Sodium channel genes and the evolution of diversity in communication signals of electric fishes: Convergent molecular evolution

Harold H. Zakon, Ying Lu, Derrick J. Zwick, and David M. Hillis

Sections of *Neurobiology and *Integrative Biology and Center for Computational Biology and Bioinformatics, University of Texas, Austin, TX 78712; and *Bay Paul Center for Comparative and Molecular Biology, Marine Biological Laboratory, Woods Hole, MA 02543

Communicated by Gene E. Robinson, University of Illinois at Urbana-Champaign, Urbana, IL, January 6, 2006 (received for review November 10, 2005)

We investigated whether the evolution of electric organs and electric signal diversity in two independently evolved lineages of electric fishes was accompanied by convergent changes on the molecular level. We found that a sodium channel gene (Na\textsubscript{1.4}a) that is expressed in muscle in nonelectric fishes has lost its expression in muscle and is expressed instead in the evolutionarily novel electric organ in both lineages of electric fishes. This gene appears to be evolving under positive selection in both lineages, facilitated by its restricted expression in the electric organ. This view is reinforced by the lack of evidence for selection on this gene in one electric species in which expression of this gene is retained in muscle. Amino acid replacements occur convergently in domains that influence channel inactivation, a key trait for shaping electric communication signals. Some amino acid replacements occur or are adjacent to sites at which disease-causing mutations have been mapped in human sodium channel genes, emphasizing that these replacements occur in functionally important domains. Selection appears to have acted on the final step in channel inactivation, but complementarily on the inactivation “ball” in one lineage, and its receptor site in the other lineage. Thus, changes in the expression and sequence of the same gene are associated with the independent evolution of signal complexity.

Recent studies have shown remarkable convergent evolution at the morphological level in a number of taxa (11–14). The extent to which morphological convergence is accompanied by convergent molecular change is an interesting question and one that can be profitably addressed by comparing the evolution of ion channel genes underlying the EOD in both lineages of electric fish.

We focused on Na\textsuperscript{+} channel genes because the EOD in at least one species of gymnotiform is shaped by the properties of the EO Na\textsuperscript{+} current (10) and because of the wealth of information on channel structure/function from site-directed mutagenesis and point mutations underlying clinical syndromes (15–20).

Results

Convergent Loss of Na\textsubscript{1.4}a from Muscle and Gain in Electric Organ. Previously we cloned portions of six sodium channel genes from the gymnotiform *Sternopygus macrurus (21) and found two orthologs (Na6 and Na1, renamed Na\textsubscript{1.4}a and Na\textsubscript{1.4}b; A. Novak, M. Jost, Y.L., A. Taylor, H.H.Z., A. Ribera, unpublished data) of a single muscle-specific mammalian Na\textsuperscript{+} channel gene (Na\textsubscript{1.4}). Sodium channels are composed of four domains (DI–DIV), each with six membrane-spanning regions (S1–S6). We cloned portions of Na\textsubscript{1.4}a and Na\textsubscript{1.4}b from two other gymnotiforms (*Apterorhous leptorhynchus and *Brachyhypopomus pinicaudatus) and two nonelectric species (*Ictalurus punctatus and *Danio rerio), all in the superorder Ostariophysi, and in a mormyrid (*Gnathonemus petersii) and two nonelectric species (*Osteoglossum bicirrhosum and *Notopterus chitala), all osteoglossomorphs, the most basal teleost lineage. The widespread occurrence of both genes and their presence in basal teleosts indicates that they duplicated early in teleost evolution, likely the result of a genomewide duplication (22, 23).

We next determined the expression pattern of these genes in muscle and EO, if present, by RT-PCR. Na\textsubscript{1.4}a and Na\textsubscript{1.4}b are generally both expressed in muscle in nonelectric species (24), although levels of Na\textsubscript{1.4}b are low in muscle from the catfish *I. punctatus (Fig. 2). We do not know whether both genes are expressed in all muscle fibers or if they are differentially expressed in different fiber types. In electric fishes, Na\textsubscript{1.4}b is expressed in muscle in all four species and in the EO in the gymnotiforms; little or no Na\textsubscript{1.4}b is expressed in the EO of *G. petersii, a mormyrid. Na\textsubscript{1.4}a is expressed solely in the EO in *S. macrurus and *B. pinicaudatus, but in the muscle of *A. leptorhynchus. Interestingly, *A. leptorhynchus possesses a larval myogenic EO that functions only early in life, then degenerates as a neurally derived EO appears (25). We presume that Na\textsubscript{1.4}a is expressed in the larval EO in this species but were unable to obtain specimens for analysis. Na\textsubscript{1.4}a is expressed only in the EO in *G. petersii, a mormyrid. Thus, Na\textsubscript{1.4}a

Conflict of interest statement: No conflicts declared.

Abbreviations: EO, electric organ; EOD, electric organ discharge.

To whom correspondence should be addressed. E-mail: h.zakon@mail.utexas.edu.

© 2006 by The National Academy of Sciences of the USA

Divergence of animal communication signals accompanies reproductive isolation and speciation (1–3). Because most communication signals are under polygenic control (4), it is difficult to identify the contributions of particular genes to signal evolution. We studied the relationship of gene expression and sequence evolution to signal divergence in weakly electric fish because of the simplicity and species diversity of their electric communication signals and the dependence of those signals on ion currents. Ion currents are generated by ion channels, the genes for which are easily identified.

Two groups of teleosts, the mormyforms of Africa and the gymnotiforms of South America, have convergently evolved electric organs (EOs) whose weak electrical emissions function as communication signals and for electrololocation. There is a 100-fold difference in electric organ discharge (EOD) duration across species (200 μsec to 20 msec) (ref. 5; Fig. 1), with both lineages showing great diversity in signal waveform. Sexual and natural selection have likely played roles in the evolution of this diversity (6–8). Variation in EOD characteristics, especially pulse duration, forms the basis for detection of species-specific signals (9). This richness in the EOD waveform is based in part on the properties of the ion channels of the cells of the EO, the electrocytes (10). The diversity in membrane excitability of electrocytes contrasts with the simple excitability of vertebrate skeletal muscle from which electrocytes evolved, highlighting ion channel genes as potential candidates for strong evolutionary change in these fishes.
has been lost from muscle and expressed exclusively in the EO, an evolutionarily novel structure, in both groups of electric fish.

**Sequence Evolution of Na\(_{1.4a}\).** To investigate changes in the selective regime acting on the Na\(_{1.4a}\) gene, we used the maximum-likelihood codon models implemented in the **CODEML** program from the **PAML** package (26). Phylogenetic analyses using a number of methods all resulted in the same tree topology (Fig. 3), which resembles the recognized phylogenetic relationships of these species based on other criteria. This topology was fixed for all **CODEML** selection analyses. We applied site models (27) to estimate variation in the pattern of substitution across sites of all 11 taxa and branch-site models (26, 28) to estimate variation in the pattern of substitution across sites on specific lineages identified *a priori.*

The site models suggest that \(\omega (\text{dN/dS or the rate of nonsynonymous substitutions/synonymous substitutions})\) varies strongly between amino acid sites (M0 vs. M3, \(P \ll 0.0001\)) and that the majority of sites are highly constrained (57% of sites, \(\omega_0 = 0.01\)) (Table I, which is published as supporting information on the PNAS web site). There was no evidence of sites under positive selection over all lineages (M7 vs. M8, \(P = 0.56\)). We applied two branch-site models to test for sites under selection on the individual lineages associated with the EO (five terminal and three internal) by using a correction for multiple tests (\(\alpha = 0.006\)). Both branch-site tests agree on the presence of sites under positive selection on the terminal lineages leading to *Electrophorus electricus* and *B. pinnicaudatus*, as well as the lineage representing the ancestor of those two species. The test results differed for the lineages leading to *S. macrurus* and *G. petersii*, with the model B test suggesting sites under positive selection that were not found by the more conservative modified model A test. Neither test suggested sites under positive selection on the lineage leading to *A. leptorhynchus*, the lineage ancestral to *S. macrurus, B. pinnicaudatus,* and *E. electricus,* or the lineage ancestral to all gymnotiforms. These results suggest that the evolution of the Na\(_{1.4a}\) gene has been shaped to some extent by positive selection in all electric species in this study with the exception of *A. leptorhynchus,* in which Na\(_{1.4a}\) still resides in muscle.

**Convergent Amino Acid Replacements in Parts of the Channel Involved in the Final Step in Inactivation.** Regions critical for fundamental properties of Na\(^+\) channels, such as ion selectivity (the pore), voltage dependence (S4), inner lining of the channel (S5 and S6), and cytoskeletal localization (ankyrin-binding motif) are highly conserved in electric fish Na\(_{1.4a}\). Amino acid replacements tended to occur in S1–S3 or intra- or extracellular loops (Fig. 3B). Because Na\(^+\) channel inactivation is likely to have a major influence on electric organ pulse duration, we focus on regions involved in this process.

Inactivation involves a number of distinct domains of the Na\(^+\) channel; the final step in fast inactivation, the rapid closing of the channel during a sustained depolarization, is accomplished by the docking of the cytoplasmic loop between DIII and DIV (LIII–IV) with the S4–S5 linkers of DIII and DIV that line the inner mouth of the channel (29–32). This interaction is largely due to hydrophobicity. The LIII–IV loop possesses a highly conserved motif centered on the hydrophobic amino acids iso-leucine-phenylalanine-methionine, experimental replacements of which profoundly alter fast inactivation (31). In the mormyrid, which makes extremely brief action potentials, there is a nonconservative replacement (I to C) and the loss of a highly conserved neighboring hydrophilic site (Fig. 4A and D). Such an I to C replacement in the human cardiac Na\(^+\) channel gene (Na,1.5) speeds recovery from inactivation (32); this naturally occurring replacement in the mormyrid Na\(^+\) channel could be important in maintaining a nondecrementing EOD pulse in the bursts of EOD pulses given during social interactions. Nearby, an S to N replacement eliminates a protein kinase C consensus site (Fig. 4E). Phosphorylation of this site in brain Na\(^+\) channels slows inactivation (34); elimination of this consensus sequence would optimize for rapid inactivation.

The inactivation loop pivots around a number of highly conserved proline residues on the C-terminal end. Nonconservative replacements of one of these prolines occurs in two gymnotiforms (Fig. 4F). Site-directed mutagenesis of this proline in an expressed rat brain gene (35) perturbs inactivation, confirming its importance. Finally, there are radical replacements of a highly conserved glutamine in the mormyrid (Q to P) and the gymnotiform *B. pinnicaudatus* (Q to L), both of which generate short EOD pulses. This site has not yet been studied experimentally, but its alteration in the two electric fish taxa with the fastest EOD pulses hints that it may play a pivotal role in influencing the speed of inactivation.

Two gymnotiform species possess a nonconservative amino acid replacement (K or R for N) in the S4–S5 linkers in DIII (Fig. 4C). This linker forms part of the receptor for the inactivation loop and is highly conserved with an N at this site from cnidaria to mammals (H.H.Z., unpublished data); a mutation at this site of the Na,1.5 gene is causal for long QT syndrome (20) (Fig. 4C, asterisk). The effect of a positively charged amino acid in this position is not known; however, site-directed replacement of N with C or S in mammalian Na,1.4 or 1.5 induces incomplete inactivation and slows recovery from inactivation (20, 30). There are no amino acid replacements here in mormyrids.

Amino acid replacements occur in the S4–S5 linker in DIII in both groups of electric fish (K to T in mormyrid and G to C in gymnotiform; Fig. 4B). Mutations flanking these sites in the S4–S5 linker DIII in human Na\(^+\) channel genes disrupt slow inactivation (the propensity for Na\(^+\) channel to inactivate after prolonged usage) and are associated with muscle diseases (16, 36).

In sum, despite strong evolutionary conservation of the domains involved in the final step of the inactivation of Na\(^+\)
channels, there are nonconservative amino acid replacements in these regions in electric fish. Most striking are the amino acid replacements that occur in and around the critical isoleucine-phenylalanine-methionine inactivation motif in mormyrids and in the receptor site for that part of the molecule in gymnotiforms. These replacements suggest that the final step of the inactivation process is under selection in both lineages, but through complementary sides of the process.

Discussion
The main finding of this study is that the evolution of signal diversity in two independently derived lineages of electric fish is accompanied by convergent alteration of expression and changes in sequence within the same or complementary functional domains of the same Na\(^+\) channel gene, Na\(_{\text{v}}\)1.4a. Site-directed mutagenesis allows us to determine how specific amino acid replacements in this gene lead to variations in species-specific communication signals. Because a number of these replacements occur at sites that are otherwise highly conserved (i.e., Q to P or L in the inactivation loop) but previously not singled out for study, our results have also pinpointed a number of amino acid sites of potential functional importance for all Na\(^+\) channels.

A genome duplication at the origin of teleosts has been suggested as a substrate for the morphological diversity and extensive speciation characteristic of teleosts (22, 23). Nevertheless, there are few concrete examples of genes whose duplication is directly related to species diversity or the emergence of novel structures in fish. Although the duplication of Na\(_{\text{v}}\)1.4 in fish is not responsible for the genesis of the EO, it allowed for the compartmentalization of one gene duplicate in the EO, providing a substrate for the evolution and divergence of species-specific signals in two lineages of fishes.

Na\(_{\text{v}}\)1.4a and Na\(_{\text{v}}\)1.4b are both expressed in the muscle in nonelectric teleosts. Like mammals, fish have a diversity of muscle fiber types, the properties of some types even varying along the body (37, 38). We do not know whether both Na\(^+\) channel genes are redundantly expressed in every muscle fiber or they are differentially expressed in particular groups or functional types of muscle fibers to “tune” the electrical responsiveness of the fiber type to its function.

Multiple lineages of teleost fish have independently evolved sonic communication systems by using a diversity of mechanisms, including the vibration of the swimbladder by specialized muscles at frequencies of hundreds of hertz (39–42). It would be intriguing to determine whether a comparable convergent compartmentalization and specialization of Na\(^+\) channels genes occurs in muscles specialized for acoustic communication to allow for rapid or prolonged firing of muscle fibers needed to generate acoustic signals. Interestingly, some mormyrid species generate sounds up to 300–400 Hz, and it is believed that their swimbladder muscles may contract at those rates (40). However, because mormyrids have already “lost” one muscle-expressing Na\(^+\) channel gene to the EO, it raises the intriguing possibility that Na\(_{\text{v}}\)1.4a might also be expressed in the swimbladder muscle of mormyrids, where it supports production of a second communication signal in another modality.

Whatever costs electric fish may have incurred for the loss of this gene from muscle are clearly outweighed by the benefits of its gain in the EO. It will be interesting to see whether there are “compensatory” changes in expression or sequence in the remaining muscle Na\(^+\) channel gene (Na\(_{\text{v}}\)1.4b) or by what alterations in transcriptional mechanisms expression was lost from muscle and gained in an evolutionarily novel structure in both electric lineages.
Na\textsuperscript{+} channels do not function alone; they are modified by beta subunits and act with other currents, such as K\textsuperscript{+} currents, to shape electrical activity. It will also be intriguing to see whether any of these other genes show comparable alterations in expression or sequence especially in those species with extremely rapid EOD pulses.

Na\textsuperscript{+} channel genes are highly constrained, as evidenced by the large number of amino acid replacements that change the properties of the Na\textsuperscript{+} current and lead to neurological, cardiac, and muscular diseases (15–20). A critical step in the evolution of electric organs is the disabling of excitation-contraction so that the organ does not twitch when it discharges. Even when expressed in a noncontractile EO, amino acid replacements in Nav1.4a would be selected against as long as Nav1.4a is also still expressed in muscle, as occurs in *A. leptorhynchus*. However, once Nav1.4a was solely expressed in a noncontractile organ, amino acid replacements might occur because of relaxation of negative selection or be selected precisely because these amino acid replacements would have generated variation in Na\textsuperscript{+} currents and, ultimately, diversity in communication signals. Thus, similar to olfactory receptor or peptide venom genes, spatially restricted expression is permissive for sequence evolution (43, 44).
We noted amino acid replacements in electric fish Na1.4a at or adjacent to amino acid sites where mutational substitutions in human Na+ channel genes are associated with a disease. Not only will study of electric fish Na+ channel genes be a window on the processes of speciation and evolution of communication signals, but identifying amino acid sites or regions of the Na+ channel that have undergone extensive evolution in electric fish may be useful for predicting amino acid changes putatively involved in channelopathies or pinpointing functionally important novel regions of Na+ channels.

**Methods**

**Sequences.** We cloned portions of Na1.4a and Na1.4b (S2 domain II to S2 domain IV) from the gold-lined black knifefish (S. macrurus: AF378144), the pin-tail knifefish (B. pinnicuda-tus: DQ351534), the brown ghost (A. leptorhynchus: DQ351533), the channel catfish (I. punctatus: AY204537), the zebrasid (D. rerio: DQ149506), the elephant nose mormyrid (G. Petersii: DQ275142), the arowana (O. bicirrhosus: DQ336343), and the clown knifefish (N. chitala: DQ336344).

For phylogenetic analyses, we used additional published sequences for the electric eel (E. electricus: M22252), the Japanese pufferfish (Tetrpodon nigroviridis: CAAE01014976), and the Japanese pufferfish (Takifugu pardalis: AB030482).

**Cloning and Sequencing.** Tissue was homogenized and RNA was isolated by the guanidinium thiocyanate method with STAT-60 (Ambion, Austin, TX). RNA was quantitated by absorbance readings, and quality was confirmed by gel electrophoreses. RNA was reverse transcribed and amplified by PCR. The PCR products were cloned into the TOPO TA vector (Invitrogen) and grown in bacteria. Inserts were sequenced in both directions with an ABI 3100 sequencer (Applied Biosystems).

**Tissue Expression.** One microgram of total mRNA from dorsal trunk muscle (taken from behind the head) in all species and EO from electric fish was reverse transcribed to cDNA with specific primers for each channel. Specific primers targeted intracellular loop DI-DIII and produced fragments of ~700–1,000 bp, depending on the species. Temperature was optimized for each primer set. PCR was carried out for 10, 20, 30, or 40 cycles. Thirty cycles was chosen as the standard for comparison because it produced a strong signal without saturation of the reaction. The use of primer sets for both Na1.4a and Na1.4b in both tissues served as an internal positive control for the quality of mRNA and the robustness of the primers.

**Codon Maximum Likelihood Analyses.** Sequences were edited and aligned. The nucleotides used corresponded to the following nucleotides from the Electrophorus sequences (starting at initial ATG): 2,149–2,882; 2,896–2,917; 2,998–3,028; 3,038–3,084; 3,151–3,195; and 3,232–4,531. Phylogenetic trees were estimated with PAUP* (45).

Analyses were performed by using the CODEML program from PAML version 3.14 (26–28, 46–49). For a given tree and codon model, CODEML finds the set of parameter values that maximize the probability of observing the data, i.e., the likelihood score. The likelihood scores of two nested models are compared via a likelihood ratio test. A significant likelihood score difference between two models that differ in the allowance of a particular feature (such as the presence of positive selection) is taken as evidence for that feature. All models were run under the F3 × 4 setting of CODEML in which the equilibrium codon frequencies are taken to be the product of empirical nucleotide frequencies at each codon position. Models M0, M3, M7, and M8 were as described in the literature (27) and are used to detect specific sites under positive selection across all lineages in a data set.

Branch-site models require a division of lineages into foreground branches (branches of interest specified a priori) and background branches (the remaining branches) and test for evidence of positive selection on specific foreground branches, each in a separate test.

Branch-site model B allows for sites that were constrained to some degree on the background branches to evolve under positive selection on the foreground branches. This model may be compared via a likelihood ratio test to site model M3 with two site categories as a test for sites under positive selection on the foreground lineage. This test was applied as in ref. 28. It has been suggested that this test can result in false positives when sites are released from evolutionary constraint on the foreground branch but none experience positive selection (48).

A recently introduced, more conservative branch-site test was also used, now termed “the branch-site test of positive selection” (49). The test uses a modified form of branch-site model A in which 0 is estimated and may take any value between zero and one. This model is compared via a likelihood ratio test to a restriction of the same model (in which 0 is fixed at 1) with 1 df. This test is less likely to mistake relaxation of constraint for positive selection and has been shown to have excellent false-positive rates but relatively low power.

Branch-site models are also able to identify individual sites on the foreground branches with a high probability of being positively selected by using an empirical Bayesian approach. It has been shown in the context of site models that the accuracy of these site predictions is low when the number of sequences in the data set is small (<15 sequences) (50). Because our data set is small at 11 sequences, we do not present the Bayesian probability data for individual sites.

This paper is dedicated to the memory of Theodore Holmes Bullock, pioneer in comparative biology, mentor, and friend (1915–2005). We thank Michael V. Bennett and Ziheng Yang for helpful discussions; Maggie Fatay for fish care; Philip Stoddard (Florida International University, Miami) for providing unpublished sequences; and Marianna Grenadier for artwork. This work was funded by National Institutes of Health Grant R01 NS025513 (to H.H.Z. and Y.L.) and National Science Foundation Integrative Graduate Education and Research Traineeship Program DGE-0114387 (to D.J.Z. and D.M.H.).