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Distinct tooth regeneration systems deploy a conserved battery of genes



Tyler A. Square^{*}, Shivani Sundaram, Emma J. Mackey and Craig T. Miller^{*}

Abstract

Background: Vertebrate teeth exhibit a wide range of regenerative systems. Many species, including most mammals, reptiles, and amphibians, form replacement teeth at a histologically distinct location called the successional dental lamina, while other species do not employ such a system. Notably, a 'lamina-less' tooth replacement condition is found in a paraphyletic array of ray-finned fishes, such as stickleback, trout, cod, medaka, and bichir. Furthermore, the position, renewal potential, and latency times appear to vary drastically across different vertebrate tooth regeneration systems. The progenitor cells underlying tooth regeneration thus present highly divergent arrangements and potentials. Given the spectrum of regeneration systems present in vertebrates, it is unclear if morphologically divergent tooth regeneration systems deploy an overlapping battery of genes in their naïve dental tissues.

Results: In the present work, we aimed to determine whether or not tooth progenitor epithelia could be composed of a conserved cell type between vertebrate dentitions with divergent regeneration systems. To address this question, we compared the pharyngeal tooth regeneration processes in two ray-finned fishes: zebrafish (*Danio rerio*) and three-spine stickleback (*Gasterosteus aculeatus*). These two teleost species diverged approximately 250 million years ago and demonstrate some stark differences in dental morphology and regeneration. Here, we find that the naïve successional dental lamina in zebrafish expresses a battery of nine genes (*bmpr1aa, bmp6, cd34, gli1, igfbp5a, lgr4, lgr6, nfatc1,* and *pitx2*), while active Wnt signaling and *Lef1* expression occur during early morphogenesis stages of tooth development. We also find that, despite the absence of a histologically distinct successional dental lamina in stickleback tooth fields, the same battery of nine genes (*Bmpr1a, Bmp6, CD34, Gli1, lgfbp5a, Lgr4, Lgr6, Nfatc1,* and *Pitx2*) are expressed in the basalmost endodermal cell layer, which is the region most closely associated with replacement tooth germs. Like zebrafish, stickleback replacement tooth germs additionally express *Lef1* and exhibit active Wnt signaling. Thus, two fish systems that either have an organized successional dental lamina (zebrafish) or lack a morphologically distinct successional dental lamina (sticklebacks) deploy similar genetic programs during tooth regeneration.

Conclusions: We propose that the expression domains described here delineate a highly conserved "successional dental epithelium" (SDE). Furthermore, a set of orthologous genes is known to mark hair follicle epithelial stem cells in mice, suggesting that regenerative systems in other epithelial appendages may utilize a related epithelial progenitor cell type, despite the highly derived nature of the resulting functional organs.

Keywords: Successional dental lamina, Tooth regeneration, Odontode, Epithelial appendage

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Background

Vertebrate teeth arose shortly after vertebrates themselves and have since greatly diversified. Vertebrate "odontodes", which include teeth and denticles, are defined by a shared basic morphology: a dentine core derived from mesenchymal cells, and an enamel or enameloid tip sheath contributed (at least in part) by epithelial cells [1-4]. The first clear evidence of dentine and enameloid-bearing structures arose at least ~ 420 million years ago in fossils of placoderms, though an even earlier origin in conodonts has been debated [5-8]. Support for the homology of odontodes comes from developmental studies that document similar histogenesis, morphological development, and shared gene expression patterns between teeth and other odontodes across large phylogenetic distances within jawed vertebrates [9–12]. Despite these deeply conserved, fundamental aspects of tooth and odontode development, teeth have radiated vastly in their arrangement [13], shape [14], size [15], placement within the body plan [16], and the regeneration system they employ [17–21]. Thus, some aspects of tooth development are staunchly conserved, while others are wildly plastic in evolution. This variation in the conservation of different aspects of tooth development presents an interesting case study in the evolution of development: what genetic signatures, if any, are common to disparate dental morphologies? Here, we assess two divergent tooth regeneration strategies found in two different ray-finned fishes: zebrafish (*Danio rerio*) and threespine stickleback (*Gasterosteus aculeatus*).

The morphogenesis of individual tooth organs has been well-studied across living jawed vertebrates [9, 22]. Teeth form like other epithelial appendages, demonstrating an intricate epithelial-mesenchymal interaction and coordinated cell motions during their early development (Fig. 1) [23]. Primary teeth first appear as a placode (thickened epithelium) underlain by a mesenchymal condensation, followed by invagination, cap formation, and eventually deposition of dentine and enamel and/or enameloid, which are composed of highly concentrated hydroxyapatite (a hard, crystalline calcium phosphate mineral that gives bones their characteristic hardness) [3, 24]. Some vertebrates, such as mice, do not replace entire teeth during their lives. By contrast, in most known extant vertebrate dental systems, once a tooth position is established, tooth replacement via regeneration can occur anywhere between one (e.g., most mammalian teeth) to potentially one hundred times or more (e.g., some sharks) at a given tooth position [25-30]. In many vertebrates with tooth replacement, this process initiates at a site called the successional dental lamina (SDL). The SDL can be identified histologically as a deep extension of epithelium connected directly to the outer dental epithelium of a predecessor tooth [9, 31]. When present, during tooth replacement the SDL first thickens, similar to the placode stage of primary teeth [9]. Subsequently, this thickened

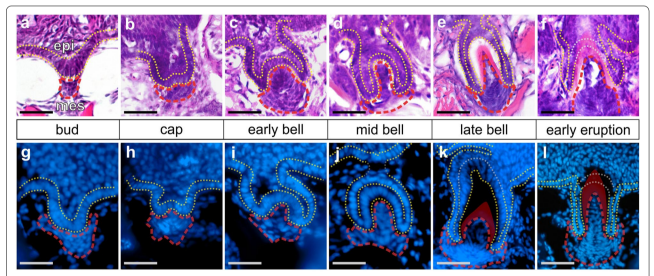


Fig. 1 Stickleback pharyngeal tooth morphogenesis. All images show sagittal or transverse sections from sticklebacks (20–40 mm standard length). The basalmost layer of epithelium is outlined with yellow dashed lines in all panels, the mesenchyme is outlined in red. During late bell stages (e, k), the epithelium towards the apex of the forming tooth becomes thickened, no longer presents a clear bilayer, and will eventually be punctured during tooth eruption (gray outlines). a-f Staging series of hematoxylin and eosin-stained transverse sections through stickleback pharyngeal tooth germs at the stages indicated. Epithelium ("epi") and mesenchyme ("mes") are labeled in a. g-l Staging series of DAPI-stained sagittal sections through stickleback pharyngeal tooth germs at the stages indicated. Gray portions of dotted epithelium lines in J indicate the epithelium/tooth connection that was present on an adjacent section. Bone matrix is false-colored red in k and l. Scale bars = 20 μm

epithelium undergoes the stereotypical tooth differentiation process outlined above (depicted in Fig. 1), and at all stages is tightly associated with underlying dental mesenchyme. The SDL was recently shown to be a key tissue in tooth regeneration: mouse molars, which do not naturally undergo regeneration, can be prompted to do so by stabilizing Wnt signaling in the rudimentary mouse SDL [32]. In most cases, a replacement tooth germ eventually prompts osteoclast activity to enzymatically dissociate the base of the predecessor tooth and remodel the underlying bone of attachment (if present) either before or during early eruption [33–37]. This osteoclast activity facilitates the shedding of the predecessor and makes space for the replacement tooth to finish mineralization and eruption, thus completing the regenerative process.

While many vertebrate tooth regeneration systems utilize an SDL, the arrangement of primary and replacement teeth is variable, both in modern species [9, 12, 17, 31, 38–44], as well as stem group jawed vertebrates [45, 46]. In some species (e.g., sharks), teeth are replaced in families from permanent, conspicuous regions in the oral jaw [12, 20]. By contrast, in other species, teeth are replaced from a relatively inconspicuous SDL that forms transiently (as in zebrafish) [42]. Some fishes such as bichir and salmonids present an even more abbreviated process, wherein the outer dental epithelium of the predecessor tooth directly relocalizes to a deeper region on the lingual side of the tooth, undergoes thickening, and eventually tooth differentiation, effectively 'skipping' the SDL phase [18, 47–49]. Cichlid oral teeth grossly undergo the same process, though interestingly their oral replacement teeth are derived from the opposite (labial) side of the predecessor [10, 50]. The patterning of tooth replacement can even vary throughout ontogeny of the same species: medaka fish initially have disorganized pharyngeal dentitions without visible tooth families, but later in life the teeth organize into highly patterned pharyngeal tooth plates with clear tooth families arranged in rows [38, 51]. Despite these clear tooth families, no discrete dental lamina was observed in medaka [38]. Together, these data show that much flexibility exists in how tooth progenitors can be arranged across dental regeneration systems [18, 21].

Despite high variation in the size, shape, and position of vertebrate teeth, gene expression and functional studies on tooth differentiation across vertebrates has revealed deeply conserved molecular processes at play [10, 21]. However, given the phylogenetic lability and morphological disparities of tooth regeneration strategies between some groups [45], it is unclear whether or not this process is homologous and uses shared cell types across vertebrates. Recent evidence suggests that not just teeth, but also other epithelial appendages (EAs) such as scales, feathers, and hair share some aspects of development, also raising the possibility of far-reaching EA homology among these organs [23, 52–59].

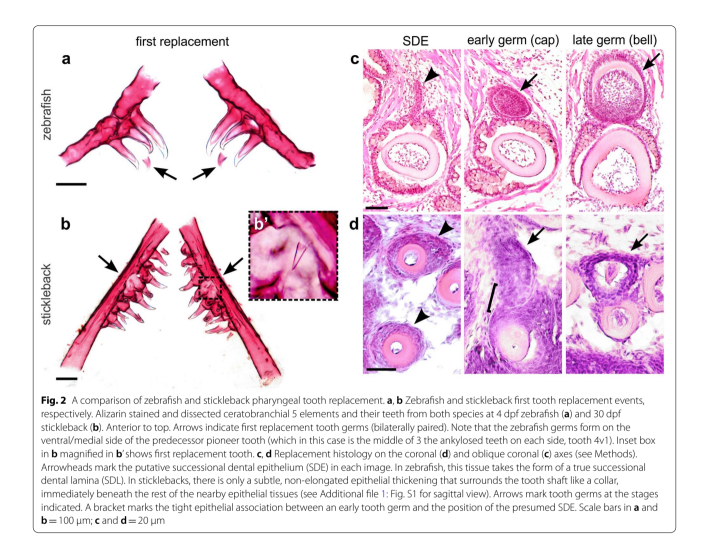
To further elucidate the developmental program underlying tooth regeneration, we compared the histology and gene expression patterns associated with the regenerative process in the pharyngeal teeth of two teleost fishes: zebrafish and threespine stickleback. These two species share a common ancestor that lived approximately 250 million years ago [60] and appear to demonstrate some stark differences in their dentitions. Sticklebacks are well documented as demonstrating differences in tooth number between populations, particularly between hightoothed freshwater populations and low-toothed marine populations [61-65]. These tooth number differences have a strong genetic basis, contributing to a significant difference in adult tooth number between certain populations, which is especially pronounced on the ventral pharyngeal jaw [63, 64]. We find that, unlike other vertebrates such as zebrafish, the threespine stickleback does not exhibit an SDL, nor does it regenerate teeth via the immediate relocalization of the deep outer dental epithelium of a predecessor tooth (as in salmonids and bichir). Furthermore, stickleback teeth appear to occasionally diverge from the common 1-for-1 mode of tooth replacement: some tooth germs appear to prompt osteoclast activity and shedding of more than one erupted tooth in concert. Despite these differences in modes of tooth regeneration, we find a suite of genes expressed in common between naïve dental epithelial cells in both fish species, suggesting that a conserved epithelial progenitor cell type underlies tooth regeneration, here referred to as the "successional dental epithelium" (SDE). We propose the use of this term to encompass not just the progenitor cells found in the histologically distinct successional dental lamina (SDL) of animals such as zebrafish, sharks, and tetrapods (where present), but also in less-pronounced epithelial regions in fishes like sticklebacks, which still undergo constant tooth regeneration across their tooth fields. Intriguingly, the genes we find localized to the SDE of both zebrafish and sticklebacks also mirror a cassette of gene expression that is well documented and functionally relevant in hair follicle epithelial stem cells (in the "bulge") [66, 67], suggesting that such a genetic module may be common across not just teeth, but also other vertebrate epithelial appendages.

Results

A brief review and comparison of the gross morphologies of stickleback and zebrafish dentitions

Tooth field position and arrangement

Zebrafish and stickleback teeth are grossly similar in tooth shape and structure (Fig. 2a and b); both species'



teeth are unicuspid and mostly conical, with a slight bend or hook towards the posterior side. However, these two species differ drastically in tooth arrangement within the body plan. Zebrafish, like other members of Cypriniformes, lack oral teeth, and possess only a single (paired) tooth field found ventrally on the last branchial arch (on ceratobranchial 5) in the pharynx (see Fig. 2a, referred to here as the ventral tooth plate or VTP, also called the lower pharyngeal jaw or LPJ in some species). Sticklebacks, like most vertebrates, have oral teeth, which are located on their premaxilla and dentary bones. In the pharynx, sticklebacks also have a single paired VTP on ceratobranchial 5 (see Fig. 2b), as well as two paired dorsal tooth plates (DTPs) [68].

Tooth number variation

Sticklebacks and zebrafish also have major differences in tooth number. In lab-reared zebrafish, each VTP typically hosts 11 tooth positions [69]. As in other cyprinids, this number is known to vary occasionally in zebrafish, occasionally observed as missing or bearing an extra tooth position [70]. Since replacement tooth mineral deposition occurs prior to the shedding of a predecessor tooth during the tooth regeneration process, an adult zebrafish typically hosts 15–17 mineralized teeth per VTP at any given point in time (i.e., there are typically 4–6 late bell stage replacement tooth germs per VTP that have not yet dislodged their predecessor). Given this relatively consistent dental arcade, zebrafish tooth positions have been named based on their location [69].

Threespine sticklebacks, on the other hand, can have up to $\sim 200-300$ + total teeth once fully grown, thus displaying both an order of magnitude more teeth than zebrafish, as well as higher within-species variation in adult tooth number. Tooth number in sticklebacks has a strong genetic basis, and varies significantly by population, with freshwater localities generally possessing more teeth than their marine counterparts [61, 63–65].

Early tooth replacement events

Sticklebacks and zebrafish both appear to replace their earliest primary teeth in a similar order, with those teeth that differentiate first also being replaced first [68, 69]. Interestingly, the tooth germs responsible for these early replacement events are essentially locationally opposite between the zebrafish and stickleback VTPs (arrows in Fig. 2a and b): zebrafish replacement tooth germs form on the medial/ventral side of each erupted tooth, towards the midline [69], while the first stickleback replacement tooth germs form on the lateral edge of the tooth plate, opposite the midline [68]. In zebrafish, this stereotypical polarized replacement direction (Fig. 2a) remains into adult stages. Conversely, as they mature, stickleback pharyngeal tooth fields do not appear to strictly adhere to the 'lateral to medial' replacement direction found in their earliest replacement events, which we detail below.

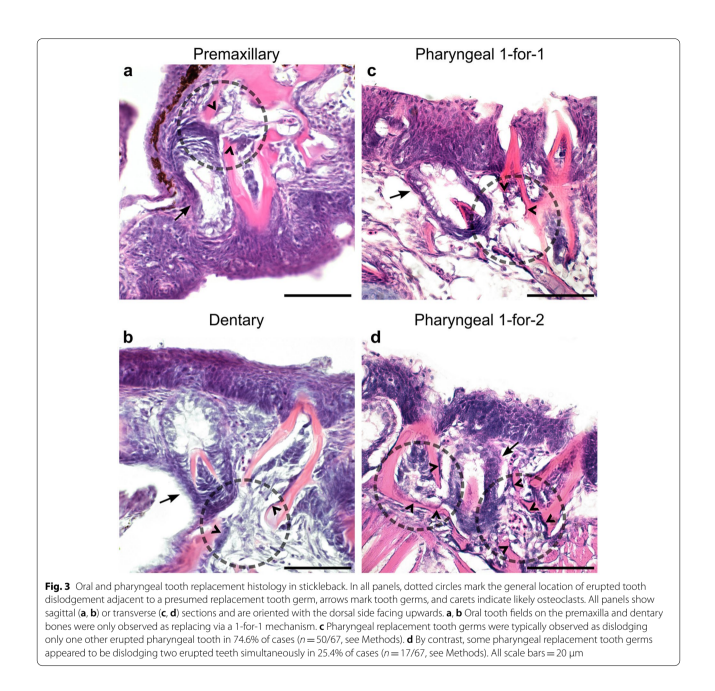
Replacement tooth histology

The histology of zebrafish tooth regeneration has been described previously in detail [42, 69]. Zebrafish tooth regeneration initiates from an SDL. In zebrafish, this region forms transiently, as an outpocketing on the ventro-medial side of each tooth, first visible as a thin projection of naïve epithelial cells stemming from the outer dental epithelium of a predecessor tooth (arrowhead in Fig. 2c) [42], protruding from the mineralized tooth and its associated epithelia deep in the tooth field. Subsequently, the SDL expands, forming a replacement tooth germ (arrows) while maintaining a close epithelial association with the predecessor tooth socket.

Stickleback pharyngeal replacement teeth arise from a more superficial region of the dentition than replacement teeth in zebrafish. After morphogenesis (Fig. 1), once ankylosis and eruption are complete, stickleback teeth do not retain a histologically distinct inner and outer dental epithelium (Additional file 1: Fig. S1). Instead, mature stickleback pharyngeal teeth retain a shallow "collar" of epithelial cells surrounding the site of eruption (arrowheads in Fig. 2d and Additional file 1: Fig. S1) that does not appear to be arranged in a clear bilayer. This collar of cells is contiguous with the basalmost epithelial sheet overlying the tooth field, juxtaposed to the pharyngeal epithelium (towards the lumen) and the basement membrane overlying mesenchymal cells (away from the lumen). It is from this collar of cells that the epithelial components of both oral and pharyngeal replacement teeth appear to be derived in sticklebacks (arrows in Fig. 2d), suggested by the tight epithelial association between early germs and the position of the putative SDE (bracket in Fig. 2d). Thus, a histologically distinct SDL is not present during oral or pharyngeal tooth regeneration in sticklebacks, a condition also reported in trout [49], the oral teeth of salmon [48], bichir [18], and cod [71], as well as pharyngeal teeth in medaka [38].

Stickleback pharyngeal teeth do not exhibit a tightly regulated 1-for-1 replacement scheme

Unlike zebrafish, stickleback tooth fields do not maintain any obvious arrangement of individual tooth positions across the entirety of each tooth field into adulthood, even within single populations. This observation initially led us to hypothesize that the stickleback tooth replacement process may introduce some spatial stochasticity of tooth position and spacing, similar to the condition in cod oral teeth, where their adult arrangement appears "haphazard" [71]. We thus sought to test the hypothesis that stickleback tooth replacement can diverge from a strict 1-for-1 tooth replacement system. We did this by analyzing H&E-stained serial sections cut in the sagittal, transverse, and coronal planes through pharyngeal tooth fields from subadult and adult fish (25-40 mm standard length; see Additional file 1: Fig. S1 for section plane orientation). Using these preparations, we identified every putative pharyngeal and oral replacement tooth germ that was in a mid- to late bell stage (the stages that coincide with bone remodeling during the regenerative cycle), allowing us to observe how many and which teeth were being actively dissociated during the replacement process. Of the few oral tooth germs we observed (n=8)total observed from two separate fish) we found that stickleback oral replacement teeth formed at the labial side of each presumed predecessor tooth, and were not observed to be clearly diverging from a 1-for-1 replacement scheme (n=0/8 tooth germs) (Fig. 3a and b). In the pharynx, most stickleback replacement tooth germs appeared to be anchoring themselves solely in the place of a single presumed predecessor tooth, which was usually the tooth to which they showed the nearest epithelial association (Fig. 3c). However, in ~ 25% of cases (17/67)tooth germs) a single replacement tooth was found to be abutting two erupted teeth that were both displaying interrupted mineralization and apparent concomitant osteoclast activity at their bases (Fig. 3d). We interpret these to be possible "1-for-2" tooth replacement events in the stickleback pharynx. These observations suggest that stickleback pharyngeal teeth deviate from a strict 1-for-1 mode of tooth replacement up to 25% of the time.



Gene expression in stickleback and zebrafish tooth germs and progenitors

Wnt signaling activation in zebrafish and stickleback replacement tooth germs

The role of Wnt signaling in tooth morphogenesis and differentiation has been well-established in mouse and zebrafish [72–76], and indicted in multiple human tooth phenotypes [77–84]. Using in situ hybridizations (ISHs) on thin sections through zebrafish and stickleback tooth plates, we compared the expression of *Wnt10a* and *Lef1* in zebrafish and sticklebacks (Fig. 4a–d). We found highly

conserved patterns of gene expression in tooth germs for both genes, with the expression of both *Wnt10a* and *Lef1* upregulated in the future inner dental epithelia (especially at the central epithelial cells of the placode, black arrows in Fig. 4a–d) and early condensing mesenchyme of bud-stage tooth germs (white arrows in Fig. 4a–d).

One major output of Wnt signaling is upregulated transcriptional activity via β -catenin and TCF/Lef transcription factors [85]. A synthetic TCF/Lef reporter construct [86] was found to be expressed in a small number of cells in the young primary teeth of zebrafish, but expression

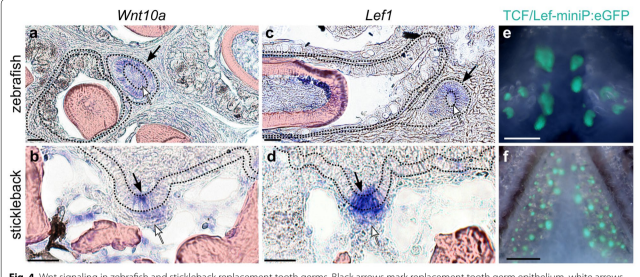


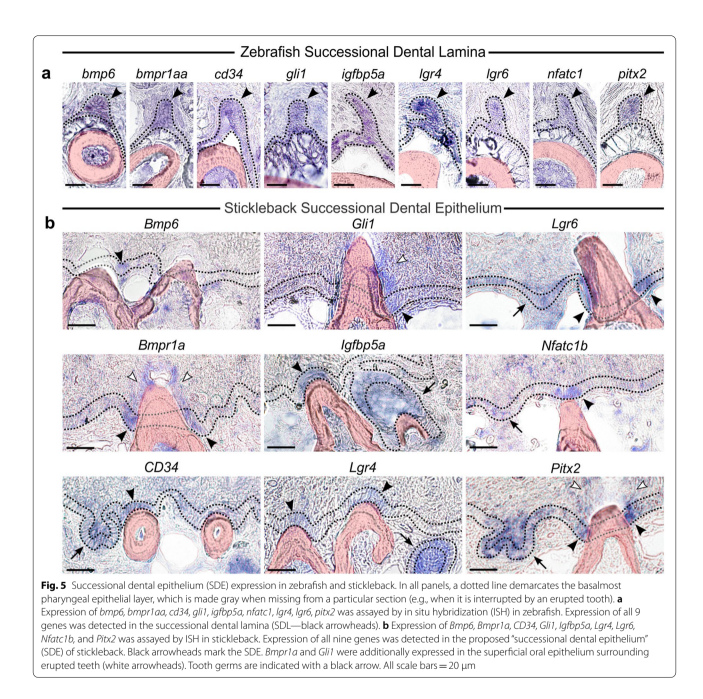
Fig. 4 Wht signaling in zebrafish and stickleback replacement tooth germs. Black arrows mark replacement tooth germ epithelium, white arrows mark replacement tooth germ mesenchyme. **a** *wnt10a* expression in a cap-stage zebrafish replacement tooth germ. **b** *Wnt10a* expression in a bud-stage stickleback replacement tooth germ. **b** *Wnt10a* expression in a cap-stage zebrafish replacement tooth germ. **b** *Wnt10a* expression in a cap-stage zebrafish replacement tooth germ. **b** *Wnt10a* expression in a cap-stage zebrafish replacement tooth germ. **d** *Lef1* expression in a bud-stage stickleback replacement tooth germ. **d** *Lef1* expression in a bud-stage stickleback replacement tooth germ. **e**, **f** TCF/Lef synthetic Wnt activity reporter is active in adult replacement tooth germs. Scale bars in $\mathbf{a} - \mathbf{d} = 20 \ \mu\text{m}$; **e** and $\mathbf{f} = 500 \ \mu\text{m}$

was not detected in the first replacement tooth [76]. This result raised the possibility of a Wnt signaling disparity between primary teeth and replacement teeth. The construct used in these assays employed a destabilized enhanced GFP (dGFP) coding region, which adds temporal resolution to the reporter construct output by increasing the degradation rate of eGFP. To address this potential primary vs. replacement tooth difference, we first compared the activity of the previously tested TCF/ Lef-miniP:dGFP reporter construct [86] in zebrafish and stickleback adults by generating novel transgenic lines in each species. We confirmed previous observations by Shim et al. (2019) that post-larval zebrafish teeth do not display detectable dGFP fluorescence (n = 4/4 juvenile zebrafish). Similarly, we also did not detect dGFP by fluorescence microscopy in stickleback teeth at any stage. Given our in situ data for Wnt10a and Lef1 expression in juvenile and adult fish (Fig. 4a-d), we suspected that downstream TCF/Lef transcriptional activity was also occurring in replacement tooth germs, but perhaps at a slightly reduced relative intensity and/or with faster turnover or signal dilution compared to the earliest zebrafish tooth germs. Thus, we re-stabilized the dGFP reporter cassette by removing the destabilization signal (see Methods) and again derived stable transgenic lines in zebrafish and sticklebacks. This change from dGFP to eGFP allowed the new TCF/Lef-miniP:eGFP cassette to be robustly detected in all developing young tooth germs in adult fish, especially tooth epithelium, and throughout the entire process of tooth morphogenesis (Fig. 3e and f). These data suggest that active Wnt signals are indeed transduced in replacement teeth of all ages in zebrafish and sticklebacks.

Expression in naïve epithelial tissues

As outlined above, sticklebacks lack a histologically defined SDL (per [31]) in their pharyngeal tooth fields. Despite this, sticklebacks must still possess epithelial replacement tooth progenitors in the pharynx, given that pharyngeal replacement teeth form in part from the endoderm adjacent to erupted teeth (Fig. 1). Here we refer to these putative epithelial progenitors collectively as the "successional dental epithelium" (SDE). We use this general term to encompass both the cell types found in permanent or transient SDLs of other vertebrates, as well as the putative epithelial tooth progenitors of less-organized systems such as stickleback pharyngeal teeth. Given our prediction regarding the location of the stickleback SDE based on histology data (arrowheads in Fig. 2d and Additional file 1: Fig. S1), we sought to assay detailed gene expression patterns in this tissue.

Previous expression assays revealed that β -catenin, Plakoglobin, and E-Cadherin are expressed in the zebrafish SDL [87, 88]. We aimed to bolster our understanding of gene expression in naïve dental epithelia by determining whether the putative stickleback SDE expresses gene orthologs that mark the zebrafish SDL. Thus, we first sought to identify a group of genes that were expressed



in the putative stickleback SDE and subsequently asked if these genes also mark the zebrafish SDL by performing (ISH) on thin sections through tooth plates of both fish species (Fig. 5). To these ends, we selected twelve "SDE candidate gene" groups based on previously published data supporting expression or function in tooth or hair regeneration, namely in the SDL or hair follicle stem cells: *Bambi* [67], *Bmi1* [50, 89], *Bmp6* [62, 90], *Bmpr1a* [91], *CD34* [92–97], *Dkk3b* [41], *Gli1* [50, 66, 98–100], *Igfbp5* [41, 50, 96, 99], *Lgr4* [50, 101], *Lgr6* [101], *Nfatc1* [95, 102], and *Pitx2* [11, 12, 47, 50, 99, 103, 104].

In stickleback, 10 of these 12 gene groups were found to have at least one ortholog gene expressed in the putative SDE, i.e., the basalmost endodermal epithelium overlying pharyngeal tooth fields, specifically marking the deep "collar" epithelial cells immediately juxtaposed to some, but not all erupted teeth, occasionally on flanking contralateral sides of a tooth (black arrowheads in Fig. 5b; see Additional file 2: Fig. S2 for DAPI counterstain of nuclei

of selected genes). Those genes we detected in the putative stickleback SDE were: Bmp6, Bmpr1a, CD34, Dkk3b, Gli1, Igfbp5a, Igfbp5b, Lgr4, Lgr6, Nfatc1b, and Pitx2 (black arrowheads in Fig. 5b and Additional file 3: Fig. S3). Of these 11 genes, Dkk3b and Igfbp5b were the least specific SDE markers, strongly marking many or most epithelial and mesenchymal cells in surrounding tissues (Additional file 3: Fig. S3), whereas the other nine genes exhibited more restricted patterns of expression (Fig. 5b). Importantly, these nine gene expression patterns in sticklebacks did not appear to collectively delineate any other cell type in the pharynx, other than early tooth germ epithelium (arrows in Fig. 5b), suggesting that this proposed "SDE genetic battery" is not common to other cell types or tissues in the pharynx. Five of these nine genes' expression patterns appeared locally restricted to the SDE (Bmp6, Gli1, Igfbp5a, Lgr4, and Nfatc1b), while the rest were found in more expansive stretches of the basalmost epithelium (CD34, Bmpr1a, Lgr6, and Pitx2). Three of these genes (Bmpr1a, Gli1, and Pitx2) had consistent detectable expression in the superficial pharyngeal epithelium surrounding erupted teeth (white arrowheads in Fig. 5). Those candidate genes not detected in the stickleback SDE were *Bmi1a*, which we did not detect in any tissue, and Bambia, which was expressed only in epithelium and mesenchyme of differentiating tooth germs, and mesenchyme of erupted teeth (Additional file 3: Fig. S3).

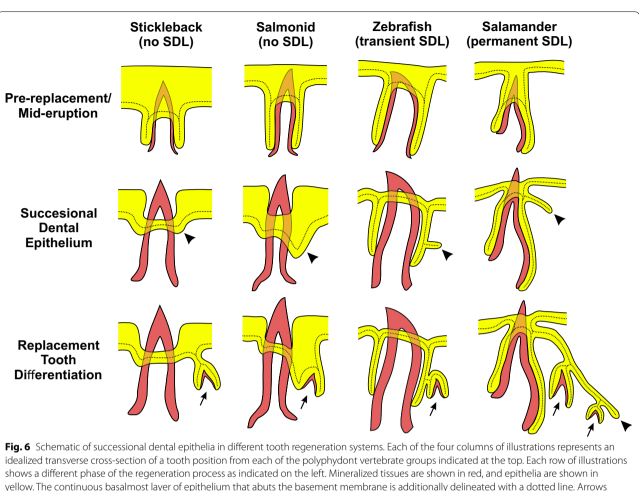
In zebrafish, we performed ISH for nearest-neighbor orthologs of the nine "SDE genetic battery" genes we observed as most specifically marking the putative stickleback SDE (bmp6, bmpr1aa, cd34, gli1, igfbp5a, *nfatc1, lgr4, lgr6,* and *pitx2*). All nine of these genes were expressed in the naïve, transiently forming SDL during its earliest stages, when it consists only of a single epithelial bilayer (Fig. 5a). The naïve SDL and its presumed daughter cells in young tooth germs were the only observed pharyngeal cell types exhibiting expression of all nine of these genes (presumably due to the maintenance of this genetic battery, consistent with our observations in sticklebacks). We did not observe *igfbp5b* expression in the zebrafish SDL, and instead detected it in a subset of dental mesenchyme (Additional file 4: Fig. S4). We did not assay bambia, bmi1a, or dkk3b expression in zebrafish.

Dental mesenchymal expression in differentiating and erupted zebrafish and stickleback odontoblasts was also observed for eight of the nine "SDE marker" genes, to the exclusion of *Pitx2*, which was not detected in pharyngeal dental mesenchyme (Additional file 5: Fig. S5). Zebrafish mesenchymal expression of these eight genes appeared widespread, strongly marking most or all odontoblasts in tooth germs and erupted teeth, with the exception of *Lgr6* which did not strongly mark mature tooth mesenchyme. Stickleback teeth showed slightly more dynamic mesenchymal expression of these genes: *Bmp6* sparsely marked odontoblasts, while *Igfbp5a* and *Lgr4* only marked younger tooth germs (Additional file 5: Fig. S5).

Discussion

Successional dental epithelia

Across vertebrates, tooth replacement systems display variable arrangements, numbers of tooth replacement cycles, and levels of anatomical conspicuousness [9]. In some vertebrates, tooth replacement is morphologically well-defined and gives rise to many teeth per tooth position, while in other vertebrates, tooth replacement events are morphologically ill-defined (e.g., they lack an SDL), and/or temporally transient, in some cases only giving rise to a single tooth (e.g., humans and most other mammals that have only one wave of tooth replacement, a.k.a. diphyodonts) [9, 18, 42, 105]. Here we asked if two vertebrates with histologically distinct tooth regeneration systems exhibit similar localizations of orthologous gene expression consistent with the presence of a conserved tooth epithelial progenitor cell type. We hypothesized that, despite the morphological differences found across regenerative systems, vertebrate teeth might deploy a homologous cell type responsible for generating the epithelial portion of a replacement tooth, but that this cell type may not always be found sequestered within a deep dental lamina. Here we show that zebrafish and sticklebacks, which are around 250 million years diverged [60], express a conserved battery of nine genes in their newly emerging replacement tooth epithelia. We propose this battery of gene expression labels a homologous, conserved dental cell type: the successional dental epithelium (SDE). Most of these nine genes have required functions in vertebrate dentitions, including Bmp6, Bmpr1a, Igfbp5, Lgr4, Nfatc1, and Pitx2 [62, 91, 106–110], as well as other epithelial appendages (discussed below). Together, these data support a model wherein the successional dental epithelia (SDE) of these two species share a common evolutionary developmental history, despite the zebrafish SDE manifesting as a true histological SDL, while the stickleback SDE is histologically minute and is not arranged as an SDL (arrowheads in Fig. 2d and Additional file 1: Fig. S1). Overall, these data suggest a common cell type (the SDE) underlies tooth regeneration in divergent tooth regeneration systems, regardless of the presence of a morphologically obvious lamina or SDL (Fig. 6). We propose that SDE cells have thus retained a distinct cellular identity despite evolved changes in conformation, such as whether or not they participate in SDL formation. More work in other vertebrates will reveal the level of gene expression domain conservation across the SDE of different vertebrate groups.



yellow. The continuous basalmost layer of epithelium that abuts the basement membrane is additionally delineated with a dotted line. Arrows indicate tooth germs, arrowheads indicate the successional dental epithelium (SDE) in each species as suggested by our model. The epithelia are depicted slightly off-plane (shallower on the z-plane) with respect to the erupted tooth (analogous to viewing a thick section). The left two columns (stickleback and salmonids) illustrate species that lack a successional dental lamina (SDL), whereas the right two columns illustrate species that do have an SDL; the zebrafish, which exhibits a transient SDL (note lack of SDL in the bottom row) vs. salamanders, which maintain a permanent SDL (arrowhead in bottom row). The stickleback drawings are representative of both oral and pharyngeal tooth regeneration and based on our own observations. The zebrafish pharyngeal tooth drawings are based on our own observations in conjunction with previous literature [42, 69]. The salmonid drawings are based on data from rainbow trout [47, 49] and Atlantic salmon [48, 136], and are based primarily on oral tooth histology (though pharyngeal teeth are described as behaving similarly in rainbow trout [49]). The salamander oral tooth drawings are based on data from the lberian ribbed newt [137, 138] and axolotl [139].

Is the lack of a distinct successional dental lamina plesiomorphic, or derived?

Micro-CT studies found that tooth plate fossils can retain signatures of past replacement events. In the fossil species *Andreolepis hedei*, a stem bony vertebrate (osteichthyan), tooth replacement did not appear to have occurred in a static 1-for-1 fashion, and likely lacked a permanent SDL [45]. These fossil data suggest that early modes of tooth replacement in ancient bony vertebrates are grossly similar to the condition in stickleback and other bony fishes such as bichir, medaka, and cod [18, 47, 71]. While some have concluded that such a regenerative system is likely secondarily derived [31], it remains possible that these species have retained an ancient, less orderly tooth regeneration system (i.e., it is plesiomorphic), and that the presence of a successional dental lamina could be convergent in certain vertebrate groups, as recently proposed [45]. By analyzing the histology and gene expression of other ray-finned fishes' dentitions, it could be possible to test if the tooth regeneration systems like those found in sticklebacks and other fishes are more likely a plesiomorphic or derived character.

Tooth number evolution in sticklebacks

We found that stickleback replacement tooth germs initiate possible "1-for-2" replacement events up to ~25% of the time. While our histological approach alone is unable to definitively determine the successional future or history at a given tooth position, our data leave open the possibility that adult tooth number variation in stickleback populations could be modulated by modifications to the replacement process, as previously posited [63]. Specifically, Ellis et al. (2015) found that marine and freshwater stickleback populations exhibited a consistent difference in adult (~40 mm standard length) bilateral ventral tooth plate number, with an average difference of around 25 teeth, an approximately 50% increase in tooth number in derived freshwater fish relative to ancestral marine fish. In this same study, tooth germ number was quantified on ventral tooth plates, and found to have an average difference of ~12 tooth germs in 40 mm adult fish, with freshwater fish from two different populations having more tooth germs than marine fish. As was previously addressed [63], these data suggest that the increased tooth numbers in freshwater stickleback populations relative to ancestral marine populations are likely not due to only differences in the appearance of new tooth germs; a difference in tooth shedding rate must also be at play. In line with these previous observations on inferred shedding rates, new histological data presented here suggest that the germ:shed ratio could be effectively doubled for some replacement events (1-for-1 vs 1-for-2) which conceivably could contribute to the overall tooth number differences seen between different populations. We hypothesize that up- or down-regulation of these "1-for-2" events may also contribute to these evolved tooth number differences in sticklebacks. Comparative studies on replacement events between multiple marine and freshwater stickleback populations could determine if "1-for-2" events differ in frequency between marine and freshwater fish, and if these differences could be related to genetic or developmental modifications to cell behavior or regeneration potential of the SDE.

Broader implications in epithelial appendage evolution and diversification

Ideas surrounding the homology of disparate epithelial appendages (EAs), like teeth and hair, have been proposed as early as Darwin's *On the Origin of Species* [111], and were thereafter bolstered by morphological comparisons and pleiotropic phenotypes affecting teeth, hair, and other epithelial organs [112–114]. Some consensus regarding the homology or shared nature of the placode stage has been established [58, 59], while other work has indicated that larger-scale patterning mechanisms (e.g., concerted activity of Wnt, Eda, FGF, and SHH pathways)

are at play across diverse EAs, including fish scales and amniote body coverings (reptile scales and feathers) [52, 115]. Distinguishing between uninterrupted 'historical homology' vs. deep homology [116, 117] of EAs like teeth and hair remains difficult to assay using only extant vertebrates; fossil evidence of intermediate forms could feasibly provide links between modernly disparate EAs. Although it is not clear whether teeth and hair are related by strict-sense historical homology or cooption, teeth, hair, and other epithelial appendages exhibit widespread similarities in their early development and regeneration [23, 57]. For example, in some rare instances, human dentitions form hair [118, 119], and *Med1* mutant mouse dentitions form hair in the mouse incisor socket [120].

Here, we tested the hypothesis that two different tooth regeneration systems (zebrafish and stickleback) deploy similar patterns of gene expression (e.g., whether putative replacement epithelia have similar patterns of gene expression). Supporting this hypothesis, we found expression of a battery of nine candidate genes present in both the stickleback and zebrafish naïve dental epithelia, which we termed the SDE. Intriguingly, all nine of these genes are also expressed in mouse hair follicle stem cells, where most of these genes are known to play important roles in hair regeneration [66, 67, 90, 94, 101, 102, 104, 121–123]. This similarity in gene expression suggests that hair and teeth may regenerate using a shared epithelial cell type, namely one that responds to Bmp and Wnt signals during the regenerative cycle, with Bmps suppressing and Wnts facilitating differentiation. This work builds on previous studies on stickleback gene expression levels in ventral tooth plates of 254 known hair follicle stem cell gene orthologs, where this gene set was found to be reduced in *Bmp6* mutants relative to wild type [62], and elevated in a high-toothed freshwater population relative to an ancestral low-toothed marine population [64]. Together, the expression profile similarities between hair follicle stem cells and dental tissues shown previously and presently will hopefully encourage further studies describing the shared and divergent developmental genetic mechanisms at play in these diverse organs. For example, the expression of the SDE markers reported here could be compared in other epithelial appendages such as amniote scales and feathers, where less is known about cellular and genetic bases of regeneration.

One curious feature of fish tooth regeneration addressed previously is the apparent lack of slow-cycling label-retaining cells (LRCs) in the bichir and salmon oral dentitions, which do not employ an SDL [18, 48]. Given that LRCs are present in other tooth regeneration systems [17, 41, 50, 124] and in the hair follicle stem cell niche [125–129], these findings could represent a major difference between some fish dentitions and other epithelial appendages, including other fish teeth [50]. Further studies on LRCs in other epithelial appendages, including other fish species' teeth, will help delineate how common and/or consequential these changes in cell behavior might be. However, regardless of whether LRCs or a true epithelial stem cell niche exists across all fish dentitions, further elucidation of the shared gene expression and cellular processes between the SDE, hair follicle stem cells, and other epithelial appendages is warranted and ongoing, including defining functional responses to Bmp and Wnt signaling inputs, and testing function of genes that mark the SDE and/or mesenchymal tooth precursors during tooth regeneration.

Methods

Animal husbandry

Animals were raised following UC Berkeley IACUC protocol AUP-2015–01-7117. Zebrafish husbandry adhered to standard methods [130]. Sticklebacks were raised in 110-L aquaria in brackish water (3.5 g/L Instant Ocean salt, 0.0217 g/L sodium bicarbonate) at 17–18 °C in 8 h of light per day. Fish were fed live artemia as young fry, live artemia and frozen daphnia as young juveniles (1–2 cm standard length), frozen daphnia and frozen bloodworms as older juveniles (2–3 cm standard length), and frozen bloodworms and *Mysis* shrimp as sub-adults and adults (3 + cm standard length).

Generation of TCF/Lef-miniP stable transgenic lines

The TCF/Lef-miniP:dGFP plasmid was a gift from Tohru Ishitani [86]. To 'restabilize' GFP, we removed the destabilization signal by digesting the previously published plasmid with HindIII and ClaI, and filling this lesion with a short double-stranded piece of DNA that comprised two synthesized (Integrated DNA Technologies), phosphorylated, and annealed oligonucleotides. Forward oligo: 5' AGCTTTAAAGCGACAT 3', reverse oligo: 5' CGATGTCGCTTTAA 3'. This change to the plasmid removed the majority of the destabilization signal, leaving just two extra amino acids (Lys, Leu) before a stop codon on the C terminus of eGFP. Zebrafish and stickleback transgenic lines were generated using both the unmodified and modified versions of the Tcf/Lef-miniP constructs according to standard methods in each species, using Tol2-driven transgenesis [131] by injecting zygotes at the one-cell stage [132, 133].

Alizarin Red staining

Alizarin Red was used to stain tooth plates as described previously [134]. In brief, whole fish were fixed in 10% neutrally buffered formalin overnight at room temperature and washed with tap water or $1 \times \text{phosphate-buffered}$ saline with 0.1% Tween-20 (PBST) for 1 h at room temperature. Samples were then washed in a 0.008% Alizarin Red S solution in 1% KOH for 24 + hours. Once adequately stained, samples were transferred to 1% KOH for 1–3 days to continue clearing. Once cleared, tooth plates were dissected, further cleared in 50% glycerol for 1–5 days, flat-mounted, and imaged on a Leica DM 2500 compound microscope.

Sectioning

Stickleback sub-adults (~2 cm) derived from marine (Rabbit Slough [RABS]) and/or freshwater (Cerrito Creek [CERC] or Fishtrap Creek [FTC]) populations, and AB strain zebrafish adults or sub-adults (1.5-2 cm) were euthanized, decapitated, and fixed overnight in 4% formaldehyde (Sigma P6148) in 1×phosphate-buffered saline (PBS) at 4° C with heavy agitation, washed 3×20 min with PBST on a nutator, then decalcified for 5-7 days (sticklebacks) or 3-5 days (zebrafish) in 20% ethylenediaminetetraacetic acid (EDTA, pH 8.0) at room temperature on a nutator. Once decalcified, specimens were again washed 3×10 min in PBST, then gradually stepped into 100% EtOH via 15-60 min washes in 30, 50, 70, and 100% EtOH in RNAse-free H₂O. Samples were sometimes washed into MeOH and stored at this stage for up to 12 weeks. Samples were then washed for 1-2 h in 50/50 EtOH/Hemo-De at room temperature, then 1 h + in 100% Hemo-De at room temperature, then 1-2 hin 50/50 Hemo-De/paraffin (Paraplast X-tra, Fisher) at 65 °C, then rinsed and washed overnight at 65 °C in 100% paraffin. Fish heads were embedded in plastic molds with 100% paraffin (heated to 65 °C), mounted, sectioned on the sagittal (stickleback), off-sagittal (~30 degree tilt on the coronal axis; zebrafish), coronal (both species) or transverse (stickleback) planes with a Microm HM 340 E microtome. Sections were captured on Superfrost Plus slides, sometimes stored for up to 3 weeks at room temperature prior to analysis. To prepare slides for histological staining or ISH, slides were de-paraffinized (5 min incubation at 65 °C, let cool, submerge for 5 then 10 min in 100% Hemo-De, 5 min 100% EtOH, 10 min 80% EtOH/ H₂O, 10 min 100% H₂O).

Hematoxylin and eosin (H&E) staining on sections

H&E staining was performed on sectioned tooth material using the following series of washes: hematoxylin solution (VWR) for 3 min, tap water for 2×20 s, bluing solution (0.1% sodium bicarbonate) for 2 min, tap water for 20 s, acid alcohol (0.32% HCl in 95% EtOH) for 20 s, tap water for 20 s, eosin solution (VWR) for 30 s, 95% EtOH for 2×2 min, 100% EtOH for 2×2 min, then Hemo-De

for 2×5 min. Slides were then immediately coverslipped with Permount (Fisher), allowed to dry overnight, and imaged on a Leica DM2500 compound microscope.

In situ hybridizations on sections

All in situ hybridization (ISH) steps were performed on freshwater (CERC) stickleback, or AB strain zebrafish sections. The following steps were performed in Lock-Mailer microscope slide jars (Sigma-Aldrich) in a volume of 9-11 mL unless stated otherwise. To begin the in situ process, slides were washed for 5 min in PBST, 5 min in proteinase K solution (15 µg/mL in PBST), rinsed briefly with PBST, then re-fixed for 20 min at room temperature in 4% formaldehyde in PBS. Fixative was then washed out with one PBST rinse and 2×10 min PBST washes before pre-hybridization. Slides were washed 2×5 min at room temperature in hybridization buffer, followed a long incubation in hybridization buffer (no probe, "prehyb" step) for 1–4 h at 67° C in a rotating hybridization oven (hybridization buffer is 50% formamide, $5 \times SSC$, 0.1% Tween, 5 mg/mL CHAPS, 1 mg/mL yeast RNA, 0.1 mg/mL heparin, pH 6.0 with citric acid). Riboprobes were generated essentially as described previously [68] against gene orthologs of interest (see Additional file 6: Table S1). Riboprobe templates were created either by PCR cloning gene fragments from cDNA or genomic DNA, or by synthesizing whole or partial inferred coding sequences (Gene Universal, Delaware, USA). See Additional file 6: Table S1 for a complete list of transcript accession numbers and sequences corresponding to exact riboprobe templates used in this study. Stickleback Bmp6 and *Pitx2* riboprobes were previously published [61]. The zebrafish *pitx2* riboprobe was previously published [135]. Zebrafish and stickleback CD34 orthologs were identified using the Genomicus synteny browser v100.01 (www. genomicus.biologie.ens.fr), which supported orthology of mammalian CD34 to the fish genes assayed here (ENSGACG00000011016 and ENSDARG00000095268). The translated products of these genes appear to code for proteins containing a partially recognizable CD34 domain (pfam06365). Riboprobes were synthesized with digoxigenin-labeled UTP and added at a concentration of ~100–500 ng/mL in 10 mL of hybridization buffer and agitated overnight in a rotating hybridization oven at 67 °C. Riboprobes in hybridization solution were stored at -80° C and reused in some cases. The following day, six pre-heated hybridization washes at 67 °C in a rotating hybridization oven were performed for 20-90 min each, totaling 5-6 h of total "hyb wash" time (hyb wash is the same recipe as hyb buffer, excluding CHAPS, RNA, and heparin). Slides were then rinsed and washed in preheated Maleic acid buffer with Tween (MABT) at 67 °C for 20 min, then washed in pre-heated MABT for 20 min at room temperature (to allow for slow cooling). Slides were then removed from slide jars, placed in a humidor (a sealed plastic container with wet paper towels for moisture and pieces of plastic to raise slides form the bottom), and blocked with 70-100 µl (enough to cover all tissue on slides) of 2% Boehringer Blocking Reagent (BBR), covered with parafilm for 1 to 3 h at room temperature. Following the block step, block was poured off each slide, and anti-digoxigenin alkaline phosphatase conjugated antibody (Roche SKU 11093274910) was added at a concentration of 1:2000 in 2% BBR, again using enough to submerge each piece of tissue on each slide $(70-100 \ \mu l)$ beneath parafilm and incubated at 4 °C overnight, or 3 h at room temperature in the dark. Importantly, tissue was not allowed to dry out beneath the parafilm during the block or antibody steps above. The following day, we performed one MABT rinse and $5 \times 20-50$ min MABT washes (back into slide jars) over the course of 3–4 h, agitated at room temperature, to wash out residual antibody, usually followed with an overnight MABT wash unagitated at 4 °C. To begin the coloration process, slides were changed into NTMT (0.1 M Tris pH 9.5, 0.05 M MgCl₂, 0.1 M NaCl, 0.1% Tween) via $3 \times 5-10$ min washes before removing the final NTMT wash and replacing it with 10 mL of coloration solution (NTMT with 25 μ g/ mL Nitro blue tetrazolium chloride [NBT] and 175 µg/ mL 5-bromo 4-chloro 3-indolyl phosphate [BCIP]). Signal development was carried out for 2-30 h to visualize mRNA localization. Once adequately developed, slides were rinsed, then washed for 10 min in PBST, fixed in 4% formaldehyde in PBS for 1-5 days at 4 °C. Slides were then washed 3×5 min in PBST. At this point, some select slides were counterstained with 0.1 µg/mL DAPI in $1 \times PBST$ for 5–30 min at room temperature (the same 10 mL of solution was used to stain all sections shown in this paper, stored at 4 °C). To prepare slides for imaging, they were rinsed then washed $3 \times 5 + \min$ with deionized H_2O , coverslipped with deionized H_2O , and imaged on a Leica DM2500 microscope.

Quantification of 1-for-1 vs 1-for-2 tooth replacement in sticklebacks

To address the possibility of 1-for-2 tooth replacement in sticklebacks, we H&E-stained serial sections (described above) of nine sticklebacks of various sizes (20–40 mm) from various backgrounds ($5 \times CERC$, $1 \times FTC$, $2 \times RABS$, $1 \times marine/freshwater$ hybrid). Across these preparations, we identified a total of 67 pharyngeal tooth germs for which we had complete histological profiles at the mid- to late bell stage, that were also present in the middle of the tooth plate (i.e., surrounded by erupted teeth), which represent presumed or potential replacement teeth. Of these 67 tooth germs, 17 (25.4%) were clearly observed to be abutting and connected to bone remodeling events at the bases of two erupted teeth. These associations also appeared to contain an abundance of likely osteoclasts surrounding the disrupted bone in the erupted teeth (carets in Fig. 4), which we interpret as evidence of possible shedding events (or at the very least, loosening events) inprogress at the time of tissue preparation. We similarly located all oral tooth germs present on these slides, though the oral tooth fields were only visible on two sectioned specimens. We did not observe oral tooth germs as diverging from a 1-for-1 replacement scheme (n = 0/8).

Supplementary information

The online version contains supplementary material available at https://doi. org/10.1186/s13227-021-00172-3.

Additional file 1: Fig. S1. Histological plane orientation and description in sticklebacks. Illustrations at top depict a tooth (red cone) and indicate section planes with dotted rectangles, which are pictured below for each respective plane of section (labeled at top). These planes of section roughly translate to whole-animal anatomical planes. Left and middle illustrations show the sagittal and/or transverse plane (because stickleback teeth are mostly conical, these two planes of section appear generally similar in section). On these planes of section, teeth often appear off their medial axis (left vs center illustration, see labels). Rightmost illustration depicts the coronal plane, where teeth appear as rings of bone. Bottom images are of H&E-stained sections representing each illustrated plane of section above. Left bottom image shows a pharyngeal tooth on the transverse plane. Center bottom image shows an oral tooth on the sagittal plane. Right bottom image shows a pharyngeal tooth sectioned on the coronal plane. An erupted tooth ("et") is indicated in the center image of an off-medial tooth (suggested with a red dotted line) sectioned on the sagittal plane. In all H&E images, tooth germs are indicated with an arrow, and the SDE is indicated with arrowheads. All scale bars $= 50 \,\mu m$.

Additional file 2: Fig. S2. DAPI counterstaining in situ hybridized sections aids in distinguishing epithelium from mesenchyme. Three gene expression patterns are shown as examples, labeled at top of columns in figure. Each column shows the same section corresponding to each gene, top two images are in brightfield, bottom two show DAPI fluorescence. 1st row of brightfield images are shown without any markup. 2nd row of brightfield images outline the basalmost epithelium with black dotted lines where possible (gray segments suggest borders out of the plane of section); bone is false-colored red. Expression domains in tooth germs are indicated with black arrows. Expression in the SDE is indicated with black arrowheads. Superficial epithelium with Gli1 expression adjacent to erupted teeth is indicated with white arrowheads. 3rd row of images shows the DAPI counterstain of each brightfield image above it, with the same epithelium outlines overlain (white dotted lines). 4th row shows DAPI images without any markup. Scale bar = $50 \,\mu\text{m}$ and applies to all panels

Additional file 3: Fig. S3. Stickleback *Dkk3b*, *Igfbp5b*, and *Bambia* ISH on sections through dental tissue. The stickleback SDE was positive for Dkk3b and Igfbp5b (black arrowheads in top images), though these genes marked many cell populations across the dental field and surrounding tissues. *Bambia* was detected in tooth germ epithelium (black arrows) and mesenchyme (white arrows), and additionally in mature tooth mesenchyme, but never in the SDE (gray arrowheads). Scale bar = 50 µm and applies to all panels.

Additional file 4: Fig. S4. Zebrafish *igfbp5b* expression in deep tooth mesenchyme. Top row shows brightfield images, bottom row shows

the corresponding DAPI counterstains to each brightfield image above. The SDL of zebrafish did not show *igfbp5b* expression (gray arrowhead). Expression was detected in a subset of the deep dental mesenchyme (white arrows). Scale bar = $50 \ \mu m$ and applies to all panels.

Additional file 5: Fig. S5. Expression of *Bmp6*, *Bmp1a*, *CD34*, *Gli1*, *Igfbp5a*, *Lgr4*, *Lgr6*, *Nfatc1*, and *Pitx2*, highlighting mesenchymal expression. All of these genes were found to mark the SDE, except for Pitx2, and are expressed in some portion of dental mesenchyme (white arrows). Species and gene assayed are indicated in the figure. All scale bars = 50 μ m.

Additional file 6: Table S1. Riboprobe sequence information. The first column (A) designates the species for which each riboprobe was designed. The second column (B) shows the gene symbol corresponding to each riboprobe. The third column (C) is the NCBI or Ensembl accession number representing the closest deposited sequence to the riboprobe template used. The fourth column (D) shows the original publication, if any, where the riboprobe activity was previously documented. The fifth column (E) contains the nucleotide sequence of the entire gene-specific portion of each riboprobe RNA sequence was the reverse complement of the sequence shown in column D).

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Authors' contributions

TAS and CTM conceived the study. TAS designed the experiments. TAS, SBS, and EJM performed the experiments and collected the data. TAS prepared the figures. TAS and CTM wrote the manuscript. All authors provided input on the contents of the final manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

The datasets used and analyzed during the current study are available from the first author on reasonable request.

Ethics approval and consent to participate

Animals were raised following UC Berkeley IACUC protocol AUP-2015-01-7117.

Consent for publication

Not applicable.

Competing interests

The authors declare no competing interests.

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