

1 **Title:** Non-lethal effects of an invasive species in the marine environment - the
2 importance of early life-history stages

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26 **Abstract**

27 Studies examining the effects of invasive species have traditionally focused on the
28 direct/lethal effects of the invasive on the native community but there is a growing
29 recognition that invasive species may also have non-lethal effects. In terrestrial
30 systems, non-lethal effects of invasive species can disrupt early life-history phases
31 (such as fertilization, dispersal and subsequent establishment) of native species but in
32 the marine environment, most studies focus on adult rather than early life-history
33 stages. Here, we examine the potential for an introduced sessile marine invertebrate
34 (*Styela plicata*) to exert both lethal and non-lethal effects on a native species
35 (*Microcosmus squamiger*) across multiple early life-history stages. We determined
36 whether sperm from the invasive species interfered with the fertilisation of eggs from
37 the native species and found no effect. However, we did find strong effects of the
38 invasive species on the post-fertilisation performance of the native species. The
39 invasive species inhibited the settlement of native larvae and, in the field, the presence
40 of the invasive species was associated with a 10-fold increase in the post-settlement
41 mortality of the native species, as well as an initial reduction of growth in the native.
42 Our results suggest that the larvae of the native species avoid settling near the
43 invasive species due to reduced post-settlement survival in its presence. Our results
44 also show that invasive species can have complex and pervasive effects (both lethal
45 and non-lethal) across the early life history stages of the native species which are
46 likely to result in its displacement and to facilitate further invasion.

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48 **Key words:** settlement, invasive species, fertilisation, postmetamorphic performance,
49 trait-mediated effects.

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Introduction

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53 Invasive species can have a range of effects on native species and lethal effects
54 are most commonly cited as the source of negative impacts on established
55 assemblages (Ruiz et al. 1999; Strayer et al. 2006). For example, invasive species can
56 prey upon native species, cause competitive displacement or modify local disturbance
57 regimes (Mack and D'Antonio 1998; Snyder and Evans 2006). Whilst the impact of
58 lethal effects on native species is becoming clear, the prevalence and role of non-
59 lethal effects in species invasions has only recently started to be considered (e.g.
60 Trussell et al. 2006). This is despite the recent recognition that non-lethal effects can
61 have major impacts on the dynamics of communities (Trussell et al. 2003, Werner and
62 Peacor 2003) and initial indications that introduced species can be a source of non-
63 lethal effects (Nystrom et al. 2001; Pangle and Peacor 2006). In terrestrial plant
64 systems, there is a growing recognition that invasive species can affect every phase of
65 the life-histories of native species. For example, high densities of flowering invasives
66 can disrupt the pollination of native species resulting in lower seed production
67 (Bjerknes et al. 2007). Invasives can also affect the dispersal syndromes of seeds,
68 disrupting frugivore mutualisms that are crucial for the effective dispersal of native
69 species (Christian 2001). Thus, the effects of invasive species can extend beyond
70 simple competitive interactions during the adult phase: non-lethal effects disrupt the
71 production and dispersal of native recruits, seriously exacerbating the effects of the
72 invasive species. This is especially important for marine sessile organisms, for which
73 “supply-side” processes can be important determinants of population dynamics
74 (Underwood and Keough 2001).

75 Many marine benthic organisms have been moved around the world's oceans
76 since ancient times by means of shipping (Carlton 1999), but the last century has seen
77 a dramatic rise in the rate of introductions of alien marine species (Cohen and Carlton
78 1998; Mack et al. 2000). As a result, non-indigenous species have been moving
79 beyond physical boundaries such as those created by ocean currents, and have spread
80 worldwide (Wonham et al. 2001). The invasion of non-indigenous species is now
81 regarded as one of the major threats to marine biodiversity and the number of studies
82 examining the effects of marine invasive species has increased dramatically (Ruiz et
83 al. 1997; Grosholz 2002; Galil 2007). Most studies examining the effects of invasive
84 species in the marine environment have focused on competitive displacement or
85 predation as the major impact of the invasive species and many have been restricted to
86 examinations of the adult phase (but see Byers and Goldwasser 2001; Trussell et al.
87 2006). More recently however, it has been recognised that invasive species in the
88 marine environment can have strong, indirect effects on native communities. For
89 example, introduced species can change trophic cascades in marine foodwebs
90 (Trussell et al. 2002, 2004; Kurle et al. 2008), reduce larval production (Gribben and
91 Wright 2006) and change the behaviour (and hence, distribution) of prey species
92 (Trussell et al. 2003). These studies strongly suggest that marine invasive species
93 have pervasive effects at a range of life-history stages and levels of community
94 organisation in the marine environment.

95 The life-history of marine organisms suggests that any non-lethal effects of
96 invasive species on the early-life-history stages of native species are likely to be
97 important. Most marine organisms are broadcast spawners, releasing eggs and sperm
98 into the water column. Due to the high rate of sperm dilution, the fertilisation of eggs
99 is rarely complete and fertilisation rates can range between 0 and 100% with mean

100 rates of ~50% in many instances (Levitan and Petersen 1995; Yund 2000).
101 Importantly, heterospecific sperm can disrupt fertilisation in broadcast spawners,
102 resulting in lower fertilisation rates (Lambert 2000; Lambert 2001). This raises the
103 possibility that marine invasive species could disrupt/reduce fertilisation success in
104 broadcast spawners analogously to pollination disruption in terrestrial systems,
105 although this possibility has not been explored. Similarly, marine invertebrate larvae
106 sometimes avoid settling near dominant competitors (Grosberg 1981; Stoner 1994;
107 but see Bullard et al. 2004). Given that marine invasive species can be competitively
108 dominant (Reusch and Williams 1999; Piazzini and Ceccherelli 2002) one might expect
109 that the larvae of native species reject settlement sites adjacent to invasive species.
110 This non-lethal effect on the dispersal of native species is analogous to the
111 disruption/reduction of frugivore mediated dispersal by invasive species in plants.
112 This potentially important effect of invasive species in the marine environment has
113 received less attention than other life-history stages. This is surprising given that the
114 supply of new recruits into marine populations can have major influences on
115 subsequent community structure (Underwood and Keough 2001) and the production
116 of zygotes has the potential, at least, to limit population growth in broadcast spawners
117 (Levitan 1995). Finally, mortality immediately following settlement can be intense in
118 sessile marine organisms and can be a major determinant of adult distributions and
119 abundance (Gosselin and Qian 1997). Given the ecological importance of the early
120 post-metamorphic period, any influence that invasive species may have during this
121 stage could have major implications for the population dynamics of native species.

122 Here we examine the effects of an introduced marine species (*Styela plicata*)
123 on a native species (*Microcosmus squamiger*) across the early life-history stages, from
124 fertilisation to larval settlement through to post-metamorphic performance. As both

125 species coexist in the studied area (SE Australia), we wanted to explore the
126 interactions between them. Given the potential for non-lethal and lethal effects to
127 interact synergistically (e.g. Meyer and Byers 2005), we investigated both types of
128 effects across different stages of the life-history. We chose solitary ascidians as our
129 study organism as they are one of the major invasive groups in marine systems
130 (Lambert 2007). We first examined whether the presence of heterospecific sperm
131 from an invasive species reduced the fertilisation success of the eggs of a native
132 species. We then examined the larval settlement responses of each species in the
133 presence and absence of heterospecific and homospecific settlers. Finally, we
134 examined the post-metamorphic survival and growth of both species in the presence
135 and absence of heterospecific recruits in the field. We found strong, non-lethal effects
136 on larval settlement and direct, lethal effects on post-metamorphic survival, as well as
137 an initial reduction in growth, suggesting that this marine invasive species has the
138 potential to dramatically change the population dynamics of native species.

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Materials and Methods

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Study site and species

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Microcosmus squamiger is native to Australia (Kott 1985; Rius et al. 2008)

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and occurs subtidally on artificial and natural substrata in sheltered areas where it can

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form dense populations (Kott 1985; and pers. obs.). *S. plicata* is considered an alien

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species in Australian waters (Hewitt 2002; Wyatt et al. 2005) and although there is no

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available information about when and where exactly this species was introduced, it

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now successfully colonizes shallow habitats in SE Australia (pers. obs.). Both species

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are solitary ascidians and they reach similar sizes (ca. 5-10 cm) as adults. At the

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Manly Marina (27°27'10"S, 153°11'22"E, Brisbane, Queensland, Australia), *S.*

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plicata is found inside the harbour attached to the floating pontoons while *M.*

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squamiger can only be found outside the harbour, with a small area at the entrance of

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the harbour where both species coexist (on the outermost pontoons). Reproductively

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mature *M. squamiger* and *S. plicata* were collected from these outer pontoons of

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Manly Marina between October and December 2006. They were then transported in

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insulated aquaria back to the laboratory (~45 min. journey) and kept in a tank with 20

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l of constantly aerated seawater at room temperature.

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General methods - production and settlement of larvae

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To extract eggs and sperm for our experiments, we used standard protocols as

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described by Marshall et al. (2000) for strip spawning solitary ascidians. To produce

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pools of fertilised eggs, we used the sperm of three individuals and the eggs of one

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individual (both species are simultaneous hermaphrodites with an almost complete

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block to self fertilisation; Rius unpubl. data). We left the gametes in contact for 45

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minutes and we then rinsed the sperm with filtered seawater and pooled the eggs from

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four individuals.

165 To produce larvae, we fertilised eggs as above and then placed the developing
166 embryos into an aerated beaker (containing ~ 500 ml of filtered seawater) in a
167 constant temperature cabinet at 20°C. In both species studied here, larvae hatch within
168 14 hours of fertilisation. Afterwards, the larvae were pipetted out and placed in the
169 experimental Petri dishes. We used pre-roughened 90mm Petri dishes that had been
170 maintained in aquaria with seawater for several days so that they could develop a
171 biofilm which facilitates larval settlement (Wieczorek and Todd 1997). After 24
172 hours, we gently rinsed the Petri dishes in seawater to remove any unattached larvae.

173 *Experiment 1: Does the presence of heterospecific sperm from an invasive reduce*
174 *fertilisation success in a native?*

175 We examined whether the prior exposure of *M. squamiger* eggs to *S. plicata*
176 sperm affected subsequent fertilization success. Eggs from a *M. squamiger* individual
177 were split in 3 groups. The 1st group was a control (i.e. no exposure to *S. plicata*
178 sperm), the 2nd group was exposed to a ‘low’ concentration ($\sim 10^5$ sperm.ml⁻¹) of *S.*
179 *plicata* sperm and the second to a ‘high’ concentration ($\sim 10^7$ sperm.ml⁻¹) of *S. plicata*
180 sperm. Sperm concentrations were estimated using three replicate counts on a
181 modified Fuchs-Rosenthal Haemocytometer. The *M. squamiger* eggs were exposed to
182 *S. plicata* sperm in a final volume of 100 ml for fifteen minutes, a period of time long
183 enough to make sure that, if there was a glycosidase release from *M. squamiger* eggs,
184 this release was completed (Lambert 2000), before being rinsed free of sperm in
185 filtered seawater. The eggs were then placed in new Petri dishes and all the eggs of
186 the 3 treatments (control, low and high) were exposed to *M. squamiger* sperm ($\sim 10^7$
187 sperm.ml⁻¹) pooled from 4 individuals for 45 minutes. We then rinsed the eggs again
188 in filtered seawater, placed them in a constant temperature cabinet at 20°C and
189 allowed the embryos to develop for fourteen hours. We then assessed fertilisation

190 success by counting the proportion of eggs that developed into unhatched embryos or
191 hatched larvae relative to unfertilised eggs. We repeated this experiment for the eggs
192 of three different individuals (i.e. 3 runs). To analyse the data, we first arcsine-square
193 root transformed the data (which was estimated as the proportion of eggs fertilised).
194 We analysed the data as an unreplicated block design where run was a random factor
195 and exposure history was a fixed factor.

196 *Experiment 2: Does the presence of recruits affect settlement?*

197 We were interested in whether the presence of heterospecific and homospecific
198 recruits affected the settlement behaviour of both species. For each species, at the 14
199 hour mark after fertilization, we gently pipetted 40 larvae into new Petri dishes. We
200 allowed them to settle (until 24 hour mark) and then gently washed off any unattached
201 larvae. We then introduced 40 homospecific or heterospecific larvae (depending on
202 the treatment) from a new fertilization event and counted how many of these new
203 larvae had attached after 24 hours. In these experiments, Petri dish was the unit of
204 replication. The experiments using still water were the only reliable way to prevent
205 the larvae to quit the system and to quantify settlement rates of a controlled larval
206 pool.

207 We examined the effect on settlement of pre-established recruits in all possible
208 combinations: the effect of *S. plicata* recruits on *M. squamiger* settlement, of *M.*
209 *squamiger* recruits on *S. plicata* settlement, of *M. squamiger* recruits on *M. squamiger*
210 settlement and, finally, the effect of *S. plicata* recruits on *S. plicata* settlement (Table
211 1). In all of these experiments, we compared settlement in treatments consisting of
212 Petri dishes with recruits to settlement in controls consisting of Petri dishes without
213 pre-established settlers and we used the same number of control and treatment

214 replicates. The number of runs and replicates, as well as the initial recruit densities in
215 the treatment dishes, are listed in Table 1.

216 Because settlement was measured as the proportion of larvae that settled, we
217 first arcsine-square root transformed the data. We analysed the effect of the presence
218 of heterospecific recruits on settlement using a two-way, mixed model ANOVA
219 where the experimental treatment was a fixed factor and experimental Run was a
220 random factor. When we examined the effect of *M. squamiger* recruits on *S. plicata*
221 settlement, we found no interaction between Run and treatment and, given that Run
222 explained little variance and was of no biological interest, it was omitted from the
223 final model (Quinn and Keough 2002). For the effect of homospecific recruits for
224 each species (one run only), we used a t-test to compare the experimental treatment
225 with the control.

226 *Experiment 3: Does the presence of competing recruits affect post-metamorphic*
227 *performance?*

228 We were interested in whether the presence of heterospecific recruits affected
229 the subsequent performance of our two focal species. Thus we settled *M. squamiger* in
230 the presence of *S. plicata* recruits and settled *S. plicata* in the presence of *M.*
231 *squamiger* as described above. Controls consisted of Petri dishes in which larvae were
232 settled in the absence of any pre-established recruits. We used 8 replicates (i.e. Petri
233 dishes) each per treatment and control for each species. The mean initial density of
234 recruits in the *M. squamiger* experiment did not differ among treatments (mixed
235 treatment mean was 16.625 (SD = 2.615) and the control was 19.375 (SD = 3.701); t-
236 test, $t = -1.716$, $n = 8$, $P = 0.108$), and the same was found for the *S. plicata*
237 experiment (mixed treatment mean was 20.375 (SD = 8.105) and the control was 14.5
238 (SD = 4.276); t-test, $t = 1.813$, $n = 8$, $P = 0.098$). We marked all the settler positions in

239 the Petri dishes, numbering them on the surface of the dishes using a pencil. We then
240 drilled an 8 mm hole in the centre of each Petri dish. The dishes were transported to
241 the field within ~45 minutes, in 20 l insulated containers. We attached the Petri dishes
242 to a Perspex backing plate (500 x 500 x 8 mm) using stainless steel screws. The Petri
243 dish positions were randomly assigned. Then, we hung the plates from the most
244 external pontoon of the Manly harbour at a depth of 2 m (the dock floated at water
245 level regardless of tide), facing down to reduce the effects of light and sedimentation
246 (following Marshall et al. 2003a). For the experiment examining the effect of *S.*
247 *plicata* recruits on the post-metamorphic performance of *M. squamiger*, we measured
248 the survival of the *M. squamiger* settlers 1, 2, 5 and 10 weeks after being deployed
249 into the field. We assessed survival as presence/absence of previously marked settlers
250 on the Petri dish, a measure that is likely to reflect survival as reattachment to surfaces
251 following removal is rare in ascidians (but see Edlund and Koehl 1998; Bullard et al.
252 2007). During each census of survival, we brought the Petri dishes back to the
253 laboratory, assessed survival and removed any additional organisms that had settled in
254 the intervening period. We also measured the size of recruits after 2, 5 and 10 weeks
255 in the field by taking digital photographs of the diameter of the settlers with a camera
256 attached to the dissecting microscope and connected to a computer. We subsequently
257 measured the photographs using Image Pro (v. 5.1.0.12, Media Cybernetics) and we
258 calibrated the measurements by taking a photograph using the haemocytometer grid.

259 For the experiment examining the effect of *M. squamiger* recruits on the post-
260 metamorphic performance of *S. plicata*, we assessed survival only 1, 2 and 4 weeks
261 after deploying the settlers in the field. This last experiment had to be halted after 4
262 weeks because the settlement plates were vandalised.

263 To analyse the survival and growth data, we used a repeated measures
264 ANOVA where Petri dish was the unit of replication. Because survival was measured
265 in proportions, we used arcsine- square root transformed data.

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Results

267 *Experiment 1: Does the presence of heterospecific sperm from an invasive reduce*
268 *fertilisation success in a native?*

269 Although the random factor Run (= individual) was significant, reflecting
270 differences in fertilization rates among individuals, there was no significant effect of
271 heterospecific sperm on the fertilisation success of the native species at either sperm
272 concentration (Table 2), nor was there any trend for a negative or positive effect.

273 *Experiment 2: Does the presence of recruits affect settlement?*

274 There was a strong effect of *S. plicata* recruits on the settlement of *M.*
275 *squamiger* (Fig. 1a). Table 3 shows that there was a strong interaction between
276 experimental Run and the treatment of interest. Because the denominator for the F
277 ratio to test the main effect is the $MS_{\text{interaction}}$, the P value for the main effect was not
278 statistically significant. However, the direction of the effect of *S. plicata* recruits on
279 *M. squamiger* settlement was consistently negative. The significant interaction was
280 simply due to the size of this effect: in Run 1, *S. plicata* had ~3-fold reduction on *M.*
281 *squamiger* settlement but in Run 2, the effect was only a ~2-fold reduction. In
282 contrast, the presence of conspecific recruits had no effect on the settlement of *M.*
283 *squamiger* (t-test, $t = 0.425$, $n = 24$, $P = 0.675$; Fig. 1a).

284 *S. plicata* settlement was lower in the presence of *M. squamiger* recruits and the
285 size of the effect was more consistent among experimental runs (Table 3; Fig. 1b).
286 The non-significant interaction term allowed us to test a reduced model in which both
287 treatment and Run proved highly significant. Again, we found no effect of
288 homospecific recruits on *S. plicata* settlement (t-test, $t = 0.159$, $n = 8$, $P = 0.879$; Fig.
289 1b).

290 *Experiment 3: Does the presence of heterospecific recruits affect post-*
291 *metamorphic performance?*

292 The proportion of *M. squamiger* recruits surviving in the field decreased over
293 time. The presence of *S. plicata* had a strong negative effect on the subsequent
294 survival of *M. squamiger* in the field (Fig. 2a). After ten weeks in the field, the mean
295 proportion of *M. squamiger* that had survived was ~33% in the absence of *S. plicata*
296 but was <5% in the presence of *S. plicata*. This difference in survival appeared to be
297 driven by the initial responses of the two treatments; there were large differences in
298 survival after the first week and they persisted through time (Table 4).

299 In contrast to the effect of *S. plicata* on *M. squamiger*, the presence of *M.*
300 *squamiger* had no effect on the subsequent survival of *S. plicata* after four weeks in
301 the field (Table 4; Fig. 2b).

302 It was impossible to photograph all *M. squamiger* recruits from the Petri dishes,
303 owing to the fact that some have settled in the corner of the dish and thus reliable
304 measurements with photographs were not possible. However, a large proportion of
305 individuals were successfully photographed (2nd week: mixed - 72.72%, control -
306 50.53%; 5th week: mixed - 66.66%, control - 93.85%; and 10th week: mixed - 100%,
307 control - 83.33%). In the 2nd week of the experiment, the *M. squamiger* recruits in
308 presence of *S. plicata* were significantly smaller than those in the controls but this
309 difference disappeared after 5 weeks (Table 5; Fig. 3). After 10 weeks no statistical
310 comparisons were possible as there was only one remaining *M. squamiger* recruit in
311 the mixed treatment.

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Discussion

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The presence of the invasive ascidian *Styela plicata* affected a number of crucial life-history stages in the native ascidian *Microcosmus squamiger* and, overall, a combination of lethal and non-lethal effects of the invasive may synergise to exclude *M. squamiger* from its native habitat. These results further expand our understanding of how sublethal effects of invasive organisms affect natives, and reaffirm the importance of such effects during early life-history stages.

We found no effect of *S. plicata* sperm on the fertilisation success of *M. squamiger* eggs. In previous studies (Lambert 2000; Lambert 2001), homologous and heterologous sperm were mixed, while in our experiment we washed the eggs before exposure to homologous (*M. squamiger*) sperm. In this way we excluded the possible negative effects of sperm competition. As a result, we restricted our observation to whether or not exposure to the sperm of the invasive was affecting fertility of the native eggs. In light of our results, we found that *S. plicata* neither activate *M. squamiger* eggs nor interfere with subsequent egg activation. The lack of interference of *S. plicata* on fertilisation of *M. squamiger* eggs may be because the two species are not closely related and thus sperm recognition proteins are highly divergent.

Alternatively, given that these species live sympatrically, there may have been a strong positive selection on sperm-egg recognition proteins to reduce costly hybridisation (Byrd and Lambert 2000; Veen et al. 2001; Harper and Hart 2005). It would be interesting to repeat our experiments in populations that are not sympatric but in our populations it appears that the invasive species does not interfere with the fertilisation success of the native species. In contrast, the effects of the invasive on the post-fertilisation performance of the native species were more dramatic.

338 Inhibition of settlement by superior competitors has been demonstrated in a
339 number of marine invertebrates (e.g. Grosberg 1981; Young and Svane 1989; Davis et
340 al. 1991) but its prevalence remains in debate (Bullard et al. 2004). In our system,
341 both species avoided settling in the presence of the other but only one species had a
342 significant, negative effect on post-metamorphic performance. The reason for the
343 negative effect of *M. squamiger* on *S. plicata* settlement remains unclear, but may be
344 due to a general avoidance response of ascidian larvae (e.g. Stoner 1994). Regardless,
345 the effect of each species on settlement of the other suggests that species recognition
346 at settlement is acting in these two species, even if *S. plicata* seems to be a relatively
347 recent introduction to Australian waters (Wyatt et al. 2005).

348 The inhibition of settlement of native larvae in the presence of the exotic is
349 analogous to the disruption of dispersal syndromes in plants whereby the presence of
350 an invasive species reduces the effective dispersal of native propagules. However, in
351 our study, the effect of inhibiting settlement may have a number of additional,
352 potentially dramatic consequences (Elkin and Marshall 2007). Inhibiting settlement
353 essentially forces larvae to continue to search for alternative suitable habitat and this
354 increase in searching time carries a number of direct and indirect costs. Mortality
355 while dispersing in the water column can be extremely high and thus any native larvae
356 that are inhibited from settling by invasive recruits may experience higher rates of
357 mortality than they would in the absence of the invasive (Morgan 1995). Furthermore,
358 in species with non-feeding larvae such as the ascidians and other marine organisms,
359 increasing the duration of the larval phase can result in reduced performance after
360 metamorphosis - larval swimming is costly and reduces the level of reserves available
361 for post-metamorphic survival and growth (Wendt 1998, Maldonado and Young
362 1999; Marshall et al. 2003b; Pechenik 2006). Thus, the post-metamorphic

363 performance of native settlers may be lower in places where the invasive species is
364 more common and inhibits settlement. Overall then, the inhibition of native larval
365 settlement by invasive recruits may negatively affect native populations in three ways:
366 decrease settlement directly, increase planktonic mortality and decrease post-
367 metamorphic performance. Previous work has shown that native species change their
368 behaviour (and thus their distribution) in response to invasive predators (Trussell et al.
369 2002, 2003). Our findings suggest that competition from invasive species can also
370 drive changes in the behaviour of native species.

371 The presence of *S. plicata* in the field increased the juvenile mortality of *M.*
372 *squamiger* by 10 fold. In addition, we found a significantly reduced growth of *M.*
373 *squamiger* in mixed treatments compared to the controls in the 2nd week. This trend
374 was not maintained in the following weeks, which is perhaps unsurprising as the
375 densities of *M. squamiger* in the mixed treatments declined dramatically over those
376 first weeks and high levels variation among the few survivors prevented a meaningful
377 comparison. Although the reason for the decreased survival and growth of the native
378 in the presence of invasive needs to be further investigated, we consider that there are
379 three (non-mutually exclusive) mechanisms for the negative effect of invasive species
380 on the survival and growth of the native species: competition for food, allelopathy or
381 indirect effects mediated by third species. We favour the first hypothesis, *S. plicata*
382 may be a better competitor for food than *M. squamiger* and thus *M. squamiger* may
383 have had higher mortality and reduced early growth due to starvation. Conversely, the
384 presence of pre-established *M. squamiger* had no effect on post-metamorphic
385 performance of *S. plicata*. Given that water flow rates were reasonably low at the
386 study site, it is possible that a better competitor could deplete the local abundance of
387 food in the boundary layer above the plates. Competition for space seems unlikely due

388 to the small size of the recruits during the first weeks, and it might have only been
389 important in the last weeks of the experiment when the animals have grown enough to
390 physically interact. However, the most drastic reduction in survival and growth of the
391 mixed treatments in comparison to the control treatments occurred in the in the first
392 few weeks. It is interesting in this sense that, in the experiment in which we analysed
393 the effect of *M. squamiger* recruits on *S. plicata* performance (and found no effect),
394 the pre-established *M. squamiger* themselves experienced high mortalities (similar to
395 those in the experiment with pre-established *S. plicata*, data not shown). In other
396 words, the presence of *S. plicata* affected the survival of *M. squamiger* even if the
397 recruits of the latter arrived before and were already in place.

398 While we believe that the most likely source of the effect of *S. plicata* on *M.*
399 *squamiger* survival in the field was competition, we must also consider other potential
400 explanations. Allelopathic effects of invasive species on natives have been found in
401 some studies (Schenk 2006; Figueredo et al. 2007), and in our study the interaction of
402 the two species might induce the production of waterborne allelopathic metabolites in
403 the introduced species that could reduce both survival and growth of the native. An
404 alternative mechanism for the negative effect of the invasive on the native species in
405 the field is that there are indirect effects via a third organism. For instance, the
406 presence of the invasive may increase predation on the native species but leave the
407 invasive unaffected. While such a scenario does not explain the early differences in
408 growth, it may still explain the differences in survival. In our experiments, the
409 experimental plates were hanging from the pontoon, which excluded benthic
410 predators, but fish could still access the experimental individuals. Although this
411 scenario seems unlikely, carefully designed predator exclusion experiments that do
412 not interfere with food supply would be necessary to rule it out. Regardless of the

413 underlying direct or indirect mechanisms, our study joins a growing list showing that
414 the presence of marine invasive species is likely to result in the reduced abundance of
415 local biota (Bando 2006).

416 The effects of *S. plicata* on the settlement and survival of *M. squamiger* and
417 the reciprocal effects of *M. squamiger* on *S. plicata* settlement have some interesting
418 implications for the dynamics of invasion in this system. We suggest that the presence
419 of the native incumbent inhibits invasion by *S. plicata*. However, if a disturbance
420 clears space for *S. plicata* to settle, then they will outcompete any newly settled *M.*
421 *squamiger* and furthermore will inhibit recolonisation by the native. We also found
422 that the presence of *S. plicata* recruits did not reduce *S. plicata* settlement success
423 suggesting that initial invasion will not interfere with further arrivals. Previous studies
424 have shown that both disturbance and prior invasion facilitate further invasion
425 (Crooks 2002; Rodriguez 2006; Altman and Whitlatch 2007), here we provide one
426 potential mechanism for such an effect. While our results appear to be a classic case
427 of a priority effect (sensu Almany 2003), interestingly, this effect is not mediated by
428 resource limitation: there was ample space for larvae to settle (only ca. 0.01 % of the
429 Petri dish surface is occupied by pre-established settlers), they are simply inhibited
430 from doing so. Whether propagule pressure can reach levels that overwhelm the
431 'biotic resistance' of the community associated to *M. squamiger* (e.g. Hollebone and
432 Hay 2007) remains unclear but at least initially, the presence of the native species
433 appears to inhibit the invasion by the introduced species (Osman and Whitlatch 1995),
434 even at different spatial scales (Stachowicz et al. 2002).

435 Overall, we found a mixture of lethal and non-lethal effect of the invasive
436 species on the native species. These effects may lead to the invasive species
437 outcompeting the native species whenever space becomes available. This study

438 suggests that invasive species can have significant non-lethal and lethal effects on
439 early life-history stages of native species in the marine environment. Further
440 experiments comparing settlement success in presence or absence of invader recruits
441 in water flow devices (see Butman et al. 1988), as well as experiments assessing the
442 interaction during adult phases will provide further understanding of the interactions
443 between invasive and native sessile marine invertebrates.

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683 **Table and figure legends**

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685 **Table 1.** Experimental treatments used to evaluate the effect on settlement of
686 pre-established recruits using all combinations of *Styela plicata* and *Microcosmus*
687 *squamiger* larvae and settlers. SD, standard deviation.

688 **Table 2.** ANOVA examining the effect on fertilisation success of pre-
689 exposing *Microcosmus squamiger* eggs to *Styela plicata* sperm. Note that the model is
690 reduced after testing for a non-significant interaction between Run and the treatment
691 of interest. Significant p values are shown in **bold**.

692 **Table 3.** ANOVA examining the effect of settled heterospecific recruits on the
693 settlement of a) *Microcosmus squamiger* larvae and b) *Styela plicata* larvae. Note that
694 model in section b is reduced after testing for a non-significant interaction. Significant
695 p values are shown in **bold**.

696 **Table 4.** Repeated measures ANOVA examining the effect of the presence of
697 one species on the survival of the other in the field. Significant p values are shown in
698 **bold**.

699 **Table 5.** Repeated measures ANOVA examining the effect of the presence of
700 *Styela plicata* on the size of the *Microcosmus squamiger* in the field. Significant p
701 values are shown in **bold**.

702 **Figure 1.** Results of experiment 2 testing whether the presence of recruits
703 affected settlement, pooling runs. Shaded bars indicate controls and open bars indicate
704 established recruits: (a) effect of *Styela plicata* and *Microcosmus squamiger* recruits
705 on the settlement of *M. squamiger*; and (b) effect of *M. squamiger* and *S. plicata*
706 recruits on the settlement of *S. plicata*. Vertical bars denote standard error.

707 **Figure 2.** Results of experiment 3 assessing if the presence of heterospecific
708 recruits affected post-metamorphic survival in the field: (a) *Microcosmus squamiger*

709 (b) *Styela plicata*. Dotted lines indicate the treatment in presence of heterospecific
710 recruits and solid lines indicates the treatment with no pre-established recruits.
711 Vertical bars denote standard error.

712 **Figure 3.** Results of experiment 3. Mean size of *Microcosmus squamiger*
713 juveniles after two and five weeks in the field. Dotted lines represent juveniles in the
714 presence of *Styela plicata*, solid lines represent control juveniles. The vertical bars
715 denote standard error and note the log scale on the y-axis.

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734 **Tables & figures**

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736 Table 1

Treatment	Run	Number of replicates	Mean number of initial recruits	SD
<i>S. plicata</i> on <i>M. squamiger</i>	1	8	10.375	1.179
	2	12	18	1.243
<i>M. squamiger</i> on <i>M. squamiger</i>	1	12	14.667	1.437
<i>M. squamiger</i> on <i>S. plicata</i>	1	8	12.750	2.455
	2	4	13.5	2.255
<i>S. plicata</i> on <i>S. plicata</i>	1	4	20.25	3.351

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751 Table 2

Source	df	MS	F	P
Experimental Run	2	0.083	16.44	0.012
Heterospecific sperm	2	<0.001	0.07	0.931
Error	4	0.005		

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772 Table 3

Source	Df	MS	F	P
a) Effect of <i>S. plicata</i> on <i>M. squamiger</i>				
Treatment	1	0.741	6.55	0.237
Experimental Run	1	0.011	1.04	0.313
Treatment x Experimental Run	1	0.113	11.18	0.002
Error	36	0.010		
b) Effect of <i>M. squamiger</i> on <i>S. plicata</i> settlement				
Treatment	1	0.212	17.79	<0.001
Experimental Run	1	0.098	8.25	0.009
Error	21	0.012		

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787 Table 4

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Source	df	MS	F	P
a) Effect of <i>S. plicata</i> on <i>M. squamiger</i>				
<i>Between Subjects</i>				
Treatment	1	3.683	14.70	0.002
Error	13	0.250		
<i>Within Subjects</i>				
Time	3	1.137	34.69	<0.001
Time x Treatment	3	0.032	0.97	0.417
Error	39	0.033		
b) Effect of <i>M. squamiger</i> on <i>S. plicata</i>				
<i>Between subjects</i>				
Treatment	1	0.005	0.05	0.823
Error	14	0.088		
<i>Within subjects</i>				
Time	2	0.217	20.48	<0.001
Time x Treatment	2	0.001	0.098	0.907
Error	28	0.011		

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795 Table 5

Source	df	MS	F	P
Treatment	1	5.65	2.79	0.1336
Error	8	2.03		
<i>Within Subjects</i>				
Time	1	281.31	179.86	<0.0001
Time*Treatment	1	9.57	6.12	0.0385
Error	8	1.56		

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816 Figure 1

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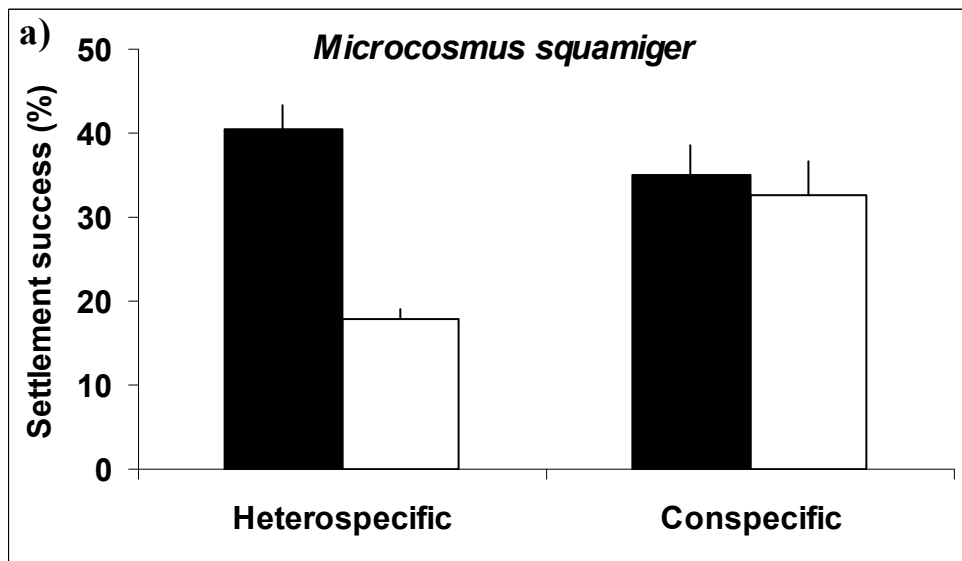
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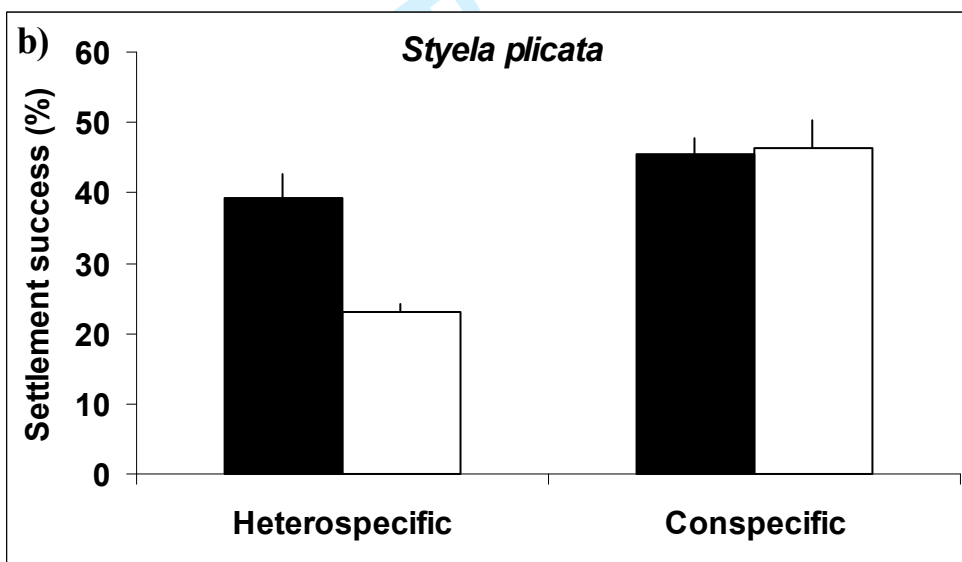
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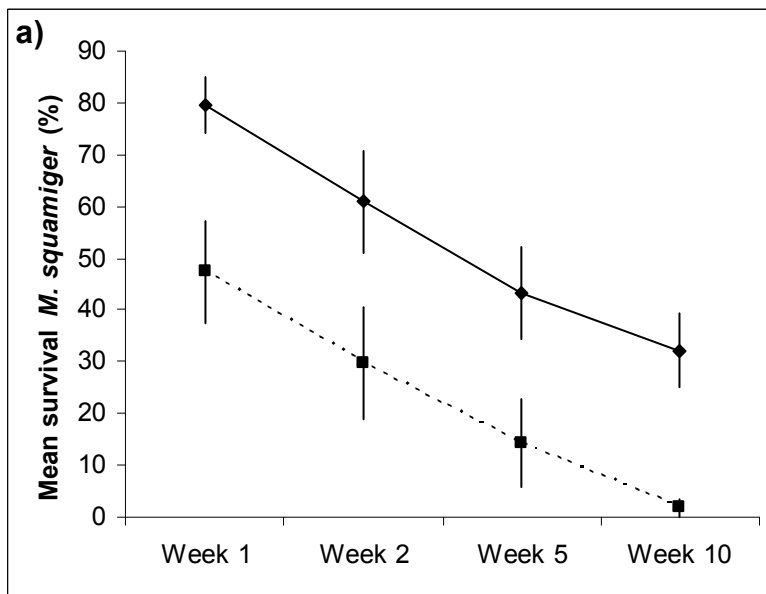
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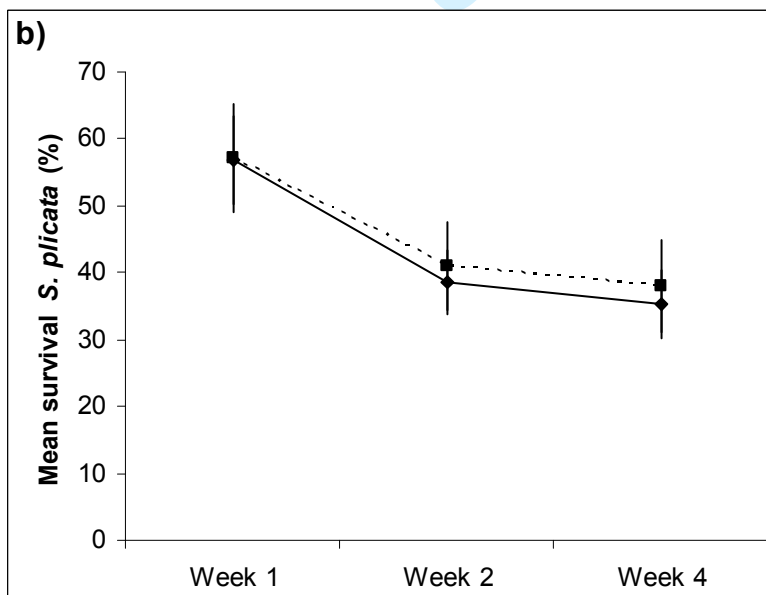
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866 Figure 3

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