PRESENCE OF CALCIFIED TISSUE IN THE HUMAN TEMPOROMANDIBULAR JOINT DISC

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Summary—Large calcified areas were observed in the articular discs of the temporomandibular joint from five patients suffering from articular dysfunctions. The calcified regions were always located inside the fibrous tissue of the discs. They had a woven bone-like morphological pattern and consisted of a compact mineralized tissue containing cells in irregular lacunae. In all the samples the calcified tissue was completely surrounded by a mineralizing border rich in cells and variously arranged collagen fibrils. Energy-dispersive spectrometry showed that mineralized regions contained large amounts of Ca and P. X-ray powder diffraction showed that mineralized regions contained large amounts of Ca and P. X-ray powder diffraction identified the crystals in these areas as hydroxyapatite.

Key words: temporomandibular joint, articular disc, calcification, TEM, EDS, XRD.

INTRODUCTION

Several studies have been performed on the microscopic and ultrastructural features of the fibrous component of the temporomandibular joint disc (Oberg and Carlsson, 1979; Bhussry et al., 1991; Berkovitz et al., 1992) and on its structural modifications following articular dysfunctions (Carlsson et al., 1967; Sharawy et al., 1987; Helmy et al., 1988, 1989; Piacentini et al., 1994; Marchetti et al., 1995). Less is known about the non-fibrous extracellular-matrix components and the possible metaplastic modifications of the tissue.

The chemical composition of the articular disc of the temporomandibular joint is modified during maturation and growth. In particular, increased amounts of chondroitin-6-sulphate and keratan sulphate/chondroitin-4-sulphate and the appearance of a cartilaginous phenotype are observed during mandibular growth. These variations were seen as a response of the articular disc to changes in the mechanical loading (Mills et al., 1988; Carvalho et al., 1993). Cartilaginous cells have also been found in discs from ageing humans (Oberg and Carlsson, 1979; Bhussry et al., 1991; Berkovitz et al., 1992). Moreover, small deposits of mineralized material in the fibrous tissue were observed among the age-related changes in the articular disc. They were described as foci of variously shaped mineral precipitates localized in the intercellular matrix (Shaw and Molineux, 1994). Our aim now was to investigate the morphological and chemical characteristics of calcification occurring in disc tissue due to articular dysfunctions.

MATERIALS AND METHODS

The study was made on five discs from individuals aged 45–53 years belonging to a group of 71 patients who underwent arthroscopy and arthrotomy because of large functional defects of the mandibular joint. To select patients for arthroscopy and surgery our diagnostic and therapeutic protocols estimate the functional damage by reference to the degree of articular pain and the clinical history. The functional damage we observed in all our patients was consistent with the presence of more than one of the defects involved in the hypothesis of internal derangement and arthrosis of the mandibular joint (Farrar, 1972; Stegenga et al., 1989). The five patients who presented calcified disc lesions had very similar symptoms and clinical histories. They all remembered having articular noise and some dynamic limitation of the condyles from many years, and they all began to suffer from articular pain more than 5 years ago. They had all been given conservative treatment (physiotherapy and bite-planes) which had produced only some small and temporary relief. The articular pain began as a small and transitory problem and increased to become a very important and disabling pathology in a variable but sudden period before we first saw them. The presence of calcifications within the disc was diagnosed radiologically (orthopantomography).
tomography, computerized tomography, three-dimensional computerized tomography), revealing the presence in each disc of large internal radiopaque areas. These individuals presented functional defects with serious limitation of motion and articular crepitation and pain, but they did not have any signs of rheumatoid or autoimmune diseases as evidenced by clinical history and clinical, radiological and blood chemistry tests.

After surgical removal the excised discs were rapidly washed in buffered saline solution and fixed in a mixture of glutaraldehyde (2.5%) and paraformaldehyde (2%) in 0.1 M Na cacodylate buffer, pH 7.4, for 5 hr at 4°C, rinsed in the same buffer solution and postfixed in 1% OsO₄ in 0.2 M collidine buffer, pH 7.4, for 2 hr at 4°C. The samples were dehydrated and embedded in epoxy resin and sectioned without mineralization. Semithin sections (0.5 μm) were stained with toluidine blue and studied by light microscopy; ultrathin sections (80 nm) were stained with uranyl acetate and lead citrate and studied by transmission electron microscopy.

Subsequently the sectioning surfaces of the epoxy resin blocks were sputtered with a carbon film and observed in a scanning electron microscope equipped with a Tracer Northern energy-dispersive X-ray spectrometer to characterize and identify the chemical composition of the samples. A qualitative analysis of a spectrum in the data memory of the chemical elements from Na to U was performed with the IDENT program. The conditions for the microprobe analysis were: accelerating voltage of 15 kV, probe current of 6 × 10⁻⁹ A, working distance of 39 mm and counting time of 30 sec.

The same samples were analysed by X-ray powder diffraction to characterize the crystalline phases.
Fig. 6. Electron micrograph of the disc region of Fig. 3. The cells are characterized by abundant cytoplasm with rough endoplasmic reticulum, Golgi apparatus and filaments. The pericellular matrix consists of amorphous material and rare and isolated collagen fibrils. Magnification x6000.

Fig. 7. Electron micrograph of a degenerated cell completely surrounded by calcified matrix. The collagen fibril banding is not completely masked by the presence of crystallites. Magnification x6500.

(Bonucci and Graziani, 1975). The hard portion of each sample was separated from the soft tissue, ground in an agate mortar and analysed in the natural state using a Philips PW 1800/10 diffractometer equipped with Digital Microvax 2000 and software APD-1700. Characteristics and variables of the diffractometer were: radiation Cu Kα, 50 kV, 30 mA; graphite monochromator range 2θ 20°.

RESULTS

Light microscopy

In all the discs an unmineralized soft tissue surrounded and limited the calcified discal area. In two discs the non-calciﬁed tissue consisted of compact bundles of collagen ﬁbres with occasional elongated ﬁbrocytes (Fig. 1). In the other three samples, collagen ﬁbre bundles were disarrayed and sometimes intermingled with deposits of dense basophilic and apparently amorphous material (Fig. 2). One of these discs had a large, superﬁcial zone consisting of a loose tissue containing fewer ﬁbres and large, basophilic and sometimes binuclear cells (Fig. 3).

In the inner part of each disc a large area (0.3–1 mm²) of hard, calcified tissue was present. It consisted of compact mineralized tissue containing cells enclosed in irregular lacunae (Figs 4 and 5). The calcified areas were variously shaped and had irregular contours. They were surrounded by an unmineralized, dense and highly basophilic layer (10–40 μm), consisting of amorphous or granular material containing numerous oval or elongated cells. This layer was continuous with the external ﬁbrous tissue of the discs containing compact bundles of ﬁbres and elongated ﬁbrocytes.

Transmission electron microscopy

The region of the disc consisting of loose tissue by light microscopy was ultrastructurally characterized by the presence of roundish or oval chondrocyte-like cells (Fig. 6). They exhibited irregular profiles with short and slender processes, abundant cytoplasm rich in rough endoplasmic reticulum, and they were sometimes binucleated. The pericellular matrix consisted of a loose, ﬁnely granular and less electron-dense material and scanty collagen ﬁbrils.

The calcified areas of the discs (Fig. 7) consisted of a highly mineralized and compact tissue where packed collagen ﬁbrils were almost completely masked by electron-dense confluent crystallites. Necrotic or severely degenerated osteocyte-like cells with scanty cytoplasm were present in this mineralized tissue. They were contained in lacunae with indented proﬁles because of a brush border of projecting crystallites. A pericellular space separated the cell surface from the mineralized matrix.

The unmineralized tissue layer outside of the calcified areas consisted of a loose matrix with bundles of variously arranged collagen ﬁbrils (Figs 8 and 9).
Fig. 8. Electron micrograph of the mineralization front close to the calcified tissue. Numerous elongated cells with irregular protrusions are distributed within an extracellular matrix rich in collagen fibrils. Magnification ×6100.

Fig. 9. Electron micrograph of the border of the mineralizing area. A cell is in contact with the calcified matrix on one side and with collagen fibrils of the mineralizing matrix on the other. At the top a blood capillary (bc) is present. Magnification ×6560; bar = 1 μm.

These fibrils were thin but fairly dense and they seemed to be continuous with those already covered by the deposition of crystallites. Numerous oval or elongated cells with thin cytoplasmic processes were contained in the matrix of this mineralization front. Blood vessels were present among the cells (Fig. 9).

Energy-dispersive X-ray spectrometry

Analysis of the chemical composition of the calcified portions of the discs showed Ca and P signals in all samples (Fig. 10). The same investigation (data not shown) performed on the outer fibrous portions of the discs gave no signals.
Calcification of human temporomandibular joint disc

Fig. 10. Energy-dispersive X-ray spectrum (EDS) of the mineralized zone of one disc. In the inset a scanning electron-micrograph of the same area is shown, photographed after EDS analysis. Vertical full scale (VFS) = 512; inset magnification x400; bar = 10 μm.

Fig. 11. X-ray powder diffraction pattern from the calcified disc of Fig. 10 showing the peaks of hydroxyapatite crystals. The graphite signal is due to the sputtering for scanning electron microscopy.

DISCUSSION

We earlier demonstrated that fibrous tissue of the articular disc may undergo structural modifications following different functional diseases (Piacentini et al., 1994; Marchetti et al., 1995). The main modifications were derangements of the fibrillar component, increased vascularization and cellular proliferation. Moreover, we observed the appearance of radio-opaque, calcified areas in some discs and in endoarticular loose bodies (Piacentini et al., 1995).

Isolated foci of mineral precipitates have been occasionally described in tissues (Shaw and Molineux, 1994). They appear as restricted and occasional features consisting of single aggregates of few crystallites without any structural arrangement. The present morphological (light and transmission electron microscopy) and chemical study demonstrates that large areas of the discs may undergo histological changes producing a woven bone-like tissue. In all samples the mineralized areas were within the discs and were surrounded by a conspicuous envelope of fibrous tissue.

The presence of cells completely enclosed in lacunae of the calcified matrix and the existence of a mineralizing border rich in cells and collagen fibres outside the calcified tissue suggests that this calcification process is not occasional and restricted. On the contrary it appears to result from a complex process that begins in the interior of the discs and subsequently extends because of aggregation of the calcifying nodules. These events result in a woven bone-like mineralized tissue enclosing the cells that had produced the extracellular matrix. These cells could be modified fibroblasts or they may have differentiated from mesenchymal cells that are usually present in the fibrous tissue. The mineralization process with deposition of crystallites in the extracellular matrix may be promoted by vascularization developed in the normally avascular fibrous tissue of the disc. The X-ray diffraction patterns showing the presence of hydroxyapatite crystals in the calcified tissues strengthen the opinion that a specific process of bone-like mineralization occurred in the articular discal tissue.

These kinds of lesions, apart from the structural peculiarities of the mandibular joint, have some clinical and morphological similarities to the production of articular osteophytes. They too are slowly and progressively formed as a result of excessive functional demands and present a structural modification of articular soft tissues (Takenoshita, 1982, 1987).

Our findings demonstrate that disc tissue may undergo profound metaplastic modifications producing bone-like calcified tissue. These morphological modifications occur after articular dysfunction producing alterations in the biomechanical forces applied to the disc. Vascularization and the possible subsequent modifications in water content and the composition of the extracellular matrix may activate mechanisms leading to the deposition of crystals, their aggregation and then the extension of the mineralized area.

Calcification of the fibrous tissue always begins inside the disc, as we have previously described (Piacentini et al., 1994; Marchetti et al., 1995). Only afterwards do these processes extend to the superficial parts of the disc. In our opinion this sequence of events suggests that the interior of the disc is most susceptible to variations in mechanical stimuli produced among the articular structures. The
morphological patterns described here demonstrate that the modifications in the articular disc result from excessive functional stress in the articular spaces. These lesions can be considered to be the result of progressive and slowly extending pathological processes in which the articular disc loses its capacity to absorb and reduce the functional stress caused by the condier movements on the functional surfaces of the condyle and glenoid.

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REFERENCES


