In the past 40 years, comparisons of developmental gene expression and mechanisms of development (evodevo) joined comparative morphology as tools for reconstructing long-extinct ancestral forms. Unfortunately, both approaches typically give congruent answers only with closely related organisms. Chordate nervous systems are good examples. Classical studies alone left open whether the vertebrate brain was a new structure or evolved from the anterior end of an ancestral nerve cord like that of modern amphioxus. Evodevo plus electron microscopy showed that the amphioxus brain has a diencephalic forebrain, small midbrain, hindbrain and spinal cord with parts of the genetic mechanisms for the midbrain/hindbrain boundary, zona limitans intrathalamica and neural crest. Evodevo also showed how extra genes resulting from whole-genome duplications in vertebrates facilitated evolution of new structures like neural crest. Understanding how the chordate central nervous system (CNS) evolved from that of the ancestral deuterostome has been truly challenging. The majority view is that this ancestor had a CNS with a brain that gave rise to the chordate CNS and, with loss of a discrete brain, to one of the two hemichordate nerve cords. The minority view is that this ancestor had no nerve cord; those in chordates and hemichordates evolved independently. New techniques such as phylostratigraphy may help resolve this conundrum.

1. Introduction

To understand the course of evolution, extinct forms at major nodes of the tree of life are typically reconstructed from commonalities shared by extant forms located on adjacent branches of the tree. This works best within phyla where body plans are similar. Problems arise when body plans are very different, such as between phyla and between rapidly evolving organisms within a phylum. In the past 30 years, phylogenetic analyses with large datasets of nuclear genes have revised many phylogenetic relationships that had been based on mitochondrial genes and/or morphology. In particular, fast-evolving groups such as tunicates and nematodes moved from basal positions to higher levels of the tree, and it has been recognized that their comparatively simple body plans are secondarily reduced.

The classical method for reconstructing ancestral forms has been comparative morphology. It has more recently been joined by comparisons of developmental gene expression and the molecular mechanisms of development (evodevo) as well as sophisticated morphological techniques such as serial transmission electron microscopy (TEM) and confocal microscopy of antibody-labelled specimens. The most recent technique is phylostratigraphy, which examines the evolutionary origins of genes that are expressed in particular structures such as the vertebrate brain [1]. This shows when the genetic framework necessary for building a structure first appeared. When all the methods agree, the hypothetical ancestor has the best chance of approximating the real one. Chordate nervous systems are good examples. All chordates (vertebrates, tunicates and cephalochordates) have dorsal hollow nerve cords. Therefore, the ancestral chordate also most probably had one. However, how much of a brain this organism probably had has been controversial. Vertebrates have a large brain, while the nerve cord in cephalochordates (amphioxus) and
tunicates has only a small anterior swelling, the cerebral vesicle or sensory vesicle. Therefore, the question was whether the vertebrate brain was a new structure or had evolved from the anterior end of an ancestral nerve cord like that of modern amphioxus. Several authors thought that the amphioxus cerebral vesicle was equivalent to the entire vertebrate brain [2–7], while Gans & Northcutt [8] argued that the amphioxus cerebral vesicle is homologous only to the vertebrate hindbrain, with the forebrain and midbrain being vertebrate inventions. Others took positions in between these two extremes [9–12]. Answers began to come in the 1990s from two lines of research: comparisons of developmental gene expression (evodevo) and three-dimensional anatomical reconstructions from serial fine sections (TEM). More recently, analyses of genome sequences (phylogenetics) and studies of the mechanisms of development have begun to address how and when in evolution vertebrate-specific structures evolved.

Understanding how the chordate central nervous system (CNS) evolved from that of the ancestral deuterostome (i.e. the ancestor of chordates, hemichordates and echinoderms) has been especially challenging. The problems are first, that the morphology of the hemichordates and echinoderms, which form a clade (Ambulacraria) basal to chordates, differs considerably between them and also differs considerably from that of the chordates. Second, the phylogenetic position of a clade uniting the acoel flatworms and xenoturbellids is very uncertain. Phylogenetic analyses with collections of nuclear genes have united acoels, nemertodermatids and xenoturbellids into a single clade, the Xenocoelomorpha, and placed it as sister group of the ambulacaria [13]. However, the acoels are very fast-evolving and, depending on which genes are used for the analyses, skew the tree; sometimes the Xenocoelomorpha are seen as basal bilateria, and sometimes only the nemertodermatids and acoels are placed basally to bilateria with Xenoturbellida as sister group of the Ambulacaria [14]. In any case, it appears that both acoels and xenoturbellids have lost a number of characters [13]. Given these considerations, it is difficult to predict with any degree of certainty the precise structure of the nervous system of the common ancestor of Ambulacaria and Chordata, let alone that of the basal deuterostome. The present review focuses on the evolution of chordate nervous systems and discusses the pros and cons of theories concerning the evolution of the chordate nervous system from that of an ancestral deuterostome.

2. How did the vertebrate brain evolve from that of an invertebrate chordate ancestor?

Although amphioxus and vertebrates split over 550 Ma, both groups are evolving relatively slowly, with the genomes of species of amphioxus evolving even more slowly than that of the slowest-evolving vertebrate known, the elephant shark [15]. The genome of the Florida amphioxus, *Branchiostoma floridae*, which was the first amphioxus genome to be sequenced, conserves a very high degree of synteny with vertebrate genomes [16]. Comparisons of the *B. floridae* and vertebrate genomes substantiated the idea first proposed by Ohno [17] that vertebrates had undergone two rounds of whole-genome duplication. Extra copies of many duplicate genes were lost, but those for developmental genes and genes coding for signalling proteins were preferentially retained [16]. It has been postulated that these extra genes gave vertebrates the genetic tool-kit to gain a large, complex brain [18]. The lack of such whole-genome duplications in amphioxus plus this slow rate of evolution support the use of amphioxus as a proxy for the ancestral chordate. Also supporting this use are fossils from the Cambrian such as *Haikouella*, which resembles modern amphioxus to a large extent but appears to have paired eyes and a larger brain and has, therefore, been proposed as a sister group of vertebrates [19,20]. Of course, modern amphioxus may well have evolved some new characters and changed some old ones over the millennia, but all available evidence indicates that it has changed relatively little.

**a) What new structures did the vertebrate brain invent?**

Comparisons of developmental gene expression together with three-dimensional reconstructions from serial TEM have shown that the amphioxus brain has homologies of most of the features of the vertebrate brain. These include a hindbrain, diencephalic forebrain with a pineal homologue, and perhaps a small midbrain (tectum), which receives input from the frontal eye [21,22]. Clear evidence for a telencephalon is lacking. Although gene markers of the vertebrate telencephalon such as BFI (FoxG1) are expressed in the anterior tip of the CNS, there is no structure comparable to the olfactory bulbs of the vertebrate telencephalon [21]. As FoxG1 is also expressed in vertebrate diencephalic structures (e.g. the optic stalks), its expression in the amphioxus CNS is not necessarily indicative of a telencephalon. The CNS of amphioxus has no gross anatomical divisions except constriction at the posterior end of the cerebral vesicle; however, as the somites extend to the anterior tip of the animal, they serve as excellent markers of anterior/posterior position. Evidence for a hindbrain comes from expression of *Hox* genes in nested patterns with the anterior limit of *Hox1* at the level of the anterior boundary of somite 2, the *Hox2* limit at the level of somite 3, that for *Hox3* at the level of somite 4, and that for *Hox6* between somites 6 and 7 [23]. In addition, motor neurons, which are located in the vertebrate midbrain and hindbrain, are located at the level of somites 2–6 in the amphioxus nerve cord [24]. They express characteristic motor neuron markers, including the estrogen-related receptor [25]. Evidence that amphioxus has a homologue of the vertebrate diencephalon is strong. Based on fine structure and histology, there is an infundibulum [26,27]. Additional evidence for a diencephalic homologue is the presence of the lamellar body, which has the same fine structure as the pineal in a larval lamprey [28,29]. In addition, consistent with the anterior tip of the amphioxus CNS being diencephalic, in both larvae and adults, the anterior-most part of the CNS includes a photoreceptor or frontal eye, which has been proposed to be homologous to the vertebrate paired eyes [22,29,30].

Evidence from gene expression indicates that amphioxus also has parts of the genetic mechanisms that specify three organizing centres in the vertebrate brain: the anterior neural ridge (ANR) zona limitans intrathalamica (ZLI) and the midbrain/hindbrain boundary (isthmic organizer) (MHB/ISO). Both the anterior tip of the amphioxus CNS and the vertebrate ANR
express Dlx5, FoxG1 (BF1) and Fgf8 [21,31–33]. However, in amphioxus, the domain of Fg8/17/18 extends to the posterior limit of the cerebral vesicle [21,32,33]. In vertebrates, the ZLI is located about midway between the anterior and posterior ends of the diencephalon where a posterior domain of Otx abuts anterior domains of Otx and Fgf [34,35]. Likewise, in amphioxus, a posterior domain of Irx8 abuts an anterior domain of Fgf in amphioxus [40], not in vertebrates. Similarly, in both amphioxus and vertebrates, an anterior domain of Otx meets a posterior domain of Glx. In amphioxus this is at the boundary between the hindbrain and cerebral vesicle and in vertebrates at the MHB. However, while engrailed is expressed at this boundary in amphioxus, it is not in amphioxus. Of the five vertebrate genes that specify neural crest are not similarly expressed between amphioxus and vertebrates [44]. However, the neural plate and the neural plate border are highly conserved between amphioxus and vertebrates [21,32,33]. In vertebrates, the ZLI is located about midway between the anterior and posterior ends of the cerebral vesicle [36]. In both amphioxus and vertebrates, Wnt8 is expressed at or near this boundary as is Fgf [37–39]. However, not all genes are identically expressed at this boundary in vertebrates and amphioxus. For example, engrailed is expressed at this boundary in amphioxus [40], but not in vertebrates. Similarly, in both amphioxus and vertebrates, an anterior domain of Otx meets a posterior domain of Glx. In amphioxus this is at the boundary between the hindbrain and cerebral vesicle and in vertebrates at the MHB. However, while engrailed is expressed at this boundary in vertebrates, it is not in amphioxus [40–42]. It is, therefore, unlikely that the amphioxus MHB and ZLI equivalents are organizers because they lack expression of some genes that are critical for organizer properties. In sum, the data from fine-structural three-dimensional reconstructions and gene expression indicate that the CNS of amphioxus, and, by extension, that of the ancestral vertebrate had a diencephalic forebrain with part of the genetic machinery for the ANR and ZLI, a small midbrain and a hindbrain, with the genetic machinery for positioning the MHB. Thus, the telencephalon is the major brain region that evolved at the base of the vertebrates.

(b) Neural crest

Another structure which the vertebrate brain but has that the amphioxus CNS lacks is neural crest—cells that migrate from the neural plate boundary and give rise to numerous cell types including pigment cells, cells of the adrenal medulla and cartilage and bone [43]. By contrast, in amphioxus, the ectoderm adjacent to the neural plate on either side migrates over it as a sheet, with the leading edge cells displaying lamellipodia (figure 1) [33]. These leading edge cells express Distalless like neural crest cells. In fact, the genes that specify the neural plate and the neural plate border are highly conserved between amphioxus and vertebrates [44]. However, the genes that specify neural crest are not similarly expressed in amphioxus and vertebrates. Notable among them is FoxD3, which is vital for neural crest migration. Of the five vertebrate FoxD genes, only FoxD3 is expressed in neural crest (figure 2). Amphioxus has a single FoxD gene, which is expressed in mesodermal tissues and the anterior neural tube, but not at the edges of the neural plate [47]. Regulatory DNA was identified that directed expression of a reporter gene to all the domains that normally expressed amphioxus FoxD [48]. While this regulatory DNA directed expression to the corresponding domains in the chick, it failed to direct expression to neural crest, demonstrating that after gene duplications in vertebrates, FoxD3 had acquired new regulatory elements [45]. Indeed, the FoxD3 enhancer directing expression to premigratory neural crest [49] has little identity with the amphioxus FoxD3 regulatory region (L. Z. Holland 2015, unpublished data). However, the amino terminal region of the FoxD3 protein also acquired new protein sequences allowing it to induce expression of neural crest genes such as HNK1 [46]. Although this is just one gene, it shows how gene duplication allowed some duplicates to retain old functions while leaving others free to gain new ones. Such acquisition of new gene regulatory elements and new protein sequences has likely occurred also for other duplicate genes during evolution of the vertebrate brain.

It has been argued that ascidian tunicate larvae may have some cells related to neural crest, but this is far from certain. Some migratory cells in the vicinity of the nerve cord were shown to migrate and develop into pigment cells, but these were not migrating from edges of the neural plate [50]. In addition, it was found that ectopic expression of Twist could induce some cells to migrate away from the Ciona neural tube [51]. However, tunicates are evolving rapidly. Their genomes are very reduced with loss of several developmental genes, and the larvae have relatively few cells. Therefore, even though tunicates are the sister group of vertebrates, it is impossible to reconstruct their common ancestor to obtain a clear idea of how many vertebrate features this ancestor had before the whole-genome duplications.

(c) Phylostratigraphy

Phylostratigraphy is a relatively new approach to investigate when particular structures evolved. This method uses comparative genomics to determine when genes expressed in a particular anatomical structure evolved (figure 3) [52]. This is not to say when the structure itself evolved, but only to predict when the genetic framework for a structure such as the brain evolved. For example, an analysis of genes involved in development of sensory structures in vertebrates showed that genes for the eyes, including the lens evolved first, with peaks for the number of new genes for the retina and eye evolving in deuterostomes and for the lens in cephalochordates [53]. By contrast, the peak appearances of new genes for the olfactory system, ear and lateral line as well as that for cranial placodes occur in tunicates while those for neural crest, adenohypophysis and trigeminal placode and ganglion are in vertebrates; however, a minor peak for the adenohypophysis occurs with the chordates [53]. When this type of analysis was applied to the brain regions, genes for the whole brain, forebrain (including diencephalon and telencephalon), midbrain and hindbrain made their peak appearance in amphioxus, although there were minor peaks for all but the midbrain genes at the base of the metazoan and in the vertebrates (figure 3) [1]. However, when the telencephalon was divided into dorsal and ventral regions, a peak for the genes of the ventral telencephalon occurred in amphioxus, but the peak for the dorsal telencephalon was in the vertebrates. There were less pronounced peaks for the ventral telencephalon in agnathans and euteleosts. When the midbrain was subdivided, there was a striking peak for the MHB in amphioxus and a minor peak in agnathans, while the tegmentum had dual peaks in amphioxus and agnathans. In addition, there are peaks of new gene appearance for the MHB and the tegmentum at the base of the Metazoa [1]. These results indicate that most of the genes involved in patterning the vertebrate brain arose in the cephalochordate ancestor. Exceptions are that most of those for the dorsal telencephalon arose in vertebrates, while those for the midbrain and optic tectum arose before evolution of eukaryotes [52]. This sheds doubt on the proposal that the larval ectoderm of direct-developing
nerve cords are not related to chordate nerve cords. Thehemichordate-like, echinoderm-like or something else and chordates may be impossible. It could be chordate-like, hemichordate-like, echinoderm-like or something else (figure 4). It is generally agreed that the adult echinoderm nerve cords are not related to chordate nerve cords. The chief evidence is that during development, the echinoderm nerve cords do not express Hex genes [56]. Whether either of the hemichordate nerve cords is homologous to the chorate nerve cord is controversial [57]. There is no evident brain in enteropneust hemichordates, although the proboscis ectoderm contains many neurons. The dorsal nerve cord does undergo a sort of neurulation in the region of the collar and has most often been proposed as homologous to the chordate nerve cord [58,59]. However, some authors could not decide which of the two nerve cords was homologous to the chordate one [60]. Relevant to this argument is that in neither indirect, nor direct-developing hemichordates does the larval nervous system contribute substantially to the adult nervous system, although in the direct-developing hemichordate, Saccoglossus kowalevskii, ectodermal neurons in the larval proboscis may carry over to the ectoderm of the adult [59,61,62]. Although expression of some nerve cell markers has been studied during hemichordate metamorphosis [59,61,62], a thorough analysis of developmental gene expression in the hemichordate nerve cord has not been done and is sorely needed. One possibility is that a nerve cord in the common ancestor of the Ambulacaria and Chordata was more like that in a modern cephahlochordate than like either of those in a modern enteropneust hemichordate and that it became secondarily reduced in hemichordates. Perhaps as the brain ceased to neurulate, it became spread out in the ectoderm of the proboscis. This would explain expression of genes such as Otx in the forebrain of chordates and in the proboscis ectoderm of hemichordates.

An alternative view is that neither hemichordate nerve cord is homologous to the chordate nerve cord; the nerve cords in the two groups evolved independently [54,63]. These authors (Pani et al. and Lowe et al.) have shown that some of the genes mediating A/P patterning of the larval ectoderm of S. kowalevskii are expressed in similar patterns in the vertebrate CNS. These include Otx, which is expressed in the proboscis ectoderm of S. kowalevskii and in the chordate forebrain. As Otx is not expressed in the ectoderm outside the CNS in chordates, its expression in the hemichordate may indicate an evolutionary relationship between the chordate forebrain and the proboscis ectoderm. However, some other

3. Can scenarios for evolution of the chordate nervous system from that of an ancestral deuterostomes be reconciled?

If reconstructing the common ancestor of tunicates and vertebrates is problematical, reconstructing the common ancestor of Ambulacaria (echinoderms and hemichordates) and chordates may be impossible. It could be chordate-like, hemichordate-like, echinoderm-like or something else (figure 4). It is generally agreed that the adult echinoderm nerve cords are not related to chordate nerve cords. The

Figure 1. Neurulation in amphioxus and vertebrates. Top: at the late gastrula stage both amphioxus and vertebrates have a neural plate with a neural plate border region. Second from top: at the early neurula stage, in amphioxus, the neural plate border region detaches from the edges of the neural plate and moves over it by lamellipodia. By contrast, in vertebrates, the neural plate border region remains attached to the neural plate as it rounds up. Third from top: at the late neurula stage, in amphioxus, the free edges of the neural plate border region fuse in the dorsal midline, and the neural plate begins to round up underneath the dorsal ectoderm. In vertebrates at a comparable stage, the neural tube has completed rounding up. Bottom: In amphioxus, the neural plate rounds up completely and detaches from the ectoderm. In vertebrates, the neural tube detaches from the ectoderm, and the neural plate border region gives rise to neural crest cells that migrate below the ectoderm and give rise to such structures as pigment cells, cells of the adrenal medulla, parts of cranial ganglia.

Figure 2. (a) Phylogenetic relations of the amphioxus FoxD gene and the five vertebrate FoxD genes that arose from whole-genome and gene duplications. (b) AmphFoxD is expressed in the forebrain, somites and notochord. In vertebrates, the ancestral FoxD expression domains have been partitioned among four of the five duplicates, FoxD1, FoxD2, FoxD4 and FoxD5. FoxD3 has acquired a new domain in neural crest. Experimental evidence has shown that FoxD3 acquired both new regulatory elements and a new amino terminal sequence, both of which are essential for its role in neural crest [45,46].
Figure 3. Summary of a phylostratographic analysis of the zebrafish (Danio rerio) CNS. The phylostrata are at the left. The vertebrate section is shaded grey. The size of the circles is proportional to the number of genes expressed in a given region of the vertebrate brain that first appeared in that phylostratum. Phylostratum 8 (ps8) includes echinoderms and hemichordates. Phylostratum 9 includes cephalochordates (amphioxus), while phylostratum 10 includes tunicates. The largest number of genes expressed in the zebrafish brain first appeared in cephalochordates. Adapted from [1].

genes expressed in the larval S. kowalevskii ectoderm, such as Hox genes, are expressed in both the CNS and in the ectoderm generally in chordates, making them poor indicators of homology between the chordate nerve cord and larval ectoderm in hemichordates. In addition, the domains of genes such as Fexp and Irx and Otx and Gbx do not abut in the hemichordate ectoderm as they do in the chordate nerve cord. Even more complicating is that the genes expressed in medio-lateral patterns in the vertebrate and amphioxus CNS are not similarly expressed in the S. kowalevskii larval ectoderm. Moreover, in amphioxus and vertebrates, opposition between posterior ventral bone morphogenic protein (BMP) and dorsal anterior nodal/vg1 signalling segregates the neuroectoderm from the remainder of the ectoderm, but this is apparently not the case in S. kowalevskii. These considerations make it all the more important for a thorough study of gene expression in the hemichordate nerve cords.

A third view is that the chordate nerve cord evolved from the ciliated bands of an auricularia-larval like adult similar to larvae of holothurians. This idea was proposed by Garstang [64] and modified by Romer [65]. Although Garstang later recanted [66], it is still current [67]. In this scheme, the common ancestor of chordates, hemichordates and echinoderms had an adoral ciliary band as well as one that extended around the mouth and anus. There was a nerve ring underlying the circumoral ciliated band. This ring evolved into the circumoral nerve ring of echinoderms, while in enteropneusts, the posterior part of the ciliated band evolved into the collar nerve cord. In chordates, the circumoral nerve ring moved dorsally to become the nerve cord. These ideas, however, do not seem viable in the light of the studies showing that except for neurons in the proboscis, little, if any, of the larval nervous system carries over into the adult in indirect developing ambulacarians [59,62]. In sum, while it is unlikely that the chordate nerve cord evolved from the larval nervous system of a hemichordate-like ancestral deuterostome, other schemes for evolving the chordate nerve cord from either the dorsal nerve cord or the ventral nerve cord are more reasonable. While the scheme for evolving the chordate nerve cord from the hemichordate ectoderm is less likely, it cannot at present be ruled out.

4. Conclusion

Evodevo studies plus modern microscopy methods have been highly successful tools for addressing the question of how the vertebrate brain evolved from the brain of an invertebrate chordate ancestor. This ancestor probably had a nerve cord with a hindbrain, diencephalic forebrain and perhaps a small midbrain. While it had the genetic scaffolds for the major organizing centres of the vertebrate brain—the ANR, ZLI and MHB—vertebrates added a number of genes to these scaffolds. The increased complexity of the gene networks in these regions of the vertebrate brain is probably correlated with the acquisition of their organizer properties. The increased complexity of the gene network operating at the edges of the chordate neural plate is due at least in part to the retention of duplicate genes deriving from the two rounds of whole-genome duplication at the base of the vertebrates. An example is FoxD3, one of the five vertebrate duplicates of a single ancestral chordate FoxD gene. Experiments showed that vertebrate FoxD3 acquired both new regulatory elements and new protein sequences allowing it to be expressed in neural crest and to induce expression of other neural crest genes. This phenomenon may explain why many duplicate genes for transcription factors and signalling pathways were retained in vertebrates and how they facilitated the evolution of new structures.
It has been far more problematic to understand where the chordate nerve cord came from. The body plans of hemichordates and echinoderms (Ambulacraria) differ both from one another and from those of vertebrates. Schemes for deriving the chordate nerve cord from the ciliated bands of larval ambulacrarians do not seem credible as their adult nervous systems develop largely independently of their larval ones. Similarly, scenarios deriving the chordate nerve cord from an adult echinoderm nerve cord are also not viable as genes are expressed very differently in nerve cords from the two groups. Opinions of the relationship between the chordate and the two hemichordate nerve cords are mixed, ranging from no relationship at all, chordate and hemichordate nerve cords evolving independently, to the collar nerve cord being homologous to the chordate nerve cord or either of the two hemichordate nerve cords being homologous to the chordate nerve cord. Without intermediate forms, it will be difficult or impossible to decide among these scenarios. The goldilocks principle—that to infer homologies, organisms must be similar, but not identical—continues to hold for evolution of the chordate nervous system.

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