

## The brittle star genome illuminates the genetic basis of animal appendage regeneration

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### Abstract

Species within nearly all extant animal lineages are capable of regenerating body parts. However, it remains unclear whether the gene expression programme controlling regeneration is evolutionarily conserved. Brittle stars are a species-rich class of echinoderms with outstanding regenerative abilities, but investigations into the genetic bases of regeneration in this group have been hindered by the limited genomic resources. Here, we report a chromosome-scale genome assembly for the brittle star *Amphiura filiformis*. We show that the brittle star genome is the most rearranged amongst echinoderms sequenced to date, featuring a reorganised Hox cluster reminiscent of the rearrangements observed in sea urchins. In addition, we performed an extensive profiling of gene expression during brittle star adult arm regeneration and identified sequential waves of gene expression governing wound healing, proliferation and differentiation. We conducted comparative transcriptomic analyses with other invertebrate and vertebrate models for appendage regeneration and uncovered hundreds of genes with conserved expression dynamics, particularly during the proliferative phase of regeneration. Our findings emphasise the crucial importance of echinoderms to detect long-range expression conservation between vertebrates and classical invertebrate regeneration model systems.

## Introduction

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Brittle stars are by far the most speciose class of echinoderms; over 2,600 extant species occupy benthic marine habitats globally (Stöhr et al. 2012; O'Hara et al. 2019). However, they remain poorly-documented from a genomic standpoint, despite their broad interest to diverse fields including marine (paleo)ecology, biodiversity monitoring, developmental biology and regenerative

55 biology (Vistisen and Vismann 1997; Vopel et al. 2003; Dupont and Thorndyke 2007; Mosher and Watling 2009; Thuy et al. 2012; Delroisse et al. 2017; Dylus et al. 2018; O'Hara et al. 2019).

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The echinoderm phylum encompasses five classes with a well-resolved phylogeny (Figure 1A; (Cannon et al. 2014; O'Hara et al. 2014; Telford et al. 2014; Mongiardino Koch et al. 2022)): brittle stars (Ophiuroidea), sea stars (Asteroidea), sea urchins (Echinoidea), sea cucumbers (Holothuroidea) and sea lilies/feather stars (Crinoidea). Genomics in this phylum began with the pioneering effort to sequence the genome of the purple sea urchin (*Strongylocentrotus purpuratus*) (Sea Urchin Genome Sequencing Consortium et al. 2006). Analysis of this genome provided broad insights into the evolution of diverse traits and biological processes, including for

65 instance biomineralization, sensory capabilities and immune recognition (Livingston et al. 2006; Raible et al. 2006; Rast et al. 2006). In recent years, the taxonomic sampling of echinoderm genomes has steadily expanded (Hall et al. 2017; Zhang et al. 2017; Davidson et al. 2020; Lawniczak et al. 2021; Davidson et al. 2022; Chen et al. 2023; Marlétaz et al. 2023). This growing wealth of genomic resources in the context of the remarkable diversity of echinoderm

70 body plans, life histories and developmental strategies, provides a unique framework to understand the evolution of novel traits. However, given the deep evolutionary divergence of the five echinoderm classes (480-500 Mya), the lack of robust genomic resources for the brittle stars represents a problematic knowledge gap.

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Adult echinoderms share a characteristic pentameral symmetry, which represents the most derived body plan amongst Bilateria (Smith 2008). Early analyses of sea urchin genomes unveiled local reorganisations within the Hox cluster, prompting speculation that they were associated with the evolution of this unique body plan (Lowe and Wray 1997; Cameron et al. 2006; Mooi and David 2008; David and Mooi 2014). However, the subsequent discovery of

80 an intact Hox cluster in the crown-of-thorns sea star revealed that these rearrangements were not instrumental in the establishment of the pentameral symmetry (Baughman et al. 2014; Byrne et al. 2016). These observations showcase the need to examine a more comprehensive sample of echinoderm whole genomes to accurately identify echinoderm-specific chromosomal rearrangements and subsequently investigate their functional

85 significance.

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Echinoderms exhibit extensive regenerative abilities. Species from each of the five classes are capable of varying levels of regeneration, including (larval) whole-body regeneration, appendage or organ regeneration (Medina-Feliciano and García-Arrarás 2021). Although species within

95 nearly all major animal groups exhibit some regenerative capacity, it is not clear whether this trait is ancestral or independently acquired through convergent evolution (Bely and Nyberg 2010; Lai and Aboobaker 2018; Srivastava 2021). A comparative analysis of whole-body regeneration across a sea star larva, planarian worm and hydra has suggested that broadly-conserved molecular pathways may mediate regeneration (Cary et al. 2019). However, given the diversity of regenerative modes, additional comparative analyses of regenerating organisms are needed to fully understand the evolution of this complex process (Lai and Aboobaker 2018; Srivastava 2021). In particular, gene expression dynamics during regeneration have not been explicitly

100 compared between invertebrates and vertebrates, partly because of the lack of gene expression  
profiling across comparable regenerating structures and of difficulties in identifying orthologs  
among distant model systems. Echinoderms are more closely related to vertebrates than other  
classical invertebrate models of regeneration, hence providing a unique phylogenetic  
perspective. Despite recent studies in sea stars and sea cucumbers (Fumagalli et al. 2018; Cary  
et al. 2019; Quispe-Parra et al. 2021), echinoderms remain largely underrepresented in  
transcriptomic assays of regeneration (Dupont and Thorndyke 2007; Goldman and Poss 2020;  
105 Bideau et al. 2021).

One highly regenerative echinoderm species is the brittle star *Amphiura filiformis*, likely related to  
their sediment dwelling lifestyle where extended arms are often severed by predators. In this  
species, fully differentiated arms regrow in a few weeks following amputation and over 90% of  
110 individuals sampled in the wild display signs of arm regeneration (Duineveld and Van Noort  
1986; Sköld and Rosenberg 1996). Consequently, *A. filiformis* is emerging as a powerful model  
for animal appendage regeneration, with a well-established morphological staging system  
(Dupont and Thorndyke 2006; Czarkwiani et al. 2013; Hu et al. 2014; Purushothaman et al.  
2015; Czarkwiani et al. 2016; Piovani et al. 2021; Czarkwiani et al. 2022). Here, we report a  
115 chromosome-scale genome assembly for the brittle star *A. filiformis* and leverage this unique  
resource to investigate the complex history of karyotypes, Hox cluster and gene family evolution  
across echinoderms. We find that *A. filiformis* displays the most rearranged echinoderm genome  
sequenced to date. Moreover, we reveal that *A. filiformis* extensive regenerative capacities  
correlate with significant expansions of genes involved in wound healing. Finally, we generate  
120 extensive transcriptomic data from regenerating brittle star arms, which we analyse in a  
comparative framework with previously generated datasets from the crustacean *Parhyale*  
*hawaiensis* (Sinigaglia et al. 2022) and the axolotl *Ambystoma mexicanum* (Stewart et al. 2013),  
to illuminate common genetic mechanisms of animal appendage regeneration.

## Results

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### The *Amphiura filiformis* chromosome-scale assembly is a key genomic resource

To address the lack of high-quality genome for the brittle stars (**Supp. Note 1**), we sequenced and assembled the genome of the brittle star *Amphiura filiformis* using high-coverage long nanopore reads assisted with proximity ligation data for scaffolding (**Methods**). The haploid assembly spans 1.57 Gb and contains 20 chromosome-size scaffolds (>60 Mb) that account for 93.5% of the assembly length (**Figure S1**, N50: 68.8 Mb). This is consistent with cytogenetic studies in two other brittle star species that identified 21-chromosome karyotypes (Colombera and Venier 1976; Saotome and Komatsu 2002). We annotated a total of 30,267 protein-coding genes (92.7% complete BUSCO score, **Methods, Table S1, Table S2**), which is in line with the predicted gene complements of other echinoderms (Hall et al. 2017; Li et al. 2020; Davidson et al. 2022; Chen et al. 2023; Marlétaz et al. 2023). In addition, we generated manually-curated lists for *A. filiformis* genes associated with immunity, stemness and neuronal function as well as transcription factors and genes involved in 19 major signalling pathways (**Table S2, Methods**). These lists allow for genome-wide interrogation of *A. filiformis* genes as a complement to gene ontology-based approaches. In summary, the *A. filiformis* genome represents the first high-quality and chromosome-scale genome assembly for the brittle star class, and fills an important knowledge gap in the echinoderm genomics landscape.

### 145 The brittle star exhibits the highest inter-chromosomal rearrangement rate amongst sequenced echinoderms

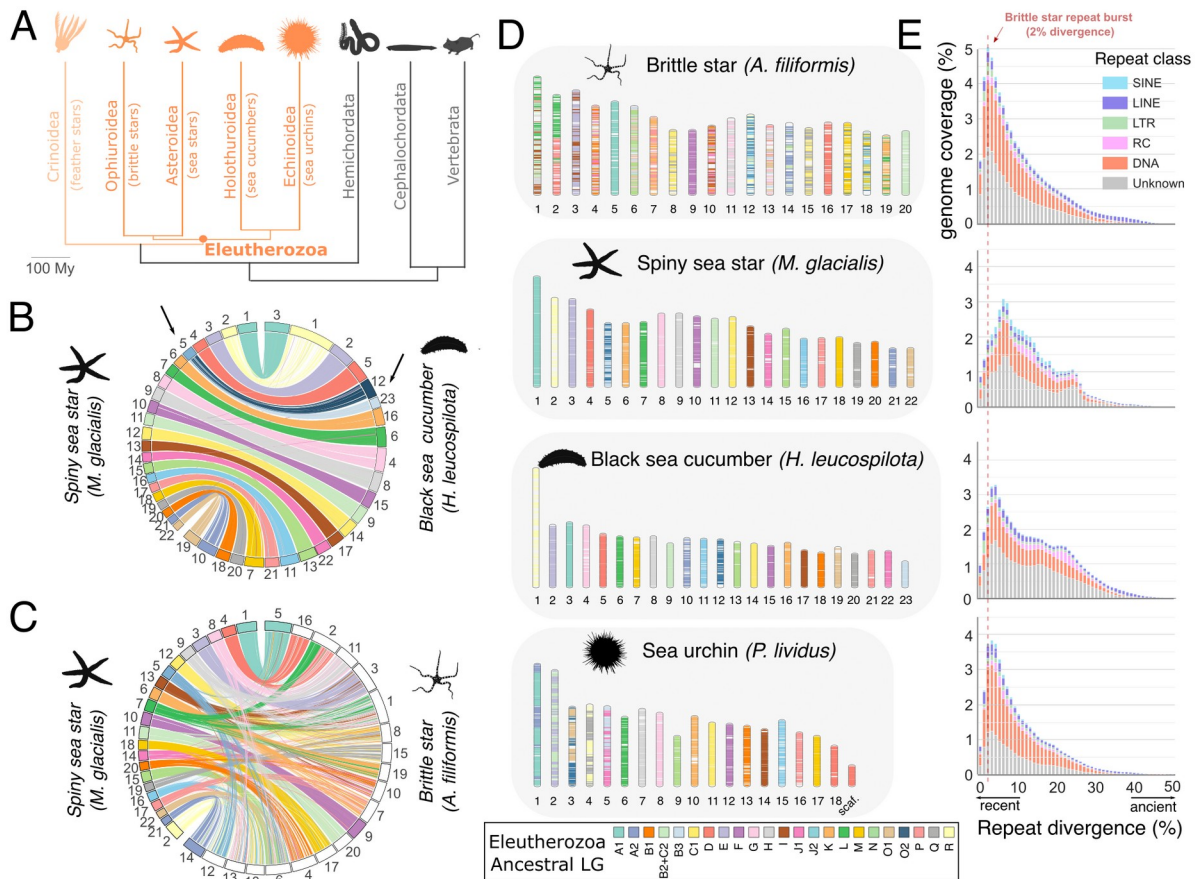
Chromosome evolution in echinoderms has primarily been investigated through the lens of sea urchin genomes, which have globally preserved the ancestral bilaterian chromosomes that were previously reconstructed based on comparisons between chordates and molluscs (Simakov et al. 2020; Simakov et al. 2022; Marlétaz et al. 2023). However, sea urchins also underwent several chromosomal fusions whose origin cannot be established without examining more echinoderm genomes. To pinpoint the timing of these fusions within the context of echinoderm evolution, and to evaluate the conservation of the ancestral chromosomal units across echinoderm lineages, we took advantage of chromosome-scale genomes released for sea stars, sea cucumbers and sea urchins (Lawniczak et al. 2021; Chen et al. 2023; Marlétaz et al. 2023). Using these genomes in combination with the brittle star genome and selected outgroups, we reconstructed the linkage groups present in their ancestor (Eleutherozoa linkage groups, or ELGs, **Figure 1A**).

Synteny comparisons between the spiny sea star (*Marthasterias glacialis*) and the black sea cucumber (*Holothuria leucospilota*) (Lawniczak et al. 2021; Chen et al. 2023) reveal that only one inter-chromosomal macrosyntenic rearrangement occurred in the 500 million years (My) of independent evolution between these two genomes (**Figure 1B, Methods**). In striking contrast, the *A. filiformis* brittle star genome is extensively rearranged: only three chromosomes have a direct one-to-one orthology relationship with spiny sea star chromosomes (**Figure 1C**, Fisher's exact test adjusted  $p$ -value <  $10^{-5}$ ). We reconstructed the ancestral ELGs based on near-perfect conservation of macrosynteny between the spiny sea star and black sea cucumber and using amphioxus (*Branchiostoma floridae*) and sea scallop (*Pecten maximus*) as outgroups to disentangle derived and ancestral chromosomal arrangements (**Figure S2**). We predicted that 23 ELGs were present in the Eleutherozoan ancestor (**Figure 1D**). The ELGs descend from the 24 bilaterian linkage groups (BLGs) (Simakov et al. 2022) through the fusion of the BLGs B2 and C2. Among echinoderms, the black sea cucumber maintained the 23 ancestral ELGs, a single chromosomal fusion took place in the spiny sea star lineage (corresponding to an inter-chromosomal rearrangement rate of 0.002 event / My), five fusions occurred in the sea urchin

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175 *Paracentrotus lividus* (0.01 event / My) and 26 inter-chromosomal rearrangements shaped the  
brittle star *A. filiformis* karyotype (0.052 event / My, the highest amongst sequenced  
echinoderms; **Figure S3**). These results indicate that, among Eleutherozoa, sea cucumbers and  
sea stars show the strongest conservation of ancestral bilaterian linkage groups, whereas the  
brittle star genome is highly reshuffled relative to the Bilaterian ancestor. Examination of  
180 additional sea stars and sea urchins genomes suggest that these trends might broadly extend to  
species within their respective classes ((Davidson et al. 2023; Liu et al. 2023; Marlétaz et al.  
2023); **Figure S3**), but, given the limited sampling, this should be re-examined as more  
chromosome-scale genome assemblies become available.

One potential driver of genomic rearrangements is transposable elements and repetitive  
185 sequences, which serve as substrates for non-allelic homologous recombination (George and  
Alani 2012; Balachandran et al. 2022). Transposable elements have also been implicated in  
driving changes in genome size (Canapa et al. 2015). Among the four echinoderm genomes  
analysed, we find that repetitive elements coverage correlates as expected with genome size but  
not with rates of rearrangements. Repeat coverage is highest in the highly-rearranged brittle star  
190 genome (1.57 Gb, repeat coverage 59.3%) and slowly-evolving black sea cucumber *H.*  
*leucospilota* (1.31 Gb, 56.0%) compared to the sea urchin *P. lividus* (927 Mb, 49.2%) and spiny  
sea star *M. glacialis* (521 Mb, 47.6%). Analysis of sequence divergence reveals that repetitive  
elements accumulated more gradually in the slowly-evolving sea star and sea cucumber  
genomes, compared to both the sea urchin and the brittle star which display recent bursts of  
195 repeat activity (**Figure 1E**). Specifically, the brittle star genome is marked by a burst of repeat  
activity 10-15 Mya, consisting mostly of DNA transposons (peak of repeats with 2% divergence  
to consensus, **Methods**). We thus speculate that the evolutionary history of *A. filiformis* includes  
at least one period of genomic instability (Belyayev 2014). Together, these data highlight  
200 contrasting trends of chromosome evolution across echinoderm classes, and indicate that *A.*  
*filiformis* is the most rearranged echinoderm genome among those sequenced to date.



**Figure 1: Chromosome evolution in echinoderms.** **A.** Phylogenetic relationships of the five echinoderm classes (orange), with the position of the Eleutherozoa ancestor highlighted, and hemichordates and chordates as outgroups. Classes with available chromosome-scale genome assembly are shown in dark orange. Divergence times amongst echinoderms and with hemichordates were extracted from (Mongiardino Koch et al. 2022), divergence with chordates from TimeTree (Kumar et al. 2017). **B.** Synteny comparison between the 22 chromosomes of spiny sea star and the 23 chromosomes of the black sea cucumber. The single macrosyntenic rearrangement between the two genomes is indicated with arrows. **C.** Synteny comparison between the 22 chromosomes of spiny sea star and the 20 chromosomes of brittle star. The three brittle star chromosomes with a 1-to-1 relationship with sea star chromosomes are shown with a colour matching its orthologous counterpart in spiny sea star. **D.** Chromosome evolution in Eleutherozoa. We named the ancestral Eleutherozoa linkage groups (ELG) using established naming conventions proposed for the 24 bilaterian ancestral linkage groups defined in (Simakov et al. 2022; Marlétaž et al. 2023). B2+C2 corresponds to a fusion of bilaterian B2 and C2 present in the Eleutherozoa ancestor. **E.** Repeat landscapes for the brittle star and the three selected echinoderm genomes, with the y-axis representing the genomic coverage and x-axis CpG-corrected Kimura divergence to the repeat consensus. Species are presented in the same order as in **D**. The dashed red line indicates the repeat burst in the brittle star.

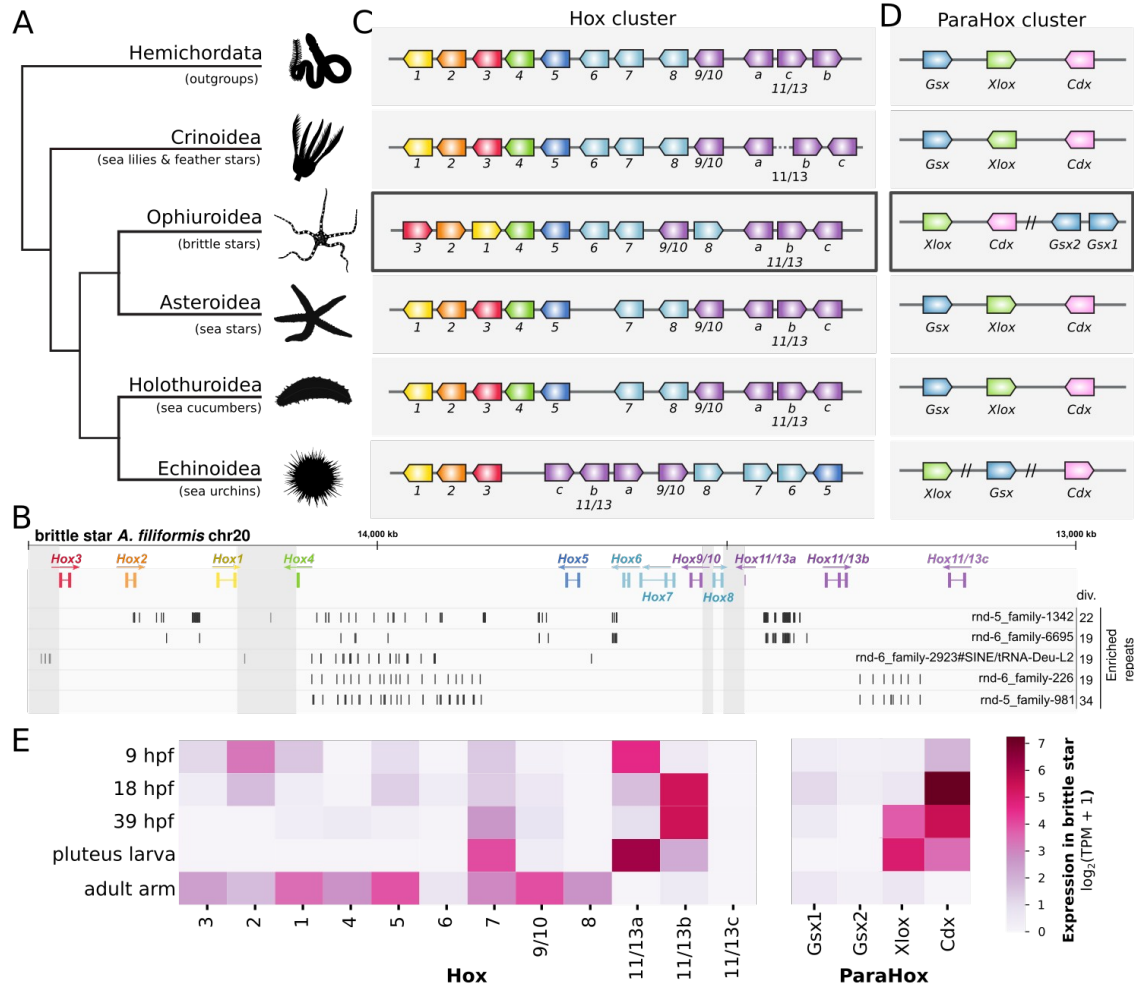
## 220 **The brittle star Hox cluster is marked by small-scale genomic rearrangements**

The organisation of the Hox and ParaHox gene clusters has been documented in each class of echinoderms with the exception of brittle stars (Cameron et al. 2006; Annunziata et al. 2013; Baughman et al. 2014; Zhang et al. 2017; Li et al. 2020). To further explore the enigmatic evolution of these developmental homeobox gene clusters in echinoderms (Byrne et al. 2016), we investigated the structure of the *A. filiformis* Hox and ParaHox clusters. Strikingly, the *A. filiformis* Hox and ParaHox clusters both exhibit genomic rearrangements (**Figure 2, Figure S4, Methods**): anterior Hox genes (*Hox1*, *Hox2* and *Hox3*) are inverted within the 3' end of the cluster and a transposition/inversion event occurred that displaced *Hox8* between *Hox9/10* and *Hox11/13a*. Five repeat families are significantly expanded within the brittle star Hox cluster, of which one is significantly associated with breakpoint locations and may have contributed to the Hox1-3 inversion through non-homologous repair (SINE/tRNA-Deu-L2, BH-corrected permutation-based p-value <0.05, **Figure 2B**). Four out of five of expanded repeats have an inferred divergence of 18-22% to their consensus, suggesting they were active approximately 100 Mya (**Methods**). While brittle star Hox reorganisation is convergent and distinct from the one observed in sea urchins, in both cases the orientation of *Hox1*, *Hox2* and *Hox3* is reversed relative to the other genes of the cluster, and, in both lineages, one of the breakpoints is located near *Hox4* (**Figure 2C**). Moreover, mirroring the evolution of the sea urchins ParaHox genes (**Figure 2D**), the brittle star ParaHox cluster also underwent disruptions, such that *Gsx* was tandemly duplicated to generate two paralogs (protein identity: 74%) located a long distance (>5 Mb) from *Xlox-Cdx*. Whereas *Xlox-Cdx* maintained close linkage in the brittle star, all three members of the ParaHox cluster are dispersed over their chromosome in sea urchins (Annunziata et al. 2013).

Hox expression throughout echinoderm embryogenesis, larval stages and metamorphosis remain largely enigmatic, such that spatio-temporal expression does not follow classical Hox collinearity rules (Arenas-Mena et al. 1998; Byrne et al. 2016). We investigated Hox and ParaHox gene expression in the brittle star using previously published datasets throughout four developmental time points (Delroisse et al. 2014; Delroisse et al. 2015; Dylus et al. 2016) (**Figure 2E, Table S1, Methods**). As in sea urchins (Arenas-Mena et al. 1998), *Hox1* and *Hox3-6* are expressed at very low levels in the brittle star embryos and pluteus larvae (normalised TPM < 2), whereas *Hox7*, *Hox11/13a* and *Hox11/13b* are highly expressed. One notable difference between the expression of Hox genes in the two lineages is seen in *Hox2*. In the brittle star, *Hox2* is expressed early in embryogenesis, with maximal expression at 9 hours post-fertilisation (developmental stage). In contrast, sea urchins *Hox2* is not expressed during early development (Arenas-Mena et al. 1998; Tu et al. 2014). Furthermore, the brittle star ParaHox genes *Xlox* and *Cdx* are each expressed during early development whereas the anterior *Gsx1* and *Gsx2* genes are not (**Figure 2E**), matching the expression patterns observed in sea stars (Annunziata et al. 2013). In contrast, dispersion of the ParaHox cluster in sea urchins is associated with the distinct temporal activation of *Gsx*, *Xlox* and *Cdx* during embryogenesis (Arnone et al. 2006).

These results highlight intriguing parallels in the reorganisation of developmental gene clusters and their expression patterns between brittle stars and sea urchins. Limited data are available on Hox gene expression in other echinoderm classes, but investigations in crinoids and sea cucumbers suggest that, even in species with an intact Hox cluster, the anterior genes (*Hox1-6*) exhibit low or no expression in early embryonic stages (Hara et al. 2006; Kikuchi et al. 2015; Li et al. 2018), and that *Hox7* and *Hox11/13b* may play an important role in embryogenesis (Annunziata et al. 2014). We therefore speculate that the relaxation of expression constraints on Hox genes during echinoderm embryogenesis may have allowed for the rearranged Hox cluster architectures seen in the sea urchins and brittle star lineages.

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**Figure 2: Hox and ParaHox clusters organisation across echinoderms. A.** Phylogenetic relationships between the five classes of echinoderms, with hemichordates as the outgroup. **B.** Genomic organisation of the brittle star *A. filiformis* Hox cluster. Significantly expanded repeats at the Hox cluster location are represented in their respective tracks below Hox genes, with the average sequence divergence to consensus indicated (div., %). Divergence to consensus is a proxy for repeat age, where higher divergence indicates older repeat insertions. Vertical grey rectangles indicate breakpoint locations. **C.** Schematic representation of Hox cluster organisation across echinoderms and outgroups, based on organisation reported in *S. kowalevskii* and *P. flava* (Freeman et al. 2012) for Hemichordata, feather star *A. japonica* (Li et al. 2020) for Crinoidea, brittle star *A. filiformis* for Ophiuroidea, crown-of-thorns sea star *A. planici* (Baughman et al. 2014; Hall et al. 2017) for Asterozoa, Japanese sea cucumber *A. japonicus* (Kikuchi et al. 2015; Zhang et al. 2017) for Holothuroidea and purple sea urchin *S. purpuratus* (Cameron et al. 2006) for Echinozoidea. **D.** ParaHox gene cluster organisation, based on the same genomes as in B. Double slashes indicate non-consecutive genes, all separated by distances > 5 Mb on the same chromosome or scaffold. **E.** Expression of Hox and ParaHox genes throughout 4 brittle star developmental time points and in the adult arm (hpf: hours post-fertilisation). Expression data from (Delroisse et al. 2014; Delroisse et al. 2015; Dylus et al. 2016) was normalised across samples using the TMM method (Robinson et al. 2010) on the full set of brittle star genes, and shown as  $\log_2(\text{TPM} + 1)$ .



## Tandem duplications of key genes contribute to brittle star larval skeleton and bioluminescence

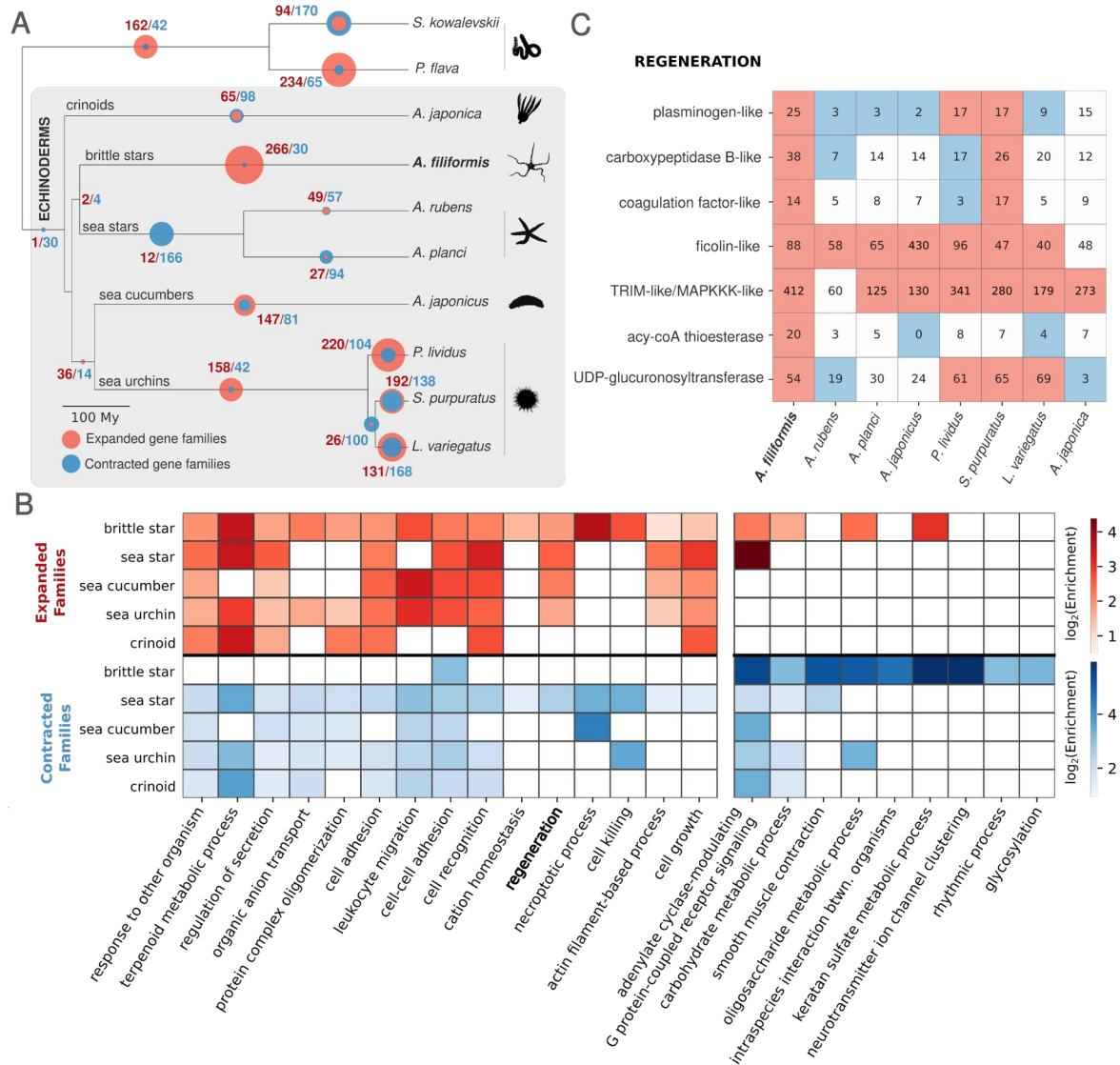
Tandem gene duplications and subsequent asymmetric divergence are widespread in the evolution of animal genomes and have been linked to the evolution of species-specific traits (Holland et al. 2017). In echinoderms, two specific gene families represent relevant examples of lineage-specific evolution through tandem duplications: *phb/pmar1* (larval skeleton) and luciferases (bioluminescence) (Dylus et al. 2016; Delroisse et al. 2017; Marlétaz et al. 2023). Pmar1 is the most upstream zygotic factor of the regulatory network controlling the specification of skeletogenic cells in sea urchins (Oliveri et al. 2003; Oliveri et al. 2008). Among Eleutherozoa, only sea urchins and brittle stars develop an elaborated larval skeleton. In sea urchins, the *pmar1* gene originated through repeated lineage-specific duplications of an ancient *phb* paired-class homeobox gene. Duplications of the *pmar1* gene have been pinpointed as important drivers for the establishment of this sea urchin-specific regulatory programme, which culminates in the formation of the larval skeleton (Dylus et al. 2016; Yamazaki et al. 2020; Marlétaz et al. 2023). In the brittle star *A. filiformis*, we identify a similar expansion of *phb* paralogs (totaling 13 *phb* genes). Phylogenetic analysis confirms that these *phb* paralogs are distant homologs of the sea urchins *pmar1*, but indicates that they are distinct from the previously described brittle star *pplx* gene (Dylus et al. 2016) (**Figure S5A**). Moreover, expression of the *A. filiformis phb* genes occur largely during early development (**Figure S5E**), as described for the *pmar1* sea urchin gene. This suggests that the convergent evolution of brittle stars and sea urchins larval skeleton may have been driven by independent duplications of *phb* genes.

Brittle stars stand out by their ability to emit light (Mallefet 2009). In *A. filiformis*, bioluminescence is mediated by a specific type of luciferase which is homologous to the well-characterised luciferase of the soft coral *Renilla reniformis* (Delroisse et al. 2017). Within the *A. filiformis* genome, we identified nine luciferase-like gene copies: seven are organised in two clusters of tandem duplicates, two are isolated copies. This corroborates and extends the previously-inferred repertoire (Delroisse et al. 2017). We find that luciferase-like genes have duplicated not only in the brittle star but also in all echinoderm lineages with the exception of sea stars (**Figure S5**). These results confirm previous propositions that echinoderms harbour multiple copies of luciferase-like genes, which likely encode diverse functions across bioluminescent and non-bioluminescent species (Delroisse et al. 2017).

## Expanded gene families in echinoderms are enriched in regeneration-related processes

To comprehensively assess the functional significance of gene complement evolution, we inferred gene family expansion and contraction events within echinoderms (**Figure 3A, Methods**). In contrast with other deuterostome lineages, which exhibit either extensive gene losses (Seo et al. 2001) or duplications (Putnam et al. 2008), we found that echinoderms harbour relatively stable gene complements, with only ~7.6% of gene families showing significant expansions or contractions in their ancestral lineage and throughout their evolution (790 of 10,367 tested families). Within the brittle star, genes in these families are enriched in GO terms that are also found in the expanded and contracted families of other echinoderm classes (**Figure 3B, Table S3, Methods**). This includes several enriched GO terms linked to immune-related processes (e.g. “response to other organisms”, “leukocyte migration”, “cell recognition”), which encompass genes known to display elevated gene birth and death rates in other animal lineages, such as Toll-Like Receptors (Nei et al. 1997; Leulier and Lemaitre 2008; Saco et al. 2023). Some GO term enrichments may reflect specific aspects of echinoderm biology. For instance, recurrent duplications of “regeneration-related” genes may underlie the remarkable regenerative capacity

of many echinoderm species. Notably, in *A. filiformis*, members of these expanded gene families (**Figure 3C**) are expressed during arm regeneration (**Figure S6**). Additionally, genes within four expanded families (*plasminogen*, *carboxypeptidase B*, *coagulation factor* and *ficolin*) directly regulate coagulation and/or clotting in vertebrates (Pryzdial et al. 2022), but may play a broader role in immune defence in echinoderms (Hanington and Zhang 2011; Loof et al. 2011).  
345 Moreover, the *ficolin* gene has also been implicated in the early stages of *A. filiformis* arm regeneration (Ferrario et al. 2018; Arenas Gómez et al. 2020). Duplications within the brittle star may thus have contributed to the evolution of a rapid and efficient wound closure process that is prerequisite to regeneration (Suárez-Álvarez et al. 2016; Ferrario et al. 2018). Finally, genes  
350 involved in keratan sulfate metabolism are over-represented in both expanded and contracted gene families in the brittle star (**Figure 3B**). Increased sulfated glycosaminoglycans production is required for proper arm regeneration in *A. filiformis* (Ramachandra et al. 2017). The numerous duplications and losses of these genes suggest that the evolution of brittle star efficient regeneration may have been accompanied by a specialisation of the glycosaminoglycan  
355 metabolism.



360 **Figure 3: Gene family evolution in echinoderms. A.** Number of significantly expanded (red) and  
 365 contracted (blue) gene families throughout echinoderm evolution, from a total of 10,367 tested gene  
 families (**Methods**). **B.** Gene ontology functional enrichment tests (Biological Process) for expanded  
 and contracted families in the different echinoderm classes. We selected the top 15 representative  
 370 GO terms enriched in the expanded brittle star gene families and 10 in contracted families (**Methods**).  
 In the heatmap, colours indicated GO terms significantly enriched in expanded or contracted families  
 in other echinoderm classes (FDR < 0.05). **C.** Gene copy number variation across echinoderms for  
 regeneration gene families with significant expansion in *A. filiformis* (>1 brittle star gene in the family  
 annotated with the GO term 'regeneration'). Gene families were named according to the *S.*  
*purpuratus* gene name. Red and blue colours denote significantly expanded and contracted families,  
 respectively.

## Gene expression during brittle star arm regeneration recapitulates major regeneration phases

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To gain insight into the transcriptional programmes that underlie brittle star arm regeneration, we profiled gene expression in seven representative regeneration stages following amputation and one non-regenerating control (**Methods**). Using soft-clustering, we classified genes into nine major temporal clusters (A1-A9; **Figure 4A**, **Figure S6**, **Methods**). Functional enrichment analysis of genes within the co-expression clusters revealed three distinct phases of arm regeneration: (1) wound healing, (2) proliferation and (3) tissue differentiation. These are consistent with morphological timelines of regeneration in the brittle star and other animal systems (Czarkwiani et al. 2016; Bideau et al. 2021; Srivastava 2021).

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Early regeneration is marked by the expression of genes involved in wound response, including immunity/wound healing (clusters A1-A2), and cell migration/tissue protection (clusters A3-A4), which are enriched in immune and kinase genes, respectively (**Figure 4B**, **4C**). Notably, the regions surrounding transcription start sites (TSS) of genes within cluster A2 are enriched for transcription factor (TF) binding motifs of NF- $\kappa$ B, a broadly conserved regulator of immune response (**Figure 4D**).

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Wound healing is followed by cell proliferation (clusters A9 and A5, A6, A7), as indicated by the overrepresentation of stemness genes and genes involved in cell proliferation, cell division and enhanced translational activity. Accordingly, binding motifs associated with several proliferation-related TF are enriched around the TSS of genes from clusters A5 and A6. This includes NRF1 and p53, which have been implicated in vertebrates in regulating (stem) cell survival and proliferation (Cui et al. 2021; Ayaz et al. 2022), PRDM14 and YY1, which regulate pluripotency (Kawaguchi et al. 2019; Dong et al. 2022), and RORa, which controls inflammation by down-regulating targets of NF- $\kappa$ B (Oh et al. 2019) and may thus play a role in the transition from wound response to proliferation (**Figure 4C**, **4D**). We also find enrichment of binding motifs corresponding to zinc-finger transcription factors that are involved in cell proliferation and pluripotency (Villot et al. 2021; Han et al. 2023). Cluster A9 encompasses genes expressed as early as 48 hours post-amputation (hpa) and active throughout regeneration, including translational regulators, cell division and vesicle transport genes (**Figure 4B**), as well as genes involved in signalling pathways that promote cell proliferation (VEGF, Akt, Insulin-like and Jak-STAT pathways, **Figure 4C**, **Figure S6**) (Xu et al. 2012; Huat et al. 2014; Apte et al. 2019; Herrera and Bach 2019). These data suggest that the signalling cascades that initiate cell proliferation are induced very early during brittle star regeneration (cluster A9); they are activated during the wound response phase and exhibit amplified expression during the peak of cell proliferation (Stage 5; **Figure 4A**). The early onset of proliferation (around 48 hpa) is consistent with previous observations of cell proliferation and expression quantification of selected marker genes (Czarkwiani et al. 2016; Czarkwiani et al. 2022).

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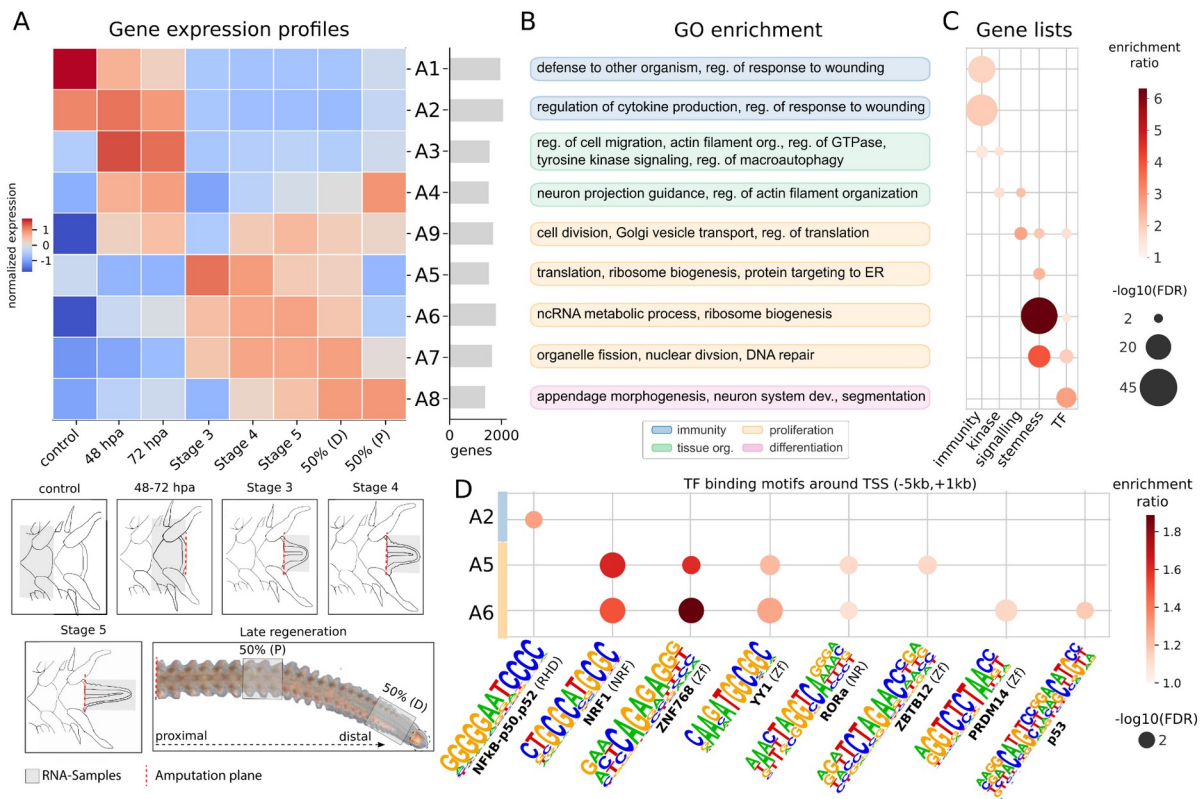
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Finally, late regeneration is characterised by the expression of genes involved in differentiation, patterning and appendage morphogenesis, with a significant over-representation of transcription factors (cluster A8, **Figure 4B**, **4C**). This cluster includes two T-box TFs that are important for axis specification in echinoderms (*tbx3-1* and *tbx3-2*) and two TFs with key roles in neurogenesis (*ngn1-like* and *hey1-like*) (Gross et al. 2003; Slota and McClay 2018; Slota et al. 2019).

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Overall, these data provide a genome-wide picture of the molecular pathways at play throughout brittle star arm regeneration and highlight three waves of gene expression that successively mobilise genes involved in wound response, cell proliferation and tissue differentiation. These general phases have been described in many regenerating animals. Consequently, this dataset

425 can be leveraged to assess the conservation of regeneration gene expression dynamics across species.



430 **Figure 4: Gene expression during brittle star arm regeneration.** **A.** Soft-clustering of gene  
 expression profiles throughout regeneration time points, yielding 9 main temporal co-expression  
 clusters (A1-A9, **Methods**, see also **Figure S6**). Co-expression clusters are temporally ordered (from  
 435 top to bottom) on the basis of their first expression time point. Barplots on the right indicate the  
 number of genes assigned to each cluster. The RNA-sampling procedure for each stage is illustrated  
 below. Early stages are sampled at 48 and 72 hpa (hours post-amputation). Subsequent stages are  
 defined by morphological landmarks: Stage 3 corresponds to the presence of the radial nerve (~6  
 days post-amputation, dpa), Stage 4 is the appearance of the first regenerated metameric units (~8  
 440 dpa), Stage 5 corresponds to advanced arm extension and differentiation onset (~9 dpa), 50% stages  
 correspond to when 50% of the regenerated arm has differentiated (~2-3 weeks post-amputation)  
 sampled at the distal (D, less differentiated) and proximal (P, more differentiated) ends (Dupont and  
 Thorndyke 2006; Czarkwiani et al. 2016). **B.** Gene ontology enrichment for each co-expression cluster  
 (**Methods**, see **Figure S7** and **Table S4** for exhaustive GO results). **C.** Curated gene lists enrichment  
 for each co-expression cluster (hypergeometric test, Benjamini-Hochberg adjusted p-values <0.05,  
 445 **Methods**, **Table S2**). **D.** TF binding motifs enriched around the TSS (5kb upstream to 1kb  
 downstream) of genes from co-expression clusters (hypergeometric test adjusted p-value <0.05,  
**Methods**).

## Animal appendage regeneration is characterised by conserved gene expression during the cell proliferation phase

450 Several key genes and pathways have been repeatedly implicated in regeneration across animal lineages (Bideau et al. 2021; Srivastava 2021), However, direct comparisons of temporal expression gene profiles throughout regeneration remain limited. Here, we investigate the conservation of the animal appendage regeneration programmes through the lens of the brittle star *A. filiformis*.

455 Using a genomic phylostratigraphy approach (Barrera-Redondo et al. 2023), we found that, overall, brittle star arm regeneration is mediated by ancient genes (i.e. metazoan or older; **Figure 5A, Methods**). The exception is the initial wound-healing phase, which is enriched in genes that are specific to the brittle star lineage (**Figure 5A**). The observation that brittle star regeneration is mostly driven by ancient genes prompted us to investigate whether these genes are similarly  
460 involved in appendage regeneration across animals, and whether they are deployed in the same temporal order. As an invertebrate deuterostome, brittle stars enable phylogenetic comparisons among vertebrate and ecdysozoa appendage regeneration models. We thus compared gene expression dynamics during appendage regeneration in *A. filiformis* with comparable datasets from the axolotl (*Ambystoma mexicanum*) (Stewart et al. 2013) and the crustacean Parhyale  
465 (*Parhyale hawaiiensis*) (Sinigaglia et al. 2022). For this analysis, we defined nine major co-expression clusters during axolotl limb regeneration (Ax1-Ax9, **Figure S8, Table S5**) that effectively recapitulate the three regeneration phases (wound healing, proliferation and differentiation), and used existing Parhyale clustering (Sinigaglia et al. 2022).

470 Pairwise comparisons and permutation tests reveal that many of the co-expression clusters employ homologous genes during appendage regeneration across the three species (**Figure 5B, Methods**). Among the nine co-expression clusters that mediate brittle star regeneration, five consist of genes that are also co-expressed during axolotl regeneration (926 genes), six clusters overlap with Parhyale (913 genes), and four clusters are consistent across the three species  
475 (154 genes) (**Figure 5B, 5C, Table S6**). Expression comparisons between the more phylogenetically distant axolotl and Parhyale identify only two conserved co-expressed gene clusters (370 axolotl genes); this direct comparison is thus considerably less informative than comparisons that include the brittle star. Most genes with conserved expression patterns in the brittle star/axolotl comparison lack identifiable homologs in Parhyale, whereas genes with a  
480 conserved expression in the brittle star/Parhyale comparison exhibit a different expression pattern in the axolotl (**Figure 5C**). This underscores the relevance of using the brittle star to bridge comparisons across established regeneration models.

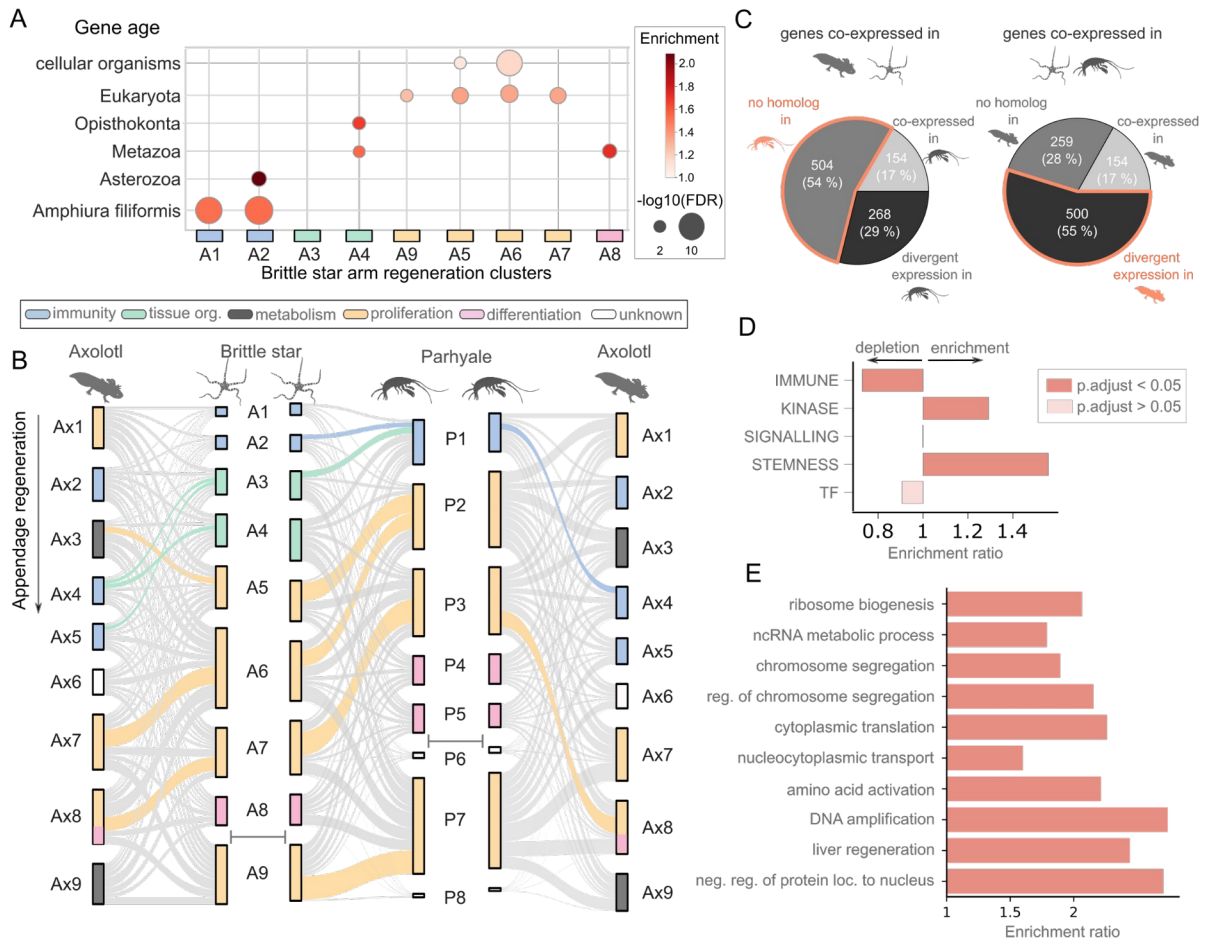
The broadly-conserved co-expression clusters largely consist of genes expressed during the proliferative phase, and, to a lesser extent, the initial wound healing phase. In contrast, the genes that comprise clusters corresponding to tissue differentiation are distinct in each species, which is consistent with the fact that the regenerating appendages are not homologous across species. Strikingly, the conserved co-expression clusters are deployed in a consistent temporal sequence in each species (**Figure 5C**). Specifically, the only identified heterochrony concerns the matching  
490 of the axolotl cluster Ax3 (peak at 0-3 hpa) with brittle star cluster A5 (peak at 6 dpa, **Figure 5C, Figure 4A, Figure S8**). Previous work suggested that similar co-expression gene modules are deployed during regeneration and development, but are activated according to distinct temporal sequences (Sinigaglia et al. 2022). To investigate if this extends across species, we compared gene expression profiles during regeneration and development from the brittle star and Parhyale,  
495 notwithstanding the fact that the indirect development of brittle stars does not allow a direct comparison (**Figure S9**). Results indicate that the order in which co-expressed gene modules are

activated is as expected more conserved within regeneration and within developmental datasets across species than between regeneration and development in individual species (**Figure S9**). Together, these results broaden previous observations of distinct expression dynamics during development and regeneration, and document conserved gene expression modules recruited for animal appendage regeneration (**Table S6**).

We further investigated the functions of brittle star genes with similar temporal expression profiles during regeneration in Parhyale and/or axolotl. Using a carefully selected background that accounts for homology-detection and functional biases of different clusters (**Methods**), we found that, within the set of genes that exhibit conserved expression in regeneration, there is a significant over-representation of kinase and stemness genes and an under-representation of immune genes (Gene lists enrichment tests, **Figure 4D**, **Figure S9**). Moreover, these genes are enriched in general biological processes related to cell proliferation, such as translation, chromosome segregation, DNA replication and intracellular transport (Gene Ontology enrichment tests, **Figure 4E**). The temporal expression patterns of genes that encode transcription factors are neither significantly more nor less conserved across species than is expected at random. Among the conservative set of 154 genes with conserved expression profiles across the three species, only two transcription factors emerge (**Table S6**): *Id2-like*, which activates regeneration-induced proliferation in mice (Kiyokawa et al. 2021) and *Wdhd1-like*, which regulates DNA replication (Zhou and Chen 2021). We thus propose that *Id2* and *Wdhd1* may play a conserved role during animal regeneration. In addition, while several TF binding motifs found in the vicinity of brittle star co-expressed genes are also over-represented near Parhyale and axolotl co-expressed genes, only YY1 and NRF1 are present in corresponding co-expression clusters (Ax7-A6, **Figure S8**), suggesting a possible conserved role for these transcription factors in regulating cell proliferation during regeneration in these distantly related organisms.

Finally, we find that two temporally-matched gene expression clusters in brittle star and Parhyale regeneration include key genes involved in repressing transposable elements (i.e. *Risc-like* [A2 - P1] and *Ago2-like* [A9 - P7]; **Table S6**). It has been proposed that transposon repression is important for proceeding from the immune response phase to regeneration (Angileri et al. 2022), by preserving genome integrity for cell proliferation and differentiation. In line with this hypothesis, we found a higher transcriptional activity of brittle star repetitive elements in the initial wound-response regeneration phase compared to the proliferative phase (**Figure S11**, **Methods**). Repetitive sequence divergence analysis indicates that these repetitive elements are significantly younger and are more frequently present in intergenic regions than those expressed during cell proliferation, suggesting a higher potential to effectively be active transposable elements.

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**Figure 5: Gene expression throughout appendage regeneration across animals. A.** Gene age enrichments for brittle star arm regeneration clusters (hypergeometric test, Benjamini-Hochberg adjusted p-values <0.05). Clusters are ordered by the time of expression onset. **B.** Comparison of co-expressed gene clusters deployed during appendage regeneration in axolotl, brittle star and Parhyale

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(left to right: axolotl vs brittle star, brittle star vs Parhyale, Parhyale vs axolotl). Clusters in Parhyale (clusters P1 to P8) correspond to the clustering reported in (Sinigaglia et al. 2022), but clusters were renamed to follow temporal activation and homogenise with respect to brittle star and axolotl clusters (**Methods**). Co-expression clusters in each species are shown in order of their temporal expression (from top to bottom), with the exception of brittle star cluster A9 and Parhyale clusters P6-P7-P8 that are expressed throughout several regeneration time points and shown at the bottom. Clusters are represented by vertical rectangles whose sizes are proportional to the number of homologous genes in the cluster, and coloured according to enriched GO terms (**Methods, Figure 4, Figure S8, see A** for legend). Links between clusters of the two compared species indicate cluster membership of homologous genes, with coloured links indicating significant overlaps (permutation-based p-values with Benjamini-Hochberg correction <0.05, **Methods**).

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The majority of genes identified as co-expressed in the brittle star - Parhyale and brittle star - axolotl comparisons are not recovered in the direct Parhyale - axolotl comparison. The majority of genes co-expressed in the axolotl and brittle star have no identified homologs in Parhyale (54%, left pie chart). Genes co-expressed in Parhyale and the brittle star have a divergent expression in the axolotl, i.e. they are not found in matched co-expression clusters (55%, right pie chart). **D.** Gene lists enrichment and depletion tests performed for the set of brittle star genes with conserved temporal expression during animal regeneration (**Methods**). **E.** Gene ontology enrichment tests, as in **D**.

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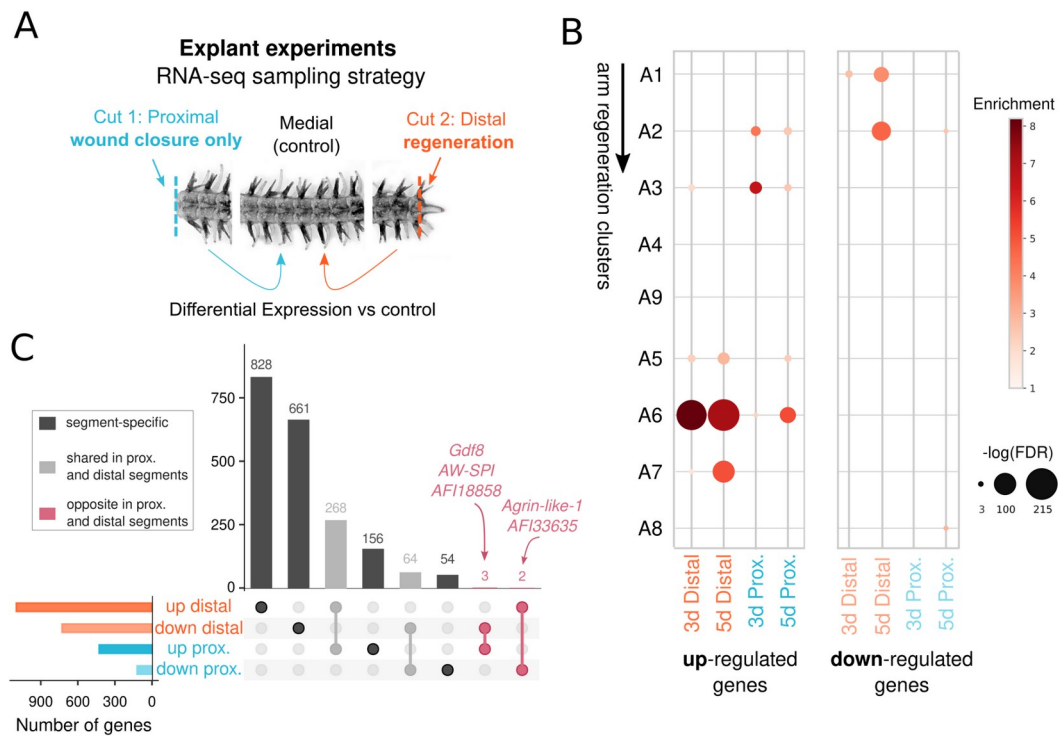


565 **Explant experiments reveal expression differences between non-regenerative and regenerative responses in the brittle star**

To define what differentiates regeneration from non-regenerative wound healing at the molecular level, we performed explant experiments in which the arm is first amputated from the body (proximal cut) and subsequently amputated a second time at the distal end (**Figure 6A**). As in whole animals, explanted brittle star arms regenerate from the distal tip, whereas the proximal end undergoes a non-regenerative wound healing response. To identify genes specifically involved in regeneration, we sampled distal, medial and proximal explant segments for RNA-seq experiments at 3 and 5 days post-amputation (dpa) when morphological differences start to become apparent (3 to 4 replicates each, for a total of 20 samples, **Figure 6A, Table S1**).

575 We tested for differential expression of genes at the distal and proximal end compared to control medial segments (**Methods, Figure 6A**). As expected, up-regulated distal genes correspond to genes expressed during the proliferative phase of the brittle star arm regeneration time-series, whereas up-regulated genes in proximal segments correspond to early response/wound closure genes (**Figure 6B**). We identified more differentially expressed genes (DEGs) in the distal regenerating samples than in proximal non-regenerating samples (distal: 595 and 828 up-regulated genes at 3 and 5 dpa respectively, 238 and 562 down; proximal: 148 and 373 up, 27 and 97 down; **Figure 6C**). Most genes differentially expressed in proximal segments are also differentially expressed in distal segments (61% of the proximal DEGs are shared with distal), whereas distal genes are largely distal-specific (82% of the distal DEGs are not shared with proximal) (**Figure 6C**). This is consistent with the expected expression patterns, as wound closure is an integral part of regeneration. Altogether, we identify hundreds of differentially expressed candidate genes, which document the genetic commonalities and differences between non-regenerative wound closure and regeneration (**Table S2**).

590 Strikingly, five genes display drastically opposite expression patterns in the wound healing and regenerating segments (**Figure 6C**) and thus are likely to contribute to distinct post-wounding outcomes. *Agrin-like-1* and *AFI33635* are significantly down-regulated during wound healing but up-regulated in regeneration (**Figure 6C**). Agrin proteins are critical for neuromuscular junction development in vertebrate embryogenesis (Hoch 1999). *AFI33635* is an uncharacterized brittle star gene with thyroglobulin and methyltransferase domains, putatively involved in regulating protease activity (Novinec et al. 2006). Conversely, the three genes *AW-SPI*, *AFI18858* and *Gdf8* are significantly up-regulated during wound healing but down-regulated in regeneration (**Figure 6C**). *AW-SPI* is an antistasin/WAP-like serine protease inhibitor, with a possible role in immune defence (Yan et al. 2016). *AFI18858* is a brittle star gene with a zf-Bbox domain, and is a member of the expanded TRIM-like gene family, broadly involved in immune responses (**Figure 3C**). Interestingly, the myostatin gene *Gdf8* is a member of the TGF-beta signalling pathway that inhibits skeletal muscle growth and regeneration in mice (McCroskery et al. 2005; Elkasrawy and Hamrick 2010). Our findings suggest that repression of *Gdf8* may similarly enable muscle regeneration in brittle stars. In summary, these five candidate genes might be tightly linked with the transition from wound healing to regeneration-induced cell proliferation, and some may have a conserved function in the brittle star and vertebrates (*Agrin* and *Gdf8*).



**Figure 6: Comparison of gene expression during wound closure and regeneration in brittle star explant experiments.** **A.** Experimental setup. Brittle star arms are amputated at the proximal (cut 1) and distal (cut 2) ends. Proximal, distal and medial (control) segments are sampled for RNA-seq at 3 and 5 days post-amputation (dpa). We identify differentially expressed genes (DEGs) in proximal (wound closure only, not followed by regeneration) segments and distal (regenerative) segments, compared to control medial segments. **B.** Comparison of DEGs from explant experiments with brittle star arm regeneration time-course clusters (see **Figure 4**, hypergeometric enrichment test, BH-corrected  $p$ -values  $<0.05$ ). **C.** Overlap between DEGs genes in distal and proximal segments. Bars in the UpSet are coloured to highlight (i) segment-specific DEGs, for DEGs unique to distal or proximal segments, (ii) shared prox. and distal segments, for DEGs shared between proximal and distal, and (iii) opposite prox. and distal segments, for DEGs up-regulated in proximal and down-regulated in distal (or vice-versa).

## Discussion

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The chromosome-scale genome of the brittle star *Amphiura filiformis* represents a critical resource for the fields of evolutionary genomics, marine ecology and regenerative biology. We leveraged this novel genome to gain fundamental insights into echinoderm chromosome evolution, Hox cluster organisation and regenerative processes across animals. Analyses of chromosome evolution have been key to understanding the basic principles of genome evolution, particularly those linked to the emergence of metazoan and bilaterian clades (Simakov et al. 2020; Simakov et al. 2022). Whereas previous studies of chromosome evolution in echinoderms were limited to sea urchins (Simakov et al. 2020; Marlétaz et al. 2023), our analyses revealed that the genomes of sea cucumbers and sea stars display even fewer rearrangements of the bilaterian ancestral chromosomal units than sea urchins. Interestingly, sea cucumbers have the lowest rate of inter-chromosomal rearrangements, yet the most derived echinoderm body plan (Rahman et al. 2019), which highlights the uncoupling of global genomic rearrangements from morphological evolution. We showed that the ‘Eleutherozoa Linkage Groups’ descend from a single fusion of ancestral bilaterian linkages (B2+C2). Chromosome-scale crinoid and hemichordate genomes will reveal whether this fusion is ancestral to Ambulacraria (the clade encompassing echinoderms and hemichordates). Critically, the fusion has not been observed in the genome of *Xenoturbella bocki*, a member of the Xenacoelomorpha lineage whose phylogenetic position is controversial, and thus cannot be used to support their proposed grouping with Ambulacraria (Philippe et al. 2019; Schiffer et al. 2022). In contrast with its sea star sister-group, the *A. filiformis* genome is highly rearranged: our analyses identified 26 inter-chromosomal rearrangements since the Eleutherozoa ancestor. A more comprehensive sampling of brittle star genomes will provide additional context toward establishing the precise timeline of chromosomal rearrangements and investigate the relative contributions of repeat expansion, chromatin architecture and population genetics dynamics to the rapid karyotype evolution in this group.

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On a more local scale, we identified convergent rearrangements in the Hox clusters of sea urchins and the brittle star, which could be hallmarks of relaxed regulatory constraints within echinoderms, perhaps resulting from the temporal decoupling of embryo and adult patterning. Specifically, Hox genes, and in particular anterior Hox, show limited expression during echinoderm embryogenesis and are mostly expressed in adults, suggesting that they are mostly used to pattern adult structures (Arenas-Mena et al. 1998; Arenas-Mena et al. 2000; Hara et al. 2006; Kikuchi et al. 2015; Li et al. 2018). In this context, we speculate that anterior and central/posterior Hox genes may belong to distinct chromatin compartments in echinoderms. Small-scale rearrangements may have occurred through elevated physical contacts at compartments boundaries (i.e. around *Hox4*), and eventually become fixed due to relaxed selection constraints on Hox expression. Further study of chromatin conformation and regulatory footprint in rearranged and non-rearranged echinoderms will make it possible to test this hypothesis. In addition, we revealed significant expansions of transposable elements in the brittle star Hox cluster ~100 Mya. In the event that Hox cluster rearrangements co-occurred with the activation of repeats, distantly related brittle star species (O’Hara et al. 2014) may exhibit distinct Hox organisations.

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The brittle star genome furthermore enables genetic characterization of the animal appendage regeneration process, and remarkably allows the detection of long-range conservation of the gene expression programme that regulates regeneration. In particular, incorporating the brittle star within a comparative transcriptomics framework extensively increased our ability to detect conserved co-expression modules between vertebrates (e.g. axolotl) and arthropods (e.g.

675 Parhyale). We revealed that the proliferative phase of regeneration displays the highest  
expression conservation across these animals, suggesting that regeneration deploys an ancient,  
evolutionarily conserved proliferation machinery. This observation ties in with the proposition that  
animal regeneration may recruit a homologous proliferating cell type (Lai and Aboobaker 2018;  
Srivastava 2021), a hypothesis that should be further explored with single-cell sequencing  
680 techniques and additional comparative analyses. The stronger conservation of gene expression  
during proliferation as opposed to the initial wound healing response is moreover consistent with  
the elevated turnover of immunity-related genes, broadly reported across animal lineages (Nei et  
al. 1997; Leulier and Lemaitre 2008; Saco et al. 2023) and which we also demonstrate here in  
echinoderms. Our results however contrast with the only previous study to have explicitly  
685 interrogated the conservation of animal regeneration gene expression programmes, which  
revealed a higher conservation of early response genes as opposed to the genes expressed  
during proliferation (Cary et al. 2019). These discrepancies might be due to asynchronous  
temporal sampling across species in the comparisons of (Cary et al. 2019), which is alleviated in  
our study through more comprehensive samplings of regeneration time points and explicit  
690 comparisons of temporal expression profiles. Alternatively, they could reflect genuine biological  
differences of (larval) whole body regeneration studied in (Cary et al. 2019) and the adult  
appendage regeneration we investigate here.

Finally, in the brittle star *A. filiformis*, we uncover significant expansions of gene families linked to  
regeneration-related processes and in particular of homologs of vertebrate coagulation regulator  
695 genes, suggesting them as relevant candidates for follow-up in-depth functional  
characterizations. We also propose a conserved role for *Gdf8* during regeneration, as it is  
repressed during regenerative proliferation in both brittle stars and mice (McCroskery et al. 2005;  
Elkasrawy and Hamrick 2010). Our findings emphasise the importance of echinoderms as a  
powerful model for regeneration owing to their unique regenerative capabilities and experimental  
700 amenability, but also to their phylogenetic position crucial for comparative analyses. The  
extensive genomic and transcriptomic resources we generated for the brittle star *Amphiura*  
*filiformis* thus represent a cornerstone to understand the evolutionary, molecular and genetic  
underpinnings of animal appendage regeneration, emergence of pentameral symmetry and  
remarkable diversity of morphologies and developmental strategies seen across echinoderm  
705 lineages.

## Data and code availability

Genome sequence and RNA-seq data have been deposited in NCBI SRA (Bioproject PRJNA1029566) and will be made publicly available upon publication. The code for the annotation workflow is publicly available on GitHub (<https://github.com/eparey/AnnotateSnakeMake>). Supplemental datasets to reproduce the results have been deposited in Zenodo (<https://zenodo.org/doi/10.5281/zenodo.10036671>, see Supplemental Material). These include the genome, gene and repeat annotations, processed gene expression tables and source data for the figures.

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## Author contributions

OOM, TL, MT, SD and KJ initiated the genome sequencing project. OOM, PO, FM and EP designed the study. FM, PO generated sequencing data, and FM and EP assembled and annotated the final genome version. LP performed proximity ligation experiments. For transcriptomic analyses, OOM and TL designed the explant study and generated RNA-seq explant data, and AC collected regeneration time course samples. EP performed the computational analysis of synteny, gene family and comparative transcriptomics of regeneration with the support of FM. DD and SA contributed in data analysis and visualisation. OOM, JD, KMB, PM and PO curated the data and assisted with interpreting the results. EP wrote the manuscript with key contributions from FM, PM and KMB. All authors commented on the manuscript and approved the final version.

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755 The authors would like to dedicate this manuscript to Michael Thorndyke and R. Andrew Cameron, who were both instrumental in the early stages of this project but, sadly, passed away before analyses were completed. Without their contributions, this work would not have been possible.

## Methods

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### Animal sampling

Adult *A. filiformis* were collected at 25-40 m depth from sediment in the Gällmarsfjord in the vicinity of Kristineberg Marine Station, Sweden, using a Petersen mud grab. Individuals were separated from the sediment by rinsing them with seawater and then maintained in natural flowing seawater at 14°C. Sperm was collected from a single-individual by dissecting the gonads from the bursae.

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### DNA extraction and sequencing

770 Sperm cells were concentrated by centrifugation, washed repeatedly, and subsequently embedded in 2% low melting agarose. Sperm cells were lysed in a solution of 1% SDS, 10mM Tris (pH 8) and 100mM EDTA and then resuspended in a solution of 0.2% N-laurylsarcosine, 2mM Tris (pH 9) and 0.13 mM EDTA. High molecular weight (HMW) DNA was released from the agarose blocks using  $\beta$ -agarase (NEB).

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Long-read sequencing was performed on six Nanopore Promethlon flowcells (vR9.4.1). Several libraries were constructed using the Ligation Sequencing Kit (Nanopore LSK109) using DNA sheared to different size using a megaRuptor (Diagenode) to optimise yield and contiguity. Bases were called from raw signal with Guppy (model "dna\_r9.4.1\_450bps\_hac\_prom", version 2.3.5). A total of 160.56 Gb nanopore reads was acquired (~100x coverage). A library of 10x linked-reads was generated using the Chromium system (10x genomics) and sequencing on a Novaseq6000 SP lane in a 2 x 150 bp layout for a total of 246M reads (86 Gb). Genome size was estimated to 1.33 Gb with a heterozygosity of 3.22% by counting k-mer (k=31) in the short read data using jellyfish2 (Marçais and Kingsford 2011) and fitted through a four-peak model using Genomescope2 (Ranallo-Benavidez et al. 2020).

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### Genome Assembly

790 We assembled Nanopore reads using flye (v2.9-b1768) (Kolmogorov et al. 2019) assuming a coverage of 30x and a genome size of 3 Gb to account for the high level of heterozygosity. We obtained a diploid assembly of 2.86 Gb (N50: ~2.78 Mb) which was subsequently polished using Racon (v1.5.0) (Vaser et al. 2017) for two iterative rounds using the nanopore reads and then for another two rounds using the short-read illumina reads that were aligned to the assembly using minimap2 (v2.24-r1122) (Li 2018). Haplotypes were then removed from the assembly using purge\_dups (v1.2.5) (Guan et al. 2020) with cutoffs visually adjusted from the coverage distribution on contigs. The resulting assembly had a total length of 1.57 Gb, with N50 and L50 of 3.2 Mb and 154 respectively, and 96.1% complete BUSCO score.

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To scaffold this assembly, we built a Hi-C library using the Omni-C kit (Dovetail) from gonadal tissue. Chromatin was fixed using PFA and digested using a sequence-independent nuclease after re-ligation and biotinylation. A sequencing library was built from purified DNA and 225M reads sequenced on a Novaseq X (~45x coverage). Hi-C reads were mapped to the polished haplopurged assembly using bwa mem (0.7.17-r1198-dirty) with options -5SP -T0 and alignment were further sanitised, sorted and duplications removed using pairtools (v1.0.2) (Open2C et al. 2023) with options `--walks-policy 5unique`, `--max-inter-align-gap 30` an a minimum MAPQ of 40. We used YAHS (v1.1a-r3) (Zhou et al. 2023) to scaffold the genomic contigs using the Hi-C read alignment as input. We obtained 20 main chromosome-scale scaffolds totalling 1.47 Gb

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corresponding to 93.5% of the total assembly length (**Figure S1**). The GC level of the final genomic sequence is 36.67 % and the N50 is 68.86 Mb.

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### Repeat annotation

We used RepeatModeler 2.0.2 to build a *de novo* repeat library for the brittle star genome and then ran RepeatMasker 4.1.2-p using this library as input to soft-mask the genome and extract repeat location (Flynn et al. 2020). We used DeepTE (Yan et al. 2020) to classify repeats that could not be classified with the homology-based repeat classification implemented in RepeatModeler. We re-trained a DeepTE model to classify metazoan repeats into 5 classes, using a balanced dataset of 12,500 distinct repeats (2,500 repeats for each of the 5 classes) from different sources including rebase (Bao et al. 2015), Dfam (Hubley et al. 2016) and homology-based classifications of repeats from 17 echinoderms and 2 hemichordates genomes (Validation accuracy=0.98 at the class probability threshold  $p \geq 0.55$ , **Figure S1 B**). On a test set of 827 brittle star repeat families that were not included in the training set and where RepeatModeler homology-based predictions serve as ground truth, this re-trained DeepTE model has higher accuracy than the default metazoa model available in DeepTE (accuracy=0.81 vs 0.67, **Figure S1 C**). Divergence to consensus (kimura %) were computed and repeat landscapes plotted using the 'calcDivergence.pl' and 'createRepeatLandscape.pl' scripts from RepeatMasker. The same methodology was applied to build repeat landscapes for *P. lividus*, *H. leucospilota* and *M. glacialis*. Repeat annotations are provided in **Dataset S1**. Repeat ages were estimated from divergence to consensus using a neutral substitution rate of  $1.885 \times 10^{-9}$  per base pair per year for *A. filiformis*, which was estimated with phyloFit from an alignment of 66,818 4-fold degenerate sites containing 17 echinoderm and 2 hemichordate genomes.

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### RNA isolation, extraction and sequencing

#### Arm regeneration RNA-seq in brittle star (time course in whole animals)

*A. filiformis* individuals were obtained in the fjord close to the Kristineberg Center for Marine Research and Innovation, Sweden, at depths of 20-60 metres. Samples of different regenerating stages were obtained as previously described in (Czarkwiani et al. 2016) for early regeneration stages (48 hpa, 72 hpa, Stages 3, 4 and 5) and in (Dupont and Thorndyke 2006) for 50% differentiation index stages (50% P and 50% D). 30 regenerates from different individuals were used per stage. Dissection for RNA-sampling was performed as follows (**Figure 4A**): (i) for the non-regenerating control, we dissected one mature arm segment, (ii) for 48 and 72 hpa samples, we dissected the last segment at the amputation site, (iii) for stages 3 to 5 we dissected the regenerative tissues, and finally (iv), for 50% regenerates, we sampled several segments of proximal and distal tissues, excluding the differentiated distal cap structure. The collected regenerates were lysed in 10 volumes of RLT (Qiagen), and total RNA extracted using RNAeasy micro RNA kit (Qiagen). RNA concentration and integrity was measured using Bioanalyzer (Agilent). Library preparation and paired-end sequencing was conducted by Novogene.

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#### Arm regeneration RNA-seq in brittle star severed arm experiments (explant)

We collected around 3,500 brittle stars with a 5-7 mm disc diameter. While animals were sedated in 3.5% w/w  $MgCl_2$  in artificial seawater, two arms from each organism were amputated by pressing a scalpel blade into the intervertebral autotomy plane. We first sectioned the arms 0.5 cm from the disc (amputation 1, **Figure 6A**) and then sectioned them again at the distal end (amputation 2, **Figure 6A**). We thus produce explants (i.e. severed brittle star arms) of 1cm in length with wound sites at the proximal and distal ends. 43 groups of 150-200 explants were cultured in flow through aquaria at 16°C. Explants were sampled at 3 and 5 days post-amputation (dpa), sedated in 3.5% w/w  $MgCl_2$  in artificial seawater for 15 minutes and then

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dissected into three sections: proximal, medial and distal (**Figure 6A**). Each explant section was flash frozen in liquid nitrogen and collected in batches of 150-200 pieces. Each batch was  
860 individually homogenised with glass pistils and RNA was extracted with the RiboPure kit (Applied Biosystems), following manufacturer's protocol. RNA concentrations were measured using a QuBit 2.0 RNA fluorometric assay (Thermo Fisher Scientific, Waltham, MA, USA) and RNA integrity was checked using 0.5% (w/v) agarose-MOPS-formaldehyde de-naturing gel electrophoresis.

865 Complementary DNA (cDNA) libraries were prepared using the Illumina TruSeq v2 mRNA sample prep kit (Illumina, San Diego, CA, USA), following a standard protocol. Briefly, mRNA was isolated with poly-A selection, followed by cDNA synthesis, Illumina standard index adapter ligation and a brief PCR reaction. Concentrations of the cDNA libraries were measured using a  
870 QuBit DNA High-sensitivity assay (Thermo Fisher Scientific, Waltham, MA, USA) and fragment length distributions were assessed using an Agilent TapeStation with a D1000 tape (Agilent, Santa Clara, CA, USA). cDNA libraries were multiplexed by equimolar pooling (5 or 6 samples/pool), and were then sent to the Swedish National Genomics Infrastructure's SNP & SEQ platform in Uppsala for Illumina HiSeq 2500 sequencing (8 lanes; 126 bp Paired-End sequencing; Illumina, San Diego, CA, USA).  
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### Gene annotation

We annotated the brittle star genome using a custom pipeline leveraging three types of evidence: (i) assembled transcriptomes from 18 brittle star developmental and arm regeneration RNA-seq  
880 (including both publicly available (Delroisse et al. 2014; Purushothaman et al. 2015; Dylus et al. 2016) and newly generated datasets, **Table S1**), (ii) similarity to proteins from 27 selected metazoa and (iii) *ab initio* predictions. We first assembled a consensus transcriptome from all RNA-seq samples with mikado (Venturini et al. 2018), combining an alignment-free transcriptome assembled with Trinity (Grabherr et al. 2011) and mapped to the genome with gmap (Wu and  
885 Watanabe 2005) with an alignment-based transcriptome mapped to the genome with STAR (Dobin et al. 2013), assembled with Stringtie (Pertea et al. 2015) and merged with taco (Niknafs et al. 2017). Second, we selected best-scoring mikado transcripts (i.e. transcripts with identified start and stop codons by TransDecoder, at least 2 exons, over 50% of the predicted coding sequence covered by a swissprot (Boutet et al. 2007) blast hit and no overlap of the coding  
890 sequence with an annotated repeat) to train a gene prediction model with AUGUSTUS (Stanke and Waack 2003). Third, we obtained similarity-based gene predictions with Metaeuk, based on proteomes from a total of 27 metazoa, including 8 echinoderms and 2 hemichordates. Fourth, we performed *ab initio* gene prediction with AUGUSTUS (Stanke and Waack 2003), using the previously trained model and providing as hints the predicted exons by mikado and Metaeuk, and  
895 curated splice junctions defined by portcullis (Mapleson et al. 2018) on the STAR transcriptome. Fifth, we filtered out all predicted gene models with over 40% of exons overlapping annotated repeats and then ran PASA (Haas et al. 2003) on retained genes to finalise models and annotate UTRs. Finally, we further filtered out 3,465 poorly supported gene models (no PFAM domain, no swissprot blast hit and maximal expression < 2 TPM), to retain 30,267 annotated gene models in  
900 the final annotation (**Table S2**). The quality and completeness of the annotation is demonstrated by a score of 92.7% complete BUSCO (C:92.7 [S:86.2%, D:6.5%], F:5.0%, M:2.3%, n:954) (Simão et al. 2015) and a total of 4,974 unique PFAM domains (Finn et al. 2014), with 76% of genes (23,047) containing a PFAM domain. Genes were named by BLAST search against the swissprot database. The names of genes specifically investigated in this study (*hox*, *phb*,  
905 *luciferase*) were further manually curated to reflect their evolutionary history. This genome annotation pipeline is implemented as a Snakemake workflow (Köster and Rahmann 2012) and

is publicly available on Github (<https://github.com/eparey/AnnotateSnakeMake>). Annotation files are provided in **Dataset S1**.

## 910 **Synteny comparisons and reconstruction of Eleutherozoa ancestral linkage groups**

For the sea urchin *P. lividus* and the black sea cucumber *H. leucospilota*, we used previously reported gene annotations (Chen et al. 2023; Marlétaz et al. 2023). We generated a draft homology-based annotation for the spiny sea star *M. glacialis* (Lawniczak et al. 2021) with MetaEuk (Levy Karin et al. 2020) using proteins of the sea urchin *S. purpuratus* (Spur\_5.0, available in Ensembl Metazoa v56, (Sea Urchin Genome Sequencing Consortium et al. 2006)), 915 the crown-of-thorns sea star *A. planci* (OKI\_Apl\_1.0, available in Ensembl Metazoa v56 (Baughman et al. 2014)) and the octopus sea star *P. borealis* (Lee et al. 2022). One-to-one orthologous genes were identified by reciprocal best blast hit between pairs of compared genomes, using diamond (Buchfink et al. 2021). We used Circos version 0.69.8 and circos-tools 920 0.23 (Krzywinski et al. 2009) to plot synteny comparisons, with the bundlelinks tool to group together neighbouring genes (with a maximum gap of 50 genes) and filter out bundles with fewer than 3 links. We then used the orderchr tool to order chromosomes so as to minimise link crossings. The ancestral Eleutherozoa linkage groups were reconstructed on the basis of synteny comparisons between the spiny sea star *M. glacialis* and the black sea cucumber *H.* 925 *leucospilota*, and with the amphioxus *B. floridae* and the scallop *P. maximus* genomes as well as previously defined bilaterian linkage groups (BLGs) to untangle derived from ancestral chromosomal arrangements (**Figure S2**). Specifically, only one macro-syntenic rearrangement occurred between the spiny sea star and the black sea cucumber: (a) spiny sea star chr5 maps to both sea cucumber chr12 and chr23. Comparisons with outgroups and ancestral BLGs 930 revealed that (a) corresponds to a derived fusion in the spiny sea star and that the black sea cucumber retained the ancestral state. Using this reconstruction, we annotated genes from matched orthologous chromosomes between sea stars and sea cucumbers with respect to their ancestral ELGs of origins and propagated annotations to orthologous genes in *P. lividus*, *A. filiformis* and other available chromosome-scale echinoderm genomes. Karyotypes were drawn 935 with RIdeograms (Hao et al. 2020): we painted genes on extant chromosomes using the ancestral chromosome colour when a significant number of genes were inferred to descend from an ancestral chromosome ( $p < 10^{-5}$ , Fisher exact tests corrected for multiple testing with the Benjamini-Hochberg procedure). Oxford grid plots of ELGs distribution in *P. lividus*, *A. filiformis* and other sequenced echinoderms (**Figure S3**) were plotted using the same statistical 940 thresholds. ELG-related data files are provided in **Dataset S2**.

## **Hox and ParaHox genes identification**

We first compiled a dataset of curated full length HOX protein sequences from *S. purpuratus* and HOX homeodomains from *B. floridae* and *S. kowaleskii* to search for homologous Hox genes in 945 the brittle star. A comprehensive list of candidate Hox genes in brittle star was then constructed using two approaches: (i) a diamond blastp (Buchfink et al. 2021) of the curated Hox dataset against brittle star predicted proteins (ii) a miniprot (Li 2023) alignment of *S. purpuratus* HOX protein sequence against the brittle star genome sequence, to recover Hox genes potentially missed by the automatic annotation process. The same approach was used to identify Hox 950 genes in *M. glacialis*. Finally, we built a molecular phylogenetic tree (**Figure S4**) with RAxML-NG (Kozlov et al. 2019) using the LG+G4+F model and 5 distinct starting parsimony trees, to reconstruct the evolutionary history of Hox genes from *S. purpuratus*, *B. floridae*, *S. kowaleskii*, *A. filiformis*, *M. glacialis* and three additional echinoderms with curated Hox genes in Echinobase (Arshinoff et al. 2022) (*L. variegatus*, *P. miniata* and *A. planci*). We extracted ParaHox 955 sequences and location in *A. japonica* from Ensembl Metazoa v56 (Yates et al. 2022), in *S. purpuratus* from Echinobase, *A. planci* from Ensembl Metazoa and similar approaches as for the

Hox to identify ParaHox genes in *A. filiformis* and *M. glacialis*. Hox and ParaHox data files are provided in **Dataset S3**.

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### Gene families expansion and contraction

We used broccoli (Derelle et al. 2020) to group proteins of 28 selected Metazoa, 10 of which were Ambulacraria, into gene families. Gene families were predicted to have originated in the last common ancestor of the species with a gene in the family. Out of the complete set of broccoli gene families, 10,367 originated before the last common ancestor of Ambulacraria (echinoderms and hemichordates outgroups). We used CAFE5 (Mendes et al. 2020) on the set of 10,367 families to identify significantly expanded and contracted gene families on each branch of the Ambulacraria phylogeny. Briefly, CAFE fits a birth-death model on a dated phylogeny from the gene count data, and tests for significant expansions/contractions on specific branches. To obtain a dated Ambulacraria phylogeny, we: (i) extracted 192 1-to-1 orthologs in Ambulacraria from broccoli gene families, (ii) built multiple sequence alignments for each orthologous group using MAFFT v7.475, (iii) concatenated alignments across orthology groups and reconstructed a Maximum Likelihood phylogeny with RAxML-NG v.1.1 (LG+G4+F model with 10 parsimony starting trees), (iv) filtered out columns of the alignment with over 15% gaps (47,520 retained sites) and (v) ran PhyloBayes v4.1b (Lartillot et al. 2009) to obtain a time-calibrated tree, with the RAxML reconstructed tree as constrained topology and selected fossil calibrations extracted from the literature ((Benton et al. 2009; Mongiardino Koch et al. 2022). The chain was run for 4,166 samples and 3,500 were retained after burn-in to estimate the posterior distributions for node ages. We next ran CAFE in 2 steps: we estimated the lambda and alpha parameters of the 2-categories CAFE GAMMA model excluding the 128 gene families with the largest copy number differential and then ran CAFE on all families with these parameters fixed to test for significant contractions and expansions (p-values <0.05). Fossil calibrations, dated species tree, gene families and CAFE output files are provided in **Dataset S4**.

### 985 Gene lists curation

We generated lists of immune genes in *A. filiformis* (**Table S2**) using a combination of PFAM domain annotation and lists of previously curated immune genes in the sea urchin *Strongylocentrotus purpuratus* (Sea Urchin Genome Sequencing Consortium et al. 2006). Specifically, we first selected *A. filiformis* genes based on their PFAM domains (e.g. TIR, IL17, Mif) and completed this list by searching for homologs (using the set of broccoli gene families) with the immune genes of the sea urchin *Strongylocentrotus purpuratus*. For the list of kinase genes, we also identified through homologies with curated kinase genes from the sea urchin genome (Sea Urchin Genome Sequencing Consortium et al. 2006). We generated a list of TFs based on the presence of DNA-binding PFAM domains. For the stemness genes, we identified putative homologues of the 180 “stemness-like” genes established by (Alié et al. 2015), that is, genes that are shared between three stem cell populations: poriferan (*Ephydatia fluviatilis*) archeocytes, *Hydra vulgaris* interstitial stem cells, and planarian (*Schmidtea mediterranea*) neoblasts. Specifically, we used the human cognates of all those genes as queries for BLAST searches (McGinnis and Madden 2004) against *A. filiformis* genes. We emphasise that, while these genes are expressed in stem cell populations, they may not all be stemness regulators. For genes involved in neuronal function, we first compiled a list of neurogenic and glial markers, TFs, cell signalling genes involved in embryonic, homeostatic, and regenerative neurogenesis in vertebrates (rodents, humans, and *Xenopus*) and invertebrates (*Caenorhabditis elegans* and *Drosophila melanogaster*). We identified putative homologues of “neuronal” genes in *A. filiformis* using a reciprocal blast approach. We generated gene lists for 19

1010 signalling pathways, in two steps: (i) manual curation of main members of selected pathways, (ii) identification of their gene ID in the *S. purpuratus* genome via echinobase gene searches (Arshinoff et al. 2022) and (iii) identification of *S. purpuratus* orthologs in *A. filiformis* using gene trees built with Generax (Morel et al. 2020) for each of our broccoli (Derelle et al. 2020) gene family. Finally, the repertoire of luciferase-like genes was identified through reciprocal BLAST searches using the reference *Renilla* luciferase (GenBank: AAA29804) as initial query (Delroisse et al. 2017).

### Gene ontology enrichment tests

1015 We used eggno-mapper (Cantalapiedra et al. 2021) to automatically annotate *A. filiformis* and *P. lividus* genes with Gene Ontology terms from the Biological Process domain. The GO annotations were then transferred to the level of gene families. Specifically, for each family, we propagated all GO annotations associated with any *P. lividus* or *A. filiformis* genes as the complete set of GO annotations for this family. Hypergeometric tests for functional enrichments were then conducted with the enricher function from the ClusterProfiler R package (Wu et al. 2021), with custom foreground and background GO annotation sets. For functional enrichment tests on expanded/contracted gene families (**Figure 3**), tests were conducted at the level of gene families with expanded or contracted families as foreground and all gene families as background, as described above. For functional enrichment tests on regeneration co-expression clusters (**Figure 4**), tests were conducted at the level of brittle star genes, using genes of a given cluster as foreground and genes of all clusters as background. We used FDR < 0.05 as significance threshold. Enrichment results were summarised with REVIGO (Supek et al. 2011), we selected top ontology terms based on the REVIGO “dispensability” score to make a representative overview of the diversity of enriched GO terms.

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### Clustering of the arm regeneration expression series

Gene expression was quantified for all samples using the alignment-free method kallisto (Bray et al. 2016). We normalised TPM values across samples using the TMM method as implemented in edgeR (Robinson et al. 2010; Robinson and Oshlack 2010) and used MFuzz (Kumar and E Futschik 2007) to perform soft-clustering of genes on the basis of their standardised expression profiles across samples. We used the minimum centroid distance method to select the optimal number of clusters (n=19, **Figure S6**). In the main text, we filtered the obtained clusters to retain clusters with > 1 enriched GO term and elevated expression in >1 regenerating sample (**Figure 4, Figure S6**). Normalised gene expression tables are provided in **Dataset S5**.

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### Transcription factor binding motif enrichment tests

1045 We used HOMER v4.11 (Heinz et al. 2010), to test for enriched transcription factor binding motifs in the proximal regulatory domains (TSS + 5 kb upstream, +1 kb downstream) of genes of each regeneration cluster. We ran the findMotifsGenome.pl script from the HOMER suite, with -h to perform hypergeometric tests, contrasting proximal regulatory domains of genes from one expression cluster as foreground with proximal regulatory domains from genes of all clusters as background.

### Axolotl limb regeneration RNA-seq time course

1050 Raw RNA-seq data for 12 limb regeneration time points from (Stewart et al. 2013) were downloaded from <https://www.axolomics.org/?q=node/2>. We used Trim Galore (<https://github.com/FelixKrueger/TrimGalore>) with default parameters to trim and quality filter raw sequencing reads via the Cutadapt tool (Martin 2011). Gene expression was quantified with kallisto (Bray et al. 2016) using the set of annotated axolotl transcripts from the latest *Ambystoma mexicanum* assembly version (AmexG\_v6.0-DD, available from <https://www.axolotl->

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1060 [omics.org/assemblies](https://www.ncbi.nlm.nih.gov/sra/omics.org/assemblies), (Schloissnig et al. 2021)). Similarly as for the brittle star regeneration series, we normalised TPM values across samples using the TMM method (Robinson et al. 2010; Robinson and Oshlack 2010) and used MFuzz (Kumar and E Futschik 2007) to cluster genes according to their expression profile (**Figure S7**). Gene ontology and TFBS motifs enrichment were performed as described for the brittle star. Normalised gene expression tables are provided in **Dataset S5**.

### 1065 **Parhyale limb regeneration RNA-seq time course**

Parhyale leg regeneration expression data were previously processed and clustered into 8 co-expression gene groups using the same approach as we used for brittle star data (Sinigaglia et al. 2022). As such, we directly used the clustering reported in (Sinigaglia et al. 2022), but renamed the clusters so that numbering follow temporal activation (P1 is R4 in the notation of Sinigaglia et al., P2 is R1, P3 is R8, P4 is R2, P5 is R6, P6 is R3, P7 is R5 and P8 is R7).

### 1070 **Comparison of gene expression dynamics during appendage regeneration**

We used broccoli (Derelle et al. 2020) to build homologous gene families encompassing genes of the brittle star *A. filiformis*, the axolotl *Ambystoma mexicanum* and *Parhyale hawaiiensis*, as well as 8 echinoderms, 6 vertebrates, 7 ecdysozoans and 12 other animal genomes. We used these gene families to identify homologous genes and compare their expression profile during appendage regeneration in a pairwise manner across the three species. For each pairwise comparison, we retained all homologous gene families with > 1 gene and < 5 genes in each of the two compared species, resulting in a total of 5,203 homologous groups retained for the axolotl (8,810 homologous genes) - brittle star (6,813 homologous genes) comparison, 3,137 for the brittle star (4,196) - Parhyale (3,617) comparison and 2,299 for the axolotl (3,903) - Parhyale (2,628) comparison (see **Dataset S5**). We next computed permutation-based p-values to test for the overrepresentation of homologous genes across co-expression clusters of the two compared species. Specifically, we generated, for each pairwise comparison, 10,000 randomizations of the gene labels of species 2, keeping clusters and orthologous gene family size constant to build a null distribution of the number of expected homologous genes shared by two clusters at random. Empirical p-values were computed from the null distribution and corrected for multiple testing using the Benjamini-Hochberg procedure. To investigate functional annotation of genes displaying co-expression across regeneration models as opposed to genes from the same clusters that do not show co-expression across species, we conducted gene list and GO enrichment tests as described previously but using carefully selected background: we used as background all brittle star genes with a homolog in either Parhyale or axolotl (i.e. whose expression conservation could be tested) and in a cluster with identified co-expressed genes in either Parhyale or axolotl (to test for the specificity of genes of a given cluster that show conservation vs those of the same cluster that do not).

### 1095 **Differential analysis of repeats transcriptional activity during brittle star arm regeneration**

We tested for differentially expressed repetitive elements in early regeneration (immune phase: 48 hpa and 72 hpa samples) versus middle regeneration (proliferation: Stage 3, Stage 4, Stage 5 samples), using our time course brittle star arm regeneration RNA-seq data. We used a conservative approach to first filter out highly duplicated genes which could have been captured in the set of repetitive elements called by RepeatModeler/RepeatMasker. We used diamond blastx (Buchfink et al. 2021) to search for homologies between repeat consensus and proteins in the swissprot database (Boutet et al. 2007) and filtered out all 'Unknown' repeat families for which the consensus sequence had a strict match in swissprot (e-value cut-off  $10^{-10}$ ) which did not correspond to transposon genes. We next use the SalmonTE pipeline (Jeong et al. 2018)

with default parameters on the full set of filtered repeat consensus (n=4,695 repeat families), followed by differential analysis with DESeq2 (Love et al. 2014) on the estimated count values, to test for differential transcriptional activity of repeats in the immune versus proliferation regeneration phases. We retained as differentially expressed the repetitive elements with an absolute  $\log_2$  fold change  $> 1$ , adjusted p-value  $< 0.001$ .

### **Differential gene expression in brittle star arm explants**

Gene expression was quantified for all samples using the alignment-free method Kallisto (Bray et al. 2016). Differential expression analyses were conducted with DESeq2 (Love et al. 2014) on count values, contrasting distal replicates against medial replicates and proximal replicates against medial replicates for each time point. All genes with an adjusted p-value  $< 0.05$  and absolute  $\log_2$  fold change  $> 1$  were retained as differentially expressed. Gene expression tables are provided in **Dataset S5**.

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