RESEARCH ARTICLE



Molecular and cellular architecture of the larval sensory organ in the cnidarian *Nematostella vectensis*

Eleanor Gilbert^{1,2,*}, Callum Teeling^{1,2,*}, Tatiana Lebedeva^{3,4}, Siffreya Pedersen¹, Nathan Chrismas¹, Grigory Genikhovich^{3,‡} and Vengamanaidu Modepalli^{1,‡}

ABSTRACT

Cnidarians are the only non-bilaterian group to evolve ciliated larvae with an apical sensory organ, which is possibly homologous to the apical organs of bilaterian primary larvae. Here, we generated transcriptomes of the apical tissue in the sea anemone Nematostella vectensis and showed that it has a unique neuronal signature. By integrating previously published larval single-cell data with our apical transcriptomes, we discovered that the apical domain comprises a minimum of six distinct cell types. We show that the apical organ is compartmentalised into apical tuft cells (spot) and larval-specific neurons (ring). Finally, we identify ISX-like (NVE14554), a PRD class homeobox gene specifically expressed in apical tuft cells, as an FGF signalling-dependent transcription factor responsible for the formation of the apical tuft domain via repression of the neural ring fate in apical cells. With this study, we contribute a comparison of the molecular anatomy of apical organs, which must be carried out across phyla to determine whether this crucial larval structure evolved once or multiple times.

KEY WORDS: Apical organ, Neuron, Evolution, Cilia, Cnidaria, Nematostella vectensis

INTRODUCTION

During early development, the majority of marine benthic invertebrates progress through a planktonic life phase, consisting of a ciliated larva with an apical organ (Marinković et al., 2020). Several behavioural studies have demonstrated that ciliated larvae use the sensory organ at the apical pole to process environmental cues and modulate their swimming behaviour (Sinigaglia et al., 2015; Iwao et al., 2002; Schmich et al., 1998). The apical pole of the larvae is enriched with flask-shaped cells, usually with an apical tuft of non-motile cilia (Fig. 1) (Marinković et al., 2020; Conzelmann et al., 2011; Verasztó et al., 2017; Goldberg et al., 2011; Garner

*These authors contributed equally to this work

[‡]Authors for correspondence (venmod@mba.ac.uk; grigory.genikhovich@univie.ac.at)

Handling Editor: Cassandra Extavour Received 12 April 2022; Accepted 18 July 2022 et al., 2016; Gruhl, 2009; Page, 2002; Chia and Koss, 1979). Besides the flask-shaped apical cells, the sensory (photosensitive and mechanosensory) and secretory/gland cells are also scattered around the apical pole and are likely associated with the sensory-ciliomotor nervous system. For instance, in bilaterian trochophore larvae, such as in larvae of the mollusc *Ischnochiton hakodadensis* (Fig. 1E) (Voronezhskaya et al., 2002; Nezlin and Voronezhskaya, 2017) and the annelid *Malacoceros fuliginosus*, the apical pole possesses several sensory cells that are positive for serotonin and the neuropeptide FMRFamide (Kumar et al., 2020). Similarly, the apical organ of the marine annelid *Platynereis dumerilii* is also equipped with photosensitive, mechanosensory and peptidergic cell types alongside an apical tuft (Marlow et al., 2014; Williams and Jékely, 2019; Randel et al., 2013; Verasztó et al., 2018).

Among non-bilaterians, a larval sensory organ with integrated neurons is only found in cnidarians (Fig. 1) (Kelava et al., 2015; Marlow et al., 2009; Layden et al., 2016). In anthozoans, like the sea anemone Nematostella, the apical pole displays several flaskshaped apical cells with a ciliated tuft and RPamide-positive sensory cells (Fig. 1D) (Zang and Nakanishi, 2020). In hydrozoans, the apical pole is highly enriched in cells expressing the neuropeptides LWamide and RFamide (Fig. 1C) (Piraino et al., 2011; Gajewski et al., 1996; Leitz and Lay, 1995; Katsukura et al., 2004). However, unlike anthozoans, the hydrozoans lack an apical organ-like sensory structure with a ciliated tuft (Pennati et al., 2013). Strikingly, animals devoid of neurons, such as the sponge Amphimedon queenslandica, also bear a set of sensory and secretory cells (globular/mucous cells) in the anterior region of the larvae (Nielsen, 2013; Brauchle et al., 2018), which are likely involved in modulating larval behaviour (Fig. 1B) (Woollacott, 1993; Richards and Degnan, 2012; Leys and Degnan, 2001; Maldonado et al., 2003). However, the advent of neuronal coordination of motile cilia is deemed to have been strategic in increasing the efficiency of sensory-to-motor transformation (Jékely, 2011).

The morphology of the apical organ in cnidarian larvae is comparable with that seen in bilaterian larvae (Sinigaglia et al., 2015; Marlow et al., 2014; Nielsen, 2015), indicating that the common ancestor of these two groups may have progressed through a free-swimming larval stage with a true larval apical organ and associated neurons (Nielsen, 2013, 2005). Earlier studies in Nematostella have identified a range of apical organ genes by blocking fibroblast growth factor (FGF) signalling during early development (Sinigaglia et al., 2015). Comparative gene expression studies between the cnidarian Nematostella and bilaterian ciliated larvae revealed a strong resemblance in the molecular topography around the apical pole (Marlow et al., 2014; Sinigaglia et al., 2013; Matus et al., 2006; Arendt et al., 2016), suggesting that the apical organ may be an evolutionarily conserved larval structure, although alternative hypotheses also suggest the convergent evolution of primary larvae (Liang et al., 2022 preprint).

¹Marine Biological Association of the UK, The Laboratory, Citadel Hill, Plymouth PL1 2PB, United Kingdom. ²School of Biological and Marine Sciences, University of Plymouth, Plymouth, PL4 8AA, UK. ³Department of Neurosciences and Developmental Biology, Faculty of Life Sciences, University of Vienna, Vienna, 1030, Austria. ⁴Doctoral School of Ecology and Evolution, University of Vienna, Vienna, 1030, Austria.

E.G., 0000-0002-7036-2133; C.T., 0000-0002-7090-1563; T.L., 0000-0003-2182-1492; S.P., 0000-0002-2932-2295; G.G., 0000-0003-4864-7770; V.M., 0000-0003-3099-4969

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution and reproduction in any medium provided that the original work is properly attributed.

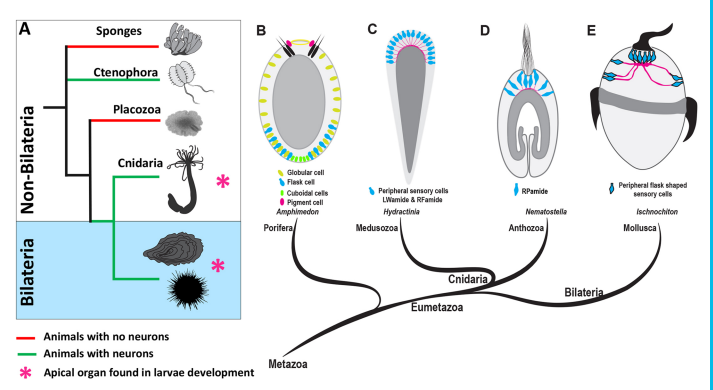


Fig. 1. Origin and evolution of the nervous system and the apical organ. (A) A brief overview of evolutionary relationships in the animal kingdom. Porifera and Placozoa do not have defined neurons. Among non-bilaterians, Cnidaria and Ctenophora have well defined neurons. (B-E) Schematic drawings of surface-contacting flask-shaped cells in the ciliated larvae of different marine phyla. Sponge larvae (B) have ciliated photoreceptor cells capable of light sensing and other peripheral cell types such as flask and cuboidal cells. Among non-bilaterians (C,D), a true larval apical organ with integrated neurons is only found in cnidarians. Schematics were drawn based on the following primary data: (B) Nakanishi et al. (2015), Richards and Degnan (2012), Ueda et al. (2016); (C) Gajewski et al. (1996), Katsukura et al. (2004); (D) Zang and Nakanishi (2020); (E) Voronezhskaya et al. (2002), Nezlin and Voronezhskaya (2017). As ctenophore aboral sensory organs, which are sometimes also termed 'apical organs', are clearly not homologous to the larval apical organs of Cnidaria and Bilateria (Tamm, 2014; Edgar et al., 2022) we did not include them in this Figure.

Cnidarians hold a key phylogenetic position for understanding nervous system evolution, and their larval sensory-ciliomotor nervous system provides a window to look into the primordial neurotransmission system. To advance understanding of this system, we aimed to map the apically enriched cell types and their gene expression profiles in the anthozoan cnidarian Nematostella vectensis. Nematostella is a well-established molecular model species that has been at the centre of fundamental discoveries in the development and evolution of the nervous system in non-bilaterian metazoans, making it a suitable model for the current study (Kelava et al., 2015; Marlow et al., 2009; Layden et al., 2016). Here, we reveal the gene expression profile of the larval apical domain (territory of the apical pole) including the sensory organ by performing transcriptomics on apical tissue that was separated from the rest of the larval body (Fig. 2A). Further, by integrating our tissue-specific (apical/body) transcriptome data with Nematostella larval single-cell RNA-sequencing (RNA-seq) data (Sebé-Pedrós et al., 2018), we identified the larval cell types enriched in the body and apical regions and their gene expression profiles. Finally, we identified ISX-like as an FGF-dependent transcription factor responsible for the definition of the apical tuft territory.

RESULTS AND DISCUSSION

Transcriptome profile of Nematostella apical sensory organ

We performed microdissections on *Nematostella* planula larvae and carefully separated the apical tissue from the rest of the larval body at the mid-planula (~50-60 h post fertilisation) and late planula (~75-85 h post fertilisation) developmental stages (Fig. 2A). We acquired

transcriptome data from both the apical tissue and the rest of the body separately to perform differential gene expression (DGE) analysis. DGE analysis showed statistically significant variations among the apical and body tissue in both the mid-planula and late planula stages (Fig. 2B,C). The late planula stage (Fig. 2C) presented a relatively large number of differentially expressed genes (DEGs) in comparison with the mid-planula stage (Fig. 2B). Furthermore, to characterise global gene expression patterns among the apical and body tissues from the mid-planula and late planula stages, we compared the transcriptomic profiles of all datasets using principal component analysis (PCA) and correlation analysis (Fig. 2D,E). The plots displayed a strong correlation among the replicates and a significant difference between the apical and body datasets from both midplanula and late planula stages. Notably, correlation analysis identified two major clusters and, as illustrated in the PCA plot, the apical datasets from both the planula and late planula formed a single cluster. Likewise, the body datasets fell under a single cluster irrespective of their developmental stage (Fig. 2F). Thus, for the downstream analysis, we pooled both apical datasets from the midplanula and late planula developmental stages; likewise, we pooled the body datasets. A DGE analysis was carried out among the apical and body datasets to identify the significantly DEGs [adjusted Pvalue or P_{adj} (false discovery rate)<0.05]. We identified 2311 DEGs, of which 1185 were enriched in the apical domain and 1126 were enriched in the body (Fig. 2G,H; Table S1).

To validate our transcriptomic data by *in situ* hybridisation (ISH), we selected a set of newly identified apical pole-enriched genes from the DGE data (Fig. 2I). ISH showed that their expression is principally

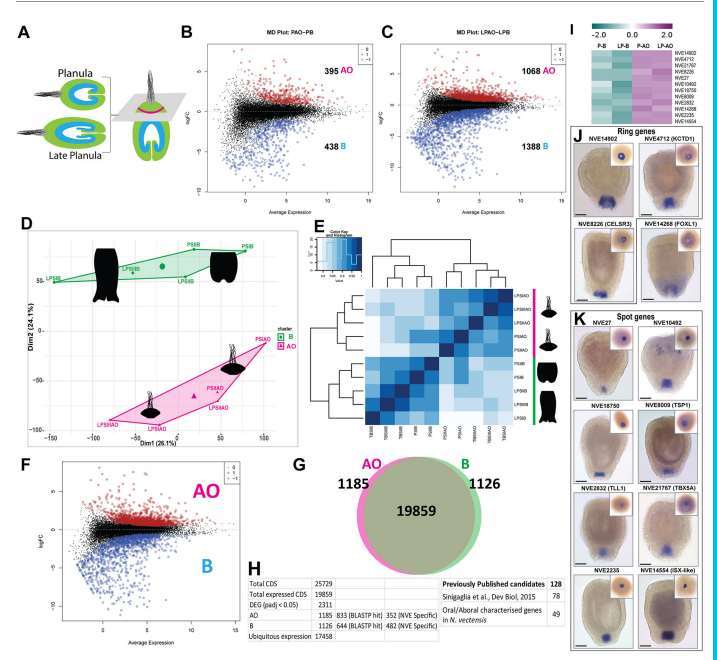


Fig. 2. Global transcriptomic analysis. (A) Schematic representation of the apical region microdissected from the rest of the larval body tissue for transcriptomics. (B-C) Mean-difference (MD) plots represent the log[fold change (FC)] ratio of differential expression between apical and body tissues from (B) planula and (C) late planula stages. The upregulated and downregulated genes are highlighted as red and blue circles, respectively. Apical organ, AO; body, B; late planula apical organ, LPAO; late planula body, LPB; planula apical organ, PAO; planula body, PB. (D) PCA plot displaying a global overview of all datasets. (E) Correlation analysis identified two major clusters, as noted in the PCA plot. (F) MD plots represent the logFC ratio of differential expression between apical and body tissues. Datasets were pooled from planula and late planula development stages. (G) Venn diagram showing the DEGs from apical and body tissues. (H) A table detailing the number of DEGs in the current study, the number of genes with homologs and genes identified in the previous apical organ study by Sinigaglia et al. (2015), and the number of genes previously shown to be associated with the oral/aboral domains in *Nematostella* planula. For additional details, refer to Table S1. (I) A heatmap displaying the gene expression of selected marker genes enriched in apical organ cells. (J,K) ISH of apical organ-enriched genes. The insets show the apical view. Images are representative of approximately 40 *Nematostella* larvae per gene. Scale bars: 50 µm.

localised to the apical organ (Fig. 2J,K). Strikingly, two distinct expression profiles were observed: probes specific to *NVE14902* (*PoxA*), *NVE4712* and *NVE8226* (*Slc*) transcripts were localised around the apical cells (Fig. 2J), whereas probes specific to *NVE27*, *NVE10492*, *NVE8009*, *NVE2832*, *NVE2235* and *NVE14554* (*ISX-like*) transcripts (Fig. 2K) were localised specifically within the apical cells. This pattern was also identified with other *Nematostella* apical organ genes and termed as 'spot' and 'ring' (Sinigaglia et al., 2015). We then overlapped our RNA-seq dataset with the previously

published list of 78 genes with confirmed aboral expression (Sinigaglia et al., 2015), and found that 71 out 78 were present among the 1185 aborally enriched transcripts we identified (Table S1).

A distinct neuronal regulatory network associated with the larval apical region

Neuropeptides in *Nematostella* may potentially be involved in modulating larval swimming behaviour like in bilaterians.

For example, in annelid and echinoderm larvae, the ciliary beating frequency is modulated by neurotransmitters, and neuropeptides actively control the swimming speed by modulating the ciliary beating frequency (Conzelmann et al., 2011; Verasztó et al., 2017; Soliman, 1983, 1984). In ciliated larvae of the annelids Platynereis and Capitella as well as bryozoan Cryptosula, the neuropeptides are expressed in the peptidergic nerves that run along with the ciliary bands (Goldberg et al., 2011; Gruhl, 2009). In cnidarians, neuropeptides are expressed at the larval stage, and it was shown in hydrozoans that neurons expressing LWamide and RFamide are enriched in the aboral (anterior) pole of larvae (Gajewski et al., 1996; Leitz and Lay, 1995; Katsukura et al., 2004). We also noted a significant difference in the spatial distribution of neuropeptide transcripts along the oral-aboral axis in Nematostella (Fig. 3A). Along with the previously identified Nv-RPamide III neuropeptide (Zang and Nakanishi, 2020), we also detected PRGamide as a

neuropeptide exclusively expressed in the apical tissue (Fig. 3A,B), whereas Nv-LWamide, Antho-RFamide neuropeptides type 2 and HIRamide were detected predominantly in the body tissue (Fig. 3A,B).

To understand the differences in neural gene expression between the apical domain and the body of the larva, we explored the tissuespecific transcriptomes using Gene Ontology (GO) analysis to identify genes related to neurotransmission. We performed a protein-protein Basic Local Alignment Search Tool (BLASTP) search in the UniProt database to identify the putative protein homologs. Additionally, we used gene functional annotation data from the published *Nematostella* single-cell transcriptome study (Sebé-Pedrós et al., 2018) (Table S1). Next, we analysed their GO terms using the David 6.7 and PANTHER 15.0 gene ontology tools. We identified several neuronal-associated genes that are differentially expressed between the apical and body regions, such

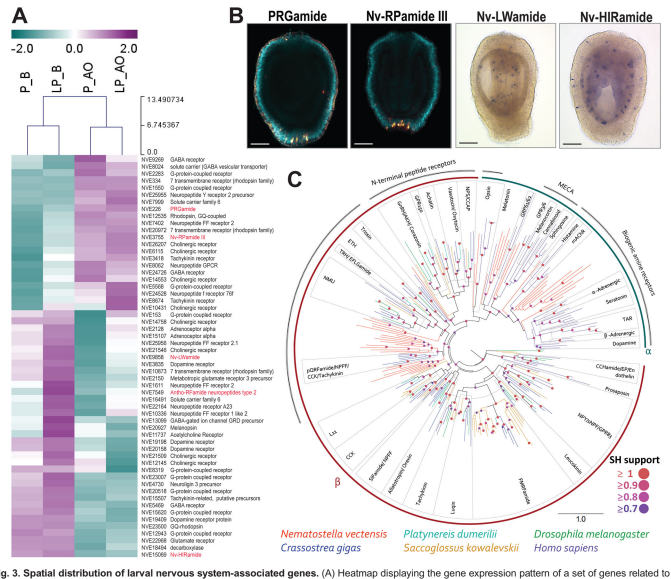


Fig. 3. Spatial distribution of larval nervous system-associated genes. (A) Heatmap displaying the gene expression pattern of a set of genes related to diverse neuronal functions. Neuropeptide genes are highlighted in red. (B) Neuropeptides *Nv-RPamide III* and *PRGamide* were exclusively expressed in the apical tissue, whereas *Nv-LWamide* and *HIRamide* were detected predominantly in the body tissue. Scale bars: 50 µm. Images are representative of approximately 40 Nematostella larvae per gene. (C) Maximum likelihood analyses of the sequences from cluster analysis Fig. S2B. Phylogeny constructed with *C. gigas, P. dumerilii, S. kovalevski, D. melanogaster* and *H. sapiens*. SH support indicates fast approximate likelihood-based measures of branch supports. Adipokinetic hormone, AKH; crustacean cardioacceleratory peptide, CCAP; cholecystokinin, CCK; excitatory peptide, EP; ecdysis triggering hormone, ETH; gonadotropin releasing hormone, GnRH; muscarinic acetylcholine receptor, mAChR; neuromedin-U, NMU; neuropeptide F, NPF; neuropeptide S, NPS; neuropeptide Y, NPY; trace amine receptor, TAR; thyrotropin releasing hormone, TRH.

as neuropeptides, G protein-coupled receptors (GPCRs), ligandgated ion channels and neurotransmitter synthesis genes (Fig. 3A) (Fig. S1). This indicates that the neuromodulation in the apical region of the *Nematostella* larva is characterised by a specific set of genes and may differ from the rest of the body.

In bilaterian models, the neurotransmitters were shown to control the ciliary beating through two broad classes of receptors: ligandgated ion channels and GPCRs (Goldberg et al., 2011; Soliman, 1983, 1984; Kuang et al., 2002; Lacalli et al., 1990). Notably, we observed a large number of GPCRs differentially expressed in the oral versus apical tissues (Fig. S1). GPCRs are the largest family of membrane receptors and mediate most of the cellular responses to hormones, neuropeptides and other neurotransmitters (Hewes and Taghert, 2001; Quiroga Artigas et al., 2020). To understand the relationship of apically enriched GPCRs with known GPCR families in Bilateria, we performed sequence-similaritybased clustering and maximum-likelihood phylogenetic analyses on apically enriched GPCRs, in parallel to BLASTP analysis. We used previously published datasets from Crassostrea gigas, P. dumerilii, Saccoglossus kovalevski and Drosophila melanogaster (Thiel et al., 2021), as well as Homo sapiens GPCRs (Ouiroga Artigas et al., 2020). The Nematostella apically enriched GPCRs clustered with a range of GPCR superfamilies from bilaterians as shown in Fig. S2. Forty-six Nematostella apically enriched GPCRs clustered with a large group of rhodopsin and four clustered with secretin (Fig. S2). Within the rhodopsin superfamily, the GPCRs formed two major groups: rhodopsin α and β . Rhodopsin α GPCRs formed a monophyletic group which included 22 Nematostella GPCRs. The remaining 24 Nematostella GPCRs fell among the rhodopsin β clades (Fig. 3C). The rhodopsin α GPCRs can be further divided into the biogenic amine receptor group and the melanocortin, endothelial differentiation sphingolipid, cannabinoid and adenosine receptor (MECA) group that includes melanocortin and opsins. Among the rhodopsin β GPCRs, 23 Nematostella GPCRs formed three clades positioned among other neuropeptide GPCRs within the rhodopsin β superfamily. In summary, the current analysis allowed us to identify several orthologue groups of GPCRs expressed in the Nematostella apical domain and their distribution across different GPCR receptor families.

Identifying the spatial distribution of *Nematostella* larval cell types by integrating tissue-specific transcriptomes with larval single-cell data

Fine morphological studies in different ciliated larvae (Chia and Koss, 1979; Pennati et al., 2013; Nakanishi et al., 2008; Martin and Chia, 1982) have revealed that, along with long ciliated apical cells, the apical region comprises other cell types such as neurons, gland cells and peripheral ciliated sensory cells. Identification of apical organ-associated cell types and their marker genes is strategic for understanding apical organ function. Single-cell RNA sequencing revealed transcriptome profiles of different cell types in ciliated larvae of Nematostella (Sebé-Pedrós et al., 2018). The larval cells were classified into 38 metacells, and each metacell represented a specific larval cell type with a unique transcriptome profile (Sebé-Pedrós et al., 2018). However, the spatial distribution of these cell types is yet to be addressed. To develop an atlas of apical organassociated cell types and their transcriptomes, we integrated the tissue-specific transcriptome data with the Nematostella whole larval single-cell RNA-seq data (Sebé-Pedrós et al., 2018).

We analysed the expression profiles of apical pole-enriched genes (1185) in each larval metacell using cluster association (Fig. 4A), PCA (Fig. 4B) and hierarchical clustering (HC) (Fig. 4C)

to define the metacells enriched in the apical domain. From the plots, we observed that the apical cell type (Apical_organ1) was clustered distinctly from other cell types and had a high expression of multiple apically enriched genes (Fig. 4A). Out of 38 metacells, the larva-specific neurons, gland/secretory-cell types 1 and 2, and undifferentiated cell types 2 and 4 also displayed specific expression of apically enriched genes (Fig. 4A), and stood out in the apical domain with minimum overlap with any other larval cell clusters (Fig. 4A-C). In parallel, we also analysed the expression profiles of the body-enriched genes (1126) in each larval metacell to identify the cells enriched in body tissue (Fig. 4D-F). In both PCA and HC, genes enriched in the body displayed a very distinctive trend from the apical tissue transcriptome (Fig. 4). We observed that the larval metacells including cnidocytes, gastrodermis and a specific gland/ secretory-cell type 4 displayed high expression of body-enriched genes (Fig. 4D-F) and clustered distinctly from other metacells (Fig. 4E,F), suggesting that these cell types are likely spatially distributed in the body and devoid from the apical region.

Flask-shaped gland/secretory cells enriched in the apical domain

Besides apical cells and larva-specific neuronal cell types, singlecell PCA revealed four gland/secretory-cell types with distinctive profiles among apical and body datasets (Fig. 4A-D). Out of four larval gland-cell types, cell types 1 and 2 were principally enriched in the apical region (Fig. 4A,C). In contrast, gland/secretory-cell type 4 was enriched exclusively in the body (Fig. 4E,F), whereas cell type 3 was expressed ubiquitously (Fig. 4). Recent studies have addressed the development of gland cells in Nematostella, mainly focusing on gland cells that develop and integrate into the pharynx and mesenteries of polyps (Steinmetz et al., 2017; Babonis et al., 2019), and ectodermal gland cells producing toxins (Sachkova et al., 2019; Columbus-Shenkar et al., 2018). However, not much focus has been given to other larval gland/secretory-cell types and their fate during development. To confirm the enrichment of gland/ secretory-cell types in the apical domain, we selected specific marker genes expressed in each of these gland-cell types and visualised their expression pattern by ISH (Fig. 5). Gland cells type 1 and 2 were enriched in the apical region (Fig. 5A-F; Movie 1). Gland-cell type 3 was presented throughout the animal (Fig. 5G,I,J; Movie 2). The marker genes for cell types 1, 2 and 3 were expressed in the outer ectoderm of the planula (Fig. 5A-J). In contrast, marker genes (NVE23810, NVE26086, NVE9234 and NVE10584) selected for gland-cell type 4 were expressed in the pharynx region and progressed into the pharyngeal/mesentery tissue of the primary polyp (Fig. 5K-O). As noted from ISH, gland-cell type 4 had a distinct expression from other peripheral gland cells and was principally restricted to pharyngeal/mesentery tissue (Fig. 5K-O). From the Nematostella larval single-cell data, we noted that the trypsin domain-containing proteins were restricted to gland-cell type 4 (Fig. 5R), suggesting that gland cell 4 is the larval cell type that develops into polyp gland/secretory-cell types in the mesenteries, where the digestive enzymes like trypsin are synthesised and secreted into the gastrovascular cavity for digestive function (Babonis et al., 2019).

To gain further insights into molecular features of the larval gland-cell types, we looked into the *Nematostella* larval single-cell data (Sebé-Pedrós et al., 2018). We noted that each of these gland cells are regulated through a set of transcription factors and some are unique to specific gland-cell types (Fig. 5P,Q). For instance, gland-cell type 1 expresses *retinal homeobox* (Rx) (v1g184843) and FoxQ2d (v1g96685), and gland-cell type 2 expresses *Emx1*

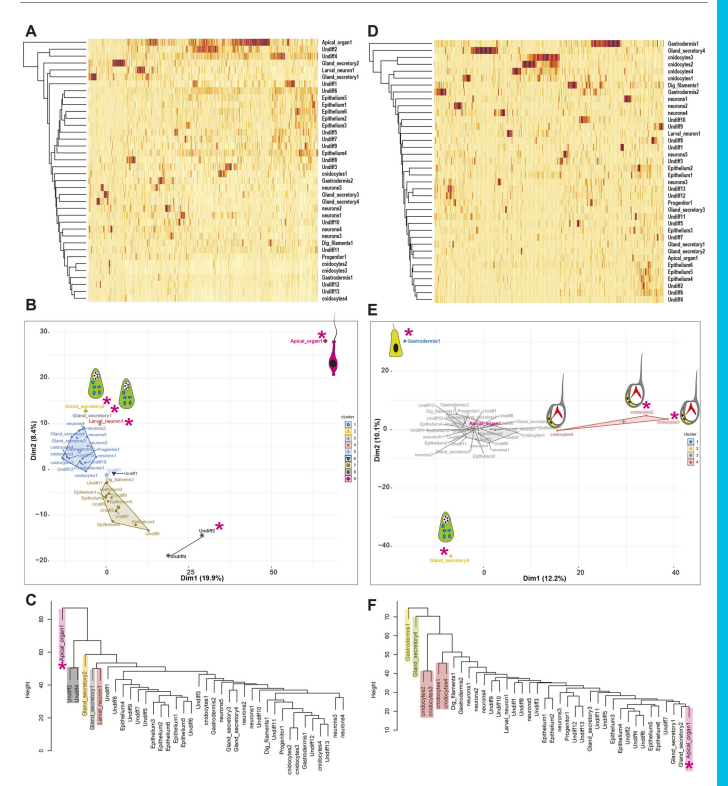


Fig. 4. Spatial distribution of *Nematostella* larval cell types. (A,D) Expression of apically (A) and body- (D) enriched genes (rows) across 38 metacells sorted by cluster association. (B,E) Single-cell PCA plot for apically (B) and body- (E) enriched genes. Pink asterisks indicate apical organ cell type. (C,F) Hierarchical clustering for apically (C) and body- (F) enriched genes. The dendrogram produced similar results as single-cell PCA.

(v1g8907). However, some of the transcription factors are expressed in several gland-cell types. For example, *PaxC* (v1g168908) is expressed both in gland-cell types 3 and 4. Similarly, *RFX4* (v1g122918) is expressed in both gland-cell types 1 and 2 (Fig. 5Q). This suggests that a different combination of transcription factors modulates the trajectory of each of these gland cells.

Nematostella apical organ is composed of apical tuft cells crowned with larval neurons

Based on the *Nematostella* single-cell transcriptome study (Sebé-Pedrós et al., 2018), two metacells were classified as larval specific with low similarity to any adult cell cluster. Based on the previously identified marker genes Fgfla (Sinigaglia et al., 2015) and Nk3, one

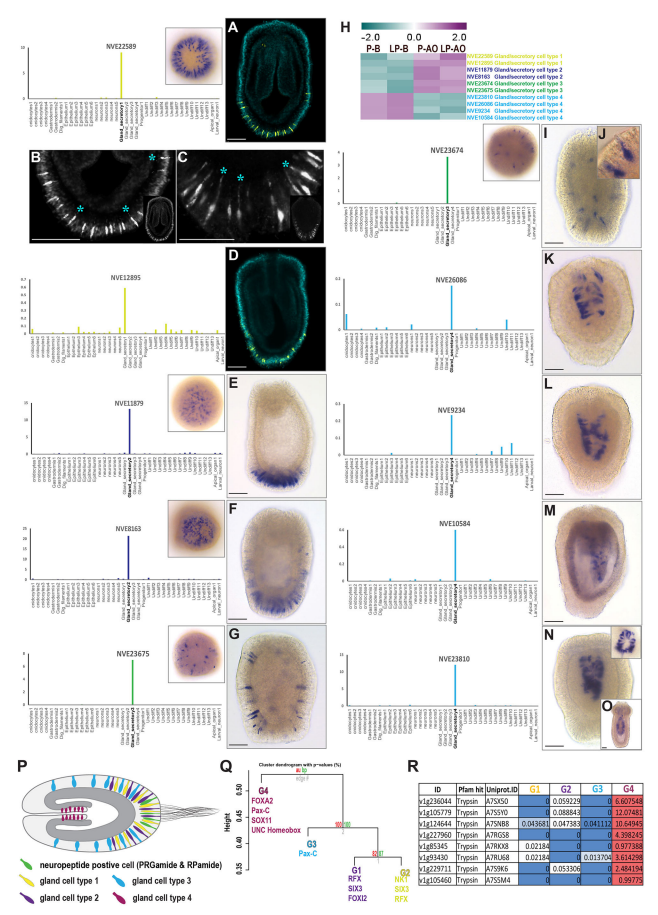


Fig. 5. See next page for legend.

Fig. 5. Spatial distribution of larval gland/secretory-cell types

visualised by whole-mount ISH. (A,D-G,I-O) ISH (right) of larval gland/ secretory-cell type marker genes. Each bar plot (left) displays the expression profile of selected marker genes in the different larval cell populations. The insets (top right) show the apical view. (A-D) ISH of larval gland/secretorycell type 1. (B,C) Monopolar sensory cells with their projection from the cell body extended towards mesoglea (asterisks); the insets show the whole animal. (E,F) ISH of larval gland/secretory-cell type 2. (G,I,J) ISH of the larval gland/secretory-cell type 3; panel J shows the ciliated gland-cell type 3 at higher magnification. (H) Heatmap displaying the gene expression pattern of selected marker genes specific for each larval gland/secretory-cell types. (K-O) ISH of the larval gland/secretory-cell type 4; panel O demonstrates the spatial expression of gland-cell type 4 concentrated in the mesentery tissue. (P) Schematic representation of the spatial distribution of larval gland/ secretory-cell types. (Q) Dendrogram displaying the correlation among gland cells and a list of a different combination of transcription factors expressed in each gland-cell type. (R) Gene expression of trypsin domain-containing proteins in gland cells from larval single-cell data. Images are representative of approximately 40 Nematostella larvae per gene. Scale bars: 50 µm.

of these metacells was recognised as an apical cell type. The second metacell showed a peculiar expression of genes with putative neuronal functions such as *cyclic nucleotide-gated*, *TrpA*, *polycystic kidney disease* and *shaker* ion channels; therefore, it was classified as an uncharacterised larval-specific neuronal cell type (Sebé-Pedrós et al., 2018).

From single-cell PCA and HC analyses, we observed that along with the apical cell type, the larva-specific neurons stood out in the apical domain (Fig. 4A-C). In our initial ISH analysis of apically enriched genes, we observed two distinct expression profiles: a set of genes expressed around the apical cells, and others expressed specifically in apical cells (Fig. 2J). The distinct expression patterns of apical organ genes suggest that apical sensory structure is composed primarily of two distinct cell types. Based on the Nematostella single-cell transcriptome (Sebé-Pedrós et al., 2018), the marker genes identified as spot specific were principally restricted to the apical cell type (Fig. 6I). However, the marker genes expressed as a ring were enriched in larval neurons (Fig. 6I). To explore this further, we carried out double fluorescence in situ hybridisation (FISH) on these marker genes (Fig. 6). The NVE8226 and NVE14902 markers, besides being expressed as a ring around the apical cells, showed expression across the whole apical region (Fig. 6A,E; Movies 3 and 4). These cells have a peculiar sensory neuronal morphology (Marlow et al., 2009) and are localised in the ectoderm with the tip pointing towards the periphery (Fig. 6B,F). Expression of these genes gradually diminishes as the larvae progress through metamorphosis (Fig. 6C,D,G,H), likely reflecting their larval-specific function.

Based on the double FISH, the apical organ gene ISX-like was found to be restricted to the apical pit and crowned with larva-specific neurons (detected by NVE8226 and NVE14902 markers) (Fig. 6J-M; Movies 5 and 6). The cells carrying the apical tuft cilia, which were visualised by immunostaining with an anti-acetylated tubulin antibody, were concentrated in the apical pit, where the spot genes like ISX-like are expressed (Fig. 6N). Our analysis allowed us to localise the previously uncharacterised larval-specific neuronal cell type to the apical organ ring. This localisation suggests that the apical cells with their ciliated tufts act as a sensory structure, and the larvaspecific neurons receive information by crowning around them and probably signalling downstream to the rest of the body (Fig. 6O). The apical organ cells lack any recognisable neuronal effector genes such as GPCRs, synaptic scaffold proteins, neurotransmitter-related enzymes and neuropeptides, suggesting a non-neuronal identity (Sebé-Pedrós et al., 2018). However, the larval-specific neuronal cell type is enriched with neuronal functional genes. To conclude, the apical tuft cells and larval-specific neurons are the principal cell types in the apical sensory organ (Fig. 6O).

ISX-like is responsible for the formation of the apical tuft cell fate

ISX-like encodes a paired-class homeodomain transcription factor with a typical spot expression in the apical organ (Figs 3 and 4). It was one of the prominent validated DGE targets in our screen and was previously described as an aborally expressed gene negatively regulated by β-catenin signalling (Lebedeva et al., 2021). Earlier studies (Mazza et al., 2010; Brauchle et al., 2018) classified ISX-like as a PRD class homeobox gene. Reciprocal BLAST search and phylogenetic analysis (Fig. S3) showed that *ISX-like* is related to the bilaterian PRD class intestine-specific homeobox (ISX). Nematostella ISX-like starts to be expressed in a broad aboral domain around 14 h post fertilisation (Fig. S4A), i.e. 2 h after the onset of the expression of the key aboral regulator Six3/6 (Sinigaglia et al., 2015; Lebedeva et al., 2021), and later becomes restricted to a spot domain in the apical tuft cells (Fig. 6J-M). As our FISH analyses showed that the ISX-like-expressing apical tuft cells and the ring of circumapical neurons formed two perfectly complementary domains, we asked whether ISX-like was involved in the regulation of the development of the spot and the ring.

RNAi-mediated, as well as antisense morpholino-mediated knockdown of ISX-like resulted in the complete loss of the apical tuft (Fig. 7A,B; Fig. S5). Analysis of the morphants showed that the fraction of larvae successfully undergoing metamorphosis was moderately reduced in comparison with controls (68% upon ISXlike knockdown versus 82% in control; unpaired, two-tailed t-test, t=3.71391, P=0.020584) (Fig. S5). ISH analyses showed that ISX*like* knockdown did not simply prevent the formation of the apical tuft cilia, but rather led to a loss of the apical tuft cell identity, i.e. the expression of the larva-specific ring neuron markers NVE8226 and NVE14902, as well as the expression of the upstream aboral regulator Six3/6, expanded into the spot domain (Fig. 7C; Fig. S5). In contrast, the spot expression of the aborally enriched neuropeptide genes PRGamide (NVE226) and RPamide III (NVE3775) was not affected at all (Fig. 7C; Fig. S5). Thus, we conclude that the formation of the apical tuft cells takes place due to ISX-like-dependent repression of the neural ring fate in these cells.

As the formation of the apical tuft was previously shown to be FGFa1-dependent (Rentzsch et al., 2008), we then tested whether ISX-like was upstream or downstream of FGF signalling. Although FGFa1 expression appeared to be only very weakly affected by ISXlike knockdown (Fig. 7C; Fig. S5), the expression of ISX-like was abolished upon incubation of the embryos in the FGF receptor inhibitor SU5402 and MEK inhibitor U0126, suggesting that ISXlike expression was positively controlled by FGF signalling (Fig. 8A). Surprisingly, the expression of the neural ring markers NVE8226 and NVE14902 reacted differently to SU5402 and U0126 treatment. Although NVE8226 expression was abolished upon treatment with both inhibitors, the ring expression of NVE14902 was abolished upon treatment with the FGF receptor inhibitor SU5402 and in approximately 61% (*n*=103) of the embryos treated with the MEK inhibitor U0126 (Fig. 8A; Fig. S6). However, NVE14902 showed spot expression in the remaining 39% of the embryos treated with U0126 (Fig. S6), which suggests a more complex, yet unclear type of regulation, possibly by non-FGFmediated MAPK signalling. Expression of the aborally expressed neuropeptide genes PRGamide and RPamide III was not abolished by treatment with either of the inhibitors, although the expression

Z

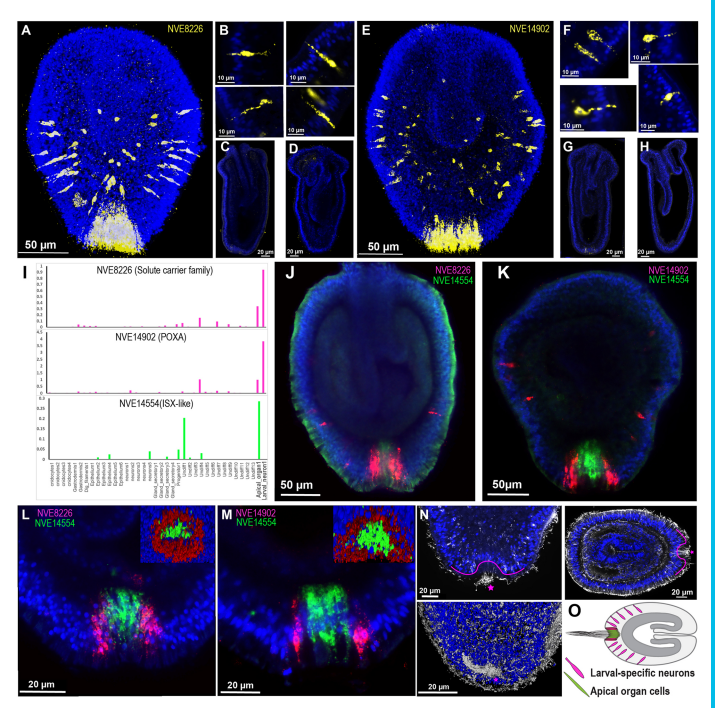


Fig. 6. Expression analysis of apical spot and neuronal ring markers. (A-D) FISH analysis of *NVE8226* (yellow). (E-H) FISH analysis of *NVE14902* (yellow). A,B,E,F show the planula, C,G show metamorphosis and D,H show the early primary polyp. (I) Bar plot displaying the expression profiles of selected marker genes. (J-M) Double FISH demonstrates mutual localisation of marker genes for the spot (*ISX-like*; green) and the ring (*NVE8226* and *NVE14902*; red). DAPI-stained nuclei are shown in blue. The insets (L,M) on the top right display the three-dimensional image from the apical view. (N) Immunostaining with the anti-acetylated tubulin antibody (white) counterstained with DAPI (blue) for nuclei; the pink line demonstrates the apical tuft cells concentrated in the apical pit. Pink stars indicate the apical tuft. J-N show the planula. Images are representative of approximately 40 *Nematostella* larvae per gene. (O) Summary diagram showing the spatial distribution of the apical organ/tuft and larval-specific neuronal cell types.

appeared weaker in U0126-treated embryos. Taken together, we conclude that *ISX-like* is an FGF signalling-dependent transcription factor responsible for the formation of the apical tuft domain acting via repression of the neural ring fate in apical cells.

Conclusions and outlook

Despite the obvious similarity to bilaterian apical organs, the *Nematostella* apical organ remains a mysterious structure.

Combined with the previously published single-cell data (Sebé-Pedrós et al., 2018), our study provides an entry point to explore its function. Contrary to a previous assumption (Marlow et al., 2014), the *Nematostella* planula single-cell transcriptome suggests that apical tuft cells lack recognisable neuronal effector transcripts, suggesting a non-neuronal identity. Conversely, the larval-specific neuronal cell cluster is enriched with neuronal transcription factors such as foxQ2b and soxB(2a), and ion channels implicated in

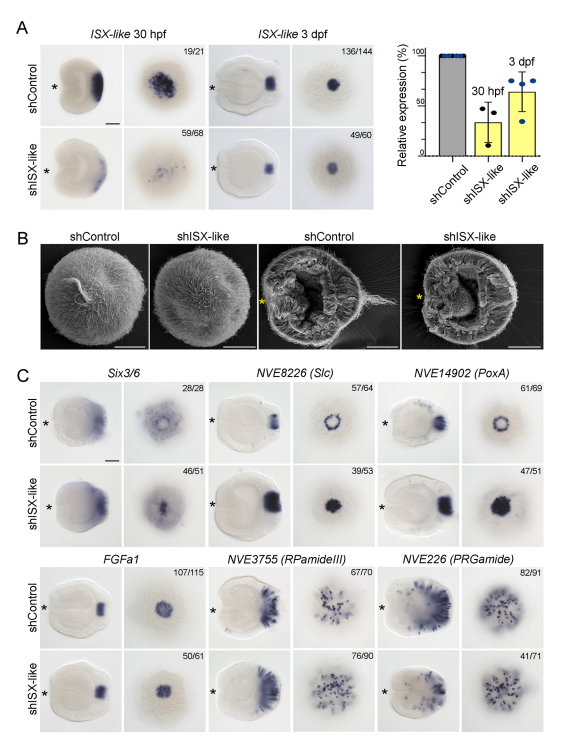


Fig. 7. *ISX-like* **RNAi** efficiency and loss-of-function analysis of *ISX-like*. (A) At 30 hpf, *ISX-like* RNAi efficiently reduces the amount of *ISX-like* mRNA as shown by ISH (left) and qPCR (right). By 3 dpf, *ISX-like* expression largely recovers. Data show the mean±s.d. (B) *ISX-like* RNAi results in loss of the apical tuft. SEM of 72 hpf embryos viewed aborally (two images on the left) or laterally (two images on the right; the embryos were cracked open to visualise the internal structures). (C) Effect of *ISX-like* RNAi on marker gene expression in the 3 dpf planula. Ring genes *NVE8226* and *NVE14902* became spot genes. *PRGamide* and *RPamide III* expression was not affected. *Six3/6* expression extended into the apical organ. *FGFa1* expression was not affected. Lateral and aboral views are shown. For lateral views, asterisks denote the oral end. In A and C, the numbers in the top right corner show the fraction of the embryo demonstrating this phenotype. B,C show the planula. Scale bars: 50 µm.

putative neuronal functions (Sebé-Pedrós et al., 2018). In this study, we revealed that apical tuft cells are crowned with larval-specific neurons, suggesting that these cell types together form the apical organ of the *Nematostella* planula. For the apical organ (i.e. the apical tuft cells plus the neural ring cells) to be considered a

true sensory organ, future research will have to demonstrate the transmission of information from the apical tuft cells to the larval nervous system. The formation of the apical organ is FGF-dependent: loss of the apically expressed *FGFa1* or its putative receptor *FGFRa*, as well as pharmacological inhibition of FGF

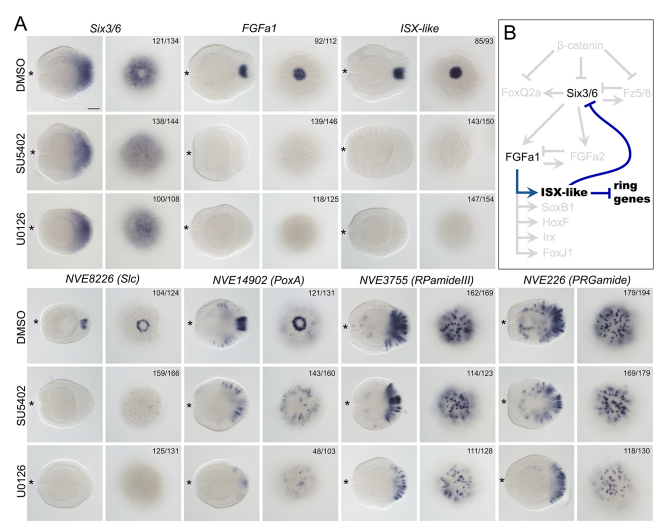


Fig. 8. *ISX-like* expression is controlled by FGF signalling. (A) The expression of the spot and ring genes is abolished upon FGF receptor or MEK inhibition, but not the expression of the aborally enriched *PRGamide* and *RPamide III. Six3/6* expression extends into the apical organ domain. Lateral and aboral views of 3 dpf planulae are shown. For lateral views, asterisks denote the oral end. The numbers in the top right corner show the fraction of the embryos displaying the phenotype shown on the image out of the total number of embryos treated and stained as indicated on the figure. Scale bar: 50 µm. (B) Genetic interactions regulating the apical domain patterning based on findings by Sinigaglia et al. (2013) and Leclère et al. (2016) (grey) and this paper (black and blue).

signalling in *Nematostella* blocks development of the apical tuft (Rentzsch et al., 2008) and the neural ring (Fig. 8). Moreover, the metamorphosis of the *Nematostella* planula into polyps is also suppressed by *FGFa1* and *FGFRa* knockdown, as well as by pharmacological inhibition of FGF signalling (Rentzsch et al., 2008). Although it is tempting to think that the loss of the apical tuft may be the cause of the failed metamorphosis, our data do not support this hypothesis: knockdown of the FGF-dependent transcription factor ISX-like results in a complete loss of the apical tuft domain in the planula, but the effect on metamorphosis is very mild (Figs. 4 and 7). In the future, it will be important to functionally test the potential role of the neural ring cells of the apical organ, as well as of the apical domain, in driving metamorphosis.

Another more profound question is the evolutionary origin of apical organs and whether the apical organs of ciliated larvae across different phyla are homologous or evolved convergently. The widespread occurrence of ciliated larvae with apical organs has prompted radically different views on their evolutionary significance. Some authors consider ciliated larvae to represent the ancestral metazoan morphology (Jagersten, 1972; Nielsen, 2012; Peterson et al., 1997; Davidson et al., 1995), whereas others strongly suggest that larval stage was intercalated independently multiple times in phylogenetically distinct lineages (Liang et al., 2022 preprint; Raff, 2008; Sly et al., 2003; Wolpert, 1999). Naturally, the homology of apical organs becomes questionable if primary larvae evolved convergently in different phyla. However, although the 'adult first versus larva first' question remains debated, a highly conserved set of genes patterning the apical/anterior ectoderm in directly and indirectly developing Bilateria and in the apical/aboral ectoderm in Cnidaria shows that these regions are very likely homologous (Marlow et al., 2014; Sinigaglia et al., 2013; Lebedeva et al., 2021; Range et al., 2013; Kitzmann et al., 2017; Santagata et al., 2012). This, in turn, makes it important to consider the possible 'deep homology' of the apical organs, even if primary larvae in different phyla evolved convergently, and to analyse the relationship between the cell types building the apical organs across phyla by using transcriptomic methods. Our study using an anthozoan cnidarian model makes a step in this direction and provides a valuable set of data from a member of the bilaterian sister group.

MATERIALS AND METHODS

Nematostella culture

Nematostella polyps were grown in 16‰ artificial seawater at 18°C in the dark and fed with freshly hatched *Artemia* nauplii. The induction of spawning was performed as previously described (Genikhovich and Technau, 2009a). After fertilisation, the gelatinous substance around the eggs was removed using 4% L-cysteine (Sigma-Aldrich) (Genikhovich and Technau, 2009a).

Microdissection of Nematostella apical organs

We performed microdissection on *Nematostella* larvae to separate the apical organ from the rest of the larval body. Two developmental stages including planula and late planula were used. Apical tissue containing the apical organ (see Fig. 2A) was isolated using 34-gauge needles under a stereomicroscope with $10 \times$ magnification. Motile larvae were placed into a fresh plastic Petri dish filled with *Nematostella* medium. The larvae tend to adhere briefly to the bottom of a fresh plastic Petri dish, allowing enough time to separate the apical tissue by cutting. Each sample was pooled from a minimum of 50 individual larvae and also included samples from a minimum of three different batches. The samples were carefully collected using glass Pasteur pipettes, excess medium was removed, and the samples were snap-frozen in liquid nitrogen and stored at -80° C until further processing.

RNA sequencing and differential gene expression

For RNA isolation, the tissue samples collected from multiple batches were combined to acquire an adequate amount of RNA for sequencing. Total RNA was isolated using the TRI Reagent (Sigma-Aldrich) according to the manufacturer's protocol. RNA quality was assessed using an Agilent RNA 6000 Nano Kit on an Agilent 2100 Bioanalyzer, and samples with RNA integrity number \geq 8.0 were used for sequencing. The SENSE mRNA-Seq Library Prep Kit (Lexogen) was used for library preparation. Before sequencing, the libraries were pre-assessed by Agilent High Sensitivity DNA Kit (Agilent) and quantified using Qubit 1× dsDNA HS Assay Kit (Invitrogen). The sequencing was outsourced (GENEWIZ Illumina NovaSeq/HiSeq 2×150 bp sequencing), generating 20 million paired end reads per replicate. Raw data were deposited at the Gene Expression Omnibus with the accession ID GSE159166. After de-multiplexing and filtering high-quality sequencing reads, the adapter contamination was removed by using Trimmomatic v0.36 (Bolger et al., 2014). The quality of the reads was verified using FastQC (http://www.bioinformatics.babraham. ac.uk/projects/fastqc). Processed reads from each sample were mapped to the *Nematostella* genome (indexed bowtie2; Langmead and Salzberg, 2012) by using STAR (Spliced Transcripts Alignment to a Reference) (Steinmetz et al., 2017). The number of reads mapping to each Nematostella (https://figshare.com/articles/Nematostella_vectensis_ gene model transcriptome_and_gene_models_v2_0/807696) was extracted using HTSeq-count v0.6 (Babonis et al., 2019). Differential expression analyses were performed using limma (Galaxy version 2.11.40.6) (Love et al., 2014) and DESeq2 (Galaxy version 2.11.40.6) (Love et al., 2014). PCA, HC and heat maps were generated using the R package in R-studio (version 1.2.5019). For functional annotation, we used BLASTP (Altschul et al., 1997) with the default curated gathering threshold to predict the protein homologs against the UniProt database. Additionally, we used gene functional annotation data from a published Nematostella single-cell transcriptome study (Sebé-Pedrós et al., 2018). GO term enrichment was performed using gene annotation tools including PANTHER Classification System (Mi et al., 2012) and DAVID (Huang da et al., 2009).

ISH and double FISH

ISH was performed according to published protocols (Wolenski et al., 2013; Genikhovich and Technau, 2009b). In brief, fixed animals were transferred into sieves and rehydrated in 1 ml 60% methanol/40% PBST [1× PBS with 0.05% (vol/vol) Tween-20] and then washed in 30% methanol/70% PBST. Samples were digested in proteinase K (80 µg/ml, Ambion) for 5 min, then blocked in glycine (4 mg/ml). Larvae were then transferred into 4% formaldehyde at room temperature (RT) for 1 h. Hybridisation was carried out with DIG-labelled probes for 48 h at 60°C. After incubation, samples were washed through serial dilutions of 25%, 50%, 75% and 100% with 2× SSCT buffer (20× concentrate, contains 0.3 M sodium citrate in 3 M NaCl, pH 7.0) at hybridisation temperature. The colour development was carried out in a 1:50 dilution of nitro blue tetrazolium/5-bromo-4-chloro-3-indolyphosphate (NBT/BCIP, Sigma-Aldrich) at RT. Stained animals were visualised with a Leica DM1000 microscope equipped with a MC190 HD microscope camera (Leica, Germany). For each gene, at least 30 specimens were tested. For double FISH after the SSCT washes, samples were blocked in 0.5% blocking reagent (FP1020, PerkinElmer) at RT for 1 h. Samples were then incubated overnight with anti-digoxigenin (1:100; Roche). After TNT (0.1 M Tris-HCl pH 7.5,0.15 M NaCl, 0.5% Triton X-100) washes, samples were incubated in Cy3 (NEL744001KT, TSA Plus Kit, PerkinElmer). To stop the peroxidase activity, the samples were washed in 0.1 M glycine, pH 2.0, and then incubated overnight with anti-fluorescein (1:250; Roche). Samples were imaged on Leica TCS SP8 DLS confocal microscope.

Whole-mount immunofluorescence and scanning electron microscopy

After 1 h of fixation in 3.7% formaldehyde in *Nematostella* medium (16‰ artificial seawater), the samples were washed five times with PBST for 10 min. The samples were blocked in 5% bovine serum albumin (BSA) in PBST for 1 h at RT. Primary antibody (1:500, mouse anti- α -tubulin, T9026, Sigma-Aldrich) incubation was performed in a blocking solution (1% BSA in PBST) for 24-36 h at 4°C. The samples were washed with PBST five times for 5 min, after which samples were incubated with secondary antibodies (1:250, goat anti-mouse IgG Alexa Fluor 594, A-11032, Thermo Fisher Scientific) diluted in blocking solution overnight at 4°C. Then, the samples were washed with PBST five times for 10 min. Imaging was performed on Leica TCS SP8 DLS and Leica DMi8 confocal microscopes. Sample preparation for scanning electron microscopy (SEM) was performed as in Kraus et al. (2016); SEM imaging was performed using the JEOL IT 300 scanning electron microscope.

GPCR clustering and phylogeny

We used previously published GPCR datasets of *Crassostrea virginica*, *P. dumerilii*, *S. kovalevski* and *D. melanogaster* (Thiel et al., 2021), as well as *H. sapiens* GPCRs (Quiroga Artigas et al., 2020). Initially, to identify the *Nematostella* GPCR candidates, grouping with bilaterians was done by clustering analysis using CLANS (Frickey and Lupas, 2004) with the BLOSUM62 scoring matrix and a *P*-value cut-off of 10^{-25} . Sequences from the main cluster were used in phylogenetic analysis. Sequences were aligned with MUSCLE (Edgar, 2004) using default settings and trimmed with TrimA1 using the automated mode (Capella-Gutierrez et al., 2009). Maximum likelihood phylogenetic analyses were completed using PhyML 3.0 online (www.atgc-montpellier.fr/phyml/) (Guindon et al., 2010). The model LG +G +F was automatically selected by the Smart Model Selection with (SH-aLRT).

Gene knockdown and inhibitor treatment

Preparation of the short hairpin RNA against the target sequence 5'-GC-GCTAGTCAACATACTGA-3' and RNAi of ISX-like was performed by electroporation as described by Karabulut et al. (2019). Microinjection of the translation-blocking morpholino oligonucleotide 5'-AATTCTCTGAT-TTTTCCATCGTGG-3' was used as the second independent approach to knock down ISX-like. As controls, shRNA against mOrange (Lebedeva et al., 2021) and a previously described control morpholino (Kraus et al., 2016) were used. RNAi efficiency was tested by quantitative PCR (qPCR) using the primers nve14554F_q (5'-ATGGAGCGAGTGTTTCTGCT-3') and nve14554R_q (5'-CTTGCGCCATTTAGCTCTTC-3'), and the activity of the morpholino was checked by co-injecting it with the wild-type or the 5-mismatch mRNA containing the morpholino recognition sequence fused in-frame to the mCherry-coding sequence (Fig. S4B). Capped mRNA was synthesised using mMessage mMachine kit (Life Technologies) and purified with the Monarch RNA clean-up kit (New England BioLabs). For the inhibitor treatment, stock solutions of U0126 (Sigma-Aldrich) and SU5402 (Sigma-Aldrich) in DMSO were diluted to a final concentration of 15 µM and 20 µM in 16‰ artificial seawater, respectively. For DMSO controls, we used the same volume as for U0126, which was diluted 3:1000 from a less concentrated stock solution, which corresponds to a final DMSO

concentration of 42.3 mM. The treatment lasted from 30 h post fertilisation (hpf) until 3 days post fertilisation (dpf).

Acknowledgements

We thank Kevin Atkins for his help with setting up the sea anemone facility. We also thank Alix Harvey and Yousef Touhami for their support in maintaining the animal facility. We are grateful to the Core Facility for Cell Imaging and Ultrastructure Research of the University of Vienna for access to the confocal and the scanning electron microscopes.

Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualisation: T.L., S.P., G.G., V.M.; Methodology: E.G., C.T., T.L., G.G., V.M.; Software: S.P., N.C.; Validation: E.G., C.T., T.L., S.P., G.G., V.M.; Investigation: E.G., C.T., T.L., S.P., G.G., V.M.; Resources: G.G., V.M.; Data curation: E.G., C.T., V.M.; Writing - original draft: G.G., V.M.; Writing - review & editing: E.G., C.T., S.P.; Visualisation: T.L., G.G., V.M.; Supervision: G.G., V.M.; Funding acquisition: G.G., V.M.

Funding

E.G. is supported by an Advanced Research and Innovation in the Environmental Sciences (ARIES) Doctoral Training Partnership (DTP) PhD studentship funded by the Natural Environment Research Council (NERC), UK. This work in the Modepalli group was supported by the Anne Warner endowed Fellowship through the Marine Biological Association, UK. The work of the Genikhovich group was funded by the Austrian Science Fund (FWF) grant P30404-B29. Open access funding provided by UK Research and Innovation. Deposited in PMC for immediate release.

Data availability

RNA-seq data have been deposited at the Gene Expression Omnibus with the accession ID GSE159166. *Nematostella vectensis* gene models data are available on Figshare (https://figshare.com/articles/Nematostella_vectensis_transcriptome_ and_gene_models_v2_0/807696).

Peer review history

The peer review history is available online at https://journals.biologists.com/dev/ article-lookup/doi/10.1242/dev.200833.reviewer-comments.pdf.

References

- Altschul, S. F., Madden, T. L., Schäffer, A. A., Zhang, J., Zhang, Z., Miller, W. and Lipman, D. J. (1997). Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. *Nucleic Acids Res.* 25, 3389-3402. doi:10.1093/nar/25.17.3389
- Arendt, D., Tosches, M. A. and Marlow, H. (2016). From nerve net to nerve ring, nerve cord and brain — evolution of the nervous system. *Nat. Rev. Neurosci.* 17, 61-72. doi:10.1038/nrn.2015.15
- Babonis, L. S., Ryan, J. F., Enjolras, C. and Martindale, M. Q. (2019). Genomic analysis of the tryptome reveals molecular mechanisms of gland cell evolution. *EvoDevo* 10, 23. doi:10.1186/s13227-019-0138-1
- Bolger, A. M., Lohse, M. and Usadel, B. (2014). Trimmomatic: a flexible trimmer for Illumina sequence data. *Bioinformatics* **30**, 2114-2120. doi:10.1093/ bioinformatics/btu170
- Brauchle, M., Bilican, A., Eyer, C., Bailly, X., Martínez, P., Ladurner, P., Bruggmann, R. and Sprecher, S. G. (2018). Xenacoelomorpha survey reveals that all 11 animal homeobox gene classes were present in the first Bilaterians. *Genome Biol. Evol.* **10**, 2205-2217. doi:10.1093/gbe/evy170
- Capella-Gutierrez, S., Silla-Martinez, J. M. and Gabaldon, T. (2009). trimAl: a tool for automated alignment trimming in large-scale phylogenetic analyses. *Bioinformatics* 25, 1972-1973. doi:10.1093/bioinformatics/btp348
- Chia, F.-S. and Koss, R. (1979). Fine structural studies of the nervous system and the apical organ in the planula larva of the sea anemone Anthopleura elegantissima. J. Morphol. 160, 275-297. doi:10.1002/jmor.1051600303
- Columbus-Shenkar, Y. Y., Sachkova, M. Y., Macrander, J., Fridrich, A., Modepalli, V., Reitzel, A. M., Sunagar, K. and Moran, Y. (2018). Dynamics of venom composition across a complex life cycle. *Elife* 7, e35014. doi:10.7554/ eLife.35014
- Conzelmann, M., Offenburger, S.-L., Asadulina, A., Keller, T., Münch, T. A. and Jékely, G. (2011). Neuropeptides regulate swimming depth of Platynereis larvae. *Proc. Natl. Acad. Sci. USA* 108, E1174-E1183. doi:10.1073/pnas.1109085108
- Davidson, E. H., Peterson, K. J. and Cameron, R. A. (1995). Origin of Bilaterian body plans: evolution of developmental regulatory mechanisms. *Science* 270, 1319-1325. doi:10.1126/science.270.5240.1319
- Edgar, R. C. (2004). MUSCLE: a multiple sequence alignment method with reduced time and space complexity. *BMC Bioinform.* 5, 113. doi:10.1186/1471-2105-5-113

- Edgar, A., Ponciano, J. M. and Martindale, M. Q. (2022). Ctenophores are direct developers that reproduce continuously beginning very early after hatching. *Proc. Natl. Acad. Sci. USA* **119**, e2122052119. doi:10.1073/pnas.2122052119
- Frickey, T. and Lupas, A. (2004). CLANS: a Java application for visualizing protein families based on pairwise similarity. *Bioinformatics* 20, 3702-3704. doi:10.1093/ bioinformatics/bth444
- Gajewski, M., Leitz, T., Schloßherr, J. and Plickert, G. (1996). LWamides from Cnidaria constitute a novel family of neuropeptides with morphogenetic activity. *Roux's Arch. Dev. Biol.* 205, 232-242. doi:10.1007/BF00365801
- Garner, S., Zysk, I., Byrne, G., Kramer, M., Moller, D., Taylor, V. and Burke, R. D. (2016). Neurogenesis in sea urchin embryos and the diversity of deuterostome neurogenic mechanisms. *Development* 143, 286-297. doi:10.1242/dev.124503
- Genikhovich, G. and Technau, U. (2009a). Induction of spawning in the starlet sea anemone Nematostella vectensis, in vitro fertilization of gametes, and dejellying of zygotes. Cold Spring Harb. Protoc. 2009, pdb.prot5281. doi:10.1101/pdb.prot5281
- Genikhovich, G. and Technau, U. (2009b). In situ hybridization of starlet sea anemone (Nematostella vectensis) embryos, larvae, and polyps. *Cold Spring Harb. Protoc.* 2009, pdb.prot5282. doi:10.1101/pdb.prot5282
- Goldberg, J. I., Rich, D. R., Muruganathan, S. P., Liu, M. B., Pon, J. R., Tam, R., Diefenbach, T. J. and Kuang, S. (2011). Identification and evolutionary implications of neurotransmitter–ciliary interactions underlying the behavioral response to hypoxia in Lymnaea stagnalis embryos. J. Exp. Biol. 214, 2660-2670. doi:10.1242/jeb.053009
- Gruhl, A. (2009). Serotonergic and FMRFamidergic nervous systems in gymnolaemate bryozoan larvae. Zoomorphology 128, 135-156. doi:10.1007/ s00435-009-0084-x
- Guindon, S., Dufayard, J.-F., Lefort, V., Anisimova, M., Hordijk, W. and Gascuel, O. (2010). New algorithms and methods to estimate maximumlikelihood phylogenies: assessing the performance of PhyML 3.0. Syst. Biol. 59, 307-321. doi:10.1093/sysbio/syq010
- Hewes, R. S. and Taghert, P. H. (2001). Neuropeptides and neuropeptide receptors in the Drosophila melanogaster genome. *Genome Res.* **11**, 1126-1142. doi:10.1101/gr.169901
- Huang Da, W., Sherman, B. T. and Lempicki, R. A. (2009). Bioinformatics enrichment tools: paths toward the comprehensive functional analysis of large gene lists. *Nucleic Acids Res.* 37, 1-13. doi:10.1093/nar/gkn923
- Iwao, K., Fujisawa, T. and Hatta, M. (2002). A cnidarian neuropeptide of the GLWamide family induces metamorphosis of reef-building corals in the genus Acropora. *Coral Reefs* 21, 127-129. doi:10.1007/s00338-002-0219-8
- Jagersten, G. (1972). Evolution of the Metazoan Life Cycle. London: Academic Press.
- Jékely, G. (2011). Origin and early evolution of neural circuits for the control of ciliary locomotion. *Proc. Biol. Sci.* 278, 914-922. doi:10.1098/rspb.2010.2027
- Karabulut, A., He, S., Chen, C. Y., McKinney, S. A. and Gibson, M. C. (2019). Electroporation of short hairpin RNAs for rapid and efficient gene knockdown in the starlet sea anemone, Nematostella vectensis. *Dev. Biol.* 448, 7-15. doi:10. 1016/j.ydbio.2019.01.005
- Katsukura, Y., Ando, H., David, C. N., Grimmelikhuijzen, C. J. P. and Sugiyama, T. (2004). Control of planula migration by LWamide and RFamide neuropeptides in Hydractinia echinata. J. Exp. Biol. 207, 1803-1810. doi:10.1242/jeb.00974
- Kelava, I., Rentzsch, F. and Technau, U. (2015). Evolution of eumetazoan nervous systems: insights from cnidarians. *Philos. Trans. R. Soc. B Biol. Sci.* 370, 20150065. doi:10.1098/rstb.2015.0065
- Kitzmann, P., Weißkopf, M., Schacht, M. I. and Bucher, G. (2017). A key role for foxQ2 in anterior head and central brain patterning in insects. *Development* 144, 2969-2981. doi:10.1242/dev.147637
- Kraus, Y., Aman, A., Technau, U. and Genikhovich, G. (2016). Pre-bilaterian origin of the blastoporal axial organizer. *Nat. Commun.* 7, 11694. doi:10.1038/ ncomms11694
- Kuang, S., Doran, S. A., Wilson, R. J. A., Goss, G. G. and Goldberg, J. I. (2002). Serotonergic sensory-motor neurons mediate a behavioral response to hypoxia in pond snail embryos. J. Neurobiol. 52, 73-83. doi:10.1002/neu.10071
- Kumar, S., Tumu, S. C., Helm, C. and Hausen, H. (2020). The development of early pioneer neurons in the annelid Malacoceros fuliginosus. *BMC Evol. Biol.* 20, 117. doi:10.1186/s12862-020-01680-x
- Lacalli, T. C., Gilmour, T. H. J. and West, J. E. (1990). Ciliary band innervation in the bipinnaria larva of Pisaster ochraceus. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* 330, 371-390. doi:10.1098/rstb.1990.0206
- Langmead, B. and Salzberg, S. L. (2012). Fast gapped-read alignment with Bowtie 2. *Nat. Methods* 9, 357-359. doi:10.1038/nmeth.1923
- Layden, M. J., Rentzsch, F. and Röttinger, E. (2016). The rise of the starlet sea anemone Nematostella vectensis as a model system to investigate development and regeneration. WIREs Dev. Biol. 5, 408-428. doi:10.1002/wdev.222
- Lebedeva, T., Aman, A. J., Graf, T., Niedermoser, I., Zimmermann, B., Kraus, Y., Schatka, M., Demilly, A., Technau, U. and Genikhovich, G. (2021). Chidarianbilaterian comparison reveals the ancestral regulatory logic of the β-catenin dependent axial patterning. *Nat. Commun.* **12**, 4032. doi:10.1038/s41467-021-24346-8
- Leclère, L., Bause, M., Sinigaglia, C., Steger, J. and Rentzsch, F. (2016). Development of the aboral domain in Nematostella requires β-catenin and the

opposing activities of Six3/6 and Frizzled5/8. *Development* 143, 1766-1777. doi:10.1242/dev.120931

Leitz, T. and Lay, M. (1995). Metamorphosin A is a neuropeptide. *Roux's Arch. Dev. Biol.* 204, 276-279. doi:10.1007/BF00208495

- Leys, S. P. and Degnan, B. M. (2001). Cytological basis of photoresponsive behavior in a sponge Larva. *Biol. Bull.* 201, 323-338. doi:10.2307/1543611
- Liang, Y., Martín-Zamora, F. M., Guynes, K., Carrillo-Baltodano, A. M., Tan, Y., Moggioli, G., Seudre, O., Tran, M., Mortimer, K., Luscombe, N. M. et al. (2022). Annelid functional genomics reveal the origins of bilaterian life cycles. *bioRxiv*, 2022.02.05.479245. doi:10.1101/2022.02.05.479245
- Love, M. I., Huber, W. and Anders, S. (2014). Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome Biol.* 15, 550. doi:10.1186/s13059-014-0550-8
- Maldonado, M., Durfort, M., Mccarthy, D. A. and Young, C. M. (2003). The cellular basis of photobehavior in the tufted parenchymella larva of demosponges. *Mar. Biol.* **143**, 427-441. doi:10.1007/s00227-003-1100-1
- Marinković, M., Berger, J. and Jékely, G. (2020). Neuronal coordination of motile cilia in locomotion and feeding. *Philos. Trans. R. Soc. B Biol. Sci.* 375, 20190165. doi:10.1098/rstb.2019.0165
- Marlow, H. Q., Srivastava, M., Matus, D. Q., Rokhsar, D. and Martindale, M. Q. (2009). Anatomy and development of the nervous system of Nematostella vectensis, an anthozoan cnidarian. *Dev. Neurobiol.* 69, 235-254. doi:10.1002/ dneu.20698
- Marlow, H., Tosches, M. A., Tomer, R., Steinmetz, P. R., Lauri, A., Larsson, T. and Arendt, D. (2014). Larval body patterning and apical organs are conserved in animal evolution. *BMC Biol.* **12**. 7, doi:10.1186/1741-7007-12-7
- Martin, V. J. and Chia, F.-S. (1982). Fine structure of a Scyphozoan Planula, Cassiopeia xamachana. *Biol. Bull.* **163**, 320-328. doi:10.2307/1541269
- Matus, D. Q., Pang, K., Marlow, H., Dunn, C. W., Thomsen, G. H. and Martindale, M. Q. (2006). Molecular evidence for deep evolutionary roots of bilaterality in animal development. *Proc. Natl. Acad. Sci. USA* **103**, 11195-11200. doi:10.1073/pnas.0601257103
- Mazza, M. E., Pang, K., Reitzel, A. M., Martindale, M. Q. and Finnerty, J. R. (2010). A conserved cluster of three PRD-class homeobox genes (homeobrain, rx and orthopedia) in the Cnidaria and Protostomia. *EvoDevo* 1, 3. doi:10.1186/ 2041-9139-1-3
- Mi, H., Muruganujan, A. and Thomas, P. D. (2012). PANTHER in 2013: modeling the evolution of gene function, and other gene attributes, in the context of phylogenetic trees. *Nucleic Acids Res.* 41, D377-D386. doi:10.1093/nar/gks1118
- Nakanishi, N., Yuan, D., Jacobs, D. K. and Hartenstein, V. (2008). Early development, pattern, and reorganization of the planula nervous system in Aurelia (Cnidaria, Scyphozoa). *Dev. Genes Evol.* 218, 511-524. doi:10.1007/s00427-008-0239-7
- Nakanishi, N., Stoupin, D., Degnan, S. M. and Degnan, B. M. (2015). Sensory flask cells in sponge larvae regulate Metamorphosis via Calcium signaling. *Integr. Comp. Biol.* 55, 1018-1027. doi:10.1093/icb/icv014
- Nezlin, L. P. and Voronezhskaya, E. E. (2017). Early peripheral sensory neurons in the development of trochozoan animals. *Russ. J. Dev. Biol.* 48, 130-143. doi:10.1134/S1062360417020060
- Nielsen, C. (2005). Larval and adult brains1. *Evol. Dev.* 7, 483-489. doi:10.1111/j. 1525-142X.2005.05051.x
- Nielsen, C. (2012). Animal Evolution: Interrelationships of the Living Phyla. Oxford: OUP.
- Nielsen, C. (2013). Life cycle evolution: was the eumetazoan ancestor a holopelagic, planktotrophic gastraea? BMC Evol. Biol. 13, 171. doi:10.1186/1471-2148-13-171
- Nielsen, C. (2015). Larval nervous systems: true larval and precocious adult. J. Exp. Biol. 218, 629-636. doi:10.1242/jeb.109603
- Page, L. R. (2002). Apical sensory organ in larvae of the Patellogastropod Tectura scutum. *Biol. Bull.* 202, 6-22. doi:10.2307/1543218
- Pennati, R., Dell'Anna, A., Pagliara, P., Scari, G., Piraino, S. and De Bernardi, F. (2013). Neural system reorganization during metamorphosis in the planula larva of Clava multicornis (Hydrozoa, Cnidaria). *Zoomorphology* **132**, 227-237. doi:10.1007/s00435-013-0188-1
- Peterson, K. J., Cameron, R. A. and Davidson, E. H. (1997). Set-aside cells in maximal indirect development: Evolutionary and developmental significance. *BioEssays* 19, 623-631. doi:10.1002/bies.950190713
- Piraino, S., Zega, G., Di Benedetto, C., Leone, A., Dell'anna, A., Pennati, R., Candia Carnevali, D., Schmid, V. and Reichert, H. (2011). Complex neural architecture in the diploblastic larva of Clava multicornis (Hydrozoa, Cnidaria). *J. Comp. Neurol.* **519**, 1931-1951. doi:10.1002/cne.22614
- Quiroga Artigas, G., Lapébie, P., Leclère, L., Bauknecht, P., Uveira, J., Chevalier, S., Jékely, G., Momose, T. and Houliston, E. (2020). A G protein–coupled receptor mediates neuropeptide-induced oocyte maturation in the jellyfish Clytia. *PLoS Biol.* 18, e3000614. doi:10.1371/journal.pbio.3000614
- Raff, R. A. (2008). Origins of the other metazoan body plans: the evolution of larval forms. *Philos. Trans. R. Soc. Lond. B Biol. Sci.* 363, 1473-1479. doi:10.1098/rstb. 2007.2237
- Randel, N., Bezares-Calderón, L. A., Gühmann, M., Shahidi, R. and Jékely, G. (2013). Expression dynamics and protein localization of rhabdomeric opsins in Platynereis larvae. *Integr. Comp. Biol.* **53**, 7-16. doi:10.1093/icb/ict046

- Range, R. C., Angerer, R. C. and Angerer, L. M. (2013). Integration of canonical and noncanonical Wnt signaling pathways patterns the neuroectoderm along the anterior-posterior axis of sea urchin embryos. *PLoS Biol.* **11**, e1001467. doi:10.1371/journal.pbio.1001467
- Rentzsch, F., Fritzenwanker, J. H., Scholz, C. B. and Technau, U. (2008). FGF signalling controls formation of the apical sensory organ in the cridarian Nematostella vectensis. *Development* **135** 1761-1769 doi:10.1242/dev.020784
- Richards, G. S. and Degnan, B. M. (2012). The expression of Delta ligands in the sponge Amphimedon queenslandica suggests an ancient role for Notch signaling in metazoan development. *EvoDevo* **3**, 15. doi:10.1186/2041-9139-3-15
- Sachkova, M. Y., Singer, S. A., Macrander, J., Reitzel, A. M., Peigneur, S., Tytgat, J. and Moran, Y. (2019). The birth and death of toxins with distinct functions: a case study in the sea Anemone Nematostella. *Mol. Biol. Evol.* 36, 2001-2012. doi:10.1093/molbev/msz132
- Santagata, S., Resh, C., Hejnol, A., Martindale, M. Q. and Passamaneck, Y. J. (2012). Development of the larval anterior neurogenic domains of Terebratalia transversa (Brachiopoda) provides insights into the diversification of larval apical organs and the spiralian nervous system. *EvoDevo* 3, 3. doi:10.1186/2041-9139-3-3
- Schmich, J., Trepel, S. and Leitz, T. (1998). The role of GLWamides in metamorphosis of Hydractinia echinata. *Dev. Genes Evol.* 208, 267-273. doi:10.1007/s004270050181
- Sebé-Pedrós, A., Saudemont, B., Chomsky, E., Plessier, F., Mailhé, M.-P., Renno, J., Loe-Mie, Y., Lifshitz, A., Mukamel, Z., Schmutz, S. et al. (2018). Cnidarian cell type diversity and regulation revealed by whole-organism singlecell RNA-Seq. *Cell* **173**, 1520-1534.e20. doi:10.1016/j.cell.2018.05.019
- Sinigaglia, C., Busengdal, H., Leclère, L., Technau, U. and Rentzsch, F. (2013). The bilaterian head patterning gene six3/6 controls aboral domain development in a cnidarian. *PLoS Biol.* **11**, e1001488. doi:10.1371/journal.pbio.1001488
- Sinigaglia, C., Busengdal, H., Lerner, A., Oliveri, P. and Rentzsch, F. (2015). Molecular characterization of the apical organ of the anthozoan Nematostella vectensis. *Dev. Biol.* **398**, 120-133. doi:10.1016/j.ydbio.2014.11.019
- Sly, B. J., Snoke, M. S. and Raff, R. A. (2003). Who came first–larvae or adults? Origins of bilaterian metazoan larvae. *Int. J. Dev. Biol.* 47, 623-632.
- Soliman, S. (1983). Pharmacological control of ciliary activity in the young sea urchin larva. Effects of monoaminergic agents. *Comp. Biochem. Physiol. C Comp. Pharmacol.* 76, 181-191. doi:10.1016/0742-8413(83)90061-0
- Soliman, S. (1984). Pharmacological control of ciliary activity in the young sea urchin larva. Studies on the role of Ca2+ and cyclic nucleotides. *Comp. Biochem. Physiol. C Comp. Pharmacol.* 78, 183-191. doi:10.1016/0742-8413(84)90067-7
- Steinmetz, P. R. H., Aman, A., Kraus, J. E. M. and Technau, U. (2017). Gut-like ectodermal tissue in a sea anemone challenges germ layer homology. *Nat. Ecol. Evol.* 1, 1535-1542. doi:10.1038/s41559-017-0285-5
- Tamm, S. L. (2014). Cilia and the life of ctenophores. *Invertebr. Biol.* 133, 1-46. doi:10.1111/ivb.12042
- Thiel, D., Yañez Guerra, L. A., Franz-Wachtel, M., Hejnol, A. and Jékely, D. (2021). Nemertean, brachiopod, and phoronid neuropeptidomics reveals ancestral spiralian signaling systems. *Mol. Biol. Evol.* 38, 4847-4866. doi:10.1093/molbev/msab211
- Ueda, N., Richards, G. S., Degnan, B. M., Kranz, A., Adamska, M., Croll, R. P. and Degnan, S. M. (2016). An ancient role for nitric oxide in regulating the animal pelagobenthic life cycle: evidence from a marine sponge. *Sci. Rep.* 6, 37546. doi:10.1038/srep37546
- Verasztó, C., Ueda, N., Bezares-Calderón, L. A., Panzera, A., Williams, E. A., Shahidi, R. and Jékely, G. (2017). Ciliomotor circuitry underlying whole-body coordination of ciliary activity in the Platynereis larva. *Elife* 6, e26000. doi:10.7554/ eLife.26000
- Verasztó, C., Gühmann, M., Jia, H., Rajan, V. B. V., Bezares-Calderón, L. A., Piñeiro-Lopez, C., Randel, N., Shahidi, R., Michiels, N. K., Yokoyama, S. et al. (2018). Ciliary and rhabdomeric photoreceptor-cell circuits form a spectral depth gauge in marine zooplankton. *Elife* 7, e36440. doi:10.7554/eLife.36440
- Voronezhskaya, E. E., Tyurin, S. A. and Nezlin, L. P. (2002). Neuronal development in larval chiton Ischnochiton hakodadensis (Mollusca: Polyplacophora). J. Comp. Neurol. 444, 25-38. doi:10.1002/cne.10130
- Williams, E. A. and Jékely, G. (2019). Neuronal cell types in the annelid Platynereis dumerilii. Curr. Opin. Neurobiol. 56, 106-116. doi:10.1016/j.conb.2018.12.008
- Wolenski, F. S., Layden, M. J., Martindale, M. Q., Gilmore, T. D. and Finnerty, J. R. (2013). Characterizing the spatiotemporal expression of RNAs and proteins in the starlet sea anemone, Nematostella vectensis. *Nat. Protoc.* 8, 900-915. doi:10.1038/nprot.2013.014
- Wolpert, L. (1999). From egg to adult to larva. *Evol. Dev.* **1**, 3-4. doi:10.1046/j.1525-142x.1999.00111.x
- Woollacott, R. M. (1993). Structure and swimming behavior of the larva of Haliclona tubifera (Porifera: Demospongiae). J. Morphol. 218, 301-321. doi:10.1002/jmor. 1052180306
- Zang, H. and Nakanishi, N. (2020). Expression analysis of cnidarian-specific neuropeptides in a sea anemone unveils an apical-organ-associated nerve net that disintegrates at metamorphosis. *Front. Endocrinol.* **11**, 63. doi:10.3389/fendo. 2020.00063

2.0	0.	.0	2.0						
BB	LPB	PAO	PAO-						
٩.	_	₽.		NVE_Geneld NVE8371	Gene ID MOUSE MGI MGI 108078 P97299	Symbol SFRP2	Gene Name Secreted frizzled-related protein 2	Protein Class transmembrane signal receptor	GO transmembrane signal receptor (PC00197)
				NVE10382 NVE19736	FELCA Ensembl ENSFCAG0000025878 M3X1M2 BOVIN Ensembl ENSBTAG0000020957 F1MGM1	HMCN1 FZD5	Uncharacterized protein Frizzled class receptor 5	transmembrane signal receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
				NVE10381	ANOCA Ensembl ENSACAG0000008287 H9GE51	NCAM1	Uncharacterized protein Uncharacterized protein		neuron projection (GO:0043005)
	_			NVE6997 NVE8133	BOVIN Gene HMCN1 E1BE11 MOUSE MGI MGI 2441950 Q80T53	AGRL3	Adhesion G protein-coupled receptor L3	tyrosine protein kinase receptor G-protein coupled receptor	transmembrane signal receptor (PC00197) transmembrane signal receptor (PC00197)
	_			NVE15843 NVE23855	MOUSE MGI MGI 1860130 Q924H0 HUMAN HGNC 13800 Q9GZZ6	NPFF2 ACH10	Neuropeptide FF receptor 2 Neuronal acety(choline receptor subunit alpha-10	G-protein coupled receptor ligand-gated ion channel	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
				NVE2185 NVE26112	CAEBR Gene CBG01491 A8WQK3 ANOCA Ensembl ENSACAG00000012709 H9GJH9	ACH1 EPHAS	Acetylcholine receptor subunit alpha-type acr-16 Uncharacterized protein	ligand-gated ion channel	neuron projection (GO:0043005) neuron projection (GO:0043005)
				NVE4515 NVE22264	HUMAN HGNC 14006 Q9H1D0 DANRE ZFIN ZDB-GENE-081022-145 B3DHM6	TRPV6 ADRB1	Transient receptor potential cation channel subfamily V member 6 Adrenergic, beta-1-, receptor	ion channel G-protein coupled receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
				NVE4175	BOVIN Gene HMCN1 E1BE11	HMCN1	Uncharacterized protein	tyrosine protein kinase receptor	transmembrane signal receptor (PC00197)
	_			NVE3518 NVE5022	BOVIN Gene SCNN1A P55270 HUMAN HGNC 11048 P23975	SCNNA NET1	Amiloride-sensitive sodium channel subunit alpha Sodium-dependent noradrenaline transporter	ion channel primary active transporter	neuron projection (GD:0043005) neuron projection (GD:0043005)
			1000	NVE5065 NVE7261	CAEEL WormBase WBGene00001519 Q9U7D5 DROME FlyBase FBgn0053517 Q8I544 ANOCA Ensembl ENSACAG00000013826 G1KPD2	ACM3 DRD2L	Muscarinic acetylcholine receptor gar-3 Dopamine D2-like receptor	G-protein coupled receptor G-protein coupled receptor	neuron projection (GD:0043005) transmembrane signal receptor (PC00197)
				NVE15552 NVE1646	ANOCA Ensembl ENSACAG00000013826 G1KPD2 BOVIN Gene CHRNA7 P54131	SYT1 ACHA7	Uncharacterized protein Neuronal acetylcholine receptor subunit alpha-7	membrane trafficking regulatory protein ligand-gated ion channel	neuron projection (GD:0043005) neuron projection (GD:0043005)
				NVE17372 NVE13969	HUMAN HGNC 5302 P34969 CAEEL WormBase WBGene00006765 P48181	SHT7R ACH2	5-hydroxytryptamine receptor 7 Acetylcholine receptor subunit beta-type unc-29	G-protein coupled receptor ligand-gated ion channel	neuron projection (GO:0043005) neuron projection (GO:0043005)
	_			NVE4045	HUMAN1HGNC1138001109GZZ6	ACH10	Neuronal acetvicholine receptor subunit alpha-10	ligand-gated ion channel	neuron projection (GO:0043005)
				NVE21238 NVE10029	BOVIN Gene HMCN1 E1BE11 BOVIN Gene BSG Q865R3	BASI	Uncharacterized protein Basigin (Fragment)	tyrosine protein kinase receptor scaffold/adaptor protein	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
_			-	NVE21441 NVE2635	ANOCA Ensembl ENSACAG00000011789 G1KLI5 HUMAN HGNC 1463 Q13555	CHRNA9 KCC2G	Uncharacterized protein Calcium/calmodulin-dependent protein kinase type II subunit gamma	ligand-gated ion channel non-receptor serine/threonine protein kinase	neuron projection (GD:0043005) neuron projection (GD:0043005)
	_			NVE5017 NVE6180	HUMAN HGNC 1396 Q9P0X4 RAT RGD 2348 Q05941	CAC1I ACHA7	Voltage-dependent T-type calcium channel subunit alpha-11 Neuronal acetylcholine receptor subunit alpha-7	voltage-gated ion channel ligand-gated ion channel	neuron projection (GO:0043005) neuron projection (GO:0043005)
	_			NVE13547	MOUSE MGI MGI 2441950 Q80TS3	AGRL3	Adhesion G protein-coupled receptor L3	G-protein coupled receptor	transmembrane signal receptor (PC00197)
	_			NVE14451 NVE1484	CHICK Ensembl ENSGAL G00000029392 P19019 DROME FlyBase FBgr0004622 P30975	GBRB3 TLR2	Gamma-aminobutyric acid receptor subunit beta-3 Tachykinin-like peptides receptor 99D	ligand-gated ion channel G-protein coupled receptor	neuron projection (GD:0043005) transmembrane signal receptor (PC00197)
				NVE7617 NVE12387	HUMAN HGNC 10588 Q99250 BOVIN Gene CAMK2D Q2HJF7	NAC2 KCC2D	Sodium channel protein type 2 subunit alpha Calcium/calmodulin-dependent protein kinase type II subunit delta	voltage-gated ion channel non-receptor serine/threonine protein kinase	neuron projection (GD:0043005) neuron projection (GD:0043005)
				NVE13402 NVE23178	CHICK [Ensembl]ENSGALG0000004466 Q75XU5 DROME [FlyBase FBgn0004168 P28285	NPFF1 SHT2A	Neuropeptide FF receptor 1 5-hydroxytryptamine receptor 2A	G-protein coupled receptor G-protein coupled receptor	transmembrane signal receptor (PC00197) somatodendritic compartment (GO:0036477)
				NVE25668	DROME FlyBase FBgn0000036 P09478	ACH1	Acetylcholine receptor subunit alpha-like 1	ligand-gated ion channel	neuron projection (GO:0043005)
				NVE22703 NVE24524	DANRE ZFIN ZDB-GENE-040801-92 Q6DC77 HUMAN HGNC 11018 Q9H598	MPRGB VIAAT	Membrane progestin receptor gamma-B Vesicular inhibitory amino acid transporter	G-protein coupled receptor	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
			_	NVE7386 NVE21442	DROME FlyBase FBgn0038880 Q8IN35 HUMAN HGNC 15781 Q494W8	SIFAR CRFM7	Neuropeptide SiFamide receptor CHRNA7-FAM7A fusion protein	G-protein coupled receptor ligand-gated ion channel	transmembrane signal receptor (PC00197) neuron projection (GC:0043005)
				NVE20517 NVE2517	MOUSE MGI MGI 1860130 (0924H0 DANRE ZFIN ZDB-GENE-070705-309 (ADAOR4IVA4	NPFF2 CC14A	Neuropeptide FF receptor 2 Dual specificity protein phosphatase CDC14AB	G-protein coupled receptor	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
				NVE8357	HUMAN HGNC 5299 Q13639	SHT4R	5-hydroxytryptamine receptor 4	G-protein coupled receptor	neuron projection (GO:0043005)
	-			NVE23188 NVE13933	ANOCA Ensembl ENSACAG00000014117 G1KPl2 MOUSE MGI MGI 2677838 A2AJ76	SYT4 HMCN2	Uncharacterized protein Hemicentin-2	membrane trafficking regulatory protein transmembrane signal receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
	_		-	NVE1327 NVE17221	FELCA Gene OPN4 Q5YKK9 MOUSE MGI MGI 104311 P32240	OPN4 PE2R4	Melanopsin Prostaglandin E2 receptor EP4 subtype	G-protein coupled receptor G-protein coupled receptor	transmembrane signal receptor (PC00197) transmembrane signal receptor (PC00197)
	_			NVE10027 NVE19228	CHICK Ensembli ENSGALG000000498481 (ADA302TXF0	CNR1 5HT4R	Cannabinoid receptor 5-hydroxytryptamine receptor 4	G-protein coupled receptor G-protein coupled receptor	transmembrane signal receptor (PC00197) somatodendritic compartment (GO:0036477)
	_			NVE15844	HUMAN HGNC 5299 Q13639 MACMU Ensembl ENSMMUG00000020241 Q9GK75	NPY1R	Neuropeptide Y receptor Y1	G-protein coupled receptor	transmembrane signal receptor (PC00197)
				NVE17029 NVE1501	MDUSE MGI MGI 2429764 P69744 CANLF Ensembl ENSCAFG00000013828 P11617	CAT2 AAZAR	Transient receptor potential cation channel subfamily V member 5 Adenosine receptor A2a	ion channel G-protein coupled receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
			_	NVE12996 NVE26217	RAT RGD 1308804 Q5XIU9 FELCA Gene QPN4 Q5YKX10	PGRC2 OPN5	Membrane-associated progesterone receptor component 2 Melanopsin	transmembrane signal receptor G-protein coupled receptor	transmembrane signal receptor (PC00197) transmembrane signal receptor (PC00197)
				NVE6519 NVE10190	FELCA Gene OPN4 Q5YKK10 CAEEL WormBase WBGene00016610 Q94177	ADRL OCTB1	Progestin and adipoQ receptor-like protein 1 Octopamine receptor beta-1R	G-protein coupled receptor G-protein coupled receptor	transmembrane signal receptor (PC00197)
-		_		NVE21472	DROME FlyBase FBgn0038980 Q9VCZ3 NEMVE EnsemblGenome NEMVEDRAFT_v1g248535 A7T1N0	TMP2L	Transient receptor potential cation channel subfamily M member-like 2	ion channel	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
				NVE24515 NVE25785	MOUSE MGI MGI 892032 1997401 MOUSE MGI MGI 99456 1001727	FRITZ MSHR	Secreted frizzled-related protein 3 Melanocyte-stimulating hormone receptor	transmembrane signal receptor G-protein coupled receptor	transmembrane signal receptor (PC00197) transmembrane signal receptor (PC00197)
	_		-	NVE6925 NVE14868	HUMAN HGNC 10585 P35498 MOUSEIMGUMGU26646681077N88	NAC1 PK1L2	Sodium channel protein type 1 subunit alpha Polycystic kidney disease protein 1-like 2	voltage-gated ion channel ion channel	neuron projection (GD:0043005) neuron projection (GD:0043005)
	_	_		NVE17821	MOUSE MGI MGI 2664668 Q7TN88 ANOCA Ensembl ENSACAG0000005650 G1KEG9	GLRA3 EPHB4	Uncharacterized protein	ligand-gated ion channel	neuron projection (GO:0043005)
_			_	NVE12212 NVE7681	BOVIN Ensembl ENSBTAG0000003446 E1BIE4 BOVIN Gene HMCN1 E1BE11	HMCN1	EPH receptor B4 Uncharacterized protein	tyrosine protein kinase receptor	neuron projection (GD:0043005) transmembrane signal receptor (PC00197)
				NVE24164 NVE25358	BOVIN Gene HMCN1 E1BE11 HUMAN HGNC 11119 Q99835	HMCN1 SMO	Uncharacterized protein Smoothened homolog	tyrosine protein kinase receptor transmembrane signal receptor	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
			_	NVE19422 NVE3573	DROME FlyBase FBgn0086906 Q917U4 MOUSE MGI MGI 97175 P20357	TITIN MTAP2	Titin Microtubule-associated protein 2	tyrosine protein kinase receptor	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
				NVE21871 NVE2492	MOUSE MGI MGI 1196627 Q9R1C8 DANRE Ensembl ENSDARG00000013649 F1R4B10	SHT6R CASR	5-hydroxytryptamine receptor 6 Calcium-sensing receptor	G-protein coupled receptor G-protein coupled receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
				NVE16130	CHICK Gene FSHR P79763	FSHR	Follicle-stimulating hormone receptor	G-protein coupled receptor	transmembrane signal receptor (PC00197)
				NVE11916 NVE19915	HUMAN HGNC 21293 Q8NDA2 CHICK Ensembl ENSGALG0000002118 093393	HMCN2 CO5	Hemicentin-2 Calcium-activated potassium channel beta subunit	transmembrane signal receptor voltage-gated ion channel	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
			-	NVE13710 NVE19835	CAEEL WormBase T23D8.1b A0A0K3AWM6 DROME FlyBase FBgn0038880 Q8IN35	MOM5 SIFAR	Protein mom-5 Neuropeptide SIFamide receptor	transmembrane signal receptor G-protein coupled receptor	transmembrane signal receptor (PC00197) transmembrane signal receptor (PC00197)
				NVE17972 NVE2156	DANRE [ZFIN ZDB-GENE-990714-19 Q9W7R4 CHICK Gene CHRNA10 Q9I8C7	TENM3 ACH10	Teneurin-3 Neuronal acetylcholine receptor subunit alpha-10	- ligand-gated ion channel	neuron projection (GO:0043005) neuron projection (GO:0043005)
				NVE20802	DANRE ZFIN ZDB-GENE-080221-1 F1Q681	NPY1R	Neuropeptide Y receptor Y1	G-protein coupled receptor	transmembrane signal receptor (PC00197)
				NVE8662 NVE19257	HUMAN HGNC 11049 Q01959 RAT RGD 1564772 F1M4Q3	SC6A3 HMCN1	Sodium-dependent dopamine transporter Hemicentin 1	primary active transporter	neuron projection (GD:0043005) neuron projection (GD:0043005)
			_	NVE275 NVE9473	RAT RGD 621570 Q9EP86 DROME FlyBase FBgn0004842 P25931	NPFF1 RYAR	Neuropeptide FF receptor 1 RYamide receptor	G-protein coupled receptor G-protein coupled receptor	transmembrane signal receptor (PC00197) transmembrane signal receptor (PC00197)
			_	NVE4599 NVE12144	DRDME FlyBase FBgn0004841 P30974 HUMAN HGNC 14079 Q9UGM1	TLR1 ACHA9	Tachykinin-like peptides receptor 86C	G-protein coupled receptor ligand-gated ion channel	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
				NVE14952	RAT RGD 1564772 F1M4Q3	HMCN1	Neuronal acetylcholine receptor subunit alpha-9 Hemicentin 1		neuron projection (GO:0043005)
				NVE8663 NVE19358	MOUSE MGI MGI 96285 Q60857 DANRE ZFIN ZDB-GENE-991117-1 093430	SC6A4 GLRA1	Sodium-dependent serotonin transporter Glycine receptor subunit alphaZ1	primary active transporter ligand-gated ion channel	neuron projection (GD:0043005) neuron projection (GD:0043005)
				NVE24392 NVE4207	HUMAN HGNC 9967 P07949 CHICK Ensembl ENSGALG00000042626 ADA1D5PCG4	RET SYT4	Proto-oncogene tyrosine-protein kinase receptor Ret Uncharacterized protein	membrane trafficking regulatory protein	neuron projection (GO:0043005) neuron projection (GO:0043005)
				NVE9028 NVE1388	DROME FlyBase FBgn0034476 Q7KIN0 CHICK Gene MC1R P55167	TOLL7 MSHR	Toll-like receptor 7 Melanocyte-stimulating hormone receptor	transmembrane signal receptor G-protein coupled receptor	transmembrane signal receptor (PC00197) transmembrane signal receptor (PC00197)
				NVE25364	RAT RGD 1564772 F1M4Q3 MOUSE MGI MGI 98328 P30875	HMCN1	Hemicentin 1 Somatostatin receptor type 2		neuron projection (G0:0043005) neuron projection (G0:0043005)
				NVE21855 NVE21136	DANRE Ensembl ENSDARG00000013649 F1R488	SSR2 CASR	Calcium-sensing receptor	G-protein coupled receptor G-protein coupled receptor	transmembrane signal receptor (PC00197)
				NVE23050 NVE20106	RAT RGD 1564772 F1M4Q3 DANRE Ensembl ENSDARG00000013649 F1R489	HMCN1 CASR	Hemicentin 1 Calcium-sensing receptor	- G-protein coupled receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
				NVE2371 NVE7629	CHICK Ensembl ENSGAL G0000002118 093393 DROME FlyBase FBgn0004622 P30975	CO6 TLR2	Calcium-activated potassium channel beta subunit Tachykinin-like peptides receptor 99D	voltage-gated ion channel G-protein coupled receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
				NVE141 NVE11735	MOUSE MGI MGI 3041203 Q80732 MOUSE MGI MGI 95749 Q91XP5	AGRD1 GLRA3	Adhesion Grpotein coupled receptor D1 Glycine receptor subunit alpha-3	G-protein coupled receptor ligand-gated ion channel	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
				NVE11735 NVE2876 NVE25008	RAT RGD 628837 P60711	ACTB NPC2	Actin, cytoplasmic 1 Stathmin-3	actin and actin related protein	neuron projection (GO:0043005)
				NVE4080	CHICK Ensembl ENSGALG0000006130 093388 RAT RGD 1564772 F1M4Q3	HMCN1	Hemicentin 1		neuron projection (GO:0043005) neuron projection (GO:0043005)
				NVE5966 NVE9440	MOUSE MGI MGI 95749 Q91XP5 RAT RGD 1305841 D4A6L0	GLRA3 GP158	Glycine receptor subunit alpha-3 Probable G-protein coupled receptor 158	ligand-gated ion channel G-protein coupled receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
				NVE18296 NVE17393	MOUSE MGI MGI 97281 P13595 MOUSE MGI MGI 1347520 Q92139	NCAM1 ROR1	Neural cell adhesion molecule 1 Inactive tyrosine-protein kinase transmembrane receptor ROR1		neuron projection (GO:0043005) neuron projection (GO:0043005)
				NVE9727 NVE20788	MOUSE MGI MGI 99402 Q60613 HUMAN HGNC 3039 O60469	AAZAR	Adenosine receptor A2a Down syndrome cell adhesion molecule	G-protein coupled receptor scaffold/adaptor protein	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
				NVE6463	DROME FlyBase FBgr0038880 Q8IN36	SIFAR	Neuropeptide SIFamide receptor	G-protein coupled receptor	transmembrane signal receptor (PC00197)
				NVE13212 NVE26151	RAT RGD 2353 P09660 MOUSE MGI MGI 1347095 Q920M6	ACHE CD97	Acetylcholine receptor subunit epsilon CD97 antigen	ligand-gated ion channel G-protein coupled receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
				NVE6241 NVE7440	HUMAN HGNC 11333 P31391 HUMAN HGNC 18992 Q8IZF6	SSR4 AGRG4	Somatostatin receptor type 4 Adhesion G-protein coupled receptor G4	G-protein coupled receptor G-protein coupled receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
				NVE8550	RAT RGD 619891 Q8VHD7	GPR83	G protein-coupled receptor 83	G-protein coupled receptor	transmembrane signal receptor (PC00197)
				NVE2244 NVE3729	CHICK Ensembl ENSGALG00000015126 ADA1D5PAY3 HUMAN HGNC 10450 014718	TRPM3 OPSX	Uncharacterized protein Visual pigment-like receptor peropsin	ion channel G-protein coupled receptor	neuron projection (GD:0043005) neuron projection (GD:0043005)
				NVE12143 NVE16819	RAT RGD 1564772 F1M4Q3 DANRE ZFIN ZDB-GENE-050913-90 A0T2N3	HMCN1 APJB	Hemicentin 1 Apelin receptor B	G-protein coupled receptor	neuron projection (GO:0043005) transmembrane signal receptor (PC00197)
				NVE12516 NVE13879	MOUSE MGI MGI 1333877 088495 DANRE ZFIN ZDB-GENE-060427-5 E7FCE4	MTR1L ROR2	Melatonin-related receptor Receptor tyrosine kinase-like orphan receptor 2	G-protein coupled receptor	transmembrane signal receptor (PC00197) neuron projection (GD:0043005)
				NVE24406	HUMAN HGNC 21293 Q8NDA2	HMCN2	Hemicentin-2	transmembrane signal receptor	transmembrane signal receptor (PC00197)
				NVE19889 NVE16011	HUMAN HGNC 4600 000222 MOUSE MGI MGI 104295 Q60898	GRM8 SLAP1	Metabotropic glutamate receptor 8 Src-like-adapter	G-protein coupled receptor	transmembrane signal receptor (PC00197) neuron projection (GO:0043005)
				NVE21548 NVE10021	MOUSE MGI MGI 99779 P49582 RAT RGD 1564772 F1M4Q3	ACHA7 HMCN1	Neuronal acetylcholine receptor subunit alpha-7 Hemicentin 1	ligand-gated ion channel	neuron projection (GD:0043005) neuron projection (GD:0043005)
				NVE2362	CHICK Ensembl ENSGALG0000002118 093393	CO6	Calcium-activated potassium channel beta subunit	voltage-gated ion channel	neuron projection (GO:0043005)
				NVE1835 NVE15043	HUMAN HGNC 4039 Q9ULW2 DANRE ZFIN ZDB-GENE-070705-262 A5WWA0	FZD10 ZNRF3	Frizzled-10 E3 ubiquitin-protein ligase znrf3	transmembrane signal receptor transmembrane signal receptor	transmembrane signal receptor (PC00197) transmembrane signal receptor (PC00197)
				NVE5945 NVE7065	RAT RGD 1564772 F1M4Q3 RAT RGD 1564772 F1M4Q3	HMCN1	Hemicentin 1 Hemicentin 1		neuron projection (GD:0043005) neuron projection (GD:0043005)

Fig. S1. Additional neuronal associated genes DE between the apical and oral regions.

Heatmap shows the relative expression of neuronal associated genes among oral or apical region of *Nematostella* larvae. The gene id and additional details are shown on the right.

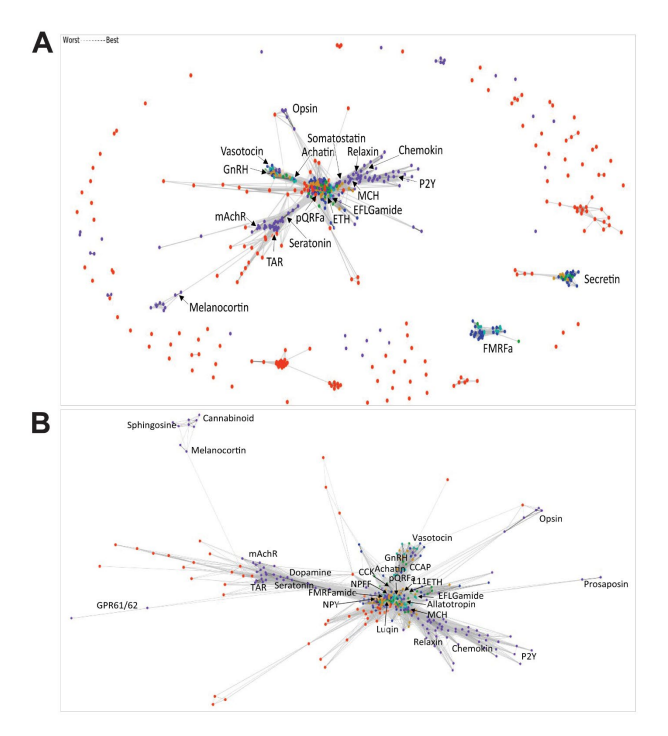


Fig.S2. (A) Sequence-similarity-based clustering of all *Nematostella* apical enriched GPCRs. Forty-six *Nematostella* GPCRs clustered with a large group of Rhodopsin and four clustered with Secretin. **(B)** Cluster map of the Rhodopsin cluster presented in A.

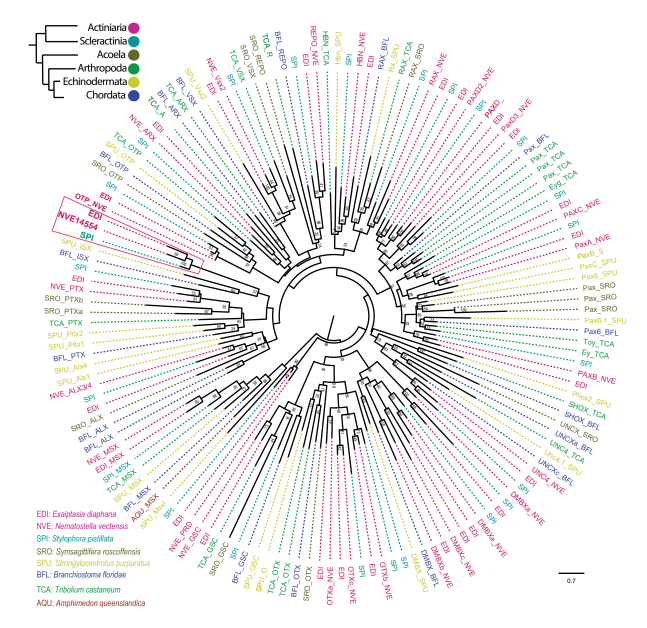


Fig. S3. Maximum likelihood phylogenetic tree of PRD class homeobox of selected animals. The homeobox sequences of a representative selection of Cnidarians and bilaterian lineages **(Supplementary Table 1)** were aligned with MUSCLE algorithm [77] in the SeaView program [78]. The SMS: Smart Model Selection in PhyML [79] was used to determine the best-fitting model for the Phylogenic reconstruction VT model (+G+F). The maximum-likelihood (ML) phylogenetic trees were constructed using PhyML v3.0 with 100 bootstrap [76]. Only bootstrap values of at least 20 at nodes are shown. The MSX family of ANTP class homeobox was used as an outgroup.

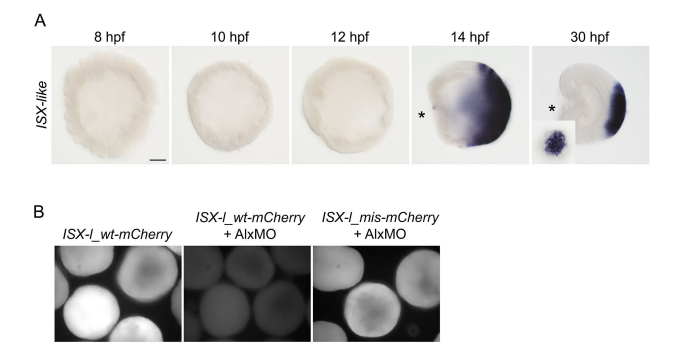


Fig. S4. RNAi and morpholino efficiency. (A) Early expression of ISX-like in *Nematostella* embryos. Inset on the 30 hpf gastrula image shows the aboral view. **(B)** Translation of the *mCherry* mRNA carrying the wild type recognition site of the ISX-like MO fused to it is efficiently suppressed by co-injecting ISX-like MO. In contrast, *mCherry* mRNA carrying the recognition site of the ISX-like MO with five introduced mismatches is efficiently translated in the presence of ISX-like MO. Lateral and aboral views, on lateral views, asterisk denotes the oral end. Scale bars 50 μm.

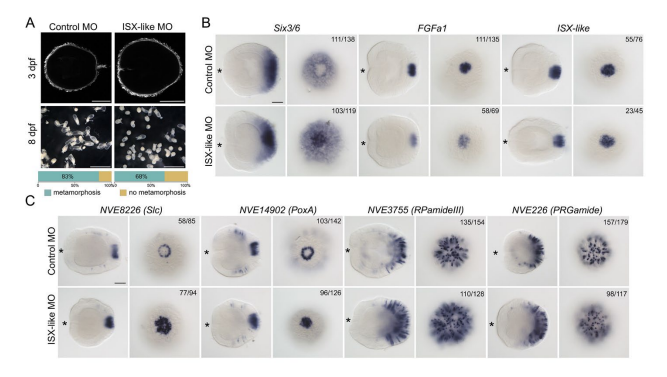


Fig. S5. Effect of the morpholino-mediated knockdown of *ISX-like* phenocopies *ISX-like* RNAi. (A) Morpholino knockdown of *ISX-like* leads to a loss of the apical tuft (anti-acetylated tubulin staining, top row) and a mild reduction in the metamorphosis rate. (B) In ISX-like morphants, Six3/6 expression expands into the apical domain, however, *FGFa1* and *ISX-like* are not strongly affected. (C) In ISX-like morphants, the expression of the ring genes extends into the pot domain, but the expression of the aborally enriched *PRGamide* (*NVE226*), and *RPamide III* (*NVE3775*) does not appear to be strongly affected. Lateral and aboral views, on lateral views, asterisk denotes the oral end. Scale bars on (A) 100 μ m, on (B) and (C) 50 μ m.

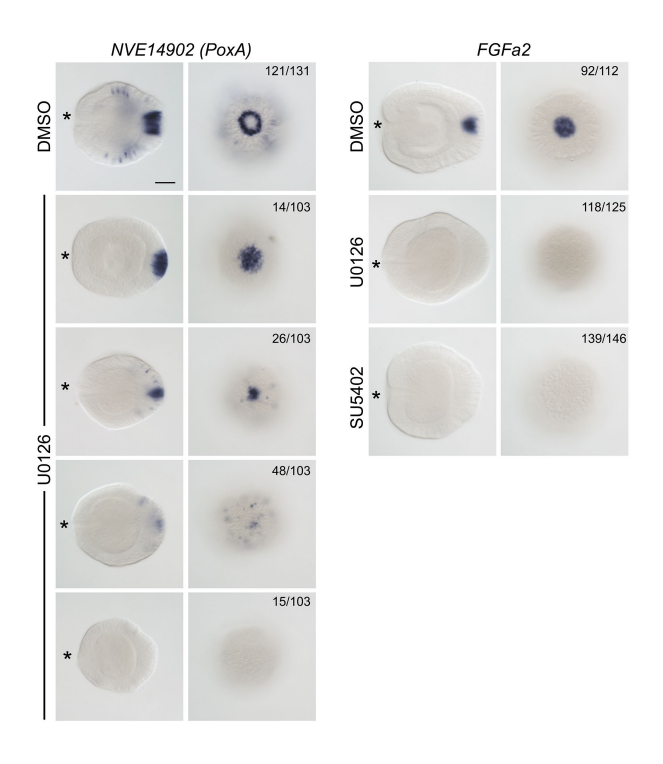
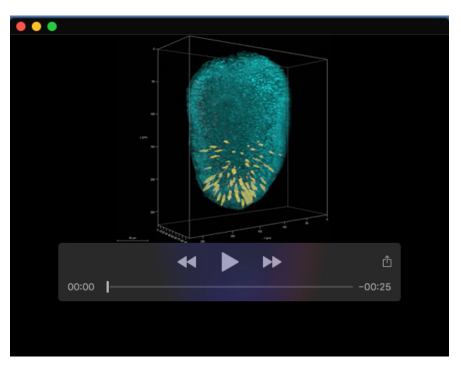


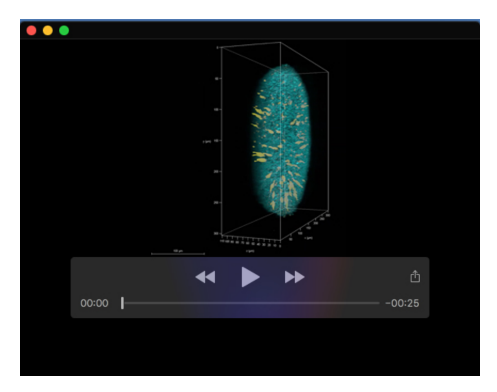
Fig. S6. Variability of the effect of the U0126 treatment on NVE14902 expression and the effect of U0126 and SU5402 on FGFa2 expression. Scale bar 50 μm.

Table S1. Data relevant to figures 2, 3, 4, S1, S2 and S3.

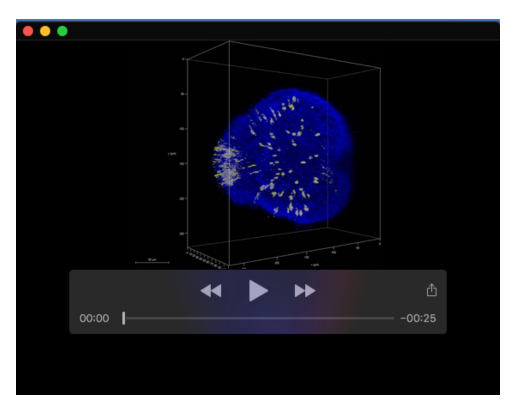
Click here to download Table S1



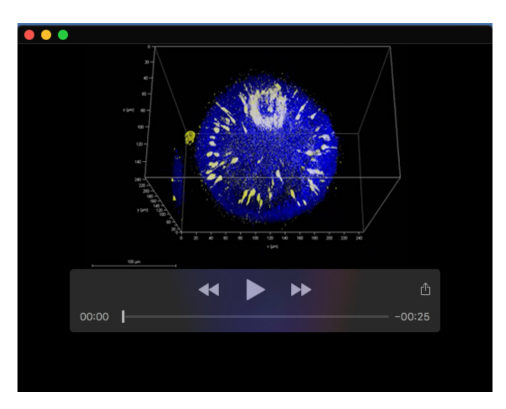
Movie 1. 3D view of fluorescence in situ hybridisation for gland/secretory cells type 1, related to Figure 5B&C. gland/secretory cells type 1 marker NVE22589 (yellow) and DAPI in cyan.



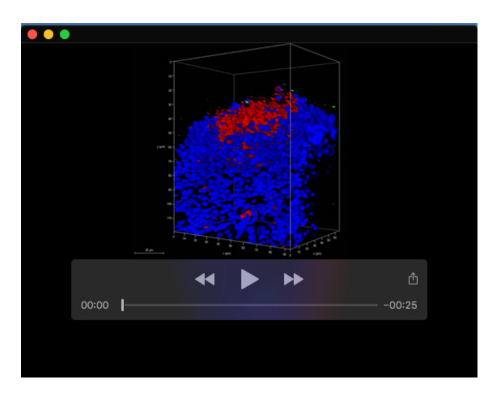
Movie 2. 3D view of fluorescence in situ hybridisation for gland/secretory cells type 3, related to Figure 5F. Gland/secretory cells type 3 marker NVE23674 (yellow) and DAPI in cyan.



Movie 3. 3D view of fluorescence in situ hybridisation for larval-specific neuron gene, related to Figure 6A. *NVE8226* mRNA probe (yellow) and DAPI in Blue.



Movie 4. 3D view of fluorescence in situ hybridisation for larval-specific neuron gene, related to Figure 6E. *NVE14902* mRNA probe (yellow) and DAPI in Blue.



Movie 5. 3D view of Double fluorescence in situ hybridisation for spot probe *NVE14554* (green) and rind NVE8226 (red) mRNA probes, related to Figure 6L. DAPI in Blue.

