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A New Pathogen Transmission Mechanism in the Ocean: The Case of Sea Otter Exposure to the Land-Parasite *Toxoplasma gondii*

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**Abstract**

*Toxoplasma gondii* is a land-derived parasite that infects humans and marine mammals. Infections are a significant cause of mortality for endangered southern sea otters (*Enhydra lutris nereis*), but the transmission mechanism is poorly understood. Otter exposure to *T. gondii* has been linked to the consumption of marine turban snails in kelp (*Macrocystis pyrifera*) forests. It is unknown how turban snails acquire oocysts, as snails scrape food particles attached to surfaces, whereas *T. gondii* oocysts enter kelp beds as suspended particles via runoff. We hypothesized that waterborne *T. gondii* oocysts attach to kelp surfaces when encountering exopolymer substances (EPS) forming the sticky matrix of biofilms on kelp, and thus become available to snails. Results of a dietary composition analysis of field-collected snails and of kelp biofilm indicate that snails graze the dense kelp-biofilm assemblage composed of pennate diatoms and bacteria inserted within the EPS gel-like matrix. To test whether oocysts attach to kelp blades via EPS, we designed a laboratory experiment simulating the kelp forest canopy in tanks spiked with *T. gondii* surrogate microspheres and controlled for EPS and transparent exopolymer particles (TEP - the particulate form of EPS). On average, 19% and 31% of surrogates were detected attached to kelp surfaces covered with EPS in unfiltered and filtered seawater treatments, respectively. The presence of TEP in the seawater did not increase surrogate attachment. These findings support a novel transport mechanism of *T. gondii* oocysts: as oocysts enter the kelp forest canopy, a portion adheres to the sticky kelp biofilms. Snails grazing this biofilm encounter oocysts as ‘bycatch’ and thereby deliver the parasite to sea otters that prey upon snails. This novel mechanism can have health implications beyond *T. gondii* and otters, as a similar route of pathogen transmission may be implicated with other waterborne pathogens to marine wildlife and humans consuming biofilm-feeding invertebrates.

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**Introduction**

*Toxoplasma gondii* is a coccidian parasite that infects humans and warm-blooded animals [1]. Infections with this terrestrial parasite have also been documented in marine mammals [2], including the southern sea otter (*Enhydra lutris nereis*). The southern sea otter is an endangered subspecies that inhabits kelp forests, including those of *Macrocystis pyrifera*, along the California coast. *Toxoplasma gondii* is recognized as a significant cause of mortality in southern sea otters, with harmful consequences for the health and recovery of this population [3]. Infected otters have been detected at several locations along the coast of California, and high-risk sites for otter exposure were described in populations from Morro Bay and Cambria, California [4,5].

The transmission mechanism of *T. gondii* in the marine food web is unknown. Sea otter exposure to this parasite is thought to occur through ingestion of oocysts that are shed by felids [6,7]. Cats can shed hundreds of millions of oocysts in their feces when infected, and only 1–10 oocysts are needed to successfully infect mammals [8,9,10]. *Toxoplasma gondii* oocysts may reach coastal waters via contaminated freshwater runoff [4], and wetland degradation may increase flux of oocysts by more than 2 orders of magnitude [11]. Oocysts introduced into coastal waters may survive for at least 24 months [12], and may adhere to aquatic aggregates or occur freely in the water column [13].

Marine invertebrates (e.g. mussels and oysters) and vertebrates (e.g. anchovies and sardines) that feed by filtering seawater through their gills may acquire *T. gondii* oocysts from seawater [14–17]. These animals may then serve as potential sources of infection of *T. gondii* to marine mammals. However, southern sea otters are diet specialists and a recent epidemiological study showed that otters that specialize on marine turban snails (*Chlorostoma brunnea*, *C. montereyi*, and *Promartynia pulligo* – formally assigned to the genus *Tegula*) in the kelp forest are 12 times more likely to acquire *T. gondii* than those consuming other prey [3]. As opposed to mussels, oysters, sardines and anchovies, which feed by filtering food particles out of the water, turban snails scrape surfaces using a radula to ingest food particles that are attached to a substrate. Thus, the mechanism by which the benthic feeding marine turban snails acquire oocysts suspended in the water is puzzling and suggests an alternate mode of feeding on small planktonic particles.
Because of their feeding strategy, turban snails are likely to consume biofilm-associated organisms that colonize kelp surfaces [18–20]. Biofilms are defined as mixed assemblages of microbes enclosed in a matrix, adhering to each other and/or to surfaces [21]. Extracellular polymeric substances (EPS) form the matrix of biofilms and are defined as a complex of high molecular weight macromolecules, mainly polysaccharides, localized outside the cell wall in the form of gels, slime or capsules [22–24]. Because of its adhesive properties, EPS is regarded as the 'biological glue' that anchors biofilm microorganisms (e.g., bacteria and microalgae) to a surface [25]. Several studies have addressed the role of EPS in the attachment of bacteria and diatoms to surfaces such as macroalgae, ice and sediments [26–28]. Microorganisms such as benthic diatoms and fungi have been observed on the surface of Macrocystis pyrifera [20,29,30]. These organisms may be trapped in a sticky EPS matrix that possibly helps them attach to the surface of the kelp blades [31–33].

A secondary source of mucilage-rich materials that may form biofilms on kelp includes waterborne TEP (transparent exopolymer particles) - a particulate form of EPS in the water column [34]. Major producers of TEP in the water column include bacteria and phytoplankton, and TEP is regarded as a major biofilm agent [35]. When water upstream of a kelp bed contains TEP, the TEP may make contact with kelp surfaces as it flows through the kelp bed, subsequently adhering directly to the kelp or to EPS-coated surfaces of the kelp. A recent study documented this phenomenon, showing colonization of TEP by bacteria and microalgae suspended in the water, and subsequently adhering to glass surfaces rapidly (e.g. 30 min), promoting biofilm development [36].

Biofilms occur on a wide range of surfaces (natural or man-made) and EPS is also a major component of diatom films found on underwater man-made surfaces [37]. The removal of pathogens from water onto biofilms that colonize the surfaces of pipes or other structures of water treatment plants has been reported [38]. Investigating the potential for T. gondii to adhere to biofilms is significant not only for understanding infection mechanisms of marine fauna, but also due to numerous reports of waterborne outbreaks of toxoplasmosis in humans worldwide [39]. To date, the adhesion of coccidian parasites’ oocysts has not been linked to EPS of biofilms that colonize natural surfaces such as kelp blades.

In this study, our major goals were to determine whether EPS on kelp surfaces and/or TEP in the water promote adhesion of T. gondii oocysts to kelp in a central California forest of Macrocystis pyrifera. We also investigated, as a secondary goal, the similarity of epiphytic organisms within the kelp blade biofilm to the organisms (or their remains) present within the feces of snails that had been feeding on those same blades. Here we hypothesized that once T. gondii are trapped in the kelp biofilm via EPS or TEP, the parasite is available for ingestion by marine turban snails and other benthic-scrapping organisms. This hypothesis could help explain why sea otters that have specialized diets on marine turban snails are more likely to be infected with T. gondii than those consuming other prey.

Methods

Two experiments were designed to: (1) provide qualitative data on snail diet and kelp biofilm composition; and (2) test whether T. gondii oocyst surrogates adhere to kelp blade surfaces via EPS on the blades and/or via TEP suspended in the water that subsequently become biofilm on the blades. Together these experiments examine the possible mechanism(s) by which T. gondii oocysts may become associated with kelp surfaces and whether snails can consume organisms entrained within the kelp biofilm: if such associations are found, then a delivery route of the planktonic oocysts to the endangered otter would be identified for the first time. We considered both EPS (the matrix of the biofilm on kelp surfaces) and TEP in the water, as potential agents that could deliver Toxoplasma to snails that graze the kelp surfaces with biofilms. Dragon green microspheres that have been previously validated as surrogate particles for T. gondii oocysts were used because they have surface properties (i.e., size, specific gravity, hydrophobicity, and surface charge) that resemble those of T. gondii oocysts [40]. Due to the biohazardous nature of T. gondii oocysts, employing surrogates in mesocosm experiments provides an alternative approach for evaluating the waterborne transport of this zoonotic pathogen. Previous studies have successfully applied these surrogates to demonstrate waterborne transport of oocysts [11] and their association with macroaggregates [13].

Experiment 1: Turban snail diet and kelp biofilm composition

On Aug 8 and 24, 2011, 30 sexually mature (shell>1.5 cm) and juvenile (shell <1.5 cm) turban snails (Chlorostoma spp.), along with the kelp blades they were associated with, were collected by hand onboard the R/V Sebastes from the canopy of kelp beds in Carmel (36°30′57″N and 121°57′16″W), approximately 44 km south of the Santa Cruz site that provided material for Experiment 2. Snails and kelp were collected under the permit ID 12119 issued by California Department of Fish and Game. Snails were transported live to the laboratory and placed in 1 L jars with 0.2 μm filtered seawater and associated kelp blades, with one snail and 1 frond placed in each jar and incubated for 16 hrs. Temperature at 15°C and 12 hrs light cycle were maintained. The snail fecal pellets produced in the jars were then recovered, their associated blade stored for later analysis of its biofilm, and pellets stored for mounting on glass slides. The biofilm present on the kelp blade from each snail-grazing container was removed using a PTFE spatula, while submerged in 0.2 μm filtered seawater. Bacteria were visualized by staining biofilm samples with 4′,6-diamidino-2-phenylindole (DAPI) (Pierce Biotechnology Inc., Rockford, IL, USA) (final concentration of 500 mg mL\(^{-1}\)). Snail pellet content and biofilm organisms from kelp blades on which they had grazed were examined using a Zeiss Axio Imager with phase contrast and a 50 W light source fitted with 2 DAPI bandpass filter sets (wavelength excitation (λem) 350 nm, (wavelength emission) λex >420 nm and λex 350 nm, λem >460 nm).

To visualize the underlying EPS matrix of the kelp biofilm, cross-sections of the kelp blades were stained using 500 μL of 4′,6-diamidino-2-phenylindole (DAPI) (Pierce Biotechnology Inc., Rockford, IL, USA) (final concentration of 500 mg mL\(^{-1}\)) with 0.02% aqueous alcian blue solution (8GX) in 0.06% of acetic acid (pH = 2.5) (ABS). EPS was visualized under bright field on the microscope described above. All micrographs were obtained using an Axio Cam HRc camera system.

Experiment 2: T. gondii surrogate adhesion to kelp blades via EPS

Seawater and kelp blades used in this laboratory experiment were collected at 0700 on July 11, 2012 from the surface of a kelp bed canopy in Santa Cruz (36°56′57″N 122°02′05″W) on board the R/V Sebastes. Six kelp blades of similar size (Mean surface area 242 cm\(^2\)±54) and ridged texture, and free of obvious macroscopic epibionts were collected from the same stipe. Surface seawater from the kelp canopy was collected with a bucket. Samples were transported to the laboratory in a cooler with ice and the
experiment was initiated immediately. Artificial kelp blades (BioModels Co., Aguanga, CA 92536) were also used in the experiments, as they had not previously been submerged in seawater, and thus provided a control substrate free of biofilm and EPS.

Four treatments were used to test whether surrogates of *T. gondii* oocysts would become associated with the EPS matrix on the surface of the field-collected kelp blades. Each treatment included 3 replicates in 4 L glass jars. All jars were pre-washed with 10% HCl. Treatment A was designed to reproduce the kelp forest canopy environment and consisted of jars with a kelp blade and unfiltered surface seawater; treatment B consisted of jars with a kelp blade and 0.2 µm filtered seawater. Filtered seawater (0.2 µm) was used to remove any TEP suspended in the water that may have been produced by phytoplankton or bacteria. Treatment C was designed as a control treatment for TEP and EPS by using 0.2 µm filtered seawater and a synthetic kelp blade, and treatment D was designed as a control for the loss of surrogates due to settling or attachment to surfaces of the jar and utilized 0.2 µm filtered seawater without kelp blades (Fig.1).

The 12 hr-controlled experiment was performed on a stir table, with paddles in each treatment jar stirring the water at 30 rpm. Water temperature was held at 12°C, and the experiment was conducted during a light cycle. At the start of the experiment, *T. gondii* surrogate microspheres labeled with a dragon green
fluorochrome (Bangs Laboratory, FC07F 5493) were added to each replicate to achieve a final concentration of 20 per mL.

**Quantifying TEP, surrogates, and chlorophyll a in water**

Aliquots of 600 mL were collected gently from each treatment at 3 time points (0, 6 and 12 hrs) to quantify the naturally occurring TEP, chlorophyll a concentrations, and spiked surrogates in the seawater. Aliquots of 550 mL for TEP and surrogate quantification were preserved with formaldehyde to achieve a 1% final concentration, and then stored at 4°C in the dark for later analysis. TEP was measured by filtering 3 replicates of 100–200 mL aliquots onto 0.4 μm polycarbonate filters. TEP concentrations were determined using a standard semiquantitative colorimetric assay [41].

Surrogates were quantified by filtering 3 replicates of 50 mL onto 0.4 μm polycarbonate filters [42]. Filters were mounted on slides and surrogates were enumerated using a Zeiss Axio Imager fitted with a FITC band pass filter set (λex 460–500 nm, λem 510–560 nm) and a 50 W light source.

Chlorophyll a was measured to verify that phytoplankton cells were absent from control treatments B, C and D, but present in treatment A. Two replicates of 25 mL were filtered onto GF/F filters and kept in a -20°C freezer. Chlorophyll a was extracted for 24 hrs in 90% acetone and subsequently analyzed on a Turner Design 10AU fluorometer [43].

**Quantifying EPS, surrogates, and the benthic diatom community on kelp blades**

At the end of the experiment, each kelp blade was removed from its jar and placed in a preservative solution consisting of 550 mL 0.2 μm filtered seawater solution and formaldehyde at 1% final concentration. Kelp blades were scraped on a glass tray while submerged in its preservative solution. Each kelp blade was photographed with a Nikon digital camera and kelp surface area was measured using the Image J image-analysis software (W.S. Rasband, Image J, U.S. national Institute of Health, http://rsb.info.nih.gov/ij/). The surfaces of kelp blades were scraped on both sides using a spatula to remove the biofilm (i.e., EPS and microorganisms) and *T. gondii* surrogates, taking care not to remove kelp tissue cells. A small PTFE spatula was used to access material on the ridges of the kelp blade. Immediately after scraping was completed, the blade was removed from the tray and the solution with the scraped material from the kelp blade was stored in 1 L glass jars in the dark at 4°C. This solution is subsequently referred to as ‘kelp extract solution’.

To quantify EPS on kelp blades, we adapted the semiquantitative colorimetric assay to quantify TEP. Both TEP and EPS may be measured with this assay [28,44]. The principle of the method lies in the staining of extracellular polysaccharides with ABS. Alcian blue has been used to stain extracellular polysaccharides in colony matrices or capsules of algae and bacteria [28,44,45]. This stain complexes carboxyl (-COO⁻) and half-ester sulfate (OSO₃⁻) reactive groups of acidic polysaccharides, the main components of EPS, allowing these otherwise transparent substances to be visualized and quantified by measuring its maximum absorbance on a spectrophotometer set at 787 nm [46].

Aliquots of 15 mL from the kelp extract solution (3 replicates per jar) were filtered onto 0.4 μm polycarbonate filters with low, constant vacuum pressure (≤150 mm of Hg). Filters were stained for 2 seconds with 500 μl of ABS, rinsed with distilled water to remove excess dye, and stored in 15 mL centrifuge tubes in a
—20°C freezer for up to 2 weeks. Extraction of polysaccharides from filters was done with 5 mL of 80% H₂SO₄ for 2 hrs and absorbance of alcian blue was measured on a UV-1201 UV-VIS spectrophotometer set at 787 nm. A calibration curve using gum xantham as the standard was constructed to generate a conversion factor (F-factor) to relate the absorbance of stained EPS to the weight of EPS [40]. Final EPS concentration is reported as μg of gum xantham equivalents per cm² of kelp blade surface by adapting the following equation from [41]:

$$\text{EPS per cm}^2 \text{of kelp blade} = \frac{(\text{absorbance} - \text{blank}) \times \text{volume filtered}}{\text{cm}^2 \text{of kelp blade area}} \times F\text{-factor}$$

To quantify surrogates on kelp blades and visualize the epiphytic diatoms within the EPS matrix, 3 replicates of 50 mL of the kelp blade extract solution were filtered onto 0.4 μm polycarbonate filters and stained for 2 sec with 500 μl ABS (see above). Filters were mounted on slides and observed with a Zeiss Axio Imager. Toxoplasma surrogates were enumerated as mentioned above. EPS was visualized with bright field and epiphytic diatoms with a chlorophyll filter set (λex 440–470 nm, λem >515 nm) and a 50 W light source.

To enumerate diatoms associated with the kelp blade, aliquots of 10 mL of kelp extract solution were settled for 24 hrs in an Utermöhl chamber following the Utermöhl method [47]. A minimum of 100 cells (benthic or planktonic diatoms genera) was counted per chamber using an inverted microscope (Olympus IMT-2).

Statistical analysis
Biostat 3.0 was used for all statistical comparisons. Mann-Whitney (or U test) was used to test whether the concentration of T. gondii oocysts surrogates decreased between two time points (6 hrs and 12 hrs) in all treatments. Mann-Whitney was also used to test whether the percentage of surrogates suspended in water and associate to kelp blades differed at the end of the experiment in each treatment. Kruskal-Wallis (or H test) tested whether the proportion of T. gondii oocysts surrogates associated with kelp blades differed among the treatment that mimicked the kelp forest canopy (A), the treatment that controlled for TEP (B) and the control treatment (C).

Results
Experiment 1: Turban snail diet and kelp biofilm composition
Turban snail (Chlorostoma spp.) fecal pellets produced over the 16-hour grazing incubation with associated kelp blades contained microorganisms similar to those of the biofilm community on the individual blade on which they had grazed. Benthic diatoms resembling Cocconeis spp. were dominant in snail pellets (Fig. 2A) and on the surface of kelp blades (Fig. 2B). Bacteria were also observed associated with the Cocconeis spp. in the kelp biofilm (Fig. 2C). Other benthic diatoms, including species within the genus Licmophora, and possibly Nastatul, were observed as part of the kelp biofilm (Fig. 2D) and ‘trapped’ in the EPS fibers (Fig. 3A, B). These genera were less abundant relative to Cocconeis and not observed in the snail’s fecal pellets.

Experiment 2: T. gondii surrogate adhesion to kelp blades via EPS
During the experiment (between 6 and 12 hrs), the number of T. gondii surrogates suspended in the water decreased significantly in all treatments (Fig. 4, Table 1). Treatment D showed the inherent loss of particles in the water column through the duration of the experiment (Fig. 4).

The percentage of surrogates attached to the kelp blades was calculated relative to the total number of surrogates in the jar (i.e., surrogates in the water + surrogates on the blades) at the end of the experiment (t = 12 hrs). After 12 hrs, the percentage of surrogates associated with kelp blade and suspended in the water significantly differed (Table 2). In the treatment that simulated the kelp forest environment (treatment A), 19% (±3.5) of T. gondii oocyst surrogates were found attached to the surfaces of the kelp (Fig. 5), with EPS being detected on the surface of the kelp blade (Fig. 6) and TEP and chlorophyll a measured in the water (Fig. 7). The highest percentage of surrogates (31%±10%) was attached to the surface of the kelp blades from treatment B (Fig. 5). In this treatment, kelp blades were covered with EPS (Fig. 6), but TEP...
and chlorophyll a were not detected in the water (Fig. 7). The lowest percentage of surrogates attached to a kelp-like surface was found in the control treatment C that housed the synthetic kelp blades and filtered seawater. In this treatment, there was no measurable TEP in the water nor EPS on the kelp blade surface (Fig. 5, 6 and 7).

Kruskal-Wallis results showed that the percentage of surrogates attached to kelp blades significantly differed between treatments with and without EPS covered blades (A and C (p < 0.05) and treatment B and C (p < 0.05)), but not between treatments A and B (p > 0.05), where real kelp blades covered with EPS were used.

The concentration of diatoms on kelp blades in samples from treatment A and B, i.e. blades incubated in unfiltered and filtered sea water, respectively, was estimated at 847 (± 138) cells per cm² and 505 (± 373) cells per cm², respectively. After 12 hrs, in treatment A, 4.2% (± 2.1) of the total number of cells associated with the kelp blade surface were diatoms from genera typically observed in the plankton (Asterionella, Chaetoceros, Cylindrotheca, Ditylum, Eucampia, Hemiaulus, Leptocylindrus, Pseudo-nitzschia, Skeletonema) (Fig. 8). The remainder included genera of several benthic diatoms. The dominant benthic genera included: Navicula, Cocconeis, Licmophora and Tabularia. No diatoms were detected on the synthetic kelp blades housed in 0.2 μm filtered seawater in treatment C.

**Table 1.** Mann-Whitney (or Test U) results shows the significant decrease in the percentage of surrogates of *T. gondii* measured in the water at t = 6 hrs and t = 12 hrs in all treatments.

| Treatment A | Treatment B | Treatment C | Treatment D |
|-------------|-------------|-------------|-------------|
| U           | 5.00        | 11          | 2           | 1           |
| p           | < 0.05      | < 0.05      | > 0.05      | > 0.05      |
| N           | 9           | 9           | 6           | 6           |

The goal of the present study was to test whether extracellular polymer substances (EPS), which form the matrix of biofilms colonizing the surfaces of *M. pyrifera* blades, play a role in the transmission of the protozoan parasite *T. gondii* to benthic feeding turban snails. Marine turban snails have been implicated in the exposure of southern sea otters to *T. gondii* [5]. As this parasite has been identified as a significant cause of mortality in endangered southern sea otter populations, it is critically important to understand the transmission mechanisms to otters, so that prevention or management strategies can be developed to reduce likelihood of exposure. Here we show that *T. gondii* surrogates may adhere to the biofilm that colonizes the surfaces of kelp blades, thereby becoming available to turban snails that feed upon organisms associated with this biofilm.

**Discussion**

The surfaces of *Macroystis pyrifera* blades in the kelp forest canopy were covered with a biofilm composed of benthic diatoms and bacteria embedded in a gel-like matrix. Microscopic observations using the alcian blue stain and quantitative measurements using the colorimetric assay from the 12 hr experiment indicate that the underlying gel-like matrix of this biofilm on the blades is composed of EPS. EPS on kelp blades possibly originates from bacteria and photosynthetic organisms present in the biofilm (i.e., pennate diatoms) and by *M. pyrifera* itself, since EPS production has been linked to photosynthesis and the presence of bacteria [48–50]. Other microorganisms (e.g., fungi) may also contribute to the kelp biofilm production system and thereby add to the EPS pool [33]. Locally, we have noted such epizootic populations in our coastal kelp forest communities [20].

The high abundance of benthic diatom frustules in fecal pellets of *Chlorostoma* spp. confirms that these subtidal snails feed upon organisms (i.e., pennate diatoms) in kelp blade biofilms. It is likely that the snails are also ingesting EPS, given their mode of food capture: grazing surfaces with a chitinous radula. Sediment dwelling animals also have been shown to ingest EPS attached to sediment particles [51]. The benefits of ingesting EPS may...
include the fact that EPS are rich in organic carbon and that it may adsorb dissolved organic matter (DOM) providing an important pool of C and N [25]. Results from Experiment 2 indicate that surrogates of \textit{T. gondii} oocysts might be entrapped in the kelp blade biofilm via EPS (Fig. 5 and 9). Therefore, we expect that once oocysts are attached to the kelp blade biofilm they can be consumed as a ‘bycatch’ item by turban snails. Indeed, preliminary findings of a study parallel to ours indicate that turban snails may acquire surrogates of \textit{T. gondii} and \textit{T. gondii} oocysts while they were kept in tanks with kelp blades exposed to surrogates and oocysts [52].

In addition to grazing benthic diatoms that are present within kelp biofilms, \textit{Chlorostoma brunnea} has been suggested to serve as a ‘farmer’ of epizooic fungi that grow on giant kelp surfaces: by wounding the kelp blade with their radula, snails promote fungal infection on the blade. Snails then consume the resultant fungi, with the infection being controlled to maintain an optimal growth rate of the fungus [20]. The presence of such fungal populations on giant kelp has been observed in the field study location, along with their associated microbial communities [20]. Fungi may also produce a large amount of EPS [33] and thus enhance biofilm formation. Although we could not confirm the presence of fungal-

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**Figure 5.** Percentages (mean ± SD) of \textit{T. gondii} surrogates suspended in water and present in kelp scrapings at the termination of Experiment 2 (12 hrs) (N = 9 per treatment).

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**Figure 6.** EPS concentrations (mean ± SD) on the kelp blades’ surface at the end of Experiment 2 (t = 12 hrs) (N = 9 per treatment).

Nd = not detected.

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like structures in the snail pellets or within the kelp biofilm in our specimens, *T. gondii*’s (and other pelagic particles) adhesion to the kelp surfaces and subsequent transmission to snails may be enhanced if fungi are part of the kelp blade biofilm.

**T. gondii** surrogate adhesion to kelp blades via EPS

Experiment 2 demonstrated a possible mechanism whereby oocysts of *T. gondii* come to coat the surfaces of kelp blades that are covered with EPS and colonized by benthic diatoms and other microorganisms (Fig. 9). At the end of the 12 hr experiment, an average of 19% and 31% of the total number of surrogates of *T. gondii* oocysts were recovered from kelp blades coated with EPS and their natural associated microorganisms in unfiltered and filtered seawater treatments, respectively. These results suggest a novel mechanism by which contaminated runoff entering the coastal ocean can deliver *T. gondii* oocysts to downstream kelp beds, with some of the oocysts adhering to kelp surfaces covered with EPS.

The mechanism of parasite adhesion to surfaces of kelp covered by EPS is likely related to hydrophobic and electrostatic attractive forces, which are influenced by the composition of the oocyst outer wall [53]. The *T. gondii* oocyst cell wall consists of a matrix of polymeric substances, mostly proteins including cysteine-rich proteins among others [54]. The cysteine-rich proteins of *T. gondii* oocyst are related to those of the walls of another coccidian parasite, *Cryptosporidium* oocyst, [55]. Interestingly, the adhesion of *Cryptosporidium* oocysts to biofilms on man-made surfaces has been verified and the roughness of the biofilm has been strongly correlated with oocyst retention [56–58]. *Giardia* cysts, which are also covered with a polymeric matrix, have also been documented to attach to biofilms [59]. Thus, our discovery that surrogates of *T. gondii* oocysts attach to biofilm that covers the surfaces of kelp is consistent with previous findings that environmentally resistant parasites can adhere to surfaces covered with sticky biofilms.

Additional evidence for a mechanism promoting adhesion of suspended particles to surfaces covered with EPS was provided by our observation that planktonic diatoms were present on surfaces of kelp blades. Planktonic diatoms, likely present in the unfiltered seawater at the start of our experiment, presumably were the source of the diatoms that we observed attached to the kelp surfaces. Thus, our study provides evidence of adhesion of planktonic diatoms and suspended particles such as *T. gondii* oocysts to surfaces covered with an EPS biofilm.

The mechanism whereby EPS serves as an adhesive may have implications in other fields of research. Biofilms occur in a variety of environments (surfaces of rocks, plants, sediments, ship hulls or wastewater treatment plants). Zoonotic pathogens other than *T. gondii* that can infect both humans and marine animals have been detected in the coastal environment. For example, (oo)cysts of *Cryptosporidium* and *Giardia* as well as enteric bacteria have been documented in the coastal ocean and found to infect marine fauna.
Although the route of infection of marine wildlife by these parasites is still unclear, it is possible that these pathogens also may become associated, via EPS, with biofilms that coat natural or man-made surfaces in the ocean. The novelty provided by these results is a proposed mechanism by which land-derived pathogens in contaminated runoff are transferred from the water column to a benthic environment, thus facilitating the exposure of benthic-feeding marine fauna to T. gondii and other pathogens.

It is unclear whether the presence of TEP in the seawater influenced the adherence of oocyst surrogates to kelp blades that had EPS coatings. TEP can be regarded as part of the particle pool of EPS [63], with both TEP and EPS containing acidic polysaccharides and possessing adhesive properties [31,64]. In a kelp forest environment, TEP may be detected in the water ([65], pers. obs.) and may serve as an agent to deliver oocysts to EPS-covered kelp blades. In our experiment, only a small difference was detected between the mean proportions of surrogates recovered from kelp in treatments with and without TEP (Fig 5., treatments A and B), with more surrogates being recovered from kelp without TEP.

The hypothesis that TEP can influence the adhesion of oocysts to surfaces should be further investigated. Biofilms of TEP origin develop quickly on surfaces submerged in filtered seawater [35]. Results from treatments C and D suggest that TEP may deliver surrogates to surfaces (Fig. 4): even though we removed all TEP producers by filtering seawater at 0.2 μm, TEP precursors in the colloidal form may have been present and could have formed TEP over the 12 hr incubation period, with concentrations possibly being below our limit of detection. Surrogates could therefore have become associated with these TEP and thereby delivered to the surfaces of the synthetic kelp blades and surfaces within the experimental containers. However, the inherent ‘loss’ of surrogates through experimental steps is also observed in experiments that use these particles in ultra purified water and recovery of all surrogates rarely occurs. The influence of TEP in oocyst delivery to kelp blades should be further investigated, perhaps in a similar experiment using a method that would detect TEP at lower concentrations.

Conclusions

Our findings suggest a novel route of exposure of sea otters to the protozoan parasite T. gondii. Although the estimated number of T. gondii oocysts that are transported to kelp forests is unknown, these experimental results provide a mechanism to explain the transmission of T. gondii oocysts to sea otters. As T. gondii oocysts are deposited in the coastal ocean via contaminated runoff, we propose that a proportion of them attach to the sticky EPS biofilms on the kelp blades, with the surface communities composed in part of benthic diatoms and bacteria. Snails, which feed by scraping these benthic diatoms from the surface of kelp blade using their radula, would ingest T. gondii oocysts as ‘bycatch’, explaining why sea otters that specialize on consuming subtidal snails are more likely to be exposed to this parasite. The route of infection of other pathogens to marine wildlife may also occur via an EPS-adhesion mechanism such as that described here. Insight into EPS-mediated pathogen transmission may also have significant implications for human public health, due to consumption of marine animals that feed on EPS-coated substances [66]. This study, therefore, suggests
Figure 9. A) Benthic diatoms scraped from the surface of kelp blades analyzed in Experiment 2 showing alcian blue staining of EPS. B) Same image as in A, but observed simultaneously with transmitted light showing both EPS, surrogates of *T. gondii* oocysts (white arrows) and benthic diatoms fluorescing red. C) Same image as in A and B, but observed using 50 W light source and a chlorophyll filter set showing surrogates (white arrows) and chlorophyll (red fluorescence) from benthic diatoms. Scale bars 100 μm.

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a new transmission route for delivering microsporidian-pelagic pathogens to higher trophic level predators in marine ecosystems.

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Author Contributions

Conceived and designed the experiments: FFMM KS MS. Performed the experiments: FFMM. Analyzed the data: FFMM KS MS. Contributed reagents/materials/analysis tools: FFMM KS. Wrote the paper: FFMM.

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