Impacts of marine invaders on biodiversity depend on trophic position and functional similarity

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ABSTRACT: Impacts of marine invaders on local biodiversity have not been analyzed across invasive species and invaded habitats. We conducted a meta-analysis of 56 field experiments published in 29 papers that examined the effects of marine invaders on local species richness, diversity, and/or evenness. We show that invaders, across studies, typically have negative effects on biodiversity within a trophic level but positive effects on biodiversity of higher trophic levels. For example, both plants and sessile filter-feeders had positive effects on richness and diversity of mobile consumers. The contrasting negative and positive effects on similar versus higher trophic levels are potentially manifested through community-wide antagonism (competition and consumption) versus facilitation (habitat and food provisioning) interactions, respectively. These relationships extended to functional interactions, as sessile invaders had negative effects on the biodiversity of sessile communities (intra-functional interactions) but positive effects on the biodiversity of mobile communities (inter-functional interactions). Our analyses highlight the importance of pairing attributes of the invader and the impacted organisms to obtain simple predictions of how the diversity of entire communities may respond to species invasions on local scales. We also note that our analysis did not require information on co-evolutionary history but that such data, coupled with long-term large-scale mensurative data, are needed to gain a comprehensive predictive insight into invasion impact.

KEY WORDS: Invasive species · Impact analysis · Diversity · Richness · Evenness · Meta-analysis · Attribute matching

INTRODUCTION

Impacts of species invasions have traditionally been studied on individuals and populations of native taxa, while overlooking how entire communities and biodiversity metrics (e.g. taxonomic richness, diversity, and evenness) have been affected (Parker et al. 1999, Byers et al. 2002, Cucherousset & Olden 2011). This lack of emphasis on community-wide effects is reflected in recent reviews of marine invasion impacts (e.g. Byers 2009, Crooks 2009, Grosholz & Ruiz 2009, Rilov 2009b, Thomsen et al. 2011b) where only 2
case-studies were reviewed, highlighting that invasive marsh plants can reduce richness of native plants (reviewed by Byers 2009) and infauna (reviewed by Grosholz & Ruiz 2009). Still, a growing number of primary studies have aimed to quantify impacts on biodiversity (e.g. Ross et al. 2007, Gribben et al. 2009, Bulleri et al. 2010, Thomsen et al. 2010), subsequently inspiring researchers to test broad questions about drivers of invasion impacts across invasive taxa and habitats using meta-analytical frameworks (i.e. calculating direction and magnitude of standardized effect sizes; Gaertner et al. 2009, Thomsen et al. 2009, Powell et al. 2011, Vilà et al. 2011). These meta-analyses show that invasive plants, across studies, have negative effects on local plant biodiversity but either negative (Vilà et al. 2011) or possible positive (Thomsen et al. 2009) effects on animal biodiversity. These conclusions build primarily on data extracted from terrestrial surveys (with the exception of Thomsen et al. 2009).

However, there are potential problems associated with reviewing effects on biodiversity from surveys. First, these data cannot separate causes and effects (Sol et al. 2008) and may therefore confound invasion impact (where the invader is the independent variable) with invasion success (where the invader is the dependent variable). Second, it is difficult to identify all relevant primary studies (because invasion impact from survey data blend into invasion success studies), thereby potentially introducing study selection bias (Englund et al. 1999). Third, surveys are often conducted on larger and longer scales (Sol et al. 2008, Powell et al. 2011) and therefore investigate different ecological mechanisms, sometimes producing contradictory results compared to experiments (Fridley et al. 2007, Powell et al. 2011). By contrast, manipulative field experiments do not confound success and impact and are conducted on relatively similar spatio-temporal scales. Thus, although existing meta-analyses have concluded that invasive plants generally have negative impacts on local plant biodiversity, these studies have not focused on (1) how biodiversity of local animals is affected or (2) how invasive animals impact local biodiversity. Furthermore, primary studies from (3) field experiments and (4) marine ecosystems have been severely underrepresented. Here we target these 4 research gaps, analyzing marine field experiments to test whether invaders occupying different trophic positions have different effects on the biodiversity of communities composed of different trophic positions. More specifically, we tested whether biodiversity effects within a trophic level are more negative than those on a higher trophic level (Fig. 1), because we expect competition to be strong on controphic communities (Gaertner et al. 2009, Thomsen et al. 2009, Powell et al. 2011, Vilà et al. 2011), whereas provision of food and habitats could facilitate higher trophic communities (Rodriguez 2006, Thomsen et al. 2010). We refer to this model as a relative trophic position hypothesis, because we examined whether knowledge about the broad trophic position of both the invader and the impacted community together can explain, and thereby predict, a significant proportion of the large variability often detected between invasion impact studies (Strayer et al. 2006, Thomsen et al. 2011b).

We note that the trophic position hypothesis is related to a broader distinctiveness hypothesis which also emphasizes that the attributes of the invader and the native species should be paired to predict impact (Ricciardi 2003, Thomsen et al. 2011a). The distinctiveness hypothesis suggests that invasion impact is large if invasive and local species are functionally and/or taxonomically different from each other (Diamond & Case 1986, Ricciardi 2003, Ricciardi & Atkinson 2004). This theory does not explicitly specify an expected effect size direction (being ‘larger’ can involve being more negative or more positive). The distinctiveness model has been supported in a meta-

Fig. 1. Conceptual model of the relative trophic position hypothesis. Sessile plants, sessile filter-feeders, and mobile consumers have negative impacts (grey solid arrows, Sub-hypotheses 1 and 2) on controphic local species through competition and consumption but positive impacts (black solid arrows, Sub-hypotheses 3 and 4) on higher trophic levels through provision of food and habitat. Dashed lines correspond to trophic position sub-hypotheses that could not be tested due to a paucity of primary studies. Images and symbols used in the figure courtesy of the Integration and Application Network (ian.umces.edu/symbols/).
analysis that tested effects of taxonomic relatedness (Ricciardi & Atkinson 2004). However, we are not aware of any studies that have tested whether pairing functional attributes can predict invasion impact on the biodiversity of local communities. We therefore repeated the trophic position test with a functional position test, using ‘mobility’ as a simple functional attribute that has often been analyzed in marine invasion studies (Sellheim et al. 2010, Thomsen et al. 2010) and that is related to trophic position (e.g. plants and most filter-feeders are sessile). In this test we compared effect sizes between mobile and sessile invaders on biodiversity of local sessile and mobile communities, expecting that effect sizes are numerically larger across than within the same functional position (Diamond & Case 1986, Ricciardi 2003, Ricciardi & Atkinson 2004).

MATERIALS AND METHODS

We located peer-reviewed studies, in which the abundance of marine invaders was controlled in replicated field experiments and which compared invaded treatments to non-invaded control plots using invader-addition or -removal manipulations (Thomsen et al. 2009), by searching ISI databases, including the Web of Science, Current Contents, and Google Scholar, using various combinations of the terms ‘experiment’, ‘invader’, ‘alien’, ‘exotic’, ‘non-indigenous’, ‘non-native’, ‘impact’, ‘effect’, ‘affected’, ‘community’, ‘diversity’, ‘richness’, ‘assemblage structure’, and ‘marine’ in ‘title’ and ‘abstract’ sections. We also identified relevant experiments by back-tracking references in previous reviews and meta-analyses on marine invasion impacts (Bruno et al. 2005, Schaffelke & Hewitt 2007, Williams 2007, Williams & Smith 2007, Byers 2009, Crooks 2009, Gribben et al. 2009, Rilov 2009a, b, Thomsen et al. 2009, 2011b). We read more than 700 abstracts and method sections and identified more than 100 relevant field experiments (see Thomsen et al. 2011a, b for preliminary data compilations). However, only 29 of these publications reported effects on biodiversity metrics, i.e. on richness, diversity, or evenness with mean values, sample sizes, and standard deviations (or other measures of data dispersion). For each of these papers, we classified manipulated invaders as plants, filter-feeders, or mobile consumers, where filter-feeders were treated as a separate intermediate trophic position that are plant-like in their low mobility and habitat provisioning but animal-like in terms of their heterotrophic diet (see Supplement 1 for details at www.int-res.com/articles/suppl/m495p039_supp.pdf). Unfortunately, we found too few studies to separate consumer effects into grazer, omnivore, and predator effects. To address the functional position hypothesis, plants and filter-feeders were merged as sessile invaders. Impacts on local communities are typically reported on coarse trophic or functional groups and could therefore only be classified as impacts on plant communities (e.g. Piazzi & Ceccherelli 2006), animal communities (e.g. Albins & Hixon 2008), mixed plant+animal communities (e.g. Griffen & Byers 2009), sessile communities (e.g. Thomsen 2010), mobile communities (Sellheim et al. 2010), or mixed mobile+sessile communities (Hollebone & Hay 2008, Thomsen 2010). We also extracted data about the experimental duration and plot sizes to examine whether effects on biodiversity were confounded by spatio-temporal co-variation in the experimental arena. We followed standard methods by only extracting the last data point from repeated-measures experiments (Salo et al. 2007, Vilà et al. 2011) and comparing controls to the highest invader density from multi-density experiments (Vilà et al. 2011), thereby avoiding strong within-experiment autocorrelation (Borestein et al. 2009). We also followed standard methods by treating the same invasive species tested in multiple studies (e.g. Parker et al. 2006, Morales & Traviset 2009) and by including orthogonal and nested experiments (e.g. Gurevitch et al. 2000, Parker et al. 2006) as ‘independent subgroups within a study’ (Borestein et al. 2009) (more conservative approaches supported our findings, see Supplement 2 at www.int-res.com/articles/suppl/m495p039_supp.pdf). A few studies reported impacts on multiple communities, e.g. on plant and animal communities (Gribben et al. 2009), which were treated here as independent observations (Kaplan et al. 2008, Vilà et al. 2011). If an experiment reported impacts on multiple diversity responses, these data were treated as non-independent ‘multiple outcomes/comparisons within study’ (Borestein et al. 2009) and averaged into a single independent effect size per experiment. Effects sizes for each experiment were standardized by calculating Hedges’ d (see Eq. 1), a common metric in ecological meta-analyses (e.g. Salo et al. 2007, Kaplan et al. 2008, Gaertner et al. 2009, Morales & Traviset 2009, Oduor et al. 2010):

$$d = \frac{(X_{\text{invaded plot}} - X_{\text{control plots}})}{SD_{\text{pooled}}} \times J \quad (1)$$

where $X_{\text{invaded plot}}$ and $X_{\text{control plots}}$ represent the mean diversity metric reported from the invaded and non-invaded plots respectively, and J is a correction factor for low sample sizes that approaches 1 for large sam-
Note that if invaders reduce biodiversity, $d$ is negative. Hedges’ $d$ is unit free and ranges from $-\infty$ to $+\infty$ where numerical values of 0.2, 0.5, 0.8, and 1 represent ‘small,’ ‘medium,’ ‘large,’ and ‘very large’ effects, respectively, corresponding to Pearson’s correlation coefficient $r$ of 0.10, 0.24, 0.37, and 0.45 (Rosenberg et al. 2000). We examined potential publication bias by correlating $d$ values with their corresponding sample sizes and variance and by calculating weighted Rosenthal’s fail-safe number (Vilà et al. 2011). Few studies reported effects on diversity and evenness, and these metrics were therefore pooled for the presented analysis.

We found few diversity/evenness data for the relative functional position analysis, and this test was therefore only conducted on richness responses. We analyzed the trophic and functional position hypotheses with unweighted random effect ANOVAs (Kaplan et al. 2008, Morales & Traveset 2009; weighted and fixed models produced similar results, M. Thomsen unpubl.) and calculated cumulative effect sizes ($d_{\text{cumulative}}$), sample heterogeneity ($Q_{\text{total}}$), and 95% bias-corrected confidence limits (CLs). All statistical analyses were carried out in Metawin 2.1 (Rosenberg et al. 2000). A $d_{\text{cumulative}}$ value was interpreted to be significantly different from 0 or another $d_{\text{cumulative}}$ if the 95% CLs did not overlap 0 or each other, respectively (as in Kaplan et al. 2008, Morales & Traveset 2009, Vilà et al. 2011).

**RESULTS**

Our meta-analysis examined impacts on biodiversity of 18 marine invasive species, including 6 plants, 7 sessile filter-feeders, and 5 mobile consumers.

**Trophic position: richness**

The overall heterogeneity of effect size was relatively small ($Q_{\text{total}} = 83$, $df = 69$, $p = 0.12$ where $Q$ approximates an asymptotic chi-squared distribution) indicating that individual effect sizes could share a common value. Importantly, $d_{\text{cumulative}}$ differed significantly among ($Q_{\text{between}} = 19$, $p(\chi^2) = 0.002$, $df = 5$), but not within ($Q_{\text{within}} = 64$, $p(\chi^2) = 0.47$, $df = 64$) the impact types examined. There was a near-significant overall negative effect of invaders on the trophic richness of local communities (Fig. 2A: ‘All’). However, we found clear contrasting positive and negative effects when the attributes of invaders and local communities were separated according to their trophic position. Effects from invaders on native communities that occupy similar trophic positions (plants on plant communities, consumers on animal communities) were statistically similar and significantly more negative than where native communities included higher trophic levels than the invader (plants on ani-

![Fig. 2. Meta-analytical evaluation of the relative trophic position hypothesis (see Fig. 1 for conceptual model). Impacts of marine invaders were evaluated on (A) local taxonomic richness, (B) diversity and evenness combined, and (C) Shannon’s diversity. Invaders (first letters) were divided into plants (P), sessile filter-feeders (F), and mobile consumers (C), and local communities (second letters) into plant- (P), animal- (A), and mixed plant+animal communities (All). Numbers in parentheses correspond to replicated experiments. S-Hyp: sub-hypothesis (see Fig. 1). $d_{\text{cumulative}}$: cumulative effect size. Fill color corresponds to hypothesized dominating interaction, where grey = competition and/or consumption, black = habitat formation and/or food provision. Empty circles correspond to mean effects pooled within and across trophic position and therefore do not have an expected interaction. Error bars are 95% confidence limits.](image-url)
mal communities, plants on mixed plant+animal communities, filter-feeders on animal communities). More specifically, we found significant negative effects of invasive plants on plant communities, and invasive consumers on animal communities, but a significant positive effect of invasive plants on mixed plant+animal communities. We also found a large (but non-significant) effect of invasive plant and filter-feeders on animal communities. Certain trophic position combinations could not be evaluated statistically because of low sample sizes, e.g. impacts of invasive consumers on plant communities \( (d_{\text{cumulative}} = 0.03, n = 1) \) and invasive consumers on plant+animal communities \( (d_{\text{cumulative}} = 0.05, n = 2) \).

**Trophic position: diversity+evenness combined**

The overall heterogeneity of effect sizes was small \( (Q_{\text{total}} = 32, df = 26, p = 0.19) \), again suggesting that effect sizes share a common value. Also, \( d_{\text{cumulative}} \) again differed significantly among \( (Q_{\text{between}} = 9, p(\chi^2) = 0.029, df = 3) \) but not within \( (Q_{\text{within}} = 23, p(\chi^2) = 0.46, df = 23) \) the examined impacts. Although fewer trophic position combinations were available for analysis, our general hypothesis was supported; effects from invaders on native communities that occupy similar trophic positions (invasive plants on plant communities, invasive consumers on animal communities) were statistically similar and significantly more negative than when native communities included higher trophic levels than the invader (plants on animal communities, filter-feeders on animal communities). Invasion impact averaged across trophic positions was not significantly different from 0 (Fig. 2B; ‘All’). However, partitioning by trophic position showed a significant negative impact of invasive plants on plant communities and a near-significant negative impact of invasive consumers on animal communities but a significant positive impact of invasive filter-feeders on animal communities. Effects of invaders on biodiversity of mixed communities were separated according to their trophic position. Effects from invaders on native communities that occupy similar trophic positions (plants on plant communities, consumers on animal communities) were statistically similar and significantly more negative than where native communities included higher trophic levels than the invader (plants on animal communities, filter-feeders on animal communities). More specifically, we found significant negative effects of invasive plants on plant communities, and invasive consumers on animal communities, but significant positive effect of invasive plants on animals and invasive filter-feeders on animal communities. Effects could not be evaluated statistically for some combinations of trophic positions due to low sample sizes or complete lack of primary data.

**Functional position: richness**

The overall heterogeneity of effect sizes was relatively small \( (Q_{\text{total}} = 85, df = 60) \) but significant \( (p = 0.02) \), suggesting that these effect sizes did not share a common value. Again, \( d_{\text{cumulative}} \) differed significantly among \( (Q_{\text{between}} = 29, p(\chi^2) < 0.001, df = 4) \), but not within \( (Q_{\text{within}} = 56, p(\chi^2) = 0.47, df = 56) \) treatments. We found no effect of invaders on the taxonomic richness of local communities when functional attributes were not paired (Fig. 3; ‘All’) but contrasting effects when the attributes were paired. Thus, effects of invaders on local communities that occupied similar functional positions were significantly negative but positive when occupying different positions. Effects of invaders on biodiversity of mixed communities containing both sessile and mobile organisms were not significantly different from 0.

**DISCUSSION**

Marine invaders influence local communities in numerous ways (Crooks 2009, Grosholz & Ruiz 2009), and their impacts are typically considered context-dependent varying with a complex mixture of invader attributes, as well as resource levels, abiotic conditions, and community structure of the invaded habitats (Strayer et al. 2006, Thomsen et al. 2011a, Ricciardi et al. 2013). This complexity is expected to be even more pronounced on aggregated community
theses correspond to replicated experiments. Invaders (first letters) were divided into sessile (S) and mobile (M) species, and local communities (second letters) into sessile (S), mobile (M), and mixed mobile+sessile communities (All). Numbers in parentheses correspond to replicated experiments. Cumulative effect size. Fill color corresponds to hypothesized dominating interaction, where grey = competition and/or consumption, black = habitat formation and/or food provision. Empty circle corresponds to mean effects pooled within and across trophic position and therefore does not have an expected interaction. Error bars are 95% confidence limits.

We suggest that the negative effects seen within trophic levels most likely are driven by competition (as suggested more specifically for plant–plant interactions by Gaertner et al. 2009, Thomsen et al. 2009, Powell et al. 2011, Vilà et al. 2011) and, for animals, a combination of competition and consumption processes. We also suggest that positive effects on higher trophic levels (e.g. invasive plants on animals) could be driven by habitat-formation and food provision. These ecological interactions are also suggested in the reviewed primary studies, e.g. for plant–plant interactions (Viejo 1997, Piazzi et al. 2005), animal–animal interactions (Albins & Hixon 2008), and plant–animal interactions (Thomsen et al. 2010). Impacted communities that included multiple trophic levels (e.g. mixed plant+animal communities) were positively affected, and the underlying facilitation mechanism could therefore again be habitat formation and modification (Gribben et al. 2009, Thomsen et al. 2010). We also found positive impacts of filter-feeders on animal biodiversity—a finding not described in previous meta-analyses. This result suggests that invasive filter-feeders, like plants, may facilitate local animal communities by providing habitat and food. These mechanisms have been demonstrated to be particularly strong for the invasive bryozoan Watersipora subtorquata (Sellheim et al. 2010). Note, however, that data variability was large, reflecting that only few primary studies existed and that invasive filter-feeders also compete with native sessile species for limited space (Blum et al. 2007).

We found contrasting effects on diversity metrics when the functional attributes of the invader and local communities were paired; effects of invaders with similar functions as the impacted community were negative, whereas effects of invaders with different functions were less negative or positive. This finding supports the distinctiveness hypothesis, because effect sizes depend on whatever invasive and local species are functionally or taxonomically different from each other or not (Diamond & Case 1986, Ricciardi & Atkinson 2004). However, our results also highlight that effect size ‘direction’ should be incorporated in the distinctiveness hypothesis to differentiate positive and negative species interactions. Our functional similarity test thereby builds on the previous meta-analysis that focused on negative effects only (Ricciardi & Atkinson 2004). We also suggest that when functionality is equated with mobility (mobile vs. sessile), competition dominates within functional groupings (Viejo 1997, Piazzi et al. 2005), whereas habitat formation and food provisioning
dominate across functional groupings (Sellheim et al. 2010, Thomsen et al. 2010).

Our analyses were likely limited by data variability, research bias, and low inference space, reflecting that the analysis is based on relatively few primary studies. However, data variability was not a major problem because we did find many significant effects to support our general hypotheses. Our analysis could also be limited by different types of auto-correlation between Hedges’ d values, but we found no correlations between effect sizes and plot sizes, experimental duration, and sample sizes (Supplement 2). Auto-correlation may also exist between studies quantifying effects on the same invasive species or under similar experimental conditions, but more conservative supplementary analyses gave similar results (Supplement 2). Finally, our results may be biased because significant results are easier to publish than non-significant results and because ‘pet-species/interactions’ are more fashionable to study than others (Rosenthal 1979, Borenstein et al. 2009). These types of bias are difficult to quantify, but by applying strict study inclusion criteria, we made our data compilation transparent (Supplement 1). It is likely that a standard research emphasis on studying a few high-impact invaders and on publishing large significant effects has inflated the effect sizes we report. However, we suggest that the above biases are less likely to affect effect size direction and that our conclusions are therefore relatively robust. Still, our conclusions should only cautiously be interpolated beyond the context of the reviewed studies (Supplement 1).

From an ecological perspective, it is important to discuss limitations associated with the few broad trophic/functional groupings analyzed. We had to focus on these groups out of necessity because diversity effects are only reported on broad types of communities and because only certain types of invasive species have been tested. The implication of analyzing coarse groupings is that many invasions may not follow the trophic/functional position rules. For example, some invasive plants and filter-feeders can, in contrast to our predictions, have positive effects on biodiversity within trophic groups, e.g. through habitat formation (Thomsen et al. 2010), and invasive plants may have both positive and negative effects on higher trophic levels depending on type and pathway of the dominant interactions, for example, through third- or fourth-order consumption cascades (Eastwood et al. 2007, Tronstad et al. 2010). We did not test how invaders affect biodiversity of lower trophic levels because our literature search did not locate enough relevant primary studies. For example, only 1 study quantified impacts of invasive mobile consumers on biodiversity of plants (Altieri et al. 2009) and none on sessile filter-feeders. Still, we expect that ‘trophic downward’ effects on biodiversity are also predictable (see Fig. 1 and Supplement 3 at www.int-res.com/articles/suppl/m495p039_supp.pdf). For example, invasive predators could have indirect positive effects on plant communities through cascading consumption (Schmitz et al. 2000, Dyer & Letourneau 2003). Net effects of invasive omnivores may be more difficult to predict as they can have negative impacts on both plant and animal diversity through consumption and competition and may also provide food to higher trophic levels, as shown for invasive freshwater omnivores (Nyström & Strand 1996, Perry et al. 1997, Parkos et al. 2003). Finally, the direction of the effect size may vary with environmental conditions, as invaders can compete with or facilitate communities within trophic/functional positions depending on abiotic stress levels (Bertness & Callaway 1994).

The previous discussion on study limitations also outlines research gaps; many types of research bias should be reduced as studies accumulate, in particular if invasive species from under-represented trophic and functional groups are targeted. It is also important that more studies report biodiversity effects on finer trophic/functional groupings. For example, separating diversity impact for the local plant, filter-feeder, grazer, omnivore, and predator communities would be an important improvement. Future syntheses should also quantify biodiversity effects on lower trophic levels and pair positions of the invader and the local species with their size structures, ontogenies, strategies, densities, and inter-versus intra-trophic functional groupings (e.g. understory, canopy, and encrusting communities responding differently to invaders, Britton-Simmons 2004). Finally, experimental data need to be compared to survey data, to better understand how impact may depend on spatio-temporal scales, invasion history, and methodological artifacts.

In conclusion, we have shown that invaders, across studies, typically have negative effects on biodiversity within a trophic level but positive effects on biodiversity of higher trophic levels. These results confirm the importance of pairing attributes of the invader and the impacted local community to provide simple predictions about how biodiversity is affected by invaders (Ricciardi 2003, Thomsen et al. 2011a). Our broad predictions do not require detailed trait data about the invader or theories about (lack of) co-
evolutionary history with local communities, but this information is important to add to gain deeper predictive insight into invasion impacts (Strayer et al. 2006, Ricciardi et al. 2013). We also note that too few studies existed to calculate effects for many combinations of invaders and community attributes. More studies are therefore needed so that high-resolution trophic/functional position models can be merged with models that pair different invader and community attributes.

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