CONTINENTAL-SCALE VARIABILITY IN THE FEEDING ECOLOGY OF JUVENILE CHINOOK SALMON ALONG THE COASTAL NORTHEAST PACIFIC OCEAN

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ABSTRACT: Trophic interactions within and among species vary widely across spatial scales and species’ ontogeny. However, the drivers and implications of this variability are not well understood. Juvenile Chinook salmon Oncorhynchus tshawytscha have a wide distribution, ranging from northern California to the eastern Bering Sea in North America, but it is largely unknown how their feeding ecology varies and changes with ontogeny across this range. We collected juvenile Chinook salmon and zooplankton using standardized protocols along the coastal Northeast Pacific Ocean. Using a combination of stomach contents and stable isotopes of nitrogen (δ¹⁵N) and carbon (δ¹³C) to characterize feeding ecology, we found regional differences in prey utilization by juvenile Chinook salmon. With growth and ontogeny, juvenile salmon in all regions became equilibrated with oceanic isotopic values. There were regional differences in the δ¹³C values of juvenile Chinook salmon that may correspond to regional differences in sea surface temperature. There were also regional differences in stable isotope-derived trophic level, and these estimates differed from those derived from stomach contents, possibly due to the different periods over which these metrics integrate. Dietary niche width, as indicated by stable isotopes, corresponded to the expected dietary diversity from stomach contents, combined with the isotopic variability seen in baseline values. Our results indicate strong geographic and ontogenetic differences in feeding ecology of juvenile Chinook salmon. These differences are likely influenced by a combination of ocean-entry date, ocean-entry size, ontogeny, growth rates and regional conditions.

KEY WORDS: Diet · Stable isotope · Trophic level · Turnover · Diet dependent discrimination factor · Oncorhynchus tshawytscha · Carbon · Nitrogen · Niche width · Ontogeny

INTRODUCTION

Understanding trophic interactions within and among species is a central theme in ecology, and is increasingly important as fisheries science moves towards ecosystem based management (Larkin 1996, Pikitch et al. 2004). Due to the complexity of modeling entire ecosystems, current approaches generally assume that trophic interactions are static, often ignoring significant spatial and temporal variability in trophic dynamics. For example, the same species can have divergent niches in different habitats.
(McCann et al. 2005), but the causes and consequences of these niche differences are largely unknown.

Further complicating the study of trophic interactions are the ontogenetic shifts in feeding ecology common in many species (Werner & Gilliam 1984). These ontogenetic niche shifts may have widespread effects on population and community dynamics (de Roos & Persson 2013), such as promoting the coexistence of competitors (Wollrab et al. 2013), reducing the stabilizing effects of ecosystem complexity (Rudolf & Lafferty 2011), and even altering the strength of trophic cascades (Rudolf & Rasmussen 2013). Variations in trophic interactions as a result of ontogenetic niche shifts are beginning to be considered in experimental and modeling food web studies (van Leeuwen et al. 2014), but there are few empirical examples of variation in species ontogeny in field settings across large geographic scales.

Ontogenetic niche shifts are prevalent in fish, especially those that are piscivorous. Juvenile fish may be gape limited, and thus must feed on small prey items at small sizes (Nunn et al. 2012). With growth, maximum and minimum prey sizes generally increase, though the rate of change of these relationships may be modulated by species and habitat (Scharf et al. 2000, Keeley & Grant 2001). These ontogenetic shifts to larger prey items may be important because growth efficiency is higher and metabolic costs are lower when feeding on larger prey (Pazzia et al. 2002), and fish prey are generally higher in caloric value (Davis et al. 1998).

Stomach content analysis and stable isotope analysis (SIA) are 2 common methods used to track ontogenetic shifts, each with its own inherent limitations and assumptions. Stomach contents can give a great deal of taxonomic resolution in diet, but only represent a snapshot of diet in time, and can be biased by differences in the digestibility of prey items (Polunin & Pinnegar 2002). Conversely, stable isotopes in the muscle tissue of an organism represent material assimilated over a period of weeks to months (Fry 2006). Stable isotopes of nitrogen ($\delta^{15}N$) generally indicate trophic position, as $\delta^{15}N$ undergoes a trophic enrichment of approximately 3.4‰ per trophic level (Post 2002), although recent studies have indicated that the value of this trophic enrichment may decrease with increasing dietary $\delta^{15}N$ (Caut et al. 2009, Hussey et al. 2014). $\delta^{15}N$ values at the base of the food web are also variable, such that estimates of consumer trophic level are often calculated relative to a baseline primary consumer (Cabana & Rasmussen 1996, Hussey et al. 2014). For stable isotopes of carbon ($\delta^{13}C$), there is a general onshore/offshore pattern in coastal waters, with onshore waters being enriched in $\delta^{13}C$ by up to 5‰ (Perry et al. 1999, Miller et al. 2008). This pattern may be due to differences in the productivity of phytoplankton (Schell 2000, Miller et al. 2008) as the fractionation of $\delta^{13}C$ in phytoplankton is related to species and growth rate, with higher $\delta^{13}C$ values associated with larger cell sizes and greater growth rates (Laws et al. 1995). Temperature can also affect $\delta^{13}C$ values, since the amount of dissolved CO$_2$ in surface waters is inversely related to sea surface temperature (SST; Weiss 1974). These higher concentrations of dissolved CO$_2$ lead to lower $\delta^{13}C$ values at lower SSTs (McMahon et al. 2013). So, while stable isotopes can indicate trophic level ($\delta^{15}N$) and source of production ($\delta^{13}C$), there is generally an overall lower taxonomic resolution than with stomach contents, and the assumptions must be made that there is a known trophic enrichment, and that the organism is at equilibrium with the isotopic baseline of the environment (Buchheister & Latour 2010). Thus overall, a powerful method to trace diet and ontogeny would be to use both stomach contents and SIA.

Along the west coast of North America, juvenile Chinook salmon *Oncorhynchus tshawytscha* are generalist predators that feed on a variety of juvenile fish and invertebrate prey (Brodeur et al. 2007, Daly et al. 2009). Chinook salmon are anadromous, and upon ocean entry begin feeding on invertebrates before shifting to preying primarily on fish (Brodeur 1991). Since mortality for juvenile Chinook salmon may be size-selective in their early marine life (Clairborne et al. 2011, Duffy & Beauchamp 2011, Woodson et al. 2013), the shift to feeding on higher-quality fish prey may be important for their overall survival rates.

Juvenile Chinook salmon show variation in diet at various spatial scales throughout their range (Brodeur et al. 2007, Davis et al. 2009, Duffy et al. 2010). In North America, the spawning range of Chinook salmon spans 3 oceanographic domains: the eastern Bering Sea shelf, the Alaska Coastal Current System, and the California Current System. In the eastern Bering Sea shelf, primary productivity and food web structure is related to ice cover (Hunt et al. 2002), with north–south and cross-shelf variations (Brown et al. 2011, Eisner et al. 2014), and a high abundance of fish prey in the pelagic zone (Farley et al. 2005). The Alaska Coastal Current is a downwelling system (Ware & McFarlane 1989), with high prey quality (Lee et al. 2006) and a fish community composed largely of salmonids (Orsi et al. 2007). The California Current System is an upwelling system
(Hickey 1979, Ware & McFarlane 1989), with relatively lower prey quality (Lee et al. 2006) and higher biomass of other fish species relative to juvenile salmon (Orsi et al. 2007). Prey quantity varies on several spatial and temporal scales throughout this range, though on an annual basis the highest primary and secondary productivity occurs off the coast of Vancouver Island at the north end of the California Current System (Ware & Thomson 2005, Hickey & Banas 2008). Combined, these regional differences have been hypothesized to affect the feeding habits of juvenile Chinook salmon (Brodeur et al. 2007).

Independent of regional oceanography and community composition, size differences of juvenile salmon may also contribute to diet differences among regions. At ocean entry, juvenile Chinook salmon range in size from an average of 75 mm on the West Coast of Vancouver Island to 160 mm in Oregon and Washington (Trudel et al. 2007). As larger juvenile salmon generally tend to be more piscivorous (Brodeur 1991), this regional variation in size may have implications for diet. The diet shift that occurs with size also varies by region, with clear ontogenetic shifts to piscivory in Oregon and Washington (Brodeur 1991, Daly et al. 2009), but little evidence of ontogenetic shifts in southeast Alaska (Weitkamp & Sturdevant, 2008). Of note, previous studies on the ontogeny of juvenile Chinook salmon have generally reported diet from only a single region or sampling program, and used only one metric of diet to observe ontogenetic niche shifts.

In this study, our main objective was to examine regional variability in the feeding ecology and ontogeny of juvenile Chinook salmon. To do so, we assembled the largest data set available on stable isotopes and stomach contents of salmon from northern California to the eastern Bering Sea. These data were collected during 1 yr and 1 season to minimize temporal variability. First, to account for variability in ontogeny, size-selective feeding, and latitude, we determined how the $\delta^{13}C$, $\delta^{15}N$, and stomach contents of juvenile salmon vary by region and body size. Because previous studies on the stomach contents of juvenile Chinook salmon have indicated that they are generally more piscivorous in the eastern Bering Sea, Southeast Alaska, and Oregon/Washington than in other regions (Brodeur et al. 2007), we hypothesized that these regions would be similarly higher in stable isotope-derived trophic levels. In contrast, because there is a general south to north decline in SST, we hypothesized that $\delta^{13}C$ values would be highest in the southern regions, and become increasingly lower in the north. We concluded the study by combining both diet approaches to examine variation on a continental scale, and specifically, to determine whether there was concordance between long-term assimilated diet as indicated by stable isotopes, and the snapshot of recent diet as indicated by stomach contents.

**MATERIALS AND METHODS**

**Field sampling and laboratory analysis**

Samples were collected off the coasts of California (CA), Oregon and Washington (ORWA), the west coast of Vancouver Island in British Columbia (WCVI), central British Columbia (CEBC), southeast Alaska (SEAK), southeastern Bering Sea (SEBS), and the northeastern Bering Sea (NEBS) (Fig. 1). NEBS and SEBS were separated along the 60°N latitude line, based on the distribution and migration routes of juvenile salmon (Farley et al. 2009). In NEBS and SEBS, we retained juveniles that were less than 325 mm, as fish larger than this in the fall are likely immature. All sampling was performed during the fall of 2007 (California: mid-August, ORWA: September, WCVI: October–November, CEBC: October–November, SEAK: October–November, SEBS: August–September, NEBS: August–September). Sampling was carried out in the fall because the stock composition of fall samples tends to better represent the region-of-capture (Tucker et al. 2011, 2012). Sampling in the fall also allows juveniles more time to move offshore (where they are available to surface trawls) and allows salmon more time to become equilibrated with oceanic baselines and lose freshwater isotopic signatures. Though they represent various freshwater life-history strategies, all juvenile Chinook salmon in this study entered the ocean sometime in the spring or summer of 2007 (Trudel et al. 2007). All programs used similar surface trawls towed behind large research or fishing vessels to collect the salmon; therefore, differences in sampling gear among regions are not expected to significantly bias our results. Once salmon were brought aboard the research vessel, they were euthanized, identified, measured, weighed and frozen for subsequent analysis.

Zooplankton samples were collected concurrently with salmon to use as an isotopic baseline in all regions but CA and ORWA. Though juvenile Chinook salmon do not typically feed directly on zooplankton prey, these organisms integrate the isotopic variability that occurs in phytoplankton, and serve as an effective baseline proxy for higher trophic levels.
In WCVI, CEBC, and SEAK, these samples were taken by vertical bongo tows (236 μm black mesh), to 150 m or within 10 m of the ocean floor. Juvenile Chinook salmon do not typically forage at these depths, but many zooplankton undergo diel vertical migrations that bring them into the range of the salmon at their highest feeding intensity periods of dawn and dusk (Benkwitt et al. 2009). Samples were size-fractionated and the smallest size fraction (0.25 to 1.7 mm) was used, due to the greater sampling coverage of this size fraction (El-Sabaawi et al. 2013). Zooplankton samples in SEBS and NEBS were sampled similarly (but using 335 μm mesh), and then filtered to same size fraction as other regions (Coyle et al. 2011). A total of 124 size-fractionated zooplankton samples were analyzed.

Stomach contents of juvenile Chinook salmon were analyzed following Brodeur et al. (2007). Briefly, stomach contents were preserved in formalin, examined under a dissecting microscope and identified to the lowest possible taxonomic level. Prey items were grouped into 17 larger categories (taxonomic groupings) for statistical analyses. The categories used were: unidentifiable fish, northern anchovy Engraulis mordax, Pacific sand lance Ammodytes hexapte-
rus, Pacific herring Clupea pallasii, capelin smelt Mallotus villossus, unidentified smelt (Osmeridae), walleye pollock Gadus chalcogrammus, sculpin (Cottidae), rockfish Sebastes spp., poacher (Agonidae), unidentified flatfish (Pleuronectidae), euphausid, decapod, amphipod, cephalopod, insect, and other. Stomach contents from CA, ORWA, SEBS and NEBS were assessed using % composition by weight at an individual level. In these regions, prey items were weighed after blotting dry. The data from these regions were then pooled by tow to prevent individual taws with large catches overwhelming any particular region and to facilitate comparison with WCVI, CEBC and SEAK, where stomach contents were assessed using % composition by volume, and pooled by tow. Stomach content data was derived from 1046 juvenile Chinook salmon, with regional sample sizes ranging from 14 for CA to 332 for WCVI (see Fig. 2).

A random subset of 949 juvenile Chinook salmon were analyzed for stable isotopes of δ13C and δ15N. Regional sample size ranged from 13 juveniles in CA to 306 in WCVI (see Table S1 in the Supplement at www.int-res.com/articles/suppl/m537p247_supp.pdf). A piece of dorsal muscle tissue was taken from each juvenile salmon and freeze-dried. Zooplankton samples were also freeze-dried. Samples were ground to a fine powder and placed into tin capsules. A Thermo Delta IV Isotope Ratio Mass Spectrometer (at the University of Victoria) was used for the determination of stable isotope values. Atmospheric nitrogen was used as the standard for δ15N and Vienna PeeDee Belemnite was used as the standard for δ13C. Stable isotope values are expressed in the delta notation:

\[ \delta^{15}N \text{ (or } \delta^{13}C) = \frac{R_{\text{sample}}}{R_{\text{standard}} - 1} \times 1000 \]  

where \( R \) is 15N:14N or 13C:12C.

We did not lipid-normalize the juvenile Chinook salmon samples or zooplankton baselines due to the possibility of taxonomic differences in the effects of lipids on the δ13C values of samples (Syväranta & Rautio 2010, Fagan et al. 2011). However, results were qualitatively similar when using mathematical lipid-corrections derived for both zooplankton and fish (not shown).

**Statistical analysis**

Juvenile Chinook salmon stomach contents

We calculated trophic level (TL) from stomach contents data using the equation in Mearns et al. (1981):
Zooplankton stable isotopes

To test for regional differences in the $\delta^{13}C$ and $\delta^{15}N$ of zooplankton, we used a Kruskal-Wallis test, since variances among groups were unequal (Levene’s test: $p < 0.05$). Differences among regions were assessed using the ‘kruskalmc’ post hoc test in the R package ‘pgirmess’ (Giraudoux 2014).

Juvenile Chinook salmon stable isotopes

Variation in baseline $\delta^{15}N$ can obscure differences in an organism’s $\delta^{15}N$ independent of the variability caused by diet (Cabana & Rasmussen 1996), so we converted all juvenile Chinook salmon $\delta^{15}N$ values to trophic positions. There appears to be an inverse relationship between $\delta^{15}N$ in the diet of an organism and the trophic enrichment that an organism experiences (Caut et al. 2009). This relationship indicates that by assuming a constant trophic enrichment, there can be an underestimation in the TL of higher TL organisms (Hussey et al. 2014). Thus, we used this scaled approach to calculate TL, rearranged from Hussey et al. (2014):

$$\text{TL}_{\text{Hussey}} = \log \left( \frac{\delta^{15}N_{\text{lim}} - \delta^{15}N_{\text{base}}}{\frac{\delta^{15}N_{\text{lim}} - \delta^{15}N_{\text{fish}}}{\delta^{15}N_{\text{lim}} - \delta^{15}N_{\text{fish}}}^{\frac{1}{k}}} \right)^{\frac{1}{k}} + \text{TL}_{\text{base}} \quad (4)$$

where $\text{TL}_{\text{Hussey}}$ is the trophic level of juvenile Chinook salmon, $\delta^{15}N_{\text{lim}}$ is the limit of $\delta^{15}N$ values as TL increases and $k$ are fitted parameters from the meta-analysis by Hussey et al. (2014), $\delta^{15}N_{\text{base}}$ is the baseline $\delta^{15}N$ value from zooplankton samples, $\delta^{15}N_{\text{fish}}$ is the $\delta^{15}N$ of the juvenile Chinook salmon sampled, and $\text{TL}_{\text{base}}$ is the trophic level of the baseline organism chosen. We assumed that the zooplankton samples from all regions were TL 2. To compare standard TL estimates to this scaled approach, we also calculated TL following Cabana & Rasmussen (1996), with a constant trophic enrichment of 3.4‰ (Post 2002).

To observe how isotopes changed with size in each region, we plotted the $\delta^{13}C$, $\delta^{15}N$ and TL values of juvenile Chinook salmon in each region as a function of mass. We used a formulation of the logistic model to model the processes of isotopic turnover and ontogenetic niche shifts in each region:

$$\delta_{xi} = \frac{\alpha}{1 + e^{-\beta - \theta \cdot W_{i}}} \quad (5)$$

where $\delta_{xi}$ are the individual isotopic values of either $\delta^{15}N$ or TL, $\alpha$ is the asymptotic value reached at equilibrium, $\beta$ is the inflection point, $W_{i}$ is the mass of each fish at capture and $\theta$ is a scaling parameter.

This formulation of the logistic model does not fit negative values well, so to fit $\delta^{13}C$ models, we first multiplied all $\delta^{13}C$ values by $-1$, then used

$$\delta_{xi} = \frac{\alpha}{1 - e^{-\beta - \theta \cdot W_{i}}} \quad (6)$$

We were interested in whether there were regional differences in the isotopic values at equilibrium, so we used a non-linear mixed effects (NLME) modeling approach (Pinheiro & Bates 2000). We compared a model with no random effects (i.e. all parameters were the same in all regions) to models where $\alpha$ or $\beta$ were the random effects (i.e. these parameters varied by region). We also compared these models to a model where all 3 parameters were random effects (i.e. all varied by region). We used Akaike’s information criterion (AIC) to determine which model was most supported by the data. Due to the lack of small fish in SEBS and NEBS, for the $\delta^{15}N$ and $\delta^{13}C$ models we simply assumed that there was a linear relationship between isotopes and size, with a slope of 0, and intercept equal to the mean. For TL models, we also had to exclude CA and ORWA, as these regions lacked the zooplankton baseline data needed to calculate TL. We used the R package ‘nlme’ to fit all models (Pinheiro et al. 2014).

Finally, recent approaches examining variation in isotopes at population-level scales have emphasized the power of simultaneously looking at variation in $\delta^{15}N$–$\delta^{13}C$ bivariate space (Layman et al. 2007, Jackson et al. 2011). However, because ocean entry size, time since ocean entry, and growth rates vary among regions (Trudel et al. 2007), the amount of tissue that has been turned over in each region is different. To minimize differences caused by these factors, we retained juvenile Chinook salmon that were predicted to be within 95% of their asymptotic weight. To do so, we set $\delta_{\text{eq}}$ to be 95% of the regional asymptotic value, and solved Eq. (6) using the parameters from our best fit NLME model for $\delta^{13}C$ (since $\delta^{13}C$ had a slower turnover time than $\delta^{15}N$; see Fig. 4), retaining all juvenile Chinook salmon larger than this value (hereafter equilibrated juvenile Chinook salmon). We also retained all SEBS and NEBS juvenile Chinook salmon for this analysis, as the lack of a size effect on the isotopic ratios suggest that they were equilibrated with their prey in these regions. Overall, there were 386 juvenile Chinook salmon that fitted our operational definition for equilibrium, with sample size of equilibrated juvenile Chinook salmon ranging from 7 in CA to 144 in SEAK (Table S1). Due to low sample sizes of equilibrated
juvenile Chinook salmon in CA, this region was not included in subsequent analyses.

In bivariate $\delta^{13}$C–$\delta^{15}$N space, we tested for differences in isotopic niche position among regions using the residual permutation procedures outlined in Turner et al. (2010). This method is based on Euclidean distance measures of each individual’s position in bivariate $\delta^{13}$C–$\delta^{15}$N space (Turner et al. 2010). We ran the procedure for 1000 iterations. We calculated $\delta^{15}$N and $\delta^{13}$C ranges for both zooplankton and equilibrated juvenile Chinook salmon following Layman et al. (2007). To test for differences in the isotopic niche breadth among regions, we used stable isotope Bayesian ellipses in R (SIBER; Jackson et al. 2011). This method constructs the parameters of ellipses from each region using Markov Chain Monte Carlo simulations, and provides an estimate of the average size of the isotopic niche that is relatively insensitive to sample size and outliers (Jackson et al. 2011). Baseline variability can also contribute to niche differences between regions (e.g. Hoeinghaus & Zeug 2008), so we also used SIBER to explore the variability present in the zooplankton baseline of different regions. All analyses were performed in the statistical language R (R Development Core Team 2013).

Environmental drivers of variability

Finally, we tested the effects of SST on the average $\delta^{13}$C values of zooplankton and juvenile Chinook salmon. Monthly average SST data were obtained in a 1° latitude × 1° longitude grid resolution from the NOAA OI.v2 SST data (Reynolds et al. 2002). These data are a combination of satellite and in situ measurements, and were obtained from the NOAA National Centers for Environmental Prediction (ftp://ftp.emc.ncep.noaa.gov/cmb/). In each region, over the latitudes where juvenile Chinook salmon were caught, we averaged the SST in the 1 × 1° block nearest to the coast over the period from May to October. In SEBS and NEBS, juvenile Chinook salmon were distributed over a broader area, therefore we extended our analysis to the maximal longitude where they were caught. The relationships between SST and zooplankton $\delta^{13}$C and equilibrated juvenile Chinook salmon $\delta^{13}$C were assessed using separate linear regressions.

RESULTS

Juvenile Chinook salmon stomach contents

Stomach content analysis showed regional differences in the feeding of juvenile Chinook salmon (Fig. 2). All pairs of regions were significantly different from each other (ANOSIM; p < 0.05) with the exception of SEBS & NEBS, and CEBC & SEAK. Juvenile salmon in ORWA, SEAK and NEBS were predominately piscivorous, with different fish prey in each region. Northern anchovy and unidentified smelt were highly consumed in ORWA, Pacific herring and Pacific sand lance in SEAK, and capelin smelt and Pacific sand lance were the dominant fish prey in NEBS (Fig. 2; SIMPER). Pacific herring was the primary fish prey in CEBC and WCVI, and capelin smelt was the common fish prey in SEBS along with walleye pollock. Juvenile rockfish and euphausiids were the top prey in CA, Pacific sand lance was consumed in each region north of WCVI, and amphipods were important in all regions except ORWA and SEAK (SIMPER). There were a large amount of unidentifiable fish in most regions, comprising 40 to 60% of total weight or volume in CA, ORWA, and SEAK (Fig. 2).

![Fig. 2. Regional composition of stomach contents of juvenile Chinook salmon for the grouped prey categories. The number of stomachs examined (N) is given at the top of each bar. See Fig. 1 legend for region abbreviations](image-url)
Stomach content data indicated that Levins’ niche breadth was largest in SEBS and WCVI (7.4 and 6.2 respectively). Niche breadth was intermediate in NEBS, CEBC and CA, with values of 5.1, 4.2 and 3.4, and was lowest in SEAK (2.7) and ORWA (2.3).

The NMDS showed that the diet compositions of SEBS and NEBS were distinct from the other regions, especially along axis 1, and encompassed a broad ordination space relative to the other regions, whereas diets in ORWA, SEAK, and most of WCVI showed much less variability and were closely grouped (Fig. 3). These patterns match closely to those shown by Levins’ niche breadth. In the NMDS, anchovy and rockfish loaded positively onto axis 1, while capelin loaded negatively on this axis. On axis 2, amphipods and pollock loaded positively, while sand lance and herring loaded negatively (Fig. 3). We found no evidence for ontogenetic niche shifts with size in any region from stomach content data, irrespective if the data were pooled by station or examined on an individual basis (p > 0.05; Fig. S1 in the Supplement at www.int-res.com/articles/suppl/m537p247_supp.pdf).

Zooplankton stable isotopes

Zooplankton δ15N was significantly different among regions (Kruskal-Wallis $\chi^2 = 63.1$, df = 4, p < 0.0001). The WCVI δ15N values were significantly lower compared to all regions except for CEBC (Table S2). Zooplankton δ13C was also significantly different among regions (Kruskal-Wallis $\chi^2 = 68.3$, df = 4, p < 0.0001). WCVI, CEBC and SEAK were generally more enriched than SEBS or NEBS (Table S2). Finally, the C:N ratios of zooplankton were significantly different between regions (Kruskal-Wallis $\chi^2 = 87.4$, df = 4, p < 0.0001) with values above 7 in SEBS and NEBS, and lowest in CEBC with a value near 4 (Table S2). The comparisons between SEAK and WCVI, and SEBS and NEBS, were the only non-significant differences.

Juvenile Chinook salmon stable isotopes

The average mass at capture of juvenile Chinook salmon was smallest in ORWA at 51.6 g (158 mm fork length) and largest in SEAK at 251.7 g (260 mm fork length) (Fig. 1). The plots of isotopes against mass show that in most areas, the juvenile salmon rapidly shift their diet and turn over tissue until they reach a plateau at approximately 85 g (roughly 200 mm) (Fig. 4; see Fig. S2 for plots by fork length). The isotopic value at which this plateau is reached appears to be different among regions; the AIC values of the NLME models confirm this, in that best model fits for each variable included $\alpha$ as a random effect (i.e. varies among regions) (Table 1). For δ13C, the best model included only $\alpha$ as a random effect, with $\beta$ and $\theta$ as fixed effects. $\alpha$ values were highest in WCVI at −15.9‰, and lowest in SEAK at −17.8‰ (Fig. 4). For δ15N and TLHussey, the best models included $\alpha$, $\beta$ and $\theta$ as random effects. For the best δ15N model, $\alpha$ values were highest in WCVI at 15.1‰ and lowest in CA at 12.8‰ (Fig. 4). For TLHussey, WCVI had the highest $\alpha$ value of 4.1, and CEB had the lowest of 3.3. For SEBS and NEBS, the mean δ15N values were more enriched and the mean δ13C values were more depleted than the predicted $\alpha$ values from all other regions (Table 2). TL estimates from stomach contents and stable isotopes were similar for NEBS and SEBS (Fig. 5). The predicted TL from SIA was lower than that predicted from stomach content analysis in CEB and SEAK, with isotopes predicting a TL approximately 0.3 to 0.5 TLs below that of the stom-
Fig. 4. Regional relationships between the stable isotopes of nitrogen and carbon ($\delta^{15}\text{N}, \delta^{13}\text{C}$) and size of juvenile Chinook salmon. See Fig. 1 legend for region abbreviations. Solid lines: the best non-linear mixed effects (NLME) model fits for each region (except for NEBS and SEBS, where averages are displayed due to the lack of a size effect); dashed lines: zooplankton baseline values in the regions where they were sampled; shaded area: ±1 SD.
ach contents. The pattern was the opposite in WCVI, where stable isotopes predicted a TL approximately 0.5 TL above stomach contents (Fig. 5).

**Equilibrated juvenile Chinook salmon stable isotopes and baseline variability**

Using the residual permutation procedure in Turner et al. (2010), we determined that the mean centroid location of each region differed from zero (p = 0.001). This suggests that each region occupied a unique area in isotopic niche space (Fig. 6c). Comparing the relative ellipses for each region using SIBER showed that the largest niche area was in NEBS (p < 0.01) (Fig. 6d). SEBS also had a significantly larger niche area than SEAK (p < 0.01), but niche area did not differ significantly among the other regions.

We also used SIBER to explore the variability present in the baseline of different regions because

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**Table 1. Results of fitting various non-linear mixed effect models on the relationships between δ¹³C, δ¹⁵N (stable isotopes of carbon and nitrogen), TLHussey (trophic level of juvenile Chinook salmon based on Hussey et al. 2014) and size. NLL: negative log-likelihood; AIC: Akaike’s information criterion; α: asymptotic value reached at equilibrium; β: inflection point; θ: scaling parameter. The best model for each variable is shown in **bold.**

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<td>-1140.5</td>
<td>2301.1</td>
<td>3.5</td>
<td>10</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Table 2. Parameter values of best fit non-linear mixed effect models for the logistic change in isotopic values of juvenile Chinook salmon. α: asymptotic value reached at equilibrium; β: inflection point; θ: scaling parameter. TLHussey: trophic level of juvenile Chinook salmon based on Hussey et al. (2014). See Fig. 1 legend for region abbreviations.**

<table>
<thead>
<tr>
<th>Isotope</th>
<th>Region</th>
<th>α</th>
<th>β</th>
<th>θ</th>
<th>NLL</th>
<th>AIC</th>
<th>ΔAIC</th>
<th>df</th>
</tr>
</thead>
<tbody>
<tr>
<td>δ¹³C</td>
<td>NEBS</td>
<td>16.3</td>
<td>27.9</td>
<td>69.6</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>α</td>
<td>SEBS</td>
<td>-20.4</td>
<td>69.6</td>
<td>69.6</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>β</td>
<td>SEAK</td>
<td>-17.8</td>
<td>69.6</td>
<td>69.6</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>α, β, θ</td>
<td>ORWA</td>
<td>-17.3</td>
<td>69.6</td>
<td>69.6</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>CA</td>
<td>CA</td>
<td>12.8</td>
<td>1.2</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>CEBC</td>
<td>CEBC</td>
<td>3.3</td>
<td>37.3</td>
<td>37.3</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>WCVI</td>
<td>WCVI</td>
<td>3.8</td>
<td>36.3</td>
<td>36.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Estimated by taking the mean value of juvenile Chinook salmon in corresponding regions

---

**Fig. 5. Regional relationships between trophic level (TLHussey: trophic level based on Hussey et al. 2014) and size of juvenile Chinook salmon. Solid lines: the best non-linear mixed effects (NLME) model fits for each region for TLHussey; dashed line: TL estimate from stomach contents (TL<sub>stom</sub>)**
baseline variability can confound large-scale comparisons. We found that the variability seen in juvenile Chinook salmon was largely similar to the regional differences in variability in zooplankton (Fig. 6a,b). Equilibrated juvenile Chinook salmon were generally enriched in $\delta^{13}$N relative to zooplankton by approximately 3 to 5‰ (Fig. 6a,c). The $\delta^{13}$C values of equilibrated juvenile Chinook salmon were generally enriched by over 2‰ relative to zooplankton $\delta^{13}$C values (Fig. 6a,c). Similar to juvenile salmon, zooplankton in NEBS and SEBS had the significantly largest niche areas, CEBC and SEAK had the smallest niche areas, and WCVI was intermediate (Fig. 6b). Interestingly, the niche variation present in zooplankton was dampened in the juvenile Chinook salmon, with each region showing much greater variation in niche areas at the zooplankton rather than the salmon level (Fig. 6a,c). The results from the $\delta^{15}$N and $\delta^{13}$C ranges showed largely the same patterns as those shown by the SIBER plots (Table S3).

**Environmental drivers of variability**

Average SST varied from lows of 6.7 and 6.6°C in NEBS and SEBS respectively, to a high of 13.8°C in ORWA. The relationship between average SST and zooplankton $\delta^{13}$C was significant ($p = 0.005$) and positive (Fig. 7). Similarly, the relationship between average SST and equilibrated juvenile Chinook salmon $\delta^{13}$C was significant ($p = 0.004$) and positive, though the relationship appears to be largely driven by the very low values in SEBS and NEBS (Fig. 7).
DISCUSSION

Our analyses showed large regional differences in the feeding ecology of juvenile Chinook salmon along the west coast of North America. The asymptotic \( \delta^{13}C \) and \( \delta^{15}N \) values of juvenile Chinook salmon varied by region, and these trends may coincide with regional variability in oceanography. Stomach contents and SIA indicated somewhat dissimilar results, especially with regards to ontogeny and TL estimates. While previous studies have assessed the stomach contents of juvenile Chinook salmon on a similar scale (Brodeur et al. 2007), this is the first study to combine multiple approaches to assess the ontogeny of feeding ecology on a continental scale.

The stomach content data for 2007 presented here matches the same general pattern as that of Brodeur et al. (2007). That is, juvenile Chinook salmon are generally piscivorous in most areas, but in coastal British Columbia and CA, other prey can make up a significant portion of their diet. In 2007, the non-fish prey that made up the remainder of the diet in CA were euphausiids, which contrasts with the period of 2000 to 2002, when juvenile Chinook salmon in this area fed mainly on cephalopods (Brodeur et al. 2007). For the eastern Bering Sea salmon we found similar patterns to Farley et al. (2009), who reported that fish were important components of the diets of juvenile Chinook salmon from 2003 to 2006. Interestingly, while Farley et al. (2009) found that the juvenile salmon from SEBS were generally more piscivorous than those from NEBS, in 2007 we found the opposite pattern. The food web structure of the Bering Sea appears to be related to SST, and 2003 to 2005 was a warm period in the Bering Sea, while 2007 was a cool year (Coyle et al. 2011). It is thus possible that the shift in ocean conditions in 2007 resulted in higher piscivory in NEBS, possibly due to shift in the distribution of age-0 walleye pollock, which were a primary prey item of juvenile salmon in warm years, but shifted their distribution offshore and deeper in cool years (Parker-Stetter et al. 2013).

The British Columbia coastal areas also showed an interesting pattern with regards to stomach contents, with salmon from both WCVI and CEB having nearly 50% of their diet made up of non-fish prey, and covering a smaller ordination space relative to those from SEBS and NEBS. Interpretation of these regional differences in stomach contents is difficult without complimentary sampling of the prey field (Brodeur et al. 2011). Furthermore, this finding may be partially due to salmon size in WCVI, where relatively small size-at-capture meant that these salmon may have been captured before the majority of them had completed their shift to piscivory. Altogether, the diet differences that we observed are probably very conservative since we had to group prey into broader categories (e.g. euphausiids, amphipods, decapods, and some of the fish groupings) due to advanced digestion in many cases, but many of these prey taxa are likely to be different between regions as they have limited geographical ranges. The advanced digestion of many prey items also calls into question the assertion that stomach contents provide a greater taxonomic resolution than stable isotopes, and echoes Baker et al. (2014) who noted that unidentifiable and inseparable digested material in stomach contents can introduce significant and unquantifiable error. Interestingly, we did not detect evidence for ontogenetic shifts in diet using stomach content data for the regions that we tested in this study. Given that previous stomach content studies with larger size ranges and sample sizes have noted strong evidence for ontogeny in ORWA (Brodeur 1991, Daly et al. 2009) and WCVI (E. Hertz et al. unpubl. data), the lack of a trend seen in our stomach content analyses is probably due to the relatively low sample sizes and smaller size ranges in this study, or the fact that we had to average stomach contents by station rather than by individual.

The relationships between size and \( \delta^{13}C \), \( \delta^{15}N \), and TL\textsubscript{Hussey} show similar asymptotic trends in most regions, and the same processes likely underlie these
trends. Chinook salmon stocks generally vary in their ocean-entry date, size and growth (Quinn 2005, Trudel et al. 2007). This means that throughout the sampling period, each individual will vary in the amount of time available for it to reach equilibrium with the marine environment, as well as have a different amount of tissue that needs to become equilibrated (Hesslein et al. 1993). These effects are likely to be larger in individuals from ORWA since this area has a larger size-at-entry, later entry date, and a wider range of size variation than other regions (Trudel et al. 2007). Overlain on this variability is the ontogenetic niche shift that occurs in juvenile Chinook salmon when they reach the marine environment. That is, as the juveniles grow, they generally become more piscivorous until they reach a region-specific plateau in the contribution of fish to the diet (Brodeur 1991, Daly et al. 2009, E. Hertz et al. unpubl. data). Furthermore, individual salmon also vary in growth rate, which can be a primary driver of tissue turnover in fast-growing juvenile fish (Buchheister & Latour 2010). Hence, this combination of date of ocean entry, amount of tissue to turn over, growth rate, and ontogeny likely determines where in this general curve each juvenile Chinook salmon would be found. On a population level, some regions with an earlier migration time (relative to the survey) may have already moved beyond this shifting isotope phase (e.g. SEAK, SEBS and NEBS). Conversely, the populations with a relatively late migration time, or large variability in release date (in the case of hatchery juvenile Chinook salmon), may not have yet reached their plateau (e.g. ORWA).

The best models for δ¹³C, δ¹⁵N, and TL_{slu}se all included α as a random effect, which indicates that there were significant differences in the regional asymptotic stable isotope values for juvenile Chinook salmon, even without including SEBS and NEBS (the 2 regions with the most unique isotopic values). We found a significant relationship between SST and the δ¹³C value for equilibrated juvenile Chinook salmon, as well as between SST and the average δ¹³C value for zooplankton. High latitude, lower-SST systems tend to have high concentrations of dissolved CO₂ due to the greater solubility of CO₂ in cooler water as well as the vertical mixing of the water column (Newsome et al. 2010). These conditions can lead to low δ¹³C values, as the fractionation associated with photosynthetic CO₂ uptake becomes greater with higher CO₂ concentrations (Laws et al. 1995, Newsome et al. 2010).

The regional differences in δ¹³C could also reflect differences in factors that affect phytoplankton growth rate, with higher δ¹³C associated with greater primary productivity (Laws et al. 1995, Schell 2000, Miller et al. 2008). This general link between primary productivity and δ¹³C is also supported by our data, as primary productivity along the coast of North America is highest off WCVI (Ware & Thomson 2005), and juvenile Chinook salmon from this area had the highest asymptotic δ¹³C values. We were unable to further directly assess this hypothesis, as chlorophyll a (chl a), the only metric of primary productivity that was available across our study range, is derived from satellite data (e.g. SeaWiFS). This is problematic for a number of reasons. Firstly, the majority of fish in WCVI, CEBC and SEAK were caught in inlets and straits, where there are no direct measures of chl a, and where satellite-derived data are unreliable. Second is the problem of cloud cover, which can be extensive in the regions of study. Finally, there is the problem of the unknown and possibly significant time lag between chl a values and the corresponding isotopic values of zooplankton and salmon. Therefore, while productivity could also be driving some of the regional variation in δ¹³C, we were unable to assess its effect in this study.

Equilibrated juvenile Chinook salmon were enriched in δ¹³C relative to zooplankton δ¹³C by values generally more than 2‰. This is greater than the typically-assumed enrichment of 0 to 1‰ (Post 2002), but is within the range of literature values (−3 to 3‰; McCutchan et al. 2003). The fact that lipids were not removed from samples may explain some of this discrepancy, since lipids are depleted in ¹³C (McConnaughey & McRoy 1979), and zooplankton had higher lipids than juvenile Chinook salmon.

Stomach contents and stable isotopes gave different estimates for a number of parameters. In 2 out of the 5 regions where we had estimates of TL from stomach contents and stable isotopes (CEBC and SEAK), SIA provided an estimate for TL that was approximately 0.3 to 0.5 TLs below the TL estimate from stomach contents. These results were qualitatively similar when we used traditional TL estimates (Cabana & Rasmussen 1996, Fig. S3 in the Supplement at [www.int-res.com/articles/suppl/m537p247_supp.pdf]). The difference in TL estimates from stomach contents and SIA may be a function of the different periods over which these metrics integrate. Stomach contents are a snapshot of diet that generally represents material consumed over the last 24 h (Polunin & Pinnegar 2002) while stable isotopes reflect a period of weeks to months (Fry 2006). As juvenile Chinook salmon generally experience ontogenetic shifts in their diet in the marine environment
(Brodeur 1991, Daly et al. 2009), the differences that we found between these approaches may be because the juvenile Chinook salmon had recently shifted their diet to fish, but due to the time lag associated with stable isotopes, the SIA still indicated a diet higher in non-fish prey.

Another explanation for the underestimated TL of juvenile Chinook salmon from SIA in some regions could come from the assumption of zooplankton being at a TL of 2. Zooplankton are rarely strictly herbivorous, and there can be significant omnivory within the zooplankton TL (Kling et al. 1992, El-Sabaawi et al. 2013). This means that the TL of the regional baselines may be appreciably higher than 2, and the estimates from stomach contents and stable isotopes may converge.

A final explanation for this disagreement between TL estimates from stomach content and stable isotope approaches concerns the differential digestibility of prey items (Polunin & Pinnegar 2002). Because fish prey are generally larger than other prey items in the stomachs of juvenile Chinook salmon, they may be expected to digest slower than other prey items, and thus end up over-represented in the stomach content data. Supporting this possibility, He & Wurtsbaugh (1993) found that in brown trout Salmo trutta, evacuation rates decreased with increasing prey size. Similarly, Tanasichuk et al. (1991) found that Pacific hake Merluccius productus digested euphausiid prey faster than fish prey. Finally, the WCVI TL estimate from stable isotopes was approximately 0.5 TL above the TL estimate from stomach contents, possibly due to the lack of a clear asymptote in this region, or because the resolution of the zooplankton baseline was insufficient. Altogether, these results again indicate the sensitivity of TL estimates to differences in discrimination values, and highlight the need to consider multiple approaches when calculating TLs at a large scale.

Because of the regional differences in δ13C and δ15N at the base of the food web, it is not surprising that equilibrated juvenile Chinook salmon from each region occupied a unique area in bivariate isotopic niche space. We found that there were no significant differences among isotopic niche breadths from ORWA, WCVI, CEBC and SEBS. Because we limited this comparison to only the large fish in each region, this finding is not particularly surprising. However, NEBS showed the significantly largest niche area, while SEAK had a niche area significantly smaller than SEBS and NEBS. For NEBS, the large niche area was likely due to a combination of the large variability in prey isotopic niche, plus the large variability in stomach contents in this region. In contrast, SEAK had significantly less baseline variability than other regions, combined with less variability in stomach contents, leading to a small isotopic niche.

One limitation of our study is that we were unable to control for stock-of-origin, as different stocks of Chinook salmon may have different diets (Schabetsberger et al. 2003). In most regions, the juvenile salmon we sampled were from a variety of stocks, with various stock-specific life-histories and migration patterns (e.g. Tucker et al. 2011, 2012). For instance, WCVI juvenile Chinook salmon were probably primarily from WCVI stocks, but Columbia River and other Washington stocks can represent over 10% of catches in some years (Trudel et al. 2009, Tucker et al. 2011, 2012). Similarly, juvenile Chinook salmon caught in SEBS in 2007 were likely a mixture of Yukon River Chinook and southern Bering Sea stocks (Murphy et al. 2009). Considering the previous studies on migration patterns, however, each region was probably primarily represented by fish that originated from that region. Future studies should look into patterns of stock-specific resource utilization across large spatial scales.

Overall, our results indicate that regardless of the approach we used, there was substantial variation in the feeding ecology of juvenile Chinook salmon from California to the eastern Bering Sea. Considering that the factors underlying the recent widespread declines of Chinook salmon survival from Alaska to California are not well understood (Schindler et al. 2013, Riddell et al. 2013), determining how differences in feeding ecology could affect the survival rates of Chinook salmon in each region is important to explore further.

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