CASE REPORT

Pseudorabies virus infection (Aujeszky’s disease) in an Iberian lynx (Lynx pardinus) in Spain: a case report

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Abstract

Background: The only natural hosts of Pseudorabies virus (PRV) are members of the family Suidae (Sus scrofa scrofa). In species other than suids infection is normally fatal. In these mammals, including carnivores, PRV typically causes serious neurologic disease. The endangered Iberian lynx (Lynx pardinus) is a wild feline endemic to southwestern Europe (Iberian Peninsula). The Iberian lynx was found to be the world’s most endangered felid species in 2002. In wild felines, PRV infection has only been previously reported once in a Florida panther in 1994. No seropositive lynxes have ever been found, nor has PRV been detected in dead Iberian lynxes to date.

Case presentation: We describe the first reported case of pseudorabies in an Iberian lynx (Lynx pardinus). Pseudorabies was diagnosed in a young wild Iberian lynx from Extremadura (SW Spain) by histopathological examination, immunohistochemistry, polymerase chain reaction (PCR) and sequence analysis. Gross lesions included alopecia of the ventral neck, bloody gastro-intestinal contents and congestion of the brain. Histopathological analysis showed a moderate nonsuppurative meningoencephalitis with diffuse areas of demyelination, necrotizing gastritis and enteritis of the small intestine. Pseudorabies virus (PRV) antigen was found in neuronal and non-neuronal cells of the brain, tonsils, and gastric glandular epithelial cells by immunohistochemical analysis. The presence of the virus in the brain was confirmed by nested PCR. The sequence analysis of the 146 bp fragment (from the viral glycoprotein B gene) showed that the amplified sequence matched (with 100% identity) the PRV genome. Furthermore, specific DNA from glycoprotein D and E encoding-genes was detected by conventional and real-time PCR, respectively, confirming the latter that this infection was produced by a wild-type PRV strain.

Conclusions: This study supports the suspicion that PRV could infect the Iberian lynx. The detection of PRV in a dead Iberian lynx suggests that the virus may have a negative impact on the survival of endangered lynxes in the wild. However, because this is the first verified instance of lynx mortality resulting from pseudorabies, its true impact on the population is unknown.

Keywords: Iberian lynx, Lynx pardinus, Pseudorabies virus, Aujeszky’s disease, Suid Herpesvirus 1, Endangered species, Case report

Background

The Iberian lynx (Lynx pardinus) is an endangered wild feline endemic to the Iberian Peninsula, located in the southwest corner of Europe. The Iberian lynx population suffered a dramatic reduction in size during the 20th century [1]. From 2002 to 2015 it was considered the world’s most endangered wild feline species. Recently, the Iberian lynx has been reclassified from a critically endangered to an endangered species [2]. The population was 406 at the 2015 census [3]. Iberian lynxes predominantly prey on wild rabbits (Oryctolagus cuniculus), although they alternatively may prey on small mammals, juvenile ungulates, partridges, other birds, reptiles and insects [4]. Moreover, Iberian lynxes occasionally consume wild boar as prey and as carrion, which puts them at risk for pseudorabies infection [5]. The species is highly solitary and territorial, although family groups
to bacterial and viral infections. A number of infectious agents have been detected in Iberian lynxes, including a variety of feline viral and bacterial pathogens [8, 9]. The causative agent of Aujeszky’s disease (AD) is Suid Herpesvirus 1 (SuHV1), which is also known as pseudorabies virus (PRV). This virus is a member of the genus Varicellovirus belonging to the subfamily Alphaherpesvirinae of the family Herpesviridae [10] that can infect a wide range of species, though it is does not affect the higher primates [5]. Pseudorabies is a viral disease which was first reported in dogs in 1902 [11], and is prevalent throughout the world. The only natural hosts of PRV are domestic and wild suids (Sus scrofa scrofa) and their hybrids [12]. Symptoms of AD depend on the age of the animal, and can include respiratory, reproductive and nervous signs [13]. PRV is a neurotropic virus which replicates in the nasopharyngeal mucosa, before taking a number of nervous pathways to reach the central nervous system (CNS). It results in a non-suppurative meningoencephalitis which is frequently fatal in piglets [14–17]. PRV seroprevalence in European wild boar varies considerably between different geographic regions. The highest seroprevalences were described in Mediterranean countries including Spain (up to 100%) [5, 12, 18–24]. In both domestic pigs and wild boar, the disease is usually subclinical because of the virus-host adaptation; however, in piglets, the infection is commonly fatal [15–17]. Like other herpesviruses, PRV usually produces latent infections in the host. PRV can infect the neurons and glial cells of apparently healthy swine over long periods. These latent infections can be reactivated, which means PRV can spread to other susceptible animals, mainly mammals, such as carnivores [14, 25], although humans seem insusceptible [26]. Species other than suids can be considered dead-end hosts because the disease is generally fatal before the virus is excreted [5]. In these mammals, including carnivores, PRV infections usually result in very severe neurological symptoms which often involve localised pruritus [26], resulting in death within hours after appearance of the first symptoms [27]. There are considerable implications for conservation, as cases have been reported in endangered carnivores such as the Florida panther [28], the wolf [25] and captive brown bears [29] after consumption of PRV-contaminated meat. No seropositive lynxes have ever been found, nor has PRV been detected in dead Iberian lynxes to date. This article is the first reported case of pseudorabies in an Iberian lynx. We describe the second reported case of pseudorabies in a wild feline considering that PRV infection has only been previously reported once in a Florida panther in 1994 [28]. PRV infection was suspected in this lynx based on histopathological findings, and because PRV infections in wild boar are endemic in SW Spain. Infection was confirmed with immunohistochemistry, PCR, and sequence analysis.

**Case presentation**

The lynx that we studied was a wild ~9-month-old male, born to a healthy 3-year-old dam. It belonged to the first two litters born in Extremadura (SW Spain) after *Lynx pardinus* was re-established in this region through the LIFE+ 10NAT/ES/ 000570 project. Using camera trapping, the estimated date of its birth was established to have been between March 8 and 12, 2015. This lynx was captured, subjected to a routine sanitary evaluation protocol, radio-collared, vaccinated against Feline Leukemia Virus (PureVax FeLV, Merial, Barcelona; Spain) and relocated back into the wild on Nov. 18, 2015. A complete blood count and plasma protein tests showed normal levels. Plasma biochemistry results were within normal ranges, although plasma concentrations of glutamyl transpeptidase and creatine phosphokinase were slightly increased. PCR tests were negative for infection with Feline Leukemia Provirus (FelV), Feline Immunodeficiency Virus (FIV) and Canine Distemper Virus (CDV) in the blood; Feline Calicivirus (FCV) and Feline Herpesvirus (FHV1) in oropharyngeal swabs; and Feline Coronavirus (FCoV) and Feline Parvovirus (FPV) in rectal swabs. The lynx was found to be antigen-ELISA (enzyme-linked immunosorbent assay) negative for FHV1, FCoV and CDV as well as negative for FCV and FPV by fluorescent antibody testing. These tests were performed as previously described [8, 30, 31]. Finally, a blocking ELISA test (CIVTEST SUIS ADV gE, Hipra, Gerona, Spain) was used to detect the presence of serum antibodies against PRV, obtaining negative results.

The lynx was found dead on Dec 1, 2015 on private land consisting of a mixture of dense scrub and open pasture in an area known as “Hornachos-Valle del Matachel” located southwest of Badajoz (Extremadura), Spain (Latitude: 38°27′10.98″ N, Longitude: 5°54′30″ W). Post-mortem examination was carried out at the Veterinary Teaching Hospital of Extremadura (Cáceres, Spain). Upon presentation for necropsy, the lynx weighed 3,060 g, and the carcass was preserved without putrefaction changes. An X-ray examination excluded general traumatisms or the presence of shotgun wounds. Gross lesions of the lynx were minimal. In agreement with our observations, AD in many cases does not develop significant macroscopic lesions in other carnivores such as dogs and cats [32, 33] and wolves [25]. The skin of the ventral neck was denuded of hair and the radio-collar appeared
torn (Fig. 1a). Intense pruritus can sometimes lead to these types of lesions due to scratching and self-mutilation as has been suggested in coyotes [34], dogs [26, 35] and cats [32]. The stomach and small intestine contained a moderate amount of partially digested blood (Fig. 1b). The large intestinal contents consisted of varying amounts of dark red to black semi-formed fecal material. The meninges were congested (Fig. 1c). These lesions are similar to those reported in the Florida panther [28], coyotes [34] and dogs [11].

Representative portions of sampled tissues were fixed in 10% neutral buffered formalin, routinely embedded in paraffin and hematoxylin and eosin (HE) stained. A histopathological analysis of the CNS showed diffuse nonsuppurative meningoencephalitis similar to that reported in domestic cats [32, 36] and other unnatural hosts such as dogs [11, 33, 35, 37], foxes [38] and coyotes [34]. Similar to that described for coyotes [34], the leptomeninges and subarachnoid space were infiltrated and expanded by slight perivascular accumulations of mononuclear cells (Fig. 2a). This meningoencephalitis was characterized by mononuclear cellular infiltrates around blood vessels (perivascular cuffs) and neuropil composed mainly of lymphocytes, as well as multifocal to diffuse microgliosis, perineuronal glial satellitosis (Fig. 2c and d), neuronal necrosis and neuronophagia (Fig. 2d). Most neurons appeared unaffected; although within damaged brain regions, several neurons showed eosinophilic intranuclear inclusion bodies could be absent in the neurons of cats [39]. Diffuse areas of demyelination and malacia were observed in sections of the cerebrum and cerebellum (Fig. 2b). These lesions have been previously described in raccoons [40]. Gastrointestinal tract lesions observed in the lynx consisted of necrotizing gastritis and enteritis of the small intestine with foci of epithelial necrosis with minimal inflammatory reactions. These lesions have been reported in cats and dogs [36, 41] and in piglets [42].

PCR tests were negative for infection with FCoV (clot, spleen, mesenteric ganglia, small intestine and intestinal scraping samples), FCV (clot samples), CDV (brain tissue, clot, mesenteric ganglia and intestinal scraping samples), FPV (clot, mesenteric ganglia and intestinal scraping samples), FHV1 (clot, spleen, mesenteric ganglia, small intestine, intestinal scraping and brain tissue samples), FIV (clot and spleen samples), FeLV provirus (clot, mesenteric ganglia, intestinal scraping and brain tissue samples), *Mycoplasma haemofelis*, *Candidatus mycoplasma haemominutum*, *Candidatus mycoplasma turicensis*, *Anaplasma phagocytophilum*, *Bartonella henselae*, *Chlamydophila felis* [8], *Cytauxzoon felis* (clot samples) [43] and *Leptospira spp.* (kidney samples) [44]. PCR tests were positive for infection with FeLV provirus (spleen and bone marrow samples) [8] and *Pasteurella spp.* (lung samples) using PrimerDesign™ genesig Kit for *Pasteurella multocida* (Genesig, Chandler’s Ford, United Kingdom). Toxicological analyses were negative for pesticides and other organic compounds (chloralose, barbiturates, and metaldehyde), anticoagulant rodenticides and anticholinesterase pesticides. No intestinal parasites were detected in the lynx feces.

The polymer detection method (PDM) to detect porcine PRV was carried out on deparaffinized tissue sections using an UltraVision Quanto Detection System HRP DAB (Thermo Scientific, Fremont, USA, #TL-060-
QHD) following manufacturer’s directions. Primary anti-serum was gE PRV-specific monoclonal antibody (Ingenasa, Madrid, Spain #M.11.ADV.B2CF2). In order to determine the specificity of the immunohistochemical reaction, primary antibody was replaced with PBS or with non-immune mouse serum (1:100). The positive control consisted of a slide containing known positive tissue (CNS of naturally PRV infected pig). Negative control slides consisted of brain sections of a PRV-free lynx. The PRV antigen was detected in the tonsils (Fig. 2f) of the Lynx. PRV has been previously detected by immunohistochemistry in the tonsils of pigs [45] and it was isolated from de tonsils of dogs [33] and racoons [40]. The replication of the virus in tonsils after its entrance via the oral route has been described in cats [32].

Specific staining was not detected in the negative control samples (Fig. 3a). The positive control, showed positive immunoreaction against the gE PRV-specific monoclonal antibody (Fig. 3b).

To assess the presence of PRV in the CNS of the affected animal, a specific nested PCR reaction (Fig. 4) designed to amplify the viral glycoprotein B gene was performed, using two pairs of primers previously described [47]. DNA from brain tissue was obtained using a commercial kit (QiAmp DNA Mini Kit®, Qiagen Ltd., Crawley, West Sussex, UK) following the manufacturer’s protocol. A total of 5 μl of the extracted DNA was used as a template for the nested PCR in a 25 μl reaction mixture containing 12.5 μl of PCR Master Mix (2x) (Green Taq, Thermo Fisher Scientific, Waltham, MA, USA) and 0.2 μM of each primer. DNA was amplified using the following amplification procedure: 1 cycle at 95 °C for 3 min, 30 cycles of denaturalization (94 °C, 45 s), annealing (62 °C, 1 min) and extension (72 °C, 1 min); and a final extension at 72 °C for 10 min. The second step of the nested PCR was performed using 0.5 μl of the first PCR product as template and following the amplification procedure previously described. PCR products were separated by 2% agarose gel electrophoresis and visualized by ethidium bromide staining under UV light.
The obtained 195 bp band was cut out of the gel, and DNA was extracted using an Ultra Clean™ GelSpin DNA Purification Kit (Mo Bio Laboratories, Inc., Carlsbad, California, 92010, USA). The 195 bp amplified product was sequenced in both directions using a BigDye Terminator v3.1 cycle sequencing kit according to the manufacturer’s instructions (Applied Biosystems). Nucleotide sequences were read on a 3730xl DNA Analyzer (Applied Biosystems). After trimming sequence primers, the 146 bp sequence was initially compared with GeneBank sequences using the Basic Local Alignment Search Tool (BLAST) (http://www.ncbi.nlm.nih.gov/blast/Blast.cgi) to verify the identity of the fragment. The BLAST analysis showed that the 146 bp sequence matched with the PRV genome. The obtained sequence and PRV available sequences were aligned by Clustal Omega [48]. The alignment showed that the analyzed fragment presented 100% sequence identity with reference PRV strains such as Kaplan or Bartha. One substitution was found in strains such as NIA3 or Becker, while 2–4 substitutions were found when isolates with an Asian origin were aligned, like HUYD (DDBJ accession number KJ526432) or Rang (DDBJ accession number KP895102) isolates. An additional PCR was carried out to amplify glycoprotein D gene with primers and conditions previously described [49]. Furthermore, in order to assess whether PRV infection was produced by a wild-type or a vaccine PRV strain, a TaqMan® base real-time PCR assay for the detection of gE gene was conducted as previously described using DNA extracted from CNS as template [50]. Specific DNA from glycoprotein D and E encoding-genes was detected confirming the latter that this infection was produced by a wild-type PRV strain.

This article describes the first reported case of pseudo-rabies in Iberian lynxes, and it confirms that they are susceptible to PRV infection. Like other felids, PRV-infected lynxes develop lethal neurological disorders. In cats, PRV infection with brief two- to four-day incubation period, produces acute encephalitis and can cause 100% mortality in experimentally infected domestic cats [32, 51]. The cats suffer from anorexia, occasional severe itching with lesions caused by scratching and self mutilation and uncoordinated movements and paralysis [32, 51]. The outcome is invariably fatal and leads to death within 12–48 h of the first appearance of clinical signs [32]. This peracute death does not allow time for a serologic response to occur [52]. If Iberian lynxes succumb as rapidly as domestic cats, detection of the clinical phase can prove to be very difficult.
The main sources of infection for cats are uncooked pig or offal [32]. Direct spread of virus from infected to non-infected carnivores likely does not occur [34]. Wild boar are a well known reservoir for PRV and hence, may pose a risk to transmit the infection to wildlife carnivores species [25, 34]. Some authors have described recently, rates of 69.70% ELISA seropositivity and 11.30% of PRV lung infections in wild boar population throughout SW Spain [53, 54]. Wild boar are not the main component of the Iberian lynx diet; however, these animals do occasionally consume wild boar as prey and as carrion [5]. Therefore, it is conceivable that some lynxes will be exposed to PRV from the ingestion of wild boar. Indirect transmission can also occur through viral excretion by pigs, without direct contact with the pigs themselves [32]. The presence of wild boar infected with the PRV could have a negative impact on conservation of wild carnivores which consume wild boar [25] such as lynxes.

Conclusions
Our findings support the suspicion that PRV could infect the Iberian lynx [5, 22]. The detection of PRV in a dead Iberian lynx suggests that the virus may have a negative impact on the endangered lynx’s survival in the wild. However, because this is the first verified instance of lynx mortality resulting from pseudorabies, its true impact on the population is unknown. A possibility for management of this species is vaccinating lynxes with an inactivated vaccine during routine captures to provide some level of immunity against this disease. This has previously been suggested for Florida panthers [28], although vaccination has not been proven efficacious. Oral vaccination of wild boar against classical swine fever virus has been shown to be successful [55], which means that oral vaccination of wild boar against PRV could be an effective strategy for protecting lynxes from the disease [25]. The effectiveness of using attenuated live vaccine for oral immunisation of wild boar against PRV has been demonstrated [56], though the safety of this technique needs to be studied in depth [25].

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Availability of data and materials
Supporting data (Additional file 1: Table S1 and Additional file 2: File S2) are included as additional files to this article.

Authors' contributions
AJM and ER carried out the necropsy, histopathology and immunohistochemistry. AJM, ER and DR were involved in the revision of literature, drafting and writing of the manuscript. MG, DR and JIN carried out the PRV detection by PCR and the sequence analysis. OM1 carried out field work, provided the case history and contributed to the necropsy. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

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Additional files

Additional file 1: Table S1. Primers used for SuHV1 amplification. (DOCX 90 kb)

Additional file 2: File S2. Partial glycoprotein B nucleotide sequence (SuHV1) isolated from Lynx pardinus. (DOCX 42 kb)

Abbreviations
AD: Aujeszky’s disease; CDV: Canine Distemper Virus; CNS: Central nervous system; DAB: Diaminobenzidine; ELISA: Enzyme-linked immunosorbent assay; FCoV: Feline Coronavirus; FCV: Feline Calicivirus; FeLV: Feline Leukemia Virus; FHV1: Feline Herpesvirus; FIV: Feline Immunodeficiency Virus; FPV: Feline Parovirus; HE: Hematoxylin and eosin; PCR: Polymerase chain reaction; PDM: Polymer detection method; PRV: Pseudorabies virus; SuHV1: Suid Herpesvirus 1.
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