Review

Regenerative Medicine of Bone and Teeth
- with special references to biological principles, problems and their indicators -*

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Abstract: So much anticipation from the side of need patients has been caused by recent advances in tissue engineering technology. However, it seems that effective results can be hardly achieved unless we establish the proper principles for this technology. In this review paper, therefore, we discuss these problems and indicators from the viewpoint of the biological principles of hard tissue regeneration, which are as follows: 1) principles of hard tissue reconstruction, 2) bio-functional nano-composites for regeneration of hard tissue, 3) industrialization of growth factors for hard tissue reconstruction, 4) history and problems of medical ceramic materials, and 5) dentin matrices as a new autograft material for osseous regeneration.

Key words: Bone, Tooth, Regenerative medicine

1) Principles of Hard Tissue Reconstruction

Aim of the session is to give a new insight into the mechanism of hard tissue formation, and thereby to reconsider the biological principles of tissue engineering. Any biocompatible substance, once if it is introduced into living tissues, is regarded as an “extracellular matrix (ECM)” by the cells there. It could be advocated from this idea that the term “biomaterials” is better referred to as “artificial ECM”, if we wish to emphasize the reaction of the material with cells and tissues. In this review paper, therefore, we discuss these problems and indicators from the viewpoint of the biological principles of hard tissue regeneration, which are as follows: 1) principles of hard tissue reconstruction, 2) bio-functional nano-composites for regeneration of hard tissue, 3) industrialization of growth factors for hard tissue reconstruction, 4) history and problems of medical ceramic materials, and 5) dentin matrices as a new autograft material for osseous regeneration.

“Geometry” is defined principally as the structure of artificial ECM at the micrometer level (roughly 0.1 - 1,000 µm), and “ECM”, in this review, means in most cases artificial ECM.

Five factors for tissue formation

We have proposed that five factors must be taken into consideration in order to understand the mechanisms of formation of hard tissues and to secure effective methods of reconstruction when they are partially defected or lost 1-5). These are [I] cells directly involved in bone formation, [II] natural and artificial extracellular matrices (ECM) produced by the cells, or “artificial
Artificial ECM is a factor we can feasibly approach experimentally and then clinically

One of the experimental systems to verify the above proposition is bone morphogenetic protein (BMP)-induced ectopic chondrogenesis and osteogenesis (6-9). In this system, body fluid and some of the mechanical stress are already available in the local area if they are in the right place at the right time. The only items we must add usually are the regulators and the artificial ECM, which correspond to BMP and its carrier, respectively. BMP is a cytokine that was originally known for its unique ability to induce bone formation when it was implanted with a certain carrier into ectopic tissues such as skin or muscle (6-7). This ability has naturally attracted the attention of scientists in the orthopedic and dental fields, including titanium implantology. One of the major problems to be overcome before clinical application is the development of the optimal carriers for this cytokine (1-5). Thus, we have developed and tested more than ten different carriers (1-5, 10-20, 22-25), including a new titanium device (titanium web) (26-28).

We have come to the conclusion that BMP-induced osteo- and chondrogenesis are highly dependent upon the nature of the carriers, notably upon their geometric properties (1-5). This is partly because the carrier of BMP functions not only as a mere drug delivery system, but also as an important artificial ECM (cell substratum) on which the cells undergo growth and differentiation (1-5). At the initial stage of research, BMP-induced bone formation was believed always to follow endochondral ossification (10). However, when new carriers such as titanium web (TW) (26-28), porous particles of hydroxyapatite (HAP) (13) or fibrous collagen membrane (FCM) (12, 14) were introduced, it was found that bone could be formed directly following the process of membranous ossification, without any detectable amount of cartilage formation. Furthermore, nonporous particles of HAP did not induce any meaningful amounts of bone or cartilage under the same conditions (13). A series of studies revealed that there are “vasculature-inducing carriers” such as porous particles or blocks of HAP (16) and “vasculature inhibiting (cartilage-inducing) carriers” such as fibrous glass membrane (16, 25). These findings led us to investigate the detailed properties of natural and artificial ECM, particularly their geometrical properties.

Table 1. Function and properties of natural and artificial ECM (1-2)

<table>
<thead>
<tr>
<th>Properties</th>
<th>Functions</th>
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<tbody>
<tr>
<td>(1) Physical</td>
<td>Mechanical support for cells, tissues, organs and body</td>
</tr>
<tr>
<td>(2) Chemical</td>
<td>Support of cells and molecules through various chemical bonds supplying and depositing of various molecules</td>
</tr>
<tr>
<td>(3) Biochemical</td>
<td>Specific interactions among adhesion molecules, signaling molecules (cytokines) and cell surface receptors</td>
</tr>
<tr>
<td>(4) Geometrical</td>
<td>Directing growth and differentiation of cells and tissues</td>
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Taxonomy of properties and function of ECM

Extracellular matrices have various properties and functions and they are categorized in many ways, but perhaps most fundamentally, they can be divided into (1) physical, (2) chemical, and (3) biochemical properties, and we have now come to a new phase where we have to add a fourth one, (4) geometry. Some of the examples of functions and properties are listed in Table 1. Among these properties, the physical, chemical and biochemical ones have been well studied and documented. However, geometrical properties of ECM have been poorly understood and studied only by a limited number of investigators.

(1) Physical  Mechanical support for cells, tissues, organs and body
(2) Chemical  Support of cells and molecules through various chemical bonds supplying and depositing of various molecules
(3) Biochemical  Specific interactions among adhesion molecules, signaling molecules (cytokines) and cell surface receptors
(4) Geometrical  Directing growth and differentiation of cells and tissues

Geometry in this table is defined as the structure at the micrometer level (0.1~1000 µm). Geometrical structures bigger than this range (macro-structure) are too large for cells to recognize. On the other hand the range smaller than this (nano-structure) is better discussed in relation to organic chemistry and structural biochemistry or, in modern terms “nano-biology”. Geometry of artificial ECM at the micrometer level is increasingly recognized for its importance in tissue engineering and regenerative medicine.
Concerning titanium as an ECM, in Table 1 (four properties and functions), it is clearly shown that the distinguishing properties of titanium are its high physical strength and chemical stability. Moreover, once we give titanium products various geometrical structures, they are maintained for very long time in vivo, as well as in vitro culture situations. No artificial ECM other than titanium possesses this character.

**Taxonomy of geometry artificial ECM**

In order to discuss the effects of geometry on cell activity and tissue formation, we attempted to classify the common geometries applied in artificial ECMs into ten categories, as shown in Table 2. It is thought that this classification is useful when we discuss any geometry of ECM and particularly when we design a new device with new geometry and specific functions. Before discussing the geometry of titanium as ECM in detail, it seems indispensable to investigate previous facts and meanings of the geometry of artificial ECM of any kinds. Thus, let us looks back briefly into the history of artificial ECM based on geometries in Table 2.

**Example of the studies in geometrical artificial ECM**

Numerous biomaterials (artificial ECM) have been devised and tested in pursuit of ideal materials as BMP carriers and other scaffolds for tissue regeneration in vivo or in vitro. During the process, a number of researchers have noticed that cells proliferate

<table>
<thead>
<tr>
<th>Groups</th>
<th>Categories</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Convex Groups</td>
<td>Fibers</td>
<td>Fibrous collagen membrane 12,14 Fibrous glass membrane 15,25,23 Titanium web 26-28 CPSA glass fibers 5</td>
</tr>
<tr>
<td></td>
<td>Particles</td>
<td>Insoluble bone matrix 18,19 Porous particles of HAP with continuous pores 16 Particles of HAP without pores Collagen beads 22,23</td>
</tr>
<tr>
<td></td>
<td>Plane</td>
<td>Conventional culture dishes</td>
</tr>
<tr>
<td>Plane Groups</td>
<td>Sheet</td>
<td>Laser-perforated membrane 17,20 (LPM, sheets with pores)</td>
</tr>
<tr>
<td></td>
<td>Block</td>
<td>Porous blocks of hydroxyapatite 16 (HAP blocks with pores)</td>
</tr>
<tr>
<td></td>
<td>Irregular pores</td>
<td>Porous particle of HAP 10,11,13 Porous block of HAP 16</td>
</tr>
<tr>
<td>Concave Groups</td>
<td>Straight pores (Tunnels)</td>
<td>Honeycomb-shaped hydroxyapatite 1-4 Honeycomb collagen Apatite-coated honeycomb collagen 22 Rat oncsior tube 23</td>
</tr>
<tr>
<td></td>
<td>Round concavities</td>
<td>Concavities on HAP monolith 37</td>
</tr>
<tr>
<td></td>
<td>Rectangular concavities</td>
<td>Micro-pits on silicon tip 30</td>
</tr>
<tr>
<td></td>
<td>Grooves</td>
<td>Microgrooves 38,40</td>
</tr>
</tbody>
</table>
and form tissues in different ways, when cells were placed upon the artificial ECM with the same substance but with different geometry,1-5, 7, 16-27, 29). We will briefly follow some parts of the history of the studies in the geometry of artificial ECM.

**Cartilage formation with close tube structure**

Reddi and Huggins were perhaps the first to notice the importance of the geometry of the ECM in cell differentiation and growth into tissues29). They constructed two geometrically different ECMS, a tube that was open at both ends and a dead-end tube, by cutting rat incisor dentin and implanting it into rat skin. After 4 weeks, they found that the open tube was filled with bone and marrow, while the dead-end tube was filled with cartilage.

The interpretation they made was that young mesenchymal cells (which the authors called fibroblasts at that time) in rat skin were induced to become chondrogenic and osteogenic cells by BMP that was contained in the decalcified dentin matrix. In the open tube, sufficient vascularization resulted in bone formation. But in the dead-end tube, particularly in the apical area, where vasculature and oxygen were scarce, young mesenchymal cells could not differentiate into bone and remained as chondrocytes, even at 100 days. Basset had already reported the phenomenon that the young mesenchymal cells were apt to form cartilage under oxygen-deficient circumstances, while they tended to differentiate into osteoblasts to form bone under oxygen-sufficient and mechanically-compressive conditions 30). Also Folkman and Greenspan31) commented that young cells can differentiate, according to a diffusion gradient, to oxygen and nutrients.

**Sizes and shape of granular type ECM**

Reddi et al. also34) reported the optimal size of granular ECM (decalcified bone powder) in BMP-induced bone formation to be around 420 – 850µ m in particle size. With smaller particles (less than 74µ m), cartilage and bone are scarcely formed34). The shape of granules is also important. Decalcified bone powders are usually irregular in shape and of porous structure, and in general are highly effective as a BMP carrier, but spherical HAP that not equipped with any porous structure is not effective as a BMP carrier35). Also, the size and shape of pores in the porous ECM are important determinant for tissue formation1-3, 16).

**Vasculature-inhibiting geometry that leads chondrogenesis**

Inspired by the experiment done by Reddi and Huggins in 1997, we proposed that by providing artificial ECM, which inhibits vasculature ingrowth, but allows the young mesenchymal cells to enter and proliferate inside, we might arrest the chondrocyte stage in endochondral ossification. One of the reported artificial ECMS with such a microspace structure (vasculature-inhibiting geometry) was fibrous glass membrane (FGM) 10, 15, 25), which was made of 1 µ m glass fibrils. Yoshimoto et al.33) constructed a device, composed of a silicon tube (6 mm length and 3 mm internal diameter) filled with FGM at a certain density together with 1 mg of recombinant human (rh)BMP-2, and implanted it into rat skin. They showed that this restricted system could maintain the cartilage stage of the endochondral process for a considerably longer period than the FGM without silicon tubing.

The results suggested that if a certain bioreabsable system of vasculature-inhibiting ECM is provided and implanted with BMP into skin or muscle, clinically auto-transplantable cartilage may be available. Following on from this work, Nawata et al.36) enclosed the cultured mesenchymal cells with BMP in a diffusion chamber, which both inhibits vasculature and prevents new cells from entering, and implanted it into the abdominal subfascial pocket of a rat. After 5 weeks, they successfully obtained mature cartilage mass within the chamber, which upon reimplantation was able to repair full-thickness cartilage defect of the same strain of rat.

**Vasculature and bone-inducing geometry**

Naturally, ECM with a permissible geometry for vasculature, can permit bone formation. Thus, vasculature-inducing geometry was proposed that promotes bone formation, in contrast to vasculature-inhibiting geometry, which promotes cartilage formation. It was found that porous particles of HAP (Table 2) that were equipped with interconnecting pores easily directed blood vessels throughout the pores of several different particles, when they were implanted into rats with BMP10, 11, 13). In the porous particles of HAP, the vasculature seems to run through neighboring particles, seeking out the orifices of pores that are oriented in the same direction in adjacent particles, just as tree roots develop, finding spaces inside rocks. Vasculature development needs appropriate spaces. This is a reason why interconnected pores are indispensable in ECMS for bone reconstruction. “Vasculature-inducing geometry” and “vasculature-inhibiting geometry” are fundamentally important concepts in ECM geometry. These concepts not only create bone and cartilage differentially, but also may open a strategy for tissue engineering. Next, let us consider vasculature-inducing geometry in more details.

**The optimal pore size for bone formation is nearly that of the Haversian system**

Tsuruga et al.19) compared bone-inducing efficacies of five different porous blocks of HAP, having pore sizes of 106-212, 212-300, 300-400, 400-500, and 500-600µ m, respectively. A block (approximately 5 x 5 x 1 mm, 40 mg) of each HAP ceramic was combined with 4µ g of rhBMP-2 and implanted into the back skin of a rat. Bone-inductive ability was estimated by alkaline phosphatase activity at 2 weeks and osteocalcin contents at 4 weeks after implantation. Results revealed that the highest amount of bone was produced in the ceramics with a pore size of 300-400 µ m.

Interestingly, histological observation of osteogenesis within the pores showed that the bone formed concentrically along the
inner walls of the pores, whose sizes were from 100 to 500 µm. However, within the pores of 500-600 µm, bone did not form concentrically along the inner wall of pores but developed irregularly inside the pores. The results indicated that osteoblasts could recognize the curvature geometry of round-shaped pores up to 400-500 µm in diameter, and construct the concentric bone along the inner wall of pores. However, osteoblasts could no longer recognize the larger curvatures above 500 µm in diameter, and could not form concentric layers along the surface of the pores in this HAP material. It was tentatively concluded from these experiments that the critical diameter of curvature that osteoblasts can recognize is approximately 500µm (250 µm in radius). Normal bone is always remodeling in a concentric manner to form Haversonian systems. It is not surprising that the optimal size of pores in artificial ECM coincides with the average diameter of the Haversonian system that is 200-300 µm. However, if we use the ECM other than pure HAP ceramics, there is a possibility that osteoblasts may behave in a somewhat different way.

**Tissue formation within the honeycomb-type straight tunnels**

As discussed above, the inner wall of a tubing structure of interconnecting pores of 300-400µm in diameter will provide the ideal geometrical environment for osteogenesis. One of the explanations of this phenomenon is that such geometry will generate a higher cell density than a convex or flat surface as shown in Figure 2. However, most of the porous ceramics so far proposed to have “interconnected” pores are, in fact, equipped with irregular pores, not straight tunnels. It was expected that honeycomb-type straight tunnels would more directly exert the geometrical effect upon the growth of bone and other tissues than irregular “interconnected” pores. Inspired by this idea we have developed honeycomb-shaped geometrical ECM made of ceramics, in collaboration with Hokkaido University and Pilot Precision Co., Japan [1,5].

The effect of tunnel size upon bone and cartilage formation was clearly shown when we compared the tissue formations between two different honeycomb apatite ceramics, those with smaller (90-110 µm) and larger tunnels (350µm) [3].

The explanation of the results is as follows: in the smaller tunnels, young mesenchymal cells can enter from the beginning, but vasculature takes time to develop in the tunnel. This situation leads to chondrogenesis and can fill the tunnels with cartilage at 2 weeks. With time, vasculature develops in the tunnels and the cartilage is gradually replaced by bone from both orifices. At 4 weeks, concentric bone fills the tunnel as vasculature runs through. In the larger 350 µm tunnels, condensation of mesenchymal cells was accompanied by vasculature at 1 week, which leads situation led to direct bone formation without any detectable cartilage formation throughout the experimental period. These results clearly indicate that the honeycomb-type geometry with straight tunnels could enhance or inhibit a certain process of tissue formation, as was shown in the examples of direct bone formation and endochondral ossification in this particular experiment.

There are several examples of honeycomb-shaped ECM, other than those made of calcium phosphate ceramics, which direct or control tissue formations. Those include a honeycomb-type insoluble collagen (Collagen Sponge Honeycomb, Koken Co., Tokyo) [21, 22, 35], a laser-perforated polymer membrane (LPM) of bio-degradable properties [17, 20], and honeycomb-shaped polymer films [56]. It was already shown that a honeycomb-type insoluble collagen promoted the differentiation of mesenchymal stem cells into osteoblasts [39], directed Haverson system formation [23], and promoted growth of peripheral nerve fibers [24]. It was reported that the LPM directed collagen fibril formation [17, 20], and honeycomb film guided differentiation and growth of nerve stem cells [56].

**Concave geometry and optimal space theory**

In Table 2, we have divided all of the geometries of ECM into three large groups: convex, plane and concave. So far, we concentrated the discussion on concave group and have concluded from the numerous previous examples that as far as bone formation is concerned, concave geometry of ECM is advantageous over convex or flat. The possible mechanisms involved are as follows: (1) concave geometry permits a higher concentration of cells, (2) it leads to effective accumulation of cytokine and other active bio-molecules (3), there is a closer chance of cell-cell communication and (4) the creation of three-dimensional (3D) environments occurs more quickly than does those of two-dimensional (2D). The optimal diameter of pores or tunnel structures was concluded to be 300-400 µm, and the effective radius of curvature up to about 250 µm, as shown in the preceding discussion. These conclusions concerning the size of concave geometry are not only applicable to completed pores or tunnels, but also may be partially eligible to open pore structures (concavities), rectangular concavities [30] and grooves [39, 40], as shown...
Hirano et al. created a tartan-patterned grooved surface with 500 mm wide grooves on titanium alloy rod. These grooves promoted faster bone ingrowth, resulting in a stronger bond with bone when implanted into the rabbit femoral condyle.

When we look at histology in many studies on BMP-induced bone formation, we find generally that most of the initial bone formation started at the concave area of ECM. Furthermore, even without the addition of BMP or cytokines, several authors have reported that osteogeneses were observed in concavities and pores of ceramics in vivo. Ripamoti and coworkers found osteogenesis in the concavities of HAP ceramics, which were implanted into baboon muscles without BMP. Endo et al. also reported bone formation in porous βTCP ceramics (porosity of 75%, pore size 100-500 µm) implanted in the muscle of beagle dogs, 56 days after implantation without BMP. Again it should be noted that initial bone formation occurred in the concave part of the ceramics in both studies.

Proposal of “optimal space theory”: these observations led us to the concept of “optimal spaces for tissue formation in artificial ECM”. By this concept we propose that each kind of tissue ECMs has an optimal geometrical shape and size, which facilitate its cells to differentiate, proliferate and develop particular tissue in a certain direction. These optimal spaces are not limited to tube- or sphere-types, but include certain spaces between the solid ECM structures, such as fibers, particles, and flat planes as shown in Table 2. The size of the optimal spaces is assumed to be approximately 200-500 µm which is dependent upon tissues.

Titanium web as an example of fiber-type ECM

The fibrous ECMs, which are classified as convex-groups in Table 2, can also compose optimal 3D spaces. However, the spaces are distorted and often collapse when the fibrous ECMs are implanted into the body. Conventional examples are fibrous membranes of polylactic acid and its copolymer with polycaprolactone, and cross-linked collagen fibrous membranes. In this regard, the titanium web (TW) construct is an extremely rigid fibrous structure. The 3D structure of TW is composed of thin titanium fibers 50-100 µm in diameter. The cross section of the fibers in this material is not round but square, due to its manufacturing process. Each titanium fiber is fused to adjacent ones by vacuum sintering, so that the network or web structure is strong and can be molded into any form indicating, disks (Figure 4A, B, C and D) and even tubing. The disk form of the titanium web has been applied to 3D cultures of osteoblasts and mesenchymal stem cells (Figure 4A, B, C and D). The results indicated that TW as 3D culture system generally increased cell proliferation due to its large surface area, and promoted differentiation due to its 3D effects on the cells, compared with
conventional cultures on a flat dish 42). There are increasing needs for appropriate ECMs for 3D culture systems. Day by day we need more information on cell behavior in a 3D environment, in addition to the 2D environments of conventional flat culture dishes 43, 44).

We should recall that five factors are required for tissue reconstruction, as discussed in the beginning. If we provide the cells with appropriate houses (matrices), nutrition (vasculature) and some of the stimulations (regulators and biomechanical dynamics), they will build up tissues and organs.

2) Bio-Functional Nano-Composites for Regeneration of Hard Tissue

Biological tissues have peculiar structures organized on a nanometer-scale. In order to synthesize such biomimetic nanostructure materials or novel functional biomaterials, interfacial interactions between constituents are important. In this section, two examples of bio-functional nano-composites are introduced for the application of clinical treatments such as osteoporosis and metastatic bony carcinoma.

Tissue engineering includes the following biological processes: (1) proliferation of functional cells mainly derived from patients themselves; (2) formation of cell-based constructs with sufficient size in vitro; (3) implantation of the constructs to the defects; and (4) regeneration of tissue in vivo. In general, bioaffinitive materials, i.e. scaffolds, are necessary to make useful cell-based constructs.

For hard tissue regeneration, possible scaffolds are ceramics, as well as bioabsorbable polymers and/or their composites. Ohgushi et al. confirmed heterotopic ossification by the implantation of osteoblasts on porous ceramics including HAp 45-47), and applied this technique to human knee joint replacement 48). In bone remodeling metabolism, scaffolds are required to have an appropriate decomposition rate; therefore, the size of HAp crystals and the state of collagen fibrillogenesis in the scaffolds should be similar to those in bone. It is therefore expected that the decomposition of such HAp/Col composite should show the same reaction as autologous transplanted to a bone defect, depending on osteoclasts.

Biomimetic synthesis of bone-like nanocomposite

In osteogenesis, osteoblasts provide raw materials, mainly collagen and HAp (calcium phosphate), which are spontaneously aligned to the nano-ordered structure of bone 49). In bone formation, osteoblasts appear to control a self-organization process of collagen and HAp as a chemical reaction. Therefore, the HAp/Col nanocomposite with similar chemical composition and nanostructure to bone should be synthesized on the basis of the self-organization mechanism of bone 50-53. The HAp/Col composite has a good cell attachment property.

The starting materials for HAp/Col composite are Ca(OH) 2, H 3PO 4 and collagen. In our studies, we have used Ca(OH) 2 prepared from CaCO 3 (alkaline analysis grade, Wako Pure Chemicals Inc.). HAp stably forms under weak basic pH 9. The collagen we
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used was atelocollagen extracted from porcine dermis (biomaterial grade, Nitta Gelatin Inc.); its isoelectric point was pH 7-9.

Self-organization of HAp and collagen are promoted at 40 °C and pH 8-9. This pH value is larger than that of animal body fluid, normally pH 7.2 to 7.4, indicating that bone forms also under such pH conditions controlled by osteoblasts. Fibrillogenesis of collagen is promoted around 40 °C and under ion strength similar to body fluid. The TEM image of the composite obtained is shown in Figure 5, together with its schematic nanostructure. The maximal length of the HAp/Col self-organized fiber was 75 µm in length; the longest fiber was synthesized in a reaction vessel with ion concentration less than 1/700 of PBS.

Dense bodies of the HAp/Col composite were obtained by filtration and dehydration of fibers under high pressure, as the HAp/Col fibers contain a large amount of water. Further, porous bodies of the HAp/Col self-organized composites were prepared from fibers by a freeze-dry method. The HAp/Col porous body demonstrated sponge-like viscoelastic property as shown in Figure 6, which increases cell migration.

Biological reaction of the HAp/Col composite

The HAp/Col composites were implanted in bone, and the biological reaction was examined. Figure 7 shows the HAp/Col composite implanted into a rat tibia for 5 days, stained with histoenzymatic tartrate resistant acid phosphatase (TRAP). TRAP active multinuclear giant cells were identified as osteoclasts, which were observed onto the HAp/Col surface. The porous HAp/Col composite was finally substituted to bone by the remodeling metabolism.

After the appearance of osteoblasts, alkaline phosphatase (AIP) positive cells were induced in the neighborhood of osteoclasts around the HAp/Col composite as shown in Figure 8, and were identified as osteoblasts that form bone. This reaction continues until the HAp/Col nanocomposite was completely absorbed, and finally the HAp/Col nanocomposites were substituted to new bone. The absorption of the composite and the bone formation reaction was almost the same reaction as in for autografted bone. This result suggests that the HAp/Col nanocomposite was recognized as “bone” by bone tissue and was incorporated into bone remodeling metabolism.

The HAp/Col porous bodies with sponge-like viscoelasticity were implanted into bone defects with appropriate deformation. The viscoelastic property was very suitable for a bone-filler and for tissue engineering of hard tissue, as it is difficult to seed cells in a 3-dimensional scaffold homogeneously. The HAp/Col porous body allowed homogeneous cell migration by repeatedly deforming the sample in a cell suspension. The porous structure of the HAp/Col composite was well-maintained after the deformation, much better than a pure collagen sponge and other biodegradable polymers even after medium exchange infiltration.

Nano-crystals for DDS of bony metastatic cancer

Recently a drug delivery system (DDS) has been studied from material-scientific points of view to improve the efficiency of drugs and/or reduce their side effects. In order to develop a DDS for the treatment of metastatic bony carcinoma, a calcium phosphate system, such as HAp and octacalcium phosphate (OCP), was adopted as a DDS carrier. Bisphosphonate Bps, such as etidronate, was controlled to into the nano-particles of HAp and OCP. It is thought that OCP forms an intercalation compound, prepared by a wet method in the presence of bisphosphonate. As shown in Figure 8, such a compound could automatically control the activity of osteoclasts, useful for the treatment of metastatic bony carcinoma.

Nano-composites such as HAp/Col and OCP/drug are responsible for osteoclasts. It was shown that this osteoclast-responsibility could be applied to new bone formation and automatic control of osteoclast activity.
3) Industrialization of Growth Factors for Hard Tissue Regeneration

In Japan, neither medical drugs nor medical devices for the reconstruction of hard tissue regeneration have been sold, while those which have been launched in USA and Europe and have contributed to cure of disease. The components in these medical products originally exist in the body and have properties to accelerate the proliferation or the differentiation of a cell. With the development of genetic technology, mass-production of these substances became possible. We have expected the application of these substances to the regeneration of skin, cornea, blood vessel, etc. as well as osteogenesis, and some substances have been used for medical purpose. In this section, we will describe the discovery history of these main factors and the situation of their industrialization.

**Discovery history of cell growth factors**

Cell growth factors were first discovered in the 1950s. The Levi-Montalcini group found Nerve Growth Factor (NGF) as the substance that is involved in the survival and growth of nerve cells. Levi-Montalcini received a Nobel Prize in 1986 together with Cohen, who discovered Epidermal Growth Factor (EGF) in 1962. In 1974, Fibroblast Growth Factor (FGF) was found by Gospodarowicz as the factor to accelerate the proliferation of 3T3 cell lines that were extracted from the bovine pituitary gland. Subsequent studies confirm that this FGF stimulates the proliferation of vascular endothelial cells, smooth muscle cells, cartilage cells, and mesenchymal stem cells. In 1984, the existence of basic Fibroblast Growth Factor (bFGF, FGF-2) and acidic Fibroblast Growth Factor (αFGF, FGF-1) was demonstrated.

Ross et al. discovered Platelet-Derived Growth Factor (PDGF) as the causal factor of arterial sclerosis. In 1965, Urist discovered a factor that induces osteogenesis and named it Bone Morphogenetic Protein (BMP). Subsequent studies demonstrate that BMP belongs to TGF-β super family. Nakamura et al. discovered Hepatocyte Growth Factor (HGF) in 1984 as the growth factor of hepatocytes. At present attention is on as the angiogenic factor. Vascular Endothelial Growth Factor (VEGF) is a strong growth factor for vascular endothelial cells, and in 1989 a grouped by Ferrara discovered it. Studies thereafter have clearly indicated that VEGF is the same as Vascular Permeability Factor (VPF) found in 1983.

**Situation of industrialization of products for hard tissue regeneration**

BMP-2 has already been launched in the US and European countries as the alternative material for bone graft in combination with collagen sponge. It has been used to meet the three indications of acceleration of spinal fusion, open tibia fracture, and the regeneration of alveolar and mandibular bone. The US is the first country in which the product was launched in 2002, followed by Europe in 2005. The product name is INFUSE® Bonegraft in the US and InductOS® in Europe.

In Japan, Yamanouchi Pharmaceuticals Co., Ltd. (now Astellas Pharma Inc.) completed clinical studies for spinal fusion and applied for manufacturing approval. It has not yet been approved, and development rights and distributorship were sold to Medtronic Japan Co., Ltd. Similarly to BMP-2, OP-1 was developed as the adjuvant therapy for spinal fusion and long bones. It has already been launched in the US, Europe, Australia, and Canada, under
the product name OP-1®. In combination with βTCP, PDGF has already been launched in the US as the medical device to repair the defect of alveolar bone due to periodontal disease. Under the product name GEM21S®. In addition, PGDF has already been sold under the name Regranex®gel in the US, Europe, Canada, Korea, etc., as a medicine for diabetic skin ulcers. In 1986, Abraham’s group in California Biotechnology Inc., USA (now Scios Inc.) reported that the group clarified the DNA sequence of human bFGF 62). Kaken Pharmaceutical Co., Ltd. contracted with Scios Inc. in 1988 and began its development in Japan. It was launched under the trade name Fiblast®Spray (Figure 10) in 2001, and the indications for bedsores and skin ulcers (burn ulcers/leg ulcers). Clinical trials of bFGF are ongoing in Japan as a drug for bone fractures and the regeneration of periodontal tissues.

Issues in the development of drugs for hard tissue regeneration

Cell growth factors that have been expected as the drugs for the regeneration of hard tissues are proteins. They are unstable materials and structural changes easily occur. This instability is a large barrier to the development of protein formulations. In-depth stabilization technology based on physicochemical studies and analysis techniques with high quality for support are required in order to overcome these issues.

When bFGF or PDGF were intravenously administered to rats, these accumulate in the liver and kidney 63,64). When bFGF was subcutaneously dosed to rats for 3 months repeatedly, the excretion of urinary proteins markedly increased65). The intravenous administration of bFGF clinically caused hypotension66) and an increase in urinary proteins67). On the other hand, studies on local application of bFGF confirmed no transfer of bFGF to the blood68). In addition, it is well known that growth factors are involved in the acceleration of growth of tumor cells. Therefore, we believe that the systemic administration of growth factors should be avoided as much as possible for safety.

4) Development of bioabsorbable and biomimetic ceramics and A big prospect for regenerative therapy-sudies in Hokkaido

In graving society, for regenerative therapy for patients, ceramics in medicine are important biomaterials and collaboration among the educational-industrial-administrative complex is an effective action. After returning the transition of bioceramics, functionally graded apatites (fg-HAp) originated from natural bone will be outlined as an interdisciplinary study focusing on biomimetic materials. The role and scheme of regenerative therapy-studies that are useful out there will be discussed. Since fg-HAp has excellent degradation-absorption, rhBMP-2-adsorption and release characteristics due to body fluid permeation and blood permeability, rhBMP-2/fg-HAp is an osteoinductive bioceramic with bone-remodeling. The fg-HAp can be designed and controlled by the calcination and dissolution-precipitation and it might be applied to higher advanced medical care by strong connection with medical and dental teams.

Biomimetic materials and collaborate study in an interdisciplinary field biomimetic ceramics for regenerative therapy

Figure 11 shows SEM photographs of three living tissues-originated materials. For starfish, calcium carbonate (CaCO₃) ceramics with homogeneous pore diameters of about 10 µm were obtained by the decomposition by enzymes and the calcination at 600°C for 2h in air. For dentin from extracted teeth, a lot of dentinal tubules with 1-2 µm were observed. For bovine bone, hexagonal grains in 1-2 µm with single phase of hydroxyapatite (HAp) were designed by the calcination at 1100°C for 24h in air. Living tissues-originated materials contain the grain morphology, the surface structure, and the chemical nature which originate from small amounts of metal ions in a living environment69). Biomimetic ceramics are functional materials that are efficient for maintaining optimal environment for cells or biopolymers. We focused on the development of degradable and bioabsorbable HAp, which can be used as chief biomimetic ceramics for regenerative therapy.

Regenerative therapy-studies recognized by the Hokkaido government through collaboration among the industrial-educational-administrative complex

In bone reconstruction or regenerative therapy, development and application of artificial organs imitating living tissues, medical devices, and medical systems, have been expected. Research and
development of biomaterials should be carried out by the fusion of many techniques, such as medical research, biotechnology, and materials engineering. Figure 12 shows collaborative studies on biomaterials and regenerative therapy in an interdisciplinary field. The title of the study is “Development of functional biomaterials and their application for regenerative therapy and advanced medical engineering”, performed as a special study in Hokkaido by seven agencies in 2006-2008. Hokkaido Industrial Research Institute (HIRI) built up a big project team, which consists of Kyoto University, Tokyo Medical and Dental University, Health Sciences University of Hokkaido and three companies. The three universities have high basic technology on biomaterials and cell engineering and the three companies possess the know-how and facilities for production and sale of medical equipment and safe treatment of animal bone. For the purpose promoting a bio-industry cluster in Hokkaido and utilizing Hokkaido resources, through the fusion of bioscience and nanotechnology, we are preparing porous composites using animal bone and developing cell culture kits or pulverizing and processing instruments to apply for regenerative therapy and advanced medical engineering.

**Invention of bioabsorbable and functionally graded apatites**

**Development of tailored ceramics**

For bone reconstruction and bone regeneration, many kinds of biomaterials have been developed. Hydroxyapatite (HAp) and β-TCP can be applied as bioactive and substituted materials for hard tissues in the dental and medical fields because of excellent biocompatibility and osteoconduction. However, HAp ceramics synthesized from reagents show very low bioabsorption characteristics in implanted regions even after several years. Therefore, commercial HAp ceramics cannot always be ceramics would be absorbed and substituted for mother tissues after operation, depending on the new bone formation and bone regeneration. We arranged the required capability of apatite products for the biomaterials early incorporated into bio-metabolic system. The new apatite products need to be bioceramics with excellent body fluid permeability, high surface area, appropriate mechanical strength, and high activity for various cells. Responding to the necessary conditions, functionally graded and porous ceramics, in which grain size and crystallinity of HAp gradually change from surface region to bulk part, were considered. We tried to control absorption and osteoinduction characteristics by loading of recombinant human bone morphogenetic protein-2 (rhBMP-2) into the ceramics.

**Biomimetic ceramics prepared using animal bone**

Safe animal bone was supplied from traceable livestock in Hokkaido Meat Packer Incorporation. To design biomimetic materials originated from animal bone, recycle techniques of apatite resources and dissolution-precipitation conditions in ceramics engineering were discussed. Bioabsorbable ceramics with spongy bone-originated skeleton structure and pore structure of nano-particles, which may promote body fluid permeability and cells-affinity, were prepared.

**Design of fg-HAp ceramics derived from bovine bone**

Spongy and cortical bovine femur bones were used as starting
materials, boiled and calcined at 800-1100°C for 24h in air to obtain crystalline HAp (b-HAp) ceramics. By the calcination process, all prion proteins and organic residues of bovine bone were completely burned out. The pulverized cortical b-HAp was completely dissolved into a HNO$_3$ solution, while the spongy b-HAp was partially dissolved into another HNO$_3$ solution. After the solutions were mixed, NH$_3$ solution was added and reprecipitated HAp (r-HAp) crystals were carefully precipitated on macro-pores and micro-pores of spongy b-HAp at pH 10.5 and 25°C$^5$. The modified spongy b-HAp was aged for 24 h to fabricate a structure of fg-HAp. The fg-HAp ceramics were filtrated and washed with distilled water and dried at 120°C. To obtain rhBMP-2/fg-HAp ceramics, rhBMP-2 (5µg) solution was added to fg-HAp ceramics in a sterilized dish.

Microstructure and crystallinity of biomimetic fg-HAp ceramics

fg-HAp had same chemical composition as b-HAp and r-HAp$^{77-79}$. The (Ca/P) ratios of fg-HAp were 1.64-1.66, which are close to the stoichiometric value of 1.67 for the standard HAp. Small amounts of Na$^+$ and Mg$^{2+}$ ions, present at less than 1%, were detected. The fg-HAp ceramics exhibited mechanical strengths sufficient for operation and handling of animal experiments. Figure 13 shows the appearance and SEM photographs of fg-HAp ceramics prepared by the calcination at 800°C and partial dissolution-precipitation. Spherical moss-like grains forming needle-like microcrystals and macro-pore sizes of 100-800 µm were observed. The specific surface areas, total pore volumes, and porosities of the ceramics were 30-40 m$^2$.g$^{-1}$, 0.35-0.40 cm$^3$.g$^{-1}$, and 60-80 %, respectively. In the curves of the pore size distribution for the ceramics, pore volumes in the micro-pore sizes of 10-160 nm were clearly recognized. The Micro-XRD analyses indicated that the degree of crystallinity of the single phase HAp was gradually distributed from the surface layer of the macro-pore wall to the bulk region of the h-HAp body structure.

Based on these results, it was found that the pore structure of the fg-HAp was classified into a macro-pore (100-800 µm) originating from spongy bone and a micro-pore (10-160 nm) prepared by the calcination and partial dissolution-precipitation$^{79}$. The fg-HAp ceramics with the specified pore structures and gradations in both crystallinity and grain size can permit body fluid to easily permeate the parts of a living body. Also, microcracks in bulk regions of the ceramics formed by partial dissolution with HNO$_3$ may be related to auto-degradation and body fluid permeation. Moreover, the fg-HAp ceramics were certificated to be abnormal prion-free materials by the enzyme antibody reaction kit (Bio-Rad) for diagnosing bovine spongiform encephalopathy in Obihiro University of Agricultural and Veterinary Medicine.

Characterization of biocompatibility for different HAp ceramics by soaking in SBF

For evaluation biocompatibility of the ceramics, each of the fg-HAp and b-HAp ceramics was soaked at 36.5°C and pH 7.4 in a simulated body fluid (SBF) for 1-90 days$^{79}$. The SBF is the solution which ion concentrations and pH nearly equal to those in human plasma at 36.5°C. Figures 14 shows photographs of the b-HAp and fg-HAp ceramics soaked at 36.5 K and pH 7.4 in SBF for 1, 8, 14, and 28 days. The b-HAp was calcined at 800°C for 24 h in air. At 14 days after the soaking, microstructure of the b-HAp became porous urchin-like grains of 5-6 µm, while at 8 days, that of the fg-HAp already changed from small grains to dense cocoon-like ones by rapid precipitation of HAp microcrystals. These significant differences in surface morphology suggest that the fg-HAp has higher bone-bonding ability than the b-HAp, and that, concerning proteins-adsorption on the two ceramics surfaces, the fg-HAp surfaces show larger amounts of proteins adsorbed and higher adsorption heats for proteins because r-HAp microcrystals on the sintered grains enhance surface roughness of the b-HAp ceramics.

Animal experiments of different HAp ceramics

Histological and immunochemical observations of explants

The fg-HAp and rhBMP-2/fg-HAp ceramics were implanted into the subcutaneous tissues of the back region in 4-week-old male Wistar rats$^{78,80-83}$. At 4, 8, and 12 weeks after implantation, these samples were explanted. The specimens were fixed in neutral buffered formalin, decalcified with formic acid, embedded in paraffin, sectioned and stained with hematoxylin and eosin (HE).
Figure 14. SEM photographs of f-HAp and b-HAp ceramics soaked at 36.5°C and pH 7.4 in SBF

They were histologically evaluated using an optical microscope. To investigate the eosinophilic areas in HE sections of the implanted ceramics, the specimens were immunostained with polyclonal rabbit antibody against rat albumin using the avidin-biotin complex method.

For the fg-HAp ceramics, at 4 weeks after the implantation, body fluid extensively permeated into the bulk regions of HAp through the nano-micro-pores of the ceramics. In the macro-pores, many multinucleated giant cell and fibroblast were observed. Surface- and bulk- degradations of the HAp proceeded, so that a total size of the HAp block remarkably decreased. In the eosinophilic region of HE sections marked in the ceramics, the immunohistochemical staining was carried out. The immunostaining of albumin in the fg-HAp explant indicated to be immunopositive.

Figure 15 shows photographs of HE sections at 4 weeks after the implantation of rhBMP-2/fg-HAp ceramics. As a whole, geometrical relation of bone-HAp was like a mosaic image. Multinucleated giant cell on the crinkle HAp surface, HAp encapsulated in new bone, and retention of body fluid in HAp observed. The HAp fragmented by body fluid permeation and new bone formation from surface and bulk regions of the ceramics were observed. The HAp, which had been cut into small pieces, was incorporated into the induced bone, suggesting that osteoinduction occurred with bone remodeling.

**In vivo-release characteristics of rhBMP-2 from different HAp ceramics**

The ¹²⁵I-labeled rhBMP-2 (0.5 µg) supported fg-HAp or b-HAp ceramics were implanted into the back subcutis of 6 week-age female ddY mice. At 1-28 day after the implantation, the mice were sacrificed. The radioactivity of the remaining HAp ceramics, excised skin, and filter paper was measured on a gamma counter. Figure 16 shows changes in the retention percentage of rhBMP-2 for rhBMP-2/fg-HAp or b-HAp ceramics. The retention percentages of rhBMP-2 decreased with increasing implantation time. At 1-28 days after the implantation, the retention percentages of rhBMP-2 for fg-HAp are higher than those for b-HAp. Even at 14 days, the value for fg-HAp was about 60%, suggesting that fg-HAp ceramics are excellent osteoinductive scaffold. The difference in rhBMP-2-release between fg-HAp and b-HAp can be caused by differences in pore structure or adsorption strength of rhBMP-2 molecules on the HAp surfaces.

**Expectation and a big prospect of regenerative therapy-studies in Hokkaido**
The fg-HAp ceramics have excellent rhBMP-2-adsorption and release characteristics, as well as degradation-absorption due to body fluid permeation and blood permeability. The crystallinity and the pore size distribution of nano-particles can be designed and controlled by the calcination and partial dissolution-precipitation method. Since it was clarified that the rhBMP-2/fg-HAp was one of osteoinductive and bioabsorbable bioceramics related to bone-remodeling system, (Japanese patent No.3718723) was purchased in 200677). Also, the fg-HAp ceramics might be applied for higher advanced medical care by strong connection with medical and dental teams. Hereafter, aiming for development and clinical application of artificial bone launching from Hokkaido into world, we are going to strengthen the alliance with the Department of Orthopedic Surgery, Graduate School of Medicine, Hokkaido University in the medical field and the Department of Oral Surgery, School of Dentistry, Health Sciences University of Hokkaido in the dental field. We would like to contribute acceleration of studies on biomaterials and development of medical industry, scoping in vivo-experiments for large animals and heightening the value of commercial biomaterials. Moreover, to improve many cases in Hokkaido and practice regenerative therapy for patients, interdisciplinary day-to-day collaboration prompting towards needs of medical locus, human network among the medical university and college in Hokkaido, establishment of Regeneration Therapy Center constituted by interdisciplinary project teams will be aspired.

5) Dentin Matrics as a New Autograft Material for Osseous Regeneration

In this section, the animal and human-derived dentin recherche and a clinical pioneering case of autogenous demineralized dentin matrices transplant are described. The bone-inducing property of rabbit dentin matrices was discovered in 1967. We confirmed that completely demineralized dentin matrices (DDM) including small patches of cementum, derived from human adult extracted teeth, induced bone and cartilage independently in subcutaneous tissues of nude mice at 4 weeks after implantation98). Until now, several dentin research papers have reported a chondro-osteoinductive potency of DDM and a BMP-like molecule in dentin matrices86-98). We confirmed that completely demineralized dentin particles derived from human adult extracted teeth, induced bone formation in the rabbit intramuscular pockets86). Until now, several dentin research papers have reported a chondro-osteoinductive potency of DDM and a BMP-like molecule in dentin matrices86-98). We confirmed that completely demineralized dentin particles derived from human adult extracted teeth, induced bone and cartilage independently in subcutaneous tissues of nude mice at 4 weeks after implantation98).

Bone induction by demineralized dentin

The history of a bone-inducing research in dentin began with a report in 1967 that rabbit demineralized dentin matrices (DDM) induced bone formation in the rabbit intramuscular pockets99). Until now, several dentin research papers have reported a chondro-osteoinductive potency of DDM and a BMP-like molecule in dentin matrices86-98). We confirmed that completely demineralized dentin particles derived from human adult extracted teeth, induced bone and cartilage independently in subcutaneous tissues of nude mice at 4 weeks after implantation98).

Basic studies of human dentin: Preparation of human demineralized dentin and non-demineralized dentin

Adult human third molar teeth were donated by outpatients at Health Sciences University of Hokkaido. The extracted third molar teeth were collected and prepared for completely demineralized dentin matrix (DDM) and non-demineralized dentin matrix, so called calcified dentin matrix (CDM). Briefly, the teeth were crushed in liquid nitrogen, washed in 1M sodium chloride, defatted with methanol/chlorophorum solution and rewashed for CDM. The CDM were demineralized completely in HCl solution (pH2.0) for DDM. The CDM and DDM particles were extensively rinsed in cold distilled water and lyophilized (Figures 17-19). The particle sizes varied from 0.4 to 0.8 mm.

Assay for biological activity and results

Seventy mg of DDM or CDM were put into cut-opened tuberculine syringes. Ten nude mice (male, 4 week-old, body weight; 50 g) were subjected to intraperitoneal anesthesia with pentobarbital sodium. A vertical incision (1 cm) was made under...
sterile conditions in the skin over the back region, and 2 subcutaneous pockets were prepared by blunt dissection. Each animal received one DDM implant and one CDM implant. The implanted materials were removed at 4 weeks, and were prepared autologous buccal mucosal graft

DDM induced bone and cartilage, independently. New bone with osteoblast lining, not osteodentin, was found directly on the surface of DDM, while CDM did not induce hard tissue formation at 4 weeks (Figure 20).

Osteo-odonto-keratoprosthesis surgery using autologous dentin-bone

A 49-year-old female lost her eyesight in end-stage Stevens-Johnson syndrome. A surprising operation for recovering the patient’s eyesight, osteo-odonto-keratoprosthesis (OOKP) surgery was first done in 2003, Japan. The OOKP procedure utilizes an autologous tooth-bone complex to mount a polymethylmethacrylate optical cylinder as an artificial cornea. The root-bone complex and the plastic lens were fixed by dental adhesive cement, and the unit was stabilized by an overlying

Dentin autograft into atrophied jaw for tooth autotransplantation

A 22-year-old woman with reversed occlusion presented with a congenital missing tooth of the mandibular right second premolar (#45) and an atrophied bone. The medical history was essentially within normal limits and noncontributory. The quantity and quality of the atrophied bone and the length of teeth were estimated by CT scan. Upper premolar teeth (#14,#24) were extracted under the orthodontic planning. The vital tooth (#14) without root canal treatment was selected and prepared for freeze-dried DDM. Microdont (#28) which root was single (root length: 10mm, long

width: 6mm) was chosen as transplanted tooth. The autogenous, immediate tooth transplantation with the freeze-dried DDM autograft was carried out in 2004 under the local anesthesia. Transplant cavity was formed using Filialit-2® dental implant system. As a limited osseous defect existed adjacent to the transplanted tooth, the defect was filled with particles of DDM, and bone augmentation on the perforated cortical bone was done by DDM (Figure 21). The mucoperiosteal flap was repositioned

Figure 17. SEM of human calcified dentin matrix (CDM)

Figure 18. SEM of human demineralized dentin matrix (DDM) use

Figure 19. Freeze-dried human DDM particles before clinical

Figure 20. Induced bone on human DDM at 4 weeks

Figure 21. Immediate tooth autotransplantation simultaneously with DDM autograft

Figure 22. Reconstituted occlusion at 40 months after operation
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The content of this paper has been described in detail to “Regenerative medicines of the bone and teeth”(100). Refer to this book for details.
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