Fatal pulmonary parafilaroidiasis in a free-ranging subantarctic fur seal (*Arctocephalus tropicalis*) coinfected with two gammaherpesviruses and *Sarcocystis* sp.

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Abstract

A juvenile subantarctic fur seal (*Arctocephalus tropicalis*) found dead in Santa Catarina state, southern Brazil, presented with disseminated verminous pneumonia due to *Parafilaroides* sp. A concomitant infection with two different gammaherpesviruses was identified by PCR in different tissues; one of them possibly a novel species (tentatively named *Otariid herpesvirus* 7). *Sarcocystis* sp. DNA was identified molecularly in skeletal muscle samples with intrasarcoplasmic bradyzoites and no apparent tissue response. All analyzed samples (mandibular, laryngeal, tracheal, and mesenteric lymph nodes, and lung) were PCR-negative for *Brucella* spp. The most likely cause of death was severe pulmonary parafilaroidiasis. The pathogenic role of the gammaherpesviruses in several of the tissues was not evident. This study describes the pathogenicity of *Parafilaroides* sp. in a subantarctic fur seal, widens the host range of herpesvirus in pinnipeds, and reports the first molecular identification of *Sarcocystis* sp. in this species.

Keywords: Herpesvirus, *Parafilaroides* sp., pneumonia, pinniped, protozoan infection, stranding.

Resumo

Um lobo-marinho-subantártico (*Arctocephalus tropicalis*) juvenil foi achado morto no Estado de Santa Catarina, sul do Brasil, apresentando pneumonia parasitária disseminada por *Parafilaroides* sp. Assinfectação concomitante por dois gammaherpesvírus diferentes foi identificada pela PCR em diversos tecidos, um desses herpesvírus possivelmente uma nova espécie (denominada provisoriamente *Otariid herpesvirus* 7). DNA de *Sarcocystis* sp. foi identificado molecularmente em amostras de músculo esquelético que apresentavam bradizoítos intrasarcoplasmáticos e sem resposta tecidual. Todas as amostras analisadas (linfonodo mandibular, laringeal, tracheal, e mesentrico lymh nodes, and lung) foram PCR-negativas para *Brucella* spp. A causa mais provável da morte do animal foi parafilaroides pulmonar severa. O papel patogênico dos gammaherpesvírus em vários tecidos não foi evidente. Este estudo descreve a patogenicidade de *Parafilaroides* sp. em um lobo-marinho-subantártico, amplia a variedade de hospedeiros de herpesvírus em pinipédeos e reporta a primeira identificação molecular de *Sarcocystis* sp. para essa espécie.


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Parasitic respiratory disease by nematodes, including *Otostrongylus circumlitus* (Metastrongyloidea: Crenosomatidae) and *Parafilaroides* spp. (Metastrongyloidea: Filaroididae), are significant causes of morbidity and mortality in free-ranging pinnipeds (DAILEY, 2001; SIEBERT et al., 2007); however, there is limited information concerning *Parafilaroides* infection and associated pathology in South American pinnipeds (JACOBUS et al., 2016).

To date, six gammaherpesvirus species have been described in Otariidae (SACRISTÁN et al., 2018), associated with different clinicopathological presentations, from asymptomatic infections to neoplasia (e.g., urogenital carcinomas) (LIPSCOMB et al., 2000; CORTÉS-HINOJOSA et al., 2016). Nevertheless, only one report describes herpesvirus infection in free-ranging pinnipeds from South America, a South American fur seal (*Arctocephalus tropicalis*) with ulcerative skin lesions (SACRISTÁN et al., 2018). Herein fatal *Parafilaroides* sp. pneumonia in a subantarctic fur seal (*Arctocephalus tropicalis*) coinfect with two different gammaherpesviruses and *Sarcocystis* sp. is described.

A 157 cm-long cachectic juvenile male subantarctic fur seal was found in a state of moderate decomposition (code 3) (GERACI & LOUNSURY, 2005) in Praia do Forte, São Francisco do Sul, Santa Catarina, Brazil (26º14'36" S, 48º38'17" W), on 19 August 2013. The carcass was stored frozen at -20 °C. After thawing at room temperature for several hours, representative tissue samples of the main organs were collected during a complete standard necropsy (GERACI & LOUNSURY, 2005) and preserved in 10% neutral buffered formalin or frozen at -80 °C. Formalin-fixed samples were routinely processed, sectioned at 5 µm and stained with hematoxylin and eosin (H&E) for histopathologic examination.

Samples of cerebral cortex, cerebellum, spinal cord, trachea, lung, heart, esophagus, thyroid, liver, spleen, pancreas, adrenal gland, kidney, testicle, lymph nodes (mandibular, laryngeal, tracheal, mesenteric), urinary bladder, and skeletal muscle were evaluated by universal PCR assays to partially amplify the herpesviral DNA polymerase and glycoprotein B genes (VANDEVANTER et al., 1996; EHLERS et al., 2008).

Samples of mandibular, laryngeal, tracheal, and mesenteric lymph nodes, and lung samples were tested for *Brucella* spp. infection by conventional PCR with primers B4 and B5 targeting a fragment of the *becp31* gene (BAILY et al., 1992). Skeletal muscle was tested for *Sarcocystis* spp. using a nested PCR technique designed to detect DNA fragments of the Toxoplasmatinae ITS-1 region (SOARES et al., 2011), that also amplifies *Sarcocystis* sp. (C. Sacrastán personal observation). Appropriate positive and negative controls (DNA from a previously herpesvirus-positive South American fur seal skin sample, *B. abortus* (vaccine RB-51) and an isolate of *Sarcocystis neurona* were used as positive controls for herpesvirus, *Brucella* spp. and *Sarcocystis* spp., respectively, while Diethylpyrocarbonate (DPEC) water was used as negative control) were selected for all PCR analyses. Positive amplicons were confirmed by direct DNA sequencing.

The obtained forward and reverse sequences were assembled using ClustalW alignment and compared with those available in GenBank/EMBL/DDBJ database using online BLASTn search (http://www.ncbi.nlm.nih.gov/blast). After the alignment of our sequences and those more similar from GenBank/EMBL/DDBJ database using ClustalW, nucleotide and amino acid identities were calculated based on the p-distance. A 1,000 bootstrap replicates maximum likelihood phylogenetic tree of amino acid sequences was generated for herpesviruses based on the MUSCLE alignment of herpesviral glycoprotein B gene from the novel sequences, and sequences published from other pinnipeds, and selected alpha-,
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Grossly, the main findings were lung edema, disseminated parenchymal and subpleural lung nodules containing *Parafilaroides* sp. nematodes (Figura 1a), and skeletal muscle wasting. Additional findings included: a complete fracture of the inferior left canine, unidentified merocercoids compatible with *Phyllobothrium* spp. in the perigenital blubber, left thyroid lobe enlargement, unidentified nematodes in bronchi, ulcerative pylocric gastritis with presence of plastic debris and small numbers of Anisakids (likely to be a species of *Contracecum* (Nematoda) in the stomach, and urinary calculi in the bladder.

Microscopically, the lung had numerous, variably large foci of alveolar emphysema with ruptured walls intercalated with atelectic, congested and edematous areas (Figura 1b), as well as mild to moderate, chronic lymphocytic bronchointerstitial pneumonia with fibrosis and remodeling. Emphysematous foci often had numerous adult and larval nematodes morphologically compatible with *Parafilaroides* sp. and little inflammatory response (Inset in Figura 1b) (GARNER et al., 1997). By contrast, there was moderate, focal inflammatory exudate in areas with L1 larvae (Inset in Figure 1b). Additional microscopic findings were: mild, multifocal, granulocytic dermatitis with mild ballooning degeneration, chronic lymphocytic tracheitis, mild acute skeletal muscle and heart rhabdomyolysis, necrohemorrhagic pylocric gastritis with submucosal microthrombi and vascular necrosis, and multinodal lymphoid reactive hyperplasia. Intrascaplocrinal cysts compatible with *Sarcozystis* sp. were observed in the longissimus dorsi muscle with no inflammatory reaction.

Two different herpesvirus sequences of 453 bp of glycoprotein B gene were obtained: one from laryngeal lymph node, adrenal gland, heart, liver, and cerebral cortex, and another from skeletal muscle, spleen and lung samples (sequences MH921583 and MH921584, respectively). Herpesviral DNA polymerase PCR produced weak bands, precluding sequencing. Upon comparison, both Subantarctic fur seal glycoprotein B gammaherpesvirus sequences presented 85.2% nucleotide and 93.3% amino acid identities among them. The gammaherpesvirus sequence MH921583 presented the highest nucleotide and amino acid identities (94.7% and 98.7%, respectively) to the *Otarid herpesvirus 5* (OtHV-5, MF496135.1) previously identified in a South American fur seal. The gammaherpesvirus sequence MH921584 had the highest nucleotide identity (85.4%) to the OtHV-5 (MF496135.1) detected in a South American fur seal, and the highest amino acid identity (94%) to a harp seal herpesvirus (KP136799.1). The amino acid phylogram indicated that our two novel herpesvirus sequences clustered with other pinniped gammaherpesvirus sequences from the genus *Percavirus* (Figure 2).

**Figure 2.** Phylogenetic analysis based on the alignment 151 deduced amino acid sequences of the herpesviral DNA glycoprotein B sequences found in the Subantarctic fur seal (*Arctocephalus tropicalis*) (circle), 29 alpha-, beta- and gammaherpesvirus sequences obtained from GenBank. An *Ictalurid herpesvirus 1* sequence was included as outgroup (GenBank accession no. NP_041137). The phylogenetic analysis was performed in MEGA7 (KUMAR et al., 2016) using the maximum likelihood method. The reliability of the tree was tested by bootstrap analyses with 1000 bootstrap replicates. Bootstrap values lower than 70% were omitted. Taxon names are presented as herpesvirus species, genus (when assigned), country of origin (for herpesvirus sequences detected in pinnipeds) and accession number.
No positive amplification of *Brucella* spp. was observed. From muscle sample, one novel ITS-1 region sequence (882 bp, excluding primers) was obtained, most similar to *Sarcocystis* spp., submitted to GenBank under accession number MH918015. The obtained ITS-1 sequence had the highest nucleotide identity (99.9%) with *Sarcocystis* spp. (HQ184185) identified in the muscle of a sperm whale (*Physeter macrocephalus*) from Washington, USA, although the sperm whale sequence was shorter than our sequence (800 vs. 882 bp). The second highest identity was to a sequence of *S. canis* (86.4%, DQ176645) obtained from the liver of a captive polar bear (*Ursus maritimus*) from Alaska, USA, that presented the same length as our sequence.

Herein, fatal pneumonia by *Parafilaroides* sp. was identified in a cachectic subantarctic fur seal coinfected by two different gammaherpesviruses and *Sarcocystis* sp. Pulmonary parafilaroidiasis cases by *Parafilaroides* sp. and *P. normani* have been reported, respectively, in South American and Antarctic fur seals (*Arctocephalus gazella*) from Brazil (JACOBUS et al., 2016), and in a subantarctic fur seal from South Africa (DAILEY, 2009). Additionally, thromboembolism associated with verminous pulmonary vasculitis by *Parafilaroides decorus* has been described in a Guadalupe fur seal (*Arctocephalus philippii townsendi*) (SEGUEL et al., 2018). Despite these previous reports, the pathology of *Parafilaroides* sp. is still poorly known. In the present case, while there was mild, chronic bronchointerstitial inflammatory response with mild remodeling, the most severe pathogenetic parasitic consequence appeared to involve airflow obstruction and alveolar emphysema with septal distortion and rupture. These findings are in agreement with previous observations of parafilaroidiasis in other pinnipeds, despite some variations among host species (MEASURES, 2001; JACOBUS et al., 2016). Our results suggest that *Parafilaroides* sp. infection is present along the distribution range of subantarctic fur seals, and that this parasite may be fatal and a significant cause of morbidity in this marine species.

Two gammaherpesvirus sequences were detected, widening the host range of herpesvirus in pinnipeds. No intranuclear inclusion bodies were observed on histopathology, similarly to other reports of herpesvirus infections in pinnipeds (MELERO et al., 2014; SACRISTÁN et al., 2018). According to the International Committee on Taxonomy of Viruses for herpesviruses (DAVISON et al., 2015), the observed nucleotide and amino acid identities suggest the gammaherpesvirus sequence MH921584 could be classified as a novel species, proposed here as *OtarIID herpesvirus* 7, while the other sequence (MH921583) was possibly a different OrHV-5 strain. The pathogenetic role of these gammaherpesviruses was not clear; both were possibly reactivated from latency due to prolonged debilitation of the studied animal, as suggested by the detection of herpesvirus in several organs, but not necessarily associated with clinical disease (GRINDE, 2013).

Based on the detection of highly similar ITS-1 *Sarcocystis* sp. sequences in muscle samples of pinniped and cetacean species from distant geographical areas (southwestern Atlantic and northeastern Pacific), one could consider the involvement of a possible *Sarcocystis* species with a marine cycle. Further research is necessary to confirm such hypothesis.

We report a case of fatal pulmonary parafilaroidiasis in a subantarctic fur seal coinfected with two different gammaherpesviruses, one of them possibly a novel species, tentatively named *OtarIID herpesvirus* 7, and *Sarcocystis* sp. Future studies are warranted to establish the potential impact of these novel herpesvirus sequences, and to identify the *Parafilaroides* and *Sarcocystis* species infecting subantarctic fur seals.

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**References**


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