

Neuronal identities derived by misexpression of the POU IV sensory determinant in a protovertebrate

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The protovertebrate *Ciona intestinalis* type A (sometimes called *Ciona robusta*) contains a series of sensory cell types distributed across the head–tail axis of swimming tadpoles. They arise from lateral regions of the neural plate that exhibit properties of vertebrate placodes and neural crest. The sensory determinant *POU IV/Brn3* is known to work in concert with regional determinants, such as *Foxg* and *Neurogenin*, to produce palp sensory cells (PSCs) and bipolar tail neurons (BTNs), in head and tail regions, respectively. A combination of single-cell RNA-sequencing (scRNA-seq) assays, computational analysis, and experimental manipulations suggests that misexpression of *POU IV* results in variable transformations of epidermal cells into hybrid sensory cell types, including those exhibiting properties of both PSCs and BTNs. Hybrid properties are due to coexpression of *Foxg* and *Neurogenin* that is triggered by an unexpected *POU IV* feedback loop. Hybrid cells were also found to express a synthetic gene battery that is not coexpressed in any known cell type. We discuss these results with respect to the opportunities and challenges of reprogramming cell types through the targeted misexpression of cellular determinants.

cell-type specification | evolutionary developmental biology | computational biology | cellular reprogramming

The specification of sensory neurons from lateral regions of the neural plate in protovertebrate *Ciona intestinalis* type A provides insights into the evolutionary origins of sensory placodes and the neural crest in vertebrates (1–10). Previous studies have identified early anteroposterior determinants that subdivide the lateral ectoderm into anterior, trunk, and tail regions (*Foxc*, *Six1/2*, and *Mxsb*, respectively) (9). These trigger regulatory networks that produce related but distinct sensory neurons, including palp sensory cells (PSCs), anterior apical trunk epidermal neurons (aATENs), and bipolar tail neurons (BTNs). All of these cell types feature robust expression of *POU IV* (Fig. 1A), a key regulatory determinant of sensory neurons in vertebrates (*Brn3*) (11). These studies suggest that *POU IV* produces a sensory “ground state” that is modulated by additional determinants such as *Six1/2* to specify distinctive neuronal cell types.

POU IV has also been implicated in the development of caudal epidermal sensory neurons (CESNs) (12, 13). These are arranged in single rows along the dorsal and ventral midlines of the *Ciona* tail, where they are thought to serve mechanosensory functions (14–16). Previous misexpression studies suggest that *POU IV* is sufficient to transform epidermal cells into supernumerary sensory neurons (13). Epidermal-specific *cis*-regulatory DNAs were used to drive expression of *POU IV* throughout the epidermis. This led to the surprising observation that the entire epidermis is neurogenic, not just the midline regions. Transformed epidermal cells have the appearance of supernumerary CESNs, suggesting that they might represent an ancestral sensory ground state. In this study, we employ single-cell RNA-sequencing (scRNA-seq) assays to examine their identities.

We used the *CesA* epidermal enhancer (17) to misexpress *POU IV/Brn3* throughout the epidermis of developing tailbud embryos. As documented previously (13), there is a striking expansion of epidermal sensory neurons. However, single-cell transcriptome analyses are not consistent with a simple transformation of epidermis into CESNs. Instead, transformed cells express many of the genes associated with a BTN sensory identity. Different subclusters of transformed epidermal cells also express varying numbers of genes associated with other sensory identities, PSCs, aATENs, and CESNs. Moreover, transformed cells also express a set of genes that are not a specific transcriptome signature of any known cell type in the *Ciona* embryo or tadpole, which we will refer to as a “synthetic gene battery.” These results give insight into the opportunities and challenges of manipulating gene regulatory networks to reprogram cell types.

Results and Discussion

POU IV is a POU/homeobox gene that is expressed in all of the sensory neurons of the *Ciona* tadpole (Fig. 1A) (12).

Significance

The protovertebrate *Ciona intestinalis* is an ideal system to investigate both gene regulatory networks that underlie cell-type specification and how cell types have evolved. In this study, we use single-cell technology, experimental manipulations, and computational analyses to understand the role of the regulatory determinant *POU IV*—a homolog of *Brn3* in vertebrates—in specifying various sensory cell types in *Ciona*. Surprisingly, the misexpression of *POU IV* throughout the epidermis led to the formation of hybrid sensory cell types, including those exhibiting properties of both palp sensory cells and bipolar tail neurons. These results demonstrate the interconnectedness of diverse sensory specification networks and give insights into the opportunities and challenges of reprogramming cell types through the targeted misexpression of cellular determinants.

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The authors declare no competing interest.

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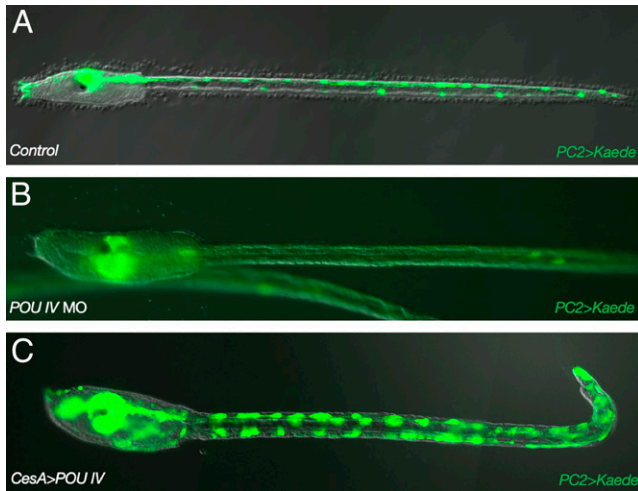


Fig. 1. *POU IV* is necessary for the specification of epidermal sensory neurons. *PC2>Kaede* transgenic larvae. (A) Kaede fluorescence in the central nervous system (CNS) and epidermal sensory neurons in control larvae. (B) Same as A except that larvae were injected with a *POU IV* MO. (C) Same as A except that *POU IV* was misexpressed throughout the epidermis by a *CesA* 5' regulatory sequence.

Homologous genes in vertebrates (*Bmn3*) are essential for the development of a variety of sensory cell types, including visual, auditory/vestibular, and somatosensory neurons (18–20). Mutant embryos injected with a *POU IV* morpholino oligonucleotide (MO) at the one-cell stage resulted in the loss of expression of a pan-neuronal reporter gene (*PC2>Kaede*) in all sensory cell types at the larval stage (Fig. 1B), demonstrating that *POU IV* is an essential sensory determinant in *Ciona*. To determine whether it is sufficient for specification, *POU IV* was misexpressed throughout the epidermis of *Ciona* tadpoles using regulatory sequences from the *CesA* gene (Fig. 1C), which encodes a cellulase synthase that is essential for the elaboration of the protective test, a defining property of all urochordates (17). The *CesA>POU IV* transgene resulted in expanded expression of *PC2>Kaede*, suggesting transformation of epidermal cells into neurons (Fig. 1C).

To determine the nature of this transformation, we performed scRNA-seq of dissociated cells from transgenic tadpoles using the 10X Chromium System (9, 21–23). All wild-type sensory neurons, wild-type epidermal cells, and transformed epidermal cells were readily identified by their expression of a variety of known marker genes (Fig. 2A and *SI Appendix, Tables S1 and S2*). Transformed epidermal cells formed a heterogeneous population that exhibited differential expression of genes associated with each sensory cell type (Fig. 2A and *SI Appendix, Fig. S1*).

This motivated us to subcluster the transformed cells into “misexpressed” (ME) clusters and model them as linear combinations of “average” wild-type cells based on their gene expression profiles (*Materials and Methods*, Fig. 2B, and *SI Appendix, Fig. S2*). We interpreted the coefficient values based on the extent to which an ME cell exhibited characteristics of the corresponding wild-type cell identity. In all subclusters, except for the BTN/CESN hybrid subcluster featured in Fig. 2D, BTN coefficients have the highest average values and right-skewed distributions, which implies that ME cells have the strongest bias toward a BTN-like identity. This is an unexpected result since previous studies suggested transformations of epidermal cells into CESNs (7, 9).

We also observed a distribution of coefficient values corresponding to all sensory cell types among ME cells, which implies “mixed” or “hybrid” sensory identities (*SI Appendix,*

Fig. S3 and Table S3). For example, the purple and pink subclusters circled in Fig. 2C consist of ME cells with high average PSC and CESN coefficient values, respectively. We hereafter refer to these cells as “BTN/PSC” and “BTN/CESN” hybrids (Fig. 2D and E). Moreover, ME cells concurrently express the PSC and BTN determinants *Neurogenin* and *Foxg*, respectively (7, 9, 10, 24) (*SI Appendix, Fig. S4*). This led us to take a particular interest in understanding the regulatory basis of the BTN/PSC hybrid identity.

While it is known that *Foxg* and *Neurogenin* directly bind and regulate *POU IV* expression (9), we explored the possibility of feedback regulation of both “upstream” determinants by *POU IV*. Thus, misexpression of *POU IV* could result in coactivation of both *Foxg* and *Neurogenin* in the transformed epidermis, a situation that is not encountered in wild-type sensory neurons. Computational analyses identified *POU IV*-binding motifs in the 5' regulatory regions of both *Foxg* and *Neurogenin* (*Materials and Methods* and Fig. 3), and reporter genes containing these binding motifs exhibit specific expression in PSCs and BTNs, respectively (Fig. 3A–D). Moreover, the *Neurogenin* enhancer contains a greater number of, and stronger, *POU IV* binding sites than *Foxg*. This correlates with altered activities of the *Neurogenin* and *Foxg* enhancers in mutant tailbud embryos injected with a *POU IV* MO at the one-cell stage (Fig. 4A and B and *SI Appendix, Fig. S5*). Expression of the *Neurogenin* enhancer is abolished, while the *Foxg* enhancer is diminished.

There is a similar loss in activity of the BTN-specific minimal *Neurogenin* enhancer upon mutagenesis of *POU IV* binding sites (Fig. 3E and F). Enhancer activity is also lost in tadpoles injected with either *POU IV* MO or *Neurogenin* MO (Fig. 4A–D). Taken together, these studies suggest feedback regulation of *Neurogenin*, and to a lesser extent *Foxg*, by *POU IV*.

To determine whether this regulatory feedback results in the coactivation of *Foxg* and *Neurogenin*, we performed targeted misexpression of *POU IV* in BTNs using the minimal *Neurogenin* enhancer and targeted misexpression of *Neurogenin* in PSCs using the minimal *Foxg* enhancer. Both cases led to hybrid neuronal cell types that express both BTN (*Asic1b*) and PSC ($\beta\gamma$ -crystallin) marker genes (Fig. 4E–H) (25, 26). Similar results were obtained upon targeted misexpression of *Foxg*, which functions upstream of *Foxg* and *POU IV* (9, 10). These findings demonstrate interconnectedness of the *Foxg*–*POU IV* and *Neurogenin*–*POU IV* regulatory networks.

We next sought to understand two key aspects of ME cell identity—their strong BTN bias, and expression of a synthetic gene battery that is not observed in normal cell populations (ME Epi, Fig. 2A and B). ME cells might exhibit a preponderance of BTN-like properties due to a more efficient induction of *Neurogenin* than *Foxg* by *POU IV*, since the *Neurogenin* enhancer is predicted to contain higher-affinity *POU IV* binding sites than its *Foxg* counterpart (Fig. 2). Consistent with this possibility is the finding of a more complete transformation of PSCs into BTNs upon targeted misexpression of both *POU IV* and *Neurogenin* (*SI Appendix, Figs. S6 and S7*). Interestingly, the combination of *Bmn3* and *Neurogenin* was found to be sufficient to reprogram human fibroblasts into different sensory cell types comprising dorsal root ganglia, a derivative of the neural crest (27). Our studies are therefore consistent with previous evidence that BTNs may correspond to derivatives of a “protoneural crest” in the *Ciona* lateral neural plate (7, 9).

To explore the characteristics of the synthetic gene battery that is deployed in hybrid sensory cell types, we performed both functional annotation and statistical overrepresentation Gene Ontology (GO) analyses of the human orthologs of these genes (*Materials and Methods*). Over 65% encode cell transporters, transmembrane signal receptors, protein-modifying enzymes, and metabolic enzymes. In addition, there is a >100-fold enrichment of various kinases, synthases, membrane

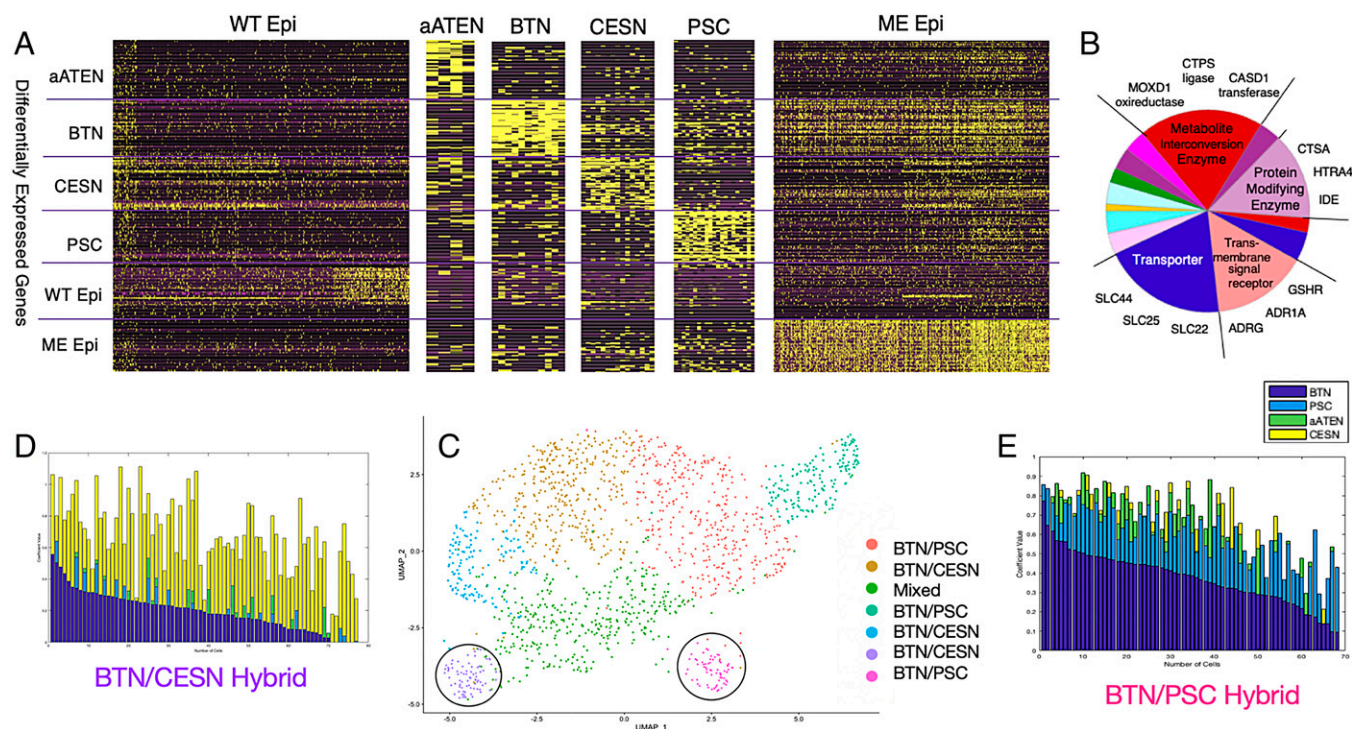


Fig. 2. Quantitative characterization of wild-type and *POU IV*-misexpressed epidermis gene expression patterns. (A) Visualization of expression levels of 300 transcriptomic profile genes of wild-type epidermis and aATEN, BTN, CESN, PSC, and *POU IV* ME cell populations. Rows are the top 50 DEGs from each cell-type population (300 transcriptomic profile). Columns are individual cells. (B) Pie chart of the main PANTHER protein classes of ME DEGs and representative genes belonging to each class that define the various subclusters. More detailed information is provided in *SI Appendix, Fig. S10*. (C) UMAP visualization of seven subclusters generated from Louvain community-based clustering of ME cells. Circled purple and pink clusters contain representative BTN/CESN and BTN/PSC hybrid cells, respectively. (D and E) Stacked barplots of cell type-specific solved coefficients (*Materials and Methods*) for each cell in BTN/PSC and BTN/CESN subclusters, respectively. Solved aATEN, BTN, CESN, and PSC coefficients are represented by green, dark blue, yellow, and light blue bars, respectively. Cells are ordered by decreasing values of BTN solved coefficients, that is, “decreasing BTN character” and “increasing PSC/CESN character.”

transporters, and cell receptor functions (Fig. 2B and *SI Appendix, Figs. S8–S12*). Some of these genes are implicated in the biogenesis of lysosomes and exosomes, such as cathepsin. These are not signatures of cell stress, senescence, or death but instead suggest hyperactive growth. This surprising, emergent property of transformed ME cells is not predicted by the features of the regulatory networks underlying the specification of the different sensory cell types.

To understand what may have caused the induction of the synthetic gene battery, we examined up to 2 kb of their upstream regulatory regions and performed motif analysis (*Materials and Methods*). Our discovered motif (*SI Appendix, Fig. S13*) had significant matches to various Forkhead transcription factors in *Ciona* and their orthologs in mouse and human (*SI Appendix, Fig. S14*). Given this analysis and the up-regulation of *Foxg* expression throughout ME cells (*SI Appendix, Fig. S1*), we postulate that *Foxg* broadly associates with Forkhead binding sites to activate the synthetic gene battery. We also noted an up-regulation of other Forkhead transcription factors (*SI Appendix, List S1*), which might lead to further activation. These observations highlight the hazards and uncertainties created by targeted misexpression of cellular determinants such as *POU IV*.

Our study provides evidence for a *POU IV* feedback loop that maintains the identities of distinct sensory neurons, including PSCs and BTNs (Fig. 5). Misexpression of *POU IV* leads to ectopic activation of upstream regulators such as *Foxg* and *Neurogenin* via these feedback loops. Normally, these determinants are separately expressed in PSCs and BTNs but are coactivated upon misexpression of *POU IV*. This results in the development

of hybrid sensory neurons exhibiting properties of both PSCs and BTNs. We consistently observe higher levels of expression of BTN identity genes as compared with PSC markers. In addition, these cells express genes not found in sensory cell types but are implicated in lysosomes, exosomes, and hyperactive growth. We postulate that they are fortuitously activated by the binding of *Foxg* and other Forkhead transcription factors.

It has been suggested that the diversification of sensory cell types in vertebrates was facilitated by the duplication and divergence of *POU IV/Bm3* paralogs (28). However, *Ciona* contains just a single *POU IV* ortholog but nonetheless produces diverse sensory cells. This study, along with several previous reports, suggests that this lone *POU IV* ortholog works in concert with regional determinants such as *Foxg*, *Six1/2*, and *Neurogenin* to specify distinct yet related sensory cell types (9). Misexpression of *POU IV* short circuits these cell-specific programs to produce a variety of hybrid cell types due to coexpression of *Foxg*, *Neurogenin*, and other regional determinants. In addition, we also observe fortuitous induction of a synthetic gene battery that is not coordinately expressed in any known natural cell type. It seems unlikely that they are induced by the combination of *Foxg* and *Neurogenin*. Instead, we found that their regulatory regions contain a significant overrepresentation of Forkhead-binding motifs. Thus, the success of future efforts to reprogram cell types with defined properties will depend on employing strategies to predict and diminish induction of unwanted secondary gene expression patterns. We believe that the strategy employed in our study—the iterative application of computational predictions and experimental manipulations of the underlying gene regulatory networks—will provide the precision required to achieve this goal.

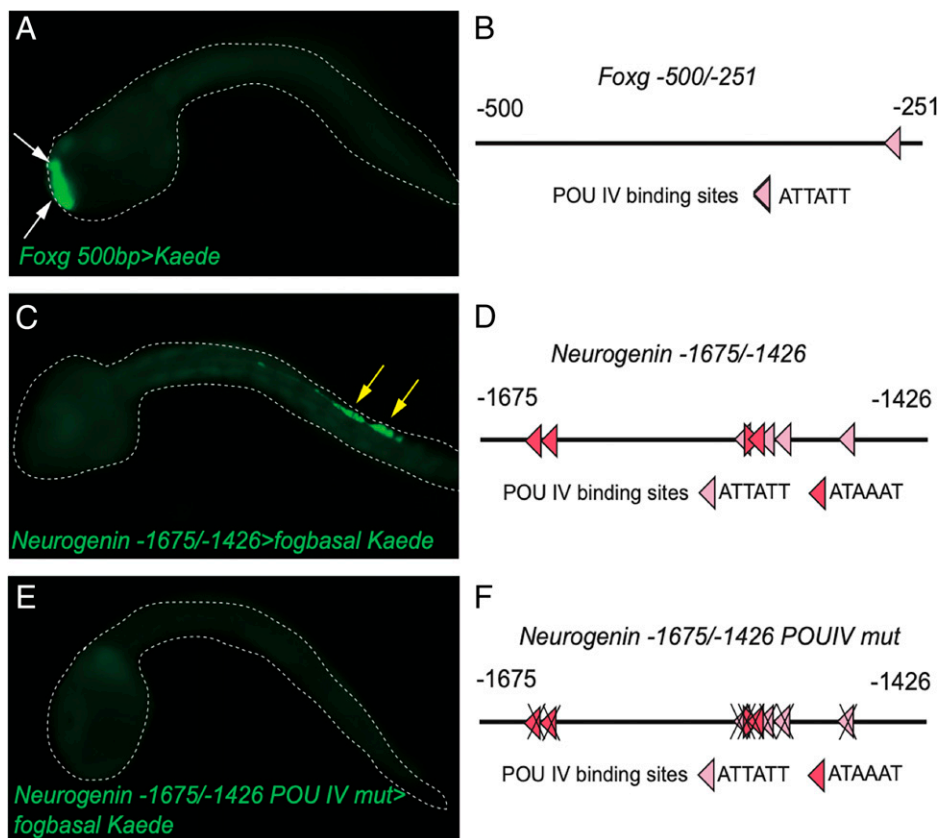


Fig. 3. POU IV binding sites are present in the PSC minimal enhancer region of *Foxg* and BTN minimal enhancer region of *Neurogenin* and are necessary for the expression of *Neurogenin* in BTNs. (A) Kaede fluorescence in *Foxg 500bp>Kaede*-injected embryos. White arrows indicate Kaede fluorescence in PSCs. (B) Schematic diagram of -500/-251 bp of the PSC-specific *Foxg* minimal enhancer. One POU IV binding site is present. (C) Kaede fluorescence of *Neurogenin -1675/-1472>Kaede*-injected embryos. Yellow arrows indicate Kaede fluorescence in BTNs. (D) Schematic diagram of -1675/-1472 bp of the BTN-specific minimal enhancer. Eight POU IV binding sites are present. (E) Mutation analysis of eight POU IV binding sites showed that these POU IV binding sites are necessary for expression of *Neurogenin* in BTNs. (F) Schematic diagram of mutation analysis in E. Dark red binding sites are the higher-affinity POU IV binding sites.

Materials and Methods

Biological Materials. Wild-type *C. intestinalis* type A (also called *Ciona robusta*) adults were obtained from M-REP and the National Bio-Resource Project for *Ciona* in Japan. Sperm and eggs were collected by dissecting the sperm and gonadal ducts. The *PC2>Kaede* transgenic lines (29, 30) were cultured and maintained in an island system (31).

Constructs. Reporter genes were designed using previously published enhancer sequences: *Asic1b* (32), β_2 -*Crystallin* (26), and *CesA* (17). The minimal enhancers of *Foxg*, *Foxg -500/-201*, and of *Neurogenin*, *Neurogenin -1675/-1426*, were isolated via PCR using sequence-specific oligonucleotide primers (SI Appendix, Table S4) and cloned into the *XhoI/BamHI* restriction site of the pSPfogbasalKaede vector. To generate pSPCiPOUIVcDNA, the coding sequence of *Ci-POU IV* was amplified by PCR with sequence-specific oligonucleotide primers (SI Appendix, Table S4). The PCR product was digested with *NotI* and inserted into the *NotI* and blunted *EcoRI* sites of pSPeGFP. To generate pSPCiCesACiPOUIVcDNA, the 5' upstream region of *Ci-CesA* was inserted into the *BamHI* site of pSPCiPOUIVcDNA. To generate pSPCineurogenin-POUIVcDNA, the *Neurogenin -1675/-1426*+fogbasal promoter was inserted into the *XhoI* and *NotI* sites of pSPCiPOUIVcDNA. To generate pSPFoxgPOUIV, *Foxg 500bp* was inserted into the *XhoI* and *NotI* sites of pSPCiPOUIVcDNA.

Microinjection of Antisense Morpholino Oligonucleotides. MOs were obtained from Gene Tools. The antisense oligonucleotide sequence of the MO against *Ci-POU IV* is 5'-gcacgttagtaaacatctgtatca-3'. MOs were dissolved in nuclease-free water (AM9930; Invitrogen) containing 1 mg/mL tetramethylrhodamine dextran (D1817; Invitrogen). The concentrations of MO and plasmid DNA in the injection medium were 0.5 mM and 2.5 to 10 ng/ μ L, respectively. Microinjections of MOs and reporter constructs were performed as described previously (9). All experiments were repeated at least twice with different batches of embryos.

Image Acquisition. Images of transgenic larvae were obtained with a Zeiss AxioPlan, AX 10 epifluorescence microscope.

Single-Cell RNA-Seq Assays. *CesA>POU IV* (2.5 ng/ μ L) + *CesA>GFP* (5.0 ng/ μ L)-injected eggs and control eggs were fertilized side by side and allowed to develop to the late-tailbud stage (13.5 h after fertilization at 18°C). For each sample, 120 morphologically normal embryos were used for scRNA-seq assays. Dissociation of the embryos and scRNA-seq assays by the 10X Genomics Chromium System were done as described previously (9).

Computational Analysis Notes. "OE cell" refers to a cell in *POU IV*-misexpressed epidermis. OEs and MEs are used interchangeably, and all code/lists use OE nomenclature. All code, analyses, and files can be accessed from the GitHub repository https://github.com/Singh-Lab/Pou4_Misexpression.

Single-Cell Data Analysis.

Data quality control and visualization. According to the Cell Ranger 3.0 cell-calling algorithm based on the EmptyDrops algorithm (33), we removed cells using a cutoff of UMIs (unique molecular identifier) 5 SDs above the mean and those expressing fewer than 1,000 genes. This rendered 5,078 total cells derived from both control and *POU IV*-misexpressed embryos with 1,163 median genes per cell. All analyses were performed using Seurat version 2.3.4. Cells were further filtered so that only cells expressing at least 200 genes that are expressed in at least three cells were considered. For each cell, gene expression measurements were normalized by total gene expression, multiplied by a factor of 10,000, and then log-transformed. Gene expression values were regressed on UMI count data to remove unwanted variation, and residuals were then scaled and centered. These gene expression values were used for downstream analyses. Variable genes were outliers on a mean variability plot, calculated, and used as inputs for principal-component analysis (PCA) dimension reduction. For cells in the control embryos, 10 PCs were used as

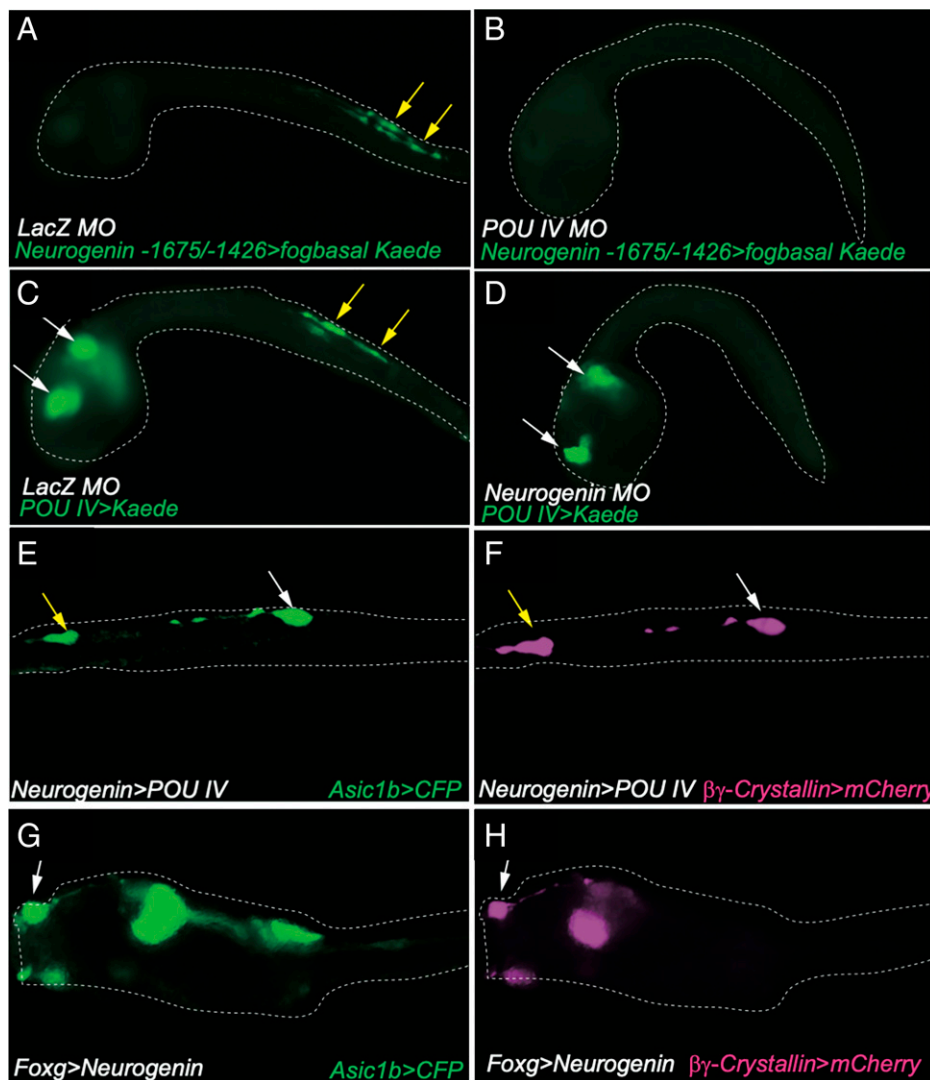


Fig. 4. Positive feedback loop between *POU IV* and *Neurogenin* in BTN and synthetic cell-type specifications. (A and B) Kaede fluorescence of *Neurogenin* -1675/-1472bp>*Kaede*-injected embryos. (A) Kaede fluorescence in BTNs in *LacZ* MO-injected embryos. Yellow arrows indicate Kaede fluorescence in BTNs. (B) Same as A except that larvae were injected with the *POU IV* MO. (C and D) Kaede fluorescence of *POU IV*>*Kaede*-injected embryos. (C) Kaede fluorescence in BTNs and CNS in *LacZ* MO-injected embryos. Yellow and white arrows indicate Kaede fluorescence in BTNs and CNS, respectively. (D) Same as C except that larvae were injected with the *Neurogenin* MO. Kaede fluorescence disappears in BTNs but is maintained in the CNS. (E and F) Misexpression of *POU IV* in BTNs leads to expression of the PSC marker gene ($\beta\gamma$ -*Crystallin*) in BTNs. *Asic1b*>*CFP* expression (E) and $\beta\gamma$ -*Crystallin*>*mCherry* (F) expression in *Neurogenin*>*POU IV*-injected larvae. (G and H) Misexpression of *Neurogenin* in PSCs leads to expression of the BTN marker gene (*Asic1b*) in PSCs. *Asic1b*>*CFP* expression (G) and $\beta\gamma$ -*Crystallin*>*mCherry* (H) expression in *Foxg*>*Neurogenin*-injected larvae.

inputs to UMAP (Uniform Manifold Approximation and Projection) dimension reduction. For cells in *POU IV*-misexpressed embryos, 16 PCs were used. The resultant PCs were subjected to the unsupervised learning Louvain modularity community detection algorithm to cluster cells, which were then visualized in UMAP plots. Quality control metrics and figures are in *SI Appendix, Figs. S15–S18*.

Identification of cell populations. Epidermal markers were used to identify 1,683 and 1,570 epidermal cell populations in both control and *POU IV*-misexpressed embryos, respectively. Four aATENs, 11 BTNs, and 21 PSCs were identified using their marker genes (*SI Appendix, Table S1*). We do not currently possess marker genes for CESNs, but 26 of them were identified as cells with the following code: *POU IV*+ *Klf*+ *Neurogenin*– β -Thymosine, in which + indicates expression and – indicates lack of expression (*SI Appendix, Table S2*). All genes in these tables and their corresponding gene IDs from the KH Assembly Model (34) (KHIDs) were obtained from the Aniseed database (35).

Calculation of differentially expressed genes and definition of transcriptomic profiles. For each cell population, the edgeR differential expression analysis package (36) was used to calculate differentially expressed genes (DEGs) between itself and select cell populations. Backgrounds were decided based on our knowledge of similarity of gene expression patterns. For

example, since CESNs and PSCs have very similar gene expression patterns, DEGs were calculated with respect to one another, to sufficiently distinguish them. aATEN DEGs were calculated with respect to CESNs and PSCs. BTN DEGs were calculated with respect to aATEN, CESN, PSC, and wild-type epidermis. Wild-type and ME epidermis DEGs were calculated with respect to all other cell-type populations. (Note: We refer to ME epidermis DEGs as “novel genes.”) All resultant DEGs were ordered by false discovery rate–corrected *P* values of less than 0.05, and then by nondecreasing log–fold change values. The top 50 most highly expressed DEGs for each cell population were identified, save for CESNs, which had 48 DEGs after this correction. We defined “transcriptomic profile” as a means of quantifying cell population identity. There were two such profiles. The first was the “300” transcriptomic profile, which consisted of the expression of all 298 DEGs. The other was the “250” transcriptomic profile, which consisted of the expression of the first 248 DEGs (all DEGs excluding those of ME cells).

Linear model to quantify OE cell identity with respect to wild-type cell population identities. This model approximates the 250 transcriptomic profile of an ME cell as a linear combination of those of average wild-type cells. According to *SI Appendix, Fig. S1*, in the 250×5 matrix (wild-type matrix), rows are 250 transcriptome profile genes and columns are “average” wild-type vectors. A

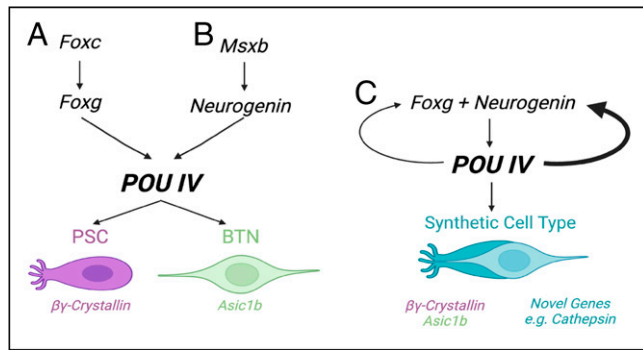


Fig. 5. Summary of gene regulatory networks underlying PSC, BTN, and synthetic cell-type specifications. The schematics pertain to (A) PSCs, (B) BTNs, and (C) synthetic cell types found in *POU IV*-misexpressed embryos.

given entry in an average vector is the average expression level of a 250 transcriptome profile gene in all cells belonging to the same cell population. The 250×1 matrix (OE vector) contains the expression levels of the 250 transcriptome profile genes for a given OE cell. The entries in the 5×1 vector comprise a nonnegative linear least-squares solution to this system of equations. Each entry is a “cell-type specific solved coefficient.” Nonnegative linear least-squares solutions (solution vector) were calculated using the `lsqnonneg` function in MATLAB (version 9.2, release name R2017a). We then wanted to retain cells that were well-described by the linear modeling. To do this, the solution vector was multiplied by the wild-type matrix, and a Spearman correlation between this resultant vector and the OE vector was calculated using the `corr` function. If the Spearman correlation had a P value < 0.05 , the corresponding solution matrix was retained for downstream analyses. Distributions of the coefficients of statistically significant solutions (cell-type specific solved coefficients) were plotted in histograms and stacked bar plots, as visualized in *SI Appendix, Fig. S2*.

Identification of *POU IV* binding sites in *Foxg* and *Neurogenin* minimal enhancer regions. *POU IV* binding sites were found from Selex data for *POU IV* (KH2012:KH.C2.42) provided by the Aniseed database. “ATAAAT” has a higher binding affinity than “ATAATT.” Instances of these motifs were found in the sequences of the minimal enhancer regions by a simple string search.

Identification of putative 5' cis-regulatory regions of novel genes. Genomic sequences for novel genes were manually obtained from the Washington Genome Browser. Their KHIDs and human orthologs are given in

“Final_OE_khids_orthologs.csv” and the genomic coordinates used are in “manual_oe_intervals.txt.” Up to 2 kb sequence upstream of the transcriptional start site was analyzed. If the sequence overlapped a gene body of a neighboring gene, it was rejected. This rendered a total of 33 sequences.

Discovery of and matches to motifs in putative 5' cis-regulatory regions of novel genes. Putative cis-regulatory regions were used as inputs to MEME (37) with these parameters: motif site distribution: ANR; maximum number of motifs: 3; minimum motif length: 5; maximum motif length: 15 (*SI Appendix, Fig. S13*). One out of three discovered motifs had a significant match (q value < 0.05) to various *Ciona* Forkhead proteins. All motifs were used as inputs to TOMTOM (38) with the CIS-BP_2.00/*Ciona intestinalis* motif database. Results are in *SI Appendix, Fig. S14 A–E*.

GO analyses of novel genes. Human orthologs of 50 novel genes were used as input to the PANTHER Classification System (39), with “*Homo sapiens*” as the selected organism. Statistical overrepresentation test of all categories (GO biological process complete, GO cellular complete, etc.) was performed using human orthologs and “*Homo sapiens* genes” as inputs. Results are in *SI Appendix, Figs. S10–S14*.

Identification of significantly higher expressed Forkhead transcription factors in ME cells. The Wilcoxon signed-rank test was performed on expression levels of each Forkhead transcription factor (given in “forkhead.csv”) between wild-type and ME cells to find those that were significantly more highly expressed in ME cells.

Data Availability. All study data are included in the article and/or *SI Appendix*. All code, analyses, and files can be accessed from the GitHub repository https://github.com/Singh-Lab/Pou4_Misexpression. In addition, the data discussed in this publication have been deposited in NCBI's Gene Expression Omnibus (Chacha et al., 2021) and are accessible through GEO Series accession number GSE192645 (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE192645>).

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1. L. Manni et al., Neurogenic and non-neurogenic placodes in ascidians. *J. Exp. Zool. B Mol. Dev. Evol.* **302**, 483–504 (2004).
2. F. Mazet et al., Molecular evidence from *Ciona intestinalis* for the evolutionary origin of vertebrate sensory placodes. *Dev. Biol.* **282**, 494–508 (2005).
3. C. Patthey, G. Schlosser, S. M. Shimeld, The evolutionary history of vertebrate cranial placodes—I: Cell type evolution. *Dev. Biol.* **389**, 82–97 (2014).
4. G. Schlosser, C. Patthey, S. M. Shimeld, The evolutionary history of vertebrate cranial placodes II. Evolution of ectodermal patterning. *Dev. Biol.* **389**, 98–119 (2014).
5. E. Wagner, A. Stolfi, Y. Gi Choi, M. Levine, Islet is a key determinant of ascidian palp morphogenesis. *Development* **141**, 3084–3092 (2014).
6. P. B. Abitua et al., The pre-vertebrate origins of neurogenic placodes. *Nature* **524**, 462–465 (2015).
7. A. Stolfi, K. Ryan, I. A. Meinertzhagen, L. Christiaen, Migratory neuronal progenitors arise from the neural plate borders in tunicates. *Nature* **527**, 371–374 (2015).
8. K. Waki, K. S. Imai, Y. Satou, Genetic pathways for differentiation of the peripheral nervous system in ascidians. *Nat. Commun.* **6**, 8719 (2015).
9. R. Horie et al., Shared evolutionary origin of vertebrate neural crest and cranial placodes. *Nature* **560**, 228–232 (2018).
10. B. Liu, Y. Satou, Foxg specifies sensory neurons in the anterior neural plate border of the ascidian embryo. *Nat. Commun.* **10**, 4911 (2019).
11. R. J. McEvilly et al., Requirement for Brn-3.0 in differentiation and survival of sensory and motor neurons. *Nature* **384**, 574–577 (1996).
12. S. Candiani et al., Ci-POU-IV expression identifies PNS neurons in embryos and larvae of the ascidian *Ciona intestinalis*. *Dev. Genes Evol.* **215**, 41–45 (2005).
13. W. J. Tang, J. S. Chen, R. W. Zeller, Transcriptional regulation of the peripheral nervous system in *Ciona intestinalis*. *Dev. Biol.* **378**, 183–193 (2013).
14. T. Horie, T. Kusakabe, M. Tsuda, Glutamatergic networks in the *Ciona intestinalis* larva. *J. Comp. Neurol.* **508**, 249–263 (2008).
15. A. Pasini et al., Formation of the ascidian epidermal sensory neurons: Insights into the origin of the chordate peripheral nervous system. *PLoS Biol.* **4**, e225 (2006).
16. K. Ryan, Z. Lu, I. A. Meinertzhagen, The peripheral nervous system of the ascidian tadpole larva: Types of neurons and their synaptic networks. *J. Comp. Neurol.* **526**, 583–608 (2018).
17. Y. Sasakura et al., Transposon-mediated insertional mutagenesis revealed the functions of animal cellulose synthase in the ascidian *Ciona intestinalis*. *Proc. Natl. Acad. Sci. U.S.A.* **102**, 15134–15139 (2005).
18. L. Gan, S. W. Wang, Z. Huang, W. H. Klein, POU domain factor Brn-3b is essential for retinal ganglion cell differentiation and survival but not for initial cell fate specification. *Dev. Biol.* **210**, 469–480 (1999).
19. D. M. Murakami, L. Erkmann, O. Hermanson, M. G. Rosenfeld, C. A. Fuller, Evidence for vestibular regulation of autonomic functions in a mouse genetic model. *Proc. Natl. Acad. Sci. U.S.A.* **99**, 17078–17082 (2002).
20. T. C. Badea et al., Combinatorial expression of Brn3 transcription factors in somatosensory neurons: Genetic and morphologic analysis. *J. Neurosci.* **32**, 995–1007 (2012).
21. T. Horie et al., Regulatory cocktail for dopaminergic neurons in a protovertebrate identified by whole-embryo single-cell transcriptomics. *Genes Dev.* **32**, 1297–1302 (2018).
22. C. Cao et al., Comprehensive single-cell transcriptome lineages of a proto-vertebrate. *Nature* **571**, 349–354 (2019).
23. L. A. Lemaire, C. Cao, P. H. Yoon, J. Long, M. Levine, The hypothalamus predates the origin of vertebrates. *Sci. Adv.* **7**, 7452 (2021).
24. K. Kim et al., Regulation of neurogenesis by FGF signaling and neurogenin in the invertebrate chordate *Ciona*. *Front. Cell Dev. Biol.* **8**, 477 (2020).
25. Y. Li et al., Conserved gene regulatory module specifies lateral neural borders across bilaterians. *Proc. Natl. Acad. Sci. U.S.A.* **114**, E6352–E6360 (2017).
26. S. M. Shimeld et al., Urochordate betagamma-crystallin and the evolutionary origin of the vertebrate eye lens. *Curr. Biol.* **15**, 1684–1689 (2005).
27. J. W. Blanchard et al., Selective conversion of fibroblasts into peripheral sensory neurons. *Nat. Neurosci.* **18**, 25–35 (2015).
28. N. Sharma et al., The emergence of transcriptional identity in somatosensory neurons. *Nature* **577**, 392–398 (2020).

29. T. Osugi, Y. Sasakura, H. Satake, The nervous system of the adult ascidian *Ciona intestinalis* type A (*Ciona robusta*): Insights from transgenic animal models. *PLoS One* **12**, e0180227 (2017).
30. T. Osugi, Y. Sasakura, H. Satake, The ventral peptidergic system of the adult ascidian *Ciona robusta* (*Ciona intestinalis* type A) insights from a transgenic animal model. *Sci. Rep.* **10**, 1892 (2020).
31. J. S. Joly et al., Culture of *Ciona intestinalis* in closed systems. *Dev. Dyn.* **236**, 1832–1840 (2007).
32. T. Coric, Y. J. Passamaneck, P. Zhang, A. Di Gregorio, C. M. Canessa, Simple chordates exhibit a proton-independent function of acid-sensing ion channels. *FASEB J.* **22**, 1914–1923 (2008).
33. A. T. L. Lun et al., Participants in the 1st Human Cell Atlas Jamboree, EmptyDrops: Distinguishing cells from empty droplets in droplet-based single-cell RNA sequencing data. *Genome Biol.* **20**, 63 (2019).
34. Y. Satou et al., Improved genome assembly and evidence-based global gene model set for the chordate *Ciona intestinalis*: New insight into intron and operon populations. *Genome Biol.* **9**, R152 (2008).
35. M. Brozovic et al., ANISEED 2015: A digital framework for the comparative developmental biology of ascidians. *Nucleic Acids Res.* **44**, D808–D818 (2016).
36. M. D. Robinson, D. J. McCarthy, G. K. Smyth, edgeR: A Bioconductor package for differential expression analysis of digital gene expression data. *Bioinformatics* **26**, 139–140 (2010).
37. T. L. Bailey, C. Elkan, Fitting a mixture model by expectation maximization to discover motifs in biopolymers. *Proc. Int. Conf. Intell. Syst. Mol. Biol.* **2**, 28–36 (1994).
38. S. Gupta, J. A. Stamatoyannopoulos, T. L. Bailey, W. S. Noble, Quantifying similarity between motifs. *Genome Biol.* **8**, R24 (2007).
39. H. Mi et al., Protocol update for large-scale genome and gene function analysis with the PANTHER classification system (v.14.0). *Nat. Protoc.* **14**, 703–721 (2019).