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## Hemichordate genomes and deuterostome origins

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### Abstract

Acorn worms, also known as enteropneust (literally, ‘gut-breathing’) hemichordates, are marine invertebrates that share features with echinoderms and chordates. Together, these three phyla comprise the deuterostomes. Here we report the draft genome sequences of two acorn worms, *Saccoglossus kowalevskii* and *Ptychodera flava*. By comparing them with diverse bilaterian genomes, we identify shared traits that were probably inherited from the last common deuterostome ancestor, and then explore evolutionary trajectories leading from this ancestor to hemichordates, echinoderms and chordates. The hemichordate genomes exhibit extensive conserved synteny with amphioxus and other bilaterians, and deeply conserved non-coding sequences that are candidates for conserved gene-regulatory elements. Notably, hemichordates possess a deuterostome-specific genomic cluster of four ordered transcription factor genes, the expression of which is associated with the development of pharyngeal ‘gill’ slits, the foremost morphological innovation of early deuterostomes, and is probably central to their filter-feeding lifestyle. Comparative analysis reveals numerous deuterostome-specific gene novelties, including genes found in deuterostomes and marine microbes, but not other animals. The putative functions

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**Online Content** Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

Supplementary Information is available in the online version of the paper.

**Author Information** Sequencing data have been deposited in NCBI BioProject under accession number PRJNA12887 (*Saccoglossus kowalevskii*) and DDBJ under accession number PRJDB3182 (*Ptychodera flava*). Readers are welcome to comment on the online version of the paper.

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of these genes can be linked to physiological, metabolic and developmental specializations of the filter-feeding ancestor.

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The prominent pharyngeal gill slits, rigid stomochord, and midline nerve cords of acorn worms led 19th century zoologists to designate them as ‘hemichordates’ and group them with vertebrates and other chordates<sup>1–4</sup>, but their early embryos and larvae also linked them to echinoderms<sup>5,6</sup>. Current molecular phylogenies strongly support the affinities of hemichordates and echinoderms as sister phyla, together called ambulacrarians<sup>7</sup>, and unite ambulacrarians and chordates within the deuterostomes (see glossary in Supplementary Note 1). Of all the shared derived morphological characters proposed between hemichordates and chordates, the pharyngeal gill slits have emerged with unambiguous morphological and molecular support, notably the shared expression of the *pax1/9* gene<sup>8–10</sup>. These structures were ancestral deuterostome characters elaborated upon the bilaterian ancestral body plan, but the gill slits were subsequently lost in extant echinoderms and amniotes<sup>11</sup>. Since extant invertebrate deuterostomes use this apparatus for efficient suspension and/or deposit feeding, the early Cambrian or Precambrian deuterostome ancestor probably also shared this lifestyle. This perspective on the last common deuterostome ancestor informs our understanding of the subsequent evolution of hemichordates, echinoderms and chordates<sup>10,12–16</sup>.

Hemichordates share bilateral symmetry, gill slits, soft bodies and early axial patterning with chordates, making them key comparators for inferring the ancestral genomic features of deuterostomes. To this end, we sequenced and analysed the genomes of acorn worms belonging to the two main lineages of enteropneust hemichordates (Supplementary Note 1): *Saccoglossus kowalevskii* (Harrimaniidae; Atlantic, North America, Fig. 1a) and *Ptychodera flava* (Ptychoderidae; Pacific, pantropical, Fig. 1b). Both have characteristic three-part bodies comprising proboscis, collar and trunk, the last with tens to hundreds of pairs of gill slits. While *S. kowalevskii* develops directly to a juvenile worm with these traits within days (Fig. 1c, e), *P. flava* develops indirectly through a feeding larva that metamorphoses to a juvenile worm after months in the plankton (Fig. 1d, e). Our analyses begin to integrate macroscopic information about morphology, organismal physiology, and descriptive embryology of these deuterostomes with genomic information about gene homologies, gene arrangements, gene novelties and non-coding elements.

## Genomes

We sequenced the two acorn worm genomes by random shotgun methods with a variety of read types (Methods; Supplementary Note 2), each starting from sperm from a single outbred diploid individual. The haploid lengths of the two genomes are both about 1 Gbp (Extended Data Fig. 1), but differ in nucleotide heterozygosity. Both acorn worm genomes were annotated using extensive transcriptome data as well as standard homology-based and *de novo* methods (Supplementary Note 3). Counting gene models with at least one detectable orthologue in another sequenced metazoan species, we find that *Ptychodera* and *Saccoglossus* encode at least 18,556 and 19,270 genes, respectively (Methods). Additional *de novo* gene predictions include divergent and/or novel genes (Extended Data Fig. 1). Despite the ancient divergence of the *Saccoglossus* and *Ptychodera* lineages (more than 370

million years ago, see below) and their different modes of development, the two acorn worm genomes have similar bulk gene content, as discussed later (Extended Data Fig. 2 and Supplementary Note 4), and similar repetitive landscapes (Supplementary Note 5).

## Deuterostome phylogeny

Deuterostome relationships were originally inferred from developmental and morphological characters<sup>2,5,17</sup> and these hypotheses were later tested and refined with molecular data<sup>6,7</sup>. Aspects of deuterostome phylogeny continue to be controversial, however, notably the position of the sessile pterobranchs among hemichordates, and the surprising association of *Xenoturbella*<sup>18</sup> and acoelomorph flatworms with ambulacrarians<sup>19</sup> proposed by some studies. We explored these issues using genome-wide analyses of the newly sequenced hemichordate genomes augmented with extensive new RNAseq from five echinoderms, three additional hemichordates (including a rhabdopleurid pterobranch) and two acoels (Fig. 2, Extended Data Fig. 3, Methods and Supplementary Note 6). We recovered the monophyly of hemichordates, echinoderms, ambulacrarians and deuterostomes, using not only amino acid characters but also presence–absence characters for introns and coding indels (Supplementary Note 4). Our analyses also placed pterobranch hemichordates as the sister-group to enteropneusts<sup>7</sup> rather than within them<sup>12</sup>. These phylogenetic analyses imply that genomic traits shared by chordates and ambulacrarians can be attributed to the last common deuterostome ancestor (see below). Using a relaxed molecular clock, we estimate a Cambrian origin of hemichordates (Methods, Extended Data Fig. 3 and Supplementary Note 6).

We also performed several analyses to assess the controversial relationships between *Xenoturbella*, acoelomorphs and deuterostomes (Supplementary Note 6). With conventional site-homogeneous models, acoels remain outside deuterostomes<sup>20–23</sup> (Fig. 2, Supplementary Figs 6.1 and 6.2). Alternative models<sup>24</sup>, however, show equivocal branching of acoels depending on the inclusion of the current sparse data for *Xenoturbella* (Supplementary Note 6). Notably, without *Xenoturbella*, acoels are positioned as a bilaterian sister group (Supplementary Fig. 6.3)<sup>20–23</sup>. Although we cannot rule out a deuterostome placement for *Xenoturbella*, our analyses generally do not support a grouping of acoels with deuterostomes<sup>19</sup>.

## The gene set of the deuterostome ancestor

By comparative analysis, we identified 8,716 families of homologous genes whose distributions in sequenced extant genomes imply their presence in the deuterostome ancestor (Methods; Supplementary Note 4). Owing to gene duplication and other processes the descendants of these ancestral genes account for ~14,000 genes in extant deuterostome genomes including human (Supplementary Table 4.1.2). The distributions of gene functions, domain compositions, and gene family sizes of hemichordates resemble those of amphioxus, sea urchin, and sequenced lophotrochozoans more than those of ecdysozoans; vertebrates also form a distinct group (Extended Data Fig. 2, Supplementary Note 4 and Supplementary Fig. 4.2).

Exon–intron structures of genes are generally well conserved among hemichordates, chordates, and many non-deuterostome metazoans, allowing us to infer 2,061 ancestral deuterostome splice sites (Supplementary Note 4). Among orthologous bilaterian genes we found 23 introns and 4 coding sequence indels present only in deuterostomes (shared between at least one ambulacrarian and chordate), suggesting that these shared derived characters may be useful to diagnose clade membership of new candidate organisms (Supplementary Note 4).

Based on whole-genome alignments, we identified 6,533 conserved non-coding elements (CNE) longer than 50 bp that are found in all of the five deuterostomes *Saccoglossus*, *Ptychodera*, amphioxus, sea urchin, and human (Methods; Supplementary Note 8). The identified CNEs overlap extensively with human long non-coding RNAs (3,611 CNE loci; 55%, Fisher's exact test  $P$  value  $< 2.2 \times 10^{-16}$ ). Those alignments usually do not exceed 250 bp (as has been reported among vertebrates<sup>25</sup>) and occur in clusters (Supplementary Note 8). Among these conserved sequences is a previously identified vertebrate brain and neural tube specific enhancer, located close to the *sox14/21* orthologue in all five species<sup>26</sup>.

## Conserved gene linkage

Ancient gene linkages ('macro-synteny'<sup>27</sup>) are often preserved in extant bilaterian genomes<sup>27,28</sup>. Comparative analysis revealed 17 ancestral linkage groups across chordates, including amphioxus and *Ciona*<sup>27</sup>. While the contiguity of the draft of the sea urchin genome assembly<sup>29</sup> is too limited to determine whether it shares this chromosome-scale organization, we find that the *Saccoglossus* genome clearly shares these chordate-defined linkage groups (Fig. 3a and Supplementary Note 7), implying that these chromosome-scale linkages were also present in the ancestral deuterostome.

On a more local scale, we find hundreds of tightly linked conserved gene clusters of three or more genes ('micro-synteny'; Methods; Supplementary Note 7) including *Hox*<sup>30</sup> and *ParaHox*<sup>31</sup> clusters in both acorn worms (Extended Data Fig. 4), as also found in echinoderms<sup>32,33</sup>. *Saccoglossus* and amphioxus share more micro-syntenic linkages with each other than either does with sea urchin, vertebrates, or available protostome genomes (Methods, Fig. 3b and Extended Data Figs 5 and 6). Conservation of micro-syntenic linkages can occur due to low rates of genomic rearrangement or, more interestingly, as a result of selection to retain linkages between genes and their regulatory elements located in neighbouring genes<sup>28</sup>.

## A deuterostome pharyngeal gene cluster

One conserved deuterostome-specific micro-syntenic cluster with functional implications for deuterostome biology is a cluster of genes expressed in the pharyngeal slits and surrounding pharyngeal endoderm (Fig. 4; Supplementary Note 9). This six-gene cluster contains four transcription factor genes in the order *nkx2.1*, *nkx2.2*, *pax1/9* and *foxA*, along with two non-transcription-factor genes *slc25A21* and *mipoll1*, whose introns harbour regulatory elements for *pax1/9* and *foxA*, respectively<sup>34–36</sup>. The cluster was first found conserved across vertebrates including humans (see chromosome 14; 1.1 Mb length from *nkx2.1* to *foxA1*)<sup>34,37</sup>. In *S. kowalevskii*, it is intact with the same gene order as in vertebrates (0.5 Mb

length from *nkx2.1* to *foxA*), implying that it was present in the deuterostome and ambulacrarian ancestors. The full ordered gene cluster also exists on a single scaffold in the crown-of-thorns sea star *Acanthaster planci*. Since these genes are not clustered in available protostome genomes, there is no evidence for deeper bilaterian ancestry. Two non-coding elements that are conserved across vertebrates and amphioxus<sup>38</sup> are found in the hemichordate and *A. planci* clusters at similar locations (A2 and A4, in Fig. 4a).

The *pax1/9* gene, at the centre of the cluster, is expressed in the pharyngeal endodermal primordium of the gill slit in hemichordates, tunicates, amphioxus, fish, and amphibians<sup>8,9</sup>, and in the branchial pouch endoderm of amniotes (which do not complete the last steps of gill slit formation), as well as other locations in vertebrates. The *nkx2.1* (thyroid transcription factor 1) gene is also expressed in the hemichordate pharyngeal endoderm in a band passing through the gill slit, but not localized to a thyroid-like organ<sup>39</sup>. Here we also examined the expression of *nkx2.2* and *foxA* in *S. kowalevskii*. We find that *nkx2.2*, which is expressed in the ventral hindbrain in vertebrates, is expressed in pharyngeal ventral endoderm in *S. kowalevskii*, close to the gill slit (Fig. 4b), and that *foxA* is expressed throughout endoderm but repressed in the gill slit region (Fig. 4b). The co-expression of this ordered cluster of the four transcription factors during pharyngeal development strongly supports the functional importance of their genomic clustering.

The presence of this cluster in the crown-of-thorns sea star, an echinoderm that lacks gill pores, and in amniote vertebrates that lack gill slits, suggests that the cluster's ancestral role was in pharyngeal apparatus patterning as a whole, of which overt slits (perforations of apposed endoderm and ectoderm) were but one part, and the cluster is retained in these cases because of its continuing contribution to pharynx development. Genomic regions of the pharyngeal cluster have been implicated in long-range promoter–enhancer interactions, supporting the regulatory importance of this gene linkage (see Supplementary Note 9)<sup>40</sup>. Alternatively, genome rearrangement in these lineages may be too slow to disrupt the cluster even without functional constraint. Here we propose that the clustering of the four ordered transcription factors, and their bystander genes, on the deuterostome stem served a regulatory role in the evolution of the pharyngeal apparatus, the foremost morphological innovation of deuterostomes.

## Deuterostome novelties

We found > 30 deuterostome genes with sequences that differ markedly from those of other metazoans, related to functional innovation in deuterostomes. Some plausibly arose from accelerated sequence change on the deuterostome stem from distant but identifiable bilaterian homologues, others represent new protein domain combinations in deuterostomes, while others lack identifiable sequence and domain homologues in other animals. In the latter group, we found over a dozen deuterostome genes that have readily identified relatives in marine microbes, often cyanobacteria or eukaryotic micro-algae, but are not known in other metazoans (Extended Data Table 1 and Extended Data Fig. 7; Supplementary Notes 10.4 and 10.5). Such genes include two of the novel deuterostome sequences associated with sialic acid metabolism (found in many microbes<sup>41</sup>, see below), enzymes that modify proteins (for example, protein arginine deiminase) and RNA (for example, FATSO

methyladenosine demethylase) as well as others that provide specialized reactions of secondary metabolism (Extended Data Table 1 and Extended Data Fig. 7; Supplementary Note 10.5). Possible explanations for the unusual phylogenetic distribution of these genes include horizontal transfer on the deuterostome stem from early marine microbes (which were plausibly commensals, pathogens, or food sources of stem deuterostomes), or convergent gene loss and/or extensive sequence divergence along five or more opisthokont lineages (Supplementary Note 10.2).

Regardless of their mechanism of origination, the various deuterostome novelties and gene family expansions of sialic acid metabolism are noteworthy. Deuterostomes are unique among metazoans in their high level and diverse linkage of addition of sialic acid (also known as neuraminic acid), a nine carbon negatively charged sugar, to the terminal sugars of glycoproteins, mucins and glycolipids<sup>42</sup>. We find expanded families of enzymes for several of these reactions in hemichordates (Fig. 5a and Extended Data Table 1). Based on the presence/absence of relevant enzymes we infer that 5 of the 11 steps of the pathways of sialic acid formation, addition to termini, and removal are not found in protostomes or other metazoans, and are deuterostome novelties (Fig. 5a and Supplementary Note 10), whereas the other steps use enzymes similar to those of the more limited pathway of some protostomes (for example, insects such as *Drosophila*)<sup>43</sup>.

The importance of glycoproteins for muco-ciliary feeding and other hemichordate activities is further supported by novel and expanded families of genes encoding the polypeptide backbones of glycoproteins, those with von Willebrand type-D and/or cysteine-rich domains (PTHR11339 classifier), including mucins, present in hemichordates and amphioxus as large tandemly duplicated clusters (with varied expression patterns as shown in Extended Data Fig. 8), but not in sea urchin, which has a different mode of feeding (Supplementary Note 10). As in amphioxus, the pharynx of *Saccoglossus* is heavily ciliated<sup>44,45</sup>, and cells of the pharyngeal walls in hemichordates and the ventral endostyle in amphioxus secrete abundant mucins and glycoproteins<sup>46</sup>. Similarly, in the deuterostome ancestor these glycoproteins probably enhanced the muco-ciliary filter-feeding capture of food particles from the microbe-rich marine environment and protected its inner and outer tissue surfaces.

## Novelty in the TGF $\beta$ signalling pathway

The signalling ligands Lefty (a Nodal antagonist) and Univin/Vg1/GDF1<sup>47</sup> (a Nodal agonist) are deuterostome innovations that modulate Nodal signalling during the major developmental events of endomesoderm induction and axial patterning in vertebrates, axial patterning in hemichordates and echinoderms, and left–right patterning in all deuterostomes<sup>48</sup> (see Fig. 5b–d and Extended Data Fig. 9a, b). *Univin* is tightly linked to the related bilaterian *bmp2/4* in the sea urchin genome<sup>49</sup> and also, we now report, in hemichordates and amphioxus, supporting its origin by tandem duplication and divergence from an ancestral *bmp2/4*-type gene, as suggested previously<sup>49</sup>.

TGF $\beta$  2 signalling (TGF $\beta$  1, 2 and 3 in vertebrates) is a deuterostome innovation that controls cell growth, proliferation, differentiation and apoptosis at later developmental stages. Accompanying the novel TGF $\beta$  2 ligand, the type II receptor has a novel ectodomain.

The extracellular matrix protein thrombospondin 1, which activates TGF $\beta$  2 in vertebrates, contains a deuterostome-unique combination of domains including three thrombospondin type 1 (TSP1) domains that bind the TGF $\beta$  2 pro-domain region. While these signalling novelties have clear sequence similarity to pan-bilaterian components, they form long stem branch clades on the phylogenetic trees, indicating extensive sequence divergence on the deuterostome stem (Supplementary Note 10). Together, these innovations appear to contribute to the increased amount and complex patterning of Smad2/3-mediated signalling in deuterostomes compared with protostomes and other metazoans.

## Conclusion

The two acorn worms whose genomes are described here represent the two main enteropneust lineages, separated by at least 370 million years and differing in their developmental modes. These analyses reveal (1) extensive conserved macro-synteny among deuterostomes; (2) a widely conserved deuterostome-specific cluster of six ordered genes, including four transcription factor genes that are expressed during the development of pharyngeal gill slits and the branchial apparatus, the most prominent morphological innovation of the deuterostome ancestor; and (3) numerous gene novelties shared among deuterostomes, many expanded into large families, with putative protein functions that imply physiological, metabolic and developmental specializations of the filter-feeding deuterostome ancestor. Some of these genes lack identifiable orthologues in other metazoans but do resemble microbial sequences and domain types. In addition to their contributions towards defining the deuterostome ancestor and illuminating chordate origins, the two genomes should inform hypotheses of larval evolution by providing a basis for future comparisons of direct-developing and indirect-developing acorn worms, which achieve remarkably similar adult forms by distinct embryological routes (Fig. 1).

## METHODS

No statistical methods were used to predetermine sample size. The experiments were not randomized and the investigators were not blinded to allocation during experiments and outcome assessment.

### Sequencing

Sperm DNA from adult males was extracted for sequencing as described in Supplementary Note 2. A single male was used for each species to minimize the impact of heterozygosity on assembly. For *Saccoglossus*, approximately eightfold redundant random shotgun coverage (totalling 8.1 Gb) was obtained with Sanger dideoxy sequencing at the Baylor College of Medicine Genome Center, including 34,279 BAC ends and 459,052 fosmid ends. For *Ptychodera*, 1.3 Gb in Sanger shotgun sequences, 15.3 Gb in Roche 454 pyrosequence reads, and 52-Gb paired-end sequences with Illumina MiSeq, along with mate-pairs, were generated at the Okinawa Institute of Science and Technology Graduate University. More sequencing details are available in Supplementary Note 2.



## Genome assemblies

We assembled the *Saccoglossus* genome with Arachne<sup>50</sup>, combined with BAC/fosmid pair information to produce the final assembly. This *Saccoglossus* assembly includes 7,282 total scaffold sequences spanning a total length of 758 Mb. The relatively modest nucleotide heterozygosity (0.5%) of *S. kowalevskii*, coupled with longer read lengths, enabled assembly of a single composite reference sequence. Half of the assembly is in scaffolds longer than 552 kb (the N50 scaffold length), and 82% of the assembled sequence is found in 1,602 scaffolds longer than 100 kb. For *Ptychodera* we used the Platanus<sup>51</sup> assembler. The resulting total scaffold length was 1,229 Mb, with half the assembly in scaffolds longer than 196 kb (N50 scaffold length). *P. flava* exhibited a notably higher heterozygosity (1.3% single nucleotide heterozygosity with frequent indels) than *S. kowalevskii*, presumably related to its pelagic dispersal and larger effective population size<sup>52</sup>. We therefore initially produced stringent separate assemblies of the two divergent haplotypes, and found that many scaffolds had a closely related second scaffold with ~94% BLASTN identity (over longer stretches, including indels). To avoid reporting both haplotypes at these loci, scaffolds with less than 6% divergence over at least 75% of their length were merged into a single haploid reference for comparative analysis. To further classify regions with 'double' depth and single haplotype regions we implemented a Hidden Markov Model classifier. We find that at least 63% of the initial Platanus assembly constitutes merged haplotypes. The inferred SNP rate for those regions is 1.3%, while for the remaining haplotype regions it is below 0.1%. Further details of assemblies are described in Supplementary Note 2.

## Gene predictions

Transcriptome data for both species were used, along with homology-guided and *ab initio* methods, to predict protein-coding genes (Supplementary Note 3). For *Saccoglossus*, 8.6 million RNAseq reads were generated from 7 adult tissues and 15 developmental stages using Roche 454 sequencing, along with previously deposited ESTs in GenBank. For *Ptychodera*, extensive EST data from egg, blastulae, gastrulae, larvae, juveniles, adult proboscis, stomochord, and gills defining 34,159 cDNA clones<sup>53</sup>, and 879,000 Roche/454 RNAseq reads from a mixed library of developmental stages<sup>54</sup> were used. The *Saccoglossus* genome was annotated using JGI gene prediction pipeline<sup>55</sup>, while Augustus<sup>56</sup> was used to produce gene models for *Ptychodera*. We find a total of 34,239 gene predictions for *Saccoglossus* (68% with transcript evidence) and 34,687 for *Ptychodera* (43% with transcript evidence), although these are overestimates of the true gene number due to fragmented gene predictions, mis-annotated repetitive sequences, and spurious predictions. As described in the main text, 18–19,000 gene models in each species have known annotations and/or orthologues in other species.

## Gene family analysis

Gene family clustering was done using a progressive (leaf to root) BLASTP-based clustering algorithm, where at a given phylogenetic node the gene families are constructed taking into account protein similarities among ingroups and outgroups<sup>57</sup>. For the inference of deuterostome gene families we use the bilaterian node of the clustering. To call gene families present in the deuterostome ancestor, we required (1) at least two ambulacrarian

orthologues out of the three available ambulacrarian genomes and at least two chordate orthologues, or (2) at least two deuterostomes (chordates and/or ambulacrarians) and two outgroups in the bilaterian level clusters.

### Transposable elements

Repetitive sequences were identified using RepeatScout<sup>58</sup>, followed by manual curation and annotation using both a Repbase release (version 20140131)<sup>59</sup> and BLASTX-based search against a custom collection of transposons, using a previously described repeat identification and annotation pipeline<sup>57</sup> (Supplementary Note 5). The assemblies were then masked with RepeatMasker version open-4.0.5<sup>60</sup>. The repetitive complements of the two hemichordate genomes are summarized in Supplementary Table 5.1.

### Phylogenetic analysis

Phylogenetic analyses were done using metazoan-level gene family clusters based on whole-genome sequences (Supplementary Note 4), selecting a single orthologue per genome with the best cumulative BLASTP to other species, and best reciprocal BLASTP hits to species with transcriptome-only information (Supplementary Note 6). Single gene alignments were built using Muscle<sup>61</sup> and filtered using Trimal<sup>62</sup> for each orthologue, and were concatenated, yielding a supermatrix of 506,428 positions with 34.9% missing data. This super-matrix was analysed with ExaML assuming a site-homogenous LG+ $\Gamma_4$  model partitioned for each gene<sup>63</sup>. A slow-fast analysis was conducted to stratify marker genes based on the length of the branch leading to acoels in individual trees. A subset of the slowest 10% of genes was analysed with the site-heterogenous CAT+ GTR+ $\Gamma_4$  model using Phylobayes<sup>24</sup>. Molecular dating was carried out using Phylobayes<sup>24</sup> using the log-normal relaxed clock model and the calibrations described in Supplementary Table 6.2.

### Synteny analysis

Macro- and micro-syntenic linkages were calculated as described in Supplementary Note 7. For Fig. 3a, we merged the amphioxus scaffolds into 17 pre-defined scaffold groups as suggested in ref. 27. These 17 merged scaffold groups represent the 17 ancestral linkage groups (ALGs) shared in chordates. Then we calculated the orthologous gene groups shared by each amphioxus *ALG-Saccoglossus* scaffold pair and generated the dot plot as described in Supplementary Note 7. For micro-synteny we required at least three genes (separated by a maximum of ten genes) to be present in pairwise comparisons. Under random reshuffling of the genome, this yields 10% false positives in pairwise genome comparisons, that is, we observe approximately one-tenth as many micro-syntenic blocks between the two genomes when gene orders are shuffled. This false-positive rate, however, falls to 1% when considering more than two species. For our inference of deuterostome ancestral and novel synteny we therefore focus on blocks present in at least three species (and both ingroup representatives, that is, ambulacrarians and chordates). This yields 698 blocks that can be traced back to the deuterostome ancestor, including 71 blocks found exclusively in deuterostome species (shared among ambulacrarians and chordates), including the pharyngeal cluster discussed in Fig. 4.

## Whole-genome alignment

Whole-genome alignments were conducted with MEGABLAST<sup>64</sup> using parameters previously reported<sup>65</sup>. We assessed the distribution of the resulting 12,722 aligned loci across known gene annotations in ENSEMBL<sup>66</sup>, previously identified conserved pan-vertebrate elements<sup>65</sup>, as well as known enhancers in human according to LBL database<sup>67</sup>.

## Gene novelties

Deuterostome gene novelties were assessed initially through bilaterian gene clusters (Supplementary Note 10) by requiring at least two species on both ambulacrarian and chordate side to be present. The novelties were further automatically subdivided into four categories: G1 (gain type I), with no BLASTP hit outside of deuterostomes; G2 (gain type II), with a novel PFAM domain present only in deuterostomes; G3 (gain type III) having a novel PFAM combination unique to deuterostomes; and G4 (gain type IV), those that do not fall under any of the G1-3 categories and define novelties due to acceleration in the substitution rate on the deuterostome stem. To confirm the novel nature, especially for G4 novelties, we have constructed phylogenies for the members and non-deuterostome BLASTP hits (up to an  $e$ -value of  $1 \times 10^{-20}$ ) using MAFFT-alignment-based FastTree calculations. The trees were assessed for the accelerated rate of evolution at the deuterostome stem (Supplementary Fig. 9.1.1). The final result is provided in the Supplementary Information.

## Curation of candidates for horizontal gene transfer on the deuterostome stem

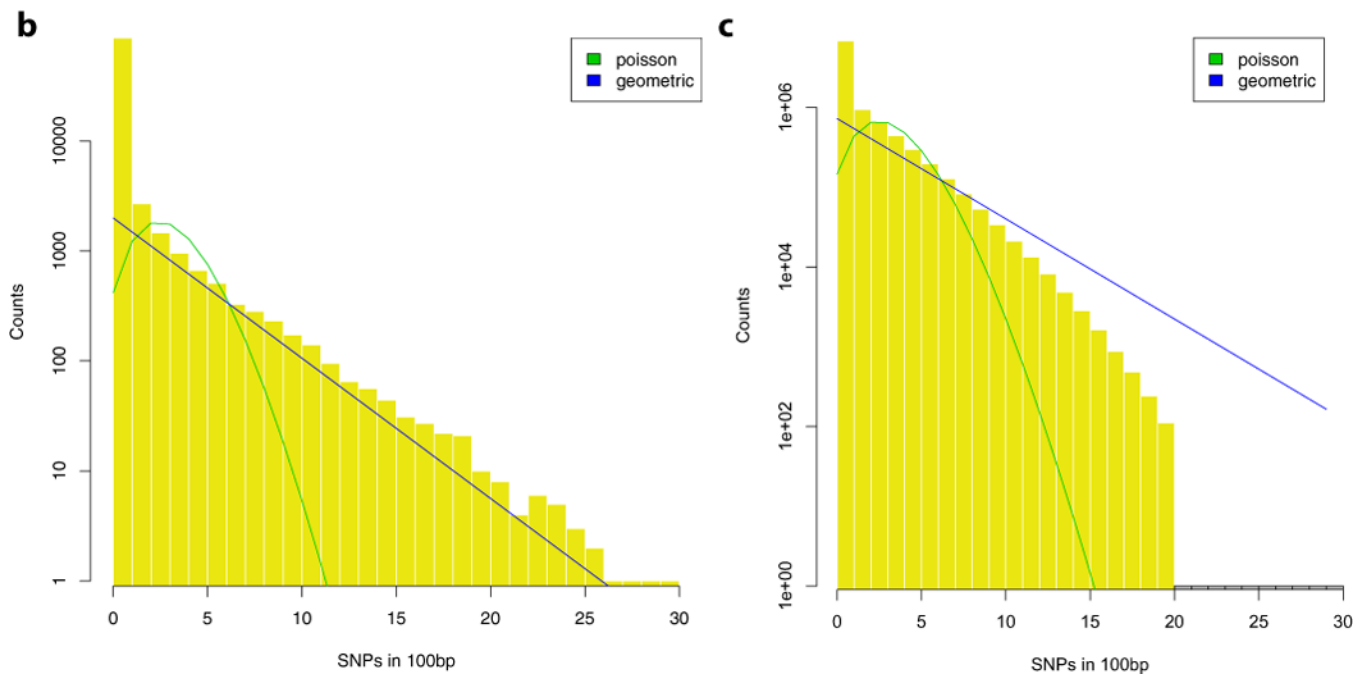
We examined in detail gene families found broadly in deuterostomes whose encoded peptides were readily alignable to microbial sequences but had no detectable similarity in non-deuterostome animals. Criteria for evaluation included: (1) the hemichordate gene matches microbial genes at least ten orders of magnitude in the  $e$ -value better than it matches sequences of non-deuterostome metazoans (most of the putative HGTs we describe have no non-deuterostome metazoan hit at all); (2) it has a defined genomic locus among bona fide metazoan genes; (3) it shares an exon-intron structure with genes of chordates and other ambulacraria; and (4) when a low bitscore match is found to a non-deuterostome metazoan sequence, that sequence is identified as containing different domains (domain structure according to CDD<sup>68</sup>) and/or different exon-intron structure, implying dubious relatedness. When phylogenetic trees are constructed for these HGT-candidate proteins, the trees contain numerous branches for microbial sequences and none for non-deuterostome metazoan sequences, or only very long branches for dubiously relatives, and hence the trees differ greatly from the metazoan species tree, except within the deuterostome clade.

## Code availability

Original data and code can be accessed at <https://groups.oist.jp/molgenu>.

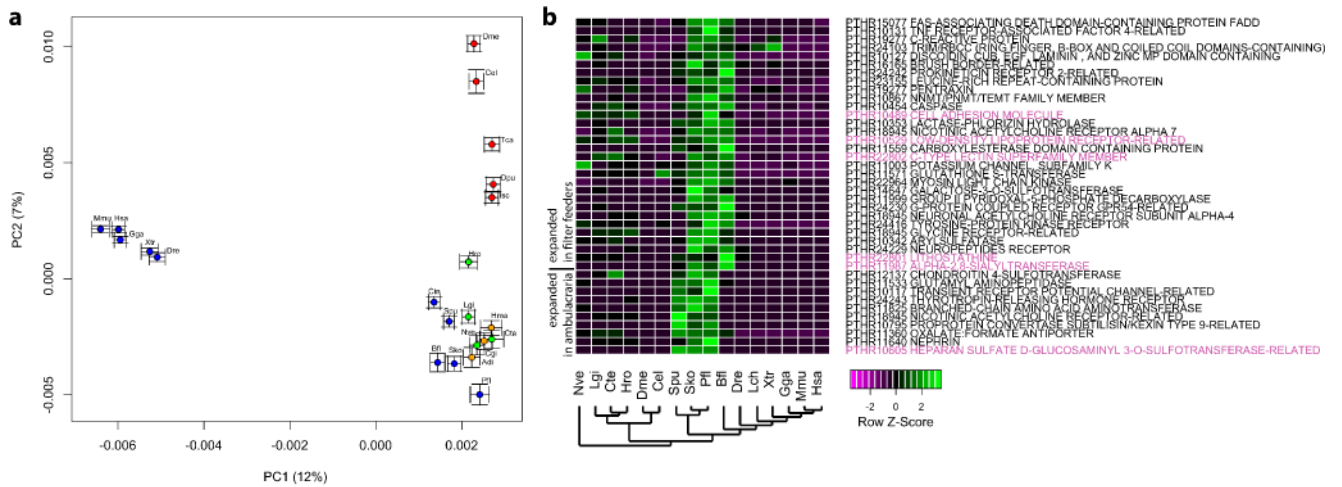
## Extended Data

| <b>a</b>                    | <i>Saccoglossus</i> | <i>Ptychodera</i>   |
|-----------------------------|---------------------|---------------------|
| Scaffold total              | 7,282               | 218,255             |
| Contig total                | 20,913              | 322,077             |
| Scaffold sequence total, Mb | 758                 | 1,229               |
| Scaffold N50, kb            | 552                 | 196                 |
| Gene models                 | 34,239              | 34,687              |
| SNP rate, %                 | 0.5                 | 1.3 (2x), 0.06 (1x) |



### Extended Data Figure 1. Summary of genome assemblies and heterozygosity distributions for *Saccoglossus* and *Ptychodera*

**a.** Genome statistics summary. **b, c.** The single nucleotide polymorphism distribution across 100-bp windows for *Saccoglossus* (**b**) and the corresponding distribution for *Ptychodera* (**c**). The distributions in **b** and **c** are fitted with a geometric (expected when high recombination rate is present) and a Poisson distribution (expected with low recombination rate). The distribution for *Saccoglossus* is fitted to windows with one or more SNPs only, as there is an excess of zero SNP windows (approximately 84% of total 94,324 selected windows). For methods refer to Supplementary Note 2.



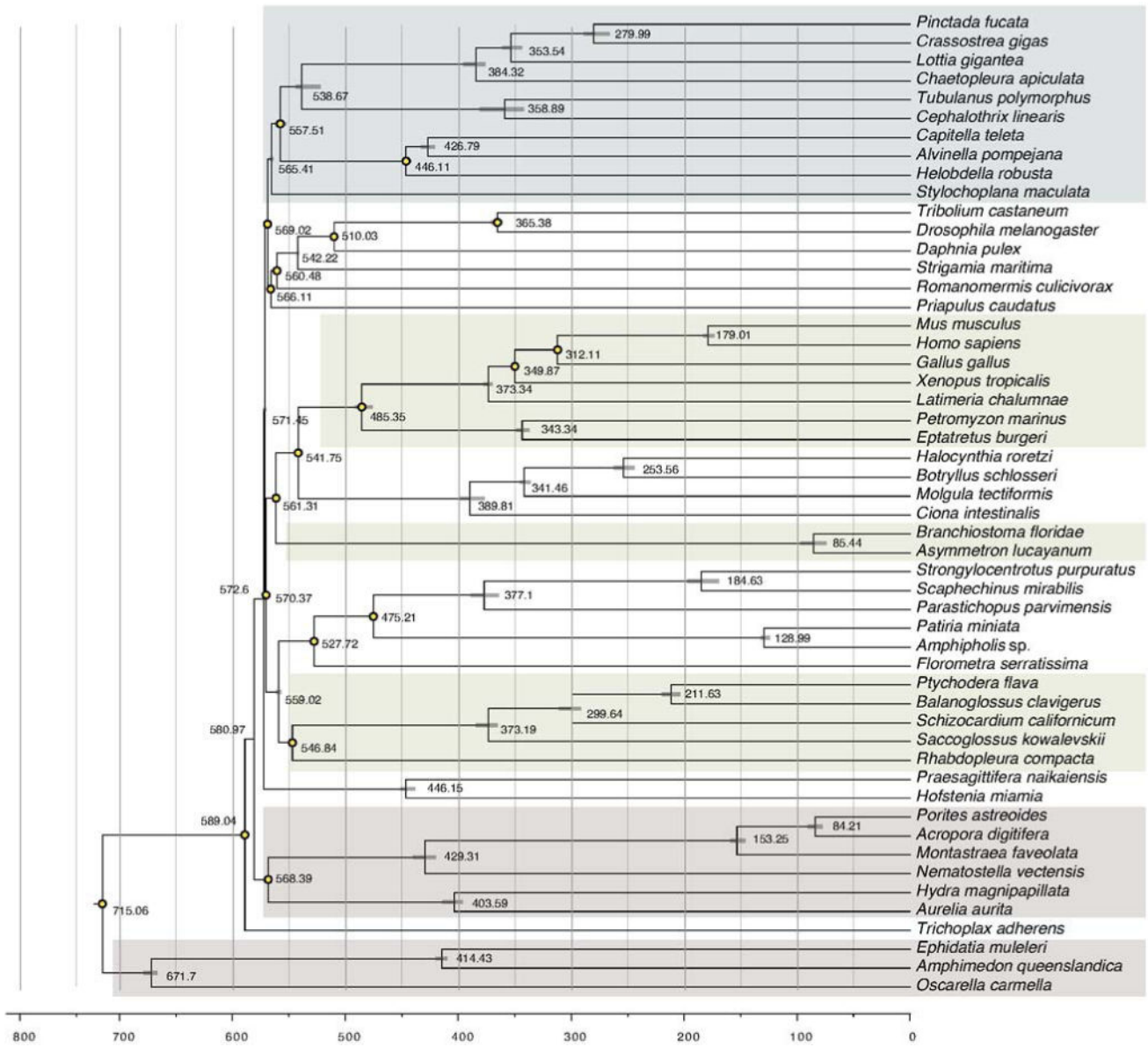
**Extended Data Figure 2. Ambulacrarians approximate the ancestral metazoan gene repertoire**  
**a**, Principal component analysis of Panther gene family sizes. Variances of the first two components are plotted in parentheses. Blue indicates deuterostomes; green indicates lophotrochozoans; red, ectodermozoans; yellow/orange, non-bilaterian metazoans. Note the clustering of the ambulacrarians Sgo, Pfl and Spu with the non-vertebrate deuterostomes Bfl and Cin in the lower right corner, also with the lophotrochozoans Cgi, Lgi, Hro, Cte and the non-bilaterians Hma, Nve and Adi. **b**, Heat map of gene family counts showing significant (Fisher’s exact test  $P$  value  $<0.01$  after Bonferroni multiple testing correction) expansion in ambulacrarians as well as in *Saccoglossus/Ptychodera/amphioxus*. The cases discussed in the main text are highlighted in red. See Supplementary Note 4 for details. Species abbreviations are defined in Supplementary Note 4.1.

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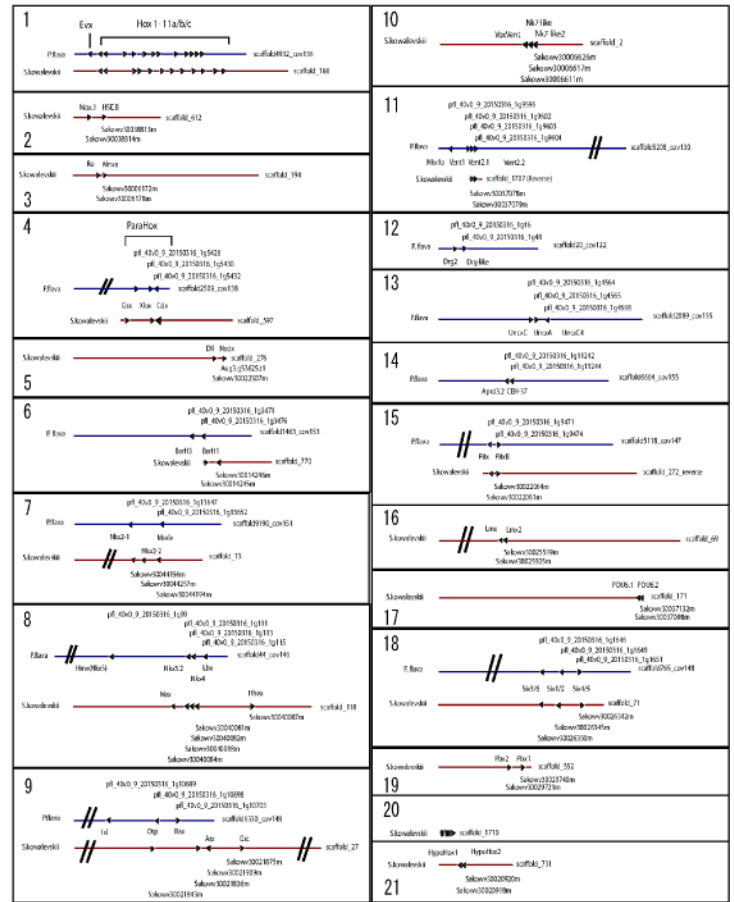
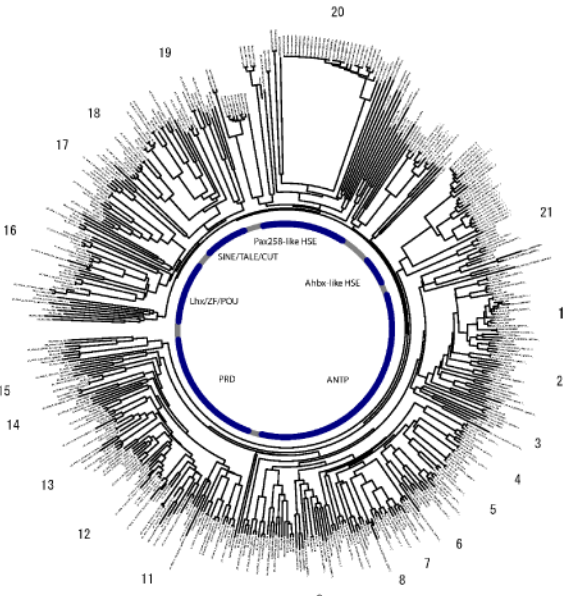
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**Extended Data Figure 3. Molecular dating of deuterostome and metazoan radiations using PhyloBayes assuming a log-normal relaxed clock model**

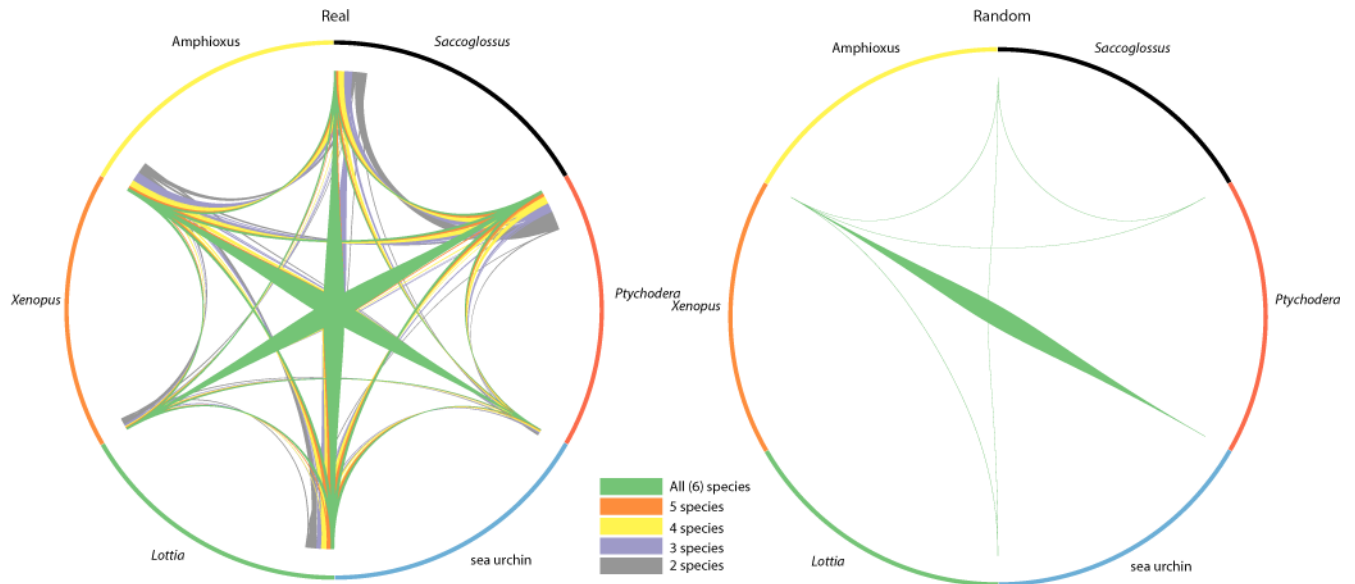
Yellow circles on particular nodes indicate the calibration dates applied from the fossil record, as indicated in Supplementary Note 6.2. Bars are 95% credibility intervals derived from posterior distributions. Note the estimated times of divergence of chordates and ambulacraria (the deuterostome ancestor) at 570 million years ago (Ma; mid-Ediacaran), hemichordates and echinoderms at 559 Ma, enteropneusts and pterobranchs at 547 Ma, and Harrimaniid and Ptychoderid enteropneusts at 373 Ma.



**Extended Data Figure 4. Homeobox gene complement of the two hemichordates in comparison to that of amphioxus. The numbers of homeobox-containing gene models are 170 in *Saccoglossus* and 139 in *Ptychodera***

These homeobox domains were aligned with 128 homeobox genes of *Branchiostoma floridae* using ClustalW2, then gaps and unaligned regions were manually removed. Since some genes have more than one homeobox domain, we kept all domains or chose the longest one according to the state of domain conservation. In total, 448 homeobox sequences were aligned. See Supplementary Information for details. The clusters of homeobox genes on scaffolds in *Saccoglossus* and *Ptychodera* were identified and drawn at positions around the tree. Conserved clusters between the two species were aligned. In addition to the well-known Hox and ParaHox cluster, 17 clusters were found in at least one of the hemichordates or some in both. Sixteen genes of the Nkx class are distributed over four clusters: (i) *nkx1a-vent1-vent2.1-vent2.2*; (ii) *nkx2.1-nkx2.2-msxlx*; (iii) *nkx5-msx-nkx3.2-nkx4-lbx-hex*; and (iv) *voxxent-nk7like-nk7like2*. The second cluster (ii) of these is part of the pharyngeal cluster (Fig. 4). Another five-gene cluster consists of one Lim class homeobox gene and four PRD class homeobox genes; *isl-otp-rax-arx-gsc*. A cluster of *six3/6-six1/2-six4/5* was found in both species, and a cluster of three *unx* genes was found only in *P. flava*. Ten more clusters were found containing two homeobox genes each. Notably, we found species-specific homeobox clusters in both species. Three remarkable clusters were found in *S. kowalevskii* in which 10, 12 and 5 homeobox-containing genes are tandem duplicated in scaffold\_1710, \_52 and \_4796, respectively. We also found such clusters in *P. flava* in which 7, 4, 8 and 10

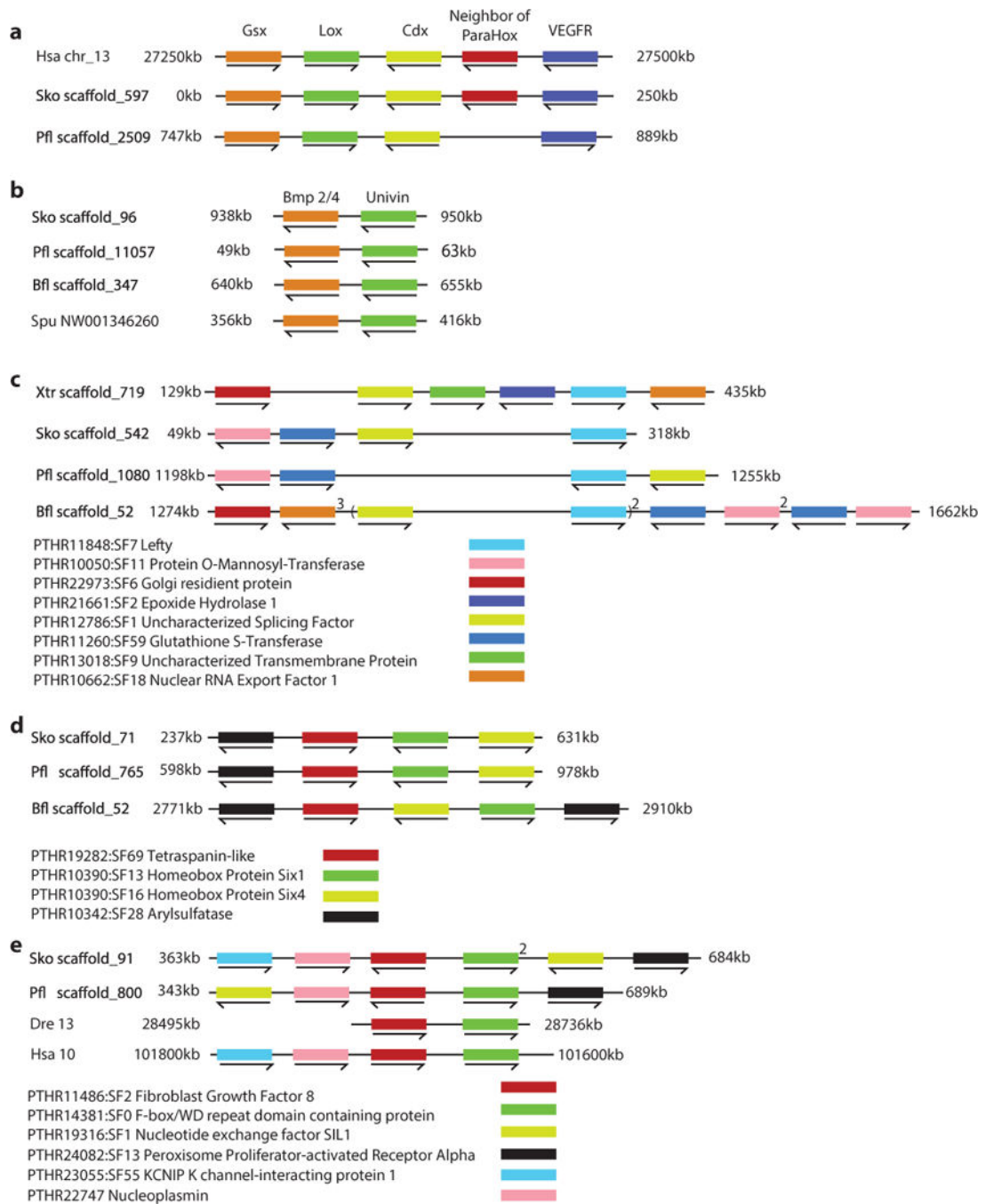
genes are aligned on scaffold 19451, scaffold 1398, scaffold 12422 and scaffold 154657, respectively. All homeobox genes identified in the genomes of the two hemichordates and amphioxus are listed in the Supplementary Table for Extended Data Fig. 4. This list includes some genes not containing a homeobox (for example, *pax1/9*) in cases where other family members do (for example, *pax2*).



**Extended Data Figure 5. High retention rate of micro-syntenic conservation in *Saccoglossus***

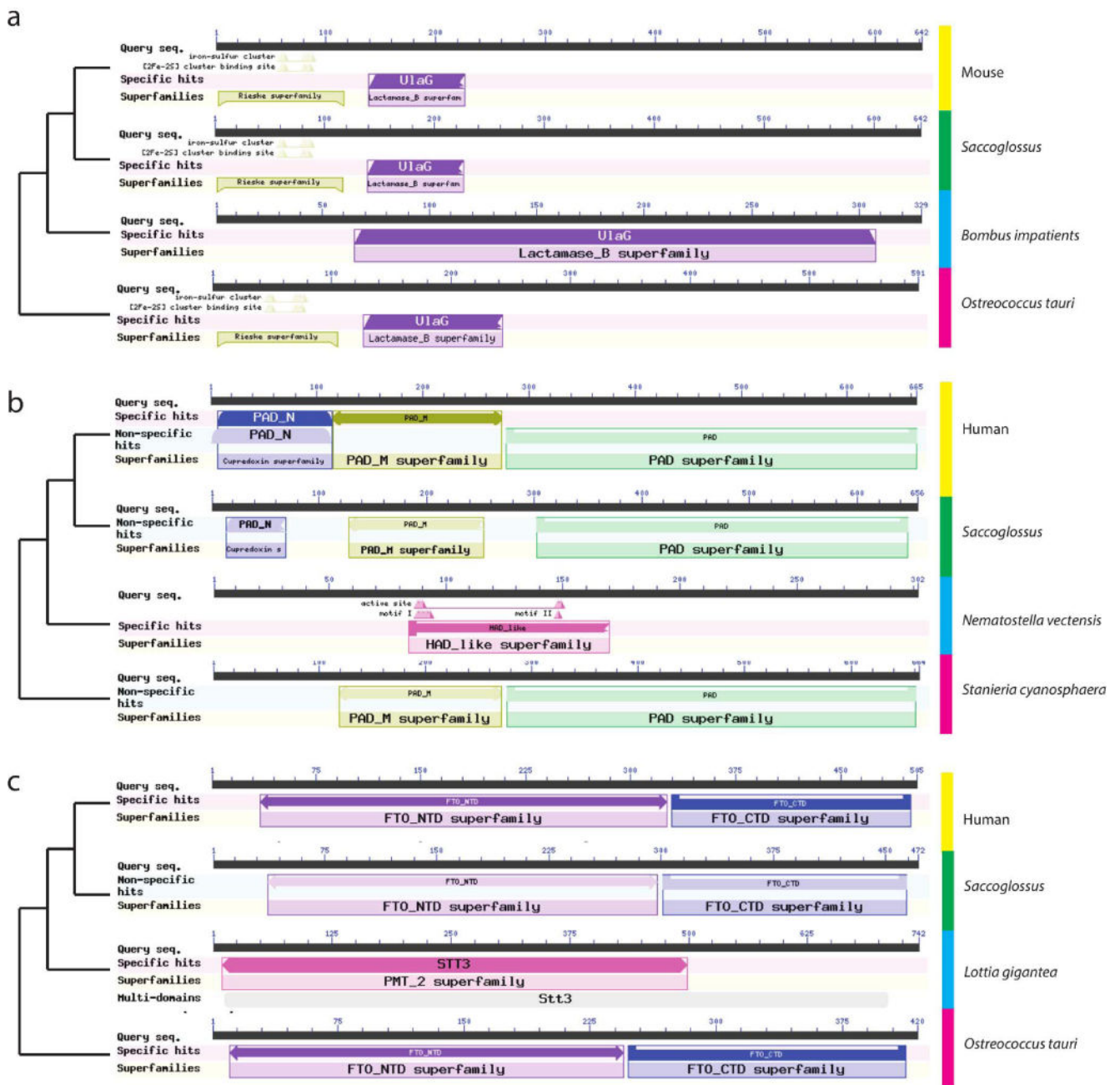
Circos plot showing micro-syntenic conservation in blocks of genes ( $n_{\max} = 10$  and  $n_{\min} = 2$ ) for six metazoan species for observed (left) and simulated (right) linkages. The width of connecting segments is proportional to the number of genes participating in the syntenic linkages (normalized by the total gene count). In this representation scaffolds are placed end-to-end, and adjacent scaffolds need not be from the same chromosome. While simulated data yields some blocks shared between pairs of species, few or no syntenic blocks can be recovered among three or more species (Methods). *Saccoglossus* shows one of the highest retentions among the selected species (and the highest among the sequenced ambulacrarians). *Xenopus* (and vertebrates in general) have lost some micro-syntenic conservation due to whole-genome duplications and differential loss of paralogues. The matching between the hemichordate *S. kowalevskii* and the chordate amphioxus is highest, consistent with the fact that neither genome has undergone extensive gene loss (as have tunicates) or pseudo-tetraploidization with extensive loss of paralogues (as have vertebrates).





**Extended Data Figure 6. Deuterostome specific micro-syntenic linkages**

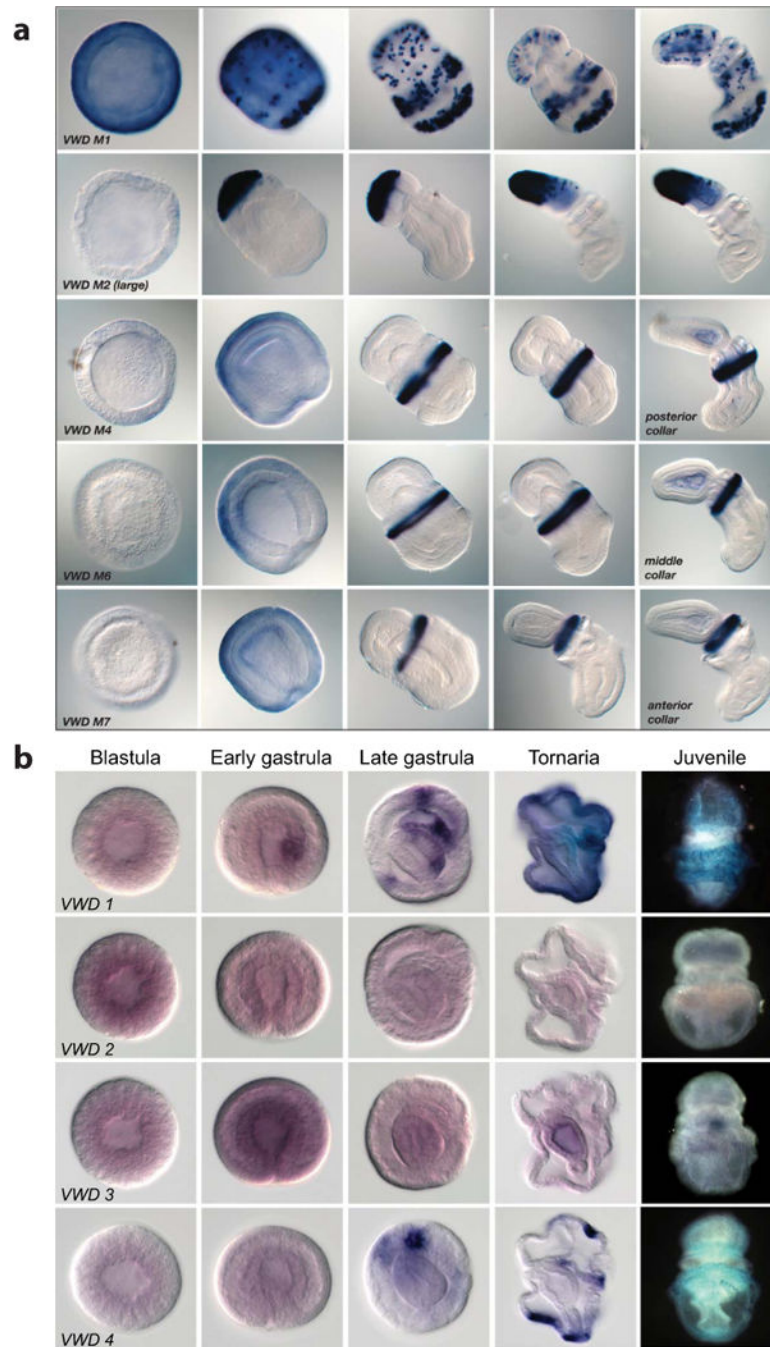
**a, b**, Very tight linkages with no intervening genes. **a**, ParaHox cluster shown in *S. kowalevskii*, *P. flava*, and human. **b**, *bmp2/4* and *univin* cluster in the hemichordates *S. kowalevskii* and *P. flava*, the sea urchin *S. purpuratus*, and the cephalochordate *B. floridae*. **c–e**, Loose micro-syntenic linkages with a maximum of five intervening genes: *lefty* (**c**), *six1–six4* (**d**), and *fgf8–fbxw* (**e**)<sup>69</sup> clusters. For **c** to **e** all species with micro-syteny are shown. Numbers above the genes indicate the copy number in the locus.



**Extended Data Figure 7. Three examples showing the domain structures of some proteins encoded by genes found in deuterostomes and marine microbes but not non-deuterostome animals**

Best BLASTP hits of the *Saccoglossus* sequence in human/mouse, as well as in non-deuterostome metazoans and in non-metazoans (such as the cyanobacterium *Staniera cyanosphaera*, or the eukaryotic micro-alga *Ostreococcus tauri*) are shown. **a**, Cytidine monophosphate-*N*-acetylneuraminic acid hydroxylase (CMAH), an enzyme of sialic acid modification; **b**, peptidyl arginyl deiminase (PAD), an enzyme of post-translational modification of proteins; **c**, FATS0-like, also called  $\alpha$ -ketoglutarate-dependent dioxygenase FTO, an enzyme that de-methylates *N*<sup>6</sup>-methyladenosine in nuclear RNA. Other analyses of

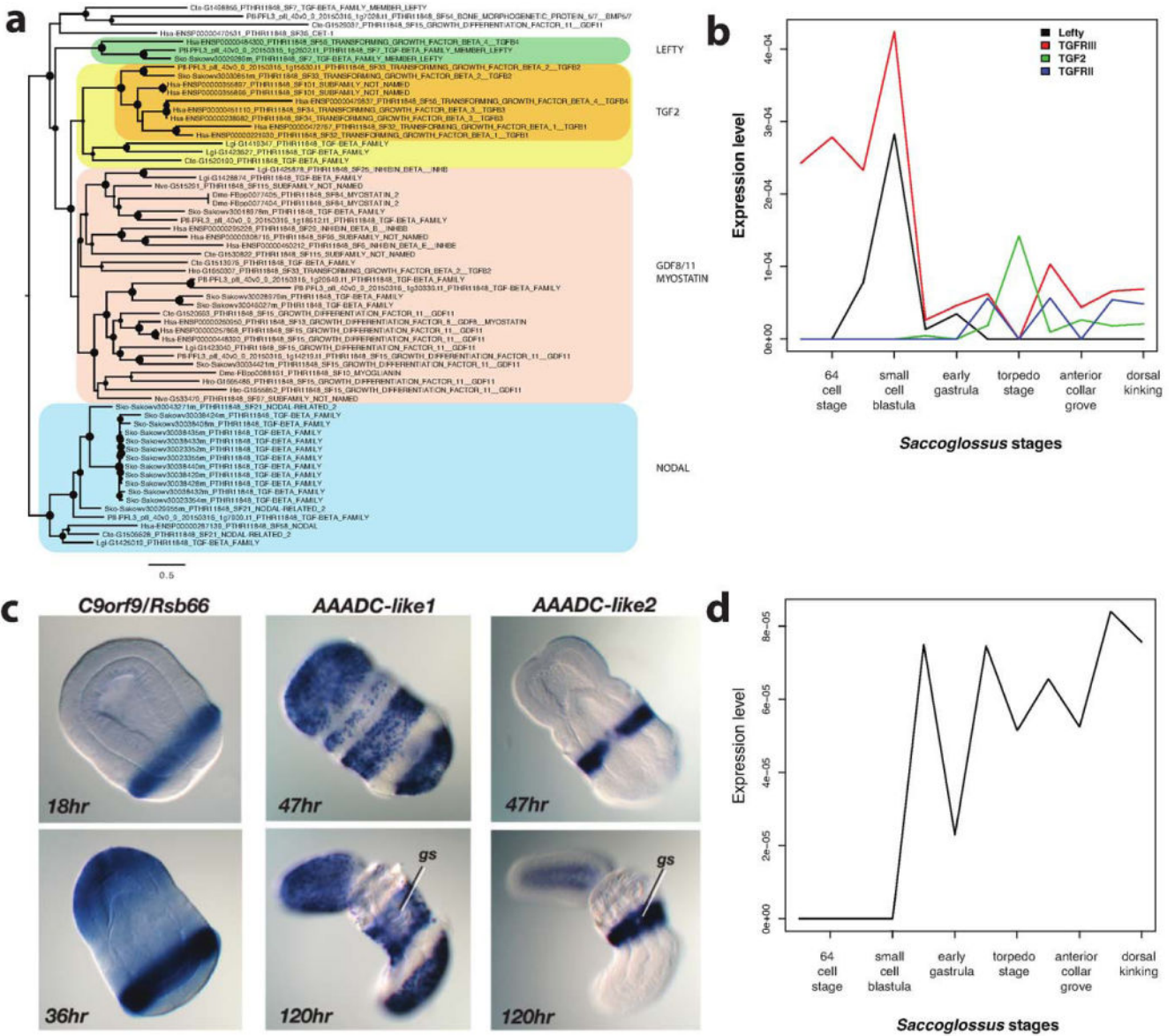
these and other genes with the unusual phylogenetic distributions can be found in Supplementary Note 10.



**Extended Data Figure 8. *In situ* hybridization demonstration of the expression of von Willebrand type D (vWD) domain-encoding genes (putative glycoproteins/mucins) in *Saccoglossus* and *Ptychodera***

**a**, In *Saccoglossus* the genes are specifically expressed in different subregions of the ectoderm of the proboscis or collar at these pre-feeding stages. **b**, In *Ptychodera*, several of

the genes are expressed in endoderm as well as ectoderm of the developing tornaria larva. The sequence IDs for the genes are provided in Supplementary Note S10.4.



**Extended Data Figure 9. Gene innovation in deuterostomes**

**a**, FastTree phylogenetic tree of the TGFβ family members Lefty, TGFβ 2, GDF8/11 and Nodal ligands (using GTR model). Bootstrap support is plotted as filled circles (size proportional to the support value) on each node. While Lefty shows deuterostome unique sequence composition, TGFβ 2 has an acceleration of sequence change at the deuterostome stem branch, compared to the GDF8/11 or Nodal groups. **b**, Temporal co-expression of Lefty and TGFβ receptor type III in *Saccoglossus* at pre-gastrulation developmental stages and of TGFβ 2 and TGFβ receptor type II at post-gastrulation stages. **c**, *In situ* hybridization demonstration of the expression in *S. kowalevskii* of one of the putative type I novelty genes (*c9orf9*, also known as *rsb66*) and of two of AAADC genes (aromatic amino acid

decarboxylases of the microbial type) of *S. kowalevskii* (also in *P. flava* and *B. floridae*), which closely resemble sequences from bacteria rather than from non-deuterostome metazoans. gs, gill slits. **d**, The temporal expression profile for *c9orf9* during *S. kowalevskii* development, taken from transcriptome data.

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## Supplementary Material

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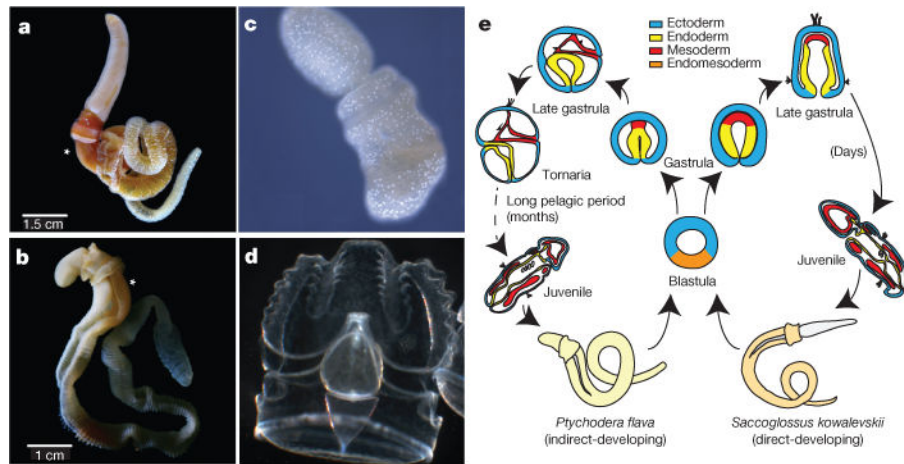
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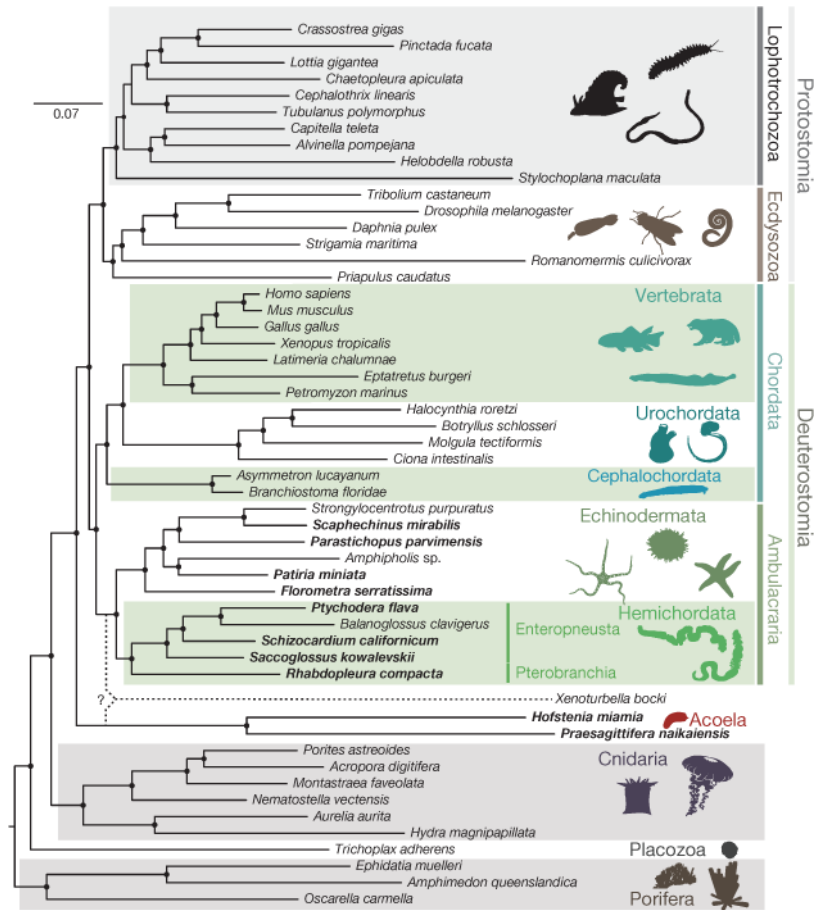
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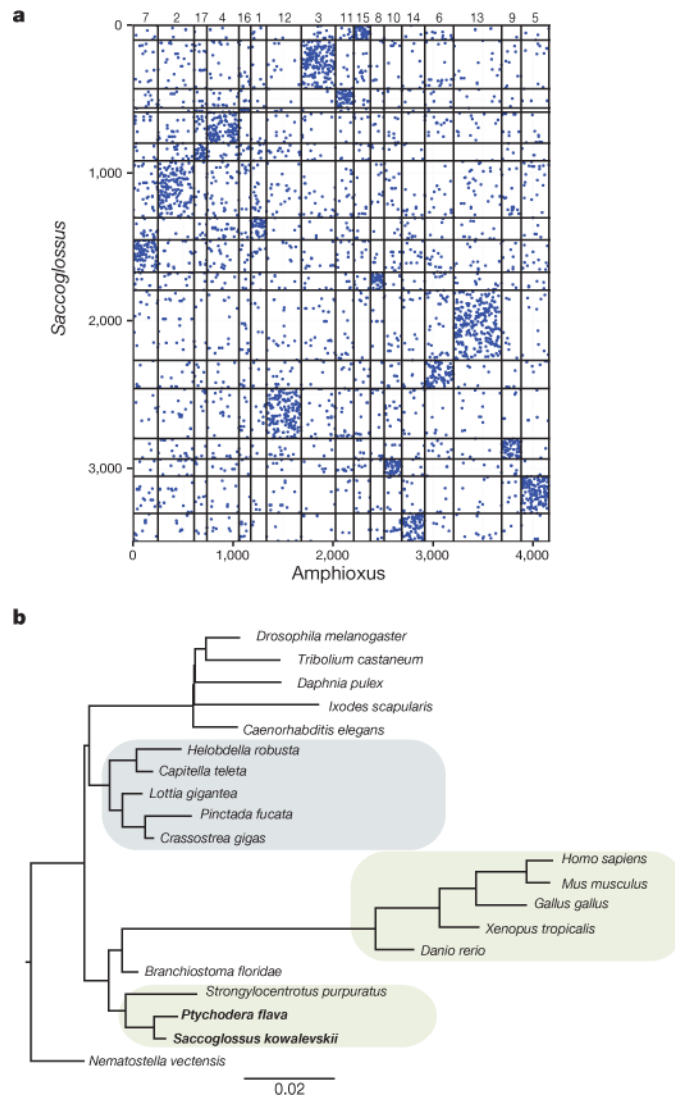


**Figure 1. Hemichordate model systems and their embryonic development**

The hemichordate phylum includes the enteropneusts (acorn worms) and pterobranchs (minute, colonial, tube-dwelling; not shown). **a, c**, *Saccoglossus kowalevskii* (Harrimaniid (direct developing) enteropneust) adult (**a**) and juvenile (**c**) with gill slits. **b, d**, *Ptychodera flava* (Ptychoderid (indirect developing) enteropneust) adult (**b**) and the tornaria stage larva (**d**). Gill slits labelled with an asterisk in **a** and **b**. **e**, Comparison of the direct and indirect modes of development of the two hemichordates, indicating the long pelagic larval period in *Ptychodera* until the settlement and metamorphosis as a juvenile.

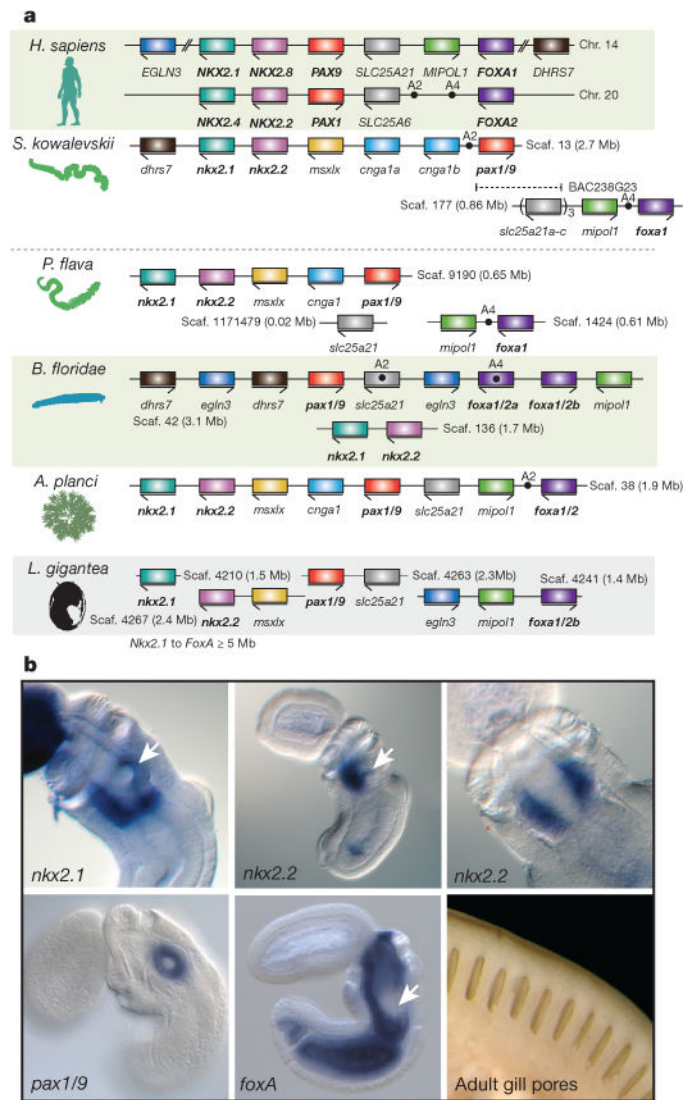


**Figure 2. Phylogenetic placement of deuterostome taxa within the metazoan tree**  
 Maximum-likelihood tree obtained with a super-matrix of 506,428 amino-acid residues gathered from 1,564 orthologous genes in 52 species (65.1% occupancy) and using a LG+Γ model partitioned for each gene. Filled circles at nodes denote maximal bootstrap support. Taxa highlighted in bold are newly sequenced genomes and transcriptomes introduced in this study. Bar indicates the number of substitutions per site.



**Figure 3. High level of linkage conservation in *Saccoglossus***

**a**, Macro-synteny dot plot between *Saccoglossus* and amphioxus; each dot represents two orthologous genes linked in the two species, and ordered according to their macro-syntenic linkage. Amphioxus scaffolds are organized according to the 17 ancestral linkage groups (ALGs) inferred by comparison of the amphioxus and vertebrate genomes<sup>27</sup>. Intersection areas of highest dot density are marked by numbers along the top of the plot, identifying each of the 17 putative ALGs. Axes represent orthologous gene group index along the genome. **b**, Branch-length estimation for loss and gain of synteny blocks with MrBayes, see Supplementary Note 7 for details. Short branches in hemichordates (in bold) indicate a high level of micro-syntenic retention in their genomes.

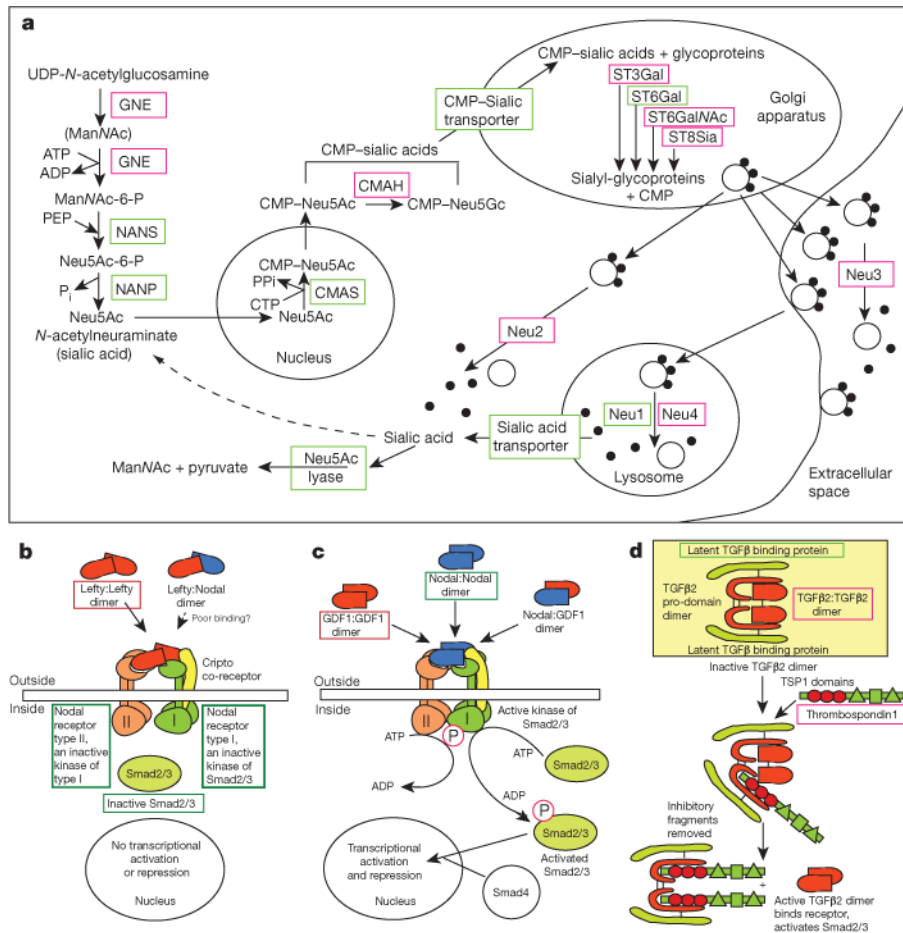


**Figure 4. Conservation of a pharyngeal gene cluster across deuterostomes**

**a**, Linkage and order of six genes including the four genes encoding transcription factors Nkx2.1, Nkx2.2, Pax1/9 and FoxA, and two genes encoding non-transcription factors Slc25A21 (solute transporter) and Mipol1 (mirror-image polydactyly 1 protein), which are putative ‘bystander’ genes containing regulatory elements of *pax1/9* and *foxA*, respectively. The pairings of *slc25A21* with *pax1/9* and of *mipol1* with *foxA* occur also in protostomes, indicating bilaterian ancestry. The cluster is not present in protostomes such as *Lottia* (Lophotrochozoa), *Drosophila melanogaster*, *Caenorhabditis elegans* (Ecdysozoa), or in the cnidarian, *Nematostella*. *SLC25A6* (the *slc25A21* paralogue on human chromosome 20) is a potential pseudogene. The dots marking A2 and A4 indicate two conserved non-coding sequences first recognized in vertebrates and amphioxus<sup>36</sup>, also present in *S. kowalevskii* and, partially, in *P. flava* and *A. planci*. **b**, The four transcription factor genes of the cluster are expressed in the pharyngeal/foregut endoderm of the *Saccoglossus* juvenile: *nkx2.1* is expressed in a band of endoderm at the level of the forming gill pore, especially ventral and posterior to it (arrow), and in a separate ectodermal domain in the proboscis. It is also

known as thyroid transcription factor 1 due to its expression in the pharyngeal thyroid rudiment in vertebrates. The *nkx2.2* gene is expressed in pharyngeal endoderm just ventral to the forming gill pore, shown in side view (arrow indicates gill pore) and ventral view; and *pax1/9* is expressed in the gill pore rudiment itself. In *S. kowalevskii*, this is its only expression domain, whereas in vertebrates it is also expressed in axial mesoderm. The *foxA* gene is expressed widely in endoderm but is repressed at the site of gill pore formation (arrow). An external view of gill pores is shown; up to 100 bilateral pairs are present in adults, indicative of the large size of the pharynx.





**Figure 5. Examples of deuterostome gene novelties**

**a**, Steps of biosynthesis of sialic acid and its addition to and removal from glycoproteins. **b–d**, Novel genes in TGFβ signalling pathways. The encoded proteins are shown and include Lefty (**b**), an antagonist of Nodal signalling, which activates Smad2/3-dependent transcription when not antagonized; Univin (**c**), an agonist of Nodal signalling, also called Vg1, DVR1, and GDF1; and TGFβ 2 (**d**), a ligand that activates Smad2/3-dependent transcription by binding to a deuterostome-specific TGFβ receptor type II, which contains a novel ectodomain (not shown). Also shown in **d** is the novel protein thrombospondin 1 that activates TGFβ 2 by releasing it from an inactive complex, by way of its TSP1 domains. Red boxes around protein names indicate their deuterostome novelty. Green boxes around the names indicate genes with pan-metazoan/bilaterian ancestry and without accelerated sequence change in the deuterostome lineage.