Nursery origin of yellowfin tuna in the Hawaiian Islands

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ABSTRACT: Stable isotopes of carbon (δ13C) and oxygen (δ18O) in otolith cores (first 2 mo of age) of young-of-the-year (YOY) yellowfin tuna Thunnus albacares were used as natural tracers to predict the nursery origin of sub-adults (age-1) collected from the Hawaiian Islands. YOY fish were first collected from nurseries throughout the western and central Pacific Ocean (WCPO) to determine whether δ13C and δ18O values of otolith cores (δ13Cotolith and δ18Ootolith) were sufficiently different to discriminate individuals from different nurseries used during the YOY period. Nurseries examined included both nearshore Hawaiian Islands and offshore Hawaiian Islands (Cross Seamount), and 4 locations in the equatorial Pacific Ocean: Line Islands, Marshall Islands, Solomon Islands, and Philippines. Significant differences existed in δ13Cotolith and δ18Ootolith among YOY yellowfin tuna from different nurseries for each year of the 2 yr study period (2008–2009). Individuals from the nearshore Hawaiian Islands were most enriched in δ18Ootolith, while samples obtained from the Philippines were most depleted in both δ13Cotolith and δ18Ootolith relative to other regions. Inter-annual variability in otolith core chemistry was minor and only observed for individuals collected from the Philippines. Overall classification success from quadratic discriminant function analysis of YOY yellowfin tuna to their respective nursery of collection was 63 and 87% for 2008 and 2009, respectively. Mixed-stock analysis indicated 91% of the sub-adult yellowfin tuna collected from the nearshore Hawaiian Islands originated from this same nursery. In addition, sub-adults from the offshore location within the Hawaiian Islands appear to originate from the nearshore Hawaiian Islands, highlighting the importance of local production and retention of yellowfin tuna to the standing stock and domestic fisheries of Hawaii.

KEY WORDS: Stable carbon and oxygen isotopes · Yellowfin tuna · Nursery origin · Otolith chemistry · Hawaiian Islands · Stock structure

INTRODUCTION

Tuna stocks of the western and central Pacific Ocean (WCPO) represent 84% of the total Pacific Ocean catch and 60% of the global tuna catch (Williams & Terawasi 2011). Exploitation rates on WCPO tuna stocks have increased steadily over the past 3 decades with mortality rates approaching maximum sustainable yield (MSY) levels. A species of particular importance in the WCPO is yellowfin tuna Thunnus albacares, which yields approximately 400,000 t yr⁻¹. Fishing mortality for juvenile and adult yellowfin tuna is estimated to have increased continuously since the beginning of industrial fishing in the region with the current stock assessment predicting a steady decline in total and spawning stock biomass (Langley et al. 2011). The variable movement parameters and recruitment variability of yellowfin tuna combined with diverse gear types and exploitation rates throughout the WCPO makes regional and international management strategies complicated. Consequently, a more refined understanding of the
movement and stock structure of yellowfin tuna within the WCPO is needed to effectively guide the management of this species.

Several complementary approaches have been used to address the movement and stock structure of yellowfin tuna in the Pacific Ocean, including molecular genetics, morphology, and tagging (conventional, electronic). Genotypic and phenotypic data of yellowfin tuna indicate limited gene flow between WCPO and eastern Pacific Ocean regions (Schaefer 1992, Ward et al. 1997) and genetic differentiation was observed between samples north and south of the equator in the eastern Pacific (Díaz-Jaimés & Uribe-Alcocer 2006). Tagging data also suggest that regional movement in the Pacific Ocean is limited, signifying that some degree of stock heterogeneity occurs within this basin (Itano & Holland 2000, Schaefer et al. 2007).

In the Hawaiian Islands, local spawning of yellowfin tuna has been documented but the degree of local recruitment remains unknown (Itano 2000). Tagging studies in waters surrounding the main Hawaiian Islands generally support the premise of retentive behaviours of yellowfin tuna and long-distance (>1000 km) movements are thought to be rare (Itano & Holland 2000). Hampton & Fournier (2001) developed a length-based and age-structured model for yellowfin tuna throughout the WCPO that assumed a large fraction of the yellowfin tuna harvested in the Hawaii-based fisheries originated from equatorial regions. Thus, the origin of recruits in the Hawaiian Islands remains uncertain and this information is critical to reduce uncertainty in population models and develop spatially explicit management strategies.

Recent studies using chemical signatures in otoliths have shown that these natural markers are valuable for evaluating the origin and movement of tunas in the Pacific Ocean (Rooker et al. 2001, Wang et al. 2009, Shi ao et al. 2010). The principal assumption underlying this approach is that the otolith accretes material as the fish grows and the chemical composition of the otolith is related to the physicochemical conditions of the water mass inhabited (Rooker et al. 2001). Therefore, material deposited in the otolith during the first weeks to months of life may serve as a natural tag of an individual’s place of origin. Previous studies have demonstrated that stable isotopes of carbon ($\delta^{13}$C) and oxygen ($\delta^{18}$O) in otolith cores (hereafter $\delta^{13}$C$_{\text{otolith}}$ and $\delta^{18}$O$_{\text{otolith}}$) can be used to determine the origin of tropical and temperate tunas (Gunn & Ward 1994, Rooker et al. 2008a,b, Schloesser et al. 2010), and thus may prove useful for determining contribution rates of yellowfin tuna recruits from different nurseries throughout the WCPO.

Here, we investigate the nursery origin of sub-adult (age-1) yellowfin tuna from 2 nurseries of the Hawaiian Islands using $\delta^{13}$C$_{\text{otolith}}$ and $\delta^{18}$O$_{\text{otolith}}$. These nurseries are referred to as ‘nearshore’ referring to multiple sampling areas within 30 km of the main Hawaiian Islands and ‘offshore’ referring to samples collected on the Cross Seamount, an isolated bathymetric feature approximately 300 km south of Oahu. We were specifically interested in determining whether sub-adult yellowfin tuna collected in nearshore and offshore nurseries of the Hawaiian Islands were locally produced (residents) or had immigrated from distant areas (transients) of the equatorial Pacific Ocean. We first assessed spatial and temporal variability in $\delta^{13}$C$_{\text{otolith}}$ and $\delta^{18}$O$_{\text{otolith}}$ in the otolith cores of young-of-the-year (YOY, age-0) yellowfin tuna to determine whether individuals from different nurseries of the WCPO could be discriminated and these data were then used to develop baseline signatures for mixed-stock analysis of sub-adult yellowfin tuna. Next, we targeted sub-adult yellowfin tuna that were recruited into the nearshore and offshore Hawaiian Island fisheries and predicted their origin by examining $\delta^{13}$C$_{\text{otolith}}$ and $\delta^{18}$O$_{\text{otolith}}$ in the otolith cores (equal to a restricted portion of the YOY period) of these individuals.

**MATERIALS AND METHODS**

YOY yellowfin tuna were collected from 6 nurseries throughout the WCPO: (1) nearshore Hawaiian Islands (ca. ≤30 km from land), (2) offshore Hawaiian Islands (Cross Seamount, 300 km southwest of the Hawaiian Islands), (3) Line Islands of Kiribati, (4) Marshall Islands, (5) Solomon Islands, and (6) Philippines (Fig. 1). Yellowfin tuna were collected over a 2 yr period (late 2007 to early 2008 and late 2008 to early 2009, hereafter 2008 and 2009, respectively) through either hook-and-line or purse seine techniques. Within each nursery, sub-samples from multiple collection dates and locations were taken to ensure that baseline $\delta^{13}$C$_{\text{otolith}}$ and $\delta^{18}$O$_{\text{otolith}}$ values were representative of each nursery. For example, YOY yellowfin tuna collected from the nearshore Hawaiian Islands were obtained from multiple collection dates and islands (Kauai, Maui, and Oahu). For purposes of this study, we define nurseries as primary regions used by YOY yellowfin tuna in the WCPO and are therefore considered as important habitat during the first year of life, while not implying
spawning location since yellowfin tuna are pelagic broadcast spawners.

Sub-adult yellowfin tuna were collected in 2009 from the nearshore Hawaiian Islands and in 2010 from the offshore Hawaiian Islands to investigate nursery-specific contribution rates (Table 1). Similar to YOY fish, sub-adults were collected over a range of dates from nearshore areas of Kauai, Maui, Oahu and the island of Hawaii. Based on previous otolith-based ageing studies in the Pacific Ocean (Uchiyama & Struhsaker 1981, Wild 1986, Lehodey & Leroy 1999), yellowfin tuna ranging in size from 21 to 59 cm fork length (FL) and 60 to 100 cm FL were classified as YOY (age-0) and sub-adults (age-1), respectively.

Sagittal otoliths were extracted from both fresh and frozen specimens, cleaned of biological residue, and stored dry in plastic vials. In the laboratory, whole otoliths were first soaked in doubly deionized water (DDIH2O), moved to a 3% hydrogen peroxide solution for 5 min to eliminate any remaining biological material, and then transferred into a new DDIH2O bath for 5 min to remove surface residue. One sagittal otolith from each yellowfin tuna was embedded in Struers epoxy resin (EpoFix) and sectioned using a low speed ISOMET saw to obtain 1.5 mm transverse sections that included the core. Following attachment to a sample plate, the portion of the otolith corresponding to approximately the first 2 mo of life was milled from the otolith section using a New Wave Research MicroMill system. A 2-vector drill path based upon otolith measurements of 10 small individual yellowfin tuna (mean ± SE fish length: 22.4 ± 0.13 cm FL, range: 21.8 to 23.0 cm FL) was created and used as the standard template to isolate core material (0 to 2 mo) for YOY and sub-adult yellowfin tuna (Fig. 2). The pre-programmed drill path was made using a 500 µm diameter drill bit and 14 passes each at a depth of 55 µm (770 µm total) were used to obtain core material from the otolith. Shiao et al. (2009) suggested that a small amount of calcium carbonate material on the outer edge may be deposited around the area surrounding the core during sub-adult and adult stages for southern bluefin tuna Thunnus maccoyii. Consequently, a small amount of outer edge material may have been incorporated into the powdered core material, but the amount of material was likely so small that it did not affect the results.

Table 1. Thunnus albacares. Summary statistics of young-of-the-year (YOY, age-0) and sub-adult (age-1) yellowfin tuna collected throughout western and central Pacific Ocean (WCPO) study areas. Mean (±1 SE) size and size range are fork lengths (FL), and collection dates given as mm/dd/yyyy.

<table>
<thead>
<tr>
<th>Study area</th>
<th>Year</th>
<th>n</th>
<th>Mean size (cm FL)</th>
<th>Size range (cm FL)</th>
<th>Collection dates</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>YOY</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Nearshore Hawaii</td>
<td>2008</td>
<td>30</td>
<td>37.5 (0.98)</td>
<td>27.9−50.0</td>
<td>10/1/2007−1/10/2008</td>
</tr>
<tr>
<td>Nearshore Hawaii</td>
<td>2009</td>
<td>20</td>
<td>35.3 (0.75)</td>
<td>29.0−41.0</td>
<td>1/26/2009−1/29/2009</td>
</tr>
<tr>
<td>Offshore Hawaii</td>
<td>2008</td>
<td>25</td>
<td>54.3 (0.78)</td>
<td>44.0−59.0</td>
<td>2/22/2008−3/16/2008</td>
</tr>
<tr>
<td>Line Islands</td>
<td>2008</td>
<td>25</td>
<td>53.3 (0.75)</td>
<td>48.0−59.0</td>
<td>4/12/2008−5/28/2008</td>
</tr>
<tr>
<td>Line Islands</td>
<td>2009</td>
<td>13</td>
<td>46.9 (1.18)</td>
<td>39.0−54.0</td>
<td>5/23/2009−10/12/2009</td>
</tr>
<tr>
<td>Marshall Islands</td>
<td>2008</td>
<td>25</td>
<td>30.7 (0.51)</td>
<td>27.0−35.0</td>
<td>1/1/2008−3/3/2008</td>
</tr>
<tr>
<td>Marshall Islands</td>
<td>2009</td>
<td>16</td>
<td>53.9 (0.74)</td>
<td>47.0−58.0</td>
<td>4/12/2009−4/18/2009</td>
</tr>
<tr>
<td>Solomon Islands</td>
<td>2008</td>
<td>25</td>
<td>35.1 (0.37)</td>
<td>32.0−37.0</td>
<td>3/2/2008−3/14/2008</td>
</tr>
<tr>
<td>Philippines</td>
<td>2008</td>
<td>25</td>
<td>26.0 (0.73)</td>
<td>21.8−33.5</td>
<td>12/22/2007−1/12/2008</td>
</tr>
<tr>
<td>Philippines</td>
<td>2009</td>
<td>19</td>
<td>30.8 (1.03)</td>
<td>24.8−37.0</td>
<td>12/22/2008−1/3/2009</td>
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<tr>
<td><strong>Sub-adult</strong></td>
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<tr>
<td>Nearshore Hawaii</td>
<td>2009</td>
<td>100</td>
<td>69.4 (0.81)</td>
<td>60.0−97.0</td>
<td>1/26/2009−3/16/2009</td>
</tr>
<tr>
<td>Offshore Hawaii</td>
<td>2010</td>
<td>25</td>
<td>79.5 (1.60)</td>
<td>61.0−98.0</td>
<td>4/1/2010−6/12/2010</td>
</tr>
</tbody>
</table>
minimal that it would not affect the results for the life stages investigated in this study. Powdered core material was transferred to silver capsules and later analyzed for δ\(^{13}\)C\(_{\text{otolith}}\) and δ\(^{18}\)O\(_{\text{otolith}}\) on an automated carbonate preparation device (KIEL-III) coupled to a gas-ratio mass spectrometer (Finnigan MAT 252) maintained at the University of Arizona’s Environmental Isotope Laboratory. The isotope ratio measurement was calibrated based on repeated measurements of National Bureau of Standards (NBS), NBS-19 and NBS-18, with 6 standards ran for every 40 samples and precision was ± 0.11‰ (SD) for NBS-19 and NBS-18, with 6 standards ran for every 40 samples and precision was ± 0.08‰ (SD) for δ\(^{13}\)C. Stable δ\(^{13}\)C and δ\(^{18}\)O isotopes are reported relative to the PeeDee belemnite (PDB) scale after comparison to an in-house laboratory standard calibrated to PDB.

Multivariate analysis of variance (MANOVA) was used to test for differences in δ\(^{13}\)C\(_{\text{otolith}}\) and δ\(^{18}\)O\(_{\text{otolith}}\) of YOY yellowfin tuna among nurseries sampled; Pillai’s trace statistic was used to test for significance. Univariate tests for both δ\(^{13}\)C\(_{\text{otolith}}\) and δ\(^{18}\)O\(_{\text{otolith}}\) were also performed using analysis of variance (ANOVA) and a posteriori differences among means were detected with Tukey’s honestly significant difference (HSD) test. The variance-covariance matrix of predictor variables was different among samples so quadratic discriminant function analysis (QDFA) was used to evaluate the classification accuracy of individual YOY yellowfin tuna to nurseries based on the jackknife reclassification (Rooker et al. 2008a). Quadratic discriminant function analysis does not have the homogeneity of covariance matrices assumption and is robust to moderate deviations from normality (McGarigal et al. 2000). Nursery-specific contribution estimates of yellowfin recruits to nearshore and offshore Hawaii were obtained using the maximum likelihood mixed-stock analysis program HISEA developed by Millar (1990). The baseline data set used for mixed-stock analysis was δ\(^{13}\)C\(_{\text{otolith}}\) and δ\(^{18}\)O\(_{\text{otolith}}\) of YOY samples collected from the 6 nurseries in 2008. Otolith cores of sub-adult samples collected from the nearshore and offshore Hawaiian Island nurseries (collected in 2009 and 2010, respectively) were used to estimate the origin of these recruits in the bootstrap mode of HISEA with 10,000 simulations, which provided non-parametric estimates of the reliability of predicted contributions from the different nurseries. All other statistical analyses were performed using SYSTAT 10.0 (SYSTAT Software) and significance for all tests was determined at the alpha level of 0.05.

RESULTS

δ\(^{13}\)C\(_{\text{otolith}}\) and δ\(^{18}\)O\(_{\text{otolith}}\) of YOY yellowfin tuna significantly differed among the nurseries sampled and showed similar spatial variation in both years of the study (MANOVA, p < 0.01) (Fig. 3). δ\(^{13}\)C\(_{\text{otolith}}\) of YOY fish was most enriched for individuals collected from the Marshall Islands (mean ± SE, 2008: −9.29 ± 0.09; 2009: −9.32 ± 0.10; p < 0.01), with mean values 0.37 to 1.27‰ (2008) and 0.42 to 0.55‰ (2009) that were significantly more enriched relative to other nurseries (Tukey HSD, p < 0.05). δ\(^{13}\)C\(_{\text{otolith}}\) values of YOY fish from the Philippines were the most depleted (2008: −10.56 ± 0.08; 2009: −9.87 ± 0.09; p < 0.01), while values for the other 4 nurseries were relatively similar with mean values within 0.08‰ of each other. δ\(^{18}\)O\(_{\text{otolith}}\) of YOY fish also varied among nurseries but spatial variation was different from δ\(^{13}\)C\(_{\text{otolith}}\). The most enriched δ\(^{18}\)O\(_{\text{otolith}}\) values were observed for YOY yellowfin tuna from the nearshore Hawaiian Islands (2008: −1.77 ± 0.03; 2009: −1.69 ± 0.03), with mean values significantly enriched relative to other nurseries (Tukey HSD, p < 0.05) by 0.33 to 0.88‰ and 0.67 to 1.13‰ in 2008 and 2009, respectively. δ\(^{18}\)O\(_{\text{otolith}}\) values of YOY fish decreased east to west and were significantly depleted in samples obtained from the Philippines in both years (2008: −2.65 ± 0.05; 2009 −2.82 ± 0.05) with the exception of similar δ\(^{18}\)O\(_{\text{otolith}}\) between YOY fish from the Philippines and Solomon Islands in 2008 (p = 0.867).

Inter-annual comparisons of δ\(^{13}\)C\(_{\text{otolith}}\) and δ\(^{18}\)O\(_{\text{otolith}}\) of YOY yellowfin tuna were possible for 4 nurseries; missing year classes for offshore Hawaiian Islands and Solomon Islands precluded assessments in these nurseries. Differences in δ\(^{13}\)C\(_{\text{otolith}}\) and δ\(^{18}\)O\(_{\text{otolith}}\) of YOY from 2008 and 2009 were only detected in the Philippines. YOY yellowfin tuna in the Philippines in 2008 had significantly depleted δ\(^{13}\)C\(_{\text{otolith}}\) and signifi-

Fig. 2. Thunnus albacares. Transverse cross section of a subadult yellowfin tuna sagittal otolith showing the primary drill path used on all samples to core out the inner region of the otolith corresponding to the first months of life to predict nursery origin.
significantly enriched δ^{18}O_{otolith} relative to 2009 by 0.69‰ (p < 0.01) and 0.17‰ (p = 0.02), respectively (Fig. 3). In contrast, no significant difference in δ^{13}C_{otolith} or δ^{18}O_{otolith} existed in samples collected between 2008 and 2009 in the nearshore Hawaiian Islands (δ^{13}C: p = 0.73; δ^{18}O: p = 0.09), Line Islands (δ^{13}C: p = 0.94; δ^{18}O: p = 0.07), and Marshall Islands (δ^{13}C: p = 0.83; δ^{18}O: p = 0.95).

Discrimination of YOY yellowfin tuna among the nurseries surveyed was relatively high using δ^{13}C_{otolith} and δ^{18}O_{otolith} with overall classification success of 63 and 87% for 2008 and 2009, respectively. Expected classification success based on random assignments for each year was 17% (2008 based on 6 nurseries) and 25% (2009 based on 4 nurseries). In 2008, classification success of YOY fish was highest for the Philippines (84%) followed by nearshore Hawaiian Islands (80%). In 2009, classification success of YOY fish from the nearshore Hawaiian Islands was 100% followed by Line Islands at 85%. Our ability to discriminate individuals from other nurseries ranged from 20% (Line Islands in 2008) to 81% (Marshall Islands in 2009) (Table 2). Using only δ^{13}C_{otolith} or δ^{18}O_{otolith} in QDFA models resulted in a reduction in overall classification success (δ^{13}C_{otolith} only: 33% in 2008, 40% in 2009; δ^{18}O_{otolith} only: 47% in 2008, 79% in 2009), suggesting that both δ^{13}C_{otolith} and δ^{18}O_{otolith} markers were useful for assigning the origin of YOY yellowfin tuna.

Table 2. Thunnus albacares. Mean (±1 SE) otolith carbon (δ^{13}C) and oxygen (δ^{18}O) stable isotope values of young-of-the-year (YOY, age-0) and sub-adult (age-1) yellowfin tuna collected among western and central Pacific Ocean (WCPO) nursery areas. Classification success (%) from quadratic discriminant function analysis is also included for each nursery area by year.

<table>
<thead>
<tr>
<th>Nursery Area</th>
<th>Year</th>
<th>δ^{13}C</th>
<th>δ^{18}O</th>
<th>Classification Success (%)</th>
</tr>
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<tbody>
<tr>
<td><strong>YOY</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nearshore Hawaii</td>
<td>2008</td>
<td>−9.70 (0.09)</td>
<td>−1.77 (0.03)</td>
<td>80</td>
</tr>
<tr>
<td></td>
<td>2009</td>
<td>−9.74 (0.09)</td>
<td>−1.69 (0.03)</td>
<td>100</td>
</tr>
<tr>
<td>Offshore Hawaii</td>
<td>2008</td>
<td>−9.68 (0.09)</td>
<td>−2.16 (0.04)</td>
<td>52</td>
</tr>
<tr>
<td>Line Islands</td>
<td>2008</td>
<td>−9.68 (0.10)</td>
<td>−2.10 (0.05)</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td>2009</td>
<td>−9.69 (0.06)</td>
<td>−2.24 (0.05)</td>
<td>85</td>
</tr>
<tr>
<td>Marshall Islands</td>
<td>2008</td>
<td>−9.29 (0.09)</td>
<td>−2.37 (0.03)</td>
<td>72</td>
</tr>
<tr>
<td></td>
<td>2009</td>
<td>−9.32 (0.10)</td>
<td>−2.37 (0.04)</td>
<td>81</td>
</tr>
<tr>
<td>Solomon Islands</td>
<td>2008</td>
<td>−9.66 (0.09)</td>
<td>−2.64 (0.03)</td>
<td>68</td>
</tr>
<tr>
<td>Philippines</td>
<td>2008</td>
<td>−10.56 (0.08)</td>
<td>−2.65 (0.05)</td>
<td>84</td>
</tr>
<tr>
<td></td>
<td>2009</td>
<td>−9.87 (0.09)</td>
<td>−2.82 (0.05)</td>
<td>79</td>
</tr>
<tr>
<td><strong>Sub-adult</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nearshore Hawaii</td>
<td>2009</td>
<td>−9.41 (0.05)</td>
<td>−1.73 (0.02)</td>
<td></td>
</tr>
<tr>
<td>Offshore Hawaii</td>
<td>2010</td>
<td>−9.93 (0.07)</td>
<td>−1.87 (0.04)</td>
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</tbody>
</table>

Fig. 3. Thunnus albacares. Otolith core carbon (δ^{13}C_{otolith}) and oxygen (δ^{18}O_{otolith}) stable isotopes of young-of-the-year (YOY, age-0) yellowfin tuna collected among nurseries in (A) 2008 and (B) 2009. Confidence ellipses represent 1 SD around the mean. Nurseries include (○) nearshore Hawaiian Islands, (∗) offshore Hawaiian Islands, (+) Line Islands, (Δ) Marshall Islands, (♂) Solomon Islands, and (♀) Philippines.
were compared to the YOY baseline samples collected during the previous year (age-class matching) to predict the origin of these fish. $\delta^{13}C_{\text{otolith}}$ (p = 0.812) and $\delta^{18}O_{\text{otolith}}$ (p = 0.900) did not significantly differ among the 4 locations and therefore samples from all 4 islands were pooled for mixed-stock analysis. $\delta^{13}C_{\text{otolith}}$ of sub-adult yellowfin tuna ranged from −8.08 to −10.59‰ (Fig. 4), closely overlapping the $\delta^{13}C_{\text{otolith}}$ of YOY fish obtained from all nurseries except the Philippines. $\delta^{18}O_{\text{otolith}}$ of sub-adults from the nearshore Hawaiian Islands ranged from −1.14 to −2.26‰ (mean: −1.73‰) and was similar to that of YOY fish from nearshore Hawaiian Islands (range: −1.22 to −2.05‰; mean: −1.77‰) (Fig. 4). Further, $\delta^{18}O_{\text{otolith}}$ of sub-adults was enriched by 0.37 to 0.92‰ relative to mean $\delta^{18}O_{\text{otolith}}$ of samples from the other 5 nurseries used for baseline contribution estimates. Results of mixed-stock analysis indicated that 90.6 ± 8.6% (mean ± SD) of sub-adults in our nearshore Hawaiian Islands sample originated from this nursery (Fig. 5), suggestive of local recruitment. Mixed-stock analysis also showed minor contributions from the 2 closest nurseries to the south of the nearshore Hawaiian Islands: Line Islands (7.3 ± 9.0%) and offshore Hawaiian Islands (2.1 ± 4.5%), with no contribution from other nurseries.

Nursery origin of sub-adult yellowfin tuna was also predicted for samples collected in the offshore Hawaiian Islands (i.e. Cross Seamount) using $\delta^{13}C_{\text{otolith}}$ and $\delta^{18}O_{\text{otolith}}$ (n = 25). $\delta^{13}C_{\text{otolith}}$ of sub-adult yellowfin tuna from this nursery ranged from −9.43 to −10.91‰, with a mean of −9.93‰ (Fig. 4). For $\delta^{18}O_{\text{otolith}}$, values of sub-adults ranged from −1.43 to −2.14‰ (mean: −1.87‰) and differed markedly from YOY collected in the offshore Hawaiian Islands (Fig. 4). Mixed-stock analysis indicated that local production in the offshore Hawaiian Islands was negligible (0.7 ± 3.4%, mean ± SD) and recruits to this nursery were derived primarily from the nearshore Hawaiian Islands (89.9 ± 12.5%), with the remaining contribution from the Line Islands (9.4 ± 12.5%) (Fig. 5).

**DISCUSSION**

$\delta^{13}C_{\text{otolith}}$ of YOY yellowfin tuna was similar among 4 of the 6 nurseries examined (nearshore Hawaiian Islands, offshore Hawaiian Islands, Line Islands, Solomon Islands), averaging less than 0.1‰, while $\delta^{13}C_{\text{otolith}}$ of yellowfin tuna from the Philippines was depleted relative to other nurseries. A positive relationship between $\delta^{13}C_{\text{otolith}}$ and salinity (~0.1‰ increase in $\delta^{13}C_{\text{otolith}}$ per 1 psu increase) combined with an inverse relationship between $\delta^{13}C_{\text{otolith}}$ and sea surface temperature (<0.1 to 0.25‰ decrease in $\delta^{13}C_{\text{otolith}}$ per 1°C increase) has been reported in other studies (Elsdon & Gillanders 2002, Kerr et al. 2007). Consequently, the low salinity (mean salinity ~34.0 psu) and high sea surface temperatures (~30°C) characteristic of the Philippines relative to other nurseries (Antonov et al. 2010, Gordon et al. 2011) may be responsible for the 0.13 to 1.27‰ depletion in $\delta^{13}C_{\text{otolith}}$ of YOY yellowfin tuna collected from this nursery. Additionally, Chen et al.
reported similar patterns of low seawater \( \delta^{13}C \) (\( \delta^{13}C_{\text{seawater}} \)) in the western Pacific with an increasing trend in \( \delta^{13}C_{\text{seawater}} \) from west to east along the tropical and sub-tropical North Pacific and linked this relationship to the zonal pattern of CO2 solubility in surface waters. The enriched \( \delta^{13}C_{\text{otolith}} \) of YOY yellowfin tuna collected in the Marshall Islands cannot be explained by the aforementioned salinity and temperature patterns alone since the Hawaiian Island nurseries have highest salinity and lowest sea surface temperatures that would lead to expected enriched \( \delta^{13}C_{\text{otolith}} \). Studies have shown over 80% of \( \delta^{13}C_{\text{otolith}} \) is derived from food and dissolved inorganic carbon (DIC) of ambient seawater (Solomon et al. 2006) and therefore the isotopic composition of the basal carbon sources and prey species specific to the Marshall Islands may account for the enriched \( \delta^{13}C_{\text{otolith}} \) of YOY yellowfin tuna in this nursery.

\( \delta^{18}O_{\text{otolith}} \) of YOY yellowfin tuna match spatial trends to seawater \( \delta^{18}O \) (\( \delta^{18}O_{\text{seawater}} \)) and salinity patterns reported for the WCPO. A decreasing trend in surface \( \delta^{18}O_{\text{seawater}} \) (upper 50 m) from the central (0.4‰, \( n = 72 \) observations) to western equatorial Pacific Ocean (0.1‰, \( n = 122 \)) reported by Schmidt et al. (1999) matched the pattern of decreasing \( \delta^{18}O_{\text{otolith}} \) from central to western nurseries examined. Mean \( \delta^{18}O_{\text{otolith}} \) differences between the Line Islands (central) and Philippines (western) nurseries were 0.55 and 0.58‰ in 2008 and 2009, respectively, slightly larger than the 0.30‰ difference in \( \delta^{18}O_{\text{seawater}} \). In the WCPO, sea surface salinity and \( \delta^{18}O_{\text{seawater}} \) are positively correlated, and both are affected by evaporation, precipitation, and atmospheric convection (Stott et al. 2004). \( \delta^{18}O_{\text{otolith}} \) is a function of evaporation (increases \( ^{18}O \) in seawater) and precipitation (decreases \( ^{18}O \) in seawater). It is also positively correlated with sea surface salinity (Campana 1999), often increasing by 0.2 (Kerr et al. 2007) to 1.4‰ (Dufour et al. 1998) per 1 psu increase. The warm, low-salinity surface waters (mean salinity: ~34.0 psu) of the western equatorial Pacific contrasts the cool and more saline surface waters of the central equatorial Pacific (Line Islands, salinity: ~34.5 psu) (Schmidt et al. 1999, Antonov et al. 2010) and matched the 0.5 to 0.6‰ difference in \( \delta^{18}O_{\text{otolith}} \) from central and western equatorial nurseries. Similar trends of decreasing \( \delta^{18}O_{\text{seawater}} \) and \( \delta^{18}O_{\text{otolith}} \) were observed north to south with nearshore Hawaiian Islands \( \delta^{18}O_{\text{seawater}} \) enriched by 0.04‰ relative to the southerly located Line Islands nursery at a similar longitude (Schmidt et al. 1999). Moreover, mean \( \delta^{18}O_{\text{otolith}} \) was enriched by 0.3‰ in 2008 and 0.6‰ in 2009 at the nearshore Hawaiian Islands (relative to the Line Islands) where mean salinity averages approximately 35.0 psu (Schmidt et al. 1999, Antonov et al. 2010), about 0.5 to 1.0 psu higher than other nurseries examined.

Inter-annual variability in \( \delta^{13}C_{\text{otolith}} \) and \( \delta^{18}O_{\text{otolith}} \) was minimal among the nurseries investigated, except the Philippines, which may have resulted from variability in climactic and oceanographic conditions. Samples collected from the Philippines nursery showed differences in both \( \delta^{13}C_{\text{otolith}} \) and \( \delta^{18}O_{\text{otolith}} \) during the 2 yr study and may be related to the El Niño Southern Oscillation (ENSO), which affected precipitation, sea surface salinity, and sea surface temperatures in our Philippines nursery. Late 2007 (August to December) and early 2008 (January to April) was a moderately strong La Niña period (Climate Prediction Center 2010), and this time period coincides with the spawning season of our 2008 YOY samples with depleted \( \delta^{13}C_{\text{otolith}} \) and
enriched δ18O

compared to 2009 samples collected the following year (late 2008 to early 2009). La Niña events generally produce heavy precipitation and cyclone activity in the western Pacific (Ropelewski & Halpert 1996); however, a seasonal reversal of the ENSO rainfall is unique to the Philippines area during summers of La Niña periods resulting in below average rainfall (Lyon et al. 2006). The seasonal southwest monsoon that usually produces high precipitation was significantly weakened resulting in fewer tropical cyclones, thereby blocking the northward migration of the inter-tropical convergence zone (Yumul et al. 2010). Consequently, anomalously low precipitation and high evaporation led to higher salinities in late 2007 to early 2008 (mean salinity: 34.1 psu) compared with late 2007 to early 2009 (mean salinity: 33.6 psu) (Gordon et al. 2011). Our 2007 to 2008 YOY Philippine samples had lower δ13C relative to the 2008 to 2009 samples despite being exposed to lower precipitation and higher salinities that would lead to increased δ13C. However, mean sea surface temperature in late 2007 to early 2008 was 2°C higher than in 2008 to 2009 (Gordon et al. 2011), which may have contributed to the 0.69‰ lower δ13C in 2007 to 2008 based on the previously reported inverse relationships between δ13C and sea surface temperature (Elson & Gillanders 2002). A direct comparison of δ13C with ambient seawater conditions may be difficult since otolith δ13C is also a product of diet, metabolism, and kinetic effects (Høie et al. 2003). With respect to δ18O, several studies have used δ18O in coral skeleton to assess inter-annual changes in sea surface salinity during ENSO events in the western Pacific and found years with decreased precipitation (and high evaporation) resulted in higher salinities and enriched δ18O in seawater and coral skeleton (δ18O of coral) (Morimoto et al. 2002, Iijima et al. 2005). Specifically, changes in sea surface salinity of approximately 0.5 resulted in a corresponding δ18O change of approximately half (0.25‰) in δ18O of seawater and δ18O of coral. Salinity differences between years in the Philippines averaged 0.5 (Gordon et al. 2011) and thus the δ18O difference of 0.2‰ was between published models using δ18O values predicting 0.1% (Kerr et al. 2007) and δ18O of coral of 0.3‰ (Morimoto et al. 2002) per 0.5 psu difference. Estimates of nursery origin using δ13C and δ18O for sub-adult yellowfin tuna collected in the nearshore Hawaiian Islands indicate these individuals were primarily derived from local production. Our estimate of over 90% of sub-adults collected in the nearshore Hawaiian Islands from the same nearshore nursery suggests that retention is high and movement of YOY yellowfin tuna away from these waters may be limited. The hypothesis of local retention or limited movement of yellowfin tuna in the Hawaiian Islands is supported by recent tagging studies. Adam et al. (2003) reported that a large percentage (97%) of yellowfin tuna tagged in the nearshore Hawaiian Islands were later recaptured in the same region, and previous work by Itano & Holland (2000) documented that displacement distances of YOY and sub-adult yellowfin tuna throughout the Hawaii exclusive economic zone (EEZ) was typically under 50 km with individuals remaining within this zone. In addition, tracks derived from archival tagged yellowfin tuna in the main Hawaiian Islands have shown localized movements around a single island or from one island to another with no significant displacements away from the main island group (K. Holland unpubl. data). Several studies have suggested that localized and restricted movement of these fish may be linked to the quality and quantity of available habitat, primarily natural bathymetric features (reef ledges, seamounts) and fish aggregation devices (FADs) (Holland et al. 1990, Brill et al. 1999, Itano & Holland 2000, Dagorn et al. 2007). In addition, prey availability found in the nearshore waters of the Hawaiian Islands is enriched by a diverse land-associated mesopelagic boundary community of vertically migrating nekton located close to the main Hawaiian Islands, supporting the expectation that these waters serve as productive foraging areas for YOY and sub-adult yellowfin tuna (Reid et al. 1991, Brock 1985). Although otolith chemistry and tagging results support the premise of local production and retention in the nearshore waters of Hawaii, both also indicate the potential for low levels of exchange with nurseries to the south. Otolith chemistry data indicated that sub-adult yellowfin tuna collected in the nearshore Hawaiian Islands had minor contribution estimates from the 2 closest nurseries to the south (offshore Hawaiian Islands, Line Islands), which is not unexpected given that limited exchange has been reported between the nearshore Hawaiian Islands and these regions (Adam et al. 2003). All potential nurseries for yellowfin tuna throughout the WCPO were not sampled for baselines; however, spatial coverage of nurseries sampled likely provided sufficient resolution of δ13C and δ18O patterns needed for estimating contribution estimates.

Enriched δ18O values in otolith cores of the sub-adult yellowfin tuna sampled from the offshore Hawaiian Island nursery closely matched the YOY nearshore Hawaiian Islands baseline, suggesting that
a large fraction (90%) of the sub-adults obtained from the offshore Hawaiian Islands originated from the nearshore Hawaiian Islands. While the majority of yellowfin tuna tagged in nearshore waters of the Hawaiian Islands have shown restricted movement, some individuals were recaptured outside of the main Hawaiian Islands farther south (ca. 200 to 500 km) from tagging locations (Itano & Holland 2000, D. G. Itano unpubl. data). Nearshore to offshore movement of yellowfin tuna in the Hawaiian Islands has been documented with fish tagged near the nearshore Hawaiian Islands later recaptured south at the Cross Seamount (offshore Hawaiian Island nursery) (Adam et al. 2003). Sibert et al. (2000) reported that migrants to the Cross Seamount were largely derived from other unknown source(s), with these individuals moving to other locations in the WCPO after frequenting this location for relatively short periods of time. Our findings clearly show that an important source of recruits to the south of the Hawaiian Islands may be the nearshore location for relatively short periods of time. Our results suggest that the contribution of equatorial nurseries to Hawaii’s domestic fisheries may be more limited (<10%) than previously assumed, but show that a south to north immigration route occurs for yellowfin tuna, which until now has only been supported by a single long distance tag recovery (Secretariat of the Pacific Community unpubl. data). Relative contribution rates of recruits from local and more distant nurseries to the south of the Hawaiian Islands may experience strong inter-annual variability and because our research was based on a single year class of sub-adults, additional research is warranted to determine whether local production and retention of yellowfin tuna in the Hawaiian Islands varies significantly across cohorts.

Acknowledgements. Funding for this work was provided by the University of Hawai‘i Pelagic Fisheries Research Program (JIMAR project 651106 to J.R.R. and D.G.I.). R.J.D.W. was supported in part by a Texas Institute of Oceanography postdoctoral research fellowship. We thank the numerous individuals for assistance in otolith collections, including G. Castrence, P. Conley, J. Dettling, B. Fukuda, D. Fuller, E-J Kim, K. Lind, J. Muir, B. Muller, K. Pollock, K. Schaefer, S. Tobiason, and T. Usu. Special thanks to R. Schloesser for assistance processing otolith samples.

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