Comparison of surface microfouling and bacterial attachment on the egg capsules of two molluscan species representing Cephalopoda and Neogastropoda

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ABSTRACT: Many organisms naturally defend themselves against microbial attachment and biofouling in the marine environment. In this study, we investigated microbial fouling on 2 molluscan egg capsules using scanning electron microscopy (SEM), two-photon laser scanning microscopy (TPLSM) with bacterial viability staining and bacterial attachment experiments with the biofilm-forming Pseudoalteromonas sp. S91 in flow chambers. Results indicated that early stage egg capsules of Dicathais orbita (Neogastropoda) are relatively free of surface microorganisms. Egg capsules during the trocophore stage had a regularly ridged microtexture, but as capsules matured, shedding of the outer wall was observed, followed by the extrusion of unidentified droplets, which then accumulated on the capsule surface in association with bacteria. By comparison, the egg capsules of Sepioteuthis australis (Cephalopoda) were found to have an irregular surface with many hills and valleys that accommodate colonization by a variety of microorganisms. At the later stages of development these squid egg capsules become heavily colonized by algal spores. Cross sections of egg capsules revealed that S. australis capsule walls were about 12 times thicker than D. orbita egg capsules. Staining the egg capsules with BacLight™ also revealed a significantly thicker biofilm, with more live and dead bacteria on S. australis capsules than on those of D. orbita (p < 0.05). Flow chamber experiments indicated that the surface of S. australis capsules provided a suitable substrate for colonization by Pseudoalteromonas sp. S91, whereas colonization was significantly less on D. orbita egg capsules after 24 and 72 h (p < 0.01). These experiments indicated that D. orbita egg capsules are better defended against fouling microbes than are the eggs of S. australis. D. orbita appears to use a combination of physical, mechanical and possibly chemical defense mechanisms to reduce fouling on their egg capsules.

KEY WORDS: Biofilm · Egg capsules · Mollusc · Scanning electron microscopy · Two-photon laser scanning microscopy · Bacterial attachment

INTRODUCTION

The economic costs of biofouling on artificial structures, such as boat hulls, nets and pipes, can be enormous. This has resulted in the use of toxic paints that have led to serious environmental problems (e.g. tributyltin, TBT; Alzieu 2000, Champ 2000, Goldberg et al. 2004). Biofilm formation also plays an important role in bacterial pathogenesis. Biofilms are typically resistant to antibiotics, providing a source for persistent infection (Hentzer & Givskov 2003) and/or re-infection from medical tubing and equipment. Understanding the mechanisms by which aquatic organisms naturally regulate fouling of their surfaces should provide a platform for ongoing developments in the control of economically and ecologically important fouling processes (de Nys & Steinburg 2002).

Surface fouling by microorganisms is one of the many threats that face organisms in the marine environment (Davis et al. 1989, Scardino et al. 2003). With
an estimated density of $5 \times 10^5$ prokaryotic cells ml$^{-1}$ seawater (Whitman et al. 1998), sessile invertebrates and algae are exposed to a constant onslaught of potentially detrimental microbes. These include biofilm-forming bacteria along with single-cell diatoms that rapidly settle, attach and form colonies on any surface placed in the marine environment (Davis et al. 1989). The formation of a microbial biofilm promotes the attachment of algal spores, protozoa, barnacle cyprids and marine fungi, followed by the settlement of other marine invertebrate larvae and macroalgae (Maki 2000, Callow & Callow 2002). Heavy surface fouling could lead to the accumulation of toxic wastes, a reduction in oxygen and nutrient availability and increased drag, which can cause sessile organisms to become dislodged from benthic substrata in strong currents. All extant marine organisms have survived for millions of years under the significant selective pressure of surface fouling and therefore must have evolved successful mechanisms for reducing these negative impacts.

The encapsulation of developing offspring within elaborate structures that are deposited on benthic substrata is a common reproductive strategy among marine molluscs. These egg capsules appear to be highly resilient multilaminate biomaterials (Rapoport & Shadwick 2002) that require substantial maternal energy investment (Stickle 1973). This implies that egg capsules must provide adaptive benefits that increase embryonic survivorship. The functional role of encapsulation is widely attributed to the protection of developing offspring from environmental stresses, as well as to deter potential predators and pathogens (Thorson 1950, Pechenik 1986, Rawlings 1999, Przeslawski 2004). However, these egg capsules can remain in the marine environment for several months, and thus would also be vulnerable to surface fouling. Nevertheless, previous studies indicate that these egg capsules remain axenic (Lord 1986) and remarkably free of surface macrofouling (Przeslawski & Benkendorff 2005). Antimicrobial activity also appears to be widespread in chemical extracts from molluscan egg masses (Benkendorff et al. 2001, Santhana Ramasamy & Murugan 2005), but it is presently uncertain if the active compounds are located on the eggs’ surface. Interestingly, a previous study on the egg cases of the dogfish Scyliorhinus canicula demonstrated that these marine egg capsules are resistant to microbial degradation and performed much better than many commercial anti-fouling treatments in the field (Thomason et al. 1994). Mechanisms for deterring fouling on these extracellular structures are not currently understood.

The functional mechanisms for preventing detrimental microbial colonization on marine organisms can include associated chemical, mechanical and/or physical defenses. Chemical extracts from a wide variety of marine invertebrates and algae have demonstrated antimicrobial activity against marine biofilm bacteria (Steinberg et al. 2001, Engel et al. 2002, Lindquist 2002, Santhana Ramasamy & Murugan 2005) and some secondary metabolites have been shown to regulate biofilm bacteria on biotic surfaces. For example, furanones produced by the red alga Delisea pulchra actively prevent bacterial and larval settlement (Dworjanyn et al. 1999, Steinberg et al. 2001). Secondary metabolites produced by sponges in the Caribbean were also shown to prevent bacterial surface colonization (Kelly et al. 2005). Mechanical methods of fouling prevention are adopted by the macroalga Dilesia carnosa, which sheds its epidermis and cuticle to remove any epibionts (Nylund & Pavia 2005). Similarly, moulting in crustaceans would facilitate the removal of epibionts (Dyrynda 1986). Relatively little is known of the role of physical defenses (Steinberg et al. 2001), though surface composition and microtexture can influence the rate of biofouling (Berntsson et al. 2000, Steinberg & De Nys 2002, Baum et al. 2003).

Biofilms cultivated on inanimate and biological surfaces under laboratory conditions have been studied extensively using a wide range of microscopic techniques (Beech et al. 2000), often in combination, to give a clear description of the biofilm. Scanning electron microscopy (SEM) allows morphological examination of biofilms established on surfaces (Beech et al. 2000). Bers & Wahl (2004) used SEM to study the surface properties of different marine organisms, such as the edible crab Cancer pagurus (Crustacea), the blue mussel Mytilus edulis (Bivalvia), the brittle star Ophinura texturata (Ophiuroidea) and the egg case of the lesser-spotted dogfish Scyliorhinus canicula (Chondrichthyes), which are all seldom associated with epibionts. These studies support the evolution of physical anti-fouling strategies, showing that homogeneous surfaces are capable of deterring attachment by limiting space available for fouling organisms.

Flow chambers such as those designed by Hale & Mitchell (2001) can also be used to study biofilm formation. Bacterial attachment to biological surfaces can be studied using Pseudoalteromonas sp. S91, a motile Gram-negative marine bacterium that secretes hydrolytic enzymes, e.g. chitinases and proteases (Techkarnjanaruks & Goodman 1999), and can utilize squid pen as a nutrient source (Techkarnjanaruks et al. 1997). The derivative strain, S91SB, has a transposon insert containing a copy of the green fluorescent protein (GFP) gene originated from the jellyfish Aequorea victoria (Chalfie et al. 1994). The use of GFP in prokaryotes has assisted the study of spatial distribution of microbial cells within a biofilm (Yoshida &
Kuramitsu 2002). In addition, the use of GFP in conjunction with two-photon laser scanning microscopy (TPLSM) can facilitate the real-time analysis of cells within a living biofilm (Stretton et al. 1998).

This study examines microbial fouling on the surface of egg capsules from 2 molluscan species: the white dogwhelk *Dicathais orbita* (Neogastropoda: Muricidae) and the southern calamari *Sepioteuthis australis* (Cephalopoda: Loliginidae) using a combination of microscopic techniques. The recent study by Przeslawski & Benkendorff (2005) on the egg masses of 18 gastropod molluscs revealed that the leathery egg capsules of neogastropods, including *D. orbita*, were significantly less fouled than a range of gelatinous egg masses. However, in Przeslawski & Benkendorff’s (2005) study, protists and algal fouling were only estimated visually under a dissecting microscope. Similarly, Steer et al. (2002) used visual observations to report high levels of electron microscopy observed at this stage—see ‘Results; Scanning electron microscopy’), and fresh squid egg capsules with undifferentiated embryos were used because these are much closer in age to the *D. orbita* trocophores than are the paralarvae.

**Scanning electron microscopy.** For each species, 5 independent egg capsules at each stage of embryonic development were fixed overnight in electron microscopy fixative (1.25% glutaraldehyde, 4% sucrose, 4% paraformaldehyde in phosphate buffered saline [PBS], pH 7.2). After 24 h, the egg capsules were removed from fixative, washed twice with PBS and cut into 0.5 cm sections. Samples were then fixed in 2% osmium tetroxide for 60 min before dehydration in increasing concentrations of ethanol (70, 90, 95 and 100% for 30 min each). Finally, the samples were dried under pressure with liquid carbon dioxide in a BalTec CPD030 Critical Point Dryer. The dried samples were placed on metal stubs using adhesive stickers and sputter coated with platinum. Egg capsules were orientated to allow morphological examination and analysis of surface bacterial load. Cross-sectional views of egg capsules were also examined. The samples were viewed using a Philips XL-30 Field Emission SEM.

Cryogenic SEM was also carried out using an Oxford Instruments CT1500 HF Cryo Preparation System attached to the SEM. Egg capsules were attached to the holder using Tissue-Tek OCT compound, then frozen in nitrogen slush, and transferred under vacuum to the preparation chamber. The temperature of the sample was raised to –92°C and held there for approximately 3 min to allow any ice on the surface to sublime. After lowering the temperature to –110°C to halt sublimation, the sample was coated with platinum, loaded onto the microscope stage and examined whilst being maintained at –150°C.

The percentage of fouling on the surface of egg capsules was determined using images taken from SEM analysis. For each egg capsule, 5 SEM images were...
examined at approximately 5000× magnification, and the percentage cover per image was estimated visually and then averaged to obtain 1 mean value per capsule. A 2-factor ANOVA was performed using the statistical package SPSS version 11.0 to determine if species and stage of development (freshly laid or developed) affected surface fouling. A separate 1-way ANOVA was also performed on *Dicathais orbita* (fresh, trochophore and late stage veliger) egg capsule images to determine if the extent of fouling at the trochophore stage differed from the other developmental stages. The assumption of equal variances was tested using Levene’s test and normality was tested using the Kolmogorov-Smirnov test in SPSS. The α-level was set at 0.05 for a significant difference, but then Bonferroni corrected to 0.0167 to account for the multiple comparisons using the same data (Quinn & Keough 2002).

**LIVE/DEAD BacLight bacterial viability assay.** A stock solution was prepared by mixing equal volumes of the reagents SYTO 9 and propidium iodide (Invitrogen, see product information, www.invitrogen.com). The working solution was then prepared by adding 3 µl of the stock solution to 1 ml of 0.85% NaCl. Positive and negative controls of live and ethanol-killed *Pseudoalteromonas* sp. S91SB cultures were independently stained to optimize the concentration of dyes and parameters for Confocal Laser Scanning Microscopy (CLSM).

Six independent capsules from each species were stained and examined. Under sterile conditions, egg capsules were excised from the cluster and rinsed with Milli-Q water before incubation in the working dye solution for 15 min in the dark. After incubation, a medial section of the egg capsule was prepared using sterile forceps and a surgical blade. The apex and basal regions of the egg capsule were removed, and a section of capsule wall (~0.5 × 0.5 cm) mounted onto a clean glass slide and viewed under an Olympus BX50F-3 epifluorescence microscope. A Bio-Rad MRC-1000UV CLSM was subsequently used to capture quality images of the stained egg capsules. Images in the green spectra (live) and red spectra (dead) were taken separately for quantification purposes.

The amount of live and dead microorganisms was quantified using Image J 1.32j software. The total area of fluorescence within xz images was quantified under a standardized area of 96.26 µm (x) by 96.26 µm (z). Paired sample *t*-tests were used to compare the amount of green and red fluorescence on the surface of egg capsules within each species (n = 6). One-way ANOVAs were then used to separately compare the amount of green and red fluorescence between species. A square-root transformation was performed on the data in order to meet the assumptions of equal variances (Levene’s test). After applying a Bonferroni correction, α was set at 0.0167.

**Bacterial preparation.** *Pseudoalteromonas* sp. Strain S91SB (hereafter S91SB) is a derivative of Strain S91 (Techarnjanaruk et al. 1997) into which a *gfp* gene has been inserted. S91SB was grown in 10 ml of marine minimal medium (MMM; Östling et al. 1991) supplemented with 0.2% glutamate at 30°C and allowed to grow overnight prior to use. Growth of S91SB was screened under a 100 W quartz halogen lamp with an infrared and 280 nm band-pass filter to check for GFP expression (Andover, part no. FS10-50) (Stretton et al. 1998). Green fluorescence indicated that bacteria were growing healthily.

**Bacterial attachment experiment using flow chambers.** Flow chamber experiments were carried out in triplicate using independent samples for 0, 24 or 72 h. Egg capsules were sterilized by dipping into 70% ethanol for 10 min, and then washed with sterile Milli-Q water before sectioning into 0.5 × 0.5 cm pieces under aseptic conditions. Examination of control samples under the Olympus BX50F-3 epifluorescence microscope confirmed that this procedure successfully destroyed the existing biofilm without structurally altering the egg capsule walls. A single piece of capsule was placed into a once-through flow chamber, with the exterior surface facing up. Flow chambers used in this experiment were as described by Hale & Mitchell (2001), and experimental conditions were maintained as described by Fitch et al. (2002). Two round glass coverslips (viewing ports) were sealed together with a rubber O-ring mounted in a stainless steel scaffold. The O-ring contained 2 needles acting as inlet and outlet ports. A piece of egg capsule was placed between the coverslips before sealing the chamber.

After assembly of the flow chamber, an 8-roller Cole-Palmer Masterflex peristaltic pump (7521-57) was used to pump 70% ethanol through the system for 15 min, followed by sterile MMM for 20 min. After flushing the system with sterile MMM, the peristaltic pump was turned off and 3 ml of a 10⁻¹ dilution of S91SB overnight culture was introduced. The chambers were left for 1 h to allow settlement and attachment of S91SB to egg capsules. Flow was resumed after 1 h. Room temperature and flow rate was maintained at 24°C and 1 ml min⁻¹, respectively (Delpin et al. 2000). Because MMM does not contain any carbon source, egg capsules were the only organic material available for bacterial attachment. After the completion of each time point (0, 24 or 72 h), the chambers were disconnected and taken to Adelaide Microscopy TPLSM for analysis.

**Microscopic analysis of biofilm production using TPLSM.** A Bio-Rad Radiance 2000MP in conjunction
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with a Nikon Eclipse TE300 inverted Multi-Photon Microscope was used to collect images of biofilms formed in the chambers. The microscope uses a 60× water immersion lens with a numerical aperture of 1.2 and a Coherent Mira900-F titanium:sapphire ultrafast laser, which has an excitation spectrum of pulsed 800 nm light equivalent to 1 photon of 400 nm light. GFP produced by S91SB was visualized using an excitation wavelength (λ) of 800 nm and an emission λ of 515 nm. The 2 photons of 800 nm light are equivalent to 400 nm but do not damage cells as does 400 nm UV light typically used to excite GFP. Auto-fluorescence produced by the egg capsule was detected as blue at 460 nm. Horizontal (xy) and vertical (xz) optical sections of biofilm were imaged; within each xy image, 3 random xz sections were collected. This procedure was repeated 3 times for each chamber, yielding 3 xy and 9 xz images. Confocal Assistant was used to analyse all images and convert them to .tif files. Image J 1.32j was used to quantify the coverage of S91SB on egg capsules by calculating the total area of fluorescence, which is then divided by the mean area for a single S91SB cell (0.56 µm²) (Everuss 2004). A standardized area of xz sections was used for quantification of fluorescence: 96 µm (x) by 52 µm (z).

One-way ANOVAs with Tukey’s Honestly Significant Difference (HSD) post hoc tests were used to compare differences in the S91SB counts among each time point for *Dicathais orbita* and *Sepioteuthis australis* egg capsules. Independent t-tests were also used to compare the difference in S91SB counts between species at 24 and 72 h. All analyses in this data set were tested more stringently (α = 0.01) because assumptions of normality were not met (Underwood 1997), even after a series of transformations were attempted on the data (e.g. log x + 1, square root and cube root).

**RESULTS**

**Scanning electron microscopy**

The SEM revealed differences in the egg capsule structure of *Dicathais orbita* and *Sepioteuthis australis* capsules. Distinct layers of *D. orbita* egg capsules can be seen in transverse sections (Fig. 1a). The capsule comprises a solid inner wall and porous sections sandwiching a thick medial section with possible layering. The capsule wall is approximately 25 µm thick (Fig. 1a). By comparison, *S. australis* capsules were observed to consist of multiple overlapping layers similar to a plywood structure (Fig. 1b). The capsule wall of fresh samples was approximately 300 µm thick, and none of the layers appeared to be porous.

Differences in the surface microtopography were observed both between the species and within species at different stages of development (Figs. 2 & 3). Freshly laid capsules (<1 wk old embryos) of *Dicathais orbita* possessed a thin layer of crust over a microtopographical structure on the outer-most surface (Fig. 2a). The surface was almost free from any bacteria and algae. As the egg capsules matured (1 to 3 wk of age), the crust began to breakdown, exposing the microtopographical features (Fig. 2b). This layer has regular homogeneous ridges separated by 1 to 5 µm (Fig. 2c). Sparse rod and cocci bacteria were observed on the walls and ridges during the second week of development. However, during the third week, densities of these bacteria increased and other fouling organisms such as filamentous algae were observed, forming a mixed biofilm community (Fig. 2d). In the later stages of development (>4 wk old veligers), the fouled outer wall structure began to dissociate and shed from the capsule, leaving behind a naked capsule without any texture (Fig. 2e). This surface was initially free from microbes, but bacteria began colonization several days
later. Unidentified droplets ranging from 8 to 20 µm appeared on the surface of mature egg capsules after the majority of the outer wall dissociated (Fig. 2f). These droplets were either solitary or clumped in association with attached bacteria. A greater density of these droplets was observed on cryostat preparations than on critically point dried capsules. However, fewer microorganisms were observed on the surface of critically point dried samples (data not shown).

The surface of Sepioteuthis australis egg capsules typically had irregular folds resembling ridges and troughs about 5 µm in width and depth (Fig. 3a). Freshly laid capsules (undifferentiated blastomere stage) collected from the sea contained fewer microorganisms on the surface (Fig. 3a,b) than did late-stage capsules containing paralarvae (Fig. 3c–f). Capsules of S. australis in the later stages of development collected from the field were covered with filamentous algae...
and spores (Fig. 3c,d), as well as an exopolymeric substance and various bacteria (rod, cocci and spiral forms, e.g. Fig. 3e) and diatoms (Fig. 3f).

Fig. 4 shows the total percentage cover by all fouling organisms on the surface of Dicathais orbita and Sepioteuthis australis egg capsules. A 2-factor ANOVA was conducted to test the effect of species and stages of development (fresh and developed) on the level of fouling (percentage cover). Percentage coverage was significantly higher in the later stages of development (fresh embryos vs. developed capsules, $F = 25.83, \text{df} = 1, p < 0.001$). However, species ($F = 1.066, \text{df} = 1, p = 0.317$) and the interaction term ($F = 2.296, \text{df} = 1, p = 0.149$) were not significant. A 1-way ANOVA for surface fouling on D. orbita capsules indicated that there was a statistically significant difference among the 3 different stages of development ($F = 16.318, \text{df} = 2, p < 0.001$). Post hoc comparisons using Tukey’s HSD test
revealed that percentage cover did not differ significantly between fresh capsules and trocophore stage capsules \( (p = 0.671) \), whereas developed capsules were significantly more fouled than both fresh and trocophore stage capsules \( (p < 0.001, p = 0.002, \text{respectively}, \text{Fig. 4a}) \).

**LIVE/DEAD BacLight bacterial viability assay**

BacLight successfully stained microorganisms on the surface of *Dicathais orbita* and *Sepioteuthis australis* egg capsules (Fig. 5). However, the unidentified droplets on the surface of *D. orbita* capsules did not stain with either of these nucleic acid dyes. Control experiments demonstrated that ethanol-killed bacteria gave the expected red fluorescence, while viable cells remained green owing to retention of SYTO9. The egg capsules of both molluscan species were found to have a mixture of live and dead bacterial cells on their surface (Figs. 5 & 6). The thickness of the biofilm on *S. australis* capsules (70 µm, Fig. 5b) was approximately 4 times thicker than the biofilm on *D. orbita* egg capsules (18 µm, Fig. 5a). Consistent with this, the total green fluorescence on the surface of *S. australis* capsules was found to be significantly higher than the green fluorescence emitted from *D. orbita* capsules \( (F = 193.61, \text{df} = \ldots) \).
Similarly, the total red fluorescence was significantly higher on *S. australis* than on *D. orbita* egg capsules ($F = 75.89, \text{df} = 1, p < 0.001$).

No significant difference was observed between the amount of live vs. dead microbes for either *Sepioteuthis australis* ($t = 1.747, \text{df} = 5, p = 0.141$) or *Dicathais orbita* ($t = 0.898, \text{df} = 5, p = 0.410$) (Fig. 6). However, the proportion of healthy bacteria was slightly higher (53%) than that of dead bacteria on surfaces of *S. australis*, whereas a slightly greater proportion of dead bacteria (52.5%) was present on the surface of *D. orbita* egg capsules. Under the CLSM, dead bacteria in association with a string of the unidentified droplets were observed on the egg capsules of *D. orbita* (data not shown). Fluorescence microscopy also revealed an association between dead bacteria and the droplets; however, small colonies of live bacteria were sometimes observed around the droplets as well.

### Bacterial attachment

*S91SB* was able to survive and attach to egg capsule surfaces within flow chambers. Autofluorescence of egg capsules was observed as blue, which allowed differentiation of GFP-fluorescing bacteria. *S91SB* did not attach well to the surface of *Dicathais orbita* egg capsules (Fig. 7), but some small micro-colonies of bacteria were observed after 24 h. Higher densities of bacteria were occasionally seen attached to the glass slide as a layer of floating GFP-producing bacteria above the surface of the capsule (data not shown). TPLSM images also showed the unidentified droplets on the surface of *D. orbita* egg capsules, but these were not associated with *S91SB* attachment. In contrast, *S91SB* was almost always observed to colonize the surface of *Sepioteuthis australis* egg capsules in high densities (Fig. 7).

The number of *S91SB* bacteria attaching to the surface of egg capsules from the 2 species was quantified by the area of fluorescence. ANOVA revealed a significant difference among the 3 time points in *Dicathais orbita* ($F = 7.717, \text{df} = 2, p = 0.005$). Post hoc analysis revealed that there were significantly fewer attached *S91SB* at 0 h than at 72 h ($p = 0.004$) (Fig. 7). The number of attached *S91SB* at 24 h did not differ significantly from that at 0 h ($p = 0.062$) or 72 h ($p = 0.370$). For *Sepioteuthis australis*, increasing numbers of *S91SB* clearly attached to the surface over the 3 d period (Fig. 7), and significant differences were observed among the time points ($F = 151.871, \text{df} = 2, p < 0.001$). However, post hoc analysis revealed that there was no significant difference between 0 and 24 h ($p = 0.146$), whereas bacterial load significantly increased after 72 h, with $p < 0.001$ for both the 0 and 24 h time points. Independent *t*-tests were performed to compare between species at 24 and 72 h. This revealed significantly higher bacterial counts on the surface of *S. australis* than on *D. orbita* egg capsules after both 24 h ($t = 9.765, \text{df} = 6.2, p < 0.001$) and 72 h ($t = 12.45, \text{df} = 5.3, p < 0.001$) in the flow chamber (Fig. 7).

### DISCUSSION

Egg capsules, being sessile in the marine environment, are susceptible to microbial assault (Shields 1990, Benkendorff et al. 2001). Therefore, they require some form of protection against fouling organisms. From this study, it is evident that the egg capsules of *Dicathais orbita* have anti-fouling defense mechanisms.
that reduce the impacts of microbial foulers on the encapsulated embryos until they are released as pelagic larvae. This was shown by comparatively low microbial densities observed on samples taken directly from the field and after experimental induction of bacterial attachment in the laboratory. By comparison, the eggs of Sepioteuthis australis appear to provide a favourable environment for fouling organisms, but may be more resistant to microbial penetration owing to the overlapping of multiple layers in the capsule wall, which is 15 times thicker than that of the D. orbita capsules (Fig. 1).

As expected from the longer exposure times, the percentage cover of fouling viewed on the egg capsules of both species increased as embryonic development progressed (Fig. 4). Percentage coverage of fouling organisms on the surface was significantly higher in developed capsules than in fresh capsules (p < 0.001). A greater range of fouling organisms was observed on the squid egg capsules (e.g. Fig. 3) than on those of Dicathais orbita, including several bacterial morphotypes as well as micro- and macroalgae. The presence of filamentous algae on well-developed Sepioteuthis australis egg capsules was observed with independent observations on a Tasmanian population of this species by Steer et al. (2002), who showed that algal fouling promotes synchronized development of S. australis egg capsules. The beneficial effects from fouling by photosynthetic algae could include increased oxygen levels for encapsulated embryos during the day (Cohen & Strathmann 1996), protection from the effects of UV radiation (Biermann et al. 1992) and possibly a reduction in settlement by other fouling organisms (Harper & Skelton 1993, Mao-che et al. 1996, Kaehler 1999, Cerrano et al. 2001). However, after removal of the crust from D. orbita capsules, a regular microtopographical feature of ridges and pores was observed throughout the trophophore stage (1 to 3 wk of age) (Fig. 2b). This surface texture is consistent with a physical defense strategy involving minimal contact points for attachment by biofouling organisms. Scardino et al. (2006) provided support for the ‘attachment point theory’, whereby reduced adhesion strength occurs for fouling organisms that are larger than the scale of microtexture. Nanoridges and pores at a scale suitable for the reduction of microbial attachment were found on the skin of pilot whales Globicephala melas (Baum et al. 2002) and bivalve shells (Scardino et al. 2003). The spacing between ridges on D. orbita capsules (1 to 5 µm, Fig. 2c) compares favorably with the relatively unfaouled shells of Mytilus galloprovincialis (1 to 2 µm, Scardino et al. 2003). Nevertheless, after only 3 wk, D. orbita egg capsules maintained in a recirculating seawater system had still become heavily fouled (Fig. 2d).

Egg capsules of Dicathais orbita in the late veliger stage (>4 wk) contained the highest percentage cover of fouling organisms (Fig. 4a) and appeared to be a lot smoother (Fig. 2e,f) than the earlier-stage capsules (Fig. 2a,b). Ablation of the micro-textured outer wall layer was observed under the SEM (Fig. 2e). Mechanical shedding of the biofilm was also reported for the alga Dilsea carnosa during late autumn and early spring (Nylund & Pavia 2005). The effectiveness of mechanical shedding in limiting surface fouling on D. carnosa was suggested to be especially important in temperate regions, where huge variations in fouling organisms arise as a result of seasonal change (Nylund & Pavia 2005). The egg capsules of D. orbita are deposited on temperate reefs in late spring to summer.
Thus, shedding of the outer capsule wall may be an effective strategy for preventing biofilm build-up at a key time in their embryonic development when there are heavy microbial loads in the water column.

After shedding of the fouled ridged layer, the surface of *Dicathais orbita* capsules was found to be relatively free of microbial fouling (Fig. 2e), but areas with increasing densities of bacteria were observed in older capsules, accompanied by large unidentified droplets (Fig. 2f). The droplets appeared to be secreted through pores in the wall as the outer capsule degrades. These droplets were clearly not cellular because they did not stain green or red when incubated with the BacLight nucleic acid probes. They also did not appear to be membrane-bound vesicles, remaining globular rather than dissolving onto the surface. This suggests that they were hydrophobic, which is further supported by the lower densities of droplets observed using standard SEM after critical point drying in ethanol compared with cryostat SEM. Soaking the egg capsules in organic solvents such as dichloromethane was also observed to remove the droplets from the surface (authors’ unpubl. data), suggesting that they may contain lipophilic compounds such as the indole dimer tyriverdin, which is a potent bacteriostatic agent previously reported from extracts of *D. orbita* egg capsules (Benkendorff et al. 2000). The droplets were observed to aggregate with bacteria on the capsule surface (Fig. 2f) and fluorescence microscopy revealed they were frequently associated with dead bacteria (data not shown), suggesting that they may have antimicrobial properties. Several species of marine macroalgae have the ability to produce secondary metabolites and release them onto the surface to prevent epibiont settlement (e.g. Ragan & Glombitza 1986, Clayton & Ashburner 1994, Dworjanyn et al. 1999). Further studies on the chemical defense of *D. orbita* capsules are clearly required.

Bacterial attachment experiments, undertaken in laboratory flow chambers, further support the idea that *Dicathais orbita* egg capsules provide a less favorable environment for biofilm formation than do the *Sepioteuthis australis* capsules. *Pseudoalteromonas* species are motile marine bacteria commonly found in biofilm communities in temperate Australian waters, and are not known to have any specific interactions with molluscan egg capsules, thus providing a good model system for random biofouling. They secrete chitinases and proteases to break down particulate organic material (POM) (Techkarnjanaruk & Goodman 1999). Since molluscan egg capsules consist mainly of proteins and carbohydrates (Hunt 1966, Flower et al. 1969, Rapoport & Shadwick 2002), it was predicted that S91SB would preferentially attach to the egg capsules and utilize them as a carbon source unless the capsules had properties that prevented bacterial attachment. However, S91SB were found to preferentially adhere to the glass coverslips than to *D. orbita* capsules (data not shown). Bacterial attachment was clearly higher at 72 h than at 0 h, however, there was no significant difference in *D. orbita* capsules between the 24 and 72 h time points, indicating that the biofilm build-up quickly becomes stagnant (Fig. 7). Conversely, bacterial attachment experiments on *S. australis* showed that S91SB was able to attach to these egg capsules and form a thick layer of biofilm (Fig. 7). A significant increase in S91SB counts at both 24 h (p < 0.001) and 72 h (p < 0.001) was observed (Fig. 7), again supporting the notion that antifouling protection is absent from *S. australis* capsules.

The removal and prevention of biofilm formation using physical, mechanical and chemical means has been described in other organisms, but typically these defense mechanisms are studied in isolation. Our present study indicates that the neogastropod *Dicathais orbita* may use a combination of all these defense mechanisms to defend its egg capsules, including a surface texture not suitable for bacterial attachment, followed by shedding of the outer layer to remove existing microbial colonization and then exudation of unidentified chemical droplets that aggregate and possibly interfere with bacterial growth on the capsules’ surface. By comparison, the egg capsules of the cephalopod *Sepioteuthis australis* had comparatively high loads of live bacteria, and their microtexture appeared to provide good habitat for microbial colonization. Our observations of differences in the surface structure and microbial fouling properties of these 2 molluscan egg capsules contribute to our understanding of antifouling defense mechanisms in marine organisms.

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