Title
Tissue-Specific Transcriptomes Reveal Gene Expression Trajectories in Two Maturing Skin Epithelial Layers in Zebrafish Embryos.

Permalink
https://escholarship.org/uc/item/5sx6n666

Journal
G3 (Bethesda, Md.), 9(10)

ISSN
2160-1836

Authors
Cokus, Shawn J
De La Torre, Maricruz
Medina, Eric F
et al.

Publication Date
2019-10-07

DOI
10.1534/g3.119.400402

Peer reviewed
Tissue-Specific Transcriptomes Reveal Gene Expression Trajectories in Two Maturing Skin Epithelial Layers in Zebrafish Embryos

Shawn J. Cokus,* Maricruz De La Torre,† Eric F. Medina,† Jeffrey P. Rasmussen,*1 Joselyn Ramirez-Gutierrez,† Alvaro Sagasti,* and Fang Wang†2

*Department of Molecular, Cell, and Developmental Biology, University of California Los Angeles and †Department of Biology, California State University, Dominguez Hills

ORCID ID: 0000-0001-6997-3773 (J.P.R.)

ABSTRACT  Epithelial cells are the building blocks of many organs, including skin. The vertebrate skin initially consists of two epithelial layers, the outer periderm and inner basal cell layers, which have distinct properties, functions, and fates. The embryonic periderm ultimately disappears during development, whereas basal cells proliferate to form the mature, stratified epidermis. Although much is known about mechanisms of homeostasis in mature skin, relatively little is known about the two cell types in pre-stratification skin. To define the similarities and distinctions between periderm and basal skin epithelial cells, we purified them from zebrafish at early development stages and deeply profiled their gene expression. These analyses identified groups of genes whose tissue enrichment changed at each stage, defining gene flow dynamics of maturing vertebrate epithelia. At each of 52 and 72 hr post-fertilization (hpf), more than 60% of genes enriched in skin cells were similarly expressed in both layers, indicating that they were common epithelial genes, but many others were enriched in one layer or the other. Both expected and novel genes were enriched in periderm and basal cell layers. Genes encoding extracellular matrix, junctional, cytoskeletal, and signaling proteins were prominent among those distinguishing the two epithelial cell types. In situ hybridization and BAC transgenes confirmed our expression data and provided new tools to study zebrafish skin. Collectively, these data provide a resource for studying common and distinguishing features of maturing epithelia.

KEYWORDS  zebrafish skin development epithelia transcriptome keratin
enveloping layer surrounding gastrulating embryos (Kimmel et al. 1990), and differentiate a few hours ahead of basal cells (O’Brien et al. 2012). In mammals, periderm differentiates from surface ectoderm in a stereotyped regional progression (Wolf 1967, 1968a; Herken and Schultz-Ehrenburg 1981; M’Boneko and Merker 1988; Hardman et al. 1999; Richardson et al. 2014). Once specified, basal and periderm cells independently proliferate (Herken and Schultz-Ehrenburg 1981; Lee et al. 2014). Basal cells are stem cells that eventually give rise to all keratinocytes of the stratified adult epidermis (Smart 1970; Fuchs and Raghavan 2002; Muroyama and Lechler 2012; Guzman et al. 2013; Lee et al. 2014). Periderm in both fish and mammals is a transient tissue that ultimately sloughs off later in development (Wolf 1967, 1968b; Holbrook and Odland 1980; M’Boneko and Merker 1988; Lee et al. 2014; Richardson et al. 2014).

Periderm and basal cells illustrate the key differences distinguishing epithelial cell types. For example, basal cells secrete a basement membrane on their basal surfaces to separate the epidermis from the mesenchymal dermis (Fuchs and Raghavan 2002; Muroyama and Lechler 2012), whereas periderm cells lack a basal ECM but display a glyocalyx and associate with mucins on their apical surfaces (Wolf 1967; Pinto et al. 2019; Pasquale 2018). Periderm cells project actin-based projections on their apical surfaces, either microvilli or elongated structures called microridges (Wolf 1967, 1968a; Lam et al. 2015; Pasquale 2018). Befitting its function as the embryo’s main barrier to the external environment, periderm cells are connected to one another by tight junctions (Morita et al. 2002; O’Brien et al. 2012; Yoshida et al. 2012; Richardson et al. 2014), whereas basal cells lack tight junctions, but have hemidesmosomes that attach them to the basement membrane (Le Guellec et al. 2004; Li et al. 2011a; O’Brien et al. 2012; Muroyama and Lechler 2012). Zebrafish basal cells have a unique interaction with the touch-sensing axon endings that innervate the zebrafish skin: initially, sensory axons innervate the region between the two cell layers, but, by 54 hr post-fertilization (hpf), the apical membranes of basal cells have begun wrapping around axons to ensheathe them in structures reminiscent of sheaths formed by non-myelinating Schwann cells (O’Brien et al. 2012; Jiang et al. 2019). Since zebrafish are externally fertilized, and transgenic reporters for basal cells and periderm cells are available, zebrafish embryos are an ideal model for studies of epithelial development.

As a foundation for understanding the maturation of epithelial cells and, more specifically, the function and development of basal and periderm cells, we determined the expression profiles of each cell layer at several early developmental timepoints. To accomplish this, we purified each cell type from transgenic zebrafish with Fluorescence-Activated Cell Sorting (FACS), and deeply profiled gene expression in each isolated population with RNA-Seq. These data identify genes that may be responsible for common and distinct features of periderm and basal cells, and provide a reference for studies of epithelial cell maturation.

**MATERIALS AND METHODS**

**Zebrafish**

Zebrafish were maintained in 28.5°C and pH 7.5 fish water. Adults were kept in a 14-hour light / 10-hour dark cycle. Tg(krt4a:dsRed), Tg(krt5:GFP), Tg(krt5:Gal4), Tg(UAS:nsf8-mCherry), and Tg(ΔNp63:Gal4) lines were previously described (Curado et al. 2007; Hu et al. 2010; O’Brien et al. 2012; Rasmussen et al. 2015). Animal care and experimental procedures were approved by the CSUDH IACUC Committee and the UCLA Chancellor’s Animal Research Committee.

**Fluorescence-Activated Cell Sorting (FACS)**

Dechorionated embryos were transferred into 1.5 ml tubes and rinsed with Ca2+-free Ringer’s solution for 15 min. During this incubation, yolk was removed by gently pipetting embryos through a 200 μl tip (with the end cut off) three to five times. Yolk-free embryos were transferred into a 35 mm petri dish with 5 ml of 0.25% Trypsin-EDTA (Sigma-Aldrich). Embryos were incubated at 28.5°C and homogenized with a 200 μl tip every 10 min until most cells were dissociated (20 to 50 min). 55 μl 100 mM CaCl2 and 350 μl FCS were added to stop digestion. Dissociated cells were transferred into a 15 ml tube and centrifuged at 300 g for 5 min at 15°C. Cells were rinsed once with 10 ml suspension solution (colorless Leibovitz medium L-15 with 0.3 g/L glutamine, 0.8 mM CaCl2, Penicillin 50 U/ml, Streptomycin 0.05 mg/ml, and 1% FCS). Dissociated cells were resuspended in suspension solution to ~107 cells/ml and immediately proceeded to cell sorting at the UCLA Broad Stem Cell Research Center Flow Cytometry Core. BD FACS ARIA II SORP instruments sorted cells, using a 488 nm laser for GFP detection and a 561 nm laser for dsRed detection.

**RNA-Seq libraries**

Sorted cells in suspension solution were immediately lysed using the Qiagen RNeasy kit. Lysis was completed within two hours of dissociating cells. Total RNA was isolated following the Qiagen RNeasy kit and stored at –80°C. Quality of all samples was assayed with an Agilent Bioanalyzer, and only samples with RNA Integrity Number > 8 were used for creating sequencing libraries. Poly-A bead-purified RNA-Seq libraries were prepared with the Illumina TruSeq RNA Library Prep Kit.

**Sequencing**

23 RNA-Seq libraries (one per experimental replicate) were run a total of 25 times in various combinations across five full and fractional single end 50 or 51 nt Illumina HiSeq 2000/2500 high output lanes (with ~1% PhiX spike-in as internal control), and demultiplexed (allowing a single mismatch to expected 7-mers) to obtain 21.2 to 45.4 million PF = 1 reads per replicate. Adapters at 3’ read ends and low quality bases were trimmed with CutAdapt 1.8.1 (Martin 2011, --m 21 --trim-n --max-n=2 –q 10,10 –a AGATCGGAAAGACACGCTACGGAAGTCGTAAGCTACGTCAGTAC); in each replicate, ≥ 98.2% of reads and ≥ 97.9% of bases survived.

**Alignment**

For cross-experiment fairness of mapping, only the first 47 nucleotides of each trimmed read were kept (retaining only trimmed reads ≥ 47 nt long). The resulting 21.0 to 45.2 million reads per replicate were aligned (with per-base quality scores) to the Ensembl release 92 (http://april2018.archive.ensembl.org/Danio_rerio/Info/Index) top-level reference Zebrafish genome (GRCz11 chromosomes 1 to 25 + mitochondrion + 847/120-sequence subset of KN/KZ scaffolds, augmented with PhiX; alternate sequences excluded) with STAR 2.5.3a (Dobin et al. 2013, in single pass mode using known junctions from the Ensembl release 92 Zebrafish top-level reference annotation set [consisting of 31,901 genes — of which 25,431 are protein_coding — and 58,867 transcripts, of which 51,233 belong to protein_coding genes], retaining multiple locations for non-unique reads; --alignEndsType EndToEnd --outFilterMultimapNMax 999 --quantMode TranscriptomeSA GeneCounts --outSAMtype BAM SortedByCoordinate --outSAM unmapped Within --outMutlimapperOrder Random --outSAMattributes All --alignIntronMin 10 --alignIntronMax 700000 --chimSegmentMin 20 --chimSegmentReadGapMax 3 --chimMainSegmentMultNmax 999 --winAnchorDistNbins 12 --winAnchorDistNbins 12 --winAnchorDistNbins 12 --winAnchorDistNbins 17 --winAnchorDistNbins 6). Per replicate, 96.4–97.6% of reads aligned, and, of those aligned, 87.5–95.5% aligned uniquely.
Quantiﬁcation
Per-library count distributions to the Ensembl release 92 top-level reference Zebraﬁsh transcripts were formed with Salmon 0.10.2 (Patro et al. 2017, submitting and including a patch that became part of Salmon 0.11.0 to ﬁx a bug discovered during this work) from the STAR transcriptome alignments (numGibbsSamples=1000 --thinningFactor=25 --useBpOpt --HybridType=U --FlDMean=200 --FlDSD=200 --rangeFactorizationBins=4 --minAssignedFrgs=1 --seqBias --noBiasLengthThreshold). For each replicate, 1,000 Gibbs bootstrap count distribution samples were retained to enable technical variance estimation in statistical analyses.

Statistical analyses
We followed the general outline of R Bioconductor package Sleuth (Pimentel et al. 2017, as if the Wasabi package (https://github.com/COMBINE-lab/wasabi) was used to import) in modeling per gene, per condition log-scale expression as normal distributions, with variance partitioned into a technical, assay component (informed by Salmon bootstraps) and a biological component (informed by replicate experiments within conditions). However, we manually conducted all analysis steps to have greater control and incorporate some features of the DESeq2 package (Love et al. 2014) that Sleuth does not support (e.g., we use p-values with 1.5x-fold change thresholds, rather than for any detectable difference); details are in supplemental methods (File S1).

Gene ranking for Figure 5
This analysis focused on genes that (1) were highly enriched in skin, and (2) had strong layer-speciﬁc expression. Using normalized transformed per-condition model mean counts (see File S1), we scored (1) as A − N, where A was the mean of the three all skin conditions, and N was the mean of the three nonskin conditions; and we scored (2) as the sum-of-absolute-deviations-from-mean for the four basal cell and periderm conditions. Genes were then ordered by descending sum of these two component scores.

In situ hybridization
In situ hybridization was performed according to the Thisse Lab In Situ Hybridization Protocol 2010 Update (updated from: Thisset and Thisset 2008). The 1,073 bp keratin 4 (krt4) probe was synthesized using primers 5′-TAAGACCTCAACACGCGT-3′ and 5′-TAATACGACTATAAGGGCTACGGTATCTGACACACC-3′. The 1,122 bp aeroysin-like protein 1 (aspl) probe was created using primers 5′-TGGGTGTGGTGGTGGATG-3′ and 5′-CATTAACTCTCAGAAAAGAGGATGTTGGTATCG-3′. The 617 bp transcupulamin beta b (tcub) probe was generated using primers 5′-GC-ACCTGAGAAGCTCTGTAAG-3′ and 5′-TTGGAGATTACATTACATGCT-GGAGA-3′. The 1,265 bp hephaestin-like 1a (hepl1) probe was constructed using primers 5′-GGACATCAGCATGCAGAGAA-3′ and 5′-GCCAAGATATTACAATGCT-3′. For each gene, 100 ng of probe was added into 400 μl of hybridization mix and hybridization incubation was conducted overnight at 70°C.

BAC transgenes
We generated translational fusion transgenes by inserting a GFP reporter gene cassette directly preceding the stop codon of target genes in Bacterial Artificial Chromosomes (BACs) (Suster et al. 2011). To create pCS2+_linkEGFP_KanR, the amino acid linker sequence 5′-GGGSGGG-3′ was added upstream of the EGFP initiation codon by PCR ampliﬁcation and the resulting linkEGFP fusion gene was cloned in place of gal4FF in plasmid pCS2+_gal4FF_KanR (Bussmann and Schulte-Merker 2011) using the restriction enzymes BamHI and XbaI. To create TgBAC(col28a1a-EGFP), the linkEGFP_KanR cassette was recombined in place of the col28a1a stop codon in BAC CH211-17D12 using the primers 5′-GCAACGGCCTTGGAAACAGGAGGATCTTGAACAGC-CTTGTGTTGACACAGGATCGTGGAGGTT-3′ and 5′-GGCTTGATTTAAAGAACAGCATTGTAAGTGGTCTTGTGCA-GAGTTGTCAGAAGAACTCGTGCAAGAAAGCG-3′. To generate TgBAC(ochb-EGFP), primers 5′-CATAACCTCCCTACAAAGAAGGGTGTACGACTAACAGACAAGACA-3′ and 5′-GTCCAATTGTTAAAACACCGATATGCTGCTGATTCTCAGGCG-3′ were used to recombine the linkEGFP_Kan cassette in place of the ochb stop codon in BACs CH73-377J and CH211-65N9. BAC transgenes were isolated with a Qiagen Midi kit and microinjected into embryos generated by two sets of transgenic lines, Tg(krt5:Gal4) crossed with Tg(UASsfsB-mCherry), and TgBAC(LNp63:Gal4) crossed with Tg(UASsfsB-mCherry) (Curado et al. 2007; Rasmussen et al. 2015) to create transient GFP expression in skin cells.

Microscopy
For confocal imaging, live zebraﬁsh embryos were mounted in 0.02% tricaine and 1% low melt agarose, and imaged with a Zeiss LSM 710 and 800 confocal microscope. For in situ hybridization images, ﬁxed and stained embryos were placed in 100% glycerol and imaged with a Leica M165 FC Stereomicroscope, with attached Leica DMC2900 camera.

Comparison to de la Garza
See File S1 for methodological details, including conversion of 85% of the pre-2009 Ensembl transcript identiﬁers involved to current genes. The modernized periderm proﬁle (converted from de la Garza et al. 2013 Supplemental Table S1) containing 1,151 genes (all but ﬁve of which are protein coding), dnrlfsb-inhibited proﬁle (converted from de la Garza et al. 2013 Supplemental Table S2) containing 344 genes (all protein coding), and the intersection of these containing 87 genes were compared to our study (see File S1 and File S7).

Data availability
Fish lines and BAC transgenes are available upon request. Reads, quantiﬁcations, and statistical intermeiates and results are available within NCBI GEO series GSE132304. File S1 contains supplemental methods. File S2 contains comprehensive heatmaps of Gene Ontol-ogy enrichments (retaining all rows and columns containing any p-value ≤ 0.0001), extending Figure 4. File S3 contains a comparison of keratin genes and chromosomal locations within our ﬁndings and Krushna Padhi et al. 2006. File S4 contains InterPro domain hit summaries used in the compilation of zebraﬁsh type I and II keratins. File S5 contains comprehensive heatmaps of expressed genes (those with normalized transformed [log2-scale] counts ≥ 2.5 [i.e., ≥ ~5.7 in linear scale] in at least one condition) that encode cell surface receptors, transcription factors, and actin-binding proteins, extending Figure 5B–D. For convenience, File S6 repeats our gene ﬂow classiﬁcations included inside NCBI GEO GSE132304. File S7 contains details of the comparison between our study and de la Garza et al. (2013). Figure S1 contains a phylogenetic tree of keratin and intermediate ﬁlament genes in zebraﬁsh. Figure S2 contains gene expression plots (in similar style as Figure 2A) for all type I and II keratin zebraﬁsh genes. A searchable website with gene expression proﬁles plotted in the style of Figure 2A is available at https://zfishskin.net. Supplemen-tal material available at FigShare: https://doi.org/10.25387/g3.8266763.
RESULTS

Developmental transcriptomes for two skin epithelial layers

To identify the genetic programs that drive maturation of periderm and basal cells, and to determine their distinguishing characteristics, we profiled gene expression at three developmental stages: the 20 somite stage (20 SS occurs at approximately 19 hpf in embryos raised at 28.5°C (Kimmel et al. 1995), when the two epithelial skin layers are not yet fully defined; 52 hpf, when the two epithelial layers are established; and 72 hpf, when both layers have matured. These timepoints encompass the development of key epithelial features, including the formation of cell–cell junctions, the production of a specialized extracellular matrix, microridge morphogenesis on periderm cells, and formation of axon sheaths by basal cells (O’Brien et al. 2012).

To isolate each cell type, we used two transgenic fish lines: krt4:DsRed and krt5:GFP (Hu et al. 2010; O’Brien et al. 2012). Expression of the krt4:DsRed reporter begins at an early embryonic stage and labels both skin cell layers, but not other tissues, at all three developmental stages (O’Brien et al. 2012; Wang et al. 2012). krt5:GFP is not significantly expressed at 20 SS, but exclusively labels periderm cells (and not basal

![Figure 1](image-url)

Figure 1 Two epithelial skin cell types isolated with FACS from transgenic fish. A–C) Confocal images of the skin covering the dorsal head region in 52 hpf embryos expressing GFP in periderm (Tg(krt5:GFP)), and RFP in both skin layers (Tg(krt4:dsRed)). D–G) Cells dissociated from 52 hpf embryos were separated by FACS. For each panel, green and red fluorescence intensities are indicated on x- and y-axes, respectively. FACS plots show gates for each cell population from wildtype animals (D), Tg(krt4:dsRed) embryos (E), Tg(krt5:GFP) embryos (F), and embryos with both transgenes (G). Cells in “All skin”, “Nonskin1”, “Basal cells”, “Periderm”, and “Nonskin2” boxes show gates that were used to collect cells into separate tubes for generating RNA-Seq samples.
cells) at 52 hpf and 72 hpf (Hu et al. 2010). By crossing these two lines, we obtained double transgenic embryos that enabled the separation of skin cells (expressing krt4:DsRed) from the rest of the embryo (no fluorescence) at 20 SS. At 52 hpf and 72 hpf, these transgenic reporters distinguished periderm cells (expressing both transgenes) from basal cells (expressing only krt4:DsRed) as well as the rest of the embryo (no fluorescence) (Figure 1A–C).

Using FACS, we separated dissociated cells into specific isolates (Figure 1D–G). This enabled construction of the following 12 experimental conditions: “all skin cells” (RFP+) at 20 SS, 52 hpf, and 72 hpf; “nonskin1” cells (RFP−) at 20 SS, 52 hpf, and 72 hpf; “periderm” (RFP+ GFP+) at 52 hpf and 72 hpf; “basal cells” (RFP+ GFP−) at 52 hpf and 72 hpf; and “nonskin2,” cells (RFP− GFP−) at 52 hpf and 72 hpf. We isolated two biological replicates for each condition, except for a single iteration of nonskin2 cells at 52 hpf.

An RNA-Seq library was prepared, sequenced, aligned, and quantified for each replicate experiment to obtain per-experiment, per-transcript estimated read count distributions (Methods and File S1); genes were handled as arithmetic sums of their transcript isoforms. The Gibbs samples of quantifications afforded estimates of per-gene technical variance, and biological replicates within conditions in combination with a shrinkage procedure enabled estimation of per-gene biological variance. Normalized expected read counts for genes (averaged over replicates within conditions), with awareness of total (technical plus biological) variance for each gene, were used for statistical analyses (e.g., for differential expression).

To visualize each gene’s expression pattern, we plotted its expression across time per tissue. For example, Figure 2A shows three previously unknown skin enriched genes: aep1, tcnbb, and hephl1a. (These three genes were selected for verification by in situ hybridization (Figure 6).) At 52 hpf and 72 hpf, aep1 was highly expressed in both layers, whereas tcnbb was specifically enriched in periderm cells. By contrast, hephl1a was expressed at much higher levels at 20 SS than at later stages. Similar plots for all genes, searchable by Ensembl identifier or gene names/keywords, are available on a website (https://zfshskin.net).

**Dynamics of gene expression in epithelial skin layers during early development**

For a high level overview of the relationships among the genetic programs of different cell types, we performed blind clustering of the experimental expression profiles (Figure 2B). Replicate-to-replicate variation was smaller than cross-condition variation, except among nonskin1 and nonskin2 experiments, which were quite similar. As expected, the transcriptomes of skin vs. nonskin populations were most different from one another. Cell populations were then divided by timepoint: for both skin and nonskin, profiles at the 20 SS stage were clearly more different from those at the 52 and 72 hpf timepoints than the latter two stages were from each other. Skin samples next diverged in expression by layer, with 52 and 72 hpf only distinct at the finest levels of comparison (apart from replicate experiments) at this scale. At 52 and 72 hpf, the expression profiles of samples involving all skin cells were between those of basal cells and periderm, consistent...
with all skin being a physical mixture of the layers. These observations illustrate that, despite their differences, distinct epithelial cell types share much in common.

To study the different dynamic patterns of gene expression, we classified every gene into categories at each timepoint (see File S1 and File S6). At 20 SS (when we did not separate the two skin layers), each gene expressed significantly more highly in all skin relative to nonskin_1 was placed into the “skin genes” (S) category, and the rest into the “nonskin genes” (N) category. At each of 52 and 72 hpf, a gene was placed into one of four categories based on the probabilities of the various orderings that periderm, basal cells, and nonskin_2 cells could have had by the gene’s expression at that timepoint, with a 1.5x-fold change threshold for significant differences. A non-negligible probability (≥ 0.02) that neither periderm nor basal cells was significantly higher than nonskin_2 cells placed the gene in the “nonskin” (N) category; otherwise, a non-negligible probability that periderm and basal cells were not significantly different placed the gene in the “general skin” (G) category; and otherwise the gene was placed in the “periderm-preferred” (P) or “basal-preferred” (B) categories, according to which layer was more likely highest. Placing genes into these categories allowed us to identify both common and distinguishing features of periderm and basal cells. Altogether, 23% of all 31,901 genes — the 7,286 “skin-enriched genes” — were in a skin category (S/G/B/P) at 52 SS, 52 hpf, and 72 hpf, in that order. Figure 3 represents all combinations by concatenating the category characters for 20 SS, 52 hpf, and 72 hpf, respectively, indicating that major aspects of cellular specification occurred between 20 SS and 52 hpf, and fewer between 52 hpf and 72 hpf. The largest changes between 52 and 72 hpf were exchanges between general skin genes and nonskin genes, but more genes changed from nonskin to general skin (~2:1), suggesting that subsequent maturation likely involves recruitment of new genes.

Basal- and periderm-preferred genes were quite stable. The four flows involving B vs. P exchanges — SBP, SPB, NBP, and NPB — had zero genes. Thus, of the 844 basal-preferred genes at 52 hpf, zero were periderm-preferred at 52 hpf while 78% remained basal-preferred (and 13% became general skin and 9% nonskin, with some fraction of these expected to be borderline cases). Similarly, of the 1,322 periderm-preferred genes at 52 hpf, zero were basal-preferred at 72 hpf while 69% remained periderm-preferred (and 21% became general skin and 10% nonskin, again with some fraction expected to be borderline cases). These results suggest that fate specification for the periderm and basal cell layers is largely irreversible by 52 hpf.

**Differential functional enrichments in two distinct epithelial skin layers during early development**

To unveil the unique characteristics of different developmental stages and epithelial layers, we used functional annotation enrichment analyses to examine every gene expression flow, as well as certain combinations of flows. An asterisk at a timepoint indicates any expression category; for instance, “BB” refers to genes that were in any category at 20 SS (N or S), but were in the basal-preferred category at both 52 and 72 hpf. Furthermore, at each of 52 hpf and 72 hpf, genes enriched in skin in any way were partitioned across three categories (B/P/G), and S (“any skin”) indicates any of these three ways. For example, “SSS” indicates genes that were skin genes at 20 SS and also belonged to any of the skin categories (B/P/G) at both 52 and 72 hpf (and not necessarily

**Figure 3** Expression dynamics of skin-enriched genes. Using RNA-Seq expression profiles, the 31,901 Ensembl release 92 zebrafish genes were classified (see File S1) at 20 SS into categories of “skin genes vs. ‘Nonskin genes, and at each of 52 and 72 hpf into categories of Periderm-preferred genes, Basal-preferred genes, General skin genes, and ‘Nonskin genes. Each gene thus belongs to exactly one of 32 possible “flows”, of which 28 have at least one gene (all of which except NNN are depicted, with width proportional to the number of genes). 15 flows have at least 70 genes, and 10 flows have at least 200 genes.
the same category at 52 hpf vs. 72 hpf). To display some of the key biological differences distinguishing cell populations, we show 15 representative flows/flow combinations and 74 representative Gene Ontology (GO) terms — 23 Cellular Component (CC) terms, 23 Molecular Function (MF) terms, and 28 Biological Process (BP) terms — in Figure 4. (Comprehensive heatmaps are available in File S2.) These analyses revealed layer-specific gene categories, and gene categories that changed as skin matured.

The basal cell-preferred (NBB, **B**, and SBB) and periderm-preferred (SPP, *PP*, and NPP) gene flows/combos were differentially enriched in several CC, MF, and BP GO terms. Basal flows/combos were enriched with GO terms indicating extracellular...
matrix (especially basement membrane) and integrin complex components; actin binding and growth factor binding functions; and cell adhesion, migration, and signaling processes. By contrast, periderm gene flows/combinations were enriched in cell–cell junction (especially tight junction) and endomembrane/GoGi components; ion, nucleotide, GTP, and ATP binding functions; and glycosylation, nucleotide-sugar, and fatty acid biosynthetic processes. Layer-specific gene flows committing to skin later (NBB and NPP) were enriched in fewer GO terms than flows for layer-specific genes committing to skin early (SBB and SPP), suggesting that later-enriched genes might be more diverse, resulting in fewer significant GO term enrichments.

Flows/combinations focusing on general skin genes (**G, **GG, and SGG) were enriched in GO terms that were mostly distinct from those for flows/combinations involving basal cell-preferred or periderm-preferred genes (although a few terms were shared with periderm), reflecting common skin cell features. For example, desmosome and cytoskeletal (actin) components, and actin filament polymerization, cell death, translation, and protein transport processes were enriched in flows/combinations focused on general skin, and thus represent common epithelial features. Flows/combinations involving any skin categories (**S, **SS, and SSS) more or less combined enrichments seen in the various more specific flows/combinations, as might be expected.

GO terms enriched at different stages revealed aspects of the maturation process. For example, genes that were skin genes at 20 SS, but were nonskin genes later (SNN) were enriched in signaling processes. These signaling processes could regulate steps in differentiation, or, since the peripheral axons of somatosensory neurons have just begun arborizing between the two skin epithelial layers at 20 SS (O’Brien et al. 2012), they might reflect interactions between skin cells and somatosensory neurons. The transient general skin gene flow (NGN) was enriched with COPI vesicle coat components, suggesting that the production of secreted or membrane proteins might be enhanced at a specific step of skin cell maturation. Chloride transmembrane transporter activity, cytosolic ribosome components, and translation were enriched in the late general skin gene flow (NNG), indicating that relatively mature epithelial cells increase active protein production.

Characterization of keratin gene expression patterns in two skin epithelial layers
Keratins are types of intermediate filaments expressed in vertebrate epithelial cells, providing them with structural integrity (Bragulla and Homberger 2009; Coulombe and Lee 2012). Although keratin filaments are abundant in all epithelial cells, specific keratin genes have restricted expression patterns, and are thus often used as diagnostic markers for specific cell types and developmental stages (Byrne et al. 1994; Conrad et al. 1998; McGowan and Coulombe 1998; Mazzalupo and Coulombe 2001; Lu et al. 2005; Krushna Padhi et al. 2006). Intracellular keratins are formed from heterodimers of type I (acidic) and type II (neutral-basic) keratin proteins. The last comprehensive analysis of keratin genes in zebrafish was reported in 2006, and identified 16 type I and seven type II zebrafish keratin genes (Krushna Padhi et al. 2006). Since the zebrafish reference genome has significantly improved since 2006, we re-surveyed its modern gene models to identify type I and type II keratins (see File S1, File S3, Figure S1, and File S4), adding seven type I keratin genes and removing one type II gene relative to the previous analysis, resulting in a total of 23 type I and six type II zebrafish keratin genes (Figure 5A). As previously noted, many of the type I genes are clustered on chromosomes 11 and 19, and five of the new type I genes added are located in these clusters. The revised keratin gene identifications provide an updated reference for the zebrafish community.

The seven added type I genes were krt17, krt18a.2, krt94, krt96, krt98, cyt11, and sick121-156i18.7 (the last being homologous to keratin19 in many organisms); the ones retained were krt15, krt18a.1, krt11b, krt91, krt92, krt93, krt95, krt97, krt99, krt1-c5, krt1-19d, krt11c19, krt11c10, krt11c19e, and cyt1. The type II genes were krt4, krt5, krt8, krt12c6, krt12c8, and krt12c22; we removed si:dkey-183i3.5 a.k.a. krt12c21 (Krushna Padhi et al. 2006) a.k.a. thread keratin alpha (Schaffeld and Schultess 2006). All of our accepted type II genes are much closer to one another in sequence similarity than any of them are to si:dkey-183i3.5, and the same holds for all of our accepted type I genes, and the InterPro domain structure details of si:dkey-183i3.5 are not like those of our accepted type I or type II keratin genes (Figure S1 and File S4). Similar remarks apply to another gene, zgc:136930, also called thread keratin gamma (Schaffeld and Schultess 2006), and a third gene, krt222. However, these three genes do appear to code for intermediate filaments (File S4).

Six type I keratin genes (krt18a.2, krt93, krt95, krt98, krt11c1, and sick121-156i18.7) had very low expression levels (mean expression level < ~10 normalized counts) in all conditions, suggesting that they do not participate in early development (Figure 5A and Figure S2). The other 17 type I keratin genes and all six type II keratin genes were expressed at higher levels, and enriched at some developmental stage in at least one type of skin cell, compared to nonskin cells (Figure 5A and Figure S2). Nine type I keratin genes (krt17, krt18a.1, krt91, krt92, krt97, krt11c9, krt11c19e, cyt1, and cyt11) were enriched in skin and highly expressed (mean expression level > ~4,000 normalized counts; in every condition, this was at least the most extreme 3% of genes). Among these nine genes, krt91 and krt11c9 were basal-preferred genes, whereas krt17, krt92, cyt1, and cyt11 were periderm-preferred genes. Furthermore, krt91 and krt11c19e were expressed at much higher levels at the two later developmental stages, whereas krt18a.1 and krt92 were expressed at higher levels at 20 SS.

Among the six type II genes, krt4, krt5, and krt8 were, by far, the most highly expressed in skin, and krt4 was the gene — in every condition involving skin — with highest normalized counts of any gene of any kind. (The gene with highest normalized counts in nonskin1, at 20 SS was an eukaryotic translation elongation factor, and in either nonskin or at 52/72 hpf were embryonic hemoglobins.) krt22c8 was strongly periderm-preferred, and the only layer-specific type II gene. krt4, krt5, and krt8 were comparably highly expressed in both layers. Interestingly, the highly periderm-specific fluorescent reporter Tg(krt5:GFP) we used for periderm FACs purification was based on an enhancer fragment from the krt5 gene (Hu et al. 2010), indicating that the elements in this fragment were not sufficient to drive expression in the native gene’s full pattern.

Skin transcriptomes reveal genes that likely promote cell type-specific characteristics
As cells differentiate and mature, they integrate signals from their environment, activate unique transcriptional programs, and undergo morphogenetic cell shape changes driven by the actin cytoskeleton. To identify specific genes that may provide insight into how periderm and basal cells adopt distinct developmental trajectories, we examined the expression patterns of genes annotated as cell surface receptors, transcription factors, and actin-binding proteins (as defined in File S1), with an emphasis on genes that (1) were highly enriched in skin, and (2) had strong layer-specific expression. We scored genes by a sum of these two criteria (see Methods) and selected for display the top 20 genes in each category (Figure 5B–D).
Among the top 20 cell surface receptors, many are involved in G protein-coupled receptor signaling pathways (Figure 5B). For example, G protein-coupled receptor 84, leukotriene B4 receptor 2a, and leukotriene B4 receptor 2b were basal cell-preferred genes, while adhesion G protein-coupled receptor F6 and adhesion G protein-coupled receptor G2a were periderm-preferred genes. Unsurprisingly, many of the top 20 transcription factors (Figure 5C) were expressed in a specific epithelial layer. Notably, tp63, a well-characterized basal cell gene (Vanbokhoven et al. 2011), was one of the top basal cell-preferred transcription factor genes. Although actin-related GO terms were generally enriched in skin cells of both layers (Figure 4), our gene-specific analysis (Figure 5D) revealed that each skin layer expressed distinct sets of actin-binding proteins. These genes (and other differentially expressed genes on the comprehensive lists; see File S5) are candidate entry points for investigating unique signaling pathways activated in each cell type, gene regulatory networks that specify these cells, and the cytoskeletal processes that endow epithelial cells with unique morphological features, such as microridges in periderm cells and ensheathment channels in basal cells.

In situ hybridization and fluorescent transgenic reporters support RNA-Seq findings

Our RNA-Seq analyses identified genes known to be expressed in skin, but also numerous genes previously unknown to be expressed in a layer- and/or developmental stage-specific manner in the embryonic skin. To confirm developmental expression patterns of representative genes, we...
Interestingly, a recent study indicated that Aep1 is an innate immune gene. With RNA-Seq analyses, we chose novel candidates suggested by our RNA-Seq analyses and examined their expression patterns with in situ hybridization. Since krt4 (Figure 6A–C) is a gene known to be strongly expressed in all skin cells at all three timepoints, it served as a positive control. The gene aerolysin-like protein (aep1) codes for a pore-forming protein, and the gene transcobalamin beta b (tcnbb) codes for a vitamin B12-binding glycoprotein; both were indicated by our RNA-Seq to be skin-enriched genes, with much higher expression levels at 52 and 72 hpf than 20 SS, whereas hephaestin-like 1a (hephl1a) codes for a metal transporter and was more highly expressed at 20 SS than 52 and 72 hpf (Figure 2A). There were limited prior expression data, primarily based on RT-PCR, for aep1, hephl1a, or tcnbb (Lam et al. 2008; Nakajima et al. 2011; Long et al. 2015; Chen et al. 2018; Benoit et al. 2018). None of these three genes had previously been suggested to be expressed in skin during early development.

We detected an in situ hybridization signal for hephl1a exclusively at 20 SS (Figure 6I–L); by contrast, aep1 and tcnbb transcripts were detected in skin cells at 52 hpf and 72 hpf, but not at 20 SS (Figure 6D–I), consistent with our RNA-Seq analyses. Interestingly, a recent study indicated that Aep1 is an innate immune molecule that inhibits bacterial infection (Chen et al. 2018). Our results are consistent with these findings and suggest that the skin is important in innate immune responses, especially during early developmental stages. In situ hybridization revealed that tcnbb had an intriguing spatiotemporal distribution pattern, which was not possible to resolve with our RNA-Seq design: in situ signal was higher in skin covering the head and yolk than in other areas at 52 hpf (Figure 6H), and further concentrated in the anterior end at 72 hpf (Figure 6I). A recent study showed that tcnbb expression is further restricted to a unique ventral-anterior area at 5 days post-fertilization (Benoit et al. 2018). These observations demonstrate that our RNA-Seq analyses were sensitive enough to detect regionally restricted genes, and suggest that the skin is regionalized early in development.

To verify layer-specific expression patterns identified by our RNA-Seq analyses, we constructed BAC transgenes with translational GFP fusions. The estimates of absolute expression that RNA-Seq provides were especially useful for these experiments, since the utility of BAC transgenes depends on enhancer strength.

We chose the periderm-preferred gene occludin b (oclnb) and basal expressing basal cells. Dotted line outlines area containing Col28a1a-GFP, which had spread beyond a single basal cell. Large images are 2-D (xy) projections from 3-D confocal z-stacks. Arrows point from apical surface (ap) to basement membrane (bm) for xz- and yz-planes. Scale bars are 20 μm long.

Figure 6 Expression patterns of four skin-enriched genes detected by in situ hybridization. A–C) krt4 was highly expressed at all three stages (20 SS, 52 hpf, and 72 hpf). The inset in (A) shows an enlarged image of krt4 expression in skin. D–F) aep1 was only expressed highly at the two later stages (52 and 72 hpf). G–I) tcnbb was only highly expressed at 52 and 72 hpf. The top inset in (H) displays tcnbb expression in skin covering the head region, while the bottom inset shows expression in skin at the trunk region. The inset in (I) shows an enlarged image of tcnbb expression in skin at the anterior end. J–L) hephl1a expression was detected at 20 SS in skin, but not later stages. All images are oriented with dorsal to the top, ventral to the bottom, anterior to the left, and posterior to the right.

Figure 7 BAC transgenes illustrate layer-specific gene expression and protein subcellular distribution. A) Confocal images of a 72 hpf Tg(krt5:Gal4;UAS:mCherry) embryo with TgBAC(oclnb-EGFP) expression. Oclnb-EGFP localized to cell junctions in periderm cells, which also expressed mCherry. B) Confocal images of a 72 hpf Tg(ΔNp63:Gal4;UAS:mCherry) embryo with TgBAC(oclnb-EGFP) expression. Oclnb-EGFP was not detected in mCherry-expressing basal cells. C) Confocal images of a 96 hpf Tg(ΔNp63:Gal4;UAS:mCherry) embryo with TgBAC(col28a1a-EGFP) expression. Col28a1a-EGFP was detected in the basement membrane, directly basal to mCherry-expressing basal cells.
cell-preferring gene collagen XXVIII alpha 1a (col28a1a), as these were both strongly layer-specific and highly expressed (https://zfishtz.com). We used two sets of transgenic lines, Tg(krt5:Gal4) crossed with Tg(UAS:nfsB-mCherry) (fluorescing only in periderm), and TgBAC(DNp63:Gal4) crossed with Tg(UAS:nfsB-mCherry) (fluorescing only in basal cells) to separately label the two skin epithelial layers (Curado et al. 2007; Rasmussen et al. 2015). Confocal microscopy of these transgenic fish microinjected with the ochnb-GFP BAC transgene revealed that ochnb was indeed highly expressed exclusively in periderm cells and localized to cell boundaries (Figure 7A–B), consistent with its role as a tight junction component. By contrast, Collagen XXVIII alpha 1a is a member of the collagen family (Gebauer et al. 2016) that has not been fully characterized. Confocal images of embryos injected with the col28a1a-GFP BAC transgene confirmed strong GFP signal just below basal cells (Figure 7C). Strikingly, the area of GFP signal was broader than the clones of cells expressing the transgene, suggesting that Collagen XXVIII alpha 1a was secreted by basal cells and diffused within the plane of the basement membrane. Collagen XXVIII alpha 1a had not previously been reported to be a basement membrane component, demonstrating that our data has the potential to reveal novel aspects of epithelial biology.

**DISCUSSION**

By combining transgenic fish lines, cell purification, and RNA-Seq, we report comprehensive transcriptomes for two epithelial skin layers at three early developmental stages in zebrafish. Our observations verified the expression patterns of genes known to be expressed in skin, and, more importantly, identified numerous novel layer- and/or developmental stage-specific genes. This investigation can thus serve as a resource for identifying epithelial layer-specific enhancers, studying the functions of layer- and/or developmental stage-specific genes, and creating tools to label specific cell types at specific developmental stages.

Our data indicated that expression of specific junctional and ECM proteins are key cellular features distinguishing periderm and basal cells. While periderm and basal cells expressed comparable levels of adherens junction and desmosomal proteins (e.g., E-cadherin and Desmoplakin a and b), only periderm cells expressed high levels of tight junction proteins, such as occludins (e.g., Figure 7); and only basal cells expressed high levels of hemidesmosome proteins, such as Integrin alpha 6b (https://zfishtz.com). These expression patterns were consistent with the periderm’s role as a barrier, and the basal cell layer’s role in attachment to the basement membrane. Secreted extracellular proteins are also major distinguishing features of periderm and basal cells. Basal cells expressed high levels of basement membrane components such as Collagen IV chains (and potentially novel basement membrane-associated proteins, such as Collagen XXVIII alpha 1a), whereas periderm cells were enriched for glycosylation enzymes that likely contribute to the formation of the apical glycocalyx. Intriguingly, periderm cells also expressed high levels of at least one mucin — Mucin 13a (https://zfishtz.com). Mucins are thought to primarily be secreted by goblet cells and diffuse onto the apical surface of periderm cells (Linden et al. 2008). Since ECM proteins are primarily located basal to the basal cell layer or apical to periderm cells, our data could provide a useful resource for studying polarization sec.

Our data also provide a foundation for better understanding the function of the periderm, an under-studied and relatively enigmatic tissue that is eventually sloughed off in both mammals and fish (Wolf 1967, 1968a, 1968b; M’Boneko and Merker 1988; Lee et al. 2014; Richardson et al. 2014). Periderm cells serve as the main barrier between embryos and the external environment, and thus contain tight junctions and display a glycocalyx on their apical surface. Defects in periderm differentiation or desquamation cause a variety of developmental abnormalities (Cui et al. 2007; Okano et al. 2012; de la Garza et al. 2013; Richardson et al. 2014; Liu et al. 2016). Our data identified specific junction genes, glycosylation factors, and transmembrane and secreted proteins that likely create these periderm-specific features. The apical surfaces of periderm cells project prominent actin-based microridges arranged in striking labyrinth-like patterns (Wolf 1967, 1968a; Lam et al. 2015; Pinto et al. 2019; Depasquale 2018). Although our study showed that actin regulatory genes as a category are enriched in both skin layers, specific actin-binding genes enriched in periderm may play specific roles in microridge morphogenesis. In mammals, the periderm disappears as the skin stratifies and the outer layers of the mature epidermis keratinize (Wolf 1968a, 1968b; Holbrook and Ollandt 1975; Hardman et al. 1999). By contrast, the basal cell-derived outer layer of the stratified adult skin does not cornify in fish, but takes on many of the characteristics of the embryonic periderm (Lee et al. 2014), reflecting the fact that fish skin is a mucosal epithelium at both embryonic and adult stages. Given their similarities, the periderm-like cells of the adult fish skin likely share much of the embryonic periderm gene expression program we report here.

A previous microarray-based study of the skin differentiation transcription factor Interferon regulatory factor 6 (Irf6) identified genes with expression enriched very early (4 hpf for basal cells develop) in wildtype zebrafish periderm cells, as well as genes inhibited at 6 hpf by dnIrf6, a dominant-negative variant of Irf6 (de la Garza et al. 2013). We converted the microarray periderm profile, dnIrf6-inhibited profile, and profile intersecting these two together to current genes to compare with our study (see Files S1 and S7). At 20 SS (our closest timepoint), all three profiles were enriched for genes we classified as ‘S’ (skin genes): 30%, 39%, and 75% of the periderm, dnIrf6-inhibited, and intersection profiles, respectively, were ‘S’ compared to 8% over all genes (4x to 10x higher than random expectation, hypergeometric p-values < 10^-35). Expanding to all three of our timepoints, the intersection profile was strikingly enriched for our periderm classifications, with our flow SPP being the most frequent in that profile (46% of its genes: 26x, p < 10^-43). In the other two profiles, SPP was the second most common flow (>7x, p < 10^-39), below flow NNN. Hence, genes in these early profiles — especially the intersection profile — may indeed be key genes in establishing periderm identity.

Basal cells are stem cells that later serve as the source for stratification and diversification of the epidermis (Muroyama and Lechler 2012; Gonzales and Fuchs 2017), and are transformed in basal cell carcinomas (White and Lowry 2015). At the stages we studied, basal cells establish a unique relationship with the endings of sensory neurons that innervate the skin (O’Brien et al. 2012). Axons first innervate the region between basal and periderm cells, but then become enveloped exclusively by basal cells in glial-like ensheathment channels. Formation of these channels requires specific lipid microdomains, association with actin, and the formation of autotypic junctions (Jiang et al. 2019), but the signals that initiate ensheathment are unknown. The basal cell-specific genes we identified suggest candidate receptors for ensheathment signals, actin regulatory proteins that could promote membrane invagination, and specific components of autotypic junctions. Basal cells also play critical roles in wound healing, proliferating and migrating to repair epidermal damage. Repeating our experiments in wound paradigms (LeBert and Huttenlocher 2014) could identify basal cell genes involved in wound healing.
The cellular and molecular similarities between zebrfish and hu-
mans make them an efficient model for studying human diseases,
including skin diseases (Li et al. 2011b; Li and Uitto 2014; Cline and Feldman 2016; Bootorabi et al. 2017). In recent years, larval zebrfish studies have shed light on the molecular causes and consequences of conditions characteristic of human skin diseases, including aber-
rant periderm development (Li et al. 2011a; de la Garza et al. 2013;
Liu et al. 2016), defective adhesion and blistering (Carney et al. 2007;
2010, Sonawane et al. 2009; Kim et al. 2010; Goonesinghe et al. 2012;
Postel et al. 2013), impaired wound healing (Niethammer et al. 2009;
Rieger and Sagasti 2011; Lisse et al. 2016), and dysregulation of epider-
cmal cell proliferation (Sonawane et al. 2005, 2009; Carney et al. 2007;
Webb et al. 2008; Dodd et al. 2009; Eisenhoffer et al. 2012; Brock et al. 2019). Our gene expression profiles provide a resource for identifying additional molecules that may contribute to these and other skin conditions.

**ACKNOWLEDGMENTS**

Funding provided by NIH R21EY024400, NIH R01GM122901, and NIH R01AR064582 to A; a CSUPERB New Investigator grant, the CSUDH Faculty Scholar Program, CSUDH RSCA grants, and a CSUDH Norris Summer grant to FW; Title V-PPOHA GWIE P031M14004 to MDLT and FW; NIH RISE R25GM62252 to EFM and JRG; NSF LSAMP HRD-1302873 to JRG; and the Jane Coffin Childs Memorial Fund to JPR. We thank the California State University Desert Studies Center for providing an excellent writing environment. We received support from the UCLA Broad Stem Cell Research Center Flow Cytometry Core Resource.

**LITERATURE CITED**

Benoit, C. R. A., E. Stanton, A. C. Tartarian, A. R. Motzer, D. M. McGaughey et al., 2018 Functional and phylogenetic characterization of noncana-
nonal vitamin B12-binding proteins in zebrfish suggests involvement in cobalamin transport. J. Biol. Chem. 293: 17606–17621. https://doi.org/10.1074/jbc.R118.005323

Brock, C. K., H. Manouchehri, R. Changizi, H. Barker, E. Palazzo et al., 2017 Zebrfish as a Model Organism for the Development of Drugs for Skin Cancer. Int. J. Mol. Sci. 18: 1550. https://doi.org/10.3390/ijms18071550

Brugga, H. H., and D. G. Homberger, 2009 Structure and functions of keratin proteins in simple, stratified, keratinized and cornified epithelia. J. Anat. 214: 516–559. https://doi.org/10.1111/j.1469-7580.2009.01066.x

Bussmann, J., and S. Schulte-Merker, 2011 Rapid BAC selection for tol2-mediated transgenesis in zebrfish. Development 138: 4327–4332. https://doi.org/10.1242/dev.068080

Byrne, C., M. Tainsky, and E. Fuchs, 1994 Programming gene expression in developing epidermis. Development 120: 2369–2383.

Conrad, M., K. Lemb, T. Schubert, and J. Markl, 1998 Biochemical identifica-
tion and tissue-specific expression patterns of keratins in the zebrfish Danio rerio. Cell Tissue Res. 293: 195–205. https://doi.org/10.1007/s004410051112

Coulombe, P. A., and C.-H. Lee, 2012 Defining keratin protein function in skin epithelia: epidermolysis bullosa simplex and its aftermath. J. Invest. Dermatol. 132: 763–775. https://doi.org/10.1038/jid.2011.450

Cui, C.-Y., M. Kunisada, D. Esibizione, S. I. Grivennikov, Y. Piao et al., 2007 Lymphotixin-beta regulates periderm differentiation during em-
byronic skin development. Hum. Mol. Genet. 16: 2583–2590. https://
doi.org/10.1093/hmg/ddm210

Curado, S., R. M. Anderson, B. Jungblut, J. Mumm, E. Schroeter et al., 2007 Conditional targeted cell ablation in zebrfish: a new tool for re-
generation studies. Dev. Dyn. 236: 1025–1035. https://doi.org/10.1002/
ddv.21100

Dapasquale, J. A., 2018 Actin Microridges: ACTIN MICRORIDGES IN EPITHELIUM. Anat. Rec. (Hoboken) 31: 81.

Dobin, A., C. A. Davis, F. Schlesinger, J. Drenkow, C. Zaleski et al., 2013 STAR: ultrafast universal RNA-seq aligner. Bioinformatics 29:
15–21. https://doi.org/10.1093/bioinformatics/bts635

Dodd, M. E., J. Hatzold, J. R. Mathias, K. B. Walters, D. A. Bennin et al., 2009 The ENTH domain protein Clnt1 is required for epidermal homeostasis in zebrfish. Development 136: 2591–2600. https://doi.org/10.1242/dev.038448

Eisenhoffer, G. T., P. D. Lofus, M. Yoshigi, H. Otsuna, C.-B. Chien et al., 2012 Crowding induces live cell extrusion to maintain homeostatic cell numbers in epidermis. Nature 484: 546–549. https://doi.org/10.1038/nature10999

Fuchs, E., and S. Raghavan, 2002 Getting under the skin of epidermal mor-
phogenesis. Nat. Rev. Genet. 3: 199–209. https://doi.org/10.1038/nrg758

de la Garza, G., J. R. Schleifarth, M. Dunnwald, A. Mankad, J. L. Weir-
ther et al., 2013 Interferon regulatory factor 6 promotes differen-
tiation of the periderm by activating expression of Grainyhead-like 3.
J. Invest. Dermatol. 133: 68–77. https://doi.org/10.1038/jid.2012.269

Gebauer, J. M., B. Kobbe, M. Paulsson, and R. Wageser, 2016 Structure, evolution and expression of collagen XXVIII: Lessons from the zebrfish. Matrix Biol. 49: 106–119. https://doi.org/10.1016/j.mabio.2015.07.001

Gonzales, K. A. U., and E. Fuchs, 2017 Skin and Its Regenerative Powers: An Alliance between Stem Cells and Their Niche. Dev. Cell 43: 387–401. https://doi.org/10.1016/j.devcel.2017.10.001

Goonesinghe, A., X.-M. Luan, A. Hurlstone, and D. Garrod, 2012 Desmosomal cadherins in zebrfish epiboly and gastrulation. BMC Dev. Biol. 12: 1. https://doi.org/10.1186/1471-213X-12-1

Guzman, A., J. L. Ramos-Balderas, S. Carrillo-Rosas, and E. Maldonado, 2013 A stem cell proliferation burst forms new layers of P63 ex-
pressing suprabasal cells during zebrfish postembryonic epidermal development. Biol. Open 2: 1179–1186. https://doi.org/10.1242/
bio.20136023

Hardman, M. J., L. Moore, M. W. Ferguson, and C. Byrne, 1999 Barrier formation in the human fetus is patterned. J. Invest. Dermatol. 113: 1106–1113. https://doi.org/10.1038/sj.jid.5527479.1999.00800.x

Herken, R., and U. Schultz-Ehrenburg, 1981 Autoradiographic investiga-
tions on the cell kinetics of epidermis and periderm of limb buds from mouse embryos in vitro. Br. J. Dermatol. 104: 277–284. https://doi.org/10.1111/j.1365-2133.1981.tb00949.x

Holbrook, K. A., and G. F. Olland, 1980 Regional development of the human epidermis in the first trimester embryo and the second trimester fetus (ages related to the timing of amniocentesis and fetal biopsy). J. Invest. Dermatol. 74: 161–168. https://doi.org/10.1111/ 1523-1747.ep1253062

Holbrook, K. A., and G. F. Olland, 1975 The fine structure of developing human epidermis: light, scanning, and transmission electron microscopy of the periderm. J. Invest. Dermatol. 65: 16–38. https://doi.org/10.1111/ 1523-1747.ep1259029

Hu, B., C. Zhang, K. Baawo, R. Qin, G. J. Cole et al., 2010 Zebrafish K5 promoter driven GFP expression as a transgenic system for oral research. Oral Oncol. 46: 31–37. https://doi.org/10.1016/j.oraloncology.2009.09.008
Lam, P.-Y., S. Mangos, J. M. Green, J. Reiser, and A. Huttonlocher, 2015 In vivo imaging and characterization of actin microridges. PLoS One 10: e0115639. https://doi.org/10.1371/journal.pone.0115639

Le Guellec, D., G. Morvan-Dubois, and J.-Y. Sire, 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: 217–231. https://doi.org/10.1515/ijdb.2004.217

Li, Q., M. Frank, M. Akiyama, H. Shimizu, S.-Y. Ho et al., 2011a Abca12-mediated lipid transport and Snap29-dependent trafficking of lamellar granules are crucial for epidermal morphogenesis in a zebrafish model of ichthyosis. Dis. Model. Mech. 4: 777–785. https://doi.org/10.1242/dmm.007146

Li, Q., M. Frank, C. I. Thisse, B. V. Thisse, and J. Uitto, 2011b Zebrafish: a model system to study heritable skin diseases. J. Invest. Dermatol. 131: 565–571. https://doi.org/10.1038/jid.2010.388

Linden, S. K., P. Sutton, N. G. Karlsson, V. Korolik, and M. A. McGuckin, 2008 Mucins in the mucosal barrier to infection. Mucosal Immunol. 1: 183–197. https://doi.org/10.1016/j.mucimm.2008.5

Lisse, T. S., B. L. King, and S. Rieger, 2016 Comparative transcriptomic analysis of nociceptive sensory neurites. eLife 5: e10345. https://doi.org/10.7554/eLife.20345

Love, M. I., W. Huber, and S. Anders, 2014 Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol. 15: 550. https://doi.org/10.1186/s13059-014-0550-8

Lu, H., M. Hesse, B. Peters, and T. M. Magin, 2005 Type II keratins precede type I keratins during early embryonic development. Eur. J. Cell Biol. 84: 709–718. https://doi.org/10.1016/j.ejcb.2005.04.001

Martin, M., 2011 Cutadapt removes adapter sequences from high-throughput sequencing reads. EMBnet.journal 17: 10–12. https://doi.org/10.14806/ej.17.1.200

Mazzalupo, S., and P. A. Coulombe, 2001 A reporter transgene based on a human keratin 6 gene promoter is specifically expressed in the periderm of mouse embryos. Mech. Dev. 100: 65–69. https://doi.org/10.1016/S0925-4773(00)00489-5

M’Bonek, V., and H. J. Merker, 1988 Development and morphology of the periderm of mouse embryos (days 9–12 of gestation). Acta Anat. (Basel) 133: 325–336. https://doi.org/10.11159/000146662

McGowan, K. M., and P. A. Coulombe, 1998 Onset of keratin 17 expression coincides with the definition of major epithelial lineages during skin development. J. Cell Biol. 143: 469–486. https://doi.org/10.1083/jcb.143.2.469

Morita, K., M. Furuse, Y. Yoshida, M. Itoh, H. Sasaki et al., 2002 Molecular architecture of tight junctions of periderm differs from that of the maculae occludentes of epidermis. J. Invest. Dermatol. 118: 1073–1079. https://doi.org/10.1046/j.1523-1747.2002.01774.x

Muroya, A., and T. Lechler, 2012 Polarity and stratification of the epidermis. Semin. Cell Dev. Biol. 23: 890–896. https://doi.org/10.1016/j.semcdb.2012.08.008

Nakajima, H., Y. Nakajima-Takagi, T. Tsuji, S.-I. Akiyama, T. Wakasa et al., 2011 Tissue-restricted expression of Nrt2 and its target genes in zebrafish with gene-specific variations in the induction profiles. PLoS One 6: e26884. Erratum 7: 10.1371/annotation.50e3e8af-3010-4c42-a130-70509a8867e. https://doi.org/10.1371/annotation.50e3e8af-3010-4c42-a130-70509a8867e

Niethammer, P., C. Grabher, A. T. Look, and T. J. Mitchison, 2009 A tissue-scale gradient of hydrogen peroxide mediates rapid wound detection in zebrafish. Nature 459: 996–999. https://doi.org/10.1038/nature08119

O’Brien, G. S., S. Rieger, F. Wang, G. A. Smolen, R. E. Gonzalez et al., 2012 Coordinate development of skin cells and cutaneous sensory axons in zebrafish. J. Comp. Neurol. 520: 816–831. https://doi.org/10.1002/cne.22797

Okano, J., U. Lichti, S. Mamiya, M. Aronova, G. Zhang et al., 2012 Increased retinoic acid levels through ablation of Cyp26b1 determine the processes of embryonic skin barrier formation and peridermal development. J. Cell Sci. 125: 1827–1836. https://doi.org/10.1242/jcs.101550

Patio, R., G. Duggal, M. I. Love, R. A. Irizarry, and C. Kingsford, 2017 Salmon provides fast and bias-aware quantification of transcript expression. Nat. Methods 14: 417–419. https://doi.org/10.1038/nmeth.4197

Pimentel, H., N. L. Bray, S. Puente, P. Melsted, and I. Pachter, 2017 Differential analysis of RNA-seq incorporating quantification uncertainty. Nat. Methods 14: 687–690. https://doi.org/10.1038/nmeth.4324

Pinto, C. S., A. Khandekar, B. Rajasekaran, P. Kiesel, G. Pigino et al., 2019 Microridges are apical epithelial projections formed of F-actin networks that organize the glycan layer. Sci. Rep. 9:12191. https://doi.org/10.1038/s41598-019-40194-2

Postel, R., C. Margadant, B. Fischer, M. Kreft, H. Janssen et al., 2013 Kindlin-1 mutant zebrafish as an in vivo model system to study adhesion mechanisms in the epidermis. J. Invest. Dermatol. 133: 2180–2190. https://doi.org/10.1038/jid.2013.154

Rasmussen, J. P., G. S. Sack, S. M. Martin, and A. Sagasti, 2015 Vertebrate epidermal cells are broad-specificity phagocytes that clear sensory axon debris. J. Neurosci. 35: 559–570. https://doi.org/10.1523/JNEUROSCI.3613-14.2015

Richardson, R. J., N. L. Hammond, P. A. Coulombe, C. Salaranta, H. O. Nousiainen et al., 2014 Periderm prevents pathological epithelial adhesions during embryogenesis. J. Clin. Invest. 124: 3891–3900. https://doi.org/10.1172/JCI71946

Rieger, S., and A. Sagasti, 2011 Hydrogen peroxide promotes injury-induced peripheral sensory axon regeneration in the zebrafish skin. PLoS Biol. 9: e1000621. https://doi.org/10.1371/journal.pbio.1000621

Schaffeld, M., and J. Schultes, 2006 Genes coding for intermediate filament proteins closely related to the hagfish “thread keratins (TK)” alpha and
gamma also exist in lampreys, teleosts, and amphibians. Exp. Cell Res. 312: 1447–1462. https://doi.org/10.1016/j.yexcr.2006.01.005

Smart, I. H., 1970 Variation in the plane of cell cleavage during the process of stratification in the mouse epidermis. Br. J. Dermatol. 82: 276–282. https://doi.org/10.1111/j.1365-2133.1970.tb12437.x

Sonawane, M., Y. Carpio, R. Geisler, H. Schwarz, H.-M. Maischein et al., 2005 Zebrafish penner/lethal giant larvae 2 functions in hemidesmosome formation, maintenance of cellular morphology and growth regulation in the developing basal epidermis. Development 132: 3255–3265. https://doi.org/10.1242/dev.01904

Sonawane, M., H. Martin-Maischein, H. Schwarz, and C. Nüsslein-Volhard, 2009 Lgl2 and E-cadherin act antagonistically to regulate hemidesmosome formation during epidermal development in zebrafish. Development 136: 1231–1240. https://doi.org/10.1242/dev.022508

Suster, M. L., G. Abe, A. Schouw, and K. Kawakami, 2011 Transposon-mediated BAC transgenesis in zebrafish. Nat. Protoc. 6: 1998–2021. https://doi.org/10.1038/nprot.2011.416

Thise, C., and B. Thise, 2008 High-resolution in situ hybridization to whole-mount zebrafish embryos. Nat. Protoc. 3: 59–69. https://doi.org/10.1038/nprot.2007.514

Vanbokhoven, H., G. Melino, E. Candi, and W. Declercq, 2011 p63, a story of mice and men. J. Invest. Dermatol. 131: 1196–1207. https://doi.org/10.1038/jid.2011.84

Wang, F., S. N. Wolfson, A. Gharib, and A. Sagasti, 2012 LAR receptor tyrosine phosphatases and HSPGs guide peripheral sensory axons to the skin. Curr. Biol. 22: 373–382. https://doi.org/10.1016/j.cub.2012.01.040

Webb, A. E., W. Driever, and D. Kimelman, 2008 Psoriasis regulates epidermal development in zebrafish. Dev. Dyn. 237: 1153–1164. https://doi.org/10.1002/dvdy.21509

White, A. C., and W. E. Lowry, 2015 Refining the role for adult stem cells as cancer cells of origin. Trends Cell Biol. 25: 11–20. https://doi.org/10.1016/j.tcb.2014.08.008

Wolf, J., 1968a Curve of development and disappearance of peridermal phase of human embryonic skin demonstrated by the replica method. Folia Morphol. 16: 36–42.

Wolf, J., 1967 Structure and function of periderm. I. Superficial structure of the peridermal epithelium. Folia Morphol. 15: 296–305.

Wolf, J., 1968b The replacement of periderm by epidermis. Folia Morphol. 16: 24–35.

Yoshida, M., Y. Shimono, H. Togashi, K. Matsuzaki, J. Miyoshi et al., 2012 Periderm cells covering palatal shelves have tight junctions and their desquamation reduces the polarity of palatal shelf epithelial cells in palatogenesis. Genes Cells 17: 455–472. https://doi.org/10.1111/j.1365-2443.2012.01601.x

Communicating editor: D. Grunwald