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Early epithelial signaling center governs tooth budding morphogenesis

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During organogenesis, cell fate specification and patterning are regulated by signaling centers, specialized clusters of morphogen-expressing cells. In many organs, initiation of development is marked by bud formation, but the cellular mechanisms involved are ill defined. Here, we use the mouse incisor tooth as a model to study budding morphogenesis. We show that a group of nonproliferative epithelial cells emerges in the early tooth primordium and identifies these cells as a signaling center. Confocal live imaging of tissue explants revealed that although these cells reorganize dynamically, they do not reenter the cell cycle or contribute to the growing tooth bud. Instead, budding is driven by proliferation of the neighboring cells. We demonstrate that the activity of the ectodysplasin/Edar/nuclear factor κB pathway is restricted to the signaling center, and its inactivation leads to fewer quiescent cells and a smaller bud. These data functionally link the signaling center size to organ size and imply that the early signaling center is a prerequisite for budding morphogenesis.

Introduction

Cellular behaviors, including cell proliferation, movement, and differentiation, are regulated in part by morphogens during embryonic development. In many organs, signaling molecules are produced by groups of specialized cells, signaling centers that control cell fates and cellular behaviors in the surrounding tissue. Well-defined examples include the apical ectodermal ridge and zone of polarizing activity of the developing limb, the isthmus regulating midbrain and hindbrain regionalization, the notochord that patterns the neural tube and the somatic mesoderm, and the enamel knot that instructs tooth morphogenesis (Jernvall and Thesleff, 2000; Partanen, 2007; Towers et al., 2012). These signaling centers share features: they are composed of a relatively small number of cells that regulate cell fates and behaviors by secreted factors, including sonic hedgehog (Shh) and members of the fibroblast growth factor (Fgf), bone morphogenetic protein (Bmp), and Wnt families. The signals expressed by the signaling centers are well characterized, but less is known about the origins and fate of signaling center cells and the cellular behaviors they regulate.

Ectodermal organs such as hair follicles, feathers, and mammary glands are initiated as a thickening of the epithelium called a placode (Pispa and Thesleff, 2003). Soon after, the underlying mesenchymal cells condense, and the epithelium invaginates to produce a bud (Jernvall and Thesleff, 2000; Schmidt-Ullrich and Paus, 2005). Like other ectodermal appendages, teeth form through epithelial–mesenchymal interactions and proceed via similar early stages (Jussila and Thesleff, 2012). In mice, tooth morphogenesis is initiated around embryonic day 11 (E11) when a horseshoe-shaped epithelial thickening, the dental lamina, appears in both jaws. Gene expression analyses indicate that the continuous dental lamina resolves into two separate domains in each jaw half, the incisor and the molar placodes, between E11 and E12 (Biggs and Mikkola, 2014). Genes that are initially expressed along the entire dental lamina and later become confined to incisor and molar primordia include Foxi3, Pitx2, and Sox2 (St Amand et al., 2000; Juuri et al., 2013; Shirokova et al., 2013). In addition, several signaling molecules such as Shh, Wnt10a/b, and Bmp2 are expressed in a more restricted manner (Dassule and McMahon, 1998; Keränen et al., 1998), suggesting the presence of a specific signaling center within the tooth placodes (Dassule and McMahon, 1998; Jernvall and Thesleff, 2000). However, the exact identity and function of this signaling center has remained obscure.

After the placode stage, tooth morphogenesis proceeds through bud, cap, and bell stages before hard tissue mineralization (Timmers and Thesleff, 2009). Immediately before the transition from the bud to the cap stage, a well-defined signaling center called the primary enamel knot (EK) forms at the tip of the bud and exhibits restricted expression of Shh, multiple members of the Bmp, Fgf, and Wnt families, and the cyclin-dependent kinase inhibitor gene p21 (Jernvall et al., 1998; Sarkar and Sharpe, 1999; Jernvall and Thesleff, 2000). In addition, expression of Edar, the receptor of the Tnf-like ligand ectodysplasin (Eda), is restricted to the EK (Tucker et al., 2000; Laurikkala et al., 2016). This article is distributed under the terms of an Attribution-Noncommercial-Share Alike-No Mirror Sites license for the first six months after the publication date (see http://www.rupress.org/terms). After six months it is available under a Creative Commons License (Attribution-Noncommercial-Share Alike 3.0 Unported license, as described at http://creativecommons.org/licenses/by-nc-3.0/).
et al., 2001). The EK is essential in controlling later stages of folding and patterning of the dental epithelium, where it stimulates proliferation of the mesenchyme and the surrounding epithelial cells, whereas cells within the EK remain nonproliferative (Jernvall and Thesleff, 2000; Jussila and Thesleff, 2012).

The molecular regulation of early tooth morphogenesis has been studied extensively, and a large number of mouse mutants displaying early developmental arrest have been described (Bej, 2009; Jussila and Thesleff, 2012; Biggs and Mikkola, 2014; Lan et al., 2014). However, the cellular behaviors contributing to dental morphogenesis are just beginning to be understood. Here, we use the mandibular incisor as a model to study the formation and function of the poorly characterized early signaling center. With the aid of Fucci cell cycle indicator mice and live laser scanning confocal tissue microscopy, we have recently shown that the cellular mechanisms driving hair placode formation include a switch to G1 phase and centripetal cell migration (Ahtiainen et al., 2014). The current study shows that in tooth placodes only a subset of cells become quiescent and form an early signaling center. These cells exhibit migratory behavior whereby they condense and form the mature signaling center. The signaling center cells remain nonproliferative, whereas a burst of cell proliferation in the neighboring dental epithelial cells leads to bud formation. We show that Eda/Edar/NF-κB signaling regulates the number of signaling center cells and thereby tooth bud size. Finally, our findings indicate that the EK is not derived from the early signaling center cells, but rather it forms de novo at the tip of the tooth bud.

Results

A subset of epithelial cells remain in G1 phase throughout embryonic mouse incisor placode and bud morphogenesis

To understand cellular mechanisms in early tooth morphogenesis, we analyzed cell cycle dynamics in developing mandibular incisors before the onset of morphological development (E10.5), at dental lamina stage (E11.0–11.5), placode stage (E12.0–12.5), and early (E13.0) and late (E13.5) bud stage. Fluorescent cell cycle indicator (Fucci) reporter mice (Sakae-Sawano et al., 2008) were used to monitor cell cycle status. At E10.5 and E11.0, G1/G0 (hereafter G1) and S/G2/M phase cells (identified by nuclear red and green fluorescence, respectively) were evenly distributed throughout the mandible (Fig. S1). At E11.5, however, a clear clustering of G1 cells was observed; the G1 focus seemed to coincide with the dental lamina. Between E11.5 and E12.0, the G1 foci in the prospective incisor and molar region were connected initially, but by E12.5 they were fully split into two separate G1 foci (Fig. S1).

To assess the location and identity of the G1 foci, we analyzed these cells in the developing incisors of transgenic mice expressing Keratin17-GFP (K17-GFP), a marker for the epithelium of ectodermal appendages (McGowan and Coulombe, 1998). K17-GFP was initially expressed in the periderm throughout the oral epithelium (Fig. 1 A). At E12.0, K17-GFP expression became more restricted to the forming placode, and a loose band of nonproliferating Fucci G1 cells was seen in the labial part of this area. At E12.5, the G1 cells had condensed into a restricted area closer to the mandibular midline (Fig. 1, A and B). At the late bud stage (E13.5), two G1 foci were evident in the incisors: one in the dental cord (junction between the bud and oral epithelium) and a new distinct cluster of cells at the tip of the bud (Fig. 1, A and B). This G1 focus corresponds to the known location of the emerging EK (Jernvall et al., 1998; Munne et al., 2009). A schematic representation illustrates the G1 cell populations in incisor morphogenesis (Fig. 1 C). To conclude, there are distinct subpopulations of epithelial cells in the forming incisor that exhibit cell cycle cessation in a specific spatiotemporal pattern.

The early G1 cell population colocalizes with signaling center markers

To characterize the G1 cell populations, mandibles were first imaged by fluorescence microscopy and then subjected to whole-mount in situ hybridization for known dental epithelial markers. Foxi3 is expressed early in the dental lamina and later at placode and bud stages; its expression spans the entire thickened dental epithelium but is absent from the oral epithelium (Shirokova et al., 2013). Dkk4 is expressed as a narrow stripe along the prospective dental epithelium from E10.5 onward (Bazzi et al., 2007; Finiaux et al., 2008). Sox2 is initially expressed in the dental lamina during E12 and becomes confined to incisor and molar regions (Juuri et al., 2013). Shh is expressed in the dental lamina and a subset of epithelial cells throughout early tooth morphogenesis and marks a putative early signaling center (Dassule and McMahon, 1998; Keränen et al., 1998; Hovorakova et al., 2011).

We found that the labial G1 population coincided with Foxi3 expression in the dental lamina at E11.5 and later in the forming placode and bud (Fig. 2 B). However, the Foxi3 expression domain was broader at all time points and colocalized with K17-GFP positivity (E12.0–E13.0; Fig. 2 A). In contrast, Sox2-expressing cells colocalized only partially with the early G1 population and K17-GFP–positive cells and mainly resided lingual to them (Fig. 2 B and Fig. S2, A and B). Dkk4 colocalized with the G1 population but was down-regulated at E13.0, whereas the G1 focus still persisted (Fig. 2 C). Interestingly, Shh expression colocalized with the G1 cell population at all observed time points (E11.5, 12.0, 12.5, and 13.5; Fig. 2 D), identifying this subset of cells as an early epithelial signaling center.

The number of G1 cells stays constant through placode and bud stages

The early focus of G1 cells was restricted to a progressively smaller area, whose shape changed from a long narrow stripe into a more triangular shape in whole-mount view (Fig. 2) and also in 3D volume renderings from fluorescence confocal z-stacks (Fig. 3 A and Video 1). To understand the mechanisms of this phenomenon, we assayed the cell number and volume in the G1 population.

G1 cells remained in the front part of the developing bud, whereas proliferating cells appeared on the lingual side as shown by Fucci S/G2/M reporter (Fig. 3 A). This was also seen by incorporation of 5-ethyl-2′-deoxyuridine (EdU) nucleoside analog delivered via in vivo injections of pregnant K17-GFP:Fucci G1 dams at E11.5, 12.5, and 13.5 (Fig. S4 A). Quantification of percentage of cells in G1 in the condensate area showed a specific increase in the proportion of G1 cells, whereas in the oral epithelium most of the cells remained proliferative (Fig. 3 B). However, the absolute number of G1 cells in the early signaling center remained constant throughout morphogenesis, whereas the number of S/G2/M cells adjacent to the early signaling center increased significantly (Fig. 3 C).
Because reduction in G1 cell number cannot explain the restriction of the early signaling center area, we next measured cell volume changes during placode formation. The cell volume in the early signaling center decreased significantly with time, whereas the volume of S/G2/M cells in the tooth epithelium and G1 cells in the oral epithelium stayed constant (Fig. 3D and Fig. S3), showing that the early signaling center cells become more compact as the early signaling center changes shape.

Differential cell migration contributes to early signaling center condensation

To study the cellular behaviors that mediate early signaling center compaction during placode morphogenesis, we used Fucci whole-mount explant cultures and live confocal fluorescence microscopy. In E12.0 mandibles, the incisor G1 focus is distinguishable as a narrow stripe. These mandibles were dissected and imaged for 24 h, and cell movement was tracked in four distinct cell populations: the medial and lateral G1 signaling center cells, the adjacent S/G2/M population, and cells of the oral epithelium not contributing to tooth formation. The cells in the lateral part of the G1 signaling center migrated directionally toward the medial midline along the dental lamina (Fig. 4A). In contrast, the cells in the proximal part of the area showed smaller overall displacement. The S/G2/M placodal cells directly adjacent to the G1 subpopulation and oral epithelial cells outside the placode area stayed localized (Fig. 4A). Quantification revealed that both the net displacement and track length were significantly higher in G1 early signaling center cells compared with the adjacent S/G2/M and oral epithelial cells (Fig. 4B). The G1 cells moved up to 35 µm from their original position, whereas the other two cell populations moved on average less than their own circumference, thus staying nonmotile (Fig. 4B). Analysis of the angle of movement of G1 cells showed that the lateral G1 cells, but not those close to the mandibular midline, displayed directional movement toward the midline (Fig. 4C).

To distinguish whether the observed cell movement results from displacement of the entire placode cell population by contraction or active migration of individual cells, we studied the movement of cell pairs with respect to each other. We traced cell pairs that were initially within close proximity and analyzed their distances in 3D 20 h later. This pairwise comparison revealed that the S/G2/M cells mostly retained their original neighbors (Fig. 4D). However, G1 signaling center cells did switch their partners frequently in both medial and lateral populations, the lateral G1 pairs ending furthest apart from each other (Fig. 4D).
We also inhibited cell movement by actin cytoskeleton disrupting agent latrunculin A (LatrA) or myosin II inhibitor blebbistatin. Inhibitors were applied at E12.0 (Fig. 4, E and F), and explants were cultured for 24 h. In contrast to control medium, both LatrA and blebbistatin prevented compaction of the G1 signaling center, which remained as a stripe similar to stage E12.0 and halted epithelial morphogenesis at placode stage (Fig. 4, E and F). Collectively, these data support directional cell migration as a mechanism for compaction of the early signaling center.

Early signaling center cells remain in G1, whereas budding morphogenesis is driven by cell proliferation in the adjacent lingual cell population

To evaluate the contribution of cell proliferation to tooth budding morphogenesis, we first followed budding morphogenesis in the very initial stages by live imaging from E12.0 to E12.0 + 22 h (Fig. S4, B and C; and Video 2). Live imaging indicated that the early signaling center cells remained in G1, whereas a wave of cell divisions occurred in the thickened placode starting at +6 h. These cells were adjacent to the G1 area on the lingual side and to a small extent on the labial side. A lower number of cytokinesis events was seen throughout the oral epithelium (Fig. S4, B and C; and Video 2).

For more detailed analysis, individual cells were traced during the extensive phase of bud growth (E12.5 + 10 h; Video 3). The G1 cells remained localized in the dental cord, close to the oral surface, in their original position and number (Fig. 5, A and B). In contrast, the S/G2/M proliferation wave in the adjacent cell population led to epithelial invagination and formation of the bud (Fig. 5, A–C). Next, we tracked individual cells, both G1 cells in the early signaling center and the proliferating S/G2/M population, to see if and
when they proceeded into mitosis. None of the followed G1 cells proceeded to S/G2/M during the time lapse; rather, they stayed localized in the early signaling center (Fig. 5 D) and thus did not contribute to the invaginating tooth bud. In contrast, the cells originally in S/G2/M phase remained in this cell cycle phase and divided during the time lapse. Their progeny contributed to the growing bud but did not exhibit obvious migratory behavior. Quantification of G1 and S/G2/M cells from confocal fluorescence live imaging samples showed an increase in the bud-forming population, whereas the number of the early signaling center cells stayed constant (Fig. 5 E). When the kinetics of the mitoses was scored, most mitotic activity was found lingual to the early signaling center (Fig. 5 F). There was no evident preference in mitotic orientation with respect to the longitudinal midline of the early signaling center (Fig. 5 G). Thus we conclude that the G1 cells of the early signaling center cells do not reenter the cell cycle and stay localized close to the oral epithelium, in the dental cord. Instead, budding morphogenesis is driven by proliferation of their epithelial neighbors.
Figure 4. Differential cell migration contributes to early signaling center condensation. Live confocal fluorescence microscopy of Fucci transgenes in mandibular whole-mount explant was used to characterize cell movement during placode formation. Explants were imaged at E12.0 + 24 h, and movement of placode cells, adjacent S/G2/M population, and cells of the oral epithelium was tracked. (A) Cell tracks in a representative incisor in four locations: (i) medial G1, (ii) lateral G1, (iii) S/G2/M in close proximity to the early signaling center, and (iv) S/G2/M in the oral epithelium. (B) Quantification of net displacement and track length showed significantly higher values in G1 early signaling center cells (medial and lateral populations together) compared with the adjacent S/G2/M and oral epithelial cells (Student’s t test, ***, P < 0.001; n_areas = 5 and n_cells = 30; data shown are means ± SD). The early signaling center G1 cells exhibited higher motility with respect to other epithelial cell populations. (C) G1 cell movement angles showed persistent directional movement by the lateral cells toward the mandible midline. Medial G1 cells, in contrast, moved toward the center of the early signaling center. (D) Movement...
The enamel knot is not a derivative of the early signaling center but is induced de novo in the posterior part of the bud

Our analysis of fixed samples showed that the two signaling centers, the early signaling center and the EK, represent two spatially separate regions of G1 cells. The signals expressed in the two signaling centers are largely the same (Jernvall and Thesleff, 2000), but it is not known whether there is a clonal relationship between the cells of the two signaling centers. The EK displays high Wnt/β-catenin signaling activity (Lammi et al., 2004; Liu et al., 2008). To gain further insight into EK induction and the origin of EK cells, we used the Tcf/Lef:H2B-GFP mouse model (Ferrer-Vaquero et al., 2010) that expresses nuclear histone 2B-GFP fusion protein as a marker of canonical Wnt signaling activity.

We first analyzed the Tcf/Lef:H2B-GFP reporter expression in fixed samples, costained with the epithelial marker EpCam. At E12.5, GFP-expressing cells were observed at the labial part of the incisor (Fig. 6 A). At E13.5, a separate area of GFP-positive cells was detected in the posterior part of the bud (Fig. 6, A and B). The GFP-positive regions corresponded with the early signaling center and the EK (Fig. 6, C and D).

Next, we imaged E13.0 Tcf/Lef:H2B-GFP;Fucci-G1 tissues up to 14 h. Initially, only one G1 focus, the early signaling center, was observed, and it overlapped with the Wnt reporter expression (Fig. 7 A). At E13.0 + 4 h, a novel cluster of Tcf/Lef:H2B-GFP cells began to emerge at the lingual part of the bud. Soon after, the first G1 cells appeared within the Wnt reporter–positive cluster. After 14 h of culture, the new signaling center (G1 focus) had further expanded. Wnt signaling activity coincided with the G1 cells but was more widespread (Fig. 7 B and Video 4). Quantification of GFP-positive and G1 cells showed an increase in number of both, but there was a higher increase in the number GFP-positive cells (Fig. 7 B).

In conclusion, live imaging of E12.5–E13.5 explants (Figs. 5 and 7) indicated that the early signaling center cells stay close to the oral surface and do not contribute to the EK. Instead, the EK appears de novo at the tip of the mature bud, as the bud cells transition into G1 phase.

The size of the early signaling center correlates with the size of the tooth bud

The Eda/Edar/NF-κB pathway is an important regulator of tooth size and shape: Eda-null mice (Tabby mice) have small molar teeth (Harjunmaa et al., 2014). These findings prompted us to analyze the function of Eda in the early signaling center, known to fully depend on Eda signaling activity in embryonic teeth (Pispa et al., 2008), was first detected at ~E11.0 in the incisor region (Fig. S5) in a manner reminiscent of the G1 focus (Fig. S1). The same was evident also at later stages (Fig. S5).

Unfortunately, the expression level of the NFκB-LacZ reporter was too low to allow reliable analysis by immunofluorescence, but overlay of Fucci-G1 expression with the whole-mount X-gal–stained E12.5/12.75 explants revealed that the Eda/Edar activity colocalized with the early signaling center (Figs. 8 A and S5 B). In addition, we used the expression of Fgf20, a target gene of Eda whose expression level is greatly reduced in developing Eda-null teeth (Häärä et al., 2012), as a proxy for Eda signaling activity. In Fgf20Gal/− mice, the LacZ gene has been inserted into the Fgf20 locus (Huh et al., 2012). β-Galactosidase staining was limited to the G1 early signaling center cells in Fgf20Gal/−:K17-GFP:Fucci-G1 embryos, further confirming that Eda signaling activity is localized to these cells (Fig. 8 B).

Next we studied the effect of the loss of Eda signaling pathway on the early signaling center G1 cell population. At E12.5, the G1 foci were less prominent in Eda-null embryos and often showed a bipartite structure (Fig. 8 C). Quantification showed a significant decrease in the number of G1 cells and in overall volume of the signaling center (Fig. 8 D). In addition, the expression domain of signaling center marker Wnt10b was reduced in Eda−/− incisors compared with controls (Fig. 8, E and F). Comparison of the bud shape and volume at E13.0 revealed that incisor buds of Eda-null embryos were smaller in size and protruded less into the mesenchyme than in wild-type (wt) littermates (Fig. 8, G and H). In conclusion, the size of the early signaling center was reduced by loss of Eda, and this correlated with the smaller size of the tooth bud.

Discussion

Here we have dissected the cellular events driving epithelial morphogenesis in early tooth development, thereby providing insights into the mechanisms driving epithelial invagination. The presence of an early signaling center has been postulated in placode-stage teeth based on the shared molecular signals with the EK, the later-forming and well-characterized signaling center (Jernvall and Thesleff, 2000). However, the exact identity, formation, and functions of the early signaling center have remained obscure. Here we show that there is a molecularly and functionally discernible cell population within the tooth placode that is characterized by cell cycle cessation and differential cell migration. The early signaling center regulates tooth budding morphogenesis through cell proliferation of the neighboring cells in the placode. Yet the signaling center remains quiescent throughout early tooth development; the cells remain of G1 cell pairs within the forming signaling center and S/G2/M cells in the placode were studied. Cells were followed by live imaging for 24 h (E12.0 onward), and cell pairs, initially ≤15 µm distance from each other, were traced (nplacodes = 32 in nplacodes = 3). The pairwise comparison revealed that placode cells switch their neighbors frequently in both medial and lateral populations (Student’s t test, *** P < 0.001; mean cell pair distances = 37.7, 25.2, and 12.6, respectively). The lateral G1 pairs end at longer distances apart from each other. In contrast, S/G2/M cells retained their original neighbors. Box and whisker plot represented as minimum, 25th percentile, median, 75th percentile, and maximum values for each dataset. (E) Latrunculin A (8 μM) was applied to K17-GFP:Fucci G1 explants at E12.0 followed by 24 h culture. Initially G1 cells were distributed along the dental lamina. After 24 h control treatment, G1 cells were observed in a condensed area focus in the anterior part of the K17-GFP positive bud, similar to E13.0 incisors (compare to Fig. S1). In latrunculin A-treated explants, morphogenesis was arrested at placode stage: the signaling center did not change shape, and no invagination of the epithelium was seen. (F) As with LatA treatment, early signaling center G1 cells stayed dispersed in samples treated with blebbistatin [100 μM], and budding morphogenesis was abrogated. Narrow vertical arrowheads indicate the position of the early signaling center and wide horizontal arrowheads show the xy optical section position in the orthogonal section. Bars, 50 μm.
Figure 5. Early signaling center G1 cells do not contribute to the tooth bud. Cell division kinetics and single-cell fates in the developing incisor were analyzed with the aid of Fucci transgenes in explants imaged at E12.5 + 10 h. (A) Volume rendering of the end point of the time lapse of E12.5 + 10 h. (B) Still images of the time lapse (volume rendering of separate channels for G1 and S/G2/M). G1 cells remained condensed and localized in the dental cord. In contrast, G2/S/M nuclei appeared in the adjacent cell population, and the number of proliferating cells consecutively increased as the bud invaginated. (C) Surface rendering of the bud at different stages of invagination. G1 early signaling center stayed localized whereas the cells adjacent to it proliferated and gave rise to the bud. (D) Tracing the contribution of individual G1 and S/G2/M cells originating from various positions of the forming bud. None of the early signaling center G1 cells \( n = 44 \) progressed into S/G2/M. Of the 31 traced S/G2/M cells, 21 proceeded to cytokinesis and divided during the time lapse. The daughter cells contributed to the growing bud locally, and the cells did not exhibit apparent migratory behavior. (E) Quantification of cells from confocal fluorescence live imaging samples. Cell number at \( t = 0 \) h was adjusted to 100%. There was an increase in the bud-forming population, whereas the number of the early signaling center cells stayed constant \( n_{\text{transition}} = 8 \); data shown are means ± SD. (F) Plot of kinetics of cell proliferation. Most mitotic activity was found in the lingual cell population adjacent to the early signaling center, where a wave of mitoses started at E12.0 + 6 h, and highest mitotic activity was seen in the next 10–18 h. (G) There was no clear preference in the mitotic orientation in the forming bud \( n_{\text{waves}} = 100, n_{\text{explants}} = 5 \). Bars: (A and B) 100 μm; (C) 50 μm. Error bars indicate ±SD.
in the dental cord that later on becomes thinner and ultimately disappears (Jussila and Thesleff, 2012). We propose that this early signaling center be called the initiation knot.

In several contexts, cell cycle arrest is important for signaling center fate and function; the mammalian ventral node and notochord are relatively quiescent (Bellomo et al., 1996), as are the most central cells of the isthmic organizer in the midbrain/hindbrain border (Trokovic et al., 2005). The latter express cyclin-dependent kinase inhibitor p21, a known regulator of G1 cell cycle arrest, and display reduced proliferation (Trokovic et al., 2005). In nascent hair follicles, the entire placode appears to act as a signaling center. Initiation of hair placode formation is marked by G1 entry, and cells remain in G1 throughout placodogenesis (Ahtiainen et al., 2014).

The early signaling center shares several features with the primary EK. Like the early signaling center, the EK shows a distinct lack of cell proliferation as indicated by absence of BrdU incorporation (Jernvall et al., 1994). Furthermore, p21 is focally expressed in the EK (Jernvall et al., 1998), as well as in a subset of tooth placode cells (Keränen et al., 1998), likely representing the G1 population identified in this study. Although comparison of published in situ hybridization data from different studies is somewhat challenging, it seems plausible that the gene expression signature of the early signaling center and the EK is largely shared. In addition to p21, shared genes include Shh, Wnt10a/b, Fgf20, Bmp2, Lrp4, and Dkk4 (Vaahtokari et al., 1996; Åberg et al., 1997; Dassule and McMahon, 1998; Keränen et al., 1998; Flinaux et al., 2008; Liu et al., 2008; Ohazama et al., 2008;
Häärä et al., 2012). This molecular similarity raises the possibility that the EK could be derived from the early signaling center cells: the early signaling center cells could be “pushed down” from their original position as the tooth bud grows. Alternatively, the early signaling center cells could exit the G1 phase and contribute to bud growth by reentering the cell cycle. We show here, however, that neither is the case. The early signaling center cells stay in G1 phase and remain in the dental cord throughout budding morphogenesis. Thus, the two signaling centers are distinct from each other, and the EK arises de novo in the tip of the growing bud at E13.5 (Jernvall and Thesleff, 2000).

It has been suggested that the early Shh+ focus, i.e., the early signaling center, represents a rudimentary, deciduous incisor (Hovorakova et al., 2011). Our data do not support this interpretation, but rather establish that the early signaling center is an integral part of the developing tooth. We propose that the early signaling center functions to regulate budding morphogenesis via several signaling pathways. First, deficiency in Eda signaling leads to a smaller incisor tooth bud (Miard et al., 1999; this study), similar to the molar bud (Pispa et al., 1999). NF-κB reporter expression is restricted to the signaling centers: it first appears in the early signaling center of the incisors and somewhat later in molars, but is not present in the intervening domain at any time point, nor in the proliferating cells of the placode. Our study shows that the number of G1 cells in the early signaling center, as well as the expression domain of a signaling center marker Wnt10b, are significantly reduced in Eda-null embryos. Given that Eda/Edar regulates the expression of multiple signaling center genes (Wnt10a/b, Fgf20, and Shh; Pummila et al., 2007; Zhang et al., 2009; Häärä et al., 2012; Voutilainen et al., 2012), it is likely that the diminished bud size is caused by a combinatorial effect of reduced expression of multiple Eda target genes. Also, the EK is smaller in Eda-null mice (Pispa et al., 1999; Harjunmaa et al., 2014), suggesting that the function of Eda is conserved in the two signaling centers. Second, we show that Shh expression correlates with the G1 cells, whereas the cells of the invaginating epithelium are in S/G2/M phase and do not express Shh. Conditional deletion of Shh, leading to loss of expression at around E12.5, leads to a flat, poorly invaginated tooth bud (Dassule et al., 2000). Furthermore, application of Shh-releasing beads on E11.5 mandibles results in ectopic epithelial invaginations, suggesting that Shh drives epithelial cell proliferation locally (Hardcastle et al., 1998). However, it is apparent that in addition to growth factors from the early signaling center, several mesenchymal cues regulate epithelial cell proliferation and bud growth, including activin A (Ferguson et al., 2000; Hosokawa et al., 2009; Veistinen et al., 2009). Mesenchymal signals may directly impact the early signaling center, such as mesenchymal Bmp4, which is required for maintenance of epithelial Shh expression (Fujimori et al., 2010).

What then causes the switch to G1 in incisor early signaling center cells, and is it sufficient for changing cell fate? Although the Eda/NF-κB pathway plays an important role, a signaling center still forms even in the absence of Eda/NF-κB signaling. Our previous study in developing skin (Ahtiainen et al., 2014) showed that in addition to Eda, stimulation of canonical Wnt activity resulted in G1 switch. This likely also happens in the tooth: forced epithelial activation of Wnt/β-cat signaling induces the formation of multiple signaling centers (and ectopic teeth) in the oral cavity (Järvinen et al., 2006; Liu et al., 2008), and conversely K14-Cre–mediated epithelial deletion of...
β-catenin leads to loss of Shh expression and a developmental arrest at the placode–bud transition (Liu et al., 2008). Further studies will be required to clarify the role of the canonical Wnt pathway in the induction of the early signaling center.

We show here that the nonproliferative early signaling center cells show differential migratory behavior toward the mandibular midline, leading to a change in the signaling center shape from a wide stripe to a compact triangle. The molecular mechanisms governing the directional movement remain to be identified. We suggest that this phenomenon is important to ensure that a single, large incisor forms, as opposed to two or three incisors typical to many mammals. Several genetic and ex vivo manipulations have shown the propensity of the incisor signaling center to disintegrate. Inactivation of mesenchymal β-catenin, leading to reduced mesenchymal Bmp4 expression, results in splitting of the incisor placode into two domains, giving rise to two adjacent incisors (Fujimori et al., 2010). Similarly, ectopic application of either Bmp antagonist noggin or exogenous activin A causes splitting of the epithelial Shh domain ex vivo (Munne et al., 2010). The disintegrated small placodes gave rise to smaller incisors, suggesting that the large mouse incisor signaling center is required for correct incisor size (Munne et al., 2010).

What then is the ultimate fate of the early signaling center cells? Studies on the EK have shown that at least some of the EK cells are cleared by apoptosis and also autophagy (Vahtokari et al., 1996; Yang et al., 2013), and this could serve as the silencing of the signaling center. The same likely applies to the EK cells identified in this study. There is reason to believe that attenuation of Wnt signaling is required for the clearance of the signaling center. In mice lacking the Wnt/Bmp inhibitor Ectodin/Sostdc1/Wise, no apoptosis was detected, but instead ectopic Wnt reporter expression was apparent in the EK without Eda signaling (Stuart et al., 2005). This phenotype was rescued by the ECT cells being removed by apoptosis and also autophagy (Vahani et al., 1996; Yang et al., 2013), and this could serve as the silencing of the signaling center.

We show here that the nonproliferative early signaling center cells show differential migratory behavior toward the mandibular midline, leading to a change in the signaling center shape from a wide stripe to a compact triangle. The molecular mechanisms governing the directional movement remain to be identified. We suggest that this phenomenon is important to ensure that a single, large incisor forms, as opposed to two or three incisors typical to many mammals. Several genetic and ex vivo manipulations have shown the propensity of the incisor signaling center to disintegrate. Inactivation of mesenchymal β-catenin, leading to reduced mesenchymal Bmp4 expression, results in splitting of the incisor placode into two domains, giving rise to two adjacent incisors (Fujimori et al., 2010). Similarly, ectopic application of either Bmp antagonist noggin or exogenous activin A causes splitting of the epithelial Shh domain ex vivo (Munne et al., 2010). The disintegrated small placodes gave rise to smaller incisors, suggesting that the large mouse incisor signaling center is required for correct incisor size (Munne et al., 2010).

Figure 8. The reduced G1 cell number in the early signaling center of Eda null incisors correlates with a smaller bud size. NF-kB reporter and Fgf20ΔGal mice were used as reporters for Eda/NF-kB signaling activity in developing incisors. (A) Whole-mount X-gal staining of an E12.75 mandible shows colocalization of NF-kB reporter activity with the Fucci G1 population. (B) Maximum-intensity projection and a single optical section (inset) of Fgf20ΔGal/+;Fucci G1;K17-GFP E12.5 mouse mandible. X-gal staining (magenta) was confined to G1 (red) early signaling center cells (merged image of β-Gal and G1 only in the inset), whereas K17-GFP was expressed throughout the bud. (inset) Narrow horizontal arrowhead indicates the position of the early signaling center and wide vertical arrowheads show the xy optical section position in the orthogonal section. (C) Expression analysis of Fucci transgenes in Eda null mandibles at E12.5 revealed that the early G1 focus often displayed a bipartite structure and was less coherent compared with the control signaling center (maximum-intensity projection). (D) Quantitation of the early signaling center G1 cell number and overall volume showed a significant decrease in the number of cells and a decrease in the overall volume of the signaling center (Student's t test, ***, P < 0.001; *, P < 0.05; n = 5 wt mice and n = 7 Eda−/− mice). (E) Similarly to the reduction seen in the Fucci G1 cell population, the Wnt10b signaling domain was reduced in E12.5 Eda−/− samples. (F) Quantification of the size of the Wnt10b expression domain in mutant versus control embryos [Student's t test, ***, P < 0.001; n = 10 ctrl mice and n = 9 Eda−/− mice]. (G) Comparison of bud shape (based on K17-GFP expression) showed a flatter shape in Eda−/− embryos compared with control at E13.0. (H) Bud volume at E13.0 was reduced in the absence of Eda signaling (Student's t test, ***, P < 0.01; n = 8 wt mice and n = 6 Eda−/− mice). All data shown are mean ± SD. Bars: (A and B) 100 μm; (C and E) 30 μm.
In conclusion, we show that growth of the tooth bud is steered by proliferation of the cell population adjacent to the nonproliferating early signaling center, the initiation knot. Directional cell migration shapes the early signaling center, which may be important in determining the number of incisors. Eda/Edar/NF-kB signaling plays a role in defining the number of G1 cells and consequently regulates the size of the growing bud. Although we did not address the role of the early signaling center in developing molars, the colocalization of Shh and other signals with the G1 focus strongly suggests that similar mechanisms account for molar budding as well. However, current evidence suggests that somewhat different mechanisms of bud formation are used in other ectodermal organs. Cell proliferation seems to have a minor contribution in the epithelial morphogenesis of developing mammary glands until the primary sprout forms. A lower mitotic index compared with the adjacent epidermal epithelium and near-absence of BrdU incorporation represent displacement length in micrometers and relative angle.

Materials and methods

Animals and preparation and culture of embryonic skin

All mouse studies were approved by the National Animal Experiment Board. Transgenic mouse lines used in this study have been described earlier: [mK17 5]-GFP mice express GFP under the Keratin 17 promoter (Bianchi et al., 2005); Fucci mice are bitransgenic mice expressing nuclear red (mKO-Ctd1) and nuclear green (mAZ-Gem) in G1 and S/G2/M phases, respectively (Sakaue-Sawano et al., 2008); Tcf/Lef:H2B-GFP mice express nuclear GFP as a marker of canonical Wnt signaling activity (#013752; The Jackson Laboratory; Ferrer-Vaquer et al., 2010); transgenic NF-kB reporter mice express β-galactosidase under an NF-kB–responsive element (Bhakar et al., 2002); FGF20β-gal mice harbor an Fgf20-β-galactosidase (β-Gal) knock-in allele (Huh et al., 2001); and SPSS Statistics (IBM) software. Differences between groups in cell volume and proliferation analyses were assessed with the parametric Student’s t test. For cell shape and condensation studies, measurements were done as described previously (Ahtiainen et al., 2014). In brief, all measurements were done in three dimensions from confocal optical stacks. For cell tracking purposes, to minimize the global motion of cells caused by tissue deformation or growth, the tissues were allowed to recover for 2 h. The explants were imaged as described previously (Ahtiainen et al., 2014) with a Leica Biosystems TCS SP5 microscope and HC PL APO 10x/0.4 (oil) objective and, for fixed samples, in addition HCX PL APO 20x/0.7 Imm Corr (water, glycerol, oil) Ltd. bl and HCX APO 63×/1.30 Corr (glycerol) CS 21 objectives. Images of fixed samples were acquired as z-stacks of 3-µm optical sections or xz-stacks of 1.5-µm optical sections. Images were acquired as z-stacks of 3-µm optical sections acquired at 20-min intervals for all time lapses. Lack of pyknotic nuclei and frequency of mitoses in every acquired z-stack, as visualized directly with the aid of the Fucci reporter, confirmed good tissue health. For determination of cell cycle status and cell quantification, only cells that were distinctly identified as either G1 or S/G2/M were scored. All results represent at least three independent experiments.

Quantitative and statistical analyses of experimental data

Images and videos were analyzed and quantitative measurements performed with Imaris 7.2.1 (Bitplane) and ImageJ software. Images were processed for presentation with Photoshop CSS and Illustrator CSS software (Adobe Systems). Statistical analysis and further graphing were done with Prism 5 (Graphpad Software), PAST (http://folk.uio.no/ohammer/past/; Hammer et al., 2001), and SPSS Statistics (IBM) software. Differences between groups in cell volume and proliferation analyses were assessed with the parametric Student’s t test. For cell shape and condensation studies, measurements were done as described previously (Ahtiainen et al., 2014). In brief, all measurements were done in three dimensions from confocal optical stacks. For cell tracking purposes, to minimize the global motion of cells caused by tissue deformation or growth, the tissues were allowed to recover after dissection and were then mounted on a solid support. They did not, therefore, show significant amounts of rotational distortion, random vibration, or distortion in arbitrary planes, but rather a constant even movement in the yz plane (initial flattening and following growth in a plane perpendicular to the epithelial sheet) or a similar movement in the xz plane. To remove this background in cell tracking experiments, global movement of the tissue was subtracted from the movement of the cell population under observation in Imaris: at least four stationary points were defined as reference points (at least 100 µm away from the target cells), and global movement was corrected accordingly to these.

For cell track plots, at least 10 representative cells from at least three placodes were selected, and cell movement was plotted on axes representing displacement length in micrometers and relative angle. Cell movement (the angle from 0° to 359° at which the cell moved with Fluor 647–conjugated secondary antibody (BD and Invitrogen). Tissues were mounted with Vectashield (Vector Laboratories). All results represent at least three independent experiments, and the number of samples for each analysis is presented in the figure legends. Whole-mount X-gal staining was performed as described previously (Pispa et al., 2008). Whole-mount in situ hybridizations with digoxigenin-labeled probes for Foxa3, Sox2, Dkk4, Shh, and Wnt10b were performed as described previously (Wang and Shackleford, 1996; Fliniaux et al., 2008; Juuri et al., 2013; Shirokova et al., 2013). Fluorescent imaging was done with a ZEISS SteREO Lumar.V12 microscope, NeoLumar S 0.8x/WD 80-mm objective, and ZEISS Axioscam MRm3 CCD camera and X-gal staining with a SZZ9 stereomicroscope (Olympus) and ColorView IIIu CCD camera (Olympus). The Wnt10b expression domain in whole-mount specimens was quantified from mutants and littermate wt or age-matched wt controls with ImageJ software (National Institutes of Health).

Fluorescence confocal microscopy and time-lapse imaging

For 3D time-lapse imaging studies, dissected tissues were allowed to recover for 2 h. The explants were imaged as described previously (Ahtiainen et al., 2014) with a Leica Biosystems TCS SP5 microscope and HC PL APO 10x/0.4 (air) objective and, for fixed samples, in addition HCX PL APO 20x/0.7 Imm Corr (water, glycerol, oil) Ltd. bl and HCX APO 63×/1.30 Corr (glycerol) CS 21 objectives. Images of fixed samples were acquired as z-stacks of 3-µm optical sections or xz-stacks of 1.5-µm optical sections. Images were acquired as z-stacks of 3-µm optical sections acquired at 20-min intervals for all time lapses. Lack of pyknotic nuclei and frequency of mitoses in every acquired z-stack, as visualized directly with the aid of the Fucci reporter, confirmed good tissue health. For determination of cell cycle status and cell quantification, only cells that were distinctly identified as either G1 or S/G2/M were scored. All results represent at least three independent experiments.
respect to the mandible medial axis used as the 0/180° line of reference) was measured for individual cells and represented on a polar plot. For 4D cell migration tracking, track length and cell net displacement were analyzed. Box-and-whiskers plots represent minimum, 25th percentile, median, 75th percentile, and maximum values for each dataset.

For analysis of cell volume, the G1 condensate volume, S/G2/M total volume in tooth epithelium, or equal volume in the oral epithelium was masked, individual nuclei were counted, and mean single-cell volume was defined as total volume divided by cell number in the respective volume.

**Sox2 immunofluorescence staining**

For whole-mount immunofluorescence staining, tissues were fixed for 2 h in 4% PFA, permeabilized, and incubated with polyclonal goat anti-Sox2 (1:100; Santa Cruz Biotechnology, Inc.) and Alexa Fluor 647-conjugated secondary antibody (Invitrogen) overnight in 0.3% Triton X-100, 0.5% BSA, and 5% normal donkey serum in PBS. Tissues were mounted with Vectashield.

**EdU in vivo injection**

For the in vivo proliferation assay, pregnant K17-GFP:Fucci G1 dams were injected at E11.5, 12.5, and 13.5 with 25 mg/kg EdU (Invitrogen) in 7.5 mg/ml sterile saline solution 2 h before sacrifice. Tissues were fixed and processed as whole mount (time points E11.5 and 12.5) or embedded in paraffin (E13.5). EdU detection was done from whole-mount samples or frontal paraffin sections as described in the Click-IT EdU Alexa Fluor 674 Imaging kit protocol (Invitrogen).

**Individual cell size measurements**

Whole-mount immunofluorescence samples stained for E-cadherin (polyclonal rat antibody; 1:300; Invitrogen) were used to visualize cell perimeters, and nuclei were visualized with Hoechst 33342 staining. For measurements of individual cell sizes, high-resolution images in both xy- and z-axis were acquired as z-stacks (xy-scan) and xy-z-stacks (vertical scanning) with a 63× high numerical aperture HCX APO 63x/1.30 Corr (glycerol) CS 21 objective. From these optical stacks, individual G1 cell sizes were defined by one of two methods: (a) cell area was measured at the center of each cell inspected from consecutive optical sections in 3D, and the section that was within equal distance from cell perimeters in horizontal, vertical, and depth measurements was chosen; or (b) cell volume was defined by measuring height (major axis), width (vertical axis), and depth (minor axis) and defining the ellipsoid volume as \( \frac{4}{3} \pi \times (\text{major} \times \text{minor} \times \text{vertical}) \).

**Online supplemental material**

Fig. S1 shows the distribution of Fucci cell cycle indicator transgene-expressing cells in the developing mouse mandible: maximum-intensity projections of confocal fluorescence microscopy z-stacks of Fucci cell cycle reporter embryonic mouse mandible whole mounts at different stages of embryonic development from E10.0 to E13.5 (related to Fig. 1). Fig. S2 shows that Sox2-positive cells reside lingually to the early signaling center. Immunofluorescence staining of Sox2 in E12.5 Fucci G1 whole-mount samples confirmed that Sox2-positive cells are distinct from the early signaling center G1 cells and reside on the lingual side of the forming epithelial bud (related to Fig. 2). Fig. S3 shows cell condensation in the early signaling center: high-resolution imaging of E-cadherin–stained incisors showing reduction in G1 cell size from E11.5 to E12.5 (related to Fig. 3). Fig. S4 shows that early signaling center cells remain in G1, whereas budding morphogenesis takes place through local cell proliferation in the adjacent lingual cell population. Incorporation of EdU nucleoside analog delivered via in vivo injections of pregnant K17-GFP:Fucci G1 dams at E11.5, E12.5, and E13.5 shows G1 cells in the front part of the developing bud, whereas proliferating cells appeared on the lingual side. Still images from time lapse of Fucci E12.0 + 22 h mandible show mitoses highlighted in the bud adjacent to the G1 area on the lingual side (related to Figs. 4 and 5). Fig. S5 shows NF-kB reporter expression in the prospective incisor region. Whole-mount X-gal staining of mandibles at E11.0–E12.5 (related to Fig. 8). Video 1 shows the contribution of cells in different cell cycle phases to the forming incisor at different stages of development. 3D surface renderings show the contribution of G1 and S/G2/M cells at E11.5, E12.0, E12.5, and E13.5 (related to Fig. 3). Video 2 shows that the S/G2/M switch is initiated in the cell population adjacent to the early signaling center. Confocal microscopy time lapse of an E12.0 Fucci mandible imaged for 22 h; individual S/G2/M cells that go into cytokinesis are highlighted (related to Fig. 5). Video 3 shows that proliferating cells next to the G1 early signaling center form the incisor tooth bud. Confocal microscopy time lapse of an E12.5 Fucci mandible imaged for 10 h; 3D volume rendering overview is followed by a closeup of the left incisor, in which individual G1 and S/G2/M cell divisions have been followed (related to Fig. 5). Video 4 shows visualization of the early signaling center and the emergence of the enamel knot by Fucci G1 and TCF/Lef:H2B-GFP reporter. Confocal time-lapse microscopy of an E13.0 TCF/Lef:H2B-GFP, Fucci G1 mandible imaged for 14 h. The cells in the early signaling center are both G1 and GFP positive and remain in the labial part, whereas the EK is initiated at the posterior part of the developing bud (related to Fig. 7). Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201512074/DC1.

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