New record and phylogenetic affinities of the oomycete *Olpidiopsis feldmanni* infecting *Asparagopsis* sp. (Rhodophyta)

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ABSTRACT: A new geographic record of the oomycete *Olpidiopsis feldmanni* infecting the tetrasporophytic stage of the red alga *Asparagopsis* sp. from the Adriatic Sea, confirmed through morphological identification, allowed us to expand previous observations of this organism. Ultrastructural investigations of environmental material showed a large central vacuole and a cell wall thicker than previously reported from other basal oomycete pathogens of algae. Phylogenetic analysis closely associates *O. feldmanni* to *O. bostrychiae* concurrent with structural observations. This constitutes the first genetic characterisation of an *Olpidiopsis* species that was initially described before 1960, adding to the genetic data of 3 other marine *Olpidiopsis* species established and genetically characterised in the last 2 decades. The paper discusses concurrences of the ultrastructural observations made here and in previous studies of the marine *Olpidiopsis* species with those made on the freshwater species.

KEY WORDS: Central vacuole · Cyochrome oxidase subunit II · COII · cox2 · SSU rRNA · *Falkenbergia* · Transmission electron microscopy · TEM · Adriatic Sea

INTRODUCTION

Much remains unknown about the eukaryotic pathogens which infect marine algae (Andrews 1976, Gachon et al. 2010). Studying the majority of these organisms can be difficult due to their often biotrophic nature, meaning that establishing and maintaining pathosystems, consisting of the host and a single pathogen, may not be possible. The eastern Mediterranean has recently been the focus of an investigation into pathogens of brown algae, specifically those of *Ectocarpus* sp. and Sphacelariales (Strittmatter et al. 2013), although previously only one protistan pathogen affecting algae, viz. *Pleotrichelus ectocarpii* Jokl, has been described from the Adriatic Sea (Jokl 1916).

While the basal oomycete *Eurychasma dicksonii* has received the most research interest in recent years (Küpper & Müller 1999, Küpper et al. 2006, Sekimoto et al. 2008a, Gachon et al. 2009, Tsirigoti et al. 2014, 2015), 3 novel species of *Olpidiopsis*, i.e. *O. bostrychiae* Sekimoto, Klochkova, West, Beakes & Honda (West et al. 2006, Sekimoto et al. 2009), *O. porphyrae* Sekimoto, Yokoo, Kawamura & Honda (Sekimoto et al. 2008b) and *O. pyropia* Kim & Klochkova (Klochkova et al. 2015), have recently been described as pathogens of red algae (Table 1). These species are regarded as the most intensively...
studied within the genus, having been investigated through molecular and ultrastructural analysis. Comparatively, these 3 marine species appear relatively diverse, with several morphological differences including the flagellation pattern of the zoospores and the presence/absence of a central vacuole (Sekimoto et al. 2008b, 2009, Klochkova et al. 2015). In comparison to other marine members of this genus, *O. bostrychiae* was noted as being morphologically similar to the little reported *O. feldmanni* Aleem, 1952, although there was no common reported host between these 2 species (Table 1; West et al. 2006, Sekimoto et al. 2009). Additionally, 4 other marine *Olpidiopsis* species have been described, though like *O. feldmanni*, they remain to be genetically surveyed (Table 1). The holotype of this genus is *O. saprolegniae* (Braun) Cornu, 1872, a freshwater pathogen of *Saprolegnia* (Oomycota) species. As with the marine *Olpidiopsis* species, freshwater *Olpidiopsis* species have been studied at the ultrastructural level (Bortnick et al. 1985, Martin & Miller 1986a,b), but to date, no freshwater *Olpidiopsis* species have been molecularly surveyed.

The present study aims to characterise a eukaryotic pathogen found on the tetrasporophytic stage of *Asparagopsis* sp. The pathogen was identified as *O. feldmanni* based on morphological observations by light microscopy and was further examined by transmission electron microscopy (TEM). Additionally, we provide the first set of molecular markers for *O. feldmanni* and explore the phylogeny of this organism.

### MATERIALS AND METHODS

#### Sample collection

*Asparagopsis* sp. tetrasporophyte (synonym *Falkenbergia*) samples were observed during routine physiological examinations of material collected in Kaštela Bay and Vira (Adriatic Sea, Croatia), at irregular intervals between December 2012 and March 2014 (Fig. 1). Samples of *Asparagopsis* sp. collected from Kaštela Bay were maintained in the aquaria of the Institute of Oceanography and Fisheries (IOF), Croatia, with a constant flow of fresh seawater. The conditions in the tank naturally replicated those of the ambient water of Kaštela Bay due to the positioning of the IOF. Average surface seawater temperature in Kaštela Bay during the survey of *Asparagopsis* sp. in 2012/2013 was between 17°C in November and 11.5°C in February (Table 2). Salinity is slightly lower compared to average salinity in the Adriatic Sea due to influence of the River Jadro, dropping from 37.0 PSU in December to 35.5 PSU in March.

Infected filaments containing pathogenic structures were isolated and stained with acetocarmine, to preferentially highlight oomycete structures; permanent microscope mounts were created using Karo™ syrup (Küpper & Müller 1999, Strittmatter et al. 2013). Parallel to this, samples were also placed in CTAB buffer for downstream molecular work (Phillips et al. 2001). Studying the micro-photographic database at the IOF allowed the investigation of previous collections to indicate previous occurrences of the infection.

#### Microscopy

Live samples and permanent mounts were imaged and processed at the IOF using an Olympus BX51™ microscope with a DP12 or Canon 550D camera. Per-
permanent mounts were viewed in Aberdeen (UK) using a ZEISS Axio imager D2™ inverted microscope (magnification 4–40×) and were imaged using a ZEISS Axiocam MRC™. Images were acquired and processed using the ZEISS Zen™ image processing software. For TEM, infected algal thalli were collected from the field and shipped in seawater from Split to Athens. The fresh material was subsequently fixed, processed and imaged as previously reported (Sekimoto et al. 2008a), with slight modifications to the initial fixative buffer; in the present study, thalli were fixed with 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer containing 0.25 M sucrose. Examination of the sections was performed with a Philips 300 transmission electron microscope.

**Molecular methods**

DNA extraction was performed on environmental samples (i.e. placed in CTAB at the time of collection) where no other eukaryotes (i.e. diatoms) or parallel infection was detected in the permanent mounts prepared, as detailed by Gachon et al. (2009). DNA extracts were then quantified and checked for quality using a spectrophotometer (260/280 nm) before being diluted to suitable concentrations for the purposes of PCR (10–50 ng µl⁻¹).

Using the oomycete sequences available in the public domain, oomycete-specific primers were designed to amplify the small subunit (SSU) rRNA marker, SFP312F (5’-ATG AAT CAA TTG AGT TTC TG-3’) and SFP1629R (5’-GAC GTA ATC AAT GCA AGC-3’) using 4.0 mM MgCl₂ and a PCR cycle of 95°C for 5 min; 35 cycles of 95°C for 30 s, 56°C for 30 s and 72°C for 60 s; and 72°C for 5 min. The cytochrome oxidase subunit II (COII) marker was amplified using the primers defined by Hudspeth et al. (2000) using 3.5 mM MgCl₂ and a PCR cycle of 95°C for 5 min; 35 cycles of 95°C for 30 s, 50°C for 30 s and 72°C for 45 s; and 72°C for 5 min. An aliquot of the PCR product was run on a 1% w/v agarose gel in 1× Tris-borate-EDTA (TBE) buffer, stained with GelRed™ (Biotium) and viewed on a UV lightbox, prior to purification (ThermoScientific GeneJet™ nucleic acid purification kit). The PCR products were sequenced using the primers, used for initial amplification, by the Eurofins Value Read™ sequencing service. The COII and SSU rRNA sequences were deposited under accession numbers KM210528 and KM210530, respectively.

The ABI files returned were initially manually trimmed and inspected for erroneous base callings using BioEdit™ (Hall 1999), and a single consensus
sequence was created for each marker. Blastn/x was used to assign preliminary identities to sequences, and alignments of oomycetes were downloaded from NCBI. Alignments were tested with the maximum likelihood (ML) algorithm using RAxML 8.1.11 (Stamatakis 2006) through the CIPRES gateway (www.phylo.org/sub_sections/portal/), using a BLOSUM62 substitution matrix for the COII amino acid alignment and a GTRCAT model for the SSU rRNA, recording branch length and allowing bootstrapping to stop automatically. Hyphochytrids were used as outgroups in every test. To ensure consistent topology, neighbour-joining (NJ) tests were also performed on the same alignments with MEGA6.0 (Tamura et al. 2013) using a p-distance (COII) and Tajima-Nei (SSU rRNA) model, with gaps treated as partial deletions above a 95% cut-off and 1000-bootstrap test of phylogeny. Alignments and the NCBI accessions of sequences used are available in the Supplement at www.intres.com/articles/suppl/d117p045_suppl.pdf.

RESULTS

Field and microscopic observations

Asparagopsis sp. tetrasporophytes infected by Olpi diopsis feldmanni were present in Vira (December 2012) and Kaštela Bay (December to January of 2012/13 and 2013/14 and March 2014; Fig. 1). They were found epiphytic on Sympodium scoparium (Linnaeus) Kützing, 1843, and Cladophora prolifera (Roth) Kützing, 1843, or epilithic between 0.5 and 1.5 m depth. The occurrence of Asparagopsis sp. tetrasporophytes in this geographic region is maximal between October and February, commonly between 0.5 and 3 m depth. The occurrence of Asparagopsis sp. tetrasporophytes in this geographic region is maximal between October and February, commonly between 0.5 and 3 m depth, and is observed to naturally decline in March. By analysing the IOF photo-microscopy data-base, outbreaks of this pathogen were also found to be present in Kaštela Bay in November 2006.

Light micrographs of fresh material and permanent mounts demonstrated a specific discolouration and enlargement of various cells of the Asparagopsis filament, inside which holocarpic parasitic thalli develop (Fig. 2A). After some time, under aquarium conditions, these thalli could be seen to have emptied and diatoms became established upon the alga (Fig. 2B). Most of the infection occurred in younger parts of the Asparagopsis filaments, while almost none occurred in the oldest cells. Infection of cells inside filaments results in formation of oblong parasitic sporangia (Fig. 2C). Pearson’s correlation determined that their dimensions corresponded to enlarged dimensions of neighbouring parallel cells and by extension, the infected Asparagopsis cell, being 39 to 76 µm in length (R = 0.8173; see the Supplement). Infection of apical cells results in the formation of spherical parasitic sporangia which are within the range of 27 to 63 µm in diameter (Fig. 2D,I−K and see Fig. 3A; additional measurement data are available in the Supplement), although in rare instances, non-spherical sporangia could be seen (Fig. 2E). We detected a significant difference between the length of the oblong sporangia and the diameter of spherical sporangia (t-test, p < 0.01; see the Supplement) as previously reported (Aleem 1952, 1953, Sparrow 1960). Occasionally exit tubes were visible on sporangia, both full and empty (Fig. 2F,G); typically, a single exit tube was visible, but in some instances multiple (up to 3) tubes could be seen (Fig. 2H). Discharge tubes are straight and measure approximately 11 to 55 µm long and 3.6 to 6.5 µm wide. The cleavage of zoospores appears before the release, and it appears that single spores can be seen as cleaved in some permanent mounts, though never in enough detail to see flagella (Fig. 2J).

TEM

A pathogenic walled thallus (Fig. 3A,B) at a stage comparable to the one shown by bright field imaging (Fig. 3A, inset), was located using TEM, allowing ultrastructural investigation of the pathogen. The walled thallus has an apical position and is illustrated here in 2 different TEM sections, both with and without a central vacuole of lipid texture (Fig. 3A−C). Numerous parasite nuclei with associated vesicles and mitochondria appear to dominate the thallus at this stage (Fig. 3A−C). The absence of membranes around the nuclei of the coenocytic cytoplasm (Fig. 3D) and of formed exit tubes implies that this phase corresponds to a pre-cleavage stage. Higher magnifications identified further ultrastructural features, such as tubular vesicles containing mastigonemes surrounded by mitochondria (Fig. 3E) and suspected kinetosomes (Fig. 3F). It should be noted that the material is taken directly from the field, a fact not allowing repetition of sampling and improvement of the fixation. Additionally a thick, electronlucent cell wall (~1.3 µm; arrowed line in Fig. 3G) surrounding the pathogen separates the contents of the parasitic thallus from the host cytoplasm and organelles which appear pushed aside (Fig. 3B,G), with chloroplasts clearly visible at high magnifications (Fig. 3G). The host cell wall remains intact, encapsulating the parasite thallus (Fig. 3G).
Fig. 2. Light micrographs of *Olpidiopsis feldmanni* infecting the tetrasporophytic stage of *Asparagopsis* sp., with specific focus on the sporangia. (A) Heavily infected material with sporangia and (B) late-stage infected material where the sporangia have been emptied, releasing infectious zoospores. The alga is vulnerable to being colonised by diatoms at this stage. (C–E) Individual sporangia, both (C) intercalated in an algal filament where the sporangium is forced into a bulging shape by the plant, and (D) infecting an apical cell where the oomycete has adopted the more common spherical sporangia, though in rare instances (E) alternative sporangium morphology is seen. (F–H) Typically 1, though (rarely) up to 3 exit tubes protrude from both forming and evacuated sporangia. (I–K) Permanently mounted preparations, where sporangia are dyed red by acetocarmine. (I) Large apical cell, swollen over 50 µm in diameter. (J) Sporangium above the algal filament, suggesting that a new branch was forming facing upwards; individual zoospores can clearly be differentiated, suggesting a maturing sporangium. (K) Spherical cells were also formed in cells that were not apical, though still young. In all images, sporangia/empty sporangia are highlighted with thick arrowheads and exit tubes with arrows. Scale bars = (A,B) 50 µm, (C–J) 20 µm.
Phylogenetic trees

All successfully amplified fragments were taken from environmental samples where corresponding permanent mounts indicated a significant infection within the red alga. No signs of a parallel infection were ever noted, so we are confident that the sequences obtained truly belong to the algal pathogen. The SSU rRNA fragment obtained (KM210530) when tested (Fig. 4) placed the Asparagopsis sp. pathogen alongside the oomycete Olpidiopsis bostrychiae (bootstrap support — ML: 99/100, NJ: 98/100) in a clade with O. porphyrae, O. pyropiae and 4 environmental sequences (Table 3; ML: 96/100, NJ: 61/100). Depending on the test of phylogeny, these environmental sequences could be seen to be closer to either O. bostrychiae and the Asparagopsis sp. pathogen (ML; Fig. 4A) or to O. porphyrae (NJ; Fig. 4B). The latter test weakly associated O. pyropiae to O. bostrychiae and our newly obtained sequence, but with very poor support (36/100; Fig. 4B). Additional to this, the relationship between genera is slightly altered depending on the test used; ML supports that the Olpidiopsis and Haliphthorales are both an equal distance from all other oomycetes (59/100; Fig. 4A), whereas the NJ test weakly supports the separation of this branch, placing the Olpidiopsis closer to the non-basal oomycetes (50/100; Fig. 4B). No other branching deviations were noted between tests. Similarly, the COII fragment obtained (KM210528) when tested (Fig. 5) placed the Asparagopsis sp. pathogen with O. bostrychiae (100/100), although the association of O. porphyrae with this clade was very weak when assessed by ML (<50) and is moved to associate within the Haliphthorales clade when tested by NJ. In addition, the close association of Halioticida sp. NJM_0034 (<50) with Olpidiopsis shown by the COII marker is not reflected in the previous SSU rRNA topology, where this abalone pathogen is seen to branch towards the non-basal oomycetes a step above Olpidiopsis in agreement with a recent large subunit rRNA phylogeny study of this genus (Macey et al. 2011). High-order branches of the COII with weak support (i.e. <70) could be collapsed by the inclusion/exclusion of certain taxa, concurrent with previous studies (Sekimoto et al. 2009).
Fig. 4. Maximum likelihood (ML) tree based on the SSU rRNA marker testing 45 sequences, including 4 environmental sequences (highlighted in red) which clade within the order Olpidiopsidales. Branches are labelled with percentage support from 1000 bootstrap tests of phylogeny for the ML test and a corresponding neighbour joining (NJ) test. ‘XX’ indicates that the node was not predicted in the NJ test. Significant differences between the ML and NJ topology are highlighted by the arrows. (A) ML predicted that the orders Olpidiopsidales and Haliphthorales branch together an equal distance from all other oomycetes and that the 4 environmental samples branch closest to *Olpidiopsis bostrychiae* and *O. feldmanni*. (B) NJ predicted that the Olpidiopsidales branch closer to the non-basal oomycetes than the Haliphthorales and also placed the 4 environmental samples closer to *O. porphyrae* than any other *Olpidiopsis* species. Alignment and accession numbers are provided in the Supplement.
**DISCUSSION**

**Olpidiopsis feldmanni:** identification and confirmation

Light microscopic investigations of the pathogen found infecting the tetrasporophytic (*Falkenbergia*) stage of *Asparagopsis* sp. are congruent to the descriptions of *Olpidiopsis feldmanni* (Sparrow 1960) originally described from the Mediterranean Sea (French coast; Aleem 1952) and Baltic Sea (Sweden; Aleem 1953). Uniquely, a rare observation of a non-spherical sporangium in the apical host cell (Fig. 2E) and a significantly larger sporangial size in apical

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**Table 3. Environmental samples which branch within the marine *Olpidiopsis* clade. GenBank accession numbers**

<table>
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<th>Tag identifier</th>
<th>Accession</th>
<th>Author</th>
<th>Water/country of discovery</th>
<th>Notes</th>
</tr>
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<td>D3P05B11</td>
<td>EF100276.1</td>
<td>Stoeck et al. (2007)</td>
<td>Baffin Bay/Greenland</td>
<td>Oxygen depleted intertidal marine sediment, upper 2 cm</td>
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<td>D3P06D06</td>
<td>EF100297.1</td>
<td>Stoeck et al. (2007)</td>
<td>Baffin Bay/Greenland</td>
<td>Coastal surface water</td>
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<tr>
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<td>EF100297.1</td>
<td>Massana et al. (2004)</td>
<td>Mediterranean Sea/Spain</td>
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<tr>
<td>H3S3Be43I</td>
<td>JQ781890.1</td>
<td>Lin et al. (2012)</td>
<td>Pacific Ocean/USA</td>
<td>Ocean waters</td>
</tr>
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**Fig. 5. Maximum likelihood (ML) tree based on the COII marker testing 42 sequences.** Branches are labelled with percentage support from 1000 bootstrap tests of phylogeny from the ML test and a corresponding neighbour joining (NJ) test, if there was >50% support at the node. Branches with <50% support are grey, and 'XX' indicates that the node was not predicted in the NJ test. Arrowed nodes were not predicted by NJ, resulting in a significant tree rearrangement; data not shown. Alignment and accession numbers are provided in the Supplement.
cells were recorded (average diameter of 40 µm, ranging from 27.33 to 62.54 µm [see the Supplement]; previously 15–30 µm; Sparrow 1960). Comparing the size of an infected cell with that of an adjacent, parallel cell provided a strong correlation, suggesting that the host may limit the expansion of the pathogenic thallus, as indicated by the shape adopted in older cells of the host alga. Comparatively, spherically shaped thalli, i.e. the natural shape of an Olpidiopsis thallus (West et al. 2006), can be seen in younger apical cells (Fig. 2). Indications of host hyperplasia are noted (Fig. 2), where the host cell appears to have become enlarged due to reproduction of the pathogen, similar to instances seen in other holocarpic oomycete pathogens, such as Eurychasma dicksonii (Sekimoto et al. 2008a) and O. antithamnionis (Whittick & South 1972).

Key ultrastructural similarities between this pathogen and other oomycetes (i.e. Olpidiopsis sp. and Eurychasma sp.) indicate that this is a correct diagnosis with the pathogen presenting a coenocytic walled thallus, containing multiple nuclei and mitochondria. A similar organization is presented by the protoplast of other oomycetes, such as in the walled vegetative thallus, containing multiple nuclei and mitochondria. The material is the enlarged cell wall of 2; Klochkova et al. 2015). It seems possible that this fusion may result in multiple, smaller vacuoles being present in a thallus, although the original study did not report this. This fusion, however, was indicated as a rationale for an unusual number of zoospores being detected in mature thalli (i.e. not to the power of 2; Klochkova et al. 2015).

A final unique ultrastructural observation from our Olpidiopsis feldmanni material is the enlarged cell wall presented by this pathogen (Fig. 3G), which has approximately 6 times the thickness of the cell wall of O. porphyrae, O. bostrychiae, O. pyropiae and E. dicksonii (~200–400 nm) at corresponding phases (Sekimoto et al. 2008b, 2009, Klochkova et al. 2015, Tsirigoti et al. 2015). Given that the latter species are imaged from established host–pathogen cultures, it may be that an environmental agent which is not present in the culture is responsible for this increase in size.

This diagnosis correlates with the phylogenetic analysis, where both markers surveyed were able to consistently associate the Asparagopsis sp. pathogen with O. bostrychiae (Figs. 4 & 5). The relative ease with which the sequences were obtained with primers shown to amplify multiple oomycete genera (Hudspeth et al. 2000) suggests that no other oomycetes were present in our samples and is supported by detailed examination of permanent mounts.

In summary, the SSU rRNA (Fig. 4) can be seen as being the key marker for the basal oomycetes. The relatively highly conserved nature of the marker appears well suited to resolving the branching order of the few sampled basal oomycete species and consistently supported the placement of the marine Olpidiopsis species into a single clade. Four environmental samples were also found to clade within this genus, which could be seen as additional marine Olpidiopsidales sequences surveyed directly from the environment (Table 3) but not morphologically investigated. Depending on the test of phylogeny used, the nearest Olpidiopsis sp. relative to these 4 environmental sequences would change (Fig. 4), although the support of the NJ topology is significantly reduced relative to the ML topology. An additional topological difference noted between these tests is the relationship between the orders Olpidiopsidales and Haliphthorales. These orders were weakly grouped together when tested by ML (Fig. 4A), i.e. branching the same distance from every other oomycete, but when tested by NJ, they were weakly supported as being differentiated (Fig. 4B) with the Olpidiopsidales branching towards the non-basal oomycetes. This uncertainty in the placement of the basal oomycete orders suggests that there may be a large amount of genetic diversity between these orders, yet to be surveyed. Additionally, the long branch distances between the Olpidiopsis species (Sekimoto et al. 2009, Klochkova et al. 2015) supports a large genetic distance between the currently surveyed Olpidiopsis species, and it is possible that there will be a large diversity of intermediate species, as yet undetected.

This weak association is further highlighted by the COI marker which poorly supports the clading of the 3 Olpidiopsis species surveyed at this marker (Fig. 5), with no corresponding sequences for O. pyropiae being available at this time. In agreement with previ-
ous findings (Sekimoto et al. 2009), we also noted that altering the taxa surveyed could significantly affect the topology, in particular the placement of *O. porphyrae* and *Halalioticida* sp.

While these analyses have shown that there is a close association morphologically, ultra-structurally and genetically between *O. feldmanni* and *O. bostrychiae*, these are still regarded as 2 distinct species. While it was not possible to evaluate the host range of *O. feldmanni*, due to the inability to bring this organism into culture, a previous host range analysis has been performed on *O. bostrychiae*, which was unable to successfully infect *Asparagopsis* sp. *in vitro* (West et al. 2006, Sekimoto et al. 2009). As a host is required to fulfil the life cycle (and the reported sexual cycle; Martin & Miller 1986a) of all *Olpidiopsis* sp., the inability to infect the same host species can be regarded as an adequate species barrier between these 2 organisms at present. It should be noted here that sex in marine *Olpidiopsis* species has not been observed to date; indeed, recent TEM investigations of fusing *O. pyropiae* thalli failed to identify the fusing of any nuclei (Klochkova et al. 2015).

This new record from the Adriatic constitutes the first published finding of *O. feldmanni* in over 60 yr and provides the first set of molecular markers for this species. Prior to this, *O. feldmanni* has only been reported from *Asparagopsis* sp. and *Bonnemaisionia hamifera* (Agardh) Silva, 1957, in the Mediterranean and Baltic Seas (Aleem 1952, 1953, Sparrow 1960). It is unclear, given that *Asparagopsis* spp. are considered invasive in the Mediterranean Sea (Garzoli et al. 2014), what the geographic origins of *O. feldmanni* are. Further studies to establish the host range are necessary to determine whether any native Mediterranean algal species are susceptible. Given the previous geographic records, it is not unimaginable that the parasite may be present in most other marine regions within western Europe where the host is present. Indeed, the regular occurrence of *O. feldmanni* infecting *Asparagopsis* sp. in the autumn on the south and west coast of the British Isles has been suggested, but no evidence was reported to substantiate this claim (Dixon & Irvine 1977). As the pathogen was found year to year during this study and was also detected in historical images, it can be supposed that this pathogen is an established member of the community and regular epidemics can be expected during the annual occurrence of the tetrasporophytic stage of *Asparagopsis* sp. The orthology of this species is retained as *O. feldmanni* in concurrence with the original description of this species (Aleem 1952), as with previous studies (West et al. 2006, Sekimoto et al. 2008b, 2009, Klochkova et al. 2015). The recently suggested reclassification of the *Olpidiopsis* species into differing genera based on host range (Dick 2001) is regarded as premature due to a lack of phylogenetic knowledge.

**Olpidiopsis sp.: a morphological overview**

The key differences between the marine and freshwater *Olpidiopsis* species, studied through electron microscopy, are summarised in Table 4. Observations by light microscopy highlight the different flagellation pattern presented by the marine species. *O. bostrychiae* presents laterally biflagellate zoospores (West et al. 2006, Sekimoto et al. 2009), which appears to be a common feature of most marine *Olpidiopsis* species (Whittick & South 1972), while *O. porphyrae* and *O. pyropiae* present sub-apically biflagellate zoospores (Sekimoto et al. 2008b). The holotype species *O. saprolegnae* also presents sub-apically biflagellate zoospores (Bortnick et al. 1985).

In addition to this, TEM of developing walled thalli reveals a single central vacuole present in the maturing thallus of *O. feldmanni* and *O. bostrychiae* (Sekimoto et al. 2009). In comparison, *O. porphyrae*, *O. pyropiae* and the freshwater species *O. varians* present multiple vacuoles of a similar density (Karling 1949, Martin & Miller 1986b, Sekimoto et al. 2008b), while *O. pyropiae* may present one or more vacuoles, as identified by light microscopy (Klochkova et al. 2012). Previous observations suggest that these vacuoles may facilitate cleavage during zoosporogenesis of holocarpic stramenopiles (Karling 1942, 1943). Subsequent ultrastructural studies from biflagellate *Petersenia palmatae*, a pathogen of *Palmaria mollis*, supported this (Pueschel & van der Meer 1985), while expulsion vacuoles have been described from germinating cysts of the biflagellate *Ectrogaella perlans* (Kumar 1980). Recently, the organization of the cytoskeleton during the process of infection and sporangium development was studied in a brown algal host cell and the parasite *E. dicksonii*. It is assumed that the cleavage procedure is directed by a modified system of microtubules and a central vacuole that appears during this phase (Tsirigoti et al. 2014). These 3 features would lend weight to a hypothesis that freshwater *Olpidiopsis* species are more closely related to *O. porphyrae* and *O. pyropiae* than the 2 other marine species genetically surveyed. With this hypothesis in mind and the large genetic difference observed between marine *Olpidiopsis* species, it
could be seen as justifiable to attempt to delineate these species into 2 separate genera. Yet as the species at present form a monophyletic clade, we believe that this would be a premature and potentially irrelevant nomenclature change. This rationale is supported by the ecology (i.e. marine) and host species (i.e. red algae) of the genetically surveyed marine Olpidiopsis species (Table 1), which should still be regarded as significant. It may be the case that freshwater Olpidiopsis, when genetically surveyed, will be further removed from all marine Olpidiopsis species. Unfortunately, with the limited molecular surveying of basal oomycetes, any tentative placement of these species based on morphological characteristics only should be viewed as highly speculative. Instead, resolving the phylogenetic position of freshwater Olpidiopsis species, through genetic screening, should be considered a key route to expand our knowledge of this highly diverse genus.

**CONCLUSION**

The present study identified *Olpidiopsis feldmanni* in nature, obtaining live environmental isolates, enabling investigation of this pathogen through molecular phylogenetics as well as light and electron microscopy, with significant evidence showing this pathogen to be genetically and morphologically closer to *O. bostrychiae* than to *O. porphyrae* or *O. pyropiae*. We note that in order to ensure the assigned sequences are correct, meticulous observation of the algae and associated material was necessary to ensure no second oomycete species was present in the DNA extraction. Additionally, as only 1 sequence was obtained with oomycete-wide directed COII primers, we are confident that only 1 oomycete species was present in our samples. This type of study is effective when expanding the knowledge of the existing catalogue of oomycetes (Sparrow 1960) for which reference molecular data exist. When molecularly surveying a new genus or a distant phylogenetic branch, where no reference data are available, then either pure cultures or fluorescent *in situ* hybridisation protocols (Langer-Safer et al. 1982) must be established.

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