RESEARCH ARTICLE

Putative parapoxvirus-associated foot disease in the endangered huemul deer (*Hippocamelus bisulcus*) in Bernardo O'Higgins National Park, Chile

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Abstract

The huemul (*Hippocamelus bisulcus*) is an endangered cervid endemic to southern Argentina and Chile. Here we report foot lesions in 24 huemul from Bernardo O’Higgins National Park, Chile, between 2005 and 2010. Affected deer displayed variably severe clinical signs, including lameness and soft tissue swelling of the limbs proximal to the hoof or in the interdigital space, ulceration of the swollen tissues, and some developed severe proliferative tissue changes that caused various types of abnormal wear, entrapment, and/or displacement of the hooves and/or dewclaws. Animals showed signs of intense pain and reduced mobility followed by loss of body condition and recumbency, which often preceded death. The disease affected both genders and all age categories. Morbidity and mortality reached 80% and 40%, respectively. Diagnostics were restricted to a limited number of cases from which samples were available. Histology revealed severe papillomatous epidermal hyperplasia and superficial dermatitis. Electron microscopy identified viral particles consistent with viruses in the Chordopoxvirinae subfamily. The presence of parapoxvirus DNA was confirmed by a pan-parapoxvirus PCR assay, showing high identity (98%) with bovine papular stomatitis virus and pseudocowpoxvirus. This is the first report of foot disease in huemul deer in Chile, putatively attributed to poxvirus. Given the high morbidity and mortality observed, this virus might pose a considerable conservation threat to huemul deer in Chilean Patagonia. Moreover, this report highlights a need for improved monitoring of huemul populations...
and synergistic, rapid response efforts to adequately address disease events that threaten the species.

**Introduction**

There is an increasing concern about the potential contribution of diseases in wildlife extinctions, particularly when they interact with other driving factors [1–5]. For example, the effects of infectious pathogens can have devastating effects when population size is small, when multi-host pathogens and reservoir hosts are available, when the infectious agent can survive in an abiotic environment or when disease transmission is influenced by environmental factors or climate change [6–8]. Furthermore, the outcome of an infectious disease depends on intrinsic characteristics of the pathogen that shape morbidity and mortality, ultimately defining severity of illness and the future of affected populations [5, 9, 10].

The huemul deer (Hippocamelus bisulcus) is a medium-sized neotropical cervid that is endemic to shrubby habitats and forests in southern Argentina and Chile [11]. Huemul were the most widespread species in Patagonian forests until the 19th century [12], but since that time their range and populations have markedly declined. Contributing factors include habitat loss, poaching, competition with introduced ungulates and susceptibility to livestock diseases [13]. Its current range is now mainly restricted to Nothophagus forests in the Andes and periglacial areas surrounding the continental icecaps in Patagonia between 36 and 52° S [14, 15]. At present, huemul are listed as endangered, and fewer than 2,500 individuals remain in fragmented populations in the wild [13, 16]. Studies on huemul health are scarce [17–23]. While most published information is dated and largely anecdotal [24], recent reports suggest disease might be of increasing conservation concern for this species [22, 23].

Bernardo O’Higgins National Park (BONP) in Chile is one of a few remaining strongholds for huemul deer in South America. Some areas of this park are home to the highest densities (4.52 deer/km$^2$) of huemul deer across its current range [25]. The remote nature and protection within the park, along with hostile weather and rugged mountainous and coastal topography, are likely significant factors that may help to buffer and protect deer in this area from threats that have led to declines in other regions [26]. Here we describe foot lesions putatively attributed to poxvirus infection and associated morbidity in huemul from BONP.

**Material and methods**

**Study area**

The study was conducted in public lands at BONP in the Magallanes Region of Chilean Patagonia. This National Park is managed by the Chilean National Forest Service (CONAF). Our study areas were located along the edge of the southern continental icecap in the Huemules (3.2 km$^2$), Katraska (5 km$^2$) and Bernardo (13.5 km$^2$) Valleys (Fig 1). Across this area, the climate is cold and wet. Mean annual precipitation is 4,000 mm and is evenly distributed throughout the year with snowfall from June through August. Annual temperatures average 7–8˚C [27]. The vegetation includes periglacial grasslands, grassland–forest ecotones, old-growth forest dominated by Nothofagus species and moorlands [26].

Human presence in the area is restricted to Puerto Edén (49°07’34”S, 74°24’48”W), an isolated coastal village with 176 inhabitants located in Wellington Island [28]. In addition, the National Park Service, CONAF maintains a field station in Témapos Fjord (48°41’33”S; 73°59’21”W). Two park guards have been based at the station throughout the year since 2002.
In 1991, 18 cattle were illegally introduced to Huemules Valley (HV), which affected both huemul abundance and habitat use patterns [26, 29]. By 2001, the cattle population had grown to 31.3 individuals/km$^2$ [30], triggering governmental control efforts. While cattle were eliminated from HV by 2004, some animals escaped to neighboring inaccessible areas and continue to be culled opportunistically. Bernardo (BV) and Katraska (KV) Valleys, on the other hand, have always been cattle-free. There are, however, no geographical barriers that prevent animal movements between HV and BV. Following cattle removal, huemul numbers increased in HV [25].

**Field data**

Park rangers in Témanos Fjord have monitored huemul deer and feral cattle presence in HV at least once weekly from 2004. Visits to KV and BV have been less frequent and limited to annual deer abundance surveys conducted once or twice a year. Individual animal identification is made by observation of natural marks, scars, and, for males, antler shape. Observation of huemul foot abnormalities, data and sample collection, and photographs of affected animals reported here were performed in the field by park rangers between 2005 and 2010.

**Laboratory analysis.** External examination, morphometric data recording, photo documentation, gross necropsy examination and tissue sample collection were performed opportunistically on dead huemul by CONAF personnel (Cases #1, #10 and #18). Tissue samples from
foot lesions and select internal organs were fixed in 10% neutral buffered formalin prior to routine histologic processing, sectioning at 5 μm, hematoxylin and eosin (HE) staining, and histologic examination. Samples from all cases were examined histologically at Facultad de Ciencias Veterinarias, Universidad de Concepción, Chillán, Chile. Additionally, in 2011, four paraffin blocks from Case #10 (male fawn) containing tissue from affected limbs, were imported to the USA (CITES # 0002245 Chile, #11US033594/9 USA) for additional diagnostics at the Wildlife Conservation Society, Bronx, New York. Following histologic examination, paraffin embedded (FFPE) lesional skin from Case #10 was further analyzed using routine Gram and silver-staining (Warthin-Starry), immunohistochemistry, polymerase chain reaction, microarray testing, and electron microscopy as described below. Laboratory sample disposal protocols at Universidad de Concepción indicate destruction of materials following examination. Therefore, only samples from Case #10 were available for exportation and ancillary diagnostics in the United States.

**Immunohistochemistry.** An immunohistochemical (IHC) assay (DAKO automatic universal staining system) using a rabbit polyclonal antibody against bovine papillomavirus, that is also broadly reactive to canine, feline and equine papillomavirus, and positive and negative controls, was performed on 5 μm sections of formalin-fixed, paraffin-embedded (FFPE) lesional skin (Case #10) (University of Pennsylvania School of Veterinary Medicine, Philadelphia, PA, USA). Briefly, 5 μm, serial sections on negatively charged glass slides were obtained from formalin FFPE tissues. Slides were then heated in a 60˚C oven for 1 hour, deparaffinized, then rehydrated with PAR clearant and progressive decreasing grades (from 100% to 95% and to water) of ethanol. Antigen retrieval in citrate buffer (pH ~9.0) and endogenous peroxidases inactivation in hydrogen peroxide (10 min) were performed. Immunolabelling was conducted with a mouse monoclonal, HPV cocktail broad spectrum primary antibody (BioCare, Pacheco, CA, USA), biotinylated goat anti-rabbit and goat anti-mouse secondary antibody, and visualization included incubation with streptavidin conjugated to horseradish peroxidase (15 min) followed incubation with AEC chromogen. Labelled slides were counterstained with hematoxylin (1 min).

**Transmission electron microscopy.** Electron microscopy (EM) was performed on a single, unstained, 5 μm section of FFPE lesional skin (Case #10) mounted on a charged glass slide (Veterinary Diagnostic Laboratory, University of Minnesota). The sample was deparaffinized, rehydrated, and post-fixed in 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer followed by a second post-fixation in 1% osmium tetroxide in 0.1 M sodium cacodylate buffer (Electron Microscopy Sciences, Hatfield, PA, USA). The sample was dehydrated using a 25%–100% ethyl alcohol gradient. It was then infiltrated and embedded “in situ” with Embed 812 resin (Electron Microscopy Sciences, Hatfield, PA, USA). Embedded tissue was sectioned on a Leica UC6 ultramicrotome (Leica Microsystems, Vienna, Austria). Thin sections (60–70 nm) were obtained and collected onto a 200-mesh nickel grid (Electron Microscopy Sciences, Hatfield, PA, USA). Grids were contrasted with 5% uranyl acetate and Santos’ lead citrate. These preparations were visualized using a JEOL 1400 transmission electron microscope (JEOL LTD, Tokyo, Japan). Images were obtained using an AMT Capture Engine Version 7.00 camera and software (Advanced Microscopy Techniques Corp. Woburn, MA, USA). Image analysis was carried out using ImageJ (NIHR public domain).

**Polymerase chain reaction (PCR).** DNA was extracted from 50 μm of FFPE sections of lesional skin (Case #10) using the QiAMP DNA FFPE tissue kit, and a protocol adapted for using deparaffinization solution (Qiagen Inc., Valencia, CA, USA). Individual pan-viral polymerase chain reactions (PCR) were performed targeting consensus regions of less than 330 bp for adenoviruses (polymerase gene), herpesviruses (polymerase gene), polyomavirus (VP-1 gene), and flavivirus (NS-5 gene) using previously described methods (Table 1), and
appropriate positive and negative controls [31, 32]. Additionally, pan-poxvirus PCR testing (192 bp) was performed on DNA extracted from 5 μm sections (30 μm total) of unstained recut FFPE tissue mounted on charged slides. The primers for this assay were designed to amplify consensus regions using an alignment of the following poxviruses: cowpox, sheeppox, goatpox, deerpox, red deer parapox, bovine popular stomatitis virus, raccoonpox, cetacean poxvirus, dolphin poxvirus, harbor seal parapox, pinniped parapox, Steller sea lion poxvirus, Steller sea lion parapox, sea otter pox, myxomavirus, avipox, canary pox, penguin pox, monkeypox and vaccinia virus as previously described [32]. Amplified products were directly sequenced in the forward and reverse directions (Eton bioscience, Union NJ, USA). All sequences were analyzed, trimmed of their primer sequences, and aligned to generate a consensus sequence that was queried against available sequences in GenBank (National Center for Biotechnology Information, Bethesda, MD, USA).

Microarray analysis. FFPE tissue sections were removed separately from five glass slides (Case #10) by wetting a scalpel with ethanol and scraping tissue from the surface. Ethanol was removed by centrifugation and samples dried. One ml xylene was added to each sample and vortexed to remove paraffin. Xylene was removed by centrifugation, and remaining tissue was washed in 100% ethanol and dried. Tissue was lysed by incubation with proteinase K at 56˚C for one hour, followed by incubation at 90˚C to reverse cross-linking. DNA was then extracted from the lysed suspension using the QIAamp cador Pathogen Mini Kit (Qiagen Inc., Valencia, CA, USA) according to the manufacturer’s recommendations. Due to low quantities of total extracted DNA, whole genome amplification was performed using the Repli-g midi kit (Qiagen Inc., Valencia, CA, USA). Samples were amplified at 30˚C for 16 hours and purified using QIAquick PCR purification columns (Qiagen Inc., Valencia, CA, USA). Extracted DNA samples were labeled with Cy3 using the NimbleGen One-Color DNA labeling kit (Roche, Inc. Madison, WI, USA). Samples were prepared for hybridization to the Lawrence Livermore Microbial Detection Array (LLMDA v5, 3 arrays x 720K probes) using the NimbleGen hybridization kit LS (Roche, Inc. Madison, WI, USA). This array includes probes designed to detect

Table 1. Primer sequences and reference methods for PCR assays conducted on samples of deer Case #10.

| PCR Assay Target         | Amplicon Size (bp) | Primer Name | Primer Sequence (5’ to 3’) | Reference and Assay Sensitivity* |
|--------------------------|--------------------|-------------|---------------------------|----------------------------------|
| Adenovirus DNA Polymerase Gene (Nested) | 330 | •pol-Fouter | •TNMGNGGGNQOMNGTGYTAYCC | [33] < 5 copies/reaction |
|                          |                    | •pol-Router | •GTDSGCAAANSNHCRTABARGMRTT |                                   |
|                          |                    | •pol-Finner | •GTNTWYATHTYGYGSHATGYCG |                                   |
|                          |                    | •pol-Rinner | •CCANCBCDRTPRGNAKNTTRA |                                   |
| Herpesvirus DNA Polymerase Gene (Nested) | 250 | •DFA | •GAHTTYGCNAGYNTTAYCRR | [34] < 5 copies/reaction |
|                          |                    | •ILK        | •TCCTGGACAACGCAARNYSNGNTAA |                                   |
|                          |                    | •KG1       | •GTCTTGCCTCACAGNTCNACNCYT |                                   |
|                          |                    | •TGV | •TGAACTCCTGTTAGGNTNTYAAACNGNT |                                   |
|                          |                    | •1YG | •CACAGATCGCTTRNCNCRTDAT |                                   |
| Polyomavirus VP1 gene | 277 | •VP1/2F-JO2F | •ATGAAATGGGGGTGCCCCCNNTGYAARG | [35] sensitivity untested |
|                          |                    | •VP1/2R-JO2R | •CCCTCAATAACCGAACYTCHACYT |                                   |
| Poxvirus DNA Polymerase Gene | 192 | •TSPoxPolF1 | •TATAGAGCGAGTACAGTCTCAAG | [32] 5 copies/reaction |
|                          |                    | •TSPoxPolF2 | •CCAGGYACCTGCTACAAG |                                   |
|                          |                    | •TSPoxPolF3 | •TAYAGACGTAGTACTTTAATAAA |                                   |
|                          |                    | •TSPoxPolF4 | •TATAGGCGAGTACKCTATTAAA |                                   |
|                          |                    | •TSPoxPolR1 | •CAACATT101GATYARACTATTATAATC |                                   |
|                          |                    | •TSPoxPolR2 | •GTT11GATYARCTGTTGTCATG |                                   |
| Flavivirus NS-5 gene | 270 | •Flavi-FWD | •TGYBTTTAYAACATAGTGGG | [36] 900 copies/reaction |
|                          |                    | •Flavi-RVS | •GTGTCCTA12CCNGNTTCTG |                                   |

*Sensitivity determined using control plasmids

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microbial species with publicly available sequence data obtainable as of December 2011. This layout of the LLMDA platform includes 1,967 bacterial species, 126 archaeal species, 136 fungal species, 94 protozoan species, and 3,111 viruses, and has been previously applied for pathogen surveillance in wildlife [37]. Approximately 15 μg labeled DNA from each sample was added to a separate array, followed by hybridization for 47 hours at 42˚C. Arrays were washed using the NimbleGen wash buffer kit and scanned with the MS200 microarray scanner (Roche, Inc. Madison, WI, USA). Data were processed, and potential microbial targets identified using Composite Likelihood Maximization Method (CLiMax) as previously described [38]. Positive probe intensity thresholds were set at the 99th percentile relative to negative control probe intensities and lowered and rerun at the 95th percentile (to broaden possible target pathogen detection). These have been applied as standard thresholds in previous studies using the LLMDA platform for a variety of sample types derived from wild and domestic animals [37, 39] as well as degraded archaeological samples and human lymphoma FFPE tissues [40, 41].

**Ethics statement.** This study was conducted within a cooperation agreement between the Chilean Forestry Agency, Corporación Nacional Forestal–CONAF, and the Wildlife Conservation Society. CONAF is the administrator of terrestrial protected areas in Chile, including Bernardo O’Higgins National Park where the huemul cases occurred. No specific permissions were required for this location or activities which were performed by government personnel following government and forestry agency regulations. As part of their duties, park rangers perform regular observations of huemul deer within the protected area and are authorized to collect samples from dead animals if observed. All animals in this study were dead at the time of necropsy by government personnel. No animals were euthanized in this study.

**Results**

Overall, 24 huemul deer with foot lesions were identified between April 2005 and August 2010. Seventy five percent (n = 18) of affected huemul were located in HV; the remainder were found in the more isolated KV (n = 1) and BV (n = 5) (Fig 1). All affected deer displayed similar but varying degrees of clinical signs. These included lameness and soft tissue swelling in one or more limbs just proximal to the hoof or in the interdigital space and, in some cases ulceration of the swollen areas. Some cases spontaneously and completely resolved while others progressed to more severe, proliferative and/or supplicative forms that caused various types of abnormal wear, entrapment, and/or displacement of the hooves and/or dewclaws. These animals showed signs of intense pain such as marked lameness, reluctance to bear any weight on the affected feet, and reduced mobility followed by loss of body condition and prolonged recumbency, which often preceded death.

Deer in HV, but not BV or KV (due to their more rugged landscapes and remoteness), were monitored periodically following the onset of disease. Details on disease evolution and outcome presented in this section are hereafter restricted to the deer in the accessible HV area. Of the 18 affected individuals in HV, four (22.2%) were adult females, ten (55.6%) were adult males, one (5.56%) was a juvenile male, and three (16.7%) were fawns: a female, a male and one of undetermined gender (Table 2). More males (n = 12, 66.7%) than females (n = 5, 27.7%) were found with lesions. Between 25 and 100% of males and 50 to 100% of females in HV were affected, respectively, depending on the year. Furthermore, all juveniles and fawns in HV during the study period were affected. Considering the minimum number of observed deer in HV, morbidity and mortality rates ranged from 40 to 80% and from 0 to 40%, respectively, during the five years of the episode (Table 3).
Regarding the distribution of lesions, one limb was affected in nine individuals (50%), two limbs in two (11%), and two fawns had either three or four limbs compromised (5.5% in each category). The number of affected limbs was undetermined in the remaining five animals (28%) (Table 2). The outcome of lesions varied. Seven individuals (38.9%) progressed to full recovery, six (33.3%) were found dead within a month of lesion detection (three females and three males), and the outcome was unknown in the remaining five deer (27.8%) because they could not be tracked. Most of the affected individuals in HV were observed during the fall (8 of 16 for which month was recorded), two and five affected deer were observed in the winter and spring, respectively, and only one case was observed in the summer.

Cases were not seen in the remote BV or KV until 2008. Between 2008 and 2010, five adult males (infection rate = 1/18 in 2008 and 4/9 in 2010, all in BV) and a single adult female (1/12 in 2010, only case in KV), were observed with lesions in these valleys. Five of the six deer with foot lesions in KV and BV were recorded in the fall; the remaining one (2008 male) was reported during the spring.

Table 2. Huemul deer with foot lesions in Huemules Valley.

| Case # | Date   | Sex and Age | Affected limbs | Fate of animal |
|--------|--------|-------------|----------------|---------------|
|        |        |             | Fore | Hind |               |
| 1      | Apr-05 | F Ad        | R    |      | Dead          |
| 2      | May-05 | M Ad        |      |      | Recovered     |
| 3      | Sep-05 | F Ad        |      |      | Dead          |
| 4      | Jun-06 | M Ad        |      |      | Recovered     |
| 5      | Jun-06 | M Juv       |      |      | Recovered     |
| 6      | Nov-06 | M Ad        |      | R    | Unknown       |
| 7      | Oct-07 | M Ad        |      | L    | Recovered     |
| 8      | Oct-07 | M Ad        |      | R    | Recovered     |
| 9      | Apr-08 | M Ad        | L    |      | Unknown       |
| 10     | Jun-08 | M Fawn      | R/L  | R/L  | Dead          |
| 11     | Aug-08 | U Fawn      | R    | R/L  | Recovered     |
| 12     | Sep-08 | F Ad        | R/L  |      | Recovered     |
| 13     | Feb-09 | F Ad        |      | L    | Unknown       |
| 14     | May-09 | M Ad        |      |      | Dead          |
| 15     | Apr-10 | M Ad        | L    |      | Unknown       |
| 16     | Apr-10 | M Ad        |      | R    | Unknown       |
| 17     | May-10 | M Ad        | L    |      | Dead          |
| 18     | Jul-10 | F Fawn      | R    | L    | Dead          |

F = female, M = male, U = unknown, Ad = adult, R = right, and L = left. Cases highlighted in bold have been described extensively in the results section.

Table 3. Morbidity and mortality rates attributed to foot disease in huemul deer observed in Huemules Valley (HV).

| Year | # of cases | # of deer in the valley | Morbidity (%) | Mortality (%) |
|------|------------|-------------------------|---------------|---------------|
| 2005 | 3          | 7                       | 42.9          | 28.6          |
| 2006 | 3          | 7                       | 42.9          | 0.0           |
| 2007 | 2          | 5                       | 40.0          | 0.0           |
| 2008 | 4          | 8                       | 50.0          | 12.5          |
| 2009 | 2          | 5                       | 40.0          | 20.0          |
| 2010 | 4          | 8                       | 80.0          | 40.0          |

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Three huemul carcasses were recovered for examination and sample collection (Cases #1, #10 and #18, respectively; Table 2) by park rangers, and included an adult female (four years old, April 2005) and two fawns (a seven month-old male in June 2008, and an eight month-old female in July 2010) from HV. In all cases where it was possible to recover affected animals or limbs, park rangers described affected tissue as having a strong, fetid smell, suggestive of possible tissue necrosis and/or bacterial or other microbial infection. Additional macroscopic details of the foot lesions presented here are based on observations from available photographs.

Case #1

The first recognized huemul with foot lesions in BONP was an adult female sighted in April 2005 (Case #1, Table 2). The deer was initially observed limping, with marked, diffuse soft tissue swelling of the right forelimb distal to the humero-radial carpal joint, including the carpo-metacarpal joint and foot. Over the next three days the deer was typically recumbent and showed increasing signs of pain (Fig 2). The deer was found drowned in a lagoon four days after initial observations; numerous culpeo fox (Pseudalopex culpaeus) prints were seen in the surrounding mud. In addition to generalized swelling of the soft tissues of the foreleg, post mortem findings included a broad based, medially located, regionally extensive area of tissue swelling just proximal to the coronary band that caused partial lateral separation of the claws, and in which the skin was partially alopecic (Fig 3). Additionally, fractures to the tips of the claws on the affected foot, and poor body condition with very small amounts of cavitary and mesenteric adipose tissue were observed.

Case #10

This was the first case of a young huemul with foot lesions recorded in June 2008 (Case #10, Table 2). This was a male fawn in which all limbs were affected. Lesions consisted of verrucous, proliferative, exophytic tissue that surrounded the base of the hoof, laterally separated the claws, and extended proximally from the level of the coronary band (Fig 4). As in the adult female (Figs 2 and 3), there was a regionally extensive area of alopecia, and there was marked swelling of the soft tissues of the leg and foot. The foot of the left forelimb was more severely affected. Circumferential, proliferative tissue was present around the claws and distal left foreleg and foot (from the claws and coronary band roughly to the mid metacarpal region). Bleeding and drainage of fluid was markedly visible. Lesions on the left rear leg was more restricted to the anterior aspect of the foot; claws on both feet appeared normally worn. The front right limb appeared swollen and draining fluid, with limited proliferative tissue growth. Initially, the fawn displayed signs of severe pain, reduced mobility, and bore most of its weight on its right legs. During the following days, the fawn spent increasing amounts of time recumbent with limited foraging periods. On the fifth successive day of observation, the rear right limb became noticeably swollen. By day ten, the general condition of the deer had declined substantially, and it spent all its time recumbent and was not seen to feed. It was found dead the following day with lesions consistent with fox predation.

Histologic examination of lesions in affected skin from Case #10 (gross lesions seen in Fig 4) consisted of moderate to severe papillomatous to papilliferous epidermal hyperplasia with moderate to severe acanthosis and mild to moderate rete peg formation with maintenance of normal epidermal stratification of the epidermis (Fig 5A). Additional epidermal changes included mild multifocal intercellular edema, mild to moderate ballooning intracellular edema (primarily along the tips of the proliferative epidermis), and moderate compact orthokeratosis and parakeratosis (Fig 5B). Multifocal aggregates of degenerate cells and clusters of mixed
bacteria (primarily short coccobacilli) were present along the surface of the thickened, compact stratum corneum. Mixed Gram positive and negative bacteria were highlighted with Gram staining; long, spirochete-like bacteria in the superficial stratum corneum were highlighted with Warthin-Starry staining (Fig 5C). In one area, fungal hyphae (Zygomycete type) were also focally present in the superficial compact stratum corneum. The cellularity of the subepidermal papillary layer of the superficial dermis was mildly increased and contained minimal, multifocal, lymphoplasmacytic inflammation, and the collagen had a slightly 'smudgy' appearance multifocally. Hypergranulosis and koilocytes were not seen. Immunohistochemical (IHC) staining for bovine papillomavirus was negative in multiple tissue sections.

Electron microscopic examination of skin tissue revealed numerous viral particles in the superficial crust and epidermis. The cytoplasm of these cells was distended and contained a large number of mature viral particles in a clear background (Fig 6A). Within the cytoplasm of epidermal cells were immature, maturing and mature virions admixed in an electron-dense accumulation of amorphous material that correspond to cytoplasmic viral factories (Fig 6A and 6B). Mature viral particles (also known as intracellular mature virus) were characterized
by the presence of an electron dense core with a distinct concave shape and lateral body covered by a layered membrane (Fig 6C). Mature virions were approximately 189.3 (SD = 14.1) x 141.6 (SD = 52.3) x 85.3 (SD = 6.8) nm. The core of mature viruses displayed a cylindrical folded morphology. The diameter of the core cylinder was approximately 56.4 (SD = 6.4) nm (Fig 6D). The mature virus membrane and core wall were approximately 29.2 (SD = 4.9) nm thick. The lateral bodies were inconspicuous. Mature virions and a mixed population of bacteria and yeast were present in the superficial crust. Virions presented morphology and dimension consistent with virus of the Chordopoxvirinae subfamily.

PCR was negative for herpesviruses, adenoviruses, polyomaviruses, and flaviviruses, and positive for a parapoxvirus (135 bp). BLASTN analysis showed 98% similarity (3 nucleotide difference) to bovine papular stomatitis virus strain BV-TX09c1 (GenBank accession number KM875472.1) and pseudocowpox virus strain FSS742 (GenBank accession number MH169576.1); the next closest match was 96% homology to another pseudocowpox virus strain VR634 (GenBank No. GQ329670.1).

Microarray analysis did not identify any microbial targets at either the 99th or 95th percentile intensity thresholds, relative to negative control probes, at the default setting of at least 20% of probes detected versus the probes expected to a given target. Hybridization data was further examined for viral identification events at low stringency thresholds (95th percentile, minimum of one detected probe); however, viral detection remained negative under these relaxed parameters.
Case #18

The third carcass recovered was a female fawn observed in July 2010 with lesions on the left rear limb (Case #18, Table 2). Nine days later, the right forelimb became affected. Eighteen days after the first observation of lesions, purulent material was seen in affected sites, and a noticeable decline in body condition was also apparent. On day 24, the deer became permanently recumbent and displayed evidence of respiratory distress. It died on the 26th day of observation. Histological changes in affected tissue from the foot lesions were similar, but perhaps somewhat more proliferative and papillomatous to those in the male fawn (Case #10).

Discussion

This is the first report of foot disease, putatively attributed to poxvirus, in huemul deer in Chile. The severity of clinical disease was variable, yet in a third of affected animals (at least six individuals), it resulted in complete incapacitation and death. That a minimum of 18 deer were affected in HV (over five years), with morbidity and mortality rates as high as 80% and
40%, respectively, denotes that foot lesions such as those reported here pose a considerable conservation threat for this species.

Fig 5. Characteristic histologic changes in affected haired skin from huemul limb (Case #10, Table 1). A) The epidermis is markedly hyperplastic and thrown up into multiple folds. Short rete pegs extend into the subjacent dermis (HE). B) Additional epidermal changes include moderate compact orthokeratotic and parakeratotic hyperkeratosis and mild to severe intracellular edema. Multifocal superficial areas containing smudgy, basophilic material contain degenerate cellular debris admixed with mixed bacteria (HE). C) Myriad long bacteria are stained dark brown and are focally present in this section of an acellular crust overlying the stratum corneum. Short bacilli and cocci that are more lightly stained are present along the base of the crust (Warthin Starry).

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Fig 6. A-D. Ultrastructure of an affected haired skin from huemul limb (Case #10, Table 1). A. The cytoplasm of the stratum granulosum cells is clear and distended by myriad poxvirus particles. Within the cytoplasm, some of these cells presented a large electron-dense aggregation consistent with a viral factory (arrows) and fragment of chromatin (arrowhead). B. Viral factories were characterized by large aggregation of electron-dense amorphous material admixed with immature (arrowhead), maturing and mature virions (arrow). Bar = 600 nm. C. A magnification showed detail of immature virions (arrowhead), which displayed a round shape lined by a radiated membrane. A black bar corresponds to the thickness of the multilayered membrane and a white bar indicates the thickness of the core of a mature virion (arrow). Bar = 200 nm. D. Different axis of mature virions. Note that the core is a folded cylinder with horizontal (D1) cross (D2 and D3), and sagittal (D4) sections. Bar = 200 nm.

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Due to limited availability of samples, a thorough investigation was only possible in one individual case (Case #10). Notwithstanding, similarities in the clinical presentation and gross lesions, as well as the progression and outcome of disease, suggest a common etiology and/or pathogenesis. Moreover, the disease behaved as a recurrent outbreak with cases observed predominantly in the fall of each year over a five-year period. Recovery in nearly half of the monitored animals suggests that clinical disease was self-limiting in some individuals. While most affected animals were adults, the increased severity of lesions and higher relative mortality in fawns and juveniles indicates that these age groups were the most likely to succumb to the disease once infected and/or were more likely to die from opportunistic predation.

Numerous foot, interdigital, and hoof diseases have been reported in domestic and wild bovidae, cervidae, and pronghorn antelope [42–52]. Those that more closely resemble the gross appearance of huemul deer cases are footrot or infectious pododermatitis caused by bacteria such as Dichelobacter nodosus and Fusobacterium necrophorum [51], and the more recently described polymicrobial, multi-treponeme infections associated with digital dermatitis (DD) (‘hairy heel wart’) in sheep and cattle, and severe Treponeme-associated hoof disease (TAHD) in elk (Cervus canadensis) [46–48, 49, 52]. Common clinical findings in many of the livestock foot diseases and TAHD of elk that were also seen in huemul are obvious pain and lameness. Grossly, the first affected female huemul (Fig 2, Case #1) presented with interdigital swelling, similar to some of the classic lesions in animals affected by DD and TAHD. However, the evolution and appearance of the lesions in this and other huemul was not typical of footrot necrosis or TAHD. In these diseases, necrosis of the interdigital skin is associated with separation of the horn and undermining of the toe or sole in both and hoof growth abnormalities in the latter. These changes were not seen in the affected huemul. In huemul, rather than necrotizing or ulcerative hoof abnormalities, lesions were primarily proliferative, and in no cases was abnormal hoof growth seen. However, chronic stages of DD can be characterized by thick granulation tissue, something that more closely resembled what was seen in some of the BNOP deer, such as Case #10 fawn (Fig 4). The huemul foot lesions we report also appear to differ from digital dermatitis in that DD most commonly presents as a circumscribed moist ulcerative erosive mass along the coronary band or interdigital space on the plantar aspect of the foot (often a rear foot, though some variation can be seen) [45]. Differences between the huemul cases and TAHD in elk were significant. In elk, a predominant feature is hoof deformation and overgrowth; underlying laminitis is also described. Hoof overgrowth was not seen in affected huemul. However, in some cases, loss of hoof structure was associated with exuberant proliferative granulation tissue. Laminitis was not grossly apparent and was not investigated as a contributing lesion in huemul due to sample availability, but more extensive examination of hooves in future cases would be of value. An additional notable difference between TAHD and huemul lesions is that the latter progressively affected several limbs, including fore and hindlimbs, something not commonly seen in TAHD of elk [49, 50, 52].

DD and TAHD are associated with the presence of multiple treponeme-like bacteria in affected tissues. However, the role of treponemes in hoof syndromes is unclear. Multiple trials have failed to fulfil Koch’s postulates for these bacteria as single causal agents, and current publications refer to a probable polymicrobial etiology, including treponemes, with variations in affected species, geographical location and environmental factors [45, 53]. Notwithstanding, there is agreement that treponemes are present in several chronic ulcerative dermatoses, suggesting common virulence factors that contribute to the development of similar clinical symptoms and lesions [45, 46, 50, 54]. In our study, spirochete-like bacteria were found superficially in the single available tissue sample from Case #10 in Fig 4. Despite several trials, we were unable to further characterize the bacteria through molecular diagnostics due to poor sample quality. However, if these were treponemes, the superficial location differs from what is seen in
DD and TAHD in that disease-associated treponemes in the latter are typically invasive and located deep within affected tissue [50]. The more superficial placement in this huemul more likely reflects opportunistic colonization by environmental contaminants rather than primary pathogens. The limitations of our investigation, however, preclude ruling out treponemes or other single or polymicrobial bacterial infections as part of the etiology of this disease.

In addition to abnormal hoof growth, a characteristic feature of TAHD disease in elk is osteomyelitis and interphalangeal osteoarthritis associated with broken and sloughed hooves [49, 50]. There is one published report of osteopathological changes in huemul limbs based on skeletal remains found in Argentina [17]. However, comparison of foot bones from BONP cases with elk was not possible because bone structures were not thoroughly inspected nor preserved in BONP diseased deer, and soft tissues were not available for examination in the skeletal specimens examined by Flueck and Smith-Flueck [17]. In Flueck’s report, examined skeletal remains spanned all age groups, without gender bias, and across a wide temporospatial range. Chronicity of lesions led the authors to favor a nutritional (e.g. selenium deficiency) versus an infectious etiology. Selenium and copper deficiency have been documented in TAHD affected and unaffected elk; however, any nutrition deficiency as a possible contributing factor in lesion development in the huemul of that or this current report, and elk remains to be further investigated. Notwithstanding, the short timeline from disease onset to death in the BONP huemul would not have allowed the development of the chronic, extensive bone remodeling that was seen in the Argentina cases described by Flueck and Smith Flueck [17], and a different underlying disease process is considered more likely.

The papillomatous appearance of huemul foot lesions is also reminiscent of those caused by several viruses including papillomavirus, pox or parapox viruses, and possibly foot and mouth disease. Though viral inclusion bodies were not seen on histological examination, electron microscopy of plastic-embedded skin preparations from the single available affected huemul (Case #10), confirmed the presence of viral particles and viral factories. In the latter, the dimensions and morphology of observed viruses and viral particles was consistent with viruses in the Chordopoxvirinae subfamily. Parapoxviruses can be morphologically distinguished from other Chordopoxvirinae in conventional negative staining electron microscopy by their ovoid appearance and the spiral tubule surrounding the virion’s surface, a distinctive diagnostic property of this genus [55]. Unfortunately, this technique can only be applied to unfixed samples, which were not available. Nevertheless, in the existing samples obtained from FFPE tissue mounted on glass slide from Case #10, intermediate stages of this poxvirus replication and assembly were identified. Stages of replication included membrane crescents surrounding a degenerated electron-dense matrix (most likely viroplasm), spherical immature virus enclosing viroplasm, and/or dense nucleoprotein surrounded by lipid bilayers and numerous mature particles viewed along their long or short axis. This is consistent with the entire spectrum of virus intermediates described in Vero cells infected with other parapoxviruses such as ORF virus and the Chordopoxvirinae prototype vaccinia virus [55]. Moreover, the presence of parapoxvirus DNA in Case #10 samples was confirmed by a pan-poxvirus PCR assay that targets shorter amplicon size ideal for recovery from FFPE tissues [32]. Recognizing that our investigation was limited to one animal due to sample accessibility constraints, testing of additional animals to better understand presence/absence and general relevance of viral infection in huemul foot lesions is warranted. Likewise, efforts to more completely characterize the viral sequence and determine whether infection crossed over from domestic cattle or potentially represents a novel virus is a high priority if additional samples become available.

Parapoxviruses, one of eleven genera within the Chordopoxvirinae subfamily, have been reported in a wide variety of wild and domestic mammals including cervids, bovids, camelids, rodents, and pinnipeds [56–59]. Of four known parapoxvirus species, the prototype is ORF
virus, which is endemic in most sheep and goat-raising countries. ORF virus has been reported in reindeer (*Rangifer tarandus*) and muskoxen (*Ovibos moschatus*) in Scandinavia [60–62] as well as in chamois (*Rupicapra rupicapra*) and ibex (*Capra ibex*) in Italy [59], and camels in Asia, Africa and the Middle East [58]. Another member of this genus, Parapoxvirus of red deer was first described in New Zealand [63] and has recently been found in Germany and Italy [59, 64]. The viral DNA in our study was closely aligned with the two additional species in the parapoxvirus genus, bovine papular stomatitis virus and pseudocowpoxvirus, which are mainly found in cattle.

Parapoxviruses are highly contagious and can be transmitted by direct contact between animals or indirectly by environmental contamination. Moreover, transmission between domestic and wild ungulates has been described, as has zoonotic transmission to humans [57, 65–67]. Proliferative pustular lesions have been attributed to parapoxviruses in several wild species. There are, however, very few reports of lesions on feet or limbs, which are considered atypical presentations in domestic animals [68]. Moreover, rarely do parapoxvirus infections become severe, extensive, and fail to spontaneously regress. The exception is disease caused by ORF virus in sheep and goats [57, 59, 68], semi-domesticated reindeer [60] and camels [58]. In cervids, most reports describe nonparapox or orthopox-like viruses in skin lesions of reindeer, mule deer (*Odocoileus hemionus hemionus*), black-tailed deer (*Odocoileus hemionus columbianus*), pudu (*Pudu puda*), and gazelle (*Gazella subgutturosa*) [69–74].

Viral DNA from Huemul Case #10 showed high identity (98%) with a bovine papular stomatitis virus isolated from cattle in the USA and a pseudocowpoxvirus isolated from human samples in Queensland, Australia. To the best of our knowledge, there are no reports of bovine papular stomatitis virus in deer or other wild ungulates, except for a suspect, non-confirmed case in a captive pudu in Chile [75]. Pseudocowpoxvirus infections have been diagnosed in Finnish reindeer (*Rangifer tarandus tarandus*) [76, 77], dromedary camels (*Camelus dromedarius*) in Sudan [58] and water buffalo (*Bubalus bubalis*) in Brazil [78]. In cattle, both papular stomatitis and pseudocowpox cause lesions on the muzzle, lips, oral mucosa and the teats. The lesions resemble those seen with vesicular stomatitis, bovine viral diarrhea and foot and mouth disease [65, 79]. Infection may also be asymptomatic. Similarly, lesions from pseudocowpoxvirus in reindeer and water buffalo are also restricted to erosions and ulcerations of the oral mucosa and tongue [77, 78]. In camels, the disease is characterized by papules that progress into scabs on the lips, muzzle, nares and eyelids and may extend into gum, palate and tongue [58]. Bovine papular stomatitis occurs worldwide in cattle and is usually of little clinical importance. Herd morbidity may be 100% but mortalities are rare. Infection can occur in animals of all ages, with higher incidence in the young. Immunity is of short duration, and reinfections can occur [79]. Pseudocowpoxvirus infections seem to be recurrent in domesticated species like reindeer and camels [58, 77]. Outbreaks in these species and in water buffalo have been associated with weather, age, husbandry, stress and overlap or contact with livestock, contaminated pastures or fomites [58, 77, 78]. Overall, huemul deer cases seem to differ from those described in cattle and semi-wild ungulates primarily in presentation (more proliferative than vesicular), body location (exclusively in feet), and severity (high morbidity-mortality).

It is possible that parapoxviruses behave differently in huemul deer than in other wild or domestic ungulates. It is also feasible that the etiopathogenesis of foot disease in huemul is similar to DD or TADH in that development may be polymicrobial, requiring two or more etiologic agents (viral and/or bacterial) to progress to severe disease. More research is needed to discern the roles of the pathogens identified in our study, as well as to investigate those that cause foot disease in other domestic and wild ruminants. For example, an investigation by Brandt et al [46] found that while bovine papilloma virus was highly prevalent in cattle with DD, the virus was unlikely to play a role in disease development and maintenance. Conversely,
the study showed that co-infecting treponemes were actively involved in disease etiology. Importantly however, many foot diseases of livestock that are prevalent worldwide and cause substantial economic losses remain poorly understood despite considerable efforts to elucidate their etiology.

Our inability to confirm viral EM and PCR findings with microarray analysis in Case #10 was unexpected. This could be related to sample quality and viral particle load. Target sequence fragments corresponding to probes present on the array, which may be distinct from the PCR targeted amplicon, might have been degraded beyond the capacity for hybridization. It is more likely, however, that presence of the pathogen was below the limit of microarray detection, which has generally been observed to be 100–1000 genomic copies [80], whereas limit of PCR detection is expected to be closer to a range of 10–100 copies. In addition, it is possible that, while sample preparation techniques sufficiently preserved particle integrity for visual identification by EM, the fixing and subsequent extraction procedures were not amenable to robust hybridization-based DNA detection.

The apparent concentration of huemul cases in HV, the only valley with a period of cattle presence, poses questions about a potential role for environmental contamination and subsequent disease transmission from feral livestock to huemul. However, cattle presence and disease development in deer did not overlap in time or space. Approximately 35 feral cattle were removed from HV between 2001 and 2004, rendering the valley cattle-free prior to the onset of foot disease in huemul deer in 2005. While specific efforts were not made to document abnormalities in culled cattle, many were slaughtered for human consumption and obvious lesions, such as those seen in huemul, were not observed. Similarly, hunters never reported cattle with mouth, face or foot lesions, nor lameness or recumbent animals. In the KV and BV, where cattle were not introduced and never seen, foot disease was documented in huemul but with lower frequency than in HV. This could reflect a true lower prevalence of disease in these valleys or, because of the low periodicity of visits, a detection bias due to decreased observation efforts in these remote locations. That infection began with huemul in HV and then spread through contact with conspecifics in the other valleys is a possibility, though movements of huemul between HV and the other valleys remain unconfirmed [25]. Notwithstanding, the distance between valleys is well within huemul deer reported movement range, 6.7 [81] to 9.0 km [82], and there are no important geographical barriers between HV and BV. Consequently, huemul movements between these valleys are feasible. Other components such as differences in home range size and patterns of movements between sexes [82], and potentially in habitat use, could also have influenced differential exposure, as male huemul were the more affected sex.

Despite significant challenges related to the remote location of BONP, the extreme environmental conditions at the site, and a restricted number of on-site staff who did as much as could be done with limited equipment, no previous necropsy or sampling training, and limited external support, we were able to identify, and in some cases monitor for the first time, the progression of life-threatening foot lesions in huemul deer in Chilean Patagonia. Even though parapoxvirus findings were limited to materials from one case, the potential implications of a disease caused by a highly-contagious and seemingly aggressive virus in this endangered deer should be readily acknowledged. Moreover, foot lesions reported here and recent Corynebacterium pseudotuberculosis infections in huemul at Cerro Castillo, Chile [22] highlight the need for improved capacity to detect, respond and potentially mitigate health risks in all remaining huemul populations. For all these reasons, we strongly recommend strengthening collaborations between government agencies, research facilities and NGOs to enable synergistic efforts and rapid response to future disease events threatening huemul deer.
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References

1. Daszak P, Cunningham AA, Hyatt AD. Emerging infectious diseases of wildlife- threats to biodiversity and human health. Science. 2000; 287: 443–449. PMID: 10642539

2. Harvell CD, Mitchell CE, Ward JR, Altizer S, Dobson AP, Ostfeld RS, et al. Climate warming and disease risks for terrestrial and marine biota. Science. 2002; 296: 2158–2162. https://doi.org/10.1126/science.1063699 PMID: 12077394

3. Lafferty K. Is disease increasing or decreasing, and does it impact or maintain biodiversity. J Parasitol. 2003; 89(Suppl.): S101–S105.

4. Smith KF, Sax DF, Lafferty KD. Evidence for the role of infectious disease in species extinction and endangerment. Conserv Biol. 2006; 20: 1349–1357. https://doi.org/10.1111/j.1523-1739.2006.00524.x PMID: 17092752

5. Smith KF, Acevedo-Whitehouse K, Pedersen AB. The role of infectious diseases in biological conservation. Anim Conserv. 2009; 12: 1–12. https://doi.org/10.1111/j.1467-1975.2008.00228.x

6. De Castro F, Bolker B. Mechanisms of disease-induced extinction. Ecol Lett. 2005; 8: 117–126. https://doi.org/10.1111/j.1461-0248.2004.00693.x

7. Timm SF, Munson L, Summers BA, Terio KA, Dubovi EJ, Rupprecht CE, et al. A suspected canine distemper epidemic as the cause of a catastrophic decline in Santa Catalina Island foxes (Urocyon littoralis catalinae). J Wildl Dis. 2009; 45: 333–343. https://doi.org/10.7589/0090-3558-45.2.333 PMID: 19395743

8. Seimon TA, Ayebare S, Sekisambu R, Muhindo E, Mitamba G, Greenbaum E, et al. Assessing the threat of amphibian chytrid fungus in the Albertine Rift: Past, present and future. PLoS One. 2015; 10. https://doi.org/10.1371/journal.pone.0145841 PMID: 26710251
9. McCallum H, Dobson A. Detecting disease and parasite threats to endangered species and ecosystems. Trends Ecol Evol.; 1995; 10: 190–194. https://doi.org/10.1016/S0169-5347(00)88050-3 PMID: 21237000

10. McCallum H. Disease and the dynamics of extinction. Philos Trans R Soc B Biol Sci. 2012; 367: 2828–2839. https://doi.org/10.1098/rstb.2012.0224 PMID: 22966138

11. Cabrera A, Yepes J. Mamíferos Sudamericanos. 2nd ed. Buenos Aires, Argentina: Ediar; 1960.

12. Díaz NI. El huemul (Hippocamelus bisulcus Molina, 1782): una perspectiva histórica. In: Díaz N, Smith-Flueck J, editors. El huemul Patagónico, un misterioso cérvido al borde de la extinción. Buenos Aires, Argentina: LOLA; 2000. pp. 1–32.

13. IUCN. IUCN Red List of Threatened Species, Version 2017 [Internet]. 2017 [cited 8 Aug 2017]. Available: http://www.iucnredlist.org.

14. López R, Serret A, Faúndez R, Palé G. Estado del conocimiento actual de la distribución del huemul (Hippocamelus bisulcus, Cervidae) en Argentina y Chile. Concepción, Chile; 1998.

15. Vila AR, Saucedo Galvez C, Aldridge D, Ramilo E, Corti González P. South Andean huemul Hippocamelus bisulcus (Molina 1782). In: Barbanti Duarte J, González S, editors. Neotropical cervidology: biology and medicine of Latin American deer. Sao Paulo, Brazil: FUNEP-IUCN; 2010. pp. 89–100.

16. Vila AR, López R, Pastore H, Faúndez R, Serret A. Current distribution and conservation of the huemul (Hippocamelus bisulcus) in Argentina and Chile. Mastozoología Neotrop. 2006; 13: 263–269.

17. Flueck WT, Smith-Flueck JAM. Age-independent osteopathology in skeletons of a South American cervid, the Patagonian huemul (Hippocamelus bisulcus). J Wildl Dis. 2008; 44: 636–648. https://doi.org/10.7589/0090-3558-44.3.636 PMID: 18689649

18. González-Acufía D, Saucedo GC, Corti P, Casanueva ME, Cicchino A. First Records of the Louse Solfenopetes binipilus (Insecta: Phthiraptera) and the Mite Psoroptes ovis (Arachnida: Acari) from Wild Southern Huemul (Hippocamelus bisulcus). J Wildl Dis. 2009; 45: 1235–1238. https://doi.org/10.7589/0090-3558-45.4.1235 PMID: 19901495

19. Chihuialaf RH, Stevenson VB, Saucedo C, Corti P, Blood mineral concentrations in the endangered huemul deer (Hippocamelus bisulcus) from Chilean Patagonia. J Wildl Dis. 2014; 50: 146–149. https://doi.org/10.7589/2013-03-063 PMID: 24171561

20. Corti P, Saucedo C, Herrera P. Evidence of bovine viral diarrhea, but absence of infectious bovine rhinotracheitis and bovine brucellosis in the endangered huemul deer (Hippocamelus bisulcus) in Chilean Patagonia. J Wildl Dis. 2013; 49: 744–746. https://doi.org/10.7589/2012-04-105 PMID: 23778636

21. Hinojosa A, Blumer E, Camacho A, Silva A, Quezada M, Brevis C. First report of fibroma in huemul (Hippocamelus bisulcus Molina 1782). Gayana. 2014; 78: 127–129. https://doi.org/10.4067/S0717-653820140000200006

22. Morales N, Aldridge D, Bbahamonde A, Carda J, Araya C, Muñoz R, et al. Corynebacterium pseudotuberculosis infection in patagonian huemul (Hippocamelus bisulcus). J Wildl Dis. 2017; 53: 621–624. https://doi.org/10.7589/2016-09-213 PMID: 28323562

23. Salgado M, Corti P, Verdugo C, Tomczowiak C, Moreira R, Durán K, et al. Evidence of Mycobacterium avium subsp. paratuberculosis (MAP) infection in huemul deer (Hippocamelus bisulcus) in patagonian fjords. Austral J Vet Sci. 2017; 49: 135–137.

24. Uhart M, Reissig E. Ungulados exóticos: amenaza sanitaria para el huemul. In: APN, editor. Reunión Binacional Argentino-Chilena sobre estrategias de conservación del huemul. 5th ed. Argentina: Administración de Parques Nacionales; 2006. pp. 25–27.

25. Briceño C, Knapp LA, Silva A, Paredes J, Avendaño I, Vargas A, et al. Detecting an increase in an Endangered huemul Hippocamelus bisulcus population following removal of cattle and cessation of poaching in coastal Patagonia, Chile. Oryx. 2013; 47: 273–279. https://doi.org/10.1017/S0030605312000014

26. Frid A. Observations on habitat use and social organization of a huemul Hippocamelus bisulcus coastal population in Chile. Biol Conserv. 1994; 67: 13–19. https://doi.org/10.1016/0012-1667(94)90035-5.

27. Carrasco JF, Casassa G, Rivera A. Climatología actual del Campo de Hielo Sur y posibles cambios por el incremento del efecto invernadero. An Inst Patagon Ser Cienc Nat. 1998; 119–128.

28. INE. Chile: Ciudades, Pueblos, Aldeas y Caseríos. Santiago, Chile, 2005.

29. Frid A. Habitat use by endangered huemul (Hippocamelus bisulcus): cattle, snow, and the problem of multiple causes. Biol Conserv. 2001; 100: 261–267. https://doi.org/10.1016/S0006-3207(01)00064-7.

30. Acosta G. Informe prospección de Huemules, sectores Fjordos Témpanos, Bahía Elizabeth y Glaciar Pío XI. P. N. Bernardo O’Higgins-XII Región. Santiago, Chile; 2001.

31. Seimon TA, Olson SH, Lee KJ, Rosen G, Ondzie A, Cameron K, et al. Adenovirus and herpesvirus diversity in free-ranging great apes in the Sangha region of the Republic of Congo. PLoS One. 2015; 10. https://doi.org/10.1371/journal.pone.0118543 PMID: 25781992
32. McAloose D, Rago MV, Di Martino M, Chirife A, Olson SH, Beltramino L, et al. Post-mortem findings in southern right whales *Eubalaena australis* at Peninsula Valdés, Argentina, 2003–2012. Dis Aquat Organ. 2016; 119: 17–36. https://doi.org/10.3354/dao02986 PMID: 27068500

33. Wellehan JFX, Johnson AJ, Harrach B, Benkó M, Pessier AP, Johnson CM, et al. Detection and analysis of six lizard adenoviruses by consensus primer PCR provides further evidence of a reptilian origin for the atadenoviruses. J Virol. 2004; 78: 13366–13369. https://doi.org/10.1128/JVI.78.23.13366-13369.2004 PMID: 15542689

34. VanDevanter DR, Warrener P, Bennett L, Schultz ER, Coulter S, Garber RL, et al. Detection and analysis of diverse herpesviral species by consensus primer PCR. J Clin Microbiol. 1996; 34: 1666–1671. PMID: 8784566

35. Johne R, Enderlein D, Nierer H, Müller H. Novel polyomavirus detected in the feces of a chimpanzee by nested broad-spectrum PCR. J Virol. 2005; 79: 3883–3887. https://doi.org/10.1128/JVI.79.6.3883-3887.2005 PMID: 15731285

36. Moureau G, Temam S, Gonzalez JP, Charrel RN, Grand G, de Lamballe X. A real-time RT-PCR method for the universal detection and identification of flaviviruses. Vector-Borne Zoonotic Dis. 2007; 7: 467–478. https://doi.org/10.1089/vbz.2007.0206 PMID: 18020965

37. Jaing C, Thissen JB, Gardner S, McLoughlin K, Slezak T, Bossart GD, et al. Pathogen surveillance in wild bottlenose dolphins *Tursiops truncatus*. Dis Aquat Organ. 2015; 116: 83–91. https://doi.org/10.3354/dao02917 PMID: 26480911

38. Gardner SN, Jaing CJ, McLoughlin KS, Slezak TR. A microbial detection array (MDA) for viral and bacterial detection. BMC Genomics. 2010; 11. https://doi.org/10.1186/1471-2164-11-668 PMID: 21108826

39. Jaing CJ, Thissen JB, Gardner SN, McLoughlin KS, Hullinger PJ, Monday NA, et al. Application of a pathogen microarray for the analysis of viruses and bacteria in clinical diagnostic samples from pigs. J Vet Diagnostic Invest. 2015; 27: 313–325. https://doi.org/10.1177/1040638715578484 PMID: 25855363

40. Devault AM, McLoughlin K, Jaing C, Gardner S, Porter TM, Enk JM, et al. Ancient pathogen DNA in archaeological samples detected with a Microbial Detection Array. Sci Rep. 2014; 4. https://doi.org/10.1038/srep04245 PMID: 24603850

41. Tellez J, Jaing C, Wang J, Green R, Chen M. Detection of Epstein-Barr virus (EBV) in human lymphoma tissue by a novel microbial detection array. Biomark Res. 2014; 2. https://doi.org/10.1186/s40364-014-0024-x PMID: 25635226

42. Edwards JF, Davis DS, Roffe TJ, Ramiro-Ibañez F, Elzer PH. Fusobacteriosis in captive wild-caught pronghorns (*Antilocapra americana*). Vet Pathol. 2001; 38: 549–552. https://doi.org/10.1354/vp.38-5-549 PMID: 11572563

43. Laurent J, Ruiz-Bascaran M, Marco I, Abarca ML, Crespo MJ, Franch J. Foot Infections Associated with *Arcanobacterium pyogenes* in Free-ranging Roosevelt Elk (*Cervus elaphus roosevelti*). J Wildl Dis. 2014; 50: 259–270. https://doi.org/10.7589/2013-07-276 PMID: 24632567

44. Wilson-Welder JH, Alt DP, Nally JE. 2015. Digital dermatitis in cattle: current bacterial and immunological findings. *Animals (Basel)*. 2015; 1114–1135. https://doi.org/10.3390/ani5040400 PMID: 26569318

45. Devault AM, McLoughlin K, Jaing C, Gardner S, Porter TM, Enk JM, et al. Microbial detection array. Vet Microbiol. 2005; 111: 171–180. https://doi.org/10.1016/j.vetmic.2005.10.016 PMID: 15731285

46. Brandt S, Apprich V, Hackl V, Tober R, Danzer M, Kainzbauer C, et al. Prevalence of bovine papillomavirus and Treponema DNA in bovine digital dermatitis lesions. Vet Microbiol. 2011; 148: 161–167. https://doi.org/10.1016/j.vetmic.2010.08.031 PMID: 20875931

47. Evans NJ, Blowey RW, Timofte D, Ischerwood DR, Brown JM, Murray R, et al. Association between bovine digital dermatitis treponemes and a range of “non-healing” bovine hoof disorders. Vet Rec. 2011; 168. https://doi.org/10.1136/vr.c5487 PMID: 21493554

48. Weaver GV, Domenech J, Thiemann AR, Karesh WB. Foot and mouth disease: a look from the wild side. *J Wildl Dis.* 2013; 49: 759–785. https://doi.org/10.7589/2012-11-276 PMID: 24502706

49. Han S, Mansfield KG. Severe hoof disease in free-ranging Roosevelt elk (*Cervus elaphus roosevelti*) in southwestern Washington, USA. *J Wildl Dis.* 2014; 50: 259–270. https://doi.org/10.7589/2013-07-168 PMID: 24484504

50. Han S, Mansfield KG, Bradway DS, Besser TE, Read DH, Haldorson GJ, et al. Treponema-associated hoof disease of free-ranging elk (*Cervus elaphus*) in Southwestern Washington State, USA. Vet Pathol. 2019; 56: 118–132. https://doi.org/10.1177/0300985818798108 PMID: 30244661

51. Cagatay IT, Hickford JGH. Update on ovine footrot in New Zealand: isolation, identification, and characterization of *Dichelobacter nodosus* strains. Vet Microbiol. 2005; 111: 171–180. https://doi.org/10.1016/j.vetmic.2005.09.010 PMID: 16280202
52. Clegg SR, Mansfield KG, Newbrook K, Sullivan LE, Blowey RW, Carter SD, et al. Isolation of digital dermatitis treponemes from hoof lesions in wild North American elk (Cervus elaphus) in Washington State, USA. J Clin Microbiol. 2015; 53: 88–94. https://doi.org/10.1128/JCM.02276-14 PMID: 25355757

53. Krull AC, Cooper VL, Coatney JW, Shearer JK, Gordon PJ, Plummer PJ. A highly effective protocol for the rapid and consistent induction of digital dermatitis in Holstein calves. PLoS ONE. 2016; 11(4): e0154481. https://doi.org/10.1371/journal.pone.0154481

54. Edwards AM, Dymock D, Jenkinson HF. From tooth to hoof: treponemes in tissue-destructive diseases. Vet Microbiol. 2003; 94: 767–780. PMID: 12694441

55. Spehner D, De Carlo S, Drillien R, Weiland F, Mildner K, Hanau D, et al. Appearance of the bona fide spiral tubule of ORF virus is dependent on an intact 10-kilodalton viral protein. J Virol. 2004; 78: 8085–8093. https://doi.org/10.1128/JVI.78.10.8085-8093.2004 PMID: 15254180

56. Robinson A, Kerr J. Poxviruses. In: Williams E, Baker I, editors. Infectious diseases of wild mammals. 3rd ed. Ames: Iowa State University Press; 2001. pp. 179–201.

57. Delhon G, Tulman ER, Afonso CL, Lu Z, de la Concha-Bermejillo A, Lehmkuhl HD, et al. Genomes of the parapox viruses orf virus and bovine papular stomatitis virus. J Virol. 2004; 78: 168–177. https://doi.org/10.1128/JVI.78.1.168-177.2004 PMID: 14671098

58. Khalafalla AI, El-Sabagh IM, Al-Busada KA, Al-Mubarak AI, Ali YH. Phylogenetic analysis of eight sudanese camel contagious ecthyma viruses based on B2L gene sequence. Virol J. 2015; 12: 124–132. https://doi.org/10.1186/s12985-015-0348-7 PMID: 26260127

59. Scagliarini A, Vaccari F, Turrini F, Bianchi A, Cordioli P, Lavazza A. Parapoxvirus infections of red deer, Italy. Emerg Infect Dis. 2011; 17: 684–687. https://doi.org/10.3201/eid1704.101454 PMID: 21470460

60. Tryland M, Josefsen TD, Oksanen A, Aschfalk A. Contagious ecthyma in Norwegian semidomesticated reindeer (Rangifer tarandus tarandus). Vet Rec. 2001; 149: 394–395. PMID: 11601519

61. Klein J, Tryland M. Characterisation of parapoxviruses isolated from Norwegian semi-domesticated reindeer (Rangifer tarandus tarandus). Virol J. 2005; 2: 79. https://doi.org/10.1186/1743-422X-2-79 PMID: 16140341

62. Vikøren T, Haugum T, Schulze J, Akerstedt J, Liliehaug A, Tryland M. A severe outbreak of contagious ecthyma (orf) in a free-ranging musk ox (Ovibos moschatus) population in Norway. Vet Microbiol. 2008; 127: 10–20. https://doi.org/10.1016/j.vetmic.2007.07.029 PMID: 17768017

63. Robinson AJ, Mercer AA. Parapoxvirus of red deer: evidence for its inclusion as a new member in the genus parapoxivirus. Virol. 1995; 208: 812–815.

64. Friederichs S, Krebs S, Blum H, Lang H, Büttner M. Parapoxvirus (PPV) of red deer reveals subclinical infection and confirms a unique species. J Gen Virol. 2015; 96:1446–62. https://doi.org/10.1099/vir.0.000080 PMID: 25701822

65. Barrett J, Grant McFadden G. Origin and Evolution of Poxviruses. In: Domingo E, Parrish C, Holland J, editors. Origin and Evolution of Viruses. 2nd ed. London, UK: Academic Press; 2008. pp. 431–446.

66. Roess AA, McCollum AM, Gruszynski K, Zhao H, Davidson W, Lafon N, et al. 2013. Surveillance of parapoxvirus among ruminants in Virginia and Connecticut. Zoonoses Public Health. 2013; 60: 543–8. https://doi.org/10.1111/zph.12036 PMID: 23398718

67. Tryland M, Klein J, Berger T, Josefsen TD, das Neves CG, Oksanen A, et al. 2013. Experimental parapoxvirus infection (contagious ecthyma) in semi-domesticated reindeer (Rangifer tarandus tarandus). Vet Microbiol. 2013; 162: 499–506. https://doi.org/10.1016/j.vetmic.2012.10.039 PMID: 23201244

68. Smith GW, Scherba G, Constable PD, Hsiao V, Behr MJ, Morin DE. Atypical parapoxvirus infection in sheep. J Vet Intern Med. 2002; 16: 287–292. https://doi.org/10.1111/j.1939-1676.2002.tb02371.x PMID: 12041659

69. Baughman B, Zhang S, Jin L, Pace LW, Cooley J, Yan L, et al. Diagnosis of Deerpox virus infection in a white-tailed deer (Odocoileus virginianus) fawn. J Vet Diagnostic Invest. 2011; 23: 965–970. https://doi.org/10.1177/1040638711416621 PMID: 21908356

70. Bracht AJ, Armien AG, Carrillo C, O’Hearn ES, Fabian AW, Moran KE, et al. Isolation and characterization of a Cervidpoxvirus from a goitered gazelle (Gazella subgutturosa) from a zoologic park in Minnesota. J Zoo Wildl Med. 2013; 44: 589–95. https://doi.org/10.1638/2012-0090R2.1 PMID: 24063086

71. Junge RE, Duncan MC, Miller RE, Gregg D, Kombert M. Clinical Presentation and Antiviral Therapy for Poxvirus Infection in Pudu (Pudu puda). J Zoo Wildl Med. American Association of Zoo Veterinarians; 2000; 31: 412–418. Available: http://www.jstor.org/stable/20996024. https://doi.org/10.1638/1042-7260(2000)031[0412:CPAAFT]2.0.CO;2 PMID: 11237153

72. McNamara T, Gregg D. A novel pox infection in pudus (Pudu puda), Proc Assoc Zoo Vet 1994: 257–264.
73. Patton JF, Nordhausen RW, Woods LW, MacLachlan NJ. Isolation of a poxvirus from a Black-tailed deer (*Odocoileus hemionus columbianus*). J Wildl Dis. 1996; 32: 531–533. https://doi.org/10.7589/0090-3558-32.3.531 PMID: 8827682

74. Williams E, Becerra V, Thorne E, Graham T, Owens M, Nunamaker C. Spontaneous poxviral dermatitis and keratoconjunctivitis in free-ranging mule deer (*Odocoileus hemionus*) in Wyoming. J Wildl Dis. 1985; 21: 430–433. PMID: 3001374

75. Paredes E, Moroni M, Verdugo C. Sospecha de infección por parapoxvirus en pudú (*Pudu pudu*) en cautiverio. Resúmenes del XX Congreso Panamericano de Ciencias Veterinarias. Santiago, Chile; 2006.

76. Tikkanen MK, McInnes CJ, Mercer AA, Büttner M, Tuimala J, Hirvelä-Koski V, et al. Recent isolates of parapoxvirus of Finnish reindeer (*Rangifer tarandus tarandus*) are closely related to bovine pseudocowpox virus. J Gen Virol. 2004; 85: 1413–1418. https://doi.org/10.1099/vir.0.79781-0 PMID: 15166423

77. Hautaniemi M, Ueda N, Tuimala J, Mercer AA, Lahdenperä J, McInnes CJ. The genome of pseudocowpoxvirus: comparison of a reindeer isolate and a reference strain. J Gen Virol. 2010; 91: 1560–1576. https://doi.org/10.1099/vir.0.018374-0 PMID: 20107016

78. Laguardia-Nascimento M, Ferreira de Oliveira AP, Fernandes FRP, Vasconcelos Rivetti Junior A, Fernandes Camargos M, Fonseca Júnior AA. Detection of pseudocowpox virus in water buffalo (*Bubalus bubalis*) with vesicular disease in the state of São Paulo, Brazil, in 2016. Vet Quart. 2017; 37: 16–22. https://doi.org/10.1080/01652176.2016.1252479 PMID: 27774853

79. Underwood W, Blauwikel R, Delano M, Gillesby R, Mischler S, Schoell A. Biology and diseases of ruminants (sheep, goats, and cattle). In: Fox J, Anderson L, Otto G, Pritchett-Corning K, Whary M, editors. Laboratory animal medicine. 3rd ed. Oxford, UK: Academic Press; 2015. pp. 623–694.

80. Thissen JB, McLoughlin K, Gardner S, Gu P, Mabery S, Slezak T, et al. Analysis of sensitivity and rapid hybridization of a multiplexed Microbial Detection Microarray. J Virol Methods. Elsevier B.V.; 2014; 201: 73–78. https://doi.org/10.1016/j.jviromet.2014.01.024 PMID: 24602557

81. Movimiento Rau J., hábitat y velocidad del huemul del sur (*Hippocamelus bisulcus*) (Artiodactyla, Cervidae). Not Mens del Mus Nac Hist Nat. 1980; 281–282: 7–9.

82. Gill R, Saucedo Galvez C, Aldridge D, Morgan G. Ranging behaviour of huemul in relation to habitat and landscape. J Zool. 2008; 274: 254–260. https://doi.org/10.1111/j.1469-7998.2007.00378.x