Antifouling character of ‘active’ hybrid xerogel coatings with sequestered catalysts for the activation of hydrogen peroxide

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Halide-permeable xerogel films prepared from sols containing 50 mol% aminopropyltriethoxysilane (APTES)/50 mol% tetraethoxysilane (TEOS) or 10 mol% APTES/90 mol% TEOS and 0.015 M selenoxide or telluride catalyst in the sol gave reduced settlement of cypris larvae of the barnacle Balanus amphitrite and larvae of the tubeworm Hydroides elegans in the presence of artificial seawater (ASW) and hydrogen peroxide (5–100 μM) relative to glass controls. Settlement of Ulva zoospores was lower on both the 50 mol% APTES/50 mol% TEOS and 10 mol% APTES/90 mol% TEOS xerogel formulations in comparison with glass controls with or without the added catalyst. The 50 mol% APTES/50 mol%TEOS xerogel containing telluride catalyst gave reduced settlement of Ulva zoospores in the presence of 100 μM H2O2 in ASW compared with the same coating without added peroxide. Scanning electron microscopy and XPS data suggest that exposure to H2O2 does not lead to chemical or morphological changes on the xerogel surface.

Keywords: active xerogels; hydrogen peroxide; biofouling; algae; Ulva; barnacles; Balanus amphitrite; tubeworms; Hydroides elegans

Introduction

Worldwide, marine biofouling is a significant economic problem costing $30–$60 million per year in transportation costs due to increased fuel consumption from added drag (Townsin 2003; Schultz 2007). The most common method to minimise biofouling is to coat the submerged surface with an antifouling (AF) paint that leaches biocide(s) to prevent marine life from attaching to the surface (Yebra et al. 2004). A number of alternative technologies are available commercially, the most notable being based on silicone elastomers, which release accumulated fouling under suitable hydrodynamic conditions (eg Kavanagh et al. 2005; Wendt et al. 2006).

A conditioning film forms within seconds of immersion and is followed by settlement of bacteria, unicellular algae and cyanobacteria (blue-green algae) to form a biofilm within hours of immersion. Surfaces bearing this biofilm rapidly become overgrown by macrofoulers such as barnacles, tubeworms and macroalgae (Callow 2000).

The barnacle Balanus amphitrite is a common member of fouling communities in coastal environments (Clare and Høeg 2008) and it is often found in great abundance on ships’ hulls and pier pilings and it has an extremely wide geographical distribution (Otani et al. 2007). The polychaete tubeworm Hydroides elegans is a common member of the hard-fouling community in tropical and sub-tropical waters and is an early colonist of submerged surfaces in Pearl Harbour (Pettingill et al. 2007). Ulva (syn. Enteromorpha) (Hayden et al. 2003) is a key macroalga that fouls ships and is tolerant of a wide range of environmental conditions and surface coating types including biocidal AF paints (Callow 1996). Dispersal of Ulva is mainly through motile, quadriflagellate zoospores, which are released in large numbers and which respond to a large number of settlement cues (Callow and Callow 2000). The settlement (attachment) of these three organisms was used to evaluate the xerogel surfaces in the present study.

An alternative to the use of biocides in the marine environment to minimise biofouling is to design a surface whose characteristics discourage either biofilm formation or the settlement of fouling organisms or whose characteristics reduce the strength of attachment of fouling organisms such that they are more easily removed by hydrodynamic forces (Callow et al. 2000; 2002; Finlay et al. 2002; Genzer and Efimenko 2006). Hybrid xerogel coatings with critical surface
energies near 20 mN m⁻¹, which is in the range where minimal bioadhesion is observed due to the formation of weak boundary layers (Baier et al. 1968; Baier 1984), reduce the settlement of cyprids of the barnacle *B. amphitrite* and, for one xerogel coating, display excellent fouling-release properties for juvenile barnacles (Tang et al. 2005).

Another approach to reduce fouling is to use naturally occurring reagents in seawater to create biocides *in situ*. Halide salts are slowly oxidised by H₂O₂ to give positive halogen species, which are known to have biocidal properties (Williams and Schroeder 2004). In the open ocean, concentrations of H₂O₂ approach 0.2 mM and can be much higher (up to 50 µM) in coastal areas where concentrations in rain water and runoff are in the range of 16–526 mM (Cooper and Zika 1983; Willey et al. 1999; Yuan and Schiller 2000, 2001; Clark et al. 2008). At these peroxide concentrations and in the presence of the 0.5 M chloride, 1 mM bromide and 1 µM iodide found in seawater, the production of positive halogen species cannot overcome the rate at which the positive halogen species decompose or are consumed.

H₂O₂ is also formed on submerged surfaces by organisms in the biofilm (Chandrasekaran and Dexter 1993; Le Bozec et al. 2001; Dexter et al. 2003). Local concentrations of H₂O₂ in the biofilm can approach 50 µM.

Diorganoselenoxides and diorganotellurides are efficient catalysts for the oxidation of halide salts with H₂O₂ to produce the corresponding hypohalous acid (Francavilla et al. 2000, 2001; Higgs et al. 2001; Drake et al. 2003; Goodman and Detty 2004; Bennett et al. 2008). If these catalysts were covalently sequestered within a porous film coating, the catalysts should react with the peroxide found in seawater (Cooper and Zika 1983; Willey et al. 1999; Yuan and Schiller 2000, 2001; Clark et al. 2008) or that is produced by the biofilm (Chandrasekaran and Dexter 1993; Le Bozec et al. 2001; Dexter et al. 2003) and with the halide salts found in seawater to create a surface that is chemically inhospitable to settlement and adhesion. Specifically, the catalysts would generate hypohalous acid or ‘bleach’ on the coating surface. The biocide(s) could inhibit settlement by either killing or deterring the attachment of larvae or spores, but if they settle the constant generation of hypohalous biocides, might be expected to kill or inhibit the growth of the attached organisms.

Xerogel films prepared by the sol–gel process are easily processed near ambient conditions, are economically and environmentally friendly, and can be applied to surfaces by a variety of means including spraying, brushing or rolling, dip coating and spin coating. In addition, the properties of the xerogel including porosity, permeability towards various ions and neutral molecules, and the extent of crosslinking can be controlled by the precursors one uses and the processing conditions (Brinker and Scherer 1990; Avnir 1995; Dave et al. 1995; Ingersoll and Bright 1997; Jordan et al. 1998). Furthermore, other reagents can be sequestered within the xerogel and these reagents can maintain their normal chemical reactivity/function found in solution (Avnir 1995; Dave et al. 1995; Pandey et al. 2000; Tang et al. 2003).

This investigation examines the performance of two-component, hybrid xerogel films that are doped with 4-(hydroxymethyl)phenyl benzyl selenoxide (Se₁) or 3-(n-hexyltelluro)-1-propanol (Te₁) (Chart 1) as ‘active’ coatings to minimise the biofouling. Specifically, settlement and growth of fouling species can be minimised by the generation of hypohalous acids from the H₂O₂ and halide salts naturally occurring in seawater. The xerogel films were formed from sols that are composed of 50 mol% aminopropyltriethoxysilane (APTES) and 50 mol% tetraethoxysilane (TEOS) or 10 mol% APTES and 90 mol% TEOS. These active coatings have been examined with respect to the settlement of cyprids (cypris larvae) of *B. amphitrite*, larvae of *H. elegans*, and *Ulva* zoospores. Glass was included as an inert standard.

**Materials and methods**

**Chemical reagents**

All reagents were used as received. Deionised water was prepared to a specific resistivity of at least 18 MΩ using a Barnstead NANOpure II system. TEOS, tetraethoxysilane (TMOS), 3-propyltrimethoxysilane (C₃TMOS) and 3-aminopropyltriethoxysilane were obtained from Gelest. Hydrochloric acid was obtained from Fisher Scientific. Ethanol was a product of Quantum Chemical. Borosilicate glass microscope slides were obtained from Fisher Scientific. Selenoxide catalyst, Se₁, was prepared according to the literature procedure (Bennett et al. 2008).

**Preparation of Te₁ catalyst**

*Preparation of 3-(tert-butyldimethylsilyloxypropyl) hexyl elluride (1)*

1-Bromohexane (0.85 ml, 6.06 mmol) was dissolved in anhydrous tetrahydrofuran (THF, 25 ml) and tert-BuLi (7.8 ml of a 1.7 M solution in THF, 13 mmol) was added dropwise via syringe at −78°C. After 1 h, Te powder (0.772 g, 6.06 mmol) was added.
and the resulting mixture was stirred for 15 min at -78°C and was then stirred at room temperature until the Te was completely consumed. The reaction mixture was cooled to -78°C, 3-bromopropoxy-tert-butylimethysilane (1.82 g, 6.06 mmol) (Francavilla et al. 2000, 2001) was added, and stirring was continued for 1 h at -78°C and for 15 h at room temperature. The reaction mixture was poured into 150 ml of water. These treated slides were dried under ambient conditions and used within 1 day.

**Preparation of 3-(n hexyltelluro)-1-propanol (Te1)**

Telluride 1 (1.04 g, 2.69 mmol) was dissolved in anhydrous THF (30 ml) and stirred for 15 min. Tetra-n-butylammonium fluoride (5.39 ml of a 1 M solution in THF, 5.39 mmol) was added dropwise over several minutes and the resulting solution was stirred for 2 h at room temperature. Excess saturated ammonium chloride (50 ml) was added and the resulting mixture was stirred for an additional 10 min. The reaction mixture was concentrated, water (50 ml) was added, and the organic products were extracted with ethyl acetate (3 x 40 ml). The combined organic extracts were dried over sodium sulphate and concentrated to give a yellow oil, which was purified on SiO2 eluting with 30% dichloromethane in hexanes to give 3-(tert-butyldimethylsilyloxypropyl)hexyltelluride (1) as a yellow oil (1.31 g, 56%). \(^1\)H NMR (300 MHz, CDCl\(_3\)) \(\delta\) 3.64 (t, 2 H, J = 6.0 Hz), 2.66 (t, 2 H, J = 7.8 Hz), 2.63 (t, 2 H, J = 7.8 Hz), 2.63 (t, 2 H, J = 7.2 Hz), 1.73 (quint, 2 H, J = 7.2), 1.27–1.35 (m, 6 H), 0.88–0.89 (m, 12 H), 0.05 (s, 6 H); \(^{13}\)C NMR (75 MHz, CDCl\(_3\)) \(\delta\) 64.2, 35.2, 32.2, 31.7, 31.2, 25.9, 22.5, 18.3, 14.0, 2.92, -1.69, -5.28; HRMS (EI) \(m/z\) 388.1431 (calcd for \(\text{C}_{13}\text{H}_{34}\text{OSiTe}\) 388.1441).

**Sol preparation**

The sol/xerogel composition is designated in terms of the mol% of Si-containing precursors. Thus, a 10/90 APTES/TEOS composition contains 10 mol% APTES and 90 mol% TEOS. The various sols and their abbreviations are summarised in Table 1.

**Sol TEOS**

TEOS (3.130 g, 15.02 mmol, 3.345 ml) was mixed with water (0.54 ml), ethanol (3.40 ml) and 15 \(\mu\)l of 0.1 N HCl (1.5 \(\mu\)mol) in a sealed glass vial and stirred at room temperature for 6 h.

**Sol APTES**

APTES (2.544 g, 11.49 mmol) was added dropwise to a stirring solution of 2 g of 6.67 N HCl and ethanol (10.56 ml). Once addition was complete, the solution was sonicated at room temperature for 40 min.

**Sol 10/90 APTES/TEOS (B6 xerogel)**

Sol APTES (5 ml) was added dropwise to a stirring mixture of Sol TEOS (16.77 ml). The resulting mixture was sonicated at room temperature for 20 min.

**Sol 10/90 APTES/TEOS with Se1 catalyst (B6Se1 xerogel)**

Selenoxide catalyst Se1 (0.088 g, 0.300 mmol) was added to Sol 10/90 APTES/TEOS (20.00 ml) and stirred at room temperature for 5 min to give 0.015 M Se1 in Sol 10/90 APTES/TEOS.

**Sol 10/90 APTES/TEOS with Te1 catalyst (B6Te1 xerogel)**

Telluride catalyst Te1 (0.096 g, 0.300 mmol) was added to Sol 10/90 APTES/TEOS (20 ml) and stirred at room temperature for 5 min to give 0.015 M Te1 in 10/90 APTES/TEOS.

**Preparation of glass control slides**

Plain 25 mm x 75 mm borosilicate glass slides were cleaned by soaking in 1 N HCl for 24 h and were then rinsed with copious amounts of deionised water and ethanol. These treated slides were dried under ambient conditions and used within 1 day.

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**Table 1. Coating abbreviations used with the various sol formulations and catalyst combinations.**

<table>
<thead>
<tr>
<th>Sol formulation</th>
<th>Catalyst</th>
<th>Coating abbreviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>50/50 APTES/TEOS</td>
<td>None</td>
<td>B5</td>
</tr>
<tr>
<td>50/50 APTES/TEOS</td>
<td>0.015 M selenoxide Se1</td>
<td>B5Se</td>
</tr>
<tr>
<td>50/50 APTES/TEOS</td>
<td>0.015 M telluride Te1</td>
<td>B5Te</td>
</tr>
<tr>
<td>10/90 APTES/TEOS</td>
<td>None</td>
<td>B6</td>
</tr>
<tr>
<td>10/90 APTES/TEOS</td>
<td>0.015 M selenoxide Se1</td>
<td>B6Se</td>
</tr>
<tr>
<td>10/90 APTES/TEOS</td>
<td>0.015 M telluride Te1</td>
<td>B6Te</td>
</tr>
</tbody>
</table>
**Sol 50/50 APTES/TEOS (B5 xerogel)**
Sol APTES (10 ml) was added dropwise to a stirring mixture of Sol TEOS (3.73 ml). The resulting mixture was sonicated at room temperature for 20 min.

**Sol 50/50 APTES/TEOS with Se1 catalyst (B5Se1 xerogel)**
Selenoxide catalyst Se1 (0.088 g, 0.30 mmol) was added to Sol 50/50 APTES/TEOS (20 ml) and stirred at room temperature for 5 min to give 0.015 M Se1 in Sol 50/50 APTES/TEOS.

**Sol 50/50 APTES/TEOS with Te1 catalyst (B5Te1 xerogel)**
Telluride catalyst Te1 (0.096 g, 0.30 mmol) was added to Sol 50/50 APTES/TEOS (20 ml) and stirred at room temperature for 5 min to give 0.015 M Te1 in 50/50 APTES/TEOS.

**Xerogel film formation**
Xerogel films were formed by spin casting onto 25 mm × 75 mm glass microscope slides. To form the film on the glass, a 0.4-ml sample of a given sol solution was evenly delivered onto the entire surface of the slide, which was then placed in the spin coater. The spin coater was then engaged and the substrate rotated at 3000 rpm for 30 s. All films were stored under ambient conditions in the dark for at least 1 week prior to submission for settlement and removal assays.

**Film thicknesses**
Film thicknesses were determined by using a Tencor instruments, alpha-step 500 profilometer with a diamond-point stylus. Triplicate samples of each film type were measured. The film thicknesses reported represent the average and associated standard deviations (SDs). An uncoated section of substratum, obtained by placing Scotch tape across the substratum edge prior to film deposition, was used to establish a baseline.

**Evaluation of halide permeability in xerogel films**
The B5 and B6 xerogels described above were prepared from sols that were doped with the luminescent reporter molecule tris(4,7-diphenyl-1,10-phenanthroline)ruthenium(II) chloride ([Ru(dpp)3]2+) and the resulting xerogels were exposed to various concentrations of dissolved I− or O2, a quencher for luminescence (Tang et al. 2003). In the simplest scenario of a luminophore sequestered within a homogeneous medium, luminophore quenching obeys the Stern–Volmer relationship (Eftink 1991; Lakowicz 1999):

\[
I_0/I_Q = 1 + K_{SV}[Q] = 1 + k_q\tau_0[Q]
\]  

(1)

In this expression, \(I_0\) and \(I_Q\) are the steady-state luminescence intensities in the absence and presence of quencher, respectively; \(\tau_0\) is the excited-state lumino- 

**Scanning electron microscope images**
Scanning electron microscope (SEM) images were recorded by using a Hitachi model S-800 field emission scanning electron microscope operating at a 25 kV acceleration voltage. The xerogel films were over-

**X-ray photoelectron spectroscopy**
X-ray photoelectron spectroscopy (XPS) spectra were recorded using a Physical Electronics Laboratories (PHI) Model 5100 spectrometer equipped with a Mg/Ti dual anode source, an Al/Be window, a hemispherical analyser, and single channel channeltron detector. An achromatic Mg K-α X-ray (1253.6 eV) source was operated at 300 W, 15 kV and 20 mA. The system base pressure was no higher than 2 × 10⁻⁶ torr, with an operating pressure that did not exceed 1 × 10⁻⁶ torr. A pass energy of 89.45 eV was used to obtain the survey spectra and 35.75 eV was used for the high resolution, multi-region scans. Spectra were obtained at 45° take-off angles. The sampling depth for C and Si is ~80–90 Å. The instrument was calibrated using Mg K-α X-radiation such that the Ag 3d₅/₂ peak of sputtered clean Ag had a binding energy of 367.9 ± 0.1 eV and the binding energy difference between the Cu 2p₃/₂ and Au 4f₇/₂ was 848.7 ± 0.1 eV. The full width at half maximum for the Ag 3d₅/₂ peak was 80.80 eV at 30,000 counts s⁻¹. Data manipulation was performed using a Perkin-Elmer 7500 professional computer running PHI ESCA Version 2.0 software.

Instrumental sensitivity allows ratios of masses of 0.01 to be measured at the surface. However, the C(1s)/Si(2p3) ratio of 0.2 measured on glass indicates
that this level of contamination with atmospheric carbon might be present in all samples and only values that are significantly larger than 0.2 for the C(1s)/Si(2p3) ratio have meaning. Background levels of N(1s)/Si(2p3) were <0.01.

Xerogel coatings were soaked in artificial seawater (ASW) without or with 0.9 M H$_2$O$_2$ for 24 h, were washed with distilled water, and air-dried for 24 h prior to XPS measurements.

**Cyprid settlement assays using B. amphitrite**

Thirty to 50 larvae were ‘drop assayed’ onto each of the replicate surfaces in ASW with and without 100 μM or 20 μM H$_2$O$_2$. Assays lasted ~48 h, although the exact duration depended on the time it took 50% of the larvae to settle (a mean settlement of 0.50) in the control standard, an uncoated glass slide. Stopping the assays after ~50% of the larvae have settled on the control surfaces provides information on AF and inductive characteristics of the experimental coatings with respect to the rate of settlement on the surfaces. At the end of the initial assay period the numbers of larvae that successfully attached and metamorphosed were counted. Larvae that did not settle by the end of the 24 h period were observed for signs of abnormal behaviour to assess any compromise to normal physiological function. Fouling resistance was estimated by determining the fraction of individuals settling on each coating.

**Leaching assays for toxic compounds during cyprid settlement**

Coatings were soaked for 6 days in 100 ml of ASW with and without added peroxide (100 μM or 20 μM) prior to settlement assays. The leachate was removed and replaced with 100 ml of seawater at 72 h intervals. The leachate from coatings was used to conduct assays of survivorship with ~200 nauplii larvae of Artemia sp. (brine shrimp). The larvae were exposed to the coating leachate and their survival was monitored for 2 days. Survival of larvae in coating leachate was compared with leachate from a glass slide control.

**Assays for metamorphosis of larvae of H. elegans**

Ten experimental coatings and uncoated glass microscope slides were rinsed with tap water and then placed in running seawater tables at the Kewalo marine laboratory for 7 days to develop a bacterial biofilm. After acquiring a biofilm, coatings were placed in plastic containers containing 19 ml of natural seawater from the Kewalo site, ASW (MBL general purpose seawater, see Cavanaugh 1975), or a 5 μM solution of H$_2$O$_2$ in ASW. One millilitre of ASW containing competent larvae of *H. elegans* was added to each dish. These larvae were allowed to settle and metamorphose, and after ~21 h the number of metamorphosed juveniles was determined for each of the coatings. Larvae were considered metamorphosed if they were permanently attached to the surface of the coatings and had begun to differentiate the branchial crown. The results of the experiment were analysed using a one-way ANOVA with Tukey’s multiple pair-wise comparisons. The percentage of metamorphosed larvae for each replicate was subjected to angular transformation in order to comply with the assumptions of parametric analysis.

**Settlement assay for Ulva zoospores**

Zoospores of *Ulva linza* were released and prepared for the assay as previously described (Callow et al. 1997). The assay followed the principles outlined in Tang et al. (2005): 10 ml aliquots [1.5 × 10$^6$ spore ml$^{-1}$] in ASW with and without 100 μM H$_2$O$_2$] were pipetted into individual compartments of polystyrene Quadrimer culture dishes (Fisher), each containing a test surface. After incubation for 1 h in darkness at ~20°C, all slides were washed in ASW to remove zoospores that had not attached. After fixation, the density of zoospores attached to the surface was counted on each of three replicate slides using an image analysis system attached to a fluorescent microscope. Means ($x = 90$) and 95% confidence limits (two SDs) were calculated and expressed as mean number of attached (settled) spores mm$^{-2}$. Data were analysed by one-way ANOVA followed by a pairwise Tukey test.

**Results**

**Characterisation of xerogel surfaces**

Key to the success of the current platform is the ability of the reagents (halide and H$_2$O$_2$) and products (hypohalous acid) to partition and diffuse within the xerogel film. Transport of small neutral molecules within these types of xerogels has been investigated previously (Tang et al. 2003; Tao et al. 2006). A screening campaign showed that there was significant $I^-$ quenching of [Ru(dpp)$_3$]$^{2+}$ within the 50% APTES/50% TEOS (B5) and the 10% APTES/90% TEOS xerogel (B6) formulations. Figure 1 compares the Stern–Volmer plots for $I^-$ quenching of [Ru(dpp)$_3$]$^{2+}$ in solution, [Ru(dpp)$_3$]$^{2+}$ sequestered within a B5 xerogel film, and [Ru(dpp)$_3$]$^{2+}$ sequestered within a hydrophobic xerogel film prepared from a 50/50 n-propyltriethoxysilane (C3TMOS)/TMOS sol, which has been described as an AF/foul-release surface
The hydrophobic 50/50 C3TMOS/TMOS formulation did not show significant quenching, suggesting that the I cannot partition/diffuse within this film. The I can clearly partition into and diffuse within the B5 xerogel. (Note: Similar behavior is seen in the B6 xerogel.)

The B5 xerogel yielded a very hydrophilic surface with water-contact angles ($\theta_w$) of 34 $\pm$ 1°. The B6 xerogel formulation was less hydrophilic with $\theta_w$ of 54 $\pm$ 1°.

Profilometry showed that the xerogel films were 1.0 $\pm$ 0.1 µm thick. Although such thin films would not likely be a ‘practical’ surface for marine deployment, the uniformity offered by the ‘spin casting method’ provides a model surface that can be reproduced lot to lot.

The class II xerogel (a metal tetraalkoxide and a metal trialkoxide covalently bound; ie TEOS and APTES for the B5 and B6 xerogels of this study) film surfaces are more uniform and uncracked in comparison with xerogels derived from pure TMOS or TEOS (Tang et al. 2003). The incorporation of organic functional groups in the hybrid xerogels reduces crosslinking in the silicate structure from Si(OSi/C17/C17)4 in the pure TMOS or TEOS to RSi(OSi/C17/C17)3 in the class II xerogels leading to a more flexible, less friable surface. SEM analysis of the B5 and B6 xerogels with and without catalysts showed them to be uniform and uncracked at magnifications up to 100,000× with resolution of 200 nm.

XPS spectra of the B6 xerogel film were recorded at a take-off angle of 45° to determine the atomic composition at the surface of the xerogel coating/film after treatment of the surface with ASW and with 0.9 M (3%) H2O2 in ASW. The results are shown in Figure 2.

In glass, the ratio of the C(1s)/Si(2p3) signals is 0.2 (Tang et al. 2005), suggesting a relatively carbon-free surface with any carbon found at the ostensibly SiO2 surface presumably arising from adventitious/adsorbed carbon-containing species from the atmosphere. The ratio of the C(1s)/Si(2p3) signals for the B6 xerogel surface is 1.75 $\pm$ 0.30 for the surface treated with ASW and 2.0 $\pm$ 0.5 for the surface treated with 0.9 M H2O2 in ASW for 24 h prior to XPS analysis. Exposure to H2O2 has little effect on the ratio of the O(1s)/Si(2p3) signals with values of 2.45 $\pm$ 0.20 in the sample exposed to ASW and 2.55 $\pm$ 0.10 in the sample exposed to 0.9 M H2O2 in ASW. The ratio of the N(1s)/Si(2p3) signals was also unchanged by the exposure of the sample to H2O2 with values of 0.20 $\pm$ 0.02 in ASW and 0.20 $\pm$ 0.05 in the sample exposed to 0.9 M H2O2 in ASW. The ratio of the N(1s)/Si(2p3) are well above the background signals of <0.01. These results suggest that carbon- and nitrogen-bearing functionalities are on the class II xerogel film surface.

Figure 3 shows high-resolution XPS spectra of the N(1s) signals of the B6 xerogel from samples exposed to ASW with and without added 0.9 M H2O2. The experimental spectra were resolved into two peaks at 404.5 eV and 402.6 eV in a ratio of 65:35 in ASW without added peroxide and 67:33 in ASW with 0.9 M H2O2. The presence of H2O2 had minimal effect on the type of nitrogen-containing functionality found on the xerogel surface.

Settlement of B. amphitrite cypris larvae

Settlement of cyprids on glass in ASW with 100 µm H2O2 was not statistically different from settlement on
glass in ASW without H$_2$O$_2$ indicating that peroxide did not have any detrimental effect on settlement. Settlement was completely inhibited on the B6, B6Se and B6Te xerogels in the presence of 100 $\mu$M peroxide in ASW. The settlement of *B. amphitrite* cypris larvae on the 50/50 APTES/TEOS (B5, B5Se and B5Te) and 10/90 APTES/TEOS (B6, B6Se and B6Te) hybrid xerogels was significantly lower in comparison with the glass controls in the presence of 100 $\mu$M H$_2$O$_2$ (Figure 4, ANOVA $p < 0.0001$). Settlement was also significantly less in comparison with the glass controls in the absence of 100 $\mu$M peroxide on the B6, B6Se and B6Te xerogels. The B5Te xerogel showed significantly lower settlement ($p < 0.05$) in the presence of 100 $\mu$M H$_2$O$_2$ when compared with ASW alone. Mortality of brine shrimp larvae exposed to the leachates from the 50/50 APTES/TEOS (B5, B5Se and B5Te) and 10/90 APTES/TEOS (B6, B6Se and B6Te) hybrid xerogels in ASW with and without added 0.9 M hydrogen peroxide was not significantly different ($p > 0.05$) in comparison with the mortality of larvae exposed to leachates from the glass controls. The glass controls showed no effect of the 100 $\mu$M H$_2$O$_2$ hydrogen peroxide on the mortality of brine shrimp nauplii larvae.

When the H$_2$O$_2$ concentration was reduced to 20 $\mu$M in ASW, larvae of *B. amphitrite* showed significantly reduced settlement ($p < 0.05$) on B5Se and B5Te xerogels in comparison with glass controls and B5 coatings without catalysts (Figure 5). In the B6 series, both the B6 and B6Te coatings had significantly lower settlement in comparison with glass controls in the presence and absence of H$_2$O$_2$ ($p < 0.05$). Settlement on the B6Se coating was not significantly different in the presence or absence of 20 $\mu$M H$_2$O$_2$ and was not significantly different from the glass controls ($p > 0.05$).

**Settlement of *H. elegans***

At peroxide concentrations of 20 $\mu$M or higher, settlement and metamorphosis of larvae of *H. elegans* were not observed on glass standards. Consequently, the H$_2$O$_2$ concentration in ASW was reduced to 5 $\mu$M. At this concentration there were no significant differences relative to peroxide-free ASW in settlement (Figure 6) or metamorphosis of *H. elegans* on uncoated glass.

A number of coatings performed as AF coatings with respect to settlement of *H. elegans* when compared with uncoated glass slides. Replicate coatings of
B5Se and B5Te performed significantly better (Tukey’s pair-wise comparisons, \( p < 0.05 \)) in comparison with uncoated glass in the ASW only treatments (Figure 6), and settlement on coating B6Te was significantly lower (\( p < 0.05 \)) in comparison with uncoated glass in the treatments with ASW containing 5 \( \mu \text{M} \text{H}_2\text{O}_2 \) (Figure 6).

**Settlement of Ulva zoospores**

*Ulva* zoospore settlement density on glass in ASW was unaffected by the presence of 100 \( \mu \text{M} \text{H}_2\text{O}_2 \). In addition, there was no subsequent effect on viability (germination and growth) by the presence of this concentration of peroxide. The presence or absence of 100 \( \mu \text{M} \text{H}_2\text{O}_2 \) had no effect on the density of settled spores on either of the control xerogel surfaces, B5 or B6 in ASW. None of the leachates from the xerogel films in ASW with or without added peroxide were toxic to *Ulva* zoospores.

Spore settlement was lower on all of the xerogel coatings in comparison with the glass standards (Figure 7). Settlement was lower on the B6 control coatings in comparison with the B5 control coatings. One-way analysis of variance showed significant differences within the xerogel data set (\( p < 0.01 \)). Comparing the samples with and without additional \( \text{H}_2\text{O}_2 \) showed no significant differences except in the case of the B5Te coating.

The inclusion of the Se1 catalyst did not significantly affect the performance of either of the control xerogel coatings (B5 and B6). However, for both coating types, the inclusion of the Te1 catalyst significantly reduced the settlement density of spores. In the case of the B5Te coating, spore density was further reduced by the presence of 100 \( \mu \text{M} \text{H}_2\text{O}_2 \).

**Discussion**

*‘Active’ xerogel surfaces*

The design of ‘active’ xerogel coatings for use as AF surfaces requires (1) the design of catalysts that activate ambient \( \text{H}_2\text{O}_2 \) and that can be covalently sequestered within the xerogel coating and (2) xerogel coatings that are permeable to the reagents necessary for activity. The APTES-containing B5 and B6 xerogels showed strong Stern–Volmer quenching of sequestered \([\text{Ru(dpp)}_3]^{2+}\) by I\(^-\). Values of \( K_{SV} \) were >200 M\(^{-1}\) for these xerogels (xerogel 2 of Figure 1). In contrast, the 50/50 C3TMOS/TMOS xerogel (Tang et al. 2005) gave \( K_{SV} \) of <20 M\(^{-1}\). Values of \( K_{SV} \) for the B5 and

![Figure 6. Effect of 5 \( \mu \text{M} \text{H}_2\text{O}_2 \) on the fraction settlement of *H. elegans*. Bars represent untransformed mean of replicate coatings. Experimental details may be found in the Materials and methods section. Error bars represent 1 SD from the mean of the untransformed data.](image)

![Figure 7. The settlement of *Ulva* zoospores on xerogel coatings in ASW and ASW + 100 \( \mu \text{M} \text{H}_2\text{O}_2 \). Each histogram bar is the mean from 90 counts, 30 from each of three replicate slides. Experimental details may be found in the Materials and methods section. Error bars represent 2 SDs from the mean (95% confidence limits).](image)
B6 xerogels suggest that the B5 and B6 coatings are much more permeable to $I^-$ and presumably other halide salts relative to the hydrophobic 50/50 C3TMOS/TMOS xerogel (Tang et al. 2005). Although the 50/50 C3TMOS/TMOS xerogel might provide better foul release, the lack of permeability to halide ions would limit the effectiveness of this material in ‘active’ coatings. The alcohol functionality of the Te1 and Se1 catalysts allows these molecules to be sequestered covalently within the B5 and B6 xerogels.

Both organotellurides and selenoxides can function as catalysts for the oxidation of halide salts by H$_2$O$_2$. Organotellurides are oxidised by H$_2$O$_2$ to the corresponding telluroxides, which are then hydrated to give the corresponding dihydroxy telluranes A (Figure 8A). The dihydroxy telluranes function as mild oxidants for the oxidation of thiols to disulphides and for the oxidation of halides to positive halogen species (Detty et al. 1994a,b, 1996; You et al. 2003). The overall process can be made catalytic in organotelluride as shown in Figure 8A for the oxidation of halides to hypohalous acids/halogens (Detty et al. 1996; Higgs et al. 2001; Abe et al. 2002; You et al. 2003). Two key intermediates are the hydroxytelluronium ion B and the halotelluronium ion C (Detty et al. 1994b), which can generate hypohalous acid or halogen, respectively, by direct nucleophilic attack of halide as shown in Figure 8A.

Subsequent studies with selenoxides demonstrated that selenoxides are catalysts for the activation of H$_2$O$_2$ for the oxidation of halides to hypohalous acids/halogen and thiols to disulphides (Drake et al. 2003; Goodman and Detty 2004). This process follows a different mechanism than that followed by the tellurides and is illustrated in Figure 8B. Just as telluroxides can add water to give dihydroxy telluranes, selenoxides can add H$_2$O$_2$ to give hydroxy perhydroxy selenanes D. In one path, halide anions can directly displace hydroxide from the perhydroxy group by attacking the OOH oxygen to give hypohalous acid and an R$_2$Se(OH)O$^-$ species, which can lose HO$^-$ to regenerate the selenoxide. In a second pathway, halide attack at the OOH oxygen of the perhydroxy selenane would generate an R$_2$Se(OH)OX species E, which can undergo attack by halide to generate halogen and an R$_2$Se(OH)O$^-$ species, which can lose HO$^-$ to regenerate the selenoxide.

Effects of H$_2$O$_2$ on the physical characteristics of xerogel surfaces

Settlement of _B. amphitrite_ cyprids, _H. elegans_ larvae and _Ulva_ zoospores was reduced relative to glass controls on the catalyst-free B6 xerogel coating in ASW without added H$_2$O$_2$. Settlement of _H. elegans_ larvae and _Ulva_ zoospores was reduced relative to glass controls on the catalyst-free B5 xerogel coating in ASW without added H$_2$O$_2$. The presence of 100 $\mu$M H$_2$O$_2$ in the ASW gave further reduction in settlement in the catalyst-free B5 and B6 xerogels for _B. amphitrite_ cyprids. It might be asked whether the H$_2$O$_2$ is causing a chemical transformation in the xerogel, which further reduces the settlement.

The APTES aminopropyl groups are the organic residues of interest within the B5 and B6 xerogel coatings for two different reasons. First, these residues could organise randomly in the sol particles/xerogels, could lie on the surface of the sol particles/xerogels, or could organise within the sol. Furthermore, interactions with H$_2$O$_2$ could alter the orientation of the aminopropyl groups in the xerogel. Second, the

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**Figure 8.** Catalytic cycles for the oxidation of halides salts with H$_2$O$_2$ using (A) organotelluride catalysts and (B) selenoxide catalysts.
ammonium groups can be protonated in an aqueous environment (pK_a of ~9–10 for a primary ammonium group) or can be oxidised to imine groups using H_2O_2 as an oxidant. As shown in Figure 2, the C(1s)/Si(2p3), O(1s)/Si(2p3) and N(1s)/Si(2p3) ratios were unchanged on the xerogel surfaces exposed to 0.9 M H_2O_2 in ASW relative to those exposed to ASW only. These data suggest that there is no significant surface reorganisation of functional groups containing carbon, oxygen or nitrogen upon exposure to H_2O_2. The high-resolution N(1s) XPS data (Figure 3) showed the presence of two peaks centered at 404.5 eV and 402.6 eV in a 65:35 ratio following exposure to ASW and in a nearly identical 67:33 ratio following exposure to ASW with 0.9 M H_2O_2. These data indicate that the oxidising environment has little effect on the ratio of the two peaks and suggest that oxidation of the amines is not occurring to a significant degree. A more likely scenario is that the two peaks represent primary ammonium (404.5 eV) and primary amino (402.6 eV) groups from the APTES residues on the xerogel surface. The 1.9 eV separation between the two N(1s) peaks is similar to the 2.5 eV separation observed between the N(1s) peaks for a protonated (402.0 eV) and unprotonated (399.5 eV) dimethylamino group in protonated 1,8-bis(dimethylamino)-naphthalene (DMAN-HBr) (Bruckner et al. 1999) (Chart 2). Unprotonated DMAN shows a single N(1s) peak at 399.5 eV.

**Chart 2.** Chemical structures of protonated (DMAN-HBr) and unprotonated (DMAN) 1,8-bis(dimethylamino) naphthalene.

Settlement on xerogel surfaces in the absence of H_2O_2

Ammonium groups from APTES residues on the xerogel surface might serve to discourage the settlement of various organisms even in the absence of H_2O_2. Ammonium compounds have been incorporated in polymers and in marine paints and have been shown to give reduced fouling (Baudrion et al. 2000; Cowie et al. 2006). Settlement on the B6 xerogels (B6, B6Se and B6Te) in the absence of peroxide was significantly less than on glass controls for both cypris larvae of *B. amphitrite* (Figures 4 and 5) and for *Ulva* zoospores (Figure 7).

In the absence of peroxide, both the B5Se and B5Te coatings showed reduced settlement (p < 0.05) of *H. elegans* relative to glass controls in ASW whereas the catalyst-free B5 xerogel did not (Figure 6). Prior to settlement studies with *H. elegans*, all of the xerogel coatings were left in natural seawater for 7 days to establish a biofilm on the surfaces prior to assay. Although the biofilm was not characterised, it is possible that peroxide-producing organisms (Chandrasekaran and Dexter 1993; Le Bozec et al. 2001; Dexter et al. 2003) might activate the B5Se and B5Te hybrid xerogels by producing hydrogen in the biofilm to discourage settlement in the ASW-only experiments whereas the catalyst-free B5 control would not be able to utilise the H_2O_2. Although the subsequent addition of 5 μM peroxide had no statistically significant effect on settlement in these coatings, local concentrations of peroxide from the biofilm could be significantly higher (Chandrasekaran and Dexter 1993; Le Bozec et al. 2001; Dexter et al. 2003).

**Effect of H_2O_2 on settlement onto xerogel surfaces**

With respect to the settlement of cypsis larvae of *B. amphitrite*, the presence of 100 μM H_2O_2 reduced settlement of both catalyst-free and catalyst-containing xerogels below that of glass controls (Figure 4). In the B5Te coating, settlement in the presence of 100 μM H_2O_2 was significantly less than settlement on the same surface in ASW without peroxide. The B6 series of coatings was surprisingly efficient at discouraging settlement in the absence and presence of 100 μM H_2O_2 with settlement on all surfaces being <20% of glass controls. In the presence of the peroxide, no settlement was observed on B6, B6Se and B6Te xerogels. The reduced settlement did not appear to be because of any adverse physiological effect upon the barnacle larvae: cyprids were active and behaving normally on all coatings for the duration of the test. This observation is further supported by the lack of mortality among brine shrimp nauplii larvae exposed to the leachates from the two series of B5 and B6 xerogel coatings in the presence or absence of 100 μM H_2O_2. Toxic substances are not leaching from the xerogel surfaces into bulk solution.

When the concentration of H_2O_2 was reduced to 20 μM in ASW, the difference between ASW only and ASW plus peroxide exposures was less pronounced (Figure 5). However, the B5Se and B5Te coatings in the presence of peroxide showed significantly reduced settlement in comparison with the B5 coating without catalyst and relative to glass controls. In the B6 series, the B6Te xerogel still showed no settlement by cypsis larvae of *B. amphitrite* in the presence of 20 μM H_2O_2, which is indicative of a highly inhibitory surface.
The settlement of *H. elegans* on the B6Te xerogel was also significantly impacted by the presence of H$_2$O$_2$ (Figure 6). In the presence of 5 μM H$_2$O$_2$, the trend shows that settlement was lower on the B6, B6Se and B6Te coatings in comparison with glass controls although only the reduced settlement on the B6Te coating in the presence of 5 μM H$_2$O$_2$ is statistically significant (*p* < 0.05). In the B5 series of xerogels, the catalyst had no significant impact on settlement relative to the B5 control in the presence of 5 μM H$_2$O$_2$.

The settlement of *Ulva* zoospores on all of the xerogel surfaces was significantly lower than on the glass standard in both the absence and presence of 100 μM H$_2$O$_2$ in ASW. However, only the B5Te xerogel in the presence of 100 μM H$_2$O$_2$ in ASW gave significantly reduced settlement (*p* < 0.05) of *Ulva* zoospores relative to settlement on the B5Te xerogel in ASW in the absence of peroxide (Figure 7).

Surface energy and surface wettability are important characteristics for settlement cues and fouling release (Rittschof and Costlow 1989; Baier and Meyer 1992; Gerhart et al. 1992; Rittschof et al. 1998; Genzer and Efimenko 2006). With the xerogels, the present authors have found that lower-energy surfaces (or surfaces with a higher water contact angle) give reduced settlement as well as increased removal of various fouling organisms (Tang et al. 2005). The APTES-containing xerogels are made more ‘hydrophobic’ by reducing the amount of APTES in the formulation. The 10% APTES/90% TEOS (B6) xerogel formulations have values of $\theta_w$ of 54° in comparison with 34° observed for the B5 coatings. The B6 coatings (B6, B6Se and B6Te) gave reduced settlement of *B. amphitrite* cyprids and *Ulva* zoospores in the absence of H$_2$O$_2$ in comparison with the B5 xerogels (B5, B5Se and B5Te) and glass controls.

Algae including *Ulva* produce reactive oxygen species including H$_2$O$_2$ as by-products of the photosynthetic electron transport system (Ross and van Alstyne 2007). Although H$_2$O$_2$ is used as an algicide (Drabkova 2007), its efficacy depends on the algal species and the ambient conditions. Green algae appear to be relatively resistant and are reportedly unaffected by concentrations found in surface waters, which is compared with the concentration used in the present experiments. In addition, green algae appear to be relatively resistant to the oxidised halide species (hypohalous acids) that are generated through the action of the catalyst. Concentration >100 mg l$^{-1}$ (0.001 M) of bleach were needed to kill fragments of the green alga *Caulerpa* (Williams and Schroeder 2004) and, as for H$_2$O$_2$, efficacy can be reduced by environmental conditions eg the presence of organics in the water.

**Conclusions**

Overall, the results indicate that hybrid class II xerogels can be used to sequester catalysts for the activation of H$_2$O$_2$ in ASW. When the coatings are ‘active’ in the presence of H$_2$O$_2$ from a biofilm either on the surface or in the surrounding seawater, reduced settlement of cypris larvae of *B. amphitrite*, larvae of *H. elegans* and zoospores of *Ulva* have been observed using the Te1 catalyst relative to glass and catalyst-free xerogel controls. The Se1 selenoxide catalyst also gave significantly reduced settlement of cypris larvae of *B. amphitrite* in ASW in the presence of 20 μM H$_2$O$_2$ when sequestered in the B5Se coating in comparison with the catalyst-free B5 coating and glass controls. Presumably, the sequestered catalysts facilitate the oxidation of halide salts to positive halogen species with H$_2$O$_2$. The ‘active’ surfaces in this study may present localised concentrations of hypohalous acids that discourage settlement of fouling organisms including cypris larvae of *B. amphitrite*, larvae of *H. elegans* and *Ulva* zoospores.

The various parameters of these coatings have not yet been optimised, but the coatings reported here illustrate the concept of ‘active’ hybrid xerogels with sequestered catalysts for the activation of H$_2$O$_2$. The B5 and B6 xerogels were selected more for halide permeability rather than for foul-release properties. In these two xerogels, the telluride (Te1) catalyst appeared to be more effective than the selenoxide (Se1) catalyst with statistically significant reductions in settlement observed with the Te1 catalyst with cyprids of *B. amphitrite*, larvae of *H. elegans* and *Ulva* zoospores in H$_2$O$_2$-containing ASW relative to catalyst-free xerogel and glass controls. Other hybrid xerogels may have relatively low surface energies, high water contact angles and permeability to both peroxide and halide salts, making them suitable alternatives to the present coatings. The loading level of catalyst has yet to be optimised. Higher loading levels relative to the 0.015 M concentrations used in the sols of this study may provide greatly improved AF properties.

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**References**


