Histochemical characterization of human osteochondral tissue: comparison between healthy cartilage,arthrotic tissues, and cartilage defect treated with MACI technique

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Abstract

Matrix-induced autologous chondrocytes implantation (MACI) is a promising technique for the treatment of articular cartilage lesions, but long term outcome have to be established. We developed and optimized specific techniques of histochemical staining to characterize healthy and pathologic osteochondral tissue. Seven different staining protocols were applied to assess tissue architecture, cells morphology, proteoglycan content, and collagen fibers distribution. Potentialities of histochemical staining and histomorphology of biopsies from second look arthroscopy will be presented.

Material and Methods

Three sample types of human osteochondral tissue were considered: healthy (lateral femoral condyle, female, 78 years), arthrotic (tibial plate, female, 84 years), and regenerated with MACI technique (lateral femoral condyle, male, 51 years, biopsy obtained during second-look arthroscopic surgery in symptomatic patient, 1 year after MACI). Tissues were fixed in 10% buffered formalin for 24h, decalcified in EDTA for 24h, embedded in paraffin, and cut transversally in 3-5um slides. Sections were subjected to one of the following protocols: Hematoxylin-Eosin Y (H&E), Safranin-O (SO), Alcian Blue pH2.5 (AB), Toluidine Blue (TB), Astra Blue pH2.5 (AsB), Azan (Az) and Picrosirius Red (PR). Stained sections were dehydrated, mounted with acrylic medium and observed at magnifications ranging from 40 to 500 times in white and polarized light. Tissue sections were evaluated for tissue regularity, proteoglycan content, spatial distribution of chondrocytes and collagen fibers orientation.

Introduction

Surgical therapies for the treatment of articular cartilage lesions can be grouped into reparative and restorative techniques. Clinical results showed reparative techniques lead to the formation of fibrocartilaginous tissue. Differently, restorative procedures can repair the osteochondral defects with a hyaline-like cartilage [1]. In last years, autologous chondrocytes implantation after in vitro expansion has been further improved by MACI (Matrix induced autologous implantation chondrocytes) technique [2], using three-dimensional scaffolds made of bioreabsorbable biomaterials. To assess the efficacy of long-term treatment, the histological evaluation of osteochondral tissue plays a key role in the distinction between hyaline cartilage and fibrocartilage. The study aimed at evaluating and optimizing histochemical staining protocols to compare healthy and arthrotic osteochondral tissue to regenerated tissue after MACI.

Results

H&E successfully characterized morphology and spatial organization of the chondrocytes. SO, AB, AsB, TB stained proteoglycans. According to its specificity [3], SO was more effective than Toluidine Blue in revealing proteoglycan depletion in pathologic tissue. AB and AsB staining can be affected by pH and temperature. Collagen fibers were stained by both Az and PR, but the latter protocol was less complex and samples can be viewed also in polarized light.

Discussion

The comparison of healthy and arthrotic tissue showed cartilage wear and surface fibrillation (fig. 1a, 1b). Regenerated tissue showed to be fibrocartilaginous in the chondral side (depletion of the cationic staining, misalignment of the chondrocytes distribution and loss of normal orientation in collagen fibers) and hyaline-like in the bone side (Fig. 1c). Moreover a deep tear, compromising
Histochemical characterization of human osteochondral tissue; comparison between healthy cartilage, arthritic tissues, was found between the two layers, showing a non complete regeneration of the lesion.

References


Figure 1. (a) healthy, (b) arthritic, and (c) regenerated osteochondral tissue. PR was obtained in polarized light. Arrows indicate tissue tear. Original magnification: 40x (a and b), 200x (c).