An inverse latitudinal gradient of diversity of peracarid crustaceans along the Pacific Coast of South America: out of the deep south

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ABSTRACT

Aim To understand the ecological and historical/evolutionary processes underlying an inverse latitudinal gradient of richness (LGR) using crustacean peracarid species as a model group.

Location The Pacific coast of South America, along the Chilean coast between 18° S and 56° S.

Methods The LGR was evaluated using a dataset including 320 marine peracarid species reported for the coasts of Chile. Five ecological hypotheses invoking a relationship between species richness and present-day conditions were tested: species–energy, species–area, Rapoport rescue effect, mid-domain geometric constraint and niche breadth. Historical/evolutionary hypotheses (i.e. biogeographic conservatism, and diversification rates) were indirectly tested by analysing the latitudinal variation in the taxonomic distinctness, the taxonomic conservatism of the midpoint of the latitudinal range and the degree of nestedness at different taxonomic levels.

Results Richness increased poleward, varying approximately eightfold, following an inverse LGR coupled with an increase in bathymetric distribution. Overall this inverse LGR seems robust to uncertainties in the completeness of the species inventory. We found support for only two of the five ecological hypotheses tested: species–area and Rapoport rescue effect. Historical/evolutionary hypotheses seemed important in structuring the richness pattern, as indicated by the higher taxonomic distinctness in the southern region, the strong taxonomic inertia in the mean range size and the high degree of nestedness of assemblages at different taxonomic levels.

Conclusions When combined, these results underscore the importance of long-term processes and historical/evolutionary explanations for the inverse LGR, conceptualized in what we term the ‘out of the deep south’ hypothesis that involves the effects of both biogeographic niche conservatism and evolutionary rates. We propose that the southern region may be a source of evolutionary novelties and/or exhibit higher diversification rates (i.e. higher speciation/lower extinction rates). Furthermore, phylogenetic conservatism of latitudinal range may limit the geographic expansion of these new taxa towards the depauperated northern region.

Keywords Amphipoda, Chile, Cumacea, Isopoda, macroecology, macroevolution, marine biodiversity, Peracarida, Tanaidacea.
INTRODUCTION

The latitudinal gradient of species richness (LGR) remains one of the most prominent patterns in macroecology and biogeography (Pianka, 1966; Willig et al., 2003; Wiens & Graham, 2005; Mittelbach et al., 2007). Several meta-analyses confirm the pervasiveness of the canonical pattern of species diversity increasing from the poles toward the tropics (Willig et al., 2003; Hillebrand, 2004a,b).

While there is an ongoing debate about the underlying mechanisms driving the LGR (Rohde, 1999; Willig et al., 2003), there is growing recognition that ultimately the richness in a region should be determined by a balance between the rates of extinction, migration (i.e. range dynamics) and origination of species (Wiens & Donoghue, 2004; Mittelbach et al., 2007). This balance can be achieved by different mechanisms that can be broadly classified into three types of hypotheses, as proposed by a recent synthesis (Mittelbach et al., 2007): ecological, historical and evolutionary. Ecological hypotheses suggest that tropical areas harbour greater species richness due to environmental effects enhancing productivity, carrying capacity or niche breadth (Currie, 1991; Rosenzweig, 1995; Evans et al., 2005). Historical explanations (also referred to as biogeographical; Mittelbach et al., 2007) invoke the tropics as sources of evolutionary novelties (i.e. clades originated in the tropics, but overall diversification rates are not variable across latitude) and that the LGR is the result of a limited dispersal towards the temperate zones (Wiens & Donoghue, 2004; Wiens & Graham, 2005; Jablonski et al., 2006). Finally, evolutionary hypotheses suggest that the LGR is the result of a latitudinal variation in the diversification rates (although novelties appear evenly across latitude), with higher speciation and/or lower extinction rates in the tropics (Evans et al., 2005; Weir & Schluter, 2007). Despite the differences between historical and evolutionary hypotheses, they both highlight the importance of long-term dynamics explaining the LGR, so we will refer to both of them as historical/evolutionary hypotheses.

Despite its generality, exceptions to the canonical LGR are not uncommon in the marine and terrestrial realms (Stevens, 1989; Stephens & Wiens, 2003; Willig et al., 2003; Smith et al., 2005; Kindlmann et al., 2007; Fernández et al., 2009; Pyron & Burbrink, 2009). The temperate Pacific coast of South America (PSA hereafter) is unusual in having many different marine taxa showing an inverse LGR, such as seaweeds (Santelices, 1982; Santelices & Marquet, 1998; Meneses & Santelices, 2000; Santelices et al., 2009), sponges (Desqueyroux & Moyano, 1987), anthozoans (Häussermann & Försterra, 2005), bryozoans (Moyano, 1991), polychaetes (Hernández et al., 2005; Moreno et al., 2006), chitons, bivalves and gastropods (Herm, 1969; Valdovinos et al., 2003; Marquet et al., 2004), and peracarid crustaceans (Lancellotti & Vásquez, 2000; González et al., 2008). The inverse LGR phenomenon holds true across more than 50 degrees of latitude along the PSA, and thus cannot be explained as an artefact of reduced spatial scale (Willig et al., 2003). In addition, analyses strongly suggest that this pattern is robust to sampling artefacts (Santelices, 1982; Valdovinos et al., 2003).

Different mutually non-exclusive ecological hypotheses have been proposed to explain the existence of the inverse LGR along the PSA, including a Rapoport rescue effect (i.e. species show shorter geographic ranges along the southern PSA; Santelices & Marquet, 1998; Santelices et al., 2009), a species–area effect (coastal shelf area and coastal length are more extensive along the southern PSA, therefore it can harbour more species; Valdovinos et al., 2003; Santelices et al., 2009) and also a mid-domain effect (the richness gradient is an artefact of the overlapping ranges of distribution within a given area; Hernández et al., 2005; Moreno et al., 2006). Other explanations, including different and more refined forms of the species–energy hypothesis (i.e. niche breadth hypothesis; Vázquez & Stevens, 2004) have not been tested yet. Recently, Fernández et al. (2009) showed that the shape of the LGR along the PSA may be influenced by the type of larval development: molluscs and crustaceans with direct development consistently display an inverse LGR, in contrast to planktotrophic species which follow the canonical pattern of decreasing species richness with increasing latitude. These authors also showed for a limited dataset of peracarids (170 species) that the pattern of richness was mainly associated with sea surface temperature, suggesting that low temperatures in southern Chile may favour population persistence and lower extinction risk due to reduced metabolic costs (Brandt, 1999; Fernández et al., 2009). All these hypotheses are based on correlations between richness and present-day environmental conditions, revealing little information about the influence of historical/evolutionary processes in shaping the LGR (Jablonski et al., 2006; Mittelbach et al., 2007; Araújo et al., 2008). Moreover, recent studies demonstrate that the ‘anomalous’ LGR of some taxa can be explained by historical hypotheses (i.e. historical biogeography and phylogenetic niche conservatism; Stephens & Wiens, 2003; Smith et al., 2005; Pyron & Burbrink, 2009).

Antarctica has been considered to act as an ‘evolutionary incubator’ or centre of origin for many taxa, including peracarids (Knox & Lowry, 1977; Watling & Thurston, 1989; Brandt, 1999; Clarke et al., 2004). Furthermore, considering that peracarid richness has been reported to be greater at high latitudes, it is relevant to examine historical/evolutionary hypotheses as a potential explanation for the inverse LGR of peracarids along the PSA. However, suitable fossil records or phylogenetic reconstructions are lacking for marine peracarids along the PSA, precluding a rigorous test of these ideas similar to what has been done for other taxa and regions (Stephens & Wiens, 2003; Smith et al., 2005; Jablonski et al., 2006; Wiens et al., 2006, 2009; Weir & Schluter, 2007; Pyron & Burbrink, 2009). Nevertheless, indirect approaches used by recent studies may also shed light on the possible role of historical/evolutionary explanations in shaping spatial gradients of species richness. For instance, Krug et al. (2008) used taxonomic species–genus ratios to infer the importance of diversification rates across a latitudinal range. In addition, Roy et al. (2009) suggested a marked phylogenetic conservatism of poleward range limits in bivalve genera, using taxonomy as a proxy for phylogeny, lending support to the idea that limited dispersion of taxa of tropical origin may lead to the canonical LGR. The existence of nested assemblages at bioge-
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Testing ecological hypotheses

A correlation approach was used to test five hypotheses that have been recurrently proposed to explain the LGR. The species–energy hypothesis (Currie, 1991; Evans et al., 2005) was tested using the mean sea surface temperature and the concentration of chlorophyll \(a\) (Chl-\(a\)) as surrogates of the environmental energy. Mean annual sea surface temperature was obtained from the World Ocean Atlas 2001 and Chl-\(a\) from SeaWiFs satellite images. The species–area hypothesis (Rosenzweig, 1995) was tested using the shelf area (< 200 m depth) as a proxy of available habitat, and it was obtained from digitalized maps (1:106 scale Lambert zenithal, azimuthal equal area) (Astorga et al., 2003). The geometric constraint hypothesis or mid-domain effect (Pineda, 1993; Colwell & Hurtt, 1994) was tested using the median number of species expected by a mid-domain null model (10,000 runs), using the module MDE (McCain, 2004). The Rapoport rescue hypothesis (Stevens, 1989) was evaluated using the mean latitudinal range size (after log\(_{10}\) transformation) of all species co-occurring in the same latitudinal bin. Finally, the niche breadth hypothesis (Vázquez & Stevens, 2004) was tested using the mean bathymetric range (after log\(_{10}\) transformation) as a rough proxy for the degree of specialization in species. The rationale behind the use of median bathymetric range is that species concentrated in a narrow bathymetric belt experience less environmental variability (i.e. in terms of temperature, water pressure, food supply and types of substrata) than species having wider bathymetric ranges. Assuming that the bathymetric ranges are under equilibrium, they may reflect the niche requirements of the species. Therefore, the species richness should be negatively related to the median bathymetric range.

The association between species richness, oceanographic and ecological variables was explored using both traditional ordinary least square (OLS) and a simultaneous autoregressive model (SAR\(_{<m}\)) (Dormann et al., 2007; Kissling & Carl, 2008). Analyses were carried out using the library spdev in the software R (R Development Core Team, 2010), following the procedures detailed by Kissling & Carl (2008). The best models were selected using the Akaike information criterion (\(\Delta\text{AIC} < 2\)).
Indirect test of historical/evolutionary hypotheses

In the absence of suitable fossil records or molecular phylogenies, we conducted indirect tests of the importance of historical/evolutionary hypotheses (biogeographic conservatism and diversification rates), using three complementary methods: (1) taxonomic distinctness, (2) taxonomic conservatism of latitudinal range midpoint, and (3) nestedness analysis. The taxonomic distinctness index, Δ+ (Clarke & Warwick, 1998), was used as a coarse proxy of phylogenetic diversity. Briefly, this index measures the degree of taxonomic inter-relatedness between species, and assumes that phylogenetic relationships are mapped in the taxonomic structure. The distinctness corresponds to the average length of the taxonomic distance between each pair of species, which included species, genera, families and orders, assuming a similar branch distance of 1. Higher Δ+ values imply a taxonomically much more diverse assemblage. The observed Δ+ values were contrasted against the predictions of a null model, built by resampling the same number of species found in each bin and recalculating the Δ+ values 10,000 times. Lineage diversity was considered statistically significant when observed values were above or below the 97.5th and 2.5th percentiles. Invariant Δ+ values (i.e. not departing from the null model) across the latitudinal gradient may imply that the inverse LGR is not directly linked to historical biogeography or evolutionary processes. In contrast, high Δ+ values (i.e. above the null model) in the southern region could be interpreted as the existence of more evolutionary novelties (i.e. clades) in the southern region, supporting historical biogeography hypotheses. Alternatively, it may imply enhanced speciation rates and/or reduced extinction rates, according to the diversification rate hypothesis.

The taxonomic conservatism of the midpoint of latitudinal range (the median between the northern and southern distribution limits of each taxon) was evaluated using taxonomic structure (orders, families, genera) as a coarse proxy of relatedness among species. For simplicity, we have used the midpoint of latitudinal range instead of the predictions of niche models based on climatic variables (Smith et al., 2005; Pyron & Burbrink, 2009; Wiens et al., 2009). This decision is justified by the prevailing north–south direction of the PSA that maps a marked latitudinal gradient in the oceanographic conditions, so each latitudinal bin would represent a unique oceanographic condition. Taxonomic conservatism was evaluated using an autocorrelogram analysis (Moran’s I) computed across different taxonomic levels. Values of Moran’s I close to 1 are indicative of taxonomic conservatism at the midpoint of latitudinal range, e.g. the mean ranges of species within a genus are more similar than would be expected by random distribution. When I = 0, no taxonomic signal is detected, whereas a taxonomic repulsion (i.e. the mean range of species within genera differs significantly from a random distribution) is indicated by values of I = −1. The existence of taxonomic conservatism at the midpoint of latitudinal range, combined with a possible southern origin of many peracarid clades (Knox & Lowry, 1977; Watling & Thurston, 1989; Brandt, 1999; Clarke et al., 2004), may support the biogeographic conservatism hypothesis. Calculations were made using the function correlogram.formula in the library ape in R (R Development Core Team, 2010).

The nestedness analysis evaluates the degree of order in the spatial distribution of taxa. Factors that assemble or disassemble species communities from a common pool in a consistent order will lead to a nested structure (Wright et al., 1998). Nestedness is often used in community ecology studies to infer the relative importance of local dispersal–extinction dynamics in shaping the spatial distribution of species (Atmar & Patterson, 1993; Wright et al., 1998). In a nested community, the poorest sites (i.e. those with fewest species) are subsets of the richest sites, and not of the entire pool. The application of nestedness analyses to biogeographic questions is straightforward (Moreno et al., 2006, 2008). In our case, nested assemblages may imply that dispersal–extinction dynamics occur over very large scales, where the poorest regions would be characterized by high extinction rates, while richness in those zones should be maintained by dispersal and colonization from species-rich regions rather than by in situ speciation or the emergence of evolutionary novelties. We used the robust and conservative discrepancy index (D) as a measure of nestedness (Brualdi & Sanderson, 1999; Ulrich & Gotelli, 2007). To evaluate the significance of the nestedness we used the ‘swap’ algorithm to produce random matrices with a fixed number of rows and columns. Analyses were carried out at species, genus and family levels. A lack of nestedness indicates that the taxonomic composition in areas of low species richness is not a mere subset of the more diverse areas. This may imply the existence of biogeographically different biotas, perhaps exhibiting different ecological and evolutionary features. On the contrary, nested assemblages may indicate the existence of biota with a shared biogeographic history. However, nested assemblages may be created by either biogeographic conservatism or latitudinal variations in diversification rates. Calculations were made using the function oecosimu implemented in the library vegan in R (R Development Core Team, 2010).

RESULTS

The species richness showed an inverse latitudinal gradient, with richness increasing towards high latitudes (Figs 1 & 2a). Richness increased about eightfold across the study region, with minimum and maximum values recorded at c. 20°S (18 species) and c. 54°S (136 species), respectively. Most of the change in richness occurred in the northern and southern areas, whereas between 34°S and 50°S the gradient was rather stable. The increase in richness is coupled with a marked poleward decline in the median latitudinal range size, from a range of c. 20° in the northern-central region to a 4° range at the southern tip of the study area (Fig. 2b). In addition, both the mean bathymetric range and the mean bathymetric midpoint of species showed a more than fivefold increase in the poleward direction (Fig. 2c); this is primarily due to the increase in the number of subtidal species with increasing latitude, while numbers of intertidal species vary little across the study area (Fig. 2d).

When the gradient was deconstructed according to the year of the first report of the species in Chile, the poleward increase in
Figure 1  Map of the study region, showing the latitudinal distribution of 320 peracarid species studied.

Figure 2  Latitudinal gradient of peracarid diversity along the Pacific coast of South America, for (a) total species richness, (b) mean latitudinal range, (c) bathymetric distribution (mean bathymetric range and mean bathymetric midpoint), and (d) species richness according to bathymetric categories (inter- and subtidal zones). Data for (b) and (c) were estimated after log_{10} transformation, but they are depicted on a linear scale to facilitate the interpretation of results.
species richness had been evident as early as 1900. The number of additional species reported between 1950 and 2007 did not change the shape of the gradient (Fig. 3a). The same pattern is also evident looking at the correlation coefficient between species richness and latitude: it has been positive and notably invariant over the last c. 100 years (Fig. 3b).

The species richness showed a variable relationship with the explanatory variables used to test the five ecological hypotheses (Table 1). In general, the residuals of the relationship showed a significant spatial autocorrelation. However, the sign of the slopes were not modified by the SARerr modelling, and qualitative similar conclusions were reached using either the traditional OLS regression or the SARerr model (Table 1). The best supported models, according to the ΔAIC, were sea surface temperature and median latitudinal range size. However, although species richness was strongly related to sea surface temperature (OLS, \( r^2 = 0.74, P < 0.05; \) Fig. 4a) and showed the lowest AIC, the sign of the coefficient was negative, against the predictions of the species–energy hypothesis. Richness was negatively related to median latitudinal range size (OLS, \( r^2 = 0.68, P < 0.05, \) Fig. 4b) according to the predictions of the Rapoport rescue hypothesis. Other models were less supported, as the ΔAIC values were greater than 2. Species richness was positively associated with shelf area (OLS, \( r^2 = 0.58, P < 0.05; \) Fig. 4c), following the predictions of the species–area hypothesis. Richness was also positively associated with the median bathymetric range (OLS, \( r^2 = 0.55, P < 0.05; \) Fig. 4d), but the direction of the relationship was opposite to the one predicted by the niche-breadth hypothesis. Richness was not significantly related to Chl-\( \alpha \) or the mid-domain effect (OLS, \( r^2 = 0.02, P > 0.05; \) respectively), indicating no support for the species–energy or geometric constraint hypotheses.

According to the results of indirect tests, historical/evolutionary processes can help to explain the observed LGR. First, the taxonomic distinctness index increased poleward (Fig. 5a). The observed \( \Delta + \) values were not different from the null expectation, except at the southern tip where \( \Delta + \) was higher than expected by chance. This trend is associated with the ordinal level patterns of richness (Fig. 5b). The latitudinal trend of species richness was mostly driven by amphipods and isopods, whereas cumaceans and tanaidaceans were present only in the southern region. Secondly, the midpoint of latitudinal range was positively autocorrelated at different taxonomic levels, according to the positive and significant values of Moran’s I (Table 2). Finally, the nestedness analyses showed differences amongst taxonomic levels. At species and genus levels, the assemblage was significantly nested (species, \( D = 496, Z\text{-score} = 5.22, P = 0.002; \) genus, \( D = 326, Z\text{-score} = 5.08, P < 0.0001 \). In contrast, no significant pattern was found at the family level (\( D = 100, Z\text{-score} = -1.32, P = 0.136 \).

**DISCUSSION**

**An inverse latitudinal gradient of richness**

Consistent with previous reports, our analyses support the existence of a strong inverse LGR of peracarid species richness along the Pacific coast of South America (Lancellotti & Vásquez, 2000; González et al., 2008; Fernández et al., 2009). Although a reverse LGR of richness challenges the apparent generality of the pattern of decreasing species diversity within increasing latitude (Willig et al., 2003; Hillebrand, 2004a,b), this result is not entirely unexpected because several exceptions to the classic pattern have been reported (Willig et al., 2003). Indeed, the inverse LGR has been previously reported for marine isopods along the entire Pacific coast of the Americas by Espinosa-Pérez & Hendrickx (2006), although their analyses were conducted at a coarser scale comparing biogeographic provinces.

Even though the species inventory in the region can be considered only as a representative fraction of the total peracarid diversity, and new surveys may add species or range extensions of known species (Thiel et al., 2003; González et al., 2008), the presented results suggest it is unlikely that the observed inverse LGR will be affected by these additions (i.e. a better sampling in the southern region). Our analysis confirmed that the inverse LGR should not be severely biased by improvements of the species inventory, as the shape of the gradient was largely independent of the inclusion of new reported species during the last century. Therefore, the latitudinal pattern reported herein appears to reflect the true biogeographic pattern for peracarid crustaceans along the PSA, validating previous results based on a much more restricted subset of peracarid species (Lancellotti & Vásquez, 2000; Fernández et al., 2009).
Ecological versus historical/evolutionary hypotheses to explain the inverse LGR

The long and lasting debate about the ultimate causes of the canonical LGR also applies to an inverse LGR: different ecological (Santelices & Marquet, 1998; Valdovinos et al., 2003; Marquet et al., 2004; Kindlmann et al., 2007; Fernández et al., 2009) and historical/evolutionary explanations have been proposed (Stephens & Wiens, 2003; Valdovinos et al., 2003; Smith et al., 2005; Pyron & Burbrink, 2009). Among the five ecological hypotheses evaluated here, we found strong support for the Rapoport rescue effect and partial support for the species–area hypothesis. These hypotheses have been previously proposed to explain the inverse gradient of richness observed in seaweeds and molluscs, respectively (Santelices & Marquet, 1998; Valdovinos et al., 2003). Other hypotheses, including the species–energy and mid-domain constraints, have been proposed previously to explain richness patterns in other invertebrates along the study region (Astorga et al., 2003; Hernández et al., 2005), but they were not supported by our analyses.

The sign of the relationship between richness, sea surface temperature and median bathymetric range was exactly the opposite of what is predicted by the species–energy (Currie, 1991; Evans et al., 2005) and niche-breadth hypotheses (Vázquez & Stevens, 2004), respectively. This sharp discrepancy may reflect the fact that the underlying processes that lead to these relationships are different from those originally proposed: sea surface temperature is not mirroring energy availability, and bathymetric range is not reflecting local species coexistence. On the one hand, the negative relationship between richness and temperature may result from the fact that many life-history

<table>
<thead>
<tr>
<th>Ecological hypothesis</th>
<th>Explanatory variable</th>
<th>OLS regression</th>
<th>SAR, regression</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species–energy</td>
<td>Sea surface temperature</td>
<td>Slope = -6.33</td>
<td>AIC = 162.6</td>
</tr>
<tr>
<td>Species–energy</td>
<td>Chlorophyll-a</td>
<td>Slope = -53.66</td>
<td>AIC = 187.4</td>
</tr>
<tr>
<td>Species–area</td>
<td>Shelf area</td>
<td>Slope = 1.93</td>
<td>AIC = 179.2</td>
</tr>
<tr>
<td>Geometric constraint</td>
<td>Mid-domain effect</td>
<td>Slope = 0.24</td>
<td>AIC = 108.4</td>
</tr>
<tr>
<td>Rapoport’s rule</td>
<td>(log_{10}) Mean latitudinal range</td>
<td>Slope = -130.5</td>
<td>AIC = 166.4</td>
</tr>
<tr>
<td>Niche breadth</td>
<td>(log_{10}) Mean bathymetric range</td>
<td>Slope = 93.2</td>
<td>AIC = 172.9</td>
</tr>
</tbody>
</table>

AIC, Akaike information criterion. Significant values (P < 0.05) are in bold.

Figure 4 Relationship between peracarid species richness and (a) sea surface temperature (SST), (b) shelf area, (c) latitudinal range size, and (d) bathymetric range size. Also shown are the values of the ordinary least squares regressions, and Moran’s I after accounting for spatial autocorrelation. ns = non-significant.

Table 1 Summary of the ordinary least squares (OLS) and simultaneous autoregressive model (SAR, ) regressions between species richness and different ecological variables aimed to test different ecological hypotheses to explain the inverse latitudinal gradient of richness.
Figure 5 Latitudinal variation of taxonomic structure: (a) taxonomic distinctness, (b) species richness by orders.

Table 2 Results of the autocorrelogram analysis (Moran’s I) used to evaluate the existence of taxonomic conservatism of the midpoint of latitudinal distribution.

<table>
<thead>
<tr>
<th>Taxonomic level</th>
<th>Rescaled Moran’s I</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species/genera</td>
<td>0.289</td>
<td>0.0004</td>
</tr>
<tr>
<td>Genera/families</td>
<td>0.345</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Families/orders</td>
<td>0.025</td>
<td>0.0062</td>
</tr>
</tbody>
</table>

traits of Peracarida show a higher performance at cooler temperatures (e.g. reduced costs of brooding, increased longevity and fecundity, and low but prolonged reproductive rates; Brandt, 1999). This k-strategy may allow peracarids to reach larger population sizes, increasing population persistence and probably species diversity by reducing extinction rates at high latitudes (Brandt, 1999; Fernández et al., 2009). On the other hand, the positive relationship between richness and bathymetric distribution may be explained by the well-known eurybathic pattern of this and other marine benthic groups in the Antarctic region (Brey et al., 1996). The wide bathymetric distribution allows species inhabiting coastal areas to survive the effects of ice scouring during glacial cycles, finding refuge in deeper areas of the shelf and slope, hence reducing their extinction risks (Brey et al., 1996; Poulin et al., 2002). Thus, rather than implying contemporaneous ecological processes, these results suggest the importance of evolutionary processes in shaping latitudinal patterns of diversity in peracarids.

When the previous evidence is combined with the results of the analyses of the taxonomic distinctness, taxonomic conservatism of latitudinal range midpoint and nestedness, a new picture of the processes involved is revealed: the poleward increase in species richness may be explained by some sort of historical/evolutionary hypothesis (Stephens & Wiens, 2003; Smith et al., 2005; Mittelbach et al., 2007; Pyron & Burbrink, 2009). On the one hand, the so-called ‘biogeographic conservatism hypothesis’ (Pyron & Burbrink, 2009), a more general version of the ‘tropical conservatism hypothesis’ (Wiens & Donoghue, 2004), predicts the origination of new clades at the higher-diversity zones (either tropical or temperate) and a recent and restricted dispersal towards low-diversity zones due to phylogenetic niche conservatism (Stephens & Wiens, 2003; Smith et al., 2005; Pyron & Burbrink, 2009). This hypothesis is conceptually similar to the ‘out of the tropics’ hypothesis proposed by Jablonski and collaborators (Jablonski et al., 2006) and is confirmed in some marine invertebrates (Martin et al., 2007; Krug et al., 2008; Roy et al., 2009). On the other hand, the evolutionary hypotheses predict latitudinal variation in evolutionary rates, e.g. higher speciation and/or lower extinction rates at the high-diversity zones (Mittelbach et al., 2007; Weir & Schluter, 2007). Although our current approach does not allow us to weigh the effects of historical and evolutionary hypotheses separately, the results highlight the potential role played by processes operating over long temporal scales in shaping the inverse LGR of peracarid species along the PSA.

The biogeographic conservatism hypothesis is supported by the larger values of the distinctness index at the southern tip, suggesting a kind of ‘museum effect’ of accumulation of older clades. This idea is further supported by the general notion of a high diversity and evolutionary success of peracarids around the southern ocean (Knox & Lowry, 1977; Watling & Thurston, 1989; Brandt, 1999; Clarke et al., 2004), and by the existence of nested assemblages at species and genus level, which suggest that clades present in the northern region are a subset of the southern biota. Indeed, there are two orders, Cumacea and Tanaiidae, that are largely absent in the northern region. However, this does not mean that evolutionary novelties are not present in the northern region. Indeed, the lack of a nested pattern at family level is explained by a considerable number of families restricted to the northern zone. Nevertheless, the restricted distribution of most of the southern clades is also reflected in a strong taxonomic conservatism of the midpoint of latitudinal range, suggesting that perhaps phylogenetic conservatism of the upper thermal tolerance of species has restricted more extensive expansion of these clades towards the northern region across evolutionary time-scales.

Latitudinal variations in evolutionary rates can also potentially explain the inverse LGR. The taxonomic distinctness may, alternatively, reflect higher diversification rates at areas harbouring more diversity, as suggested in marine bivalves inhabiting tropical areas (Krug et al., 2008). This idea is also supported by accumulation of short-ranged species in the southern region (the Rapoport rescue effect), as the result of the high habitat fragmentation induced by glacial–interglacial cycles occurring in the region since the Pliocene (Valdovinos et al., 2003; Marquet et al., 2004). Furthermore, the existence of a positive relationship between richness, shelf area (Valdovinos et al., 2003; Marquet et al., 2004; Fernández et al., 2009) and habitat
An inverse gradient of latitudinal diversity

heterogeneity (Santelices et al., 2009) provides support for this hypothesis. Speciation rate should increase in larger areas (Losos & Schluter, 2000), because larger areas may harbour more habitats and also increase the probabilities of disrupting genetic flow among populations.

In parallel to a reduced speciation rate, local and regional-scale extinctions may also be higher in the northern region. We hypothesize that the existence of a shallow oxygen minimum zone (OMZ) and cyclic El Niño (EN) events would be major mechanisms increasing extinction probabilities in this region. A very shallow OMZ has been described for the Peru–Chile region up to c. 37°S (Morales et al., 1999; Helly & Levin, 2004), the upper boundary of the OMZ can reach depths as shallow as 20 m (Helly & Levin, 2004) and the anoxic conditions recorded within the OMZ suppress macrobenthic diversity in the region (Levin et al., 2002, 2003; Palma et al., 2005). The importance of the OMZ in shaping large-scale diversity patterns in the northern region is suggested by the fact that bathymetric distributions of species are mostly restricted to inter-tidal habitats (Fig. 2d). Notably, the prevalence of intertidal over subtidal species is reversed at c. 37°S, precisely the southern limit of the shallow OMZ in the south-eastern Pacific (Helly & Levin, 2004). The shallow OMZ has operated at least since the early Pleistocene along the PSA (Emeis et al., 1991), so it is possible that the low incidence of species inhabiting deeper waters may have existed since several million years ago. Additionally, the strong impact of EN at physiological, population and ecosystem levels (Thiel et al., 2007) may also suppress the intertidal species richness north of 35°S (Fig. 2d), particularly via thermal stress. EN may have existed in the region since the Pliocene (Mohtadi et al., 2006; Ragaini et al., 2008), and therefore it may be forcing the shallow water diversity not only at ecological but also at evolutionary time-scales. In the southern region, the advances of ice lobes should have covered the entire coast during glacial times (Glasser et al., 2008), forcing the extinction of many shallow water benthic species (Rivadeneira & Marquet, 2007); only eurybathic species may have endured during the harsher glacial times, finding refuge in deeper areas of the shelf or the slope. This idea is further supported by two findings: (1) deeper bathymetric distributions in the southern region (Fig. 2c) and the prevalence of subtidal forms (Fig. 2d), and (2) the positive relationship between richness and the mean bathymetric range of species (Fig. 3d).

CONCLUDING REMARKS

When studied in isolation, traditional hypotheses and approaches used to explain the LGR phenomenon offer little insight about the processes shaping spatial biodiversity patterns. The application of indirect tests evaluating the importance of ecological and historical/evolutionary hypotheses, coupled with information describing environmental variations over geological time-scales, offer a new integrative picture of the processes moulding the inverse gradient of peracarid species richness. We suggest that the inverse LGR of peracarid species may be explained by processes operating at long temporal scales, mixing elements of historical and evolutionary hypotheses (sensu Midtibach et al., 2007) that we term the ‘out of the deep south’ hypothesis, which combines three key elements: (1) a high-latitude origin for many clades of peracarids with deep bathymetric distributions, (2) a strong taxonomic conservatism of the latitudinal distribution, restricting the dispersal of taxa towards lower latitudes, and (3) latitudinal variations in the evolutionary rates, with higher speciation rates and/or lower extinction rates towards higher latitudes.

Future phylogenetic and phyleogeographic analyses may validate these ideas. For instance, phylogenetic information of monophyletic taxa may be used to: (1) reconstruct the ancestral region of origin (Clark et al., 2008), validating the idea of a high-latitude (i.e. Antarctic) origin of the group; (2) evaluate the existence of phylogenetic conservatism in the geographic niche of the species, which can explain the limited dispersal towards lower latitudes (Wiens & Graham, 2005); (3) evaluate the timing of clade dispersal and its coincidence with the late Neogene global cooling; and (4) estimate diversification rates across the latitudinal gradient (Ricklefs, 2007). Since many of the patterns reported here may also hold in other taxa and regions exhibiting the inverse latitudinal gradient, the processes inferred may provide a general framework to understand the processes shaping large-scale patterns of marine biodiversity.

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REFERENCES


BIOSKETCHES

The authors are interested in understanding the processes that shape marine biodiversity. They have used the Chilean coast as model system. Using a range of approaches, including palaeontology, phylogeography, ecology, ethology and traditional taxonomy, they explore historical and current processes. Their studies incorporate a wide diversity of taxa in the quest to uncover the factors influencing the distribution of these organisms along the Chilean coast.

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