FEATURE ARTICLE

Influences of biofilm-associated ciliates on the settlement of marine invertebrate larvae

Jeff Shimeta*, Justin Cutajar, Matthew G. Watson, Thelma Vlamis

School of Applied Sciences, RMIT University, Melbourne, Victoria 3083, Australia

ABSTRACT: Settlement of benthic marine invertebrate larvae often limits recruitment, influencing the structure and dynamics of natural populations as well as biofouling of marine infrastructure, ship hulls, and aquaculture operations. Certain microbial components of substratum biofilms influence settlement (e.g. bacteria, diatoms), but the importance of biofilm protozoa has been unknown. We tested for effects of ciliates by comparing settlement and survival of common fouling invertebrates among 3 biofilm conditions: no biofilm, a purely bacterial biofilm, and a biofilm of bacteria and ciliates. With an assemblage of 7 ciliates (from Hypotrichia, Haptoria, and Scuticociliatia), the serpulid polychaete Galeolaria caespitosa showed a 44 to 49% average reduction in settlement rate compared to the purely bacterial biofilm, and post-settlement mortality increased 7-fold to 34%. In contrast, settlement and survival of the bryozoan Bugula neritina were unaffected. With a partially different assemblage of 11 ciliates (from Hypotrichia, Stichotrichia, Haptoria, Colpodida, and Scuticociliatia), settlement of the serpulid Pomatoceros taeniata more than doubled, whereas that of the blue mussel Mytilus galloprovincialis was reduced by 54% compared to the purely bacterial biofilm. The results could not be explained by ciliates changing the total abundance of biofilm bacteria. We hypothesize that mechanisms could include direct interactions between larvae and ciliates (physical interactions, interference from ciliates’ feeding currents, or responses to chemicals from ciliates), or indirect effects from ciliates altering the bacterial assemblage or its settlement cues. Such large and species-specific effects of ciliates on larval settlement and post-settlement mortality might impact invertebrate recruitment rates and species assemblages, especially because biofilm ciliates are highly variable over time and space.

*Email: jeff.shimeta@rmit.edu.au

KEY WORDS: Larval settlement · Ciliates · Biofilm · Biofouling · Galeolaria caespitosa · Pomatoceros taeniata · Mytilus galloprovincialis · Bugula neritina

INTRODUCTION

Most marine benthic invertebrates have a larval stage that disperses in the water column before settling onto a substratum, metamorphosing, and recruiting into the population. The process of larval settlement can be a limiting step for recruitment, ultimately influencing the structure and dynamics of adult populations (Underwood & Keough 2001). Studies of larval settlement have become key approaches for explaining and predicting patterns and processes in marine assemblages. Particularly
for sessile species, larval settlement can be the primary determinant of adult distributions (e.g. Raimondi 1991). Understanding the constraints on larval settlement in sessile, hard-substratum species is important not only for studies of general ecology but also in efforts to increase yield in aquaculture industries and to prevent biofouling of marine infrastructure and boat hulls (Qian et al. 2007).

Larvae settle in response to a variety of factors that affect them passively or to which they respond behaviorally. Some of these factors are substrate composition and texture (Berrntsson et al. 2000, Qian et al. 2000), hydrodynamic forces (Walters et al. 1997), sediment parameters (Butman 1987), presence of conspecifics (Bryan et al. 1997), presence of a food source or habitat indicator (Daume et al. 2000, Swanson et al. 2006), and microbial biofilms (Qian et al. 2007).

Biofilms are complex 3-dimensional structures composed of microorganisms and a matrix of their extracellular polymers that cover nonliving substrata as well as macrofauna and macroflora (Dobretsov 2010). The development of a biofilm typically follows a succession from adsorption of dissolved chemicals to colonization by prokaryotes and eventually eukaryotes (Wahl 1989, Arndt et al. 2003). When fully developed, the microorganisms in biofilms typically include bacteria, diatoms, thraustochytrids, fungi, and protozoa such as heterotrophic flagellates, amoebae, foraminifers, heliozoans, and ciliates.

Varied and complex interactions between invertebrate larvae and biofilms have been discovered in recent years (reviewed by Qian et al. 2007). Natural biofilms influence larval settlement in numerous species, having either inductive or inhibitory effects. Early studies showed that these effects vary with general biofilm factors such as age, location, or season of development, but often the biofilm organisms responsible for the influences on settlement were not identified (e.g. Todd & Keough 1994, Keough & Raimondi 1995, Wieczorek et al. 1996). More recently, settlement has been shown to relate to the presence of certain taxa of microorganisms, by either correlation or experimental demonstration. These taxa include bacteria (Dobretsov & Qian 2006, Bao et al. 2007, Hung et al. 2007), diatoms (Harder et al. 2002a, Lam et al. 2003), and thraustochytrids (Raghukumar et al. 2000). Some metabolites and extracellular polymers of bacteria and diatoms have been isolated and identified as the cues to which various invertebrate larvae respond (Harder et al. 2002b, Lau et al. 2003, Lam et al. 2005, Patil & Anil 2005, Hung et al. 2009).

Despite increasing recognition that larvae respond to specific microorganisms and cues in biofilms and the fact that protozoa are ubiquitous and influential in biofilms, no published studies have investigated the effects of protozoa on larval settlement. Protozoans in biofilms can have strong impacts on bacterial abundances and spatial heterogeneity, bacterial and microalgal species assemblages, biofilm architecture, and sloughing dynamics (Jackson & Jones 1991, McCormick 1991, Arndt et al. 2003, Parry 2004, Parry et al. 2007). The diverse protozoans in biofilms have complex trophic interactions, seasonal variations, successional dynamics, and constraints from the flow regime and other environmental factors (Franco et al. 1998, Sekar et al. 2002, Arndt et al. 2003, Gong et al. 2005, Wey et al. 2008, Norf et al. 2009, Risse-Buhl & Küsel 2009). Although the literature on biofilm protozoa is considerable in size, it deals almost entirely with freshwater systems, and relatively little is known about the dynamics of protozoans in marine biofilms. Nonetheless, it is conceivable that biofilm protozoa could influence marine larval settlement, either indirectly by affecting other microorganisms or biofilm conditions that serve as settlement cues, or directly by interacting behaviorally or chemically with larvae.

Other than some general acknowledgments that protozoa are part of the typical biofilm community, the literature on larval settlement includes few remarks on the possible roles of protozoa. In studying the attachment of planula larvae of the sea jelly Cyanea capillata, Brewer (1976) noted that his settlement dishes unintentionally contained variable densities of protozoans, which correlated directly with planula attachment rate. However, he speculated that the correlation was spurious in that protozoan abundance probably reflected the abundance of their bacterial prey, and that bacteria were the more likely influence on planula attachment. After observing high post-settlement mortality of didemnid ascidians on settling panels deployed in the field, Todd & Keough (1994) noted that the dead juveniles were infested with ciliates and bacteria, which they speculated might explain the subsequent inhibition of further larval settlement. Wieczorek et al. (1995) found the effects of biofilms to shift from inhibitory to stimulatory for barnacle settlement as the biofilms aged, corresponding to an increase in overall microbial diversity and abundances of protozoans, but they did not attempt to isolate the effects of the protozoans themselves.

This is the first study to test for effects of biofilm protozoa on invertebrate larval settlement. We conducted still-water, no-choice laboratory settlement
assays to compare settlement and survival rates on biofilms in the presence and absence of a mixed-species assemblage of ciliates. We tested a range of common, hard-substratum, fouling invertebrate species from 3 phyla. The serpulid polychaete tube-worms Galeolaria caespitosa and Pomatoceros taeniata have not been studied previously for biofilm cues in larval settlement, although biofilms are known to stimulate settlement in the congener P. lamarckii (Hamer et al. 2001), and the related serpulid Hydrodides elegans is among the most extensively studied marine invertebrate in this regard (e.g. Lam et al. 2005 and references therein). G. caespitosa is a gregarious settler that forms dense reef-like mats on intertidal rocks (Minchinton 1997), whereas populations of P. taeniata consist of sparser, prostrate tubes. The blue mussel Mytilus galloprovincialis (previously considered a subspecies of M. edulis, Gosling 1994) is a common aquaculture species, and its larval settlement is stimulated by biofilms (Bao et al. 2007). The cosmopolitan, arborescent bryozoan Bugula neritina is also known to respond to biofilm cues (Dobretsov & Qian 2006), but it differs from the other species we tested by having a briefer, nonfeeding larval stage that is considered a relatively indiscriminate settler (Dahms et al. 2004).

MATERIALS AND METHODS

Biofilm bacteria and ciliates

Bacteria and ciliates were cultured from natural biofilms that were grown in the subtidal zone at Williamstown, Port Phillip Bay, Australia. Plastic petri-dish microscope slides (Analyslide™, Pall) were deployed for 2 wk on a panel hung under a pier, after which they were capped and returned to the laboratory for immediate processing.

A presumably mixed assemblage of bacteria was isolated from the slides by scraping off the biofilm, dispersing bacteria by sonicating for 5 s (Branson Ultrasonics 200 W at 27% intensity), and passing the suspension through a 0.8 µm pore-size filter to remove debris and larger organisms. These bacteria were then cultured at 4°C in 0.2 µm filtered and autoclaved sea water (FSW) with 0.005% yeast extract.

The abundance of motile, non-stalked ciliates on the biofilms grown in the field was determined by counting cells under a dissecting microscope (352 ± 17 cells cm⁻², n = 10). Surface-associated, motile ciliates were isolated by micropipette into monoclonal cultures and inoculated with the mixed assemblage of biofilm bacteria as a food source. Thus, all cultured ciliates were bacterivorous. Ciliates were cultured at 20°C in FSW with 0.005% yeast extract and rice grains. Ciliates were identified after preserving in Bouin’s fixative and staining with protargol (Montagnes & Lynn 1987), following keys in Lee et al. (2000) and Carey (1992). Seven species were established in culture and used in experiments with Galeolaria caespitosa and Bugula neritina (Table 1). Subsequently, the 2 hypotrich cultures collapsed, and a second field collection yielded 5 more ciliate isolates that were added to the collection for use in experiments with Pomatoceros taeniata and Mytilus galloprovincialis (Table 1).

Invertebrate larvae

Adults of the serpulid polychaete Galeolaria caespitosa were collected from the intertidal zone at Williamstown, Port Phillip Bay. Spawning was stimulated in the laboratory by gently crushing the tubes in FSW. Eggs from 2 to 3 females were combined with a diluted suspension of sperm from 4 to 5 males. After 24 h, the trochophore larvae were rinsed over a 38 µm sieve and transferred to fresh FSW, fed the flagellate Isochrysis sp. (Australian National Algae Culture Collection CS-177, cultured on F/2 medium), at 1 × 10⁴ cells ml⁻¹, and maintained at 20°C on a 15:9 h light:dark cycle. On a daily basis, larvae were rinsed gently over a sieve, transferred to new FSW, and fed. The settlement assay was begun when larvae displayed behavior of settlement competency (i.e. demersal swimming and exploring) at 7 d of age, when they were 200 µm in length. A final rinse with FSW was done over a 63 µm sieve before the settlement assay.

Colonies of the gymnolaemate bryozoan Bugula neritina were collected from settling plates under the pier at Williamstown and held in an aquarium at 20°C in the dark. After 48 h, the colonies were exposed to bright light, which stimulated them to release larvae within 1 h. The nonfeeding, coronate larvae (175 µm) were competent for settlement upon release from the colonies. The larvae were immediately rinsed over a 63 µm sieve with FSW and used in the settlement assay.

Adults of the serpulid polychaete Pomatoceros taeniata were collected from settling plates hung under St. Kilda Pier in Port Phillip Bay. Worms were spawned in several petri dishes of FSW, each containing 2 females and 1 male that released gametes when removed from their tubes. After 15 min, the fer-
tilized eggs were rinsed over a 25 µm mesh sieve to remove sperm and transferred to fresh FSW with gentle aeration. After 24 h, larvae reached the trophophore stage and were fed a mixture of Isochrysis sp. (described above) and Pavlova lutherii (Australian National Algae Culture Collection CS-182, cultured on F/2 medium), at 1.5 × 10^5 cells ml⁻¹. Larvae were maintained at 20°C on a 15:9 h light:dark cycle with gentle aeration and daily feeding and changes of water (using a 63 µm sieve after Day 5). The settlement assay was begun at 14 d of age, when larvae were in the metatrochophore stage (cf. McDougall et al. 2006) and were 250 µm long.

Pediveliger larvae of the mussel Mytilus galloprovincialis were obtained at an age of 22 d from the Victorian Shellfish Hatchery, Dept. of Primary Industries, Queenscliff. The larvae were held at 20°C on a 15:9 h light:dark cycle and fed a mixture of algae obtained from the hatchery (Isochrysis sp., Pavlova lutherii, Chaetoceros muelleri, and C. calcitrans) at 5 × 10^5 cells ml⁻¹. The larvae were rinsed over a 125 µm sieve and transferred to new FSW every second day, and their food was replenished twice daily. The settlement assay was begun at 25 d of age, when the larvae were 350 µm long.

**Larval settlement assays**

Settlement assays were run separately for each invertebrate species. Assays were run in 55 mm diameter polystyrene petri dishes with 3 treatments and 8 replicate dishes of each: (1) untreated (i.e. no biofilm), (2) a thin biofilm of bacteria in a monolayer on the dish, and (3) the same monolayer bacterial biofilm plus a mixed assemblage of ciliates. Comparison of larval settlement between the bacterial biofilm treatment and the same biofilm plus ciliates revealed whether there was an effect of ciliates on settlement. Comparison of settlement in the absence of a biofilm and in the presence of the purely bacterial biofilm revealed any influence of the bacteria on settlement. This latter comparison was included in the design to evaluate whether any decline of bacterial density during the assays caused by ciliate grazing could explain observed effects on settlement. We did not include a treatment of ciliates without a bacterial biofilm because many of these surface-associated bacterivorous ciliates (Table 1) would require substratum bacteria in order to engage in normal behaviors of movement and foraging. Therefore, bacteria were included to induce normal ciliate behavior and were not intended to represent a natural bacterial biofilm; indeed, they comprised only cultivable bacteria rather than a complete assemblage of all bacteria from the field.

The monolayer bacterial biofilm was grown by incubating the bottom halves of the petri dishes for 24 h at 20°C in an aerated bath of FSW with 0.001% yeast extract and 0.001% bacto-peptone, inoculated with the mixed assemblage of biofilm bacteria isolated from the field. These biofilms developed consistently with an abundance of (2 to 3) × 10^4 cells cm⁻².

### Table 1. Ciliates isolated into pure culture from field-grown biofilms and used in larval settlement assays. Classification follows Lee et al. (2000). Two stichotrichians could not be identified further

<table>
<thead>
<tr>
<th>Subclass or Order</th>
<th>Species</th>
<th>Cell size (µm)</th>
<th>Used in experiment with:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Galeolaria caespitosa</td>
</tr>
<tr>
<td>Hypotrichia</td>
<td>Diophrys oligothrix</td>
<td>150</td>
<td>•</td>
</tr>
<tr>
<td></td>
<td>Euplotes arenularum</td>
<td>50</td>
<td>•</td>
</tr>
<tr>
<td></td>
<td>Euplotes elegans</td>
<td>60</td>
<td>•</td>
</tr>
<tr>
<td></td>
<td>Euplotes minuta</td>
<td>50</td>
<td>•</td>
</tr>
<tr>
<td>Stichotrichia</td>
<td>Amphisiella sp.</td>
<td>50</td>
<td>•</td>
</tr>
<tr>
<td></td>
<td>Unidentified spp. (2)</td>
<td>60</td>
<td>•</td>
</tr>
<tr>
<td>Haptoria</td>
<td>Litonotus sp.</td>
<td>105</td>
<td>•</td>
</tr>
<tr>
<td>Colpodida</td>
<td>Colpodia sp.</td>
<td>35</td>
<td>•</td>
</tr>
<tr>
<td>Scuticociliatia</td>
<td>Cyclidium sp.</td>
<td>40</td>
<td>•</td>
</tr>
<tr>
<td></td>
<td>Protocruzia sp.</td>
<td>40</td>
<td>•</td>
</tr>
<tr>
<td></td>
<td>Pseudocohnilembus sp.</td>
<td>35</td>
<td>•</td>
</tr>
<tr>
<td></td>
<td>Uronema sp.</td>
<td>40</td>
<td>•</td>
</tr>
</tbody>
</table>
as measured by epifluorescence microscopy (see below). After growing the biofilms, all petri dishes (as well as untreated dishes) were gently rinsed and filled with FSW. A mixture of cultured ciliates (Table 1) was added to the ciliate treatment dishes to yield 350 cells cm$^{-2}$ on the bottom (matching the abundance measured on the 2 wk old biofilms collected from the field). The mixture included equal parts from each ciliate culture, although the cultures differed in their cell abundances. The total volume of liquid in each dish was 12 ml.

Settlement assays were begun by adding 20 larvae to each dish of every treatment immediately after adding ciliates to the ciliate treatment dishes. The dishes were randomly intermixed in an array on a benchtop during the assay. Each assay was run at the same temperature and light:dark cycle at which the larvae were held prior to the experiment. Dishes with 

\textit{Galeolaria caespitosa} and \textit{Pomatioceros taeniata} were placed on white paper to stimulate settlement, but dishes with \textit{Mytilus galloprovincialis} were not. Dishes with \textit{Bugula neritina} were kept in the dark. At 2 time points, all dishes were examined under a dissecting microscope, and larvae were scored as either settled, not settled, or dead. Larvae of \textit{G. caespitosa}, \textit{P. taeniata}, and \textit{B. neritina} were scored as settled if an attachment to the dish was evident. Larvae of \textit{M. galloprovincialis} were considered settled if they were motionless on the bottom, no velum was visible through the shell, and they did not move when the dish was gently agitated. The total length of the assays varied according to the rapidity with which each species settled. The time points for examination were 48 h and 72 h for \textit{G. caespitosa} and \textit{P. taeniata}, 1 h and 24 h for \textit{B. neritina}, and 24 h and 48 h for \textit{M. galloprovincialis}. After the second time point, the abundance of ciliates on the bottoms of the dishes was determined by counting them under a dissecting microscope. The contents of all dishes were then gently decanted and replaced with 1% formalin in FSW to preserve the biofilm bacteria.

Bacterial densities on the bottoms of the petri dishes were determined by removing the formalin, adding 50 µl of DAPI solution (0.02 mg ml$^{-1}$) and a 22 × 40 mm coverglass, and counting cells at 1000× magnification using oil immersion on an epifluorescence microscope with UV excitation (340–380 nm).

Settlement rates and mortality rates (both as arcsine square-root transformed proportions) were tested at each time point by an \textit{a priori} test of main effects (for the effect of the treatments, i.e. the between-subjects factor) using the MSE from a repeated-measures analysis of variance (ANOVA). If the effect of the treatment was significant at $\alpha = 0.05$, Tukey pairwise comparisons were run among the treatments at that time point using the MSE from the test of main effects. The biofilm bacterial densities at the end of each assay were tested for differences among the 3 treatments by 1-way ANOVA, followed by Tukey pairwise comparisons. All statistical tests were run on Systat v5.2.1 for Macintosh.

\section*{RESULTS}

\textit{Galeolaria caespitosa}

After 48 h, the proportion of larvae that had settled and survived differed significantly among the treatments ($p < 0.01$; Fig. 1A). Settlement was enhanced in the presence of a pure bacterial biofilm compared to the untreated dishes ($p < 0.01$). Settlement in the presence of ciliates and bacteria was 49% lower than in the pure bacterial biofilm ($p = 0.041$) and not different from the untreated dishes ($p = 0.64$). There was ca. 5% post-settlement mortality (proportion of settled larvae that had died), but it did not differ among treatments ($p = 0.88$; Fig. 1A). The total proportion of settled larvae (including live and dead) therefore had a similar pattern to the settled live larvae ($p = 0.013$ for the main effect), although the 48% decline in the presence of ciliates compared to the pure bacterial biofilm was only marginally close to significant at $p = 0.081$ (Fig. 1A).

After 72 h, the settled live larvae showed a similar enhancement in the pure bacterial biofilm compared to the untreated dishes ($p < 0.001$ for both the main effect and the comparison; Fig. 1B). There was also a 44% reduction of live settlers in the biofilm with ciliates compared to the pure bacterial biofilm ($p < 0.001$). Post-settlement mortality in the treatment with ciliates, however, increased dramatically to 34%, whereas it remained at ca. 5% in the other treatments ($p < 0.01$ for the main effect). Thus, there was a 7-fold increase in mortality rate associated with the presence of ciliates. As a result, the total proportion of settlers (live and dead) differed among the treatments ($p < 0.01$): it was similar in the 2 biofilm treatments (with and without ciliates, $p = 0.41$), and approximately twice the settlement in the untreated dishes ($p < 0.01$ for the pure bacterial biofilm and $p = 0.042$ for the bacteria+ciliates treatment).

At the end of the assay (72 h), there was no difference in the abundance of biofilm bacteria between
the bacteria treatment and the bacteria+ciliates treatment (p = 0.80; Table 2). The untreated dishes had some bacterial growth on the bottom, but it was only 28% of the abundance in the 2 biofilm treatments (p < 0.001 in both cases). The bacteria+ciliate dishes had 378 ± 30 ciliates cm$^{-2}$, whereas no ciliates were detected in the other treatments.

**Bugula neritina**

Many of the *Bugula neritina* larvae settled on the sides of the petri dishes. Because the dishes had been fully submerged in the bacterial bath for growing the biofilms, we assume that bacteria grew on the sides as well as on the bottoms. We examined the sides of the dishes for ciliates in the bacteria+ciliates treatment, finding ciliates to be in similar abundance on the side and the bottom.

After 1 h, settlement was significantly lower in the 2 biofilm treatments (with and without ciliates) compared to the untreated dishes (p < 0.001 for the main effect and for both comparisons; Fig. 2A). There was no difference between the 2 biofilm treatments (p = 0.14), i.e. the presence of ciliates had no significant effect on settlement. After 24 h, the pure bacterial biofilm still had a significantly lower settlement than the untreated dishes (p < 0.01 for both the main effect and the comparison), but no other comparisons were statistically significant (p > 0.05). Thus, the treatment with ciliates was not different from any other treatments. There was no mortality in the assay.

At the end of the assay (24 h), there was no difference in the abundance of biofilm bacteria between the bacteria treatment and the bacteria+ciliates treatment (p = 0.34; Table 2). The abundance of bacteria in the untreated dishes was an order of magnitude lower than that in the 2 biofilm treatments (p < 0.001 in both comparisons). The bacteria+ciliate dishes had 342 ± 30 ciliates cm$^{-2}$, whereas no ciliates were detected in the other treatments.

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**Fig. 1.** *Galeolaria caespitosa*. Settlement assay with larvae. 'Settled live' is the proportion of attached, live larvae among the total number of larvae added to the dish. 'Settled total' is the analogous proportion including all attached larvae, both live and dead. 'Mortality' is the proportion of attached, dead larvae among the total number of attached larvae (hence, post-settlement mortality). Bars show mean ± SE, and shading indicates treatments of the petri dishes ('Untreated' had no biofilm; 'Bacteria' had only a bacterial biofilm; 'Bacteria + Ciliates' had the same bacterial biofilm plus ciliates). Within each category on the abscissa, means with different letters are significantly different at $\alpha = 0.05$. (A) 48 h tests of main effects (for the treatments); settled live p = 0.0044, settled total p = 0.013, mortality p = 0.88. (B) 72 h tests of main effects: settled live p < 0.001, settled total p = 0.0022, mortality p = 0.0011

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**Table 2.** Biofilm bacterial densities (mean ± SE cells cm$^{-2}$) at the end of each larval settlement assay. 'Untreated' petri dishes were not subjected to biofilm growth prior to the assay. ‘Bacteria’ and ‘Bacteria + Ciliates’ petri dishes were incubated in a bacterial culture prior to the assay. Only ‘Bacteria + Ciliates’ petri dishes had ciliates added at the start of the assay. Means with different superscripts were significantly different by post hoc multiple comparison tests following a significant 1-way analysis of variance ($\alpha = 0.05$)

<table>
<thead>
<tr>
<th>Larval assay</th>
<th>Untreated (×10$^3$)</th>
<th>Bacteria (×10$^3$)</th>
<th>Bacteria + Ciliates (×10$^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Galeolaria caespitosa</em></td>
<td>7.82 ± 0.61$^{a}$</td>
<td>2.91 ± 0.20$^{b}$</td>
<td>2.62 ± 0.21$^{b}$</td>
</tr>
<tr>
<td><em>Bugula neritina</em></td>
<td>1.92 ± 0.41$^{a}$</td>
<td>1.48 ± 0.09$^{b}$</td>
<td>1.63 ± 0.08$^{b}$</td>
</tr>
<tr>
<td><em>Pomatoceros taeina</em></td>
<td>6.97 ± 0.56$^{a}$</td>
<td>2.52 ± 0.15$^{b}$</td>
<td>1.89 ± 0.15$^{c}$</td>
</tr>
<tr>
<td><em>Mytilus galloprovincialis</em></td>
<td>2.94 ± 0.20$^{a}$</td>
<td>2.19 ± 0.10$^{b}$</td>
<td>1.96 ± 0.12$^{b}$</td>
</tr>
</tbody>
</table>
At both time points, the proportion of larvae that had settled was significantly different among the treatments (p < 0.001; Fig. 2B). There was negligible mortality at 72 h, and it did not differ among treatments (p = 0.61, data not shown). At 48 h, settlement on the pure bacterial biofilm was significantly greater than on the untreated dishes (p = 0.034), but at 72 h they were not significantly different (p = 0.26). At both time points, the presence of ciliates increased settlement significantly compared to the pure bacterial biofilm (p = 0.038 at 48 h and p < 0.001 at 72 h), by an average of 2.4-fold.

By the end of the assay, the abundance of biofilm bacteria was 25% lower in the presence of ciliates than in the pure bacterial biofilm treatment (p < 0.01; Table 2), presumably as a result of ciliate grazing. There was slight growth of bacteria in the untreated dishes, but it was significantly less than the bacteria in the other treatments (p < 0.001 in both cases). The bacteria+ciliate dishes had 356 ± 29 ciliates cm\(^{-2}\), whereas no ciliates were detected in the other treatments.

**Mytilus galloprovincialis**

At both time points, the proportion of settled larvae was significantly different among all treatments (p < 0.001), showing the same pattern in each case (Fig. 2C). The presence of a pure bacterial biofilm enhanced settlement by an average of 9.8-fold compared to the untreated dishes (p < 0.001 at both time points). Settlement in the presence of ciliates was reduced to 46% of settlement in the pure bacterial biofilm (p < 0.001 at both time points). There was no mortality in any of the treatments.

At the end of the assay (48 h), there was no difference in the abundance of biofilm bacteria between the bacteria treatment and the bacteria+ciliates treatment (p = 0.19; Table 2). The slight growth of bacteria in the untreated dishes was an order of magnitude less than in the 2 biofilm treatments (p < 0.001 in both cases). The bacteria+ciliate dishes had 432 ± 28 ciliates cm\(^{-2}\), whereas no ciliates were detected in the other treatments.

**DISCUSSION**

**Effects of ciliates on larval settlement**

We observed a variety of responses among invertebrate species to the presence of ciliates in the biofilm, including inhibition of settlement, facilitation of settlement, enhanced post-settlement mortality, and no response. There are several possible mechanisms that could be involved in these effects. Here we present hypotheses that require further investigation to substantiate.
A possible mechanism for the 50% reductions of settlement rate in *Galeolaria caespitosa* and *Mytilus galloprovincialis* is direct interference from ciliates near the substratum, e.g. through behavioral interactions or if the ciliates' feeding currents interrupt the swimming or crawling of larvae as they test the substratum. The ciliates in these experiments (Table 1) included strongly thigmotactic species that crawl and cling tightly to the substratum (e.g. hypotrichs such as *Diophrys oligotheix* and *Euplotes* spp.) and epibenthic species that glide on the substratum and periodically swim above it (e.g. scuticociliates such as *Cyclidium* sp. and *Uronema* sp.; Patterson et al. 1989). These ciliates graze on attached bacteria and/or suspension feed. The largest ciliates in our experiments were up to 30% to 75% of the size of larval *G. caespitosa* and *M. galloprovincialis*, which might allow the ciliates' movements to interfere with larval feeding currents of these types of ciliates allow the ciliates' movements to interfere with larval swimming or crawling as they test the substratum. The ciliates in these experiments (Table 1) included strongly thigmotactic species that crawl and cling tightly to the substratum (e.g. hypotrichs such as *Diophrys oligotheix* and *Euplotes* spp.) and epibenthic species that glide on the substratum and periodically swim above it (e.g. scuticociliates such as *Cyclidium* sp. and *Uronema* sp.; Patterson et al. 1989). These ciliates graze on attached bacteria and/or suspension feed. The largest ciliates in our experiments were up to 30% to 75% of the size of larval *G. caespitosa* and *M. galloprovincialis*, which might allow the ciliates' movements to interfere with larvae. The feeding currents of these types of ciliates can be up to 0.2 mm s⁻¹ in the vicinity of the cells and 1 mm s⁻¹ near their cilia (Fenchel 1986, Shimeta et al. 2001), which might disrupt slowly swimming pediveligers moving at speeds of 0.6 mm s⁻¹ (Cragg 1980). Polychaete larvae are less likely to be disrupted because of their faster swimming speeds, e.g. 4 mm s⁻¹ in metatrochophores of *G. caespitosa* (Bolton & Havenhand 1997). Once larvae are crawling on the substratum, they could be more easily affected by ciliate movements and currents. Another possibility for a direct interaction is the release of a chemical cue by ciliates that deters settlement, similar to negative cues released by some bacteria and algae (Lau & Qian 1997, Rao et al. 2007).

Another possible mechanism for the settlement inhibition in *Galeolaria caespitosa* and *Mytilus galloprovincialis* is indirect via an influence of ciliates on biofilm bacteria, because the pure bacterial biofilm did stimulate settlement in these invertebrates compared to the untreated dishes. Ciliates did not change the total bacterial abundance, however, so the settlement inhibition cannot be explained as a response to total bacterial density. It is possible that grazing altered the abundances of certain bacterial species that serve as settlement cues, either enhancing species that are negative cues or reducing species that are positive cues. We cannot evaluate this hypothesis from our results, however, because we did not characterize the bacterial assemblage and therefore do not know if it shifted in response to the presence of ciliates. It is well known that some bacteria serve as settlement inducers while others are inhibitors (Qian et al. 2007), and that settlement responses of invertebrates can vary depending on the bacterial species assemblage in a biofilm (Lau et al. 2005). Protozoans have large grazing impacts on biofilm bacterial abundances and biofilm structure, although this is primarily known for freshwater systems rather than marine (Arndt et al. 2003, Parry 2004), and there is little information on grazing effects on the bacterial species assemblages in biofilms. Another possible indirect effect from protozoan grazing could be that larval settlement is inhibited by bacterial metabolites released as a chemical defense against grazing (Matz et al. 2008). Finally, chemicals released by ciliates might cause changes in the bacterial species assemblage, similar to algal metabolites that have been shown to influence the bacterial community on their surfaces (Steinberg et al. 2002). We note that the mechanism of settlement inhibition was not necessarily the same for *G. caespitosa* and *M. galloprovincialis*, particularly because these species were exposed to partially differing assemblages of ciliates.

The second type of negative effect from ciliates on larval settlement was post-settlement mortality in *Galeolaria caespitosa*, which affected 30% of the settlers. Possible mechanisms include a direct physical interference from ciliates or a noxious chemical released by ciliates or bacteria in the biofilm. We observed dense aggregations of ciliates around dead larvae, similar to Todd & Keough’s (1994) observations around dead ascidian settlers, which could reflect a direct interference from ciliates that killed the settled larvae. We do not know, however, whether these ciliates caused the mortality or were attracted to the carcasses afterward. Early post-settlement mortality can be a large and important factor determining recruitment success in marine invertebrates (Hunt & Scheibling 1997). It has a variety of known causes, but this is the first demonstration of mortality associated with the presence of ciliates.

The positive influence of ciliates on settlement rate in *Pomatoceros taeniata* (more than a doubling) could have been caused by a stimulatory cue, e.g. larvae responding to the presence of ciliates as a food source or to a stimulatory chemical released by ciliates. We observed numerous ciliates crawling over the tubes of newly settled *P. taeniata*, which might reflect a facilitative relationship. In this assay, the ciliates reduced the bacterial abundance presumably by grazing, but since the bacteria had a stimulatory effect on settlement (at one of the 2 time points), a loss of total bacterial abundance by grazing cannot account for the increased settlement rate. We cannot exclude, however, that grazing altered the bacterial assemblage, which in turn stimulated settlement.
We caution against concluding from our results that the 2 serpulid polychaetes (Galeolaria caespitosa and Pomatoceros taeniata) have opposite responses to ciliates in general, because they were tested using partially differing assemblages of ciliates. *P. taeniata* and *Mytilus galloprovincialis*, however, were tested with the same assemblage of ciliates and showed opposite responses (facilitation and inhibition, respectively). These results could reflect differential responses to one or more ciliate species, but we cannot exclude that any differences in the bacterial assemblage between these assays could have induced different larval responses through indirect effects of ciliates on settlement cues from bacteria.

We found no evidence for an effect of ciliates on the settlement or survival of the bryozoan *Bugula neritina*. Although bryozoan larvae do respond to settlement cues from biofilms, they are considered to be relatively indiscriminate settlers compared to other invertebrates, attributed to their short planktonic phase (Dahms et al. 2004, Dobretsov & Qian 2006). Unwettable surfaces such as polystyrene dishes are known to stimulate settlement of *B. neritina* larvae (Mihm et al. 1981), which could limit the potential to detect stimulatory cues from ciliates if they exist. However, the significant reduction in settlement caused by the presence of a bacterial biofilm in our experiments would have at least partially removed such a masking effect, and the addition of ciliates to the bacterial biofilm showed no further stimulation of settlement. Taking into account these considerations, the most conservative conclusion is that ciliates caused no inhibitory effects on settlement or survival in *B. neritina*.

**Implications**

Effects of ciliates on larval settlement and survival appear to be highly varied and species specific. Having observed them in 3 of 4 species tested, including mussels and polychaetes, they may be widespread among taxa. Because several mechanisms could be involved, many variables might mediate these interactions in the field to determine their impact on invertebrate population dynamics.

Biofilms in the field are more complex than those we created for our experiments, which included only cultivable strains of bacteria grown in a sparse monolayer. Natural biofilms have more bacterial diversity as well as microalgae and other types of protozoa, and complex 3-dimensional architecture associated with the matrix of bacterial extracellular polymers and microcolonies of cells (Arndt et al. 2003, Dobretsov 2010). Therefore, if any influences of ciliates on larval settlement in our experiments were through an indirect effect on bacteria, further investigations would be needed to determine how these interactions function in a more complex, natural microbial biofilm community.

The nature of the ciliate fauna in a biofilm could also play an important role in the effects on settlement. We used a ciliate assemblage and total density from 2 wk old biofilms from the field; however, the assemblage composition and density are likely to be different in younger or older biofilms. Larvae might respond to certain ciliate species more than others, due to differences in ciliate behavior, grazing impacts, etc. Indeed, the settlement responses we measured might be caused by only certain ciliates in the assemblages we used. Furthermore, these assemblages represented only a subset of ciliates that grew on our settlement slides in the field, and there may be other types that influence settlement differently. We only isolated mobile, easily cultured, bacterivorous species, not herbivores, predators, or attached species such as the stalked peritrichs or suctorians which create strong feeding currents. Ciliates with these different trophic modes could have other effects on the microbial biofilm community and larval settlement, e.g. impacting microalgae that serve as settling cues (e.g. Harder et al. 2002a) or altering the boundary-layer flow field (Fenchel 1986). Any flow-mediated interactions between ciliates and larvae might be influenced by the surrounding flow field in nature, which should be considered in future investigations, whereas our experiments were done in still water.

Elucidating the mechanisms by which ciliates influence larval settlement may help to predict settlement patterns in the field and to interpret studies on settlement responses to biofilms. Although ciliates are ubiquitous in marine biofilms, their densities and species assemblages vary seasonally as well as during succession after space is exposed by a disturbance or when a new substratum is introduced to the environment (Anderson 1995, Arndt et al. 2003, Gong et al. 2005). Their species assemblages correlate with environmental parameters (e.g. temperature, salinity, and nutrients; Gong et al. 2005) and therefore are likely to vary spatially among locations or on substrata of different materials (Coppellotti & Matarazzo 2000), although these aspects of their distribution have received little attention in marine habitats. Based on our results, we hypothesize that recruitment of certain invertebrates onto an available
patch of substratum can depend on the ciliate fauna present at that point in the succession of the biofilm community. This phenomenon might explain some of the variations of recruitment onto settlement panels in the field in previous studies that neglected to examine protozoa in the biofilms. For example, settlement of various invertebrates has been found to differ among natural biofilms of different ages (Keough & Raimondi 1995, Bao et al. 2007), grown in different seasons (Wieczorek et al. 1996, Bao et al. 2007), and grown in different locations (Dobretsov & Qian 2006, Hung et al. 2007). In these studies and others, such effects have been attributed to assumed differences in the general microbial film community or specifically to measured differences in bacteria, but the ciliate assemblages might also have played important roles.

Furthermore, because the settlement responses to ciliates in our experiments differed among invertebrate species, these interactions could play a role in shaping the species assemblage of invertebrates at a site. For example, settlement of Pomatoceros taeniata doubled, whereas that of Mytilus galloprovincialis was halved when exposed to the same assemblage of ciliates (although the bacterial assemblage in the biofilm was not necessarily identical). Based on these results, we hypothesize that a substratum with these ciliates should favor colonization by P. taeniata rather than M. galloprovincialis.

Effects of ciliates on settlement rates also have implications for aquaculture and for efforts to reduce invertebrate fouling of marine infrastructure, because of the known importance of biofilms in these systems (Qian et al. 2007). For example, the reduced settlement of Mytilus galloprovincialis suggests that elimination of ciliates from aquaculture ropes could improve settlement rates of cultivated mussels. If an aquaculture species is stimulated to settle by ciliates, seeding the substratum with ciliates might enhance yield. In the field of anti fouling research, efforts are increasingly directed toward harnessing natural antifoulant chemicals (Qian et al. 2010), where further research on the mechanisms of settlement inhibition by ciliates might have application.

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