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Phylogenetic and ecological factors affecting the sharing of helminths between native and introduced rodents in Central Chile

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Abstract

In order to analyse the effect of hosts' relationships and the helminthic load on the switching of parasites between native and introduced hosts, we sampled rodents belonging to two suborders from Central Chile. We compared the number of helminthic species shared between murids (introduced) and cricetid (native, same suborder) rodents to those shared between murids and hystricomorphs (native, different suborder), and we assessed the association between parasitic presence, abundance and geographical dispersion in source hosts to the presence and abundance in recipient hosts. Introduced rodent species shared more helminth species with cricetid rodents than with non-cricetids. Presence and abundance in recipient hosts was not associated with the prevalence and mean abundance in source hosts' population. The mean abundance of parasites in source hosts throughout the territory and wider dispersion was positively associated with the likelihood of being shared with a recipient host. Closer relationships between native and introduced hosts and high parasitic abundance and dispersion could facilitate host switching of helminths between native and introduced rodents. This work provides the first documentation of the importance of parasitic abundance and dispersion on the switching of parasites between native and introduced hosts.

Introduction

Biological invasions, including parasites as invaders, have occupied an important position in conservation biology due to their importance in the processes of species loss (Wilcove and Master, 2005; Taraschewski, 2006). The process of host switching of introduced parasites or pathogens from introduced hosts is known as parasite spillover (Grabner et al. 2015; Morand et al. 2015), and has been observed in helminths transmitted by rodents (Smith and Carpenter, 2006; Romeo et al. 2015; Loxton et al. 2017). The consequences of the spillover of parasites and pathogens have been studied not only in native hosts (Barrett et al. 2002; Tompkins et al. 2003), but in human populations as well (Bordes et al. 2015). Spillover can also occur in the opposite direction, from native to introduced host (Barton, 1997). In this case, parasites can be amplified by introduced hosts and then transmitted back to native hosts, which is known as spillback (Kelly et al. 2009; Mastitsky and Veres, 2010). Alternatively, native parasites may not replicate in introduced hosts, which is known as the dilution effect (Johnson and Thieltges, 2010). Thus, several studies have focused on the effect of introduced parasites and hosts on native parasites and hosts (Macneil et al. 2003; Taraschewski, 2006; Paterson et al. 2011; Young et al. 2017). However, few surveys have studied factors that favour host switching of parasites between native and introduced hosts (e.g. Landaeta-Aqueveque et al. 2014). Regarding phylogenetic factors, although generalist native parasites have been found in introduced rodent species belonging to different families (Pisanu et al. 2009), some studies show that there is a decrease in the probability of transmission as an effect of increase in the taxonomic distance between hosts (Wells et al. 2015; Young et al. 2017). However, these predictions have seldom been quantitatively studied. Therefore, in this paper, we hypothesize that introduced host species share a larger number of parasite species with native species to which they are more closely related.

Many ecological factors can also facilitate host switching of parasites between native and introduced hosts. In the context of parasitism, propagule pressure has been defined as the number of parasites that arrive with the introduced host to the new territory (MacLeod et al. 2010). However, recognizing that spillover of parasites is a subsequent step in the invasion process, propagule pressure can also be understood as the parasite load in the source host population (Hatcher et al. 2012). However, the importance of this factor has not been studied. Thus, we hypothesize that the higher the prevalence and mean abundance of parasites in the

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source host population, the higher the probability of parasitic presence and the higher the abundance of parasites in the recipient host population.

In continental Chile, a region of a substantially isolated nature (see Landaeta-Aqueveque et al. 2014), there is evidence of parasite transmission between native and introduced rodents (Landaeta-Aqueveque et al. 2007a,b). Rodents in Central Chile belong to two suborders, Myomorpha, including the families Muridae (introduced species) and Cricetidae (native species), and Hystricomorpha (native species, hereafter non-cricetid) including the families Octodontidae and Abrocomidae (Muñoz-Pedreros, 2009). We studied host switching of parasites between native and introduced rodents with two aims. The first was to compare the number of parasite species that introduced rodents (murids) shared with cricetid rodents to the number of parasite species that introduced rodents shared with non-cricetid rodents. Thus, we quantitatively analysed the importance of host relatedness. The second aim was to assess the association between the prevalence and mean abundance of parasites in source host populations and the presence and abundance of parasites in the recipient hosts. Thus, this is the first work to study the importance of these variables in the sharing of parasites between native and introduced mammal hosts.

Materials and methods

From 2002 to 2011, we sampled adult rodents in 11 localities in Chile, from 31°S to 33°S, including protected and non-protected wild areas, agricultural areas and an urban settlement, all at altitudes lower than 1100 m.a.s.l. (see details of trapping localities in Fig. 1 and Table 1). Rodents were caught with live traps and killed with an isoflurane overdose. Viscera and cavities were examined for the presence of helminths, which were examined under a light microscope. Nematodes were cleared with lactophenol or ethanol-glycerin, and cestodes were stained with carmine-HCl and were identified using Anderson et al. (2009) keys for nematodes and Khalil et al. (1994) keys for cestodes, and published descriptions of helminths of rodents. The Comité de Ética of the Facultad de Ciencias Veterinarias y Pecuarias (Ethics Committee of the Faculty of Veterinary and Animal Sciences) at the Universidad de Chile approved and certified the study (certificate without number, 15 April 2011), and the Servicio Agrícola y Ganadero (Agricultural and Livestock Service) of Chile authorized trapping (resolution certificates 2041 and 6652 to C.L.-A.).

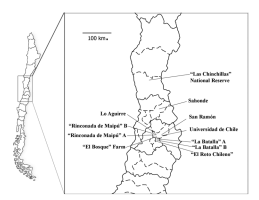


Fig. 1. Map of Chile with localities where rodents were trapped. Details are given in Table 1.

Each set of host specimens obtained from the same locality and over a period of <31 days was considered a study unit (SU). The terminology used to describe parasitic assemblages (locality, prevalence, abundance, mean abundance) follows Bush *et al.* (1997); 'presence' as variable refers to the dichotomous 'presence/absence' variable.

Fisher's exact test was used to compare the proportion of species of parasites shared with murid (introduced) rodents between cricetid and non-cricetid native rodents. To avoid consideration of parasitic species that did not have the opportunity for host switching, we included only species of parasites found in SUs that harboured both native and introduced rodents.

Simple binomial negative regressions were performed to assess the association between the mean abundance and prevalence of a parasite species in a source host population with the abundance of this species in individuals of the recipient population in the same SU. Thus, the abundance of the parasite in the recipient host was considered to be the dependent variable, and the mean abundance and prevalence of this parasite in the source host population in the SU in which this recipient host was found were considered to be the independent variables. Similarly, we used logistic regression to assess the association between the prevalence and the mean abundance of a parasite species in the source host population with the presence of this species in individuals of the recipient population in the same SU. For simplicity, we will use the terms source and recipient population even when the sources or the recipients could be a community.

Table 1. Details of localities where rodents were trapped

| Locality | District | Region | Details (Coordinates) |
|----------------------------------|-----------------|---------------------|---------------------------------------------------------------------------------------|
| Las Chinchillas National Reserve | Illapel | Coquimbo Region | Protected area (31°30′31.76″S; 71°6′21.93″W) |
| Sahonde | Putaendo | Valparaíso Region | Agricultural landscape (32°36′52.33″S; 70°40′50.18″W) |
| Lo Aguirre | Pudahuel | Metropolitan Region | Non-protected wild area (33°26′56.63″S; 70°50′5.78″W) |
| 'El Bosque' Farm | Maipú | Metropolitan Region | Agricultural landscape (33°32′43.23″S; 70°48′4.64″W) |
| 'Rinconada de Maipú'-A | Maipú | Metropolitan Region | Agricultural landscape (33°29′35.41″S; 70°48′54.75″W) |
| 'Rinconada de Maipú'-B | Maipú | Metropolitan Region | Non-protected wild area (33°29′11.81″S; 70°49′8.43″W) |
| 'La Batalla'-A | Calera de Tango | Metropolitan Region | Agricultural landscape (33°39′10.42″S; 70°47′14.63″W) |
| 'La Batalla'-B | Calera de Tango | Metropolitan Region | Non-protected wild area (33°40′9.47″S; 70°48′4.82″W) |
| 'El Roto Chileno' | Talagante | Metropolitan Region | Non-protected wild area close to agricultural landscape (33°41′1.21″S; 70°49′40.30″O) |
| Universidad de Chile | La Pintana | Metropolitan Region | Agricultural landscape (33°34′19.71″S; 70°37′53.39″W) |
| San Ramón | San Ramón | Metropolitan Region | Urban house (33°32′21.89″S; 70°39′2.48″W) |

Finally, the associations of the next independent variables were assessed by means of simple logistic regressions: (i) 'the number of localities in which a parasite species is present', (ii) 'the mean abundance of a parasite species in its source hosts among all of the SUs in which it was found' and (iii) 'the prevalence of a parasite species in its source hosts among all of the SUs in which it was found', with 'the odds of being or not being shared with a recipient host' (the dependent variable). These mean abundances and prevalences were also compared between shared and not shared parasite species by means of Wilcoxon rank-sum (Mann–Whitney) test. In both cases, only parasites of myomorph rodents were considered in order to control for host relatedness. Stata/SE 11.1 software (StataCorp LP) was used to perform the statistical analyses.

Results

Overall results

A total of 353 host individuals, belonging to nine species, were studied. The native species and their number of individuals (in parentheses) were the following: Abrocoma bennetti Waterhouse, 1837 (5) – Abrocomidae; Octodon degus Molina, 1782 (31) – Octodontidae; Abrothrix longipilis Waterhouse, 1837 (4); Abrothrix olivaceus Waterhouse, 1837 (119); Oligoryzomys longicaudatus Bennett, 1832 (10); and Phyllotis darwini Waterhouse, 1837 (49) – Cricetidae. The introduced species and the number of individuals (in parentheses) were: Mus musculus Linnaeus, 1758 (84); Rattus norvegicus Berkenhout, 1769 (25); and Rattus rattus Linnaeus, 1758 (26) – Muridae (see abundance of rodents by species, SU and locality in Table 2).

A total of 8141 specimens of parasites, belonging to 29 taxa, were found (Table 3). Eleven parasite taxa were considered to be native species because they were originally described

parasitizing rodents native to the Neotropical region (Babero and Cattan, 1975; Babero *et al.* 1975; Quentin, 1975; Sutton, 1989; Robles *et al.* 2006; Notarnicola and Navone, 2011; Digiani *et al.* 2017). In addition, we determined 10 helminth taxa to genus or family level. We considered these taxa, with the exception of *Pterygodermatites* (*Paucipectines*) sp. 1 and *Capillaria* sens. lat. sp., to be native species because they were found only in native rodents.

On the other hand, we considered eight species as introduced species because they had been recorded mainly in one of the introduced rodent species in Chile and elsewhere (Harkema, 1936; Tena et al. 1998; Pisanu et al. 2001; Marangi et al. 2003; Milazzo et al. 2003; Asakawa, 2005; Kataranovski et al. 2011) (Table 3). Given the lack of evidence, *Pterygodermatites* (*Paucipectines*) sp. 1 and *Capillaria* sens. lat. sp. were classified neither as native nor as introduced and were not included in the analyses.

The following parasite taxa were found in SUs that were inhabited by both native and introduced rodents: Anatrichosoma sp.; Andrya octodonensis Babero and Cattan, 1975; Anoplocephalidae sp.; Aspiculuris tetraptera Nitzsch, 1821; Capillaria sens. lat. sp.; Graphidioides taglei Babero and Cattan, 1975; Heligmonellidae sp. 2.; Helminthoxys gigantea Quentin, Courtin and Fontecilla, 1975; Heterakis spumosa Schneider, 1866; Heteroxinema chilensis Quentin, 1975; Hydatigera taeniformis Batsch, 1786; Hymenolepis cf. diminuta Rudolphi, 1819; Litomosoides pardinasi Notarnicola and Navone, 2011; Longistriata degusi Babero and Cattan, 1975; Nippostrongylus brasiliensis Travassos, 1914; Physaloptera calnuensis Sutton, 1989; Pterygodermatites (Paucipectines) sp. 1; Pterygodermatites (Paucipectines) sp. 2; Hymenolepis (syn. Rodentolepis sensu Khalil et al. 1994) sp.; Syphacia muris Yamaguti, 1941; Syphacia obvelata Rudolphi, 1802; Syphacia sp.; cf. Trichuris pardinasi Robles, Navone and Notarnicola, 2006; and Trichuris muris Schrank, 1788. These taxa, with the exception

Table 2. Abundance of rodents by species, study unit and locality of Central Chile

| | | | | | | | Roo | dents* | | | | |
|------------|----------------------------------|----------------|---------|---------|-------|-------|--------|----------|-------|---------|-------|-------|
| | | | | | | | | Myomorph | a | | | |
| | | | Hystric | omorpha | | Crico | etidae | | | Muridae | | |
| Study unit | Locality | Date (mm-yyyy) | A. b. | 0. d. | A. l. | А. о. | O. l. | P. d. | М. т. | R. n. | R. r. | Total |
| 1 | Universidad de Chile | May 2002 | | | | 13 | | | 27 | 6 | | 46 |
| 2 | El Bosque Farm | July 2002 | | | | 9 | | | 13 | 12 | 3 | 37 |
| 3 | Lo Aguirre | August 2002 | | 6 | | 4 | | 26 | | | | 36 |
| 4 | Universidad de Chile | September 2002 | | | | 38 | 1 | | 9 | | 1 | 49 |
| 5 | El Bosque Farm | October 2002 | | | | 8 | | | 16 | | 3 | 27 |
| 6 | El Roto Chileno | February 2008 | | 3 | | | | 1 | | 1 | | 5 |
| 7 | Las Chinchillas National Reserve | May 2008 | | 11 | 1 | 1 | | 13 | | | | 26 |
| 8 | Universidad de Chile | January 2009 | | | | 45 | | | 18 | | 1 | 64 |
| 9 | El Roto Chileno | April 2009 | 1 | 5 | | | | 3 | | 1 | 8 | 18 |
| 10 | Sahonde | May 2009 | 2 | 4 | | | | 4 | | 3 | 3 | 16 |
| 11 | San Ramón | March 2008 | | | | | | | | | 5 | 5 |
| 12 | Rinconada de Maipú-A | November 2010 | | | | 1 | 1 | | 1 | | 1 | 4 |
| 13 | Rinconada de Maipú-B | November 2010 | | | | | | 2 | | | | 2 |
| 14 | La Batalla-A | March 2011 | | | | | 1 | | | 1 | | 2 |
| 15 | Rinconada de Maipú-A | August 2011 | | | | | 7 | | | 1 | 1 | 9 |
| 16 | La Batalla-B | December 2010 | 2 | 2 | 3 | | | | | | | 7 |

^{*}Abbreviation of the host species names: A. b., Abrocoma bennetti; A. l., Abrothrix longipilis; A. o., Abrothrix olivaceus; M. m., Mus musculus; O. d., Octodon degus; O. l., Oligoryzomys longicaudatus; P. d., Phyllotis darwini; R. n., Rattus norvegicus; R. r., Rattus rattus.

Table 3. Origin (native, introduced), hosts and parasitic loads of helminths found in rodents of Central Chile

| the polition of the politic | | | | | | my child | | | | | |
|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------|-----------------------|------------|--------------------------------------------|-------------------------|----------------------|--------|------------------|--------------------------------|-------------------|----------------------------------------------------|
| Direct Native Ab, Odd Pal' Ao Ao Ao Ao Ao Ao Ao A | Helminth | lype of life cycle | Origin | Hystricomorpha Non-Cricetidae ^a | Cricetidae ^a | Muridae ^a | number | (n) ^b | Mean abundance ^c | Mean intensity | Proportion of SU with transmission ^d |
| Indirect Native Oct Ab., Oct Ac. | cf. Anatrichosoma sp. ^e | Direct | Native | | A.o. | | 87 | 3.4 (119) | 1.7 | 21.8 | |
| Indirect Native O.d. P.d.¹ 94 513 (85) 0.5 Direct Native O.d. P.d.¹ P.d.¹ 94 5.5 (31) 15.7 2.5 Direct Native O.d. P.d. P.d.¹ P.d. P.d. | Anoplocephalidae sp. | Indirect | Native | A.b., O.d. | | | 4 | 40 (5) | 0.4 | 2.0 | |
| Direct Native Od. Od. 152 127 127 127 127 127 127 127 127 127 12 | Andrya octodonensis | Indirect | Native | 0.4. | P.d. ^f | | 34 | 31.3 (80) | 0.5 | 1.4 | |
| Direct Native O.d. A.b. A.b | Graphidioides taglei | Direct | Native | O.d. | | | 47 | 6.5 (31) | 15.7 | 23.5 | |
| Direct Native Ab. Ab. Ac. Ac. | Pudica cattani | Direct | Native | 0.0. | | | 125 | 3.2 (31) | 12.5 | 125.0 | |
| Direct | Heligmonellidae sp. | Unknown | Native | A.b. | | | 33 | 40 (5) | 16.5 | 16.5 | |
| Direct Native O d O d. O d. | Helmithoxys gigantea | Direct | Native | 0.d. | | | 272 | 54.8 (31) | 10.9 | 16.0 | |
| blict Native OLI, P.d. Rr. 548 129 (85) 559 4 bliect Native O.d. A.O., O.L.* M.m., R.n.*, R.n.* 447 265 (31) 1.1 cetines) 9.2 Indirect Native O.d.* A.O., O.L.* M.m., R.n.*, R.n.* 447 20.5 (324) 3.4 3 Indirect Native O.d.* A.O., O.L.* M.m., R.n.*, R.n.* 447 20.5 (324) 3.4 3 Indirect Native A.O. A.O. 1.0 4.1 (49) 0.3 3 Indirect Native A.O. A.O. A.O. 4.0 4.1 (49) 0.5 4 Native A.O. A.O. A.O. A.O. 4.1 (49) 0.5 1.0 5 Indirect Native A.O. A.O. A.O. A.O. A.O. A.O. 4.1 (49) 0.5 1.0 0.5 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 <td< td=""><td>Heteroxinema chilensis</td><td>Direct</td><td>Native</td><td>O d.</td><td></td><td></td><td>49</td><td>3.3 (31)</td><td>1.6</td><td>4.9</td><td></td></td<> | Heteroxinema chilensis | Direct | Native | O d. | | | 49 | 3.3 (31) | 1.6 | 4.9 | |
| birect Native O.d. A.L. M.m., R.n.', R.r.' 447 265 (34) 3.4 octines) Sp. 2 Indirect Native O.d.' A.L. 1 26 (4) 3.4 3 Indirect Native O.d. A.L. 1 25 (4) 0.3 3 Indirect Native A.O., O.L. A.O., O.L. 1 25 (4) 0.3 4 Indirect Native A.O., O.L. A.O., O.L. 1 25 (129) 1.2 3 Indirect Native A.O. A.O. 4.O. 9 4.1 (49) 0.6 4 Native A.O. A.O. A.O. A.O. 1.29 2.5 (129) 0.5 5 Intect Native A.O. | Litomosoides pardinasi | Indirect | Native | | 0.l., P.d. | R.r. | 548 | 12.9 (85) | 55.9 | 49.8 | 2/3 |
| profilect Native Od,¹ A.o., O.I.¹ M.m., R.n.¹, R.r.¹ 447 205 (564) 344 actines) Sp. 2 Indirect Native A.o., O.I. 129 225 (129) 1.2 3 Indirect Native Ao., O.I. 129 22.5 (129) 1.2 3 Indirect Native Ao. Ao. 129 37,119 6.6 1.2 3 Indirect Native Ao. Ao. 264 4.1, (49) 0.6 1.2 4 Direct Native Ao. Ao. Ao. 147 (68) 4.6 1.0 5 Direct Native Ao. Ao. Ao. 14.7 (68) 9.8 1.0 6 Direct Native Ao. M.m., R.r. 18 15.4 (13) 0.8 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1 | Pudica degusi | Direct | Native | 0.0. | | | 10 | 6.5 (31) | 1.1 | 5.0 | |
| p Al. Al. 1 25 (4) 0.3 setines) Sp. 2 Indirect Native Ao, O.I. 129 22.5 (129) 1.2 3 Indirect Native Ao, O.I. 40 619 41 (49) 0.6 1 Indirect Native Al, O.I. P.d. A. 264 147 (68) 0.6 1 Direct Native Al, O.I. P.d. A. 14 16.1 (31) 0.6 1 Direct Native O.d. A. A. 14 16.1 (31) 0.8 1 Direct Native O.d. A. A. 1.4 1.0 0.8 1 Direct Native A. A. A. A. 1.4 1.0 0.8 2 Direct Introduced A. A. A. A. 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 | Physaloptera calnuensis | Indirect | Native | 0.d.f | A.o., O.l. ^f | R.n. ^f , | 447 | 20.5 (264) | 3.4 | 8.3 | 2/2 |
| sectines) Sp. 2 Indirect Native Ao, O.I. 129 2.5. (129) 1.2 3 Indirect Native Ao, O.I. P.d. 619 4.1 (49) 0.6 3 Indirect Native Ao, O.I. P.d. 264 147 (68) 4.6 1.0 5 Direct Native Ao, O.I. P.d. 264 147 (68) 4.6 1.0 6 Direct Native Ao, O.I. P.d. AI, O.I. P.d. 315 (54) 10 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 <td< td=""><td>Protospirura cf. numidicola</td><td>Indirect</td><td>Native</td><td></td><td>A.I.</td><td></td><td>1</td><td>25 (4)</td><td>0.3</td><td>1.0</td><td></td></td<> | Protospirura cf. numidicola | Indirect | Native | | A.I. | | 1 | 25 (4) | 0.3 | 1.0 | |
| 3 Indirect Native P.d. P.d. 9 4.1 (49) 0.6 Indirect Native A.o. 619 37(119) 6.2 1 Direct Native A.l., O.l., P.d. 264 147 (68) 4.6 1 Direct Native O.d. A.l., O.l., P.d. 114 16.1 (31) 0.8 1 Direct Native O.d. A.g. A.g. 114 16.1 (31) 0.8 1 Direct Native O.d. A.g. A.g. 1.0 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.8 1 0.9 1 0.9 1 0.9 1 0.9 1 0.0 0.9 0.9 0.9 0.0 | Pterygodermatites (Paucipectines) sp. 2 | Indirect | Native | | | | 129 | 22.5 (129) | 1.2 | 4.4 | |
| Indirect Native Ao. 619 37(119) 62 1 Direct Native Al., Ol, P.d. 264 147 (68) 62 3 Direct Native P.d. Al., Ol, P.d. 311 31.5 (54) 10 1 Direct Native O.d. P.d. A.A. 31 13.5 (54) 10 1 Direct Native P.d. A.A. A.A. 33 37 (54) 9.8 1 Direct Introduced A.A. M.m., R.r. 18 4.5 (110) 0.4 1 12.2 5 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | Pterygodermatites (P.) sp. 3 | Indirect | Native | | P.d. | | 6 | 4.1 (49) | 9.0 | 4.5 | |
| Direct Native Al, Ol, Pd Ab 41, Ol, Pd 46 47 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 46 47 46 46 47 47 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 <th< td=""><td>Hymenolepis sp.</td><td>Indirect</td><td>Native</td><td></td><td>A.o.</td><td></td><td>619</td><td>37(119)</td><td>6.2</td><td>14.1</td><td></td></th<> | Hymenolepis sp. | Indirect | Native | | A.o. | | 619 | 37(119) | 6.2 | 14.1 | |
| Direct Native P.d. P.d. 311 315 (54) 10 10 Direct Native Od. P.d. 14 16.1 (31) 0.8 10 Direct Native Od. Mm., R.r. 18 3.7 (54) 9.8 12 Mohrown Unknown A.o. Mm., R.r. 18 4.5 (110) 0.4 12 Siget Introduced A.o. Mm., R.r. 10 15.7 (51) 0.1 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 12 | Syphacia sp. | Direct | Native | | A.l., O.l, P.d. | | 264 | 14.7 (68) | 4.6 | 24.5 | |
| Direct Native O.d. P.d. 14 16.1 (31) 0.8 Direct Native P.d. M.m., R.r. 18 4.5 (110) 0.4 9.8 1 Unknown Unknown A.o. M.m., R.r. 220 7.8 (51) 12.2 5 Sist Introduced A.o. M.m., R.r. 10 15.7 (51) 0.1 1 is Direct Introduced R.r. 26 13.7 (51) 0.1 2 ectines) Sp. 1 Indirect Unknown P.d.* R.r., R.r. 26 13.7 (51) 0.1 2 loiect Introduced P.d.* R.r., R.r. 264 19 (100) 127.1 10 loiect Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 ectines) Sp. 1 Indirect Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 | Trichostrongylidae sp. | Direct | Native | | P.d. | | 311 | 31.5 (54) | 10 | 18.3 | |
| Direct Native P.d. 39 3.7 (54) 9.8 1 Direct Introduced M.m., R.r. 18 4.5 (110) 0.4 0.4 Direct Unknown Unknown A.o. M.m., R.r. 220 7.8 (51) 12.2 5 is Direct Introduced A.o. M.m., R.r. 10 15.7 (51) 0.1 2 ectines) Sp. 1 Indirect Unknown P.d.* R.r., R.r. 3 3.8 (26) 0.3 3 ectines) Sp. 1 Indirect Introduced P.d.* R.r., R.r. 2054 19 (100) 127.1 10 birect Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 Indirect Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 Indirect Introduced A.o. M.m. 1 4 (25) 1 1 | Trichuris bradleyi | Direct | Native | 0.d. | | | 14 | 16.1 (31) | 0.8 | 2.8 | |
| Direct Introduced M.m., R.r. 18 4.5 (110) 0.4 Unknown Unknown Rn, Rr. 220 7.8 (51) 12.2 5 Introduced A.o. M.m., R.r. 374 8.3 (254) 4.3 1 is Introduced Rn, R.r. 26 13.7 (51) 0.1 2 ectines) sp. 1 Indirect Unknown Rn, R.r. 3 3.8 (26) 0.3 3 birect Introduced Rn, R.r. 2054 19 (100) 127.1 10 birect Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 Indirect Introduced R. n. R. n. 1 4 (25) 1 1 | cf. Trichuris pardinasi | Direct | Native | | P.d. | | 39 | 3.7 (54) | 9.8 | 19.5 | |
| Unknown Unknown Unknown A.o. R.n., R.r. 220 7.8 (51) 12.2 5 Indirect Introduced A.o. M.m., R.n., R.r. 374 8.3 (254) 4.3 1 is Indirect Introduced R.n., R.r. 10 15.7 (51) 0.1 2 ectines) sp. 1 Indirect Unknown R.n., R.r. 3 3.8 (26) 0.3 1 Direct Introduced A.o. M.m. 2267 19 (100) 127.1 1 Indirect Introduced A.o. R.n. R.n. 1 4 (25) 1 | Aspiculuris tetraptera | Direct | Introduced | | | M.m., R.r. | 18 | 4.5 (110) | 0.4 | 3.6 | |
| Direct Introduced A.o. M.m., R.n., R.r. 374 8.3 (254) 4.3 1.3 is Indirect Introduced R.n., R.r. 10 15.7 (51) 0.1 0.1 sectines) sp. 1 Indirect Unknown R.r. 26 13.7 (51) 2 2 Pedf R.n., R.r. 3 3.8 (26) 0.3 1 10 Direct Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 Indirect Introduced R.n. R.n. 1 4 (25) 1 | Capillaria sens. lat. sp. | Unknown | Unknown | | | R.n., R.r. | 220 | 7.8 (51) | 12.2 | 55.0 | |
| is Direct Introduced Rn., Rr. 10 15.7 (51) 0.1 ectines) sp. 1 Introduced Rr. 26 13.7 (51) 2 Ectines) sp. 1 Indirect Unknown Rn., Rr. 3 3.8 (26) 0.3 Direct Introduced A.o. M.m. 2267 19 (100) 127.1 10 Indirect Introduced R. n. 1 4 (25) 1 5 | Heterakis spumosa | Direct | Introduced | | A.o. | M.m., R.n., R.r. | 374 | 8.3 (254) | 4.3 | 17.8 | 1/4 |
| is Direct Introduced R.r. 26 13.7 (51) 2 ectines) sp. 1 Indirect Unknown R.n., R.r. 3 3.8 (26) 0.3 P.d.* R.n., R.r. 2054 19 (100) 127.1 10 Direct Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 Indirect Introduced R.n. 1 4 (25) 1 1 | Hymenolepis cf. diminuta | Indirect | Introduced | | | R.n., R.r. | 10 | 15.7 (51) | 0.1 | 1.3 | |
| ectines) sp. 1 Indirect Unknown R.n., R.r. 3 3.8 (26) 0.3 Direct Introduced R.n., R.r. 2054 19 (100) 127.1 10 Direct Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 Indirect Introduced R.n. 1 4 (25) 1 | Nippostrongylus brasiliensis | Direct | Introduced | | | R.r. | 26 | 13.7 (51) | 2 | 3.7 | |
| Direct Introduced P.d.* R.n., R.r. 2054 19 (100) 127.1 10 Direct Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 Indirect Introduced R. n. 1 4 (25) 1 1 | | Indirect | Unknown | | | R.n., R.r. | ю | 3.8 (26) | 0.3 | 3.0 | |
| Direct Introduced A.o. M.m. 2267 21.2 (203) 19.7 5 Indirect Introduced R. n. 1 4 (25) 1 | Syphacia muris | Direct | Introduced | | P.d. ^f | R.n., R.r. | 2054 | 19 (100) | 127.1 | 108.1 | 1/2 |
| Indirect Introduced | Syphacia obvelata | Direct | Introduced | | A.o. | M.m. | 2267 | 21.2 (203) | 19.7 | 52.7 | 9/9 |
| | cf. Hydatigera taeniformis | Indirect | Introduced | | | R. n. | 1 | | 1 | 1.0 | |
| Trichuris muris ^e Direct Introduced M.m. 123 7.1 (84) 4.6 20.5 | Trichuris muris ^e | Direct | Introduced | | | M.m. | 123 | 7.1 (84) | 4.6 | 20.5 | |

of *Pterygodermatites* (*Paucipectines*) sp. 1 and *Capillaria* sens. lat. sp., were those used to analyse the effect of the relatedness of hosts in the sharing of parasites. Native and introduced hosts shared only five parasite species (prevalences in the source host community are given in parentheses): *L. pardinasi* (90%), *P. calnuensis* (27%), *H. spumosa* (13%), *S. muris* (31%) and *S. obvelata* (49%). These five species were shared between cricetid and introduced rodents. Only one species was shared between non-cricetid and introduced rodents, *P. calnuensis*, which was also shared with cricetid rodents.

The relatedness of the hosts

In SUs that harboured both native and introduced rodents, the proportion of parasite species that introduced rodents shared with native rodents was higher when native rodents were cricetids (5/16) than when they were non-cricetids (1/17); taken as a whole, this difference was non-significant (Fisher one-tail test, P = 0.074). However, this difference was significant after excluding the most generalist species (P. calnuensis) (P = 0.043).

The parasitic loads in source hosts

We studied the native parasite species *P. calnuensis* and the introduced species *S. obvelata* to assess the association of the prevalence and mean abundance of parasites in source host populations with the presence and abundance in the recipient host populations in the same SU, because these were the species shared in the largest number of SUs: *S. obvelata* was shared in SUs 1, 2, 4, 5 and 8; *P. calnuensis* was shared in SUs 1, 2, 4, 5 and 15. The source host species of *P. calnuensis* was *A. olivaceus* and the recipient species were *M. musculus*, *R. rattus* and *R. norvegicus*. The source host species of *S. obvelata* was *M. musculus* and the recipient species was *A. olivaceus*.

Neither mean abundance nor prevalence of *P. calnuensis* in source host populations showed significant association with its abundance in the recipient hosts (mean abundance: coefficient = 0.09, standard error (s.e.) = 0.1, Z = 0.83, P = 0.41; prevalence: coefficient = 1.83, s.e. = 4.43, Z = 0.41, P = 0.68) nor with its presence in the recipient hosts [mean abundance: odds ratio (OR) = 1.09, s.e. = 0.07, Z = 1.38, P = 0.167; prevalence: OR = 1.86, s.e. = 4.2, Z = 0.28, P = 0.78]. Similarly, the mean abundance and the prevalence of *S. obvelata* in source host populations also did not show significant association with its abundance in the recipient hosts (mean abundance: coefficient = 0.02, s.e. = 0.03, Z = 0.76, P = 0.45; prevalence: coefficient = 2.58, s.e. = 2.89, Z = 0.89, Z = 0.89,

Considering all parasite species of myomorph rodents, the mean abundance of parasites among all source host populations was not significantly associated with higher odds of being shared with a recipient host (OR = 1.19; confidence interval = 0.959, 1.494; P = 0.11). However, shared parasites had higher mean abundances in their source populations than not shared parasites (Z = 2.314, P = 0.02). The analogous analyses (logistic regression and Wilcoxon test) for the prevalence did not show significant associations with being shared (P > 0.23 in both cases). On the other hand, the number of localities in which the parasite was present was significantly associated with the odds of being shared (OR = 3.00; confidence interval = 1.207, 7.474; P = 0.018, respectively).

Discussion

The proportion of shared species between murid and cricetid rodents was higher than with non-cricetid rodents. Although

this difference was not clearly significant, it shows an agreement with what was expected (Klimpel et al. 2007; Wells et al. 2015). The only species shared between non-cricetid and introduced rodents was a generalist nematode, P. calnuensis. In addition, P. calnuensis was also found in cricetid rodents, demonstrating low host specificity. The other less generalist species were not transmitted between suborders, and, considering only these, there is a significant association of the relatedness of the hosts with the sharing of parasites, suggesting that the relatedness of hosts, combined with the level of specialization of parasites are two forces that drive the sharing of parasites. This result complements that of MacLeod et al. (2010), who found that the host specificity of the parasite affects the persistence of the parasite after arrival. Our results are also consistent with the previous studies (Pisanu et al. 2009; Wells et al. 2015) and reinforce quantitatively the importance of the relatedness of the hosts in the sharing of parasites between native and introduced hosts. Thus, our results enable hypotheses regarding which native species will more likely share parasites with a particular invasive species, and which parasites will more likely be shared.

Other attributes of the life history of the parasites must be studied in order to assess how they interact with the phylogenetic distance. For instance, the existence of unknown ecto-parasites as vectors (e.g. *L. pardinasi*) or unknown intermediate hosts present in the cycle of some parasites (e.g. *P. calnuensis*) can affect the likelihood of parasite transmission. In this regard, it has been reported that the diet of the host and its position in the food web can also affect parasite acquisition (Locke *et al.* 2014).

In general terms, the parasitic loads of P. calnuensis and S. obvelata in source host populations did not show significant association with the presence and abundance in their recipient populations. Thus, the overall results suggest that the presence and the abundance of a parasite in a recipient host are not affected by the prevalence and mean abundance of this parasite in the source host population. This can be explained by the fact that parasites have been transmitted from sources to recipient host for hundreds of years - R. rattus and M. musculus from the 1600s, R. norvegicus from the 1800s (Jaksic, 1998) - in such a way that the transmission dynamics, especially transmission between recipient hosts, could make the importance of the parasitic loads in the source hosts irrelevant. In other words, our results suggest that S. obvelata and P. calnuensis did not seem to require the source populations to persist in the recipient hosts. This is similar to what was observed in the California Channel Islands, where T. muris persists in Peromyscus maniculatus Wagner (1845) even on islands where R. rattus was eradicated (Smith and Carpenter, 2006). Future studies must be performed to confirm this hypothesis. Our results can also be interpreted to mean that other factors not considered in this study associated to the susceptibility of the recipient hosts can also affect their infection rates. More studies are necessary to control for these factors.

On the other hand, the higher mean abundance of shared parasites among source hosts than that of not shared parasites can be considered the first evidence in support of the hypothesis that the number of parasites in the source host population within a large territory could favour the sharing of parasites between native and introduced hosts in at least one locality. Thus, our results suggest the importance of the abundance but not the prevalence of parasites in the source populations. This is consistent with the hypothesis that, given the aggregated dispersion of parasites (i.e. most parasites colonize few hosts), those few hosts with a high number of parasites, and not all infected hosts, are responsible for most parasite transmission (Woolhouse *et al.* 1997). In addition, the number of localities in which the parasites are present is also associated with being shared with a recipient

host, which enhances the likelihood of contact with recipient host in at least one locality.

For free-living organisms, the propagule pressure is one of the most important factors in the success of the invasion process (Lockwood et al. 2005). However, in the context of parasitism, propagule pressure, understood as the number of parasites that arrived with a host, is not a major factor in the persistence of parasites in the introduced host population (MacLeod et al. 2010). Hatcher et al. (2012) mentioned that the spillover propagule pressure includes two parts, the propagule size and the number of propagules, i.e. the number of spillover events. This latter is very difficult to assess in the context of parasitism due to the possible transmission events of a parasite without successful persistence among recipient hosts, which therefore makes it difficult to prove. Thus, an acceptable equivalence for the number of propagules could be the territorial dispersion and abundance of parasites, which could be associated with the chances for a parasite to contact a recipient host and achieve a spillover event. In this context and consequent with our results, it is possible that the larger the abundance of a parasite species among its source hosts in a large territory and the larger the number of localities in which the parasite is present, the higher the probability of the parasite being shared in at least one locality. Thus, if the goal of a management is the prevention of the spillover of parasites from invasive hosts, the control of the dispersion of this invasive host is a core aim.

Concluding remarks

The foregoing allows us to suggest that the relatedness of the hosts combined with the low host specificity of parasites favour the spillover of parasites between native and introduced hosts. The prevalence of parasites found in source host populations is not as significant as a factor in parasite spillover. On the contrary, the abundance and dispersion of parasites in source hosts may affect parasite sharing between native and introduced hosts. Thus, the association of the abundance and the dispersion of parasites with host switching are proposed as factors driving the sharing of parasites between native and introduced rodent hosts.

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Ethical standards. We have strictly conformed to relevant ethical standards, involving the use of the minimum number of animals necessary to produce statistically reproducible results. The Comité de Ética of the Facultad de Ciencias Veterinarias y Pecuarias (Ethics Committee of the Faculty of Veterinary and Animal Sciences) at the Universidad de Chile approved and certified the study (certificate without number, 15 April 2011), and the Servicio Agrícola y Ganadero (Agricultural and Livestock Service) of Chile authorized trapping (resolution certificates 2041 and 6652 to C.L.-A.).

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Conflict of interest. None.

References

- **Anderson RC, Chabaud AG and Willmott S** (2009) *Keys to the Nematode Parasites of Vertebrates.* CAB International, Wallingford.
- Asakawa M (2005) Perspectives of host-parasite relationships between rodents and nematodes in Japan. Mammal Study 30, S95–SS9.

Babero BB and Cattan PE (1975) Helmintofauna de Chile: III. Parasitos del roedor degu, Octodon degus Molina, 1782, con la descripcion de tres nuevas especies. Boletin Chileno de Parasitologia 30, 68–76.

- Babero BB, Cattan PE and Cabello C (1975) Trichuris bradleyi sp. n., a whipworm from Octodon degus in Chile. The Journal of Parasitology 61, 1061–1063
- Barrett JL, Carlisle MS and Prociv P (2002) Neuro-angiostrongylosis in wild black and grey-headed flying foxes (*Pteropus* spp). *Australian Veterinary Journal* 80, 554–558.
- Barton DP (1997) Introduced animals and their parasites: the cane toad, Bufo marinus, in Australia. Australian Journal of Ecology 22, 316–324.
- Bordes F, Blasdell K and Morand S (2015) Transmission ecology of rodentborne diseases: new frontiers. *Integrative Zoology* **10**, 424–435.
- Bush AO, Lafferty KD, Lotz JMShostak AW (1997) Parasitology meets ecology on its own terms: Margolis, et al. Revisited. The Journal of Parasitology 83, 575–583.
- Digiani MC, Landaeta-Aqueveque C, Serran PC and Notarnicola J (2017) Pudicinae (Nematoda: Heligmonellidae) parasitic in endemic Chilean rodents (Caviomorpha: Octodontidae and Abrocomidae): description of a new species and emended description of *Pudica degusi* (Babero and Cattan) n. comb. *The Journal of Parasitology* 103, 736–746.
- Grabner DS, Weigand AM, Leese F, Winking C, Hering D, Tollrian R and Sures B (2015) Invaders, natives and their enemies: distribution patterns of amphipods and their microsporidian parasites in the Ruhr Metropolis, Germany. Parasites & Vectors 8, 1–15.
- Harkema R (1936) The parasites of some North Carolina rodents. Ecological Monographs 6, 151–232.
- Hatcher MJ, Dick JTA and Dunn AM (2012) Disease emergence and invasions. Functional Ecology 26, 1275–1287.
- Jaksic F (1998) Vertebrate invaders and their ecological impacts in Chile. Biodiversity & Conservation 7, 1427–1445.
- Johnson PTJ and Thieltges DW (2010) Diversity, decoys and the dilution effect: how ecological communities affect disease risk. The Journal of Experimental Biology 213, 961–970.
- Kataranovski M, Mirkov I, Belij S, Popov A, Petrović Z, Gačić Z and Kataranovski D (2011) Intestinal helminths infection of rats (*Ratus norvegicus*) in the Belgrade area (Serbia): the effect of sex, age and habitat. *Parasite: Journal de la Société Française de Parasitologie* 18, 189.
- Kelly DW, Paterson RA, Townsend CR, Poulin R and Tompkins DM (2009)
 Parasite spillback: a neglected concept in invasion ecology? *Ecology* 90, 2047–2056.
- **Khalil LF, Jones A and Bray TA** (1994) Keys to the Cestode Parasites of Vertebrates. Wallingford: CAB International.
- Klimpel S, Förster M and Schmahl G (2007) Parasites of two abundant sympatric rodent species in relation to host phylogeny and ecology. *Parasitology Research* 100, 867–875.
- Landaeta-Aqueveque C, Robles MDR and Cattan PE (2007a) The community of gastrointestinal helminths in the housemouse, *Mus musculus*, in Santiago, Chile. *Parasitología Latinoamericana* **62**, 165–169.
- Landaeta-Aqueveque C, Robles MDR and Cattan PE (2007b) Helmintofauna del roedor Abrothrix olivaceus (Sigmodontinae) en áreas sub-urbanas de Santiago de Chile. Parasitología Latinoamericana 62, 134–141.
- Landaeta-Aqueveque C, Henríquez A and Cattan PE (2014) Introduced species: domestic mammals are more significant transmitters of parasites to native mammals than are feral mammals. *International Journal for Parasitology* 44, 243–249.
- Locke S, Marcogliese D and Tellervo Valtonen E (2014) Vulnerability and diet breadth predict larval and adult parasite diversity in fish of the Bothnian Bay. *Oecologia* 174, 253–262.
- Lockwood JL, Cassey P and Blackburn T (2005) The role of propagule pressure in explaining species invasions. *Trends in Ecology & Evolution* 20, 223–228.
- Loxton KC, Lawton C, Stafford P and Holland CV (2017) Parasite dynamics in an invaded ecosystem: helminth communities of native wood mice are impacted by the invasive bank vole. *Parasitology* 144, 1476–1489.
- MacLeod CJ, Paterson AM, Tompkins DM and Duncan RP (2010) Parasites lost do invaders miss the boat or drown on arrival? *Ecology Letters* 13, 516–527.
- Macneil C, Fielding NJ, Dick JT, Briffa M, Prenter J, Hatcher MJ and Dunn AM (2003) An acanthocephalan parasite mediates intraguild predation between invasive and native freshwater amphipods (Crustacea). *Freshwater Biology* **48**, 2085–2093.

- Marangi M, Zechini B, Fileti A, Quaranta G and Aceti A (2003) Hymenolepis diminuta infection in a child living in the urban area of Rome, Italy. Journal of Clinical Microbiology 41, 3994–3995.
- Mastitsky S and Veres J (2010) Field evidence for a parasite spillback caused by exotic mollusc *Dreissena polymorpha* in an invaded lake. *Parasitology Research* 106, 667–675.
- Milazzo C, de Bellocq JG, Cagnin M, Casanova JC, di Bella C, Feliu C and Santalla F (2003) Helminths and ectoparasites of *Rattus rattus* and *Mus musculus* from Sicily, Italy. *Comparative Parasitology* **70**, 199–204.
- Morand S, Bordes F, CHEN HW, Claude J, Cosson JF, Galan M, Czirjak GA, Greenwood AD, Latinne A, Michaux J and Ribas A (2015) Global parasite and Rattus rodent invasions: the consequences for rodent-borne diseases. Integrative Zoology 10, 409–423.
- Muñoz-Pedreros A. (2009). Orden rodentia. In Muñoz-Pedreros A., Yáñez J. (eds). Mamíferos de Chile, Valdivia: Editorial CEA, pp. 93–157.
- Notarnicola J and Navone G (2011) Litomosoides pardinasi n. sp. (Nematoda, Onchocercidae) from two species of cricetid rodents in northern Patagonia, Argentina. Parasitology Research 108, 187–194.
- Paterson RA, Townsend CR, Poulin R and Tompkins DM (2011) Introduced brown trout alternative acanthocephalan infections in native fish. *Journal of Animal Ecology* 80, 990–998.
- Pisanu B, Chapuis JL and Durette-Desset MC (2001) Helminths from introduced small mammals on Kerguelen, Crozet, and Amsterdam Islands (Southern Indian Ocean). The Journal of Parasitology 87, 1205– 1208
- Pisanu B, Lebailleux L and Chapuis JL (2009) Why do Siberian chipmunks Tamias sibiricus (Sciuridae) introduced in French forests acquired so few intestinal helminth species from native sympatric Murids? Parasitology Research 104, 709–714.
- Quentin JC (1975) Oxyure de Rongeurs: II. Essai de classification des oxyures Heteroxynematidae. Memoires Du Museum National D'histoire Naturelle, Zoologie Serie A 94, 51–96.
- Robles MDR, Navone GT and Notarnicola J (2006) A new species of *Trichuris* (Nematoda: Trichuridae) from Phyllotini rodents in Argentina. *Journal of Parasitology* **92**, 100–104.

- Romeo C, Ferrari N, Lanfranchi P, Saino N, Santicchia F, Martinoli A and Wauters LA (2015) Biodiversity threats from outside to inside: effects of alien grey squirrel (*Sciurus carolinensis*) on helminth community of native red squirrel (*Sciurus vulgaris*). Parasitology Research 114, 2621–2628.
- Smith KF and Carpenter SM (2006) Potential spread of introduced black rat (Rattus rattus) parasites to endemic deer mice (Peromyscus maniculatus) on the California Channel Islands. Diversity and Distributions 12, 742–748.
- Sutton C (1989) Contribution to the knowledge of Argentina's parasitological fauna XVII. Spirurida (Nematoda) from Neotropical Cricetidae: Physaloptera calnuensis n. sp. and Protospirura numidica criceticola Quentin, Karimi and Rodríguez De Almeida. Bulletin Du Museum National D'histoire Naturelle, Paris, 4° Série 11, 61–67.
- Taraschewski H (2006) Hosts and parasites as aliens. *Journal of Helminthology* **80**, 99–128.
- Tena D, Simón MP, Gimeno C, Pomata MTP, Illescas S, Amondarain I, González A, Domínguez J and Bisquert J (1998) Human infection with Hymenolepis diminuta: case report from Spain. Journal of Clinical Microbiology 36, 2375–2376.
- **Tompkins DM, White AR and Boots M** (2003) Ecological replacement of native red squirrels by invasive greys driven by disease. *Ecology Letters* **6**, 189–196.
- Wells K, O'Hara RB, Morand S, Lessard JP and Ribas A (2015) The importance of parasite geography and spillover effects for global patterns of host-parasite associations in two invasive species. *Diversity and Distributions* 21, 477–486
- Wilcove DS and Master LL (2005) How many endangered species are there in the United States? Frontiers in Ecology and the Environment 3, 414–420.
- Woolhouse ME, Dye C, Etard JF, Smith T, Charlwood JD, Garnett GP, Hagan P, Hii JLK, Ndhlovu PD, Quinnell RJ and Watts CH (1997) Heterogeneities in the transmission of infectious agents: implications for the design of control programs. *Proceedings of the National Academy of Sciences* 94, 338–342.
- Young H. S., Parker I. M., Gilbert G. S., Sofia Guerra A. and Nunn C. L. (2017). Introduced species, disease ecology, and biodiversity-disease relationships. *Trends in Ecology & Evolution* 32, 41–54.