameworks. The difference in 237 the intra/inter ratio that we found should therefore lie elsewhere, which requires some explanation. 238 One could argue that such high intra/inter ratio arises because we consider the genus as our baseline 239 taxonomic unit, rather than the species. It is logical that niche differentiation increases as one gets up the phylogenetic tree, and that getting down to the species level could slightly decrease 241 that ratio (but see Narwani et al., 2017, in which phylogenetic closeness decreases competition strength). However, taxonomic resolution is unlikely to be the sole explanation for the high intra/inter 243 ratio interaction strength found here, for two reasons. First, phytoplankton species belonging to 244 different genera are often found to compete in experiments (Titman, 1976; Tilman et al., 1982; Descamps-Julien & Gonzalez, 2005). In the field-based dataset studied here, the same genera that 246 are considered in experiments are found not to compete (or only weakly), hence there must be some

niche differentiation occurring in the field but not in the lab. Second, the only other study that managed to provide MAR(1) estimates down to the species level for phytoplankton, that of Huber & Gaedke (2006), provides an intra/interspecific strength ratio similar to ours (point 7a in Fig. 4).

Strong self-regulation seems therefore a genuine feature of field phytoplanktonic communities.

Another main finding of our study is the large frequency of positive interactions, with 30% truly 252 mutualistic (+/+) interactions and between 40 and 70% facilitative effects. Although a seasonal 253 environment can generate some positive covariation between taxa, those effects have already been 254 filtered out by the inclusion of our 2 abiotic covariates (Fig. S4). The facilitative effects shown 255 here are therefore residual effects, once abiotic trends are accounted for. Between 40 and 70% 256 facilitation can be compared to the meta-analysis by Adler et al. (2018b) who also found facilitative 257 interactions, but less than here ($\approx 30\%$). However, Adler et al. (2018b)'s review contains many 258 experiments while the plant literature is replete with field examples of facilitation (Brooker et al., 259 2008; McIntire & Fajardo, 2014), so that plant facilitation could be higher in the field. At the 260 moment, it is therefore unknown how the predominance of facilitative interactions that we found 261 in phytoplankton compares to facilitation in terrestrial plants. We note that several authors using 262 MAR(1) models previously forbade positive interactions within the same trophic level, so that the 263 fraction of facilitative interactions in plankton cannot be computed from literature-derived MAR(1) estimates. 265

The large niche differences and facilitative interactions that arise when considering a single 266 trophic level are an emergent property, resulting from hidden effects of resource or predator 267 partitioning/sharing (Chesson, 2018). In our previous publication investigating in detail the Arcachon 268 study sites (Barraquand et al., 2018), we have argued that for phytoplankton, the strong intragroup 260 density-dependence could arise from effects of natural enemies (Haydon, 1994). Natural enemies 270 could also very well create apparent mutualism between prey species (Abrams et al., 1998; de Ruiter 271 & Gaedke, 2017). We believe this to be likely for the present study, given that the study regions 272 (Arcachon, Oléron, Brittany, Mediterranean) have similar predators (zooplankton, e.g., Jamet et al., 2001; Modéran et al., 2010; Tortajada et al., 2012) and parasites (viruses, e.g., Ory et al., 2010; 274 fungi). Though natural enemies are good candidates to explain the observed niche differences and

emerging facilitation, one must bear in mind that other known drivers of phytoplankton dynamics such as allelopathy (Felpeto et al., 2018), auxotrophy (Tang et al., 2010) or hydrodynamics (Lévy et al., 2018) can all, in theory, help create different niches and an emerging facilitation (see last subsection of the Discussion). Finally, resources that are usually considered limiting for all species might in fact not always be: Burson et al. (2018) show that phytoplanktonic taxa specialize on different components of the light spectrum. This constitutes an example of fine-scale resource partitioning of one resource, light, that all species and genera are usually thought to compete for.

No complexity-stability relationship but connections between self-regulation and intergroup interactions

There was no relation between the complexity of the communities (measured as either the weighted 285 connectance or the interaction coefficient variance) and their stability (measured by the largest 286 modulus of the eigenvalues, which quantifies the return time to a point equilibrium, i.e., resilience). 287 This result is conditional upon our model being a good approximate description of the system 288 (i.e., no multiyear limit cycles or chaotic attractors as the mapping between eigenvalues and actual 289 stability is distorted in that case, Certain et al., 2018). However, we already showed on a subset of 290 this data that a fixed point in a MAR(1) model, perturbed by seasonality and abiotic variables, is 291 an accurate description of the system (Barraquand et al., 2018). Therefore, we are confident that 292 the absence of complexity-resilience relationship found here is not a mere artefact of an inadequate 293 model. This absence of direct link between complexity and stability could be an actual feature of empirical systems, as shown previously by Jacquet et al. (2016) using a different technique. This 295 result seems to contradict theory based on random matrices, especially for competitive and/or mutualistic networks (Allesina & Tang, 2012). However, one must bear in mind that such result could 297 also be generated by the limited size of our networks, as random matrix theory relies on asymptotics 298 (Allesina & Tang, 2015). We should also mention that our interaction matrices (discrete-time model) 290 are not strictly analogous to the ones used most frequently in theoretical ecology (continuous-time 300 model), though the spectral radius (largest modulus) can be tightly related to the real part of the lead eigenvalue. Thus the jury is still out regarding the absence of complexity-resilience relation

found here, though it may well be a genuine absence. In addition to complexity metrics, we also found that the percentage of mutualistic interactions, that is thought to affect the stability of a network, either positively or negatively (Mougi & Kondoh, 2012; Coyte et al., 2015; García-Callejas et al., 2018), does not in fact have a major impact on our networks' resilience.

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In addition to weighted connectance and interaction variance, indices at the genus level (vulnera-307 bility and impact) approximate the average effects exerted and sustained by any given taxa in the 308 different study sites. While, at the network level, network structure (either complexity measures 309 or the percentage of mutualistic interactions) did not affect resilience, a relation emerged between 310 self-regulation, necessary for coexistence, and genus-level indices. We found that the more a genus is 311 self-regulated, the more it tends to be vulnerable to other genera's impacts and the less it impacts 312 other genera. We examined whether vulnerability and impact could be affected by phylogenetic 313 correlations; they were not, as on Fig. 3, points were not clustered according to genus, family or 314 phylum. High self-regulation usually indicates large niche differences with the rest of the community, 315 and it makes therefore sense that a species/genus whose needs strongly differ from the others only 316 marginally impacts the resources of the other coexisting species. This is what we expect under strong 317 niche partitioning. A low self-regulation was also correlated with high average abundance, which 318 echoes findings by Yenni et al. (2017) who demonstrated that rare species usually show stronger 319 self-regulation. This correlation between relative rarity and self-regulation could explain the lesser 320 impact of highly self-regulated species/genus: a taxon which dominates the community composition 321 can have a major effect on the others, especially as they usually cover more space, while it is harder 322 for the less common taxa to have large impacts. In contrast, it was more difficult to explain the 323 relationship between self-regulation and vulnerability: a genus that is more self-regulated and less 324 common was found here to be on average more vulnerable to other genera's increases in densities. 325 Such relation implies greater stability (sensu resilience, Ives et al. 2003, and also invariability, 326 Arnoldi et al. 2019) for the network as a whole, because the taxa that are the more vulnerable 327 to other taxa's impacts are also those whose dynamics are intrinsically more buffered. By which mechanisms this could happen is so far unclear and open to speculation. It could just be a "mass 329 effect": common taxa are in high enough numbers to deplete resources or change the environment in

ways that affects the less common ones, but the reverse is not true. As a final note on relationships
between interaction matrix coefficients, we caution that the trends evidenced are all relatively weak:
considerable stochasticity still dominates the distribution of interaction matrix coefficients.

334 Ghosts of competition past and present

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Overall, the dominance of niche differentiation in observational plankton studies – based on our 335 analysis of the REPHY dataset and re-analysis of the MAR(1) literature – is similar to what has been recently found in plant community studies (Volkov et al., 2007; Adler et al., 2018b) or empirically 337 parameterized food webs including horizontal diversity (Barabás et al., 2017). Large niche differences might be due to the ghost of competition past, i.e., competition has occurred in the past, leading to 339 strong selection and subsequent evolution, and then to progressive niche separation. In this scenario, species have evolved niches that allow them not to compete or to interact only weakly (very strong 341 facilitative effects might be likewise destabilizing, Coyte et al., 2015). The likely predator effects that we highlighted above could be comprised within such niche differentiation sensu largo: specialized 343 predators can make strong conspecific density-dependence emerge (Bagchi et al., 2014; Comita et al., 344 2014), while switching generalists can also promote diversity (Vallina et al., 2014). Both predators 345 and resources have often symmetrical effects and can therefore contribute almost equally to such 346 past niche differentiation (Chesson, 2018). 347

An intriguing new possibility, dubbed the "ghost of competition present" (Tuck et al., 2018), suggests by contrast that spatial distributions in relation to abiotic factors might have a large impact on the interaction strengths inferred from temporal interaction models such as ours. Recent combinations of model fitting and removal experiments have shown that model fitting usually underestimates the effect of competitors that are uncovered by removal experiments (Tuck et al., 2018; Adler et al., 2018a). This could occur for instance if species are spatially segregated (at a small scale) because each species only exists within a domain where it is relatively competitive (Pacala's spatial segregation hypothesis, Pacala & Levin 1997), while a focal species could spread out if competitors were removed. This means that a species can be limited by competitors, but act so as to minimize competition (a little like avoidance behaviour in animals) and maximize opportunities

for positive interactions, which implies that competition is in effect hard to detect when all species are present. This mechanism would require spatial segregation between phytoplankton species 359 at the scale of interactions, i.e., at the microscale. At the moment, it is known that the fine-360 scale hydrodynamics generate inhomogeneities at the microscale (Barton et al., 2014; Breier et al., 361 2018) but it is quite unclear how microscale hydrodynamics affects multivariate spatial patterns 362 of species distributions (sensu Bolker & Pacala 1999, or Murrell & Law 2003). Moreover, even if microscale hydrodynamics generates spatial structure with segregation between species, the "ghost of 364 competition present" mechanism might not work for phytoplankton as in terrestrial plants, because turbulence rather than active organism movement dictates where the phytoplankton patches can or 366 cannot appear.

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375 Supporting Information

This article contains supporting information.

377 Authors' contributions

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CP and FB contributed equally to the project design. CP wrote the code for the analyses. FB and CP interpreted the results and wrote the manuscript.

Data accessibility

- The REPHY dataset has already been published (REPHY, 2017) and all scripts for MAR and
- subsequent network analyses are available online in a GitHub repository (https://github.com/
- 384 CoraliePicoche/REPHY-littoral). This repository will be made public upon acceptance and
- codes can be shared with referees should they wish to access them.

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