

Developmental Biology

Transcriptomic profiling of tissue environments critical for post-embryonic patterning and morphogenesis of zebrafish skin

Reviewed Preprint

Revised by authors after peer review.

About eLife's process

Reviewed preprint version 3

August 3, 2023 (this version)

Reviewed preprint version 2

June 30, 2023

Reviewed preprint version 1

April 25, 2023

Sent for peer review

February 23, 2023

Posted to bioRxiv

February 4, 2023

Andrew J. Aman, Lauren M. Saunders, August A. Carr, Sanjay R. Srivatsan, Colten D. Eberhard, Blake Carrington, Dawn Watkins-Chow, William J. Pavan, Cole Trapnell, David M. Parichy

Department of Biology, University of Virginia, Charlottesville, VA • Department of Genome Sciences, University of Washington, Seattle, WA • National Human Genome Research Institute, National Institutes of Health, Bethesda, MD • Department of Cell Biology, University of Virginia, Charlottesville, VA

 (https://en.wikipedia.org/wiki/Open_access)

 (<https://creativecommons.org/licenses/by/4.0/>)

Abstract

Pigment patterns and skin appendages are prominent features of vertebrate skin. In zebrafish, regularly patterned pigment stripes and an array of calcified scales form simultaneously in the skin during post-embryonic development. Understanding mechanisms that regulate stripe patterning and scale morphogenesis may lead to discovery of fundamental mechanisms that govern development of animal form. To learn about cell types and signaling interactions that govern skin patterning and morphogenesis we generated and analyzed single cell transcriptomes of skin from wild-type fish as well as fish having genetic or transgenically induced defects in squamation or pigmentation. These data reveal a previously undescribed population of epidermal cells that express transcripts encoding enamel matrix proteins, suggest hormonal control of epithelial-mesenchymal signaling, clarify the signaling network that governs scale papillae development, and identify a critical role for the hypodermis in supporting pigment cell development. Additionally, these comprehensive single-cell transcriptomic data representing skin phenotypes of biomedical relevance should provide a useful resource for accelerating discovery of mechanisms that govern skin development and homeostasis.

eLife assessment

This study provides a clearly presented and thoughtfully analyzed single cell-resolution dataset of gene expression in wildtype and mutant zebrafish skin. These data are used by the authors to develop and test hypotheses about cell lineage relationships and signaling interactions between cell types in the skin, allowing them to identify roles for several signaling pathways and the hypodermis in scale and pigment cell development. These findings constitute a **fundamental** contribution to the field, and the rigor of the analyses make this manuscript **compelling**.

Introduction

The steady-state chemistry of life on earth occurs within compartments bounded from the rest of the cosmos. Providing this boundary function and serving as the primary interface between organisms and their environments are sophisticated integuments that, in vertebrates, comprise marvelous and varied skins, decorated with patterns of pigmentation and arrayed appendages including feathers, fur or scales. Understanding the mechanistic underpinnings of animal form and phenotypic diversity is an enduring goal of basic biology and studying skin patterning and morphogenesis can advance that goal. Additionally, while human skin is a major contributor to our outward appearance, bears all our physical interactions, and detects all our tactile sensations, it remains a failure-prone organ system with numerous poorly understood and debilitating pathologies.

Studying the skin of research organisms chosen based on phylogeny or experimental exigency can improve our understanding of regulatory mechanisms underlying integumental patterning and morphogenesis. Comparing developmental mechanisms across species can provide clues to the origin and evolution of this important organ system and may also reveal fundamental mechanisms relevant to human health and disease. To these ends, the development of skin, and cell types within the skin, have been studied across a variety of research organisms, yielding insights into both general and species-specific mechanisms (Duverger and Morasso, 2009; Chen et al., 2015; Patterson and Parichy, 2019; Aman and Parichy, 2020).

Zebrafish (*Danio rerio*) is an outstanding research organism for studying vertebrate skin patterning and morphogenesis. Zebrafish skin, like all vertebrate skin, has a superficial epidermis composed of ectoderm-derived epithelial cells and an underlying dermis composed of mesoderm-derived mesenchymal cells and collagenous stromal matrix (Le Guellec et al., 2004; Aman and Parichy, 2020). During post-embryonic development, zebrafish skin simultaneously develops arrays of calcified scales and pigmented stripes. Both form superficially on the surface of the animal and are dispensable for survival in the laboratory, making them readily amenable to imaging and experimental perturbation, and enabling analyses of underlying cellular dynamics and molecular mechanisms (Lee and Kimelman, 2002; Aman et al., 2018; Cox et al., 2018; Iwasaki et al., 2018; Rasmussen et al., 2018; Patterson and Parichy, 2019; De Simone et al., 2021).

To reveal potentially rare cell populations important for zebrafish scale development and pigment patterning we used unbiased single-cell transcriptional profiling and live imaging of skins undergoing post-embryonic morphogenesis. Additionally, to gain insights into molecular mechanisms underlying human skin pathologies we profiled skins from *ectodysplasin a* (*eda*) mutants, *basonuclin 2* (*bnc2*) mutants, and hypothyroid fish (hypoTH; **Figure 1A**). Eda-Edar-NF- κ B is a conserved signaling pathway that is necessary for normal skin appendage development in all vertebrates examined to date (Kere et al., 1996; Srivastava et al., 1997; Kondo et al., 2001; Houghton et al., 2005; Harris et al., 2008; Di-Poi and Milinkovitch, 2016). Mutations in the signaling ligand Eda-A (Ectodysplasin-A), its receptor Edar, or downstream signal transduction molecules in the NF- κ B (nuclear factor- κ B) pathway underlie human ectodermal dysplasias, hereditary disorders defined by loss of skin appendages and teeth (Cluzeau et al., 2011). Similarly, *eda* mutant zebrafish completely lack scales, though specific mechanisms linking Eda signaling to scale formation remain unclear (Harris et al., 2008). To learn more about potential interactions between pigment cells and their skin microenvironment, we profiled a mutant for *bnc2*, a conserved zinc finger containing protein implicated in human pigment variation that acts through the tissue environment to promote pigment cell development in zebrafish (Lang et al., 2009; Patterson and Parichy, 2013; Visser et al., 2014; Endo et al., 2018; Ayoola et al., 2021). Finally, we profiled

skins from hypothyroid fish (hypoTH) that are unable to synthesize thyroid hormone owing to transgene-mediated ablation of the thyroid gland (McMenamin et al., 2014). Thyroid hormone (TH) is a potent regulator of vertebrate skin development, and thyroid dysfunction underlies debilitating skin pathologies (Mancino et al., 2021). We have shown that TH is necessary for dermal morphogenesis as well as pigment cell maturation and pattern formation though the underlying mechanisms remain elusive (McMenamin et al., 2014; Saunders et al., 2019; Aman et al., 2021).

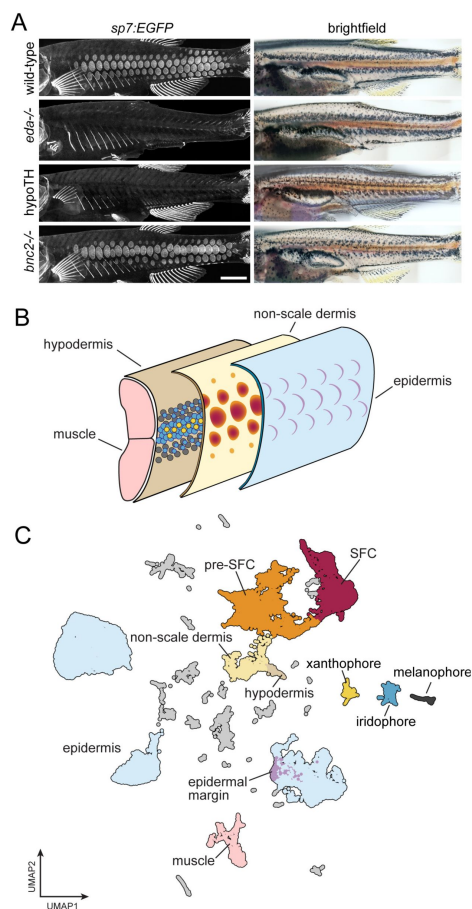


Figure 1.

A whole-skin single cell transcriptome from zebrafish undergoing skin patterning. **(A)** Confocal images illustrate scale forming cells (SFC) expressing *sp7:EGFP* and brightfield images of the same fish show pigment pattern. At 9.2 SSL, approximately three weeks post fertilization under standard conditions, wild-type zebrafish have all steps of scale development represented and are developing a pigment pattern of dark stripes, with melanophores and sparse iridophores, alternating with light interstripes of densely packed iridophores and yellow xanthophores. Wild-type individuals of 9.6 SSL having precisely four rows of scales were selected for preparation of nuclei to be used in single cell indexed RNA-Seq (sci-RNA-Seq); *eda* mutants and hypoTH fish, devoid of scales, as well as *bnc2* mutants having fewer, dysmorphic scales, were likewise reared to 9.6 SSL for isolation of nuclei. **(B)** Schematic representation of zebrafish skin at 9.2 SSL. The outermost layer of skin is epidermis (blue), which develops crescent shaped placodes (epidermal margin, lavender,) above each scale. In the dermis (yellow), scale-forming cells (SFCs) differentiate (orange → red). In the hypodermis (brown), dark melanophores, yellow xanthophores and iridescent iridophores (grey, yellow and blue circles, respectively) organize into alternating stripes. **(C)** UMAP visualization of 35,114 transcriptomes from nuclei of wild-type fish, colored by cell type assignments inferred by marker gene enrichment. Colors correspond to the schematic in (B). SFC, Scale forming cell. Scale bar, 500 μm (A).

Our analyses, using single cell transcriptomics supplemented by histological analyses of gene expression, fate mapping and experimental manipulations revealed a previously undescribed epidermal cell type that expresses transcripts encoding enamel matrix proteins, relevant to understanding the ancient origins of calcified tissues, and have clarified the position of *Eda* within the signaling network that governs scale papilla induction. We also discovered a novel regulatory pathway that connects globally circulating TH to local epithelial-mesenchymal interactions during dermal development. We further identify the hypodermis as a crucial pigment cell supporting tissue that provides a permissive environment for the self-organizing interactions of adult stripe formation. Lastly, by comparing analyses of single cell transcriptomes with spatial analyses of gene expression and cell-type differentiation we uncover instances in which single cell bioinformatic inferences represent, and fail to represent, true cell state transformations. Together, these analyses provide new insights into the development and evolution of vertebrate skin and highlight the importance of validating inferences of differentiation and lineage from single cell transcriptomics with paradigms for assessing developmental events *in vivo*.

Results

sci-RNA-seq of whole skin reveals cell type diversity during post-embryonic development

Skin is a large and complex organ system, with contributions from multiple embryonic germ layers and a variety of distinct cell types. Likely due to a shared requirement for TH, pigment pattern formation and squamation occur simultaneously during post-embryonic development, in different layers of the skin ([Figure 1B](#)) (McMenamin et al., 2014; Saunders et al., 2019; Aman et al., 2021). To capture individual transcriptomes from a minimally biased sampling of skin cells, we performed single nucleus combinatorial indexing (sci)-RNA-seq (Cao et al., 2017; Cao et al., 2019) on nuclei from pooled, fresh-frozen whole skins at 9.6 mm standardized standard length [9.6 SSL (Parichy et al., 2009)], a key developmental stage of skin patterning and morphogenesis. In 9.2 SSL wild-type fish, all steps of scale morphogenesis are represented and a ‘primary’ pigment pattern is apparent with secondary pattern elements just beginning to form ([Figure 1A,B](#); [Figure 1—figure supplement 1](#)) (Aman et al., 2019; Parichy et al., 2009).

To better understand the contributions of individual cell types and key factors involved in the major skin patterning events at this stage, we included three additional backgrounds representing distinct developmental perturbations: *eda* mutants, hypoTH fish and *bnc2* mutants (Harris et al., 2008; Lang et al., 2009; McMenamin et al., 2014). We processed all tissue in a single sci-RNA-seq experiment, barcoding each genotype by reverse transcription index during library preparation. In total, we recovered high quality transcriptomes from 144,466 individual nuclei with an average of 1,300 unique molecular identifiers (UMIs) and 720 genes detected per cell (~60% duplication rate). Cell recovery, UMIs per cell, and numbers of genes detected were consistent across all sample groups ([Figure 1—figure supplement 2](#)). We removed likely multiplets (11%) using Scrublet (Wolock et al., 2019) and further processed the data and performed dimensionality reduction and clustering with Monocle3 (Cao et al., 2019). To characterize cell types and developmental trajectories, we focused on the wild-type data alone (35,114 cells). We classified cells into major cell types by assessing expression of published markers for different skin and skin-associated cell types ([Figure 1C](#); [Figure 1—figure supplement 3](#); [Supplementary file 1—Table 1](#); [Supplementary file 2—Table 2](#)). The majority of cells were from epidermal and dermal populations, where we recovered transcripts from relatively rare subsets, including the *edn3b/bnc2*+ hypodermal monolayer that forms the deep limit of the dermis and *shha*+ epidermal placode cells at the posterior scale margin (Sire and Akimenko, 2004; Lang et al., 2009; Budi et al., 2011; Patterson and Parichy, 2013; Aman et al., 2018). It is likely that additional cell-state heterogeneity exists within these major cell types beyond what we have annotated here ([Figure 1—figure supplement 4](#); [Supplementary file 2—Table 5](#)). As expected, based on our stage selection, we recovered dermal scale forming cells (SFCs) and their progenitors (pre-SFCs) ([Figure 1—figure supplement 1](#)). We also recovered less abundant cell types including pigment cells, lateral line cells, goblet cells, ionocytes, glia and immune cells ([Figure 1C](#); [Figure 1—figure supplement 3](#)).

Decoupling of transcriptional dynamics and inferred lineage relationships during scale development and epidermal maturation

One goal of our study was to resolve transcriptional dynamics during differentiation of cell lineages that underlie skin patterning and morphogenesis. Adult zebrafish are adorned with

a full coat of partially overlapping elasmoid scales, thin plates of calcified ECM that grow between the dermis and epidermis during post-embryonic development (**Figure 1; Figure 1—figure supplement 1**) (Sire et al., 1997; Aman et al., 2018). Scale development is associated with a population of dermal cells—referred to here as SFCs—that express genes encoding transcription factors, including *sp7* (also known as *osterix*, *osx*) and *runx2a/b*, that are necessary for differentiation of osteoblasts and odontoblasts (Sire et al., 1997; Komori, 2010; Li et al., 2011; Zhang, 2012; Aman et al., 2018; Bae et al., 2018; Cox et al., 2018; Iwasaki et al., 2018; Rasmussen et al., 2018). Given the presence of all steps of scale development in our sampled timepoint, we captured single cells along the continuum of SFC differentiation from pre-SFCs to mature, matrix-secreting cells. Scale development also coincides with expansion of the three major epidermal cell types – periderm (also known as superficial epidermal cells), suprabasal cells, and basal cells – of which basal cells give rise to both of the other cell types (Lee et al., 2014). Our transcriptomic analyses confirmed that SFCs express *sp7* and suggested that they likely differentiate from *runx2b*+ progenitors (**Figure 2A**). Comparing *runx2b* and *sp7* mRNA distributions during scale development revealed that *runx2b* is consistently more broadly expressed than *sp7*, suggesting the existence of a restricted halo of SFC progenitors surrounding the growing scale (**Figure 2B,D**). To confirm that these cells represent SFC progenitors, we exploited the differential localization of two transgenic reporters, cytosolic *ET37:EGFP*, labeling all dermal cells but having markedly reduced expression in SFCs, and photoconvertible, nuclear-localizing *sp7:nEOS*, restricted to differentiated SFCs (Parinov et al., 2004; Aman et al., 2021): if peripheral *runx2b*+ cells give rise to SFCs, initially cytosolic reporter expression should transition to nuclear reporter expression. We excluded previously differentiated *sp7:nEOS*+ SFCs from consideration by photoconversion (green → red), and looked for *ET37:EGFP*+ cells (green cytosol) newly expressing *sp7:nEOS* (green nucleus, without red). As expected, over three days of scale development, we observed numerous cells acquire nuclear nEOS expression as they were incorporated into growing scales, supporting the inference that peripheral, presumptively *runx2b*+ dermal cells differentiate as SFCs (**Figure 2C,D**).

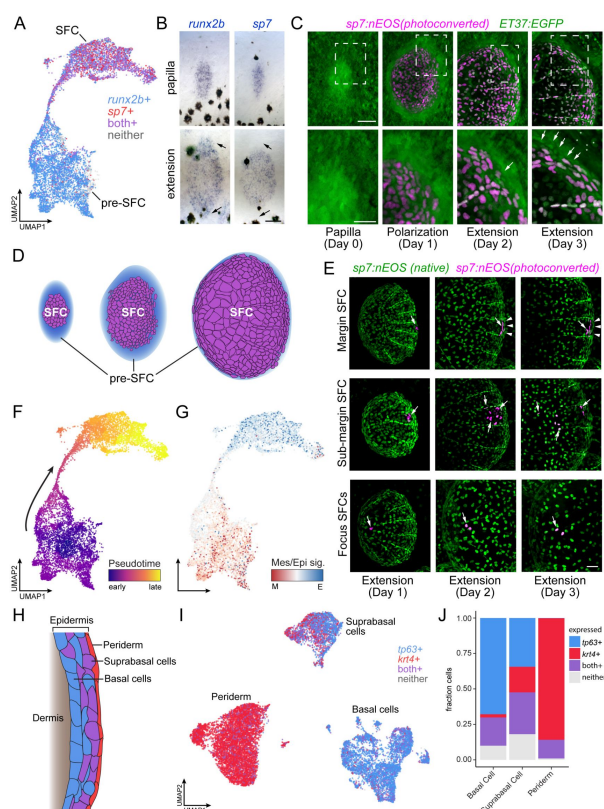


Figure 2.

Postembryonic skin cell lineage relationships are not reflected in UMAP space. **(A)** UMAP visualization showing distribution of differentiated SFC expressing *sp7* and pre-SFC progenitors expressing *runx2b*. **(B)** *In-situ* hybridization of *sp7* and *runx2b* shows that a halo of pre-SFC progenitors surround the growing scale (arrows). **(C)** *sp7:nEOS*-expressing differentiated SFC (magenta), were labelled by photoconversion on Day 1. Over the following two days, newly differentiated, un-photoconverted SFC appeared at the scale margin (arrows; $n = 5$ fish). **(D)** Schematic representation of differentiated SFC (purple) and the associated halo of pre-SFC (blue). **(E)** Photoconversion of small groups of SFC in the scale margin and sub-margin; and single-cell photoconversion of focus SFCs (arrows) showed that SFC are progressively displaced toward the scale focus and that SFC in all these regions are capable of cell division (arrows, $n \geq 4$ fish for each region tested). Margin SFCs were displaced towards the posterior by newly differentiated, un-photoconverted SFCs (arrowheads). **(F)** SFCs in UMAP space colored by “pseudotime” rooted in the SFCs. **(G)** SFCs in UMAP space colored by the ratio of a mesenchymal (migratory) signature to an epithelial signature (**Supplementary file 2—Table 3**). **(H)** Schematic

representation of epidermis with major substrata. **(I)** UMAP visualization of wild-type epidermis, subclustered independently of other cell types and displaying expression of the epidermal basal cell marker *tp63* (blue) and the periderm marker *krt4* (red). **(J)** The fraction of cells from panel **H** that pass a minimum threshold for expression of *tp63*, *krt4* or both genes. Scale bars, 50 μm (B,C,E); 25 μm , (C, lower).

Because the stage at which cells were collected contains SFCs at all steps of differentiation, and because the transcriptomes of SFCs displayed a continuous path between pre-SFC and differentiated SFC in UMAP space ([Figure 2A](#)), we anticipated that the cell fate transition during SFC differentiation would be captured by pseudotemporal ordering, which can reveal cell state transitions in asynchronous populations of differentiating cells (Trapnell et al., 2014). To test this idea, we compared gene expression in UMAP space to spatial patterns of gene expression as revealed by *in-situ* hybridization, using a combination of new staining and previously published staining (Aman et al., 2018; Cox et al., 2018; Iwasaki et al., 2018; Rasmussen et al., 2018). We predicted that cells having the intermediate state predicted by pseudotemporal ordering should be present circumferentially, between the halo of *runx2*⁺ dermal cells and differentiated *sp7*⁺ SFCs. Instead, these pseudotemporally intermediate cells were located at scale radii, a relatively late-appearing structure comprising cells unlikely to be in a transitional state of SFC differentiation ([Figure 2—figure supplement 1G](#)). To resolve the true SFC differentiation trajectory, we fate-mapped individual SFCs *in-vivo* by photoconverting *sp7:nEOS*⁺ SFCs within specific scale regions and following them over several days ([Figure 2E](#), [Figure 2—figure supplement 2A](#)). These analyses showed that SFCs at the posterior margin are progressively displaced towards the scale focus, presumably by addition of newly differentiated SFCs from *sp7*-negative progenitors ([Figure 2E](#), top). SFCs in the sub-marginal region contributed to elongated marginal cells, scale radii cells, and an SFC subset that rapidly displaced toward the scale focus ([Figure 2E](#), middle). Lastly, cells at the focus in a nascent scale remained in the focus as the scale grew ([Figure 2E](#), bottom). These results demonstrate that radii cells descend from marginal SFCs, consistent with previous live imaging results (Cox et al., 2018; Iwasaki et al., 2018; Rasmussen et al., 2018), confirming that SFCs do not pass through an intermediate state as scale radii cells during differentiation, which a facile interpretation of pseudotemporal ordering might suggest. Thus, although cell state transitions inferred from transcriptomes alone can suggest cell lineage relationships and cell states along a differentiation continuum, we find that the pseudotemporal ordering, while continuous from pre-SFC to differentiated SFC, did not faithfully represent the cell state path during SFC differentiation. Instead, the continuity of cell states revealed by pseudotemporal ordering could reflect continuous variation in other biological processes apart from differentiation, like states of the cell cycle or migration (Buettner et al, 2015; McFaline-Figueroa, et al. 2019). Given the mesenchymal appearance of pre-SFC dermal cells and the epithelial appearance of differentiated SFCs ([Figure 2C](#); [Figure 1—figure supplement 1](#)) (Iwasaki et al., 2018; Aman et al., 2021), we predicted that pseudotemporal order in this dataset might reflect differences in gene expression associated with these different cellular morphologies. To test this possibility, we constructed gene expression signatures for mesenchymal and epithelial states from the literature and mapped the ratio of epithelial-to-mesenchymal scores on cells in UMAP space, which revealed an overall correspondence of these scores with pseudotemporal order ([Figure 2F,G](#)). Finally, our observations also provided an opportunity to resolve a controversy as to whether differentiated SFCs are capable of proliferating or whether scale growth occurs exclusively by hypertrophic growth of individual cells (Cox et al., 2018; Iwasaki et al., 2018): both transcriptomic analyses and live imaging experiments showed that differentiated SFCs remain proliferative ([Figure 2—figure supplement 2B–D](#)).

In addition to SFC differentiation, we sought to understand the transitional dynamics of epidermal differentiation during post-embryonic skin maturation. During this stage of

development, basal epidermal cells are the stem cell population that differentiate into both suprabasal and periderm cells, and each of the three major epidermal cell types are well represented in our dataset ([Figure 2H,I](#); [Figure 1—figure supplement 3](#)) (Guzman et al., 2013; Lee et al., 2014). While periderm cells at the sampled stage are likely of dual origin, representing a mixture of early embryonic and stem cell derived cells, suprabasal cells are entirely derived from basal cells (Kimmel et al., 1990; Guzman et al., 2013; Lee et al., 2014). Given the known lineage relationships, we predicted that the single cell transcriptome data would reveal a continuum of cell state transitions during the differentiation of both cell types. While the established cell type markers for basal cells and periderm cells, *tp63* and *krt4*, displayed a clear transition between basal cells (*tp63*⁺, *krt4*⁻), suprabasal cells (*tp63*⁺, *krt4*⁺), and periderm (*tp63*⁻, *krt4*⁺), the trajectories were discontinuous in UMAP space ([Figure 2I,J](#)). These discontinuities were present in both global and tissue-specific UMAP projections and across a wide range of UMAP parameters ([Figure 2I](#); [Figure 1—figure supplement 3](#)). Moreover, the number of differentially expressed genes between each of the epidermal types was about twice as large as between cell clusters having continuous trajectories (basal cell vs. periderm, 7341 DEGs; basal cell vs. suprabasal, 4024 DEGs; pre-SFC vs. SFC, 2373 DEGs; all $q < 0.01$), suggesting that discontinuities among epidermal cell subtypes reflect abrupt transcriptional changes during differentiation, rather than artifacts of the dimensionality reduction itself. Together, our results highlight the importance of coupling true lineage information with well sampled, high-resolution single cell transcriptomes in order to understand complex transcriptional dynamics over the course of differentiation *in vivo*.

A heterogenous population of epidermal and dermal cells contributes to scale plate formation

Zebrafish elasmoid scales represent one of several forms of calcified skin appendages that cover the bodies of most non-tetrapod fish species (hereafter referred to as ‘fish’). Calcified skin appendages are an ancient vertebrate trait with a robust fossil record, first appearing in the Ordovician 450 million years ago, prior to the appearance of paired fins, jaws and teeth (Smith et al., 2002; Märss, 2006; Sire et al., 2009; Fraser et al., 2010; Turner et al., 2010; Janvier, 2015). These ancestral skin appendages were morphologically and compositionally similar to modern-day teeth, with a hypercalcified, enamel-like matrix capping collagen-rich calcified matrices resembling dentin, bone or both (Sire, 1990; Smith et al., 2002; Märss, 2006; Sire et al., 2009) ([Figure 3A](#)). While certain extant non-teleost fishes, like sharks, bichir and gar, have scales that resemble skin appendages of ancient fishes and modern teeth, the flattened morphology and elastic flexibility of elasmoid scales typical of most extant fish is highly derived (Sire, 1990; Sire et al., 2009). Histological and ultrastructural studies have shown that the elasmoid scales of zebrafish and other teleosts are composed of weakly calcified collagenous matrix, known as elasmoidin, capped by a collagen-free, hypermineralized limiting layer that forms in close proximity to basal epidermal cells with an overtly secretory morphology ([Figure 3A](#)) (Sire, 1990; Sire et al., 1997; Quan et al., 2020). In vertebrate teeth and tooth-like scales of non-teleost fish, layers of calcified matrix are deposited by two cooperating cell types, mesenchymal cells of dermal or neural crest origin that form collagen-rich calcified matrix like dentin, bone or both, and overlying epithelial cells of epidermal or endodermal origin that produce hypermineralized matrices like enamel (Sire, 1990; Sire and Huysseune, 2003; Oralová et al., 2020; Kawasaki et al., 2021).

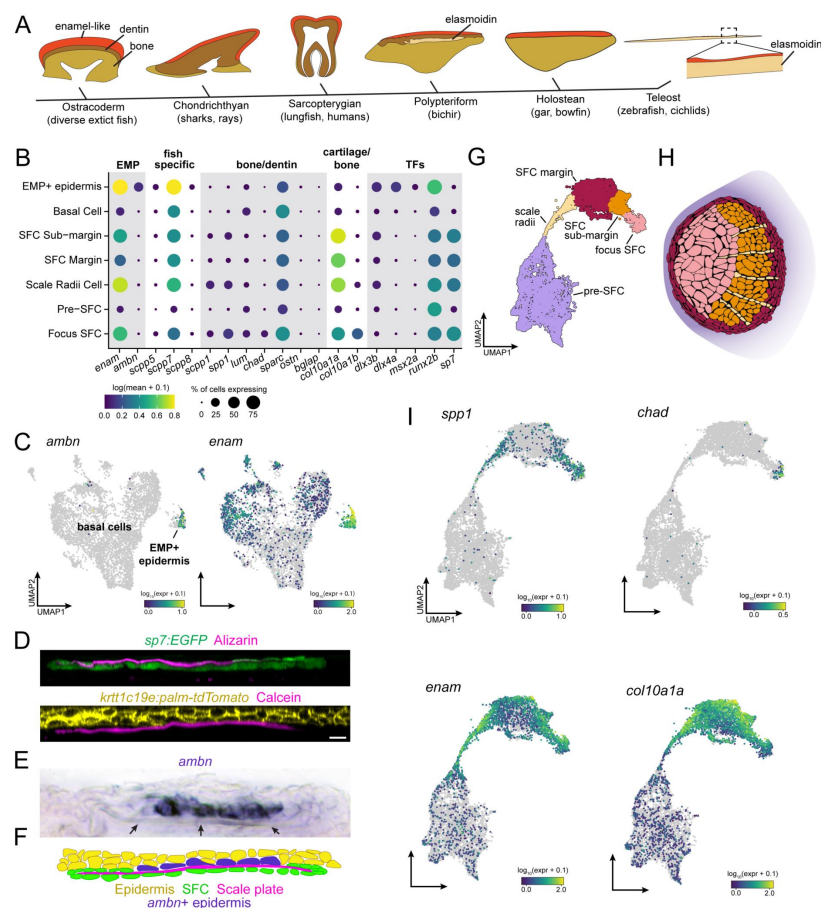


Figure 3.

Evidence for epidermal and dermal contributions to scale plate ECM. (A) Simplified vertebrate phylogeny with schematic depictions of calcified appendages. Phylogeny based on (Near et al., 2012; Betancur et al., 2013); schematics drawn after (Sire et al 2009). (B) Dot plot visualization of transcripts encoding EMPs, fish-specific SPP tandem duplicates, non-collagen matrix proteins associated with bone and dentin, collagen associated with cartilage and transcription factors that regulate osteoblast and ameloblast differentiation. (C) UMAP visualization of epidermal basal cells showing distribution of transcripts encoding enamel matrix proteins. (D) Optical sections of growing scale in live animals showing the relative position of calcified matrix dyed with Alizarin Red S (ARS) or Calcein, and dermal SFC visualized with *sp7:EGFP* transgene and epidermis visualized with *krt1c19e:palm-tgTomato* transgene. (E) *In-situ* hybridization of *ambn*, encoding the enamel matrix protein Ameloblastin. Arrows point to the calcified scale plate. (F) Schematic representation of epidermal *ambn* expressing cells

(blue), the calcified scale plate (magenta), dermal SFC (green) and epidermis (yellow). (G) UMAP visualization of dermal SFC and pre-SFC. (H) Position of SFC sub-types within a scale inferred from *in-situ* hybridization assays (Figure 2—figure supplement 1). (I) UMAP visualization of transcripts encoding non-collagen matrix proteins associated with bone (*spp1*, *chad*), enamel (*enam*) and cartilage (*col10a1a*). Scale bar, 10 μ m.

Hypothesizing that epidermis of zebrafish retains this ancient enamel deposition function, we predicted that a subset of epidermal basal cells would express transcripts encoding enamel matrix proteins (EMPs). EMPs were originally discovered in human patients suffering from Amelogenesis Imperfecta, a congenital condition characterized by defective enamel formation (Smith et al., 2017). EMPs are part of family of proteins know as secretory calcium-binding phosphoproteins (SCPPs) that are critical for calcification of bone, dentin and enamel (Kawasaki, 2009). The zebrafish genome harbors two previously identified EMP-gene orthologs, including orthologs of human Enamelin (encoded by *enam*) and Ameloblastin (encoded by *ambn*) in addition to fish-specific orthologs that likely originated by tandem duplication (Qu et al., 2015; Braasch et al., 2016; Kawasaki et al., 2017). We therefore analyzed the distribution of SCPP-encoding transcripts, predicting that EMP-transcript expressing cells would be found among basal epidermal cells. Consistent with our prediction, we identified a transcriptionally distinct population of basal epidermal cells, separated from other basal epidermal cells in UMAP projections, that expressed EMP genes (Figure 3B,C). Strikingly, transcripts of the conserved EMP gene *ambn* was found almost exclusively in this subpopulation of basal cells, which we refer to as EMP+ epidermal cells (Figure 3B,C). A previous ultrastructure study revealed epidermal basal cells with a secretory morphology in contact with the calcified scale matrix (Sire et al., 1997). To test

correspondence of that population with EMP+ epidermal cells identified in our transcriptomic data, we visualized *ambn* expression by *in-situ* mRNA hybridization, which revealed that EMP+ epidermal cells are positioned precisely where epidermal cells contact the scale matrix (**Figure 3D-F**).

Although enamel-like capping matrix is common in phylogenetically diverse vertebrates (**Figure 3A**) (Sire et al., 2009), it is possible that epidermal expression of EMP transcripts evolved convergently in zebrafish. If epidermal EMP+ basal cells are homologous with mammalian ameloblasts, we reasoned that in addition to genes encoding matrix proteins, these cells should express transcription factors that regulate mammalian ameloblast differentiation. Indeed, we find that *dlx3a*, *dlx4a*, *msx2a* and *runx2b* are expressed in these cells (**Figure 3B**) (Pemberton et al., 2007; Debais-Thibaud et al., 2011; Urzúa et al., 2011; Babajko et al., 2015; Chu et al., 2018; Woodruff et al., 2022). Together, these results suggest that the enamel-like, hypermineralized limiting layer of the scale is produced by deeply conserved, ameloblast-like epidermal cells.

In teeth and tooth-like skin appendages, enamel-like matrix caps more collagen-rich calcified matrices such as dentin or bone that are deposited by condensed mesenchymal cells (Sire et al., 2009; Fraser et al., 2010; Murcia et al., 2016; Arola et al., 2018). Although dermal SFCs are frequently referred to as ‘osteoblasts’ due to their association with calcified matrices and their expression of conserved transcription factor genes *sp7* and *runx2a/b*, the elasmoidin matrix they deposit, characterized by weakly calcified, plywood-like layers of hydrated collagen fibrils, is materially distinct from bone or dentin (Sire et al., 2009; Metz et al., 2012). We therefore hypothesized that SFCs would express a distinct complement of SSCP transcripts. To elucidate this repertoire and compare dermal SFCs with osteogenic cell types like osteoblasts, odontoblasts and ameloblasts, we assessed the expression SSCP transcripts. For these analyses, we subclustered the SFCs/pre-SFCs and identified five major cell states based on gene expression from sci-RNA-seq and *in-situ* hybridization (**Figure 3G,H; Figure 2—figure supplement 1**). Consistent with an overall similarity between osteoblasts and SFCs, we detected transcripts of Osteopontin (*spp1*) (**Figure 3B,I**) and Chondroadherin (*chad*) (**Figure 3B,I; Figure 2—figure supplement 1J**), encoding bone matrix proteins, in addition to Secretory-calcium-binding Phosphoprotein 1 (*scpp1*), which is homologous to human Dentin-Matrix-Acidic Proteins that form a component of both bone and dentin (**Figure 3B,I**) (Larsson et al., 1991; Raouf et al., 2002; Hessel et al., 2013; Venkatesh et al., 2014; Braasch et al., 2016; Kawasaki et al., 2017). Nevertheless, we failed to detect robust expression of transcripts encoding bone matrix proteins Osteocrin (*ostn*) or Osteocalcin (Bone gamma-carboxyglutamate protein, *bglap*) in SFCs (**Figure 3B**) (Thomas et al., 2003). In addition to a subset of bone-specific transcripts, SFCs also expressed EMP genes and genes encoding fish-specific SCPPs (**Figure 3B,I**). Expression was spatially restricted among SFCs, with the bone-associated transcripts, *spp1* and *chad* primarily expressed in the scale focus and radii SFCs, while EMP genes were restricted to SFCs at the scale margin (**Figure 3B,G,H,I; Figure 2—figure supplement 1J**).

Scale elasmoidin is a flexible, collagenous ECM, material properties that are similar to cartilage (Quan et al., 2020). We therefore wondered whether dermal SFCs express matrix proteins associated with cartilage formation. Col10a1 and Col2a1 are major structural molecules in collagen, although *col10a1* transcription its expression has also been documented in osteoblasts (Gu et al., 2014; Yang et al., 2014; Kawasaki et al., 2021). The zebrafish genome harbors genes encoding two Col10a1 orthologs (*col10a1a* and *col10a1b*) and we found both transcripts in SFCs representing distinct steps of maturation, however, transcripts of Col2a1 genes (*col2a1a* and *col2a1b*) were not robustly detected in these cells (**Figure 3B,I; Figure 2—figure supplement 1F,I**). Transcripts of genes encoding additional factors associated with mineralized matrix formation, such as Osteonectin (*sparc*), were

expressed broadly in skin (**Figure 2A,H**) raising the possibility of additional roles beyond assembly of calcified matrix (Rosset and Bradshaw, 2016).

Together, these analyses revealed the presence of epidermal EMP expressing cells and support a hypothesis of ancient homology between ameloblast-like cells in fish skin and the mammalian dental lamina, and between the enamel-like materials that coat zebrafish scales and tetrapod teeth. Furthermore, our observations that dermal SFCs express a subset of genes associated with bone development, as well as genes encoding EMPs and cartilage proteins, suggest that the distinct properties of elasmoid matrix are due to a distinct complement of matrix proteins. While these conclusions are correlational, this work will facilitate functionally testing the role of candidate matrix proteins in the material properties of calcified matrices.

Eda-Edar-NF- κ B and TH regulate distinct stages of dermal SFC development

Eda-Edar-NF- κ B signaling plays conserved roles in regulating the patterning and morphogenesis of vertebrate skin appendages (Cui and Schlessinger, 2006; Lefebvre and Mikkola, 2014), whereas TH regulates multiple aspects of skin development and homeostasis (Mancino et al., 2021). Both *eda* mutants and hypoTH fish completely lack scales at the stage sampled, allowing us to compare transcriptomic signatures and cell type complements associated with scale loss in these different backgrounds (**Figure 1A**). Analyses of cell type abundance revealed that both scale-free conditions were characterized by a complete lack of fully differentiated dermal SFCs (**Figure 4A,B**). These analyses further showed that *eda* mutants—despite homozygosity for a presumptive null allele—retained a small subset of dermal pre-SFC progenitors, whereas hypoTH skins lacked pre-SFC entirely. We have previously shown that hypoTH fish lack superficial dermal cell types at the stage sampled for sequencing, though these cell types do appear later in development (Aman et al., 2021). To confirm the presence of residual, pre-SFC in *eda* mutants, we imaged live fish expressing *ET37:EGFP* (Parinov et al., 2004; Aman et al., 2021). Indeed, *eda* mutants exhibited a population of *ET37:EGFP*⁺ dermal cells—presumptive pre-SFC—beneath the epidermis that was not present in hypoTH fish (**Figure 4C,D**).

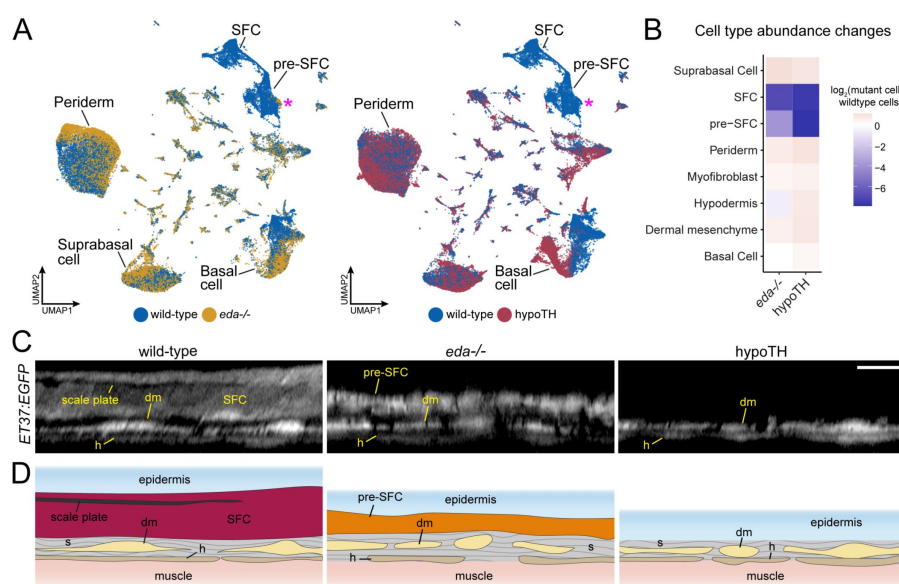


Figure 4.

Eda and TH regulate signaling ligand transcription in basal epidermal cells. (A) UMAP visualizations with wild-type cells in blue, *eda* mutant cells in yellow and hypoTH cells in red highlighting the absence of SFC and residual pre-SFC in skins of *eda* mutants and the absence of both populations in skins of hypoTH fish. Magenta asterisks mark differences in pre-SFC complements between *eda* mutant and hypoTH; other cell types designated in **Figure 1C**. (B) Heatmap visualization of cell type abundance shows that *eda* mutants retain more pre-SFCs than hypoTH

fish. (C) Optical sections and (D) schematic representations of wild-type, *eda* mutant and hypoTH skin, visualized in

ET37:EGFP transgenics, showing abundance of SFC in wild-type, a thin layer of pre-SFC in *eda* mutants, and lack of pre-SFC in hypoTH skin. dm, dermal mesenchyme; h, hypodermis; s, stromal collagen. Scale bar, 10 μ m (C).

In previous work, we found that *Eda-Edar-NF- κ B* signals are transmitted from the dermis to epidermis during scale morphogenesis (Aman et al., 2018). The apparently monogamous receptor for *Eda*, encoded by *edar*, is expressed in the epidermis during scale morphogenesis. The signaling ligand, encoded by *eda*, has a dynamic expression pattern that shifts from broad expression in unspecified epidermis to localized expression in dermal papillae during scale morphogenesis. These expression dynamics are strikingly similar during mouse hair follicle and chicken feather patterning (Montonen et al., 1998; Headon and Overbeek, 1999; Houghton et al., 2005). We confirmed that these expression domains were reflected in our sci-RNA-seq data, in which *edar* is detected primarily in basal epidermal cells, with most intense expression in the *shha*⁺ epidermal placode (Figure 5A) (Aman et al., 2018). *eda* transcripts were detected at much lower levels and were much more dispersed across cell types than *edar* transcripts but were nevertheless enriched in pre-SFC as expected (Figure 5A).

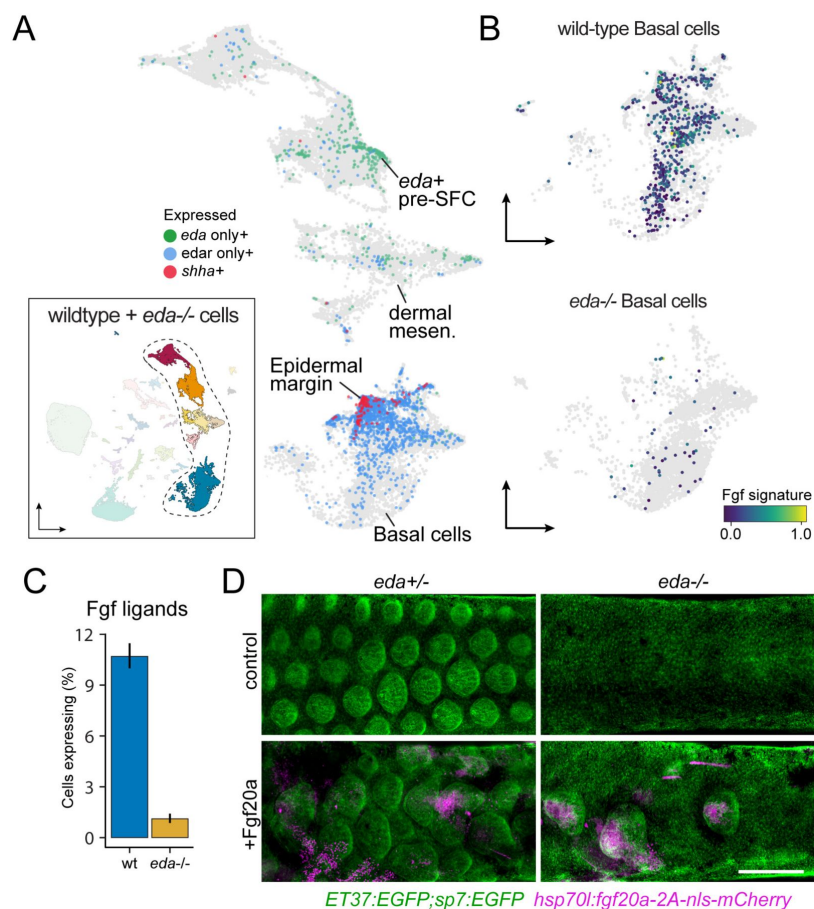


Figure 5.

Eda regulates SFC differentiation via transcriptional regulation of basal epidermal Fgf ligands. (A) Wild-type dermal and basal epidermal cells plotted in UMAP space and colored by whether they are expressing *eda*, *edar*, or *shha* (*shha*⁺ cells also include cells expressing both *edar* and *shha*). (B) Wild-type and *Eda* mutant basal cells plotted in UMAP space and colored by the expression of the signature score of Fgf ligands that are specifically expressed in basal cells (*fgf24*, *fgf20a*, *fgf20b*; specificity score > 0.1). (C) Percent of cells expressing the Fgf ligand signature between wild-type and *Eda* mutant basal cells (error bars estimated via bootstrapping (*n* = 100)). (D) Scales and dermis visualized in *sp7:EGFP; ET37:EGFP* double transgenics (green). Heat-shocked control *eda*^{+/+} larvae developed well-patterned, uniformly shaped scales (*n* = 6) whereas heat-shocked control *eda*^{-/-} larvae developed no scales (*n* = 6). Mosaic heat-shock induction of *Fgf20a*, stringently selected for expression in epidermal cells (magenta) caused mis-patterned and dysmorphic scales to grow in wild-type *eda*^{+/+}

larvae (*n* = 25 epidermal clones in 6 fish) and rescued scale formation in *eda*^{-/-} larvae (*n* = 36 epidermal clones in 6 fish). Scale bar, 500 μ m (D).

Since Eda-Edar-NF- κ B pathway activation occurs exclusively in epidermis but drives morphogenesis of dermal cells, we predicted the existence of an Eda-dependent signaling ligand expressed in basal epidermal cells that would complete a presumptive epithelial-mesenchymal signaling loop (Aman et al., 2018). We previously observed that global misexpression of an Fgf ligand rescued scale development in *eda* mutant fish, and that Edar expression in epidermal cells, but not dermal cells, was sufficient to drive scale formation (Aman et al., 2018). Those results suggested that Eda signaling drives SFC differentiation via epidermal expression of one or more Fgf ligands. Our transcriptomic analysis showed that, indeed, epidermal expression of Fgf ligands is Eda-dependent (Figure 5B,C). Among Fgf ligand transcripts detected in epidermis was that of *fgf20a*, which plays a conserved role in regulating dermal morphogenesis in amniote skin appendage development and has been implicated in scale development and regeneration in zebrafish (Huh et al., 2013; Daane et al., 2015). If Fgf20a functions downstream of epidermal Eda-Edar-NF- κ B signaling, we predicted that experimental restoration of *fgf20a* expression in *eda* mutant skin should bypass the requirement for Eda in scale formation, thereby rescuing scales even in the absence of Eda function. Indeed, heatshock-driven expression in F0 mosaics stringently selected for basal epidermal expression of Fgf20a in the skin of *eda* mutants led to localized rescue of scales where transgene expression was detectable (Figure 5D). Notably, *fgf20a* mutant zebrafish do not have a squamation phenotype unless present in an *fgfr1a* mutant background, suggesting functional redundancy among Fgf ligands (Daane et al., 2015). Those results and our present findings together suggest that Eda-Edar-NF- κ B signaling regulates SFC differentiation via multiple epidermal Fgf ligands, including Fgf20a.

Unlike Eda signaling, very little is known of potential TH targets in the skin. Therefore, we compared between wild-type and hypoTH backgrounds the expression of transcripts within each of 5 major cell types. This analysis revealed substantial differences in gene expression between backgrounds in dermal and epidermal cell types, and particularly in basal cells of the epidermis (Figure 6A). Given the absence of superficial pre-SFC in hypoTH skin (Figure 4B–D), we hypothesized that TH regulates the expression of cues in epidermal basal cells that recruit dermal cells to the most-superficial layer of the dermis, subjacent to the epidermis, prior to scale papilla formation (Aman et al., 2021). To test this idea we examined ligand genes with detectable expression across each of 7 major pathways between wild-type and hypoTH basal cells, which revealed markedly reduced expression for several non-FGF MAPK ligand genes, including PDGF α orthologs (*pdgfaa*, *pdgfab*) that are known in amniotes to regulate mesenchymal cell motility and proliferation (Figure 6B–D) (Karlsson et al., 1999). If PDGF α ligands are responsible for recruiting dermal cells similarly in zebrafish skin, then restoring expression of *pdgfaa* in basal cells of the epidermis in hypoTH fish should rescue the formation of superficial dermal cells in this background. When we forced expression of *Pdgfaa* in basal cells of epidermis by heatshock induction and stringent selection of basal epidermal expression in F0 mosaics, we found, as predicted, a recruitment of dermal cells in hypoTH skin, leading to a locally stratified dermis (Figure 6E) similar to that of the wild-type (Figure 4C). Early, heatshock-induced *Pdgfaa* expression also led to precocious dermal stratification in wild-type and *eda* mutant fish (Figure 6—figure supplement 1A). *Pdgfaa* expression did not, however, rescue the onset of squamation in hypoTH fish, which begins at a larger size and only after about twice the time as in wild-type fish (14 SSL vs. 9 SSL, 40 vs. 21 days post-fertilization in our rearing conditions) (Aman et al., 2021). Nor did *Pdgfaa* lead to mis-patterned and dysmorphic scales in wild-type fish, or a rescue of squamation in *eda* mutants, as we observed for Fgf20a (Figure 5D). Together these observations suggest that *Pdgfaa*-dependent stratification of dermis and Fgf-dependent differentiation of SFC are functionally decoupled processes that occur sequentially during skin morphogenesis (Figure 6—figure supplement 1B). Because *Pdgfaa* rescued dermis stratification, but not scale development in hypoTH skin, we predict that additional TH transcriptional targets regulate skin morphogenesis. Indeed, differential expression analyses suggested several excellent

candidates for mediating additional signals from epidermal basal cells to pre-SFC (**Figure 6—figure supplement 1C**).

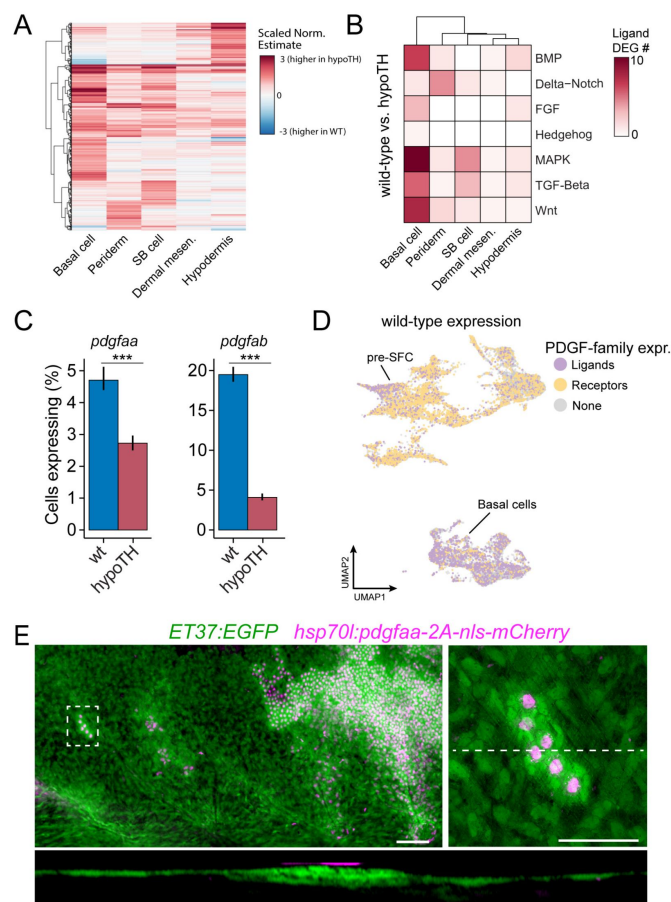


Figure 6.

TH drives dermal stratification via transcriptional regulation of epidermal *Pdgfa* expression. **(A)** Differential gene expression analysis between cell types of wild-type and hypoTH fish revealed extensive changes in expression across dermal and epidermal cell types ($n = 836$ genes, q -value < 0.01 , normalized effect > 2) (**Supplementary file 2—Table 4**). SB cell, suprabasal cell. **(B)** Of the differentially expressed genes, ligands of major signaling pathways involved in morphogenesis are also enriched in basal cells. **(C)** Both *pdgfaa* and *pdgfab* ligands are differentially expressed (*** q -value $< 1e-10$) between wild-type and hypoTH basal cells of the epidermis [error bars estimated via bootstrapping ($n = 100$)]. **(D)** Wild-type dermal and basal cells of epidermis plotted in UMAP space and colored by whether they express *pdgfaa*, *pdgfab* or both (ligands) as well as *pdgfra*, *pdgfrb* or both (receptors). **(E)** Upper left: Mosaic heat-shock induction of *Pdgfaa* (magenta), stringently selected for expression in epidermal basal cells, rescued stratification of hypoTH dermis, visualized with *ET37:EGFP* (green) ($n = 65$ clones in 8 fish). Upper right: Higher magnification of boxed area showing accumulation of dermal cells underneath *Pdgfaa*+ epidermal cells. Bottom: Optical cross section of boxed area reveals multiple dermal layers only in proximity to *Pdgfaa*+ epidermal cells. Scale bars, 50 μ m (E, upper left panel), 10 μ m

(E, enlarged region, upper right and lower panels).

These observations from scaleless skins indicate that epidermal basal cells are critical targets for TH and Eda-Edar-NF- κ B signals, and that epidermally expressed *Pdgfaa* and *Fgf* ligands link TH and Eda signaling to dermal cell recruitment and dermal papilla development, respectively (**Figure 7**). These findings are of potential clinical significance as the pathophysiology underlying TH skin diseases remain unclear (Mancino et al., 2021).

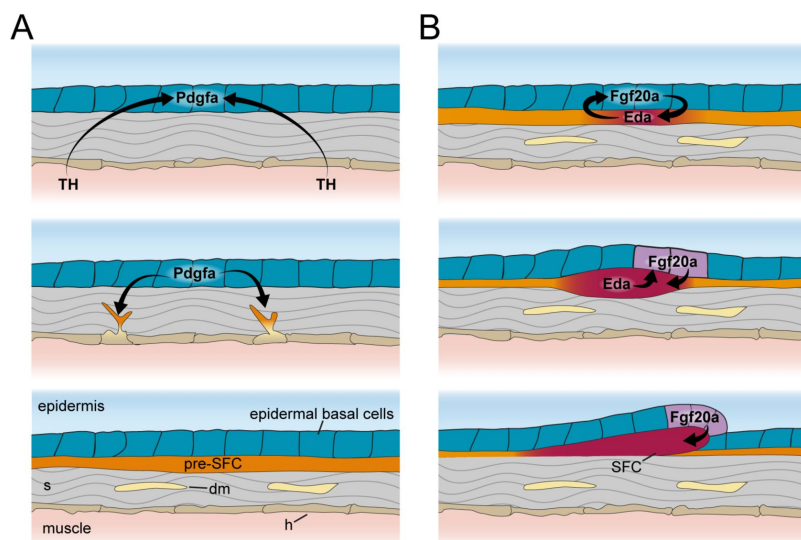


Figure 7.

Schematic representation of signaling interactions that regulate dermal morphogenesis. **(A)** Globally circulating TH (thyroid hormone) stimulates expression of transcripts encoding Pdgfa ligand in basal epidermal cells (blue). Epidermally expressed Pdgfa ligand regulates migration and differentiation of dermal mesenchyme in the stroma and pre-SFCs that accumulate just beneath the epidermis prior to scale development. **(B)** Epidermally expressed Fgf20a ligand stimulates differentiation of SFCs in the superficial pre-SFC population. Subsequently, SFCs express Eda ligand which, in turn, maintains expression of Fgf20a. The mechanisms that initiates squamation and

regulates hexagonal scale patterning remain unknown.

Hypodermal contribution to the microenvironment of stripe-forming pigment cells

Zebrafish pigment patterning is a useful study system for elucidating principles that govern developmental patterning and post-embryonic developmental progression (Patterson and Parichy, 2019). The alternating dark stripes of melanophores with sparse blue-tinted iridophores and light interstripes of yellow-tinted iridophores with orange xanthophores form deep in the skin (Hirata et al., 2005; Gur et al., 2020), which remains remarkably transparent throughout the life of the animal (Figure 1A,B). Although pigment cells are an integral part of the skin and comprise its major visual element, we know little about how these cells interact with other cell types in this microenvironment.

To better define how pigment cells are integrated with other skin cell types and identify tissue environmental factors that may influence pigment patterning, we included in our study the *bonaparte* mutant, homozygous for a presumptive loss of function mutation in *basonuclin 2* (*bnc2*) (Lang et al., 2009). *bnc2* mutants have a very sparse complement of pigment cells as adults owing to progressive pigment cell death during the larva-to-adult transition, with iridophores especially affected (Lang et al., 2009; Patterson and Parichy, 2013).

At the stage of tissue collection, fewer iridophores and melanophores were evident in *bnc2* mutants compared to wild-type controls, mirroring prior quantitative comparisons (Figure 8A). Previous analysis of genetic mosaics demonstrated that Bnc2 function is required in the hypodermis for survival and patterning of pigment cells (Lang et al., 2009). Accordingly, we predicted that wild-type and *bnc2* mutant fish should have substantial differences in gene expression within the hypodermal cell population. Yet alignment of UMAP projections for wild-type and *bnc2* mutant cells revealed instead a profound deficiency of hypodermal cells themselves (Figure 8B). We quantified the proportion of various dermal cell types and pigment cells relative to wild-type controls in our sci-RNA-Seq dataset and found marked deficits in hypodermal cells, as well as iridophores, melanophores, and xanthophores,

whereas numbers of dermal mesenchyme cells were relatively unchanged between genotypes (**Figure 7C**).

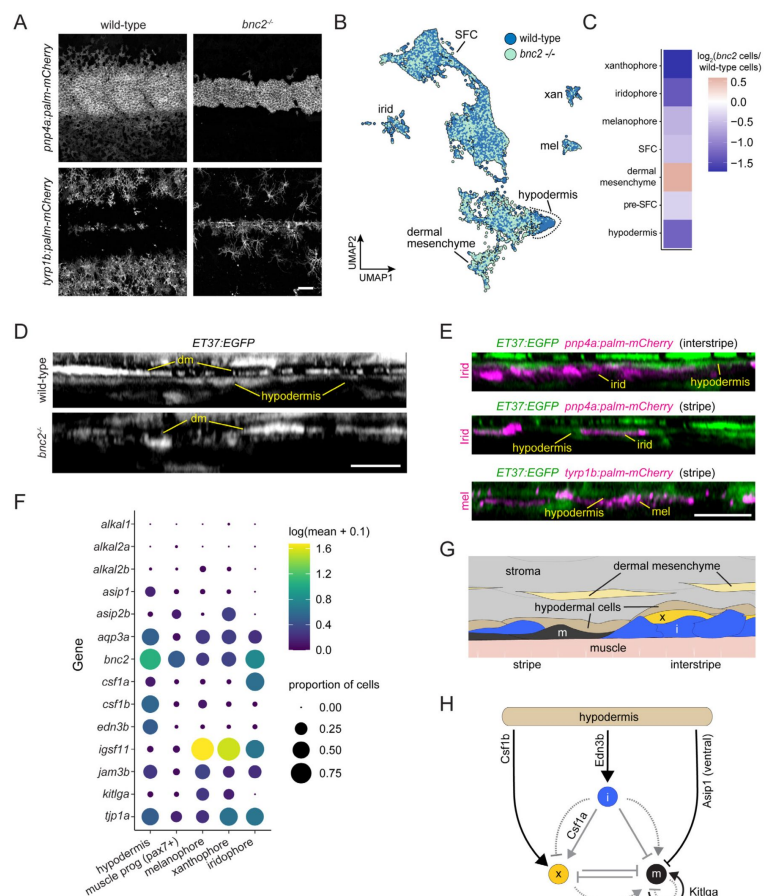


Figure 8.

Hypodermis is a pigment cell support cell population. **(A)** At 9.6 SSL, *bonaparte* mutants have grossly fewer iridophores (*pnp4a:paln-mCherry*) and melanophores (*tyrp1b:paln-mCherry*). **(B)** UMAP visualization of dermal cells and pigment cells with wt in blue and *bnc2* mutant in red shows specific deficiency in hypodermal cells. **(C)** Heatmap showing the log2 proportion of dermal cell subtypes and pigment cells. **(D)** Orthogonal projections of live, super-resolution imaging of dermal cells in wild-type and *bnc2* mutants expressing *ET37:EGFP*. Wild-type hypodermis is a thin, confluent cell layer underneath the more brightly labeled dermal mesenchyme (dm). Stage-matched *bnc2* mutant dermis had dermal mesenchyme, but lacked a hypodermal layer. **(E)** Live imaging of fish doubly transgenic for *ET37:EGFP* to visualize hypodermis, and *pnp4a:paln-mCherry* to visualize iridophores (irid) or *tyrp1b:paln-mCherry* to visualize melanophores (mel). Both pigment cell types reside in close contact with hypodermal cells. **(F)** Dot-plot heatmap showing expression level of known pigment cell trophic factors. **(G)** Schematic representation of pigment cell microenvironment, greatly expanded along its deep-to-superficial axis

to better illustrate organization of the very flat pigment and hypodermal cells. Xanthophore location inferred from (Hirata et al., 2005). **(H)** Potential regulatory linkages between hypodermis and pigment cell types, inferred from expression (black arrows). Previously documented interactions among the pigment cells represented by grey arrows. Scale bars, 100 μ m (A), 10 μ m (D,E).

Based on these findings, we predicted that hypodermis would be malformed or missing in live *bnc2* mutants. To test this prediction and validate inferences from the single cell transcriptome data, we imaged the dermis of live fish expressing *ET37:EGFP* (Parinov et al., 2004; Aman et al., 2021). In wild-type controls, the hypodermis appeared as a thin, nearly confluent cell layer deep in the dermis, whereas in *bnc2* mutants no such layer was apparent; instead the deepest dermis appeared to contain only dermal mesenchyme (**Figure 8D**; **Figure 8—figure supplement 1A**; **Figure 1—figure supplement 1B**). Consistent with the hypodermis having roles in supporting pigment cells, we observed iridophores, expressing *pnp4a:paln-mCherry* and melanophores, expressing *tyrp1b:paln-mCherry* in close proximity to *ET37:EGFP*⁺ hypodermal cells in wild-type skin (**Figure 8E**) (McMenamin et al., 2014; Lewis et al., 2019). It is possible that hypodermal cells physically persist in *bnc2* mutants but have sufficiently altered transcriptional profiles such that they no longer cluster together with wild-type hypodermal cells or express the *ET37:EGFP* transgene. Nevertheless, these analyses suggest that *ET37:EGFP*⁺ hypodermal cells likely play a role in pigment pattern development.

Given the close contact of wild-type melanophores and iridophores with hypodermal cells, together with loss of hypodermis and pigment cells in *bnc2* mutant fish, we hypothesized that factors important for pigment cell development and pattern formation are provided particularly by hypodermal cells, as opposed to other non-pigment cells of the local tissue environment (i.e., non-hypodermal stromal cells, developing SFCs, or superficially located muscle progenitor cells, also present in our data set). To test this idea, we examined the expression of genes encoding signaling ligands and adhesion factors implicated previously in pigment pattern development.

These analyses revealed that hypodermal cells are likely to be an important driver of iridophore development as they were the principal cells to express *endothelin 3b* (*edn3b*) (Figure 8F; Figure 8—figure supplement 1B), encoding a secreted protein that is processed to an active 21 amino acid peptide required by iridophores. Diminished expression of *edn3b* is associated with reduced numbers of iridophores and attenuated interstripes and stripes both in zebrafish mutants and in the naturally occurring pattern of the zebrafish relative, *D. nigrofasciatus* (Spiewak et al., 2018). Edn3 acts directly on iridophores but only indirectly on melanophores, through an interaction between these cells and Edn3-dependent iridophores (Parichy et al., 2000; Krauss et al., 2014; Spiewak et al., 2018).

More direct roles for hypodermal cells in regulating melanophore development were suggested by expression of other factors, including *agouti signaling protein 1* (*asip1*), which encodes a secreted protein that represses melanophore differentiation in ventral regions of the flank (Cal et al., 2019), though our dataset does not allow us to segment cells spatially. Interestingly, *kit ligand a* (*kitlga*), encoding a melanogenic factor that promotes survival, migration, and differentiation, was expressed at only moderate levels by hypodermal cells at this stage, despite its broad expression in the much simpler skin of embryos and early larvae (Hultman et al., 2007; Budi et al., 2011; Dooley et al., 2013). Nevertheless, *kitlga* was expressed at higher levels by melanophores themselves, whereas *junctional adhesion molecule 3b* (*jam3b*), encoding a 2-Immunoglobulin like domain adhesion receptor, was expressed by both hypodermal cells and melanophores. Jam3b mediates homophilic and heterophilic adhesive interactions, and is required autonomously by melanophores for an adherent phenotype; a Jam3b fusion protein accumulates at sites of overlap between mature melanophores (Ebnet, 2017; Eom et al., 2021). In the absence of Jam3b, melanophores tend to be hypopigmented, fail to acquire their mature, well-spread morphology and orderly arrangement, and many die. Together with prior studies, our observations suggest a model in which Jam3b facilitates interactions between immature melanophores and hypodermis, with subsequent Jam3b-mediated interactions between melanophores facilitating Kitlga-dependent maturation and survival (Figure 8H). That a second factor likely repressive for melanogenesis, *asip2b*, was expressed by xanthophores further suggests a mechanism for preventing differentiation of new melanophores within the interstripe.

Our analyses point to a role for hypodermal cells in regulating xanthophore differentiation as well. Xanthophores require signaling through Colony stimulating factor-1 receptor-a (*Csf1ra*) for migration, survival and differentiation (Parichy and Turner, 2003; Patterson and Parichy, 2013). One source of Csf1 is iridophores, which express *csf1a* (Figure 8F): in *bnc2* mutants xanthophores are tightly associated with residual iridophores. Nevertheless, xanthophores eventually cover the flank of *bnc2* mutants, as well as other iridophore-deficient mutants in which xanthophore development is delayed. This recovery presumably reflects expression of *csf1b*, encoding a ligand that is similarly potent to Csf1a in its ability to induce xanthophore differentiation, by a previously undefined population of cells in the skin (Patterson and Parichy, 2013). Our sci-RNA-seq dataset shows hypodermal cells to be the presumptive major source of Csf1b, though its transcripts were detected at lower levels in other dermal mesenchyme, superficial pre-SFCs and other cell types as well (Figure 8F-H; Figure 8—figure supplement 1C). To test requirements for Csf1 genes in adult pigmentation

we generated alleles having premature termination codons. As late juveniles (~16 SSL), fish homozygous for *csf1a* mutations had overtly normal pigment patterns on the trunk but less regular patterns of pigment cells on the fins as compared to wild-type ([Figure 8—figure supplement 2](#)). By contrast, fish homozygous for a *csf1b* mutation were deficient for pigmented xanthophores, evident particularly on the dorsum, which lacks *csf1a*-expressing iridophores in the hypodermis, though some xanthophores persisted along scale margins where a few iridophores occur (e.g., Supplementary Figure 4 of Gur et al., 2020). Fish doubly homozygous for *csf1a* and *csf1b* mutations lacked virtually all xanthophores, recapitulating the phenotype of *csf1ra* mutants. These observations support a model in which hypodermally derived Csf1b promotes xanthophore differentiation during normal development, and can substitute for iridophore-derived Csf1a in backgrounds deficient for iridophores; we presume that eventual recovery of xanthophores in *bnc2* mutants deficient for both iridophores and hypodermis reflects residual Csf1b availability from other dermal cell types.

Finally, genes that affect each of the major classes of pigment cell were expressed by hypodermal cells, yet *bnc2* mutants are particularly deficient for iridophores at early stages (Patterson and Parichy, 2013). We therefore predicted that transcriptomic states of iridophores would be more severely affected by loss of *bnc2* than transcriptomic states of melanophores or xanthophores. Consistent with this idea and a greater impact of *bnc2* loss on iridophores than other pigment cells, we found more differentially expressed genes in iridophores than xanthophores or melanophores (~1000 cells of each cell type) ([Figure 8—figure supplement 1D](#)). This disproportionality may be a consequence of the markedly fewer hypodermal cells and attendant loss of Edn3b or other iridogenic signals. Alternatively, *bnc2* may have more pronounced activities within iridophores, as these cells express *bnc2* and at levels greater than melanophores or xanthophores ([Figure 8F](#)), in addition to its non-autonomous functions in pattern formation through the hypodermis (Lang et al., 2009). Though further manipulative analyses will be needed to test these several interactions, our analyses of gene expression and cell-type abundance identify hypodermal cells as a key source of factors permissive, and possibly instructive, for adult interstripe and stripe development.

Discussion

Skin is a large, heterogenous and biomedically important organ and the skin of zebrafish is a useful system in which to elucidate mechanisms of skin patterning and morphogenesis. We have generated a minimally biased single-cell resolution transcriptional atlas of zebrafish skin at a key stage during the larva-to-adult transition, during squamation and pigment patterning. These data include transcriptomes for all major epidermal and dermal skin cell types in addition to numerous skin-associated cell types including pigment cells.

Zebrafish skin is endowed with an array of elasmoid scales, thin plates of calcified extracellular material deposited in the skin from dermal papillae that aggregate at the interface of dermis and epidermis. Calcified skin appendages in extant fish are diverse, encompassing various and sundry spines, plates, odontodes and scales. These forms are composed of extracellular matrices that range from among the hardest material in biology to some of the most flexible (Sire and Huyseune, 2003). We systematically assessed the expression of genes encoding non-collagen calcified matrix proteins throughout the skin during squamation, leading to the discovery of a transcriptionally distinct population of basal epidermal cells that express EMP transcripts, likely corresponding to epidermal secretory cells proposed to participate in scale matrix formation based on ultrastructure (Sire et al., 1997). These cells also express *dlx3a*, *dlx4a*, *runx2b* and *msx2a* but not *sp7*, a

transcription factor suite that is shared with ameloblasts that form tooth enamel. While these transcription factors are not exclusive to ameloblasts and have been reported in osteoblasts and odontoblasts, in addition to cell types that do not produce calcified matrix, such as neurons, their co-expression along with EMP-encoding transcripts in basal epidermal cells is consistent with a common origin of ameloblasts and the EMP+ epidermal cells reported here. One alternative hypothesis is that co-expression of these gene products arose convergently and can be explained by mechanistic linkages among them. Future work aimed at functionally dissecting the regulatory mechanisms that govern EMP gene expression in a variety of organisms may clarify these issues either by providing evidence of additional commonalities, supporting a shared ancestor, or by revealing diverse, lineage-specific regulatory architectures, supporting convergent evolution of superficial enamel deposition in teeth and fish skin appendages.

Additionally, the complement of genes encoding matrix proteins expressed by dermal SFC suggests that although these cells may share fundamental regulatory machinery with mammalian osteoblasts and odontoblasts, including regulation by Runx2 and Sp7 transcription factors, they likely produce a derived form of calcified matrix, fibrillary plate elasmoidin, which is distinct from bone or dentin. Independent patterning of epidermal EMP-expressing cells and dermal SFCs might underlie some of the morphological diversity among fish skin appendages. For example, it is possible that hard spines, as in pufferfish and armored catfish, are formed by patterned expression of epidermal EMP gene products. Manipulation of matrix protein expression in zebrafish SFC and EMP+ epidermal cells may provide a powerful new system to elucidate general mechanisms of biomineralization and genetic encoding of material properties that could, in turn, inform future research in clinical dentistry.

Scales develop from dermal papillae that form under the epidermis. The regulatory underpinnings of scale papillae patterning and morphogenesis depend on reciprocal epithelial-mesenchymal signaling interactions, including contributions of Eda-A-Edar-NF- κ B signaling, that are widely conserved across vertebrate skin appendages (Cui and Schlessinger, 2006; Harris et al., 2008; Aman et al., 2018). Analysis of scale development therefore affords a relatively accessible approach to understanding epithelial mesenchymal signaling interactions that underlie dermal morphogenesis. To this end, we generated and analyzed single cell transcriptomes for two scaleless conditions, *eda* mutants and hypoTH fish (Harris et al., 2008; McMenamin et al., 2014). Eda is a paracrine factor that binds receptors expressed in the epidermis and thyroid hormone is an endocrine factor with potential to regulate transcription in any cell (Cui and Schlessinger, 2006; Brent, 2012; Aman et al., 2018). Despite widely different spatial ranges over which signals are transmitted, our transcriptomic analysis suggests that both molecules regulate transcription of signaling ligands in basal epidermal cells that ultimately affect dermal morphogenesis. We further showed that Eda signaling indirectly regulates SFC differentiation by triggering expression of Fgf ligands. TH is implicated in human dermatopathies; myxedema, characterized by dry, waxy skin, is clinically synonymous with hypothyroidism (Safer, 2011). Yet we know remarkably little about cutaneous transcriptional targets of TH. Our analyses show that genes encoding PDGF α ligands are transcriptionally regulated by TH and can themselves regulate dermal-epidermal morphogenesis. This finding may be of relevance to understanding and potentially treating skin conditions associated with treatment-resistant TH insensitivity. Indeed, PDGF α and TH gain and loss of function studies in mouse yield similar hair cycle phenotypes (Safer et al., 2001; Tomita et al., 2006; Contreras-Jurado et al., 2015; González et al., 2017).

The eponymous striped pattern of zebrafish arises from neural crest derived pigment cells that reside deep within the skin, beneath the concurrently forming scales (Le Guellec et al., 2004; Hirata et al., 2005), and depends on interactions among all three pigment cell types

(Patterson and Parichy, 2019). Although much has been learned about stripe pattern formation from analyses of mutants lacking one or more pigment cell types, much less is known about how pigment cells integrate into the skin microenvironment. Analyses of genetic mosaics have hinted at an important role for skin cells (Lang et al., 2009; Krauss et al., 2014; Patterson et al., 2014; Eskova et al., 2017); still, this aspect of pigment patterning remains largely unexplored empirically or computationally (Volkening and Sandstede, 2015; Watanabe and Kondo, 2015; Volkening and Sandstede, 2018; Owen et al., 2020). Using single cell transcriptomics and live imaging of wild-type and *bnc2* mutant fish, we have identified a discrete population of dermal cells express genes that regulate differentiation and morphogenesis of pigment cells, including *edn3b*, which is required for iridophore population expansion.

Our analyses have focused on scales and pigmentation, however, zebrafish skin is a fruitful study system for many areas of biology including regeneration and wound healing (Richardson et al., 2016; Cox et al., 2018; Iwasaki et al., 2018; Morris et al., 2018; Pfalzgraff et al., 2018); innate immunity (Lü et al., 2015; Wurster et al., 2021), stem cell regulation (Lee et al., 2014; Chen et al., 2016; Brock et al., 2019), sensory physiology and developmental neuroscience (Rasmussen et al., 2015; Rasmussen et al., 2018; Peloggia et al., 2021), and human disease modeling (Feitosa et al., 2011; Li et al., 2011). Additionally, considerable insight into general mechanisms of development can emerge from comparing developmental mechanisms across species or between organ system. We expect the transcriptomic data presented here will help in identifying useful markers for cross-species comparisons, enabling a deeper understanding of the molecular and cellular bases of phenotypic evolution.

Materials and Methods

Zebrafish lines and husbandry

Fish were maintained in the WT(ABb) background at 28.5°C. Lines used were: *Tg(sp7:EGFP)^{b1212}* abbreviated *sp7:EGFP* (DeLaurier et al., 2010), *Et(krt4:EGFP)^{s4et37}* abbreviated *ET37:EGFP* (Parinov et al., 2004), *Tg(sp7:nEOS)^{vp46rTg}* (Aman et al., 2021), *bnc2^{utr16e1}* (Lang et al., 2009), *eda^{dt1261}* (Harris et al., 2008), *Tg(tg:nVenus-v2a-nfnB)^{wprt8Tg}* abbreviated *tg:Venus-NTR*, *Tg(tyrb1b:palm-mCherry)^{wprt11Tg}* (McMenamin et al., 2014), and *Tg(pnp4a:palm-mCherry)^{wprt10Tg}*. Thyroid ablation and rearing of hypoTH fish were done as previously described (McMenamin et al., 2014). *csf1a* and *csf1b* mutant fish were generated by injecting CRISPR/Cas9 reagents (PNAbio) including synthetic single-guide RNA targeting the genomic sequences (*csf1a*: 5'-GCGGCATTCCTCACATAC; *csf1b*: 5'-GGCATGTTTGCAAGGACCG) into zygotes, selecting phenotypic F0 animals, and repeat outcrossing to generate F2 families (Hwang et al., 2013). Recovered alleles contained premature termination codons owing to frame shift mutations (*csf1a*: 10 bp insertion; *csf1b*: 2 or 5 bp deletions having phenotypes indistinguishable from one another).

Imaging

Alizarin-Red-S vital dye, MS-222 anesthesia and mounting for microscopy were performed as previously described (Aman et al., 2021). Images in Figure 1—figure supplement 1B, Figure 2C,D, Figure 4C, Figure 5E and Figure 8A,D,E were acquired on a Zeiss LSM880 in fast Airyscan mode. Images in Figure 2—figure supplement 1A and Figure 4A were acquired on a Zeiss LSM880 in conventional confocal mode. Images in Figure 1A, Figure 1—figure supplement 1C, Figure 3C,E, Figure 5D, Figure 6E, Figure 6—figure supplement 1A,B, were

acquired on a Zeiss Observer equipped with Yokogawa CSU-X1 spinning disc. Images in Figure 7—figure supplement 2 were acquired on a Zeiss SteREO Discovery.V12 stereomicroscope. Orthogonal views were produced using FIJI (Schindelin et al., 2012). Brightness and contrast were adjusted in Adobe Photoshop and non-linear gamma adjustments were applied to images when necessary to highlight relevant cell types. Photoconversion of nuclear EOS was done using a 405 nm laser on a Zeiss LSM800.

mRNA *in-situ* hybridization

All in-situ hybridization probe templates were amplified using PrimeStar-GXL (Takara) from cDNA prepared with SSIII (ThermoFisher) with the following primers: ambn 5'-TGATGATCGTGTGCTTTCTTGCTG, 5'-aaaaTAATACGACTCACTATAGCATTGCCCCCTGTTGTGGTCTTG; itga5 5'-AGGAAGGAAGTGTACATGGGTGA, 5'-aaaaTAATACGACTCACTATAGgatccagttttgtcccagatgac; itgb3b 5'-TGGACCTGTCCTACTCCATGAAT, 5'-aaaaTAATACGACTCACTATAGacactgtcttttagcgtgtcc; col10a1a 5'-gaaccaagtatgccgatttgacc, 5'-aaaaTAATACGACTCACTATAGgttttgatgtgatgtggatgggt; col10a1b 5'-gcttagcttcagaaaATGGACCTCA, 5'-aaaaTAATACGACTCACTATAGTGGTTGTCCTTTTTCACCTGGATA; tcf7 5'-CCAACAAGGTGTCGGTGGT, 5'-'aaaaTAATACGACTCACTATAGACCAGTCCGTCTGttggttcag; jag1a 5'-CCCTTGACCAAACAAATGACAA, 5'-aaaaTAATACGACTCACTATAGGCTGTGTTTCTTCAGGTGTGG. chad 5'-AGACCAAACATCCAGACAGCAA, 5'-aaaaTAATACGACTCACTATAGGCAATTGCATCATCCTTCACAT. *In-situ* hybridization probes and tissue were prepared as described previously (Quigley et al., 2004), with hybridization and post-hybridization washes performed on a BioLane HTI 16Vx platform (Intavis Bioanalytical Instruments) and post-staining vibratome sectioning in some cases as described (Aman et al., 2018).

Heatshock transgene cloning and expression

Full length coding sequences were amplified from SSIII cDNA (ThermoFisher) using PrimeStar-GXL polymerase (Takara) and the following primers: *fgf20a* 5'-AAGCAGGCTCACCATGGGTGCAGTCGGCGA; 5'-GACTGCACCCATGGTGAGCCTGCTTTTTGTACAACTTGG; *pdgfaa* 5'-GCAGATATAAGGTGCGCCAGCGTCACCCA, 5'-CGCGTTCTCATGGTGAGCCTGCTTTTTGTACAACTTGG. Coding sequences were cloned into a hsp70l heatshock misexpression vector from (Aman et al., 2018) using NEBuilder HiFi DNA Assembly Master Mix (NEB). Zygotes were injected and raised to 8.5 SSL and given 6 x 1 hour 41°C heatshocks per day for 7 days in a modified Aquaneering rack. Only individuals with epidermal basal cell expression were selected for analysis.

Tissue dissection and storage

Fish were staged according to (Parichy et al., 2009) and 9.6 SSL individuals were selected for dissection, euthanized with MS-222 and processed immediately. Following removal of the head and fins, skins of wild-type controls, *eda* mutant homozygotes, *bnc2* homozygotes and hypoTH fish in an *sp7:EGFP* transgenic background zebrafish were removed with forceps and immediately flash frozen in liquid nitrogen then stored at -80°C prior to isolation of nuclei (*n* = 300 fish skins total).

Nuclei isolation and sci-RNA-seq2 library preparation

Separately for each background, frozen skins ($n = \sim 60$) were thawed over ice in cold lysis buffer (10 mM Tris-HCl, pH 7.4, 10 mM NaCl, 3 mM MgCl₂, 0.1% IGEPAL CA-630) (Cao et al., 2019) supplemented with 5% Suprase RNA Inhibitor and minced with a razorblade until no visible pieces remained (< 1 min). The cell suspension was then pipetted a few times and put through a 50 μ M filter into 10 ml of fixation buffer (5% Paraformaldehyde, 1.25X PBS) (Srivatsan et al., 2020). Nuclei were fixed on ice for 15 minutes then centrifuged at 700 $\times g$ for 10 minutes. Fixed nuclei were subsequently rinsed twice with 1 ml of nuclei resuspension buffer (NSB: 10 mM Tris-HCl, pH 7.4, 10 mM NaCl, 3 mM MgCl₂, 1% Suprase RNA Inhibitor, 1% 0.2 mg/ml Ultrapure BSA), spun down at 750 $\times g$ for 6 minutes and incubated in 400 μ L of permeabilization buffer (NSB + 0.25% Triton-X) for 3 minutes on ice. Permeabilized nuclei were spun down, resuspended in 400 μ L of NSB and sonicated on 'low' for 12 seconds. Following sonication, nuclei were spun down once more, resuspended in 400 μ L of NSB, and nuclei from each sample were DAPI-stained and counted on a hemocytometer. Sci-RNA-seq2 libraries were then prepared as previously described (Cao et al., 2017). Briefly, 1,200 nuclei in 2 μ L of NSB and 0.25 μ L of 10 mM dNTP mix (Thermo Fisher Scientific, cat no. R0193) were distributed into each well of 12 96-well plates – 4 per background (LoBind Eppendorf). Then 1 μ L of uniquely indexed oligo-dT (25 μ M) (Cao et al., 2017) was added to every well, incubated at 55°C for 5 minutes and placed on ice. 1.75 μ L of reverse transcription mix (1 μ L of Superscript IV first-strand buffer, 0.25 μ L of 100 mM DTT, 0.25 μ L of Superscript IV and 0.25 μ L of RNaseOUT recombinant ribonuclease inhibitor) was then added to each well and plates incubated at 55°C for 10 minutes and placed on ice. Wells were pooled, spun down and resuspended in 500 μ L NSB and transferred to a flow cytometry tube through a 0.35 μ m filter cap; DAPI was added to a final concentration of 3 μ M. Pooled nuclei were then sorted on a FACS Aria II cell sorter (BD) at 300 cells per well into 96 well LoBind plates containing 5 μ L of EB buffer (Qiagen). After sorting, 0.75 μ L of second strand mix (0.5 μ L of mRNA second strand synthesis buffer and 0.25 μ L of mRNA second strand synthesis enzyme, New England Biolabs) were added to each well, second strand synthesis performed at 16°C for 150 minutes. Tagmentation was performed by addition of 5.75 μ L of tagmentation mix (0.01 μ L of a N7-only TDE1 enzyme (in-house) in 5.7 μ L 2x Nextera TD buffer, Illumina) per well and plates incubated for 5 minutes at 55°C. Reaction was terminated by addition of 12 μ L of DNA binding buffer (Zymo) and incubated for 5 minutes at room temperature. 36 μ L of Ampure XP beads were added to every well, DNA purified using the standard Ampure XP clean-up protocol (Beckman Coulter) eluting with 17 μ L of EB buffer and DNA transferred to a new 96-well LoBind plate. For PCR, 2 μ L of indexed P5, 2 μ L of indexed P7 (Cao et al., 2017) and 20 μ L of NEBNext High-Fidelity master mix (New England Biolabs) were added to each well and PCR performed as follows: 75°C for 3 minutes, 98°C for 30 seconds and 19 cycles of 98°C for 10 seconds, 66°C for 30 seconds and 72°C for 1 minute followed by a final extension at 72°C for 5 minutes. After PCR, all wells were pooled, concentrated using a DNA clean and concentrator kit (Zymo) and purified via an additional 0.8X Ampure XP cleanup. Final library concentrations were determined by Qubit (Invitrogen), libraries visualized using a TapeStation D1000 DNA Screen tape (Agilent) and libraries sequenced on a Nextseq 500 (Illumina) using a high output 75 cycle kit (Read 1: 18 cycles, Read 2: 52 cycles, Index 1: 10 cycles and Index 2: 10 cycles).

Pre-processing of sequencing data

Sequencing runs were demultiplexed using bcl2fastq v.2.18 and expected PCR barcode combinations. The backbone computational pipeline for read processing was previously published (Cao et al., 2017). Following assignment of RT indices, reads were trimmed using trim-galore and mapped to the zebrafish transcriptome (GRCz11 with extended 3' UTRs)

(Saunders et al., 2019) using the STAR aligner (Dobin et al., 2013). Reads were then filtered for alignment quality, and duplicates were removed. Non-duplicate genes were assigned to genes using bedtools (Quinlan and Hall, 2010) to intersect with an annotated gene model. Cell barcodes were considered to represent a real cell if the number of UMIs was greater than 600, a number chosen based on a user-defined threshold on the knee plot. Cells with greater than 6000 UMIs were also discarded as likely multipliers. Reads from cells that passed the UMI thresholds were aggregated into a count matrix and then loaded and saved as a CDS object for downstream analysis with Monocle3 (Cao et al., 2019).

Dimensionality reduction, alignment and background correction

The wild-type only ($n = 35,114$) and all-background (wild-type, *eda* mutant, *bnc2* mutant, hypoTH; $n = 144,466$) CDS objects were processed separately. Cells were assigned to their background by matching recovered RT barcode information to the original plate loadings. For each dataset, the standard monocle3 processing workflow was followed (estimate_size_factors(), detect_genes(), preprocess_cds()) and the top 50 PCs were retained, and PCA was calculated using all genes as input. A few corrections were then made on the original PCA matrix. First, to account for possible cytoplasmic RNAs in the supernatant of each sample that could contribute to “background” in the resulting transcripts assigned to individual cells, we performed a sample-specific background correction as previously described (Packer et al., 2019). Briefly, the background distribution of RNA from was calculated from “cells” that had less than 15 UMIs and we used this to compute a “background loading.” Next, a linear regression model was fit using these background loadings (real cell PCA matrix ~ cell background loadings), and its residuals were considered the “background corrected PCA matrix.” This corrected PCA matrix was then subject to Mutual Nearest Neighbor (MNN) alignment (Haghverdi et al., 2018) by sample using the “align_cds” function in monocle3. The background corrected, MNN-aligned PCA matrix was then used as input for Uniform Manifold Approximation and Projection (UMAP) (Becht et al., 2018) dimensionality reduction using the “reduce_dimension” function and default settings (except `umap.min_dist = 0.15`, `umap.n_neighbors = 20L`). Clustering was performed with “cluster_cells” (wild-type resolution = $2e-4$; all-background resolution = $1e-4$), which uses the Leiden community detection algorithm (Traag et al., 2019). Clustering resolution was selected manually based on clear distinction of non-adjacent groups of cells and a reasonable recovery of overall UMAP structure.

Cell type classification and trajectory analysis

For each cluster in the wild-type dataset, the most specific genes were calculated using the “top_markers” function. These genes were sorted by specificity and clusters were annotated by comparing genes to published studies and *in-situ* hybridization databases. We assigned 43 clusters to 33 unique cell types and one “unknown” group when the cell type was not able to be determined based on gene expression. To annotate cells from the *eda* mutant, *bnc2* mutant and hypoTH backgrounds, we built a marker-free cell-type classifier with the wild-type cell annotations using Garnett (Pliner et al., 2019) and applied it to the remaining cells. Trajectory analysis was performed on a sub-setted and re-processed set of cells from the wild-type dermis. These cells form the developing scales and we chose a single stage that contained scale-forming cells along the entire developmental trajectory. After repeating dimensionality reduction, we applied the monocle3 function “learn_graph” and “order_cells” to root the graph and calculate “pseudotime” values for each cell that increased as a function of principal-graph distance from the root.

Analysis of cell type abundance differences

To compute cell type abundance differences across genotypes, cell counts for each type were normalized by the sample's size factor (the total cell counts from each sample divided by the geometric mean of all sample's total cell counts). The abundance difference was then calculated as $\log_2(\text{normalized query cell count} / \text{normalized reference cell count})$ for each cell type relative to wild-type for all backgrounds.

Differential expression analysis

Differentially expressed genes were computed by fitting the size-factor normalized UMI counts for each gene from each individual nucleus with a generalized linear model using the “fit_models” function in monocle3. To fit the regression model for each background's effect on each gene in each cell type, we first selected the pair-wise backgrounds and cells that were relevant for the model (i.e. Basal cells from wild-type and *eda*). Then we filtered for genes expressed in at least 10 cells, and used background as the model formula (model_formula_str = “~genotype”) and extracted the coefficient table, p-values and multiple testing corrected q-values with the “coefficient_table” function. Genes were considered significantly background-dependent differentially expressed (DEGs) if their q-value was less than 0.05. For analysis of specific pathways, we used *D. rerio* gene-pathway associations from WikiPathways (Martens et al., 2021).

Data availability

The datasets generated and analyzed during the current study are available in the NCBI Gene Expression Omnibus (GEO) repository: GSE224695.

Acknowledgements

Supported by NIH R35 GM122471 and NIH R01 AR078320 (DMP) as well as NIH U54 HL145611, NIH UM1 HG011586, NIH R01 HG010632 to CT and the Paul G. Allen Frontiers foundation (Allen Discovery Center) (CT). Thanks to Amber Schwindling, other Parichy lab members, and Raman Sood for assistance and oversight of fish care.

Author contributions

Andrew J Aman, Conceptualization, Resources, Data curation, Formal analysis, Funding acquisition, Validation, Investigation, Visualization, Methodology, Writing—original draft, Writing—review and editing; Lauren M Saunders, Conceptualization, Resources, Data curation, Formal analysis, Validation, Investigation, Visualization, Methodology, Writing—original draft, Writing—review and editing; August A Carr—Investigation; Sanjay R Srivatsan, Methodology; Colten D Eberhard, Resources; Blake Carrington, Resources; Dawn Watkins-Chow, Resources; William J Pavan, Resources; Cole Trapnell, Conceptualization, Resources, Funding acquisition; David M Parichy, Conceptualization, Resources, Data curation, Formal analysis, Funding acquisition, Visualization, Writing—original draft, Writing—review and editing

Ethics

Animal experimentation: This study was performed in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. All of the animals were handled according to approved institutional animal care and use committee (IACUC) protocol 4170 of the University of Virginia. For imaging and other procedures animals were anesthetized with MS222 or euthanized by overdose of MS222 and every effort was made to minimize suffering.

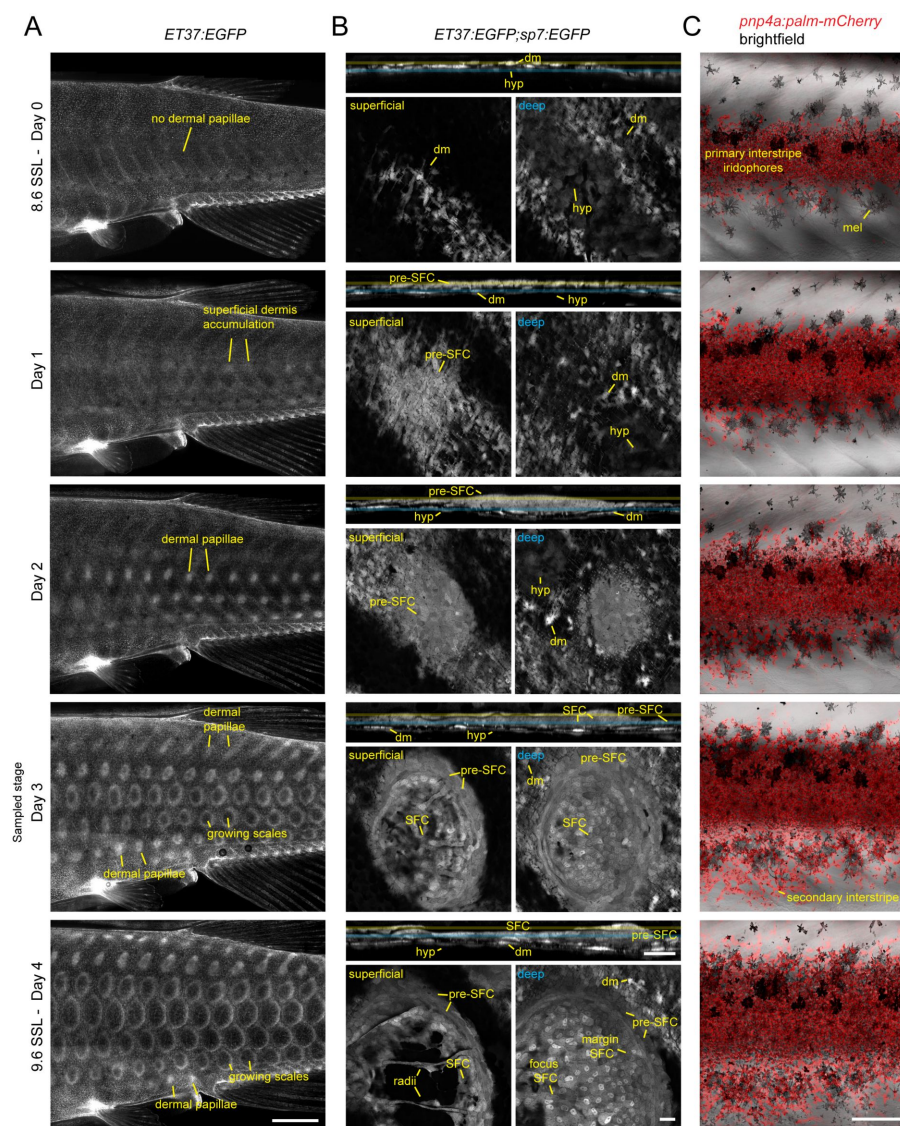


Figure 1—figure supplement 1.

Post-embryonic skin morphogenesis. (A) Overview of dermal development between 8.6 and 9.6 SSL, visualized by imaging fish transgenic for *ET37:EGFP*, which labels most dermal cells (Aman et al, 2021)(Parichy et al., 2009). Morphology on Day 3 of this series corresponds to the stage at which fish were selected for skin dissection and isolation of nuclei. (B) Development of an individual scale in *ET37:EGFP;sp7:EGFP* double transgenics. Each timepoint shows an orthogonal projection through the entire dermis above single confocal slices of superficial dermis (yellow) and deep dermis (blue). (C) Stripe and interstripe reiteration occurs during the same timeframe, with secondary interstripe initiation occurring at the sampled stage. Each series was made by repeated imaging of individual specimens. hyp, hypodermis; dm, dermal mesenchyme; mel, melanophore. Scale bars, 1 mm (A), 10 μ m (B), 500 μ m (C).

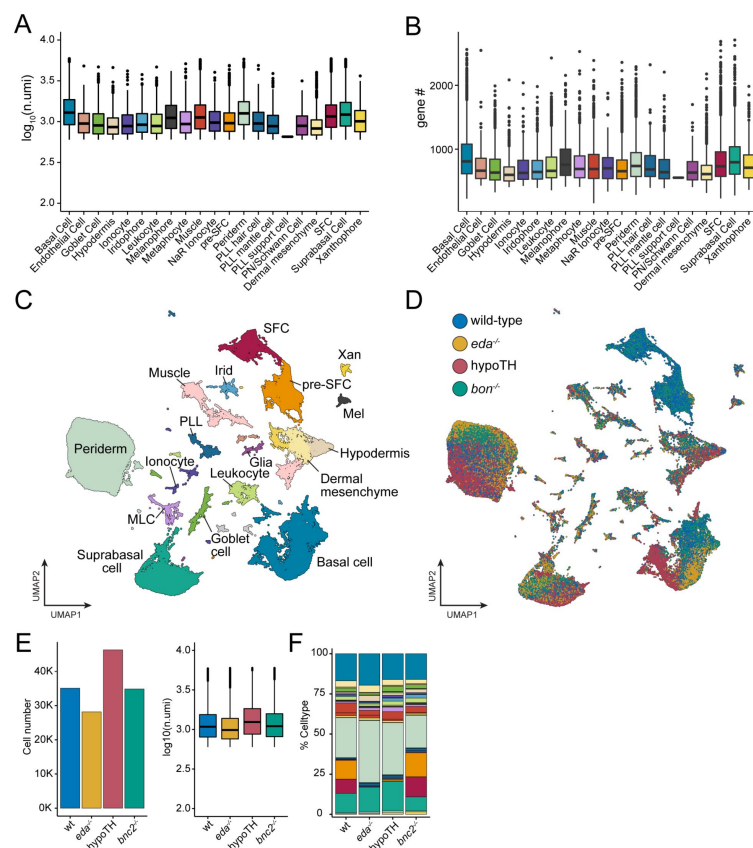


Figure 1—figure supplement 2.

sci-RNA-seq data quality metrics across back-grounds. **(A, B)** Counts of unique molecular identifiers (UMIs) and unique genes expressed across annotated cell types (shown are medians with boxes spanning interquartile ranges; vertical lines indicate farthest observations of data with outlier samples shown individually). **(C)** UMAP plot with cells from all genetic backgrounds, colored by cell type annotation with major groups labelled (cell $n = 144,466$). **(D)** The same UMAP plot as (C) with cells colored by their genetic background of origin (wild-type, *eda*^{-/-}, *hypoTH*, and *bnc2*^{-/-}). **(E)** Recovered cell counts from each genetic background and boxplots displaying UMI counts across backgrounds. **(F)** Cell type percentages across backgrounds. Colors correspond to cell types in (C).

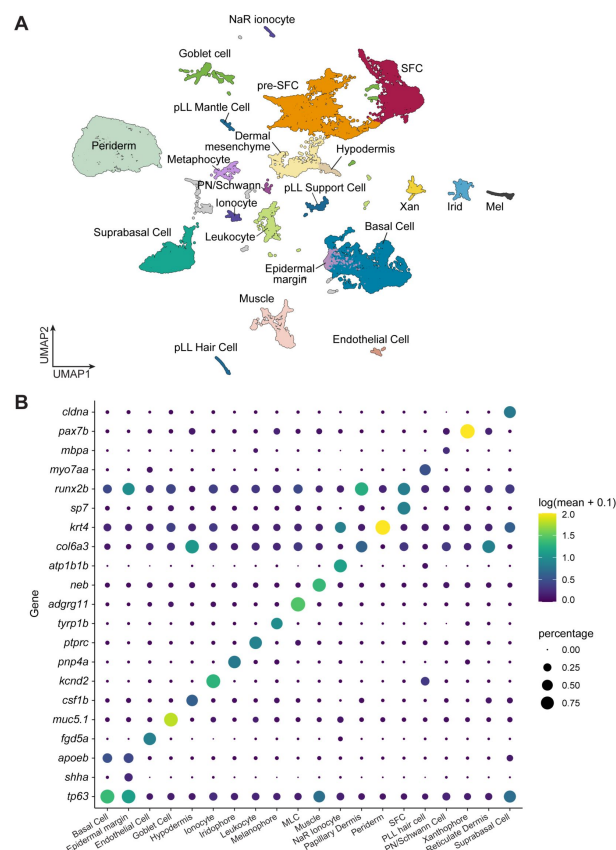


Figure 1—figure supplement 3.

A transcriptome atlas identifies major cell types in post-embryonic skin. **(A)** A UMAP plot for wild-type cells, colored and labelled by annotated cell types (cell $n = 35,114$ cells). **(B)** Dot-plot heatmap displaying known cell type marker genes and new candidate markers.

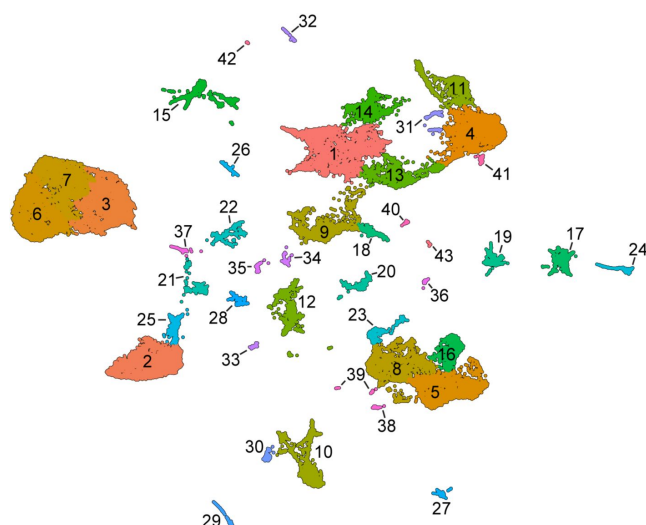


Figure 1—figure supplement 4.

Unsupervised Leiden clustering of wild-type skin cells. This analysis reveals additional, potentially biologically relevant cell-states nested within the cell types annotated by analysis of marker genes.

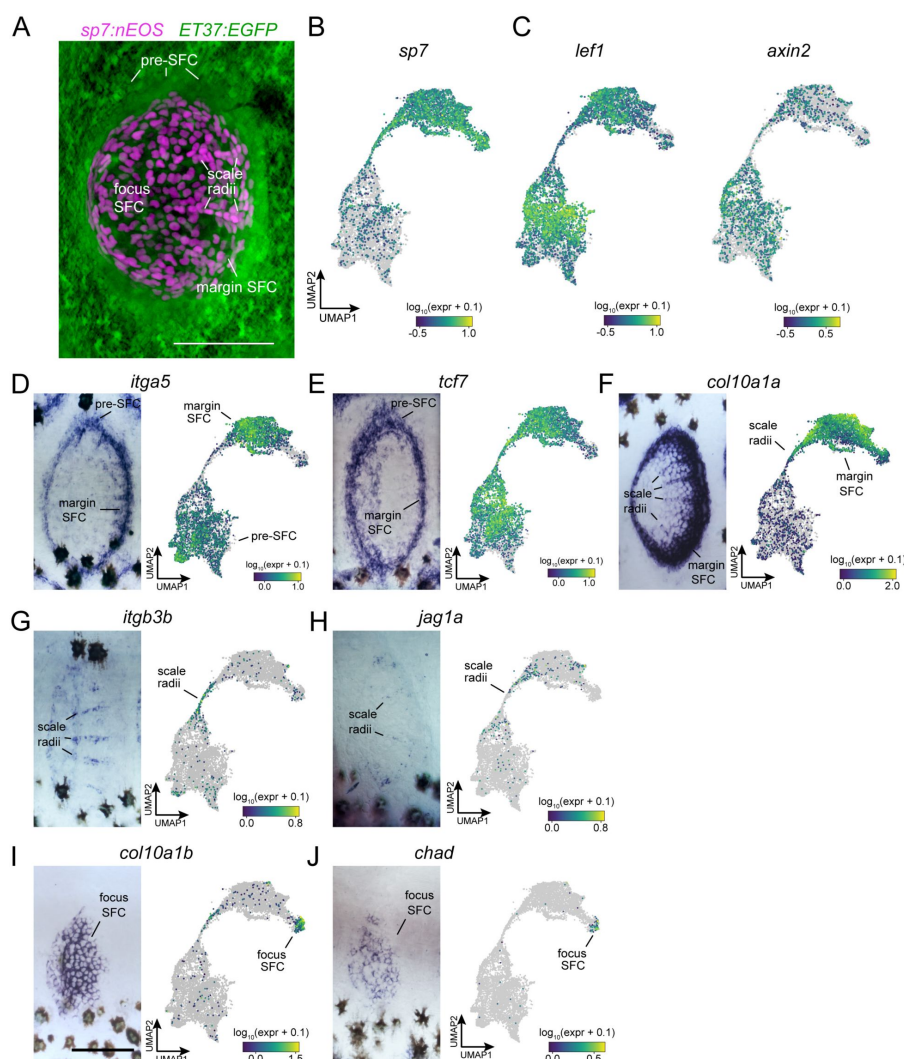


Figure 2—figure supplement 1.

In-situ hybridization using probes against specific transcripts localizes heterogeneous SFC in the developing scale. (A) Live imaging of differentiated SFC expressing *sp7:nEOS* (green) surrounded by a halo of pre-SFC labelled with *ET37:EGFP* (magenta). (B) UMAP visualization of *sp7*+ differentiated SFC and *sp7*- pre-SFC. (C) UMAP visualization of *lef1* and *axin2*, transcripts localized to pre-SFC and margin SFC in previously published *in-situ* hybridization (Aman et al 2018). (D,E) *itga5* and *tcf7* were expressed in pre-SFC and SFC at the scale margin. (F) *col10a1a* was highly expressed in scale margin SFCs and radii SFCs but not pre-SFCs. (G,H) *itgb3b* and *jag1a* localized to radii SFCs. (I,J) *col10a1b* and *chad* were expressed predominantly in SFC at the scale focus. Scale bars, 50 μ m.

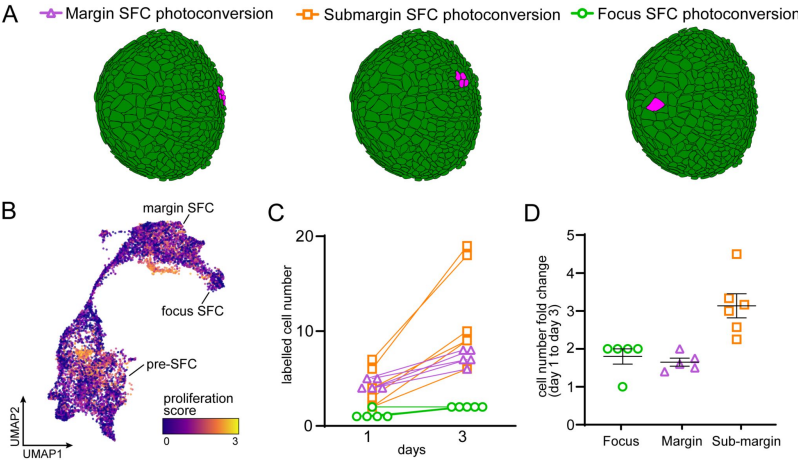


Figure 2—figure supplement 2.

SFC lineage and proliferation. (A) Schematic representation of photoconversion experiments in (Figure 3C) showing representative photoconversion regions. (B) UMAP visualization of transcripts associated with high rates of cell proliferation. (C) The number of labelled cells increased in every specimen examined. (D) Fold-change of labelled cell number (mean±SEM) over two days of scale growth (margin and submargin, $n = 5$ each; focus $n = 6$).

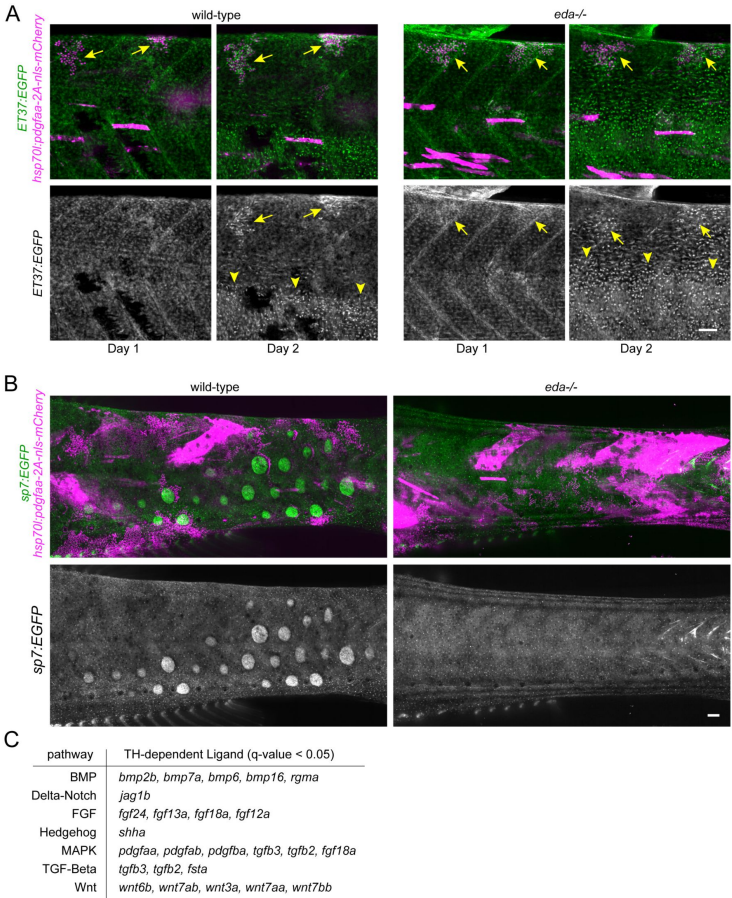


Figure 6—figure supplement 1.

Excess *Pdgfra* expression led to precocious dermal stratification in wild-type and *eda* mutant fish. (A) *Pdgfra* expression (magenta), stringently selected for basal epidermal expression, caused precocious dermal stratification, labelled with *ET37:EGFP* (green) in wild-type and *eda* mutants ($n = 8$ clones in 4 fish for wild-type; 9 clones in 4 fish for *eda* mutant). Arrows indicate regions of precocious dermal stratification. Arrowheads indicate regions where normal dermal stratification is occurring. (B) *Pdgfra* expression did not lead to dysmorphic and mis-patterned scales, visualized with *sp7:EGFP* (green) in wild-type and did not rescue scale formation in *eda* mutants ($n = 54$ clones in 8 wild-type fish, 70 clones in 10 *eda* mutant fish). Scale bars, 50 μ m (A and B). (C) Differentially expressed signaling ligands downregulated in hypoTH basal epidermal cells.

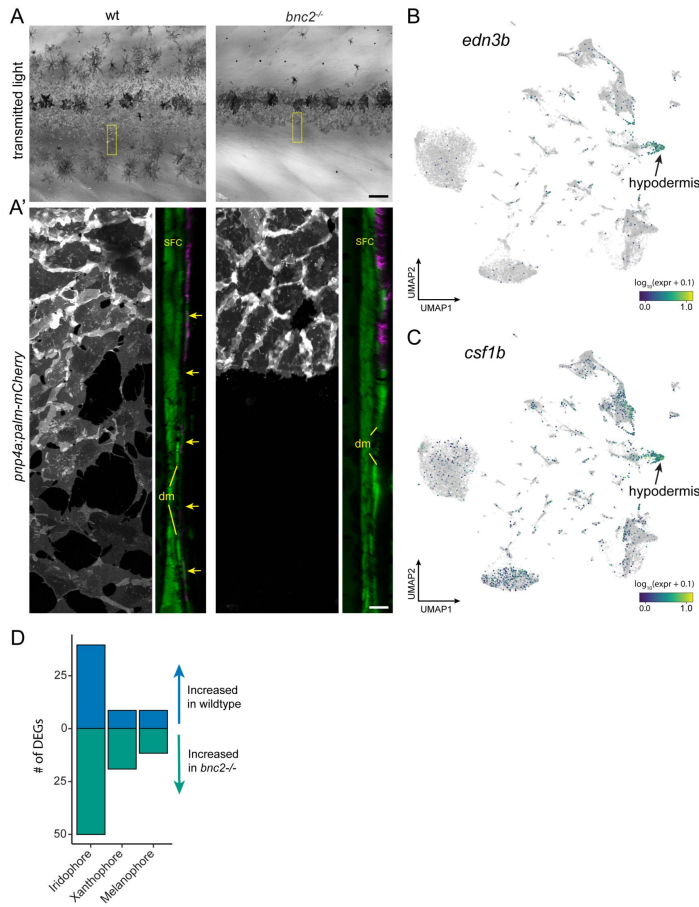


Figure 8—figure supplement 1.

Hypodermis supports pigment cells. **(A)** Iridophores and melanophores in wild type and *bnc2* mutant skins highlighting the relative dearth of pigment cells in the mutant. **(A')** Super-resolution images of boxed region in **(A)**, iridophores expressing *pnp4a:paln-mCherry* and orthogonal projection showing both iridophores and dermal cells expressing *ET37:EGFP*. *bnc2* mutant dermis is bounded by dermal mesenchyme (dm) and lacks the thin, confluent hypodermis seen in wild-type individuals (arrows). **(B,C)** UMAP visualizations of pigment cell trophic factors, *edn3b* and *csf1b*. Compare to cell types illustrated in **Figure 1C**. **(D)** *bnc2*-dependent loss of hypodermis is associated with a greater number of differentially expressed genes (DEGs) in iridophores than in other pigment cell types. Scale bars, 100 μ m (**A**), 10 μ m (**A'**).

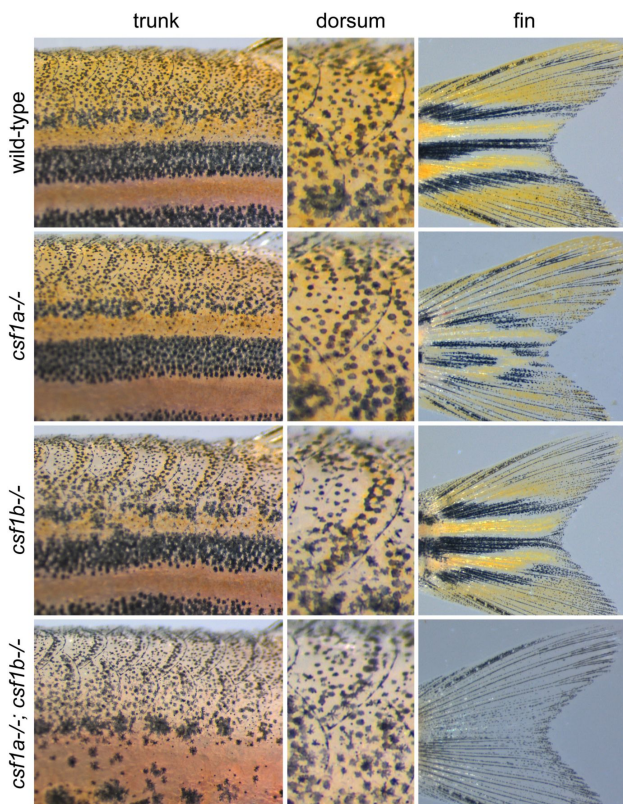


Figure 8—figure supplement 2.

Csf1 requirements for xanthophore pigmentation. Fish that were wild-type ($n = 23$), homozygous mutant for *csf1a* ($n = 15$), *csf1b* ($n = 9$) or both *csf1a* and *csf1b* ($n = 10$) were examined 4–12 weeks post-fertilization, with representative phenotypes at 6 weeks shown here. *csf1a* mutants had disrupted patterns and reduced xanthophore pigmentation in fins, whereas *csf1b* mutants had fewer pigmented xanthophores in the hypodermis and on scales, deficiencies especially apparent on the dorsum of the fish. Fish doubly mutant for both Csf1 loci lacked virtually all xanthophores and resembled *csf1a* mutants (Parichy and Turner, 2003).

References

- Aman AJ , Fulbright AN , Parichy DM (2018) **Wnt/beta-catenin regulates an ancient signaling network during zebrafish scale development** *Elife* **7**
<https://doi.org/10.7554/eLife.37001>
- Aman AJ , Kim M , Saunders LM , Parichy DM (2021) **Thyroid hormone regulates abrupt skin morphogenesis during zebrafish postembryonic development** *Dev Biol* **477**:205–18
<https://doi.org/10.1016/j.ydbio.2021.05.019>
- Aman AJ , Parichy DM , Cartner SC , Eisen JS , Farmer SC , Guillemin KJ , Kent ML , Sanders GE (2020) **Chapter 8 - Zebrafish Integumentary System** *The Zebrafish in Biomedical Research* :91–6
- Arola D , Murcia S , Stossel M , Pahuja R , Linley T , Devaraj A , Ramulu M , Ossa EA , Wang J (2018) **The limiting layer of fish scales: Structure and properties** *Acta Biomaterialia* **67**:319–30
<https://doi.org/10.1016/j.actbio.2017.12.011>
- Ayoola AO , Zhang BL , Meisel RP , Nneji LM , Shao Y , Morenikeji OB , Adeola AC , Ng'ang'a SI , Ogunjemite BG , Okeyoyin AO , Roos C , Wu DD (2021) **Population Genomics Reveals Incipient Speciation, Introgression, and Adaptation in the African Mona Monkey (*Cercopithecus mona*)** *Mol Biol Evol* **38**:876–90
<https://doi.org/10.1093/molbev/msaa248>
- Babajko S , de La Dure-Molla M , Jedeon K , Berdal A. (2015) **MSX2 in ameloblast cell fate and activity** *Frontiers in Physiology* **5**
<https://doi.org/10.3389/fphys.2014.00510>
- Bae JM , Clarke JC , Rashid H , Adhami MD , McCullough K , Scott JS , Chen H , Sinha KM , de Crombrughe B , Javed A. (2018) **Specificity Protein 7 Is Required for Proliferation and Differentiation of Ameloblasts and Odontoblasts** *J Bone Miner Res* **33**:1126–40
<https://doi.org/10.1002/jbmr.3401>
- Becht E , McInnes L , Healy J , Dutertre CA , Kwok IWH , Ng LG , Ginhoux F , Newell EW (2018) **Dimensionality reduction for visualizing single-cell data using UMAP** *Nat Biotechnol*
<https://doi.org/10.1038/nbt.4314>
- Bendall SC , Davis KL , David Amir E , Tadmor MD , Simonds EF , Chen TJ , Shenfeld DK , Nolan GP , Pe'er D (2014) **Single-cell trajectory detection uncovers progression and regulatory coordination in human B cell development** *Cell* **157**:714–25
- Betancur RR , Broughton RE , Wiley EO , Carpenter K , Lopez JA , Li C , Holcroft NI , Arcila D , Sanciangco M , Cureton Ii JC , Zhang F , Buser T , Campbell MA , Ballesteros JA , Roa-Varon A , Willis S , Borden WC , Rowley T , Reneau PC , Hough DJ , Lu G , Grande T , Arratia G , Orti G (2013) **The tree of life and a new classification of bony fishes** *PLoS Curr* **5**
<https://doi.org/10.1371/currents.tol.53ba26640df0ccaee75bb165c8c26288>

Braasch I, Gehrke AR, Smith JJ, Kawasaki K, Manousaki T, Pasquier J, Amores A, Desvignes T, Batzel P, Catchen J, Berlin AM, Campbell MS, Barrell D, Martin KJ, Mulley JF, Ravi V, Lee AP, Nakamura T, Chalopin D, Fan S, Wcisel D, Canestro C, Sydes J, Beaudry FEG, Sun Y, Hertel J, Beam MJ, Fasold M, Ishiyama M, Johnson J, Kehr S, Lara M, Letaw JH, Litman GW, Litman RT, Mikami M, Ota T, Saha NR, Williams L, Stadler PF, Wang H, Taylor JS, Fontenot Q, Ferrara A, Searle SMJ, Aken B, Yandell M, Schneider I, Yoder JA, Volff J-N, Meyer A, Amemiya CT, Venkatesh B, Holland PWH, Guiguen Y, Bobe J, Shubin NH, Di Palma F, Alföldi J, Lindblad-Toh K, Postlethwait JH (2016) **The spotted gar genome illuminates vertebrate evolution and facilitates human-teleost comparisons** *Nat Genet* **48**:427–37
<https://doi.org/10.1038/ng.3526>

Brent GA (2012) **Mechanisms of thyroid hormone action** *The Journal of Clinical Investigation* **122**:3035–43
<https://doi.org/10.1172/JCI60047>

Brock CK, Wallin ST, Ruiz OE, Samms KM, Mandal A, Sumner EA, Eisenhoffer GT (2019) **Stem cell proliferation is induced by apoptotic bodies from dying cells during epithelial tissue maintenance** *Nature Communications* **10**
<https://doi.org/10.1038/s41467-019-09010-6>

Budi EH, Patterson LB, Parichy DM (2011) **Post-embryonic nerve-associated precursors to adult pigment cells: genetic requirements and dynamics of morphogenesis and differentiation** *PLoS Genet* **7**
<https://doi.org/10.1371/journal.pgen.1002044>

Cal L, Suarez-Bregua P, Comesana P, Owen J, Braasch I, Kelsh R, Cerda-Reverter JM, Rotllant J (2019) **Countershading in zebrafish results from an Asip1 controlled dorsoventral gradient of pigment cell differentiation** *Scientific reports* **9**
<https://doi.org/10.1038/s41598-019-40251-z>

Cao J, Packer JS, Ramani V, Cusanovich DA, Huynh C, Daza R, Qiu X, Lee C, Furlan SN, Steemers FJ, Adey A, Waterston RH, Trapnell C, Shendure J (2017) **Comprehensive single-cell transcriptional profiling of a multicellular organism** *Science* **357**:661–7
<https://doi.org/10.1126/science.aam8940>

Cao J, Spielmann M, Qiu X, Huang X, Ibrahim DM, Hill AJ, Zhang F, Mundlos S, Christiansen L, Steemers FJ, Trapnell C, Shendure J (2019) **The single-cell transcriptional landscape of mammalian organogenesis** *Nature* **566**:496–502
<https://doi.org/10.1038/s41586-019-0969-x>

Chen CF, Foley J, Tang PC, Li A, Jiang TX, Wu P, Widelitz RB, Chuong CM (2015) **Development, regeneration, and evolution of feathers** *Annu Rev Anim Biosci* **3**:169–95
<https://doi.org/10.1146/annurev-animal-022513-114127>

Chen CH, Puliafito A, Cox BD, Primo L, Fang Y, Di Talia S, Poss KD. (2016) **Multicolor Cell Barcoding Technology for Long-Term Surveillance of Epithelial Regeneration in Zebrafish** *Dev Cell* **36**:668–80
<https://doi.org/10.1016/j.devcel.2016.02.017>

Chu Q , Gao Y , Gao X , Dong Z , Song W , Xu Z , Xiang L , Wang Y , Zhang L , Li M , Gao Y (2018) **Ablation of Runx2 in Ameloblasts Suppresses Enamel Maturation in Tooth Development** *Scientific Reports* **8**
<https://doi.org/10.1038/s41598-018-27873-5>

Cluzeau C , Hadj-Rabia S , Jambou M , Mansour S , Guigue P , Masmoudi S , Bal E , Chassaing N , Vincent MC , Viot G , Clauss F , Maniere MC , Toupenay S , Le Merrer M , Lyonnet S , Cormier-Daire V , Amiel J , Faivre L , de Prost Y , Munnich A , Bonnefont JP , Bodemer C , Smahi A (2011) **Only four genes (EDA1, EDAR, EDARADD, and WNT10A) account for 90% of hypohidrotic/anhidrotic ectodermal dysplasia cases** *Hum Mutat* **32**:70–2
<https://doi.org/10.1002/humu.21384>

Contreras-Jurado C , Lorz C , García-Serrano L , Paramio JM , Aranda A (2015) **Thyroid hormone signaling controls hair follicle stem cell function** *Mol Biol Cell* **26**:1263–72
<https://doi.org/10.1091/mbc.E14-07-1251>

Cox BD , De Simone A , Tornini VA , Singh SP , Di Talia S , Poss KD. (2018) **In Toto Imaging of Dynamic Osteoblast Behaviors in Regenerating Skeletal Bone** *Curr Biol* **28**:3937–47
<https://doi.org/10.1016/j.cub.2018.10.052>

Cui CY , Schlessinger D (2006) **EDA signaling and skin appendage development** *Cell Cycle* **5**:2477–83
<https://doi.org/10.4161/cc.5.21.3403>

Daane JM , Rohner N , Konstantinidis P , Djuranovic S , Harris MP (2015) **Parallelism and Epistasis in Skeletal Evolution Identified through Use of Phylogenomic Mapping Strategies** *Molecular Biology and Evolution*
<https://doi.org/10.1093/molbev/msv208>

De Simone A , Evanitsky MN , Hayden L , Cox BD , Wang J , Tornini VA , Ou J , Chao A , Poss KD , Di Talia S. (2021) **Control of osteoblast regeneration by a train of Erk activity waves** *Nature* **590**:129–33
<https://doi.org/10.1038/s41586-020-03085-8>

DeLaurier A , Eames BF , Blanco-Sanchez B , Peng G , He X , Swartz ME , Ullmann B , Westerfield M , Kimmel CB (2010) **Zebrafish sp7:EGFP: a transgenic for studying otic vesicle formation, skeletogenesis, and bone regeneration** *Genesis* **48**:505–11
<https://doi.org/10.1002/dvg.20639>

Di-Poi N , Milinkovitch MC (2016) **The anatomical placode in reptile scale morphogenesis indicates shared ancestry among skin appendages in amniotes** *Science Advances* **2**
<https://doi.org/10.1126/sciadv.1600708>

Dobin A , Davis CA , Schlesinger F , Drenkow J , Zaleski C , Jha S , Batut P , Chaisson M , Gingeras TR (2013) **STAR: ultrafast universal RNA-seq aligner** *Bioinformatics (Oxford, England)* **29**:15–21
<https://doi.org/10.1093/bioinformatics/bts635>

Dooley CM , Mongera A , Walderich B , Nusslein-Volhard C (2013) **On the embryonic origin of adult melanophores: the role of ErbB and Kit signalling in establishing melanophore stem cells in zebrafish** *Development* **140**:1003–13
<https://doi.org/10.1242/dev.087007>

Duverger O , Morasso MI (2009) **Epidermal patterning and induction of different hair types during mouse embryonic development** *Birth Defects Res C Embryo Today* **87**:263–72
<https://doi.org/10.1002/bdrc.20158>

Ebnet K (2017) **Junctional Adhesion Molecules (JAMs): Cell Adhesion Receptors With Pleiotropic Functions in Cell Physiology and Development** *Physiol Rev* **97**:1529–54
<https://doi.org/10.1152/physrev.00004.2017>

Endo C , Johnson TA , Morino R , Nakazono K , Kamitsuji S , Akita M , Kawajiri M , Yamasaki T , Kami A , Hoshi Y , Tada A , Ishikawa K , Hine M , Kobayashi M , Kurume N , Tsunemi Y , Kamatani N , Kawashima M (2018) **Genome-wide association study in Japanese females identifies fifteen novel skin-related trait associations** *Scientific reports* **8**
<https://doi.org/10.1038/s41598-018-27145-2>

Eom DS , Patterson LB , Bostic RR , Parichy DM (2021) **Immunoglobulin superfamily receptor Junctional adhesion molecule 3 (Jam3) requirement for melanophore survival and patterning during formation of zebrafish stripes** *bioRxiv:2021.03.01.433381*
<https://doi.org/10.1101/2021.03.01.433381>

Eskova A , Chauvigné F , Maischein H-M , Ammelburg M , Cerdà J , Nüsslein-Volhard C , Irion U (2017) **Gain-of-function mutations in Aqp3a influence zebrafish pigment pattern formation through the tissue environment** *Development (Cambridge, England)* **144**:2059–69
<https://doi.org/10.1242/dev.143495>

Feitosa NM , Richardson R , Bloch W , Hammerschmidt M (2011) **Basement membrane diseases in zebrafish** *Methods Cell Biol* **105**:191–222
<https://doi.org/10.1016/b978-0-12-381320-6.00008-4>

Fraser GJ , Cerny R , Soukup V , Bronner-Fraser M , Streelman JT (2010) **The Odontode Explosion: The origin of tooth-like structures in vertebrates** *BioEssays: news and reviews in molecular, cellular and developmental biology* **32**:808–17
<https://doi.org/10.1002/bies.200900151>

González R , Moffatt G , Hagner A , Sinha S , Shin W , Rahmani W , Chojnacki A , Biernaskie J (2017) **Platelet-derived growth factor signaling modulates adult hair follicle dermal stem cell maintenance and self-renewal** *NPJ Regen Med* **2**
<https://doi.org/10.1038/s41536-017-0013-4>

Gu J , Lu Y , Li F , Qiao L , Wang Q , Li N , Borgia JA , Deng Y , Lei G , Zheng Q (2014) **Identification and characterization of the novel Col10a1 regulatory mechanism during chondrocyte hypertrophic differentiation** *Cell Death Dis* **5**
<https://doi.org/10.1038/cddis.2014.444>

Gur D , Bain EJ , Johnson KR , Aman AJ , Pasoili HA , Flynn JD , Allen MC , Deheyn DD , Lee JC , Lippincott-Schwartz J , Parichy DM (2020) **In situ differentiation of iridophore crystalloids underlies zebrafish stripe patterning** *Nat Commun* **11**
<https://doi.org/10.1038/s41467-020-20088-1>

Guzman A , Ramos-Balderas JL , Carrillo-Rosas S , Maldonado E (2013) **A stem cell proliferation burst forms new layers of P63 expressing suprabasal cells during zebrafish postembryonic epidermal development** *Biol Open* **2**:1179–86
<https://doi.org/10.1242/bio.20136023>

Haghverdi L , Lun ATL , Morgan MD , Marioni JC (2018) **Batch effects in single-cell RNA-sequencing data are corrected by matching mutual nearest neighbors** *Nat Biotechnol* **36**:421–7
<https://doi.org/10.1038/nbt.4091>

Harris MP , Rohner N , Schwarz H , Perathoner S , Konstantinidis P , Nüsslein-Volhard C (2008) **Zebrafish eda and edar Mutants Reveal Conserved and Ancestral Roles of Ectodysplasin Signaling in Vertebrates** *PLoS Genet* **4**
<https://doi.org/10.1371/journal.pgen.1000206>

Headon DJ , Overbeek PA (1999) **Involvement of a novel Tnf receptor homologue in hair follicle induction** *Nat Genet* **22**:370–4
<https://doi.org/10.1038/11943>

Hessle L , Stordalen GA , Wenglén C , Petzold C , Tanner EK , Brorson S-H , Baekkevold ES , Önnérjörð P , Reinholt FP , Heinegård D (2013) **The Skeletal Phenotype of Chondroadherin Deficient Mice** *PLOS ONE* **8**
<https://doi.org/10.1371/journal.pone.0063080>

Hirata M , Nakamura K-I , Kondo S (2005) **Pigment cell distributions in different tissues of the zebrafish, with special reference to the striped pigment pattern** *Developmental Dynamics* **234**:293–300
<https://doi.org/10.1002/dvdy.20513>

Houghton L , Lindon C , Morgan BA (2005) **The ectodysplasin pathway in feather tract development** *Development* **132**:863–72
<https://doi.org/10.1242/dev.01651>

Huh SH , Narhi K , Lindfors PH , Haara O , Yang L , Ornitz DM , Mikkola ML (2013) **Fgf20 governs formation of primary and secondary dermal condensations in developing hair follicles** *Genes Dev* **27**:450–8
<https://doi.org/10.1101/gad.198945.112>

Hultman KA , Bahary N , Zon LI , Johnson SL (2007) **Gene Duplication of the zebrafish kit ligand and partitioning of melanocyte development functions to kit ligand a** *PLoS Genet* **3**
<https://doi.org/10.1371/journal.pgen.0030017>

Hwang WY , Fu Y , Reyon D , Maeder ML , Tsai SQ , Sander JD , Peterson RT , Yeh JR , Joung JK (2013) **Efficient genome editing in zebrafish using a CRISPR-Cas system** *Nat Biotechnol* **31**:227–9
<https://doi.org/10.1038/nbt.2501>

Iwasaki M , Kuroda J , Kawakami K , Wada H (2018) **Epidermal regulation of bone morphogenesis through the development and regeneration of osteoblasts in the zebrafish scale** *Dev Biol*
<https://doi.org/10.1016/j.ydbio.2018.03.005>

Janvier P (2015) **Facts and fancies about early fossil chordates and vertebrates** *Nature* **520**:483–9
<https://doi.org/10.1038/nature14437>

Karlsson L , Bondjers C , Betsholtz C (1999) **Roles for PDGF-A and sonic hedgehog in development of mesenchymal components of the hair follicle** *Development* **126**:2611–21
<https://doi.org/10.1242/dev.126.12.2611>

Kawasaki K (2009) **The SCPP gene repertoire in bony vertebrates and graded differences in mineralized tissues** *Development Genes and Evolution* **219**:147–57
<https://doi.org/10.1007/s00427-009-0276-x>

Kawasaki K , Keating JN , Nakatomi M , Welten M , Mikami M , Sasagawa I , Puttick MN , Donoghue PCJ , Ishiyama M (2021) **Coevolution of enamel, ganoin, enameloid, and their matrix SCPP genes in osteichthyans** *iScience [Internet]* **24**
<https://doi.org/10.1016/j.isci.2020.102023>

Kawasaki K , Keating JN , Nakatomi M , Welten M , Mikami M , Sasagawa I , Puttick MN , Donoghue PCJ , Ishiyama M (2021) **Coevolution of enamel, ganoin, enameloid, and their matrix SCPP genes in osteichthyans** *iScience* **24**
<https://doi.org/10.1016/j.isci.2020.102023>

Kawasaki K , Mikami M , Nakatomi M , Braasch I , Batzel P , Postlethwait J H. , Sato A , Sasagawa I , Ishiyama M (2017) **SCPP Genes and Their Relatives in Gar: Rapid Expansion of Mineralization Genes in Osteichthyans** *Journal of Experimental Zoology Part B: Molecular and Developmental Evolution* **328**:645–65
<https://doi.org/10.1002/jez.b.22755>

Kere J , Srivastava AK , Montonen O , Zonana J , Thomas N , Ferguson B , Munoz F , Morgan D , Clarke A , Baybayan P , Chen EY , Ezer S , Saarialho-Kere U , de la Chapelle A , Schlessinger D. (1996) **X-linked anhidrotic (hypohidrotic) ectodermal dysplasia is caused by mutation in a novel transmembrane protein** *Nat Genet* **13**:409–16
<https://doi.org/10.1038/ng0895-409>

Komori T (2010) **Regulation of osteoblast differentiation by Runx2** *Adv Exp Med Biol* **658**:43–9
https://doi.org/10.1007/978-1-4419-1050-9_5

Kondo S , Kuwahara Y , Kondo M , Naruse K , Mitani H , Wakamatsu Y , Ozato K , Asakawa S , Shimizu N , Shima A (2001) **The medaka rs-3 locus required for scale development encodes ectodysplasin-A receptor** *Current Biology* **11**:1202–6
[https://doi.org/10.1016/S0960-9822\(01\)00324-4](https://doi.org/10.1016/S0960-9822(01)00324-4)

Krauss J , Frohnhöfer HG , Walderich B , Maischein H-M , Weiler C , Irion U , Nüsslein-Volhard C (2014) **Endothelin signalling in iridophore development and stripe pattern formation of zebrafish** *Biology Open* **3**:503–9
<https://doi.org/10.1242/bio.20148441>

Lang MR , Patterson LB , Gordon TN , Johnson SL , Parichy DM (2009) **Basonuclin-2 requirements for zebrafish adult pigment pattern development and female fertility** *PLoS Genet* **5**
<https://doi.org/10.1371/journal.pgen.1000744>

Larsson T , Sommarin Y , Paulsson M , Antonsson P , Hedbom E , Wendel M , Heinegård D (1991) **Cartilage matrix proteins. A basic 36-kDa protein with a restricted distribution to cartilage and bone** *J Biol Chem* **266**:20428–33

Le Guellec D , Morvan-Dubois G , Sire JY. (2004) **Skin development in bony fish with particular emphasis on collagen deposition in the dermis of the zebrafish (Danio rerio)** *Int J Dev Biol* **48**:217–31
<https://doi.org/10.1387/ijdb.031768dg>

Lee H , Kimelman D (2002) **A dominant-negative form of p63 is required for epidermal proliferation in zebrafish** *Dev Cell* **2**:607–16

Lee RT , Asharani PV , Carney TJ (2014) **Basal keratinocytes contribute to all strata of the adult zebrafish epidermis** *PLoS One* **9**
<https://doi.org/10.1371/journal.pone.0084858>

Lefebvre S , Mikkola ML (2014) **Ectodysplasin research--where to next?** *Semin Immunol* **26**:220–8
<https://doi.org/10.1016/j.smim.2014.05.002>

Lewis VM , Saunders LM , Larson TA , Bain EJ , Sturiale SL , Gur D , Chowdhury S , Flynn JD , Allen MC , Deheyn DD , Lee JC , Simon JA , Lippincott-Schwartz J , Raible DW , Parichy DM (2019) **Fate plasticity and reprogramming in genetically distinct populations of Danio leucophores** *Proceedings of the National Academy of Sciences of the United States of America* **116**:11806–11
<https://doi.org/10.1073/pnas.1901021116>

Li Q , Frank M , Thisse CI , Thisse BV , Uitto J. (2011) **Zebrafish: a model system to study heritable skin diseases** *J Invest Dermatol* **131**:565–71
<https://doi.org/10.1038/jid.2010.388>

Li S , Kong H , Yao N , Yu Q , Wang P , Lin Y , Wang J , Kuang R , Zhao X , Xu J , Zhu Q , Ni L. (2011) **The role of runt-related transcription factor 2 (Runx2) in the late stage of odontoblast differentiation and dentin formation** *Biochem Biophys Res Commun* **410**:698–704

<https://doi.org/10.1016/j.bbrc.2011.06.065>

Lü AJ , Hu XC , Wang Y , Zhu AH , Shen LL , Tian J , Feng ZZ , Feng ZJ (2015) **Skin immune response in the zebrafish, *Danio rerio* (Hamilton), to *Aeromonas hydrophila* infection: a transcriptional profiling approach** *J Fish Dis* **38**:137–50

<https://doi.org/10.1111/jfd.12214>

Mancino G , Miro C , Di Cicco E , Dentice M. (2021) **Thyroid hormone action in epidermal development and homeostasis and its implications in the pathophysiology of the skin** *Journal of Endocrinological Investigation*

<https://doi.org/10.1007/s40618-020-01492-2>

Märss T (2006) **Exoskeletal ultrasculpture of early vertebrates** *Journal of Vertebrate Paleontology* **26**:235–52

[https://doi.org/10.1671/0272-4634\(2006\)26](https://doi.org/10.1671/0272-4634(2006)26)

Martens M , Ammar A , Riutta A , Waagmeester A , Slenter DN , Hanspers K R AM , Digles D , Lopes EN , Ehrhart F , Dupuis LJ , Winckers LA , Coort SL , Willighagen EL , Evelo CT , Pico AR , Kutmon M (2021) **WikiPathways: connecting communities** *Nucleic Acids Res* **49**:D613–d21

<https://doi.org/10.1093/nar/gkaa1024>

McMenamin SK , Bain EJ , McCann AE , Patterson LB , Eom DS , Waller ZP , Hamill JC , Kuhlman JA , Eisen JS , Parichy DM (2014) **Thyroid hormone-dependent adult pigment cell lineage and pattern in zebrafish** *Science* **345**:1358–61

<https://doi.org/10.1126/science.1256251>

Metz JR , de Vrieze E , Lock E-J , Schulten IE , Flik G. (2012) **Elasmoid scales of fishes as model in biomedical bone research** *Journal of Applied Ichthyology* **28**:382–7

<https://doi.org/10.1111/j.1439-0426.2012.01990.x>

Montonen O , Ezer S , Saarialho-Kere UK , Herva R , Karjalainen-Lindsberg M-L , Kaitila I , Schlessinger D , Srivastava AK , Thesleff I , Kere J (1998) **The Gene Defective in Anhidrotic Ectodermal Dysplasia Is Expressed in the Developing Epithelium, Neuroectoderm, Thymus, and Bone** *Journal of Histochemistry & Cytochemistry* **46**:281–9

<https://doi.org/10.1177/002215549804600301>

Morris JL , Cross SJ , Lu Y , Kadler KE , Lu Y , Dallas SL , Martin P (2018) **Live imaging of collagen deposition during skin development and repair in a collagen I - GFP fusion transgenic zebrafish line** *Developmental biology* **441**:4–11

<https://doi.org/10.1016/j.ydbio.2018.06.001>

Murcia S , Lavoie E , Linley T , Devaraj A , Ossa EA , Arola D (2016) **The natural armors of fish: A comparison of the lamination pattern and structure of scales** *J Mech Behav Biomed Mater*

<https://doi.org/10.1016/j.jmbbm.2016.09.025>

Near TJ , Eytan RI , Dornburg A , Kuhn KL , Moore JA , Davis MP , Wainwright PC , Friedman M , Smith WL (2012) **Resolution of ray-finned fish phylogeny and timing of diversification** *Proceedings of the National Academy of Sciences* **109**:13698–703
<https://doi.org/10.1073/pnas.1206625109>

Oralová V , Rosa JT , Larionova D , Witten PE , Huysseune A (2020) **Multiple epithelia are required to develop teeth deep inside the pharynx** *Proc Natl Acad Sci U S A* **117**:11503–12
<https://doi.org/10.1073/pnas.2000279117>

Owen JP , Kelsh RN , Yates CA (2020) **A quantitative modelling approach to zebrafish pigment pattern formation** *eLife* **9**
<https://doi.org/10.7554/eLife.52998>

Packer JS , Zhu Q , Huynh C , Sivaramakrishnan P , Preston E , Dueck H , Stefanik D , Tan K , Trapnell C , Kim J , Waterston RH , Murray JI (2019) **A lineage-resolved molecular atlas of C. elegans embryogenesis at single-cell resolution** *Science* **365**
<https://doi.org/10.1126/science.aax1971>

Parichy DM , Elizondo MR , Mills MG , Gordon TN , Engeszer RE (2009) **Normal table of postembryonic zebrafish development: Staging by externally visible anatomy of the living fish** *Developmental Dynamics* **238**:2975–3015
<https://doi.org/10.1002/dvdy.22113>

Parichy DM , Mellgren EM , Rawls JF , Lopes SS , Kelsh RN , Johnson SL (2000) **Mutational analysis of endothelin receptor b1 (rose) during neural crest and pigment pattern development in the zebrafish Danio rerio** *Dev Biol* **227**:294–306
<https://doi.org/10.1006/dbio.2000.9899>

Parichy DM , Turner JM (2003) **Temporal and cellular requirements for Fms signaling during zebrafish adult pigment pattern development** *Development* **130**:817–33

Parinov S , Kondrichin I , Korzh V , Emelyanov A (2004) **Tol2 transposon-mediated enhancer trap to identify developmentally regulated zebrafish genes in vivo** *Dev Dyn* **231**:449–59
<https://doi.org/10.1002/dvdy.20157>

Patterson LB , Bain EJ , Parichy DM (2014) **Pigment cell interactions and differential xanthophore recruitment underlying zebrafish stripe reiteration and Danio pattern evolution** *Nat Commun* **5**
<https://doi.org/10.1038/ncomms6299>

Patterson LB , Parichy DM (2013) **Interactions with iridophores and the tissue environment required for patterning melanophores and xanthophores during zebrafish adult pigment stripe formation** *PLoS Genet* **9**
<https://doi.org/10.1371/journal.pgen.1003561>

Patterson LB , Parichy DM (2019) **Zebrafish Pigment Pattern Formation: Insights into the Development and Evolution of Adult Form** *Annual Review of Genetics* **53**:505–30
<https://doi.org/10.1146/annurev-genet-112618-043741>

Peloggia J , Münch D , Meneses-Giles P , Romero-Carvajal A , Lush ME , Lawson ND , McClain M , Pan YA , Piotrowski T (2021) **Adaptive cell invasion maintains lateral line organ homeostasis in response to environmental changes** *Developmental Cell*
<https://doi.org/10.1016/j.devcel.2021.03.027>

Pfalzgraff A , Barcena-Varela S , Heinbockel L , Gutschmann T , Brandenburg K , Martinez-de-Tejada G , Weindl G (2018) **Antimicrobial endotoxin-neutralizing peptides promote keratinocyte migration via P2X7 receptor activation and accelerate wound healing in vivo** *Br J Pharmacol* **175**:3581–93
<https://doi.org/10.1111/bph.14425>

Pliner HA , Shendure J , Trapnell C (2019) **Supervised classification enables rapid annotation of cell atlases** *Nat Methods* **16**:983–6
<https://doi.org/10.1038/s41592-019-0535-3>

Qu Q , Haitina T , Zhu M , Ahlberg PE (2015) **New genomic and fossil data illuminate the origin of enamel** *Nature* **526**:108–11
<https://doi.org/10.1038/nature15259>

Quan H , Yang W , Lapeyriere M , Schaible E , Ritchie RO , Meyers MA (2020) **Structure and Mechanical Adaptability of a Modern Elasmoid Fish Scale from the Common Carp** *Matter* **3**:842–63
<https://doi.org/10.1016/j.matt.2020.05.011>

Quigley IK , Turner JM , Nuckels RJ , Manuel JL , Budi EH , MacDonald EL , Parichy DM (2004) **Pigment pattern evolution by differential deployment of neural crest and post-embryonic melanophore lineages in Danio fishes** *Development* **131**:6053–69
<https://doi.org/10.1242/dev.01526>

Quinlan AR , Hall IM (2010) **BEDTools: a flexible suite of utilities for comparing genomic features** *Bioinformatics* **26**:841–2
<https://doi.org/10.1093/bioinformatics/btq033>

Raouf A , Ganss B , McMahon C , Vary C , Roughley PJ , Seth A (2002) **Lumican is a major proteoglycan component of the bone matrix** *Matrix Biol* **21**:361–7
[https://doi.org/10.1016/s0945-053x\(02\)00027-6](https://doi.org/10.1016/s0945-053x(02)00027-6)

Rasmussen JP , Sack GS , Martin SM , Sagasti A (2015) **Vertebrate epidermal cells are broad-specificity phagocytes that clear sensory axon debris** *J Neurosci* **35**:559–70
<https://doi.org/10.1523/jneurosci.3613-14.2015>

Rasmussen JP , Vo NT , Sagasti A (2018) **Fish Scales Dictate the Pattern of Adult Skin Innervation and Vascularization** *Dev Cell* **46**:344–59
<https://doi.org/10.1016/j.devcel.2018.06.019>

Richardson R , Metzger M , Knyphausen P , Ramezani T , Slanchev K , Kraus C , Schmelzer E , Hammerschmidt M (2016) **Re-epithelialization of cutaneous wounds in adult zebrafish combines mechanisms of wound closure in embryonic and adult mammals** *Development* **143**:2077–88

<https://doi.org/10.1242/dev.130492>

Rosset EM , Bradshaw AD (2016) **SPARC/osteonectin in mineralized tissue** *Matrix Biol* **52**:78–87

<https://doi.org/10.1016/j.matbio.2016.02.001>

Safer JD (2011) **Thyroid hormone action on skin** *Dermatoendocrinol* **3**:211–5

<https://doi.org/10.4161/derm.3.3.17027>

Safer JD , Fraser LM , Ray S , Holick MF (2001) **Topical triiodothyronine stimulates epidermal proliferation, dermal thickening, and hair growth in mice and rats** *Thyroid* **11**:717–24

<https://doi.org/10.1089/10507250152484547>

Saunders LM , Mishra AK , Aman AJ , Lewis VM , Toomey MB , Packer JS , Qiu X , McFaline-Figueroa JL , Corbo JC , Trapnell C , Parichy DM (2019) **Thyroid hormone regulates distinct paths to maturation in pigment cell lineages** *eLife* **8**

<https://doi.org/10.7554/eLife.45181>

Schindelin J , Arganda-Carreras I , Frise E , Kaynig V , Longair M , Pietzsch T , Preibisch S , Rueden C , Saalfeld S , Schmid B , Tinevez J-Y , White DJ , Hartenstein V , Eliceiri K , Tomancak P , Cardona A (2012) **Fiji: an open-source platform for biological-image analysis** *Nature Methods* **9**:676–82

<https://doi.org/10.1038/nmeth.2019>

Sire J-Y , Allizard F , Babiar O , Bourguignon J , Quilhac A (1997) **Scale development in zebrafish (*Danio rerio*)** *Journal of Anatomy* **190**:545–61

<https://doi.org/10.1046/j.1469-7580.1997.19040545.x>

Sire J-Y , Donoghue PCJ , Vickaryous MK (2009) **Origin and evolution of the integumentary skeleton in non-tetrapod vertebrates** *Journal of Anatomy* **214**:409–40

<https://doi.org/10.1111/j.1469-7580.2009.01046.x>

Sire JY (1990) **From Ganoid to Elasmoid Scales in the Acinopterygian Fishes** *Netherlands Journal of Zoology* **40**:75–92

<https://doi.org/10.1163/156854289X00192>

Sire JY , Akimenko MA (2004) **Scale development in fish: a review, with description of sonic hedgehog (shh) expression in the zebrafish (*Danio rerio*)** *Int J Dev Biol* **48**:233–47

Sire JY , Huysseune A (2003) **Formation of dermal skeletal and dental tissues in fish: a comparative and evolutionary approach** *Biol Rev Camb Philos Soc* **78**:219–49

Sire JY , Quilhac A , Bourguignon J , Allizard F (1997) **Evidence for participation of the epidermis in the deposition of superficial layer of scales in zebrafish (*Danio rerio*): A SEM and TEM study** *J Morphol* **231**:161–74
[https://doi.org/10.1002/\(sici\)1097-4687\(199702\)231:2<161::aid-jmor5>3.0.co;2-h](https://doi.org/10.1002/(sici)1097-4687(199702)231:2<161::aid-jmor5>3.0.co;2-h)

Smith CEL , Poulter JA , Antanaviciute A , Kirkham J , Brookes SJ , Inglehearn CF , Mighell AJ (2017) **Amelogenesis Imperfecta; Genes, Proteins, and Pathways** *Front Physiol* **8**
<https://doi.org/10.3389/fphys.2017.00435>

Smith MP , Donoghue PCJ , Sansom IJ (2002) **The spatial and temporal diversification of Early Palaeozoic vertebrates** *Geological Society, London, Special Publications* **194**:69–83
<https://doi.org/10.1144/gsl.sp.2002.194.01.06>

Spiewak JE , Bain EJ , Liu J , Kou K , Sturiale SL , Patterson LB , Diba P , Eisen JS , Braasch I , Ganz J , Parichy DM (2018) **Evolution of Endothelin signaling and diversification of adult pigment pattern in Danio fishes** *PLOS Genetics* **14**
<https://doi.org/10.1371/journal.pgen.1007538>

Srivastava AK , Pispas J , Hartung AJ , Du Y , Ezer S , Jenks T , Shimada T , Pekkanen M , Mikkola ML , Ko MS , Thesleff I , Kere J , Schlessinger D. (1997) **The Tabby phenotype is caused by mutation in a mouse homologue of the EDA gene that reveals novel mouse and human exons and encodes a protein (ectodysplasin-A) with collagenous domains** *Proc Natl Acad Sci U S A* **94**:13069–74

Srivatsan SR , McFaline-Figueroa JL , Ramani V , Saunders L , Cao J , Packer J , Pliner HA , Jackson DL , Daza RM , Christiansen L , Zhang F , Steemers F , Shendure J , Trapnell C (2020) **Massively multiplex chemical transcriptomics at single-cell resolution** *Science* **367**:45–51
<https://doi.org/10.1126/science.aax6234>

Thomas G , Moffatt P , Salois P , Gaumond MH , Gingras R , Godin E , Miao D , Goltzman D , Lanctôt C (2003) **Osteocrin, a novel bone-specific secreted protein that modulates the osteoblast phenotype** *J Biol Chem* **278**:50563–71
<https://doi.org/10.1074/jbc.M307310200>

Tomita Y , Akiyama M , Shimizu H (2006) **PDGF isoforms induce and maintain anagen phase of murine hair follicles** *J Dermatol Sci* **43**:105–15
<https://doi.org/10.1016/j.jdermsci.2006.03.012>

Traag VA , Waltman L , van Eck NJ. (2019) **From Louvain to Leiden: guaranteeing well-connected communities** *Scientific reports* **9**
<https://doi.org/10.1038/s41598-019-41695-z>

Trapnell C , Cacchiarelli D , Grimsby J , Pokharel P , Li S , Morse M , Lennon NJ , Livak KJ , Mikkelsen TS , Rinn JL (2014) **The dynamics and regulators of cell fate decisions are revealed by pseudotemporal ordering of single cells** *Nat Biotechnol* **32**:381–6
<https://doi.org/10.1038/nbt.2859>

Turner S , Burrow CJ , Schultze H-P , Blicek A , Reif W-E , Rexroad CB , Bultynck P , Nowlan GS (2010) **False teeth: conodont-vertebrate phylogenetic relationships revisited** *Geodiversitas* **32**:545–94
<https://doi.org/10.5252/g2010n4a1>

Urzúa B , Ortega-Pinto A , Morales-Bozo I , Rojas-Alcayaga G , Cifuentes V (2011) **Defining a new candidate gene for amelogenesis imperfecta: from molecular genetics to biochemistry** *Biochem Genet* **49**:104–21
<https://doi.org/10.1007/s10528-010-9392-6>

Venkatesh B , Lee AP , Ravi V , Maurya AK , Lian MM , Swann JB , Ohta Y , Flajnik MF , Sutoh Y , Kasahara M , Hoon S , Gangu V , Roy SW , Irimia M , Korzh V , Kondrychyn I , Lim ZW , Tay BH , Tohari S , Kong KW , Ho S , Lorente-Galdos B , Quilez J , Marques-Bonet T , Raney BJ , Ingham PW , Tay A , Hillier LW , Minx P , Boehm T , Wilson RK , Brenner S , Warren WC (2014) **Elephant shark genome provides unique insights into gnathostome evolution** *Nature* **505**:174–9
<https://doi.org/10.1038/nature12826>

Visser M , Palstra R-J , Kayser M (2014) **Human skin color is influenced by an intergenic DNA polymorphism regulating transcription of the nearby BNC2 pigmentation gene** *Human Molecular Genetics* **23**:5750–62
<https://doi.org/10.1093/hmg/ddu289>

Volkening A , Sandstede B (2015) **Modelling stripe formation in zebrafish: an agent-based approach** *Journal of The Royal Society Interface* **12**
<https://doi.org/10.1098/rsif.2015.0812>

Volkening A , Sandstede B (2018) **Iridophores as a source of robustness in zebrafish stripes and variability in Danio patterns** *Nat Commun* **9**
<https://doi.org/10.1038/s41467-018-05629-z>

Watanabe M , Kondo S (2015) **Is pigment patterning in fish skin determined by the Turing mechanism?** *Trends Genet* **31**:88–96
<https://doi.org/10.1016/j.tig.2014.11.005>

Wolock SL , Lopez R , Klein AM (2019) **Scrublet: Computational Identification of Cell Doublets in Single-Cell Transcriptomic Data** *Cell systems* **8**:281–91
<https://doi.org/10.1016/j.cels.2018.11.005>

Wurster S , Ruiz OE , Samms KM , Tatara AM , Albert ND , Kahan PH , Nguyen AT , Mikos AG , Kontoyiannis DP , Eisenhoffer GT (2021) **EGF-mediated suppression of cell extrusion during mucosal damage attenuates opportunistic fungal invasion** *Cell Reports* **34**
<https://doi.org/10.1016/j.celrep.2021.108896>

Yang L , Tsang KY , Tang HC , Chan D , Cheah KS (2014) **Hypertrophic chondrocytes can become osteoblasts and osteocytes in endochondral bone formation** *Proc Natl Acad Sci U S A* **111**:12097–102
<https://doi.org/10.1073/pnas.1302703111>

Zhang C (2012) **Molecular mechanisms of osteoblast-specific transcription factor Osterix effect on bone formation** *Beijing Da Xue Xue Bao Yi Xue Ban* **44**:659–65

Author information

Andrew J. Aman

Department of Biology, University of Virginia, Charlottesville, VA

Lauren M. Saunders

Department of Genome Sciences, University of Washington, Seattle, WA

ORCID iD: [0000-0003-4377-4252](https://orcid.org/0000-0003-4377-4252)

August A. Carr

Department of Biology, University of Virginia, Charlottesville, VA

Sanjay R. Srivatsan

Department of Genome Sciences, University of Washington, Seattle, WA

Colten D. Eberhard

National Human Genome Research Institute, National Institutes of Health, Bethesda, MD

Blake Carrington

National Human Genome Research Institute, National Institutes of Health, Bethesda, MD

Dawn Watkins-Chow

National Human Genome Research Institute, National Institutes of Health, Bethesda, MD

William J. Pavan

National Human Genome Research Institute, National Institutes of Health, Bethesda, MD

Cole Trapnell

Department of Genome Sciences, University of Washington, Seattle, WA

David M. Parichy

Department of Biology, University of Virginia, Charlottesville, VA, Department of Cell Biology, University of Virginia, Charlottesville, VA

Editors

Reviewing Editor

Alvaro Sagasti

University of California, Los Angeles, United States of America

Senior Editor

Didier Stainier

Max Planck Institute for Heart and Lung Research, Germany

Reviewer #1 (Public Review):

In their study, Aman et al. utilized single cell transcriptome analysis to investigate wild-type and mutant zebrafish skin tissues during the post-embryonic growth period. They identified new epidermal cell types, such as ameloblasts, and shed light on the effects of TH on skin morphogenesis. Additionally, they revealed the important role of the hypodermis in supporting pigment cells and adult stripe formation. Overall, I find their figures to be of high quality, their analyses to be appropriate and compelling, and their major claims to be well-supported by additional experiments. Therefore, this study will be an important contribution to the field of vertebrate skin research.

Reviewer #2 (Public Review):

This work describes transcriptome profiling of dissected skin of zebrafish at post-embryonic stages, at a time when adult structures and patterns are forming. The authors have used the state-of-the-art combinatorial indexing RNA-seq approach to generate single cell (nucleus) resolution. The data appears robust and is coherent across the four different genotypes used by the authors.

The authors present the data in a logical and accessible manner, with appropriate reference to the anatomy. They include helpful images of the biology and schematics to illustrate their interpretations.

The datasets are then interrogated to define cell and signalling relationships between skin compartments in six diverse contexts. The hypotheses generated from the datasets are then tested experimentally. Overall, the experiments are appropriate and rigorously performed. They ask very interesting questions of interactions in the skin and identify novel and specific mechanisms. They validate these well.

The authors use their datasets to define lineage relationships in the dermal scales and also in the epidermis. They show that circumferential pre-scale forming cells are precursors of focal scale forming cells while there appeared a more discontinuous relationship between lineages in the epidermis.

The authors present transcriptome evidence for enamel deposition function in epidermal subdomains. This is convincingly confirmed with an ameloblastin in situ. They further demonstrate distinct expression of SSCP and collagen genes in the SFC regions.

The authors then demonstrate that Eda and TH signalling to the basal epidermal cells generates FGF and PDGF ligands to signal to surrounding mesenchyme, regulating SFC differentiation and dermal stratification respectively.

Finally, they exploit RNA-seq data performed in parallel in the *bnc2* mutants to identify the hypodermal cells as critical regulators of pigment patterning and define the signalling systems used.

Whilst these six interactions in the skin are disparate, the stories are unified by use of the sci-RNA-seq data to define interactions. Overall, it's an assembly of work which identifies novel and interesting cell interactions and cross-talk mechanisms.

The paper provides robust evidence of cell interrelationships in the skin undergoing morphogenesis and will be a welcome dataset for the field.