

# 1        **Molecular evidence of anteroposterior patterning in adult echinoderms**

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## 30 Abstract

31 The origin of the pentaradial body plan of echinoderms from a bilateral ancestor is one of the most  
32 enduring zoological puzzles<sup>1,2</sup>. Since echinoderms are defined by morphological novelty, even the most  
33 basic axial comparisons with their bilaterian relatives are problematic. To revisit this classical question,  
34 we used conserved antero-posterior (AP) axial molecular markers to determine whether the highly  
35 derived adult body plan of echinoderms masks underlying patterning similarities with other  
36 deuterostomes. We investigated the expression of a suite of conserved transcription factors with well-  
37 established roles in the establishment of AP polarity in deuterostomes<sup>3-5</sup> and other bilaterians<sup>6-8</sup> using  
38 RNA tomography and *in situ* hybridization in the sea star *Patiria miniata*. The relative spatial expression  
39 of these markers in *P. miniata* ambulacral ectoderm shows similarity with other deuterostomes, with the  
40 midline of each ray representing the most anterior territory and the most lateral parts exhibiting a more  
41 posterior identity. Strikingly, there is no ectodermal territory in the sea star that expresses the  
42 characteristic bilaterian trunk genetic patterning program. This suggests that from the perspective of  
43 ectoderm patterning, echinoderms are mostly head-like animals, and provides a developmental rationale  
44 for the reevaluation of the events that led to the evolution of the derived adult body plan of echinoderms.

## 45 Introduction

46 Echinoderms, defined by their calcitic endoskeleton, unique water vascular system, and perhaps most  
47 strikingly by their pentaradial body plan<sup>1,2</sup>, are among the most enigmatic animal phyla. Since  
48 echinoderms are phylogenetically nested within the deuterostomes<sup>9,10</sup> (echinoderms, hemichordates, and  
49 chordates), their pentaradial organization was evidently derived from a bilateral ancestor. Yet, despite a  
50 rich fossil record, comparative morphological studies have come to conflicting conclusions regarding  
51 the axial transformations that led to pentamery from the ancestral bilaterian state<sup>2,11</sup>. Among bilaterians,  
52 the deployment of the gene regulatory network that specifies ectoderm AP polarity is highly conserved<sup>3-8</sup>  
53 and represents a suite of characters that is often more conserved than the body plans they regulate<sup>4,5</sup>  
54 (Fig. 1a). The deployment of this AP gene regulatory network could therefore provide an alternative  
55 way to test hypotheses of axial homology in cases such as echinoderms where morphological characters  
56 are too divergent to reconstruct ancestral states<sup>12</sup>.

57 Detailed comparisons between chordate and hemichordate axial patterning have established the  
58 exquisite conservation of the deuterostome ectodermal AP patterning program<sup>4,5</sup>, which provides a  
59 phylogenetically robust framework for addressing the evolution of echinoderm axial properties. We  
60 considered two contrasting scenarios. First, the ancestral deuterostome AP patterning network could  
61 have been dismantled and reassembled into novel conformations during the radical body plan  
62 modifications along the echinoderm stem lineage. In this scenario, expression of transcription factors in  
63 derived morphological structures without extensive conservation of relative spatial expression would  
64 imply co-option into novel developmental roles<sup>13,14</sup>. Alternatively, conservation of spatially coordinated  
65 expression of this network during the elaboration of the echinoderm adult body plan would provide a  
66 molecular basis for testing hypotheses of axial homology with bilaterians, and establish regional  
67 homologies masked by divergent anatomies<sup>12</sup> (Fig. 1b).

68 Under the second scenario, four main hypotheses have been proposed to relate the echinoderm body  
69 plan to other bilaterians that explicitly consider axial homology (Fig.1c). The bifurcation<sup>15</sup> and the  
70 circularization<sup>15,16</sup> hypotheses can be ruled out, since they require a unique molecular identity for each  
71 of the five echinoderm rays that is inconsistent with molecular data<sup>17</sup>. In the duplication hypothesis<sup>15,18</sup>,  
72 each of the five echinoderm rays is a copy of the ancestral AP axis joined anteriorly in the disk, and in  
73 the stacking hypothesis<sup>2,11,19</sup> the oral-aboral axis of adult echinoderms is homologous to the ancestral  
74 AP axis. Although broad bilaterian comparisons of AP axis patterning are typically based on ectodermal  
75 expression domains, the stacking hypothesis was proposed largely on the basis of nested posterior Hox  
76 gene expression in the posterior mesoderm of the bilateral larval stages in holothuroids (sea cucumbers),  
77 crinoids (sea lilies), and echinoids (sea urchins)<sup>20-26</sup>. While initially restricted to coelomic  
78 compartments<sup>2,19</sup>, some iterations of the stacking hypothesis have also included the oral ectoderm as the

79 most anterior compartment<sup>11</sup>. However, recent analyses suggest that the anterior identity of the oral  
80 ectoderm should be considered independent of mesoderm derivatives<sup>27</sup>.

81 Here, we tested these classical hypotheses by examining gene expression in the asteroid (sea star) *Patiria*  
82 *miniata* (Extended Data Fig. 1a,b), using a statistical approach based on spatial transcriptomics and *in*  
83 *situ* hybridization. Our data provide evidence for unexpected conservation of the ancestral AP  
84 ectodermal patterning network during ambulacrum development in juveniles. Importantly, our results  
85 do not fit with any of the established hypotheses and instead suggest a new ambulacral-anterior model  
86 to relate echinoderm pentaradial symmetry to the ancestral bilateral symmetry.

## 87 **Results and discussion**

### 88 RNA tomography in adult *Patiria miniata*

89 The manifestation of pentaradial symmetry in asteroids is simpler than in other echinoderm classes  
90 because it follows a planar organization with the rays extending around the oral-aboral axis through  
91 easily distinguishable alternating ambulacral territories (where the grooves bearing the tube feet are  
92 located) and interradial territories (Fig. 2a). This makes asteroids particularly well-suited for  
93 investigating potential axial homologies between the pentaradial echinoderm body plan and closely  
94 related bilateral deuterostomes using the molecular anatomy coordinates offered by the conserved,  
95 ectodermal AP patterning network. The adult body plan of asteroids is defined by the endoskeleton,  
96 which underlies the body wall in the aboral and interradial areas, the digestive tract and pyloric caeca,  
97 the main body muscles, the water vascular system along each ambulacrum, and the central nervous  
98 system that includes five radial nerve cords and the circumoral nerve ring. We used micro computed  
99 tomography (micro-CT) to show the exact spatial arrangement of the principal anatomical features in *P.*  
100 *miniata* juveniles along each organizational plane of the juvenile arms (Fig. 2a, Extended Data Fig. 1c,d;  
101 Supplementary Fig. 1; Supplementary Video 1).

102 Within this anatomical framework, we began investigating the transcriptional landscape of *P. miniata*  
103 along its body axes with an unbiased spatial transcriptomics approach using RNA tomography<sup>28</sup>. We  
104 cryo-sectioned three arms from *P. miniata* juveniles at a stage that matched the micro-CT scanned  
105 specimen, along three different dimensions: from the proximal to the distal part of the arm (P-D), from  
106 the oral to the aboral side (O-A), and from the left to the right side (hereafter referred to as medio-lateral;  
107 M-L) (Fig. 2b; Extended Data Fig. 1c, Supplementary Video 1; Supplementary Fig. 2a-c). Sections were  
108 barcoded and pooled for single molecule real time sequencing with PacBio IsoSeq (Supplementary Fig.  
109 2d,e; Supplementary Fig. 3a-d; Supplementary Tables 1-4), yielding a three-dimensional atlas of 25,794  
110 gene expression profiles along the P-D, O-A and M-L dimensions of *P. miniata* arms (Fig. 2c). Gene  
111 clustering based on the similarity of RNA tomography expression profiles highlighted seven principal  
112 patterns of gene expression (Fig. 2c). We confirmed that transcriptional landscape determined by RNA  
113 tomography was consistent with the anatomy of the animal by considering principal component analyses  
114 and Spearman correlations between sections along each dimension and analyzing the expression profile  
115 of marker genes known to be expressed in specific tissues (Extended Data Fig. 1e-g). To aid in this  
116 analysis we generated a new *P. miniata* genome assembly (Supplementary Fig. 2e; Supplementary Fig.  
117 3d,e; Supplementary Tables 4,5).

118 To consider possible molecular anatomical homologies across deuterostomes we identified 36 conserved  
119 molecular markers in the RNA tomography dataset that define specific ectoderm territories along the  
120 AP axis in hemichordates and chordates (Fig. 1a; Supplementary Fig. 4) and retrieved their expression  
121 profiles (Fig. 2d). These marker genes included transcription factors (*fezF*, *six3/6*, *nkx2.1*, *foxG*, *lhx2/9*,  
122 *otp*, *dmbx*, *tlx*, *emx*, *barH1*, *pax6*, *irx*, *dbx*, *otx*, *lhx1/5*, *engrailed*, *gbx*, *pax2/5/8*, *hox1*, *hox2*, *hox3*, *hox4*,  
123 *hox5*, *hox7*, *hox8*, *hox9/10*, *hox11/13b*); members of the Wnt signaling pathway (*sfrp1/5*, *sfrp3/4*, *fzd5/8*,  
124 *wnt3*); and the ligand *hedgehog*. Four additional transcription factor markers (*rx*, *dlx*, *hox11/13a* and  
125 *hox11/13c*) were excluded from the computational analyses because of low expression levels  
126 (Supplementary Fig. 3f). *Hox6* was absent in our Hox cluster assembly (Supplementary Fig. 5), as

127 reported previously in the closely related species *P. pectinifera*<sup>29</sup> and in the crown of thorns sea star  
128 *Acanthaster planci*<sup>30</sup>, consistent with its loss in valvate asteroids.

129 The duplication and stacking hypotheses would be supported by staggered expression of these AP  
130 patterning markers along the P-D or O-A dimensions, respectively. We ranked the AP patterning  
131 markers from the anterior to posterior using the hemichordate *Saccoglossus kowalevskii* as a template,  
132 since it is the most closely related bilateral species with a comprehensive expression pattern dataset for  
133 these markers (Supplementary Fig. 6). We then tested the Spearman correlation between the ranking of  
134 the genes and their position along the P-D and O-A dimensions. In both cases, we found moderate but  
135 not statistically significant correlations ( $\rho=0.25$ ,  $p=0.16$  and  $\rho=0.27$ ,  $p=0.13$ , respectively) (Fig. 2e).  
136 Organizing the AP patterning genes into seven groups based on their expression profiles (Supplementary  
137 Fig. 6b), we found that the moderate correlation with the O-A dimension was mostly explained by the  
138 Hox genes alone (Fig. 2f). This was expected, since the stacking hypothesis was primarily informed by  
139 the sequential expression of Hox genes in mesoderm derivatives<sup>11,19</sup>.

140 Unexpectedly, we found a much stronger correlation ( $\rho=0.62$ ,  $p=1.4\times 10^{-4}$ ) between gene order and the  
141 medio-lateral axis (M-L) (Fig. 2e). The most anterior genes appeared to be largely expressed close to  
142 the midline of the arm, while more posterior genes were expressed more laterally on either side of the  
143 midline. To confirm that the observed correlations were robust and not the result of sensitivity to the AP  
144 patterning gene ranking assigned from the *in situ* hybridization expression data, we simultaneously  
145 shuffled the gene ranking within each of the 7 groups and probabilistically sampled the position for each  
146 gene, using the expression z-score as a law of probability, over  $10^6$  replications. For all the simulated  
147 replications, the distribution of M-L correlations was significantly superior to the P-D and O-A  
148 correlations, with a two-sided Wilcoxon-rank test  $p$ -value inferior to  $10^{-4}$  in both cases (Fig. 2e). This  
149 suggests that neither the duplication nor the stacking hypotheses accurately describe the deployment of  
150 AP-related patterning genes in *P. miniata*, and that most of the underlying patterning logic of the  
151 pentaradial plan is not explained by existing models.

## 152 M-L deployment of AP patterning genes

153 While RNA tomography provides coarse average axial positional information, these patterns cannot be  
154 directly linked to germ layers or anatomical structures. To investigate axial patterning in more detail,  
155 we examined the expression pattern of the 36 marker genes using *in situ* hybridization chain reactions  
156 (HCR, Supplementary Fig. 7) on post-metamorphic juveniles. This representative stage follows the  
157 resorption of larval structures into the rudiment during metamorphosis (Fig. 3a). The elaboration of the  
158 pentaradial symmetry is initiated earlier within the larval mesoderm, but the mechanistic basis of this  
159 process is likely distinct from those that give their regional identity to the final axes of the pentaradial  
160 body plan. Classical ectoderm AP patterning genes such as *six3/6*, *nkx2.1*, *dmbx*, *pax6* and *otx* initiated  
161 expression at low levels in the presumptive adult tissues at the onset of metamorphosis, but only reached  
162 robust levels of expression that were relatable to ambulacral polarity in post-metamorphic juveniles,  
163 when the definitive adult body plan is elaborated (Fig. 3b; Supplementary Fig. 8). We focused on early  
164 juvenile stages as a readout of early establishment of axial patterning. As in the adult, the juvenile body  
165 plan is organized into ambulacral territories on the oral side and interradial territories at the edge of the  
166 ambulacra and extending around to envelope the entire aboral side. The ambulacral ectoderm is divided  
167 in two main regions: the medial ambulacral ectoderm comprising the radial nerve cords (RNCs) and the  
168 circumoral nerve ring (CNR), and more laterally the epidermis covering the podia (Fig. 3c,d).

169 We analyzed the expression of four groups of patterning genes expressed along the AP axis in both  
170 hemichordates and chordates. The first group (*fzd5/8*, *nkx2.1*, *rx*, *sfrp1/5*, *foxG*, *six3/6*, *hedgehog*) has  
171 strong anterior ectodermal localizations in the proboscis of *S. kowalevskii*<sup>4,5</sup> and in the forebrain of  
172 vertebrates<sup>31,32</sup> (Fig. 1a,b). In *P. miniata* we found similarly overlapping patterns of regional expression,  
173 mostly restricted to the developing CNR, the RNCs, and in the case of *six3/6* and *hedgehog*, in repeated  
174 domains where lateral nerves from the RNCs connect with each pair of secondary podia (Fig. 3e-l;  
175 Supplementary Fig. 9a-e). This region corresponded to the most medial part of the ambulacral ectoderm

176 (Fig. 3d). These findings are inconsistent with the duplication hypothesis, which predicts a staggered  
177 expression of anterior to posterior markers along each RNC.

178 The next group (*lhx1/5*, *dmbx*, *tlx*, *irx*, *fezF*, *dbx*, *otx*, *barH* and *pax6*) overlaps with the previous anterior-  
179 class genes in *S. kowalevskii* but with a more caudal distribution in the posterior proboscis and into the  
180 collar<sup>4</sup>; in chordates, these genes are primarily expressed either in the forebrain or the midbrain<sup>3,31</sup> (Fig.  
181 1a,b). In *P. miniata*, expression of this group of genes overlaps in the most medial territory with the  
182 most anterior class genes, but with expanded lateral domains on either side of the RNCs (Fig. 3d, m-s;  
183 Supplementary Fig. 9f-h), into the epidermis covering the podia. *FezF* and *dbx* were only expressed in  
184 a limited number of cells in the ambulacral ectoderm (Supplementary Fig. 10a-c). Only *pax6* was not  
185 expressed in the medial ambulacral ectoderm and was restricted to the podia epidermis, as previously  
186 reported in other echinoderm species<sup>33,34</sup> (Fig. 3t). Thus, the lateral ambulacral ectoderm appears to have  
187 a more posterior molecular identity, similar to that of the hemichordate collar or the vertebrate midbrain.

188 The third category (*gbx*, *wnt3*, *hox1*, *pax2/5/8*) includes genes that have more posterior expression  
189 patterns in hemichordates and chordates (Fig. 1a,b). *Gbx*, *hox1* and *pax2/5/8* are all expressed in  
190 hemichordates in the anterior trunk, close to the boundary with the collar, and in chordates in the  
191 hindbrain and into the midbrain/hindbrain boundary<sup>5,35,36</sup>. *Wnt3* is expressed at the far posterior end of  
192 the AP axis in both phyla<sup>37,38</sup>. In *P. miniata*, we found that these four genes are expressed at the boundary  
193 between the ambulacral ectoderm and the interradial territory (Fig. 3d, u-β; Supplementary Fig. 9i-l).  
194 *Gbx* and *wnt3* were expressed in the outer part of the ambulacral ectoderm, establishing a mutually  
195 exclusive boundary with more anterior genes like *six3/6* (Fig. 3u-x). *Hox1* and *pax2/5/8* were expressed  
196 more laterally compared to *gbx* and *wnt3*: *hox1* outlined the entire ambulacral area, while *pax2/5/8* had  
197 a more complex expression pattern and was primarily expressed between the ambulacra (Fig. 3y-β). We  
198 suggest that in *P. miniata* these genes marked the outer limit of an anterior compartment.

199 Finally, Hox genes are expressed in hemichordates and chordates posteriorly to the collar/trunk and  
200 midbrain/hindbrain boundary, respectively, and are involved in trunk patterning<sup>4,35,39</sup> (Fig. 1a,b). In *P.*  
201 *miniata*, only *hox1* was detected in the ectoderm. *Hox3*, *hox5*, *hox8* and *hox9/10* were expressed in  
202 mesoderm derivatives (Fig. 3γ-ζ). *Hox4* expression was barely above the detection threshold but was  
203 found in the pharynx (Fig. 3η), while *hox7*, *hox11/13a* and *hox11/13b* were expressed in the developing  
204 intestinal tract (Fig. 3θ-κ). These observations are in line with previous reports that Hox expression in  
205 echinoderms is largely restricted to internal germ layers<sup>20-26</sup> and consistent with the Hox-driven O-A  
206 correlation observed in our RNA tomography dataset. In hemichordates and other bilaterians, Hox gene  
207 domains are intercalated during trunk development between the anterior domains and the posterior end  
208 of the animal, which expresses posteriorizing factors such as *wnt3*<sup>35,40</sup>. We propose that there is no  
209 ectoderm equivalent to a trunk region in *P. miniata*, because *wnt3* is expressed at the edge of the  
210 ambulacral region, and because *hox1* is the only Hox gene expressed in the ectoderm. Therefore, the  
211 deployment of the AP patterning system in *P. miniata* seems to be limited to the ambulacral region and  
212 its boundary.

213 We also found that in *P. miniata* some ancestral deuterostome AP patterning genes did not exhibit  
214 conserved relative expression relative to hemichordates and vertebrates, either because they were not  
215 detected (*dlx*), not expressed in the ectoderm (*engrailed*, *sfrp3/4*), or exhibited different relative spatial  
216 arrangements (*lhx2/9*, *emx*, *otp*) (Supplementary Fig. 10d-i). Some well-established boundaries in  
217 hemichordates and vertebrates like the abutting *otx* and *gbx* domains were also not observed in *P.*  
218 *miniata*. This presumably reflects the plasticity of the AP patterning system and its adaptation to the  
219 radically different pentaradial body plan. Despite these discrepancies, germ-layer specific expression  
220 patterns corroborate that the M-L dimension revealed by RNA tomography, is the main and unexpected  
221 driver of the ectoderm AP patterning logic of the pentaradial body plan in *P. miniata*.

222 If the ancestral deuterostome AP patterning system is used to remodel the sea star body plan at  
223 metamorphosis, are other correlated bilaterian axial patterning systems contributing to the patterning of  
224 the adult body plan as well? In bilaterians, the dorso-ventral (DV) axis is specified transiently during  
225 embryogenesis by conserved BMP2/4 and Chordin gradients<sup>41,42</sup>. Later on, the neuroectoderm and



226 mesoderm derivatives are patterned along the DV axis via the deployment of transcription factors such  
227 as *pax3/7*, *msx* and *tbx2/3* in response to BMP signaling<sup>43-45</sup> (Extended Data Fig. 2a). To investigate  
228 whether the BMP/Chordin axis is involved in the formation of the adult body plan in *P. miniata*, we first  
229 looked at the expression of genes involved in DV axis specification through metamorphosis and in post-  
230 metamorphic juveniles (Extended Data Fig. 2b-s; Supplementary Fig. 4). While *chordin* was not  
231 detected by *in situ* hybridization, we found that *BMP2/4*, *BMP1* and *ADMP1* were consistently  
232 expressed in the distal part of the developing tube feet mesoderm, starting early in metamorphosis and  
233 into the juvenile (Extended Data Fig. 2c-s). In addition, we looked at known target genes of BMP  
234 signaling (Extended Data Fig. 2b,t-v; Supplementary Fig. 4). While *pax3/7* is lost in echinoderms<sup>46</sup>,  
235 *tbx2/3* and *msx* were co-expressed with *BMP2/4* in the tube feet, and *tbx2/3* also expressed in the  
236 overlying tube feet ectoderm, suggesting a similar relationship to BMP signaling than in vertebrates  
237 (Extended Data Fig. 2t-v). These expression data are consistent with previous studies from a direct-  
238 developing echinoid<sup>47</sup> and support a role for BMP signaling in tube feet development, more equivalent  
239 to the later patterning role of BMP in dorsal midline patterning in hemichordates and vertebrates<sup>43,48</sup>.  
240 However, our results do not indicate BMP/Chordin signaling as defining an anatomical axis in the  
241 pentaradial body plan of *P. miniata*.

242 Echinoderm rays have also been proposed<sup>49,50</sup> to be homologous to bilaterian appendages<sup>49,50</sup>. In principle,  
243 this scenario could be compatible with our expression data, with the rays being appendages of an anterior  
244 territory. To assess this possibility, we considered conserved markers for proximo-distal appendage  
245 development, which involves polarized expression of *meis*, *pbx*, *dlx* and *sp8-9* in vertebrates, arthropods  
246 and cephalopods<sup>51</sup> (Extended Data Fig. 2w,x; Supplementary Fig. 4). Although we found the expression  
247 of *Pbx* and particularly *sp8-9* to be compatible with this idea, *dlx* was not detected and *meis* had an  
248 inconsistent expression domain (Extended Data Fig. 2x-a). The extension of the echinoderm ambulacra  
249 likely involves substantial developmental novelty. The RNA tomography dataset will be an invaluable  
250 resource for hypothesis generation and identification of candidate genes to form the basis of a more in  
251 depth developmental investigations of this process.

## 252 Evolution of axial properties in echinoderms

253 The organizational modifications to the ancestral bilateral deuterostome body plan during early  
254 echinoderm evolution are so profound that even basic axial comparisons with other deuterostome taxa  
255 have been problematic at the morphological level<sup>2</sup>. Here we investigated this fundamental question by  
256 spatially mapping the deployment of the ancestral bilaterian ectodermal AP patterning system in the  
257 pentaradial body plan of *P. miniata*. Since this patterning system is largely conserved between  
258 hemichordates and chordates, we can confidently reconstruct its ancestry in early deuterostomes and at  
259 the base of ambulacrarians<sup>4,5</sup>.

260 We found that that much of the ancestral anterior patterning network is spatially deployed in a manner  
261 incompatible with previously proposed hypotheses of echinoderm axial homologies<sup>11,15</sup>. Rather,  
262 expression patterns map onto a novel coordinate system that we call the “ambulacral-anterior” model of  
263 echinoderm body plan evolution (Fig. 4a,b). In this model, the midline of each ambulacrum expresses  
264 the most anterior bilaterian molecular identity, equivalent to the forebrain and proboscis in vertebrates  
265 and hemichordates, respectively. The mid-lateral regions on either side of the nerve cords, including the  
266 ectoderm wrapping the podia, share patterning similarities with more caudal ectodermal territories of  
267 hemichordates and chordates, down to the collar and midbrain, respectively. Finally, the boundary at  
268 the edge of the ambulacral ectoderm displays the most posterior molecular profile corresponding to the  
269 collar/trunk boundary of hemichordates and the midbrain/hindbrain boundary of vertebrates. According  
270 to our ambulacral-anterior model, echinoderms are the first example of bilaterians in which the  
271 “anterior” identity is located at the center of a sheet of tissue, rather than being located at an extremity.  
272 The anatomical outputs of this anterior domain, however, share similarities across all deuterostome  
273 phyla (including echinoderms) in that they include neural condensations and an extensive array of  
274 sensory structures.

275 Strikingly, despite the presence of a genomic Hox cluster, ectodermal Hox gene expression is largely  
276 absent in the sea star (except for *hox1*), suggesting a loss of the ancestral ectodermal trunk regulatory  
277 program, which is consistent with previous observations in the echinoid *Peronella japonica*<sup>27</sup>. Yet, Hox  
278 genes are expressed in the mesoderm and endoderm, displaying a marked uncoupling of germ layer AP  
279 patterning as recognized previously<sup>27,52</sup>. The mesoderm and endoderm trunk programs are wrapped in  
280 an ectoderm with two clear territories, including (1) an ambulacral anterior-like domain that is the main  
281 focus of our work, and unexpectedly (2) an extensive interradian domain that extends around the aboral  
282 side of the animal and displays uncertain axial identity, without any detectable readout of the ancestral  
283 AP patterning program, as previously observed in echinoids<sup>27</sup>. The uncoupling of an ectodermal head  
284 and trunk programs is not unique to *P. miniata* and has been demonstrated in both larval echinoderms  
285 and hemichordates<sup>35,53</sup>, and more recently in protostome larvae<sup>40,54</sup>, suggesting that these regulatory  
286 programs can be uncoupled over macro-evolutionary time frames.

287 Previous studies<sup>13,14</sup> proposed that the deployment of transcription factors to pattern adult echinoderm  
288 body plans could be the result of co-option, meaning recruitment of an existing gene or regulatory  
289 module (or duplication thereof) to fulfill a new developmental role<sup>55</sup>. Here, using a much larger set of  
290 genes, we demonstrate that the patterning of the ambulacral ectoderm in *P. miniata* involves a collection  
291 of interacting modules that define distinct regions along the AP axis of bilateral animals rather than a  
292 single regulatory program. Cooption of this network would require redeployment of these interacting  
293 modules into a novel role in stem echinoderms. Thus, while we acknowledge that co-option remains a  
294 possible scenario, we propose that the observed pattern in extant asteroid ambulacra is a result of  
295 modification of an ancestral deuterostome axial program during echinoderm evolution. Documented  
296 cases of co-option such as the deployment of posterior Hox genes during limb patterning<sup>56</sup> involve the  
297 (re)use of an existing regulatory network to pattern a new structure, without affecting the ancestral role  
298 of the system. To our knowledge there is no example of co-option that involves dismantling of the  
299 ancestral patterning networks following their recruitment into the development of a novel structure. A  
300 co-option scenario in echinoderms would imply that the ancestral function of the AP patterning network  
301 was lost in the adult body plan, after co-option into an evolving ambulacral territory, which in our  
302 opinion is less parsimonious than proposing descent with modification of an ancestral AP axis.  
303 Interestingly, a route to resolve this issue might come from some interpretations of homalozoan and  
304 helicoplacoid echinoderms that propose these groups to be transitional forms between bilateral and  
305 pentaradial symmetry<sup>57</sup>. Re-interpreting these fossils in light of new patterning datasets could allow us  
306 to discriminate between co-option with loss of the ancestral axial registry, or descent with modification.

307 Did the reorganization of the ancestral AP axis documented here occur during the early stem evolution  
308 of echinoderms as a defining regulatory feature of crown group echinoderms, or during the later  
309 diversification of asteroids? Our model inferred from analysis of asteroid data is consistent with  
310 published findings from two other major echinoderm clades, crinoids and echinoids. In crinoids, the  
311 expression of *six3/6*, *otx*, and *pax6* has been reported in the ambulacral ectoderm<sup>58</sup>. In echinoids, AP  
312 patterning related genes have been surveyed in various species including *Strongylocentrotus purpuratus*  
313 and *Heliocidaris erythrogramma*<sup>20,33,34</sup>, and most extensively in *Peronella japonica*<sup>25,27</sup> where the  
314 anterior identity of the ambulacral ectoderm was already reported<sup>27</sup>. In both crinoids and echinoids, most  
315 of the anterior patterning related genes studied show expression patterns compatible with the  
316 ambulacral-anterior model. Furthermore, Hox gene expression has been described in echinoids,  
317 holothuroids, and crinoids<sup>20-26</sup> and is also largely congruent with the expression patterns that we  
318 observed in *P. miniata*. This suggests that based on limited available comparative data, the ambulacral-  
319 anterior model could be a general development feature of all extant echinoderm classes and that our  
320 observations reflect early regulatory changes that occurred during stem echinoderm evolution (Fig.4c).  
321 Further comprehensive comparative analyses will be required to test this hypothesis.

322 The evolution of body plan variations across echinoderm classes has proven challenging to reconstruct  
323 based on morphological features alone<sup>59</sup>. The ambulacral-anterior model offers a powerful tool to  
324 establish robust regional homologies between echinoderm classes. Most importantly, the new axial  
325 paradigm established here can be integrated with the exquisite fossil record of the phylum to  
326 reinvestigate key morphological transformations in light of regulatory changes.

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462 **Figure legends**

463 Figure 1: Deployment of the antero-posterior patterning system in deuterostomes

464 **a**, Expression map of the conserved transcription factors and signaling ligands involved in ectoderm  
465 patterning along the AP axis, as observed in the hemichordate *Saccoglossus kowalveskii*. **b**, Previous  
466 work in chordates and hemichordates has demonstrated extensive regulatory conservation in ectodermal  
467 AP patterning, establishing the ancestral regulatory characteristics of early deuterostomes. How this  
468 system is deployed in echinoderms remains unclear. **c**, Four hypotheses have been proposed for the  
469 deployment of the AP patterning system in echinoderm adult body plan: bifurcation, circularization,  
470 duplication and stacking.

471 Figure 2: RNA tomography reveals the medio-lateral dimension of the arms as the main driver of antero-  
472 posterior patterning system deployment in *Patiria miniata*

473 **a**, Reconstructions of a young juvenile *P. miniata* scanned by micro-CT and segmented to highlight the  
474 endoskeleton (grey), the digestive tract (yellow), the main body muscles (red), the water vascular system  
475 (purple) and the central nervous system (blue). The ambulacral areas correspond to the grooves  
476 harboring the podia and are separated by interradial areas. Scale bar = 1mm. **b**, Experimental design of  
477 the RNA tomography with the cutting plans along the proximo-distal (P-D), oral-aboral (O-A) and  
478 medio-lateral (M-L) dimensions. **c**, Heatmap of gene expression z-scores along the P-D, O-A and M-L  
479 dimensions of the RNA tomography. **d**, Expression profiles of AP patterning related genes along the P-  
480 D, O-A and M-L dimensions of the RNA tomography. For the M-L dimension, the dotted line indicates  
481 the midline. Genes are ranked from the most anterior to the most posterior based on their expression  
482 patterns in *Saccoglossus kowalevskii*. **e**, Spearman correlations between the ranking of the AP patterning  
483 related genes and their position along the three dimensions of the RNA tomography. Dots indicate the  
484 raw correlation values. Boxplots indicate the distribution of correlation values when determining the  
485 gene position along each dimension probabilistically, and simultaneously shuffling the ranking of the  
486 AP patterning related genes for each group ( $n = 10^6$  independent samples). The distribution of  
487 correlations for the M-L dimension is significantly higher than for the P-D and O-A dimensions (Two-  
488 sided Wilcoxon-rank test;  $p < 10^{-4}$ ). Centre lines: median; box: interquartile range (IQR); whiskers:  
489 highest and lowest values at  $\pm 1.5 \times \text{IQR}$ . **f**, Relative contribution of each group of AP patterning related  
490 genes to the Spearman correlation between the ranking of the AP patterning related genes and their  
491 position along the three dimensions of the RNA tomography.

492 Figure 3: Gene expression data reveals the deployment of the antero-posterior patterning system in  
493 *Patiria miniata* ambulacral ectoderm

494 **a**, Metamorphosis in *P. miniata*. The anterior part of the brachiolaria larva (cyan asterisk) is resorbed  
495 into the rudiment. **b**, HCRs showing *pax6* (red) and *otx* (cyan) starting to be expressed during  
496 metamorphosis in the developing adult pentaradial body plan. Specimens are counterstained with DAPI  
497 (blue). **c**, Main anatomical features of post-metamorphic juveniles, imaged from the oral side: the  
498 nervous system (cyan) stained with antibodies against acetylated-tubulin highlighting the nerve tracts  
499 and the neuronal marker *elav* highlighting the cell bodies, the hydrocoel stained with the marker *patched*,  
500 the digestive tract stained with the marker *lox* and the endoskeleton stained using calcein. RNC: radial  
501 nerve cord; CNR: circumoral nerve ring; RaC: radial canal; RiC: ring canal; St: stomach **d**, Schematics  
502 showing the oral-aboral position of the HCR z-projections shown in **e-k** (top) and the main anatomical  
503 regions visible on the post-metamorphic juveniles oral side (bottom). The white square on the bottom  
504 panel indicates the position of the close-ups in **l,t,w,x, $\alpha,\beta$** . **e-k**, HCRs of *P. miniata* juveniles imaged  
505 from the oral side (**e- $\eta$** ) or the aboral side ( **$\theta$ - $\kappa$** ). In **e-k,m-s,u,v,y,z, $\gamma$ - $\kappa$** , specimens are counterstained  
506 with DAPI (blue). **l,t,w,x, $\alpha,\beta$** , magnification of a single ambulacrum. **e-l**, Genes primarily expressed in  
507 the medial ambulacral ectoderm. Colocalization with *elav* indicates expression in the CNR and the  
508 RNCs. **m-t**, Genes primarily expressed in the podia epidermis. **u- $\beta$** , Genes primarily expressed in the  
509 ambulacral boundary. In  **$\alpha,\beta$** , orange dotted lines outline the ambulacral ectoderm.  **$\gamma$ - $\zeta$** , Hox genes

510 primarily expressed in the coeloms.  $\eta$ - $\kappa$ , Hox genes primarily expressed in the digestive tract. In  $\theta$ - $\kappa$ ,  
511 red asterisks indicate the position of the developing intestinal tract. Scale bars = 100 $\mu$ m.

512 Figure 4: The ambulacral-anterior model of echinoderm body plan evolution

513 **a**, Expression map of the conserved transcription factors and signaling ligands involved in the  
514 ambulacral ectoderm patterning in *Patiria miniata*, organized from the midline of the ambulacrum (left)  
515 towards the interradius (right). **b**, Diagram of the ambulacral-anterior model in a generalized asteroid.  
516 Anterior patterning genes are expressed in the oral ambulacral ectoderm, shown through a cut-away of  
517 the forward part of the animal and a cross-section through the forward arm. Only genes expressed in the  
518 ectoderm are shown. **c**, Ambulacral-anterior model applied to echinoderm classes with available gene  
519 expression data, echinoids (left) and crinoids (right). In echinoids, the ambulacral ectoderm is  
520 internalized and shown through a cut-away of the forward part of the test, while in crinoids the  
521 ambulacral ectoderm is embedded in the oral epidermis, similar to asteroids.

522



## 523 **Methods**

### 524 Animal husbandry and tissue preparation

525 All animal collection and handling was done in compliance with US ethical regulations. Adult and young  
526 (~1-2cm wide) *Patiria miniata* specimens were collected off the coast of Monterey bay, California, US,  
527 and kept in circulating sea water tanks. Young specimens were used for X-ray micro-computed  
528 tomography (micro-CT), extract genomic DNA and RNA tomography (see below). For RNA  
529 tomography datasets and genomic DNA (see below), two young specimens of the same class of size  
530 than those used for micro-CT (referred to as specimens #1 and #2) were used. They were anesthetized  
531 in filtered sea water and 7.5% MgCl (1:1), and then four arms were dissected out. One of the arm was  
532 proceeded for DNA extraction, and the three others were included in HistoPrep embedding medium  
533 (ThermoFischer). Finally, to generate fixed material for HCRs and immunohistochemistry (see below),  
534 several batches of gravid adults were spawned by injecting 1ml of 1 $\mu$ M 1-methyladenine (Acros  
535 Organics) in each of the gonads. Sperm and mature oocytes were released by the animals about 45' and  
536 90' after the injection, respectively. Following *in vitro* fertilization, embryos were cultured at 14°C in  
537 UV-sterilized filtered seawater, first at a density of about 100 embryos per mL and within 3L glass jars  
538 oxygenated by a motorized paddle. At 48 hours post fertilization, the culture concentration was adjusted  
539 to about 1 larva per mL, and from that point about 95% of the seawater was renewed every 2 or 3 days.  
540 Following water renewal, the larvae were fed *ad libitum* with freshly grown *Rhodomonas lens*  
541 microalgae. Brachiolaria larvae started to settle on the glass jars and to undergo metamorphosis between  
542 1 and 2 months post fertilization. After metamorphosis, juveniles were collected from the glass jars by  
543 being first relaxed in a 1:1 mix of 7.5% MgCl<sub>2</sub> and filtered seawater and then being gently detached  
544 using a paintbrush.

### 545 X-ray micro-computed tomography

546 For X-ray micro-CT analyses, *P. miniata* young juveniles of the same size than those used for RNA  
547 tomography were relaxed in a 1:1 mix of 7.5% MgCl<sub>2</sub> and filtered seawater for 10', and then fixed in a  
548 modified PHEM buffer<sup>60</sup> (90mM PIPES, 37mM HEPES, 14mM EGTA, 30mM MgCl<sub>2</sub>, 9% fructose)  
549 overnight at 4°C. The samples were then washed in modified PHEM buffer, before being counterstained  
550 in 2% osmium tetroxide in deionized water for two hours. Following staining, the samples were washed  
551 extensively in deionized water, gradually dehydrated in ethanol and stored in ethanol at 4°C. Prior to  
552 imaging, samples were dried and then pinned on their aboral side to an imaging support. Scanning was  
553 carried out using a Zeiss Xradia 520 Versa 3D X-ray microscope. Specimens were scanned at a  
554 resolution of 12.138 $\mu$ m per voxel. In order to identify different tissues in the scans, regions of interest  
555 were segmented using Dragonfly v.2022.2. For segmentation, the deep learning tool was used to train a  
556 3D U-net model, which has been shown to be ideal for rapid and accurate automatic biomedical image  
557 segmentation<sup>61</sup>. The U-net model had nine classes, corresponding to different tissue types, and was  
558 trained for 75 epochs (15.5 hours) using a training set of fifteen manually segmented slices from the  
559 dataset. During model training, a data augmentation factor of 2 was used, including horizontally and  
560 vertically slipping the data, rotation, shearing, and zoom. Following model training, this U-net was used  
561 to automatically segment and identify our nine different tissue classes. Following the initial deep  
562 learning-based segmentation, the nine classes were reduced to the five regions of interest shown in the  
563 3D model (endoskeleton, digestive tract, musculature, water vascular system, and central nervous  
564 system) *via* the merging of classes relating to similar tissue types (for instance, combining classes  
565 pertaining to different types of endoskeletal tissue). At this point, further segmentation was carried out  
566 manually in Dragonfly to clean up regions of interest and to produce 3D models and mesh files for  
567 visualization. After segmentation, the consistency between the size of the anatomical features  
568 reconstructed from the micro-CT scans and the extent of tissue marker expression profiles from the  
569 RNA tomography was checked manually. The samples used for micro-CT were registered at the Natural  
570 History Museum (London, UK) under the registration number NHMUK 2023.263. The segmented scans  
571 and reconstructed mesh files were deposited on Morphosource under the project number 000529415.

572 Genomic DNA isolation

573 Genomic DNA (gDNA) for genome sequencing was isolated from a dissected arm of the young *P.*  
574 *miniata* specimen #1. Using an extended handle conical tip pestle (Bel-Art Proculture), the arm was  
575 homogenized in the presence of the extraction buffer and of proteinase K. Genomic DNA was then  
576 isolated using the DNeasy Blood and Tissue Kit (Qiagen) following the manufacturer's instructions.

577 Genome sequencing

578 *Ultra-Low Input HiFi library preparation*

579 Using gDNA from the arm of specimen #1, we generated a HiFi genome of *P. miniata*. The general  
580 workflow is described in Supplementary Fig.2e. As the gDNA isolated from the arm was predominantly  
581 shorter than the 10-15 kb which is recommended size for HiFi genomic library creation, a size selection  
582 was performed prior to doing an Ultra-Low Input (ULI) amplification and library preparation to remove  
583 fragments <7kb. The size selection was done on a SAGE BluePippin system using the 0.75% Agarose  
584 Dye-free Gel Cassette and the S1 Marker (SAGE). Approximately 100ng of DNA was recovered post  
585 size selection and used as input for the ULI PCR-based HiFi library protocol. The sample was amplified  
586 using the SMRTbell gDNA Amplification Kit (PacBio) and a HiFi SMRTbell library was constructed  
587 using the SMRTbell Express Template Prep Kit 2.0 (PacBio) following manufacturer's recommended  
588 protocol. After library construction, a final size selection was performed on the SAGE BluePippin as  
589 previously described using a size cut-off of 7kb. Library size was characterized on an Agilent 2100  
590 BioAnalyzer using the DNA 12000 kit (Agilent). The additional size selection ensured having a final  
591 library with a fragment size range greater than 7kb.

592 *HiFi reads sequencing*

593 Sequencing reactions were performed on the PacBio Sequel II System with the Sequel Sequencing Kit  
594 2.0 (PacBio). The kit uses a circular consensus sequencing (CCS) mode which provides >99.5% single  
595 molecule read accuracy<sup>62</sup>. The samples were pre-extended without exposure to illumination for 2 hours  
596 to enable the polymerase enzymes to transition into the highly progressive strand-displacing state and  
597 sequencing data was collected for 30 hours to ensure maximal yield of high-quality CCS reads. CCS  
598 reads were generated from the data using the SMRT Link Version 9.0 (PacBio). For the genomic HiFi  
599 sequencing, the library was bound to the sequencing enzyme using the Sequel II Binding Kit 2.2 and  
600 the Internal Control Kit 1.0 (PacBio). The HiFi reads generated 15,614,751 HiFi reads with a mean read  
601 length of 9,039 bp  $\pm$  1,671bp (Supplementary Fig. 3e).

602 *Genome assembly*

603 Prior to *de novo* genome assembly, the reads were trimmed on the ends to remove any PCR primer  
604 sequences from the ultra-low amplification process using lima v.2.2 (PacBio). The forward sequence of  
605 the amplification adapter used was AAGCAGTGGTATCAACGCAGAGTACT. Once the HiFi reads  
606 were trimmed, a filtering step was performed to remove duplicate reads from the PCR step using SMRT  
607 link v.9.0. After removing duplicate reads, the HiFi reads (~100X genomic coverage) were used to  
608 generate a draft diploid assembly using Hifiasm v.0.15<sup>63</sup>. This resulted in two highly contiguous  
609 haplotype primary assemblies of 680 Mb and 674 Mb respectively (Supplementary Table 5). Assembly  
610 completeness was assessed with the Benchmarking Single Copy Ortholog<sup>64</sup> (BUSCO v.3.0) gene set for  
611 Metazoa at 94.8% for each of the individual haplotype primary assemblies and 96.2% overall for the  
612 diploid assembly as a whole (Supplementary Fig.3d; Supplementary Table 4). The two haplotypes were  
613 deposited at DDBJ/ENA/GenBank under the accession number [JAPJSQ000000000](#) and  
614 [JAPJSR000000000](#).

## 615 RNA tomography section preparation and RNA extraction

616 The three arms embedded in HistoPrep medium were cryosectioned using a Leica CM3050-S microtome  
617 along the appropriate dimensions: proximo-distal (P-D), oral-aboral (O-A) or medio-lateral (M-L)  
618 (Supplementary Fig.2a-c). The blocks used for the P-D and O-A dimensions came from the specimen  
619 #1, while the block used for the L-R dimension came from the specimen #2. Slice thickness was set to  
620 25µm for the P-D and O-A dimensions and to 30µm for the M-L dimension, resulting in total in 430,  
621 160 and 160 slices for the three respective dimensions. While sectioning the blocks, every 20 (P-D,  
622 O-A) or 10 (M-L) contiguous slices were pooled together into 1.5mL tubes. These resulted in a total of  
623 22 tubes for the P-D dimension, 8 tubes for the O-A dimension and 16 tubes for the L-R dimension  
624 (Supplementary Fig.2d). Each tube was then processed for RNA extraction using a modified  
625 Trizol/RNeasy RNA extraction protocol<sup>65</sup>. In each tube the slices were homogenized in 1mL of Trizol  
626 using an extended handle conical tip pestle (BelArt Proculture). After vigorously mixing the Trizol  
627 homogenate with chloroform, each tube was centrifuged at 10,000xg RCF for 18' at 4° C. The aqueous  
628 phase containing the RNA was carefully removed and the RNA was further purified using the RNeasy  
629 Plus Micro Kit (Qiagen) following the manufacturer's instructions.

## 630 RNA tomography

### 631 *Barcoded cDNA IsoSeq SMRTbell library preparation*

632 Using RNA isolated from the three sets of cryosections, we generated a RNA tomography<sup>28,66</sup> dataset  
633 for *P. miniata* juveniles. The general workflow is described in Supplementary Fig.2e. Barcoded PacBio  
634 IsoSeq SMRTbell libraries were constructed using the SMRTbell Express Template Prep Kit 2.0  
635 (PacBio) following the manufacturer's instructions. We used a set of 22 barcoded sequences  
636 (Supplementary Table 1) which were combined with each RNA extracts (Supplementary Table 2).  
637 Typically, 12-14 PCR amplification cycles were used to generate enough barcoded double-stranded  
638 cDNA for the library preparation and subsequent sequencing runs (Supplementary Table 2).

### 639 *Library sequencing*

640 For the IsoSeq transcript libraries, library was bound to the sequencing enzyme using the Sequel II  
641 Binding Kit 2.1 and Internal Control Kit 1.0 (PacBio) and the sequencing reactions were performed on  
642 the PacBio Sequel II System. The three dimensions were sequenced independently, since there were  
643 shared barcodes between the different libraries. We obtained a total of 71,582,642 reads (Supplementary  
644 Table 3) with a mean read length of 3,843bp, 3,152bp and 2,450bp for the P-D, O-A and L-R  
645 dimensions, respectively (Supplementary Fig.3a). In addition, HiFi read length distributions were  
646 consistent across each barcode within the respective RNA tomography dimension (Supplementary  
647 Fig.3b). Sequence read archives were deposited at DDBJ/ENA/GenBank under the bioproject  
648 [PRJNA873766](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA873766) and individual accession numbers for each barcode are provided in Supplementary Table  
649 3. Recent advancements in full length transcript concatenation protocols<sup>67</sup>, as well as higher multiplexed  
650 SMRT sequencing flow cells have increased throughput with a concomitant decrease in sequencing cost  
651 over a single 24 hour run. Genomic, epigenetic and transcript data sets can now all originate from the  
652 same individual greatly improving the mappability and subsequent analysis of transcripts in highly  
653 polymorphic non-model organisms where gene sequences can differ by >5% in coding regions and as  
654 much as 40% in UTR sequence which complicate short read transcript alignment and reliable  
655 quantitation.

### 656 *IsoSeq reads demultiplexing and refining*

657 For each HiFi read file generated, the data was demultiplexed into barcode specific read files using lima  
658 v.2.2 (PacBio) and the barcodes listed in Supplementary Table 1. Once the data were demultiplexed  
659 each read file was refined to include only full length non-chimeric reads using isoseq3 v.3.4.0<sup>68</sup>.  
660 Chimeras were identified by inclusion of 5' or 3' RT-PCR primer sequences internal to the initial 'full  
661 length' HiFi read. The primer sequences used were NEB\_5P (GCAATGAAGTCGCAGGGTTGGG),

662 Clontech\_5P (AAGCAGTGGTATCAACGCAGAGTACATGGGG), and NEB\_Clontech\_3P  
663 (GTACTCTGCGTTGATACCACTGCTT). Transcript clusters were identified using Cupcake  
664 v.25.2.0<sup>69</sup> which leverages a genomic reference alignment-based strategy to identify redundant  
665 isoforms/transcripts for gene loci. We leveraged the diploid assembly to minimize dropouts that might  
666 be caused by haplotype specific null alleles and/or poor mapping between haplotypes for the same locus.  
667 Comparative alignment of the two haplotype derived transcriptomes facilitated further collapse of the  
668 gene set to one representative per loci.

#### 669 *Transcriptome generation and curation*

670 Using each haplotype primary assembly, the complete set of full length non-chimeric transcript reads  
671 (FLNC) were clustered and collapsed to reduce gene redundancy while maintaining the highest possible  
672 level of gene completeness. The complete set of FLNC reads across all three combined RNA  
673 tomography datasets were clustered from a list of input bam files based on each of the haplotype specific  
674 primary assemblies. Minimap2 v.2.21<sup>70</sup> was used to align the FLNC reads to each primary assembly and  
675 Sqanti2 v.7.4<sup>71</sup> was used to cluster and filter redundancies. Once minimal transcript sets were obtained  
676 for each haplotype, a comparative alignment between the sets was performed using Minimap2 v.2.21  
677 aligner to find unique transcripts between the two. From the output paf files, transcripts unique to  
678 haplotype 2 which were not present in the haplotype 1 were identified, filtered out from the haplotype 2  
679 set, and added to the Haplotype 1 set to obtain a more complete single copy transcript set.

680 Manually curated developmental genes of interest were identified (see below). All sequence in our  
681 “single copy” transcript set were aligned using TBLASTX to the manually curated set to (1) remove  
682 any duplication of these key transcripts in our reference transcriptome (e.g. the high polymorphism rate  
683 resulted in both copies of some of the manually curated genes being present) and (2) remove partial  
684 duplications (non-full length transcripts matching the manually curated set). The result of this curation  
685 step was to ensure that for every gene of interest, we had only one sequence in our reference  
686 transcriptome to maintain accurate downstream quantification. Given the occasional duplication of  
687 transcripts in our reference transcriptome (eliminated for our genes of interest), it is assumed that the  
688 un-curated transcripts have some level of duplication.

689 Once we derived a near complete curated transcriptome for *P. miniata*, the barcode specific  
690 demultiplexed RNA tomography datasets were aligned to the transcriptome reference using Minimap2  
691 v.2.21. The final refined *P. miniata* transcriptome comprised 25,794 transcripts and represented a nearly  
692 complete (91.5%) set of metazoan BUSCO genes (Supplementary Fig.3d; Supplementary Table 4).  
693 Transcript expression counts for each of the 25,794 gene model in our reference transcriptome were  
694 tallied using a simple Perl v.5.30.1 script that only counted primary alignments (no supplementary  
695 alignments) with a quality value of 15 or greater. Each section was tallied independently and the data  
696 merged. The alignment counts for each section were normalized to the total reads for each barcode in  
697 order to allow spatial comparisons to be made across each dimension of the RNA tomography dataset.  
698 This was done to account for variable recovery of total RNA in each barcoded section. (Supplementary  
699 Fig.3c).

#### 700 Orthologues identification

701 *P. miniata* orthologues of developmental genes of interest, which included 36 AP patterning related  
702 genes, 6 DV specification and patterning related genes, 3 limb proximo-distal patterning related genes,  
703 the pan-neuronal marker *elav*, the gut marker *lox* and the hydrocoel marker *patched* were identified from  
704 the FLNC reads by reciprocal best blast hit and validated by phylogenetic trees (Supplementary Fig.4).  
705 Nucleotide sequences for these transcripts were deposited at DDBJ/ENA/GenBank and accession  
706 numbers are provided in Supplementary Table 6. Trees were calculated with both the Maximum  
707 likelihood and Bayesian inference methods. Maximum likelihood trees were calculated in MEGA  
708 v.7.0.26<sup>72</sup> with the robustness of each node being estimated by bootstrap analyses (in 1000  
709 pseudoreplicates). Bayesian inference trees were calculated using MrBayes v.3.1.2<sup>73</sup> in 1,000,000  
710 generations with sampling of trees every 100 generations and a burn-in period of 25%. The branching



711 pattern of the ML tree was retained in the final tree figure, displaying, at each node, the bootstrap support  
712 of the ML analysis as well as the posterior probability support of the BI analysis.

### 713 RNA tomography analyses

714 RNA tomography analyses were performed in R v.4.1.2 using custom-written code. For downstream  
715 analyses, the 22 sections of the P-D were merged pairwise by simple addition of the read counts mapping  
716 for each transcripts, bringing the final number of sections in the P-D dimensions to 11 (Supplementary  
717 Fig.2d). In addition, section 14 of the M-L dataset was removed because it yielded a total read count  
718 77% lower than the average total read count per section for the M-L dimension, which could have biased  
719 the quantification analyses. To maintain the symmetry in the M-L dimension, section 3 was also  
720 removed, bringing the final number of sections in the M-L dimension to 14 (Supplementary Fig.2d).

721 Individual read counts from each section of the three RNA tomography dimensions were normalized  
722 against the total read count of the section in order to account for sequencing depth differences between  
723 the three dimensions and for geometrical disparities between the different sections of a single dimension.  
724 Because we were interested in the profile of highly expressed and variable genes, whereas genes with a  
725 uniform expression across the sections were poorly informative, a cutoff was applied to discard genes  
726 which consistently had the 20% lowest average expression level or 20% lowest variability in all of the  
727 three dimensions, resulting in a final set of 21,847 gene models (Supplementary Fig.3f). Of the 36 AP  
728 patterning related genes investigated in this study, four of them fell below the cutoff and were excluded  
729 from the computational analyses because of their low expression levels: *rx*, *dlx*, *hox11/13a*, and  
730 *hox11/13c* (Supplementary Fig.3f). For further analyses, the expression levels of each gene along each  
731 of the three dimension sections was transformed into z-score.

732 For the clustering analysis, the dimensionality of the dataset was first reduced using a principal  
733 component analysis (PCA) performed simultaneously on the three dimensions of the RNA tomography.  
734 Kaiser-Guttman's criterion<sup>74</sup> was used to select the significant PCs of the PCA. Then, the coordinates  
735 of the transcripts along the retained PCA axes were used to compute the Euclidean distance matrix and  
736 hierarchical agglomerative clustering using Ward's aggregation to produce an expression profile  
737 dendrogram<sup>75</sup>. Following the methodology established in previous RNA tomography studies<sup>28</sup>, we  
738 estimated a maximal number of 12 clusters based on discernible anatomical features in the Spearman  
739 correlation matrices of the sections (Extended Data Fig. 1f). However, the relevant number of clusters  
740 was likely lower than 12 due to the lack of independence of gene expression between the three  
741 dimensions. We thus used a silhouette index<sup>76</sup> to evaluate the consistency of clusters when choosing  
742 between 2 and 12 clusters. The silhouette index indicated that 7 was the most appropriate number of  
743 clusters, and this number also minimized redundancy between the different dimensions.

744 For the hypothesis testing, we determined the ranking of the 36 investigated AP patterning related genes  
745 based on their expression profiles in the hemichordate *S. kowalevskii*, which is the closest bilateral  
746 echinoderm-relative with an extensive dataset of AP patterning gene expression profiles<sup>4,5,37,48,77-80</sup>. Gene  
747 expression patterns in other hemichordate studies are largely consistent with those observed in *S.*  
748 *kowalevskii*<sup>35,81</sup>. The 36 genes were divided into seven groups based on their expression domains  
749 (anterior proboscis (A), proboscis (B), anterior collar (C), posterior proboscis and collar (D), collar-  
750 trunk boundary (E), trunk (F), posterior tip of the trunk (G)) (Supplementary Fig.6). Within each group,  
751 genes were ranked based on their expression pattern, and, when required, tied based on the available  
752 gene expression patterns in other closely related species. We were confident that the assignment of each  
753 gene to the groups A to G based on their expression profile in *S. kowalevskii* was robust to interpretation  
754 biases. On the other hand, given that the genes expressed in these groups are more or less coincident in  
755 parts of the AP axis of *S. kowalevskii*, we recognized that the internal gene ranking within each group  
756 was more subject to possible interpretation biases.

757 To test the Spearman correlation between the AP patterning related gene ranking and the gene  
758 expression profiles along the three dimensions of the RNA tomography, the symmetrical sections of the  
759 M-L were averaged pairwise (1 and 14; 2 and 13; and so on). We then used the highest z-score as a

760 readout for the position of each gene in each dimension, and correlated it to the ranking of the AP genes  
761 as described above. The significance of this raw correlation was assessed by a two-sided Spearman  
762 correlation test. In addition, each of the seven group of genes was independently removed to assess its  
763 contribution to the correlation. However, this approach is subjected to two potential sources of errors,  
764 resulting from (1) choosing the single highest z-score value for the position of genes with multimodal  
765 expression profiles, and (2) and as mentioned above from biases in determining the ranking of the AP  
766 genes within each of the seven gene groups, as mentioned above. To account for these biases, we also  
767 ran  $10^6$  independent correlations, with each correlation (1) probabilistically determining the position of  
768 each gene by using the z-score as a law of probability, and (2) randomly shuffling the ranking of the  
769 genes within the seven gene groups. The distribution of the  $10^6$  correlation values was then compared  
770 across the three dimensions of the RNA tomography using a two-sided Wilcoxon rank test.

### 771 *In situ* hybridization

772 Short *in situ* hybridization antisens DNA probes were designed based on the split-probe design of HCR  
773 v.3.0<sup>82</sup> using HCR 3.0 Probe Maker<sup>83</sup> with adjacent B1, B2 or B3 amplification sequences depending on  
774 the genes (Supplementary Table 7). Between 14 and 33 probe pairs were designed for each gene  
775 including the CDS, and for some of them adjacent 5' and 3' UTRs. The probe pairs were then ordered  
776 as oligo pools (Integrated DNA Technology) and suspended in nuclease-free water at a concentration of  
777 0.5 $\mu$ M.

778 For *In situ* hybridization, *P. miniata* brachiolaria larvae, metamorphosing larvae and juveniles were  
779 incubated in fixation buffer (1X phosphate buffered saline (PBS), 0.1M MOPS, 0.5M NaCl, 2mM  
780 EGTA, 1mM MgCl<sub>2</sub>) containing 3.7% formaldehyde overnight at 4°C. Fixed samples were then  
781 dehydrated in methanol and stored at -20°C for at least 24 hours and up to several months. The samples  
782 were progressively rehydrated in PBS containing 0.1% Tween-20 (PBST). They were permeabilized in  
783 detergent solution (1.0% SDS, 0.5% Tween-20, 150mM NaCl, 1mM EDTA (pH 8), 50mM Tris-HCl at  
784 pH 7.5) 30' for larvae and 2 hours for juveniles. For juveniles, this was step was followed by an extra  
785 permeabilization step in 4 $\mu$ g/mL proteinase K (Sigma-Aldrich) for 10' at 37°, and a postfixation in 3.7%  
786 formaldehyde for 25'. The samples were then extensively washed in PBST, and then in 5X saline sodium  
787 citrate buffer containing 0.1% Tween-20 (SSCT), before being pre-hybridized in hybridization buffer  
788 (Molecular Instruments) for 1h at 37°C. The probes were then added to the hybridization buffer at a  
789 final concentration of 0.05 $\mu$ M and the samples were let to hybridize at 37°C overnight under gentle  
790 agitation. Following hybridization, the samples were washed 4 times 30' in probe wash buffer  
791 (Molecular instruments) at 37°C and then in 5X SSCT at room temperature. They were then pre-  
792 amplified in amplification buffer (Molecular Instruments) for 30'. Meanwhile, H1 and H2 components  
793 of the HCR hairpins B1, B2 or B3 coupled either to Alexa546 or Alexa647 fluorophores (Molecular  
794 Instruments) were incubated separately at 95°C for 90", cooled down to room temperature in the dark  
795 and then pooled together before being added to the amplification buffer at a final concentration of 60nM.  
796 The amplification was then performed overnight at room temperature. The samples were subsequently  
797 washed 4 times 30' in 5X SSCT and incubated in PBST containing 1:1000 DAPI (Invitrogen) for three  
798 hours. Finally, the samples were cleared in a series of 20%, 40%, 60%, and 80% fructose diluted in  
799 NaCl-PBS. Each fructose bath was carried out for at least 1 hour. Clarified samples were mounted in  
800 80% fructose diluted in PBS for imaging, which was done using a Zeiss LSM700 confocal microscope.  
801 For each sample, series of optical sections were taken with a z-step interval of 2-3 $\mu$ m Multichannel  
802 acquisitions were obtained by sequential imaging. Confocal optical sections spanning regions of interest  
803 along the oral-aboral axis were compiled into maximum intensity z-projections using ImageJ v.1.52g<sup>84</sup>  
804 and assemble into figure using Adobe Illustrator v.15.0.0.

805 The specificity of the antisens DNA probes and amplification hairpins was validated by running the  
806 protocol without hairpins and probes or with hairpins alone, and comparing antisens and sense probe  
807 sets for *nkx2.1* (Supplementary Fig.7). The consistency of the expression patterns obtained through this  
808 method was further validated by comparisons of *elav*, *nkx2.1*, *dmbx* and *otx* with colorimetric whole-  
809 mount *in situ* hybridization using single antisens RNA probes as described previously<sup>85</sup> (Supplementary  
810 Fig.7). To ensure reproducibility of the *in situ* hybridization expression patterns, each gene was surveyed

811 in at least two independent experiments, and for each experiment at least 10 samples were analyzed and  
812 gave consistent results.

813 Immunohistochemistry

814 Immunofluorescence stainings were performed as described previously<sup>86</sup> using an anti-acetylated  
815 tubulin antibody produced in mouse (Sigma aldrich, Reference #T7451-100UL, Lot #036M4856V)  
816 diluted at 1:200. Secondary antibody used was a goat anti-mouse IgG (H+L) secondary antibody coupled  
817 to Alexa Fluor 647 (Invitrogen, Reference #A21235, Lot 2270554) diluted at 1:500. For endoskeleton  
818 stainings, larvae reaching the brachiolaria stage were incubated in seawater supplemented with 5mL per  
819 1L of saturated calcein solution (Sigma-Aldrich), a fluorescent calcium analogue that is incorporated  
820 into the endoskeleton<sup>87</sup>, until the completion of the metamorphosis. The post-metamorphic juveniles  
821 were then fixed, counter-stained with DAPI and cleared following the same procedure than for  
822 immunofluorescence stainings. Imaging was done following the same procedure than for *in situ*  
823 hybridizations.

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891

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904 **Authors contribution**

905 Experiments were designed by L.F., P.P., D.R.R. and C.J.L. Preliminary data were acquired by I.K.,  
906 J.M. and K.R.U. Genome sequencing was done by P.P. and D.R.R. RNA tomography sectioning and  
907 sequencing was done by P.P., M.P., D.R.R. and C.J.L. RNA tomography analyses were performed by  
908 L.F. and D.S.R. Micro-computed tomography preparation was done by L.F. and segmentation of the  
909 scans was done by J.R.T. Immunofluorescence, HCRs and imaging were performed by L.F. Data were  
910 analyzed by L.F, D.S.R. and C.J.L. The manuscript was written by L.F., D.S.R., and C.J.L. with input  
911 from all authors.

912 **Conflict of interests**

913 P.P. and D.R.R. are employees and shareholders of Pacific Biosciences. Other authors declare no  
914 conflict of interests.

915 **Supplementary information**

916 Supplementary Information is available for this paper. Correspondence and requests for materials should  
917 be addressed to L.F. Reprints and permissions information is available at [www.nature.com/reprints](http://www.nature.com/reprints).

918 **Data availability**

919 Specimens used for Micro-CT analyses are registered at the Natural History Museum (London, UK)  
920 under the registration number NHMUK 2023.263. The segmented scans and reconstructed mesh files  
921 are available in Morphosource under the project number [000529415](https://morphosource.org/project/000529415). Genome haplotypes are available  
922 at DDBJ/ENA/GenBank under the accession number [JAPJSQ000000000](https://www.ncbi.nlm.nih.gov/nuccore/JAPJSQ000000000) and [JAPJSR000000000](https://www.ncbi.nlm.nih.gov/nuccore/JAPJSR000000000).  
923 Sequence read archives for RNA sequencing are available at DDBJ/ENA/GenBank under the bioproject  
924 [PRJNA873766](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA873766). RNA tomography dataset is available at Zenodo under DOI [10.5281/zenodo.8327479](https://doi.org/10.5281/zenodo.8327479).

925 **Code availability**

926 Custom code used for RNA tomography analyses is available at Zenodo under DOI  
927 [10.5281/zenodo.8327479](https://doi.org/10.5281/zenodo.8327479).

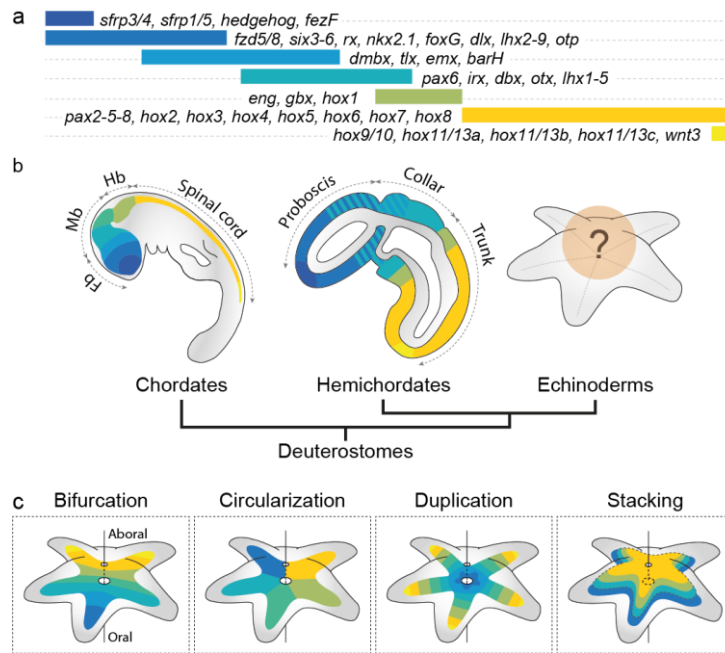
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929 Extended Data Figure 1: *Patiria miniata* anatomy is reflected by RNA tomography

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948 Extended Data Figure 2: Dorso-ventral and appendage patterning in *Patiria miniata*

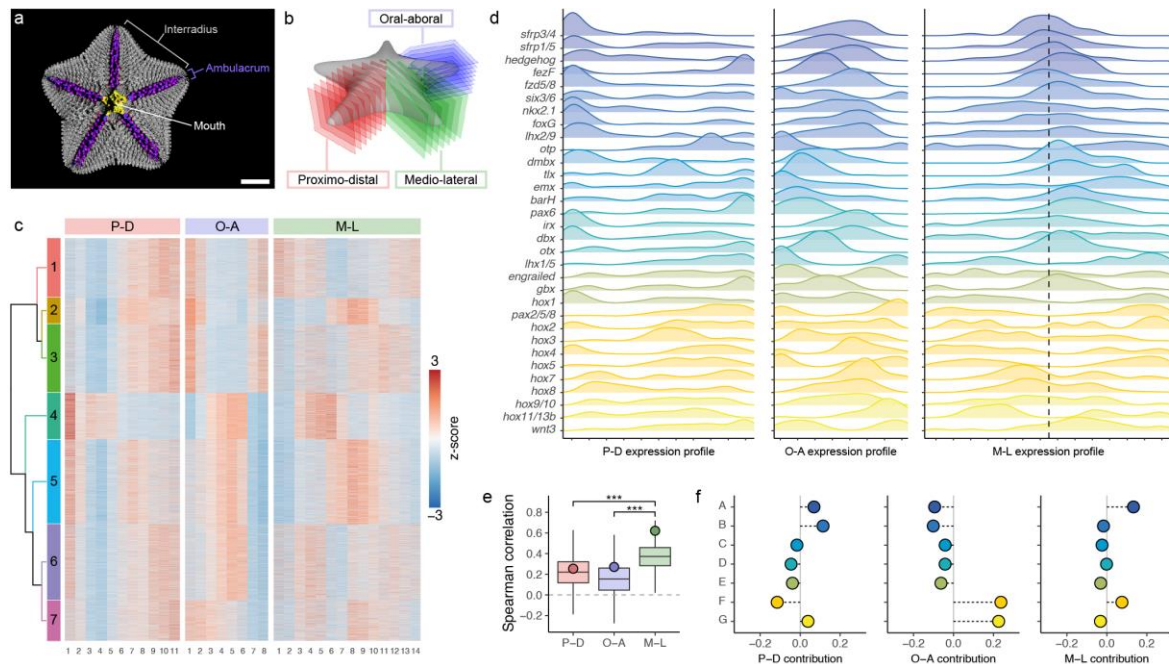
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952 metamorphosis (**d,i,n**), late metamorphosis (**e,j,o**) and post-metamorphic juveniles (**f,g,k,l,p-v, y- $\alpha$** )  
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954 Magnification of a single ambulacrum. **c-v**, Expression patterns of genes involved in DV axis  
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956 bilaterians. **x**, Expression profile of limb proximo-distal patterning genes along the P-D dimension of  
957 the RNA tomography. **y- $\alpha$** , Expression patterns of genes involved in limb proximo-distal patterning.  
958 Scale bars = 100 $\mu$ m.



490

491 Figure 1: Deployment of the antero-posterior patterning system in deuterostomes

492 **a**, Expression map of the conserved transcription factors and signaling ligands involved in ectoderm  
 493 patterning along the AP axis, as observed in the hemichordate *Saccoglossus kowalveskii*. **b**, Previous  
 494 work in chordates and hemichordates has demonstrated extensive regulatory conservation in ectodermal  
 495 AP patterning, establishing the ancestral regulatory characteristics of early deuterostomes. How this  
 496 system is deployed in echinoderms remains unclear. **c**, Four hypotheses have been proposed for the  
 497 deployment of the AP patterning system in echinoderm adult body plan: bifurcation, circularization,  
 498 duplication and stacking.

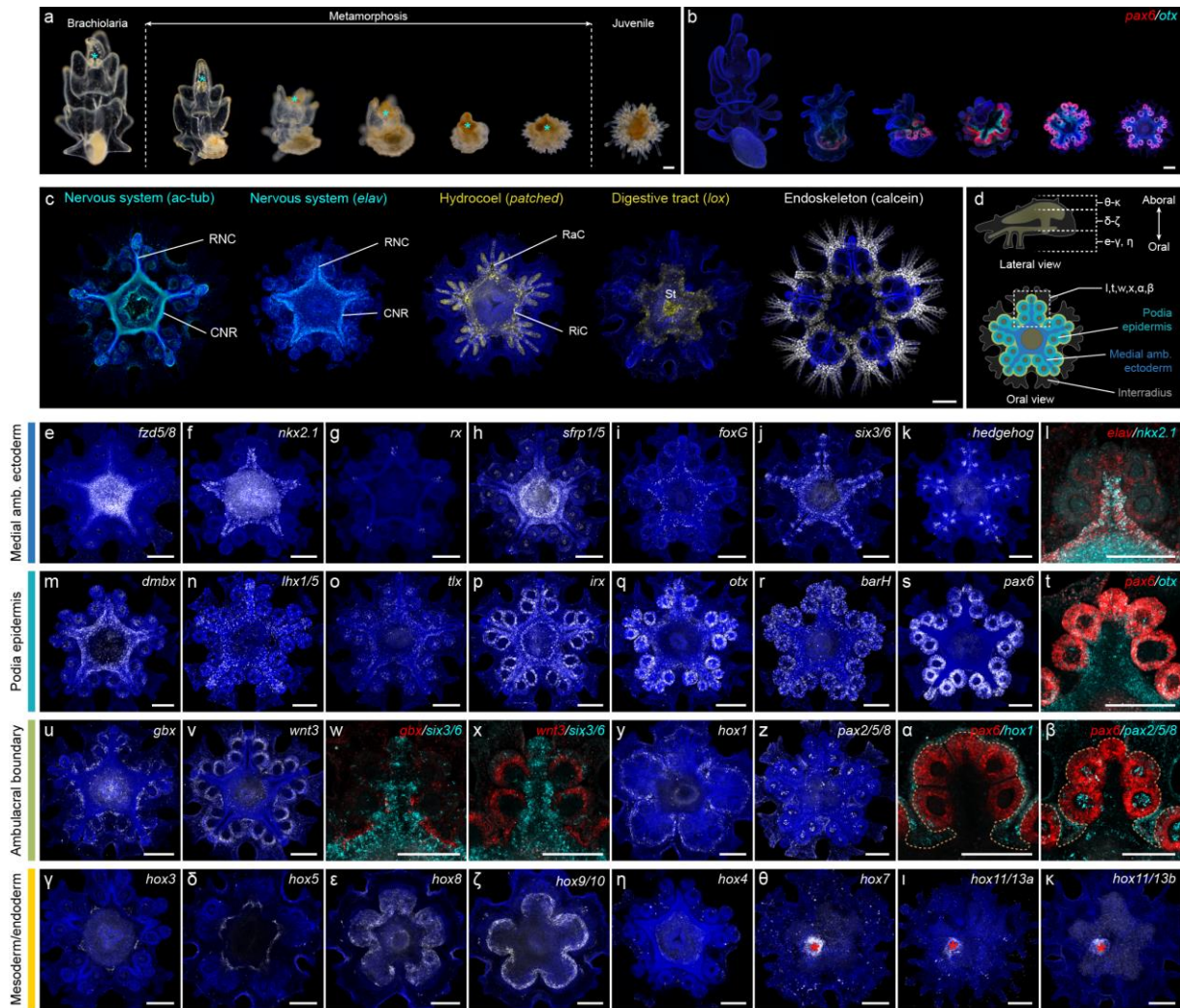


499

500 **Figure 2: RNA tomography reveals the medio-lateral dimension of the arms as the main driver of antero-**  
 501 **posterior patterning system deployment in *Patiria miniata***

502 **a**, Reconstructions of a young juvenile *P. miniata* scanned by micro-CT and segmented to highlight the  
 503 main anatomical features of the animal, including the endoskeleton (grey), the digestive tract (yellow),  
 504 the main body muscles (red), the water vascular system (purple) and the central nervous system (blue).  
 505 The ambulacral areas correspond to the grooves harboring the podia and are separated by interradial  
 506 areas. Scale bar = 1mm. **b**, Experimental design of the RNA tomography with the cutting plans along  
 507 the proximo-distal (P-D), oral-aboral (O-A) and medio-lateral (M-L) dimensions. **c**, Heatmap of gene  
 508 expression z-scores along the P-D, O-A and M-L dimensions of the RNA tomography showing seven  
 509 main trends of spatial expression. **d**, Expression profiles of AP patterning related genes along the P-D,  
 510 O-A and M-L dimensions of the RNA tomography. For the M-L dimension, the dotted line indicates the  
 511 midline. Genes are ranked from the most anterior to the most posterior based on their expression patterns  
 512 in *Saccoglossus kowalevskii*. **e**, Spearman correlations between the ranking of the AP patterning related  
 513 genes and their position along the three dimensions of the RNA tomography. The dots indicate the raw  
 514 correlation values obtained by using the highest z-score as a readout of gene position along each  
 515 dimension, and by ranking the AP patterning related genes following observations in *S. kowalevskii*.  
 516 The boxplots indicate the distribution of correlation values when determining the gene position along  
 517 each dimension probabilistically, and simultaneously shuffling the ranking of the AP patterning related  
 518 genes for each group. The distribution of correlations for the M-L dimension is significantly higher than  
 519 for the P-D and O-A dimensions (Two-sided Wilcoxon-rank test). Centre lines: median; box:  
 520 interquartile range (IQR); whiskers: highest and lowest values at  $\pm 1.5 \times \text{IQR}$ . **f**, Relative contribution  
 521 of each group of AP patterning related genes to the Spearman correlation between the ranking of the AP  
 522 patterning related genes and their position along the three dimensions of the RNA tomography. Anterior  
 523 gene groups drive the P-D and M-L correlations, while the Hox genes are the main drivers of the O-A  
 524 correlation.

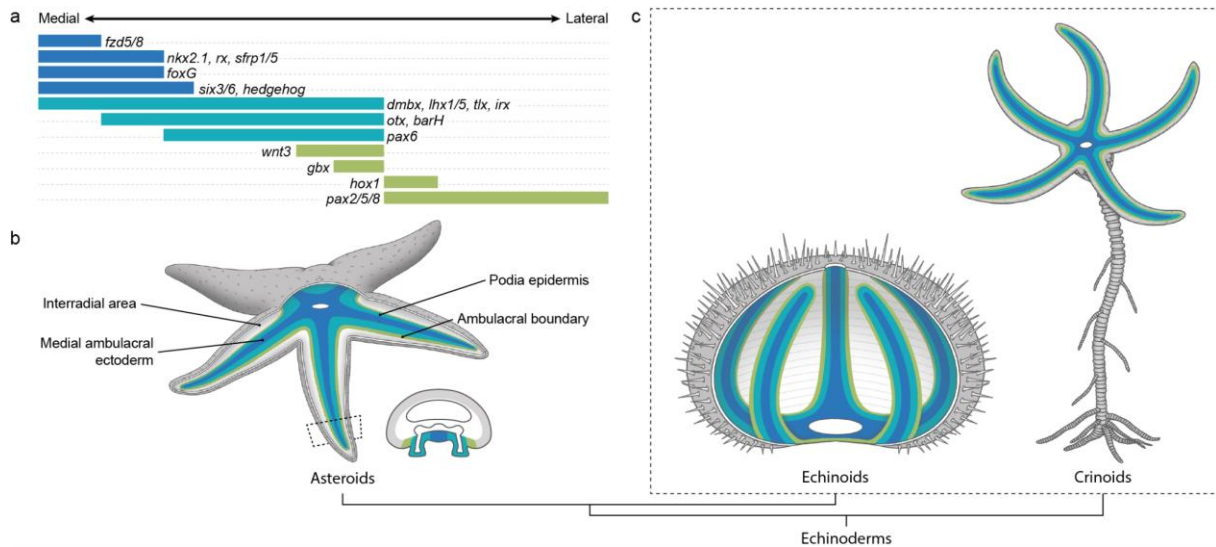




525

526 Figure 3: Gene expression data reveals the deployment of the antero-posterior patterning system in  
 527 *Patiria miniata* ambulacral ectoderm

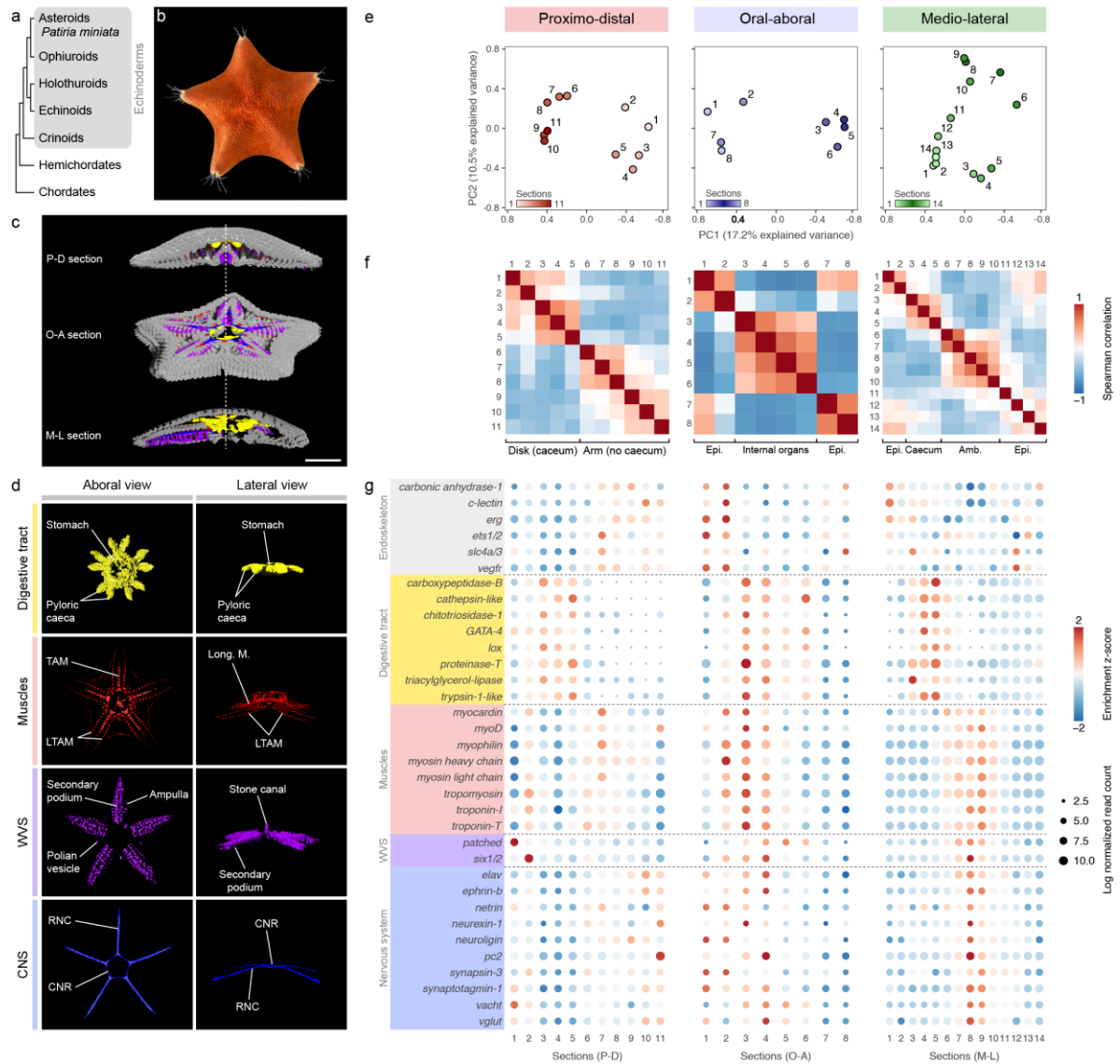
528 **a**, Metamorphosis in *P. miniata*. The anterior part of the brachiolaria larva (cyan asterisk) is resorbed  
 529 into the rudiment. **b**, HCRs showing *pax6* (red) and *otx* (cyan) starting to be expressed during  
 530 metamorphosis in the developing adult pentaradial body plan. Specimens are counterstained with DAPI  
 531 (blue). **c**, Main anatomical features of post-metamorphic juveniles, imaged from the oral side: the  
 532 nervous system (cyan) stained with antibodies against acetylated-tubulin highlighting the nerve tracts  
 533 and the neuronal marker *elav* highlighting the cell bodies, the hydrocoel stained with the marker *patched*,  
 534 the digestive tract stained with the marker *lox* and the endoskeleton stained using calcein. RNC: radial  
 535 nerve cord; CNR: circumoral nerve ring; RaC: radial canal; RiC: ring canal; St: stomach **d**, Schematics  
 536 showing the oral-aboral position of the HCR z-projections shown in **e-κ** (top) and the main anatomical  
 537 regions visible on the post-metamorphic juveniles oral side (bottom). The white square on the bottom  
 538 panel indicates the position of the close-ups in **l,t,w,x,α,β**. **e-κ**, HCRs of *P. miniata* juveniles imaged  
 539 from the oral side (**e-η**) or the aboral side (**θ-κ**). In **e-k,m-s,u,v,y,z,γ-κ**, specimens are counterstained  
 540 with DAPI (blue). **l,t,w,x,α,β**, magnification of a single ambulacrum. **e-l**, Genes primarily expressed in the  
 541 medial ambulacral ectoderm. Colocalization with *elav* indicates expression in the CNR and the  
 542 RNCs. **m-t**, Genes primarily expressed in the podia epidermis. **u-β**, Genes primarily expressed in the  
 543 ambulacral boundary. In **α,β**, orange dotted lines outline the ambulacral ectoderm. **γ-ζ**, Hox genes  
 544 primarily expressed in the coeloms. **η-κ**, Hox genes primarily expressed in the digestive tract. In **θ-κ**,  
 545 red asterisks indicate the position of the developing intestinal tract. Scale bars = 100μm.



546

547 Figure 4: The ambulacral-anterior model of echinoderm body plan evolution

548 **a**, Expression map of the conserved transcription factors and signaling ligands involved in the  
 549 ambulacral ectoderm patterning in *Patiria miniata*, organized from the midline of the ambulacrum (left)  
 550 towards the interradius (right). **b**, Diagram of the ambulacral-anterior model in a generalized asteroid.  
 551 Anterior patterning genes are expressed in the oral ambulacral ectoderm, shown through a cut-away of  
 552 the forward part of the animal and a cross-section through the forward arm. Only genes expressed in the  
 553 ectoderm are shown. **c**, Ambulacral-anterior model applied to echinoderm classes with available gene  
 554 expression data, echinoids (left) and crinoids (right). In echinoids, the ambulacral ectoderm is  
 555 internalized and shown through a cut-away of the forward part of the test, while in crinoids the  
 556 ambulacral ectoderm is embedded in the oral epidermis, similar to asteroids.

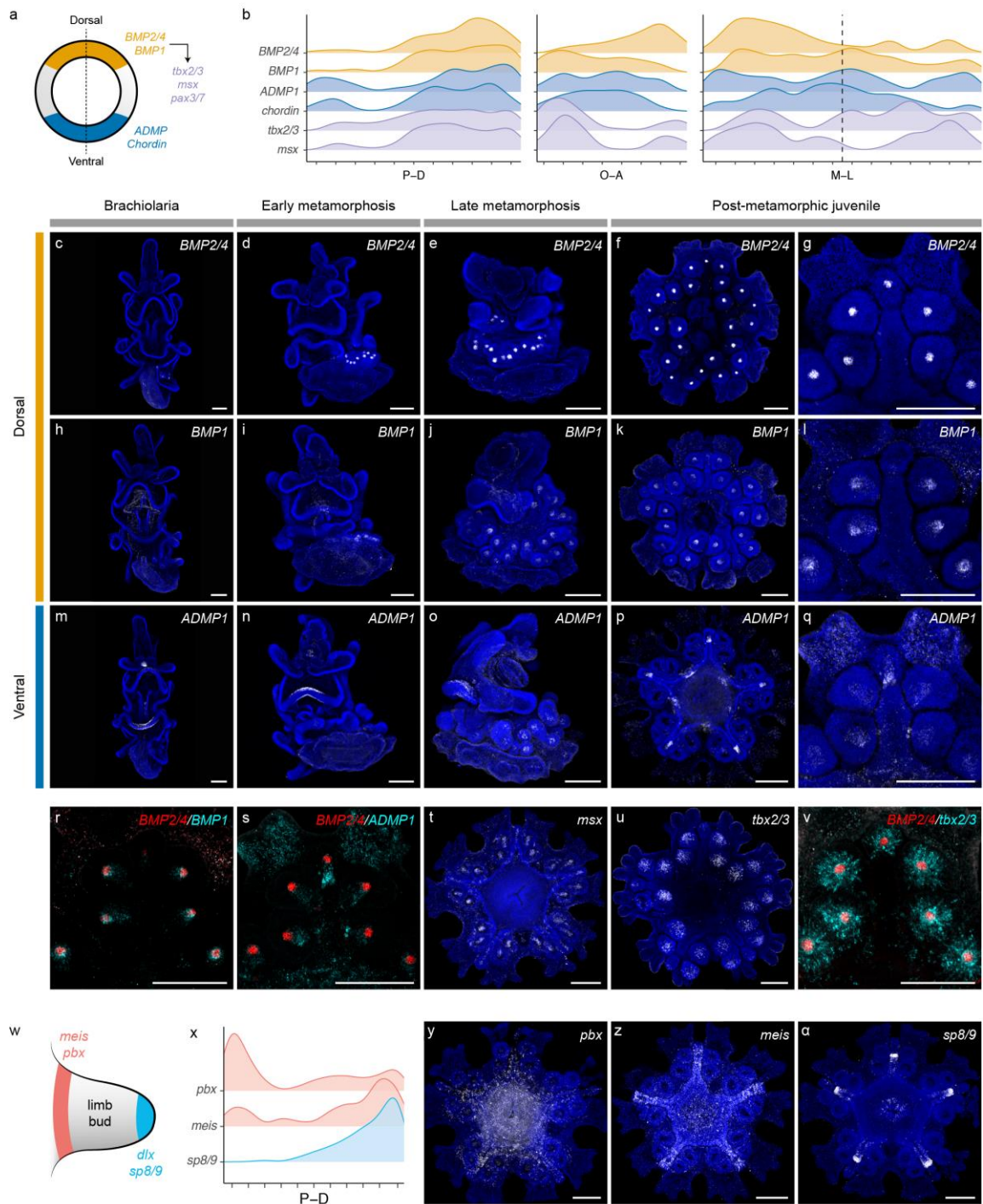


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