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Migration history and stock structure of two putatively diadromous teleost fishes, as determined by genetic and otolith chemistry analyses

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Abstract: Migratory life cycles and population structure of 2 putatively diadromous Australian fishes were examined using otolith chemistry ($^{87}$Sr/$^{86}$Sr) and genetics (microsatellites and mitochondrial deoxyribonucleic acid [mtDNA]). Australian whitebait (Lovettia sealii) is widely considered to be one of only a few anadromous fish species in the southern hemisphere. The congolli or tupong (Pseudaphritis urvillii) is reported to undertake an unusual form of sexually segregated catadromous migration, where females switch habitats between marine and freshwater, while males remain in marine or estuarine environments. Sr-isotope profiles of L. sealii showed this species does not move into fully freshwater habitats during its life cycle, suggesting it should be considered semi-anadromous or estuarine-dependent, rather than truly anadromous. This life-history strategy is unique among the Galaxiidae. Lovettia sealii is regionally divided into at least 3 well differentiated genetic stocks: northern and southern Tasmanian coasts and mainland Australia. Sr-isotope profiles of P. urvillii showed that females are catadromous, with the early life history spent in the marine environment and a single migratory transition from marine to freshwater occurring at an early point in the life history. Lack of bidirectional adult migration between freshwater and the sea suggests that female P. urvillii are semelparous, returning to the marine habitat to mate with resident males after an extended period of freshwater residence. Pseudaphritis exhibit weak genetic structure across their mainland range. An isolation-by-distance relationship describes the genetic structure of this species, a pattern it shares with several other nearshore-restricted catadromous fishes.

Key words: Microsatellites, mitochondrial DNA, otolith chemistry, Sr isotopes, migration, standardized FST, Pseudaphritidae, Galaxiidae

Diadromy is a habitat-switching life-history strategy involving migration between freshwater and marine waters for completion of the life cycle (Myers 1949). Diadromous migration provides opportunity for connectivity among subpopulations via the ocean, but the magnitude and spatial configuration of this connectivity is variable and influenced by many factors. Some of these include the form of diadromy exhibited by a species (e.g., anadromy, catadromy, amphidromy; McDowall 1988), the existence of philopatric behavior (e.g., natal homing; Vaha et al. 2007), environmental and ecological variables (e.g., coastal distance and temperature regime; Dionne et al. 2008), and historical events (e.g., glaciation and sea-level fluctuation may subdivide a species into demographically independent stocks; King et al. 2001). The widespread distribution of many diadromous taxa is attributable to oceanic connectivity (McDowall 2008). More-limited distributions of some species may be explained, at least in part, by shorter larval durations in the ocean and distance travelled from shore (Sorensen and Hobson 2005, Lord et al. 2012).
The Australian native fish fauna includes ~33 diadromous species, most of which are poorly understood in terms of migratory movements and population structure (McDowall 1988, Miles 2007). Catadromy and amphidromy are the most common forms of diadromy in Australian fishes, whereas the dominant form in the northern hemisphere (anadromy; McDowall 2008) is recognized in only 4 Australian species (Lovettia sealii, Retropinna tasmanica, Geotria australis, and Mordacia mordax; McDowall 1988, Miles 2007). Catadromy involves migration of adults from freshwater reaches to marine habitat for the purpose of reproduction. Juveniles then return to freshwater for growth and development (McDowall 1988). Anadromy is the reverse of catadromy, so that adults reside in the marine environment and migrate to freshwater for breeding (McDowall 1988). In amphidromy, the switch between habitats is not for the purpose of reproduction. Instead, adults reside and reproduce in freshwater and larvae migrate passively to the marine habitat for a period of early growth before returning to freshwater as juveniles (McDowall 1988).

The current deficient state of knowledge of Australian diadromous fishes, at least in part, reflects the difficulty of monitoring movements of small larval and juvenile fish that can travel potentially great distances between different habitats. Otolith chemistry is an indirect method well suited to overcome this problem and can be used to estimate the movement of an individual fish across different chemical environments throughout its lifetime (Gillanders 2005, Walther and Limburg 2012). Otoliths (earstones) grow continuously and provide a permanent record of environmental conditions experienced by a fish throughout its life that can be recovered by measuring particular elemental (e.g., Sr/Ca, Ba/Ca) or isotopic (87Sr/86Sr) ratios that differ among habitats (Gillanders 2005, Walther and Limburg 2012). Therefore, otolith chemistry is useful for discerning whether diadromy occurs in a species or population, for determining which form of diadromy occurs, and for assessing whether variation exists among individuals or populations in the incidence of diadromous migration (Milton and Chenery 2005, Crook et al. 2008, Chapman et al. 2012).

Otolith chemistry can reveal within-lifetime migration patterns, but estimates of population connectivity require methods that integrate movements of successful dispersers over many generations. Genetic data is complementary to otolith chemistry in this regard. Inferences from genetic analyses may be applied over large spatial scales to infer stock structure and evolutionary history (Waples et al. 2008, Griffiths et al. 2010, Feutry et al. 2013) or over small scales to infer demographic parameters, such as population size and migration rate (Lowe and Allendorf 2010). Genetic studies of anadromous fishes indicate that most species exhibit an isolation-by-distance (IBD) pattern of genetic connectivity at scales of up to several thousand kilometers of coastline (Bradbury and Bentzen 2007). IBD is a positive association between genetic and geographic distance in which subpopulations are linked by dispersal in a stepping-stone fashion. This pattern probably is a product of limited dispersal resulting from natal homing, which is almost ubiquitous among anadromous fishes (McDowall 2001). Lampreys are one of the few exceptions to natal homing in this group, and they exhibit low levels of genetic structure and very weak IBD relative to other anadromous taxa (Waldman et al. 2008, Spice et al. 2012).

Catadromous fish encompass several patterns of genetic connectivity including panmixia, the absence of genetic structure because of random gene flow among demes. Such a pattern exists in anguillid eels, where all spawning in a species occurs within a single region of the ocean and the return juvenile migration may involve distances of >5000 km (Als et al. 2011). Catadromous fishes that breed in estuaries or inshore waters often show low levels of structuring with an IBD pattern of population connectivity (Keenan 1994, Shaddick et al. 2011). This pattern probably indicates that juvenile migration back into rivers is not random but is spatially restricted to some extent, either by limited pelagic larval duration or retention of juveniles within the natal estuary (see also Feutry et al. 2013). Amphidromous taxa show the weakest level of population structure among diadromous species. Most are either panmictic or have very low levels of population structure, regardless of whether they occur in the tropics or have temperate continental distributions (Crandall et al. 2010, Schmidt et al. 2011, Cook et al. 2012). Genetic structuring detected at large spatial scales for some widespread amphidromous species is attributed to biogeographic history and does not reflect contemporary patterns of migratory behavior in these taxa (e.g., Waters et al. 2000, Lord et al. 2012).

Coupling otolith chemistry with genetic analysis is a promising approach for understanding the ecological, evolutionary, and conservation implications of diadromous migration. These methods were applied to 2 diadromous fish species endemic to temperate southeastern Australia. Our goals were to assess: 1) whether transects of Sr isotope (87Sr/86Sr) ratios through otoliths of Lovettia sealii (Johnston) and Pseudaphritis urvillii (Valenciennes) (Pseudaphritidae) are consistent with habitat switching between freshwater and marine environments by these species, 2) whether migratory behavior varies among or within populations, 3) whether multiple stocks can be discriminated within each species using genetic data, and 4) to define a spatial pattern of genetic connectivity among subpopulations.

METHODS

Study species

The Australian whitebait L. sealii is a small, slender-bodied monotypic member of the Galaxiidae, formerly only known from coastal rivers of Tasmania until recently discovered across Bass Strait at a single locality on mainland

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Australia (Allen et al. 2002, Raadik 2008a). Lovettia sealii is classified as anadromous (McDowell 1988, Allen et al. 2002, Gillanders 2005, Miles 2007), but this classification is based on a single study (Blackburn 1950). Current understanding is that sexually maturing adults migrate upstream from the sea during the late austral winter to early summer, spawning occurs in upper estuarine reaches after which most adults deteriorate in condition and die, and larvae are then transported back toward the ocean (Blackburn 1950). Whether all larvae enter the open ocean and whether growth and development occur in the ocean or closer to shore have not been verified. However, an unpublished study suggested that some larvae remain within the estuary in Tasmania (Fulton and Pavuk 1988), and juveniles (<45 mm total length) have been recorded in April from within a coastal inlet on mainland Australia by one of us (TR, personal observation). Blackburn (1950) proposed that L. sealii consists of 2 independent stocks in northern and southern Australia based on differences in pigmentation, growth, timing of spawning runs, and distribution. Support for multiple stocks was provided by unpublished allozyme studies that resolved a well differentiated northern and southern region (Pavuk 1994). Fulton and Pavuk (1988) and Pavuk (1994) also proposed several other genetic stocks. However, these results may have been disproportionately influenced by a single locus (PEP A), and it is possible that this estimate of genetic structure was biased by selection. A significant commercial whitebait fishery existed in Tasmania from 1941 to 1974, and L. sealii made up ~95% of the catch (Blackburn 1950, Fulton 2000). Catches changed by the 1950s, leading to closure of the fishery, which reopened in 1990 for recreational purposes under limited seasonal operation (Fulton 2000). Many diadromous fishes have suffered similar precipitous declines over the last century because of overfishing, barriers to migration, and pollution (Limburg and Waldman 2009). A better understanding of population structure and migratory behavior of L. sealii is warranted for more informed management and conservation of the species, particularly considering its recently discovered trans-Bassian distribution.

Pseudaphritis urvillii (commonly known as congolli or tupong) is a non-Antarctic member of the icefish suborder Notothenioidei (Near et al. 2004) and is endemic to and common in coastal rivers of southeastern Australia, including Tasmania (Allen et al. 2002, Raadik 2008b). Pseudaphritis urvillii is currently classified as catadromous (McDowell 1988, Gillanders 2005, Miles 2007). An unusual sexually dimorphic residence behavior appears to exist where adult females reside in freshwater habitats and undertake downstream migration to the sea where spawning occurs (Crook et al. 2010). Males reside either in estuaries or the sea (Hortle 1979, BZ, personal observation). The spatial scale of movements made during the juvenile phase is unknown, and no information exists on population structure. Population declines may have occurred at range margins of the mainland distribution in South Australia and New South Wales (Miles 2007, Hammer et al. 2009). Barriers that inhibit migratory movements required to complete the life cycle are implicated in these declines. These barriers may include infrastructure, such as dams and weirs, or decreases in water level caused by drought and water abstraction (Rolls 2011, Wedderburn et al. 2012). For example, dramatic population declines of juvenile P. urvillii migrating upstream were observed during a recent drought and were attributed to failed recruitment caused by limited access of reproductively mature females to estuarine/marine spawning habitats (Zampatti et al. 2010).

Field sampling

Samples of L. sealii were collected using hand-held dip nets from the upper estuarine reaches of 5 coastal rivers, including 3 from northern Tasmania and 2 from southern Tasmania, and by trawl from 1 mainland site, Anderson Inlet (= Tarwin River), Victoria (used only for genetic analysis) (Fig. 1A, Table 1). The sample from Black River consisted of a single individual, which was used for otolith analysis but not for genetics. Female P. urvillii were sampled by electrofishing from the freshwater reaches of 8 coastal rivers upstream of tidal influence, including 7 mainland sites and 1 from northern Tasmania (Fig. 1B, Table 1). The sample from Hopkins River was used for otolith analysis but not for genetics. Where possible, 30 individuals of each species were sampled per site. All procedures were carried out according to Australian Ethics Commission protocol number ENV/01/09/AEC.

Water collection and analysis

Water samples were collected from the lower freshwater reaches of each study river for analysis of 87Sr/86Sr. At least 1 sample was collected for each location included in the otolith analysis and multiple (2–3) samples were analyzed from several localities to check for temporal variation in 87Sr/86Sr (Victoria: summer 2008/2009, winter 2009; Tasmania: autumn 2009, spring 2009). Based on repeat sampling of sites in our study and from other Tasmanian rivers (DC, unpublished data), temporal variation in 87Sr/86Sr was minor relative to among-site variation (Table 1), so a single water sample from each site was regarded as sufficient for the purpose of our study. Samples were collected in 125-mL acid-washed polyethylene bottles, refrigerated at 4°C, and transferred to the School of Earth Sciences, University of Melbourne. Twenty-milliliter aliquots of each water sample were filtered through a 0.2-μm Acrodisc syringe-mounted filter (Pall Corporation, Ann Arbor, Michigan) into a clean polystyrene beaker and dried overnight in a high efficiency particulate air (HEPA)-filtered fume cup.
board. Previous analyses have shown that filtering after transfer to the laboratory, rather than after sample collection in the field, has no influence on measurement of $^{87}\text{Sr}/^{86}\text{Sr}$ (e.g., Palmer and Edmond 1989).

Sr was extracted with a single pass over 0.15-mL ($4 \times 12$ mm) beds of EICHROM™ Sr resin (50–100 $\mu$m; Eichrom Technologies, Lisle, Illinois). Following Pin et al. (1994), matrix elements were washed off the resin with 2M and 7M HNO$_3$, followed by elution of clean Sr in 0.05M HNO$_3$. The total blank, including syringe-filtering, was $\leq 0.1$ ng, implying sample to blank ratios of $\geq 4000$, so no blank corrections were deemed necessary. Sr-isotope anal-

Figure 1. Sampling locations (black bolded waterways) and mitochondrial deoxyribonucleic acid (mtDNA) data summary for *Lovettia seallii* (A) and *Pseudaphritis urvillii* (B). Frequency pies depict relative frequency of mtDNA haplotypes in 3 arbitrary groups shaded black, grey, and white in network diagrams. Rivers without frequency pies were included in the otolith chemistry analysis but not genetics. Haplotype codes in mtDNA network diagrams correspond with data presented in Tables S1, S2.
Table 1. Summary of water, otolith, and genetic samples analyzed and genetic variation in populations of *Lovettia sealii* and *Pseudaphritis urvillii* based on mitochondrial deoxyribonucleic acid (mtDNA) and microsatellite markers. Water $^{87}$Sr/$^{86}$Sr values are point estimates or mean (± SD) where replicate samples were taken. $H_O$ = observed heterozygosity, $H_E$ = expected heterozygosity, $NA$ = total number of alleles, $AR$ = allelic richness (value in brackets is the standardized sample size).

<table>
<thead>
<tr>
<th>Species, river</th>
<th>Latitude, longitude</th>
<th>Water $^{87}$Sr/$^{86}$Sr</th>
<th>Otolith sample size</th>
<th>mtDNA Sample size</th>
<th>Haplotype diversity</th>
<th>Microsatellites Sample size</th>
<th>$H_O$</th>
<th>$H_E$</th>
<th>NA</th>
<th>AR (n = 14)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Lovettia sealii</em></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leven</td>
<td>41°09.941'S, 146°04.926'E</td>
<td>0.708642</td>
<td>10</td>
<td>23</td>
<td>0.846 ± 0.071</td>
<td>27</td>
<td>0.711</td>
<td>0.769</td>
<td>74</td>
<td>9.579</td>
</tr>
<tr>
<td>Rubicon</td>
<td>41°24.074'S, 146°36.349'E</td>
<td>0.705645</td>
<td>10</td>
<td>22</td>
<td>0.602 ± 0.121</td>
<td>30</td>
<td>0.694</td>
<td>0.762</td>
<td>80</td>
<td>9.773</td>
</tr>
<tr>
<td>Huon</td>
<td>42°59.909'S, 146°55.673'E</td>
<td>0.713497</td>
<td>10</td>
<td>19</td>
<td>0.614 ± 0.130</td>
<td>23</td>
<td>0.661</td>
<td>0.767</td>
<td>70</td>
<td>9.580</td>
</tr>
<tr>
<td>Derwent</td>
<td>42°45.544'S, 147°00.480'E</td>
<td>0.709885</td>
<td>10</td>
<td>13</td>
<td>0.641 ± 0.150</td>
<td>18</td>
<td>0.670</td>
<td>0.741</td>
<td>58</td>
<td>8.858</td>
</tr>
<tr>
<td>Black</td>
<td>40°50.801'S, 145°16.92'E</td>
<td>0.711765 ± 0.001138</td>
<td>1</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Tarwin</td>
<td>38°41.160'S, 145°49.86'E</td>
<td>–</td>
<td>–</td>
<td>6</td>
<td>0.800 ± 0.172</td>
<td>15</td>
<td>0.641</td>
<td>0.649</td>
<td>41</td>
<td>6.676</td>
</tr>
<tr>
<td><strong>Total/average</strong></td>
<td></td>
<td></td>
<td></td>
<td>41</td>
<td>83</td>
<td>0.865 ± 0.027</td>
<td>113</td>
<td>–</td>
<td>0.738</td>
<td>126</td>
</tr>
<tr>
<td><em>Pseudaphritis urvillii</em></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(n = 12)</td>
</tr>
<tr>
<td>Goolwa</td>
<td>35°31.574'S, 138°48.518'E</td>
<td>–</td>
<td>–</td>
<td>20</td>
<td>0.732 ± 0.077</td>
<td>30</td>
<td>0.845</td>
<td>0.857</td>
<td>107</td>
<td>10.730</td>
</tr>
<tr>
<td>Darlots</td>
<td>38°13.186'S, 141°46.233'E</td>
<td>0.708339 ± 0.000159</td>
<td>10</td>
<td>20</td>
<td>0.768 ± 0.062</td>
<td>20</td>
<td>0.824</td>
<td>0.848</td>
<td>90</td>
<td>10.465</td>
</tr>
<tr>
<td>Tarwin</td>
<td>38°39.861'S, 145°56.651'E</td>
<td>0.705729 ± 0.000017</td>
<td>5</td>
<td>14</td>
<td>0.824 ± 0.070</td>
<td>24</td>
<td>0.869</td>
<td>0.863</td>
<td>95</td>
<td>11.201</td>
</tr>
<tr>
<td>Tarwin (Ruby Creek)</td>
<td>–</td>
<td>0.705549</td>
<td>9</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Thomson</td>
<td>38°08.399'S, 147°04.738'E</td>
<td>–</td>
<td>–</td>
<td>21</td>
<td>0.710 ± 0.096</td>
<td>21</td>
<td>0.871</td>
<td>0.880</td>
<td>90</td>
<td>10.848</td>
</tr>
<tr>
<td>Snowy</td>
<td>37°42.618'S, 148°27.125'E</td>
<td>–</td>
<td>–</td>
<td>28</td>
<td>0.722 ± 0.059</td>
<td>35</td>
<td>0.872</td>
<td>0.856</td>
<td>113</td>
<td>10.825</td>
</tr>
<tr>
<td>Bemm</td>
<td>37°36.462'S, 148°54.061'E</td>
<td>0.718793 ± 0.000215</td>
<td>18</td>
<td>10</td>
<td>0.711 ± 0.118</td>
<td>21</td>
<td>0.850</td>
<td>0.861</td>
<td>88</td>
<td>10.345</td>
</tr>
<tr>
<td>Great Forster</td>
<td>41°02.766'S, 147°37.516'E</td>
<td>–</td>
<td>–</td>
<td>22</td>
<td>0.784 ± 0.053</td>
<td>30</td>
<td>0.852</td>
<td>0.866</td>
<td>107</td>
<td>10.429</td>
</tr>
<tr>
<td>Hopkins</td>
<td>38°23.10'S, 142°35.28'E</td>
<td>0.709457 ± 0.000191</td>
<td>2</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td><strong>Total/average</strong></td>
<td></td>
<td></td>
<td></td>
<td>44</td>
<td>135</td>
<td>0.761 ± 0.025</td>
<td>181</td>
<td>–</td>
<td>0.855</td>
<td>150</td>
</tr>
</tbody>
</table>
yses were carried out on a Nu Plasma multicollector inductively coupled plasma mass spectrometer (Nu Instruments, Wrexham, UK) interfaced with an Aridus™ desolvating nebulizer (CETAC, Omaha, Nebraska), operated at an uptake rate of ~40 μL/min (Maas et al. 2005). Mass bias was corrected by normalizing to 88Sr/86Sr = 8.37521, and results were reported relative to a value of 0.710230 for the SRM987 Sr isotope standard. Internal precisions (±2 SE) based on at least thirty 10-s integrations averaged ±0.00002 and average reproducibility (±2 SD) was ±0.00004.

Otolith dissection and preparation

Sagittal otoliths were dissected and mounted individually, proximal surface downward, on an acid-washed glass slide in thermostatic plastic glue (Crystalbond™; Aremco Products Inc., New York) and polished down to the primordium with a graded series of wetted lapping films (9.5, 3 μm) and alumina slurry (0.5 μm). The slide was then reheated, and the polished otolith was transferred to a master slide, on which otoliths from all collection sites were combined and arranged randomly to remove any systematic bias during analysis. The samples were rinsed several times in Milli-Q water (Millipore Corporation, Billerica, Massachusetts) and air dried overnight in a class-100 laminar flow cabinet at room temperature.

Sr isotope analysis

Laser ablation-inductively coupled plasma mass spectrometry (LA-ICPMS) was used to measure 87Sr/86Sr in the otoliths with a Nu Plasma multi-collector LA-ICPMS (Nu Instruments) coupled to a HeEx laser ablation system (Laurin Technic, Canberra, Australia, and the Australian National University) built around a 193-nm Compex 110 excimer laser (Lambda Physik, Gottingen, Germany). Otolith mounts were placed in the sample cell and the primordium of each otolith was found visually with a video imaging system (GeoStar version 6.14; Resonetics, Nashua, New Hampshire). The intended ablation transect on each sample was plotted digitally and each otolith was then ablated along this transect from the primordium to the dorsal margin at the widest radius using a 6 × 100-μm rectangular laser slit. The laser was operated at 90 mJ, pulsed at 10 Hz, and scanned at 3 μm/s across the sample. Ablation was done under an atmosphere of pure He to minimize redeposition of material, and the sample was then rapidly entrained into the Ar carrier gas flow. A preablation step using reduced energy (50 mJ) was conducted along each transect to remove any surface contaminants, and a 20- to 60-s background was measured before acquiring data for each sample. Potential Kr interferences were corrected by subtracting baselines on peak before each analysis, and corrections for 87Rb interferences were made following the method of Woodhead et al. (2005), assuming a natural 87Rb/86Rb ratio of 0.3865. Mass bias was then corrected by reference to an 86Sr/88Sr ratio of 0.1194. Iolite (version 2.13; Paton et al. 2011) was used to process data offline, and possible interferences from Ca argide/dimer species were corrected during this step. A modern marine carbonate (MMC) standard composed of mollusk shells was analyzed after every 10 otolith samples. Mean 87Sr/86Sr from 22 analyses of the MMC standard was 0.70917 ± 0.00011 (±2 SD), identical within error to the accepted modern seawater value of 0.70916 (McArthur and Howarth 2004).

Genetic data collection

Genomic deoxyribonucleic acid (DNA) was extracted from fin tissue using a standard phenol-chloroform extraction (Sambrook et al. 1989). For the L. sealii samples, a fragment of mitochondrial DNA (mtDNA) was amplified by polymerase chain reaction (PCR) using primers HYPSL510 and PH15938 for the 3’ end of cytochrome b (cyt b) (Thacker et al. 2007). These primers yielded poor amplification success, so a fragment spanning adenosine triphosphate (ATP) synthase subunit 6 and ATP synthase subunit 6 genes was amplified and sequenced using primers and protocols described by Woods et al. (2010). For the P. urvillii samples, mtDNA data were generated using cyt b primers as described above. Sequencing was done on a 3130xl Genetic Analyser (Applied Biosystems, Foster City, California) at Griffith University DNA sequencing facility. Sequences were edited and aligned using Sequencher (version 4.1.2; (GeneCodes Corp., Ann Arbor, Michigan) and deposited in GenBank under accession numbers KC992794–KC992815 (Tables S1, S2).

Microsatellite markers developed for L. sealii and P. urvillii were amplified and genotyped using primers developed by Schmidt et al. (2013). Six loci were used for L. sealii (lsea009, lsea012, lsea013, lsea019, lsea021, lsea023), and 7 loci were used for P. urvillii (purv002, purv004, purv005, purv008, purv010, purv011, purv013). Markers were labeled for fragment analysis using the multitalied primer tagging method of Real et al. (2009). Fragment analysis was conducted on an ABI 3130 sequencer (Applied Biosystems) and data were scored using GENEMAPPER (version 4.0; Applied Biosystems).

Genetic data analyses

Genetic diversity of subpopulations was explored by estimating observed (Ho) and expected (He) heterozygosity using ARLEQUIN (version 3.5.1.2; Excoffier and Lischer 2010). Tests for deviation from Hardy–Weinberg Equilibrium (HWE) for each locus–population combination were carried out using exact tests implemented in ARLEQUIN. Genetic structure was quantified by estimating pairwise and global FST values in ARLEQUIN. These values were tested for significant deviation from panmictic expectations by 10,000 permutations of individuals among populations. Because of high levels of within-population diversity, a
standardized value for $F_{ST}$ was obtained by expressing $F_{ST}$ as a proportion of the maximum value ($F_{STmax}$) that could be obtained (i.e., $F_{ST} = F_{ST}/F_{STmax}$; Meirmans and Hedrick 2011). The required estimate of $F_{STmax}$ was made indirectly by recoding the data file using RecodeData, (Meirmans 2006) followed by processing in ARLEQUIN. IBD was tested by correlating genetic distance (Slatkin’s Linearized $F_{ST}$) and geographic distance (coastline distance between rivers in km) for each pairwise test. The correlation was tested for significance with a Mantel test in ARLEQUIN using 1000 permutations. The critical value ($a$) was corrected for multiple tests using the BY False Discovery Rate method (BY-FDR) which controls experiment-wide Type I error without the loss of power associated with the Bonferroni adjustment (Benjamini and Yekutieli 2001, Narum 2006).

For each species, the existence of distinct genetic groups in the set of individual multilocus genotypes was tested using a model-based Bayesian clustering method. The probability of an admixture model was tested for up to 8 clusters ($K$) using STRUCTURE (version 2.3.1; Pritchard et al. 2000). Models were tested using 8 independent Markov Chain Monte Carlo (MCMC) simulations, each consisting of $1 \times 10^6$ iterations after a burn-in of $5 \times 10^5$ iterations. The most likely number of homogeneous clusters was assessed with the $2^{nd}$-order rate of change $L'' (K)$ following Evanno et al. (2005) using the online application STRUCTURE HARVESTER (Earl and Vonholdt 2012). Genealogical relationships of sampled mtDNA haplotypes were estimated using statistical parsimony in TCS (version 1.21; Clement et al. 2000).

RESULTS

Otolith Sr isotopes, _L. sealii_

Otolith transect measurements of $^{87}\text{Sr}/^{86}\text{Sr}$ were obtained from 41 _L. sealii_ individuals sampled from 5 rivers in Tasmania. All individual life-history profiles displayed low variability from the juvenile phase at the otolith core to the adult stage at the otolith edge and did not show evidence of a transition between isotopically distinct phases that would indicate habitat switching between marine and freshwater habitats (Fig. 2A). Individual profiles overlapped the marine standard $^{87}\text{Sr}/^{86}\text{Sr}$ value of 0.70916 with only 4 exceptions. These exceptions from the Huon River all had Sr isotope ratios marginally, but consistently, above the marine value (see Fig. 2A individual profile b). However, the mean (SD) ratio of 0.70981 (0.00021) for these 4 individuals was distinctly lower than the Huon river freshwater ratio of 0.71397 (Fig. 2A), suggesting that these fish did not at any stage inhabit full freshwater. A clear pattern evident from the summary of individual profiles presented in Fig. 2A is that most otolith Sr isotope ratios did not overlap with the corresponding river-water value at any point in the profile. Three individuals from the Derwent River had maximum values that encompassed their river-water value (Fig. 2A), but in each case, the maximum value was a short-lived signal spike rather than a representative trend in the $^{87}\text{Sr}/^{86}\text{Sr}$ transect (see Fig. 2A individual profile a, representative example of signal spike).

Genetic structure, _L. sealii_

Null alleles, large allele drop-out, stuttering artefacts, and linkage disequilibrium were not detected in the _L. sealii_ microsatellite data set (Schmidt et al. 2013). Three significant deviations from HWE were observed from a total of 30 locus–population combinations (Table S3). Microsatellite genetic diversity was high. A total of 126 alleles were identified from 6 loci genotyped in 113 individuals with expected heterozygosity averaged across all loci and populations of ~0.74 (Table 1). Comparison of microsatellite diversity among populations based on allelic richness standardized to a sample size of 14 showed that the 4 Tasmanian populations shared relatively high diversity compared to the Tarwin site on the mainland (Table 1). The edited alignment of 83 _L. sealii_ ATP sequences was 573 base pairs (bp) in length with 37 variable positions and 32 haplotypes (Fig. 1A). Haplotype diversity was moderate to high in all populations and ranged from ~0.60 to ~0.85 (Table 1). The sampled haplotypes formed a shallow network spanning a maximum of 7 steps (Fig. 1A). Two haplotypes (1 and 18, Fig. 1A) composed 52% of the total sample, whereas the remaining 30 haplotypes occurred at low frequencies of <5% (Table S1).

The global $F_{ST}$ estimate for microsatellite data was relatively low but significantly higher than the random expectation ($F_{ST} = 0.036, p < 0.0001$) and standardization for within-population genetic diversity produced a value ~4 times higher ($F_{ST} = 0.141$). Locus-by-locus estimates are provided in Table S4. For mtDNA data, the global $F_{ST}$ estimate was moderately high and significant ($F_{ST} = 0.242, p < 0.0001$) and the corresponding standardized measure was ~2 times as high ($F_{ST} = 0.514, p < 0.0001$). Microsatellite and mtDNA data produced concordant estimates of pairwise differentiation that reflected the geographic position of sampling sites. The mainland site on the Tarwin River was significantly differentiated from all other sites for both marker types, and no mtDNA haplotypes were shared with other populations (Tables S1, S5). Samples from Tasmania’s northern (Leven and Rubicon) and southern coasts (Huon and Derwent) were significantly differentiated from one another and only a single low-frequency haplotype was shared between north and south (haplotype 10, Fig. 1A; Tables S1, S5). No differentiation was found within the northern or southern groups for microsatellites or mtDNA (Table S5). Therefore, the pattern of genetic structure among the 5 river samples of _L. sealii_ follows the pattern expected from the species’ distribution, with 3 well differentiated groups: northern Tasmania, southern Tasmania, and...
Figure 2. Summary of individual life-history profiles based on otolith Sr isotope transects in *Lovettia sealii* (A) and *Pseudaphritis urvillii* (B). Bar plots summarize otolith data obtained for each individual grouped according to river sample. Grey bars represent the $^{87}\text{Sr}/^{86}\text{Sr}$ range (maximum–minimum) over the full transect. Open circles represent the average $^{87}\text{Sr}/^{86}\text{Sr}$ value at the otolith core, taken from the first 5 point measurements spanning ∼26 μm from the otolith primordium. Closed diamonds represent the average $^{87}\text{Sr}/^{86}\text{Sr}$ value at the otolith edge, taken from the last 5 point measurements spanning ∼26 μm from the otolith outer edge. Dashed line denotes the seawater $^{87}\text{Sr}/^{86}\text{Sr}$ ratio (0.70916). Solid lines denote the river water $^{87}\text{Sr}/^{86}\text{Sr}$ ratio for each sampling location. Four representative profiles are provided for each species. Sample codes for these fish are: *Lovettia sealii* 5007 (a), 4000 (b), 1005 (c), 3002 (d), and *Pseudaphritis urvillii* 5034 (a), 5134 (b), 7037 (c), 7139 (d).
Otolith Sr isotopes, _P. urvillii_

Otolith transect measurements of $^{87}$Sr/$^{86}$Sr were obtained from 44 _P. urvillii_ females sampled from 4 rivers: Bemm, Hopkins, Fitzroy (Darlrots Creek), and Tarwin. The Tarwin sample included individuals from the main channel and from a tributary (Ruby Creek), which differed slightly in river-water Sr isotope ratio (Table 1; Fig. 2B). Most individual life-history profiles showed a clear modulation in Sr isotope ratio, starting from the otolith core at values close to the marine value, then shifting at some intermediate point in the profile to values close to the river-water Sr isotope ratio (see Fig. 2B individual profiles a, b, c, d). This pattern is consistent with most individuals spending a significant period of their early life in marine water before migrating into freshwater. No evidence was found for multiple movements between marine and freshwater environments, suggesting that the downstream migration to the sea described by Crook et al. (2010) using acoustic telemetry occurs only once during life.

Some exceptions to the clear migratory pattern observed in most profiles included samples from Darlrots Creek and Hopkins River, where the river-water Sr isotope ratios ($0.708369, 0.707927$) were very close to the marine ratio ($0.70916$). This similarity made any transition between the freshwater and marine $^{87}$Sr/$^{86}$Sr values along the profile difficult to discern (Fig. 2B). The profiles of 4 individuals from the Tarwin sample and 1 individual from the Bemm sample did not overlap with the marine Sr ratio at any point in the transect (Fig. 2B). This result may indicate that some individuals spend their early life histories in brackish rather than marine habitats. It also is possible that the marine phase was very short-lived in these fish and that our $^{87}$Sr/$^{86}$Sr transect analyses failed to detect the marine phase.

Genetic structure, _P. urvillii_

Null alleles, large allele drop-out, stuttering artefacts, and linkage disequilibrium were not detected in the _P. urvillii_ microsatellite data set (Schmidt et al. 2013). Two significant deviations from HWE were observed from a total of 49 locus–population combinations (Table S6). Microsatellite genetic diversity was very high. A total of 150 alleles were identified from 7 loci genotyped in 181 individuals with expected heterozygosity averaged across all loci and populations of ~0.85 (Table 1). All of the 7 populations were similar in terms of microsatellite diversity based on allelic richness standardized to a sample size of 12 (Table 1). The edited alignment of 135 _P. urvillii_ cyt b sequences was 601 bp in length with 23 variable positions and 21 haplotypes (Fig. 1B). Haplotype diversity was moderate to high in all populations and ranged from ~0.71 to ~0.82 (Table 1). The sampled haplotypes formed a shallow network spanning a maximum of 8 steps, similar in structure to the _L. sealii_ network (Fig. 1B). As in _L. sealii_, 2 haplotypes occurred at relatively high frequency (33%; haplotype 3, 36%; haplotype 4; Fig. 2B, Table S2), with the remaining 19 haplotypes at relatively low frequencies.

The global $F_{ST}$ estimate for microsatellite data was relatively low but significantly higher than the random expectation ($F_{ST} = 0.0094, p < 0.0001$), and standardization for within-population genetic diversity produced a value ~7× higher ($F_{ST} = 0.067$). Locus-by-locus estimates are provided in Table S4. For mtDNA data, the global $F_{ST}$ estimate was low and nonsignificant ($F_{ST} = 0.020, p = 0.099$), and the corresponding standardized measure was nearly 3× higher and significant ($\Phi_{ST} = 0.058, p = 0.0197$). Pairwise estimates of differentiation were not concordant between microsatellite and mtDNA data sets. Five pairwise comparisons were significant (out of 21 tests) for microsatellites and one was significant for mtDNA (Table S7). A significant correlation between geographic distance and genetic distance was found among the 7 sampling sites for microsatellite data but not for mtDNA data ($R^2_{\text{micro}} = 0.26, p = 0.0177; R^2_{\text{mtDNA}} = 0.0009, p = 0.38$; Fig. S1). Bayesian clustering of individual microsatellite genotypes returned the highest mean likelihood score for $K = 1$ group, and therefore, no population structure was detectable in the microsatellite data set with this clustering approach.

**DISCUSSION**

_L. sealii_

Classification of _L. sealii_ as an anadromous fish has been widely accepted, and _L. sealii_ is one of the few proposed representatives of this form of diadromy in the southern hemisphere (McDowall 1988, Allen et al. 2002, Gillanders 2005, Miles 2007, Helfman et al. 2009). However, the Sr isotope profiles presented here are not consistent with habitat switching between marine and freshwater, but instead suggest that most individuals complete their life cycle in an environment that is equal or very similar in Sr isotope composition to marine water (i.e., saline or brackish water). Thus, if our results are general-
ized to the species, they suggest *L. sealii* is neither anadromous nor diadromous in the strict sense because residence in pure freshwater does not appear to occur during the life cycle. This life-history trait is unique among the Galaxiidae where all other diadromous members complete an obligate period of residence in freshwater. Extensive sampling of *L. sealii* during the peak of the Tasmanian whitebait spawning season (Blackburn 1950) at a salinity of 0.5% (≈ 0.1‰ chlorinity of Blackburn 1950) and above. Some spent fish may subsequently move into pure freshwater, but because most individuals live only 1 year, this short postreproductive freshwater period would not contribute to completion of the life cycle. The otolith chemistry data do not offer good resolution on whether the prespawning phase of the life cycle occurs in fully marine habitat or in estuaries because $^{87}$Sr/$^{86}$Sr ratios asymptotically approach marine values very quickly across a salinity gradient, so that mid-salinity estuarine habitats are difficult to distinguish from fully marine ones (Walther and Limburg 2012). Some evidence indicates that the habitat residence of the prespawning phase of the *L. sealii* life cycle is variable, ranging from fully marine, with anecdotal accounts of schools of whitebait observed several miles out to sea ~1 month before they begin to migrate into estuaries (Blackburn 1950), to mid-estuarine, with juveniles recorded in mid-austral autumn from within a coastal inlet on mainland Australia (TR, personal observation). In functional terms, *L. sealii* might best be regarded as semianadromous or even as estuarine-dependent rather than truly anadromous (Elliott et al. 2007).

Genetic diversity of the *L. sealii* sample was not as high as that of the *P. urvillii* sample but was in the range typical of marine rather than freshwater fishes (McCusker and Bentzen 2010). Unlike *P. urvillii*, genetic variation was strongly structured among subpopulations, with 3 well differentiated stocks consisting of northern Tasmania, southern Tasmania, and mainland Victoria. These results agree with Blackburn’s (1950) proposal that Australian whitebait were divided into northern and southern populations based on body size and pigmentation, and they support the similar conclusions reached by Pavuk (1994) based on analysis of allozymes. Blackburn (1950) also inferred that northern Tasmanian populations were homogenous because fishery-induced declines in abundance occurred simultaneously across the whole region despite differences in fishing pressure. Our results agree with this inference because our 2 northern population samples (Leven and Rubicon) were genetically homogenous.

Genetic differentiation between southeastern Australian and Tasmanian populations of freshwater or diadromous fishes has been observed in some species (e.g., *Retrospina* spp.; JMH, unpublished data) but not others (e.g., *Macquaria colonorum*, Shaddick et al. 2011). Some taxa show patterns inferred to represent drainage connections along the continental shelf that joined the 2 land masses during periods of Pleistocene lowered sea level (e.g., *Nannoperca australis*; Unmack et al. 2011), whereas others do not follow these predictions (e.g., *Galaxiella pusilla*; Unmack et al. 2012). Our finding of moderate genetic divergence between Tasmania and the mainland adds to this complex biogeographic picture. It remains to be determined whether the isolated mainland population of *Lovettia* is a relic of a former Pleistocene distribution connected via Lake Bass or was more recently founded by dispersal from Tasmania.

Regional patterns of genetic structure evident from pairwise *F*$_{ST}$ analyses of mtDNA and microsatellites were not supported by clustering analyses and hierarchical partitioning of variation. The Bayesian clustering method STRUCTURE was unable to recover any substructure within the microsatellite data set. This result is probably a consequence of very high microsatellite diversity that limited the amount of differentiation that could be expressed between subsets of individuals. Latch et al. (2006) showed that the minimum *F*$_{ST}$ at which STRUCTURE could correctly recover a priori groups was 0.03, and this value standardized for genetic diversity was *F*$_{ST}$ = 0.28. After standardizing for genetic diversity, the overall level of differentiation for the *L. sealii* data set (*F*$_{ST}$ = 0.14) was well below the detectable limit of STRUCTURE. The performance of STRUCTURE under conditions of high genetic variation and variable sample size has already been questioned and warrants further investigation (Latch et al. 2006, Kalinowski 2011). Hierarchical partitioning of genetic variation using AMOVA also produced a nonsignificant result when populations were arranged into northern, southern, and mainland groups. However, the explanation for this result is more straightforward. It is impossible to arrange 5 populations into a hierarchical structure and obtain a significant AMOVA result using permutation testing (Fitzpatrick 2009). Sampling more rivers is the solution to this issue, but even based on our limited sampling of 5 rivers, our results are consistent with previous research suggesting that *L. sealii* is subdivided into ≥2 stocks around the coast of Tasmania (Blackburn 1950, Pavuk 1994) in addition to our new finding that the recently discovered isolated population on mainland Australia may represent an independent genetic subpopulation within this species.

**P. urvillii**

Otolith $^{87}$Sr/$^{86}$Sr profiles confirm that *P. urvillii* females switch from marine to freshwater habitat during their early life history. This transition was most clear when divergence in Sr isotope ratio between freshwater and ma-
The genetic diversity of the *P. urvillii* sample was very high. Expected heterozygosity of microsatellites and haplotype diversity of mtDNA were both similar to average values for marine fish species (McCusker and Bentzen 2010). Genetic variation was only weakly structured among subpopulations, indicating that the sampled rivers were connected by some degree of dispersal and gene flow. The observation that some pairwise comparisons were significant for microsatellites but not for mtDNA is unlikely to be biologically significant. This difference probably is a consequence of the larger number of alleles and greater sample size of microsatellite data relative to mtDNA data, which leads to higher power to detect low levels of differentiation in microsatellite data (Larsson et al. 2009). An IBD relationship was detected in the microsatellite data, so this genetic connectivity probably is mediated by a stepping-stone pattern of dispersal along the coastline. Such a pattern could arise if the dispersal kernel of juveniles is centered mostly on rivers close to the parents’ point of origin, with a lower probability of long-distance dispersal. Alternatively, the IBD pattern may be mediated by limited dispersal of adults along the coastline. Similar IBD relationships have been reported for other coastline-restricted catadromous fishes (Keenan 1994, Jerry and Baverstock 1998, Shaddick et al. 2011). Therefore, it seems likely that the apparent absence of genetic structure (panmixia) observed in anguillid eels (Als et al. 2011) is an exception to the more-common pattern of weakly structured genetic connectivity (IBD) in other catadromous fishes. Generalizations about patterns of connectivity in diadromous organisms require further refinement to account for differences observed among the various modes of diadromy (Hughes et al. 2013).

An IBD pattern also could be produced by secondary intergradation of 2 differentiated genetic stocks (Turgeon and Bernatchez 2001) or from a hierarchical pattern of genetic structure (Meirmans 2012). Neither of these explanations seems likely for *P. urvillii* because clustering analyses did not detect hierarchical structure and no evidence was found of phylogeographic breaks in the mtDNA data that are commonly seen in obligate freshwater taxa subdivided into independent stocks along a coastline (Cook et al. 2014, Huey et al. 2014). Our study showed no evidence for multiple stocks across the sampled portion of the distribution of *P. urvillii*, but the potential for the existence of additional stocks around the coast of Tasmania, which was represented here by a single sample, cannot be ruled out. The northern part of the species’ distribution in New South Wales and the western extent past Spencer Gulf in South Australia also were not sampled as part of our study.

In summary, our results provide new information and independent confirmation of previous research on the characteristics of diadromy by *P. urvillii* and information that contradicts previous suggestions that *L. sealii* is anadromous. Our findings also suggest that genetic structuring of *P. urvillii* conforms to an IBD model, whereas *L. sealii* are structured across broad geographic regions. This fundamental information on ecology and population structure is critical as a basis for formulating effective management and conservation strategies for the species. In particular, our results highlight the importance of maintaining access to upper estuarine reaches for completion of the life cycle and for maintenance of population connectivity in both species.

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