Binding specificities of mono- and polyclonal antibodies to the protozoan oyster pathogen

*Perkinsus marinus*

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**ABSTRACT:** In an effort to develop direct detection methods for *Perkinsus marinus* cells in host tissue and environmental samples, the production of both mono- and polyclonal antibodies specific for this apicomplexan oyster pathogen was attempted. A particulate hypnospore immunogen was isolated from infected oyster hemolymph following incubation in Ray's fluid thioglycollate medium. Polyclonal antisera produced in both mice and rabbits following immunization with this preparation exhibited high antibody titers for pathogen cell epitopes, but not for host oyster tissue epitopes. At least some antibodies to hypnospore antigenic determinants recognized common epitopes on trophozoite cells in infected oyster tissues, as well as on zoospores and other proliferative cells produced by incubation of infected oyster hemolymph in sterile seawater. Polyclonal antibodies also recognized a diffusible, noncellular substance present in host oyster tissues in areas surrounding focal lesions. Rabbit polyclonal antibodies did not recognize *Dermocystidium* species infecting salmonid fishes or *Haplosporidium nelsoni* infecting eastern oysters *Crassostrea virginica*. These antibodies did recognize many, but not all, *Perkinsus* species infecting selected molluscan hosts worldwide. Monoclonal antibodies produced to date apparently recognize epitopes unique to the hypnospore immunogen.

**INTRODUCTION**

Perkinsiosis caused by the apicomplexan protozoan *Perkinsus marinus* is currently the most widespread and lethal infectious disease of the eastern oyster, *Crassostrea virginica*. In spite of many investigations spanning some 40 yr, neither the pathogenesis of the disease, the life cycle of the pathogen, nor the dynamics and mechanisms of its dissemination are completely understood (Andrews 1988). At least 3 distinct cell types (trophozoites or aplanospores, prezoosporangia or hypnospores, and planonts or zoospores) are recognized as stages in the pathogen's life cycle. Not all stages are obligatory to maintenance or transmission of infections (Perkins 1988), and experimental laboratory infections have been reported following exposure of uninfected oysters to both the trophozoite stage (Mackin 1962) and the zoospore stage (Perkins 1988). Transmission of the disease to uninfected oysters has also been documented following exposure to infected *Boonea impressa*, a gastropod ectoparasite which may function as a vector (White et al. 1987).

Histological examination of fixed host tissues, or incubation of fresh tissues in a fluid thioglycollate-based medium, followed by staining and microscopic observation (Ray 1963), are the only currently available methods for detection of *Perkinsus marinus*. Of these 2 methods, some variation of Ray's fluid thioglycollate medium (RFTM) incubation method is usually favored as a diagnostic tool due to its greater sensitivity and relative ease by comparison to histological methods. Recent modifications of this assay which utilize a small hemolymph tissue sample provide a novel opportunity for nonacute determination of the disease status of live oysters (Gauthier & Fisher 1990).

The utility of Ray's method is limited to detection of pathogen cells which will enlarge in RFTM. In the absence of independent verification methods, Ray's (1952) hypotheses that (1) all *Perkinsus marinus* cells enlarge, but (2) do not proliferate, during incubation have not been rigorously tested. Therefore, estimates of parasite tissue burdens by enumeration of RFTM-induced hypnospores in host tissue samples (Mackin 1962, Choi et al. 1989, Gauthier & Fisher 1990) remain tenuous.
In the higher latitude portions of the pathogen’s range, overwintering tissue stages are thought to be refractile to enlargement induction in RFTM, and seasonal variability in the sensitivity of the assay is thought to underestimate population disease prevalences from samples taken during winter months (Andrews 1988). The efficacy of Ray’s method for detection of Perkinsus marinus zoospores, or other dispersal stages thought to occur in enzootic waters, has not been established; no method exists for direct detection and enumeration of infectious particles in environmental samples.

In an effort to overcome the limitations of existing methodologies, the production of antibodies for direct detection of this pathogen was attempted. It was anticipated that at least some pathogen-specific antigenic determinants would be expressed by all stages and pathogen cell types, and that antibodies binding to common epitopes would universally detect Perkinsus marinus cells, cell fragments, or extracellular products. The availability of specific antibodies is expected to permit definitive studies of the pathogen life cycle and the mechanisms of disease transmission by providing a direct means for localization, quantification, and purification of pathogen cells or components in experimental systems. Such antibodies are also expected to provide the basis for development of sensitive diagnostic immunocassays for direct detection and enumeration of pathogen cells in both host tissue and environmental samples.

Development of monoclonal antibodies (MABs) with specificity for universal epitopes common to all Perkinsus marinus cell types will provide investigators with unlimited quantities of standardized immunodetection reagents. Development of MABs specific for unique epitopes present on specific P. marinus cell types or subcellular components will provide useful reagents for studies of biochemical composition, gene expression, and physiology of various cell types postulated to function in the pathogen life cycle.

One recent report describes the failure of antiserum produced by immunization of rabbits with a soluble Perkinsus marinus hypnospore fraction to recognize in vivo pathogen cell types (Choi et al. 1991). The present report describes a method for isolation of a particulate hypnospore immunogen from the hemolymph of infected oysters, and its use in production of both mouse and rabbit polyclonal antibodies which bind to all known P. marinus cell types. Results of attempts to produce murine monoclonal antibodies with similar binding specificities are presented. The binding specificity of rabbit polyclonal antibodies for selected protozoan pathogens infecting a variety of aquatic hosts worldwide is also described.

**MATERIALS AND METHODS**

**Oysters.** Adult Crassostrea virginica used in this study were collected from natural populations around Chesapeake Bay, USA, during the course of annual disease surveys conducted by the Maryland Department of Natural Resources from 1989 to 1991. Random, 30-oyster subsamples from all populations were assayed for the presence of Perkinsus marinus infections by independently incubating both rectal and hemolymph tissue samples from each sampled individual in RFTM, followed by observation for enlarged pathogen cells.

Uninfected oysters were collected from a tributary of the Potomic River which had been historically free of Perkinsus marinus infections. Ten numbered oysters were aseptically dissected and several types of samples retained. Rectal and hemolymph tissues were assayed for P. marinus, as described above. A sample for histological examination was processed by standard methods. Approximate 1.0 g portions of ventral mantle tissue were individually frozen. Tissues from individual oysters in which no evidence of P. marinus infection was detected were employed as negative controls in assays described below.

**Pathogen isolation and purification.** Oysters from populations with high prevalences of heavily infected individuals were numbered, drilled at the valve margins adjacent to the adductor muscle, and placed in recirculating aquaria for 24 h prior to sampling. After dipping the drilled valve margin in a 10% (v/v) commercial bleach solution, a 1.0 to 3.0 ml hemolymph sample was taken from the adductor muscle sinus of each individual, using a sterile syringe fitted with an 18 gauge needle.

Hemolymph was distributed at 0.5 ml well⁻¹ into sterile, 24-well tissue culture plates. To the hemolymph sample in each well was added 1.5 ml of RFTM supplemented with 200 units ml⁻¹ of nystatin, 500 units ml⁻¹ of penicillin, and 500 μg ml⁻¹ of streptomycin. Plates were sealed with sterile mylar plate sealers, incubated at 25°C for 5 to 7 d, and all wells were observed daily with an inverted microscope. Wells containing large numbers of Perkinsus marinus hypnospores, and which remained free of contaminating protozoans, fungi or bacteria, were selected for harvesting.

Mylar plate sealers were wiped with 95% EtOH. Hypnospores were then harvested by puncturing the mylar film with an 18 gauge needle mounted on a sterile syringe, and using the plunger to triturate the contents of the wells. The resulting suspensions were transferred to sterile, centrifuge tubes. Harvested wells were washed once with a sterile centrifugation buffer (EPBS, pH 7.5) containing 0.1 M sodium phosphate, 0.2 M sodium chloride, and 1.0 mM disodium ethy-
lenediaminetetraacetate (EDTA). Cells harvested by washing were pooled with other harvested cell suspensions.

Hypnospores were separated from host oyster components by repeated differential centrifugation at decreasing centrifugal force, using a modification of the methods of Chu & Greene (1989). Supernates were always discarded, and cell pellets were resuspended in approximately 20 volumes of sterile EPBS. Briefly, the suspension harvested from wells was pelleted for 8 min at 750 × g. The pellet was then washed 3 times at 475 × g for 5 min, 3 times at 280 × g for 3 min, and 3 times at 150 × g for 3 min.

The washed pellet was resuspended in 20 volumes of EPBS containing 2.0% (w/v) paraformaldehyde and fixed at 4 °C overnight. Following fixation, cells were pelleted at 240 × g for 3 min, and washed 3 times in EPBS at 150 × g for 3 min. Prior to the last wash, the number of hypnospores in suspension was estimated by a hemacytometer count. Cells were washed twice in sterile, 0.85% NaCl for 3 min at 150 × g. The pellet was resuspended in sterile 0.85% NaCl to give a final concentration of 1 × 10^6 or 1 × 10^7 cells ml^{-1}, aliquoted in sterile 1.0 ml cryotubes, and frozen at −70 °C until used for immunizations. For staining, samples of hypnospore immunogen cells were allowed to settle onto the surface of poly-L-lysine-coated microscope slides, fixed, washed, and stored at 4 °C in EPBS.

Hypnospores for use as ELISA plate-coating antigen were recovered from oysters whose sampled hemolymph contained high concentrations of pathogen cells. These oysters were aseptically separated from their valves, dipped in 10% (v/v) commercial bleach for surface decontamination, immersed whole in 20 ml of antimicrobial-supplemented RFTM in sterile, 50 ml, polyethylene tubes, and incubated at 20 °C for 10 to 14 d. Oyster meats and culture medium were then pooled and homogenized for 15 s in a sterile blender. The homogenate was passed through 3 sterile, stainless steel screens (500, 250, and 150 µm mesh openings, respectively), and the filtrate processed as for hypnospores from hemolymph, with the following differences. Following fixation, hypnospores were washed 3 times in EPBS at 150 × g, quantified, aliquoted at 1 × 10^6 cells ml^{-1}, and stored at 4 °C in EPBS containing 0.02% (w/v) sodium azide.

For production of motile zoospores, as well as non-motile proliferative cells, 200 to 500 µl of infected oyster hemolymph were added to tissue culture flasks containing 5.0 ml of 25 ppt, sterile, artificial seawater (SASW) fortified with 500 units ml^{-1} of penicillin and 500 µg ml^{-1} of streptomycin. Flasks were incubated unsealed at 25 °C. Motile zoospores and other non-motile proliferative cells were harvested by removal and replacement of SASW in flasks. Harvested cells were immobilized by filtration onto the surface of a poly-L-lysine-coated, 1.0 µm pore size, black polycarbonate membrane filter (Poretics, Livermore, CA, USA) and fixed with 2.0% (w/v) paraformaldehyde in EPBS. Alternatively, culture water containing motile cells was placed in a poly-L-lysine-coated chamber slide and held at 4 °C overnight to immobilize cells prior to fixation.

**Immunization and antiserum production.** The hypnospore immunogen was thawed, vortexed, and either diluted with sterile 0.85% NaCl, flocculated with Al(OH)_3 (Harlow & Lane 1988), or emulsified with one volume of Freund's Complete Adjuvant (FCA) before injection. Two New Zealand white rabbits were immunized with *Perkinsus marinus* hypnospores as follows:

**Day 1:** Prebleed, 10 ml. Inject with 1.0 ml of FCA/hypnospore emulsion containing 3.0 × 10^5 cells, and administered in 2 scapular subcutaneous (sc) injections, and 2 hindquarter intramuscular (im) injections.

**Day 14:** Boost, sc, with 0.5 ml Al(OH)_3-flocculated hypnospores at 6.0 × 10^5 cells ml^{-1}.

**Day 42:** Boost, sc, with 0.5 ml Al(OH)_3-flocculated hypnospores at 6.0 × 10^5 cells ml^{-1}.

**Day 47:** Test bleed, 25 ml.

**Day 48:** Test bleed, 25 ml.

Rabbits were boosted and bled on a monthly cycle until 300 ml of serum had been collected and stored at −70 °C. Serum antibody titers for both *P. marinus* hypnospore and normal oyster tissue epitopes were determined for each serum sample by endpoint dilution ELISA assays. Preimmune sera were used as controls.

**Monoclonal antibody production.** BALB/c mice were hyperimmunized with the hypnospore immunogen until serum antibody titers to the immunogen exceeded 1 : 5000 in the hybridoma screening assay. Initial intraperitoneal (ip) injection of 1.0 × 10^5 cells in FCA, followed by 2 ip boosts with 1.0 to 3.0 × 10^5 cells in sterile saline, failed to produce the desired serum titers. Subsequent sc boosts, both with and without FCA, resulted in elevated antibody titers. A prefusion, sc/ip boost of 4.0 × 10^5 cells was administered 4 d prior to each of 2 separate fusions. Blood collection from splenocyte donor mice was maximized at splenectomy, and recovered serum was employed at a 10^{-3} dilution as a positive control in hybridoma screening assays. Preimmune sera were employed, at the same dilution, as negative controls.

Murine splenocyte/plasmacytoma hybridomas were generated by standard methods (Harlow & Lane 1988).
Splenocytes were fused with either X63-Ag8.653 or SP2/O-Ag14, BALB/c-derived, mouse plasmacytoma cells using polyethylene glycol (PEG 1500) as the fusogen, and a splenocyte:plasmacytoma ratio of 1:1. Fusion products were plated in RPMI-1640 medium supplemented with 20% (v/v) fetal bovine serum, 10% (v/v) Origen\(^*\) (IGEN, Rockville, MD), 1.0 mM oxaloacetate, 0.2 U ml\(^{-1}\) insulin, 0.45 mM pyruvate, 100 \(\mu\)M hypoxanthine, 16 \(\mu\)M thymidine, and 0.4 \(\mu\)M aminopterin, at a density calculated to generate less than one viable hybridoma per well in 96-well tissue culture plates, and incubated in a 5.0% CO\(_2\) atmosphere. Splenocyte feeder cells were not employed in the first fusion (X63-Ag8.653 fusion partner), but were employed in the second fusion (SP2/O-Ag14 fusion partner).

Wells with positive hybridoma growth were screened for the presence of *Perkinsus marinus*-specific monoclonal antibody using a hypnosporae-based ELISA assay. Positive hybridomas were expanded and cloned at least twice by limiting dilution in 96-well tissue culture plates. Cloned hybridomas were expanded and cultured continuously until sufficient conditioned medium had been acquired for evaluation of MABs.

**ELISA assays.** Rabbit serum antibody titers against both normal oyster tissue antigens and *Perkinsus marinus* hypnosporae antigens were estimated by endpoint dilution ELISA assays in which antigens were adsorbed to polystyrene ELISA plates. Normal oyster tissue antigen-coated plates were prepared by homogenizing 1.0 g of normal oyster mantle tissue in 20 \(ml\) of 0.15 M phosphate-buffered saline (PBS), and allowing 50 \(\mu\)l of the homogenate to adsorb to virgin wells for 1 h. To enhance particle retention, hypnosporae-coated plates were precoated for 30 min with 1.0 mg ml\(^{-1}\) poly-L-lysine (molecular weight = 100 000), washed once with PBS, and 5.0 \(\times\) 10\(^3\) hypnosporae in 50 \(\mu\)l of PBS were adsorbed to each well for 1 h.

Following antigen adsorption, unoccupied binding sites on the support plastic were blocked for 1 h with PBS containing 2.0% (w/v) bovine serum albumin (BSA), 0.05% (v/v) Tween-20 and 0.02% (w/v) sodium azide. Plates were washed once with PBS containing 0.05% Tween-20 (PBS-T), and 50 \(\mu\)l of standard rabbit serum dilutions from all test bleeds were applied to wells. Standard rabbit serum dilutions representing \(\log_{10}\) serum dilution factors of \(-2.0, -3.0, -4.0, -4.3, -4.6, -4.9, -5.2,\) and \(-5.5\) were made in PBS containing 1.0% (w/v) BSA (PBS-T/BSA). Dilutions of preimmune sera and serum diluent alone were employed as controls. Serum dilutions were incubated in wells for 1 h at 20 \(^\circ\)C. Wells were washed 3 times with PBS, and bound antibody immunoglobulin was labeled using 50 \(\mu\)l of an affinity-purified, horseradish peroxidase antibody conjugate [0.4 \(\mu\)g Ig ml\(^{-1}\)] incubated for 1 h at 20 \(^\circ\)C. Following 3 washes with PBS, the enzyme conjugate was detected using 0.5 mg ml\(^{-1}\) of \(\alpha\)-phenylenediamine (OPD) and 0.015% (v/v) \(H_2O_2\) in 0.15 M citrate-phosphate buffer (pH 5.0). Color development was stopped after 10 min with 1 volume of 2.0 \(N\) \(H_2SO_4\), and absorbances read at 492 nm. Titers were recorded as the \(-\log_{10}\) of the endpoint serum dilution.

Hybridomas were screened using an antigen/antibody-capture assay in which the capture antigen was *Perkinsus marinus* hypnosporae immobilized as described above. Isotyping of MABs was performed using an antibody/antibody-capture ELISA format and a commercial subisotyping kit (Hyclone, Logan, UT, USA) according to the manufacturer's directions. Monoclonal antibody titers in hybridoma culture media were quantified relative to that of the 10\(^{-3}\) dilution of mouse polyclonal antiserum employed as a positive control. Two-fold dilutions of hybridoma culture media and of the polyclonal control serum were applied to the same capture antibody-coated ELISA plates used in subisotyping assays. Assay conditions were identical to those employed for subisotyping assays, except that a polyclonal enzyme conjugate with specificity for all murine immunoglobulin heavy and light chains was employed for detection. In all MAB ELISA assays, hybridoma-conditioned culture medium was incubated for 12 to 16 h at 4 \(^\circ\)C in ELISA plate wells coated with the capture antibody or antigen, prior to application of 2\(^o\) antibodies.

**Immunofluorescence and immunohistochemical assays.** Tissues from both infected and uninfected oysters were fixed in Davidson's AFA fixative, embedded in paraffin, and sectioned by standard methods. Prior to immunostaining, sections were deparaffinized and rehydrated through a graded ethanol series. After equilibration in PBS, sections were blocked for 30 min. Blocked sections were incubated for 30 to 60 min in rabbit or mouse antiserum diluted to either 10\(^{-2}\) or 10\(^{-3}\) in PBST/BSA, or in undiluted hybridoma culture medium (RPMI-20). The same culture medium, conditioned by the parent plasmacytoma cells used in fusions, was employed as a negative control reagent in MAB assays. Pre-immune sera were employed as negative control reagents in polyclonal antibody assays. Uninfected oyster tissue sections were employed as negative control samples in assays of both types of 1\(^o\) antibodies.

Sections were washed with PBST and incubated with conjugated, species immunoglobulin-specific 2\(^o\) antibodies for 30 min. Incubation of duplicate sections in which blocking buffer was substituted at the 1\(^o\) antibody incubation step was always employed as a control for spurious binding of conjugated 2\(^o\) antibodies, autofluorescence, or endogenous enzyme activity.

For immunofluorescence, FITC-conjugated 2\(^o\) antibodies were used after dilution to 10 \(\mu\)g Ig ml\(^{-1}\) with
PBST/BSA. The FITC-stained slides were washed with PBST, counterstained 30 s with 0.01 % (w/v) Evan’s blue, mounted in buffered glycerol, and observed with an epifluorescence microscope.

For immunohistochemical analyses, horseradish peroxidase-conjugated 2° antibodies were used after dilution to 0.4 μg l mg−1 with PBST/BSA. Sections were washed with PBST and the reporter enzyme detected using cobalt-enhanced diaminobenzidine-4HCl (DAB) at 0.5 mg ml−1 in 0.1 M Tris buffer (pH 7.6) containing 0.01 % H2O2 and 8.0 mM Co(C2H3O2)2 (KPL, Gaithersburg, MD) for 5 min at 20°C. Immunostained sections were counterstained with Mayer’s hematoxylin and eosin (H & E) and coverslipped as permanent preparations.

Cytological preparations of hypnospores, zoospores, and proliferating cells immobilized on membranes or slides were fixed, washed with PBST, blocked, and fluorescence or enzyme immunostained as described above, using similar controls. Membrane-immobilized cells were fixed, rinsed, blocked, and stained by sequential passage of solutions through membranes clamped in the same holders used in the sample collection. Stained membranes were removed from holders and mounted in buffered glycerol on slides for microscopic observation.

Antibody binding to related pathogens. Paraffin blocks containing fixed tissues of aquatic hosts infected with selected protozoan pathogen species were solicited from investigators worldwide. Upon arrival, blocks were sectioned by standard methods and enzyme immunostained as described above, using DAB-cobalt as the reporter chromophore. In addition to the use of known infected and uninfected Crassostrea virginica sections as batch controls, duplicate sections of each solicited sample were processed identically, except that one member of each pair was not exposed to the 1° rabbit antibody. Pathogens in each sample were scored as reactive or nonreactive with the 1° rabbit antibody after comparison of immunostained sections with all positive and negative controls. Host species, pathogen species, references, and sample sources for this material are listed in Table 1.

**RESULTS**

Pathogen isolation and purification

Yields of Perkinsus marinus hypnospores from pooled hemolymph samples averaged 2.2 × 105 hypnospores ml−1, with a range of 1.0 to 2.9 × 105 hypnospores ml−1 of hemolymph, among 5 different groups of oysters which were used. The double harvesting of 24-well plates resulted in an estimated harvest efficiency greater than 95 % of all hypnospores present. Hypnospores adhered tenaciously to both glass and plastic surfaces, in spite of the chelation of divalent cations, and some processing losses resulted. Microscopic observation of purified hypnospore preparations revealed that the only particulates present in the suspension were P. marinus hypnospores. A low proportion of hypnospores harvested from hemolymph incubations was capable of zoosporulation when a sample taken prior to fixation was incubated in 25 ppt SASW at 25°C. Germ tube formation was frequently observed, and actively moving zoospores within sporangium walls were less frequently observed.

<table>
<thead>
<tr>
<th>Pathogen species</th>
<th>Host species</th>
<th>Geographic source</th>
<th>Sample source</th>
<th>Reference</th>
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<tr>
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Motile zoospores were never observed to escape the confining hypnospore wall, and their motility ceased after approximately 10 d. The average yield of *Perkinsus marinus* hypnospores from whole animal tissue homogenates was \(4.4 \times 10^6\) hypnospores oyster\(^{-1}\). Microscopic observation of supernates discarded during centrifugation revealed that a large, but unquantified, percentage of hypnospores remained in suspension, and losses resulted. Microscopic examination of processed hypnospore suspensions revealed that oyster gill bars were present as a minor particulate contaminant in an otherwise pure preparation.

**Immunization and antiserum production**

Serum antibody titers for both *Perkinsus marinus* hypnospore and normal oyster tissue epitopes, which were typical of rabbits injected with the hypnospore immunogen, are shown in Fig. 1. \(\log_{10}\) titers of antibodies binding to pathogen hypnospores increased rapidly and stabilized at 4.6 by the third sample (60 d). Apparent titers for antibodies binding to normal oyster tissue epitopes were measured in preimmune sera, but did not increase with repeated exposure to the hypnospore immunogen, and are interpreted as a nonspecific binding artifact. These data indicate that such rabbit antisera will reliably detect pathogen hypnospores in ELISA assays.

**Monoclonal antibody production**

Of 2 fusion protocols employed, that which utilized the SP2/0-Ag14 parent and splenocyte feeder cells in plating the fusion gave superior results. While the first fusion produced only 10 viable hybridomas, 1 of which was positive in the screening assay, the second fusion produced 512 hybridomas, 25 of which were positive in the screening assay. MABs secreted by stable hybridomas were predominantly of IgM isotype. Despite their consistent ability to bind to pathogen hypnospores in ELISA assays, and immunoglobulin concentrations 1 to 32 times that of a \(10^{-3}\) dilution of the murine polyclonal antiserum successfully used as a positive control in both ELISA and immunohistochemical assays, none of the MABs bound to *in vivo* pathogen cell types when tested by immunostaining of histological sections, and are presumed to recognize epitopes unique to the hypnospore immunogen.

**Immunostaining of *Perkinsus marinus* cells**

Both murine and rabbit polyclonal antisera raised against *Perkinsus marinus* hypnospores specifically bound to *in vivo* pathogen stages in histological sections from infected oysters. A large, mature trophozoite within mantle tissue is shown fluorescence immunostained in Fig. 2. In addition to the pathogen cytoplasm, antibodies strongly labeled the pathogen nuclear membranes and endosome. The eccentric vacuole characteristic of mature trophozoite cells was unstained. Fig. 3 shows a single trophozoite cell which demonstrates the relative intensity of antibody labeling of the pathogen nucleus, and which is internalized within a circulating hemocyte. Fig. 4 shows a 4-cell

![Fig. 1 Typical serum titers of rabbit antibodies binding to Perkinsus marinus hypnospores and to normal Crassostrea virginica tissue epitopes induced by regular injection of a hypnospore immunogen. Titers to hypnospore epitopes rise rapidly and stabilize without evidence of tolerance induction. Apparent low titers to oyster tissue epitopes are confirmed as spurious by their detection in preimmune serum and their failure to increase with exposure to the immunogen. Titers are expressed as \(-\log_{10}\) of the endpoint serum dilution.](image-url)
cluster of pathogen cells which appear to have recently completed plasmotomy within a circulating hemocyte, and in which antibody labeling was confined to cell surfaces. In Fig. 5, immature trophozoite cells forming a rosette in mantle epithelium show paired, fluorescent structures suggestive of karyotomy, within their nuclei.

Low-intensity background staining of normal oyster components in these and other micrographs is due to the red, nonspecific fluorescence of the Evan's blue counterstain, which is easily distinguished from the higher intensity, green fluorescein signal. Remarkably, considering that hemocyte cellular antigens were the

Figs. 2 to 5. Crassostrea virginica infected by Perkinsus marinus. Fluorescence immunostained histological sections of infected oyster tissues counterstained with Evan's blue. Fig. 2. Mature trophozoite in mantle tissue showing antibody binding to pathogen cell cytoplasm, nuclear membranes, and endosome. Typical eccentric vacuole remains unstained. Scale bar = 10 μm. Fig. 3. Trophozoite within a circulating hemocyte showing relative intensity of antibody binding to pathogen nucleus and cytoplasm. Arrow indicates hemocyte nucleus. Scale bar = 5 μm. Fig. 4. Surface-immunostained cluster of pathogen cells within one circulating hemocyte appear to have recently divided. Arrow indicates hemocyte nucleus. Scale bar = 5 μm. Fig. 5. Four-cell rosette of immature trophozoites in mantle epithelium. Paired nuclear endosomes (arrows) in several cells suggest incipient karyotomy. Scale bar = 5 μm.
most likely host component to have copurified with the hypnospore immunogen, no evidence of antibody binding to hemocytes was observed.

In addition to these pathogen cell types, which have been previously described (Mackin 1962, Perkins 1988), one or more epitopes which occurred as small (ca 0.5 μm) particles or as a diffusible, nonparticulate substance were localized in areas immediately surrounding focal lesions, and were stained by rabbit polyclonal antibodies. Whether immunoreactive, noncellular small particulates represent soluble substances coagulated during fixation is not known.

Fig. 6 shows a cross-sectioned digestive gland distributing duct with a focal epithelial lesion and a diffuse infection of the connective tissue at the epithelial basal lamina. Small fluorescent particles were associated especially with the cytoplasm and apical surfaces of epithelial cells involved by the focal lesion. The concentration of these particles in the duct epithelium decreased with distance from the focal lesion, and they were almost entirely absent from the uninvolved epithelium on the opposite side of the duct lumen.

Fig. 7 shows a cross-section of oyster intestine in which the epithelium of one arm was the site of a large focal lesion, while the epithelium of the opposite arm was uninvolved. Small particulate and diffuse nonparticulate fluorescence was seen in epithelial and connective tissues adjacent to the lesion, as well as within the intestinal lumen. This diffuse fluorescent signal decreased with distance from the lesion, although several fluorescent pathogen cells were seen free in the lumen of the uninvolved arm of the intestine. Similar observations of pathogen cells free within the gonoduct lumen (not pictured) suggested several possible routes of pathogen dissemination from infected hosts which would be coupled to normal excretory or spawning activities.

Small fluorescent particles were seen associated with epithelial cells and their apical cilia in a focal lesion of stomach epithelium shown in Fig. 8. Apical cilia and cells near the site of heaviest pathogen concentration showed evidence of degeneration, while cells and their cilia a short distance from the lesion appeared unaffected. The epithelium of the longitudinally sectioned and branching digestive gland distributing duct shown in Fig. 9 was heavily infected, yet the epithelia of absorptive tubules which surrounded the duct, and to which the duct communicates, showed few pathogen cells. The distribution of diffuse fluorescence was limited to connective tissues interstitial between absorptive tubules.

Rabbit antisera also labeled *Perkinsus marinus* cells produced in vitro. Fig. 10 shows immunoperoxidase staining of the hypnospore cell type used as the immunogen in this study. Epitopes labeled by rabbit antibodies appear black in this preparation by the precipitation of DAB-cobalt at sites of antibody/enzyme localization. An enzyme-linked reporter system was chosen over a fluorescent reporter system for this sample because *P. marinus* hypnospores possessed several autofluorescent components. Low-intensity, yellow-green autofluorescence of the cell wall occurred upon excitation with blue light. A higher-intensity autofluorescence by large droplets or granules occurring within the cytoplasm and the eccentric vacuole was common. This second autofluorescent source was excited by a range of wavelengths spanning the spectrum from ultraviolet to green, and its emission wavelength (color) varied as a function of excitation wavelength. Results reported from studies utilizing fluorescent reporter molecules without controls for endogenous sources of sample fluorescence are suspect (e.g. Choi et al. 1991).

Fig. 10 demonstrates that rabbit antibodies bound to the external surface of the cell wall, to the plasma membrane and cell cytoplasm, and to the nuclear membranes and endosome of hypnospore immunogen cells. The strong correlation between dominant structures immunostained in RFTM-incubated hypnospores and in trophozoites present in host tissues suggested a close structural and biochemical homology between these cell types. Variable permeability of the thick hypnospore wall to immunoglobulin molecules was indicated in our preparations by the occurrence of some hypnospore cells showing an absence of immunostained internal elements, but typical immunostaining of the external wall surface.

Figs 6 to 9. *Crasostrea virginica* infected by *Perkinsus marinus*. Fluorescence immunostained histological sections of infected oyster tissues counterstained with Evan's blue. Fig. 6. Cross-sectioned digestive gland distributing duct with a focal, epithelial lesion (arrow). Noncellular immunofluorescence is associated with epithelial cells involved by the lesion, but is absent from uninvolved epithelium opposite the lesion. Scale bar = 50 μm. Fig. 7. Cross-sectioned intestine where the epithelium of one arm is heavily infected, while that of the other arm is uninvolved. Noncellular, diffuse immunofluorescence is seen in the epithelia, intestinal lumen, and connective tissues involved by the lesion. The signal decreases with distance from the lesion. Several fluorescent pathogen cells are free in the intestinal lumen (arrow) away from the lesion. Scale bar = 50 μm. Fig. 8. Focal lesion of stomach epithelium. Only involved epithelial cells and their apical cilia show degenerative changes and minute, noncellular, immunofluorescent particles (arrow). Scale bar = 50 μm. Fig. 9. Longitudinally sectioned and heavily infected digestive gland distributing duct. Epithelia of absorptive tubules surrounding the duct, and with which the duct communicates, are only lightly involved. Scale bar = 100 μm.
Figs. 10 to 13. *Perkinsus marinus* in vitro. Figs. 10. Enzyme immunostained *P. marinus* hypnospore in vitro demonstrating antibody binding to the external wall surface, plasma membrane, cytoplasm, nuclear membranes (arrow), and endosome of an immunogen cell. Immunostained elements are black. Scale bar = 10 µm. Figs. 11 to 13. *P. marinus* proliferative cells produced in vitro following incubation of infected oyster hemolymph in SASW for 30 d. All cells were fluorescence immunostained following immobilization on polycarbonate membranes. Fig. 11. Several 2- to 4-cell division figures showing surface immunostaining and shared membrane systems (arrow). Scale bar = 10 µm. Fig. 12. Multicellular division figure which is surface-immunostained and apparently bound by a common membrane. Scale bar = 10 µm. Fig. 13. Two-cell division figure and an immunostained biflagellate zoospore. Scale bar = 5.0 µm.

Fluorescence immunostained proliferative cells produced in vitro by incubation of infected oyster hemolymph for 30 d in 20 ppt SASW are shown immobilized on polycarbonate membranes in Figs. 11 to 13. Fig. 11 shows several 2-cell and 4-cell figures comprised of variably sized cells which appeared to be
actively dividing. One apparent result of this division process was the production of multiple small cells from single large cells. Presumably, replication and division of genetic material preceded cell division. The presence of an apparent shared external membrane which was immunostained and enveloped several pairs of 2-cell figures suggested a common progenitor cell.

Fig. 12 shows an immunostained multicellular figure (schizont) bound by a common membrane. The restriction of immunostaining to cell surfaces in these preparations suggested a low permeability of cell membranes to immunoglobulin molecules.

Fig. 13 shows a large cell bisected by a division furrow and a small, biflagellate zoospore. Zoospore flagellar morphology was not well demonstrated by either method used for zoospore immobilization. However, numerous similar flagellated cells were present in our immunostained preparations, and were presumed to represent motile cells observed in the SASW sample which was immobilized. These immunostained flagellated cells had dimensions (1.3 × 2.0 μm) which were smaller than those described by Perkins & Menzel (1967) (2–3 × 4–6 μm) for zoospores produced in sporangia following RFTM-incubation of pathogen cells. Our data showed that flagellated cells recognized by rabbit antibodies raised against Perkinsus marinus hypnospores were produced following simple incubation of axenic, infected oyster hemolymph in SASW. If proliferative cells shown immunostained in Figs. 11 & 12 gave rise to motile zoospores, the developmental sequence may differ from that described by Perkins & Menzel (1967), since sporangia were not observed in our preparations.

Table 2. Pathogen species, host species, sample source, and Rabbit α Perkinsus marinus binding reaction for samples of aquatic host tissues infected with selected protozoan pathogen species. For full genus names see Table 1

<table>
<thead>
<tr>
<th>Pathogen species</th>
<th>Host species</th>
<th>Geographic source</th>
<th>Rabbit α P. marinus antibody binding</th>
</tr>
</thead>
<tbody>
<tr>
<td>H. nelsoni</td>
<td>C. virginica</td>
<td>MD, USA</td>
<td>–</td>
</tr>
<tr>
<td>P. marinus</td>
<td>C. virginica</td>
<td>AL, USA</td>
<td>+</td>
</tr>
<tr>
<td>P. marinus</td>
<td>C. virginica</td>
<td>MD, USA</td>
<td>+</td>
</tr>
<tr>
<td>P. marinus</td>
<td>C. virginica</td>
<td>MA, USA</td>
<td>+</td>
</tr>
<tr>
<td>Perkinsus sp.</td>
<td>M. balthica</td>
<td>HI, USA</td>
<td>+</td>
</tr>
<tr>
<td>Perkinsus sp.</td>
<td>M. arenaria</td>
<td>MD, USA</td>
<td>+</td>
</tr>
<tr>
<td>P. atlanticus</td>
<td>R. decussatus</td>
<td>Portugal</td>
<td>+</td>
</tr>
<tr>
<td>P. olseni</td>
<td>H. laevigata</td>
<td>Australia</td>
<td>+</td>
</tr>
<tr>
<td>Perkinsus sp.</td>
<td>T. maxima</td>
<td>Australia</td>
<td>+</td>
</tr>
<tr>
<td>Perkinsus sp.</td>
<td>T. crocea</td>
<td>Australia</td>
<td>+</td>
</tr>
<tr>
<td>Perkinsus sp.</td>
<td>K. rhytiphora</td>
<td>Australia</td>
<td>–</td>
</tr>
<tr>
<td>Perkinsus sp.</td>
<td>C. pacificus</td>
<td>Australia</td>
<td>–</td>
</tr>
<tr>
<td>Perkinsus sp.</td>
<td>A. trapezia</td>
<td>Australia</td>
<td>–</td>
</tr>
<tr>
<td>Perkinsus sp.</td>
<td>A. irritans</td>
<td>NB, Canada</td>
<td>–</td>
</tr>
<tr>
<td>Dermocystidium sp.</td>
<td>S. salar</td>
<td>CA, USA</td>
<td>–</td>
</tr>
<tr>
<td>D. salmonis</td>
<td>O. tshavrytscha</td>
<td>OR, USA</td>
<td>–</td>
</tr>
</tbody>
</table>
Immunostained flagellated cells produced by our methods (1.3 × 2.0 μm) were smaller than those described by Perkins & Menzel (1967) (2–4 × 4–6 μm). Whether this difference in dimensions reflects distinctions in function or ontogeny of motile Perkinsus marinus cells, or whether it is due to differences in nutritional conditions under which zoospore differentiation occurred, is not known. The appearance of nonmotile dividing cells in our preparations as they approach zoospore dimensions suggests that the end result of the proliferative process documented here may be the production of motile cells by a pathway which is distinct from that described by Perkins & Menzel (1967), and which does not require exposure to RFTM. Alternatively, the nonmotile proliferative cells occurring together with motile zoospores in our seawater cultures of infected oyster hemolymph may represent a separate capacity for heterotrophic pathogen proliferation outside the host environment. Whether these cells are infective has not been tested.

Previous attempts to develop Perkinsus marinus-specific rabbit antisera have been made by at least 2 groups. In one effort, the immunogen used was a detergent-solubilized extract of sonicated P. marinus hypnospores, which had been purified from RFTM incubated with infected oyster tissues. Rabbit antisera produced using this immunogen bound to host tissue epitopes, but not to pathogen cells in cryosections of infected host tissues (Choi et al. 1991). A separate effort used whole, washed, but unfixed P. marinus hypnospores isolated from RFTM-incubated oyster tissues following trypsin digestion. Rabbit antisera harvested after multiple injections of 1.0 to 5.0 × 10^6 pathogen cells showed no detectable antibody titer to pathogen epitopes when tested by immunostaining of fixed, infected tissues (E. M. Burreson pers. comm.).

Differences between the results obtained in previous efforts and those reported for the present study may reflect important differences in the methodology used to isolate and purify the immunogen. Since oyster hemocytes die and lyse over several days in RFTM, alternatively the nonmotile proliferative cells are infective has not been tested. Progress towards identifying the antigens of Perkinsus marinus specific rabbit antisera have been made by at least 2 groups. In one effort, the immunogen used was a detergent-solubilized extract of sonicated P. marinus hypnospores, which had been purified from RFTM incubated with infected oyster tissues. Rabbit antisera produced using this immunogen bound to host tissue epitopes, but not to pathogen cells in cryosections of infected host tissues (Choi et al. 1991). A separate effort used whole, washed, but unfixed P. marinus hypnospores isolated from RFTM-incubated oyster tissues following trypsin digestion. Rabbit antisera harvested after multiple injections of 1.0 to 5.0 × 10^6 pathogen cells showed no detectable antibody titer to pathogen epitopes when tested by immunostaining of fixed, infected tissues (E. M. Burreson pers. comm.).

Differences between the results obtained in previous efforts and those reported for the present study may reflect important differences in the methodology used to isolate and purify the immunogen. Since oyster hemocytes die and lyse over several days in RFTM, immunogen pathogen cells were obtained during the present study without recourse to sonication, detergent extraction, or enzymatic digestion. The use of such procedures may affect removal or alteration of important antigenic determinants. The use of infected oyster hemolymph as a source of pathogen cells may allow more thorough separation of pathogen and host components by minimizing both the relative proportion and the diversity of host determinants copurifying with pathogen cells.

Presentation of the immunogen in particulate form was expected to be immunostimulatory. That intracellular antigenic determinants were processed and presented by immunized animal macrophages is evi-
enced by strong binding affinities of polyclonal antisera for intracellular pathogen epitopes. Variability in immunostaining of intracellular epitopes both among the different cell types tested, as well as within single cell types, suggested variable permeability of pathogen cell membranes or walls to immunoglobulins. Staining of intracellular epitopes was most consistent in cells which had been processed histologically. The enhanced availability of intracellular epitopes in histological samples may derive from differences in fixation, organic solvent extraction, exposure to heat, or physical breach.

In the present effort, *Perkinsus marinus* hypnospores were sedimented extensively at low relative centrifugal force, allowing smaller host cell particulates and soluble components to be discarded with supernates. That some normal host components copurified with the hypnospore immunogen is suggested by the measurable titers of rabbit antibodies binding to oyster tissue epitopes in ELISA assays. However, the measurement of identical titers in preimmune sera, and their failure to increase with exposure to the immunogen both confirm that they represent spurious antibody binding at high serum concentrations. While minor antibody binding to apical surfaces of mucus-secreting epithelia was infrequently observed in immunostained histological sections of oyster tissues, antibody binding to hemocytes or soluble hemolymph components was never detected.

Although polyclonal antisera from hyperimmunized splenocyte donor mice used in hybridoma production showed strong binding specificities for both surface and intracellular epitopes of pathogen cells in histological sections, cloned murine hybridomas were found to secrete MABs which recognized epitopes which were apparently unique to the hypnospore immunogen. The binding specificities of these MABs suggest limited utilities in detection of a range of pathogen cell types. Since binding specificities of polyclonal antisera produced in both mice and rabbits indicate the presence of common epitopes shared by *Perkinsus marinus* hypnospores and other pathogen cell types, development of hybridoma screening assays which selectively detect specificities for these common epitopes will permit the generation of MABs with broader utilities.

Results of the present study suggest that several subcellular components of the *Perkinsus marinus* hypnospore cell are good immunogen candidates for further antibody production efforts. Highly immunogenic hypnospore epitopes which appear to be shared by other *P. marinus* cell types are associated with the cell and nuclear membranes, cytoplasmic elements, and the nuclear endosome. Refinement of methods for pathogen cell fractionation will facilitate the development of selective antibody screening assays, and will also permit needed studies of pathogen biochemical composition, physiology, and genetics. Development of methods for *in vitro* propagation of *P. marinus* cells will similarly support such efforts.

Antibody binding specificity is not an absolute quality. Failure of our polyclonal antisera to recognize any epitopes on some tested protozoan pathogens suggests, but does not confirm, complete dissimilarity with *Perkinsus marinus*. Positive binding reactions indicate the presence of one or more epitopes common to *P. marinus* hypnospores. The broad range of *Perkinsus* species with which our antibodies reacted serves to substantiate taxonomic distinction of the group based on morphology and their unique response to RFTM exposure. However, the failure of our antibodies to recognize any epitopes on several *Perkinsus* species indicates that significant antigenic heterogeneity exists among *Perkinsus*-like organisms infecting aquatic molluscan hosts.

Binding specificities of rabbit polyclonal antibodies for various protozoan pathogens infecting molluscan hosts in Chesapeake Bay indicate some degree of serological homogeneity among local *Perkinsus* species. Although morphological (Perkins 1988, Valiulis & Mackin 1969) and epizootiological (Ray 1954) differences have been cited in efforts to differentiate these pathogens, our data do not support a distinction. Failure of our antisera to bind to *Haplospundium nelsoni*, another significant pathogen of *Crassostrea virginica* in Chesapeake and Delaware Bay waters, indicates that these antibodies will be useful in selectively detecting *P. marinus* in diagnostic samples from this geographic area.

Polyclonal antibodies to *Perkinsus marinus* recognized both *P. atlanticus* and *P. olseni*, but failed to bind to *P. karlssonii*, indicating a degree of similarity among the first 3 *Perkinsus* species that is not shared by *P. karlssonii*. Failure of our antibodies to recognize 2 *Dermocystidium* species infecting salmonid fishes further supports other evidence (Perkins 1988, Olson et al. 1991) that at least some *Dermocystidium* species infecting fishes are unrelated to apicomplexans infecting molluscan hosts. Variability in antibody binding among *Perkinsus* species infecting a variety of Australian molluscs indicates that diverse parasite taxa were represented in the samples we examined.

Throughout this report, historical terminology for *Perkinsus marinus* cell types has been used for clarity, in spite of our recognition that many of these terms are incorrectly applied and potentially misleading. In the absence of full knowledge of the *P. marinus* life cycle, and of genetic and functional characteristics of its various cell types, adoption of terminology consistent with the apicomplexan affinities of this organism involves embracing untested assumptions. It is our hope that,
through development of a specific probe for *P. marinus*, we have facilitated the resolution of a comprehensive knowledge of this pathogen’s life cycle, and promoted adoption of a consistent terminology for its stages.

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