First report of pre-Hispanic Fasciola hepatica from South America revealed by ancient DNA

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Introduction

Fasciolasis is a zoonotic parasitic disease caused by the liver flukes Fasciola hepatica and Fasciola gigantica (Trematoda: Digenea). This helminthic disease is of worldwide distribution (Mas-Coma et al., 2009) and is considered as an important veterinary health problem due to the substantive economic losses it gives rise to livestock husbandry. Moreover, it is of great public health importance in some countries, due to its high pathogenicity (Mas-Coma et al., 2014) and is therefore included within the group of Foodborne Trematodiases among Neglected Tropical Diseases (NTDs) by the World Health Organization (WHO, 2013).

Fasciola hepatica has a worldwide distribution and F. gigantica is found in tropical climates and restricted to Africa, the Middle East and South and East Asia. In America, fasciolasis is caused by F. hepatica and transmitted by many different intermediate snail hosts belonging to the family Lymnaeidae, mainly species included within the GalbaLlFlexis group. In South America, human endemic areas have been described in Andean regions, mainly in highlands of Bolivia, Peru and Chile, and sporadic cases are reported in other countries (Mas-Coma, 2003; World Health Organization, 2013; Carmona and Tort, 2016).

It is generally assumed that entry of F. hepatica to America coincided with the first arrival of the Europeans and their associated livestock in the late 15th century. Throughout the 500 years since its introduction, the parasite gained new definitive hosts among native species. The South American camelids – llamas, alpacas, vicuñas and guanacos – the natural livestock of the Andean region, might have represented the first to be parasitized, since these species would graze along with the introduced livestock (Mas-Coma et al., 2009; Bargues et al., 2017). The parasite is now widespread in livestock and can be mapped across the whole South America and certain regions of North America.

Argentina has a large livestock production, where sheep and cattle constitute important economic sources. Animal fasciolasis is currently found in spots across the country, according to official slaughterhouses, and the most important hosts are cattle and sheep. Goats, horses, pigs and some wild native and non-native mammals (deer, vicuña, guanacos, llamas, rabbits, hares and capybaras) are also found infected by F. hepatica. Despite the fragmented and anecdotal nature of several reports of liver flukes in South American wildlife, it is evident that diverse species can host the parasite and eventually act as reservoirs (Issia et al., 2009; Carmona and Tort, 2016).

Given the medical and veterinary importance of fasciolasis, there are several multidisciplinary studies investigating parasite origins and dispersals across the world; shared objectives include elucidation of palaeobiogeographical origins and colonization within and between continents (Mas-Coma et al., 2009).
Palaeoparasitology is the study of parasite remains from archaeological and palaeontological sites (Ferreira, 2014), focused on the knowledge of parasite-induced illness of humans in the past and on the palaeoecological knowledge of the environment, ecology, settlement, diet, hygiene and health in the antiquity (Reinhard, 1992). In a previous palaeoparasitological study, digenetic eggs similar to *F. hepatica* were found in coprolites from one of the two endemic species of deer inhabiting the narrow Andean-Patagonian temperate forest strip in the west of southern America, the southern pudú (*Pudu puda*) and the hue-mul (*Hippocamelus bisulcus*). The samples were obtained from an archaeological site of Patagonia named ‘Cueva Parque Diana’ (CPD) and were dated around 2300 years B.P. (Late Holocene) (Beltrame et al., 2017). In order to contribute with the study of the palaeobiogeographical origins of *F. hepatica* and their settlement across America, the main objective of the present study is to identify the digenesean eggs found from CPD from an ancient DNA (aDNA) study.

**Material and methods**

**Sample collection**

In a previous study (Beltrame et al., 2017), 34 coprolites were processed for palaeoparasitological purposes. Coprolites belonged to native deer identified as *P. puda* or *H. bisulcus*. Samples were collected from the CPD archaeological site, Lanín National Park, North Patagonia, Argentina (40°19′93″S, 71°20′74″W). This site is a rock shelter part of the archaeological locality named Meliquina. It is located at 964 m.a.s.l. and 50 m close to the Hermoso River. The archaeological sequence was divided into three components representing different hunter–gatherer occupation processes. The Upper Component was dated between 760 ± 60 and 580 ± 60 14C years B.P. (vegetal charcoal), the Middle Component was dated between 990 ± 60 and 900 ± 60 14C years B.P. (vegetal charcoal) and the Lower Component was dated at 2370 ± 70 14C years B.P. (vegetal charcoal). The site was occupied by hunters–gatherers and fishermen along the late Holocene (Pérez, 2010; Pérez et al., 2015). The weather in the area is cold and wet, with annual precipitations around 1500–2000 mm.

Eighteen of the 34 samples were positive for digenesean eggs. Positive coprolites were found in the upper, middle and lower components. The analysed eggs (Fig. 1 in this study belong to two positive samples studied in Beltrame et al. (2017) from the lower component. The eggs were identified under a light microscope (100× magnification) and were manually isolated by the use of a micropipette and stored in PCR tubes with phosphate-buffered saline. The isolated eggs were used for aDNA extraction, amplification and sequencing of partial regions from the cytochrome c oxidase subunit 1 (COI) and the nicotinamide adenine dinucleotide dehydrogenase subunit 1 (NADHI) mitochondrial genes.

**aDNA extraction and polymerase chain reaction amplification**

Before DNA extraction, 30 eggs were washed three times in ultrapure water (Invitrogen by Thermo Fisher Scientific, Waltham, MA, USA). Once washed, a first disruption step was performed by five continuous cycles of freezing (immersion in liquid nitrogen for 10 s) and heating (immersion of the tube in boiling water for 2–3 s).

DNA was extracted using the ZR Fecal DNA Miniprep kit (Zymo Research, Irvine, CA, USA), following the manufacturer’s instructions. PCR reactions were individually carried out for each partial fragment of the *mtDNA* genes coding for NADHI and COI. Each reaction was constituted into a final volume of 50 μL, containing 37.5 μL of ultrapure distilled water (Invitrogen), 5 μL of 10× buffer, 4.5 μL of PCR mix containing 1 mmol of MgCl₂, (2 μL) 0.2 mM of DNTPs (1 μL), 50 pmol of each primer (1 μL) and 2.5 U of *Taq* Polymerase (0.5 μL) (all reagents from Invitrogen), plus 3 μL of each template. PCR primers and temperature settings were previously described by Ichikawa and Itagaki (2012) for NADHI and COI. The primers’ sequences were 5′-AAGGATGTTGCTTGTGCTGG-3′ (forward), 5′-GGAGTACGGTTACATTACAATTCACA-3′ (reverse), for NADHI, and 5′-ACGGTGTGCAATTAGGCACATTG-3′ (forward), 5′-CTGATACACATAACCTCT-3′ (reverse), for COI. Negative controls (ultrapure water and PCR reagents) were also ran along with templates amplification.

Amplicons were visualized in 2% agarose gels stained with SybrSafe (Invitrogen) using a blue light transilluminator (Safe Imager, Invitrogen), and purified by a commercial kit (DNA Clean & Concentrator-5, Zymo Research), according to the manufacturer’s instructions. Purified PCR products were electrophoresed into 10 μL and submitted to Macrogen’s sequencing service (Seoul, Korea), for capillary electrophoretic sequencing using the same primers as for the PCR reactions. Once available, the sequences were submitted to the GenBank database (NCBI).

**Data analysis**

Electropherograms were scored and analysed using Chromas2.01 (Technelysium, Helensvale, QLD, Australia) and aligned using CLUSTALW2 (Larkin et al., 2007) in MEGA 7.0.26 (Kumar et al., 2016) using default settings. Homologies were performed using the BLASTN programme from the National Center for Biotechnology Information website (http://www.ncbi.nlm.nih.gov/BLAST).

In order to prove the phylogenetic identity, our mitochondrial partial sequences of aDNA from COI and NADHI were compared with the corresponding mitochondrial sequences of COI and NADHI of current individuals of *F. hepatica*. These last mitochondrial sequences of both molecular markers correspond to the
following GenBank accession numbers: KR422380–KR422388, MG870561, MG870563–MG870566, MG870568–MG870570, MG987190, MG9871902, LC273097, LC273100, LC273110, LC273111 and LC273113 from COI; and LC273198–LC273202, MG972375–MG972379, MG972405–MG972409, LC076246, LC076249, LC076255, LC076259, LC076261, KR422389–KR422393 and KJ852771 from NADH. Both COI and NADHI parasites were selected from several continental regions in order to include a broad representation of genetic diversity of these parasites characterized by a wide distribution worldwide. As out-groups we used mitochondrial sequences from two species of the family Fasciolidae (*F. gigantica* and *Fascioloides magna*); with the GenBank accession numbers AB385622, AB385621, AB385620, AB207176, AB207181, EF534996, EF534997 and EF534998 from COI; and MG972405, MG972406, MG972407, MG972408, MG972409, EF534999 and EF535000 from NADHI.

**Phylogenetic inferences**

We used jModelTest (Darriba *et al*., 2012) to infer the best-fit substitution model for both sets of data (COI and NADH). To assess the robustness of parameter estimates, four independent chains were run with identical settings. Log-files were analysed to assess the robustness of parameter estimates, four independent chains were each adequately sampling the same probability distribution and effective sample sizes for all parameters of interest were >500, conditions suggested by the authors for the proper functioning of the analysis.

A phylogenetic tree was also constructed using a maximum likelihood approach (ML) implemented in MEGA7 (Kumar *et al*., 2016), considering the same out-groups used in the Bayesian phylogenetic inference. The programme implements simultaneous Nearest Neighbor Interchanges (NNIs) to improve the phylogenetic inferences (ML and Bayesian) strongly support the hypothesis that our target sequences of deer effectively correspond to *F. hepatica* (Fig. 2). In our phylogenetic tree, these two sequences of mitochondrial aDNA were clearly included within the clade of *F. hepatica*, showing high node supports using both phylogenetic inferences. Also, these sequences showed high genetic distances relative to *F. gigantica* and *F. magna* within the phylogenetic tree.

**Discussion**

The sequences of the mitochondrial aDNA used in this study demonstrate that the trematode eggs from native deer coprolites from the CPD archaeological site belong to *F. hepatica*. The results confirm the presence of this trematode in South America from at least 2300 years B.P. This is the first report and the first aDNA study of *F. hepatica* in South America prior to the arrival of the European cattle in the 15th century. Previous palaeoparasitological studies have reported the presence of this parasite in Europe and Asia (e.g. Bouchet, 1995; Bouchet *et al*., 2003; Dittmar and Teegen, 2003; Askari *et al*., 2018). The first study that identifies aDNA of *Fasciola* sp. was made by Soe *et al.* (2015). In this study, *Fasciola* sp. eggs were recovered from environmental samples collected at a Viking-age settlement in Viborg, Denmark, dated at 1018–1030 A.D. Ancient DNA studies also were performed from European archaeological sites such as Côte *et al.* (2016) and Soe *et al.* (2018). In a recent study, Le Bailly *et al.* (2019) found one trematode egg in domestic camelds which could belong to *Fasciola* recovered from the pre-Hispanic Chimú culture site of Huanchaquto-Las Llamas, Peru. Although its diagnosis has not been confirmed yet, this would correspond to an additional evidence of the presence of the liver fluke genus *Fasciola* in pre-Hispanic times of America.

The current knowledge of the presence of *F. hepatica* on current native deer from Patagonia is limited. Some studies reported the presence of *F. hepatica* in the southern pudu (*Cortés, 2006; Bravo Antilef, 2013*) and in the huemul (*Díaz and Smith-Flueck, 2000; Serret, 2001*). It was also registered in the introduced red deer (*Cervus elaphus*) (*Flueck and Smith-Flueck, 2012*). Our COI sequence of aDNA was mostly related with contemporary South American DNA sequences (Ecuador and Peru), and was most distant to DNA sequences from Asia, Africa and Europe. However, the NADH sequence not only showed a great phylogenetic affinity with sequences of South America, but with Armenia and Egypt (see Fig. 2). The objective of this work was to identify digenean eggs found from deer coprolites from an aDNA study, without a deeper more conclusive evaluation of their possible palaeobiogeographical origin or colonization routes within this continent. In this sense, there are many methodological limitations that hinder the inference about a
possible geographical origin of these aDNA sequences reported here, beyond including contemporary DNA samples from various regions of the world in our phylogenetic analyses. In principle, knowledge about the dynamics of migration and colonization of this parasite among different continents seems to be very complex. Demographic patterns of *F. hepatica* have not only been conditioned by the regional history of domestic animal translocations performed by humans in recent decades, but also to their associations with populations of native mammal species. The latter can lead to confusing the biogeographic and demographic history of *F. hepatica* at the continental level, especially when the ancient material is compared with current DNA samples. The need for future palaeoparasitological studies around the world is evident in order to contribute to the palaeobiogeographical origin of this trematode.

The fact that *F. hepatica* was present in America before the arrival of the Europeans in the 15th century indicates that the species was not first introduced by the European cattle during this time, but there was another alternative route of prior entry. At the moment our data do not allow us to propose a plausible
hypothosis about the possible entry of *F. hepatica* to the American continent prior to this period. Future palaeoparasitological studies are needed which should consider the different migratory routes into the American continent in pre-Hispanic times. Evolutionary relationship among *F. hepatica* and American native hosts, both definitive and intermediate, is an interesting point to be studied in future studies. The pre-Hispanic presence of *F. hepatica* in South America brings new insights to the common assumption on the palaeobiogeography and settlement of this species.

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None.

**Conflict of interest**
None.

**Ethical standards**
Not applicable.

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