Cartilage-Bone Interface Features, Scaffold and Cell Options for Regeneration

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Introduction

Tissues with different material and biological properties are connected to one another through interfaces, which can be generally categorized as soft-to-soft tissue interfaces (muscle-tendon, etc.), soft-to-hard tissue interfaces (cartilage-bone, tendon-bone, etc.) and hard-to-hard tissue interfaces (dentin-enamel, etc.). Since these interfaces merge biological materials, i.e., tissues, having distinct composition, structure and function, they possess complexities associated with their hierarchical structures, and when injured their healing/regeneration pathways follow more intricate phenomena compared to single tissues making up the interfaces. Findings reveal that injuries related to tissues connected in series occur mostly at the interfaces due to the mismatch between material properties of individual tissues. Therefore, interface tissue engineering has recently attracted significant attention from academia to be able to understand the mechanism of cell-materials interactions relevant to interfaces. This paper reviews the structure, composition and function of cartilage-bone interface in conjunction with the scaffold and cell options for its regeneration.

Cartilage-Bone Interface

Osteoarthritis (OA) is a degenerative joint disease seen at the cartilage-bone interface in the knee. It is characterized by the lesions in the articular cartilage at early stages and complete loss of function of the knee joint at advanced stages [1]. OA is known as one of the most common joint-related traumas, and it is reported that, between 2010 and 2012, around 52.5 million adults (18 years and over) were diagnosed with OA and that every 1 of 2 adults over 65 years suffered from OA in the US, only [2]. Unfortunately, widely used clinical approaches (lavage, periosteal grafts, subchondral drilling, microfracture, mozaicplasty, etc.) are far from integrating cartilage tissue to bone biologically. Especially, following mozaicplasty, a gold standard for the repair of large size articular cartilage defects, subchondral bone remains exposed to cartilage tissue through the gaps between inserted plugs. This allows for the advancement of capillaries from bone tissue to cartilage, which eventually leads to formation of a fibrovascular tissue that is mechanically inferior to and biologically different from articular cartilage. The native cartilage-bone interface, on the other hand, was designed to prevent progression of capillaries to cartilage zone by a tiny membrane called tide-mark. Therefore, due to insufficiency of current procedures to form a tide-mark-like zone, repair/regeneration of OA remains as a challenge and new approaches are needed to remedy this problem.

Function, Composition and Structure

One of the critical functions of native cartilage-bone interface is to minimize the stress concentrations, which may occur between the two tissues when load is applied to one, to form a smooth transition region in order not to overload either tissue. This function is a direct consequence of hierarchical organization of the extracellular matrix (ECM) components forming the interface. Another function of the interface is not to allow capillary development from vascularized bone tissue to avascular cartilage tissue [3], which is accomplished by the tide-mark.

Cartilage-bone interface at the knee joint has a thickness of around 100-200 μm [4] and structurally contains three different yet uninterrupted compartments, namely, cartilage, mineralized cartilage, and bone, containing various cell phenotypes such as chondrocytes, hypertrophic chondrocytes and osteoblasts, respectively [5]. The major ECM components contained in these sub-tissues are collagen type II and glycosaminoglycans; collagen type I, glycosaminoglycans, minerals; and collagen type I and minerals, respectively (Table 1).

| Tissue         | Cell Type             | Components                  |
|----------------|-----------------------|-----------------------------|
| Cartilage      | Chondrocyte           | - Collagen type II          |
|                |                       | - Glycosaminoglycan         |
| Mineralized cartilage | Hyperthrophic chondrocyte | - Collagen Type I, X         |
|                |                       | - Glycosaminoglycan         |
|                |                       | - Mineral                   |
| Bone           | Osteoblast            | - Collagen type I           |
|                |                       | - Mineral                   |

Table 1: Compositional distribution in cartilage-bone interface.

Another school of thought for the hierarchical organization of the cartilage-bone interface is the belief of graded change in the composition of ECM components [6-9]. Although the compartmental organization approach for cartilage-bone interface components has been dominant for years [5,10], recent studies characterizing cartilage-bone, as well as tendon-bone, interface at microscopic dimensions found that mineral composition is changing gradually across the interface [4,11,12].

With the recent increased interest in the field of interface tissue characterization, we are now more equipped with the structure and composition of cartilage-bone interface, which could be translated into design and fabrication of scaffolds for such applications.

Scaffolds for Cartilage-Bone Interface

One of the methods utilized for the treatment of damages associated with the cartilage-bone interface at the knee is cell therapy (Table 2), applied either directly or in conjunction with scaffolds [13]. Even though cell therapy is a well-recognized and a versatile technique used for regenerative purposes, it has its own drawbacks such as difficulties...
associated with cell isolation, proliferation, storage, transfer, cost (for allogenic sources) and immune system reactions [14,15]. In addition, this technique has limitations due to potential infection, pathogen transfer and tumor growth. Therefore, cellular (with prementioned risks) or acellular scaffolds remain as an attractive alternative for direct cell therapy.

| Therapy       | Cell          | Scaffold |
|---------------|---------------|---------|
| Cell-based    | Direct cell transplantation | ✓ ✓ |
|               | Cellular scaffolds | ✓ ✓ |
| Cell-free     | Acellular scaffolds | ✓ ✓ |

Table 2: Options for regeneration.

As the name implies, cellular scaffolds are structured and shaped biomaterials incorporated with cells. Acellular scaffolds, on the other hand, contain no cells and may be incorporated with chemotactic agents to recruit endogenous cells into the scaffold to initiate regeneration. Cell recruitment represents a more recent development in the regenerative engineering, and is seen as a promising approach. This method was successfully used with 3D printed polycaprolactone (PCL) scaffolds in a sheep model for meniscus regeneration [16]. However, cell recruitment has not been utilized in the regeneration attempts for cartilage-bone interface at a scale of 100-200µm thickness.

Scaffolds utilized for cartilage-bone applications were traditionally designed either as unitary homogeneous [17] or layered configurations [10,18]. However, recent investigations at microscopic levels showed that the osteochondral interface exhibits a gradual change in the composition of the matrix components [12], leading to a paradigm shift in the understanding of scaffold design. Therefore, more recent investigations focused on the design and fabrication of graded scaffolds for osteochondral interface applications [6,7,19]. In one of these studies, tricalcium phosphate (TCP) mineral was embedded in PCL nanofibers to fabricate scaffolds with gradually changing TCP concentrations [6]. The same group of researchers produced functionally graded PCL scaffolds incorporated with insulin and beta-glycerophosphate (beta-GP) biomolecules with varying concentrations in opposite direction [7]. Human adipose derived stem cells (hADSCs) were seeded on the graded scaffolds to form osteochondral-like structures. In a similar study, poly (D,L-lactide-co-glycolic acid), PLGA, microspheres were enriched with growth factors, stimulating cartilage and bone formation, in a gradually changing fashion, and their capacity were investigated for the treatment of osteochondral defects in a rabbit femoral condile model [9]. Putting together, these investigations demonstrate that scaffolding for osteochondral tissue engineering is still under development, and that progress in technology will certainly open up new avenues for the design and fabrication of more realistic scaffolds [20].

Cell Options

Commonly used cell options for the osteochondral interface tissue engineering include but are not limited to chondrocytes, osteoblasts and stem/progenitor cells. Chondrocytes and osteoblasts could be harvested through surgical procedures, proliferated, and used as cell sources to be seeded on scaffolds to form cartilage and bone regions, respectively. Nevertheless, limited availability of these cells due to donor shortages, and morphological changes associated with some cell types, especially for chondrocytes, when cultured under in vitro conditions, stem/progenitor cells remain as a more favourable cell sources.

Chondrocytes harvested from sheep [21] and bovine [22] are commonly used as cell sources for osteochondral regeneration. In addition, co-culture of chondrocytes and osteoblasts obtained from similar sources is also an attractive strategy to form osteochondral-like structures. In this regard, Cao et al. seeded chondrocytes on one side of the 3D printed PCL scaffold and osteoblast on the other side, and observed a mixture of the two distinct cell phenotypes in the middle zone [23].

In the category of stem/progenitor cells, bone marrow derived stem cells, adipose derived stem cells, synovial stem cells and embryonic stem cells were demonstrated to be appropriate cell choices for osteochondral regeneration [24,25]. Specific differentiation of these cells into proper lineages is achieved by incorporating relevant biomolecules into scaffolds in a specially controlled manner. For example, transforming growth factor beta1 (TGF-beta1) and transforming growth factor beta3 (TGF-beta3) are known to trigger these cells to differentiate into chondrogenic lineage, while bone morphogenic protein 2 (BMP2) can lead to osteogenic differentiation [26-29]. Therefore, incorporating TGF-beta1 or TGF-beta3 into one side and BMP2 into the other, also coupled with the stem cells may form appropriate conditions to generate structures resembling the osteochondral interface.

Regeneration Attempts

Biomaterial/scaffold and biomolecule selection, as well as employment of appropriate cells could play significant roles in osteochondral regeneration efforts. Biomaterials are required to be biologically compatible. They should not create any adverse effects in terms of cell attachment, proliferation, morphology, membrane characteristics and cellular activity such as ECM production and expression of relevant markers. Biomolecules should contribute to the cellular activities in the direction observed in their native environment. Similarly, selected cells are expected to have properties or should have capacities to perform activities of cells present in the target tissues. Tissue regeneration may be possible at best only if these three parameters are chosen appropriately, and combined to create a synergistic effect to be able to generate structures, compositions and functions observed in native osteochondral interface.

Biomaterial/scaffold design targeting osteochondral interface regeneration could be classified as homogeneous (first generation), compartmentalized (second generation), and graded (third generation) scaffolds. Although, the first two generation scaffolds are still widely investigated due to their ease of design/manufacture and simplicity in terms of composition and structure, the third generation graded scaffolds are now more prevalent due to their biomimicry. These scaffolds could be designed to contain multiple biomolecules positioned into the scaffold in a controlled fashion, and to release these biofactors in a time-dependent manner to also serve as controlled delivery devices. Such scaffolds were successfully employed for interface regenerative engineering studies both in vitro and in vivo [6,7,28]. These investigations revealed that when the right cells are used in conjunction with an appropriate scaffold and/or scaffold biofactors combination choice, native osteochondral interface could be approximated both biologically and physically.
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