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Critical Review

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Abstract
Phylogenetic reconstructions provide a means of inferring the branching relationships among members of multigene families that have diversified via successive rounds of gene duplication and divergence. Such reconstructions can illuminate the pathways by which particular expression patterns and protein functions evolved. For example, phylogenetic analyses can reveal cases in which similar expression patterns or functional properties evolved independently in different lineages, either through convergence, parallelism, or evolutionary reversals. The purpose of this article is to provide a robust phylogenetic framework for interpreting experimental data and for generating hypotheses about the functional evolution of globin proteins in chordate animals. To do this, we present a consensus phylogeny of the chordate globin gene superfamily. We document the relative roles of gene duplication and whole-genome duplication in fueling the functional diversification of vertebrate globins, and we unravel patterns of shared ancestry among globin genes from representatives of the three chordate subphyla (Crania, Urochordata, and Cephalochordata). Our results demonstrate the value of integrating phylogenetic analyses with genomic analyses of conserved synteny to infer the duplicative origins and evolutionary histories of globin genes. We also discuss a number of case studies that illustrate the importance of phylogenetic information when making inferences about the evolution of globin gene expression and protein function. Finally, we discuss why the globin gene superfamily presents special challenges for phylogenetic analysis, and we describe methodological approaches that can be used to meet those challenges.

Keywords: cytoglobin, gene family evolution, hemoglobin, myoglobin, neuroglobin

INTRODUCTION
During the last half century, hemoglobin (Hb) and myoglobin (Mb) played starring roles in research efforts to understand relationships between protein structure and function, and in efforts to identify molecular mechanisms of adaptation and pathophysiology. In the past decade, investigations into the structure, function, and evolution of globin proteins have been reinvigorated by the discovery of new members of the globin protein superfamily in humans and other vertebrates. Of these recently characterized globin proteins, neuroglobin (Ngb) and cytoglobin (Cygb) have received the most attention (1–9). These two proteins are highly conserved and are possessed by all jawed vertebrates (gnathostomes) examined to date. In gnathostomes, the physiological functions of Ngb and Cygb are not yet clearly understood, but investigations into their structures, ligand reactivities, biochemical activities, and expression patterns are gradually yielding clues (6–10). In jawless fishes (cyclostomes, represented by lampreys and hagfish), homologs of the Ngb, Mb, and Hb genes appear to have been secondarily lost, and the Cygb homolog evolved a specialized respiratory function in blood-oxygen transport (11).

In addition to Ngb and Cygb, recent surveys of vertebrate genome assemblies and sequence data bases have led to the discovery of several other globin genes with much more restricted phyletic distributions. The Globin X (GbX) gene is the product of a duplication event that predates the divergence between protostomes and deuterostomes, and among vertebrates it has only been found in the genomes of teleost fish and Xenopus (12). The Globin Y (GbY) gene appears to be restricted to gnathostome vertebrates and has thus far only been found in the genomes of Xenopus, anole lizard, bearded dragon lizard, and platypus (13–16). Finally, the Globin E (GbE) gene has thus far only been found in the genome of birds (16–19). Expression patterns of these recently discovered globin genes have been characterized, but the physio-
logical functions of the encoded proteins remain a mystery. Ongoing experimental studies can be expected to yield new and surprising insights in coming years.

To interpret the results of functional experiments, and to generate informed hypotheses about the functional evolution of globin proteins, it is important to have a correct understanding of phylogenetic relationships. Phylogenetic reconstructions allow us to infer the branching relationships among members of a multigene family that have diversified via successive rounds of duplication and divergence. In comparisons among different species, phylogenetic reconstructions provide a means of distinguishing between paralogous genes (which trace their common ancestry to duplication events) and orthologous genes (which trace their common ancestry to speciation events—that is, they descend from a common ancestral gene by phylogenetic splitting at the organismal level). The reconstruction of phylogenetic relationships among homologous members of a multigene family is also essential for understanding the pathways by which various functional properties evolved. For example, it is possible to reconstruct the history of evolutionary change in particular structural and functional features of proteins by “mapping” character states onto a phylogenetic tree that is estimated using independent data. This mode of inference can reveal whether certain physiological functions of modern-day globins represent derived, “repurposed” modifications of distinct ancestral functions. Phylogenetic reconstructions are also essential for identifying cases in which similar features evolved independently in different lineages, either through convergence, parallelism, or evolutionary reversals.

The purpose of this paper is to provide a robust phylogenetic framework for interpreting experimental data and for generating hypotheses about the functional evolution of globin proteins in chordate animals. To do this, we present a consensus phylogeny of the globin gene superfam-

### CONSENSUS PHYLOGENY OF CHORDATE GLOBINS

To reconstruct the phylogeny of chordate globins, we assembled a dataset that included the complete set of globins from representatives of all major vertebrate lineages, in addition to representatives of the two other chordate subphyla: the sea squirt (Ciona intestinalis, a urochordate, (20)) and amphioxus (Branchiostoma floridae, a cephalochordate, (21)). To compile the vertebrate globin dataset, we interro-

clauding teleost fish, amphibians, squamate reptiles, birds, and mammals), and we used bioinformatic tools to annotate the entire globin gene repertoire of each species. In addition, we compiled globin sequences from three representative cartilaginous fish (red stingray, gummy houndshark, and Port Jackson shark) and three representative cyclo-
stomes: Arctic lamprey, sea lamprey (subclass Hyperartia), and hagfish (subclass Myxini). Even though most of the gnathostome species included in this study possess multiple paralogous copies of the α- and β-like globin genes, we only included a representative subset of sequences from each species in our analyses, because monophyly of the α- and β-like globin genes has been well established (22, 23).

Amino acid sequences were aligned using the E-INS-i, GINS- i, and L-INS-i strategies from Mafft version 6.8 (24, 25), and for each alignment, we estimated phylogenetic relationships using maximum likelihood and Bayesian approaches. Maximum likelihood searches were carried out using Treefinder version October 2008 (26) using a mixed model of amino acid substitution, which was identified as the best-fitting model of amino acid substitution with the “propose model” subroutine from Treefinder. Support for the nodes was evaluated with 1,000 bootstrap pseudoreplicates. Bayesian analyses were conducted in MrBayes v.3.1.2 (27) running six simultaneous chains for 10,000,000 gener-

ations, sampling every 1,000 generations, under a mixed model of amino acid substitution and using default priors. Support for the nodes and parameter estimates were derived from a majority rule consensus of the last 2,500 trees. The average standard deviation of split frequencies remained less than 0.01 after the burn-in threshold.

Bayesian and maximum likelihood phylogenies arranged chordate globins into four distinct clades (Figure 1). The first clade contains the complete set of vertebrate-specific globins (Cygb, Cyclostome Hbs, Mb, GhE, the α- and β-Hbs of gnathostomes, and GbY) plus Gbs-7, 10, 11, and 15 from amphioxus. The second clade includes the complete globin repertoire of the sea squirt plus Gbs-1, 2, 5, and 9 from amphioxus, the third clade includes vertebrate GbX plus Gbs-3, 6, 12, 13 and 14 from amphioxus, and the fourth clade includes vertebrate Ngb and its putative ortholog in amphioxus, Gb-4 (21). The correct phylogenetic placement of an additional amphioxus globin, Gb-8, is unclear. Consistent with results of previous studies (11–13, 18, 21), our phylogenetic reconstruction reveals that two of the vertebrate globin paralogs, GbX and Ngb, derive from independent duplication events that occurred before the deuterostome/protostome divergence. The remaining members of the vertebrate globin gene repertoire are all products of vertebrate-specific duplication events (18). Amphioxus possesses an especially diverse repertoire of globin genes, as sequences from this taxon are represented in each of the different globin gene lineages that trace back to the chordate common ancestor. In the absence of gene losses, each clade of orthologous sequences should independently recapitulate the expected organismal phylogeny. For example, a hypothetical clade of orthologous globin genes from representatives of the three chordate subphyla (Craniata, Urochordata, and Cephalochordata) would be expected to
Figure 1. Maximum likelihood phylogram describing relationships among globin genes from representative chordates: 11 jawed vertebrates (Gnathostomata), three jawless fishes (Cyclostomata), the sea squirt [*Ciona intestinalis* (Urochordata)], and amphioxus [*Branchiostoma floridae* (Cephalochordata)]. Numbers above the nodes correspond to maximum likelihood bootstrap support values, and those below the nodes correspond to Bayesian posterior probabilities. The inset tree depicts the organismal phylogeny and the timing of two successive whole-genome duplications (the 1R and 2R duplications) in the stem lineage of vertebrates.
group the sea squirt with vertebrates to the exclusion of amphioxus (28, 29). Contrary to these expectations, the phylogenetic reconstruction shown in Figure 1 indicates that orthologs of GbX, Ngb, and the pro-ortholog of vertebrate-specific globins have all been secondarily lost from the sea squirt genome (18). Likewise, the pro-ortholog of the sea squirt globins was lost in the stem lineage of vertebrates.

The set of vertebrate-specific globin genes fall into four main clades: (i) Cygb and cyclostome Hbs; (ii) Mb + GbE; (iii) the α- and β-chain Hbs of gnathostomes; and (iv) GbY (Figure 1; (11, 18)). As reported previously (11), the “Hb” genes of cyclostomes are clearly orthologous to the Cygb gene of gnathostome vertebrates. The relationships depicted in Figure 1 indicate that progenitors of the four main vertebrate-specific globin lineages were present in the vertebrate common ancestor (11, 18). At face value, the fact that four clades of vertebrate-specific globins are sister to a single clade of amphioxus globins is consistent with the hypothesis that those four clades represent the paralogous products of two successive rounds of whole-genome duplication (WGD) in the stem lineage of vertebrates (the “1R” and “2R” WGDs). If the four main clades of vertebrate-specific globins represent products of the 1R/2R WGDs in the vertebrate common ancestor, then representatives of the four vertebratespecific gene lineages should be embedded in unlinked chromosomal regions that share similar, interdigitated arrangements of paralogous genes (paralogons). The flanking tracts of paralogous duplicates may not contain identical subsets of genes, but the globin-defined paralogons should be united by gene families that trace their duplicative origins to the stem lineage of vertebrates. For example, we would expect that a number of globin-linked genes are members of “4:1” gene families—that is, quartets of paralogs that coduplicated with the globin genes such that each of the four duplicate copies are located on a different globin-defined paralogon. The problem with identifying 4:1 gene families is that only a small subset of gene families would be expected to retain all four of the resultant paralogs following two rounds of WGD, and subsequent gene turnover via smallscale duplications and deletions would further obscure the signal of WGD (30). However, the same globin-defined paralogons that are united by 4:1 gene families would also be united—in various combinations—by 3:1 and 2:1 gene families. Members of such gene families are located on either three or two of the four globin-defined paralogons, respectively (the implication is that the missing members of the expected gene quartet were deleted after the first or second rounds of WGD). Finally, if the four clades of vertebrate-specific globins are products of two rounds of WGD in the vertebrate common ancestor, then the globin-defined paralogons should all derive from a single linkage group of the ancestral chordate protokaryotype (29).

We tested each of the above predictions by examining the genomic map positions of the vertebrate-specific globin genes, by characterizing large-scale patterns in the physical locations of paralogous gene duplicates in the flanking chromosomal regions, and by reconstructing the phylogenetic relationships of the globin-linked genes (18). Results of this analysis revealed that the Cygb gene, the Mb/GbE gene pair, and the Hb/GbY gene pair are each embedded in clearly identifiable paralogons (18, 19). The Hb paralogon is defined by the α-globin gene cluster of amniotes and is defined by the tandemly linked α and β-globin gene clusters in teleost fishes and amphibians. In the human genome, the “Cygb” and “Hb” paralogons correspond to large segments of Chromosome 17 and 16, respectively, and the “Mb” paralogon is partitioned among segments of Chromosomes 7, 12, and 22. We also identified a fourth set of linked genes on human Chromosome 19 that coduplicated with the Cygb, Mb, and Hb paralogons, but the associated globin paralog has been secondarily lost (Figure 2). Synteny comparisons revealed that this segment of Chromosome 19 represents the fourth WGD-derived paralogon—the inference is that it once harbored a globin gene that was coparalogous to the proto-Cygb, Mb, and Hb genes (and it is possible that this globin gene lineage has been retained in the genomes of early branching vertebrate lineages like cartilaginous fish). We henceforth refer to this fourth paralogon as the “globin minus” (Gb”) paralogon. As predicted by the genome duplication model, the 4:1, 3:1, and 2:1 gene families that unite the Cygb, Mb, Hb, and GbY paralogons all trace their duplicative origins to the stem lineage of vertebrates (Figure 3; (18)).

Integrating the phylogenetic reconstructions 5 with synteny comparisons revealed that the GbY gene and the proto-Hb gene represent paralogous products of an ancient tandem gene duplication that occurred prior to the two rounds of WGD in the stem lineage of vertebrates (18). The ancestral linkage arrangement of these two genes is still retained in the genomes of Xenopus, anole lizard, and platypus, as GbY is located at the 3′ end of the α-globin gene cluster in each of these taxa (13-14,16). Similarly, the GbE and Mb genes represent the paralogous products of a tandem gene duplication that occurred in the common ancestor of gnathostome vertebrates, and their ancestral linkage arrangement is still retained in the genomes of all birds examined to date (chicken, turkey, mallard duck, and zebra finch; (18, 19)).

By comparing the complete sequences of vertebrate genomes to those of nonvertebrate chordates like amphioxus, it is possible to reconstruct the protokaryotype of the chordate ancestor (29). As a result of the 1R/2R WGDs, each of the protochromosomes in the chordate ancestor would have been quadruplicated in the stem lineage of vertebrates. This is reflected by the well-documented “tetra-paralogon” structure that is detectable in the genomes of contemporary vertebrates (30). Consistent with the genome duplication hypothesis, the synteny analysis of Hoffmann et al. (18) revealed that the Cygb, Mb, and Hb “paralogons all descend from a single ancestral linkage group in the reconstructed chordate protokaryotype (linkage group 15 of Putnam et al. 29). This provides conclusive evidence that the Cygb, Mb, and Hb gene lineages represent the paralogous products of WGD, not just large-scale segmental duplications (18).
**Figure 2.** Graphical depiction of gene duplicates that are shared between the three globin-defined paralogs (*Cygb*, *Mb*, and *Hb*) and the *Gb*− paralogon in the human genome. There are seven 4:1 gene families that unite the *Gb*− paralogon with the *Cygb*, *Mb*, and *Hb* paralogs, there are seven 3:1 gene families that unite the *Gb*− paralogon with two of the three globin-defined paralogs, and there are four 2:1 gene families that unite the *Gb*− paralogon with a single globin-defined paralogon. The shared paralogs are depicted in colinear arrays for display purposes only, as there is substantial variation in gene order among the four paralogs. For clarity of presentation, genes that are not shared between the *Gb*− paralogon and any of the three globin-defined paralogs are not shown. In the human genome, the *Gb*− paralogon on Chromosome 19 shares multiple gene duplicates with fragments of the *Hb* paralogon on Chromosomes 16 and 7, and it shares multiple gene duplicates with fragments of the *Mb* paralogon on Chromosomes 12 and 22.
The traditional textbook account of the origins of vertebrate globins is that the protoMb and Hb genes were produced by duplication of an ancestral, single-copy globin gene in the vertebrate common ancestor. In the Hb gene lineage, a subsequent tandem gene duplication gave rise to the progenitors of the α and β-globin gene subfamilies. This scenario is correct in broad outline, but we now know that the progenitors of the Cygb/ GbE/ Mb gene lineage and the α/β-Hb gene lineage originated via WGD in the stem lineage of vertebrates (Figure 4; 18, 19).
CONVERGENT EVOLUTION OF GLOBIN GENE EXPRESSION AND PROTEIN FUNCTION

Phylogenetic analyses of the vertebrate α- and β-chain Hb genes have revealed a number of examples of convergent evolution in the developmental timing of gene expression. In all tetrapod vertebrates that have been examined to date, developmentally regulated members of the α- and β-globin gene families direct the synthesis of functionally distinct Hb isoforms in primitive (embryonic) erythrocytes derived from the yolk sac and in definitive (adult) erythrocytes derived from the bone marrow (31–35). In the α-globin gene cluster, the physiological division of labor between early- and late-expressed genes was established in the common ancestor of tetrapod vertebrates and it appears to have been retained in nearly all descendant lineages. The ancestral arrangement of the tetrapod α-globin gene cluster is 5′-α²-α³-3′ (16, 36). In the stem lineage of tetrapods, the α²- and α³-globin genes originated via tandem duplication of an ancestral proto α-globin gene and the α²/α³-globin gene originated subsequently via tandem duplication of the proto α²-globin gene (36). In all tetrapods that have been examined, the α²-globin gene is exclusively expressed in larval/embryonic erythroid cells, and the α³-globin gene is expressed in definitive erythroid cells during later stages of prenatal development and postnatal life. The α³-globin gene does not appear to be expressed in mammalian erythrocytes, but it is expressed in both primitive and definitive erythroid cells in all birds and nonavian reptiles that have been examined to date (34, 35).

In contrast to the ancient functional diversification of the α-like globin genes, the developmental regulation of gene expression in the β-globin gene cluster evolved independently in several different tetrapod lineages (16). For example, in mammals and birds, the β-like globin genes that are expressed during the earliest stages of embryogenesis are not 1:1 orthologs, as they are independently derived from lineage-specific duplications of the same proto β-globin gene (16, 23, 37). Even within mammals, embryonic β-like globin genes appear to have originated independently as the products of lineage-specific duplication events in monotremes (platypus and echidnas) and in the common ancestor of marsupials and placental mammals (37). Likewise, fetally expressed β-like globin genes originated independently in anthropoid primates and in some artiodactyls such as goats and cows. In anthropoid primates (including humans), duplicate copies of the embryonic γ-globin genes have been coopted for fetal expression, whereas in artiodactyls, duplicate copies of the adult β-globin gene were co-opted for fetal expression (23). These observations indicate that genes with similar stage-specific expression patterns do not necessarily descend from a homologous gene copy that was inherited from a common ancestor. Instead, the genes represent the paralogous products of independent, lineage-specific duplication events, and their similar patterns of stage-specific expression are attributable to convergent evolution.

An especially remarkable case of convergence involves the independent evolution of erythroid-specific, oxygen-transport Hbs from different ancestral precursor proteins in gnathostomes and cyclostomes (11). A comprehensive phylogenetic analysis of vertebrate globins revealed that the erythroid Hbs of cyclostomes are orthologous to the Cygb protein of gnathostome vertebrates, a hexacoordinate globin that has no oxygen-transport function. The independent evolution of oxygen-transport Hbs in cyclostomes and gnathostomes represents an example of “cooptive convergence” where paralogous members of the same gene family independently evolve the same specialization of function in different lineages. During their independent acquisition of oxygen-transport functions, the two paralogous globins convergently evolved distinct forms of both homotropic and heterotropic cooperativity from different ancestral precursor proteins that lacked cooperativity. In both cases, multisubunit quaternary structures provided the basis for cooperative oxygenbinding and allosteric regulation, but the underlying structural mechanisms are quite distinct.

It has been suggested that globins in the common ancestor of eukaryotes may have performed functions unrelated to oxygen-transport (31, 38, 39). It is not known whether the progenitor of eukaryotic globins had a “2-on-2” or “3-on-3” tertiary structure, or whether it had pentacoordinate or hexacoordinate heme chemistry. In pentacoordinate globins such as Hb and Mb, the heme iron is coordinated by four nitrogen atoms in the porphyrin ring and the proximal histidine in the F helix (HisF8). In the deoxy state, the sixth coordination site of the ferrous (Fe²⁺) iron atom is accessible to oxygen-binding. In hexa-

Figure 4. Cladogram describing phylogenetic relationships among chordate globins. The products of whole-genome duplications (the 1R and 2R duplications) are indicated in the clade of vertebrate-specific globins.
coordinate globins such as Ngb and Cygb, the distal histidine in the E helix (HisE7) is bound to the sixth coordination position of both Fe\(^{2+}\) and Fe\(^{3+}\) (4). The binding of oxygen or other exogenous ligands therefore requires the displacement of the distal histidine. Given that heme coordination chemistry is such an important determinant of ligand-binding dynamics (and hence, physiological function), it might be supposed that pentacoordinate and hexacoordinate globins represent two highly distinct and ancient phylogenetic lineages. However, this is not the case, as phylogenetic reconstructions have revealed that the alternative heme coordination chemistries have evolved multiple times independently (8, 10, 12). In fact, phylogenetic reconstructions of metazoan globins suggest that hexacoordination actually evolved multiple times from different pentacoordinate ancestral states (10).

**CHALLENGES ASSOCIATED WITH THE PHYLOGENETIC ANALYSIS OF GLOBIN GENE FAMILY EVOLUTION**

Historically, much of the uncertainty about the phylogenetic relationships among vertebrate globins was attributable to inadequate sampling of taxa and/or paralogous gene lineages. For example, it was previously assumed that \(Hb\) and \(Mb\) originated via duplication of an ancestral, single-copy globin gene before the cyclostome/gnathostome divergence, such that each of these two vertebrate lineages inherited orthologous copies of the same “proto-\(Hb\)” gene (22, 23). According to this scenario, the \(Hbs\) of cyclostomes would be sister to the clade of gnathostome \(\alpha\) and \(\beta\)-chain \(Hbs\): [\(Mb\) (cyclostomes \(Hb\), gnathostome \(Hb\)]. Until gnathostome Cygb sequences were included in phylogenetic reconstructions—which revealed the [gnathostome \(Hb\) (cyclostome \(Hb\), gnathostome Cygb)] topology (11)—the phylogenetic affinities between cyclostome and gnathostome \(Hbs\) could not be correctly inferred.

The availability of complete genome assemblies for an exponentially growing list of taxa not only enhances the density of “taxon” sampling in phylogenetic analysis—it also increases the sampling of paralogous gene lineages, because some globin genes have a very limited phyletic distribution. Despite the increasing availability of sequence data, reconstructing the phylogeny of chordate globins remains a challenging proposition. Phylogenetic uncertainty can stem from several sources, including ambiguities in the sequence alignment (40–42) and uncertainty about the best-fitting model of nucleotide or amino acid substitution (43), and of course, there is a vast literature on the computational and statistical challenges associated with exploring the vast universe of all possible phylogenetic tree topologies (44). These inherent problems in phylogeny estimation are exacerbated by the fact that globin proteins are typically less than 200 amino acids in length, so there are a rather limited number of potentially informative sites. Finally, as discussed above, many globin gene lineages trace their origins to extremely ancient duplication events. For example, all vertebrate-specific globins shown in Figure 1 originated via WGDs or subsequent small-scale duplication events that occurred before the split between cartilaginous fish and the common ancestor of teleost fish and tetrapods, over 500 million years ago. Over a broad range of timescales, lineage-specific gene duplications and deletions can also complicate phylogenetic inference (see below). Finally, because of the complex duplicative history of animal globins, the choice of appropriate outgroups is not always straightforward.

There are a number of steps that can be taken to minimize sources of uncertainty in phylogenetic reconstructions. First, it is highly advisable to conduct sensitivity analyses in which the same set of sequences is aligned using different algorithms, and the resultant alignments are then analyzed using different methods of phylogeny estimation under a range of different substitution models. For example, statistically robust inferences about the relationship between cyclostome and gnathostome Hbs required a comprehensive sensitivity analysis to evaluate how phylogeny estimates were affected by the use of different alignment algorithms, the use of different amino acid substitution models, and the use of different outgroup sequences (11). In the study by Hoffmann et al. (11), phylogenetic searches were performed on 10 alternative alignments under three different substitution models, and topology tests were then used to test alternative phylogenetic hypotheses.

In the case of Bayesian analyses, it is necessary to confirm the convergence of Markov chain Monte Carlo simulations to ensure an efficient exploration of parameter space. In the case of maximum likelihood analyses, competing phylogenetic hypotheses can be tested statistically using topology tests such as the Kishino–Hasegawa test (45), the Shimodaira–Hasegawa test (46), the approximately unbiased (AU) test (47), and the SOWT test (48, 49). These approaches have been used to test alternative phylogenetic hypotheses in a number of studies of globin gene family evolution (11, 19, 36, 37). Finally, it is often useful to incorporate additional sources of information to resolve phylogenetic relationships. As illustrated by the diversification of vertebrate-specific globin genes discussed above, tree topologies and patterns of conserved synteny provide reciprocally illuminating sources of information about the duplicative origins of globin genes.

**CONCERTED EVOLUTION AND BIRTH-AND-DEATH EVOLUTION**

In addition to the methodological issues discussed above, there are also certain modes of gene family evolution that can greatly complicate efforts to decipher the correct branching history of gene duplication and species divergence. Within the \(\alpha\) and \(\beta\)-globin subfamilies of vertebrates, tandemly linked genes are often identical or nearly identical in sequence. For example, most mam-
mals possess two to three tandemly linked copies of the adult α-globin gene that have identical coding sequences (50, 51). This pattern is typically attributable to a history of gene conversion—a form of nonreciprocal recombinational exchange between duplicated genes. Recurrent gene conversion results in the gradual homogenization of sequence variation among paralogous members of the same gene family, giving rise to a pattern referred to as “concerted evolution.” Concerted evolution complicates phylogenetic reconstructions because the homogenization of sequence variation between paralogous genes erases phylogenetic history and creates the false appearance of recent common ancestry. Specifically, concerted evolution can create a confusing situation where paralogous genes within the genome of a single species are more similar to one another than they are to their orthologous counterparts in closely related species.

In the α- and β-globin gene clusters of mammals, there are a number of cases, in which the sequence similarity between tandemly duplicated globin genes is only partly attributable to concerted evolution between pre-existing paralogs—instead, it is often attributable to recent ancestry between the products of de novo gene duplications that occurred independently in different lineages (37, 50–54). To distinguish between the effects of concerted evolution and gene turnover (birth-and-death evolution), it is necessary to integrate phylogenetic information from multiple partitions of genomic sequence alignments. Because interparalog gene conversion is largely restricted to the coding regions of globin genes (55–58), orthologous and paralogous relationships can typically be determined by analyzing variation in flanking sequence and/or intronic sequence (37, 50, 52–54, 59).

CONCLUSIONS

The phylogenetic reconstruction presented in Figure 4 provides a framework for interpreting experimental data and for generating hypotheses about the functional evolution of globin proteins in chordates. As more complete genome sequences become available in coming years, it will be possible to trace the ancestry of some chordate globins back to more ancient branch points in the metazoan phylogeny. To reconstruct these ancient pathways of gene family evolution, it will be necessary to use rigorous, model-based methods of phylogeny estimation that account for sources of uncertainty in the sequence alignments, the substitution models, and the choice of outgroup sequences that are used to root the tree. As it becomes possible to probe deeper into the evolutionary history of metazoan globin genes, it may become especially important to complement phylogenetic analyses with comparisons of conserved synteny. Unraveling phylogenetic relationships among the diverse globin genes in protostome taxa—and documenting their affinities with deuterostome globins—may require a combined approach that integrates molecular phylogenetics, alignment-free computational methods, and genomic analyses of conserved synteny.

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