228

MINI-REVIEW

Branching morphogenesis in the fetal mouse submandibular gland is codependent on growth factors and extracellular matrix

Edward W Gresik¹, Noriko Koyama², Toru Hayashi², and Masanori Kashimata²

¹Department of Cell Biology and Anatomy, Sophie Davis School of Biomedical Education, City University of New York, NY, USA ; and ²Department of Pharmacology, Asahi University School of Dentistry, Mizuho, Japan

Abstract : Branching morphogenesis (BrM) is a basic developmental process for the formation of the lung, kidney, and all exocrine glands, including the salivary glands. This process proceeds as follows. An epithelial downgrowth invaginates into underlying mesenchyme, and forms a cleft at its distal end, which is the site of dichotomous branching and elongation ; this process of clefting and elongation is repeated many times at the distal ends of the invading epithelium until the desired final extent of branching is reached. The distal ends of the epithelium differentiate into the secretory endpieces, and the elongated segments become the ducts. This presentation is a brief historical review of studies on BrM during the development of the submandibular gland (SMG). J. Med. Invest. 56 Suppl. : 228-233, December, 2009

Keywords : Submandibular, branching morphogenesis, growth factors, salivary gland

All exocrine glands and several other organs, such as the lung and kidney, develop by a basic developmental process known as branching morphogenesis (BrM) (Fig. 1). In this process, a lining epithelium grows into mesenchyme, which condenses around the epithelium as a capsule. The epithelium elongates to form a stalk that ends in a rounded endpiece. A cleft or groove indents the distal end of the epithelium, marking the site where the epithelium will elongate into two branches, ending in rounded endpieces. The original elongated stalk acquires a lumen to become a duct. A cleft forms at the distal ends of the new endpieces, and the process of branching, lumen formation, and further clefting is repeated over and over again until the desired size of the organ is achieved.

The SMG rudiment arises on the 12th day of fetal life, referred to as E12, as a downgrowth of the oral epithelium, and by E13 the epithelium has branched into 3-5 endpieces surrounded by a capsule of condensed mesenchyme. The study of BrM was greatly facilitated by the work of Elio Borghese, who cultured rudiments of mouse fetal SMGs and showed that BrM could be maintained *in vitro* (1, 2). BrM proceeded well with E14 or older SMGs,

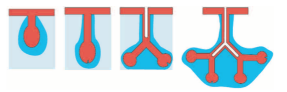


Figure 1. Diagram illustrating steps in branching morphogenesis. Epithelium in red, mesenchyme in blue. See text for explanation.

Received for publication October 16, 2009 ; accepted October 23, 2009.

Address correspondence and reprint requests to Edward W Gresik, Ph.D., Department of Cell Biology and Anatomy, Sophie Davis School of Biomedical Education, City University of New York, New York, NY, USA and Fax : +01-212-650-6812.

but rudiments from mice E13 of younger branched poorly. The extent of BrM was dependent on the amount of mesenchyme taken out with the epithelium, implying that reciprocal interactions between the epithelium and the mesenchyme are needed for BrM. Grobstein then improved culture methods and E13 SMGs also branched in vitro (3, 4). Moreover, if the mesenchyme were separated from the epithelium by trypsin, the epithelium failed to branch, but BrM resumed if the epithelium was recombined with mesenchyme (5). When denuded epithelium was cultured across a filter from SMG mesenchyme, abnormal BrM occurred with many long branches ending in a terminal swelling (5). Grobstein proposed that the epithelium required both direct contact with mesenchyme and interaction with diffusible substances of mesenchymal origin for BrM to proceed (6). He then showed that SMG rudiments treated with collagenase lose their branches, but resume BrM if the enzyme is removed, demonstrating the importance of collagen (7). Later on, working with Rutter and Wessells, he concentrated more on the development of the epithelium in the SMG and in other tissues (8).

Wessells trained a number of talented scientists, including Ken Yamada, Brian Spooner and Merton Bernfield (9-11). Bernfield's group focused on the role of the epithelium, and defined the critical role of the basal lamina for BrM to proceed (12-16). By using agents that specifically interfered with the synthesis of collagen or glycosaminoglycans, Spooner's group showed that both of these ECM components are needed for BrM (17-20).

Nogawa and Nakanishi, working independently or in collaboration, intensively studied the role of collagens in BrM of the SMG, and began to introduce specificity into the analysis of the mesenchyme by defining the spatial distribution of different types of collagen, and showed that while collagen I was widely dispersed in the mesenchyme, type III collagen preferentially localized at the points of clefting, and at the constrictions between the stalk and the endpiece (21-27). They also established that BrM could proceed in the absence of cell proliferation (28).

Kadoya and Yamashina very thoroughly characterized the fine structure of the basal lamina of the SMG, and studied the epithelial synthesis of two of its components, laminin and collagen IV (29, 30). They then demonstrated the presence of the alpha-6 subunit of the integrin receptor for laminin on the basal surface of the fetal SMG epithelium (31). Kadoya and his coworkers then showed that laminins and alpha-6 and beta-1 integrin subunits are needed for BrM, and defined the roles of specific laminin chains, and even specific domains in these laminins that are required for BrM (32-35). This and much more of his work is summarized in their excellent review article (36).

In 1991 Nogawa and Takahashi revolutionized the research on BrM in the SMG by showing that clefting and endpiece formation take place in epithelium stripped of its mesenchyme, and then covered with the basal lamina equivalent, Matrigel, and EGF. If either of these were left out, the epithelium formed a rounded cyst and did not branch (37-38). Nakanishi and coworkers noted that the epithelium formed endpieces, but not elongated stalks that would form the ducts (39). Morita and Nogawa then showed that elongation and stalk formation are driven by the FGF system (40).

Kashimata and Gresik showed that EGF is actually physiologically important, since the mRNAs for EGF and the EGF receptor are expressed during fetal development of the SMG, and that EGF promotes the synthesis of the alpha-6 integrin subunit (41). They also showed that the receptor is localized mainly in the epithelium (42). Hieda and coworkers then established that other ligands related to EGF and other ErbB receptors are also important for BrM, and not only EGF, TGF-alpha and EGFR (ErbB1), namely HB-EGF, Neuregulins, and ErbB2 and 3 (43-44). They have also studied the first signs of cytodifferentiation by defining the roles of junctional proteins in lumen formation (45-47).

Akamatsu and Hosoi and coworkers characterized cytodifferentiation of the rat SMG by following the expression of aquaporin-5 in the epithelium (48), and of the proprotein convertase PACE4, which they showed is also needed for BrM to proceed (49).

Kashimata and Koyama then demonstrated that EGF regulates BrM by activating intracellular signaling cascades involving tyrosine phosphorylation of the EGFR itself, the MAPK Erk-1/2, PLCgamma1, and PI3K (Akt), and showed that the pattern of activation of these signaling pathways varies with age (50-51).

Larsen and Sakai and Yamada confirmed the role of PI3K (52) and have extensively studied the interaction of specific mesenchymal components and integrins important for BrM, such as fibronectin and the alpha-5 integrin subunit (53-55). By laser capture microdissection Sakai demonstrated that fibronectin mRNA is localized in the epithelium, that fibronectin is deposited at the cleft site and in the deepening cleft whene it interacts with it receptor on the epithelial cells, alpha-5-beta-1 integrin. Antibodies against either of these two components interfere with BrM. Many others have contributed importantly to further progress in this field, but space does not allow me to give them proper consideration.

Hoffman and his coworkers at the NIH have extensively documented the critical and wide ranging influence of the FGF signaling system for SMG development and BrM (56-60).

Melnick and Jaskoll have demonstrated that FGF and several other signaling systems are involved in SMG BrM and have emphasized that regulation of development of this gland is not linear, but rather results from multiple interactions; their extensive and important work is summarized in two excellent reviews (61-62).

Recently Nogawa and his coworkers showed that in E12 SMGs FGF induces the epithelium to be able to respond to EGF (63). Mesenchyme-free E12 epithelium exposed first to FGF, and then to EGF, forms endpieces. If it is exposed to FGF and then only to more FGF, it forms elongated stalks, as expected. If it is not exposed to FGF and then exposed to EGF, it does not form endpieces. Koyama and coworkers recently showed that although EGF induces strong phosphorylation of Erk-1/2, FGFs are weak inducers, highlighting the complexity of these interactions (64).

There is a great deal of interest in the study of BrM in the salivary glands and in other organs (e.g. mammary gland, kidney, lung, etc). This knowledge will help not only to elucidate mechanisms of normal development, but will also lay a foundation for understanding abnormal growth, including tumor formation, allowing for new therapeuric approaches. Moreover, the emerging world of tissue engineering would benefit from increased understanding of the factors driving cytodifferentiation and the establishment of tissue architecture.

Given the constraints of space, we apologize to those scientists whose work was not cited.

REFERENCES

- 1. Borghese E : The development *in vitro* of the sub-mandibular and sub-lingual glands of Mus musculus. J Anat 84 : 287-302, 1950
- 2. Borghese E : Explanation experiments on the

influence of the connective tissue capsule on the development of the epithelial part of the submandibular gland of Mus musculus. J Anat 84:303-318, 1950

- 3. Grobstein C : Analysis *in vitro* of the early organization of the rudiment of the mouse submandibular gland. J Morph 93 : 19-44, 1953
- Grobstein C : Epithelio-mesenchymal specificity in the morphogenesis of mouse submandibular rudiments *in vitro*. J Exp Zool 124 : 383-414, 1953
- 5. Grobstein C : Morphogenetic interactions between mouse tissues separated by a membrane filter. Nature 172 : 869-871, 1953
- 6. Grobstein C : Tissue interactions in the morphogenesis of mouse embryonic rudiments in vitro. In : G Rudnick, eds. Aspects of Synthesis and Order in Growth Princeton U Press. Princeton Inc, NJ, 1954, pp.233-256
- 7. Grobstein C, Cohen J : Collagenase : effect on the morphogenesis of embryonic salivary epi-thelium *in vitro*. Science 150 : 626-628, 1965
- 8. Rutter W, Wessells NK, Grobstein C : Control of specific synthesis in the developing pancreas. Natl Cancer Inst Monogr 13 : 51-65, 1964
- 9. Bernfield MR, Wessells, NK : Intra- and extracellular control of epithelial morphogenesis. Symp Soc Dev Biol 29 : 195-249, 1970
- 10. Spooner BS, Yamada KM, Wessells NK : Microfilaments and cell locomotion. J Cell Biol 49 : 595-613, 1971
- Spooner BS, Wessells NK : An analysis of salivary gland morphogenesis : role of cytoplasmic microfilaments and microtubules. Dev Biol 27 : 38-54, 1972
- 12. Bernfield M, Banerjee S : Acid mucopolysaccharide (glycosaminoglycan) at the epithelialmesenchymal interface of mouse embryo salivary glands. J Cell Biol 52 : 64-673, 1972
- Bernfield M, Banerjee S, Cohn R : Dependence of salivary epithelial morphology and branching morphogenesis upon acid mucopolysaccarideprotein (proteoglycan) at the epithelial surface. J Cell Biol 52 : 674-689, 1972
- 14. Banerjee SD, Cohn RH, Bernfield MR : Basal lamina of embryonic salivary epithelia. Production by epithelium and role in maintaining lobular morphology. J Cell Biol 73 : 445-463, 1977
- 15. Cohn RH, Banerjee SD, Bernfield MR : Basal lamina of embryonic salivary epithelia. Nature of glycosaminoglycan and organization of extracellular materials. J Cell Biol 73 : 464-478, 1977

- 16. Bernfield M, Banerjee S : The turnover of basal lamina glycosamino-glycan correlates with epithelial morphogenesis. Dev Biol 90 : 291-305, 1982
- 17. Spooner B, Faubion JM : Collagen involvement in branching morphogenesis of embryonic lung and salivary gland. Dev Biol 77 : 84-102, 1980
- 18. Thompson H, Spooner B : Inhibition of branching morphogenesis and alteration of glycosaminoglycan biosynthesis in salivary glands treated with β -D-xyloside. Dev Biol 89 : 417-424, 1982
- 19. Thompson H, Spooner B : Proteoglycan and glycosaminoglycan synthesis in embryonic mouse salivary glands. Effects of β -D-xyloside, an inhibitor of branching morphogenesis. J Cell Biol 96 : 1443-1450, 1983
- 20. Spooner B, Bassett K and Stokes B : Sulfated glycosaminoglycan deposition and processing at the basal epithelial surface in branching and β -D-xyloside-inhibited embryonic salivary glands. Dev Biol 109 : 177-183, 1985
- 21. Nogawa H, Mizuno T : Mesenchymal control over elongating and branching morphogenesis in salivary gland development. J Embryol Exp Morphol 66 : 209-221, 1981
- 22. Nogawa H : Determination of the curvature of epithelial cell mass by mesenchyme in branching morphogenesis of mouse salivary gland. J Embryol Exp Morphol 73 : 221-232, 1983
- 23. Nakanishi Y, Sugiura F, Kishi J, Hayakawa T : Collagenase inhibitor stimulates cleft formation during early morphogenesis of mouse salivary gland. Dev Biol 113 : 201-206, 1986
- 24. Nakanishi Y, Sugiura F, Kishi J, Hayakawa T : Scanning electron microscopic observation of mouse embryonic submandibular glands during initial branching : preferential localization of fibrillar structures at the mesenchymal ridges participating in cleft formation. J Embryol Exp Morphol 96 : 65-77, 1986
- 25. Nogawa H, Nakanishi Y : Mechanical aspects of the mesenchymal influence on epithelial branching morphogenesis in salivary gland development. Development 101 : 491-500, 1987
- 26. Fukuda Y, Masuda Y, Kishi J, Hashimoto Y, Hayakawa T, Nogawa H, Nakanishi Y: The role of interstitial collagens in cleft formation in mouse embryonic submandibular gland during initial branching. Development 103 : 259-267, 1988
- 27. Nakanishi Y, Nogawa H, Hashimoto Y, Kishi J, Hayakawa T : Accumulation of collagen III at

the cleft points of developing mouse submandibular epithelium. Development 104 : 51-59, 1988

- 28. Nakanishi Y, Morita T, Nogawa H : Cell proliferation is not required for the initiation of early cleft formation in mouse embryonic submandibular epithelium *in vitro*. Development 99 : 429-437, 1987
- 29. Kadoya Y, Yamashina S : Reconstruction of the basement membrane in a cultured submandibular gland. Anat Embryol 183 : 491-499, 1991
- 30. Kadoya Y, Yamashina S : Ultrastructure of the basement membrane and its precursor in developing rat submandibular gland as shown by alcian blue staining. Cell Tiss Res 268 : 233-238, 1992
- Kadoya Y, Yamashina S: Distribution of α6 integrin subunit in developing mouse submandibular gland. J Histochem Cytochem 41: 1707-1714, 1993
- 32. Kadoya Y, Kadoya K, Durbeej M, Holmvall K, Sorokin L, Ekblom P : Antibodies against domain E3 of laminin-1 and integrin α6 subunit perturb branching epithelial morphogenesis of submandibular gland, but by different modes. J Cell Biol 129 : 521-534, 1995
- 33. Kadoya Y, Salmivirta K, Talts JF, Kadoya K, Mayer U, Timpl R, Ekblom P : Importance of nidogen binding to laminin γ1 for branching epithelial morphogenesis of the submandibular gland. Dev 124 : 683-691, 1997
- 34. Kadoya Y, Nomizu M, Sorokin LM, Yamashina S, Yamada Y: Laminin α1 chain G domain peptide, RKRLQVQLSIRT, inhibits epithelial branching morphogenesis of cultured embry-onic mouse submandibular gland. Dev Dyn 212: 394-402, 1998
- 35. Kadoya Y, Mochizuki M, Nomizu M, Sorokin L, Yamashina S : Role for laminin α 1 chain LG4 module in epithelial branching morphogenesis. Dev Biol 263 : 153-164, 2003
- 36. Kadoya Y, Yamashina S : Salivary gland morphogenesis and basement membranes. Anat Sci Internat 80 : 71-79, 2005
- 37. Takahashi Y, Nogawa H : Branching morphogenesis of mouse salivary epithelium in basement membrane-like substratum separated from mesenchyme by the membrane filter. Development 111 : 327-335, 1991
- 38. Nogawa H, Takahashi Y : Substitution for mesenchyme by basement-membrane-like substratum and epidermal growth factor in inducing

branching morphogenesis of mouse salivary epithelium. Development 112: 855-861, 1991

- Mori Y, Yoshida K, Morita T, Nakanishi Y: Branching morphogenesis of mouse embryonic submandibular epithelia cultured under three different conditions. Dev Growth Differ 36: 529-539, 1994
- 40. Morita K, Nogawa H : EGF-dependent lobule formation and FGF7-dependent stalk elongation in branching morphogenesis of mouse salivary epithelium *in vitro*. Dev Dyn 215 : 148-154, 1999
- 41. Kashimata M, Gresik EW : Epidermal growth factor system is a physiological regulator of development of the mouse fetal submandibular gland and regulates expression of the α 6 integrin subunit. Dev Dyn 208 : 149-161, 1997
- 42. Gresik EW, Kashimata M, Kadoya Y, Mathews R, Minami N, Yamashina S : Expression of epidermal growth factor receptor in fetal mouse submandibular gland detected by a biotinyltyramide-based catalyzed method. J Histochem Cytochem 45 : 1651-1657, 1997
- 43. Umeda Y, Miyazaki Y, Shiinoki H, Higashiyama S, Nakanishi Y, Hieda Y : Involvement of heparin-binding EGF-like growth factor and its processing by metalloproteinases in early epithelial morphogenesis of the submandibular gland. Dev Biol 237 : 202-211, 2001
- 44. Miyazaki Y, Nakanishi Y, Hieda Y : Tissue interaction mediated by neuregulin-1 and ErbB receptors regulates epithelial morphogenesis of mouse embryonic submandibular gland. Dev Dyn 230 : 591-596, 2004
- 45. Hieda Y, Iwai K, Morita T, Nakanishi Y: Mouse embryonic submandibular gland epithelium loses its tissue integrity during early branching morphogenesis. Dev Dyn 207 : 395-403, 1996
- 46. Hashizume A, Ueno T, Furuse M, Tsukita S, Nakanishi Y, Hieda Y : Expression patterns of claudin family of tight junction membrane proteins in developing mouse submandibular gland. Dev Dyn 231 : 425-431, 2004
- 47. Hashizume A, Hieda Y: Hedgehog peptide promotes cell polarization and lumen formation in developing mouse submandibular gland. Biochem Biophys Res Commun 339 : 996-1000, 2006
- 48. Akamatsu T, Parvin MN, Murdiastuti K, Kosugi-Tanaka C, Yao C, Miki O, Kanamori N, Hosoi K : Expression and localization of

aquaporins, members of the water channel family, during development of the rat submandibular gland. Pflügers Arch Eur J Physiol 446 : 641-651, 2003

- 49. Akamatsu T, Azlina A, Purwanti N, Karabasil MR, Hasegawa T, Yao C, Hosoi K : Inhibition and transcriptional silencing of a subtilisin-like proprotein convertase, PACE4/SPC4, reduces the branching morphogenesis of and AQP5 expression in rat embryonic submandibular gland. Dev Biol 325 : 434-443, 2009
- 50. Kashimata M, Sayeed S, Ka A, Onetti-Muda A, Sakagami H, Faraggiana T, Gresik EW : The ERK-1/2 signaling pathway is involved in the stimulation of branching morphogenesis of fetal mouse submandibular glands by EGF. Dev Biol 220 : 183-296, 2000
- 51. Koyama N, Kashimata M, Sakashita H, Sakagami H, Gresik EW : EGF-stimulated signaling by means of PI3K, PLCγ1, and PKC isozymes regulates branching morphogenesis of the fetal mouse submandibular gland. Dev Dyn 227 : 216-226, 2003
- 52. Larsen M, Hoffman MP, Sakai T, Neibaur JC, Mitchell JM, Yamada KM : Role of PI3-kinase and PIP3 in sumandibular gland branching morphogenesis. Dev Biol 255 : 178-191, 2003
- 53. Sakai T, Larsen M, Yamada KM : Fibronectin requirement in branching morphogenesis. Nature 423 : 876-881, 2003
- 54. Larsen M, Wei C, Yamada K : Cell and fibronectin dynamics during branching morphogenesis. J Cell Sci 119 : 3376-3384, 2006
- 55. Larsen M, Artym VV, Green JA, Yamada KM : The matrix reorganized : extracellular matrix remodeling and integrin signaling. Curr Opin Cell Biol 18 : 463-471, 2006
- 56. Hoffman MP, Kidder B, Steinberg ZL, Lakhani S, Ho S, Kleinman HK, and Larsen M : Gene expression profiles of mouse submandibular gland development : FGFR1 regulates branching morphogenesis *in vitro* through BMP- and FGF-dependent mechanisms. Development 129 : 5767-78, 2002
- 57. Steinberg Z, Myers C, Heim VM, Lathrop CA, Rebustini IT, Stewart JS, Larsen M, Hoffman MP : FGFR2b signaling regulates *ex vivo* submandibular gland epithelial cell proliferation and branching morphogenesis. Development 132 : 1223-1234, 2005
- 58. Patel VN, Rebustini IT, Hoffman MP : Salivary gland branching morphogenesis. Differentiation

74:349-64,2006

- 59. Rebustin IT, Hoffman MP : ECM and FGF-dependent assay of embryonic SMG epithelial morphogenesis : investigating growth factor/ matrix regulation of gene expression during submandibular gland development. Methods Mol Biol 522 : 319-330, 2009
- 60. Makarenkova HP, Hoffman MP, Beenken A, Eliseenkova AV, Meech R, Tsau C, Patel VN, Lang RA, Mohammadi M : Differential interactions of FGFs with heparan sulfate control gradient formation and branching morphogenesis. Sci Signaling 2 (88), ra55 [DOI : 10.1126/ scisignal 2000304], 2009
- 61. Melnick M, Chen H, Zhou YM, Jaskoll T: The functional genomic response of developing embryonic submandibular glands to NF-kappaB

inhibition. BMC Dev Biol 1: 15, 2001

- 62. Melnick M, Phair RD, Lapidot SA, Jaskoll T: Salivary gland branching morphogenesis : a quantitative systems analysis of the Eda/Edar/ NFKB paradigm. BMC Dev Biol 9 : 32, 2009
- 63. Nitta M, Kume T, Nogawa H : FGF alters epithelial competence for EGF at the initiation of branching morphogenesis of mouse submandibular gland. Dev Dyn 238 : 315-323, 2009
- 64. Koyama N, Hayashi T, Ohno K, Siu L, Gresik EW, Kashimata M : Signaling pathways activated by epidermal growth factor receptor or fibroblast growth factor receptor differentially regulate branching morphogenesis in fetal mouse submandibular glands. Dev Growth Differ 50 : 565-576, 2008