Molecular Evolution of the Neural Crest Regulatory Network in Ray-Finned Fish

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Abstract

Gene regulatory networks (GRN) are central to developmental processes. They are composed of transcription factors and signaling molecules orchestrating gene expression modules that tightly regulate the development of organisms. The neural crest (NC) is a multipotent cell population that is considered a key innovation of vertebrates. Its derivatives contribute to shaping the astounding morphological diversity of jaws, teeth, head skeleton, or pigmentation. Here, we study the molecular evolution of the NC GRN by analyzing patterns of molecular divergence for a total of 36 genes in 16 species of bony fishes. Analyses of nonsynonymous to synonymous substitution rate ratios (dN/dS) support patterns of variable selective pressures among genes deployed at different stages of NC development, consistent with the developmental hourglass model. Model-based clustering techniques of sequence features support the notion of extreme conservation of NC-genes across the entire network. Our data show that most genes are under strong purifying selection that is maintained throughout ray-finned fish evolution. Late NC development genes reveal a pattern of increased constraints in more recent lineages. Additionally, seven of the NC-genes showed signs of relaxation of purifying selection in the famously species-rich lineage of cichlid fishes. This suggests that NC genes might have played a role in the adaptive radiation of cichlids by granting flexibility in the development of NC-derived traits—suggesting an important role for NC network architecture during the diversification in vertebrates.

Key words: Actinopterygii, teleost, development, ontogeny, gene regulatory network, cichlid, NC-genes, hourglass model.

Introduction

The vertebrate-specific neural crest (NC) is a transient population of migrating multipotent cells that gives rise to various, highly diverse, tissues and cell types (fig. 1A). The NC is considered a key innovation in vertebrate evolution (Gans and Northcutt 1983; Shimeld and Holland 2000). Due to its broad developmental potential, the NC has been even referred to as the fourth germ layer (Hall 2000). Its derivatives provide the developmental basis for the vast morphological and physiological diversification that has occurred during vertebrate evolution (Betancur et al. 2010; Bronner and LeDouarin 2012; Simões-Costa and Bronner 2015). NC cells contribute to evolutionary novelties, such as the cranium, the branchial skeleton, and peripheral nervous system (Shimeld and Holland 2000; Santagati and Rijli 2003). Many of the distinctive phenotypic differences found in bony fishes are largely influenced by the NC cell lineage (Kimmel et al. 2001; Donoghue et al. 2008; Braasch et al. 2009). Most prominently, NC-derived mesenchyme shapes the astonishing diversity of head and jaw morphologies. Also, pigment cells are NC-derived, and populate the whole body axis to generate the extraordinary diversity of coloration and pigmentation patterns found in the most species-rich vertebrate lineage, the ray-finned fish (Actinopterygii) (Sugie et al. 2004; Salzburger et al. 2007; Braasch et al. 2009; Kelsh and Barsh 2011; Renz et al. 2011; Roberts et al. 2011; Powder et al. 2014; Kratochwil et al. 2015). However, up to now, surprisingly little is known about the patterns of molecular evolution of the NC of actinopterygians.

To get a better grasp on the molecular evolution of the NC regulatory network, it is important to know during what ontogenetic phase and how the specific genes of the GRN act and interact. NC genes have been intensively studied in model systems, such as the zebrafish (Betancur et al. 2010; Bronner
and LeDouarin 2012; Simões-Costa and Bronner 2015). The aspects of development that are NC-specific are initiated during gastrulation. Cells of the NC first start to appear at the interface of neural plate and surface ectoderm, the so-called neural plate border (NPB) (Knecht and Bronner-Fraser 2002; Basch et al. 2006). When the neural tube invaginates and closes during neurulation, the NC is located at its dorsal side (Mayanil 2013). Subsequently, NC cells begin to migrate along predefined paths to colonize various parts of the body (Kulesa and Gammill 2010). Here, the final differentiation is taking place (fig. 1A) (Simões-Costa and Bronner 2015). Induction, specification, and differentiation of NC cells are controlled by a gene regulatory network (GRN) (Sauka-Spengler and Bronner-Fraser 2008). During the last decades, both the genetic components and the regulatory interactions of NC GRN genes have been intensively studied in various vertebrate model organisms (Betancur et al. 2010; Minoux et al. 2013; Simões-Costa and Bronner 2015).

Alterations in GRNs can result in phenotypic and physiological changes (Crombach and Hogeweg 2008; Van Otterloo et al. 2013). To understand the underlying evolutionary dynamics of GRNs, biochemical pathways (Rausher et al. 1999; Lu and Raisher 2003), signal transduction pathways (Riley et al. 2003; Alvarez-Ponce et al. 2009), and developmental pathways (Davila-Velderrain et al. 2014) have been previously analyzed by examining patterns of sequence variation within an evolutionary framework. However, a consensus on general patterns of developmental GRN evolution is still lacking. The topology of GRNs (i.e., the arrangement and connectivity of the GRN elements) has been suggested to influence the rates of molecular evolution within the different members of the network (Cork and Purugganan 2004). Additionally, developmental constraints shape the degree of sequence conservation in developmental GRN. The sequence conservation has been suggested to be higher either during early development (early conservation model or funnel-like model) or at the phylotypic stage, a time point during midembryogenesis when the embryo most closely resembles that of other species (hourglass model)—both models have been controversially discussed in the recent years (Prud’homme and Gompel 2010; Kalinka and Tomancak 2012; Piasecka et al. 2013).

The range of taxa analyzed in this study covers species that diverged between approximately 4 Ma—for the most recently diverged cichlid fishes—and approximately 380 Ma—for the divergence between the coelacanth and ray-finned fish (Amemiya et al. 2013). Furthermore, to study the role of NC genes in cases of extreme phenotypic divergence and adaptive radiations, we included six cichlid species (family Cichlidae) for which genomes have recently become available (Brawand et al. 2014; Elmer et al. 2014). Cichlids have undergone extremely rapid speciation and phenotypic diversification that is often associated with NC-derived traits, such as craniofacial morphology and coloration. This is especially true for cichlid

![Diagram](image)

**Fig. 1.** Classification of neural crest-related genes in respect to developmental expression subunits. (A) Thirty-one genes from the NC GRN are classified into four groups depending on their deployment during development, after Betancur et al. (2010): Genes responsible for early induction of future NC (1), genes defining the NPB that lines the neural plate before neurulation (2), specifiers of the NC itself as well as regulators of their migration (3), and genes determining the fate of migrated multipotent NC cells (4). (B) Genes in this study sorted by developmental subunit. Five housekeeping genes (group C) were added to the analyses as a control group. (C) Regulatory connections between network genes without distinction regarding directionality and mode of interaction.
species that arose in the African Great Lakes (Neolamprologus brichardi, Astatotilapia burtoni, Metriaclima zebra, and Pundamilia nyererei) that are one of the best examples for adaptive radiation and fast phenotypic diversification in vertebrates (Turner 2007; Brawand et al. 2014; Henning and Meyer 2014).

The aim of our study was to ascertain whether sequence conservation among the components of the NC GRN is linked to network topology or developmental sequence and how patterns of natural selection might have changed during evolution. Here, we studied the molecular evolution of genes, most of which are transcription factors and guidance molecules (fig. 1) that are part of the NC GRN. We analyzed their molecular divergence in 16 species of bony fishes: 15 actinopterygians (ray-finned fish) and 1 sarcopterygian (lobe-finned fish) that we used as outgroup (fig. 2). Furthermore, we looked for evidence of positive selection or relaxation of purifying selection in these genes among the species-rich and phenotypically diverse clade of African cichlids, where five genomes have become recently available (Brawand et al. 2014).

Materials and Methods

Data Set

A total of 31 genes associated with the development of the NC in bony fish were included in this study. The genes were chosen based on prior characterizations of NC GRN (Sauka-Spengler and Bronner-Fraser 2008; Betancur et al. 2010; Simões-Costa and Bronner 2015). Only genes that could be found in greater than 80% of our selected species were analyzed. Depending on the time of deployment during development (Sauka-Spengler and Bronner-Fraser 2008; Betancur et al. 2010; Simões-Costa and Bronner 2015), genes were grouped into one of four developmental groups: Induction genes, NPB specification genes, NC specification genes, and NC differentiation genes (fig. 1 and supplementary table S1, Supplementary Material online). Additionally, we analyzed a control group of five housekeeping genes that are not directly related to NC formation, but for which we expected a very high conservation and comparable dN/dS ratios (see below). We downloaded gene sequences of 11 of the 16 species from Ensembl release 75 (Flicek et al. 2014) (fig. 2). After identifying orthologous loci using the EnsemblCompara pipeline (Vilella et al. 2009), we extracted the longest transcript coding sequence (CDS) of each gene. Sequences for cichlid species other than the Nile Tilapia, Oreochromis niloticus, were obtained from the deposited genome assemblies (Brawand et al. 2014; Elmer et al. 2014) (fig. 2). Orthologous loci in cichlid genomes were identified by BLASTN searches using Nile Tilapia sequences as queries and an e-value threshold of $10^{-4}$. The integrity of obtained sequences was double-checked by BLASTN searches against available transcriptomes to ensure correct reading frames. Gene orthology was confirmed using phylogenetic analysis (see below). In total, this study contains 483 sequences and 36 genes, each containing at least 13 of the 16 species.

Multiple Sequence Alignment and Phylogeny Reconstruction

Multiple sequence alignments of individual genes were constructed with translatorX (Abascal et al. 2010): After translation, sequences are aligned at the amino acid level using MUSCLE (Edgar 2004), unreliable amino acid positions are removed with Gblocks (Castresana 2000) under the least stringent parameters, and the corresponding nucleotide alignments are created guided by trimmed amino acid alignments. Maximum-likelihood phylogenetic reconstruction was performed with PhyML version 3.1 (Guindon et al. 2010) after best-fit models of evolution were chosen based on AICc (Akaike information criterion with correction) scores (Hurvich and Tsai 1989) calculated in jModeltest version 2.1.5 (Posada 2008). Due to the low sequence divergence among cichlids, single gene alignments contained too few informative sites to permit reconstructing reliable phylogenetic trees. Therefore, we used the previously reported species phylogeny (Brawand et al. 2014) for further analyses of cichlid data.

Saturation of nonsynonymous substitutions in gene alignments could in principle bias dN/dS estimates, even though probabilistic methods have been proven robust in simulation studies (Yang 2006). To exclude this possibility, we built plots of uncorrected p-distances against maximum-likelihood distances for both synonymous and nonsynonymous sites. MEGA version 6 (Tamura et al. 2013) was used to calculate genetic distances, using the parameters estimated by jModeltest for maximum-likelihood distances. Saturation plots did not show noticeable sequence saturation upon visual inspection, supporting the notion that dN/dS calculations are reliable.

Computation of dN/dS Ratios

We estimated the effects of selection on genes by calculating ratios between nonsynonymous and synonymous substitution rates (dN/dS). Using yn00 of the PAML package version 4.7 (Yang 2007) we calculated dN and dS values for pairwise comparisons between all 16 species. Divergence times used for mapping the values onto the phylogeny were taken from the TimeTree database (Hedges et al. 2006).

Using codeml (within the PAML package) we inferred dN/dS values using maximum likelihood. Under neutrality, both synonymous and nonsynonymous changes are expected to accumulate at equal rates (dN/dS = 1). An excess of nonsynonymous over synonymous substitutions is viewed as evidence of positive selection (dN/dS > 1), whereas the opposite is true for negative selection (dN/dS < 1). Measures of dN/dS for each alignment provided information of selective pressures across the different genes and developmental groups (fig. 1), based
on the M0 model that averages $dN/dS$ across codons and sequences. Three more sophisticated models were used to examine changes in $dN/dS$ between codon positions (site models), lineages (branch models), and codon positions in particular lineages (branch-site models). 1) Site models can detect specific codons that evolved under positive selection by comparing the M2a and M8 models of positive selection against the nearly neutral models M1a and M7, respectively. The M7/M8 comparison is similar to M1a/M2a, but assumes $dN/dS$ to follow a beta distribution, which can take a variety of shapes (e.g., uniform, linear, exponential, bell) depending on two shape parameters that appear as exponents of the random variable. In this sense, M7 is a more flexible null model than M1a (Yang 2006). 2) In branch models, $dN/dS$ values are calculated for a predefined lineage or lineages (foreground) separately from the rest (background), allowing the examination of changes in selection in specific lineages. In this case, the alternative model assuming different $dN/dS$ across lineages is compared against the M0 model. 3) The branch-site test of positive selection compares model A (that allows $dN/dS$ to vary among sites in predefined branches) and a null model assuming nearly neutral evolution (codons can only have $dN/dS \leq 1$). For each of the three model types, the relative fit of nested pairs of models was compared by differences...
in log-likelihood against a chi-square distribution (i.e., likelihood ratio test), whereas the fit of nonnested models was assessed by differences in AIC scores (Akaike 1974). In both site and branch-site models, positively selected codons were estimated by using a Bayes empirical Bayes (BEB) procedure (Deely and Lindley 2012). Codons under positive selection were only considered, if they had a posterior probability $> 0.95$. To discard suboptimal results by maximum likelihood getting stuck in local maxima, computations were repeated with different initial $d_{N}/d_{S}$ values (0.05, 0.4, 1.5, and 10) (Bielawski and Yang 2004). All calculations used a F3×4 codon substitution model, where expected codon frequencies are derived from three sets of nucleotide frequencies for the three codon positions. Further scripts used for computational analyses were generated in R (http://www.r-project.org, last accessed October 28, 2015), or Bioconductor (http://www.bioc conductor.org, last accessed October 28, 2015), using R packages cluster, ggbiplot, ggplot2, ggplots, grDevices, mclust, and RColorBrewer.

Multivariate Cluster Analysis

The clustering analysis of sequence conservation features was performed as previously described for the floral organ specification GRN (Davila-Velderrain et al. 2014). The four variables used were 1) the coefficient of variation in protein sequence lengths, 2) mean pairwise DNA and 3) protein distances, and 4) $d_{N}/d_{S}$ ratios. Genetic distances were calculated in MEGA 6. A principal component analysis—based on the different sequence conservation features 1–4—was performed in R using the princomp and ggbiplot libraries and normal data ellipses were generated with the standard probability (0.69). The cluster analysis itself was conducted with the package mclust version 4 using the implemented function “Mclust.” Mclust uses the Bayesian Information Criterion (BIC) to identify the best-fit model for constructing covariance matrices that define the clusters within the data set. To validate the clustering results, we calculated the similarity of the clustering decisions to the four a priori conceived developmental subgroups of the NC GRN (fig. 18) using the Rand index (Rand 1971). Through bootstrap resampling of 100,000 random clusters of similar composition, we determined whether the Rand index of our clustering is significantly higher than the Rand index of randomly generated clusterings with the same number of clusters compared with the four developmental groups.

Results

Changes in Selective Pressure Follow an Hourglass Pattern along Developmental Timing

Genes were grouped into the following four developmental groups: 1) Induction genes, 2) NPB specification genes, 3) NC specification genes, and 4) NC differentiation genes (fig. 1 and supplementary table S1, Supplementary Material online).

The four groups (although partially overlapping) act in chronological order within the NC GRN, with the induction genes (1) being the first genes active and NC differentiation genes (4) being active during the last steps of NC development (fig. 1). Since specification genes (3) regulate NC development around the phylotypic stage, they would be expected to exhibit the strongest conservation according to the hourglass model. On the contrary, following the funnel-like model, the early expressed Induction genes (1) should be most conserved (fig. 1A). We added five housekeeping genes as a control group. Here, we expected consistently high conservation throughout all analyzed species thereby serving as comparative base line for highly conserved genes. All analyzed genes—both from NC group and control group—show signs of strong purifying selection, with $d_{N}/d_{S}$ values well beneath what would be considered neutral evolution ($d_{N}/d_{S} = 1$). For ray-finned fish, ratios among NC-associated genes range from 0.024 in zic1 to 0.150 in bmp7a (mean: 0.072 ± 0.030) (fig. 3B, table 1, and supplementary table S2, Supplementary Material online). Values in the control group of housekeeping genes are consistently low and range from 0.021 to 0.057 (mean: 0.040 ± 0.015) (fig. 3B, table 1, and supplementary table S2, Supplementary Material online).

In an effort to discover differences in selective regimes between genes deployed at different developmental points, we analyzed the distribution of $d_{N}/d_{S}$ values across the four developmental groups (fig. 1A–C) to determine whether the distribution of molecular constraints follows an hourglass (fig. 3A, left) or a funnel-like distribution (fig. 3A, right). To compare and evaluate the fit of the data to these alternative hypotheses, we performed a linear model fit (fig. 3C and supplementary fig. S1, Supplementary Material online). AIC values for the different linear fit curves favor a quadratic fit over a simple linear model (fig. 3C and supplementary fig. S1, Supplementary Material online). A quadratic fit supports the idea of an hourglass type of conservation across the NC GRN versus a funnel-like or random distribution. In fact, the two central groups (2 and 3) have significantly lower $d_{N}/d_{S}$ values (and therefore higher constraints) than the early and late groups (1 and 4) (t-test; $P=0.0486$; fig. 3D), whereas the differences between early (1 and 2) and late groups (3 and 4) are not significant (t-test; $P=0.5973$; fig. 3D).

Evolutionary Conservation Features Are Highly Homogeneous across the NC GRN

In an attempt to find groups within the NC GRN that display similar evolutionary patterns, we conducted an unbiased clustering of genes based on different conservation features. This method has previously been reported to yield clear results for another developmental GRN (Davila-Velderrain et al. 2014). An initial clustering of NC GRN genes using four conservation features ($d_{N}/d_{S}$, DNA and protein sequence distances and coefficient of variation of protein length) resulted in a
classification of genes into ten different clusters (supplementary fig. S2 and tables S3 and S4, Supplementary Material online). Ten clusters is more than any initially anticipated biological classification of genes (such as GRN topology; fig. 1C). The members of the NC GRN are highly conserved at the sequence level, which is supported by the comparison with a previously analyzed developmental GRN that shows more variation and therefore a clearer clustering (Davila-Velderrain et al. 2014) (supplementary fig. S3, Supplementary Material online). We repeated the clustering analyses for modified species-sets for which we removed phylogenetically very distant (spotted gar, coelacanth) or very close (cichlids) species, in order to exclude the possibility that our clustering results could be biased by too many or too few differences, respectively (supplementary tables S5 and S6, Supplementary Material online). Clustering analyses on these modified data sets rendered the same clustering results.

The best-fit model obtained in the clustering analysis (EEV) utilizes covariance matrices of equal shape and volume (BIC = 627.46) (supplementary fig. S2, Supplementary Material online). The ten identified groups show a Rand similarity index of 0.71 when compared with the four initial function-related developmental subgroups. The clustering with Rand = 0.71 has a lower similarity to the predefined developmental groups than 15% of random clusterings of ten clusters. A principal component analysis of all NC genes including all conservation features reveals a rather homogeneous cluster of genes (fig. 4A). However, although the four developmental groups are overlapping, centroids of the central groups (2 and 3) separate from the early and late groups (1 and 4) on the first PC axis (73% of variance) (fig. 4A). The grouping between middevelopment versus early/late groups further supports an hourglass pattern based on more similar evolutionary conservation. In fact, when comparing the two pairs of groups solely...
on the first principal component (PC 1) the difference between groups 1/4 and 2/3 (t-test; \( P = 0.005 \)), but not between groups 1/2 and 3/4 (t-test; \( P = 0.654 \)) is statistically significant (fig. 4B).

### Molecular Constraints of the Majority of NC Genes Are Constant throughout Ray-Finned Fish Evolution

In an effort to gain insight into if and how molecular constraints changed during the evolution of ray-finned fish, we calculated \( dN/dS \) for pairwise comparisons between all species and plotted them against their estimated divergence times (Hedges et al. 2006; Genner et al. 2007; Near et al. 2013; Brawand et al. 2014) for both single genes and gene groups (fig. 4C and supplementary figs. S4 and S5; Supplementary Material online). Additionally, we performed comparisons centered on single species to emphasize differences of comparisons to particular species (supplementary fig. S6, Supplementary Material online). For stable selective constraints, synonymous and nonsynonymous substitutions are expected to accumulate at constant rates resulting in \( dN/dS \) values that do not change with evolutionary time. Indeed, both nonsynonymous (supplementary fig. S7, Supplementary Material online) and synonymous substitution rates (supplementary fig. S8, Supplementary Material online) can be described as a linear function of evolutionary distance (Nei et al. 2010). Consequently, \( dN/dS \) values for genes of the NC GRN were almost constant, with only a slight decrease in \( dN/dS \) ratios from long to short evolutionary distances (\( \Delta dN/dS = -0.006 \) per 380 Ma; linear regression; \( P = 0.015 \)) (fig. 4C and supplementary figs. S4 and S5, Supplementary Material online). To account for possible effects due to phylogenetic nonindependence and species selection biases, we repeated the analyses with two differently composed reduced data sets (supplementary figs. S9 and S10, Supplementary Material online) and two phylogenetically completely independent data sets of pairwise-comparisons (supplementary figs. S11 and S12, Supplementary Material online). The analyses of these data sets suggest that \( dN/dS \) values are rather constant over evolutionary time (linear regression; \( P > 0.05 \) for all data sets).

Additionally, pairwise comparisons (fig. 4D and supplementary figs. S5–S15, Supplementary Material online) provide further evidence for the particular high selective constraints among NC specification genes. In almost half (48.5%) of all 66 pairwise comparisons (without radiating cichlids) of the group of NC specification genes, \( dN/dS \) values of NC specifiers are smaller than the average value of the housekeeping gene group. Furthermore, all 66 (100%) pairwise \( dN/dS \) values of group 3 are below the average of the remaining NC groups. This is further evidence for the strong constraints of the NC specification genes that are active around the phylotypic stage and provides additional support for the hourglass model of molecular constraint in the NC GRN.

### Selective Constraints of Late NC Genes Increased after the Split between Ray- and Lobe-Finned Fish

To investigate whether there are changes in selective constraints across time within specific groups of the NC GRN, estimated pairwise \( dN/dS \) values were plotted as a function of evolutionary time, separately for the five groups of genes (fig. 4C). The NC differentiation genes (group 4) were the only group of genes that showed a significant change in \( dN/dS \) values (fig. 4C). This suggests an increase of molecular constraints throughout ray-finned fish evolution within the NC differentiation genes of group 4 (\( \Delta dN/dS = -0.043 \) per 380 Ma; linear regression; \( P < 0.0001 \)). We confirmed this result by analyzing the reduced data sets that account for taxon sampling biases (i.e., compensating for the presence of many closely related cichlids or equilibrating lineage representation), which equally showed no significant change for groups 1–3, and significant change for group 4 (supplementary figs. S9 and S10, Supplementary Material online). Significant changes in group 4 could be also observed in two phylogenetically independent data sets of species pairs (supplementary figs. S11 and S12, Supplementary Material online) as well as for all of the genes of group 4 analyzed individually (supplementary fig. S4, Supplementary Material online). To get a better grasp on the distribution of \( dN/dS \) values over all pairwise comparisons, we visualized all developmental groups as heatmaps (fig. 4D and supplementary figs. S13–S15, Supplementary Material online). In group 1, high \( dN/dS \) values are distributed rather randomly and apparently without a strong correlation to evolutionary distance of the respective species pair (fig. 4D and supplementary fig. S13, Supplementary Material online). In contrast, for group 4 genes there is a trend of an increase in selective constraint. High \( dN/dS \) values are consistently found within the comparisons against coelacanth, spotted gar, and zebrafish. The highest \( dN/dS \) values are found among the pairwise comparisons among NC specification genes.

| Group 1–4 | 0.072 ± 0.030 | 0.086 ± 0.038 | 0.066 ± 0.023 | 0.060 ± 0.031 | 0.082 ± 0.022 | 0.040 ± 0.015 |
|---|---|---|---|---|---|---|
| Group 1 | 0.064 | 0.090 | 0.065 | 0.053 | 0.090 | 0.046 |
| Group 2 | 0.024 (zic1) | 0.033 (wnt6b) | 0.024 (zic1) | 0.027 (twist2) | 0.059 (neurog1) | 0.021 (rpl8) |
| Group 3 | 0.150 (bmp7a) | 0.150 (bmp7a) | 0.094 (rsx1) | 0.121 (snai1b) | 0.108 (mitfa) | 0.057 (g6pd) |
that involve the coelacanth, however the observed trend is still significant, if the coelacanth is removed ($\Delta d_{N/dS} = -0.022$ per 380 Ma; linear regression; $P = 0.0001$). Interestingly, groups 2 and 3 as well as the control group have no clear trend; however, some species stand out more clearly than in group 1 (coelacanth and pufferfish for group 2, spotted gar and zebrafish for group 3, and blind cave fish for the control group; supplementary fig. S4, Supplementary Material online). The high values for the species pairs with blind cave fish and pufferfish are however due to a small number of genes with high $d_{N/dS}$ values ($hprt1$ and $rpl8$ for blind cave fish; $tflap2a$ and $id3$ for pufferfish) and disappear if one visualizes medians of groups instead of means (supplementary figs. S13–S15, Supplementary Material online).

Positive Selection on $rhob$ and $dct$ Genes
After assessing the evolution of the NC GRN as a whole, we investigated how selection regimes in single genes changed during ray-finned fish evolution. Codeml site models found clear evidence for positive selection in two genes. In $rhob$ (Ras homolog gene family member b), likelihood ratio tests were significant for the codeml models $M1a/M2a$ ($LR = 16.48$, $P=2.6 \times 10^{-4}$) and $M7/M8$ ($LR = 31.41$, $P=1.51 \times 10^{-7}$) comparisons (supplementary table S7, Supplementary Material online).
Material online). In both cases, BEB analyses identified amino acid position 186 (based on the Nile Tilapia sequence) as positively selected. Position 186 in Nile Tilapia is a proline located close to the C-terminal end of the protein. It is a site within a highly variable region at the C-terminus of the protein among the six cichlid species. Amino acid position 186 is three amino acids away from a highly conserved C-terminal motif (CVNCKVL in the coelacanth/CFKCCVL in the African cichlids) that is involved in the localization and degradation of the protein (Pérez-Sala et al. 2009). The second gene with positively selected codons was dct (dopachrome tautomerase), where the test comparing M7/M8 models was significant (LR = 16.68, \( P = 2.3 \times 10^{-5} \)) and BEB analysis showed that amino acid position 7 (in Nile Tilapia) has evolved under positive selection (supplementary table S7, Supplementary Material online). The amino acid at position 7 is a glycine in all cichlids. This glycine is shared with the flatfish Xiphophorus maculatus, but it is otherwise variable among the rest of species. The N-terminus of the encoded enzyme, the dopachrome tautomerase (also TRP-2 for tyrosine related protein 2), is involved in its catalytic reaction (Hiroshi Sugimoto et al. 1999). However, both sites appear to lie within regions of strong variability of their respective amino acid sequences.

Differences in Selective Pressures Arose during Cichlid Evolution

To test whether the NC GRN evolved under different selective regimes in lineages that are highly diverse in NC-derived traits, we analyzed the family of cichlid fishes (fig. 2). Cichlids underwent exceedingly rapid speciation often associated with astonishing diversification of NC-derived traits (Albertson and Kocher 2006). This is especially true for species that arose in the African Great Lakes (Staňny and Meyer 1999; Albertson and Kocher 2006; Renz et al. 2011; Albertson et al. 2014; Powder et al. 2014). Therefore, we grouped the five cichlid species into two separate groups: One including cichlids associated with adaptive radiations (N. brichardi, A. burtoni, M. zebra, and P. nyererei) and the other including the rest of species (O. niloticus and Amphilophus citrinellus). Using branch models in codeml, we found significant differences in \( dN/dS \) ratios between these two groups in 7 of 31 genes (fig. 5A and supplementary table S8, Supplementary Material online). Most estimated \( dN/dS \) ratios were lower than 1, implying prevalence of purifying selection in both groups of species. Six of these genes showed a \( dN/dS \) that was at least 2.3 times higher in radiating cichlids than in nonradiating cichlid lineages: bmp7a, dla, oig2, rhob, sox10, and wnt8a (fig. 5A and supplementary table S8, Supplementary Material online). In addition, rhob had a \( dN/dS \) > 1 in radiating cichlids. This gene had already shown evidences of positive selection in site model analyses (see above). Interestingly, neurog1 was the only gene that showed the reverse pattern of a lower \( dN/dS \) in radiating cichlids (fig. 5A and supplementary table S8, Supplementary Material online).

To maximize information content in the comparisons between cichlids associated or not with evolutionary radiations, all branch models were repeated on alignments containing only the six cichlid species. The removal of more divergent ray-finned fish allowed establishing more reliable codon homologies within cichlids. Additionally, resulting alignments were longer after Gblocks filtering, thus including more informative codon sites. In this case, the four radiating African cichlids were compared with the Nile Tilapia (O. niloticus) and the Neotropical Midas cichlid (Am. citrinellus). Results are consistent with already reported analyses: Six of the genes from the earlier analysis show significantly higher \( dN/dS \) in radiating cichlids (fig. 5B and D and supplementary table S9, Supplementary Material online). Five additional genes display significant differences in \( dN/dS \) between radiating and nonradiating cichlids: ents1, fgf2, id3, mitfa, and snai1b (fig. 5B and D and supplementary table S9, Supplementary Material online). Except for snai1b, all of them have an at least 2.3-fold higher \( dN/dS \) in radiating cichlids.

To ascertain whether the changes in \( dN/dS \) inferred for radiation-associated cichlids were specific to these species or rather characteristic of the whole cichlid lineage, we further studied differences in \( dN/dS \) using branch models comparing all cichlids (as a foreground group) and all other fish species (as background) (fig. 5C and supplementary table S10, Supplementary Material online). In these tests rhob and wnt8a, two of the genes that appeared in the earlier analyses, also showed a higher \( dN/dS \) in cichlids (fig. 5C and supplementary table S10, Supplementary Material online). The comparison of the relevant models by AIC suggests that the increase in \( dN/dS \) in wnt8a most likely occurred in radiating cichlids rather than in the whole cichlid lineage (\( \Delta AIC = 11.03 \)). Aside from that, we observed significant changes in selective pressure in eight additional genes in cichlids. Of these, two show an increase of \( dN/dS \) in cichlids (bmp2b and dct), whereas six show a decrease (foxd3, gbx2, pax3a, neurog1, sox9a, and zic5) (fig. 5C and supplementary table S10, Supplementary Material online).

Natural selection is expected to act on particular amino acid positions of functional relevance, and during specific time periods after which selection turns into purifying to maintain adaptations (Zhang et al. 2005). Branch models average \( dN/dS \) over the alignment, and thus can have low power to detect positive selection if it has occurred in only a few localized positions (Yang 2006). Therefore, we used branch-site models that allow detecting positive selection in specific codons in particular branches. Based on the same comparisons between species groups as described above, branch-site
models detected position 186 in rhob to have evolved under positive selection in radiating cichlids (supplementary table S11, Supplementary Material online), as was already found by site models (supplementary table S7, Supplementary Material online). The comparison of the relevant models favored the hypothesis that position 186 in rhob is under positive selection on all studied fish species rather than on cichlids specifically (best was M8 site model, with ΔAIC > 25.9 to M2a, branch and branch-site models assuming dN/dS changes in either radiating or all cichlids).

**Discussion**

The NC GRN Is under Strong Selective Constraint

The NC is a transient developmental unit of migrating multipotent cells that contributes to many different tissues within the vertebrate body. NC development is directed by an intricate GRN of transcription factors and signaling molecules. Here, we analyzed the molecular evolution of the CDSs of involved genes to gain insights on how the genes within this network change over evolutionary times and look for patterns...
of changes in selective pressures. Our results show that almost all genes of the NC GRN are under strong selective constraint across ray-finned fish, as evidenced by the values of $d\text{N}/d\text{S} < 1$ (Yang 2006). These overall low $d\text{N}/d\text{S}$ values found across the whole NC GRN are typical for a regulatory module that is active in such early developmental stages (Garfield and Wray 2009). Additionally, pleiotropic effects that affect the development of other tissues and cell types that changes in these genes might create, are an explanation for the strong purifying selection they evolve under (Wagner and Zhang 2011).

Molecular Evolution of NC GRN Modules Supports the Developmental Hourglass Model

Our data show a distribution of $d\text{N}/d\text{S}$ values across the four developmental groups that support an hourglass model of conservation, where intermediate stages (groups 2 and especially 3) are under stronger purifying selection than earlier and latter stages (groups 1 and 4) (fig. 3C). This result is in line with previous studies that support the hourglass model (Hatzkani Covo et al. 2005; Roux and Robinson-Rechavi 2008; Piascecka et al. 2013). There is increasing evidence that the phylotypic stage (i.e., the time point in development when phylogenetically diverse species resemble most closely to each other) can be indeed visualized at the genomic and transcriptomic level by increased constraints (Hatzkani Covo et al. 2005; Roux and Robinson-Rechavi 2008; Levin et al. 2012; Piascecka et al. 2013; Irie and Kuratani 2014). A recent comparison of $d\text{N}/d\text{S}$ ratios of genes expressed at different developmental stages in zebrafish (Piascecka et al. 2013) did not show significant differences between the studied stages—however, the distribution of means suggested that the highest degree of sequence conservation occurred around the phylotypic stage.

It has been hypothesized that the mechanisms behind the phenomenon of high selective constraint during middevelopment (phylotypic stage) is their strong internal recursive wiring within these central nodes of the regulatory network (Davidson and Enwin 2006) (fig. 1C). This is valid for many of the highly conserved genes that can be found among the NC and plate border specifiers (fig. 1B) including msxe, pax3a/b, tfap2, zic1/5, foxd3, snail1b, and sox9/10 (Betancur et al. 2010; Simões-Costa and Bronner 2015) and would explain the occurrence of stronger purifying selection in this subset of genes. The relatively relaxed selective pressure in differentiation genes is expected under all the above models of GRN evolution. Differentiation genes are most divergent in pairwise $d\text{N}/d\text{S}$ comparisons over evolutionary time and have some of the highest overall $d\text{N}/d\text{S}$ ratios (codeml M0 model). This agrees with predictions that genes that are more downstream would also experience less severe selective constraints. In this regard, it is interesting to note that group 4 shows lower $d\text{N}/d\text{S}$ values in pairwise $d\text{N}/d\text{S}$ comparisons between more recently diverged species (supplementary fig. S12, Supplementary Material online). An increase of genes depending on the NC differentiation genes (i.e., further downstream in different GRNs) might have led to a gradual increase in selective pressure throughout ray-finned fish evolution.

Clustering Does Not Link Conservation Features to Developmental Groups or Network Topology

Clustering the NC GRN genes without prior hypotheses resulted in a classification into ten groups. This number of clusters is much higher than any anticipated biological grouping of the genes. Also, the principal component analysis of gene conservation features did not reveal a clear aggregation pattern of the genes into one of the preconceived classifications (supplementary fig. S2, Supplementary Material online). However, comparing the PC1 of the central groups (2 and 3) with early and late groups (1 and 4) reveals a significant difference ($P=0.005$; fig. 4B), further supporting an hourglass pattern of conservation. Additionally, the results reveal that the variation in protein size is a minor contributing factor in the variation of conservation in NC GRN genes, as it is the variable that constitutes most of the second principal component (fig. 4A).

A comparison of the first two principal components from our data set combined with a previous study of another developmental GRN (Davila-Velderrain et al. 2014) reveals that the NC GRN is more conserved along the two axes (supplementary fig. S3, Supplementary Material online). Clustering techniques based on molecular conservation features might thus not be a viable method for GRNs that are extremely conserved at the sequence level.

Specific Codons Are under Positive Selection in rhôB and dct

Positively selected codons across the studied species could be detected in two genes: rhôB and dct. Interestingly, both rhôB and dct are deployed rather late in development. RhôB is grouped within the NC specifiers, but it is associated with the later steps of this phase including delamination and migration (Liu and Jessell 1998). The dct gene is directly regulated by mitfa, that is a direct part of the enzymatic pathway for melanin production (Potterf et al. 2001). The fact that positive selection was only found in late genes and not in any of the earlier genes shows that gene deployed earlier during development are under stronger selective constraints. This observation, taken together with the pattern of $d\text{N}/d\text{S}$ values between developmental groups, lends further credence to the assertion that the evolution of the NC GRN follows an hourglass model.

Release of Constraint in the Radiating African Cichlid Lineages

Periods of increased or decreased selective pressure can be detected through statistical testing of branch models.
Our results show that six genes (bmp7a, dla, olig2, rhob, sox10, and wnt8a) showed higher dN/dS ratios in the radiation-associated and phenotypically diverse cichlids, both when compared with nonradiating cichlids or with all other fish. This suggests that selective pressure acting on these genes was relaxed in cichlids associated with evolutionary radiations (fig. 5A, C, and D). Five more genes (fgf2, id3, ets1, snail1b, and mitfa) showed significantly different dN/dS between radiating and nonradiating cichlids (fig. 5B and D). Overall, there is statistical evidence for relaxation of purifying selection in 10 of 31 genes in the NC GRN. One might therefore speculate that this release of constraint in cichlids might be in part responsible for the enormous diversity in NC-derived traits displayed by this family, in particular a number of trophic and phenotypic adaptations that enabled the astonishing cichlid radiations (Trainor et al. 2003; Albertson and Kocher 2006; Clabaut et al. 2007). For two of the above mentioned genes (id3 and rhob), a dN/dS > 1 is inferred for the lineage of radiating cichlids, which is evidence for positive selection (fig. 5A and B). The fact that these two genes show such high dN/dS values in cichlids is attributable to the very low amount of synonymous substitutions, which are outnumbered by nonsynonymous changes. Surprisingly, even though we found a dN/dS > 1 for id3 in cichlids associated with adaptive radiations, no specific codon sites could be identified to have undergone positive selection (estimated in either the site test or the branch-site test).

Tests using branch models were also used to compare all cichlids against the rest of fish species. Only three of the genes that already appeared in the above analysis also showed significantly different dN/dS values in this case: neurog1, rhob, and wnt8a (fig. 5D). For rhob and wnt8a, the comparison of AIC scores between relevant branch models favored the scenario of relaxation of purifying selection in the lineage of radiating cichlids rather than in the whole cichlid lineage (ΔAIC = 2.7 and ΔAIC = 11.03, respectively), whereas for rhob both scenarios had a similar fit (ΔAIC = 0.8). Notably, the other nine genes that displayed higher dN/dS in radiating cichlids did not show significant differences in the comparisons with all cichlids, which is further evidence that these changes in purifying selection have indeed arisen in the radiating clade. It may also be of note that all genes showing higher dN/dS in cichlids are either genes that are deployed very early (bmp2b and wnt8a) or rather late (rhob and dcl) in development whereas the other genes are more central and have lower dN/dS in cichlids. Nevertheless, the remaining genes that do not show significant differences in dN/dS do not follow a congruent pattern with this observation and thus this conclusion should be taken with caution. The interpretation of lower dN/dS found in cichlids compared with all other lineages must be taken with precaution, as it might be due to other important changes in selective regions of genes in species outside of the cichlid lineage.

Positive Selection at Specific Sites in Radiating Cichlids

Site model tests can detect the effect of selection at specific sites and ignore among-branch differences, whereas branch models allow detecting periodic changes in selective pressure and ignore among-codon differences. Overall, both types of models are highly conservative (Yang 2006). Branch-site models account for both effects and are in principle more sensitive than branch or site models alone. Tests of branch-site models have proven to yield insightful results in other studies, even involving some of the species we have included in this study (Fan et al. 2011; Amemiya et al. 2013). For the NC GRN, branch-site tests of positive selection applied to the comparisons with either radiating or all cichlids as foreground branches only detected positive selection rhob. The comparison among different scenarios of dN/dS change for rhob suggests that the positively selected site 186 is most likely not exclusive of cichlids, but rather shared with other ray-finned fish.

Conclusions

Ray-finned fish exhibit an enormous diversity in NC-derived traits, and yet the NC GRNs and most of its genes have so far not been analyzed from an evolutionary perspective. Analyses of protein CDSs support an hourglass model of protein CDS conservation in the different subunits of the GRN. The common morphological developmental pattern of an “hourglass” is reflected and possibly caused not or not only by developmental constraints but rather by constraints at the underlying genetic or genomic, in the sense of gene network interactions, levels. Cluster analysis revealed the extreme degree of sequence conservation throughout the NC GRN, illustrating its deep roots within vertebrate evolution (Gans and Northcutt 1983; Shimeld and Holland 2000). Values of dN/dS are constant throughout long evolutionary time spans demonstrating the strong, obligatory integration of the NC GRN within ray-finned fish. Interestingly, NC differentiation genes, the last group to be deployed in development, is an exception to this pattern, as it shows increasingly stronger purifying selection in ray-finned fish after the split from lobe-finned fish. The genes identified to have released selective constraint in radiating cichlids (bmp7a, dla, ets1, fgf2, id3, mitfa, olig2, and sox10) could be considered for future studies involving NC-derived traits. Further analyses of the molecular evolution of regulatory interactions will continue to shed light on the evolution of the NC network (Kratochwil and Meyer 2015a, 2015b, 2015c).

Supplementary Material

Supplementary figures S1–S15 and tables S1–S11 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org). Alignments are available at Dryad, doi:10.5061/dryad.75427.
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