Activin is an essential early mesenchymal signal in tooth development that is required for patterning of the murine dentition

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Development of the mammalian tooth has been intensively studied as a model system for epithelial/mesenchymal interactions during organogenesis, and progress has been made in identifying key molecules involved in this signaling. We show that activin is expressed in presumptive tooth-germ mesenchyme and is thus a candidate for a signaling molecule in tooth development. Analysis of tooth development in activin mutant embryos shows that incisor and mandibular molar teeth fail to develop beyond the bud stage. Activin is thus an essential component of tooth development. Development of maxillary molars, however, is unaffected in the mutants. Using tissue recombination experiments we show that activin is required in the mesenchyme prior to bud formation and that although activin signaling from mesenchyme to epithelium takes place, mutant epithelium retains its ability to support tooth development. Implantation of beads soaked in activin A, into developing mandibles, is able to completely rescue tooth development from E11.5, but not E12.5 or E13.5, confirming that activin is an early, essential mesenchyme signal required before tooth bud formation. Normal development of maxillary molars in the absence of activin shows a position specific role for this pathway in development of dentition. Functional redundancy with activin B or other TGFβ family members that bind to activin receptors cannot explain development of maxillary molars in the mutants since the activin-signaling pathway appears not to be active in these tooth germs. The early requirement for activin signaling in the mesenchyme in incisor and mandibular molar tooth germs must be carried-out in maxillary molar mesenchyme by other independent signaling pathways.

[Key Words: Activin; tooth; follistatin; patterning]

Received January 22, 1998; revised version accepted May 28, 1998.

Development of the mammalian tooth has been intensively studied as a model system for epithelial/mesenchymal interactions during organogenesis (for review, see Thesleff et al. 1995; Thesleff and Sharpe 1997). Reciprocal epithelial–mesenchymal signaling has been shown to be involved at different stages of tooth development. Initiation of tooth germs has been shown to require epithelial signals, epithelial cell differentiation in tooth buds requires mesenchymal signals and signals between epithelial and mesenchymal cells are required for differentiation of ameloblasts and odontoblasts at later stages. Recently, progress has been made in identifying key molecules involved in these signaling pathways at different stages of early tooth development. Thus, the transcription factors, MSX1 and LEF1 have been shown to be required for tooth development to proceed beyond the epithelial bud stage with BMP4 being an important component of the signaling pathway (Satokata and Maas 1994; van Genderen et al. 1994; Chen et al. 1996). Signaling molecules belonging to the TGFβ, WNT, FGF, and HH families are also all expressed in cell-restricted fashions during different stages of tooth development and may play key roles in setting up patterning and initiating tooth development (Vainio et al. 1993; Heikinheimo et al. 1994; Bitgood and McMahon 1995; Iseki et al. 1996). In this paper we focus on the role of activin, a member of the TGFβ family of growth factors.

Activin proteins are produced from two gene-products, activin βA and activin βB which dimerize to form activin A (βA:βB), activin B (βB:βB) and activin AB (βA:βB). The closely related inhibins, inhibin A and inhibin B, are dimers consisting of an activin βA or βB subunit linked to an inhibin-specific α subunit (Vale et al. 1990; Roberts et al. 1991; Roberts and Barth 1994). Activin has been shown to have an essential role in mesoderm and neural induction in Xenopus development. Activin protein is
present in the egg and is later expressed zygotically in the Spemann’s organizer (Dohrmann et al. 1993). Micro-injection of activin mRNA causes partial axis duplication (Thomsen et al. 1990). Furthermore, treatment of prospective ectodermal tissue with activin causes the induction of mesodermal cell types (Asashima et al. 1990). Different concentrations of activin specify at least five different cell types, including the most dorsal mesodermal cell type, notochord (Green and Smith 1990; Green et al. 1992). The role of activin in early Xenopus development has also been demonstrated using over-expression of truncated activin receptors. Inhibition of activin signaling using these receptors causes a loss of mesodermal derivatives and an increase in neural differentiation (Hemmadi-Brivanlou and Melton 1992, 1994). Recently, a novel design of truncated receptor which, unlike previous designs, specifically inhibits activin but does not affect Vg1 or any other known mesoderm inducer, was also reported to disrupt mesoderm formation, thus confirming earlier findings (Dyson and Gurdon 1997).

Targeted mutation of either activin βA or βB genes or both genes together does not affect mesoderm development or neural differentiation in mice (Matzuk et al. 1995a), implying either that activin signaling has no obvious role in mesoderm and neural induction in the mouse or that there is functional redundancy with other as yet undiscovered activins, or other TGFβ family members. Nodal, a member of the TGFβ family has already been shown to be required for correct mesoderm formation in the mouse (Zhao et al. 1993; Conlon et al. 1994). A third possibility is that activin A produced in the maternally derived decidua is able to rescue embryonic development. However, the lack of defects in the mesoderm in activin receptor type II knockouts argues against this (Matzuk et al. 1995b).

We report here that activin βA has an essential role in tooth development, being required for development of incisor and mandibular molar tooth germs to progress beyond the bud stage. Development of maxillary molars in activin βA mutants is normal, producing normally mineralised teeth at birth. Thus, the essential role for activin in tooth development is restricted to specific teeth, suggesting a role for activin signaling in patterning of the dentition.

Here, we show that mandibular molar tooth development in mutant tissues can be rescued by addition of exogenous activin A into mesenchyme prior to bud formation or by recombination of mutant dental epithelium with wildtype dental mesenchyme. Development cannot, however, be rescued at the bud stage either by exogenous activin or by recombination of wild-type epithelium with mutant mesenchyme. From these experiments, we can conclude that activin A has an essential early role in cell signaling from presumptive tooth-bud mesenchyme at a time when it is believed that signals from the epithelium are essential for tooth development. We further show that the existence of this essential mesenchyme signal is not inconsistent with data showing that oral epithelium is instructive for tooth development at this stage because activin expression requires a signal, identified as FGF8, which is restricted to the oral epithelium to maintain its expression.

activin βA is normally expressed in the condensing mesenchyme surrounding the epithelial tooth bud in all developing teeth so the patterning phenotype cannot be easily explained by spatial differences in activin activity. We show here that follistatin is a downstream target of activin signaling and that its expression is down-regulated in the epithelium of all teeth in the mutants, including the maxillary molars, indicating that neither activin βA nor follistatin are required for maxillary molar tooth development and suggesting that maxillary molar development involves activin-independent signaling pathways.

Results

Expression of activin βA in early tooth development

Expression of activin βA was analyzed in the developing jaws of embryos between days 10.5 and 14.5 of gestation (E10.5–E14.5) using radioactive in situ hybridization. Weak expression could be first detected between E10.5 and E11 in mesenchymal cells of the mandibular process of the first branchial arch (data not shown). At this stage, no transcripts were detected in the maxillary process of the first branchial arch, which is slightly delayed in its development with respect to the mandibular process. However, activin βA expression was clearly visible at E11.5 in both the mandible and maxilla in the mesenchymal cells underlying the epithelial thickenings that will give rise to the future molars and incisors (Fig. 1A,B). At later stages activin βA-expressing cells comprised a distinct band of mesenchymal cells condensing beneath the epithelial tooth buds (Fig. 1C–H). In addition, activin βA expression could be detected in the mesenchyme surrounding the nasal epithelium and the palatal shelves (Fig. 1C,D).

Tooth development in activin βA and activin βA/βB double mutants

Frontal sections of newborn activin βA mutant mice were compared with those of heterozygous and wild-type littersmates (Fig. 2). The sections reveal the absence of both maxillary and mandibular incisors as well as mandibular molars in the mutants, the expected positions of these teeth being occupied by bone (Fig. 2, B,D,F vs. A,C,E). The maxillary molar teeth, however, were present in their correct positions in the mutants and had a normal histological appearance (Fig. 2 E,F). The tooth phenotype in activin βA mutants thus represents a patterning abnormality, where development of only certain teeth are affected. Patterning of the dentition is also affected in Dlx1/1/2 double mutants that have a reciprocal phenotype of activin βA mutant embryos, where only development of maxillary molars is affected (Qiu et al. 1997; Thomas et al. 1997).

To establish whether odontogenesis of incisors and mandibular molars is initiated in the mutants, frontal
sections of mutant embryos at E12.5 (early bud stage of tooth development), E13.5 (late bud stage), and E14.5 (cap stage) were examined. Development of maxillary incisors and all mandibular teeth was found to be initiated but arrested at the bud stage, whereas maxillary molar tooth germs were present at the correct stage in all of the embryos examined (Fig. 3). The mesenchymal component of the mutant teeth appeared to be unaffected at a histological level, since condensations of mesenchymal cells were found to underlie the epithelial buds as in wild-type tooth buds.

To investigate whether activin βB activity might compensate for the loss of activin βA within the unaffected upper molars in the mutants, the newborn phenotype of activin βA/βB double-knockout mice was analyzed. As in the activin βA mutants, the maxillary molars were present and had a normal histological appearance (Fig. 2G). All other teeth were absent, showing that there is no functional redundancy between the two activins for maxillary molar development.

Rescue of tooth development by exogenous activin activin βA expression first becomes clearly visible in presumptive tooth mesenchyme at E11.5 and, because tooth development is arrested in the mutants at the bud stage (E13.5), the requirement for activin must be between these times. To pinpoint the time when activin is required, beads soaked in activin A protein were implanted into the mesenchyme of mutant mandible explants at E11.5 and E13.5. In situ hybridization was used to confirm that activin is unable to induce its own expression. Therefore the bead was acting as the only source of activin in these experiments (Fig. 4F,G). Explants were cultured for 24–72 hr and then transplanted into kidney capsules. Table 1 shows the numbers of teeth obtained, and Figure 4, B–E, shows typical histology of the resulting tissue. At E11.5 6 molars were formed from a total of 12 explants with implanted activin beads, whereas at E13.5 no teeth were formed.

Figure 1. The expression pattern of activin βA during early tooth development. Radioactive in situ-hybridization on frontal sections showing activin βA expression in the developing jaws of embryos at E11.5 (A,B), E12.5-13 (C–E), and E14.5 (F–H). Tooth germ epithelium is outlined in red. (A,B) Teeth at the epithelial thickening stage of development. The arrow indicates the lower incisor region. (B) A more proximal section than shown in A, cut through the molar region. (C–E) Teeth at the early bud stage of development, showing expression in the upper incisors (C), lower incisors (D), and molars (E). The arrow in C indicates the nasal process; the arrow in D indicates the palatal shelves. (F–H) Teeth at the late bud stage of development, showing expression in the upper incisors (F), lower incisors (G), and molars (H).

Figure 2. The tooth phenotypes of wild-type, activin βA mutant, and activin βA/βB mutant newborn mice. Frontal sections show that upper and lower incisors are present in wild-type mice (A,C), but missing in activin βA mutants (B,D). Maxillary and mandibular molars are present in wild-type mice (E); mandibular molars are missing in both the activin βA (F) and activin βA/βB mutants (G). (inc) Incisor; (mb) mandibular bone; (Mc) Meckels’ cartilage; (ns) nasal septum; (tg) tongue.

Figure 3. The tooth phenotype of activin βA heterozygous and homozygous mutant embryos at E14.5. (A) Frontal sections showing molar tooth germs at the cap stage of development in activin βA heterozygotes. (B) In activin βA homozygous mutant embryos the upper molars are at the cap stage, whereas the lower molars are arrested at the bud stage. Note condensation of mesenchyme still occurs around the buds in the mutant (arrow).
stead of teeth, the four explants gave rise to four keratinizing epithelial cysts embedded in alveolar bone. The control experiments showed that implanting BSA beads had no effect on the explants, the mutant explants giving rise to cysts and wild-type explants giving rise to teeth (see Table 1). Exogenous activin thus enables completion of tooth organogenesis and terminal cytodifferentiation in mutant mesenchyme when delivered at E11.5 but is unable to rescue at E13.5. These results indicate that activin is required at the epithelial thickening stage, but before the full bud stage of tooth development.

When performing these experiments we observed that although there is no visible difference between wild-type and mutant embryos in the appearance of the epithelial thickenings at E11.5, there is a clear difference in morphology at E13.5. At this stage, the depth of the epithelial invaginations that form a bud is greatly reduced in the mutant compared to those in wild-type embryos (data not shown). This suggests that the tooth defect in the mutants is present by E13.5 and therefore too late to be rescued. To determine whether there was any degree of rescue on addition of activin beads at E12.5, beads were implanted at E12.5 and E13.5 which were then cultured for 6 days and sectioned. Of the 12 mutant tooth primordia (incisors and molars) examined, which had access to an activin bead, all had tooth germs arrested at the bud stage, whereas tooth primordia in wild-type control cultures all reached the cap stage (Table 2; Fig. 5). These results indicate that rescue of the mutant phenotype is restricted to E11.5, as addition of beads at E12.5 and E13.5 is too late to rescue tooth development.

Table 1. Teeth formed in kidney capsules by mutant tooth anlagen following implantation of activin-soaked beads

| No. of teeth | E11.5  | E13.5 |
|--------------|--------|-------|
| Mutant molar + activin A bead | 6/12  | 0/4   |
| Mutant molar + BSA bead        | 1*/6  | 0/5   |
| Wild-type molar + BSA bead    | 9**/6 | 5/6   |

(*) A small misshapen molar developed, reflecting that the mutant phenotype of absent mandibular molars is not quite 100% penetrant. (**) The appearance of primary and secondary molars from primary tooth anlagen accounts for the high number of teeth obtained from these explants.

Figure 4. Tooth development in activin βA mutant molar primordia treated with exogenous activin A protein at E11.5 and E13.5. (A) Schematic diagram showing procedures involved in the bead rescue experiments. Mandibular molar anlagen were dissected away from surrounding oral and aboral tissues. Beads soaked in activin A protein or BSA were embedded in the dental mesenchyme in close proximity to the invaginating dental epithelium. (i) Incisor; (m) molar. (B–E) Representative sections of E11.5 (B,C) and E13.5 (D,E) explants after culture as subrenal grafts. Note that beads became dissociated from explants during transfer to the kidneys. (B) Fully developed molar resulting from the "rescue" of E11.5 mutant molar tissue cultured in the presence of an activin A bead. Note the cytodifferentiation of odontoblasts (od) secreting dentine (de) and ameloblasts (am). (C) A keratinizing-epithelial cyst arising from E11.5 mutant molar tissue cultured in the presence of a BSA bead. (D) Activin A failed to rescue E13.5 mutant molar anlagen, resulting in keratinizing epithelial cysts. (E) Control E13.5 explants cultured in the presence of BSA beads gave rise to keratinizing epithelial cysts. Note that the tissues shown in B–E were removed from surrounding alveolar bone before being processed for routine histology. (F,G) Control experiment to show that activin does not induce its own expression in the mesenchyme. Activin beads were added to the mesenchyme of mandible explants at E11.5. After 2 days in culture the explants were fixed, sectioned, and analyzed using radioactive in situ hybridization. (F) Light-field photomicrograph showing the position of the activin bead in the mesenchyme. Incisor tooth buds are marked by arrows. (G) Corresponding dark-field photomicrograph showing expression of activin βA beneath incisor buds, but not around the bead.
Activin is unable to rescue the tooth development when added to mutant tooth primordia at E12.5. (Figure 5.)

Recombination experiments carried out by several different groups have established that between E10.5 and E12, tooth development can only be initiated by oral epithelium (Kollar and Baird 1969; Mina and Kollar 1987; Lumsden 1988). The existence of an essential mesenchymal signal at this time therefore appears inconsistent with this data. To resolve this we investigated the expression of activin βA in the absence of epithelium by culturing mandibles explanted at E11.5 for 1–2 days with the epithelium removed. Removal of epithelium resulted in loss of activin βA expression indicating that signals from the epithelium are required to maintain expression (Fig. 7C,D). A likely candidate for such an epithelial signal is FGF8, which is expressed in oral epithelium between E9 and E13 (Heikinheimo et al. 1994; Crossley and Martin 1995). In Figure 7, A and B, consecutive serial sections of an E11.5 embryo show the expression of Fgf8 in the epithelium and activin βA expression in the adjacent underlying mesenchyme. Thus, the coexpression of Fgf8 and activin βA at this time is consistent with FGF8 being a candidate molecule involved in the regulation of activin βA. The ability of exogenous FGF8 to maintain activin βA expression was therefore assayed by implanting beads into mesenchyme explants without epithelium and culturing for 1 day. In situ hybridization revealed activin βA expression around the FGF8 beads but not around control BSA beads, indicating that FGF8 secreted from the epithelium is a likely endogenous signal for maintenance of activin βA expression (Fig. 7E,F). The requirement for activin as an essential early mesenchymal signal is thus consistent with the instructive role of oral epithelium for tooth germ initiation at this stage, as signals from the epithelium are required to maintain activin expression in the mesenchyme.

Recently, Neubüser et al. (1997) showed that antagonistic interactions between FGF8 and BMP4 serve to regulate the expression of Pax9, which encodes a member of the family of paired-box-containing transcription factors and is expressed in the odontogenic mesenchyme from E10. Given the similar expression patterns of activin βA and Pax9 (e.g., cf. Figs. 1E and 8D, which are consecutive sections from a wild-type embryo at E12.5–E13), we were prompted to investigate whether BMP4 had any effect on activin βA expression. The action of BMP4 on activin βA expression in E11.5 explants was tested in three ways: (1) To test whether BMP4 has an inhibitory action, BMP4 beads were implanted into

Table 2. Cap stage tooth germs observed following implantation of activin or control BSA beads

| N. of tooth germs | −/− | +/+ |
|-------------------|-----|-----|
| + Activin bead    | 0/12| 10/12|
| + BSA bead        | 0/4 | 7/8  |

Implantation at E12.5 and culture for 6 days.

Tooth development is dependent on activin signaling within the dental mesenchyme.

Because exogenous activin could not rescue tooth development at E12.5 and E13.5, this meant that the essential activin signaling identified at E11.5 must be inducing changes that are required for progression beyond the bud stage. To determine which tissue was being targeted by the activin signal, epithelial–mesenchymal recombinations were carried out between mutant and wild-type tissues. Mandibular molar region oral epithelium was removed and recombined with mesenchyme from the same region at E13.5. Mutant epithelium was recombined with wild-type mesenchyme and vice versa, and the explants cultured for 24–48 hr before being transplanted under kidney capsules. Table 3 shows the numbers of teeth obtained and Figure 6 shows typical morphology. The combination of wild-type mesenchyme and mutant epithelium resulted in eight morphologically normal teeth from nine explants. Wild-type epithelium and mutant mesenchyme produced eight normal teeth from eight explants. Mandibular molar region oral epithelium was therefore assayed by implanting beads into mesenchyme explants without epithelium and culturing for 1 day. In situ hybridization revealed activin βA expression around the FGF8 beads but not around control BSA beads, indicating that FGF8 secreted from the epithelium is a likely endogenous signal for maintenance of activin βA expression (Fig. 7E,F). The requirement for activin as an essential early mesenchymal signal is thus consistent with the instructive role of oral epithelium for tooth germ initiation at this stage, as signals from the epithelium are required to maintain activin expression in the mesenchyme.

Regulation of activin βA expression

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Figure 5. Activin is unable to rescue the tooth development when added to mutant tooth primordia at E12.5. (A) Representative section showing a cap stage incisor in a wild-type explant after culture in the presence of an activin A-loaded bead. (B,C) Representative sections of bud stage incisors in mutant explants after 6 days of culture in the presence of activin A beads (B) and BSA beads (C). Tooth germ epithelium is outlined in white. (VL) Vestibular lamina.
with epithelium removed; and (3) to test whether BMP-4 planted alongside FGF8 beads into mandible explants the maintenance activity of FGF8, BMP4 beads were intact mandibles; (2) to test whether BMP4 could inhibit activin βA expression: BMP4 could neither maintain or inhibit activin βA expression (data not shown) nor inhibit the maintenance activity of FGF8 (Fig. 7G,H).

Table 3. Teeth formed in kidney capsules after recombination of E13.5 dental epithelium and mesenchyme

| Type                | Teeth Formed |
|---------------------|--------------|
| Wild-type epithelium| 11/10        |
| Wild-type mesenchyme| 8/9          |
| Mutant epithelium   | 1*/9         |
| Wild-type mesenchyme| 0/2          |

(*) The mutant phenotype is not 100% penetrant, which was confirmed by control cultures. The incisors of the mice were cultured intact and then transferred to kidney capsules as control cultures. Five epithelial cysts and one tiny, underdeveloped incisor were retrieved from the five pairs of mutant incisors explanted.

Gene expression in activin βA mutant teeth

To identify genes downstream of activin signaling in tooth development, the expression of well-characterized mesenchymal and epithelial marker genes was examined in mutant embryo teeth. The expression patterns in mutant mandibular molars were compared with those in the normal maxillary tooth germ. Expression in the mutant incisors was compared with expression in incisors of wild-type or heterozygous littermates (Fig. 8). Radioactive in situ hybridization was performed on wild-type, heterozygous, and homozygous mutant embryos up to the bud stage. The expression patterns of genes examined included Barx1, Mxs1, Dlx2, Pax9, Gli3, Lef1, syndecan-1, Tgfβ1, Tgfβ3, Bmp4, Bmp7, Sonic hedgehog (Shh), CD44, and Otx2.

In wild-type teeth at the bud stage, Barx1 gene expression is restricted to the molar region of the mandible and maxilla but is present in a broad field of neural crest derived mesenchymal cells rather than being restricted to dental mesenchyme (Fig. 8A; Tissier-Seta et al. 1995). Mxs1, Lef1, and Bmp4 are expressed in the dental mesenchyme (i.e., the condensing mesenchymal cells associated with invaginating incisor and molar epithelial tooth buds) in response to epithelial signaling (Fig. 8B; Jacobson et al. 1991; Vainio et al. 1993; Kratochwil et al. 1996). It is thought that these genes are then involved in reciprocal signaling from mesenchyme to epithelium, which is required for epithelial cell differentiation including the formation of the enamel knot. Dlx2 expression is restricted to mesenchymal cells immediately surrounding the epithelial bud, but is also present in the dental epithelium on the buccal side of the buds (Fig. 8C; Thomas et al. 1995; Qui et al. 1997). Pax9 is expressed in early tooth mesenchyme prior to bud formation and subsequently in condensing mesenchyme at the bud stage (Fig. 8D; Nübler et al. 1997). Gli3 is expressed in the mesenchyme from E10.5. At the bud and cap stage Gli3 expression is slightly more localized than Pax9 expression, and is restricted to the dental papilla and dental follicle (Fig. 8E; Hardcastle and Sharpe 1998). Syndecan-1, a cell surface heparin-sulfate proteoglycan is transiently expressed in the dental mesenchyme and is thought to regulate dental mesenchymal cell condensation beneath the invaginating dental epithelium (Fig. 8G; Thesleff et al. 1996). Tgfβ1 is found in the dental mesenchyme and weakly in the epithelium of the incisors and only appears in the molars in the dental epithelium at the cap stage (Fig. 8H; Vahtokari et al. 1991). Tgfβ3 expression is widespread in the mesenchyme of the face but, interestingly, its expression appears to be excluded from the condensin mesenchymal cells immediately adjacent to the epithelial buds of incisors and molars (Fig. 8I; Chai et al. 1994).

Analysis of genes known to be expressed in tooth germ epithelium included Bmp7, Shh, CD44, and Otx2 genes. In wild-type embryos, Bmp7 is initially expressed in the dental epithelium, but expression shifts to the mesenchyme around the tooth buds from E13.5 (Åberg et al. 1997). At E13.5 mesenchymal Bmp7 expression was

Figure 6. Tooth development in recombinations of dental tissues from wild-type and activin βA mutant embryos at E13.5. (A) Schematic diagram illustrating the procedures involved in the tissue recombination experiments. (Black) Wild-type tissue; (white) mutant tissue. (E) Dental epithelium; (M) dental mesenchyme. (B) Representative molars derived from the combination of mutant epithelium with wild-type mesenchyme. (C) Typical cysts derived from the combination of wild-type epithelium with mutant mesenchyme. Histological analysis of these cysts revealed keratinizing stratified epithelium, as shown in Figs. 4C–E.
homolog of the human gene, which when mutated, causes the disease known as Rieger syndrome in which teeth are absent (Semina et al. 1996).

Expression of these genes was found to be normal in activin βA mutant tooth buds (Fig. 8A–N), showing that at the level of transcription, the essential role for activin in tooth development does not involve any of these molecules.

Role of follistatin in tooth development

Follistatin is an activin-binding protein that has been shown to inhibit the activity of activin (Michel et al. 1993; De Winter et al. 1996). To assess whether follistatin might interact with activin during early tooth development, the expression patterns of follistatin and activin βA were compared on consecutive frontal sections of wild-type embryos by in situ hybridization analysis (cf. Figs. 1, C, D, and E, and 9, A, C, and E). Follistatin expression was found in tooth germ epithelial cells immediately adjacent to activin βA-expressing cells from E11.5 (data not shown). At later stages, follistatin expression was found to be restricted to the columnar-shaped cells that form the outermost layer of the epithelial bud, while the central core of epithelial cells were follistatin-negative (Fig. 9 C,E). Follistatin was also found to be expressed in the nasal epithelium, in the epithelium of the palatal shelves, and strongly in the tongue.

Because follistatin is expressed in the tooth epithelium adjacent to and in a complementary pattern to activin βA in the tooth mesenchyme, it is a potential target of activin signaling. The expression of follistatin was therefore analyzed in activin βA mutant embryos and compared with wild-type embryos. No expression could be detected in the tooth epithelium of the mutants, or in the nasal epithelium and the palatal shelf epithelium, but normal levels of expression were evident in the tongue where activin is not strongly expressed. Loss of expression in the dental epithelium was found in all teeth (Fig. 9B,D,F), indicating that follistatin is not required for maxillary molar tooth development.

Activation of follistatin expression by activin

The epithelial expression of follistatin is down-regulated in activin βA mutant embryos which indicates that follistatin expression is downstream of activin signaling in developing teeth. In Xenopus, activin has been shown to activate the expression of follistatin in an animal cap assay (Hemmati-Brivanlou et al. 1994). To investigate whether activin can regulate follistatin expression in epithelium derived from the first branchial arch in the mouse, epithelium was isolated from E11.5 mandibles and cultured for 24 hr in the presence of beads soaked in activin A protein. Explants were fixed, sectioned and examined for follistatin expression using in situ hybridization. Follistatin was found to be expressed around the activin beads, whereas no expression was detected around BSA control beads (Fig. 10A–E). The induction of
follistatin expression by activin was confirmed as being specific to the epithelium by implanting activin beads into intact mandible explants at E11.5, which showed that follistatin expression was not induced in the mesenchyme (data not shown).

To confirm that activin is an endogenous inducer of follistatin expression we recombined activin A mutant epithelium (follistatin-negative) with wild-type mesenchyme (activin-positive). follistatin expression was found to be induced in the mutant epithelium indicating that activin A is a likely endogenous factor responsible for induction of follistatin gene expression in dental epithelium (Fig. 10F,G). These results demonstrate a mesenchyme to epithelium component to the activin signaling pathway.

Discussion

We have analyzed the expression patterns of activin βA and its antagonist follistatin during the early stages of odontogenesis using radioactive in situ hybridization. We detected weak expression of activin βA in presumptive tooth mesenchyme between E10.5 and E11 and strong expression of both activin βA and follistatin at E11.5 in developing tooth germs. Our results demonstrate earlier expression in the branchial arches than those of Fälgen et al. (1994), who mapped the expression of activin βA and follistatin in postimplantation mouse embryos between E6.5 and 12.5. They detected the first expression of activin βA in the dental mesenchyme at E12.5 but did not report the presence of follistatin transcripts in the tooth epithelium at this stage. Recently, Heikinheimo et al. (1997) reported the expression of activin βA and follistatin in the developing molars of embryos between E14 and E19. They suggested a role for the activin-follistatin pathway in the terminal differentiation of odontoblasts. Our results support an earlier and more fundamental role for the activin signaling pathway in the patterning and morphogenesis stages of tooth development.

The restricted expression of activin βA in presumptive tooth mesenchyme and the induction of follistatin in tooth epithelium suggested a role for this signaling pathway in epithelial-mesenchymal interactions known to occur during odontogenesis. We examined tooth development in activin βA mutant mice and found that incisor and mandibular molar tooth development was arrested at the bud stage but, surprisingly, that maxillary...
molar teeth developed normally. This phenotype indicates a role for activin signaling in patterning of the dentition, being required for development of incisors and mandibular molar teeth but not of maxillary molars. This is the second tooth patterning phenotype to be described in mutant mice. Mutations in Dlx1 and Dlx2 homeobox genes result in specific arrest of maxillary molar development but incisors and mandibular molars are normal (Thomas et al. 1997). Thus, the tooth phenotype in the activin bA mutants is the reciprocal of the Dlx1/2 mutant phenotype. These phenotypes differ from those of Msx1, Lef1, and Pax9 mutant mice (Satokata and Maas 1994; Kratochwil et al. 1996; Neubüser et al. 1997), in which all of the teeth are affected. Activin bA, like DLX1 and DLX2, appears to have an essential role in the patterning of the dentition.

There are thus two aspects of activin signaling in tooth development that require explanation: an essential role in development of incisors and mandibular molars so that their development proceeds beyond the bud stage and a non-essential role in development of maxillary molars.

Activin is an early mesenchymal signal

To address the role of activin signaling in development of incisors and mandibular molars, we used bead rescue and recombination experiments to identify the stage at which activin signaling is critical, and which tissues are responsive to activin signaling. It is well established that tooth development is controlled by interactions between epithelial and mesenchymal cells (for review, see Thesleff and Sharpe 1997). Oral epithelium was originally identified as the source of tooth inducing signals from tissue recombination experiments in which early oral epithelium (taken at stages prior to tooth bud formation) was shown to induce tooth development when recombined with non-oral mesenchyme, as long as the mesenchyme was neural crest-derived. The reciprocal recombination of non-oral epithelium with mandibular arch mesenchyme did not produce teeth (Mina and Kol lar 1987; Lumsden 1988). Thus between E10.5 and E11.5,
signals pass from the dental epithelium to the underlying mesenchyme to stimulate tooth development. By the early bud stage (E12) the direction of interactions is reversed and signals pass from the mesenchyme to the tooth germ epithelium. The detection of activin βA expression in presumptive tooth mesenchyme between E10.5 and E11.5 and its ability to induce follistatin expression in tooth germ epithelium suggested activin was involved in a key early mesenchyme-to-epithelium signaling event. This was investigated by transient application of activin A-soaked beads to the mesenchyme to determine whether mutant molar tooth development could be rescued. Remarkably, we found that the program of tooth development could be rescued completely from E11.5 by a single activin-soaked bead, but not from E12.5 or E13.5, at which times activin beads fail to rescue progress beyond the bud stage. These experiments show clearly that activin is required for mandibular molar tooth development before E12.5 and that these developing teeth reach the early bud stage in the absence of activin βA, the cannot be rescued subsequently. This suggests that activin might be signaling to the epithelium at E11.5 and inducing a response that permits development to proceed to the bud stage. Therefore, if activin was signaling to the epithelium we expected wild-type E13.5 epithelium to support tooth development beyond the bud stage when recombined with activin βA mutant mesenchyme and that no rescue would be seen with mutant epithelium, which would not have received a signal, when recombined with wild-type mesenchyme at E13.5. However, wild-type epithelium was unable to rescue tooth development, whereas the recombinations including mutant epithelium were able to rescue tooth development. These results, together with the rescue achieved by implanting activin beads at E11.5 but not at E12.5 or E13.5, lead to several possible interpretations of the mechanism of the early essential activin signaling. Either activin signals between mesenchymal cells or activin signals to the epithelium and there follows an unknown reciprocal signal back from the epithelium to the mesenchyme by E13.5, or signaling involves both mesenchyme-to-mesenchyme as well as mesenchyme-to-epithelial elements. Analysis of the expression patterns of activin receptors IIA and IIB in E11.5 embryos reveals low levels of expression in both epithelium and mesenchyme (data not shown), indicating that the activin signal can be transduced in both of these tissues.

Nevertheless, as far as tooth development is concerned, the results obtained appear more consistent with an essential role for activin in mesenchymal–mesenchymal cell interactions rather than mesenchymal–epithelial cell interactions. To elaborate, epithelium from activin mutants at E13.5 is able to support tooth development when recombined with wild-type mesenchyme. Thus, if essential mesenchyme-to-epithelial signals are involved before E13.5, these signals must have no effect on the ability of epithelium to support tooth development. Moreover, it seems unlikely that such an elaborate reciprocal signaling exchange would be active with no discernible affects on the epithelium.

There is no doubt, however, that activin does signal to the epithelium at E11.5 to induce expression of follistatin. Because activin is continuously produced by tooth mesenchymal cells it needs to be continuously removed for a temporal sequence of signaling to continue to be instructive. Expression of follistatin in adjacent epithelial cells may thus act as a continuous “sink” for activin protein to remove it from the mesenchyme.

The question of whether activin plays a role in signaling to the epithelium after E13.5 in addition to its essential early signaling is not addressed by the bead rescue data. It is unlikely, however, that such late signaling is important for tooth development. We have shown that the implanted beads act as the only source of activin in the rescue experiments because exogenous activin does not cause the induction of its own expression. It is improbable, therefore, that the teeth derived from mutant explants in the bead rescue experiment at E11.5 developed because of a sustained activin signal throughout the period of culture. The stability of the activin protein in a mandible explant is difficult to assess experimentally, but it is unlikely that the activin protein diffusing from a single activin bead implanted at E11.5 could remain stable and sustain tooth development for 12 days in culture. Furthermore, because we have shown that activin βA induces follistatin expression in the epithelium, follistatin would subsequently bind activin and further reduce its activity in the explants.

A large number of genes have been identified as being expressed at the bud stage of tooth development in either epithelial or mesenchymal cells, or both (http://www.honeybee.helsinki.fi/toothexp). Using in situ hybridization we have shown that expression of a number of the most significant of the mesenchymal and epithelial expressed genes is unaffected in activin βA mutant tooth germs. Thus, the expression of Msx1, Lef1, Bmp4, Dlx2, and Pax9, all of which have been shown to have an essential role for tooth development, is the same in tooth bud mesenchyme of activin βA mutant incisor and mandibular molars as in wild-type buds and the unaffected maxillary molar buds in the mutants. Therefore, at a transcriptional level, none of these genes is involved in the activin signaling pathway downstream of activin A binding to its receptors, and this expression analysis thus gives no further clue as to whether activin is signaling within the mesenchyme or via the epithelium. Clearly, activin signaling acts on as yet unidentified molecules in the odontogenic tissues, which points to the existence of a previously unidentified pathway acting within the developing teeth.

We believe the simplest explanation that is consistent with the results is that activin is an essential early mesenchymal signal acting at E11.5 and before E12.5, which is required to induce changes in the mesenchyme before E13.5 so that tooth development can proceed beyond the early bud stage.

The finding that the maintenance of activin βA expression requires signals from oral epithelium is consistent with the early instructive role of oral epithelium for tooth development. The identification of FGF8 as a fac-
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tor that is able to substitute for oral epithelium in maintaining activin expression adds activin to a growing list of genes, including Pax9, Lhx6, Lhx7, and Msx1 expressed in facial mesenchyme that are regulated by FGF8 (Neubüser et al. 1997; Grigoriou et al. 1998; Kettunen and Thesleff 1998).

The role of activin in dental patterning

There are two possible explanations for why maxillary molar teeth develop in the absence of activin. The most obvious possibility is that activin is involved in development of all teeth and in its absence another TGFβ-like signaling molecule compensates for loss of activin βA in maxillary molar mesenchyme. The gene product of activin βA contributes to homodimeric activin A, heterodimeric activin AB and inhibin A, which are missing in the activin βA mutants. Activin βA could be functionally redundant for maxillary molar development with another TGFβ-family member. The most likely candidates for this role would be activin/hibinin or TGFβ-family molecules that are able to bind to activin receptors and thus activate the same downstream genes as activin. There are three pieces of evidence that suggest this is an unlikely explanation. First, activin βB activity is unable to compensate for activin βA within the unaffected upper molars in the mutants because in activin βA/βB double-knockout mice maxillary molar development is normal (Fig. 2G). Absence of both βA and βB subunits abolishes all activin and inhibin activity in these mice. Second, we found that epithelial expression of the follistatin gene was down-regulated in all teeth in activin βA mutant embryos, but expression was unaffected in some other facial tissues such as the tongue where activin βA is not expressed (Fig. 9). Because follistatin gene expression is lost in vivo in the unaffected maxillary molars of the mutants, we conclude that it is not required for maxillary molar development. This implies that maxillary molar development would be expected to be normal in follistatin mutant mice, and examination of follistatin mutant embryos confirms this (Matzuk et al. 1995c; M. Matzuk and P.T. Sharpe, unpubl.). A role for follistatin in tooth development may only be evident in the presence of activin A because follistatin functions to inhibit activin. Follistatin mutant embryos do have a weakly penetrant lower incisor development phenotype that is dependent on genetic background (Matzuk et al. 1995c). The fact that follistatin expression is downstream of activin, yet is absent from unaffected maxillary molars in the mutant, suggests that the mesenchyme-to-epithelium component of the activin signaling pathway is not activated by a compensatory molecule in the maxillary molars of mutants. This does not exclude the possibility that the mesenchymal activin pathway is activated by such a molecule. Finally, preliminary analysis of activin type IIA and IIB receptor double mutant embryos indicates that loss of these receptors can produce a similar tooth patterning phenotype to loss of activin βA (P. Oh, E. Li, and P.T. Sharpe, unpubl.).

An alternative possibility to explain why maxillary molars are unaffected in activin βA mutants is that development of these teeth is fundamentally different from all other teeth and does not require activin. Development of maxillary molar teeth certainly appears to involve a pathway that is independent of other tooth development. Mutations in the Dlx1 and Dlx2 homeobox genes result in a tooth phenotype where maxillary molars fail to develop, but all other teeth develop normally (Qiu et al. 1997; Thomas et al. 1997). Thus, this phenotype is the reciprocal of the activin βA mutant phenotype. It is possible that DLX1 and DLX2 may activate the same essential genes that are downstream in the activin pathway in maxillary molars, thus negating the need for activin during the development of these teeth. In turn, maxillary molar tooth development might be fundamentally different from that of all other teeth.

Materials and methods

Production and analysis of mutant embryos

Activin βA mutant and activin βA/βB double mutant embryos were produced as described by Matzuk et al. (1995a) using homologous recombination in ES cells. Animals were maintained as heterozygotes and mated to generate homozygous mutant embryos. Day E0 was taken to be midnight prior to finding a vaginal plug. Embryos were harvested at the appropriate time and genotyped using PCR and Southern blot analysis of genomic DNA extracted from unused embryonic or extraembryonic tissue. The following primers were used for the PCR assays: 5’-TGATGAAAGCCGTCGGTTAA-3’ and 5’-TTTTACA-GACATTATTGCTGCTGCT-3’, which amplified a 1.8-kb fragment containing exon 2 in wild-type and heterozygote littersmates. When PCR was used, the embryos that were identified by the absence of the 1.8-kb PCR fragment were classified as potential homozygous mutants. The genotypes were then confirmed by Southern blot hybridization as described.

For basic histology embryos were fixed in 4% paraformaldehyde, wax embedded, serially sectioned, and stained with malachite green. Newborn mice were decalcified in 0.5M EDTA (pH 7.6) following fixation, double embedded, that is, by use of a combination of paraffin wax and nitricelulose as embedding media according to Page (1990), serial sectioned, and stained with alcan blue/chloroquine fast red.

In situ hybridization

Radioactive in situ hybridization procedures were carried out on embryos at E10.5–E14.5 as described by Wilkinson (1995). The radioactive antisense probes used were generated from mouse cDNA clones that were gifts from several laboratories: activin βA and follistatin (A.J.M. van den Eijnden-van Raaij, Hubrecht Laboratory, Netherlands Institute for Developmental Biology, Utrecht, The Netherlands), Barx1 (J.-F. Brunet, Developmental Biology Institute of Marseille, France), Bmp4 (B. Hogan, Vanderbilt University Medical School, Nashville, TN), Bmp7 (V. Rosen, Genetics Institute, Inc., Cambridge, MA), CD44 (C. Isacke, Imperial College of Science, Technology, and Medicine, London, UK), Dlx2 (J. Rubenstein, University of California, San Fransisco), Fgf8 (I. Mason, United Medical and Dental Schools, Guy’s Hospital, London, UK), Gli3 (C.C. Hui, Harvard University, Cambridge, MA), Otx2 (C. Goridis, Developmental Biology Institute of Marseille), Pax9 (R. Balling, Institut für Säuger-
Activin bead rescue experiments

Mandibles from embryos at E11.5, E12.5, and E13.5 were dissected in DMEM with glutamak-1 (GIBCO BRL). The rest of the embryo was used for genotyping. For cultures at E11.5 and E13.5, individual molar tooth anlagen were isolated from surrounding oral and aboral tissue, whereas for cultures at E12.5, whole mandibles were used. These were placed with oral surfaces facing upwards, on membrane filters supported by metal grids following the Trowell technique as modified by Saxén (Trowell 1959; Saxén 1966). Affi-gel agarose beads (Bio-Rad) were washed several times in PBS and then dried before being added to either recombinant activin A protein (gift from Mike Jones, NIMR, London, UK) at 1 µg/ml, a concentration that induced mesoderm formation in Xenopus animal cap assays (not shown), or BSA at the same concentration for 1 hr at 37°C. The beads were pushed into the mesenchyme so that they lay in close proximity to the developing tooth germs as indicated in Figure 4A. The mandible explants at E11.5 and E13.5 were cultured with FGF8/BMP4 bead experiments

Mandibles were dissected at E11.5. Where indicated, epithelium was removed after incubation in dispase (2 U/ml) for 10 min at 37°C. For the application of FGF8 protein, heparin acrylic beads (Sigma) were washed and then incubated overnight in 1 mg/ml FGF8 protein (recombinant mouse FGF8b; R&D Systems, Abingdon, UK) at 4°C. For the application of BMP4 protein, Affi-gel agarose beads were washed and soaked in the protein (recombinant human BMP4; gift from the Genetics Institute, Cambridge, MA) at 1 µg/ml for 1 hr at 37°C. A concentration of 100 µg/ml BMP4 was used, as this concentration has been shown to inhibit Pax9 expression in a similar assay (Nebüser et al. 1997). After 24–48 hr in culture the explants were fixed and processed for in situ hybridization. Digoxigenin (DIG) whole-mount in situ hybridization was carried out as described by Pownall et al. (1996).

Epithelial cultures

Oral epithelium was isolated after incubation of E11.5 mandibles in Dispase (2 U/ml) for 10 min at 37°C. The epithelium was cultured in Matrigel (Collaborative Biomedical Products) on membrane filters supported by metal grids as described above and indicated in Figure 10A. Matrigel is a solubilized basement membrane extracted from the Engelbreth-Holm-Swarm mouse sarcoma cell line and provides a matrix in which epithelial cells can develop. The gel sets rapidly and irreversibly at temperatures between 22°C and 35°C. Therefore, the product must be kept on ice and precooled pipettes used. The filters were covered with Matrigel and set at 37°C before the epithelium were pipetted on top. To visualize the epithelium they were weakly dyed with neutral red before being placed onto the Matrigel. Affi-gel agarose beads (Bio-Rad) soaked in recombinant activin A protein (1 mg/ml) or BSA were prepared as described above. Beads were placed on top of the epithelia, which were then topped with more Matrigel so that the cultures were surrounded. The epithelium with beads were cultured for 48 hr in DMEM with 10% FCS. Cultures were washed in ice-cold methanol for 1 min and fixed in fresh 4% paraformaldehyde for 1 hr at room temperature. Cultures were then prepared for 35S section in situ hybridization.

Acknowledgments

This research was funded by grants from the Medical Research Council and the Human Frontier Science Program Organization (to P.T.S.) and from the National Institutes of Health (grant HD32067 to M.M.M.). A.L.L. is supported by the National Science Foundation Developmental Biology Training grant BIR-9413237. We thank Bethan Thomas for critical reading of the manuscript, Klaus Kratochwil for teaching us the kidney capsule culture technique, and En Li and Paul Oh for allowing us to quote unpublished results.

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*Genes Dev.* 1998, 12:
Access the most recent version at doi:10.1101/gad.12.16.2636

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