Genome duplication and multiple evolutionary origins of complex migratory behavior in Salmonidae

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1. Introduction

Multiple rounds of whole genome duplication have characterized the evolution of vertebrates (Jaillon et al., 2004). Specifically, in actinopterygians (ray-finned fishes), genome duplication events correlate with bursts of morphological change in extant lineages (Ohno, 1970; Taylor et al., 2001, 2003; Van de Peer et al., 2003, 2009; Braasch et al., 2009a,b) but see (Heimberg et al., 2008). However, incorporating fossil taxa into these analyses demonstrates that morphological complexity accumulated more continuously through time (Donoghue and Purnell, 2005). Whether episodic or periodic, little is known about the behavioral consequences of genome duplication in vertebrates, and if duplication events coincide with transitions in behavior and physiology (Novak et al., 2006).

The ecophysiologicl consequences of genome duplication remain understudied (Mable, 2004; Soltis et al., 2010; Mable et al., 2011), but evidence from plants suggests that polyploids are able to inhabit a greater diversity of environments compared to their diploid progenitors (Otto and Whitton, 2000). Generally speaking, duplication events trigger morphological and physiological changes in size and ecological tolerance (Adams and Wendel, 2005; Hu et al., 2012). For example, recent work suggests that newly formed polyploid populations are more resistant to parasites (Oswald and Nuismer, 2007). Furthermore, polyploidy affects cellular physiology by altering cell size and enhancing photosynthetic capacity (Warner and Edwards, 1993); it may even change cellular metabolic rates (Comai, 2005). Less is known about the ecophysiologicl consequences of genome duplication in vertebrates. In one notable example, mating calls in gray tree frogs follow directly from changes in ploidy, which leads to reproductive isolation via female mate choice (Mable et al., 2011). The link between genome duplication and the evolution of complex traits has also been hypothesized for salmonid fishes (Ohno, 1970).

Salmonids (salmon, trout, char, whitefishes, and graylings) are one of the most well studied vertebrate groups that have undergone genomic duplication. Their migratory behaviors require major physiological plasticity, which are known popularly in Atlantic and Pacific salmon. These fishes are capable of overcoming tremendous obstacles to return from distant oceans to natal streams for spawning. This behavior has inspired centuries of research into the evolution of salmon migrations and the origin(s)
of this life history strategy (McDowall, 2002). Molecular and morphological evidence places the freshwater Esocidae (pikes) as the closest relatives of salmonids, which suggests that salmonids arose in freshwater environments (Stearley, 1992; Ishiguro et al., 2003; Ramsden et al., 2003; Nelson, 2006; Broughton, 2010; Wilson and Williams, 2010). Furthermore, the Eocene salmon id Eosalmo driftwoodensis is preserved in all life stages within freshwater deposits, offering strong support for non-migratory behavior in stem-salmonids as well. In addition to complex migratory behavior, all salmonids (estimated to have originated 63.2–58.1 MYA. (Crete-Lafreniere et al., 2012)) share a tetraploid genome that arose between 100 and 25 My. (Johnson et al., 1987; Sakamoto et al., 2000; Angers et al., 2002).

Beginning in the 1970s (Ohno, 1970), authors speculated that the salmonid duplication event may have provided the genetic material necessary for the evolution of migratory behavior (Allendorf and Thorgaard, 1984; Phillips and Rab, 2001; Koop et al., 2008; Norman et al., 2011, 2012). The hypothesis suggests that duplicate genes (resulting from whole genome duplication) could provide flexibility for differential optimization, as salmonids transitioned to a multi-modal life history. Recent evidence suggests that copies of the FXYD isotypes (Tipsmark, 2008; Tipsmark et al., 2008a,b,c), ATP1a1b loci (Norman et al., 2011) and other genes involved in osmoregulation (Gharbi et al., 2004, 2005; Woram et al., 2004; Dannmann et al., 2005; Norman et al., 2012) may have gained new functions following duplication events, and permitted an increased tolerance to salinity. A comprehensive analysis of salmon and pike transcriptomes has revealed an asymmetrical relaxation of selection on paralogous salmonid genes, allowing individual paralogs to evolve at faster rates (Leong et al., 2010). Many paralogs were lost, but remaining paralogs began to diverge (Leong et al., 2010). This divergence may have been a crucial factor leading to complex migratory behavior and the resulting access to marine niches with greater resources and more space. However, the absolute timing of genomic duplication in relation to the evolution of anadromy remains unknown.

Despite investigations into the consequences of genome duplication and the origin(s) of migratory behavior in salmonids, at least two important questions remain unanswered: (1) did complex migratory behavior evolve multiple times in salmonids? (2) Does the evolution of salmonid migratory behavior coincide with genome duplication? These are both phylogenetic questions. Thus, migratory behavior and ploidy can be treated as traits and studied using a time calibrated phylogenetic tree. However, one problem with this approach is that anadromy is highly variable within the Salmonidae, even at the intraspecific level. Traits with rapid evolutionary rates are difficult to reconstruct because their histories can be quickly lost (Schulter et al., 1997; Cunningham et al., 1998; Oakley and Cunningham, 2000; Oakley et al., 2005). Furthermore, there is only one temporal estimate for genomic duplication in salmonids, but confidence intervals span 75 MYA (Allendorf and Thorgaard, 1984; Crete-Lafreniere et al., 2012; Shedko et al., 2012). Despite improvement in this latter approximation, no study thus far has estimated explicitly the absolute timing of the transition from diploidy to tetraploidy.

To improve upon previous analyses and test long-standing questions about salmonid evolution, we utilized new statistical tools and copious data to evaluate the relationships among phylogeny, migratory behavior, and genome duplication. We reconstructed a phylogenetic tree of salmonids and their close relatives, and analyzed the character matrix using a mixed-model multi-partition dataset (DNA and morphology) under Bayesian and Maximum Likelihood statistical frameworks. Furthermore, we constructed a series of fossil-calibrated trees by concatenating extinct and extant taxa, thereby significantly improving upon previous divergence time estimates. We included a key freshwater stem-salmonid (E. driftwoodensis), which offered unique insight into the transition to migratory behavior. We then mapped the history of migratory behavior and genome duplication onto the resulting phylogeny using Bayesian methods (Pagel et al., 2004) and stochastic character mapping (Bollback, 2006). We estimated the absolute timing of character transitions with a new method that uses ancestral character simulation data to plot the frequency of character transitions per branch on a time-calibrated phylogeny. This method improves on previous approaches because it estimates character transitions along branches instead of only at nodes. These methods allowed consideration of uncertainty in rates of trait evolution, a factor that importantly impacts ancestral state estimations (Schultz et al., 1996). Our analyses improve upon most of the limitations of previous studies and strongly support the parallel evolution of anadromy in salmonid fishes. Results indicate that complex migratory behavior evolved at least twice in salmonids (in salmon and whitefishes), but it is unclear how many times large-scale anadromy evolved, likely due to the rapid evolution of migratory behavior in crown-Salmoninae. Furthermore, we provide new estimates of ancestral tetraploidization that suggest genome duplication significantly preceded the transition to complex migratory behavior in whitefishes.

2. Materials and methods

2.1. Data collection

We constructed a data matrix from 97 species or subspecies-level salmonids and related outgroups. Our data set included 41 data partitions. We collected nucleotide sequence data from GenBank using 16 nuclear and 18 mitochondrial genes (Phillips and Oakley, 1997; Crespi and Fulton, 2004). We also assembled seven categorical data sets, including morphological data that were compiled from three sources, excluding overlapping data (Smith, 1992; Stearley and Smith, 1993; Wilson and Li, 1999). Besides morphology, other categorical data sets included rDNA RFLPs (Phillips et al., 1992, 1994, 1995), mitochondrial RFLPs (Nielsen et al., 1998; Carrera et al., 1999), chromosome number (Phillips and Rab, 2001), Short Interspersed Nuclear Elements (SINES) (Takasaki et al., 1997; Hamada et al., 1998), microsatellites (Olsen et al., 2000), and allozymes (Allendorf and Seeb, 2000).

To fossil-calibrate the root of the tree, we extended sampling beyond Salmonidae to include Esociformes (pikes such as Novumaba hubsi and Esox lucius), Osmeriformes (smelts such as Hypomesus olidus, Plecoglossus altivelis and Argentina sialis), Conorychiformes (milkfishes such as Chanos chanos), and Cypriniformes (zebrafishes such as Danio rerio).

2.2. Phylogenetic analyses

We excluded data sparse taxa from further analyses. In fact, not all species-level taxa have data for all or even most data partitions: of the 97 × 41 = 3977 possible combinations of species taxa × data partition, the complete initial data matrix was 80% missing. With the exception of fossil taxa – for which only morphological data are available – we excluded from further analyses species and subspecies-level taxa containing fewer than three data partitions. Many excluded taxa were subspecies for which only chromosome numbers were available. After excluding extremely sparse taxa, the resulting matrix contained 58 taxa and was 26.6% complete as data partitions available per taxon (Supplementary Table 1). Nucleotide data collected from GenBank were aligned using the default parameters of MUSCLE (Edgar, 2004), with the exception of growth hormones and intron c; these were aligned manually (Oakley and Phillips, 1999). Several mitochondrial genomes are available for
salmonid taxa; these were separated into individual genes for alignment before concatenating with the entire dataset.

After obtaining an aligned matrix, we conducted numerous phylogenetic analyses, using Bayesian Inference (BI) and Maximum Likelihood (ML) approaches. We divided genetic data into partitions by gene and categorical data as binary or multistate. Using MrBayes 3.1 (Huelsenbeck and Ronquist, 2001), we assumed GTR + G + I for DNA partitions, which was the best-fit of 56 model comparisons implemented in jModeltest (Posada, 2008). For all categorical data, we assumed the Mk model of Lewis (Lewis, 2001). Only branch length parameters were linked across data partitions and we assumed default priors for all parameters. To estimate the posterior probability distribution of phylogenetic trees and rate parameters, we used Metropolis-coupled MCMC with proposal mechanisms described by Nylander et al. (Nylander et al., 2004). We conducted a run of 10,000,000 generations, sampling every 100th generation. We excluded the first 100,000 generations as “burn-in” using Tracer v1.5 (Rambaut and Drummond, 2004), after which the Markov chain was stationary. Using RAxML (Stamatakis, 2006), we did a ML search with a mixed partition model. Random starting trees were used and topological robustness was evaluated using 100 non-parametric bootstrap replicates.

We used r8s v 1.7 (Sanderson, 2003) to construct an ultrametric tree with the best scoring tree from RAxML. To this end, we used a semi-parametric Penalized Likelihood approach with a truncated Newton algorithm and bound constraints. We then used BEAST v 1.7.2 (Drummond et al., 2012) to conduct a fully parametric Bayesian analysis on the same data. With BEAST we used several different tree priors to compare the outcomes, namely the traditional Yule model (Gernhard, 2008), the standard Birth–Death model (Gernhard, 2008), and the Birth–Death model with incomplete sampling (Stadler, 2009). Furthermore, all categorical data were included in the BEAST analyses by coding it into BEAST XML input (following Pyron, 2011) using a custom R function. Extinct species taxa were included as dated, non-contemporaneous OTUs (See Table 1 for tip dates and calibrations) using the tip-dates function in BEAUTI v 1.7.2 (Drummond et al., 2012), with the morphological data encoded BEAST’s XML format as in Pyron (2011) using a custom R script. The R script converts standard NEXUS-formatted morphological data (including ambiguities) to BEAST’s format which requires a single alphanumeric code for each character state and each combination of ambiguous states (e.g., “?” is encoded as “Z”); the ambiguity codes are user-determined and also generated by the script. The script is available at http://oakley-web.eemb.uchicago.edu/ASTOMO13/. Uncertainty in the dating of fossil OTUs was accounted for with a uniform distribution covering the published age estimate of the fossil, as done in Wood et al., 2012. All BEAST runs were fully partitioned, with unlinked substitution-rates, linked trees, and clock models. Uncorrelated lognormal relaxed clocks and random starting trees were used for all analyses. Each analysis was run for 30,000,000 generations, sampled every 1000th, and checked for stationarity using Tracer v 1.5 (Rambaut and Drummond, 2004). Consensus trees with mean node values were constructed using TreeAnnotator v 1.7 (Drummond et al., 2012) after having determined the appropriate burn-in.

2.3. Trait mapping

We assigned five states to the character of migratory behavior for mapping onto our phylogenetic trees: state (0) freshwater – taxon members spend their entire lives in freshwater; state (1) anadromous – taxon members migrate between freshwater and marine stages, but migrations are not directly coupled to spawning; state (2) small-scale anadromy – taxon members are born in freshwater before migrating to estuaries or other coastal environments (upriver migrations are made shortly before spawning); state (3) large-scale anadromy – taxon members are born in freshwater before migrating far out to sea, often following ocean currents for well over a year, rapid upriver migrations are then made shortly before spawning; and state (4) marine – taxon members are born and live entirely in the sea. These character states reflect our attempt to allow for specific definitions, as well as to arrive at a consensus of previous authors’ opinions (see Supplementary Table 1). We are the first to use the terms small and large-scale anadromy: many previous authors do not discriminate between these categories (McDowall, 2002). By distinguishing them, we are stating our agreement with Stearley (Stearley, 1992) that the long migrations to sea by Atlantic and Pacific salmon, which often follow ocean currents, deserve their own category. However, we disagree with Stearley (Stearley, 1992) that other salmonids are not anadromous: he used the term anadromous for taxa that conduct migrations a short distance to sea. Therefore, our categories of small and large-scale anadromy are consistent with most authors given that they consider many salmonids to be anadromous. Yet we also recognize that large-scale migrations of some salmon represent a different life history strategy. For character state simulations of ancestral tetraploidy (see below), we coded all diploids as 0 (outgroups) and all tetraploid as 1 (salmonids) (Allendorf and Thorgaard, 1984).

2.4. Ancestral state estimation

We used Bayesian statistics for ancestral state reconstructions, implemented in the software Bayes-Multistate (Pagel et al., 2004). A Bayesian approach aims to estimate the distribution of a particular parameter, such as the probability of a character state at a certain node. We chose this approach because it considers uncertainty in both phylogenetic tree topology and in rates of migratory evolution to reconstruct the most likely condition at the base of crown-salmonids.

### Table 1: Fossil dates and calibrations.

<table>
<thead>
<tr>
<th>Fossil</th>
<th>Formation</th>
<th>Reference</th>
<th>Age (MYA)</th>
<th>Calibration prior</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multiple taxa</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+Oldenomeres sp.</td>
<td>Obere solnhofener schichten Germany</td>
<td>Arratia (1997)</td>
<td>165.2–149.85</td>
<td>Ln offset = 149.85 (mean 0, std 0.5)</td>
</tr>
<tr>
<td>+Eosalmo driftwoodensis</td>
<td>Driftwood creek formation, alleny formation, Klondike mountain formation Canada</td>
<td>Wilson and Li (1999)</td>
<td>52–51.5</td>
<td>Tip date mean 51.75 uniform distribution 51.5–52</td>
</tr>
<tr>
<td>+Salvelinus larsoni</td>
<td>Lower chalk hills formation, idaho USA</td>
<td>Smith (1981) and Smith (1992)</td>
<td>9.8–7.4</td>
<td>Tip date mean 8.6 uniform distribution 9.8–7.4</td>
</tr>
<tr>
<td>+ Oncorhynchus lacustris</td>
<td>Glenns ferry formation, idaho USA</td>
<td>Smith (1981)</td>
<td>5–2.5</td>
<td>Tip date mean 3.75 uniform distribution 5–2.5</td>
</tr>
<tr>
<td>+Oncorhynchus keta</td>
<td>Chalk hills formation, malheur co., oregon USA</td>
<td>Smith (1992) and Eiting and Smith (2007)</td>
<td>6.6–4.8</td>
<td>Ln offset = 6.6 (mean 1, std 1)</td>
</tr>
</tbody>
</table>

Fossil data used to calibrate the r8s and BEAST trees showing taxa, formations, references, ages, and parameters used for calibration.
2.5. Absolute timing of character transitions

To estimate the absolute timing of character state changes, we performed stochastic character mapping (SCM) (Nielsen, 2002; Huelsenbeck et al., 2003) on fossil-calibrated time trees. We abbreviate this approach as time tree stochastic character mapping (ttSCM). We performed ttSCM for two characters: (1) ploidy (states = tetraploid and diploid); and (2) migratory behavior (states described above under “trait mapping”). We used 150 time calibrated branches pulled from posterior distributions of MCMC chains that assumed a Yule model tree prior (details described above). For ploidy levels (ancestral tetraploidization) we used all available taxa. However, for migratory behavior we analyzed salmonids separately from their outgroups, since: (1) outgroup migratory traits are irrelevant to the question of whether anadromy evolved independently within salmonids; (2) outgroups are undersampled; and (3) outgroups are represented by additional character states that do not occur in salmonids. On each tree, we conducted 100 SCM simulations for each character. We chose the best-fit model for each character using standard Likelihood Ratio Tests to compare nested models (e.g. symmetric vs. asymmetric rate matrices) (Moors and Schluter, 1999). We then performed SCM using make.simmap of the phytools v0.2-90 R package (Revell, 2012). Make.simmap, like simmap itself (Bollback, 2006), generates calibrated character histories with the constraint that tip character states match observed tip states. Make.simmap first uses ML to determine parameter values for the model of character evolution. Next, make.simmap generates samples from a posterior distribution conditional on the ML-fitted model to reconstruct character histories on trees. This ML parameter fitting approach is different from simmap (Bollback, 2006), which samples from a joint posterior of character histories plus substitution rates. The make.simmap approach is similar in spirit to the empirical Bayes approach for ASR introduced by Pagel (Pagel et al., 2004), in which character rate models are set to ML estimates. We wrote custom R-scripts to plot histograms of the number of state changes in different time bins. In addition to estimating the timing of migratory transitions across the whole tree, we used ttSCM to demonstrate its capability to estimate character transitions on branches of interest. For this analysis, we established support for the homology of whitefish anadromy and the non-homology of whitefish and salmon anadromy. We then coded for all taxa a binary character for whitefish anadromy, with an anadromous whitefish state (1), and a state (0) for all other taxa. By doing so, we were able to estimate the absolute difference in timing between tetraploidy and a transition to anadromy (observed within the Coregoninae).

3. Results

3.1. Data collection

Taxa with the most available data included Oncorhynchus mykiss (data available for every partition) and Salmo salar (data available for 88% of partitions). The most complete data partitions included morphological data (available for 95% of taxa), chromosomal data (two characters, available for 69% of taxa), and ITS-1 (59% of taxa). The final nucleotide alignment consisted of 32,029 bps, which was concatenated with 275 categorical characters over 58 taxa.

3.2. Phylogenetic results

Bayesian and Maximum Likelihood analyses produced phylogenetic trees with strong support for most nodes (Fig. 1). The results of our supermatrix approach have implications for resolving the tree of life, because we obtained strong phylogenetic signal from a data matrix that was only 26.6% complete in terms of data partitions available. This is consistent with other studies (Sanderson et al., 2011). The placement of three freshwater salmoniform (Fig. 1) clades is particularly relevant to the multiple origins of anadromy. The nearest salmonid outgroup is a largely freshwater clade containing Esox and Novumbra. This topology is supported by other studies based on mitochondrial genome data (Ishiguro et al., 2003), and other molecular (Lopez et al., 2004) and morphological characters (Williams, 1987; Wilson and Williams, 2010). However, the position of the second freshwater clade is more surprising. The freshwater Thymallinae (graylings) emerged as sister to all other salmonids, with a posterior probability of 1.0. The Coregoninae are usually considered to be sister to other salmonids, including thymallins (Stearley and Smith, 1993). The third freshwater lineage that impacts the freshwater ancestry hypothesis is Driftwoodensis. This taxon is abundant in the Eocene Green Lake formation, which preserves all its life history stages. The stem-position of *Eosalmo* in our tree as sister to crown-group Salmoninae is consistent with previous phylogenetic analyses (Wilson and Li, 1999). Our results, therefore, indicate multiple transitions in salmonins and coregonins from a freshwater ancestry to complex migratory behavior.

By combining molecular and morphological characters, we present the first fossil-calibrated phylogenetic framework for salmonid diversification. Including fossils in the time-calibrated tree permitted us to incorporate geological ages into the analysis, instead of simply using prior probability distributions on the dates of nodes. The tip-dating method allows more accurate placement of temporal constraints, thereby yielding more reliable estimates of divergence times (Pyron, 2011). The disadvantages of using fossil taxa to constrain nodes instead of tip dates are apparent in the case of *Eosalmo*. In contrast to previous analyses that used *Eosalmo* to constrain the root of the Salmonidae (Shedlock et al., 2012), we placed *Eosalmo* within our phylogeny and applied the constraint directly to the *Eosalmo* branch. Another recent approach used *Eosalmo* to constrain Coregoninae + Salmoninae (Crette-Lafreniere et al., 2012), which led to a more recent divergence of crown-Salmonidae. Furthermore, our approach differs from past studies in that we included dates from six fossils, three of which are OTUs that were incorporated directly into the analysis.

Trees recovered using BEAST were nearly identical to those recovered using RAXML and MrBayes (see Figs. 1 and 2), though support values for the BEAST trees were slightly higher for all nodes. One topological difference included the position of the fossil *Salvelinus larsoni*, which was recovered as sister to the extant *Salvelinus* in BEAST, but which emerged as sister to *Parahucho perryi* in the RAXML and MrBayes analyses. This indicates that the inclusion of dating information in a total-evidence analysis might occasionally change the inference of tree topology, as suggested by Ronquist et al. (2012). This should not be surprising, since more information is being considered in an analysis with dated fossil OTUs relative to an undated analysis. The ages of fossils might constrain their possible positions in a dated tree. The phylogenetic position of *S. larsoni* remains uncertain because of low support from all utilized methods. Divergence time estimates were similar between the r8s analysis and all BEAST trees, with the latter yielding slightly older dates (Fig. 2). Overall, the three BEAST trees (Yule, birth–death, and birth–death with incomplete sampling) produced nearly identical topologies and dates, with no obvious differences. The root of Salmonidae (using mean values from all separate BEAST analyses; Yule tips, BD tips, BD Inc. tips) was estimated at 69.6 Ma with highest posterior density (HPD) values ranging between 78.3 and 60.8 MYA across all separate speciation models. This represents a vast improvement in confidence compared to previous estimates.
3.3. Ancestral state estimation

With a one-rate model, patterns of trait evolution were similar whether using a flat prior distribution or a prior based on the likelihood of the single rate parameter. In both cases, the ancestral states of Salmonidae and Salmoninae were reconstructed as “freshwater” with posterior probabilities greater than 0.95. Neither analysis could unambiguously reconstruct the migratory behavior of the most recent common ancestor of Atlantic and Pacific salmon (the node of common ancestry between Salmo and Oncorhynchus). See Fig. 1 for ancestral state estimates of major nodes.

3.4. Absolute timing of character transitions

The absolute timing of salmonid tetraploidization was estimated using time tree stochastic character mapping (ttSCM). We used an equal rates (ER) model with coequal rates between diploid and tetraploid states.

Assuming an equal rates model, we plotted a histogram of transitions per branch across the tree in 10 million year time bins (Fig. 3A and B). The time bin with the highest number of transitions per branch is 80–70 MYA, with a distribution of bins ranging from 80–50 MYA. This estimate is consistent with a previous estimate of 100–25 MYA (Allendorf and Thorgaard, 1984), and with the results of a recent salmonid phylogeny (Crette-Lafreniere et al., 2012). As expected, it is also slightly older than our estimates for the root of crown-group Salmonidae (78.3–60.8 MYA), which previous approaches (those that constrained character changes to nodes) would perceive as the original timing of tetraploidization. However, because ttSCM permits character changes along branches, it provides older estimates relative to methods that constrain only nodes.

We also estimated the timing of migratory transitions over salmonid evolution. We plotted transitions per branch for three character states: freshwater, small scale anadromy, and large-scale anadromy (Fig. 4). Using a Likelihood-Ratio Test we found that a symmetric model fits migratory patterns better than ER or ARD models (p-value rejecting ARD = 0. 0.2460857, p-value rejecting AR = 1). The highest rate of transitions per branch to freshwater habitats occurred during the early history of salmonids (75–70 MYA); transitions to small-scale anadromy per lineage peaked 30–20 MYA; and transitions to large-scale anadromy peaked 10–5 MYA. Although it is possible to identify maximum time bins, the nature of the data and the Markov models leads to large numbers of character transitions through time, and therefore the time bins across the tree are very long. Early transitions to freshwater and late transitions to large-scale anadromy are consistent with an ancestral freshwater environment, and the independent origins of migratory behavior. Moreover, the intermediate timing of maximal transitions to small-scale anadromy is consistent with the possibility of common, ordered character transitions from freshwater to small-scale anadromy to large-scale anadromy.

We also estimated the timing of the transition from freshwater to small-scale anadromy on the branch leading to Coregoninae (Fig. 3C and D). The maximum time bin was 25–20 MYA, with a distribution spanning the entire length of the branch. Coupled with estimates of the transition to tetraploidy, this result further supports the hypothesis that genome duplication significantly preceded the evolution of complex migratory behavior by 55–50 MYA.

All custom scripts used in this study have been made available for download at http://oakley-web.eemb.ucsb.edu/ASMO13/.
A couple of recent studies used fossil constraints to calibrate the salmonid molecular clock (Allendorf and Thorgaard, 1984; Crete-Lafreniere et al., 2012; Shedko et al., 2012). The topology of the trees recovered in these studies is similar to our own, however, the divergence times differ significantly. Shedko et al. (2012) used two calibration points, one for Oncorhynchus (11.6–5.3 MYA) and one for Salmonidae (55.8–33.9 MYA), which led to the more recent divergence estimates. Our results also differ from Crete-Lafreniere et al. (2012). Despite the use of the same fossil taxa and a very similar dataset, their divergence estimates were older than the results presented here. The primary source of these differences seems to be the placement of Eosalmo, and the choice of which node to constrain. We overcome this problem by including morphological data from Eosalmo (and other fossils) in order to date them as tips in our tree. We also extended our sampling significantly outside of salmonids to include additional calibration points for salmoniform and euteleostean fishes.

Including fossils in a phylogeny is important because they represent distinct tips in a tree with individual branch lengths (Pyron, 2011). However, most studies use fossil information to constrain nodes among extant taxa. We used a combination of these
calibration strategies (as in Wood et al., 2012), including prior distributions on the dates of particular nodes representing more complete fossils, and dated fossil taxa as terminal tips. The phylogenetic and ecological histories of \( C_{160} \)Eosalmo (as a stem-Salmoninae that inhabited lacustrine habitats) contribute strongly to the freshwater signal in crown-Salmoninae. Other fossil taxa included in the phylogeny (\( C_{160} \)S. larsoni and \( C_{160} \)Oncorynchus lacustris) are ambiguous in terms of migratory behavior. Nevertheless, using fossil taxa as tips in a phylogeny offers various advantages. Namely, the molecular clock can be calibrated automatically while co-estimating the phylogeny using extinct taxa of known age (Pyron, 2011; Heikkila et al., 2012). This approach sets our calibrated phylogeny apart from the surprisingly few other attempts that estimate divergence times in salmonids (Allendorf and Thorgaard, 1984; Crete-Lafreniere et al., 2012; Shedko et al., 2012).

4.2. Multiple origins of anadromy

The evolution of salmonid migratory behavior has interested biologists for centuries (Pennant, 1776; Fleming, 1828; Günther, 1866; Buckland, 1873; Day, 1887; Meek, 1916; Tchernavin, 1939; Hoar, 1976; Thorpe, 1982; Stearley, 1992; McDowall, 2002). Migration may present advantages such as increased access to resources and tolerance to environmental changes. Considering the benefits, it is not surprising that migratory behavior could have evolved multiple times, resulting in different variations on the same theme. We classify these variations as large-scale anadromy and small-scale anadromy, and consider them to be distinct behaviors based on the travel distance from spawning grounds to the sea. This distinction between different forms of anadromy leads to a question of whether transitions to anadromy occurred through gradual steps (i.e. from freshwater to small-scale and then large-scale anadromy), or whether the large-scale anadromy evolved directly from a freshwater ancestor. Furthermore, this remains unclear partly due to the homoplasious nature of the character, and to limitations of existing methods used to trace the evolutionary histories of rapidly evolving characters.

4.3. Genome duplication and anadromy

Cytogenetic evidence suggests that tetraploidy likely arose at the root of Salmonidae (Allendorf and Thorgaard, 1984; Phillips and Rab, 2001; Ramsden et al., 2003). Our results concur, and further show that genome duplication significantly preceded at least one origin of complex migratory behavior within salmonids. However, at least two competing hypotheses may explain the time discordance between genome duplication and migration. The first hypothesis suggests that the genome duplication event may have

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**Fig. 3.** Migratory character state transitions. Histograms of character simulation results showing transitions to different character states per branch. The axes have been standardized across graphs to show the number of transitions to each state on the y-axis and the time after the root on the x-axis.
provided the genetic material necessary for the evolution of migratory behavior (Ohno, 1970; Allendorf and Thorgaard, 1984; Phillips and Rab, 2001; Koop et al., 2008; Norman et al., 2011, 2012). Duplicated genes can gain new functions over time and affect complex morphological, behavioral, and physiological traits (Lynch and Conery, 2000; Semon and Wolfe, 2007; Mable et al., 2011). In the case of salmonids, a whole genome duplication event generated a full set of new genes, providing novel gene copies with potential adaptive qualities. Some of these new gene copies could have gained new functions, optimized via selection to allow species to physiologically adapt to different environments. This optimization could have resulted in increased salinity tolerance and plasticity in osmoregulatory functions, allowing salmonids to extend out of their ancestral freshwater habitats into more productive habitats such as estuaries and oceans. Furthermore, the observed lag between genome duplication and transitions to anadromy (55–50 MYA) may have been necessary to allow time for paralogs to diverge significantly and gain new function or sub-functionalize. However, genomic evidence suggests that most duplicated genes are lost or silenced within a few million years after duplication (Lynch and Conery, 2000). Additionally, many genes that are duplicated during whole-genome duplication events become deleterious and cause epigenetic instability via non-additive gene regulation (Comai, 2005).

The second hypothesis proposes that there is no causal relationship between genome duplication and the evolution of migratory behavior. Some evidence suggests that duplicate copies of salmonid genes involved in osmoregulation may have gained new functions and permitted an increased tolerance to salinity (Charbi et al., 2004, 2005; Woram et al., 2004; Danzmann et al., 2005; Tipsmark, 2008; Tipsmark et al., 2008a,b,c; Norman et al., 2011, 2012). Nonetheless, these duplicated genes may have arisen post genome duplication as independent events that did not result directly from tetraploidy. Additional comparative genomic data will be required to estimate the timing of adaptive gene duplications. Our results provide an estimate of the timing of tetraploidy in salmonids that can be used for such comparisons.

4.4. Time tree stochastic character mapping (ttSCM)

The ability to calculate the absolute timing of character transitions was crucial in testing whether genome duplication coincided or preceded transitions to migratory behavior. Although multiple previous authors have estimated character histories on fossil-calibrated time trees (e.g. Wiens et al., 2006; Syme and Oakley, 2012), traditional ancestral state estimation methods (e.g., maximum likelihood) usually provide probabilities of different character states at the nodes of trees, although this is mostly a matter of computational convenience and graphical display, and not intrinsic to the statistical method. In contrast, SCM simulates character histories, providing a ready-made probabilistic display of the character state histories along branches of the tree. We are unaware of previous authors using SCM on a time tree for absolute estimates of character state timing. Estimates of the absolute timing of state changes can allow comparisons of character evolution to geological events, and to the timing of other characters.

We report not the total number of state changes per time bin, but the state changes per branch during each time bin. This is because SCM simulates numerous changes on each branch that stem from the rapid rates of character evolution estimated by the Markov model. The high numbers of inferred changes often seem counter-intuitive, and may indicate that homogeneous models poorly describe the evolutionary process of many characters. Instead, models that permit various rates across a tree may better represent
character evolution (Skinner, 2010). Thus, in our analyses, when rates of evolution were high, we noted that raw histograms mainly reflected the number of lineages through time: when state changes occur on all branches, more branches yield more total changes. In this way, measuring state changes per branch reveals a more refined understanding of character history.

5. Conclusions

We present the first phylogenetic analysis to support multiple evolutionary transitions to migratory behavior in salmonids. We also support the idea that raw histograms mainly reflect the number of lineages through time, when state changes occur on all branches, more branches yield more total changes. In this way, measuring state changes per branch reveals a more refined understanding of character history.

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Appendix A. Supplementary material

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References


