

NOTE

Kudoa alliardia in flesh of Argentinian hoki *Macruronus magellanicus* (Gadiformes; Merlucciidae)

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ABSTRACT: Myxozoans of the genus *Kudoa* are widespread parasites of marine fishes and primarily infect the body musculature of their hosts. Although *Kudoa* species are not usually associated with host mortality, some do form macroscopic cysts in the tissue and some are associated with post mortem tissue degradation. This is of concern to commercial fisheries as fillets may be unmarketable due to these infections. Because different species of *Kudoa* have different effects on their hosts, it is important to correctly identify species with epidemiological relevance, distinguishing those that are benign from those that are associated with these detrimental effects. Using morphological and molecular analyses, we identified *K. alliardia* infecting Argentinian hoki *Macruronus magellanicus*. Comparisons of the small subunit ribosomal DNA sequence revealed that *K. alliardia* is genetically very similar to *K. rosenbuschi*. Furthermore, there is significant overlap in myxospore dimensions between descriptions of these 2 *Kudoa* species as well as those of other Patagonian fishes. Thus, without careful examination of the myxospore dimensions, it may be difficult to identify these species on a routine basis. It is critical to accurately identify *K. alliardia* as, unlike *K. rosenbuschi*, it is not associated with tissue degradation. Ambiguities in some species descriptions highlight the need for thorough morphological analyses accompanied by molecular comparisons to clarify the species boundaries between *Kudoa* parasites of Patagonian fishes.

KEY WORDS: *Kudoa alliardia* · *Kudoa rosenbuschi* · Myxospore morphology · Patagonia · Argentina · Small subunit ribosomal DNA

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INTRODUCTION

Myxozoan parasites of the genus *Kudoa* (Multi-valvulida: Kudoidae) primarily infect the muscle tissue of their fish hosts (Moran et al. 1999, Swearer & Robertson 1999). Although *Kudoa* species may not be associated with host mortality, the manifestation of these infections by some species (macroscopic cysts, post-mortem tissue degradation) is of concern to commercial fisheries as fillets may be unmarketable. In fishes of Patagonia (southern Chile and Argentina), 5 *Kudoa* species have been identified: *K. alliardia*, *K. clupeiidae*, *K. peruvianus*, *K. ramsayi*, and *K. rosenbuschi* (see

Moran et al. 1999, Kalavati et al. 2000). Post mortem tissue degradation in the musculature of the fish host has been observed for *K. rosenbuschi* (see Sardella et al. 1987) and *K. peruvianus* (see Mateo 1972). Macroscopic cysts have been observed in the muscle of hosts infected with *K. alliardia* and *K. ramsayi* (Kovaleva et al. 1979, Kalavati et al. 2000). *K. clupeiidae* has even been implicated in host mortality (Lom & Dyková 1992).

With the exception of *Kudoa ramsayi*, which has stellate spores (Kalavati et al. 2000), the remaining species bear diminutive quadrate spores that are very similar to one another (Swearer & Robinson 1999). Given that myxozoan spores, when compared to other metazoan

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phyla, possess relatively few features from which identifications and descriptions can be made, it has been suggested that molecular characterizations should accompany descriptions of new species (Kent et al. 2001). Recently, Abollo et al. (2005) sequenced the small subunit ribosomal DNA (SSU rDNA) gene of *K. rosenbuschi*. Based on a phylogenetic analysis of this sequence, Abollo et al. (2005) found that *K. rosenbuschi* was closely related to the morphologically similar parasites *K. clupeidae* and *K. funduli*. In 2002, we received specimens of Argentinian hoki *Macruronus magellanicus* infected with a *Kudoa* species. It was our goal to identify this parasite by both morphological and molecular analyses.

MATERIALS AND METHODS

In August 2002, we received 3 specimens of frozen muscle tissue of Argentinian hoki *Macruronus magellanicus* containing cysts. The tissues had been frozen for up to 2 mo at -20°C . After a brief cytological examination, the specimens were fixed in 95% ethanol and stored for 3 yr before the following more detailed microscopic and molecular investigations were undertaken.

For morphological analyses, digital images were taken of myxospores in wet mount preparations of ethanol-fixed muscle tissue using the SPOT version 3.5.5 for Windows camera and software (SPOT Diagnostic Instruments). Following the guidelines of Lom & Arthur (1989), spore length, width and thickness, and polar capsule width and thickness were measured from multiple images with at least 10 measurements for each dimension.

Sequencing of the partial SSU rDNA employed existing polymerase chain reaction primers, parameters, and sequencing protocols as described previously (Whipps et al. 2003). This sequence was compared to those of other related *Kudoa* species as determined from previous analyses (Blaylock et al. 2004, Whipps et al. 2004, Abollo et al. 2005, Diamant et al. 2005) and using the basic local alignment search tool (BLAST) on GenBank (Altschul et al. 1990). Sequences were aligned with Clustal X (Thompson et al. 1997) and pairwise sequence similarities were generated with BioEdit version 7.0.5 (Hall 1999).

RESULTS AND DISCUSSION

Upon defrosting there was no evidence of myoliquefaction in the affected tissues. Specimen 1 consisted of 3 brown objects, likely cysts, 3 to 6×1 to 2 mm in dimension. Specimen 2 consisted of 4 or 5 sections of

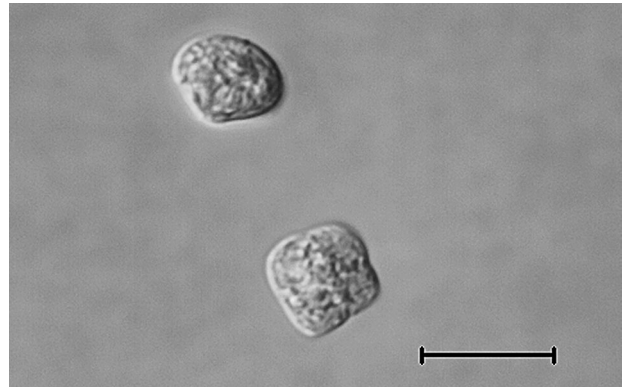


Fig. 1. *Kudoa alliaris*. Spores in lateral (top) and apical (bottom) aspect. Photographed following fixation in ethanol after being frozen at -20°C for up to 2 mo. Scale bar = 10 μm

muscle tissue with numerous yellow to cream, round to oval-shaped cysts, 2 to 3 mm diameter, and 2 to 6 mm long, distributed amongst the muscle fibres. Specimen 3 consisted of 4 or 5 sections of muscle tissue with numerous white, oval-shaped cysts, 2 to 3 mm diameter, and 2 to 10 mm long, distributed amongst the muscle fibres. Although morphology of the cysts in each sample varied to some degree, myxozoan spores consistent with those of the genus *Kudoa* (4 polar capsules and valves) were observed upon microscopic examination of the cyst contents.

Spores were quadrate in apical view, rounded in lateral view, with visible valve sutures (Fig. 1). Mean (range) dimensions of spores from ethanol-fixed tissues were 6.35 (5.84 to 6.86) μm in length, 8.51 (7.84 to 9.25) μm wide, and 7.08 (6.46 to 7.55) μm thick (Table 1). Polar capsules were equal, pyriform, 2.60 μm (2.27 to 2.82) long, and 1.61 μm (1.40 to 1.87) wide. Some spore features (i.e. sporoplasm, polar filaments) were difficult to discern, most likely due to fixation; ethanol fixation is known to cause some shrinkage of myxospores, reducing spore and polar capsule length by approximately 5%, but spore width is less likely to be distorted (Parker & Warner 1970). Thus, measurements made from ethanol-fixed material will be smaller, but very close to those made from fresh material.

Even with the assumption that there was some reduction in spore length due to alcohol fixation, the spores were morphologically most consistent with those of *Kudoa alliaris* (Table 1). Both the locality (Argentina) and host (*Macruronus magellanicus*) were consistent for this species and pseudocysts were white or yellow as described by Kovaleva et al. (1979). The pseudocysts from Specimen 1 were probably brown as a result of the accumulation of melanised breakdown products as the cysts were killed by the host immune response, as seen for other *Kudoa* species (Moran et al. 1999).

Table 1. *Kudoa* spp. Comparison of myxospore dimensions for *Kudoa* species of Patagonian fishes. Mean measurements in μm , range in parentheses

Species	Host species	Locality	Length	Thickness	Width	Source
<i>K. alliaris</i>	<i>Macruronus magellanicus</i>	Argentina	6.35 (5.84–6.86)	7.08 (6.46–7.55)	8.51 (7.84–9.25)	Present study
	<i>Micromesistius australis</i>	Argentina	(7–8)	(8–9)	(9–10)	Kovaleva et al. (1979)
	<i>Patagonotothen ramsay</i>	Argentina	(6.3–7.4)	(6–6.3)	(6.3–7.4)	Kovaleva et al. (1979)
<i>K. clupeiidae</i>	<i>Brevoortia tyrannus</i>	Eastern USA	5.25 (5.0–5.5)	6 (5.5–6.5)	6.5	Reimschuessel et al. (2003)
	<i>Brevoortia tyrannus</i>	Eastern USA	5.1	6.4	–	Meglitsch (1947)
<i>K. peruvianus</i>	<i>Merluccius gayi</i>	Chile	(4.65–5.10)	–	(5.58–6.51)	Mateo (1972)
<i>K. ramsayi</i>	<i>Patagonotothen ramsayi</i>	Argentina	(8.0–10.4)	(2.8–4.8)	(4.8–8.0)	Kalavati et al. (2000)
<i>K. rosenbuschi</i>	<i>Merluccius hubbsi</i>	Argentina	(6–7)	6.5	5.5	Kovaleva et al. (1979)
	<i>Merluccius gayii</i> (sic)	Argentina	–	–	7	Gelormini (1944)

In the original description of *Kudoa alliaris*, Kovaleva et al. (1979) identified 4 different fish species as hosts: *Micromesistius australis*, *Patagonotothen ramsay*, *Notothenia conina*, and *Macruronus magellanicus*. Measurements were only provided for *K. alliaris* from 2 of these hosts (Table 1) and there are large differences in spore dimensions between the two. Kovaleva et al. (1979) suggested that these distinctions were probably connected with various conditions of existence in different hosts. Thus, some of the differences in dimensions observed here for *K. alliaris* from *M. magellanicus* could also be accounted for by variation between hosts. Similar morphological variation between hosts and locations has been shown for *Kudoa iwatai* (Diamant et al. 2005). However, as other studies indicate, it is equally likely that morphologically distinct spores from different hosts can represent separate *Kudoa* species (Whipps et al. 2003, Yokoyama et al. 2004).

Four other *Kudoa* species have been reported from fishes off the coasts of southern Argentina and Chile with morphological traits similar to those of *K. alliaris* (Table 1). *K. peruvianus* possesses smaller spores than *K. alliaris* and is found in waters off of Chile and Peru, but not Argentina. Because this species has significant overlap in dimension with *K. rosenbuschi*, *K. clupeiidae*, (Table 1) and *K. miniauriculata* (see Whitaker et al. 1996), conspecificity of *K. peruvianus* with these species cannot be ruled out. *K. ramsayi* is likely a distinct species as its spores were described as stellate, not quadrate (Kalavati et al. 2000). Nonetheless, *K. ramsayi* and *K. alliaris* share the same host, *Patagonotothen ramsay*, and therefore further study is warranted.

Kudoa clupeiidae is usually found in fishes of the western North Atlantic Ocean (Moran et al. 1999). However, there is a report of this species from *Merluccius hubbsi* in Argentina (Di Antonio & Cenci Goga 1993). Unfortunately, in this report, measurements of spores are not provided and mention is not made of other parasites more commonly observed in this region (*K. alliaris* or *K. rosenbuschi*). Given that this is the

type host and locality for *K. rosenbuschi*, which is morphologically similar to *K. clupeiidae*, it is conceivable that Di Antonio & Cenci Goga (1993) misidentified this parasite.

Such reports highlight the need for detailed descriptions for new species, and thorough identification of existing species when encountered. This is especially important for myxozoans, as spores possess a limited number of characteristics upon which identifications can be made. Furthermore, from reports published prior to the guidelines of Lom & Arthur (1989) it is sometimes unclear as to exactly which spore dimension measurements are based upon. To overcome the limitations of morphology alone, it has been suggested that studies on myxozoan parasites incorporate DNA sequence analyses (Kent et al. 2001). Here, we sequenced most of the SSU rDNA of *Kudoa alliaris*. We obtained a 1680 bp sequence and submitted this to GenBank (accession number DQ182561).

Based on pairwise comparisons, the SSU rDNA sequence of *Kudoa alliaris* was most similar to that of *K. rosenbuschi* (Table 2). In fact, these sequences only differed by 2 nucleotides across the entire sequence length. Other similar sequences were from *K. funduli*, a *Kudoa* sp., and *K. clupeiidae* (all within 2% sequence

Table 2. Identity of *Kudoa alliaris* small subunit ribosomal DNA (SSU rDNA) (GenBank accession number DQ182561) to other *Kudoa* spp. SSU rDNA sequences available on GenBank. Sequence ambiguities were removed prior to calculation

Species	DNA sequence identity to <i>K. alliaris</i>	GenBank acc. no.
<i>K. rosenbuschi</i>	1677/1679 (99.9%)	AY623795
<i>K. funduli</i>	704/713 (98.7%)	AY312279
<i>Kudoa</i> sp.	1542/1564 (98.6%)	AY302723
<i>K. clupeiidae</i>	1291/1311 (98.5%)	AY197771
<i>K. miniauriculata</i>	1525/1564 (97.5%)	AF034639
<i>K. diana</i>	1514/1564 (96.8%)	AF414692
<i>K. paniformis</i>	1505/1564 (96.2%)	AF034640

difference; Table 2). Molnár et al. (2002) also reported high sequence similarities between *Myxobolus* species, with <1% sequence difference between SSU rDNA sequences from the morphologically similar *Myxobolus pseudodispar*, *M. cyprini*, and *M. musculi*. Likewise, *K. minithyrsites* and *K. thyrsites* exhibit only a 1.5% SSU sequence difference from one another (Whipps et al. 2003), and *M. pendula* differs from *M. pellicides* by only 8 bases across approximately 2000 bp of SSU rDNA sequence (Kent et al. 2001).

Conversely, intraspecific SSU sequence differences have also been observed in geographically distant (allopatric) representatives of both *Kudoa amamiensis* and *K. thyrsites*, which have slightly different SSU rDNA sequences (Whipps et al. 2003). Intraspecific SSU sequence variation has even been observed for myxozoans from different host species living in sympatry, such as *Myxidium lieberkuehni* (see Schlegel et al. 1996) and *M. pseudodispar* (see Molnár et al. 2002). Thus, the high sequence similarity of *K. alliarina* to *K. rosenbuschi* does not necessarily equate to conspecificity, nor do the minor sequence differences mean they are separate species.

The distinction between *Kudoa rosenbuschi* and *K. alliarina* is somewhat unclear, especially given the variability between spore dimensions for *K. alliarina* observed here and those of Kovaleva et al. (1979) (Table 1). Both species possess quadrate spores, but *K. rosenbuschi* is reported to have more of a quadrangular platform at the apex of the spore, whereas the apex of *K. alliarina* spores is rounded off (Kovaleva et al. 1979). Thus, the species we observed here had spores most consistent with those of *K. alliarina* (Fig. 1). It is conceivable that such a minor morphological feature could be overlooked, perhaps more so if spores are not viewed laterally. Spores of *K. alliarina* are reported to be wider than those of *K. rosenbuschi* (Table 1). However, the resemblance of *K. alliarina* spores (Fig. 1) to those of *K. rosenbuschi* provided in figures by Sardella (1988) and Abollo et al. (2005) is striking. This is not to say that these authors made a misidentification, but simply illustrates some of the morphological similarities between these parasites.

There are currently no universal criteria, whether they are morphological or molecular, for determining boundaries between myxozoan species. However, consistent differences in these parameters observed in multiple specimens lend strong support to species distinctions. Genetic differences in a few specimens from a conserved gene such as the SSU rDNA are suggestive of an ecological separation, but not necessarily speciation (i.e. different sequences may represent multiple alleles of the same gene). It is important to identify genetically distinct parasite populations and species, especially where parasites are used for identi-

fication of fish stocks (Sardella & Timi 2004). Severity of infection, susceptibility of hosts, and distribution may vary between species. Among the *Kudoa* species discussed here, *K. peruvianus* and *K. rosenbuschi* are known to cause post mortem myoliquefaction of host tissues, whereas *K. alliarina* is not (Moran et al. 1999). *K. clupeiidae* may even be associated with host mortality (Lom & Dyková 1992). Modes of transmission may vary between species as both direct (Diamant 1997) and indirect (Koie et al. 2004) life cycles have been reported for marine myxozoans. Thus, although these species have very similar SSU rDNA sequences and are morphologically similar, their epidemiological relevance to fisheries differs.

It is critical that future studies and species descriptions employ careful morphological analysis of spores for identification of species. Molecular data should be included wherever possible, and may be needed from multiple genes, and/or ethanol-fixed samples should be deposited to a public database for use in future research. Only with thorough analysis of both morphological and molecular characteristics can the ambiguities between the *Kudoa* species of Patagonian fishes be clarified.

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