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Sensory cells in tunicates: insights into mechanoreceptor evolution

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Tunicates, the sister group of vertebrates, offer a unique perspective for evolutionary developmental studies (Evo-Devo) due to their simple anatomical organization. Moreover, the separation of tunicates from vertebrates predated the vertebrate-specific genome duplications. As adults, they include both sessile and pelagic species, with very limited mobility requirements related mainly to water filtration. In sessile species, larvae exhibit simple swimming behaviors that are required for the selection of a suitable substrate on which to metamorphose. Despite their apparent simplicity, tunicates display a variety of mechanoreceptor structures involving both primary and secondary sensory cells (i.e., coronal sensory cells). This review encapsulates two decades of research on tunicate mechanoreception focusing on the coronal organ's sensory cells as prime candidates for understanding the evolution of vertebrate hair cells of the inner ear and the lateral line organ. The review spans anatomical, cellular and molecular levels emphasizing both similarity and differences between tunicate and vertebrate mechanoreception strategies. The evolutionary significance of mechanoreception is discussed within the broader context of Evo-Devo studies, shedding light on the intricate pathways that have shaped the sensory system in chordates.

KEYWORDS

mechanoreceptor, evolution, placode, chordates, hair cells, primary sensory cells, secondary sensory cells

1 Introduction

Twenty years ago, a paper provocatively titled "Novel, secondary sensory cell organ in ascidians: in search of the ancestor of the vertebrate lateral line" by Burighel and others (Burighel et al., 2003), provided evidence that the tunicate ascidian *Botryllus schlosseri* possessed a complex mechanosensory organ, the coronal organ. Unlike the previously characterized multicellular mechanoreceptor organs of adult tunicates (Manni and Pennati, 2015), this novel organ was not composed of peripheral neurons (*i.e.*, primary sensory cells) but showed dedicated axonless secondary receptor cells. These secondary receptor cells were contacted at their base by neurites coming from brain neurons, forming both afferent and efferent synapses with the sensory cells (Figure 1A). This discovery also revealed that the adult tunicate brain possessed sensory neurons, since then not considered, for the elaboration of afferent information from the coronal sensory cells and their control by means of efferent inputs. Moreover, in *B. schlosseri*, coronal sensory cells showed an apical bundle with a cilium accompanied by microvilli and/or stereovilli. They were aligned on the



(A) Illustration of a primary and a secondary receptor cell in tunicates. The primary receptor is a peripheral neuron, whose soma (indicated by the blue nucleus) is in the epidermis. The secondary receptor (orange) is, vice versa, a dedicated receptor that transmits its input to a brain sensory neuron. (B) Chordate evolutionary tree. * The monophyly of Phlebobranchia is disputed [see (DeBiasse et al., 2020)]. Stolidobranchia species are defined as Pleurogona (with gonads in the lateral body wall), whereas Phlebobranchia and Aplousobranchia are defined as Enterogona (with gonads close to the gut), (C) Adult individual of the ascidian Molaula socialis (right view), (D) Young colony of Botryllus schlosseri composed by two adult zooids and their buds. Dorsal view. (E) Ciona robusta larva at stage 28, 18 h post fertilization at 20°. Arrowheads: two anterior papillae. Ascidian larvae are composed of an anterior cephalenteron, i.e., a body part including both head structures (such as the brain) and trunk structures (such as the gut), and a posterior tail. The cephalenteron is usually called a "trunk".

oral siphon tentacles and exposed to the incoming seawater. In many aspects, these cells resembled vertebrate hair cells of lateral line organs (Manni et al., 2004). These features, combined with the evolutionary proximity between tunicates and vertebrates, considered sister groups (Delsuc et al., 2018), initiated a controversial yet exciting debate on the homology of coronal sensory cells and hair cells. This discussion extended to the homology of the embryonic territories from which they originate. Vertebrate hair cells derive from neurogenic placodes (namely, from the otic and the lateral line placodes) that, together with the neural crest, were at that time considered exclusive to vertebrates (Manni et al., 2001; 2004). Therefore, the discovery of the coronal organ (with sensory cells hypothesized homologous to vertebrate hair cells), together with the publication of the first data on the presence of placodal area and neural crest-like cells in tunicates (Manni et al., 2001; Jeffery et al., 2004; Mazet and Shimeld, 2005), challenged the foundation of the main theory of vertebrate evolution, the so-called "New head hypothesis" (Gans and

Northcutt, 1983). This theory proposed that neurogenic placodes and neural crest cells were, with respect to non-vertebrate chordates, novel cell populations that contributed to the success of vertebrates and their development of complex nervous systems.

In the past 20 years since the discovery of the coronal organ, numerous aspects of its morphology, physiology, and development have been elucidated (Burighel et al., 2011; Manni and Pennati, 2015). The organ has been found in all the tunicate taxa (except for Salps, see below) so far examined. Its mechanoreceptive function and synaptic connectivity has been established and some key developmental genes studied. Nevertheless, many questions remain unanswered, making the investigation of tunicate mechanoreception an intriguing question in evolutionary developmental biology research.

Tunicates constitute a diverse group of marine invertebrates, including both pelagic and sessile animals with different behavior and motility, thus having varied sensory requirements. Traditionally, tunicates were classified into three classes: the sessile Ascidiacea and



the pelagic Thaliacea and Larvacea. However, molecular phylogenies suggest that ascidians are a paraphyletic group and support the monophyly of thaliaceans (Delsuc et al., 2018; Kocot et al., 2018; DeBiasse et al., 2020) (Figures 1B-D). The tunicate tadpole swimming larva exhibits a typical chordate body plan which is lost during metamorphosis in ascidians and thaliaceans. At this stage the latter adopts a sac-like body with two apertures, the oral and the atrial siphons, for seawater circulation and filtration. The sessile ascidians, the most extensively studied group, exhibit a larva with a tripartite brain derived from the dorsal nerve tube. The larva also possesses numerous primary mechanosensory cells scattered in the monolayered epidermis, allowing the detection of a suitable substrate for metamorphosis (Wakai et al., 2021; Sakamoto et al., 2022). The sessile adult has a ganglionic brain, and its mobility is limited to the siphon and body wall contraction as defensive responses (Mackie and Burighel, 2005).

Here we review research on mechanoreception in tunicates. We begin by examining mechanoreceptor cells and organs based on primary receptors in the ascidian larva (Section 2). Then, we describe the diversity of mechanoreception structures (including both single or clustered cells and multicellular organs) based on primary receptors exhibited by adult tunicates (Section 3). Lastly, we consider 20 years of research on the coronal organ from a morphological, physiological and developmental point of view, describing similarities and differences between coronal sensory cells and vertebrate hair cells (Section 4).

2 Putative mechanosensory cells of the ascidian larva

As the larvae are primarily responsible for ascidian dispersal, their simple swimming behavior is modified by environmental stimuli. This is likely to increase the odds of escaping predation and settling in a suitable location for metamorphosis. For instance, mechanical stimulation of the adhesive/sensory papillae, the three sensory organs (two dorsal and one ventral) located in the anterior larval region (Figures 1E, 2A–B) is sufficient and necessary to trigger metamorphosis in *Ciona* (Wakai et al., 2021; Sakamoto et al., 2022). While mechanosensitive modulation of swimming has yet to be definitively shown in ascidian larvae, startle-like behaviors have been described in *Ciona* (Athira et al., 2022). Here we discuss what is known about the development and function of the candidate primary mechanosensory cell types that have been identified in these larvae.

2.1 Caudal epidermal neurons

Sometimes referred to as Caudal Epidermal Sensory Neurons, these primary receptors are a subset of a broader class of tail epidermal neurons and were initially described in *Diplosoma listerianum* (previously named *D. macdonaldi*) (Torrence and Cloney, 1982). They have since been reported in numerous other

species spread across both enterogonid (e.g., Ciona, Phallusia) and stolidobranch (e.g., Halocynthia, Molgula) ascidians (Torrence and Cloney, 1982; Takamura, 1998; Imai and Meinertzhagen, 2007; Terakubo et al., 2010; Ohtsuka et al., 2014; Ryan et al., 2018). Although the CENs have yet to be conclusively shown to be mechanosensory, Torrence and Cloney proposed this based on ultrastructural similarities to cupular organ mechanoreceptors of the adult (Torrence and Cloney, 1982). (Figure 1E) CENs are found embedded in the epidermis of the larval tail, along both the dorsal and ventral midlines (Figure 1E). Therefore they are sometimes divided into Dorsal CEN (DCEN) and Ventral CEN (VCEN) subtypes (Ryan et al., 2018) (Figure 2A). CENs occur in regularly interspersed pairs, but their number is variable, with an average of 14 pairs per larva reported in Ciona (Pasini et al., 2006). Each neuron bears a single cilium that projects into the overlying extracellular tunic. In the tunic these cilia form a branched network termed the ASNET (Ascidian Dendritic Network In Tunic) (Torrence and Cloney, 1982; Konno et al., 2010; Terakubo et al., 2010; Yokoyama et al., 2014). While the base of the cilia are formed by microtubules and are clearly stained by anti-acetylated tubulin immunofluorescence, their distal portions in the tunic do not contain ordered microtubule arrays (Torrence and Cloney, 1982; Terakubo et al., 2010). CENs possess short axons that form contacts primarily between each other and a few putative relay neurons, such as the Bipolar Tail Neurons (BTNs) (Stolfi et al., 2015; Ryan et al., 2018). The BTNs have been proposed to be homologous to vertebrate cranial sensory neurons (Papadogiannis et al., 2022), though their sensory capabilities are entirely unknown. Like all epidermal neurons, CENs express Slc17a6/7/8, encoding Vesicular glutamate transporter (Vglut) and are therefore likely glutamatergic (Horie et al., 2008).

Extensive work in Ciona has revealed the embryonic origins of the CENs and the molecular pathways regulating their specification. CENs arise from neurogenic midlines of the tail epidermis, both dorsally and ventrally. Both midlines are derived from b-lineage blastomeres that also give rise to epidermal cells and BTNs (Pasini et al., 2006). While induction of the dorsal and ventral neurogenic midlines occurs independently through different signals, they converge on a shared gene regulatory network for sensory neurogenesis (Pasini et al., 2006; 2012; Tang et al., 2013; Waki et al., 2015). Both midlines express Msx, which in turn activates the expression of the proneural bHLH gene, Achaete-Scute-like.a (Ascl.a, though sometimes referred to as Ascl2 or Ascl.b previously). However, in the dorsal midline, Msx is activated by Otx and Nodal, while in the ventral midline it is activated instead by Tbx2/3 (Pasini et al., 2006; Waki et al., 2015). Otx and Nodal expression in the dorsal midline in turn depends on FGF signaling, while Tbx2/ 3 expression in the ventral midline is induced by ADMP/BMP signaling instead (Pasini et al., 2006; Waki et al., 2015). Downstream of Ascl.a, both dorsal and ventral networks appear to function through a series of transcription factors, especially conserved neuronal selectors such as Pou4 and Myt1 (Tang et al., 2013). While all the cells in these neurogenic midlines express Ascl.a and thus likely have the potential to give rise to CENs, the final number of neurons is limited by typical lateral inhibition via the Delta/Notch pathway and the microRNA miR-124 (Chen et al., 2011; Tang et al., 2013).

Given that the dorsal and ventral neurogenic midlines are induced by different mechanisms, it has been proposed that one may have evolved as a co-option of the other (Waki et al., 2015). More specifically, it was proposed that the ventral midline is the ancestral one, as induction of ventrolateral sensory neurons by BMP is observed in cephalochordates as well (Lu et al., 2012). The neurogenic dorsal midline of tunicates and vertebrates would therefore represent a co-option of this neurogenic program in the last common olfactorian ancestor (last common ancestor of tunicates and vertebrates). In vertebrates, the dorsal neurogenic domain would have allowed for the emergence of neural crestderived neurons and other sensory neuron types, like Rohon-Beard cells of anamniote larvae. Alternatively, the ventral midline may have evolved specifically in tunicates as a co-option of an ancestral Msx-dependent neural plate border program for sensory neuron specification. Complementary to these scenarios, it has also been suggested that both dorsal and ventral midlines were neurogenic in the chordate ancestor, and that vertebrates lost the ventral one (Pasini et al., 2006). However, it was reported that another tunicate species, Halocynthia roretzi, has only a small number of ventral CENs near the tail tip, and that its dorsal neurogenic midline depends on FGF, Nodal, and BMP combined, along with yetundiscovered inductive signals (Ohtsuka et al., 2014). Although the midline neurogenic programs are deeply conserved across tunicates (Coulcher et al., 2020), it is clear that additional work on diverse tunicate species will be required to better refine our evolutionary models.

2.2 Trunk epidermal neurons (RTENs, aATENs, and pATENs)

In addition to putative mechanosensory neurons of the tail, there are three epidermal neuron subtypes found in the dorsal areas of the epidermis of the larval "trunk". These primary receptors were defined as Trunk Epidermal Neurons. These are, from anterior to posterior, Rostral Trunk Epidermal Neurons (RTENs), Anterior Apical Trunk Epidermal Neurons (aATENs), and Posterior Trunk Epidermal Neurons (pATENs) (Imai and Meinertzhagen, 2007; Ryan et al., 2018) (Figure 2A). In Ciona, there are 7 RTENs on either left/right side (14 total neurons) of the dorsolateral epidermis between the papillae and the sensory vesicle (Ryan et al., 2018). The aATENs occur as two left/right pairs on either side of the dorsal midline (4 total neurons), while the four pATENs appear to lie directly on the midline (Ryan et al., 2018). Although all trunk epidermal neurons contribute to the larval ASNET, like CENs their mechanosensory abilities have never been tested (Abitua et al., 2015; Poncelet et al., 2022; Hoyer et al., 2024). The three subtypes all have well-developed axons but show different connectivity within the PNS network, hinting at distinct functions. RTENs form extensive chemical synapses onto a few different interneurons in the larval brain including the Eminens cells, which are GABAergic (Cao et al., 2019). pATENs on the other hand form extensive inputs onto the Ascending Motor Ganglion (AMG) complex, especially the sole cholinergic (i.e., excitatory) AMG neuron, AMG5 (Ryan et al., 2018; Ryan et al., 2016; Ryan et al., 2017; Kourakis et al., 2019; Popsuj and Stolfi, 2021). Downstream connections even suggest opposite effects on swimming behavior, either arresting swimming (RTENs) or triggering swimming (pATENs). In contrast, the aATENs do not appear to form very many chemical synapses, at least at the relatively early larval stage documented by the connectome studies (Ryan et al., 2018). This may support its proposed role as a neurosecretory cell, and potentially homologous to both olfactory neurons and Gonadotropin-releasing hormone (GnRH) neurons in vertebrates (Abitua et al., 2015; Okawa et al., 2020).

Much less is known about the development of the different Trunk Epidermal Neurons, compared to the CE Ns. In *Halocynthia*, RTENs are specified from anterior neural plate lateral border cells by FGF, Nodal, and BMP signaling (Ohtsuka et al., 2014), while excess BMP signaling appears to suppress the formation of the oral siphon placode, or stomodeum, which gives rise to the aATENs in Ciona (Abitua et al., 2015). A similar FGF/Nodal/BMP combination is required for CEN specification in Halocynthia (see above), though this may be different in *Ciona* and other tunicate families. This suggests that a common gene regulatory network might be shared between Trunk and Caudal subsets of ESNs. The development of the pATENs has not been studied at all, to our knowledge. In sum, much work remains to be done on both the function and development of these different Trunk Epidermal Neurons.

2.3 Papilla neurons

Despite our current knowledge of the Ciona larval connectome and the regulation of caudal and Trunk Epidermal Neuron development, there is little direct evidence supporting their mechanosensitive nature. There is no evidence directly refuting that CENs and assorted Trunk Epidermal Neurons are mechanosensory cells, either. However, the larval neuron most widely accepted as a mechanosensitive cell type is the Papilla Neuron (PN) (Figure 2A) (Manni et al., 2021). In Ciona, PNs (sometimes called Papilla Sensory Neurons or Primary Sensory Neurons of the Papillae) are found surrounding the three adhesive/sensory papillae at the very anterior end of the larva (Zeng et al., 2019). There are two dorsal papillae (one left, one right) and one medial ventral papilla (Figures 2A, B). Each papilla contains exactly 4 PNs, and additional cell types with proposed adhesive and/or sensory functions (Zeng et al., 2019; Johnson et al., 2023b). PNs are also found in species with complex eversible papillae (e.g., Diplosoma spp.) and even in those without overtly protrusive papillae, (e.g., Molgula spp) (Torrence and Cloney, 1982; Vorontsova et al., 1997). Larval metamorphosis in Ciona depends on mechanosensation, as larvae attach to a solid substrate to initiate tail regression and the transition to the post-metamorphic, sessile stage. Mechanical stimulation of the papillae were shown to be sufficient and necessary for triggering metamorphosis, while impairing PN development or function can block metamorphosis (Wakai et al., 2021; Sakamoto et al., 2022; Hoyer et al., 2024).

Like all epidermal neurons in the *Ciona* larva, PNs have apical cilia and axons. Their axons continue to extend posteriorly towards the larval brain during the swimming period, and these potentially late connections coincide with the competence period (Johnson et al., 2023b). Swimming larvae are not immediately competent to initiate tail regression and metamorphosis immediately after hatching, and competence to settle is acquired only after a few

hours of swimming (Nakayama-Ishimura et al., 2009), presumably while PN axons are still growing. Unfortunately, the *Ciona* larva connectome was described in a relatively early larval specimen, and these later connections have not been documented at the synaptic level (Ryan et al., 2016). Little else is known about how PNs might regulate tail regression and metamorphosis downstream of mechanical stimulation. It is known that GnRH is important for tail regression, while GABA appears to regulate GnRH release and other processes in metamorphosis, such as body rotation (Hozumi et al., 2020). However, it is unclear where and how these neurotransmitters act, in the absence of PN synaptic connectivity data.

The PNs develop from an anterior neurogenic territory surrounding the central cells of the papillae that shows many similarities to the neurogenic midlines of the tail (Johnson et al., 2023b; Roure et al., 2023). This territory expresses Ascl.a, and later on Delta/Notch signaling limits the number of Pou4+ cells that will differentiate into PNs (Johnson et al., 2023b). The papilla territory in turn is specified by Foxc and Foxg orthologs (Horie et al., 2018; Liu and Satou, 2019), which suggests an evolutionary connection to anterior placodes of vertebrate embryos. However, the cells that give rise to PNs appear to downregulate Foxg, while sustained Foxg expression is associated with the more central papilla cell types, like the Axial Columnar Cells (ACCs) (Johnson et al., 2023b). Knocking out Pou4 blocks PN differentiation and metamorphosis (Sakamoto et al., 2022; Johnson et al., 2023b). Similarly, using chemogenetics to inhibit PN function also inhibits metamorphosis (Hoyer et al., 2024). Candidate effectors of PN functions have also been knocked down/out, resulting in similar loss of metamorphosis. For instance, morpholino knockdown and TALEN-mediated knockout of a gene encoding the TRP channel family member PKD2 reduced the incidence of mechanically-induced Ca2+ transients in PNs and moderately inhibited metamorphosis (Sakamoto et al., 2022). Similarly, CRISPR/Cas9-mediated knockout of Vamp1/2/3, encoding synaptic vesicle protein Synaptobrevin, also modestly inhibited metamorphosis (Johnson et al., 2023a). However, the exact mechanotransduction channel in the PNs has yet to be identified. Based on Ca2+ imaging, PNs also respond to chemical cues, suggesting they may be polymodal sensory cells (Hoyer et al., 2024). Certain chemicals can promote or inhibit tunicate settlement and metamorphosis, suggesting that the larvae rely on both biotic and abiotic cues for optimal settlement site selection (Durante, 1991; Rae Flores and Faulkes, 2008; Hoyer et al., 2024). Interestingly, the ACCs at the very center of the papillae also respond to mechanical stimuli (Hoyer et al., 2024), which may reflect independent mechanosensory ability, or local communication between PNs and ACCs. In sum, although substantial work is still needed to better understand the development and function of the PNs, they represent a promising model for the study of tunicate mechanosensation thanks to the clear metamorphosis defects associated with their loss or perturbation.

2.4 Otolith and antenna cells

Most ascidian larvae have an otolith/statocyst, which is most frequently a single, rounded melanin-containing cell suspended in the lumen of the sensory vesicle (Torrence, 1986; Jiang et al., 2005) (Figures 2A–C). *Ciona* larvae exhibit strong geotactic behavior, preferring to settle on the dark underside of obstacles in the



water (Jiang et al., 2005), such as floating docks and ship hulls. A light- and gravity-dependent circuit has been proposed for ensuring such behavior, as larvae will swim up (or position themselves upwards if facing down) when drifting under a shaded area (Bostwick et al., 2020). Key to this behavior are the Antenna Cells, a pair of neurons that make contact with the otolith and presumably detect its displacement in the sensory vesicle via mechanotransduction (Torrence, 1986; Sakurai et al., 2004). However, little is known about these neurons aside from their characterization by the connectome studies, in which they were shown to make extensive synapses onto a handful of relay neurons in the larval brain (Ryan et al., 2016). Therefore, their mechanosensory nature is by far the most speculative and poorly documented out of all the candidate mechanoreceptors of the tunicate larva. A variant of the typical larvae of some colonial species belonging to the taxon Styelidae have only one sensory organ, the photolith, which is thought to function in both gravity and light reception (Sorrentino et al., 2001). In B. schlosseri, it consists of a unicellular statocyst, formed by an expanded pigment cup, which receives extensions from six photoreceptor cells (Sorrentino et al., 2001).

3 Mechanosensation in adult tunicates

Mechanoreception in adult tunicates is well developed, relying on both primary sensory cells, which are either scattered, organized in small clusters, or in specific organs (Figure 3), and the secondary

sensory cells of the coronal organ (as discussed in Section 3.3). However, information on primary sensory cells and/or organs containing primary sensory cells in adult tunicates is quite limited in comparison to that regarding the ascidian larva. In most cases only morphological data are available with occasional supplementation from the results obtained by neurophysiologists who worked in the field in the 70-90 s of the last century. Even though no developmental data are currently available for these primary sensory cells, developmental data is available for the oral siphons, a very sensitive region where many primary sensory cells are located. Specifically, the oral siphon primordium expresses anterior placode markers Pitx and Dlx, indicating that oral siphon primordia express genes shared with vertebrate placodes (Boorman and Shimeld, 2002; Irvine et al., 2007; Graham and Shimeld, 2013). The comparative morphology of the coronal organ has been deeply analyzed in several tunicate species and some aspects of its development and physiology have been studied in a select few ascidians.

3.1 Scattered or clustered primary mechanoreceptor cells in adult tunicates

Isolated primary cells (or small clusters of 2-3 primary sensory cells) have been described in the vicinity of the siphons, the most responsive regions of adult ascidian and thaliaean tunicates [reviewed in (Mackie and Burighel, 2005; Manni and Pennati, 2015)] (Figures 3A, B). These cells are peripheral neurons whose

somata are in the monolayered epidermis that both delimits the animal body and descends into the siphons. A long axon extends from the soma base to the central nervous system, whereas a dendrite, represented by a single cilium, projects apically. Since the epidermis is covered by the tunic, these intraepithelial sensory cells are not directly exposed to the external seawater. They react to stimuli through the tunic, which has a different thickness and elasticity depending on species and body region. In the ascidians Corella inflata and Corella eumyota, each siphon has approximately 8,000 primary sensory neurons that have been revealed by immunohistochemical labeling (Mackie et al., 2006). Physiological tests have shown that these cells are tactile and vibration sensors. In the same siphon region, rounded, axonless cells were also frequently seen, along with cells with very short axons; both have been suggested to be early stages in the formation of the sensory neurons. In the colonial ascidian B. schlosseri, the oral siphon primary sensory cells have been analyzed in relation to the blastogenetic cycle and to colony aging (Anselmi et al., 2022) (Figure 3B). The number of these mechanoreceptor cells varies along the cycle, increasing from early-to mid-cycle, before decreasing in late-cycle. This dynamic pattern parallels zooid sensitivity to stimuli, which is greatest when the number of mechanoreceptors is highest. Similarly, both the number of oral siphon primary sensory cells and zooid sensitivity to stimulation are higher in zooids belonging to young colonies than in those belonging to old colonies. In B. schlosseri, the ability of these mechanoreceptors to respond to stimuli has been detected using a sensitive behavioral test, the siphon stimulation test (Anselmi et al., 2022) (Figure 3C). This test assesses the ability of the oral siphon to close after stimulation with a waterjet whose pressure is controlled by a microinjector. The waterjet pressure is increased progressively, and the minimum pressure needed to induce a siphon contraction is recorded as a parameter indicating zooid sensitivity.

Primary ciliated sensory neurons, presumed mechanoreceptors, have also been reported in thaliaceans and appendicularians and were described using conventional light microscope staining techniques or Nomarski microscopy [reviewed in (Bone, 1998)]. Unfortunately, detailed morphological observations from transmission electron microscopy (TEM) are not available for these sensory cells. In *Pyrosoma*, those around the inhalant siphon were investigated experimentally, finding that their delicate touch evokes siphon contraction, whereas a stronger stimulation evokes a siphon contraction by branchial ciliary arrest (Mackie and Bone, 1978). In salps, sensory cells with a long cilium were also reported on the mouth lips, sometimes also organized in small groups. However, no detailed or physiological information is available on them (Bone, 1998).

3.2 Multicellular mechanoreceptor organs based on primary sensory cells in adult tunicates

A number of multicellular organs with putative mechanoreceptive function have been described morphologically in tunicates, both at light and electron microscopy [reviewed in (Bone, 1998; Mackie and Burighel, 2005; Caicci et al., 2013; Manni and Pennati, 2015)]. In some cases, their mechanoreceptive function has been determined by means of experimental studies; in other cases, it has been inferred on the basis of organ morphology and position. No data are available on their development. Among tunicates, ascidians have been more extensively studied than thaliaceans and larvaceans. The variety of multicellular organs, probably evolved from clusters of simple ciliated mechanoreceptors (Mackie and Singla, 2003), underlines the importance of mechanoreception and its behavioral integration.

The first organs to be described using scanning and transmission electron microscopy were the cupular sense organs (75-100 per individual) located in the atrial mantle epithelium of the adult ascidian Ciona intestinalis (Bone and Ryan, 1978) (Figure 3D). They are composed of groups of supporting cells flanking 15-20 ciliated neurons whose sensorial cilia are embedded in a gelatinous cupula, probably produced by the supporting cells. The cupula gives the name to the organs. These organs are able to detect near field vibrations as well as local water movements that displace the cupula and the cilia within it, resulting in electrical responses in the sensory cells. For their overall morphology and physiology, shared with the neuromasts of the lateral line organ and the hair cells of the vertebrate inner ear, the cupular sense organs were suggested by the authors to be evolutionarily linked to the vertebrate mechanosensory organs. From a cellular point of view, however, the cupular sense organs comprise primary sensory cells, whereas the vertebrate counterparts comprise secondary sensory cells, making the hypothesized homology inconsistent.

For many years the cupular sense organs were the only multicellular mechanoreceptor organs known in adult ascidians, until Mackie and Singla described in the atrial wall of the branchial sac of the solitary ascidian Chelyosoma productum the capsular organs at light and electron microscopy (Mackie and Singla, 2003) (Figure 3E). In the latter, the sensory cells are grouped in a macula and are characterized by a group of short microvilli surrounding a long cilium projecting into a small cavity (the "capsule"). The capsule cavity is delimited by supporting cells, is filled with a fluid and has an acellular diaphragm spanning an opening in the top. Each sensory cell has an axon reaching the brain via the visceral nerve, the nerve connecting the brain to the visceral organ (branchial sac, gut and heart). By means of electrophysiological recordings and tests aimed to determine their sensitivity, the authors concluded that these organs are vibrational-sensing and are adaptive in detecting the movements of objects in the vicinity.

The same authors described also in the genus *Corella* other organs based on primary sensory cells (Mackie and Singla, 2005). Using immunocytochemical analyses, they found in *C. eumyota* structures resembling the cupular sense organs of *C. intestinalis*, but located on the atrial surface of the branchial sac. Moreover, they recognized in *C. inflata*, using both immunocytochemistry and electron microscopy, a novel sense organ, the cupular strand (Figure 3F) which is a very elongated cupular organ located in the atrial surface of the branchial sac. Axons from the sensory cells enter the cerebral ganglion through the visceral nerve. Neither the cupular sense organs nor the cupular strand have been studied physiologically. However, by analogy with such structures in other metazoans, cupular organs were supposed to be hydrodynamic sensors registering local disturbances or changes in water flow through the atrial cavity.

A similar function was hypothesized for primary sensory cells of the colonial ascidian *Polyandrocarpa misakiensis* (Koyama, 2008). These cells form small corpuscles located in epidermal pockets filled with tunic at the base of the oral and atrial siphons and have been called "oral tentacular sensory cells" and "atrial tentacular sensory cells", respectively. The sensory cells, both isolated and forming small clusters, have axons joining a nearby nerve located at the base of the siphons (Figure 3G). Their apical apparatus is composed of a long, modified cilium projecting into the tunic, accompanied by a ring of microvilli of equal length. Supporting cells delimit the small cluster of sensory cells or are located between the isolated sensory cells. Some oral tentacular sensory cells are also found associated with neurosecretory cells.

Cupular sense organs have also been described in the thaliacean Pyrosoma atlanticum (Pyrosomatida), in a study aimed at describing at electron transmission microscopy the oral sensory structures of this tunicate (Caicci et al., 2013). The organs, previously mentioned by (Fedele, 1923), are scattered on the rounded flaps of the oral siphon and are composed of pyriform sensory cells accompanied by supporting cells (Figure 3H). The sensory cell apical plasmalemma exhibits a long cilium surrounded by 50-60 microvilli and is embedded in a tunic-like cupula secreted by supporting cells. An axon emerges from the sensory cell basal side. The organ function has not been investigated. However, displaying strong morphological resemblance with the ascidian cupular organs, it was supposed they play a similar mechanosensory role, probably in relation to reflex patterns involved in swimming control. Indeed, when the oral siphon is stimulated by touching, or by the collision of large particles or their entry into the gill, the animal responds by arresting ciliary beating and contracting the siphon (Bone and Ryan, 1978). It has been suggested that ascidian and thaliacean cupular organs are the result of evolutionary convergence (Caicci et al., 2013).

In the thaliacean *Doliolum nationalis* (Doliolida), triads of sensory cells, have been described in whole mount preparations (Bone, 1959) and by transmission electron microscopy (Caicci et al., 2013). These are a dozen groups of three sensory cells (supporting cells are not present) regularly arranged around the oral siphon, covered by the tunic. Each sensory cell has an apical long cilium projecting into the tunic and extends an axon from its base (Figure 3I). The triads are stimulated by the deformation of their apical cilia when water flows through the oral siphon as the animal swims (Bone, 1959). The oozooid stage of doliolids displays also an otocyst, but no detailed information is available on its morphology (Bone, 1998).

Apart from a statocyst containing a statolith, located on the left part of the Oikopleuridae brain, whose mechanosensory function has not been studied (Bone, 1998), no other multicellular mechanosensory organs based on primary sensory cells have been found in larvaceans. The ventral organ, a sensory structure below the mouth constituted of about 30 primary ciliated receptors, is considered a chemosensor (Bollner et al., 1986).

3.3 Secondary sensory cells in tunicates

The coronal organ has been found in all examined tunicates except salps (see paragraph 4.1) (Burighel et al., 2011; Caicci et al., 2013; Rigon et al., 2013). Positioned at the outer edge of the velum and the tentacles, at the base of the oral siphon, this organ comprises a continuous row of secondary sensory cells (Figure 4A). These cells are characterized by the presence of numerous stereovilli or microvilli and nonmotile cilia (a single cilium or multiple cilia) composed of 9 + 2 microtubules. The secondary sensory cells form a ring at the base of the oral siphon exposed to incoming water. Indeed, they function as mechanoreceptors involved in detecting variation in water flowing inside the oral siphon and possibly dangerous particles (Mackie et al., 2006).

Both afferent and efferent synapses are found between the base of coronal sensory cells and the peripheral axons of sensory neurons whose cell bodies lie on the brain (Burighel et al., 2003). The innervation pattern of the coronal organ has been studied through immunochemistry (Mackie et al., 2006; Gasparini et al., 2013a; Anselmi et al., 2022), and synaptic connectivity has been established using transmission electron microscopy (Burighel et al., 2003; Manni et al., 2004; 2006; Caicci et al., 2010b; Caicci et al., 2013) and in situ hybridization (ISH) experiments (Rigon et al., 2018). Each tentacle contains nerve fibers (from the subcoronal nerve) located at the base of the ciliated cells branching from the pericoronal nerve (Figures 4A, 5A, B), a nerve that encircles the oral siphon and originates from the anterior nerve brain. Synaptic contacts have been identified, using TEM, based on the presence of small presynaptic vesicles on one or both sides of the synaptic cleft and the characteristic thickening of the postsynaptic membrane. Frequently, sensory coronal cells make synapses with multiple neurites (Burighel et al., 2011). Glutamate (which mediates afferent hair cell inputs), acetylcholine, GABA and serotonin (which is involved in efferent stimulation to hair cells) are expressed in the coronal organ (Rigon et al., 2018).

The sensory cells are flanked on both sides by supporting cells and, in some species, by secretory cells. Typically, supporting cells extend apically a cytoplasmic crest delimiting the nearby sensory bundle (Figure 4A) and are connected to neighboring cells through tight junctions. There is no gap junction: signal transmission to the central nervous system is solely mediated by neurons located in the brain (Burighel et al., 2011). Secretory cells, when present, face towards the middle of the tentacles and do not form synapses with the nerve that contacts the sensory cells. Their function is not known, however the abundance of ER and the extended Golgi apparatus suggest that they are involved in protein synthesis. Their vicinity to the sensory cells suggests a secretory mechanism activated by the stimulation of the sensory cells (Manni et al., 2006). Sensory cells, supporting cells and, if present, secretory cells are all supported by a basal lamina that consists of a layer of fibers that merge and surround with the nerve fibers.

3.4 Variability of coronal sensory cells

The coronal organ exhibits a remarkable degree of diversity among the different tunicate species and even within the same species (Table 1). The diversity of the coronal organ is correlated to the variability of the apical structure and the presence or absence of secretory cells (Figure 4B). In Stolidobranch ascidians, three types of sensory cells have been identified based on the organization of their apical structure: a central cilium surrounded by microvilli (type 1),



FIGURE 4

Secondary sensory cells in the adult tunicates. (A) Location and main features of the coronal organ in tunicates. The organ is composed of a continuous row of cells on the oral tentacles and the velum (orange). Each sensory cell makes synapses with the subcoronal nerves (two per tentacle, close to the coronal organ) that are branches of the pericoronal nerve (green). The latter is a mixed nerve, connected to the brain through the anterior nerves. Sensory cells (pink) are flanked by supporting cells (grey); in some enterogona species, also secretory cells (violet) can be recognised. Stereovilli are apical, finger-like, long structures, composed of parallel actin filaments connected to the coronal organ variability in some representatives of tunicate groups. Stolidobranchia ascidians display the greatest complexity in the sensory apical bundle, which can be composed of microvilli or stereovilli, the latter also graded in length. * The monophyly of Phlebobranchia is disputed [see (DeBiasse et al., 2020)]. (C) Responses obtained after a strong (upper) and a light (bottom) stimulation of the coronal cells. The latter response is detected in the "tentacle stimulation test".

two long cilia and graded-height stereovilli (type 2), a complete ring of stereovilli surrounding two cilia (type 3). Interestingly, some species like *Styela plicata* can exhibit both type 1 and type 2 sensory cells (Manni et al., 2004). *Molgula socialis* presents a particularly complex condition since displaying all three types of sensory cells with types 2 and 3 predominantly located towards the proximal side of tentacles where they are exposed to inflowing water, while type 1 is located more peripherally (Figures 5C–F). It is worth noting that in Stolidobranch ascidians, extracellular radial filaments connecting the cilia to the surrounding stereovilli have been described (Burighel et al., 2003; Caicci et al., 2010a), even though the precise mechanism of signal transduction is not yet fully understood. In Phlebobranchiata and Aplusobranchiata some species have short microvilli (*Ciona robusta*), while others lack them (*Phallusia mammillata*). Additionally, some species such as *Ascidiella aspersa* and *C. inflata* possess secretory cells. Secretory granules have been found not only in both type 2 and type 3 sensory cells of Pleurogona (Manni et al., 2004; Manni et al., 2006; Caicci et al., 2007) but also in some sensory cells of Enterogona species (Manni et al., 2006).

In larvaceans the ultrastructure of the coronal organ was studied in two species of *Oikopleura* (Rigon et al., 2013). These animals have a single type of secondary sensory cells with numerous cilia of the same length, microvilli in multiple lines. They are flanked by nonciliated supporting cells forming a crest alongside the coronal organ. A recent study on the mesopelagic giant appendicularian

TABLE 1 Table summarizing the principal findings on the tunicate secondary sensory cells.

| Taxon | Species | Location | Proposed function | Behavioural test | Hair bundles | Cytoplasm of sensory cell | Radial filament connecting the cilia | Supporting cells | Accessory secretory cells | References |
|-------------------------------|--|-----------------------------|---|------------------------------|--|--|---|--|---------------------------------|--|
| Pleurogona stolidobranchia | Botryllus schlosseri | tentacles of oral siphon | Sensitivity to contact of inflowing particles | Tentacle stimulation test | Single cilium and stereovilli | Accessory centriole | Loose fibrillar matrix generally present among microvilli and cilia | Supporting cells form a wall or crest | | Burighel et al. (2003) |
| | Botrylloides leachi, B. violaceus | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | Single cilium and stereovilli | Accessory centriole | Loose fibrillar matrix generally present among microvilli and cilia | Supporting cells form a wall or crest | | Burighel et al. (2003), Burighel et al. (2008) |
| | Molgula socialis | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | Three types (type 1,2 and 3). Stereovilli | Electron dense granules, accessory centriole in sensory cells | Extracellular radial filaments connecting the cilium or cilia to the surrounding stereovilli | Supporting cells form a wall or crest | | Caicci et al. (2007) |
| | Pyura stolonifera, P.haustor | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | A pair of cilia surrounded by a crescent ring of stereovilli graded in length | Accessory centriole in sensory cells, electron dense granules | Extracellular radial filaments connecting the cilium or cilia to the surrounding stereovilli | Supporting cells form a wall or crest | | Caicci et al. (2010a) |
| | Styela plicata. S. montereyensis, S. gibsi | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | A pair of cilia surrounded by a crescent ring of stereovilli graded in length | Accessory centriole in sensory cells, electron dense granules | Extracellular radial filaments connecting the cilium or cilia to the surrounding stereovilli | Supporting cells form a wall or crest | | Manni et al. (2004), Caicci et al. (2010a) |
| | Polyandrocarpa zorritensis | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | A pair of cilia surrounded by a crescent ring of stereovilli graded in length | Accessory centriole in sensory cells, electron dense granules | Extracellular radial filaments connecting the cilium or cilia to the surrounding stereovilli | Supporting cells form a wall or crest | | Caicci et al. (2010a) |
| Enterogona aplousobranchia | Clavelina lepadiformis | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | More than two cilia of same length that constitute an oriented rows parallel to coronal organ; microvilli | | | Supporting cells form a wall or crest | | Manni et al. (2006) |
| | Diplosoma listerianum | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | More than two cilia of same length that constitute an oriented rows parallel to coronal organ; microvilli | | | Supporting cells form a wall or crest | | Manni et al. (2006) |

(Continued on following page)

| Taxon | Species | Location | Proposed function | Behavioural test | Hair bundles | Cytoplasm of sensory cell | Radial filament connecting the cilia | Supporting cells | Accessory secretory cells | References |
|------------------------------|------------------------------------|---|---|------------------------------|---|--|--|--|---------------------------------|---|
| Enterogona phlebobranchia | Ciona robusta | tentacles of oral siphon | Sensitivity to contact of inflowing particles | Tentacle stimulation test | More than two cilia of same length that constitute an oriented rows parallel to coronal organ; microvilli | Accessory centriole in sensory cells | | | | Mackie et al. (2006), Manni et al. (2006) |
| | Ascidiella aspersa | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | More than two cilia of same length that constitute an oriented rows parallel to coronal organ; microvilli | | | | Accessory secretory cells | Mackie et al. (2006), Manni et al. (2006) |
| | Phallusia mammillata | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | More than two cilia of same length that constitute an oriented rows parallel to coronal organ, no microvilli/stereovilli | Electron dense granules in sensory cells | | | | Mackie et al. (2006), Manni et al. (2006) |
| | Chelyosoma productum | tentacles of oral siphon | Sensitivity to contact of inflowing particles | | More than two cilia of same length that constitute an oriented rows parallel to coronal organ; microvilli | | | | Accessory secretory cells | Mackie et al. (2006), Manni et al. (2006) |
| | Corella inflata, C. willmeriana | tentacles of oral siphon | Sensitivity to contact of inflowing particles | Tentacle stimulation test | More than two cilia of same length that constitute an oriented rows parallel to coronal organ; no microvilli/stereovilli | | | | Accessory secretory cells | Mackie et al. (2006), Manni et al. (2006) |
| Appendicularia | Okopleura dioica, O. albicans | lower lip and pharynx | Monitoring particle flow into pharynx | | More than two cilia different in lengths and shorter toward the cell edges; microvilli | | | Supporting cells form a wall or crest | | Bone, 1998; Rigon et al. (2013) |
| Thaliacea | Pyrosoma atlanticum | flaps and a single ventral tentacle | Sensitivity to contact of inflowing particles | | Single cilium, stereovilli | | | | | Caicci et al. (2013) |
| | Doliolum nationalis | flaps | Sensitivity to contact of inflowing particles | | Single cilium, stereovilli | | | | | Caicci et al. (2013) |
| Salpe | Thalia democratica | absent | | | | | | | | Rigon et al. (2013) |

Bathchordeaus stygius has revealed the presence of three pairs of oral sensory organs within the mouth cavity, hypothesized to be homologous to the coronal organ. Each of these organs is composed of sensory cells with an apical cilium, innervated by brain nerves and surrounded by non-ciliated epidermal cells that nearly cover the organ (Le et al., 2023). In addition to the coronal organ, appendicularians possess the so-called Langerhans cells, which are secondary mechanoreceptors located in the posterior of the "trunk" epidermis and connected with afferent neurites through gap junctions. When stimulated, Langerhans cells trigger the escape response of the animal (Bone and Ryan, 1978).

In thaliaceans, the coronal organ has been studied in Pyrosoma atanticum and D. nationalis (Caicci et al., 2013). These animals have a single type of secondary sensory cell possessing a cilium accompanied by microvilli. Instead of tentacles, at the base of the oral siphon thaliaceans have flaps (D. nationalis), or single ventral tentacles with dozens of flaps (P. atlanticum). Notably, the coronal organ is absent in Thalia democratica, a salp. This absence is likely a derived condition evolved in parallel with the different feeding system adopted by this group of animals. Indeed a cladistic analysis, performed using 19 morphological characters in 16 tunicates species, and a cephalochordate and three vertebrate species as outgroups, revealed that the putative ancestral coronal cell in tunicates was a simple monociliated cell, that successively differentiated into the current variety of oral mechanoreceptors in the various tunicate lineages. The evolutionary changes in sensory cells may correspond to different feeding strategies (Rigon et al., 2013).

3.5 Physiology of the coronal organ

Studies aimed to elucidate the function of the coronal organ have primarily focused on two species: the solitary ascidian C. inflata (Mackie et al., 2006) and more recently the colonial ascidian B. schlosseri (Manni et al., 2018; Anselmi et al., 2022; Thompson et al., 2022). Behavioral experiments aimed to manipulate water flow patterns and observe siphon closure responses demonstrated that the secondary sensory cells are mechanoreceptors (Mackie et al., 2006). In C. inflata, a pioneering study showed that stimulating the oral tentacles with a glass needle caused the atrial siphon to contract to less than half its resting diameter, with no change in the diameter of the oral siphon. This response was named "crossed response" (Figure 4C). Depending on the stimulus strength and duration, the degree of atrial siphon closure during the crossed response varied. While gentle stimulation of the inner surfaces of the siphon or oral tentacles elicited varying degrees of the crossed response, stronger stimulation induced "squirts", characterized by a robust, synchronous contraction of both siphons and adjacent regions of the body wall (Figure 4C). This was accompanied by arrest of the cilia activity in the branchial stigmata responsible for creating the water current. Notably, a single stimulation could evoke not just one but a series of contractions suggesting coordination through a pacemaker (Mackie et al., 2006). These responses were lost after tentacle amputation. Electrophysiological recordings on the oral siphon were conducted to measure the electrical activity of the secondary sensory cells when exposed to specific stimuli in order to understand how sensory cells are activated and transmit signals. The results confirmed that crossed responses and squirts are centrally mediated reflexes but local conduction pathways also exist and persist after brain removal (Mackie et al., 2006).

Further insights have emerged from a different type of behavioral experiment, the tentacle stimulation test, conducted in B. schlosseri to assess animal performance under different conditions (Manni et al., 2018; Anselmi et al., 2022). This test aims to record the minimum pressure applied to the tentacle required to trigger the crossed reflex. Controlled and quantifiable pressure was applied through a water jet flow directly to the tentacles. Results showed that stage of adult individuals, the age of the colonies, and their overall condition (e.g., exposure to drug) significatively influence the zooids performance. Specifically, a higher threshold for response is observed in case of lower numbers of brain neurons, as in old colonies and zooids approaching their resorption, or in case of coronal organ impairment following drug treatment. In this regard, is it important to mention that the coronal sensory cells, like vertebrate hair cells, are damaged by gentamicin (an ototoxic drug) treatment resulting in a loss of coronal sensory cell continuity along the organ (Manni et al., 2018). This leads to a significant decrease in the percentage of responsive zooids to the tentacle stimulation test compared to the same colonies before treatment. Interestingly, fenofibrate has been found to have a strong protective effect on coronal sensory cells against the gentamicin-induced toxicity, similar to what occurs in vertebrate hair cells (Park et al., 2017; Manni et al., 2018).

Additionally, experiments involving stimulation of the oral siphon with ultrasound were conducted on three solitary ascidians. These experiments revealed that the coronal organ plays a role in perceiving ultrasounds, exhibiting a frequency-dependent behavioral response. Higher sensitivity was observed at the highest frequency tested (Varello et al., 2023).

3.6 Secondary sensory cell development

In tunicates, the coronal organ develops during embryogenesis from a thickened ectodermal epithelium known as the "anterior proto-placode". This tissue eventually gives rise to the oral siphon, tentacles and velum (Manni et al., 2004; Gasparini et al., 2013a; Manni et al., 2018). Importantly, the anterior proto-placode expresses homologues of some placodal genes (Patthey et al., 2014), (Figure 6).

The development of the coronal organ has been studied using TEM in *C. robusta* and *B. schlosseri* (Manni et al., 2004; Gasparini et al., 2013a). In *Ciona*, coronal cells become morphologically recognizable during the early juvenile stage when they appear as cells with short cilia and occasional microvilli. Over time, these cells progressively develop hair cell-like features, including microvilli containing actin and myosin, and associated with supporting cells. Importantly, the coronal organ continues to grow throughout the entire lifespan of the animal, parallel to the growth of the tentacles. Immunohistochemistry has revealed higher mitotic activity in the coronal organ of adults, with proliferating coronal sensory cells and supporting cells detected using Histone H3 antibody (Gasparini et al., 2013a). ISH has shown that orthologs of genes such as *Atoh1*, *Notch*, *Delta-like*, *HES*, and *Musashi* are expressed during the development of the coronal organ



FIGURE 5

(A,B) Confocal pictures of the *B. schlosseri* oral siphon and tentacles stained with anti-alpha tubulin (green) labelling nerves, phalloidin (red) labelling cytoplasmatic actin and dapi (blue) labelling cell nuclei. (C,D) Scanning electron microscopy showing the coronal organ of *Molgula socialis*. Squared area in C is enlarged in D. The organ is composed of a row of 1-2 sensory cells (recognisable by their hair bundle) flanked by supporting cells characterized by an apical cytoplasmic crista (arrowhead). Two types of sensory cells can be recognised: with a couple of cilia surrounded by graded stereovilli (type 3), and with a single cilium surrounded by microvilli (type 1). (E,F) Transmission electron microscopy showing a transverse section of the coronal organ of *M. socialis*. Squared area in E is enlarged in F to show the different apical bundle structure: two sensory cells at left display microvilli (type 1), whereas the sensory cell at right possesses stereovilli (type 2 or 3).

(Rigon et al., 2018) mirroring gene expression patterns seen in vertebrate neural and hair cell differentiation (Fritzsch and Elliott, 2017).

The development of the coronal organ during asexual reproduction has been investigated in *B. schlosseri* (Manni et al., 2018). The coronal cells undergo cyclical development on a weekly basis becoming first recognizable in the buds during mid-cycle due to the differentiation of their apical bundle and basal synapses. Their definitive configuration is reached when their parents are in late-cycle which coincides with the degeneration of both the parent brain and sensory system.

4 Evolutionary relationships between vertebrate and invertebrate mechanoreceptor cells

In addition to primary mechanosensitive sensory neurons, vertebrates possess specialized secondary mechanosensory cells, including the hair cells of the inner ear and lateral line and Merkel cells of the skin. Vertebrate hair cells share several developmental, morphological, and functional similarities with tunicate primary and secondary mechanoreceptor cells. These similarities and differences are discussed below.

4.1 The development of vertebrate hair cells from cranial placodes

Towards the end of gastrulation, the vertebrate neural plate arises from the most dorsal population of embryonic ectoderm in response to a variety of organizing signals (Grocott et al., 2012; Groves and LaBonne, 2014; Thawani and Groves, 2020). The border between the developing neural plate and the future epidermis gives rise to two distinct multipotential lineages - neural crest cells and cranial placodes. The cranial placodes are ectodermal thickenings that give rise to (1) cranial sensory neurons of the trigeminal ganglia, (2) cranial sensory neurons of the epibranchial ganglia, (3) the otic placode which will form the entire inner ear including hair cells and sensory neurons, (4) the olfactory epithelium, (5) the lens of the eye, and (6) the anterior pituitary gland (Singh and Groves, 2016; Streit, 2018); (Figure 6A). In aquatic vertebrates, lateral line placodes will give rise to the hair cell-bearing neuromasts located along the head and trunk of the animal. The anterior and posterior lateral line neuromasts of teleosts, named for the direction in which their cells migrate, arise from a lateral line placode in the vicinity of the otic placode. Some aquatic species, like axolotls and paddlefish, have ampullary organs containing specialized electroreceptor cells that also arise from a lateral line placode (Baker et al., 2013; Modrell et al., 2017).

Cells destined to give rise to cranial placodes first express members of the AP2a and FoxI families, which distinguish nonneural ectoderm from the adjacent neural ectoderm (Ohyama and Groves, 2004; Grocott et al., 2012; Khatri et al., 2014; Pla and Monsoro-Burq, 2018); (Figure 6B). Some cells within this region then express both Six homeobox transcription factors and Eya coregulators in what has been termed the pre-placodal domain (Streit, 2004; 2007; Sato et al., 2010). Six1 is expressed throughout the preplacodal domain, posterior placodes like the otic placode express Eval and Six4, and anterior placodes like the olfactory placode express Eya2 and Six3 (Streit, 2004; 2018; Sato et al., 2010). Locally acting signals then divide this pre-placodal region into distinct placodal territories along its anterior-posterior axis. Members of the Pax gene family play an important role in this process: Pax6expressing progenitors give rise to the olfactory and lens placodes, Pax3-expressing progenitors give rise to the trigeminal placode, and Pax2/8-expressing progenitors give rise to the otic placode and the epibranchial placodes (Baker and Bronner-Fraser, 2001); (Figure 6B).

Once the otic placode has formed, the tissue transforms by invagination into the otic vesicle or otocyst. The otocyst co-opts dorso-ventral and anterior-posterior signals used to pattern the central nervous system (Groves and Fekete, 2012; Wu and Kelley, 2012) to form a series of prosensory patches expressing the *Sox2* transcription factor (Raft and Groves, 2015). These prosensory patches will give rise to the auditory and vestibular epithelium containing hair cells and supporting cells. The surrounding nonsensory epithelium of the ear expresses the transcription factor *Lmx1a* and will give rise to structures such as the semicircular canals of the vestibular system (Gu et al., 2016; Żak and Daudet, 2021). Mutual antagonism between *Sox2* and *Lmx1a*, driven in part by Notch pathway signaling, leads to the correct positioning and distribution of the prosensory patches (Mann et al., 2017). Hair cells and supporting cells develop from each sensory patch through a

process of Notch-mediated lateral inhibition. Differentiating hair cells express Notch ligands to suppress a hair cell fate in neighboring cells, which then differentiate as supporting cells (Basch et al., 2016; Brown and Groves, 2020). Below, we discuss evidence for conservation of these developmental events in the formation of coronal sensory cells in tunicates.

4.2 What elements of vertebrate placode development are shared in tunicates?

The presence of thickened, placode-like structures in tunicates was first suggested by a study of the neurohypophysial duct, which generates the neural gland rudiment and migratory cells that contribute to the cerebral ganglion (Manni et al., 2005; 2001; 1999). The discovery of secondary hair cell-like cells in the coronal organs of Botryllus and Ciona (Burighel et al., 2003; Manni et al., 2005) suggested they may also derive from placodal structures. Subsequent analyses identified four thickened placodallike structures or "proto-placodes" in tunicate larvae: (1) the rostral proto-placode which will form the larval adhesive organ and its associated sensory neurons, and the adult stolon (Section 2.4), (2) the stomodeal proto-placode that will give rise to the oral siphon including the coronal organ, (3) the afore-mentioned neurohypophysial proto-placode, and (4) the atrial proto-placode which will form the atrial siphon and atrial chamber wall, containing primary mechanosensory cells of the cupular organs (Manni et al., 2004; Gasparini et al., 2013b); (Figure 6A). Subsequent studies analyzed the expression of vertebrate placodal patterning gene orthologues in the developing tunicate proto-placodes at the embryonic, larval, and juvenile stages of Ciona and Botryllus (Mazet and Shimeld, 2005; Gasparini et al., 2013b). The Ciona orthologs of AP2a and Sox2/3, Tfap2-r.b and Sox1/2/3, respectively promoted epidermal and neural fate of ectoderm cells (Imai et al., 2017). Members of the Six and Eya families were expressed in both the anterior (stomodeal/neurohypophysial) and posterior (atrial) proto-placodes, with Six1/2 marking both structure placodes and Six3/6 being confined to the anterior proto-placode. Tunicate FoxI orthologues were expressed in the posterior atrial proto-placode (Mazet and Shimeld, 2005; Gasparini et al., 2013b); (Figure 6B).

Based on these studies of two evolutionarily distant tunicate species, it has been proposed that the tunicate anterior protoplacodes resemble the vertebrate olfactory/lens/hypophyseal placodes, and the tunicate posterior proto-placode resembles the vertebrate otic/epibranchial/lateral line placodes (Gasparini et al., 2013b). However, in vertebrates only the posterior otic and lateral line placodes produce hair cells (Groves and LaBonne, 2014; Piotrowski and Baker, 2014), whereas in tunicates coronal sensory cells that most closely resemble vertebrate hair cells are derived from the anterior proto-placode. Only the primary mechanosensory cells of the cupular organs are derived from the posterior placode (Gasparini et al., 2013b). Since Six and Eya genes are initially expressed throughout the vertebrate pre-placodal domain (Streit, 2007; Schlosser, 2014), it is likely that additional transcription factor combinations are required to divide this domain more precisely into individual placodes, or that individual placodes are specified at different times. In this regard, it is interesting to note that vertebrate Foxi1/3 genes are initially expressed throughout the



FIGURE 6

Comparison of vertebrate placodal and tunicate proto-placodal development and vertebrate hair cell and tunicate coronal sensory cell structures. (A) Schematic of vertebrate placodes compared to tunicate proto-placodes. The anterior placodes include the olfactory, anterior pituitary, and lens placodes. The posterior placodes include the trigeminal, epibranchial, and otic placodes. Tunicates have three anterior proto-placodes:, the rostral, stomodeal, and neurohypophysial placodes. Tunicates have two posterior atrial proto-placodes. Following metamorphosis, the stomodeal proto-placode will give rise to the oral siphon and the atrial proto-placodes will fuse to form the atrial siphon. (B) Conservation of genes expressed during vertebrate placodal and tunicate proto-placodal development. Several key genes involved in placode development appear to be conserved. (C) Comparison of hair cells from vertebrates and coronal sensory cells from tunicates. Vertebrate hHair cells (tan) are flanked by supporting cells (gray). Sensory cells possess kinocilium (red) and stereovili (purple) that are connected together by different links.

pre-placodal domain, at a time when all parts of the pre-placodal domain are competent to generate the otic placode (Solomon et al., 2003a; 2003b; Ohyama and Groves, 2004; Birol et al., 2016) and are gradually downregulated in an anterior-posterior direction (Khatri et al., 2014; Birol et al., 2016). However, Foxi1/3 appear to be required for the development of only the posterior otic, epibranchial and lateral line placodes in vertebrates (Solomon et al., 2003a; Hans et al., 2004; Birol et al., 2016). Rigon and others (Rigon et al., 2018) speculated that the common ancestor of vertebrates and tunicates may have generated mechanosensory cells from both anterior and posterior placode-like structures, with the ability to generate such cells being lost from the anterior placodes in vertebrates and from the posterior ptoto-placodes in tunicates. One possible explanation for this difference is that the evolution of the tunic covering the exterior surface of the animal placed constraints on where mechanosensory cells could function, restricting them to oral structures on the interior of the body that do not have a tunic covering (Manni et al., 2006).

It should be stressed that the putative homology between a Foxi1/3 and Pax2/8-expressing atrial primordium and the vertebrate "otic-epibranchial progenitor domain" is still far from settled (Graham and Shimeld, 2013; Patthey et al., 2014). A n alternative explanation for this paradox is that the posterior region of the tunicate larval head that expresses Six and FoxI genes does not give rise to sensory structures at all. Vertebrate Six and Foxi1/3 genes are also expressed in the developing pharyngeal arch region of vertebrates at a slightly later stage than their expression in the pre-placodal domain (Ohyama and Groves, 2004; Khatri and Groves, 2013; Edlund et al., 2014; Birol et al., 2016; Ankamreddy et al., 2023), and they are required for correct formation of the pharyngeal arch structures (Solomon et al., 2003a; Nissen et al., 2003; Edlund et al., 2014). Interestingly, FoxI and Six orthologues are also expressed in the branchial fissures (stigmata) of the tunicate atrium where peribranchial and branchial epithelium contact each other and fuse (Gasparini et al., 2013b). This expression pattern is reminiscent of the requirement for Foxi1/3 in the vertebrate pharyngeal pouches and clefts that form by fusion of pharyngeal ectoderm and endoderm (Edlund et al., 2014; Hasten and Morrow, 2019). Thus, while Six and FoxI genes mark the posterior atrial proto-placode in both Ciona and Botryllus, it is possible that these genes are acting to regulate formation of the atrium itself, rather than the cupular mechanosensory cells in the atrial walls. It may be possible to test the function of tunicate Six and FoxI orthologues by CRISPR-based loss-of-function studies to determine if they are necessary for the formation of primary cupular mechanosensory cells in the atrium or only for the formation of the branchial fissures.

As discussed above, vertebrate placodes acquire their unique identity by expression of different Pax family genes. *Ciona* has six Pax family genes, and, although several are expressed in regions of the larval central nervous system, most do not appear to be expressed in any of the proposed proto-placode structures identified in tunicate larvae (Mazet et al., 2003; Imai et al., 2004; Hudson and Yasuo, 2005). Of the Pax genes in *Ciona*, *Pax2/5/8.a* is expressed in the larval atria and stomodeum cavities, and *Pax2/5/8.b* is expressed weakly in the invaginating stomodeum (Mazet et al., 2003; Mazet and Shimeld, 2005). It remains an open question whether any other Pax genes play a role in the formation or patterning of other placode-like structures in tunicates. For

example, Pax6 gene family members are well known to regulate eye development across most animal phyla (Kozmik, 2008). In vertebrates, Pax6 has also been co-opted to regulate the formation of the lens through its expression in the lens placode (Cvekl and Ashery-Padan, 2014). Ciona Pax6 is expressed in parts of the brain associated with the photoreceptive ocellus and has the three "lens" cells lying above the ocellus, although they are not believed to be homologous to vertebrate lens cells. These "lens" cells do not express Pax6, nor do they express beta-crystallin (Shimeld et al., 2005). Moreover, vertebrate Pax6 genes have a lens-specific enhancer that is not present in tunicates (Irvine et al., 2008), suggesting that the co-option of Pax6 to regulate development of a lens structure occurred after vertebrates and tunicates diverged. Clarification of the role of other Pax genes in tunicate placode derivatives will require more sensitive tools to localize their transcripts, such as single cell RNA-seq and in situ hybridization, and to test gene function using loss of function approaches such as CRISPR-Cas9.

4.3 What elements of vertebrate hair cell development are shared in tunicates?

As described above, vertebrate inner ear and lateral line hair cells develop from patches of prosensory tissue marked by Sox2, a member of the SoxB transcription family (Neves et al., 2013). SoxC family members, such as Sox4 and Sox11 then act within these patches to provide competence for differentiation of hair cells and supporting cells (Gnedeva and Hudspeth, 2015; Wang et al., 2023). Cells within this expression domain upregulate the proneural transcription factor Atoh1, which initially marks the progenitors of both hair cells and supporting cells (Yang et al., 2010; Li et al., 2022). ATOH1, which is both necessary and sufficient for hair cell fate, is quickly restricted to differentiating hair cells through Notchmediated lateral inhibition (Jarman and Groves, 2013; Cai and Groves, 2015). ATOH1 regulates other transcription factors, such as GFI1 and POU4F3, to establish a hair cell gene regulatory network (Iyer and Groves, 2021; Iyer and Groves, 2021). GFI1 can act with ATOH1 to positively promote the expression of hair cell genes and can also act alone to inhibit expression of neuronal genes (Jen et al., 2022; Jen et al., 2022). Since ATOH1 is also responsible for the differentiation of neurons in the cerebellum, brainstem, and spinal cord (Ben-Arie et al., 1997; Lai et al., 2011; Wu et al., 2023), it is possible that secondary mechanosensory receptor cells co-opted Gfi1 to repress neuronal gene networks during evolution. POU4F3 also promotes hair cell differentiation by acting as a feed-forward pioneer factor: it is first induced by ATOH1 and then binds to many other ATOH1 target genes to make them transcriptionally accessible (Yu et al., 2021). The combined action of ATOH1, GFI1, and POU4F3 promotes a hair cell fate, and these transcription factors are also capable of reprogramming embryonic stem cells, fibroblasts, or non-sensory cells of the ear to a hair cell fate (Costa et al., 2015; Menendez et al., 2020; Iyer et al., 2022).

Several lines of evidence suggest that these three transcription factors have an evolutionarily conserved role in the differentiation of mechanosensory cells. *Drosophila* orthologues of *Atoh1*, *Gfi1*, and *Pou4f3* (*atonal*, *senseless* and *acj6* respectively) are expressed in

developing chordotonal organs that have mechanosensory functions in proprioception, hearing, and balance (Jarman et al., 1993; 1995; Nolo et al., 2000; Lee and Salvaterra, 2002). Significantly, Atoh1 and Gfi1 can functionally replace atonal and senseless in Drosophila, and atonal can functionally replace Atoh1 in mice (Ben-Arie et al., 2000; Wang et al., 2002; Acar et al., 2006). Orthologues of Atoh1, Gfi1 and Pou4f3 have also been identified in Caenorhabditis elegans (Atonh1/ lin-32, Gfi1/pag-3, and Pou4/unc-86) and are necessary for the formation of AVM/PVM mechanosensory neurons (Baumeister et al., 1996; Zhao et al., 2020). A Class IV POU gene orthologue has also been shown to be necessary for the development of cnidarian (sea anemone) hair cell-like cells, although it is not known whether atonal-like factors regulate this gene in sea anemones (Ozment et al., 2021). Finally, Atoh1, Gfi1, and Pou4f3 are all expressed in the touch-sensitive Merkel cells of the skin (Lumpkin et al., 2003; Haeberle et al., 2004; Yu et al., 2021), and Atoh1 and Pou4f3 are both necessary for the differentiation of these cells (Maricich et al., 2009; Yu et al., 2021). Significantly, the inductive and pioneer feed-forward relationship between Atoh1 and Pou4f3 seen in hair cells is also conserved in Merkel cells, even though they regulate overlapping but distinct sets of genes in these two different mechanosensory cell types (Yu et al., 2021).

It is interesting to speculate on what gene networks are regulated by Atoh1, Gfi1, and Pou4f3 orthologues in the different kinds of mechanosensory cells described in the previous paragraph. In a very simplified view, a mechanosensory cell requires (1) membrane specializations to detect mechanical force (such as vertebrate stereovilli or arthropod ciliated dendrites); (2) membrane components to develop a receptor or axon potential; (3) a synaptic apparatus to allow propagation of the mechanosensory stimulus to downstream neurons. These functional modules are created by gene networks expressed during development and then homeostasis. When comparing the molecular identity of cell types, it is important to functionally contextualize homologous genes across species. For example, gene networks regulating synaptic specializations are likely to be more highly conserved between different mechanosensory cells compared to networks regulating the more varied types of force-detecting machinery in these different cell types. Supporting this idea, a recent study comparing vertebrate hair cells and Merkel cells found that genes directly regulated by ATOH1 and POU4F3 in both cell types tended to be associated with synapses, cation channels and potassium channels (Yu et al., 2021). While some modules expressed by an ancestral mechanosensitive cell type may have been conserved, it is also possible that comparable modules were convergently evolved. At present, we have little on how the development information of tunicate mechanosensory cells is regulated. There is currently no evidence that the coronal sensory cellscells and supporting cells of the coronal organ derive from a SoxB/SoxC-expressing domain analogous to the prosensory patches of vertebrates. Atonal and Pou4 orthologues are present in Ciona and are expressed in larval ciliated epidermal sensory neurons; in these cells CiAtonal has been reported to be epistatic to CiPouf4 (Tang et al., 2013). The coronal organs of Ciona express an Atonal orthologue, as well as members of the Notch pathway (Rigon et al., 2018), but these genes have yet to be definitively localized to coronal sensory cells or supporting cells. With the advent of molecular techniques such as single cell RNA-seq and CUT&RUN/CUT&Tag, it will become feasible to identify gene networks expressed in mechanosensory cells of different species and to identify the direct targets of transcription factors such as ATOH1, GFI1, and POU4F3 within these networks.

As discussed above, the development of tunicate coronal organ sensory cells from the anterior, stomodeal proto-placode differs from that of hair cells of the vertebrate inner ear and lateral line, which develop from posterior (otic and lateral line) placodes. Although evolutionary scenarios have been proposed to account for this difference (Rigon et al., 2018), the limited data on expression of downstream placodal and prosensory genes in tunicates makes it hard to define the pathways by which coronal sensory cells form in the developing coronal organ. Indeed, it is possible that expression of Atonal orthologues in the coronal organ epithelium is sufficient to generate coronal sensory cells and supporting cells without the need to pass through a pre-placodal, placodal or prosensory state. In support of this idea, the chordotonal organs of Drosophila are generated by upregulation of *atonal* in embryonic ectoderm to form sensory organ precursors, and over-expression of atonal or Atoh1 is sufficient to generate ectopic chordotonal organs in embryonic ectoderm (Jarman et al., 1993; 1995; Ben-Arie et al., 2000). Merkel cells of the vertebrate skin are generated directly from keratin-expressing epidermis without passing through a Sox2+ prosensory phase; here SOX2 appears to control the maturation of Merkel cells, rather than their specification (Lesko et al., 2013; Perdigoto et al., 2014). Finally, activation of Atoh1, Gfi1, and Pou4f3 in primary mouse fibroblasts is sufficient to induce many aspects of the hair cell gene regulatory network without prior activation of SoxB or SoxC factors (Menendez et al., 2020). Localization of Atoh1 and Sox2/SoxB orthologues in developing and mature coronal organ tentacles may help to address some of these questions and to more accurately identify the stages of differentiation of these cells.

4.4 What elements of vertebrate hair cell regeneration are shared in tunicates?

Many vertebrate inner ear and lateral line hair cells undergo gradual turnover and replacement in mature animals, and nonmammalian vertebrates can also robustly regenerate new hair cells after the endogenous hair cells are killed (Stone and Cotanche, 2007; Kniss et al., 2016). In non-mammalian vertebrates, new hair cells are generated by the upregulation of *Atoh1* in supporting cells, which then trans-differentiate to a hair cell fate (Stone and Cotanche, 2007). This can occur with or without supporting cell division, but ultimately leads to full replacement of hair cells and functional recovery. The one exception to this is mammals, where the cochlea is unable to regenerate new hair cells after the onset of hearing and the vestibular system is capable of only a modest amount of turnover and regeneration (Groves, 2010; Bucks et al., 2017). Given the ability of other vertebrates to regenerate hair cells, it is possible that with the ancestral form that gave rise to hair cells also had the capacity to regenerate. If tunicate coronal sensory cells and vertebrate hair cells have a shared evolutionary origin, do tunicate coronal sensory cells regenerate? As adult tunicates mature, the tentacles of the oral siphon continue to grow, implying that there must be some postmetamorphic mechanism to generate new coronal sensory cells and supporting cells. Transmission electron microscopy has revealed rare instances of apparently dividing sensory cells in Pyura haustor

(Caicci et al., 2007), and analysis of mitosis by PH3 staining in the coronal organs of adult and juvenile *C. intestinalis* indicates that both supporting cells and coronal sensory cells are capable of creating new coronal sensory cells (Gasparini et al., 2013a). Recent work suggests that exposure of tunicate coronal sensory cells to the ototoxic aminoglycoside gentamicin leads to an apparent loss of some sensory cells from the tentacles and impairs responsiveness of the coronal organ tentacles to touch (Manni et al., 2018). However, it is not known whether coronal sensory cells can be regenerated after such damage, nor whether any new coronal sensory cells are generated by neighboring supporting cells. Further studies are required to explore the potential for tunicate coronal sensory cell regeneration, and whether genes associated with hair cell regeneration in vertebrates like *Atoh1* play a role in this process.

4.5 A consideration of mechanotransduction in vertebrate hair sells and tunicate coronal sensory cells

Vertebrate hair cells are exquisitely sensitive mechanoreceptors; the human ear can detect sounds that vibrate the eardrum by one picometer. Hair cells have a hair bundle protruding from their apical surface consisting of a graded, stair-case-like array of long modified microvilli termed stereocilia or stereovilli (Vélez-Ortega and Frolenkov, 2019); (Figure 6C). Vertebrate hair cells develop with a single true cilium or kinocilium that migrates to an eccentric position on one side of the apical surface of the hair cell as the hair bundle develops (Frolenkov et al., 2004). The kinocilium persists in most mature vertebrate hair cells but degenerates in mammalian cochlear hair cells prior to the onset of hearing (Wang and Zhou, 2021). The apical tips of all but the longest stereovilli are joined to the next tallest stereovillus by a tip link consisting of a heterodimer of a protocadherin, PCDH15, and a cadherin, CDH23 (Vollrath et al., 2007). A mechanotransduction complex (Qiu and Müller, 2018; Holt et al., 2021), is present in all but the tallest stereovilli and this complex consists of pore-forming cation channels, TMC1 and/ or TMC2, and two other membrane proteins, TMIE and TMHS/ LHFPL5 which help modulate the pore properties of the channel [TMIE; (Zhao et al., 2014; Cunningham et al., 2020)] and bind to PCDH15 [TMHS; (Xiong et al., 2012; Zhao et al., 2014; Ge et al., 2018)]. Loss of any of these proteins compromises hair cell function and causes severe hearing loss. Deflection of the hair bundle applies force to each tip link, leading to an extremely fast (~10 µs) gating of the mechanotransduction channel (Gillespie and Müller, 2009). An array of accessory proteins (such as MYOSIN7A, harmonin, and sans) inside the stereovilli anchors the mechanotransduction complex to the actin core of each stereovillus (Schwander et al., 2010), and mutations in these proteins, or in CDH23 or PCDH15, lead to hereditary deaf-blindness known as Usher syndrome (Cosgrove and Zallocchi, 2014; Whatley et al., 2020). In addition, a second mechanosensitive ion channel, PIEZO2, lies at the base of the hair bundle and is responsible for what have been termed reverse-polarity currents (Beurg and Fettiplace, 2017; Wu et al., 2017), although precise function of PIEZO2 in hair cell mechanotransduction and bundle integrity is still unclear (Qiu and Müller, 2018).

As discussed in Section 3.4, the coronal sensory cells of tunicates show a far greater degree of diversity in different taxa than those of vertebrates (Manni et al., 2006; Caicci et al., 2010a; Rigon et al., 2013). This diversity is seen in the number of cilia, which can vary from just one or two in some groups, to multiple cilia that can be present in single or multiple rows (Table 1; Figure 4B; Figure 6C). The cilia can be located centrally or eccentrically as in vertebrates. Short microvilli can be present or can be elongated to appear more like stereovilli. In most tunicate taxa the stereovilli are of the same length, but in some groups the stereovilli have a more graded morphology reminiscent of a vertebrate hair bundle. Multiple different morphologies of sensory cells can occur in the coronal organs of some taxa, again reminiscent of the different hair cell types seen in vertebrate sensory organs, such as inner and outer hair cells of the mammalian cochlea, type I or type II vestibular hair cells, or the tall and short hair cells of the bird hearing organ, the basilar papilla. Unlike vertebrates, tunicate sensory cells do not appear to have clear tip links connecting their stereovilli, but some taxa show evidence of lateral connections between stereovilli, or between stereovilli and cilia (Burighel et al., 2003; Caicci et al., 2007; Rigon et al., 2013); (Table 1). Such links have some resemblance to the side links, ankle links, shaft connectors and top connectors that are present between stereovilli and between stereovilli and the kinocilium (Richardson and Petit, 2019).

What types of stimulus gate tunicate sensory cells? As discussed in Section 3.5 above, gentle stimulation of the oral tentacles by direct touch, vibration, or electrical shocks can lead to contractions of the atrial and oral siphons known as the crossed response, with stronger stimuli evoking a squirt response caused by strong contractions of both siphons and the body wall (Mackie et al., 2006; Manni et al., 2018). Similar responses can be evoked by particulate matter such as polystyrene beads or ground vegetable matter (Mackie et al., 2006), suggesting that at least one function of the coronal organ is to mediate particle expulsion in response to direct mechanical stimulation. This does not preclude other functions for sensory cells; in this regard it is intriguing that the bundle morphology of some tunicate sensory cells resembles that of electroreceptors seen in many fish and some amphibians (Baker, 2019). Elucidating the types of stimuli that tunicate coronal sensory cells respond to requires more electrophysiology studies such as whole-cell voltage clamp recordings from sensory cells or using fluid jet stimulation to evoke and measure mechanotransduction currents. Additionally, CRISPR may be used to create transgenic tunicates expressing a membranelocalized calcium sensor to detect mechanotransduction and presynaptic activity in hair cells.

It is currently unknown how coronal sensory cells respond to mechanical force, nor the range of forces that can evoke synaptic release. The wide variety of tunicate sensory bundle types, together with the absence of apical tip links in coronal sensory cells suggests it is unlikely that CDH23/PCDH15-mediated gating of a mechanotransduction channel of the sort seen in vertebrates is occurring in tunicates. However, the presence of side links between stereovilli and between stereovilli and cilia suggest an alternative method of mechanoreceptor gating. Indeed, such to kinociliary links have been shown mediate mechanotransduction in developing zebrafish hair cells before being replaced by stereovilli-based mechanotransduction in mature hair cells (Kindt et al., 2012). Insect chordotonal organs

facilitate mechanotransduction with members of the TrpN and TrpV channel family (Li et al., 2018). Although vertebrate hair cells seem to use TMC and PIEZO2 channels for mechanotransduction (see above), TrpN channels may also be required for mechanosensation in some cases (Sidi et al., 2003). The TRPA1 channel was originally proposed as candidates for the vertebrate mechanotransduction channel (Corey et al., 2004), but data from knockout mice suggests that neither TRPA1 nor 32 other Trp channels are necessary for mechanotransduction in mouse hair cells (Kwan et al., 2006; Wu et al., 2016). A Ciona orthologue of TrpA1 is expressed in coronal sensory cells (Rigon et al., 2018), but its role in mechanotransduction has yet to be tested. PIEZO2 is another possible candidate for the tunicate mechanotransduction channel; it is located at the base of the hair bundle in vertebrate hair cells (Wu et al., 2017) and therefore does not require tip-link based mechanotransduction. PIEZO2 also mediates Merkel cell mechanotransduction without the need for elaborate stereovilli or tip link-based machinery (Maksimovic et al., 2014; Woo et al., 2014; Nakatani et al., 2015).

4.6 Are vertebrate hair cells and tunicate coronal sensory cells homologous?

During chordate evolution, some cell types remain tightly conserved while others have been either lost or convergently evolved across different species. The concept of a "core regulatory complex" (CoRC) of transcription factors has been useful in devising evolutionary scenarios for cell types (Arendt et al., 2016) and as discussed above, mechanosensory cells across vertebrate and invertebrate taxa appear to share factors such as atonal/Atoh1, senseless/Gfi and Pou4 factors. Several models for the evolution of chordate mechanosensory cells have been proposed (for example, Schlosser, 2021). These models propose some form of basal primary sensory cell giving rise to two distinct cell types: a primary sensory neuron that is not mechanosensitive and defined by neurogenin-like transcription factors, and a mechanosensitive cell defined by atonal-like transcription factors and which either lacked an axon altogether (hair cells and coronal sensory cells) or just a short axon (caudal epidermal neurons; see 2.1 above).

At present, only atonal/Atoh1 expression has been characterized in the tunicate coronal organ and has not yet been localized to the coronal sensory cells. Nevertheless, the presence of both hair celllike cells adjacent to supporting cells, the expression of Notch pathway genes in these cell types and their derivation from proto-placodal structures make a reasonable case for homology between these cell types. However, this conclusion is complicated by the fact that tunicates undergo metamorphosis, which prevents a clear visualization of the transition from tunicate "proto-placodal" structures to a sensory organ. This transition can be readily observed in vertebrates as the pre-placodal domain gives rise to individual placodes, some of which produce hair cells.

Resolving the question of homology between vertebrate hair cells and tunicate coronal sensory cells will be helped by three recent technical advances. First, single cell transcriptional analysis will be able to determine whether the CoRC transcription factors present in vertebrate hair cells and supporting cells are also expressed in coronal sensory cells and their associated supporting/accessory cells. Second, the advent of CRISPR has facilitated loss-offunction studies in many new model and non-model organisms, and disruption of tunicate CoRC mechanosensory transcription factors will allow testing of their necessity for coronal sensory cell differentiation. Finally, it may be possible to perform lineage tracing experiments to determine tunicate proto-placodal cells do indeed contribute to coronal sensory cells following metamorphosis. Resolving these questions could elucidate the ancestral mechanosensory hair cell gene regulatory network or could uncover novel mechanisms of creating mechanosensitive hair cell-like cells in different species.

Author contributions

CA: Conceptualization, Data curation, Investigation, Writing-original draft, Writing-review and editing. GF: Data curation, Investigation, Writing-original draft, Writing-review and editing. AS: Investigation, Writing-original draft, Writing-review and editing. AG: Investigation, Writing-original draft, Writing-review and editing. LM: Conceptualization, Investigation, Supervision, Writing-original draft, Writing-review and editing.

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Conflict of interest

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References

Abitua, P. B., Gainous, T. B., Kaczmarczyk, A. N., Winchell, C. J., Hudson, C., Kamata, K., et al. (2015). The pre-vertebrate origins of neurogenic placodes. *Nature* 524, 462–465. doi:10.1038/nature14657

Acar, M., Jafar-Nejad, H., Giagtzoglou, N., Yallampalli, S., David, G., He, Y., et al. (2006). Senseless physically interacts with proneural proteins and functions as a transcriptional co-activator. *Development* 133, 1979–1989. doi:10.1242/dev.02372

Ankamreddy, H., Thawani, A., Birol, O., Zhang, H., and Groves, A. K. (2023). Foxi3GFP and Foxi3CreER mice allow identification and lineage labeling of pharyngeal arch ectoderm and endoderm, and tooth and hair placodes. *Dev. Dyn.* 252, 1462–1470. doi:10.1002/dvdy.645

Anselmi, C., Kowarsky, M., Gasparini, F., Caicci, F., Ishizuka, K. J., Palmeri, K. J., et al. (2022). Two distinct evolutionary conserved neural degeneration pathways characterized in a colonial chordate. *Proc. Natl. Acad. Sci. U. S. A.* 119, e2203032119. doi:10.1073/pnas.2203032119

Arendt, D., Musser, J. M., Baker, C. V. H., Bergman, A., Cepko, C., Erwin, D. H., et al. (2016). The origin and evolution of cell types. *Nat. Rev. Genet.* 17, 744–757. doi:10.1038/nrg.2016.127

Athira, A., Dondorp, D., Rudolf, J., Peytral, O., and Chatzigeorgiou, M. (2022). Comprehensive analysis of locomotion dynamics in the protochordate *Ciona intestinalis* reveals how neuromodulators flexibly shape its behavioral repertoire. *PLoS Biol.* 20, e3001744. doi:10.1371/journal.pbio.3001744

Baker, C. V., and Bronner-Fraser, M. (2001). Vertebrate cranial placodes I. Embryonic induction. *Dev. Biol.* 232, 1–61. doi:10.1006/dbio.2001.0156

Baker, C. V. H. (2019). "The development and evolution of lateral line electroreceptors: insights from comparative molecular approaches," in Electroreception: fundamental Insights from comparative approaches *springer* handbook of auditory research. Editors B. A. Carlson, J. A. Sisneros, A. N. Popper, and R. R. Fay (Cham: Springer International Publishing), 25–62. doi:10.1007/978-3-030-29105-1_2

Baker, C. V. H., Modrell, M. S., and Gillis, J. A. (2013). The evolution and development of vertebrate lateral line electroreceptors. *J. Exp. Biol.* 216, 2515–2522. doi:10.1242/jeb.082362

Basch, M. L., Brown, R. M., Jen, H.-I., and Groves, A. K. (2016). Where hearing starts: the development of the mammalian cochlea. *J. Anat.* 228, 233–254. doi:10.1111/joa. 12314

Baumeister, R., Liu, Y., and Ruvkun, G. (1996). Lineage-specific regulators couple cell lineage asymmetry to the transcription of the *Caenorhabditis elegans* POU gene unc-86 during neurogenesis. *Genes Dev.* 10, 1395–1410. doi:10.1101/gad.10.11.1395

Ben-Arie, N., Bellen, H. J., Armstrong, D. L., McCall, A. E., Gordadze, P. R., Guo, Q., et al. (1997). Math1 is essential for genesis of cerebellar granule neurons. *Nature* 390, 169–172. doi:10.1038/36579

Ben-Arie, N., Hassan, B. A., Bermingham, N. A., Malicki, D. M., Armstrong, D., Matzuk, M., et al. (2000). Functional conservation of atonal and Math1 in the CNS and PNS. *Development* 127, 1039–1048. doi:10.1242/dev.127.5.1039

Beurg, M., and Fettiplace, R. (2017). PIEZO2 as the anomalous mechanotransducer channel in auditory hair cells. *J. Physiol. (Lond)* 595, 7039–7048. doi:10.1113/JP274996

Birol, O., Ohyama, T., Edlund, R. K., Drakou, K., Georgiades, P., and Groves, A. K. (2016). The mouse Foxi3 transcription factor is necessary for the development of posterior placodes. *Dev. Biol.* 409, 139–151. doi:10.1016/j.ydbio.2015.09.022

Bollner, T., Holmberg, K., and Olsson, R. (1986). A rostral sensory mechanism in *Oikopleura dioica* (appendicularia). *Acta Zool.* 67, 235–241. doi:10.1111/j.1463-6395. 1986.tb00868.x

Bone, Q. (1959). Observations upon the nervous systems of pelagic tunicates. J. Cell Sci. S3-100, 167–181. doi:10.1242/jcs.s3-100.50.167

Bone, Q. (1998). The biology of pelagic tunicates (Oxford University PressOxford). doi:10.1093/oso/9780198540243.001.0001

Bone, Q., and Ryan, K. P. (1978). Cupular sense organs in ciona (tunicata: Ascidiacea). J. Zool. 186, 417–429. doi:10.1111/j.1469-7998.1978.tb03931.x

Boorman, C. J., and Shimeld, S. M. (2002). Pitx homeobox genes in Ciona and amphioxus show left-right asymmetry is a conserved chordate character and define the ascidian adenohypophysis. *Evol. Dev.* 4 (5), 354–365. doi:10.1046/j.1525-142X.2002.02021.x

Bostwick, M., Smith, E. L., Borba, C., Newman-Smith, E., Guleria, I., Kourakis, M. J., et al. (2020). Antagonistic inhibitory circuits integrate visual and gravitactic behaviors. *Curr. Biol.* 30, 600–609. doi:10.1016/j.cub.2019.12.017

Brown, R., and Groves, A. K. (2020). Hear, hear for notch: control of cell fates in the inner ear by notch signaling. *Biomolecules* 10, 370. doi:10.3390/biom10030370

Bucks, S. A., Cox, B. C., Vlosich, B. A., Manning, J. P., Nguyen, T. B., and Stone, J. S. (2017). Supporting cells remove and replace sensory receptor hair cells in a balance organ of adult mice. *eLife* 6, e18128. doi:10.7554/eLife.18128

Burighel, P., Caicci, F., and Manni, L. (2011). Hair cells in non-vertebrate models: lower chordates and molluscs. *Hear. Res.* 273, 14–24. doi:10.1016/j.heares.2010.03.087

Burighel, P., Caicci, F., Zaniolo, G., Gasparini, F., Degasperi, V., and Manni, L. (2008). Does hair cell differentiation predate the vertebrate appearance? *Brain Res. Bull.* 75, 331–334. doi:10.1016/j.brainresbull.2007.10.012

Burighel, P., Lane, N. J., Fabio, G., Stefano, T., Zaniolo, G., Carnevali, M. D. C., et al. (2003). Novel, secondary sensory cell organ in ascidians: in search of the ancestor of the vertebrate lateral line. *J. Comp. Neurol.* 461, 236–249. doi:10.1002/cne.10666

Cai, T., and Groves, A. K. (2015). The role of atonal factors in mechanosensory cell specification and function. *Mol. Neurobiol.* 52, 1315–1329. doi:10.1007/s12035-014-8925-0

Caicci, F., Burighel, P., and Manni, L. (2007). Hair cells in an ascidian (Tunicata) and their evolution in chordates. *Hear. Res.* 231, 63–72. doi:10.1016/j.heares.2007.05.007

Caicci, F., Degasperi, V., Gasparini, F., Zaniolo, G., Del Favero, M., Burighel, P., et al. (2010a). Variability of hair cells in the coronal organ of ascidians (Chordata, Tunicata). *Can. J. Zool.* 88, 567–578. doi:10.1139/z10-036

Caicci, F., Gasparini, F., Rigon, F., Zaniolo, G., Burighel, P., and Manni, L. (2013). The oral sensory structures of Thaliacea (Tunicata) and consideration of the evolution of hair cells in Chordata. *J. Comp. Neurol.* 521, 2756–2771. doi:10.1002/cne.23313

Caicci, F., Zaniolo, G., Burighel, P., Degasperi, V., Gasparini, F., and Manni, L. (2010b). Differentiation of papillae and rostral sensory neurons in the larva of the ascidian Botryllus schlosseri (Tunicata). *J. Comp. Neurol.* 518, 547–566. doi:10.1002/cne.22222

Cao, C., Lemaire, L. A., Wang, W., Yoon, P. H., Choi, Y. A., Parsons, L. R., et al. (2019). Comprehensive single-cell transcriptome lineages of a proto-vertebrate. *Nature* 571, 349–354. doi:10.1038/s41586-019-1385-y

Chen, J. S., Pedro, M. S., and Zeller, R. W. (2011). miR-124 function during *Ciona intestinalis* neuronal development includes extensive interaction with the Notch signaling pathway. *Development* 138, 4943–4953. doi:10.1242/dev.068049

Corey, D. P., García-Añoveros, J., Holt, J. R., Kwan, K. Y., Lin, S.-Y., Vollrath, M. A., et al. (2004). TRPA1 is a candidate for the mechanosensitive transduction channel of vertebrate hair cells. *Nature* 432, 723–730. doi:10.1038/nature03066

Cosgrove, D., and Zallocchi, M. (2014). Usher protein functions in hair cells and photoreceptors. Int. J. Biochem. Cell Biol. 46, 80-89. doi:10.1016/j.biocel.2013.11.001

Costa, A., Sanchez-Guardado, L., Juniat, S., Gale, J. E., Daudet, N., and Henrique, D. (2015). Generation of sensory hair cells by genetic programming with a combination of transcription factors. *Development* 142, 1948–1959. doi:10.1242/dev.119149

Coulcher, J. F., Roure, A., Chowdhury, R., Robert, M., Lescat, L., Bouin, A., et al. (2020). Conservation of peripheral nervous system formation mechanisms in divergent ascidian embryos. *Elife* 9, e59157. doi:10.7554/eLife.59157

Cunningham, C. L., Qiu, X., Wu, Z., Zhao, B., Peng, G., Kim, Y.-H., et al. (2020). TMIE defines pore and gating properties of the mechanotransduction channel of mammalian cochlear hair cells. *Neuron* 107, 126–143. doi:10.1016/j.neuron.2020.03.033

Cvekl, A., and Ashery-Padan, R. (2014). The cellular and molecular mechanisms of vertebrate lens development. *Development* 141, 4432–4447. doi:10.1242/dev.107953

DeBiasse, M. B., Colgan, W. N., Harris, L., Davidson, B., and Ryan, J. F. (2020). Inferring tunicate relationships and the evolution of the tunicate hox cluster with the genome of Corella inflata. *Genome Biol. Evol.* 12, 948–964. doi:10.1093/gbe/evaa060

Delsuc, F., Philippe, H., Tsagkogeorga, G., Simion, P., Tilak, M.-K., Turon, X., et al. (2018). A phylogenomic framework and timescale for comparative studies of tunicates. *BMC Biol.* 16, 39. doi:10.1186/s12915-018-0499-2

Durante, K. M. (1991). Larval behavior, settlement preference, and induction of metamorphosis in the temperate solitary ascidian Molgula citrina Alder and Hancock. *J. Exp. Mar. Biol. Ecol.* 145, 175–187. doi:10.1016/0022-0981(91)90174-U

Edlund, R. K., Ohyama, T., Kantarci, H., Riley, B. B., and Groves, A. K. (2014). Foxi transcription factors promote pharyngeal arch development by regulating formation of FGF signaling centers. *Dev. Biol.* 390, 1–13. doi:10.1016/j.ydbio.2014.03.004

Fedele, M. (1923). Le attivitá dinamiche ed i rapporti nervosi nella vita dei Dolioli. *Pubbl Staz Zool. Napoli* 4, 129–240.

Fritzsch, B., and Elliott, K. L. (2017). Gene, cell, and organ multiplication drives inner ear evolution. *Dev. Biol.* 431, 3–15. doi:10.1016/j.ydbio.2017.08.034

Frolenkov, G. I., Belyantseva, I. A., Friedman, T. B., and Griffith, A. J. (2004). Genetic insights into the morphogenesis of inner ear hair cells. *Nat. Rev. Genet.* 5, 489–498. doi:10.1038/nrg1377

Gans, C., and Northcutt, R. G. (1983). Neural crest and the origin of vertebrates: a new head. *Science* 220, 268–273. doi:10.1126/science.220.4594.268

Gasparini, F., Caicci, F., Rigon, F., Zaniolo, G., Burighel, P., and Manni, L. (2013a). Cytodifferentiation of hair cells during the development of a basal chordate. *Hear. Res.* 304, 188–199. doi:10.1016/j.heares.2013.07.006

Gasparini, F., Degasperi, V., Shimeld, S. M., Burighel, P., and Manni, L. (2013b). Evolutionary conservation of the placodal transcriptional network during sexual and asexual development in chordates. *Dev. Dyn.* 242, 752–766. doi:10.1002/dvdy.23957

Ge, J., Elferich, J., Goehring, A., Zhao, H., Schuck, P., and Gouaux, E. (2018). Structure of mouse protocadherin 15 of the stereocilia tip link in complex with LHFPL5. *eLife* 7, e38770. doi:10.7554/eLife.38770

Gillespie, P. G., and Müller, U. (2009). Mechanotransduction by hair cells: models, molecules, and mechanisms. *Cell* 139, 33-44. doi:10.1016/j.cell.2009.09.010

Gnedeva, K., and Hudspeth, A. J. (2015). SoxC transcription factors are essential for the development of the inner ear. *Proc. Natl. Acad. Sci. U. S. A.* 112, 14066–14071. doi:10.1073/pnas.1517371112

Graham, A., and Shimeld, S. M. (2013). The origin and evolution of the ectodermal placodes. J. Anat. 222 (1), 32–40. doi:10.1111/j.1469-7580.2012.01506.x

Grocott, T., Tambalo, M., and Streit, A. (2012). The peripheral sensory nervous system in the vertebrate head: a gene regulatory perspective. *Dev. Biol.* 370, 3–23. doi:10. 1016/j.ydbio.2012.06.028

Groves, A. K. (2010). The challenge of hair cell regeneration. Exp. Biol. Med. (Maywood) 235, 434-446. doi:10.1258/ebm.2009.009281

Groves, A. K., and Fekete, D. M. (2012). Shaping sound in space: the regulation of inner ear patterning. *Development* 139, 245-257. doi:10.1242/dev.067074

Groves, A. K., and LaBonne, C. (2014). Setting appropriate boundaries: fate, patterning and competence at the neural plate border. *Dev. Biol.* 389, 2–12. doi:10. 1016/j.ydbio.2013.11.027

Gu, R., Brown, R. M., Hsu, C.-W., Cai, T., Crowder, A. L., Piazza, V. G., et al. (2016). Lineage tracing of Sox2-expressing progenitor cells in the mouse inner ear reveals a broad contribution to non-sensory tissues and insights into the origin of the organ of Corti. *Dev. Biol.* 414, 72–84. doi:10.1016/j.ydbio.2016.03.027

Haeberle, H., Fujiwara, M., Chuang, J., Medina, M. M., Panditrao, M. V., Bechstedt, S., et al. (2004). Molecular profiling reveals synaptic release machinery in Merkel cells. *Proc. Natl. Acad. Sci. U. S. A.* 101, 14503–14508. doi:10.1073/pnas.0406308101

Hans, S., Liu, D., and Westerfield, M. (2004). Pax8 and Pax2a function synergistically in otic specification, downstream of the Foxi1 and Dlx3b transcription factors. *Development* 131, 5091–5102. doi:10.1242/dev.01346

Hasten, E., and Morrow, B. E. (2019). Tbx1 and Foxi3 genetically interact in the pharyngeal pouch endoderm in a mouse model for 22q11.2 deletion syndrome. *PLoS Genet.* 15, e1008301. doi:10.1371/journal.pgen.1008301

Holt, J. R., Tobin, M., Elferich, J., Gouaux, E., Ballesteros, A., Yan, Z., et al. (2021). Putting the pieces together: the hair cell transduction complex. *J. Assoc. Res. Otolaryngol.* 22, 601-608. doi:10.1007/s10162-021-00808-0

Horie, R., Hazbun, A., Chen, K., Cao, C., Levine, M., and Horie, T. (2018). Shared evolutionary origin of vertebrate neural crest and cranial placodes. *Nature* 560, 228–232. doi:10.1038/s41586-018-0385-7

Horie, T., Kusakabe, T., and Tsuda, M. (2008). Glutamatergic networks in the *Ciona intestinalis* larva. *J. Comp. Neurol.* 508, 249–263. doi:10.1002/cne.21678

Hoyer, J., Kolar, K., Athira, A., van den Burgh, M., Dondorp, D., Liang, Z., et al. (2024). Polymodal sensory perception drives settlement and metamorphosis of Ciona larvae. *Curr. Biol.* doi:10.1016/j.cub.2024.01.041

Hozumi, A., Matsunobu, S., Mita, K., Treen, N., Sugihara, T., Horie, T., et al. (2020). GABA-induced GnRH release triggers chordate metamorphosis. *Curr. Biol.* 30, 1555–1561. doi:10.1016/j.cub.2020.02.003

Hudson, C., and Yasuo, H. (2005). Patterning across the ascidian neural plate by lateral Nodal signalling sources. *Development* 132, 1199–1210. doi:10.1242/dev.01688

Imai, J. H., and Meinertzhagen, I. A. (2007). Neurons of the ascidian larval nervous system in *Ciona intestinalis*: II. Peripheral nervous system. *J. Comp. Neurol.* 501, 335–352. doi:10.1002/cne.21247

Imai, K. S., Hikawa, H., Kobayashi, K., and Satou, Y. (2017). Tfap2 and Sox1/2/ 3 cooperatively specify ectodermal fates in ascidian embryos. *Development* 144, 33–37. doi:10.1242/dev.142109

Imai, K. S., Hino, K., Yagi, K., Satoh, N., and Satou, Y. (2004). Gene expression profiles of transcription factors and signaling molecules in the ascidian embryo: towards a comprehensive understanding of gene networks. *Development* 131, 4047–4058. doi:10.1242/dev.01270

Irvine, S. Q., Cangiano, M. C., Millette, B. J., and Gutter, E. S. (2007). Non-overlapping expression patterns of the clustered Dll-A/B genes in the ascidian Ciona Intestinalis. *J. Exp. Zool Part B Mol. Dev. Evol.* 308 (4), 428–441. doi:10.1002/jez.b.21169

Irvine, S. Q., Fonseca, V. C., Zompa, M. A., and Antony, R. (2008). Cis-regulatory organization of the Pax6 gene in the ascidian *Ciona intestinalis*. *Dev. Biol.* 317, 649–659. doi:10.1016/j.ydbio.2008.01.036

Iyer, A. A., and Groves, A. K. (2021). Transcription factor reprogramming in the inner ear: turning on cell fate switches to regenerate sensory hair cells. *Front. Cell. Neurosci.* 15, 660748. doi:10.3389/fncel.2021.660748

Iyer, A. A., Hosamani, I., Nguyen, J. D., Cai, T., Singh, S., McGovern, M. M., et al. (2022). Cellular reprogramming with ATOH1, GFI1, and POU4F3 implicate epigenetic changes and cell-cell signaling as obstacles to hair cell regeneration in mature mammals. *eLife* 11, e79712. doi:10.7554/eLife.79712

Jarman, A. P., Grau, Y., Jan, L. Y., and Jan, Y. N. (1993). Atonal is a proneural gene that directs chordotonal organ formation in the Drosophila peripheral nervous system. *Cell* 73, 1307–1321. doi:10.1016/0092-8674(93)90358-w

Jarman, A. P., and Groves, A. K. (2013). The role of Atonal transcription factors in the development of mechanosensitive cells. *Semin. Cell Dev. Biol.* 24, 438–447. doi:10.1016/j.semcdb.2013.03.010

Jarman, A. P., Sun, Y., Jan, L. Y., and Jan, Y. N. (1995). Role of the proneural gene, atonal, in formation of Drosophila chordotonal organs and photoreceptors. *Development* 121, 2019–2030. doi:10.1242/dev.121.7.2019

Jeffery, W. R., Strickler, A. G., and Yamamoto, Y. (2004). Migratory neural crest-like cells form body pigmentation in a urochordate embryo. *Nature* 431, 696–699. doi:10. 1038/nature02975

Jen, H.-I., Singh, S., Tao, L., Maunsell, H. R., Segil, N., and Groves, A. K. (2022). GFI1 regulates hair cell differentiation by acting as an off-DNA transcriptional coactivator of ATOH1, and a DNA-binding repressor. *Sci. Rep.* 12, 7793. doi:10.1038/ s41598-022-11931-0

Jiang, D., Tresser, J. W., Horie, T., Tsuda, M., and Smith, W. C. (2005). Pigmentation in the sensory organs of the ascidian larva is essential for normal behavior. *J. Exp. Biol.* 208, 433–438. doi:10.1242/jeb.01420

Johnson, C. J., Kulkarni, A., Buxton, W. J., Hui, T. Y., Kayastha, A., Khoja, A. A., et al. (2023a). Using CRISPR/Cas9 to identify genes required for mechanosensory neuron development and function. *Biol. Open* 12, bio060002. doi:10.1242/bio. 060002

Johnson, C. J., Razy-Krajka, F., Zeng, F., Piekarz, K. M., Biliya, S., Rothbächer, U., et al. (2023b). Specification of distinct cell types in a sensory-adhesive organ for metamorphosis in the *Ciona* larva. *BioRxiv*. doi:10.1101/2023.05.02.539060

Khatri, S. B., Edlund, R. K., and Groves, A. K. (2014). Foxi3 is necessary for the induction of the chick otic placode in response to FGF signaling. *Dev. Biol.* 391, 158–169. doi:10.1016/j.ydbio.2014.04.014

Khatri, S. B., and Groves, A. K. (2013). Expression of the Foxi2 and Foxi3 transcription factors during development of chicken sensory placodes and pharyngeal arches. *Gene Expr. Patterns* 13, 38–42. doi:10.1016/j.gep.2012.10.001

Kindt, K. S., Finch, G., and Nicolson, T. (2012). Kinocilia mediate mechanosensitivity in developing zebrafish hair cells. *Dev. Cell* 23, 329–341. doi:10.1016/j.devcel.2012.05.022

Kniss, J. S., Jiang, L., and Piotrowski, T. (2016). Insights into sensory hair cell regeneration from the zebrafish lateral line. *Curr. Opin. Genet. Dev.* 40, 32–40. doi:10. 1016/j.gde.2016.05.012

Kocot, K. M., Tassia, M. G., Halanych, K. M., and Swalla, B. J. (2018). Phylogenomics offers resolution of major tunicate relationships. *Mol. Phylogenet. Evol.* 121, 166–173. doi:10.1016/j.ympev.2018.01.005

Konno, A., Kaizu, M., Hotta, K., Horie, T., Sasakura, Y., Ikeo, K., et al. (2010). Distribution and structural diversity of cilia in tadpole larvae of the ascidian *Ciona intestinalis*. *Dev. Biol.* 337, 42–62. doi:10.1016/j.ydbio.2009.10.012

Kourakis, M. J., Borba, C., Zhang, A., Newman-Smith, E., Salas, P., Manjunath, B., et al. (2019). Parallel visual circuitry in a basal chordate. *eLife* 8, e44753. doi:10.7554/ eLife.44753

Koyama, H. (2008). Sensory cells associated with the tentacular tunic of the ascidian Polyandrocarpa misakiensis (Tunicata: Ascidiacea). *Zool. Sci.* 25, 919–930. doi:10.2108/ zsj.25.919

Kozmik, Z. (2008). The role of Pax genes in eye evolution. Brain Res. Bull. 75, 335–339. doi:10.1016/j.brainresbull.2007.10.046

Kwan, K. Y., Allchorne, A. J., Vollrath, M. A., Christensen, A. P., Zhang, D.-S., Woolf, C. J., et al. (2006). TRPA1 contributes to cold, mechanical, and chemical nociception but is not essential for hair-cell transduction. *Neuron* 50, 277–289. doi:10.1016/j.neuron.2006.03.042

Lai, H. C., Klisch, T. J., Roberts, R., Zoghbi, H. Y., and Johnson, J. E. (2011). *In vivo* neuronal subtype-specific targets of Atoh1 (Math1) in dorsal spinal cord. *J. Neurosci.* 31, 10859–10871. doi:10.1523/JNEUROSCI.0445-11.2011

Le, M.-L. V., Müller, L.-M., and Stach, T. (2023). The oral sensory organs in Bathochordaeus stygius (Tunicata Appendicularia) are unique in structure and homologous to the coronal organ. *Res. Sq.* doi:10.21203/rs.3.rs-3167810/v1

Lee, M.-H., and Salvaterra, P. M. (2002). Abnormal chemosensory jump 6 is a positive transcriptional regulator of the cholinergic gene locus in Drosophila olfactory neurons. *J. Neurosci.* 22, 5291–5299. doi:10.1523/JNEUROSCI.22-13-05291.2002

Lesko, M. H., Driskell, R. R., Kretzschmar, K., Goldie, S. J., and Watt, F. M. (2013). Sox2 modulates the function of two distinct cell lineages in mouse skin. *Dev. Biol.* 382, 15–26. doi:10.1016/j.ydbio.2013.08.004

Li, S., Fan, T., Li, C., Wang, Y., Li, J., and Liu, Z. (2022). Fate-mapping analysis of cochlear cells expressing Atoh1 mRNA via a new Atoh13*HA-P2A-Cre knockin mouse strain. *Dev. Dyn.* 251, 1156–1174. doi:10.1002/dvdy.453

Li, T., Bellen, H. J., and Groves, A. K. (2018). Using Drosophila to study mechanisms of hereditary hearing loss. *Dis. Model. Mech.* 11, dmm031492. doi:10.1242/dmm.031492

Liu, B., and Satou, Y. (2019). Foxg specifies sensory neurons in the anterior neural plate border of the ascidian embryo. *Nat. Commun.* 10, 4911. doi:10.1038/s41467-019-12839-6

Lu, T.-M., Luo, Y.-J., and Yu, J.-K. (2012). BMP and Delta/Notch signaling control the development of amphioxus epidermal sensory neurons: insights into the evolution of the peripheral sensory system. *Development* 139, 2020–2030. doi:10.1242/dev.073833

Lumpkin, E. A., Collisson, T., Parab, P., Omer-Abdalla, A., Haeberle, H., Chen, P., et al. (2003). Math1-driven GFP expression in the developing nervous system of transgenic mice. *Gene Expr. Patterns* 3, 389–395. doi:10.1016/s1567-133x(03)00089-9 Mackie, G. O., and Bone, Q. (1978). Luminescence and associated effector activity in pyrosoma (tunicata: pyrosomida). *Proc. R. Soc. B Biol. Sci.* 202, 483–495. doi:10.1098/ rspb.1978.0081

Mackie, G. O., and Burighel, P. (2005). The nervous system in adult tunicates: current research directions. *Can. J. Zool.* 83, 151–183. doi:10.1139/z04-177

Mackie, G. O., Burighel, P., Caicci, F., and Manni, L. (2006). Innervation of ascidian siphons and their responses to stimulation. *Can. J. Zool.* 84, 1146–1162. doi:10.1139/z06-106

Mackie, G. O., and Singla, C. L. (2003). The capsular organ of Chelyosoma productum (Ascidiacea: corellidae): a new tunicate hydrodynamic sense organ. *Brain Behav. Evol.* 61, 45–58. doi:10.1159/000068878

Mackie, G. O., and Singla, C. L. (2005). Cupular organs in two species of Corella (tunicata: Ascidiacea). *Invertebr. Biol.* 123, 269–281. doi:10.1111/j.1744-7410.2004. tb00161.x

Maksimovic, S., Nakatani, M., Baba, Y., Nelson, A. M., Marshall, K. L., Wellnitz, S. A., et al. (2014). Epidermal Merkel cells are mechanosensory cells that tune mammalian touch receptors. *Nature* 509, 617–621. doi:10.1038/nature13250

Mann, Z. F., Gálvez, H., Pedreno, D., Chen, Z., Chrysostomou, E., Żak, M., et al. (2017). Shaping of inner ear sensory organs through antagonistic interactions between Notch signalling and Lmx1a. *eLife* 6, e33323. doi:10.7554/eLife.33323

Manni, L., Agnoletto, A., Zaniolo, G., and Burighel, P. (2005). Stomodeal and neurohypophysial placodes in *Ciona intestinalis*: insights into the origin of the pituitary gland. *J. Exp. Zool. B Mol. Dev. Evol.* 304, 324–339. doi:10.1002/jez.b.21039

Manni, L., Anselmi, C., Burighel, P., Martini, M., and Gasparini, F. (2018). Differentiation and induced sensorial alteration of the coronal organ in the asexual life of a tunicate. *Integr. Comp. Biol.* 58, 317–328. doi:10.1093/icb/icy044

Manni, L., Caicci, F., Anselmi, C., Vanni, V., Mercurio, S., and Pennati, R. (2021). Morphological study and 3D reconstruction of the larva of the ascidian Halocynthia roretzi. *JMSE* 10, 11. doi:10.3390/jmse10010011

Manni, L., Caicci, F., Gasparini, F., Zaniolo, G., and Burighel, P. (2004). Hair cells in ascidians and the evolution of lateral line placodes. *Evol. Dev.* 6, 379–381. doi:10.1111/j. 1525-142X.2004.04046.x

Manni, L., Lane, N. J., Burighel, P., and Zaniolo, G. (2001). Are neural crest and placodes exclusive to vertebrates? *Evol. Dev.* 3, 297–298. doi:10.1046/j.1525-142x.2001. 01040.x

Manni, L., Lane, N. J., Sorrentino, M., Zaniolo, G., and Burighel, P. (1999). Mechanism of neurogenesis during the embryonic development of a tunicate. J. Comp. Neurol. 412, 527-541. doi:10.1002/(sici)1096-9861(19990927)412:3<527:: aid-cne11>3.0.co;2-u

Manni, L., Mackie, G. O., Caicci, F., Zaniolo, G., and Burighel, P. (2006). Coronal organ of ascidians and the evolutionary significance of secondary sensory cells in chordates. *J. Comp. Neurol.* 495, 363–373. doi:10.1002/cne.20867

Manni, L., and Pennati, R. (2015). "Tunicata," in *Structure and evolution of invertebrate nervous systems* (Oxford University Press), 699-718. Available at: https://doi.org/10.1093/acprof:oso/9780199682201.003.0031 (Accessed May 25, 2021).

Maricich, S. M., Wellnitz, S. A., Nelson, A. M., Lesniak, D. R., Gerling, G. J., Lumpkin, E. A., et al. (2009). Merkel cells are essential for light-touch responses. *Science* 324, 1580–1582. doi:10.1126/science.1172890

Mazet, F., Hutt, J. A., Millard, J., and Shimeld, S. M. (2003). Pax gene expression in the developing central nervous system of *Ciona intestinalis*. *Gene Expr. Patterns* 3, 743–745. doi:10.1016/s1567-133x(03)00137-6

Mazet, F., and Shimeld, S. M. (2005). Molecular evidence from ascidians for the evolutionary origin of vertebrate cranial sensory placodes. *J. Exp. Zool. B Mol. Dev. Evol.* 304, 340–346. doi:10.1002/jez.b.21054

Menendez, L., Trecek, T., Gopalakrishnan, S., Tao, L., Markowitz, A. L., Yu, H. V., et al. (2020). Generation of inner ear hair cells by direct lineage conversion of primary somatic cells. *eLife* 9, e55249. doi:10.7554/eLife.55249

Modrell, M. S., Lyne, M., Carr, A. R., Zakon, H. H., Buckley, D., Campbell, A. S., et al. (2017). Insights into electrosensory organ development, physiology and evolution from a lateral line-enriched transcriptome. *eLife* 6, e24197. doi:10.7554/eLife.24197

Nakatani, M., Maksimovic, S., Baba, Y., and Lumpkin, E. A. (2015). Mechanotransduction in epidermal Merkel cells. *Pflugers Arch.* 467, 101–108. doi:10.1007/s00424-014-1569-0

Nakayama-Ishimura, A., Chambon, J., Horie, T., Satoh, N., and Sasakura, Y. (2009). Delineating metamorphic pathways in the ascidian *Ciona intestinalis. Dev. Biol.* 326, 357–367. doi:10.1016/j.ydbio.2008.11.026

Neves, J., Vachkov, I., and Giraldez, F. (2013). Sox2 regulation of hair cell development: incoherence makes sense. *Hear. Res.* 297, 20–29. doi:10.1016/j.heares.2012.11.003

Nissen, R. M., Yan, J., Amsterdam, A., Hopkins, N., and Burgess, S. M. (2003). Zebrafish foxi one modulates cellular responses to Fgf signaling required for the integrity of ear and jaw patterning. *Development* 130, 2543–2554. doi:10.1242/dev.00455

Nolo, R., Abbott, L. A., and Bellen, H. J. (2000). Senseless, a Zn finger transcription factor, is necessary and sufficient for sensory organ development in Drosophila. *Cell* 102, 349–362. doi:10.1016/s0092-8674(00)00040-4

Ohtsuka, Y., Matsumoto, J., Katsuyama, Y., and Okamura, Y. (2014). Nodal signaling regulates specification of ascidian peripheral neurons through control of the BMP signal. *Development* 141, 3889–3899. doi:10.1242/dev.110213

Ohyama, T., and Groves, A. K. (2004). Expression of mouse Foxi class genes in early craniofacial development. *Dev. Dyn.* 231, 640–646. doi:10.1002/dvdy.20160

Okawa, N., Shimai, K., Ohnishi, K., Ohkura, M., Nakai, J., Horie, T., et al. (2020). Cellular identity and Ca2+ signaling activity of the non-reproductive GnRH system in the *Ciona intestinalis* type A (Ciona robusta) larva. *Sci. Rep.* 10, 18590. doi:10.1038/ s41598-020-75344-7

Ozment, E., Tamvacakis, A. N., Zhou, J., Rosiles-Loeza, P. Y., Escobar-Hernandez, E. E., Fernandez-Valverde, S. L., et al. (2021). Cnidarian hair cell development illuminates an ancient role for the class IV POU transcription factor in defining mechanoreceptor identity. *eLife* 10, e74336. doi:10.7554/eLife.74336

Papadogiannis, V., Pennati, A., Parker, H. J., Rothbächer, U., Patthey, C., Bronner, M. E., et al. (2022). Hmx gene conservation identifies the origin of vertebrate cranial ganglia. *Nature* 605 (7911), 701–705. doi:10.1038/s41586-022-04742-w

Park, C., Ji, H.-M., Kim, S.-J., Kil, S.-H., Lee, J. N., Kwak, S., et al. (2017). Fenofibrate exerts protective effects against gentamicin-induced toxicity in cochlear hair cells by activating antioxidant enzymes. *Int. J. Mol. Med.* 39, 960–968. doi:10.3892/ijmm.2017. 2916

Pasini, A., Amiel, A., Rothbächer, U., Roure, A., Lemaire, P., and Darras, S. (2006). Formation of the ascidian epidermal sensory neurons: insights into the origin of the chordate peripheral nervous system. *PLoS Biol.* 4, e225. doi:10.1371/journal.pbio. 0040225

Pasini, A., Manenti, R., Rothbächer, U., and Lemaire, P. (2012). Antagonizing retinoic acid and FGF/MAPK pathways control posterior body patterning in the invertebrate chordate *Ciona intestinalis*. *PLoS ONE* 7, e46193. doi:10.1371/journal.pone.0046193

Patthey, C., Schlosser, G., and Shimeld, S. M. (2014). The evolutionary history of vertebrate cranial placodes--I: cell type evolution. *Dev. Biol.* 389, 82–97. doi:10.1016/j. ydbio.2014.01.017

Perdigoto, C. N., Bardot, E. S., Valdes, V. J., Santoriello, F. J., and Ezhkova, E. (2014). Embryonic maturation of epidermal Merkel cells is controlled by a redundant transcription factor network. *Development* 141, 4690–4696. doi:10.1242/dev.112169

Piotrowski, T., and Baker, C. V. H. (2014). The development of lateral line placodes: taking a broader view. *Dev. Biol.* 389, 68–81. doi:10.1016/j.ydbio.2014.02.016

Pla, P., and Monsoro-Burq, A. H. (2018). The neural border: induction, specification and maturation of the territory that generates neural crest cells. *Dev. Biol.* 444 (Suppl. 1), S36–S46. doi:10.1016/j.ydbio.2018.05.018

Poncelet, G., Parolini, L., and Shimeld, S. M. (2022). A microfluidic device for controlled exposure of transgenic *Ciona intestinalis* larvae to chemical stimuli demonstrates they can respond to carbon dioxide. *BioRxiv*. doi:10.1101/2022.08.15. 492342

Popsuj, S., and Stolfi, A. (2021). Ebf activates expression of a cholinergic locus in a multipolar motor ganglion interneuron subtype in ciona. *Front. Neurosci.* 15, 784649. doi:10.3389/fnins.2021.784649

Qiu, X., and Müller, U. (2018). Mechanically gated ion channels in mammalian hair cells. *Front. Cell. Neurosci.* 12, 100. doi:10.3389/fncel.2018.00100

Rae Flores, A., and Faulkes, Z. (2008). Texture preferences of ascidian tadpole larvae during settlement. *Mar. Freshw. Behav. Physiol.* 41, 155–159. doi:10.1080/10236240802360914

Raft, S., and Groves, A. K. (2015). Segregating neural and mechanosensory fates in the developing ear: patterning, signaling, and transcriptional control. *Cell Tissue Res.* 359, 315–332. doi:10.1007/s00441-014-1917-6

Richardson, G. P., and Petit, C. (2019). Hair-bundle links: genetics as the gateway to function. *Cold Spring Harb. Perspect. Med.* 9, a033142. doi:10.1101/cshperspect.a033142

Rigon, F., Gasparini, F., Shimeld, S. M., Candiani, S., and Manni, L. (2018). Developmental signature, synaptic connectivity and neurotransmission are conserved between vertebrate hair cells and tunicate coronal cells. *J. Comp. Neurol.* 526, 957–971. doi:10.1002/cne.24382

Rigon, F., Stach, T., Caicci, F., Gasparini, F., Burighel, P., and Manni, L. (2013). Evolutionary diversification of secondary mechanoreceptor cells in tunicata. *BMC Evol. Biol.* 13, 112. doi:10.1186/1471-2148-13-112

Roure, A., Chowdhury, R., and Darras, S. (2023). Regulation of anterior neurectoderm specification and differentiation by BMP signaling in ascidians. *Development* 150, dev201575. doi:10.1242/dev.201575

Ryan, K., Lu, Z., and Meinertzhagen, I. A. (2016). The CNS connectome of a tadpole larva of *Ciona intestinalis* (L.) highlights sidedness in the brain of a chordate sibling. *eLife* 5, e16962. doi:10.7554/eLife.16962

Ryan, K., Lu, Z., and Meinertzhagen, I. A. (2017). Circuit homology between decussating pathways in the ciona larval CNS and the vertebrate startle-response pathway. *Curr. Biol.* 27, 721–728. doi:10.1016/j.cub.2017.01.026

Ryan, K., Lu, Z., and Meinertzhagen, I. A. (2018). The peripheral nervous system of the ascidian tadpole larva: types of neurons and their synaptic networks. *J. Comp. Neurol.* 526, 583–608. doi:10.1002/cne.24353

Sakamoto, A., Hozumi, A., Shiraishi, A., Satake, H., Horie, T., and Sasakura, Y. (2022). The TRP channel PKD2 is involved in sensing the mechanical stimulus of adhesion for initiating metamorphosis in the chordate Ciona. *Dev. Growth Differ.* 64, 395–408. doi:10.1111/dgd.12801

Sakurai, D., Goda, M., Kohmura, Y., Horie, T., Iwamoto, H., Ohtsuki, H., et al. (2004). The role of pigment cells in the brain of ascidian larva. *J. Comp. Neurol.* 475, 70–82. doi:10.1002/cne.20142

Sato, S., Ikeda, K., Shioi, G., Ochi, H., Ogino, H., Yajima, H., et al. (2010). Conserved expression of mouse Six1 in the pre-placodal region (PPR) and identification of an enhancer for the rostral PPR. *Dev. Biol.* 344, 158–171. doi:10.1016/j.ydbio.2010.04.029

Schlosser, G. (2014). Early embryonic specification of vertebrate cranial placodes. Wiley Interdiscip. Rev. Dev. Biol. 3, 349–363. doi:10.1002/wdev.142

Schlosser, G. (2021). Evolutionary origin of sensory and neurosecretory cell types: vertebrate cranial placodes, volume 2. 1st Edn. Boca Raton: CRC Press. doi:10.1201/9781003160625

Schwander, M., Kachar, B., and Müller, U. (2010). Review series: the cell biology of hearing. J. Cell Biol. 190, 9–20. doi:10.1083/jcb.201001138

Shimeld, S. M., Purkiss, A. G., Dirks, R. P. H., Bateman, O. A., Slingsby, C., and Lubsen, N. H. (2005). Urochordate betagamma-crystallin and the evolutionary origin of the vertebrate eye lens. *Curr. Biol.* 15, 1684–1689. doi:10.1016/j.cub.2005.08.046

Sidi, S., Friedrich, R. W., and Nicolson, T. (2003). NompC TRP channel required for vertebrate sensory hair cell mechanotransduction. *Science* 301, 96–99. doi:10.1126/science.1084370

Singh, S., and Groves, A. K. (2016). The molecular basis of craniofacial placode development. Wiley Interdiscip. Rev. Dev. Biol. 5, 363-376. doi:10.1002/wdev.226

Solomon, K. S., Kudoh, T., Dawid, I. B., and Fritz, A. (2003a). Zebrafish foxi1 mediates otic placode formation and jaw development. *Development* 130, 929–940. doi:10.1242/ dev.00308

Solomon, K. S., Logsdon, J. M., and Fritz, A. (2003b). Expression and phylogenetic analyses of three zebrafish FoxI class genes. *Dev. Dyn.* 228, 301–307. doi:10.1002/dvdy.10373

Sorrentino, M., Manni, L., Lane, N. J., and Burighel, P. (2001). Evolution of cerebral vesicles and their sensory organs in an ascidian larva. *Acta Zool.* 81, 243–258. doi:10. 1046/j.1463-6395.2000.00054.x

Stolfi, A., Ryan, K., Meinertzhagen, I. A., and Christiaen, L. (2015). Migratory neuronal progenitors arise from the neural plate borders in tunicates. *Nature* 527, 371–374. doi:10.1038/nature15758

Stone, J. S., and Cotanche, D. A. (2007). Hair cell regeneration in the avian auditory epithelium. *Int. J. Dev. Biol.* 51, 633–647. doi:10.1387/ijdb.072408js

Streit, A. (2004). Early development of the cranial sensory nervous system: from a common field to individual placodes. *Dev. Biol.* 276, 1–15. doi:10.1016/j.ydbio.2004.08.037

Streit, A. (2007). The preplacodal region: an ectodermal domain with multipotential progenitors that contribute to sense organs and cranial sensory ganglia. *Int. J. Dev. Biol.* 51, 447–461. doi:10.1387/ijdb.072327as

Streit, A. (2018). Specification of sensory placode progenitors: signals and transcription factor networks. *Int. J. Dev. Biol.* 62, 195–205. doi:10.1387/ijdb.170298as

Takamura, K. (1998). Nervous network in larvae of the ascidian *Ciona intestinalis*. *Dev. Genes Evol.* 208, 1–8. doi:10.1007/s004270050147

Tang, W., Chen, J. S., and Zeller, R. W. (2013). Transcriptional regulation of the peripheral nervous system in *Ciona intestinalis. Dev. Biol.* 378, 183–193. doi:10.1016/j. ydbio.2013.03.016

Terakubo, H. Q., Nakajima, Y., Sasakura, Y., Horie, T., Konno, A., Takahashi, H., et al. (2010). Network structure of projections extending from peripheral neurons in the tunic of ascidian larva. *Dev. Dyn.* 239, 2278–2287. doi:10.1002/dvdy.22361

Thawani, A., and Groves, A. K. (2020). Building the border: development of the chordate neural plate border region and its derivatives. *Front. Physiol.* 11, 608880. doi:10.3389/fphys.2020.608880

Thompson, S. H., Anselmi, C., Ishizuka, K. J., Palmeri, K. J., and Voskoboynik, A. (2022). Contributions from both the brain and the vascular network guide behavior in the colonial tunicate Botryllus schlosseri. *J. Exp. Biol.* 225, jeb244491. doi:10.1242/jeb.244491

Torrence, S. A. (1986). Sensory endings of the ascidian static organ (Chordata, Ascidiacea). Zoomorphology 106, 61–66. doi:10.1007/BF00312108

Torrence, S. A., and Cloney, R. A. (1982). Nervous system of ascidian larvae: caudal primary sensory neurons. *Zoomorphology* 99, 103–115. doi:10.1007/BF00310303

Varello, R., Asnicar, D., Boaga, J., and Cima, F. (2023). Behavioural responses to ultrasound antifouling systems by adult solitary ascidians. *JMSE* 11, 1115. doi:10.3390/jmse11061115

Vélez-Ortega, A. C., and Frolenkov, G. I. (2019). Building and repairing the stereocilia cytoskeleton in mammalian auditory hair cells. *Hear. Res.* 376, 47–57. doi:10.1016/j. heares.2018.12.012

Vollrath, M. A., Kwan, K. Y., and Corey, D. P. (2007). The micromachinery of mechanotransduction in hair cells. *Annu. Rev. Neurosci.* 30, 339–365. doi:10.1146/ annurev.neuro.29.051605.112917

Vorontsova, M. N., Nezlin, L. P., and Meinertzhagen, I. A. (1997). Nervous system of the larva of the ascidian *Molgula citrina* (alder and hancock, 1848). *Acta Zool.* 78, 177–185. doi:10.1111/j.1463-6395.1997.tb01004.x

Wakai, M. K., Nakamura, M. J., Sawai, S., Hotta, K., and Oka, K. (2021). Two-Round Ca2+ transient in papillae by mechanical stimulation induces metamorphosis in the ascidian *Ciona intestinalis* type A. *Proc. Biol. Sci.* 288, 20203207. doi:10.1098/rspb.2020. 3207

Waki, K., Imai, K. S., and Satou, Y. (2015). Genetic pathways for differentiation of the peripheral nervous system in ascidians. *Nat. Commun.* 6, 8719. doi:10.1038/ ncomms9719

Wang, D., and Zhou, J. (2021). The kinocilia of cochlear hair cells: structures, functions, and diseases. *Front. Cell Dev. Biol.* 9, 715037. doi:10.3389/fcell.2021. 715037

Wang, V. Y., Hassan, B. A., Bellen, H. J., and Zoghbi, H. Y. (2002). Drosophila atonal fully rescues the phenotype of Math1 null mice: new functions evolve in new cellular contexts. *Curr. Biol.* 12, 1611–1616. doi:10.1016/s0960-9822(02)01144-2

Wang, X., Llamas, J., Trecek, T., Shi, T., Tao, L., Makmura, W., et al. (2023). SoxC transcription factors shape the epigenetic landscape to establish competence for sensory differentiation in the mammalian organ of Corti. *Proc. Natl. Acad. Sci. U. S. A.* 120, e2301301120. doi:10.1073/pnas.2301301120

Whatley, M., Francis, A., Ng, Z. Y., Khoh, X. E., Atlas, M. D., Dilley, R. J., et al. (2020). Usher syndrome: genetics and molecular links of hearing loss and directions for therapy. *Front. Genet.* 11, 565216. doi:10.3389/fgene.2020.565216

Woo, S.-H., Ranade, S., Weyer, A. D., Dubin, A. E., Baba, Y., Qiu, Z., et al. (2014). Piezo2 is required for Merkel-cell mechanotransduction. *Nature* 509, 622–626. doi:10. 1038/nature13251

Wu, D. K., and Kelley, M. W. (2012). Molecular mechanisms of inner ear development. *Cold Spring Harb. Perspect. Biol.* 4, a008409. doi:10.1101/cshperspect. a008409

Wu, S.-R., Butts, J. C., Caudill, M. S., Revelli, J.-P., Dhindsa, R. S., Durham, M. A., et al. (2023). Atoh1 drives the heterogeneity of the pontine nuclei neurons and promotes their differentiation. *Sci. Adv.* 9, eadg1671. doi:10.1126/sciadv.adg1671

Wu, X., Indzhykulian, A. A., Niksch, P. D., Webber, R. M., Garcia-Gonzalez, M., Watnick, T., et al. (2016). Hair-cell mechanotransduction persists in TRP channel knockout mice. *PLoS ONE* 11, e0155577. doi:10.1371/journal.pone.0155577

Wu, Z., Grillet, N., Zhao, B., Cunningham, C., Harkins-Perry, S., Coste, B., et al. (2017). Mechanosensory hair cells express two molecularly distinct mechanotransduction channels. *Nat. Neurosci.* 20, 24–33. doi:10.1038/nn.4449

Xiong, W., Grillet, N., Elledge, H. M., Wagner, T. F. J., Zhao, B., Johnson, K. R., et al. (2012). TMHS is an integral component of the mechanotransduction machinery of cochlear hair cells. *Cell* 151, 1283–1295. doi:10.1016/j.cell.2012.10.041

Yang, H., Xie, X., Deng, M., Chen, X., and Gan, L. (2010). Generation and characterization of Atoh1-Cre knock-in mouse line. *Genesis* 48, 407–413. doi:10.1002/dvg.20633

Yokoyama, T. D., Hotta, K., and Oka, K. (2014). Comprehensive morphological analysis of individual peripheral neuron dendritic arbors in ascidian larvae using the photoconvertible protein Kaede. *Dev. Dyn.* 243, 1362–1373. doi:10.1002/dvdy.24169

Yu, H. V., Tao, L., Llamas, J., Wang, X., Nguyen, J. D., Trecek, T., et al. (2021). POU4F3 pioneer activity enables ATOH1 to drive diverse mechanoreceptor differentiation through a feed-forward epigenetic mechanism. *Proc. Natl. Acad. Sci.* U. S. A. 118, e2105137118. doi:10.1073/pnas.2105137118

Żak, M., and Daudet, N. (2021). A gradient of Wnt activity positions the neurosensory domains of the inner ear. *eLife* 10, e59540. doi:10.7554/eLife.59540

Zeng, F., Wunderer, J., Salvenmoser, W., Hess, M. W., Ladurner, P., and Rothbächer, U. (2019). Papillae revisited and the nature of the adhesive secreting collocytes. *Dev. Biol.* 448, 183–198. doi:10.1016/j.ydbio.2018.11.012

Zhao, B., Wu, Z., Grillet, N., Yan, L., Xiong, W., Harkins-Perry, S., et al. (2014). TMIE is an essential component of the mechanotransduction machinery of cochlear hair cells. *Neuron* 84, 954–967. doi:10.1016/j.neuron.2014.10.041

Zhao, D., Chen, S., Horie, T., Gao, Y., Bao, H., and Liu, X. (2020). Comparison of differentiation gene batteries for migratory mechanosensory neurons across bilaterians. *Evol. Dev.* 22, 438–450. doi:10.1111/ede.12331