INTRODUCTION

The enormous geographic ranges of some species that would otherwise be expected to have low dispersal abilities have intrigued biologists for well over a century (see Darwin 1845, Johannesson 1988). The acceptance of the plate tectonic theory in the late 1960s inspired largely vicariant explanations of such broad distributions (e.g. Garbary 1987, Chin et al. 1991), with many historical biogeographers tending towards the view that species distributions are driven more by long-term geological processes than by dispersal events (Nelson & Platnick 1981, Humphries & Parenti 1999, Ebach & Tangney 2007). The increasing use of molecular data in biogeographical research has, however, demonstrated that general (multitaxon) biogeographical patterns can be generated via long-distance oceanic dispersal events, if the mechanism allowing for dispersal operates steadily over long periods (see review by Sanmartín et al. 2007).

In light of new molecular phylogenetic evidence for passive oceanic rafting as a mechanism for historic long-distance dispersal (Waters & Roy 2004, de Queiroz 2005, Donald et al. 2005, Gordillo 2006, Waters 2008), we predict that the potential of rafting macroalgae to shape composition of island biotas may be substantial. By examining patterns of intraspecific genetic variation among communities that have a biological propensity...
for passive dispersal, and no means for active dispersal, it may be possible to elucidate the ecological and evolutionary importance of long-distance rafting (Edgar 1987, Helmuth et al. 1994, Thiel & Haye 2006).

Ecological (Ingolfsson 1995, Edgar & Burton 2000, Macaya et al. 2005) and genetic (Coyer et al. 2001, Muhlin et al. 2008, Muhlin & Brawley 2009) studies of buoyant macroalgae suggest that macroalgal rafting may be an important process in the generation and distribution of biodiversity in marine systems. Long-distance dispersal of intertidal organisms by rafting on detached macroalgae may be especially significant in the subantarctic marine regions, where a series of oceanic and continental islands separated by hundreds to thousands of km of open ocean lie amid the powerful Antarctic Circumpolar Current (ACC; Fig. 1). At speeds measured for the ACC (20 to 40 cm s⁻¹; Hoffmann 1985), an 8000 km rafting journey from Campbell Island to Falkland Islands, for example, would take ca. 1 yr.

*Durvillaea antarctica* (Chamisso) Hariot (Phaeophyceae) or southern bull kelp (‘bull kelp’ for brevity) presents an ideal system for studies of rafting biology. Exposed subantarctic coastlines are characterised by dense beds of this robust and highly buoyant kelp. Detached *D. antarctica* have been inferred to float for at least 5000 km in the ACC (Dartnall 1974). The ACC has been estimated to have 20 million *D. antarctica* plants adrift with intact holdfasts at any one time (Smith 2002). Detached *D. antarctica* plants cannot re-attach, but likely continue to grow and remain reproductively viable while afloat, as observed for other buoyant phaeophycean taxa (e.g. *Macrocystis*, Macaya et al. 2005). The large, domed holdfasts of *D. antarctica* (Fig. 2a) support high biodiversity when attached to the rocky substrate (Morton & Miller 1968, Smith & Simpson 2002), with many crustacean, molluscan, annelid and echinoderm species exploiting them as a habitat and food source (Edgar & Burton 2000). Modification of a solid holdfast into a habitat for invertebrates often begins with the creation of intricate chambers by *Limnoria* isopods (Fig. 2a). Members of several other widespread crustacean (e.g. *Parawaldeckia*, *Hyale*; Smith & Simpson 2002) and molluscan (e.g. *Kerguelenella*, *Margarella*, *Onchidella*; Morton & Miller 1968) genera may then enter and hollow out the holdfast further, reducing the extent of attachment to the rocky substrate, and presumably increasing the probability of holdfast detachment and subsequent drifting.

Although dispersal via rafting is by no means a new idea (e.g. Wheeler 1916, Heatwole & Levins 1972, Highsmith 1985), molecular genetic markers have not been previously used to assess connectivity in macroalgal epifauna on a circumpolar scale. We here examine phylogeographic structuring across the subantarctic in 2 widespread peracarid crustacean taxa that dwell within bull kelp holdfasts: *Limnoria stephenseni* Menzies, 1957 (Isopoda: Limnoriidae), and *Parawaldeckia kidderi* (Smith, 1876) (Amphipoda: Lysianassidae) (Fig. 2b,c). We compare and contrast their mtDNA phylogeography to previously published data (Fraser et al. 2009) on their macroalgal host, *Durvillaea antarctica*.

The 2 invertebrate species analysed here are abundant and widespread members of the subantarctic epifaunal community of *Durvillaea antarctica* (Edgar 1987, Smith & Simpson 2002). As they are sedentary and produce direct-developing offspring, rafting is presumably their primary means of dispersing to and between remote oceanic islands. Indeed, individuals of these taxa are unlikely to move more than a few metres unassisted (Eltringham & Hockley 1961,
Rafting, on the other hand, provides potential for long-distance dispersal, and both genera have been observed in the holdfasts of beach-cast specimens of *D. antarctica* in New Zealand (J. M. W. pers. obs.). Both of these epifaunal species can also exploit holdfasts of other large buoyant seaweeds such as *Macrocystis pyrifera* (Edgar 1987) and *Lessonia* spp. (Cookson 1991). While the other presumptive macroalgal host species are similarly broadly distributed across the subantarctic (Dhargalkar & Verlecar 2009), *D. antarctica* is likely to play a key role in rafting as its rafts are the most abundant and longest-lived at sea (Smith 2002). Unlike some of its congeners, *Limnoria stephenseni* has never been found on other buoyant substrates (e.g. wood).

A recent global phylogeographic survey of *Durvillaea antarctica* inferred that most subantarctic populations of the species were re-established after the Last Glacial Maximum, most plausibly following elimination by sea ice that scoured the intertidal zone (Fraser et al. 2009). Obligate epifauna of intertidal macroalgae are predicted to have experienced a similar demise and to have relied on rafting on macroalgae in their recolonization of the subantarctic islands. These historical hypotheses would be supported by obligate kelp-holdfast dwelling invertebrates (*Limnoria stephenseni, Parawaldeckia kinderi* showing (1) little genetic variation across the far-flung subantarctic intertidal habitats, and (2) phylogeographic structuring across the subantarctic similar to that of their macroalgal host (*D. antarctica*).

**MATERIALS AND METHODS**

**Sample collection.** Holdfast epifaunal samples were collected from bull kelp beds on 6 oceanic islands located in ‘subantarctic’ waters south of 46° S in the South Atlantic, South Indian and South Pacific Oceans: Falkland Islands, Marion Island (Prince Edward Island group), Possession Island (in the Crozet Archipelago), and Kerguelen, Macquarie and Campbell Islands (Fig. 1, Tables 1 & 2). Animals were collected from whole holdfasts that were detached from their rocky substrate during low tide using a hatchet. The numbers of *Parawaldeckia kinderi* in a typical holdfast were generally substantially larger than numbers of
Limnoria stephensi. Samples were preserved directly in 95% ethanol and stored at –20°C prior to analysis. Samples from different holdfasts were usually stored separately; the exceptions, where some specimens came from pooled samples of fauna from several holdfasts, are indicated in Table 1. Individuals included in genetic analyses were typically selected from different holdfasts to minimise the sampling of closely related individuals (e.g. siblings from the same mother). In some cases, however, 2 or 3 different size classes of individuals from within a single holdfast were sampled to ensure adequate sample sizes.

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<table>
<thead>
<tr>
<th>Island/archipelago</th>
<th>Collection locality</th>
<th>Min. age of the island(s)</th>
<th>Acc. nos. <em>Parawaldeckia kidderi</em></th>
<th>Acc. nos. <em>Limnoria stephensi</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Falkland Islands</td>
<td>Kidney Island and Cape Pembroke</td>
<td>2500 Myr^a^</td>
<td>FJ608920, FJ608921^b^</td>
<td>FJ608831–FJ608833^b^</td>
</tr>
<tr>
<td></td>
<td>New Island, SE Point</td>
<td></td>
<td>FJ608834, FJ608835^b^</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sea Lion Island</td>
<td></td>
<td>FJ608836–FJ608838</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Surf Beach</td>
<td></td>
<td>FJ608931–FJ608934^b^</td>
<td>FJ608839–FJ608840^b^</td>
</tr>
<tr>
<td>Marion Island, Prince Edward Islands</td>
<td>Boulder Beach, Transvaal Cove</td>
<td>450 kyr</td>
<td>FJ608935–FJ608942</td>
<td>FJ608847–FJ608855</td>
</tr>
<tr>
<td></td>
<td>King Bird Head</td>
<td></td>
<td>FJ608943–FJ608945</td>
<td>FJ608856–FJ608863</td>
</tr>
<tr>
<td>Possession Island, Crozet Archipelago</td>
<td>Baie du Marin</td>
<td>8.1 Myr</td>
<td>FJ608946–FJ608957</td>
<td>FJ608864–FJ608874</td>
</tr>
<tr>
<td></td>
<td>Jardin Japonais</td>
<td></td>
<td></td>
<td>FJ608875</td>
</tr>
<tr>
<td>Kerguelen Islands</td>
<td>Port-Aux-Français</td>
<td>1840 Myr^a^</td>
<td>FJ608958–FJ608966</td>
<td>FJ608876–FJ608889</td>
</tr>
<tr>
<td>Macquarie Island</td>
<td>Garden Cove</td>
<td>80–300 kyr</td>
<td>FJ608967, FJ608968^b^</td>
<td>FJ608890–FJ608903</td>
</tr>
<tr>
<td></td>
<td>ANARE Station</td>
<td></td>
<td>FJ608969–FJ608981^b^</td>
<td>FJ608904</td>
</tr>
<tr>
<td>Campbell Island</td>
<td>Perserverance Harbour</td>
<td>16 Myr</td>
<td></td>
<td>FJ608905–FJ608919</td>
</tr>
</tbody>
</table>

^a At the time of their formation, the land masses of these islands were not in their present day positions owing to plate tectonic dynamics.

^b Some or all of the analyzed specimens were selected from pooled faunal samples from several holdfasts. Details given in GenBank entries.

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**Table 2.** *Parawaldeckia kidderi* and *Limnoria stephensi*. Bull kelp epifauna cytochrome oxidase subunit (COI) sequence sample sizes (N), and numbers of holdfasts (N_HF) from which the sequenced animals were taken. Length of analysed mtDNA COI fragment is given. FI: Falkland Islands. MI: Marion Island, PO: Possession Island, KE: Kerguelen Islands, MQ: Macquarie Island, CA: Campbell Island. See Fig. 1 for geographical locations of the islands.

<table>
<thead>
<tr>
<th>Taxon</th>
<th>COI bp</th>
<th>FI</th>
<th>MI</th>
<th>PO</th>
<th>KE</th>
<th>MQ</th>
<th>CA</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Limnoria stephensi</em></td>
<td>912</td>
<td>N</td>
<td>16</td>
<td>17</td>
<td>12</td>
<td>14</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td></td>
<td>N_HF</td>
<td>≥6</td>
<td>7</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td><em>Parawaldeckia kidderi</em></td>
<td>980</td>
<td>N</td>
<td>15</td>
<td>11</td>
<td>12</td>
<td>9</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td></td>
<td>N_HF</td>
<td>≥7</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>2</td>
<td>2</td>
</tr>
</tbody>
</table>

^a P. kidderi was not found in Campbell Island; another *Parawaldeckia* species occupied the sampled holdfasts. Uncorrected COI sequence divergence between the 2 species was 8.8%.^b^ Analysis samples were picked from 2 samples that together contained animals from >10 different holdfasts.
Preparation of data. We extracted total genomic DNA from leg or pleotelson muscle tissue of the invertebrates using a standard Chelex method (Walsh et al. 1991). A fragment of the mitochondrial gene coding for cytochrome oxidase subunit I (COI) was amplified using the universal invertebrate primers LCO1490 (5’-GGT CAA CAA ATC ATA AGA TAT TGG-3’) (Folmer et al. 1994) and H7005mod1 (5’-ART GNG CNA CNA CRT ART ANG TRT CRT G-3’) (Donald et al. 2005), under the following conditions in an Eppendorf Mastercycler: initial denaturation phase of 2 min at 95°C; 40 cycles of 50 s at 95°C, 1 min 30 s at 48°C, 1 min 30 s at 72°C; final extension step of 10 min at 72°C. PCR products were purified using the Invitrogen Purelink PCR purification kit, and sequenced by the Allan Wilson Centre Genome Service. The primer LCO1490 was used for sequencing COI of Parawaldeckia kidderi. Sequencing of Limnoria stephenseni COI was done with an internal forward primer (5’-TCATTC GGC TTG AGC TAG GGC A-3’) that was designed based on preliminary sequences obtained using the universal primers (above). Numbers of sampled individuals and holdfasts per island and species are listed in Table 2. The sequences were deposited in GenBank under accession numbers FJ608831–FJ608981. Details of sampling localities and GenBank accessions per species per locality are presented in Table 1.

Data analysis. Sequence chromatograms were inspected, edited, aligned and the amino acid translation of the sequences checked using Sequencher 4.8 software (GeneCodes). The lengths of the homologous sequence fragments, qualified for final analysis by having complete data across all studied individuals, were 912 and 980 nucleotide base pairs in having complete data across all studied individuals, were (GeneCodes). The lengths of the homologous sequences checked using Sequencher 4.8 soft-
suspected, edited, aligned and the amino acid translation per locality are presented in Table 1.

RESULTS

Diversity and distribution of mtDNA COI haplotypes

We detected a single haplotype with a circumpolar distribution in both epifaunal crustacean species (Fig. 3a,b). In each case, the circumpolar haplotype also occupied a central position in the inferred statistical parsimony network and was the most common haplotype, carried by 38% of Parawaldeckia kidderi and 67% of the Limnoria stephenseni specimens. In both species, within-island polymorphism consisted mostly of low-frequency haplotypes inferred to have descended directly from the ancestral circumpolar haplotypes, producing a star-like network. All but one of the rare haplotypes were restricted to single islands, and they were seldom more than 2 mutational steps divergent from the common circumpolar haplotype. Overall diversity was higher in P. kidderi than in L. stephenseni, as shown by the haplotype networks (Fig. 3a,b) and molecular diversity indices (Table 3). Haplotypic diversity was higher in the epifaunal invertebrate samples than previously observed in their host Durvillaea antarctica.

The phylogeographic patterns of the 2 invertebrate species differed substantially in the case of the Falkland Islands, where Limnoria stephenseni had 3 rare haplotypes along with the circumpolar haplotype, but Parawaldeckia kidderi samples completely lacked the circumpolar haplotype. Instead, the sampled Falkland P. kidderi haplotypes were between 2 and 13 mutational steps divergent from the circumpolar haplotype, and a total of 4 distinct haplotype lineages were present in the sample. Of these lineages, 2 were closely related to the circumpolar haplotype and to the rare haplotypes found on the other islands, whereas the other 2 were ~1.3% different from it and from each
other (uncorrected COI divergence). Notably, both of these divergent \textit{P. kidderi} lineages were found in samples collected from the ACC-exposed Sea Lion Island, whereas the lineages closer to non-Falkland haplotypes dominated the more sheltered locations (Surf Beach, Cape Pembroke). In contrast to the broad circumpolar haplotype sharing, \textit{L. stephenseni} specimens from Campbell Island did not share haplotypes with other subantarctic island samples, but instead, had a unique haplotype group that was one mutational step away from the circumpolar haplotype; the same feature was found in \textit{Durvillaea antarctica} (Fig. 3c).


differentiation between islands and indicators of population expansion

Fixation index values and their tests of significance (Table 4) reveal the Falkland Islands’ \textit{Parawaldeckia kidderi} sample as significantly different from all other island samples; no statistically significant differentiation was found among any other island samples in \textit{P. kidderi}. In the \textit{Limnoria stephenseni} data set, the Kerguelen and Campbell Islands’ samples were significantly different from those of all other islands and from one another, whereas for \textit{Durvillaea antarctica}, only Campbell Island populations were significantly different from all others. No significant differentiation was inferred for any of the 3 species among samples from Marion or Possession and Macquarie Islands (distance ≥7400 km) or among samples from Marion and Possession islands (distance ~1000 km) (Table 4).

According to the hierarchical analysis of molecular variance (Table 3), the majority of the mtDNA variability of \textit{Limnoria stephenseni} (65\%) was distributed among islands. In contrast, most genetic variability in \textit{Parawaldeckia kidderi} (85\%) was attributable to within-island variation, reflecting the relatively high within-island polymorphism in this species. All AMOVA results were statistically highly significant [p (random value
of among-population variation ≥ observed value) < 0.0001]. The highly divergent lineages encountered only in the Falkland Islands accounted for all among-island variability in *P. kidderi*, as shown by the effect of their exclusion from analysis of molecular variance (AMOVA) (Table 3). This finding conforms with the *F*_{ST} analysis that found no significant differentiation among island-pairs that did not include Falkland Islands. The high proportion of among-island variability in *L. stephenseni* was due to the haplotype group exclusive to the Campbell Island sample and to the relatively low frequency of the circumpolar haplotype in the Kerguelen Island as compared to the other populations carrying it.

Fu’s *F*_{S} statistic gave statistically significant negative values for both the epifaunal species and their host when estimated across all subantarctic samples (Table 3). *Parawaldeckia kidderi* samples from the Falkland Islands, however, yielded a positive *F*_{S} statistic value when analysed separately. Thus, with the exception of *P. kidderi* from the Falkland Islands, mtDNA diversity patterns of the kelp holdfast-dwelling species are consistent with the scenario of historical population expansion.

### DISCUSSION

Our chief finding of circumpolar haplotypes in *Durvillaea antarctica* and its direct-developing epifaunal invertebrates, across some of the world’s largest longitudinal stretches of open ocean, argues in favour of macroalgal rafting in the ACC as an important long-distance dispersal mechanism for intertidal kelp epifauna in the subantarctic. More broadly, macroalgal rafting may be a significant ecological factor explaining similarities in species composition of marine intertidal communities across the subantarctic landmasses affected by the ACC.

#### Comparison of phylogeographic patterns in southern bull kelp and its epifaunal invertebrates

The level of mtDNA diversity detected among populations of *Parawaldeckia kidderi* and *Limnoria stephenseni* is consistent with each taxon representing a single species across the subantarctic, considering the distribution of COI divergences observed between congeneric crustacean species (Lefebure et al. 2006). Indeed, subantarctic populations of both of these epifaunal invertebrate taxa display high frequencies of a single circumpolar haplotype. Across 5 of the 6 studied islands, the invertebrates also exhibit low phylogenetic diversity that consists of rare haplotypes descended from the common circumpolar haplotype. With the sole exception of Falkland Islands *P. kidderi*, these epifaunal phylogeographic patterns are highly similar to that observed for their macroalgal host (Fraser et al. 2009). An additional concordant pattern detected across taxa is the presence of a distinct Campbell Island haplotype group in both bull kelp and the isopod *L. stephenseni* (Fig. 3a,c). Finally, all 3 species exhibit genetic signatures of population expansion (neutrality tests), supporting the hypothesis of rapid historical population growth in the circumpolar region.

The strong phylogeographic similarities evident for subantarctic epifaunal species and bull kelp imply that all 3 taxa experienced similar postglacial recolonization histories in at least 5 out of the 6 studied subantarctic islands. Although alternative possible mechanisms exist, e.g. rafting on other substrates (reviewed by Thiel & Gutow 2005) and anthropogenic introductions, they are relatively unlikely. For instance, *Limnoria stephenseni* and *Parawaldeckia kidderi* are not capable of fouling ship hulls, and their successful anthropogenic introduction would likely have required shipping of whole macroalgae with holdfasts. Moreover, *Durvillaea antarctica* eggs are unlikely to survive more than a few days at sea (Hay 1994), so would be unable to complete ocean crossings in ship ballast; additionally, historic records of bull kelp from the subantarctic islands predate modern ships using

<table>
<thead>
<tr>
<th>Falkland</th>
<th>Marion</th>
<th>Possession</th>
<th>Kerguelen</th>
<th>Macquarie</th>
<th>Campbell</th>
</tr>
</thead>
<tbody>
<tr>
<td>Falkland</td>
<td>0.00 ns</td>
<td>0.00 ns</td>
<td>0.00 ns</td>
<td>0.00 ns</td>
<td>0.87***</td>
</tr>
<tr>
<td>Marion</td>
<td>0.00 ns</td>
<td>0.00 ns</td>
<td>0.00 ns</td>
<td>0.00 ns</td>
<td>0.86***</td>
</tr>
<tr>
<td>Possession</td>
<td>−0.01 ns</td>
<td>0.00 ns</td>
<td>0.00 ns</td>
<td>0.00 ns</td>
<td>0.84***</td>
</tr>
<tr>
<td>Kerguelen</td>
<td>0.44***</td>
<td>0.64***</td>
<td>0.61**</td>
<td>0.00 ns</td>
<td>0.81***</td>
</tr>
<tr>
<td>Macquarie</td>
<td>−0.01 ns</td>
<td>0.01 ns</td>
<td>−0.01 ns</td>
<td>0.51***</td>
<td>0.81***</td>
</tr>
<tr>
<td>Campbell</td>
<td>0.73***</td>
<td>0.90***</td>
<td>0.88***</td>
<td>0.79***</td>
<td>0.80***</td>
</tr>
</tbody>
</table>

Table 4. *Durvillaea antarctica*, *Limnoria stephenseni* and *Parawaldeckia kidderi*. Pairwise *F*_{ST} values between island samples of *D. antarctica* (unshaded), *L. stephenseni* (light-shaded) and *P. kidderi* (dark-shaded) calculated from nucleotide differences. Significance level of values against 1023 random permutations of sequence samples among islands is given (*p < 0.05, **p < 0.01, ***p < 0.001, ns = not significant). The significance (+) and nonsignificance (–) of pairwise differentiation in island samples of *D. antarctica/L. stephenseni/P. kidderi* are summed above diagonal in the lower part of the table; nd = no data.
ballast water (e.g. Choris 1822). On the other hand, as these epifaunal invertebrates are obligate kelp-dwellers, macroalgal rafting is clearly the most plausible mechanism to explain their recent circumpolar dispersal. The probability of an individual rafting event following kelp detachment may be low (Miranda & Thiel 2008), but the apparent abundance of rafts in the subantarctic (Smith 2002) may make such passive dispersal events inevitable over long timescales.

Coalescence theory predicts that populations that have a long uninterrupted history exhibit higher mtDNA nucleotide diversity (further differentiated haplotypes) than populations with short histories (Tajima 1983). Therefore, low genetic diversity and broad genetic similarity of subantarctic bull-kelp populations is thought to reflect postglacial recolonization (Fraser et al. 2009). Although the epifaunal data presented here broadly support postglacial recolonization, the relatively high mtDNA diversity detected in Falkland Islands *Parawaldeckia kidderi* suggests that these islands may have represented a glacial refuge for this species. The common ancestor of the lineages found at Falkland Islands can be inferred to predate the Last Glacial Maximum unless substitutions in the COI-gene of *P. kidderi* occur at a rate 3 times faster than the fastest short-term COI substitution rate suggested for crustaceans (Audzijonyte & Väinölä 2006).

For the subantarctic islands whose bull kelp and invertebrate COI variation have both been assessed, the recorded haplotype and nucleotide diversity values are lowest in bull kelp (5 island samples fixed for the circumpolar haplotype), whereas most epifaunal invertebrate samples include a few rare haplotypes (Fig. 3a–c). The singleton kelp COI haplotype FJ550100 reported from Possession (Crozet) Island by Fraser et al. (2009) was found to be an artefact introduced by erroneous base calling, indicating that only the Campbell Island kelp samples exhibit mtDNA diversity. Under neutrality, and assuming that the subantarctic bull kelp and epifaunal populations are of the same postglacial age, this discrepancy in diversity between species might simply be explained by different circumpolar effective population sizes of the taxa. Population genetic theory predicts that, in the absence of selective sweeps, nucleotide diversity (π) of a population at mutation-drift equilibrium is the product (π = 4N_eμμ; Tajima 1983) of the mutation rate of DNA (μ) and the effective population size (N_e). Alternatively, the discrepancy might reflect variation in COI mutation rate between the species.

**Subantarctic biogeography**

Intraspecific genetic diversity of Southern Hemisphere biota remains an understudied topic (Behere-garay 2008), and the subantarctic intertidal zone is particularly poorly studied. Nevertheless, this southern region has distinctive features (sparse intertidal habitat, powerful circumpolar current) that distinguish it from its Northern Hemisphere (Arctic) equivalent, offering a unique setting for studies of long-distance dispersal. In the current study, the genetic similarity of isolated subantarctic populations clearly contrasts with the relatively old geological ages suggested for many of these islands (see Table 1), and apparently conflicts with ‘multiregional’ models proposed for subantarctic biota (e.g. Greve et al. 2005). On the other hand, the common circumpolar genetic similarity of the bull kelp community supports Udvardy’s (1987) hypothesis of a circumpolar ‘Insulantarctica’ province.

Comparisons with taxa that live in terrestrial habitats — species that are clearly not prone to dispersal by macroalgal rafting — can provide an informative context within which to interpret the findings of the current study. On the one hand, the circumpolar haplotype sharing and low nucleotide diversity of the bull kelp community in most of the studied islands contrasts with the relatively high between-island genetic differentiation and within-island diversity documented for subantarctic populations of weevils (Grobler et al. 2006), springtails (Stevens et al. 2006), and Antarctic hairgrass (van de Wouw et al. 2008). On the other hand, the widespread distribution of a single mitochondrial haplotype in the subantarctic bull kelp and its epifaunal species resembles that of anthropogenically introduced terrestrial species on subantarctic islands (Myburgh et al. 2007). This similarity apparently emphasizes the effectiveness of kelp rafts as dispersal vectors. It appears that natural macroalgal rafting was able to induce an early postglacial genetic homogeneity across the circumpolar region, comparable to human-mediated homogeneity brought about by recent global-scale introductions via ship traffic in other parts of the globe (e.g. Voisin et al. 2005). Also, the contemporary mtDNA haplotype sharing patterns in *Limnoria stephenseni* and *Parawaldeckia kidderi* are similar to a ribbon worm, *Parborlasia corrugatus*, whose long-lived planktonic larvae presumably travel along the ACC between subantarctic islands (Thornhill et al. 2008).

Hypothetically, postglacial recolonization of the subantarctic islands by the bull kelp community could have been mediated by rafting from a single geographical source with subsequent ‘stepping-stone’ dispersal eastward via the ACC, from one island to the next. Such a history would yield broad genetic homogeneity, as observed here in the form of geographically widespread, numerically dominant mtDNA haplotypes. Specifically, the essentially ubiquitous circumpolar haplotypes observed for each species we looked
at suggest that their postglacial maternal ancestries can be traced back to single source populations. Although we have sampled most of the known distribu-
tional ranges of *Limnoria stephensi* and *Parawadal
dekia kidderi*, neither of which occurs in mainland New Zealand, we are presently unable to confidently identify a source region for postglacial recolonization.

Acknowledgements. We are grateful to the persons and organisations that helped us in sample collection, including: I. Ansorge (University of Cape Town) and the South African National Antarctic Program; M. Lebourvier, S. Gutjahr, S. Mal-
loj, J. P. Orts and D. Renault (French Polar Institute, IPY pro-
gramme 136); A. Wiebekin, J. Doube and R. Clifton (Australian Antarctic Division); P. Brickle and J. Brown (Department of Fisheries, Falkland Islands) and I. Strange; S. Banks (Depart-
ment of Conservation, New Zealand) and G. Wilson (Univer-
sity of Otago); Heritage Expeditions. We thank L. J. Cookson and G. Penwick for taxonomic identifications of isopods and amphipods, respectively. T. King is acknowledged for assis-
tance and advice in the laboratory. K. Miller kindly compiled Fig. 2 from photographs taken by C.I.F. We thank anonymous reviewers for their helpful comments on the manuscript. Resources for our work were provided by the Marsden Fund (contract 07-UOO-099), Shackleton Scholarship Fund, Department of Zoology in the University of Otago and Allan Wilson Centre for Molecular Ecology and Evolution.

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Submitted: September 29, 2009; Accepted: January 1, 2010

Proofs received from author(s): April 22, 2010