Perkinsus mediterraneus n. sp., a protistan parasite of the European flat oyster Ostrea edulis from the Balearic Islands, Mediterranean Sea

Sandra M. Casas1, Amalia Grau2, Kimberly S. Reece3, Kathleen Apakupakul3, Carlos Azevedo4, Antonio Villalba1,*

1Centro de Investigación Mariñas, Consellería de Pesca e Asuntos Marítimos, Xunta de Galicia, Apartado 13, 36620 Vilanova de Arousa, Spain
2Estación de Acuicultura, D. G. Pesca, Govern Balear, Port d’Andratx, Mallorca, 07158 Spain
3School of Marine Science, Virginia Institute of Marine Science, College of William and Mary, Gloucester Point, Virginia 23062, USA
4Department of Cellular Biology, Institute of Biomedical Sciences, University of Oporto, Porto 4099-003, Portugal

ABSTRACT: A new species, Perkinsus mediterraneus, a protistan parasite of the European oyster Ostrea edulis (L.), farmed along the coast of the Balearic Islands, Mediterranean Sea, is described. Morphological examinations with light and transmission electron microscopy, DNA sequence-analysis and enlargement in Ray’s fluid thioglycollate medium (RFTM) confirmed that this parasite belongs to the genus Perkinsus. Specific morphological and genetic characteristics indicated that it should be considered a new species in the genus. Sequencing of the small subunit ribosomal (ssu rRNA) gene confirmed that the parasite belongs to the genus Perkinsus, and sequences of the internal transcribed spacer (ITS) were distinct from any Perkinsus ITS sequences previously published and/or deposited in the GenBank. Phylogenetic analysis revealed that the ITS sequences of the new species formed a monophyletic group comprising a sister clade to the *P. atlanticus/olseni* group. In addition, morphological differences were observed between the new species and the other described Perkinsus spp.. After incubation in RFTM for 1 wk, the prezoosporangium had reached an extremely large size (97.4 ± 1.99 µm) (mean ± SE), and after 2 wk incubation had again almost doubled in size (167.1 ± 8.09 µm). The discharge-tube length was one sixth the diameter of the zoosporangium, i.e. a ratio of 17.36:97.38, the lowest ratio observed for any Perkinsus species. At the ultrastructural level, zoosporangia and zoospores exhibited some differences compared to other Perkinsus species.

KEY WORDS: Perkinsus mediterraneus n. sp. · Ostrea edulis · Oyster parasite · Ultrastructure · Zoosporulation · ssu rRNA gene · Internal transcribed spacer · Perkinsus atlanticus

INTRODUCTION

that Perkinsus species are dinoflagellates, or are closely related to them. Other studies have placed the genus Perkinsus within the alveolates, in a new phylum Perkinsozoa, together with the phyla Dinoflagellata, Apicomplexa and Ciliophora (Norén et al. 1999).

To date, the genus Perkinsus has comprised 6 species: *P. marinus* in the eastern oyster *Crassostrea virginica* (Mackin et al. 1950) along the Atlantic and Gulf coasts of the USA, *P. olsenii* in the blacklip abalone *Haliotis ruber* from Australia (Lester & Davis 1981), *P. atlanticus* (Azevedo 1989) in the carpet-shell clam *Tapes decussatus* from the Iberian Peninsula, *P. qugwadi* in the Japanese scallop *Patinopecten yessoensis* (Blackbourn et al. 1998) from British Columbia, Canada, *Perkinsus chesapeaki* (McLaughlin et al. 2000) in the softshell clam *Mya arenaria* from Chesapeake Bay, USA, and *P. andrewsi* (Coss et al. 2001b) in *Macoma balthica* from Delaware Bay.

In European waters, Perkinsus-like organisms were detected in the clam *Tapes decussatus* and the flat oyster *Ostrea edulis* in 1978 (da Ros & Canzonier 1985). *P. atlanticus* is the only species of the genus identified along the European coasts to date. It has been found in *T. decussatus* from the Iberian Peninsula: Algarve, South Portugal (Azevedo 1989), Huelva, South Spain (de la Herrán et al. 2000) and Galicia, NW Spain (Robledo et al. 2000, Casas et al. 2002a) and in *T. philippinarum* from Cataluña (NE Spain) (Sagristà et al. 1996).

The occurrence of Perkinsus-like organisms in the Mediterranean Sea has been documented in *Ostrea edulis, Crassostrea gigas, Tapes decussatus, T. philippinarum, Venerupis secta*, Venus verrucosa, Callista chione, Cerastoderma edule, *Mytilus galloprovincialis*, Chamelea gallina and *Musculista senhousia* from Italian coastal waters (da Ros & Canzonier 1985, Ceschia et al. 1991, Berilli et al. 1998, Canestri-Trotti et al. 1999, 2000a,b), and in bivalves from the French Mediterranean coasts (Goggin 1992), but none of those studies identified the parasite to species level.

Cases of infection by Perkinsus-like organisms were observed in the tissues of the European flat oyster *Ostrea edulis* cultivated in the Balearic Islands, Mediterranean Sea. This article describes the morphological characteristics of different life stages and DNA sequences of the rRNA complex of the oyster parasite from Balearic Islands, and identifies a new species of the genus Perkinsus, *P. mediterraneus* n. sp.

MATERIALS AND METHODS

Oyster sampling. European flat oysters infected by a Perkinsus-like organism were found during an historical survey of a commercial oyster-growing site in close proximity to Maó Port (Isle of Menorca, Balearic Islands, Spain), in the western Mediterranean Sea. The survey involved monthly harvesting of oyster samples, which were processed by standard histological techniques. Briefly, an approximately 5 mm thick section of tissue containing gills, visceral mass, and mantle lobes was excised, fixed in Davidson’s solution (Shaw & Battie 1957) and embedded in paraffin. Sections of 5 µm thickness were stained with Harris-hematoxylin and eosin. Histological sections of infected oysters were examined by light microscopy to determine the morphology of the Perkinsus-like organism.

Isolation of prezoosporangia and induction of zoosporulation. A preliminary trial was performed in November 1998. The digestive gland, gills and rectum of 25 oysters collected from the same site were incubated in Ray’s fluid thioglycollate medium (RFTM) (Ray 1966) to induce transformation of trophozoites of the Perkinsus-like parasite into prezoosporangia. After 1 wk in RFTM at room temperature in the dark, the tissues were homogenised (8000 l min⁻¹, 30 s, ultraturrax T-25, IKA Labortechnik), and the prezoosporangia were isolated (de la Herrán et al. 2000, Casas et al. 2002a) as follows: oyster tissues were trypsinised (0.25%) for 60 to 90 min, and prezoosporangia were separated by passing the suspension through sieves of 425, 225, 140 and 30 µm. Only material retained on the 30 µm sieve was processed further, since no significant numbers of prezoosporangia were detected in the other sieve sizes. Prezoosporangia in the 30 µm sieve were washed with sterile filtered (0.22 µm) seawater (35‰) plus antibiotics (200 U ml⁻¹ Penicillin G and 400 µg ml⁻¹ streptomycin) (SFSWA). Prezoosporangia were incubated at 22°C in the dark in Petri plates with SFSWA to induce zoosporulation. Plates were checked daily by light microscopy to monitor cell morphology until zoosporulation.

A second trial was performed in November 1999. The digestive gland, gills and rectum of 30 oysters were incubated for a longer period (2 wk) in RFTM at room temperature. The incubated tissues were homogenised and trypsinised as described above. Because of the increased size of prezoosporangia after extended incubation, only the 425 and 30 µm sieves were used to avoid loss of prezoosporangia in discarded 225 and 140 µm sieve material. Prezoosporangia retained in the 30 µm sieve were processed as described above to induce zoosporulation.

Transmission electron microscopy. Zoosporangia and zoospores obtained in the second trial were washed and centrifuged. The pellets were fixed in 2.5% glutaraldehyde in 0.2 M sodium cacodylate buffer, pH 7.2, for 2 h at 4°C. The pellets were then washed twice with the same buffer, and postfixed in buffered 2% OsO₄ for 2 h at 4°C. The cells were
immersed in 1.5% agar solution, and the agar blocks were dehydrated in an ethanol series and embedded in Epon. Ultrathin sections were double-stained with uranyl acetate and lead citrate, and observed in a JEOL 1010 transmission electron microscope (TEM) operated at 60 kV.

**DNA-sequencing. Genomic DNA isolation:** Zoosporangia and zoospores obtained from infected oysters (N = 20) in the second trial were washed and centrifuged. Pellets were covered with tissue-storage buffer (0.25 M ethylenediamine tetraacetate [EDTA], 20% dimethylsulfoxide [DMSO], saturated NaCl [pH 8.0]) (Seutin et al. 1991) and shipped to the Virginia Institute of Marine Science for further processing. Before DNA extraction, the zoosporangia and zoospores were again pelleted, and washed with phosphate-buffered saline. DNA was extracted using a CTAB (1% N-cetyl,N,N,N-trimethylammonium bromide)-based DNA extraction method (Carlini & Graves 1999) with twice the published concentration of proteinase-K in the lysis buffer.

**PCR amplification:** The small subunit ribosomal RNA (ssu rRNA) gene and the internal transcribed spacer region (ITS) of the ribosomal RNA gene unit were amplified in the polymerase chain reaction from the genomic DNA. Amplification of the ssu RNA gene required a *Perkinsus*-specific primer that annealed within the ITS region that was paired with a ‘universal’ ssu primer (16S-A, Medlin et al. 1988). The *Perkinsus*-specific ITS primer, ‘PerkITS320comp’ (5’-CCTTTGTCAGTWGCGTTG-3’) was designed to be used in conjunction with 16S-A and to preferentially amplify the genomic DNA. Amplifications were performed with 10 to 50 ng of genomic DNA in 25 μl reactions, using the reaction conditions recommended for the BRL PCR Reagent System (Life Technologies) with twice the published concentration of proteinase-K in the lysis buffer.

**Phylogenetic analysis:** ssu sequences obtained from the *Ostrea edulis* *Perkinsus* sp. were aligned to available *Perkinsus* spp. ssu sequences (Fong et al. 1993, GenBank Accession No. X75762; Goggin & Barker 1993, Accession No. L07375; S. I. Kotob et al. unpubl., Accession Nos. AF042708 and AF042707; Robledo et al. 2000, Accession No. AF140295; C. A. Coss et al. unpubl., Accession No. AF252288; Coss et al. 2001b, Accession No. AF102171), and to those of 2 *Perkinsus* host species, *Crassostrea virginica* (Littlewood et al. 1991, Accession No. X60315) and *Tapes decussatus* (B. Novoa & A. J. Figueras unpubl., Accession No. AF295121) using the CLUSTAL-W algorithm (Thompson et al. 1994) in the MacVector 7.0 DNA Sequence Analysis Software package (Oxford Molecular). Using the same method, ITS sequences obtained in this study were aligned to those of other *Perkinsus* spp. (Goggin 1994, Accession Nos. PAU07697, PMU07700, POU07701, PSU07698, PSU07699; S. I. Kotob et al. unpubl., Accession Nos. AF091541, AF091542, AF126022, AF150988, AF150989, AF150990; G. D. Brown et al. unpubl., Accession Nos. AF149876, AF150985, AF150986, AF150987; Robledo et al. 2000, Accession No. AF140295; Coss et al. 2001b, Accession Nos. AF252288, AF102171; D. Hervio et al. unpubl., Accession No. AF151528; Casas et al. 2002a, Accession Nos. AF369967 to AF369979; Dungan et al. 2002, Accession Nos. AF440464 to AF440471; Casas et al. 2002b, Accession Nos. AF441207 to AF441218). Parsimony jackknife analysis of each set of sequences was performed using PAUP* 4b8.0 (Swofford 2001) with 100 random-addition replicates.

**Influence of length of incubation in RFTM on size of *Perkinsus atlanticus* prezoosporangium.** An experiment was performed to test the influence of length of incubation in RFTM on *Perkinsus atlanticus* prezoosporangium size, in order to evaluate the taxonomic significance of the large size of the *Perkinsus*-like prezoosporangia isolated from the oysters. *P. atlanticus* was chosen because it is the only species of the genus described from European waters thus far. Carpet-shell clams *Tapes decussatus* (> 40 mm) were collected from an intertidal bed infected by *P. atlanticus*, located in Vilalonga (Ría de Arous, Galicia, NW Spain). We
selected 27 clams with the highest numbers of whitish pustules on their gill surfaces (a conspicuous sign of infection by *P. atlanticus* for *P. atlanticus* isolation, and divided them into 3 groups (A, B and C) of 9 clams each. Gills of the 9 clams in Group A were cut into small fragments, one-third of which was incubated in RFTM for 7 d (Replicate 7A), another third was incubated in RFTM for 14 d (Replicate 14A), and the remaining one-third was incubated in RFTM for 21 d (Replicate 21A). The gills of clams in Groups B and C were processed similarly. Thus, 3 replicates (A, B, and C) were performed for each of the 3 incubation periods (7, 14 and 21 d).

After incubation, gill fragments of each replicate were separately trypsinised (0.25%) for 1 h, a 1 ml sample was taken from each replicate, the diameter of 50 cells was measured, and cellular viability was determined with neutral red (10 mg l⁻¹).

**Statistical analysis.** Differences in *Perkinsus atlanticus* prezoosporangium size among the 3 incubation periods were compared by 1-way ANOVA followed by a Fisher test, using MINITAB 13 software.

**RESULTS**

**Morphology of parasitic stages in oyster tissues**

Different stages corresponding to the life cycle of a *Perkinsus*-like parasite were observed in histological sections of *Ostrea edulis* (Figs. 1 to 4). Trophozoites occurred mostly in the connective tissue of different organs (visceral mass, gills, labial palps, mantle lobes) and also inside gonad follicles invaded by haemocytes. Trophozoites were spherical cells (mean diameter ± SE = 7.9 ± 0.34 µm; N = 50; range = 4 to 19 µm), each containing a large vacuole that occupied most of the cell volume, conferring a ‘signet ring’ appearance to the cell (Figs. 1 & 2). A vacuoplast was evident within the vacuole (Fig. 2). The nucleus was located in the periphery of the cell, and sometimes contained a nucleolus (Figs. 1 & 2). The cell was surrounded by a wall-like structure (Figs. 1 & 2). Multicellular stages, indicating vegetative multiplication of a mother cell, were also observed, with daughter cells in a rosette-like arrangement inside the wall-like structure of the mother cell (Figs. 3 & 4).

Only light-to-moderate infections were visible in the histological sections. The infection evoked haemocytic infiltration of the connective tissue in the proximity of the parasite (Figs. 1 to 4). Trophozoites were frequently observed in the cytoplasm of haemocytes and, less frequently, free in the connective tissue or surrounded by a few haemocytes. Partially degraded trophozoites were visible both inside and outside the haemocytes.

**Cellular morphology during zoosporulation**

Under light microscopy, prezoosporangia of the *Perkinsus*-like parasite isolated from *Ostrea edulis* in the first trial appeared spherical, with a large vacuole that displaced the nucleus to the periphery of the cell (Fig. 5). The mean diameter (±SE) of the prezoosporangia was 97.4 ± 1.99 µm (N = 53, range = 62 to 135 µm). Zoosporulation occurred in prezoosporangia incubated in seawater (Figs. 6 to 8). It began with the formation of a discharge tube, progressing through successive karyokinesis and cytokinesis of the cell, and producing hundreds of zoospores. The mean length of the discharge tube was one-sixth (17.36:97.38 µm) the diameter of the zoosporangial body. Zoosporangia with 2 discharge tubes were not observed. Biflagellated zoospores were liberated through the discharge tube (Fig. 8) 5 to 6 d after transferring the prezoosporangia into seawater. Mean (± SE) zoospore length was 4.4 ± 0.18 µm (N = 50, range = 2.9 to 5.8 µm).

The mean diameter of prezoosporangia of the *Perkinsus*-like parasite isolated from *Ostrea edulis* in the second trial was 167.1 ± 8.09 µm (N = 24, range = 110 to 317 µm). In addition to its larger size, the cell wall was thicker than in the first trial, and 4% (N = 250) of the zoosporangia had 2 discharge tubes (Figs. 9 & 10) instead of a single tube. The distance between the 2 tubes was variable and both tubes were similar in length. As in the first trial, zoosporulation progressed through successive karyokinesis and cytokinesis, and produced hundreds of zoospores that were liberated 5 to 6 d after transfer of the prezoosporangia into seawater.

Ultrastructural aspects of the zoosporulation sequence and zoospore morphology are shown in Figs. 11 to 17. In the first stages of zoosporulation, zoosporangia contained few and large prezoospores (Figs. 11). Prezoospore cytoplasm had numerous vacuoles, vacuoplasts, mitochondria and lipid droplets (Fig. 11). A vacuoplast was frequently visible in the vacuoles of prezoospores and zoospores (Figs. 11 to 17). Asynchrony was observed in the zoosporulation process, with large prezoospores that had not completed the division process and mature zoospores occurring simultaneously in zoosporangia (Fig. 12). Zoosporangia had a thick cell wall (9 to 10 µm), with 2 layers. The outer layer had an external glycolycal-like structure and 2 thin, parallel, electron-dense layers (Fig. 13), the inner layer being the more electron-dense (Fig. 14). Zoosporangia plugs and discharge tubes were formed from the inner layer of the wall (Fig. 14). Lomosomes were not observed in the zoosporangia wall.

Zoospores were uninucleated and biflagellated cells (Figs. 15 to 17). Zoospore shape was ellipsoidal to
Figs. 1 to 4. Light micrographs of oyster ostrea edulis tissues infected by Perkinsus mediterraneus n. sp. Fig. 1. Trophozoite covered by wall-like structure (arrowheads) in connective tissue of oyster visceral mass (N: nucleus; V: vacuole). Fig. 2. Trophozoite in area of oyster connective tissue infiltrated by haemocytes, showing parasite surrounded by wall-like structure (arrowhead) and vacuoplast (Va) inside large vacuole. Fig. 3. Multicellular stage resulting from vegetative multiplication of a mother cell with daughter cells in a rosette-like arrangement inside wall-like structure covering mother cell. Fig. 4. Later stage of vegetative multiplication, with individual daughter cells inside structure covering mother cell (arrowheads). All scale bars = 5 µm.
ovoid, with tapered anterior and rounded posterior ends. Both flagella were inserted laterally in the anterior part of the cell (Fig. 15) and had the typical eukaryotic microtubular arrangement of 9 doublets + 2. There was an electron-dense body inside the basal bodies (Fig. 15). A large nucleus was located in the posterior part of the cell, with the chromatin condensed at the periphery and lacking a patent nucleolus (Figs. 16 & 17). The mitochondria (or mitochondrion) extended from the apical to the posterior part of the cell in a lateral position (Figs. 16 & 17). Mitochondria were of a bar-like shape, with tubular cristae in the periphery and a wide matrix (Fig. 17). Several vacuoles were scattered throughout the cytoplasm. Other zoospore structures described in *Perkinsus* spp. also were found. The apical ribbon was open-sided and located in the apical part of the cell (Fig. 17). The ribbon-associated vesicles were moderately electron-dense vesicular structures deployed in bundles with a wavy appearance (Figs. 16 & 17). The ribbon-associated vesicles extended from the apical ribbon to the posterior region of the zoospores, wrapped around the nucleus, and then extended anteriorly. Toxicysts comprised a bundle of vesicular structures that extended from the apical ribbon to the posterior portion of the cell, maintaining a straight appearance and ending in a knob-like structure (Figs. 15 to 17). The rhoptries were vaso-shaped electron-dense structures, and extended from the apical ribbon area to the posterior portion of the zoospores (Figs. 15 to 17). Cortical alveoli were observed in the anterior part of the zoospore surface (Fig. 17).

**DNA-sequencing**

A total of 9 ssu/ITS and 10 ITS DNA clones were sequenced. BLAST searches demonstrated that all the sequences obtained by PCR amplification with both 16S-A and PerktSS320comp and the *Perkinsus*-specific ‘D’ ITS primers most closely matched *Perkinsus* sequences in GenBank. The closest matches were to the *P. atlanticus* and *P. olseni* sequences. The sequence of the ITS portion of the ssu/ITS amplifica-
Figs. 11 to 15. *Perkinsus mediterraneus* n.sp. Transmission electron micrographs showing zoosporulation process. Fig. 11. Zoosporangium in early stage of zoosporulation containing large prezoospores with mitochondria (M), lipid droplets (L) and numerous vacuoles (V) and vacuoplasts (Va) (scale bar = 2 µm). Fig. 12. Co-occurrence of prezoospores (Pz) and zoospores (Z) inside a zoosporangium, denoting asynchronous division (scale bar = 2 µm). Fig. 13. Enlargement of zoosporangium wall (W), showing outer layer and glycocalix-like structure (G) (scale bar = 0.5 µm). Fig. 14. Enlargement of discharge-pore area of zoosporangium, showing a plug blocking the pore; wall is divided into outer layer (OW) and electron-dense inner layer (IW); zoospores are visible inside zoosporangium (scale bar = 2 µm). Fig. 15. Zoospores inside zoosporangium showing nucleus (N), flagella (F), basal bodies (BB) and other zoospore structures (DB: electron-dense body; R: rhoptry; T: toxicyst) (scale bar = 1 µm)
tion product matched the ITS sequences obtained in the 'D' primer amplifications.

We found 10 polymorphic sites within the 1786 bp ssu rRNA gene fragment among the DNA clones for the Ostrea edulis Perkinsus species. Among the ssu sequences available for all Perkinsus species, only 26 sites were parsimony-informative. Parsimony jackknife analysis of the ssu rRNA gene resulted in a tree (Fig. 18) that clearly separated the Crassostrea virginica and Tapes decussatus sequences from Perkinsus spp. sequences. ssu sequences from the O. edulis Perkinsus sp. did not group with host sequences, but with those of Perkinsus spp. with 100% jackknife support, indicating that the parasite DNA was targeted for amplification. There was relatively poor resolution, however, among Perkinsus ssu sequences. The Perkinsus sp. from Mercenaria mercenaria grouped with P. chesapeakei and P. andrewsi with 90% jackknife support. The Perkinsus sp. from Mya arenaria and P. marinus grouped together with 96% jackknife support. The Perkinsus sp. from Anadara trapezia, P. atlanticus and the O. edulis Perkinsus sp. were unresolved by the ssu analysis.

There were 12 polymorphic sites among the 10 DNA clones for the 643 bp ITS region fragment obtained for the Ostrea edulis Perkinsus sp.; 144 sites were parsimony-informative for the phylogenetic analysis based on Perkinsus spp. ITS sequences. All the O. edulis Perkinsus sp. ITS sequences formed a monophyletic group with 100% jackknife support (Fig. 19) comprising a sister group to a clade containing ITS region sequences from P. atlanticus, P. olseni, and Perkinsus sp. from Anadara trapezia and Perkinsus sp. from Chama pacificus. P. marinus sequences formed a monophyletic group with 100% jackknife support that comprised a sister group to the larger clade containing the P. atlanticus/P. olseni and the O. edulis Perkinsus sp. sequences.

**Influence of RFTM incubation period on Perkinsus atlanticus prezoosporangium size**

The size increases of Perkinsus atlanticus cells in RFTM and prezoosporangium viability are shown in Table 1. Prezoosporangium size after 7 d incubation.
Casas et al.: *Perkinsus mediterraneus* n. sp. parasite of *Ostrea edulis* was significantly smaller than after the other 2 longer (14 and 21 d) incubation periods, but mean prezoosporangium size did not reach 40 µm during any incubation period. Mortality was 100% for each replicate at the longest incubation period.

**DISCUSSION**

A *Perkinsus*-like parasite has been detected in tissues of *Ostrea edulis* from the Balearic Islands, in the Mediterranean Sea. The morphology of the different stages of the parasite and of its zoosporulation stages resemble those of other *Perkinsus* species (Mackin et al. 1950, Lester & Davis 1981, Azevedo 1989, Blackbourn et al. 1998, McLaughlin et al. 2000, Coss et al. 2001a). The parasitic cells in the oyster tissues increased in size after incubation in RFTM. Incubation of host tissues in RFTM is a widely used diagnostic technique for *Perkinsus* spp. (Ray 1966, Bushek et al. 1994, Rodríguez & Navas 1995). One of the authors (A. G.) has used it as a routine technique to estimate the prevalence of this parasite in oysters at growing sites in the study area, and, using light microscopy, has observed the genus typical dark spheres that become visible after immersion of RFTM-incubated tissue in Lugol's solution. She found that up to 70% of the oysters were infected in autumn samples. Some morphological characters and the sequence of the analysed DNA regions of the European flat oyster parasite are, however, different from those of the species currently included in the genus *Perkinsus*.

Trophozoite morphology is of limited taxonomic value, because it can be influenced by the host (Goggin & Lester 1995), the time of year (Ray & Chandler 1955, Bushek et al. 1994) and nutrient availability (La Peyre et al. 1993, La Peyre & Faisal 1996, Casas et al. 2002b).

Some differences have been reported in the morphological characters of the prezoosporangium/zoosporangium among *Perkinsus* spp., such as prezoosporangium size, ratio between discharge tube length and zoosporangial body diameter, and number of discharge tubes. The morphology of the prezoosporangium/zoosporangium of the European flat oyster parasite did not match with any group of

<table>
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<th>Incubation (d)</th>
<th><em>P. atlanticus</em> Viability %</th>
<th><em>P. mediterraneus</em> Size (µm)</th>
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<tr>
<td>7</td>
<td>82</td>
<td>35.2 ± 0.85</td>
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<tr>
<td>14</td>
<td>5</td>
<td>39.8 ± 0.68</td>
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<td>21</td>
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<td>39.4 ± 1.37</td>
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Table 1. *Perkinsus* spp. Mean size (± SE) and viability of *P. atlanticus* prezoosporangia after incubation of *Tapes decussatus*-infected gills in RFTM for 7, 14 and 21 d. Size of *P. mediterraneus* prezoosporangia included for comparison. nd: no data.
features described for other *Perkinsus* spp. The mean diameter of the prezoosporangium of the European flat oyster parasite (97 ± 1.99 µm; range = 62 to 135 µm) after 1 wk incubation in RFTM was much greater than that of *P. olseni* (mean = 74 µm; range = 56 to 94 µm) (Lester & Davis 1981), *P. marinus* (‘often in the 30 to 80 µm range’) (Perkins 1996), *P. chesapeakei* (mean = 69.3 ± 29.3 µm; range = 20 to 135 µm) (McLaughlin et al. 2000) and *P. andrewsi* (mean = 67 ± 12 µm) (Coss et al. 2001a). Ray (1952) reported individual *P. marinus* prezoosporangium diameters of 100 to 150 µm after an RTM incubation of 1 wk or slightly longer, with values of up to 280 µm being recorded. The maximum prezoosporangium diameter of the European flat oyster parasite incubated for 2 wk in RFTM was 317 µm. Although there are differences in prezoosporangium size between the European flat oyster parasite and other *Perkinsus* spp., the taxonomic value of these differences has not been established. Host and incubation conditions could affect prezoosporangium size. A study is necessary to evaluate the taxonomic value of this feature by comparing the various *Perkinsus* spp. under identical experimental conditions.

The discharge-tube length is one-sixth the zoosporangial body diameter in the European flat oyster parasite, one-fifth in *P. marinus*, one-quarter in *P. olseni* from *Haliotis laevigata*, one-third in *Perkinsus* spp. from *Tridacna gigas*, *Anadara trapezia* and *Macoma balthica* (Valiulis & Mackin 1969, Goggin & Lester 1995), and one-third (0.30) in *P. atlanticus* (Casas et al. 2002a), the former being the lowest described for any *Perkinsus* species.

Increasing the length of incubation in RFTM had different effects on the *Perkinsus*-like parasite of *Ostrea edulis* and on *P. atlanticus*, (the latter being the only species of the genus *Perkinsus* found in European waters thus far). Substantial cell-size increases, cell-wall thickening, and higher cell viability were seen in the former, while a significant but smaller cell-size increase and low or zero viability were observed in the latter.

Zoosporangia of the European flat oyster parasite had 1 or (less frequently) 2 discharge tubes. Two tubes
have also been observed in *Perkinsus marinus* (Perkins & Menzel 1966) and *P. chesapeaki* (McLaughlin et al. 2000). Zoosporangia with 2 discharge tubes occurred only after prolonged incubation (2 wk) in RFTM, when the European oyster-parasite cells had reached a large size.

At the ultrastructural level, the zoosporangium of the European flat oyster parasite had a very thick cell wall. This cell consisted of 2 layers, as in *Perkinsus atlanticus* (Azevedo et al. 1990) and *P. chesapeaki* (McLaughlin et al. 2000), whereas that of *P. marinus* has 3 layers (Perkins & Menzel 1967). Lomosomes associated with the zoosporangial wall have been described in all *Perkinsus* species. (Perkins 1969, Azevedo et al. 1990, McLaughlin et al. 2000) except *P. qugwadi* and *P. andrewsi* (Blackbourn et al. 1998, Coss et al. 2001a), but these structures were not observed in the zoosporangium of the European flat oyster parasite. However, a glycocalyx-like structure was observed on the cell-wall surface of the zoosporangium of the latter, a type of structure that has never been reported for *Perkinsus* spp.

The zoospore lengths of the *Ostrea edulis* parasite (4.4 ± 0.18 µm, range = 2.9 to 5.8 µm) was within the range of *Perkinsus* spp.: 4 to 6 µm in *P. marinus* (Perkins & Menzel 1967), 4.5 ± 0.6 µm in *P. atlanticus* (Azevedo 1989), 4.5 ± 1.0 µm in *P. qugwadi* (Blackbourn et al. 1998), 4.4 ± 0.6 µm in *P. andrewsi* and 3.73 ± 0.48 µm in *P. chesapeaki* (McLaughlin et al. 2000). The fine structure of the zoospores of the *O. edulis* parasite also corresponded to that of *Perkinsus* spp. The angle forming the flagella at their insertion (which has received taxonomic attention: Perkins & Menzel 1967, Azevedo 1989, Blackbourn et al. 1998, McLaughlin et al. 2000) could not be measured because of the lack of satisfactory images. An electron-dense body in the basal body that has been described in all *Perkinsus* spp. (Perkins & Menzel 1967, Azevedo 1989, Blackbourn et al. 1998, McLaughlin et al. 2000) could not be resolved further, supporting the recently proposed synonymy of these 2 species (Murrell et al. 2002). In addition, *P. chesapeaki* and *P. andrewsi* ITS sequences, together with sequences from clonal cultures of *Perkinsus* sp. isolated from *Mya arenaria* and *Tagelus plebeius*, form a monophyletic clade, and a recent study has suggested that these 2 species may also be synonymous (Dungan et al. 2002). The 10 ITS region sequences obtained in our study suggest, however, that the *Perkinsus* sp. from *O. edulis* is closely related to, yet distinct from, *P. atlanticus* and *P. olseni*. Although we combined parasite zoosporangia and zoospores from 20 different host-individuals for the DNA analysis, all 10 ITS sequences grouped together, while none of these ITS sequences grouped with sequences from already described *Perkinsus* species, strongly suggesting that only 1 *Perkinsus* species was isolated from the infected *O. edulis* used in this study. Although our relatively small dataset of ITS sequences from the *O. edulis* parasite may have missed some intra-specific variation, there is a very large dataset of ITS sequences for *P. olseni*/*atlanticus* and none of those sequences overlapped with our new ITS sequences. Reciprocal monophyly of the clades based on the molecular data supports our claim that this is a new *Perkinsus* species.

Gross morphological and ultrastructural examination revealed a few differences between *Perkinsus* sp. from *Ostrea edulis* and the other *Perkinsus* spp., with the molecular study revealing that *Perkinsus* sp. from *Ostrea edulis* is different from *P. marinus*, *P. olseni*/*atlanticus*, *P. chesapeaki*/*andrewsi*, and *P. qugwadi*. Thus, we conclude that the parasite of *Ostrea edulis* is a new species of the genus *Perkinsus*, and propose to name it *Perkinsus mediterraneus* n. sp.

The response of the *Ostrea edulis* cellular defence system against the trophozoites involved haemocytic infiltration of affected tissue areas and phagocytosis of parasite cells. Parasite encapsulation by a few oyster haemocytes was observed less frequently. Similarly, the most evident defence mechanism of another oyster mollusc, *Crassostrea virginica*, against *Perkinsus marinus* was phagocytosis, although attempts at encapsulation of trophozoites by oyster haemocytes were occasionally seen (Mackin 1951, La Peyre et al. 2000). Zoosporangia with 2 discharge tubes occurred only after prolonged incubation (2 wk) in RFTM, when the European oyster-parasite cells had reached a large size.
Nos. for ssu rRNA gene sequences: AY486139, AY486140, AY486141, AY487831, AY487832, AY487833, AY517645, AY517646, AY517647. Accession Nos. for ITS region sequences: AY487834 to AY487843.

**Etymology.** Specific epithet derives from geographical area where the parasite was found, the Mediterranean Sea.

**Acknowledgements.** E. Penas, M. I. Meléndez and L. Corral provided technical assistance, J. M. Valencia sampled the oysters. This work was partially supported by funds of the Secretaría Xeral de Investigación e Desenvolvemento Tecnolóxico da Xunta de Galicia, through the project PGIDT-CIMA 99/10, and by funds of the Planes Nacionales JACUMAR 1999. S.M.C. was supported by a scholarship from the Consellería de Pesca e Asuntos Marítimos da Xunta de Galicia. This is VIMS contribution number 2579.

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Editorial responsibility: Albert Sparks,
Seattle, Washington, USA

Submitted: March 19, 2003; Accepted: October 10, 2003
Proofs received from author(s): January 19, 2004