NOTE

Enhanced detection of *Enterocytozoon salmonis* (Microspora), an intranuclear microsporean of salmonid fishes, with the Warthin-Starry stain combined with hematoxylin and eosin

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ABSTRACT: *Enterocytozoon salmonis*, an intranuclear microsporean of salmonid fishes, is often difficult to visualize by light microscopy. In paraffin sections stained with hematoxylin and eosin (H&E), the parasite appears as a light staining sphere within the nucleus, and can easily be confused with nucleoli or nuclear vacuoles. Furthermore, the spores of *E. salmonis* are very small and do not form large aggregates, and thus are not easily identified in sections or wet mounts. The Warthin-Starry stain combined with H&E dramatically enhances the detection of the parasite in tissue sections. Using this method, prespore stages stain light brown to black, and spores stain dark brown or black.

KEY WORDS: Microsporea · *Enterocytozoon* · Warthin-Starry stain

The genus *Enterocytozoon* (Enterocytozooidae: Apansporoblastina) contains 2 described species, 1 from humans and 1 from fishes. *E. bieneusi* infects enterocytes of humans with AIDS (Weber et al. 1994). *E. salmonis* infects the nuclei of immature leucocytes of salmonid fishes and is associated with anemia (Elston et al. 1987) and lymphoproliferation (Hedrick et al. 1990, Morrison et al. 1990). We often see the parasite within the nuclei of plasmablasts from chinook salmon *Oncorhynchus tshawytscha* with plasmacytoid leukemia (Kent et al. 1990). Most reports of *E. salmonis* have been from chinook salmon from the Pacific region of North America. The organism has also been found in rainbow trout *O. mykiss* in France (Chilmonczyk et al. 1991), and we have detected *E. salmonis* in paraffin sections of coho salmon *O. kisutch* from Chile. A similar microsporean infects the nuclei of leucocytes of lumpfish *Cyclopterus lumpus*, and also has been associated with lymphoproliferation (i.e. lymphosarcoma) (Mullins et al. 1994).

The spores of *Enterocytozoon* species are very small (about 1 to 2 μm) and do not form large aggregates (e.g. in xenomas). Therefore, detection of these parasites in sections using routine stains such as hematoxylin and eosin (H&E) or in wet mount preparations is difficult. *E. salmonis* is particularly difficult to detect in histological sections because prespore stages appear as light staining spheres within the nucleus and can easily be confused with nucleoli or nuclear vacuoles.

The Warthin-Starry stain, a stain usually used to visualize certain bacteria, has been recommended by a number of researchers for the detection of *E. bieneusi* in intestinal biopsies (Field et al. 1993a, b). We have found that the Warthin-Starry stain combined with Harris' H&E (regressive method) (see Appendix) is an excellent method for enhancing the contrast between *E. salmonis* and nuclear material (Fig. 1). Using this method, prespore stages stain brown, spores are dark brown or black, while nucleoli retain their characteristic basophilic (blue) staining and nuclear spaces remain unstained.

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Fig. 1. *Oncorhynchus tshawytscha*. Plasmeblasts infected with *Enterocytozoon salmonis* in retrobulbar tissue. All photographs are from the same area. Note that many spores (arrows) and more prespore stages (arrowheads) are visible with the Warthin-Starry/H&E (A and B), compared to sections stained only with H&E (C). Scale bar = 10 μm
Kent et al.: Detection of Enterocytozoon salmonis

Appendix. Warthin-Starry stain (adapted from Field et al. 1993b) combined with Harris' H&E regressive method (from Luna 1968)

1. Place paraffin sections on slides.
2. Dry slides, deparaffinize and rehydrate.
3. Rinse unstained slides thoroughly with distilled water, and place slides into freshly prepared 1% silver nitrate for 30 min at 37°C.
4. Wash in distilled water, 3 changes.
5. Place sections in developing solution (see below) in a 60°C water bath until the sections turn light brown (i.e. approximately 2 to 3 min).
6. Rinse thoroughly in distilled water. Then rinse in hot running tap water, then rinse in distilled water.
7. Fix sections in 5% sodium thiosulphate for 3 min.
8. Counter stain with Harris' H&E using the regressive method (Luna 1968). However, decolorize the haematoxylin in acid alcohol (0.1% HCl) for 1 min and stain in eosin for only 15 s.
9. Dehydrate, clear and mount sections.

To minimize background and other staining problems, do not pretreat slides with adhesives, ensure all glassware (including slides) is clean, and use analytical grade reagents.

Reagents

1% Silver nitrate. Dilute 1:10 from stock 10% silver nitrate with acidified water at pH 4.0.

Developing solution. Mix 30 ml water, pH 4.0, with 36 ml 5% gelatin. Preheat above mixture to 60°C. Add 3 ml 10% silver nitrate and 2 ml 1.5% quinol immediately before developing. Quinol solution should be no older than 1 wk.

LITERATURE CITED


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