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Author manuscript *Nature*. Author manuscript; available in PMC 2016 March 17.

Published in final edited form as:

Nature. 2015 August 13; 524(7564): 220-224. doi:10.1038/nature14668.

# The octopus genome and the evolution of cephalopod neural and morphological novelties

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# Abstract

Coleoid cephalopods (octopus, squid, and cuttlefish) are active, resourceful predators with a rich behavioral repertoire<sup>1</sup>. They have the largest nervous systems among the invertebrates<sup>2</sup> and present other striking morphological innovations including camera-like eyes, prehensile arms, a highly derived early embryogenesis, and the most sophisticated adaptive coloration system among all animals<sup>1,3</sup>. To investigate the molecular bases of cephalopod brain and body innovations we sequenced the genome and multiple transcriptomes of the California two-spot octopus, *Octopus bimaculoides*. We found no evidence for hypothesized whole genome duplications in the octopus lineage<sup>4–6</sup>. The core developmental and neuronal gene repertoire of the octopus is broadly similar to that found across invertebrate bilaterians, except for massive expansions in two gene families formerly thought to be uniquely enlarged in vertebrates: the protocadherins, which regulate neuronal development, and the C2H2 superfamily of zinc finger transcription factors. Extensive mRNA editing generates transcript and protein diversity in genes involved in neural excitability,

contributed equally

Author contributions

#### **Competing financial interests**

The authors declare no competing financial interests

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

The Chicago and the OIST/Berkeley groups initiated their transcriptome and genome projects independently. In the subsequent collaboration, both groups worked closely on every aspect of the project. Chicago group: C.B.A., Z.Y.W., J.R.P., C.W.R; OIST/ Berkeley group: O.S., T.M., E.E.-G., S.B., D.S.R.

as previously described<sup>7</sup>, as well as in genes participating in a broad range of other cellular functions. We identified hundreds of cephalopod-specific genes, many of which showed elevated expression levels in such specialized structures as the skin, the suckers, and the nervous system

expression levels in such specialized structures as the skin, the suckers, and the nervous system. Finally, we found evidence for large-scale genomic rearrangements that are closely associated with transposable element expansions. Our analysis suggests that substantial expansion of a handful of gene families, along with extensive remodeling of genome linkage and repetitive content, played a critical role in the evolution of cephalopod morphological innovations, including their large and complex nervous systems.

Soft-bodied cephalopods such as the octopus (Fig. 1a) show remarkable morphological departures from the basic molluscan body plan, including dexterous arms lined with hundreds of suckers that function as specialized tactile and chemosensory organs, and an elaborate chromatophore system under direct neural control that enables rapid changes in appearance<sup>1,8</sup>. The octopus nervous system is vastly modified in size and organization relative to other molluscs, comprising a circumesophageal brain, paired optic lobes, and axial nerve cords in each arm<sup>2,3</sup>. Together these structures contain nearly half a billion neurons, more than six times the number in a mouse brain<sup>2,9</sup>. Extant coleoid cephalopods show extraordinarily sophisticated behaviors including complex problem solving, task-dependent conditional discrimination, observational learning and spectacular displays of camouflage<sup>1,10</sup> (Supplementary Videos 1, 2).

To explore the genetic features of these highly specialized animals, we sequenced the *Octopus bimaculoides* genome by a whole genome shotgun approach (Supplementary Note 1) and annotated it using extensive transcriptome sequence from 12 tissues (Methods; Supplementary Note 2). The genome assembly captures more than 97% of expressed protein coding genes and 83% of the estimated 2.7 Gb genome size (Methods; Supplementary Notes 1–3). The unassembled fraction is dominated by high-copy repetitive sequences (Supplementary Note 1). Nearly 45% of the assembled genome is composed of repetitive elements, with two bursts of transposon activity occurring ~25 and ~56 mya (Supplementary Note 4).

We predicted 33,638 protein-coding genes (Methods, Supplementary Note 4) and found alternate splicing at 2,819 loci, but no locus has an extraordinary number of splice variants (Supplementary Note 4). A-to-G discrepancies between the assembled genome and transcriptome sequences provided evidence for extensive mRNA editing by adenosine deaminases acting on RNA (ADARs). Many candidate edits are enriched in neural tissues<sup>7</sup> and are found in a range of gene families, including "housekeeping" genes such as the tubulins, which suggests that RNA edits are more widespread than previously appreciated (Extended Data Fig. 1, Supplementary Note 5).

Based primarily on chromosome number, several researchers proposed that whole genome duplications were important in the evolution of the cephalopod body plan<sup>4–6</sup>, paralleling the role ascribed to the independent whole genome duplication events that occurred early in vertebrate evolution<sup>11</sup>. While this is an attractive framework for both gene family expansion and increased regulatory complexity across multiple genes, we found no evidence for it. The gene family expansions present in octopus are predominantly organized in clusters along the

paleopolyploid<sup>12,13</sup> (Supplementary Note 6.2). While genes that regulate development are often retained in multiple copies after paleopolyploidy in other lineages, they are not generally expanded in octopus relative to limpet, oyster, and other invertebrate bilaterians<sup>11,14</sup> (Table 1; Supplementary Notes 7.4, 8).

While Hox genes are commonly retained in multiple copies following whole genome duplication<sup>15</sup>, we found only a single Hox complement in *O. bimaculoides*, consistent with the single set of Hox transcripts identified in the bobtail squid *Euprymna scolopes* with PCR<sup>16</sup>. Remarkably, octopus Hox genes are not organized into clusters as in most other bilaterian genomes<sup>15</sup>, but are completely atomized (Extended Data Fig. 2; Supplementary Note 9). While we cannot rule out whole genome duplication followed by considerable gene loss, the extent of loss needed to support this claim would far exceed that which has been observed in other paleopolyploid lineages, and it is more plausible that chromosome number in coleoids increased by chromosome fragmentation.

Mechanisms other than whole genome duplications can drive genomic novelty, including expansion of existing gene families, evolution of novel genes, modification of gene regulatory networks, and reorganization of the genome through transposon activity. Within the *O. bimaculoides* genome, we found evidence for all of these mechanisms, including expansions in several gene families, a suite of octopus- and cephalopod-specific genes, and extensive genome shuffling.

In gene family content, domain architecture, and exon-intron structure, the octopus genome broadly resembles that of the limpet *Lottia gigantea*<sup>17</sup>, the polychaete annelid *Capitella teleta*<sup>17</sup>, and the cephalochordate *Branchiostoma floridae*<sup>14</sup> (Supplementary Note 7, Extended Data Fig. 3). Relative to these invertebrate bilaterians, we found a fairly standard set of developmentally important transcription factors and signaling pathway genes, suggesting that the evolution of the cephalopod body plan did not require extreme expansions of these "toolkit" genes (Table 1; Supplementary Note 8.2). Statistical analysis of protein domain distributions across animal genomes did, however, identify several notable gene family expansions in octopus, including protocadherins, C2H2 zinc-finger proteins (C2H2-ZNFs), interleukin 17-like genes (IL17-like), G-protein coupled receptors (GPCRs), chitinases, and sialins (Figs. 2–3; Extended Data Figs 4–6; Supplementary Notes 8 and 10).

The octopus genome encodes 168 multi-exonic protocadherin genes, nearly three-quarters of which are found in tandem clusters on the genome (Fig. 2b), a dramatic expansion relative to the 17–25 genes found in *Lottia, Crassostrea,* and *Capitella* genomes. Protocadherins are homophilic cell adhesion molecules whose function has been primarily studied in mammals, in which they are required for neuronal development and survival as well as synaptic specificity<sup>18</sup>. Single protocadherin genes are found in the invertebrate deuterostomes *Saccoglossus kowalevskii* and *Strongylocentrotus purpuratus,* indicating that their absence in *Drosophila melanogaster* and *Caenorhabditis elegans* is due to gene loss. Vertebrates also show a remarkable expansion of the protocadherin repertoire, which is generated by complex splicing from a clustered locus rather than tandem gene duplication (reviewed

A search of available transcriptome data from the longfin inshore squid *Doryteuthis* (formerly, *Loligo*) *pealeii*<sup>20</sup> also demonstrated an expanded number of protocadherin genes (Supplementary Note 8.3). Surprisingly, our phylogenetic analyses suggest that the squid and octopus protocadherin arrays arose independently. Unlinked octopus protocadherins appear to have expanded ~135 mya, after octopuses diverged from squid. In contrast, clustered octopus protocadherins are much more similar in sequence, either due to more recent duplications or gene conversion as found in clustered protocadherins in zebrafish and mammals<sup>21</sup>.

The expression of protocadherins in octopus neural tissues (Fig. 2) is consistent with a central role for these genes in establishing and maintaining cephalopod nervous system organization. Protocadherin diversity provides a mechanism for regulating the short-range interactions needed for the assembly of local neural circuits<sup>18</sup>, which is where the greatest complexity in the cephalopod nervous system appears<sup>2</sup>. The importance of local neuropil interactions, rather than long-range connections, is likely due to the limits placed on axon density and connectivity by the absence of myelin, since thick axons are then required for rapid high fidelity signal conduction over long distances. The sequence divergence between octopus and squid protocadherin expansions may reflect the dramatic differences between decapodiforms and octopuses in brain organization, which have been most clearly demonstrated for the vertical lobe, a key structure in cephalopod learning and memory circuits<sup>2,22</sup>. Finally, the independent expansions and nervous system enrichment of protocadherins in coleoid cephalopods and vertebrates offers a striking example of convergent evolution between these clades at the molecular level.

As with the protocadherins, we found multiple clusters of C2H2-ZNF transcription factor genes (Fig. 3a; Supplementary Note 8.4). The octopus genome contains nearly 1,800 multiexonic C2H2-containing genes (Table 1), more than the 200–400 C2H2-ZNFs found in other lophotrochozoans and the 500–700 found in eutherian mammals, in which they form the second largest gene family<sup>23</sup>. C2H2-ZNF transcription factors contain multiple C2H2 domains that, in combination, result in highly specific nucleic acid binding. The octopus C2H2-ZNFs typically contain 10–20 C2H2 domains but some have as many as 60 (Supplementary Note 8.4). The majority of the transcripts are expressed in embryonic and nervous tissues (Fig. 3b). This pattern of expression is consistent with roles for C2H2-ZNFs in cell fate determination, early development, and transposon silencing, as demonstrated in genetic model systems<sup>23</sup>.

The expansion of the *O. bimaculoides* C2H2-ZNFs coincides with a burst of transposable element activity at ~25 mya (Fig. 3c). The flanking regions of these genes show a significant enrichment in a 70–90 bp tandem repeat (31% for C2H2 genes vs. 4% for all genes; Fisher's exact test p-value  $< e^{-16}$ ), which parallels the linkage of C2H2 gene expansions to beta-satellite repeats in humans<sup>24</sup>. We also found an expanded C2H2-ZNF repertoire in amphioxus (Table 1), showing a similar enrichment in satellite-like repeats. These parallels

suggest a common mode of expansion of a highly dynamic transcription factor family implicated in lineage-specific innovations.

To investigate further the evolution of gene families implicated in nervous system development and function, we surveyed genes associated with axon guidance (Table 1) and neurotransmission (Table 2), identifying their homologues in octopus and comparing numbers across a diverse set of animal genomes (Supplementary Notes 8–10). Several patterns emerged. The gene complements present in the model organisms *D. melanogaster* and *C. elegans* often showed dramatic departures from those seen in lophotrochozoans and vertebrates (Table 2; Supplementary Note 10). For example, *D. melanogaster* encodes one member of the discs, large (DLG) family, a key component of the postsynaptic scaffold. In contrast, mammals have four *DLGs*, which (along with other observations) led to suggestions that vertebrates possess uniquely complex synaptic machinery<sup>25</sup>. We find, however, three *DLGs* in both octopus and limpet, suggesting that vertebrate-fly gene number differences are not necessarily diagnostic of exceptional vertebrate synaptic complexity (Supplementary Note 10.6).

Overall, neurotransmission gene family sizes in the octopus were very similar to those seen in other lophotrochozoans (Table 2, Supplementary Note 10), except for a few dramatically expanded gene families such as the sialic acid vesicular transporters (sialins) (Supplementary Note 10.2). We did find variations in the sizes of neurotransmission gene families between human and lophotrochozoans (Table 2, Supplementary Note 10), but no evidence for systematic expansion of these gene families in vertebrates relative to octopus or other lophotrochozoans. While some gene families were larger in mammals or absent in lophotrochozoans (*e.g.*, ligand gated 5-HT receptors), others were absent in mammals and present in invertebrates (*e.g.*, anionic glutamate and acetylcholine receptors). The complement of neurotransmission genes in octopus may be broadly typical for a lophotrochozoan, but our findings suggest it is also not obviously smaller than what is found in mammals.

Among the octopus complement of ligand-gated ion channels, we identified a set of atypical nicotinic acetylcholine receptor-like genes, most of which are tandemly arrayed in clusters (Extended Data Fig. 7). These subunits lack several residues identified as necessary for the binding of acetylcholine<sup>26</sup>, so it is unlikely that they function as acetylcholine receptors. The high levels of expression of these divergent subunits within the suckers raises the interesting possibility that they act as sensory receptors, as do some divergent glutamate receptors in other protostomes<sup>27</sup>. In addition, we identified 74 *Aplysia*-like and 11 vertebrate-like candidate chemoreceptors among the octopus GPCR superfamily of ~320 genes (Extended Data Fig. 6).

We found, amid extensive transcription of octopus transposons, that a class of octopusspecific SINEs is highly expressed in neural tissues (Supplementary Note 4, Extended Data Fig. 8). While the role of active transposons is unclear, elevated transposon expression in neural tissues has been suggested to serve an important function in learning and memory in mammals and flies<sup>28</sup>.

Transposable element insertions are often associated with genomic rearrangements<sup>29</sup> and we found that the transposon-rich octopus genome displays substantial loss of ancestral bilaterian linkages that are conserved in other species (Supplementary Note 6; Extended Data Fig. 9). Interestingly, genes that are linked in other bilaterians but not in octopus are enriched in neighboring SINE content. SINE insertions around these genes date to the time of tandem C2H2 expansion (Extended Data Fig. 9d), pointing to a crucial period of genome evolution in octopus. Other transposons such as Mariner show no such enrichment, suggesting distinct roles for different classes of transposons in shaping genome structure (Extended Data Fig. 9c).

Transposable element activity has been implicated in the modification of gene regulation across several eukaryotic lineages<sup>29</sup>. We found that in the nervous system, the degree to which a gene's expression is tissue-specific is positively correlated with the transposon load around that gene (r-squared values ranging from 0.49 in optic lobe to 0.81 in subesophageal brain; Extended Data Fig. 8; Supplementary Note 4). This correlation may reflect modulation of gene expression by transposon-derived enhancers or a greater tolerance for transposon insertion near genes with less complex patterns of tissue-specific gene regulation.

Using a relaxed molecular clock, we estimate that the octopus and squid lineages diverged ~270 mya, emphasizing the deep evolutionary history of coleoid cephalopods<sup>8,30</sup> (Supplementary Note 7.1; Extended Data Figure 10a). Our analyses found hundreds of coleoid- and octopus-specific novel genes, many of which were expressed in tissues containing novel structures, including the chromatophore-laden skin, the suckers, and the nervous system (Extended Data Fig. 10, Supplementary Note 11). Taken together, these novel genes, the expansion of C2H2-ZNFs, genome rearrangements, and extensive transposable element activity yield a new landscape for both *trans-* and *cis-* regulatory elements in the octopus genome, resulting in changes in an otherwise "typical" lophotrochozoan gene complement that contributed to the evolution of cephalopod neural complexity and morphological innovations.

# **METHODS**

#### Data access

Genome and transcriptome sequence reads are deposited in the SRA as BioProject PRJNA270931. The genome assembly and annotation are linked to the same BioProject ID. A browser of this genome assembly is available at http://octopus.metazome.net/.

#### Genome sequencing and assembly

Genomic DNA from a single male *Octopus bimaculoides*<sup>31</sup> was isolated and sequenced using Illumina technology to 60-fold redundant coverage in libraries spanning a range of pairs from ~350 bp to 10 kb. These data were assembled with meraculous<sup>32</sup> achieving a contig N50-length of 5.4 kb and a scaffold N50-length of 470 kb. The longest scaffold contains 99 genes and half of all predicted genes are on scaffolds with 8 or more genes (Supplementary Note 1).

# Genome size and heterozygosity

The *O. bimaculoides* haploid genome size was estimated to be ~2.7 gigabases (Gb) based on fluorescence (2.66–2.68 Gb) and k-mer (2.86 Gb) measurements (Supplementary Notes 1 and 2), making it several times larger than other sequenced molluscan and lophotrochozoan genomes<sup>17</sup>. We observed nucleotide-level heterozygosity within the sequenced genome to be 0.08%, which may reflect a small effective population size relative to broadcast-spawning marine invertebrates.

#### Transcriptome sequencing

Twelve transcriptomes were sequenced from RNA isolated from ova, testes, viscera, posterior salivary gland (PSG), suckers, skin, developmental stage 15 (St15)<sup>33</sup>, retina, optic lobe (OL), supraesophageal brain (Supra), subesophageal brain (Sub), and axial nerve cord (ANC) (Supplementary Note 2). RNA was isolated using Trizol (Invitrogen) and 100bp paired-end reads (insert size 300bp) were generated on an Illumina HiSeq2000 sequencing machine.

#### De novo transcriptome assembly

Adapters and low quality reads were removed before assembling transcriptomes using the Trinity *de novo* assembly package [version r2013-02-25<sup>34,35</sup>]. Assembly statistics are summarized in Table S2.2. Following assembly, peptide-coding regions were translated using TransDecoder in the Trinity package. We compared the *de novo* assembled RNA-Seq output to the genome to evaluate the completeness of the genome assembly. To minimize the number of spuriously assembled transcripts, only transcripts with ORFs predicted by TransDecoder were mapped onto the genome with BLASTN. Only 1,130 out of 48,259 transcripts with ORFs (2.34%) did not have a match in the genome with a minimum identity of 95%.

#### Annotation of transposable elements

Transposable elements were identified with RepeatModeler<sup>36</sup> and Repeatmasker<sup>37</sup>, as outlined in Supplementary Note 4.2. The most abundant transposable element is a previously identified octopus-specific SINE<sup>38</sup> that accounts for 4% of the assembled genome.

# Annotation of protein coding genes

Protein-coding genes were annotated by combining transcriptome evidence with homologybased and *de novo* gene prediction methods (Supplementary Note 4). For homology prediction we used predicted peptide sets of three previously sequenced molluscs (*L. gigantea*, *C. gigas*, and *A. californica*) along with selected other metazoans. Alternative splice isoforms were identified with PASA<sup>39</sup>. Annotation statistics are provided in Table S4.1.1. Genes known in vertebrates to have many isoforms, such as ankyrin, *TRAK1*, and *LRCH1*, also show alternative splicing in octopus but at a more limited level. Octopus genes with ten or more alternative splice forms are provided in Table S4.1.2.

# Calibration of sequence divergence with respect to time

The divergence between squid and octopus was estimated using  $r8s^{40}$  by fixing cephalopod divergence from bivalves and gastropods to 540 mya<sup>8</sup>. Our estimate of 270 mya for the squid-octopus divergence corresponds to mean neutral substitution rate of dS ~2 based on the protein-directed CDS alignments between the species (Supplementary Figure S6.1.2) and a dS estimation using the yn00 program<sup>41</sup>. Throughout the manuscript we convert from sequence divergence to time by assuming that dS ~1 corresponds to 135 million years. For example, unlinked octopus protocadherins appear to have expanded ~135 mya based on mean pairwise dS~1, after octopuses diverged from squid. In contrast, clustered octopus protocadherins are much more similar in sequence (mean pairwise dS ~0.4, or ~55 mya).

#### Quantifying gene expression

Transcriptome reads were mapped to the genome assembly with TopHat 2.0.11<sup>42</sup>. A range of 76–90% of reads from the different samples mapped to the genome. Mapped reads were sorted and indexed with SAMtools<sup>43</sup>. The read counts in each tissue were produced with bedtools multicov program<sup>44</sup> using the gene model coordinates. The counts were normalized by the total transcriptome size of each tissue and by the length of the gene. Heatmaps showing expression patterns were generated in R using the heatmap.2 function.

## Gene complement

Gene families of particular interest, including developmental regulatory genes, neuralrelated genes, and gene families that appear to be expanded in O. bimaculoides, were manually curated and analyzed. We searched the octopus genome and transcriptome assemblies using BLASTP and TBLASTN with annotated sequences from human, mouse, and D. melanogaster. Bulk analyses were also performed using Pfam<sup>45</sup> and PANTHER<sup>46</sup>. We also used BLASTP and TBLASTX to search for specific gene families in deposited genome and transcriptome databases for L. gigantea, A. californica, C. gigas, C. teleta, T. castaneum, D. melanogaster, C. elegans, N. vectensis, A. queenslandica S. kowalevskii, B. floridae, C. intestinalis, D. rerio, M. musculus, and H. sapiens. Candidate genes were verified with BLAST<sup>47</sup> and Pfam<sup>45</sup> analysis. Genes identified in the octopus genome were confirmed and extended using the transcriptomes. Multiple gene models that matched the same transcript were combined. The identified sequences from octopus and other bilaterians were aligned with either MUSCLE<sup>48</sup>, CLUSTALO<sup>49</sup>, MacVector 12.6 (MacVector Inc, North Carolina), or Jalview<sup>50</sup>. Phylogenetic trees were constructed with FastTree<sup>51</sup> using the Jones-Taylor-Thornton model of amino acid evolution, and members of each family were counted.

## Synteny

Microsynteny was computed based on metazoan node gene families (Supplementary Note 7). We used Nmax 10 (maximum of 10 intervening genes) and Nmin 3 (minimum of three genes in a syntenic block) according to the pipeline described in Simakov et al. (2013) (Supplementary Note 6). To simplify gene family assignments we limited our analyses to 4,033 gene families shared among human, amphioxus, *Capitella*, *Helobdella*, *Octopus*, *Lottia*, *Crassostrea*, *Drosophila*, and *Nematostella*. We required ancestral bilaterian syntenic

blocks to have a minimum of one species present in both ingroups, or in one ingroup and one outgroup. To examine the effect of fragmented genome assemblies, we simulated shorter assemblies by artificially fragmenting genomes to contain on average 5 genes per scaffold (Supplementary Note 6).

In comparison with other bilaterian genomes, we find that the octopus genome is substantially rearranged. In looking at micro-syntenic linkages of genes with a maximum of 10 intervening genes, we found that octopus conserves only 34 out of 198 ancestral bilaterian microsyntenic blocks; the limpet *Lottia* and amphioxus retain more than twice as many such linkages (96 and 140, respectively). This difference remains significant after accounting for genes missed through orthology assignment as well as simulations of shorter scaffold sizes (Methods; Supplementary Note 6; Extended Data Fig. 9b). Scans for intragenomic synteny, and doubly conserved synteny with *Lottia*, were performed as described in Supplemental Note 6.

#### Transposable elements and synteny dynamics

5kb up- and down-stream regions of genes were surveyed for transposable element (TE) content. For genes with non-zero TE load, their assignment to either conserved or lost bilaterian synteny in octopus was done using the micro-synteny calculation described above. The number of genes for each category and TE class were as follows: 484 genes for retained synteny and 1,193 genes in lost synteny for all TE classes; 440 and 1,107, respectively, for SINEs; and 116 and 290, respectively, for Mariner. Wilcoxon-U-tests for the difference of TE load in linked vs. non-linked genes were conducted in R.

To assess transposon activity we assigned transcriptome read aligned to 5,496,558 annotated transposon loci using bedtools<sup>44</sup>. Of these, 2,685,265 loci showed expression in at least one of the tissues.

#### **RNA** editing

RNA-Seq reads were mapped to the genome with TopHat<sup>52</sup>, and SAMtools<sup>43</sup> was used to identify SNPs between the genomic and the RNA sequences. To identify polymorphic positions in the genome, SNPs and indels were predicted using GATK HaplotypeCaller version 3.1-1 in discovery mode with a minimum Phred scaled probability score of 30, based on an alignment of the 350 bp and 500 bp genomic fragment libraries using BWA-MEM version 0.7.6a. Using bedtools<sup>44</sup>, we removed SNPs predicted in both the transcriptome and the genome and discarded SNPs that had a Phred score below 40 or were outside of predicted genes. SNPs were binned according to the type of nucleotide change and the direction of transcription. Candidate edited genes were taken as those having SNPs with A-to-G substitutions in the predicted mRNA transcripts.

# Cephalopod-specific genes

Cephalopod novelties were obtained by BLASTP and TBLASTN searches against the whole NR database<sup>53</sup> and a custom database of several mollusc transcriptomes (Supplementary Note 11.1). To ensure that we had as close to full-length sequence as possible, we extended proteins predicted from octopus genomic sequence with our *de novo* assembled

transcriptomes, using the longest match to query NR, transcriptome, and EST sequences from other animals. Gene sequences with transcriptome support but without a match to noncephalopod animals at an evalue cutoff of 1E-3 were considered for further analysis. Octopus sequences with a match of 1E-5 or better to a sequence from another cephalopod were used to construct gene families, which were characterized by their BLAST alignments, HMM, PFAM-A/B, and UNIREF90 hits. The cephalopod-specific gene families are listed in the source data file "cephalopodNovelties.xls". Octopus-specific novelties were defined as sequences with transcriptome support but without any matches to sequences from any other animals (evalue <1e-3), including nautiloid and decapodiform cephalopods.

# Extended Data



# Extended Data Figure 1. RNA editing in octopus

a, Approximate maximum likelihood tree of adenosine deaminases acting on RNA (ADARs) in bilaterians. *ADAR1*, *ADAR2*, *ADAR-like/ADAD*, and *ADAT* (t-RNA specific adenosine deaminase) were identified in Hsa, Mmu, Cin, Dme, Cte, Lgi, *D. opalescens* (Dop<sup>54</sup>), and Obi with Shimodaira-Hasegawa-like support indicated at the nodes. b, *O. bimaculoides* ADAR1, ADAR2 and ADAR-like proteins contain one or two double-stranded RNA binding domains (dsRBD) as well as an adenosine deaminase domain. ADAR1 also has a z-alpha domain. c, Expression profiles of the three *ADAR* genes found in 12 *O. bimaculoides* tissues by RNA-Seq profiling. d, DNA-RNA differences in *O. bimaculoides* show prominent A-to-G changes. Histogram illustrates the number of DNA-RNA differences detected between coding sequences in the genome and 12 *O. bimaculoides* transcriptomes after filtering out polymorphisms identified in genomic sequencing. Differences were binned by the type of change (see key) in the direction of transcription. A-to-G changes are the most prevalent, particularly in neural tissues and during development,

paralleling the expression of octopus *ADARs* in c. Other types of changes were also detected at lower levels, possibly resulting from uncharacterized polymorphisms.



Extended Data Figure 2. Local arrangement of Hox gene complement in *O. bimaculoides* and selected bilaterians

At the top, the four compact Hox clusters of *H. sapiens* and the single *B. floridae* cluster are depicted. The *D. melanogaster* Hox complex is split into two clusters. We included genes in the *D. melanogaster* locus that are homologues of Hox genes but have lost their homeotic function, such as *fushi tarazu (ftz)*, *bicoid*, *zen* and *zen2* (the latter three are represented as overlapping boxes). Hox genes in *C. teleta* are found on three scaffolds<sup>17</sup>. *L. gigantea* has a single cluster with the full known lophotrochozoan gene complement. In *O. bimaculoides* many of the scaffolds are several hundred kb long, and no two Hox genes are on the same scaffold. The positions of *O. bimaculoides* genes approximate their locations on scaffolds. Dashed lines indicate that the scaffold continues beyond what is shown. Scaffold length is depicted to scale with size noted on the left. Genes are positioned to illustrate orthology, which is also highlighted by color.

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#### Extended Data Figure 3. Gene complement and gene architecture evolution in metazoans

a, Principal component analysis of gene family counts. *O. bimaculoides* highlighted in green. Deuterostomes are indicated in blue, ecdysozoans in red, lophotrochozoans in green, and sponges and cnidarians in orange. Xtr: *Xenopus tropicalis*, Gga: *Gallus gallus*, Tca: *Tribolium castaneum*, Dpu: *Daphnia pulex*, Isc: *Ixodes scapularis*, Ava: *Adineta vaga*, Spu: *S. purpuratus*, Hma: *Hydra magnipapillata*, Adi: *Acropora digitifera*. For methods, see Supplementary Note 7.4. b–d, MrBayes<sup>55</sup> tree (constrained topology) on binary characters of presence or absence of Pfam domain architectures (b), introns (c), or indels (d); scale bar represents estimated changes per site. For methods, see Supplementary Note 7.3.





Extended Data Figure 4. Protocadherin genes within a genomic cluster are similar in sequence and sites of expression

a, Expression profile of the 31 protocadherin genes located on Scaffold 30672 in 12 octopus transcriptomes. Over three-quarters of the protocadherins are highly expressed throughout central brain, OL, and ANC, while the others show more mixed distributions. b, Phylogenetic tree highlighting Scaffold 30672 protocadherins in grey bars. c, Expression profile of the 17 protocadherin genes located on Scaffold 9600. Almost all of these protocadherins are most highly expressed in nervous tissues, with the exception of Ocbimv220039316m, which is most highly expressed in the St15 sample. d, Phylogenetic tree highlighting Scaffold 9600 protocadherins in grey bars. As seen in b, protocadherins of the same scaffold tend to cluster together on the tree. Order of the genes in the heatmaps (a, c) follows the ordering on the corresponding scaffold.



# Extended Data Figure 5. Expansion of Interleukin (IL) 17-like genes

a, Phylogenetic tree of interleukin genes in Obi, Cte, Cgi, and Lgi. Mammalian IL1a, IL1b, and IL7 used as outgroups. Human and mouse IL17s branch from other members of the IL family. Octopus ILs (as well as all identified invertebrate ILs) group with the mammalian IL17 branch and are named "IL17-like". The 31 octopus genes are distributed across 5 scaffolds: Scaffold A (Obi\_A), 23 members; Scaffold B (Obi\_B), 4 members; Scaffold C (Obi\_C), 2 members; Scaffolds D (Obi\_D) and E (Obi\_E), 1 member each. b, Expression profile of 31 octopus IL17-like genes. Heatmap rows are arranged by order on each scaffold. Blank rows indicate genes not expressed in our transcriptomes. The 27 genes found in our transcriptomes have strong expression in the suckers and skin. The Scaffold C genes are enriched in the PSG and the Scaffold D gene is enriched in the viscera. c, Conserved cysteine residues in human IL17 and invertebrate IL17-like proteins. The human IL17 proteins share a conserved cysteine motif comprising 4 cysteine residues, which may form interchain disulfide bonds and facilitate dimerization<sup>56</sup>. Octopus IL17-like proteins also contain this 4-cysteine motif, highlighted in yellow. One octopus sequence encodes only 3 of these highly conserved cysteine residues. These four cysteines are also present to varying degrees in Lottia, Capitella, and Crassostrea sequences. Two additional conserved cysteine residues were found in the octopus sequences and are highlighted in red. The first cysteine residue is found in all invertebrate sequences examined, and none of the mammalian IL17 sequences.



#### Extended Data Figure 6. G protein-coupled receptors

GPCRs, also known as 7-transmembrane (7TM) or serpentine receptors, form a large superfamily that activates intracellular second messenger systems upon ligand binding. This figure considers a subset of the 329 GPCRs we identified in *O. bimaculoides*. The full complement of GPCRs is presented in Supplementary Note 8.5. a and b, As reported for other lophotrochozoan genomes, the octopus genome contains chemosensory-like *GPCRs*: 74 GPCRs are similar to the *Aplysia* chemosensory *GPCRs*<sup>57</sup> and 11 *GPCRs* are similar to vertebrate olfactory receptors. c, We identified 4 opsins in the octopus genome (from top to bottom): rhodopsin, rhabdomeric opsin, peropsin, and retinochrome. d, The octopus Class F *GPCRs* comprises 6 genes: 5 Frizzled genes and 1 Smoothened gene (\*). e, Thirty octopus genes show similarity to vertebrate adhesion GPCRs.



#### Extended Data Figure 7. O. bimaculoides acetylcholine receptor (AchR) subunits

a, Phylogenetic tree of AchR subunit genes identified in Hsa, Mmu, Dme, Cte, Lgi, and Obi. Black asterisk indicates a Dme sequence that groups with alpha 1-4-like subunits despite lacking two defining cysteine residues. b, Expression profiles of octopus AchR subunits. Genes ordered as in the tree (a), starting from the gray arrow and continuing counterclockwise. Putative non-Ach binding subunits are highly expressed in the suckers. One sequence was not detected in our transcriptome datasets. In a and b, red asterisks indicate subunits with the substitution known to confer anionic permissivity<sup>58</sup>. c, Divergent octopus subunits lack nearly all residues necessary for Ach binding. Alignment of sequence flanking the cysteine loop (yellow) of the *L. stagnalis* Ach binding protein (Lst\_AchBP), the human and octopus alpha-7 receptor subunits (Has\_AchR7, Obi\_10697+), and the 23 divergent AchR subunits. Essential Ach-binding residues on the primary (pink) and complementary (blue) side of the ligand-binding domain are indicated<sup>26</sup>, with conservative substitutions in a lighter shade. Outside of the binding residues, residues shared between the alpha 7 subunits are shaded in light grey, with bold letters for conservative substitutions.



Extended Data Figure 8. Active transposable elements and gene expression specificity

a, Transposable element expression across 12 tissues. b, Correlation between the total TE load (in bp) in the 5kb regions flanking the gene and the fraction of genes with tissue-specific expression (defined as having at least 75% of expression in a single tissue; Source Data: TELoadAndTissueExpression.xls). p-value indicates F-statistic for the significance of linear regression (H0: r-squared=0), with tissues with a p-value 0.05 indicated in pink.



# Extended Data Figure 9. Synteny dynamics in octopus and the effect of transposable element (TE) expansions

a, Circos plot showing shared synteny across 6 genomes. Individual scaffolds are plotted according to bp length; scaffolds with no synteny are merged together (lighter arcs). Despite the large size of the octopus genome, only a small proportion of the scaffolds show synteny. b, Synteny reduction in octopus quantified based on synteny inference using gene families with at least one representative in human, amphioxus, *Capitella, Helobdella, Octopus, Lottia, Crassostrea, Drosophila,* and *Nematostella. Drosophila, Helobdella,* and *Octopus* show the highest synteny loss rates. Branch lengths, estimated with MrBayes<sup>55</sup>, reflect extent of local genome rearrangement (Supplementary Note 6). c, Enrichment of overall and specific TE classes (base pairs masked) around genes from ancient bilaterian synteny blocks, including those absent in octopus (see key). Asterisks indicates Mann-Whitney U test with p-value < 0.02. d, Transposable element insertion history (Jukes-Cantor distance adjusted, see text) into the vicinity of genes from 'lost' synteny blocks. Notice that only one SINE peak is present; a more recent peak (visible in "All genomic SINEs") cannot be recovered from those insertions.



#### Extended Data Figure 10. Cephalopod phylogeny and novelties

a, Whole-genome-derived phylogeny of molluscs and select other phyla showing the relative position of octopus at the base of the coleoid cephalopods. For methods see Supplementary Note 7.1. Members of the cephalopod class are indicated in blue, scale indicates number of substitutions per site. b, Phylogenetic tree of reflectin genes. Reflectins are cephalopod-specific genes that allow for rapid and reversible changes in iridescence. Six reflectin genes were identified in the octopus genome. c and d, Novel gene expression across multiple tissues. Bars depict all cephalopod novelties; dark grey indicates sequences with no similarity to non-cephalopod genes using HMM searches (Source Data: CephalopodNovelties.xls). c, Counts of tissue-specific novelties in a given tissue. d, Proportion of expression of novel genes versus total expression in individual tissues. CNS (central nervous system) combines Supra, Sub, OL and ANC expression data.

# **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

# Acknowledgments

We thank Titus Brown and Josh Rosenthal for making *Doryteuthis* RNA-Seq data available before publication, Cindy Ha, Jeffrey Orenstein, Jonas Brandenburger, Magdalena Glotzer and Holly Gui for bioinformatic assistance, Shuichi Shigeno for help with tissue dissection, Christine Huffard and Roy Caldwell for providing the *O. bimaculoides* specimen used for genomic DNA isolation, and Emina Begovic for genomic DNA preparation. This work was supported by the Molecular Genetics Unit of the Okinawa Institute of Science and Technology Graduate University (S.B. and D.S.R.) and by funding from the NSF (IOS-1354898) and NIH (R03 HD064887) to C.W.R. and from the NSF (DGE-0903637) to Z.Y.W. This work employed the Vincent J. Coates Genomics Sequencing Laboratory at UC Berkeley, supported by NIH S10 Instrumentation Grants S10RR029668 and S10RR027303, and the University of Chicago Genomics Facility, supported by NIH Grant UL1 TR000430.

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#### Figure 1. Octopus anatomy and gene family representation analysis

a, Schematic of *Octopus bimaculoides* anatomy, highlighting the tissues sampled for transcriptome analysis: viscera (heart, kidney, and hepatopancreas), yellow; gonads (ova or testes), peach; retina, orange; optic lobe (OL), maroon; supraesophageal brain (Supra), bright pink; subesophageal brain (Sub), light pink; posterior salivary gland (PSG), purple; axial nerve cord (ANC), red; suckers, grey; skin, mottled brown; stage 15 (St15) embryo, aquamarine. Skin sampled for transcriptome analysis included the eyespot, shown in light blue. b, C2H2 and protocadherin domain-containing gene families are expanded in octopus. Enriched Pfam domains were identified in lophotrochozoans (green) and molluscs (yellow), including O. bimaculoides (light blue). For a domain to be labeled as expanded in a group, at least 50% of its associated gene families need a corrected p-value of 0.01 against the outgroup average. Some Pfams (e.g., Cadherin and Cadherin\_2) may occur in the same gene, however multiple domains in a given gene were counted only once. Abbreviations used throughout: Obi: O. bimaculoides, Lgi: Lottia gigantea, Pfu: Pinctada fucata, Cgi: Crassostrea gigas, Aca: Aplysia californica, Cte: Capitella teleta, Hro: Helobdella robusta, Dme: Drosophila melanogaster, Cel: Caenorhabditis elegans, Bfl: Branchiostoma floridae, Dre: Danio rerio, Lch: Latimeria chalumnae, Xtr: Xenopus tropicalis, Gga: Gallus gallus, Mmu: Mus musculus, Hsa: Homo sapiens.

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#### Figure 2. Protocadherin expansion in octopus

a, Phylogenetic tree of cadherin genes in Hsa (red), Dme (orange), *Nematostella vectensis* (Nve, mustard yellow), *Amphimedon queenslandica* (Aqu, yellow), Cte (green), Lgi (teal), Obi (blue), and *Saccoglossus kowalevskii* (Sko, purple). I, Type I classical cadherins; II, calsyntenins; III, octopus protocadherin expansion (168 genes); IV, human protocadherin expansion (58 genes); V, dachsous; VI, fat-like; VII, fat; VIII, CELSR; IX, Type II classical cadherins. Asterisk denotes a novel cadherin with over 80 extracellular cadherin domains found in Obi and Cte. b, Scaffold 30672 and Scaffold 9600 contain the two largest clusters of protocadherins, with 31 and 17, respectively. Clustered protocadherins vary greatly in genomic span and are oriented in head-to-tail fashion along each scaffold. c, Expression profiles of 161 protocadherins and 19 cadherins in 12 octopus tissues; 7 protocadherins were not detected in the tissues sampled. Cells are colored according to number of standard deviations from the mean expression level. Protocadherins have high expression in neural tissues. Cadherins generally show a similar expression pattern, with the exception of a group of sucker-specific cadherins.



#### Figure 3. C2H2-ZNF expansion in octopus

a, Genomic organization of the largest C2H2 cluster. Scaffold 19852 contains 58 C2H2 genes that are transcribed in different directions. b, Expression profile of C2H2 genes along Scaffold 19852 in 12 octopus transcriptomes. Neural and developmental transcriptomes show high levels of expression for a majority of these C2H2 genes. In a and b, arrow denotes scaffold orientation. c, Distribution of fourfold synonymous site transversion distance (4DTv) distances between C2H2 domain containing genes.

#### Table 1

# Metazoan developmental control genes

	Obi	Lgi	l Cte	Г Dme	Cel	Bfl	
Ligands							
Fibroblast Growth Factor	3	2	1	3	3	8	22
Wnt	12	10	12	7	5	17	19
TGFβ/BMP	12	9	14	6	5	22	33
Delta/Jagged	4	1	1	2	4	2	7
Hedgehog	1	1	1	1	0	1	3
Axon Guidance	10	9	9	6	8	23	33
<b>Transcription Factors</b>							
C2H2 Zinc Finger	1790	413	222	326	211	1338	764
Homeodomain	114	121	111	104	99	133	333
High Mobility Group	23	15	14	13	16	51	125
Helix Loop Helix	50	63	64	59	42	78	118
Nuclear Hormone Receptor	40	44	45	16	274	33	48
Fox	16	28	26	17	18	42	43
Tbox	9	9	7	8	21	9	18

Number of members of developmental ligand and transcription factor families from *O. bimaculoides*, and selected other taxa. Dendrogram above species names reflects their evolutionary relationships.

#### Table 2

#### Ion Channel Subunits

	<b>—</b>						
	Obi	Aca	Lgi	Cte	Dme	Cel	Hsa
Voltage Gated Calcium Channels	8	8	6	10	9	10	10
Voltage Gated Sodium Channels	3	2	3	2	4	0	13
Transient Receptor Potential Channels	36	45	40	43	13	23	29
K+ Channels							
Voltage Gated	30	23	29	20	10	51	40
Calcium Activated, Small/Large Conductance	12	8	9	6	3	6	8
Inward Rectifying	3	4	5	6	4	3	16
2 Pore	12	9	12	14	11	47	15
Non-Voltage Gated	27	21	26	26	18	72	39
Cys-Loop Receptors							
Glutamate	21	15	47	36	30	15	18
Nicotinic Acetylcholine	53	16	52	77	10	88	16
Inhibitory Acetylcholine	3	2	5	2	0	4	0
5-HT3	0	0	0	0	0	1	5
GABA	6	5	4	9	3	7	19
Glutamate Gated Chloride Channels	7	5	8	5	1	6	0
Glutamate Gated Chloride Channels	7	5	8	5	1	6	0

Number of subunits of representative ion channel families in *O. bimaculoides* and across examined taxa. Dendrogram above species names shows their evolutionary relationships.