Whole-Body Single-Cell Sequencing Reveals Transcriptional Domains in the Annelid Larval Body

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Abstract

Animal bodies comprise diverse arrays of cells. To characterize cellular identities across an entire body, we have compared the transcriptomes of single cells randomly picked from dissociated whole larvae of the marine annelid Platynereis dumerilii. We identify five transcriptionally distinct groups of differentiated cells, each expressing a unique set of transcription factors and effector genes that implement cellular phenotypes. Spatial mapping of cells into a cellular expression atlas, and wholemount in situ hybridization of group-specific genes reveals spatially coherent transcriptional domains in the larval body, comprising, for example, apical sensory-neurosecretory cells versus neural/epidermal surface cells. These domains represent new, basic subdivisions of the annelid body based entirely on differential gene expression, and are composed of multiple, transcriptionally similar cell types. They do not represent clonal domains, as revealed by developmental lineage analysis. We propose that the transcriptional domains that subdivide the annelid larval body represent families of related cell types that have arisen by evolutionary diversification. Their possible evolutionary conservation makes them a promising tool for evo–devo research.

Key words: evo–devo, single-cell transcriptomics, expression atlas, transcriptional domain, cell type family, Platynereis dumerilii.

Introduction

The organization of cells into tissues and organs represents a fundamental feature of animal bodies. Recently, a number of studies have applied single-cell transcriptomics to assess cellular diversity within tissues, such as the vertebrate pancreas (Muraro et al. 2016), intestine (Grun et al. 2015), and different parts of the brain, including cortex, basal ganglia, and hypothalamus (Zeisel et al. 2015; Tasic et al. 2016; Romanov et al. 2017). This approach allows the molecular characterization of cell types within a given tissue, as well as an assessment of their heterogeneity. However, the sheer number of cells in conventional vertebrate or insect model systems has so far limited the comparison of cellular identities to single tissues or organs, and hampered comparisons across entire bodies.

Other laboratories have focused on the molecular comparison of different tissues, referred to as tissue transcriptomics. This approach determines and compares the expression profiles of entire tissues via bulk RNAseq (Henrichsen et al. 2009; Ramskold et al. 2009; Brawand et al. 2011; Breschi et al. 2016), with the potential to compare across the entire body. However, this approach requires defining tissues a priori by morphology or marker gene expression. This is problematic because a given tissue may be composed of very different cell types, and similar cell types can be found in distinct tissues as is the case for vertebrate ciliary photoreceptors, which populate both the retina and pineal organ (Arendt 2008). Therefore, this approach lacks the resolution necessary for understanding how cell types are distributed across entire organisms, and how this relates to morphologically defined tissues.

To advance on this, we have established the marine annelid Platynereis dumerilii, a molecular model species for development, evolution, and neurobiology (Fischer and Dorresteijn 2004; Williams and Jekely 2016), as an experimental paradigm to explore how cellular identities compare and relate to each other across an entire animal body. The Platynereis larval body is ideally suited for investigating the transcriptional organization of an entire organism as it harbors a variety of morphologically distinct cell types, but has a relatively low number of cells...
Identification of robust cell groups within the data. The clustering approach enables identification of genes that characterize each cell type. Finally, we used hierarchical clustering to investigate the similarity between the identified cell clusters.

(Fischer et al. 2010), making it amenable for applying single-cell RNAseq to the whole organism. Further, Platynereis develops highly stereotypically, which allows for the construction of a cellular atlas onto which single-cell transcriptomes can be spatially mapped (Tomer et al. 2010; Asadulina et al. 2012; Vergara et al. 2016).

Here, we apply single-cell RNAseq to randomly sampled cells from the dissociated whole larvae at 48-h postfertilization (hpf). Our whole-body analysis reveals that, at this stage, the larval annelid body comprises five well-defined groups of differentiated cells with distinctive expression profiles. Cells in each group share expression of a unique set of transcription factors together with effector genes encoding group-specific cellular structures and functions. To correlate these groups with larval morphology, we establish a gene expression atlas for 48 hpf Platynereis larvae using the recent “Profiling by Signal Probability mapping” (ProSPr) pipeline (Vergara et al. 2016). For each group, we then locate individual cells in this atlas using an established algorithm for spatial mapping of single cells (Achim et al. 2015). The spatial distribution of each group was further validated by conducting wholomount in situ hybridization of selected group-specific genes. We thus reveal that the five distinct groups of differentiated cells spatially subdivide the larval body into coherent and nonoverlapping transcriptional domains that comprise (1) sensory-neurosecretory cells located around the apical tip of the larva, (2) peptidergic prospective midgut cells, (3) somatic myocytes, (4) cells with motile cilia constituting the larval ciliary bands, and (5) larval surface cells with epidermal and neural characteristics. We also show that these domains do not reflect developmental lineage, as they unite cells of distinct clonal origin.

To characterize the remaining six groups further, we identified differentially expressed genes (see Materials and Methods). The largest group of cells, which resulted from combining the two closely related groups, was characterized by the specific expression of genes known to be active in developmental precursors, such as DNA replication (DNA topoisomerase, Rfc3, Rfc5, Mcm3, Mcm4, Mcm7), cell proliferation (Pena, Md21I, Apc7, supplementary fig. 2i, Supplementary Material online), or chromatin remodeling (Nucleoplasmin, Bptf), suggesting this group was primarily composed of undifferentiated cells. Cells in the other five groups showed significantly lower expression of these markers.
Mapping Sequenced Cells to a Cellular Expression Atlas

As a prerequisite for localizing the five distinct groups of differentiated cells in the larval body, we built a gene expression atlas with cellular resolution for the 48 hpf larva, using our recently established Profiling by Signal Probability mapping (ProSPr) pipeline (Vergara et al. 2016). This involved whole-mount in situ hybridization for 176 genes, automated acquisition of up to 40 expression images per gene, and registration of these images to a reference larva to construct an expression average for each gene (fig. 2 and supplementary fig. 4, Supplementary Material online).

Taking advantage of this atlas, we used a dual strategy to investigate the spatial distribution of sequenced cells. First, we located individual cells in the atlas via the comparison of complete, specificity-weighted mRNA profiles for each cell with the positional gene expression vectors derived from the atlas (Achim et al. 2015). Following this strategy, the vast majority (95%) of cells could be mapped to distinct positions within the larva (fig. 3; left panels show the single most likely position for each cell in each group of differentiated cells, right panels show the highest scoring mapping positions for one selected cell in each group). Notably, cells belonging to the same group showed spatially correlated mapping to distinct body regions, as apparent from the overlap with the expression of one marker gene specific for each group. Remarkably, looking at the highest scoring mapping positions for an individual cell, we consistently detected pairs of bilateral coordinates, reflecting the bilateral symmetry of the animal. We also noted different degrees of spatial acuity: While some cells mapped to unique (or closely adjacent) left/right pairs of coordinates (fig. 3b and d, right panels), other cells mapped to segmentally iterated positions, reflecting the segmental nature of the larval trunk with iterated occurrence of cell types (fig. 3f and h, and j, right panels).

Transcriptional Domains Subdivide the Larval Body

Complementing the spatial mapping of individual cells, we also determined the overall spatial distribution for each group of differentiated cells in the larval body. To this end, we identified transcripts that showed maximum specificity for each group (fig. 4, left panels) and determined their expression pattern in the larval body using wholemount in situ hybridization (WMISH) (fig. 4, right panels). We reasoned that in combination, the expression patterns of the genes most specific for each group should approximate the overall spatial distribution of cells belonging to that group in the larval body. If not yet included, we fed group-specific genes into the expression atlas to accurately determine expression overlap (and to refine the spatial mapping of individual cells).

The two approaches, that is, the mapping of individual cells (see above) and the mapping of the entire group via
In particular, group 1 cells mapped to the larval brain around the apical tip of the larva, where Prohormone Convertase 2 (Phc2) is expressed, a marker for this domain (fig. 3a and b). Consistent with this, expression of another domain-specific marker gene, Phosphodiesterase-9 (Pde9), was restricted to the most apical territory of the larva (fig. 4a). Group 2 cells mapped to the midgut primordium alongside the yolk-rich macromeres, as confirmed by the spatially highly restricted expression of Hnf4 (fig. 3c and d and fig. 4b). Cells of group 3 mapped to the differentiating striated myocytes that consistently expressed the Striated muscle myosin heavy chain (St-mhc) gene (fig. 3e and f and fig. 4c); and group 4 cells populated the ciliary bands, matching the expression of Foxj (fig. 3g and h) and of Radial spoke head protein homolog 4 (Rsph4) (fig. 4d) (Fischer et al. 2010). Finally, group 5 cells covered large part of the larval surface, reflected by Uncx4 expression (fig. 3i and j) and by the broad surface expression of the Metabotropic glutamate receptor 7 (Grm7) in the larval episphere and hyposphere (fig. 4e).

Although the transcriptional domains for groups 2, 3, and 4 matched known larval tissues (early midgut, striated myocytes, and ciliary bands), no obvious morphological features or boundaries corresponded to, or delimited groups 1 and 5 transcriptional domains. This is remarkable because the topology of the hierarchical tree (fig. 4 and supplementary fig. 3, Supplementary Material online) supported a deep transcriptional divide between the two groups. Transcriptionally, group 5 cells were more similar to the ciliary bands (group 4) and to striated myocytes (group 3), than to group 1 cells, with high bootstrap support.

To locate the boundary between transcriptional domains 1 and 5 more precisely in the larval body, we investigated the expression of group-specific marker genes taking advantage of our expression atlas. We thus determined that the group 1 transcriptional domain comprised a cruciform patch of differentiated apical cells, deeply embedded in the anterior neuroectoderm and covered by surface cells (fig. 5). In contrast, the expression of group 5-specific genes consistently remained restricted to the larval surface. Coexpression analysis revealed virtually no expression overlap between genes specific for the one or the other group, corroborating their strict separation as distinct transcriptional domains.

### The Group 1 Transcriptional Domain Represents the Apical Nervous System

To explore the molecular nature of the groups/transcriptional domains, we investigated their distinctive gene sets. Figure 6 shows the most specific and informative regulatory and effector genes that were found to be specific for each group. Illustrating the power of this approach, the genes specific for groups 2, 3, and 4 cells clearly identify these as differentiated midgut cells, striated myocytes, and the multiciliated cells of the ciliary bands, respectively (fig. 6b, c, and d). Further inspection of the group 1-specific genes (fig. 6a) revealed that many of these encoded neuropeptides, such as the broadly expressed GNXQNpeptide or GGNamide, two lophotrochozoan neuropeptides (Conzelmann et al. 2013), and the neuropeptide-cleaving Prohormone convertase 2 (Phc2)
(fig. 5), indicating neurosecretory release from these cells (Tessmar-Raible et al. 2007). In addition, we found noncalcium-binding members of the synaptotagmin family implicated in the generation and fusion of large dense core vesicles for neurosecretion, such as Synaptotagmin17/B/K (Syt17), Sytz, and Syt4 (figs. 6a and 7) (Gustavsson et al. 2012; Moghadam and Jackson 2013; Park et al. 2014). This is in line with electron microscopy data, which show that cells in the dorsomedian Platynereis brain are rich in dense core vesicles while synapses are often sparse or absent (Williams et al. 2017). Group 1 cells also specifically expressed genes associated with photoreception, such as the bistable photopigment Peropsin (Nagata et al. 2010) and components of cGMP-based signal transduction such as Pde9 and the CNG channel components Cngb and Cnga-α (fig. 6a). Individual cells expressed ciliary photoreceptor-specific genes and
were thus identified as the brain ciliary photoreceptors (Neuropeptide Y [Npy] in combination with c-opsin or c-opsin2; Arendt et al. 2004), or as photosensory-vasotocinergic cells (expressing Vasotocin–neurophysin, Proenkephalin, and c-opsin; Tessmar-Raible et al. 2007).

Interestingly, we also found that neuroglobin was specific for this domain, encoding a monomeric globin evolutionarily older than hemoglobin or myoglobin known to reversibly bind oxygen (Burmester et al. 2000). Taken together, the nature of group 1-specific genes is strong evidence that this domain represents the apical nervous system (ANS) as characterized previously (Tosches and Arendt 2013; Tosches et al. 2014; Williams et al. 2017).

**Group 5 Transcriptional Domain Cells Show Epidermal and Neural Characteristics**

The ectoderm-derived Group 5 cells expressed genes are indicative of mixed epithelial and neural characteristics (fig. 6e). Expression of some of these genes is shared with both multiciliated cells and striated myocytes, consistent with the close relationship we found between these groups in our hierarchical clustering analysis (compare fig. 4). Among the most broadly expressed genes shared with multiciliated cells and myocytes were LaminA/C, a fibrous protein of the nuclear lamina (Kim et al. 2012) (supplementary fig. 5, Supplementary Material online), and the related Neurofilament-70 (NF70), a neural intermediate filament found in Lophotrochozoans; Extended Synaptotagmin 2 (Esyt2) has a reported function in membrane lipid composition dynamics, extracellular signal transduction, and cell adhesion (Herdman and Moss 2016); and P4ha2 encodes a component of prolyl 4-hydroxylase, a key enzyme in collagen synthesis. The genes specifically shared between group 5 and the ciliary bands included Anoctamin, a transepithelial chloride channel (Pedeomonte and Galietta 2014), Protocadherin-15 (Pcdh15), an atypical cadherin mediating structural integrity of ciliated sensory receptors (Seiler et al. 2005) (supplementary fig. 5, Supplementary Material online), and Semaphorin 2 (Sema2), a secreted extracellular guidance molecule involved in axon pathfinding (Bates and Whitington 2007). In line with an overall epithelial identity of these cells, they expressed the epidermis-specific transcription factor Grainyhead, known for its conserved role in maintaining the epidermal barrier (Boglev et al. 2011), and the ETS-homologous factor that likewise specifies epithelial cell type identities (Feldman et al. 2003).

Most noticeably, however, we detected a set of genes representing an extracellular matrix-remodeling module (fig. 8). These genes encoded matrix-modifying proteases such as a serine protease Plasminogen related to vertebrate
FIG. 6. Tissue specific marker genes reflect cellular functions. For each of the differentiated tissues, we show a heatmap of tissue specific gene expression: (a) expression profile of group 1 apical nervous system-specific genes; (b) expression profile of group 2 midgut genes; (c) expression profile of group 3 myocyte-specific genes; (d) expression profile of group 4 multiciliated cell-specific genes; and (e) expression profile of group 5 neural/epithelial surface-specific genes. Functionally related groups of genes are highlighted. TF: transcription factor, td: transcriotional domain.
Plasmin (Sonderegger and Matsumoto-Miyai 2014) and Hepatocyte Growth Factor (HGF) (Tydall and Walikonis 2006) (supplementary table 2, Supplementary Material online); metalloproteinases such as Matrix Metalloproteinase (Mmp9) (Michaluk et al. 2011) together with its specific substrate Cd109 (Vadon-Le Goff et al. 2015); and the cysteine proteinase Cathepsin L (Felbor et al. 2002) (fig. 8b). Many of these protease systems have been implicated in the modification of dendritic spines (Michaluk et al. 2011) and synapses (Sonderegger and Matsumoto-Miyai 2014), suggesting nervous system-related functions of this module.

In line with this, the genes most specific for group 5 (not shared with the ciliary bands or myocytes) abundantly encoded proteins with reported roles in neurite outgrowth and synaptic plasticity. For example, we identified Neurotactin, which encodes a cell surface glycoprotein of the serine esterase superfamily (Speicher et al. 1998), and its specific binding partner Amalgam (Zeev-Ben-Mordehai et al. 2009); both involved in axonal pathfinding in Drosophila (Speicher et al. 1998; Fremion et al. 2000). We also found two orthologs of Neurotrypsin (fig. 8a), a matrix-modifying serine protease, together with the extracellular proteoglycan Agrin (Stephen et al. 2008), indicating that the Neurotrypsin-Agrin system is active in these cells (Matsumoto-Miyai et al. 2009) (fig. 6e). The Hspg2 gene, encoding a conserved proteoglycan related to Agrin, was also part of this set. Furthermore, the specific expression of the nicotinic acetylcholine receptor alpha-9 is noteworthy. In vertebrates, this unconventional receptor plays restricted roles in the mechanosensory hair cells, where it is involved in the reception of efferent signals (Murthy et al. 2009), and in epidermal keratinocytes, where it triggers epithelialization via local, nonneuronal acetylcholine cytotransmission (Chernyavsky et al. 2007). Finally, smaller subsets of group 5 cells also expressed Wnt ligands such as Wnt4 (Pruitt et al. 2014), and the bHLH HES transcription factors Hes1, Hes2, Hes6, and Hes11; and homeodomain transcription factors with conserved roles in neuronal specification such as Uncx4, Tbx20, or Phox2 (Denes et al. 2007; Rabe et al. 2012; Nomaksteinsky et al. 2013) (fig. 6e).

In conclusion, a closer inspection of group 5-specific genes suggests both epithelial and neural properties. However, since none of the group 5 cells showed a clear and distinct neuronal profile as observed in the apical nervous system (supplementary fig. 6, Supplementary Material online), the possible classification of (subsets of) group 5 cells as neuronal remains preliminary. We therefore used our ProSPr 48hpf cellular expression atlas to test for co-expression of pan-neuronal markers with group/domain-specific genes. Supporting neural characteristics of group 1 (apical nervous system, fig. 9a) and of at least some group 5 cells (figure 9b), our cellular expression atlas revealed extensive coexpression of group 1 markers (Vat1l, Syt4; compare fig. 6a) and restricted but clear coexpression of group 5 markers (Tbx20, uncx4, LaminAC; compare fig. 6e) with the pan-neuronal marker Rab3 (fig. 9).

Incongruence of Transcriptional Domains and Developmental Lineage

Finally, we set out to explore a possible correlation between the five transcriptional domains of the larval body and the Platynereis developmental lineage (Ackermann et al. 2005; Vopalensky et al. 2018), to find out whether there was congruence between transcriptional and clonal domains. We reasoned that common origin from a shared developmental precursor would be one possible explanation for the observed transcriptional similarity between cells (Scialdone et al. 2016; Kee et al. 2017). Facilitating this analysis, the lineage of the Platynereis episphere has recently been solved up to 30 hpf by in vivo imaging, and individual cells can be identified in any embryo up to 15 hpf due to the highly stereotypical embryonic development (Vopalensky et al. 2018). Building on these resources, we investigated the clonal origin of all Phc2+ cells of the apical nervous system (fig. 10). We found them to derive from multiple different sublineages of the Platynereis larval episphere, thus representing a multiclonal assembly. Likewise, the group 5 neural-epithelial surface cells and the motile ciliated cells are developmentally heterogeneous, as they both comprise episphere and hyposphere cells that derive from distinct parts of the Platynereis lineage (Ackermann et al. 2005). The cellular lineage of the midgut and somatic myocytes is yet unknown. We can thus conclude that the groups and transcriptional domains for which cellular lineage information is available are polyclonal; and that developmental lineage cannot account for their expression similarity.

To further explore the developmental fate of the transcriptional domains, we performed whole-mount in situ hybridization of apical nervous system and neural/epithelial surface marker genes for stages earlier than 48 hpf and for later lifecycle stages (supplementary fig. 7, Supplementary Material online). In the absence of whole-body single-cell data for these stages, these preliminary data suggest that the transcriptional domains remain stable over time and may thus represent an inherent and constant property of the animal body.

Discussion

A New Molecular Subdivision of the Platynereis Larval Body

For this study, we analyzed the transcriptomes of cells randomly collected from dissociated, entire larvae of the marine annelid Platynereis dumerilii. Previous single-cell studies had captured cells from specific body parts and tissues only (such as the mouse forebrain; Zeisel et al. 2015; Tasic et al. 2016). Our data are unique as they allow single-cell level comparison of differentiated cells across the entire body. We were able to sort all differentiated cells into five major groups, based on similarities and differences in gene expression profiles. Spatial mapping of sequenced cells into the 48 hpf ProSPr cellular-resolution expression atlas, in combination with wholemount in situ hybridization of group-specific marker genes, revealed five coherent transcriptional domains in the 48 hpf Platynereis body. In contrast to an expression domain, which delimits expression in the body for a single gene, a transcriptional domain delimits the expression of many genes to a specific
part of the body. For the first time, the discovery of transcriptional domains opens up the possibility to understand animal organization that does not rely on predefined morphological boundaries or a priori gene selection, but instead reflects the molecular boundaries that correspond to the most pronounced differences in overall gene expression.

We probed the molecular nature of these transcriptional domains and found they were united by the shared expression of specific sets of transcription factors with known roles in cell fate selection, such as *Hnf4* for midgut (Palanker et al. 2009), or *Grainyhead* for epithelial cells (Boglev et al. 2011). Although we did not test the function of these transcription factors in this study, it is plausible to hypothesize that they play a role in regulating domain-specific effector genes. In line with this, we found each domain shared a coherent set of effector genes representing distinct cellular modules, as exemplified for the apical nervous system (fig. 7) and the neural/epithelial surface cells (fig. 8). Beyond that, however, we also observed considerable differences in transcription factor and effector gene expression within a given group (as evident from fig. 6), suggesting the presence of several related but distinct cell types. For example, cells of the apical nervous system differ with regard to opsin expression, neuropeptide content, and signaling components—indicating that each, or almost each, cell might represent a separate type. These
differences are unlikely to reflect technical noise, as we find known cell type markers coexpressed within individual cells. It is important to stress that, although our sample size is large enough to resolve transcriptional domains, it is too small for the full characterization of individual cell types, which in some cases may be represented by only a single, bilateral pair of cells (Vergara et al. 2016). It is thus plausible that the number of sequenced cells would have to be increased substantially to fully resolve the individual cell types within each given domain.

Despite our inability to resolve individual cell types, the comparison of these transcriptional domains to larval morphology and developmental lineage revealed several notable results. First, whereas some domains clearly delineate distinct morphological tissues, such as midgut, striated muscles, and ciliary bands, we also uncovered a fundamental divide between apical nervous system and neural/epithelial surface ectoderm, which was emphasized by the topology of the tree that placed both far apart from each other. This divide was unexpected because the boundary between both domains does not coincide with a clear morphological boundary, and both domains coexpress pan-neuronal markers (fig. 9). Second, we found that the transcriptional domains were largely incongruent with clonal domains (fig. 10). This is clear evidence that their transcriptional similarity is not due to shared developmental lineage, and accords with recent findings for the vertebrate retina (Macosko et al. 2015), which found differentiated rods and cones more closely related to each other transcriptionally than to other retinal cell types, even though they come from different developmental progenitors (Cepko 2015). If not developmental lineage, what then accounts for the similarity of cells and cell types found within the transcriptional domains?

The Concept of Cell Type Families

One plausible explanation for the observed similarities in expression profile between cells belonging to the same group/domain is evolutionary relatedness; or, in other words, that their constituting cell types derive from shared ancestral cell types. We have recently proposed that the diversity of cell types in animals evolved via a sequence of cellular diversification events, producing pairs of sister cell types in diverging lineages. With continued diversification, ancient cell types thus give rise to cell type families. Reflecting their common ancestry, related cell types would share the expression of some transcription factors involved in their combinatorial specification and of (subsets of) cellular effector genes implementing their similar phenotypes (Hobert 2011; Arendt et al. 2016).

The somatic myocytes (fig. 6c) exemplify the concept of cell type families. Although the transcription factor Mef2 is expressed in all cells of the group, the expression of other factors is likewise robust but restricted to few cells in the groups, in- including MyoD, Hand, Max, Paraxis, and Twist, consistent with more specific roles as myoblast identity genes as reported, for example, for Drosophila (Dobi et al. 2015). This would suggest that the Platynereis myocyte family contains several distinct myocyte types representing sister cell types (Brunet et al. 2016).

Evolutionary Conservation of the Apical Nervous System

Our hypothesis that the transcriptional domains represent cell type families can be tested via comparison to other
species. Given that the Platynereis larval body is composed of five transcriptional domains; and assuming that the cell types of each domain indeed diversified from single founder cell types in animal evolution, then these five founder cell types must have existed early in animal evolution. This would make the transcriptional domains that we have identified in Platynereis very ancient and imply that they should also exist in other remote animal phyla. Formal testing of this hypothesis will require sampling additional species for whole-body single-cell profiling in diverse phyla. However, several initial
lines of evidence already support our hypothesis. In particular, it has previously been proposed that the annelid apical nervous system and the vertebrate hypothalamus are homologous. This was based on the conserved developmental expression of transcription factors such as siX3 and foxq (Marlow et al. 2014), on the shared presence of a neurosecretory center releasing conserved neuropeptides such as oxytocin-vasopressin (Tessmar-Raible et al. 2007; Williams et al. 2017), on conserved nonvisual ciliary photoreceptors employing ciliary opsin and a vertebrate-type phototransduction cascade (Arendt et al. 2004), and on the shared presence of an active center of melatonin synthesis in control of circadian behavior (Tosches et al. 2014). These similarities have been taken as evidence to propose evolutionary conservation of the apical nervous system as a distinct part of the bilaterian brain (Tosches and Arendt 2013), which may trace back to early metazoan ancestors (Marlow et al. 2014).

Our new data allow us to further challenge and validate this hypothesis, via the comparison of new apical nervous system domain-specific markers identified in Platynereis with expression data available for the vertebrates. And indeed, this comparison is highly supportive of evolutionary conservatism. For example, in vertebrates the noncalcium-binding and neurosecretion-specific Syt4 is specifically active in the neuroendocrine hypothalamus, where it is known to regulate the release of the neuropeptide Oxytocin (Zhang et al. 2011). In addition, vertebrate Syt17 has initially been isolated from the vasopressin-secreting supraoptic and paraventricular nuclei and likewise shows strong expression in the hypothalamus (Lee et al. 2001), as does Neuroglobin (Hundahl et al. 2008, 2012; Fabrizius et al. 2016). Finally, the T-box transcription factor T-brain is specifically expressed in the developing para- and periventricular and supraoptic hypothalamic nuclei (Dominguez et al. 2015). These observations suggest that transcriptional domains and the corresponding cell type families are conserved in animal evolution and that their comparative study represents a highly promising new direction in evolutionary research.

Materials and Methods

Cell Dissociation of Platynereis Whole Larvae

Platynereis dumerilii larvae were incubated in filtered natural seawater (FNSW), at 18°C for 47 h to reach the desired stage. For single-cell dissociation, larvae were concentrated using a 40 μm size mesh, picked, washed in Ca, Mg-free artificial seawater (CMF-SW), and incubated for 10 min in 0.5% Pronase (Roche cat # 10165921001), 1% sodium thioglycolate (Sigma T0632) in CMF-SW, at room temperature. Larvae were then rinsed in 1× PBS, and incubated in 500 μl of 150 μg/ml Liberase (Roche, cat # 05401119001) in 1× PBS, for 1 min. Larvae were gently triturated by pipetting, centrifuged at 200×g for 3 min, and reconstituted in 600 μl of 1× PBS. Cells were then dissociated by trituration using a syringe. Dissociated cells were washed twice in 200 μl of 1× PBS and concentrated by centrifuging (1 min, 200×g). Cells were resuspended in 20 μl of 1× PBS, of which 12 μl was loaded on the Fluidigm C1 capture chip.

Cell Capture and Sequencing

Platynereis larval cells were captured, and single-cell cDNA synthesis was performed using the Fluidigm C1 Single-Cell Auto Prep System, and the sequencing libraries were prepared as previously described (Achim et al. 2015). We used Fluidigm IFCs for mRNA-seq with capture sites for 5–10 μm and 10–17 μm to collect the cells, allowing the capture of a full range of cell sizes in 48hf stage of Platynereis. In total, nine IFC-s, each representing an independent collection of cells derived from a different batch of animals, were processed for this study. The information about the samples distribution on chips, the chip sizes, cell number per sample, and the pooling of samples on sequencing lanes is described in supplementary table 3, Supplementary Material online. 100-bp paired-end sequences were generated using the Illumina HiSeq2000 platform. For one library (CN62), 75-bp paired-end sequences were generated using the Illumina NextSeq500 platform. Libraries were sequenced to an average depth of 2.1 million reads.

Data Availability

The raw sequence data for the cells analyzed in this study are available from ArrayExpress, accession numbers E-MTAB-2865 (see supplementary table 5, Supplementary Material online, for matching the cell ID-s between the E-MTAB-2865 and this study) and E-MTAB-5953. The R code used for analysis is available at: https://github.com/MarioniLab/Platynereis2017.

Read Alignment

FastQC (Andrews 2010) was used for the quality trimming of the raw sequencing reads using default settings. For read mapping, the quality-filtered sequencing reads were mapped, using bowtie2, against a Platynereis reference transcriptome combined from the two previously published assemblies (Conzelmann et al. 2013; Achim et al. 2015). Briefly, the transcriptome assemblies were concatenated and contigs showing >94% identities were removed using CD-HIT (Li and Godzik 2006), leaving 44977 transcripts. In the generated reference, each gene should be represented by one transcript, reducing the problem of multiple mapping of the reads. Expression counts for each transcript were obtained using HTSeq (Anders et al. 2015) while only considering uniquely mapped reads.

Gene Annotation

To annotate genes of the two transcriptome assemblies (Conzelmann et al. 2013; Achim et al. 2015), reciprocal BLAST comparison of genes sequences against the Uniref90 protein database (Arendt assembly) or against Swissprot (Jekely assembly) was performed. For each transcript, the BLAST hit with highest E-value was selected for annotation.

Quality Control

Low-quality cells were removed from the data set based on the following filtering criteria. Visual inspection of capture sites on the IFC revealed empty wells, wells with multiple cells
or debris captured. Only wells with positive cell capture were further processed for sequencing. For downstream analysis, libraries marked as single cells were chosen. Cells with <1,000 reads mapping to ERCC spike-ins, <100,000 mapped transcriptomic reads or <60% of mapped reads being allocated to the transcriptome were removed. Additionally, cells that express unusually small number of genes (<2000) were removed. Prior to normalization, genes with >1,000,000 reads or <10 reads across all cells were removed. A tsne plot showing batch labels for each cell is presented in supplementary figure 1l, Supplementary Material online.

Normalization
The BASiCS package (Vallejos et al. 2015) was used to normalize read counts by incorporating ERCC spike-ins for technical noise estimation. Specific ERCC spike-ins were removed if not detected in the data set. Posterior estimates for model parameters were computed by Markov chain Monte Carlo (MCMC) simulation with 40,000 iterations. The BASiCS_DenoisedCounts function was utilized to compute normalized read counts.

Running the MCMC independently on the different batches confirmed similar small levels of technical noise in the data (supplementary fig. 1h, Supplementary Material online).

Spatial Mapping
Spatial mapping of the sequenced single-cell transcriptomes onto the Platynereis reference atlas was performed as described previously (Achim et al. 2015). For the reference atlas construction, we performed whole-mount in situ hybridization for 176 genes in 48 hpf larvae and produced up to 53 expression images per gene via automated microscopy. We then registered these images to a reference larva and used ProSPR to reconstruct an average expression pattern for each gene that achieves cellular resolution (Vergara et al. 2016). Plotting patterns for all genes onto the same larval average resulted in a cellular resolution atlas (supplementary fig. 4, Supplementary Material online and fig. 3), to which we then correlated the expression vectors of sequenced cells (Achim et al. 2015). The Platynereis reference atlas is provided as supplementary table 4, Supplementary Material online, and the spatial mapping results for all the sequenced cells are provided in supplementary table 5, Supplementary Material online. We defined functional regions within the embryo based on spatial expression patterns of known marker genes. The region in which Phc2 is expressed comprises the apical nervous system. The muscle region of the animal is defined by the expression of St-mhc. Foxj expression defines the ciliated cells and Hnf4 expression the midgut region. The trunk ectoderm can be described by Uncx4 expression. To assign mapped cells to particular regions, we first performed graphical clustering on the voxels for each cell. Second, we identified the voxel cluster with highest mapping confidence for each cell (right hand side of fig. 3). For visualizing the mapping of individual cells, the centroids of all voxel clusters were plotted (left hand side of fig. 3). To compute the overlap of mapped cells to spatial expression patterns, we focused on the cluster with highest mapping confidence for each cell. If more than 50% of these voxels also show a particular gene expression (e.g., Syt1), we consider the cell as falling into this gene region.

Detecting Highly Variable Genes
The BASiCS_DetectHVG function in BASiCS allows the detection of highly variable genes by incorporating spike-ins to estimate expected technical variability. Testing was done after MCMC simulation with a variance threshold of 0.98 and an evidence threshold of 0.7.

Clustering
To detect clusters of cells based on the expression of highly variable genes, we used a sparse K-means clustering approach taken from the sparcl package in R (Witten and Tibshirani 2010; Team 2014). We clustered the data using $K = 2, \ldots, 10$ and chose $K = 7$ since only small subpopulations emerged at $K = 8, \ldots, 10$ (supplementary fig. 1j, Supplementary Material online). An elbow plot showing the averaged within-cluster sum of squares is noninformative for the choice of $K$ (supplementary fig. 1i, Supplementary Material online). AWSS = $\frac{1}{C_1 \cdots C_K} \sum_{i=1}^C \sum_{j=1}^N (x_{ij} - \mu_j)^2 / m$, where $x_{ij}$ is the log$_{10}$-transformed transcript count of gene $i$ in cell $j$ and cluster $k$, $\mu_j$ the log$_{10}$-transformed mean expression of gene $i$ in cluster $k$ and $m$ the number of clusters. The parameters were tuned with $2 < \text{wbounds} < 100$ using 15 permutations. Genes with weights $> 0.03$ define the cluster-characteristic features of the data. We compared this clustering strategy to one that does not require an a priori definition of $K$. The $\text{dynamicTreeCut}$ package in R (Langfelder et al. 2008) performs a flexible branch cutting for hierarchical clustering approaches. The $\text{mclust}$ package in R (Fraley et al. 2012) performs clustering based on finite mixture modeling of Normals. Comparisons of the cell labels for each K using the sparse clustering approach and the labels detected by $\text{dynamicTreeCut}$ and $\text{mclust}$ shows a good overlap at $K = 7$ (supplementary fig. 1K, Supplementary Material online). The following parameters were used for $\text{dynamicTreeCut}$ clustering: minClusterSize = 5, deepSplit = 0 and the dendrogram was build using the spearman dissimilarity (see Tree Building) and using ward.D2 as linkage method. The parameters for $\text{mclust}$ were chosen by calculating the BIC with the default function $\text{mclustBIC}$. We used the rand index calculated by the $\text{cluster_similarity}$ function of the $\text{cluster}$ package in R to compare the labels.

Removing Cell Doublets
Possible cell doublets were removed after clustering and marker gene detection. The percentage of group specific marker gene expression (# marker genes expressed/# all genes) was calculated for each cell. Cells with unusually high expression of markers nonspecific for their group were removed from the data set.
Tree Building
To investigate the transcriptional similarity between cell types, we hierarchically clustered cell types before and after averaging gene expression across all cells within each group. The averaging strategy increases stability of the hierarchical clustering algorithm by averaging out dropout events in single-cell data. Distances were calculated based on the Spearman dissimilarity: $\sqrt{\frac{1}{2}(1-r_s)}$ (Van Dongen and Enright 2012). To compute bootstrapped P values, we used the pvclust package in R (Ryota Suzuki 2006). Hierarchical clustering was performed on the log_{10}-transformed expression counts before and after averaging within each group using the spearman dissimilarity as distance, complete linkage clustering, and 1000 bootstrap iterations to evaluate cluster stability.

Differential Expression
To identify cell group specific enriched marker genes, we used two approaches. First, we used the sce package (Kharchenko et al. 2014) to compare each of the identified cell group against all the remaining samples in order to identify novel group specific genes from the unannotated portion of the Platynereis transcriptome. These genes were then annotated, added to the reference, and the clustering was repeated. Second, differential expression analysis was performed using the findMarkers function from the scran package. This approach uses limma (Ritchie et al. 2015) on the log_{2}-transformed, normalized counts. Group specific marker genes were defined as differentially expressed genes with a FDR < 0.1 and log_{2}FC > 0 for all pairwise comparisons. Initially, differential expression was tested between all seven detected groups. Groups 6 and 7 consistently grouped together when hierarchical clustering (default options from the hclust package) was performed on the log_{2} fold changes in expression between one group and all others (supplementary fig. 2, Supplementary Material online). We therefore merged groups 6 and 7 and tested differential expression between the now largest group and all other groups (supplementary table 1, Supplementary Material online). In the next step, we tested differential expression pairwise between all differentiated cell groups (supplementary table 2, Supplementary Material online).

In Situ Hybridization
For in situ hybridization (ISH), we collected 48 hpf or 72 hpf Platynereis larvae. Animals were fixed overnight in 4% PFA/1.5× PBS and ISH was performed as described previously (Tessmar-Raible et al. 2005).

Cloning of Platynereis Genes and ISH Probe Synthesis
For the cloning of P. dumerilii cDNA sequences, wild-type Platynereis RNA was isolated by Trizol/phenol/chloroform extraction method, reverse transcribed using SuperScriptIII reverse transcriptase (Life Technologies, cat. # 18080044), and amplified by PCR using Takara ExTag DNA polymerase (Clontech, cat. # RR001A) and respective gene-specific primers. The resulting PCR fragments were cloned into pCRII-TOPO vector (Life Technologies, cat # K4610-20). The cloned gene sequences and the gene specific primer sequences are available upon request.

For the synthesis of ISH probes, cDNA plasmids were linearized and antisense RNA probes were transcribed using SP6 or T7 RNA polymerase (Roche, cat. # 11487671001 and 10881775001, respectively) and DIG RNA-labeling mix (Roche, cat. # 11277073910) or Fluorescein RNA labeling mix (Roche, cat. # 11685619910).

Microscopy and Image Processing
For imaging of WMISH samples for ProSPr, samples were mounted in 97% 2,2′-thiodiethanol (Sigma, cat. # 166782) and imaged on a Leica TCS SP8 confocal microscope, using a combination of fluorescence and reflection microscopy (Jekely and Arendt 2007). The colocalization analyses and image postprocessing was performed using Fiji (Schindelin et al. 2012) software. The colorimetric WMISH samples were imaged on Zeiss AXIO Imager M1 microscope. The figure panels were compiled using Adobe Illustrator and Adobe Photoshop software.

Supplementary Material
Supplementary data are available at Molecular Biology and Evolution online.

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Author Contributions
K.A. and N.E. planned and conducted experiments, interpreted the data, designed the study, and wrote the manuscript. H.M.V., P.B., and P.C. conducted experiments. T.B. and J.C.M. interpreted the data, designed the study, and wrote the manuscript. H.M.V., P.B., and P.C. conducted experiments. T.B. and J.C.M. interpreted the data, designed the study, and wrote the manuscript. K.A. and N.E. planned and conducted experiments, interpreted the data, designed the study, and wrote the manuscript.

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