

Parasites of the Brazilian flathead *Percophis brasiliensis* reflect West Atlantic biogeographic regions

PAOLA E. BRAICOVICH^{1*}, CAMILA PANTOJA², ALDENICE N. PEREIRA², JOSE L. LUQUE² and JUAN T. TIMI¹

¹ *Laboratorio de Ictioparasitología, Instituto de Investigaciones Marinas y Costeras (IIMyC), Facultad de Ciencias Exactas y Naturales, Universidad Nacional de Mar del Plata- Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET), Funes 3350, (7600) Mar del Plata, Argentina*

² *Curso de Pós-Graduação em Ciências Veterinárias. Departamento de Parasitologia Animal, Universidade Federal Rural do Rio de Janeiro, Caixa Postal 74-540, 23851-970, Seropédica, RJ, Brasil*

(Received 18 February 2016; revised 2 May 2016; accepted 4 May 2016; first published online 3 November 2016)

SUMMARY

With the aim of evaluating the utility of marine parasites as indicators of zoogeographical regions in the South West Atlantic, we analyzed data on assemblages of long-lived larval parasites of 488 specimens of *Percophis brasiliensis* distributed in 11 samples from nine localities covering the entire distribution of the species in the Argentine biogeographical Province. Near half a million long-lived parasite individuals belonging to 17 species present in the whole sample displayed clear latitudinal patterns. Data for parasite assemblages at infracommunity and component community levels were analysed in relation to the geographical distance. Significant similarity decay of parasite assemblages over distance was observed, with those based on abundances and mean abundances showing departures from predicted values of regressions. These departures were represented by higher dissimilarities between samples coming from different zoogeographical regions than between those caught within the same region, independently of the distance separating them. Consequently, zoogeographical regions were identified in a distance-decay context. Multivariate analyses corroborated a close fit of similarity between assemblages to existing zoogeographical classifications. Regressions representing distance decay of similarity, and the identification of their outliers, can therefore shed light on the existence of discontinuities or uniformities in the geographic distribution of parasite assemblages and, in turn, in the zoogeography of their fish hosts.

Key words: parasite assemblages, parasite tags, distance decay, zoogeography.

INTRODUCTION

The use of parasites as biological markers of fish stocks or populations, a methodology coined and developed in the field of fishery sciences, is ultimately based on the geographical distribution of parasites (Timi and MacKenzie, 2015). Indeed, fish can become infected with a parasite only when they come within the endemic area of that parasite, which is that geographic region where the conditions are suitable for a successful transmission and the completion of its life cycle (MacKenzie and Abaunza, 2013). These suitable conditions include environmental factors, mainly the temperature and salinity profiles, for marine organisms (Esch and Fernández, 1993), as well as the presence of all hosts involved in the parasite life cycle, which also should be at population densities large enough to sustain parasite populations and ensure transmission. A change in one of these factors, or of any combination of them along a geographic or environmental gradient, will produce a gradual shift of parasite

prevalence or abundance and, if several species are considered simultaneously, of parasite assemblage structure and composition. This means that, from an ecological perspective, the reliability of parasites as biological tags is underpinned by the almost universal decay in the similarity of the species composition of assemblages as a function of increasing distance between them (Poulin and Kamiya, 2015).

Although the relationship between population structure studies, in a fishery context, and similarity decay with increasing distance, in a zoogeographical context, have been acknowledged in some papers on parasite tags (Timi *et al.* 2010; Vales *et al.* 2011; Braicovich *et al.* 2012), these two aspects of parasite distribution have usually been addressed independently in the literature. This is probably due to the spatial scales considered by each kind of study. Research on parasite tags tends generally to discriminate between neighbouring fish populations, and very few studies have used parasites as indicators of large scale zoogeographical patterns (Rohde, 2002). On the other hand, research on distance decay of similarity has focused on large spatial scales and in the factors influencing its strength (Soininen *et al.* 2007), especially when applied to marine fish (Poulin, 2003; Oliva and González, 2005; Pérez-del-Olmo *et al.* 2009; Timi *et al.* 2010).

* Corresponding author: Laboratorio de Ictioparasitología, Instituto de Investigaciones Marinas y Costeras (IIMyC), FCEyN, Universidad Nacional de Mar del Plata-CONICET, Funes 3350, (7600) Mar del Plata, Argentina. E-mail: braicovi@mdp.edu.ar

Recently, based on studies on parasite tags over large spatial scales along south western Atlantic coasts and on their own data, Pereira *et al.* (2014) proposed that the scope of parasite tags studies can be expanded from local to regional scales as indicators of ecoregions and of higher levels of biogeographic classifications. Similarly, Cantatore and Timi (2015) proposed that shifting the focus from fish/parasite populations to communities can provide information to delineate ecosystem boundaries for host communities. Here we analyze the usefulness of parasites of the Brazilian flathead *Percophis brasiliensis* (Quoy and Gaimard, 1824) as indicators of zoogeographical regions. This demersal species inhabits coastal waters in the south-western Atlantic, at depths generally not exceeding the 50 m isobath (Milessi and Marí, 2012). Its distribution extends from 23°S (Rio de Janeiro, Brazil) to 47°S (north of Santa Cruz Province, Argentina) (Cousseau and Perrotta, 2004), a range over which the distribution patterns and endemisms of the biota have been utilized to describe a series of biogeographical units (realms, provinces, and ecosystems), although with different degrees of geographic resolution depending on the study (Spalding *et al.* 2007; Menni *et al.* 2010; Miloslavich *et al.* 2011; Briggs and Bowen, 2012).

The study area of the present work covers near 3300 km of the coastal region of South America, between Rio de Janeiro and the northern Argentine Sea, which shows a continuous distribution of warm temperate fauna and has historically been characterized as the Argentine Biogeographical Province. It was recently renamed as the Temperate Western South Atlantic Province (Menni *et al.* 2010) or Warm Temperate Southwestern Atlantic (Spalding *et al.* 2007) (Fig. 1).

Percophis brasiliensis in the northern Argentine Sea shows a clear latitudinal population structure as indicated by its parasites (Braicovich and Timi, 2008, 2010), as well as for other methodologies (Avigliano *et al.* 2015). In the present work, the distribution of parasites of *P. brasiliensis* is analysed in this region, where oceanographic conditions, dominated by strong gradients of temperature and salinity that determine fish parasite distributions (Cantatore and Timi, 2015), provide a suitable scenario for studies on similarity decay over distance, a pattern already observed for a sympatric fish species in the region (Timi *et al.* 2010). Similarity decay of parasite assemblages over distance is here analysed in relation to existing zoogeographical classifications derived from biological systems other than parasites.

The aim of this study is, therefore, to evaluate the utility of marine long-lived larval parasites as indicators of zoogeographical regions in the South West Atlantic, following the more recent zoogeographical classifications. We predict a similarity decay of parasite assemblages over distance, with departures from predicted values of similarity represented by more

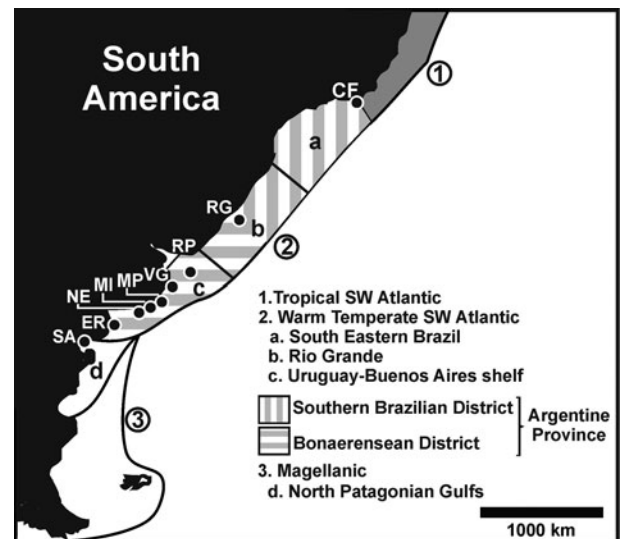


Fig. 1. Map showing the sampling localities in the Southwestern Atlantic coast and their zoogeographical schemes. Numbers (1–3) and letters (a, b, c) show the biogeographic provinces and ecoregions, respectively, according to Spalding *et al.* (2007). Striped area shows the districts of the Argentine province according to Menni *et al.* (2010). CF, Cabo Frio; ER, El Rincón; MP, Mar del Plata; MI, Miramar; NE, Necochea; RG, Rio Grande; RP, Río de la Plata; SA, San Antonio Oeste; VG, Villa Gessel.

pronounced dissimilarities between samples coming from different zoogeographical regions than those caught within the same region, independently of the distance separating them; therefore zoogeographical regions may be identifiable in a distance-decay context.

MATERIALS AND METHODS

Fish and parasites sampling

The dataset comprises the parasite communities of 11 samples of *P. brasiliensis* from nine localities in the south-western Atlantic, covering the entire distribution of the species in the Argentinean Zoogeographical Province (Menni *et al.* 2010) (Fig. 1) and including a total of 488 fish. Data from southern zones, in Argentine and Uruguayan waters, were obtained from previous studies, which aimed to use parasites as indicators for discriminating among fish populations (Braicovich and Timi, 2008, 2010); therefore, there are significant differences among some of them, whereas samples caught seasonally in the same locality were homogeneous in terms of parasite loads. Four new samples were included in the analyses, two from Brazilian waters and two from Argentine waters. The datasets are summarized in Table 1.

Fish were either kept fresh or deep frozen in plastic bags at -18°C until examination. After thawing, the total length was measured (TL, cm). Parasites sampling procedures were as indicated in Braicovich and Timi (2008, 2010).

Table 1. Composition of samples of *Percophis brasiliensis* in nine localities, ordered by latitude, of the South Western Atlantic

Locality	Locality code	Latitude, longitude	Date of capture	<i>n</i>	Total length (cm) \pm s.d.	Reference
Cabo Frio	CF	23°00'S, 41°30'W	06/2012	40	40.0 \pm (36.5–45)	Present study
Rio Grande	RG	32°40'S, 51°00'W	02/2013	47	48.1 \pm (39.5–64.5)	Present study
Río de la Plata	RP	35°30'S, 54°30'W	10/2005	35	49.4 \pm (31–59)	Braicovich and Timi (2008)
Villa Gessel	VG	37°50'S, 56°15'W	04/2010	51	55.3 \pm (49–70)	Present study
Mar del Plata	MP1	38°20'S, 56°40'W	07/2005	49	55.2 \pm (42–73)	Braicovich and Timi (2010)
Mar del Plata	MP2	38°20'S, 56°40'W	10/2005	48	57.3 \pm (44–71)	Braicovich and Timi (2010)
Mar del Plata	MP3	38°20'S, 56°40'W	02/2006	50	55.3 \pm (46–67)	Braicovich and Timi (2010)
Miramar	MI	38°45'S, 57°10'W	04/2007	40	53.7 \pm (45–65)	Present study
Necochea	NE	39°20'S, 58°45'W	05/2006	45	56.4 \pm (49–64)	Braicovich and Timi (2010)
El Rincón	ER	40°00'S, 61°20'W	10/2005	51	49.5 \pm (44–58)	Braicovich and Timi (2008)
San Antonio Oeste	SA	42°00'S, 64°00'W	01/2005	32	52.9 \pm (41–73)	Braicovich and Timi (2008)

According to their geographic location, samples were assigned to different biogeographic districts of the Argentine Province: those from Cabo Frio (CF) belong to the Southern Brazilian district, although located in the northern boundary with the Brazilian Province. All samples from the Argentine Sea belong to the Bonaerensean district, with the southern one coming from an ecotonal zone in the limit with the Magellanic Province. The sample from Rio Grande do Sul corresponds to the boundary between the Bonaerensean and Southern Brazilian districts (Menni *et al.* 2010) or to the ecoregion named Rio Grande (RG) by Spalding *et al.* (2007).

Population and community descriptors

Prevalence and mean abundance for each parasite species in each locality, as well as species richness at infracommunity level were calculated following Bush *et al.* (1997). Species richness also was calculated after excluding short-lived parasites (i.e. larval stages living in the gut, adult endoparasites and ectoparasites), with long-lived species being represented by larval species found in tissues or body cavity. Long-lived parasites were selected for all subsequent analyses because samples were caught at different seasons and years and these parasite species, that persist for long periods or even for the entire life of hosts, are more reliable for proper comparisons, diminishing possible effects of seasons or other short-term temporal variations on parasite loads and allowing the locality effects to be more evident.

Similarity analysis

Similarity in parasite species composition was computed at component community and infracommunity levels. Both the Jaccard and Bray-Curtis similarity indices (qualitative and quantitative, respectively) (Magurran, 1988) were calculated

between all possible pairs of hosts (infracommunities) from different zones, and those between each pair of localities were averaged. At the level of component communities, Bray-Curtis similarity indices were calculated on values of both prevalence and mean abundance between all possible pairs of samples.

Distance between pairs of host populations was calculated as the shortest linear distance between sampling localities obtained from a map, following the coast line below the 50 m isobath. For samples caught during research cruises and comprising several trawls, a central position was selected as locality; in the case of commercial catches, with no precise information on the capture sites, localities were selected as the point on the 50 m isobath closer to the landing port. Each parasite assemblage was used in more than one pairwise comparison, and consequently was not truly independent from distance in a statistical sense; furthermore, the most recent sample (Cabo Frio) was also the most distant from southern ones and it was composed of the smallest fishes. Therefore a possible relationship between distance and both date of capture and host size could result in an artefact due to these factors. For these reasons, the associations between similarity and geographic distance were tested using Partial Mantel tests implemented in the R package Ecodist package (Goslee and Urban, 2007). Separate models, including temporal and fish size differences as covariates, were constructed, each using 10 000 permutations of the dataset, this was because preliminary Mantel tests showed no relationship between temporal differences in captures and differences in fish size ($r = 0.13$, $P > 0.05$), however both variables were significantly related to geographic distance, with a higher correlation coefficient for size differences ($r = 0.77$, $P < 0.01$) than for temporal ones ($r = 0.51$, $P = 0.03$). For temporal variability, the number of months from the first sampling date was counted, whereas for differences in fish size between samples, Euclidean distances were

calculated between sizes of all possible pairs of hosts from different samples and then data were averaged.

The decay in similarity of parasite communities as a function of geographic distances was assessed in all cases by plotting log-transformed similarity values against untransformed distances (Poulin, 2003) between all possible pairs of localities. Since the values of Jaccard and Bray-Curtis are different, similarity was converted to relative similarity, by expressing each value as a percentage of the maximum of all values obtained in each analysis. This way, the *y*-axis on all plots had the same scale, which allowed comparisons of decay relationships among dissimilarity indices as well as between infracommunities and component communities.

Departures from predicted regressions on distance decay of similarity are expected between parasite assemblages for two opposite reasons: (1) samples from different zoogeographical regions, especially those from close ones should be more different than predicted by the distance separating them and (2) samples from the same zoogeographical region, especially distant ones, should be more similar than predicted by the distance separating them. As a measurement of these departures, the residuals of each point (the difference between the observed value of the dependent variable and the predicted value) were calculated. All those observed values with residuals larger than 10 (an arbitrarily defined threshold, representing 10% of possible variation of similarity, which was converted to relative similarity, with a maximum 100%) were identified individually.

To corroborate the effectiveness of departures from expected values in the regressions as indicators of unexpected high similarity or dissimilarity in relation to the distance separating samples, nonmetric multidimensional scaling analyses (MDS) (Clarke and Gorley, 2006) were performed to visualize the geographic patterns in the composition of parasite assemblages for the four datasets. In all cases, the fit of the MDS ordinations was quantified by a value of stress. Hierarchical agglomerative clustering of samples were applied to both averaged infracommunity similarities and component community similarities using group-average linking and the resemblance levels were overlaid on the MDS plots (Clarke and Gorley, 2006).

All similarity and distance measures, as well as MDS and clustering analyses were implemented in PRIMER V6 and PERMANOVA+ for PRIMER package (Anderson *et al.* 2008).

RESULTS

General results

The parasite fauna of the four new samples of *P. brasiliensis* comprised 30 parasite species (Table 2),

whereas a total species richness of 33 was recorded considering all the samples. A total of 471 144 metazoan parasites were counted, with a high proportion of them being represented by long-lived larval stages (439 211 specimens) belonging to 17 species; a smaller proportion, 30 455 individuals, were members of a short lived species complex, namely *Scolex polymorphus*, living in the gut; finally comparatively minor numbers of adult endoparasites (1344 specimens, 11 species) and ectoparasites (134 specimens, 4 species) were registered. These proportions were quite constant across samples, with the exception of flatheads from CF, where short-lived larval stages predominated numerically (Fig. 2).

Parasite burdens, in terms of prevalence and mean abundance, varied among the component communities from the considered zones as exemplified by the most prevalent and abundant long-lived species (Table 2), with several species reaching higher loads in the northern Argentine waters, but other increasing either southwards (i.e. *A. simplex s.l.*) or northwards (i.e. *Grillotia (Christianella) carvajalregororum* and *Hysterothylacium* sp.) mainly in terms of abundance; fish from CF showed generally lower values of prevalence and abundance (Fig. 3).

This spatial distribution of parasite species resulted in a bell-shaped distribution of species richness along the latitudinal range (Fig. 4a), a pattern also identifiable for long-lived species, although with more stable values across samples in the northern region of the Argentine Sea. The mean total abundance, irrespective of parasite species, showed a similar pattern with the exception of notably high values in RG (Fig. 4b). In this locality exceptionally high abundances were recorded for *G. (C.) carvajalregororum* and *Hysterothylacium* sp., for which maximum values of 16 604 and 3054 larvae were harboured respectively, by a single host.

Similarity analyses

Significant decays in similarity of parasite assemblages over distance were recorded for all communities analysed using both similarity indices and after controlling for temporal and fish size effects (Table 3, Fig. 5). Higher correlation coefficients were observed after controlling for the second variable as a consequence of the stronger correlation between distances separating samples and the differences in host size regarding those of date of capture. Comparing among the four distance-decay regressions, those based on qualitative similarity of infracommunities (Fig. 5a) and on Bray-Curtis similarity of prevalence (Fig. 5b) showed a more gradual slope, representing a lower decay rate, than those based on quantitative similarity of infracommunity abundances (Fig. 5c) and of component community mean abundances (Fig. 5d). Despite their steeper slopes, these regressions showed lower

Table 2. Taxonomic composition, prevalence (*P*) and mean abundance \pm standard deviation (MA \pm S.D.) of parasites of *Percophis brasiliensis* from four newly sampled localities in the South West Atlantic. Long-lived larval species are identified with an asterisk

	CF		RG		VG		NE	
	<i>P</i>	MA \pm S.D.	<i>P</i>	MA \pm S.D.	<i>P</i>	MA \pm S.D.	<i>P</i>	MA \pm S.D.
DIGENEA								
<i>Aponurus laguncula</i>	–	–	–	–	–	–	2.2	0.02 \pm 0.1
<i>Cardicola ambrosioi</i>	10	0.5 \pm 1.9	4.3	0.06 \pm 0.3	–	–	2.2	0.02 \pm 0.1
<i>Derogenes varicus</i>	–	–	–	–	–	–	2.2	0.09 \pm 0.6
Didymozoidae gen. sp.*	5	0.05 \pm 0.2	–	–	–	–	–	–
<i>Elytrophalloides oatesi</i>	–	–	–	–	–	–	2.2	0.02 \pm 0.1
<i>Lecithochirium microstomum</i>	–	–	17.0	0.9 \pm 2.5	27.5	1.4 \pm 3.9	33.3	3.1 \pm 7.2
<i>Lecithocladium cristatum</i>	–	–	–	–	3.9	0.05 \pm 0.3	–	–
<i>Parahemiurus merus</i>	–	–	–	–	–	–	2.2	0.02 \pm 0.1
<i>Prosorhynchus</i> sp.*	–	–	–	–	9.8	0.09 \pm 0.3	6.7	0.1 \pm 0.5
CESTODA								
<i>Scolex polymorphus</i>	40	20.3 \pm 55.8	44.7	1.1 \pm 1.9	47.1	34.9 \pm 73.0	73.3	76.6 \pm 118.0
<i>Grillotia</i> (<i>Christianella</i>) <i>carvajalregorum</i> *	90	7.35 \pm 9.9	100	1925.7 \pm 3219.5	100	491.2 \pm 386.0	100	719.9 \pm 538.3
<i>Callitetrarhynchus gracilis</i> *	2.5	0.03 \pm 0.2	–	–	2.0	0.02 \pm 0.1	13.3	0.1 \pm 0.3
<i>Pterobothrium</i> sp.	10	0.1 \pm 0.3	–	–	–	–	–	–
Tentaculariidae gen. sp.*	17.5	0.3 \pm 0.6	4.3	0.04 \pm 0.2	54.9	1.9 \pm 3.9	46.7	1.0 \pm 1.5
<i>Pseudophyllidea plerocercoid</i> *	–	–	–	–	2.0	0.02 \pm 0.1	–	–
NEMATODA								
<i>Anisakis simplex</i> s.l.*	–	–	6.4	0.1 \pm 0.4	15.7	0.3 \pm 0.7	57.8	2.8 \pm 4.9
<i>Contracecum</i> sp.*	–	–	12.8	0.03 \pm 1.2	11.8	0.1 \pm 0.3	2.2	0.04 \pm 0.3
<i>Cucullanus</i> sp.	–	–	–	–	2.0	0.01 \pm 0.1	–	–
<i>Hysterothylacium</i> sp.*	87.5	5.3 \pm 5.0	100	374.3 \pm 518.8	100	172.4 \pm 144.0	100	68.3 \pm 64.5
<i>Hysterothylacium fortalezae</i> *	12.5	0.2 \pm 0.6	–	–	–	–	–	–
<i>Hysterothylacium aduncum</i>	–	–	–	–	2.0	0.02 \pm 0.1	–	–
<i>Moravecchia argentinensis</i>	22.5	0.3 \pm 0.5	6.4	0.6 \pm 0.2	43.1	1.1 \pm 2.1	35.6	0.6 \pm 1.0
<i>Terranova</i> sp.*	5.0	0.05 \pm 0.2	–	–	–	–	2.2	0.02 \pm 0.1
<i>Raphidascaris acus</i> *	17.5	0.2 \pm 0.4	4.3	0.2 \pm 0.9	–	–	–	–
ACANTHOCEPHALA								
<i>Bolbosoma turbinella</i> *	7.5	0.08 \pm 0.3	8.5	0.4 \pm 1.8	13.7	0.2 \pm 0.5	–	–
<i>Corynosoma australe</i> *	2.5	0.03 \pm 0.2	76.6	25.0 \pm 57.1	100	110.5 \pm 197.0	100	167.4 \pm 164.3
<i>Corynosom cetaceum</i> *	2.5	0.03 \pm 0.2	12.8	0.2 \pm 0.6	62.7	1.4 \pm 1.8	75.6	2.8 \pm 3.3
<i>Serrasentis</i> sp.*	2.5	0.03 \pm 0.2	–	–	–	–	–	–
ISOPODA								
Gnathiidae gen. sp.	–	–	2.1	0.02 \pm 0.1	39.2	0.8 \pm 1.4	13.3	0.3 \pm 1.1
COPEPODA								
<i>Caligus</i> sp.	2.5	0.03 \pm 0.2	–	–	–	–	–	–

CF, Cabo Frio; RG, Rio Grande; VG, Villa Gessel; NE, Necochea.

correlation coefficients due to a number of similarities between pairs of samples departing considerably from the expected function. Indeed, residual values >10 were observed only for those regressions concerning infracommunity abundance and mean abundance. Some of these points were considerably more similar than expected given distance separating them, whereas the opposite situation was observed for other pairs of samples. The identification of such values evidenced that, among those points representing pairs of samples more similar than expected (those over the regression line) in the case of infracommunity similarity (Fig. 5c), the most extreme value was the similarity between the northern and southern boundaries of the Argentine

province (CF-San Antonio (SA), square over the regression line); the others were those including RG and several localities in the northern Argentine Sea and two including Río de la Plata (RP) and the two southernmost samples El Rincón (ER) and SA (white circles over the regression line). Contrarily, among localities evidencing a higher dissimilarity than expected by the distance between them (those below the regression line), were all represented by pairs of samples including CF (white circles below the regression line), with the most extreme dissimilarity occurring between CF and RG (square below the regression line). A similar picture was observed for analyses based on mean abundances (data not shown).

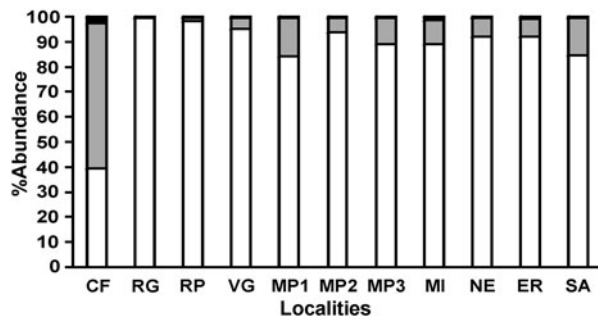


Fig. 2. Proportional contribution in terms of abundance of different parasite guilds to the assemblages parasitizing *Percophis brasiliensis* in 11 samples distributed along the south-western Atlantic coasts. Black: short-lived parasites (adult endoparasites + ectoparasites); grey: short-lived larval endoparasites (those living in the gut, such as larval cestodes); white: long-lived larval endoparasites. Locality codes as in Fig. 1.

MDS and cluster analyses on the four datasets were congruent in their results, revealing apparent patterns of separation between samples following a latitudinal pattern (Fig. 6), which was in all cases substantially different from random as shown by their low stress levels (0.01–0.02). Assemblages in fish from CF were the most different and branched first in the clusters, the second site differing from the rest was SA in all cases. In the case of intra-community qualitative similarity (Fig. 6a) the third branching region was ER followed by RG, but this order was inverted in the other three analyses (Fig 6b–d); on the other hand, all samples from northern Argentine Sea clustered together at the higher similarity levels in all cases.

DISCUSSION

Studies on variability of similarity in parasite assemblages of fish with increasing distance have usually focused on comparisons of distance-decay rates between kinds or species of hosts (Poulin, 2003; Oliva and González, 2005), guilds of parasites (Fellis and Esch, 2005) or hierarchical community levels (Timi *et al.* 2010), as well as being used to compare alternative hypotheses on biogeographical gradients in parasite species diversity (Seifertová, *et al.* 2008). The present results evidence that, beyond assessing the influence of distance *per se* on the similarity of parasite communities, distance decay regressions, and the identification of their outliers, can also shed light on the existence of discontinuities or uniformities in the geographic distribution of parasite assemblages and, in turn, in the zoogeography of their fish hosts.

Parasites have been considered as excellent indicators in present day and historical biogeography at different temporal and spatial scales (Brooks and Hoberg, 2000; Waltari *et al.* 2007). However, few studies have attempted to use parasites as indicators

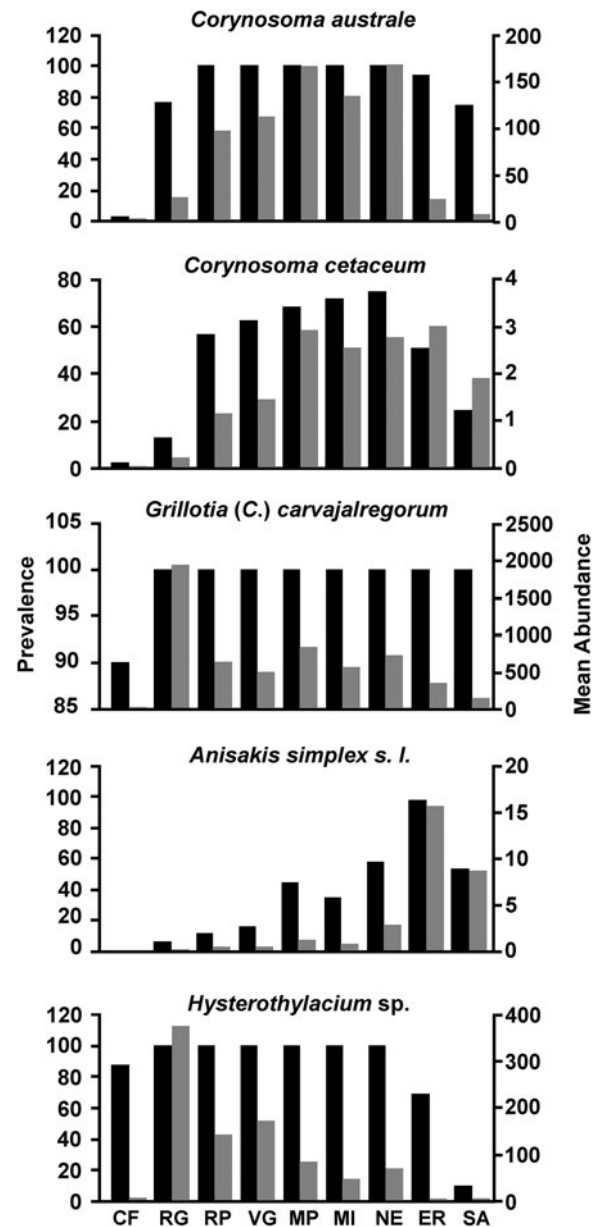


Fig. 3. Prevalence (black column) and mean abundance (grey column) of the most prevalent parasites of *Percophis brasiliensis* in nine samples (three samples from Mar del Plata averaged) distributed along the South Western Atlantic coasts. Locality codes as in Fig. 1.

of zoogeographical regions in the marine realm (Rohde, 2002).

Here we tested the potential of parasites as zoogeographical indicators by analysing long-lived parasites of a single species of marine fish, distributed along a unique zoogeographical province to assess their value as markers at lower levels of zoogeographical resolution. This guild was dominant in most parasite assemblages of *P. brasiliensis* over members of all other guilds, namely ectoparasites, larval gut parasites and adult endoparasites, which is a generalized pattern previously recorded across fish species in the northern Argentine Sea (Cantatore and Timi, 2015), and is here mainly

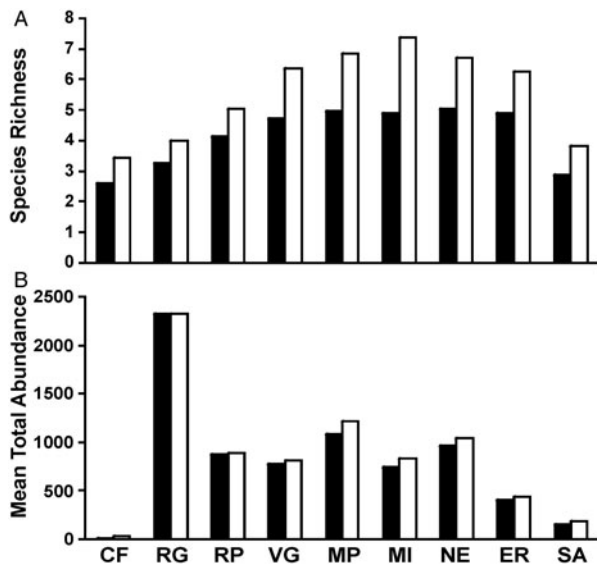


Fig. 4. Averaged descriptors of parasite infracommunities of *Percophis brasiliensis* in nine samples (three samples from Mar del Plata averaged) distributed along the south-western Atlantic coasts. (a) Species richness; (b) total abundance. White bars: all parasites; black bars: long-lived larval parasites. Locality codes as in Fig. 1.

due to the high prevalence and abundance of a few dominant species. Samples from CF, on the other hand, were the exception. Lower values of abundance were observed in this locality for those species, which were dominant in southern localities (*Corynosoma australe*, *G. (C.) carvajalregorum* and *Hysterothylacium* sp.), which represent a major difference of this locality regarding southern ones. The lower burdens of these parasites in CF may be due, at least in part, to the smaller size of fish in the sample, since the long temporal persistence of these larval parasites results in cumulative patterns of abundance as fish grow (Timi and Lanfranchi, 2013). Indeed, *Hysterothylacium* sp. has often been recorded as more abundant in CF than in southern localities in comparative studies of other fish species (Timi *et al.* 2005, 2010; Braicovich *et al.* 2012; Alarcos *et al.* 2016). However, for the other two parasite species, the abundances were always lower in the region of Rio de Janeiro (RJ) in all these hosts. Furthermore, juvenile *P. brasiliensis* from waters of the northern Argentine Sea, which were smaller than those from RJ (37.9 ± 4.3 , $n = 48$), harboured notably higher abundances of *C. australe* (mean abundance = 9.1) and *G. (C.) carvajalregorum* (mean abundance = 255.6) (unpublished own data), indicating that a locality effect, rather than host size, is determining the abundance of parasite populations. As a result, the sample from the Southern Brazilian district was the most different, probably also influenced by its position in the border with the northern Brazilian zoogeographical province.

In the Bonaerensean district, including fish from RG at the limit with Southern Brazilian district,

parasite species showed different patterns of latitudinal variation in their loads. These patterns are attributable to the latitudinal gradient in oceanographic conditions (mostly of temperature and salinity) characteristic of the region (Cantatore and Timi, 2015), along which a cline of combinations of biotic and abiotic factors provides sequential sets of optimal conditions where the reproduction and transmission of each parasite species are differentially favoured. These patterns are analogous to those expected according to the 'favourable centre' hypothesis, which predicts a unimodal distribution of parasites abundance, peaking at the centre of the geographical range of the host and declining toward the margins (Poulin and Dick, 2007; Seifertová *et al.* 2008). This seems to be the pattern for some parasites in *P. brasiliensis*, and responsible for the higher species richness of its infracommunities in the central zone of the Argentine Province. However, as only larval parasites were included in the analyses, the distribution of definitive, intermediate and other paratenic hosts surely play a major role in their distribution. The apparent 'favourable centre' in the central area of the study region is probably determined mainly by environmental conditions rather than by the distribution of *P. brasiliensis*. As mentioned above, these parasites appear to display similar distributions on other fish species; therefore the confirmation of the existence of repeatable distribution patterns across different host species will provide further evidence of their utility as zoogeographic indicators in the region. Furthermore, as several of the parasites analysed are distributed in other oceans (Sardella *et al.* 2005; Mattiucci and Nascetti, 2008; Beveridge and Campbell, 2010) they are potentially applicable as markers of different environmental conditions, and therefore as zoogeographical indicators in other regions.

The gradual variation of parasite species in terms of prevalence and abundance along the latitudinal and environmental gradient resulted in significant decays in similarity of parasite assemblages over distance. Qualitative similarity of infracommunities and quantitative similarity of prevalence decayed at a lower rate than those based on quantitative similarity of abundances (at both infracommunity and component community levels), an expectable result given that compositional variations are thought to be less pronounced within a zoogeographical province. On the other hand, higher variability in the abundance of different parasites among localities is a predictable response to environmental heterogeneity among them; variability probably amplified by the differences of three and four magnitude orders in the abundance of dominant species across sampling sites. These variations in abundance were also responsible for the lower correlation coefficients of abundance-based regressions, especially because

Table 3. Results of similarity decay analyses over distance of parasite assemblages of *Percophis brasiliensis* in the South West Atlantic, computed at component community and infracommunity levels, controlling for temporal and fish size effects

Similarity index	Community level	Response matrix data	Controlling for host size		Controlling for date	
			<i>r</i>	<i>P</i>	<i>r</i>	<i>P</i>
Jaccard	Infracommunity	Presence/absence	-0.89	<0.01	-0.94	<0.01
Bray-Curtis	Infracommunity	Abundance	-0.59	<0.01	-0.82	<0.01
Bray-Curtis	Component community	Mean abundance	-0.67	<0.01	-0.84	<0.01
Bray-Curtis	Component community	Prevalence	-0.89	<0.01	-0.94	<0.01

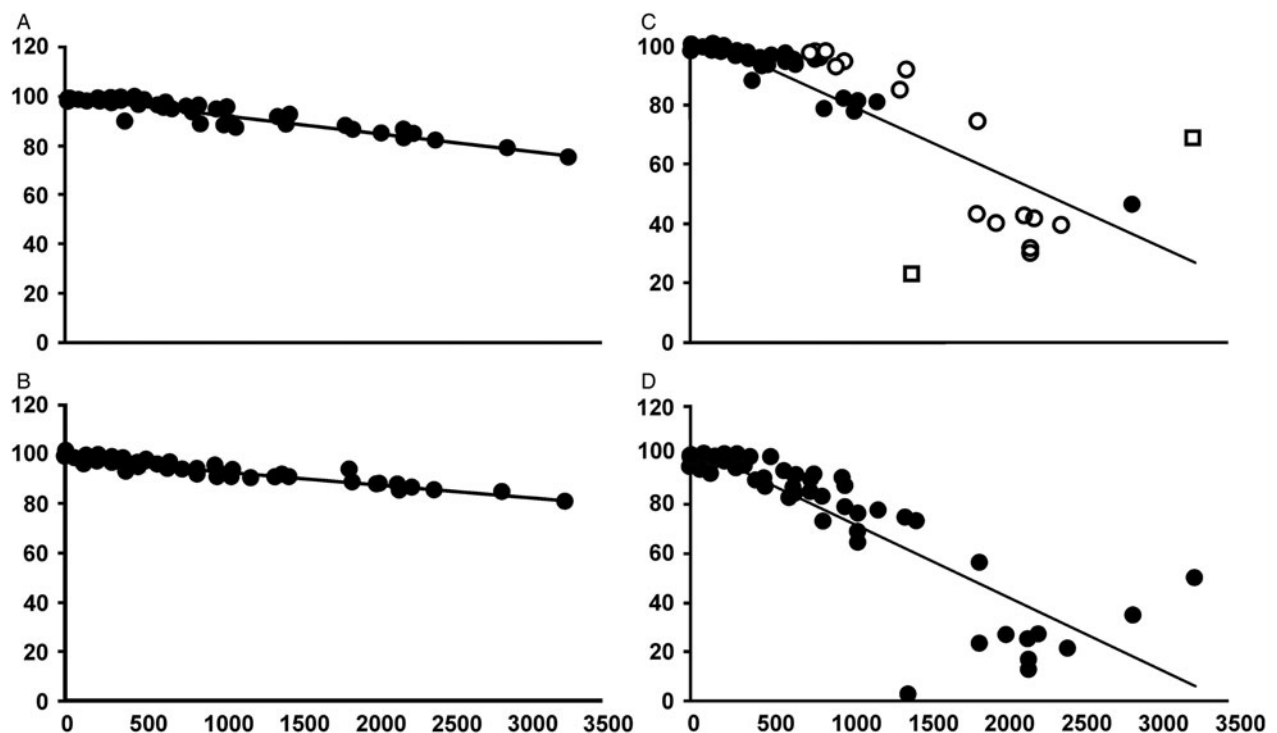


Fig. 5. Relationship between log-transformed similarity and untransformed geographic distance (km) in parasite assemblages of 11 samples of *Percophis brasiliensis* caught along the south-western Atlantic coasts. Similarities between samples are shown as a percentage of the maximum value of similarity observed. (a) averaged Jaccard similarity of infracommunities, (b) Bray-Curtis similarity of prevalence, (c) averaged Bray-Curtis similarity of infracommunity abundances, (d) Bray-Curtis similarity of component community mean abundance. (c) Different symbols are used to identify pairwise comparisons with residuals >10%; square over the regression line: Cabo Frio-San Antonio; white circles over the regression line: Rio Grande and several localities in the northern Argentine sea and two points including Rio de la Plata and the two southernmost samples (El Rincón and San Antonio); square below the regression line: Cabo Frio-Rio Grande; white circles below the regression line: Cabo Frio and several localities in the northern Argentine sea.

similarities between several pairs of samples departed considerably from the expected function, indicating that a border between zoogeographical units could occur when pairs of samples are more different than expected given the short distance separating them. This was the case between the two Brazilian localities, whereas distant localities unexpectedly more similar than predicted reflect their belonging to the same zoogeographical unit, for example RG-SA. The high similarity between both extremes of the study region (CF-SA) is, on the other hand, a consequence

of the lower burdens of most parasite species and therefore of the lower infracommunity species richness relative to the central localities. The lower prevalence and abundance recorded for all the species in both extremes of the study area can be attributed to the effect of opposite oceanographic conditions at these extremes, namely tropical waters in the north and subantarctic in the south, therefore similarity in distance-decay should be cautiously interpreted in the light of the knowledge of the environmental conditions characteristic of each region.

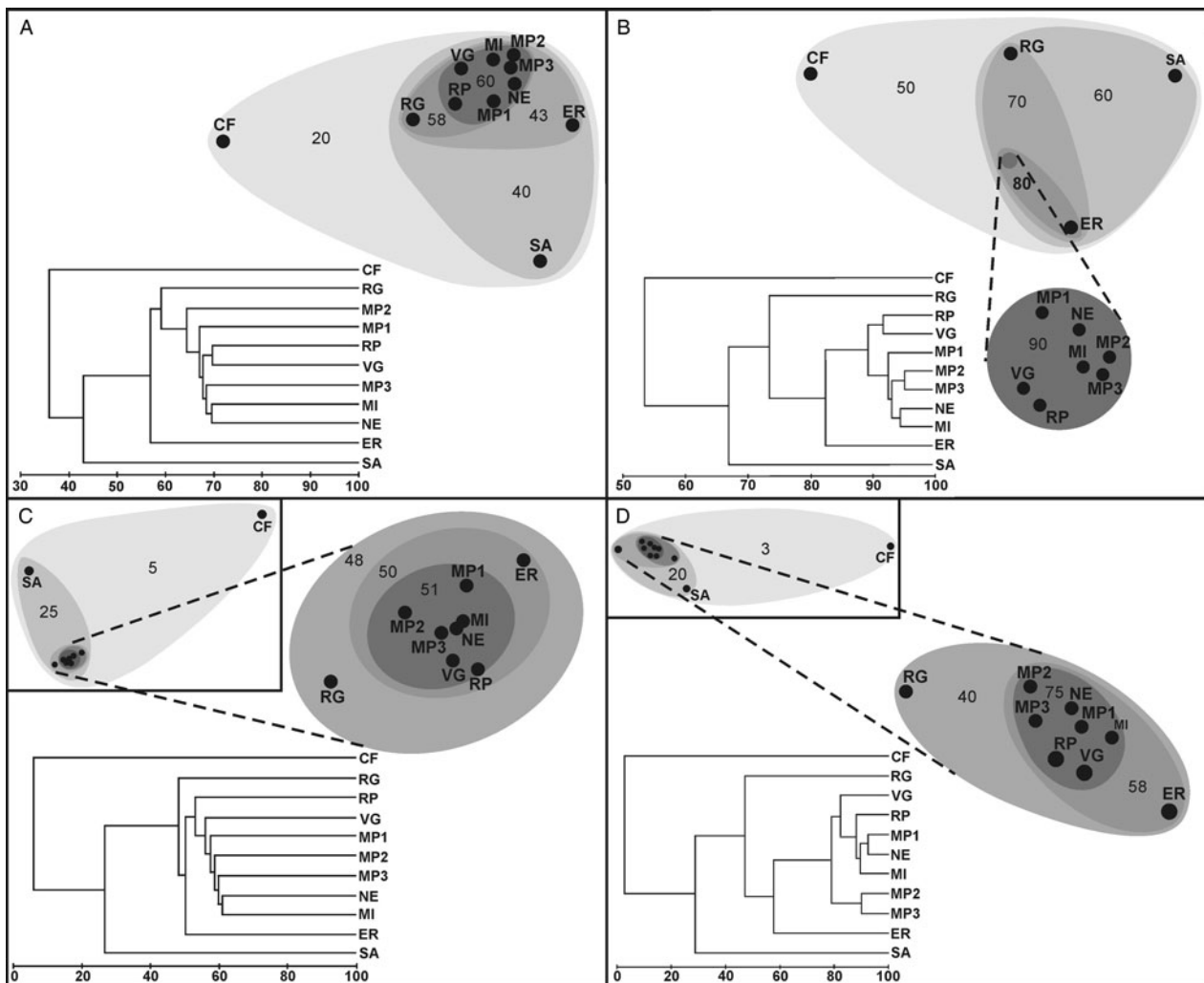


Fig. 6. Nonmetric two-dimensional ordination plot and cluster analyses of parasite communities in 11 samples of *Percophis brasiliensis* along the south-western Atlantic coasts. (a) averaged Jaccard similarity of infracommunities, (b) Bray-Curtis Similarity of Prevalence, (c) averaged Bray-Curtis similarity of infracommunity abundances, (d) Bray-Curtis similarity of component community mean abundance. Results of a hierarchical agglomerative clustering are shown as dendrograms and overlaid on the MDS plot with similarity levels represented by a grey scale, with its value given as a number inside each grey area. Locality codes as in Fig. 1.

These patterns were corroborated by multivariate analyses, which showed recurrent zoogeographical patterns in their results. Indeed, MDS and cluster analyses evidenced a clear first separation between samples from both districts, the Southern Brazilian and the Bonaerensean, with samples from RG clearly belonging to the second one. Among the Bonaerensean samples, SA was clearly different from the rest due to their location in the southern limit of the Argentine province bordering with the Magellanic province, and the ecotonal characteristics of the San Matias Gulf, which is considered as belonging to a different ecoregion (North Patagonian Gulfs) of the Magellanic Province (Spalding *et al.* 2007). Brazilian flatheads from RG were also different from southern samples, due mostly to their highest abundance of *G. carvajalregorum* and *Hysterothylacium* sp., although more

similar to them in their compositional structure, indicating their location on or near to the northern boundary of the Bonaerensean district or, following the scheme by Spalding *et al.* (2007), their membership to the ecoregion named RG. In synthesis, multivariate analyses revealed the value of larval parasites, recognized previously as suitable markers for flathead stock assessment (Braicovich and Timi, 2008, 2010), as valuable tools as zoogeographical indicators, corroborating a close fit of similarity between assemblages to existing zoogeographical classifications.

Finally, in light of the observed distributional patterns of parasites of *P. brasiliensis*, it is concluded that at least two discrete populations of this species inhabits in Brazilian waters, clearly differentiable from those three previously identified in Argentine and Uruguayan seas.

ACKNOWLEDGEMENTS

The authors wish to thank to Instituto Nacional de Investigación y Desarrollo Pesquero, Mar del Plata, and to Instituto de Biología Marina y Pesquera Almirante Storni, San Antonio Oeste, for providing some of the samples from Research Cruises.

FINANCIAL SUPPORT

Financial support to P. E. Braicovich and J. T. Timi provided by grants from CONICET (PIP # 112–201101-00036) and ANPCYT (PICT 2012 # 02094). Financial support to C. Pantoja, A.N. Pereira and J. L. Luque provided by research fellowships from Conselho Nacional de Desenvolvimento Científico e Tecnológico do Brazil (CNPq).

REFERENCES

- Alarcos, A. J., Pereira, A. N., Taboada, N. L., Luque, J. L. and Timi, J. T.** (2016). Parasitological evidence of stocks of *Paralichthys isosceles* (Pleuronectiformes: Paralichthyidae) at small and large geographical scales in South American Atlantic coasts. *Fisheries Research* **173**, 221–228.
- Anderson, M. J., Gorley, R. N. and Clarke, K. R.** (2008). *PERMANOVA+ for PRIMER: Guide to Software and Statistical Methods*. PRIMER-E, Plymouth.
- Avigliano, E., Saez, M. B., Rico, R. and Volpedo, A. V.** (2015). Use of otolith strontium: calcium and zinc: calcium ratios as an indicator of the habitat of *Percophis brasiliensis* Quoy & Gaimard, 1825 in the south western Atlantic Ocean. *Neotropical Ichthyology* **13**, 187–194.
- Beveridge, I. and Campbell, R. A.** (2010). Validation of *Christianella* Guiart, 1931 (Cestoda: Trypanorhyncha) and its taxonomic relationship with *Grillotia* Guiart, 1927. *Systematic Parasitology* **76**, 111–129.
- Braicovich, P. E. and Timi, J. T.** (2008). Parasites as biological tags for stock discrimination of the Brazilian flathead, *Percophis brasiliensis* in the south-west Atlantic. *Journal of Fish Biology* **73**, 557–571.
- Braicovich, P. E. and Timi, J. T.** (2010). Seasonal stability in parasite assemblages of the Brazilian flathead, *Percophis brasiliensis* (Perciformes: Percophidae): predictable tools for stock identification. *Folia Parasitologica* **57**, 206–212.
- Braicovich, P. E., Luque, J. L. and Timi, J. T.** (2012). Geographical patterns of parasite infracommunities in the rough scad, *Trachurus lathami* Nichols, in the southwestern Atlantic ocean. *Journal of Parasitology* **98**, 768–777.
- Briggs, J. C. and Bowen, B. W.** (2012). A realignment of marine biogeographic provinces with particular reference to fish distributions. *Journal of Biogeography* **39**, 12–30.
- Brooks, D. R. and Hoberg, E. P.** (2000). Triage for the biosphere: the need and rationale for taxonomic inventories and phylogenetic studies of parasites. *Comparative Parasitology* **67**, 1–25.
- Bush, A. O., Lafferty, K. D., Lotz, J. M. and Shostak, A. W.** (1997). Parasitology meets ecology on its own terms: Margolis *et al.* Revisited. *Journal of Parasitology* **83**, 575–583.
- Cantatore, D. M. P. and Timi, J. T.** (2015). Marine parasites as biological tags in South American Atlantic waters, current status and perspectives. *Parasitology* **142**, 5–24.
- Clarke, K. R. and Gorley, R. N.** (2006). *PRIMER v6: User Manual/Tutorial*. PRIMER-E, Plymouth.
- Cousseau, M. B. and Perrotta, R. G.** (2004). *Peces marinos de Argentina. Biología, distribución, pesca*. Publicaciones especiales INIDEP, Mar del Plata, Argentina.
- Esch, G. W. and Fernández, J. C.** (1993). *A Functional Biology of Parasitism*, 1st Edn. Chapman & Hall, London, UK.
- Fellis, K. J. and Esch, G. W.** (2005). Variation in life cycle affects the distance decay of similarity among bluegill sunfish parasite communities. *Journal of Parasitology* **91**, 1484–1486.
- Goslee, S. C. and Urban, D. L.** (2007). The ecodist package for dissimilarity-based analysis of ecological data. *Journal of Statistical Software* **22**, 1–19.
- MacKenzie, K. and Abaunza, P.** (2013). Parasites as biological tags. In *Stock Identification Methods*, 2nd edition (ed. Cadrin, S., Kerr, L. and Mariani, S.), pp. 185–204. Elsevier, San Diego, USA.
- Magurran, A. E.** (1988). *Ecological Diversity and its Measurement*. Princeton University Press, Princeton, New Jersey.
- Mattiucci, S. and Nascetti, G.** (2008). Advances and trends in the molecular systematic of anisakid nematodes, with implications for their evolutionary ecology and host-parasite co-evolutionary processes. *Advances in Parasitology* **66**, 47–148.
- Menni, R. C., Jaureguizar, A. J., Stehmann, M. F. W. and Lucifora, L. O.** (2010). Marine biodiversity at the community level: zoogeography of sharks, skates, rays and chimaeras in the southwestern Atlantic. *Biodiversity and Conservation* **19**, 775–796.
- Milessi, A. C. and Mari, N. R.** (2012). Ecología trófica del pez palo, *Percophis brasiliensis* (Quoy y Gaimard, 1825) en el ecosistema costero Argentino-Uruguayo (34° S- 41° S). *Revista de Investigación y Desarrollo Pesquero* **21**, 61–72.
- Miloslavich, P., Klein, E., Diaz, J. M., Hernandez, C. E., Bigatti, G., Campos, L., Artigas, F., Castillo, J., Penchaszadeh, P., Neill, P., Carranza, A., Retana, M., Diaz de Astarloa, J. M., Lewis, M., Yorio, P., Piriz, M., Rodriguez, G., Yoneshigue-Valentin, Y., Gamboa, L. and Martín, A.** (2011). Marine biodiversity in the Atlantic and Pacific coasts of South America: knowledge and gaps. *PLoS ONE* **6**, e14631.
- Oliva, M. E. and González, M. T.** (2005). The decay of similarity over geographical distance in parasite communities of marine fishes. *Journal of Biogeography* **32**, 1327–1332.
- Pereira, A. N., Pantoja, C., Luque, J. L. and Timi, J. T.** (2014). Parasites of *Urophycis brasiliensis* (Gadiformes: Phycidae) as indicators of marine ecoregions in coastal areas of the South American Atlantic with the assessment of their stocks. *Parasitology Research* **113**, 4281–4292.
- Pérez-del-Olmo, A., Fernández, M., Raga, J. A., Kostadinova, A. and Morand, S.** (2009). Not everything is everywhere: the distance decay of similarity in a marine host-parasite system. *Journal of Biogeography* **36**, 200–209.
- Poulin, R.** (2003). The decay of similarity with geographical distance in parasite communities of vertebrate hosts. *Journal of Biogeography* **30**, 1609–1615.
- Poulin, R. and Dick, T. A.** (2007). Spatial variation in population density across the geographical range in helminth parasites of yellow perch *Perca flavescens*. *Ecography* **30**, 629–636.
- Poulin, R. and Kamiya, T.** (2015). Parasites as biological tags of fish stocks: a meta-analysis of their discriminatory power. *Parasitology*, **142**, 145–155.
- Rohde, K.** (2002). Ecology and biogeography of marine parasites. *Advances in Marine Biology* **43**, 1–86.
- Sardella, N. H., Mattiucci, S., Timi, J. T., Bastida, R., Rodriguez, D. and Nascetti, G.** (2005). *Corynosoma australe* Johnston, 1937 and *C. ceta-ceum* Johnston & Best, 1942 (Acanthocephala: Polymorphidae) from marine mammals and fish in Argentinean waters: genetic differentiation and taxonomic status. *Systematic Parasitology* **61**, 143–156.
- Seifertová, M., Vyskočilová, M., Morand, S. and Šimková, A. S.** (2008). Metazoan parasites of freshwater cyprinid fish (*Leuciscus cephalus*): testing biogeographical hypotheses of species diversity. *Parasitology* **135**, 1417–1435.
- Soininen, J., McDonald, R. and Hillebrand, H.** (2007). The distance decay of similarity in ecological communities. *Ecography* **30**, 3–12.
- Spalding, M. D., Fox, H. E., Allen, G. R., Davidson, N., Ferdaña, Z. A., Finlayson, M., Halpern, N. S., Jorge, M. A., Lombana, A., Lourie, S. A., Martin, K. D., McManus, E., Molnar, J., Recchia, C. A. and Robertson, J.** (2007). Marine ecoregions of the world: a bioregionalization of coastal and shelf areas. *BioScience* **57**, 573–583.
- Timi, J. T. and Lanfranchi, A. L.** (2013). Ontogenetic changes in heterogeneity of parasite communities of fish: disentangling the relative role of compositional versus abundance variability. *Parasitology* **140**, 309–317.
- Timi, J. T. and MacKenzie, K.** (2015). Parasites in fisheries and mariculture. *Parasitology* **142**, 1–4.
- Timi, J. T., Luque, J. L. and Sardella, N. H.** (2005). Parasites of *Cynoscion guatucupa* along South American Atlantic coasts: evidence for stock discrimination. *Journal of Fish Biology* **67**, 1603–1618.
- Timi, J. T., Lanfranchi, A. L. and Luque, J. L.** (2010). Similarity in parasite communities of the teleost fish *Pinguipes brasilianus* in the southwestern Atlantic: infracommunities as a tool to detect geographical patterns. *International Journal of Parasitology* **40**, 243–254.
- Vales, D. G., García, N. A., Crespo, E. A. and Timi, J. T.** (2011). Parasites of a marine benthic fish in the Southwestern Atlantic: searching for geographical recurrent patterns of community structure. *Parasitology Research* **108**, 261–272.
- Waltari, E., Hoberg, E. P., Lessa, E. P. and Cook, J. A.** (2007). Eastward Ho: phylogeographical perspectives on colonization of hosts and parasites across the Beringian nexus. *Journal of Biogeography* **34**, 561–574.