The knee joint loose body as a source of viable autologous human chondrocytes

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Abstract

Loose bodies are fragments of cartilage or bone present in the synovial fluid. In the present study we assessed if loose bodies could be used as a source of autologous human chondrocytes for experimental purposes. Histological examination of loose bodies and differential enzymatic digestions were undertaken, the isolated cells were cultured in alginate bead microspheres and immunolocalisations were undertaken for chondrogenic markers such as aggrecan, and type II collagen. Isolated loose body cells had high viability (>90% viable), expressed chondrogenic markers (aggrecan, type II collagen) but no type I collagen. Loose body cells may be a useful source of autologous chondrocytes of high viability.

Introduction

Loose bodies (also known as rice bodies and knee joint mice) are fragments of cartilage or bone that float freely within the synovial fluid component of the knee joint. They may occur in single or multiple forms but generally only affect a single knee. Loose bodies are classified as either stable or unstable. The former are located in fixed positions in the knee and are generally well tolerated and asymptomatic while the latter are free to move around the knee joint and may be the cause of pain, knee joint swelling, joint weakness and may cause the knee to lock abruptly. Loose bodies have a traumatic origin such as dislocation of the patella or a complication arising from an orthopaedic procedure and their occurrence is more likely in patients affected by osteoarthritis (OA) or rheumatoid arthritis (RA). Smooth bodies are classified as fibrous, cartilaginous and osteocartilaginous.

Fibrinous loose bodies result from intra-articular bleeding or by death of synovial tissue associated with tuberculosis, OA or RA. Cartilaginous loose bodies are caused by traumatic injury to the OA joint. Osteocartilaginous loose bodies are caused by fractures, osteochondritis dissecans, inflammation, synovial chondromatosis or tumours (osteochondromas).5,6 Loose bodies are normally small, but cases of giant loose bodies have also been reported.8

In the present study histological examination of loose bodies showed to our surprise that they were highly cellular containing large numbers of viable chondrocytes and suggested that they may be a potential source of autologous human chondrocytes for repair strategies. We subsequently went on to isolate this chondrocyte population, they displayed high viability and were highly proliferative compared to chondrocytes isolated from residual tibial and femoral articular cartilage from the same knee joint replacements. The loose body chondrocytes expressed type II collagen, aggrecan and 5D4 KS in 3D alginate bead culture but no detectable type I collagen.

Materials and Methods

All chemicals and supplier details are as previously indicated described.10,11 Monoclonal antibodies to aggrecan (clone 4D11 2A9), type I collagen (clone I-8H5), type II collagen (clone II-4C11), biotinylated anti-mouse IgG and anti-rabbit IgG secondary antibody, horseradish peroxidase conjugated streptavidin have been described previously.10,11

Tissues and cells

Loose bodies harvested from total knee replacement surgical discards from 6 patients (median age 56 years, 3 male, 3 female) of the Orthopaedics Clinic at North Shore Private Hospital, St. Leonards, were obtained with informed consent under ethical approval of the human ethics and care committee of the Royal North Shore Hospital who approved all procedures.

Isolation and culture of chondrocytes from the loose bodies

Chondrocytes were enzymatically isolated from loose bodies by sequential digestion with: i) 0.1% (w/v) pronase in DMEM-F12 media containing 10% (v/v) FCS for 2 h at 37°C; ii) 0.05% (w/v) clostridial collagenase in media for 4 h; and iii) overnight digestion with collagenase.11 The cells were spun down (10 min x 800 g) and cell viability and numbers determined on a haemocytometer using trypan blue exclusion. Examination of the front and side scatter characteristics of these cells by flow-cytometry was similar to results obtained earlier with only one major cell population evident.11 The cells were established in monolayer culture in 75 cm² canted neck flasks at a density of 100,000 cells/mL in DMEM-F12 + 10% FCS + antibiotics under an atmosphere of 5% CO2 and 98% humidity at 37°C. After cellular attachment overnight, the flasks were rinsed in PBS to remove non-adherent cells and cultured in DMEM-F12 media + 10% FCS + antibiotics, with media changes every 3 days. The cells became confluent on day 3-6, the cells were sub-cultured up to passage 9.

Preparation and culture of loose body cells in calcium alginate beads

The loose body cells from one 75 cm² canted neck flask were detached using trypsin-EDTA and pelleted (800 g x 10 min) then washed in sterile DMEM-Hams F12 + 10% v/v FCS and resuspended in 2 mL of DMEM-Hams F12-FCS. Cell numbers and viability assays were measured on an aliquot using a haemocytometer and trypan blue exclusion. A known number of cells (95% viability) were spun down again and
dispersed at a density of $3 \times 10^6$ cells/mL alginate in sterile isotonic saline. This mixture was loaded into a 2 mL syringe and extruded drop-wise through a 23 guage needle into an agitated solution of sterile CaCl$_2$ (102 mM) in a laminar flow hood to maintain sterility. After 10 min curing time the calcium alginate beads (~10 mL/30,000 cells) were established in culture in small petri dishes (100 beads/plate) in DMEM-F12 + 10% FCS + 50 µg/mL ascorbic acid (5 mL media/plate). The plates were incubated at 37°C, in an atmosphere of 5% CO$_2$ in air, with a humidity of 98% and the medium replenished every 48 h. The loose body cells were cultured up to 4 weeks and samples of beads collected after 2, 3 and 4 weeks of culture as indicated earlier.$^{11,13,14}$ The remaining beads were rinsed in isotonic saline and solubilized in 55 mM trisodium citrate in 150 mM NaCl (2 mL media/plate) and the cells spun down at 800 g x 10 min. The cells were then either cryopreserved in liquid nitrogen or were re-established in monolayer culture.

**Histological processing of alginate beads**

Beads were fixed 3 h in 10% (v/v) formalin, 85% (v/v) ethanol, 5% (v/v) acetic acid transferred overnight into 70% (v/v) ethanol and embedded in paraffin then sectioned at 4 µm thickness and attached to SuperFrost ultraPlus positively charged microscope slides. The slides were de-paraffinised in xylene (2 changes x 5 min), re-hydrated through graded ethanol washes (100-70% v/v) to water.

**Histochemistry**

Bead and loose body sections (4 µm) were stained for 10 min with 0.04% (w/v) toluidine blue in 0.1 M sodium acetate buffer, pH 4.0 followed by a 2-min counterstain in 0.1% (w/v) fast green FCF. Sections were also stained in Mayer’s Haematoxylin (5 min), rinsed in tap water blued in Scotts Blueing solution (1 min) and counterstained in 0.0001% (w/v) eosin (5 min).

**Immunolocalisation of type I and type II collagen, aggrecan and keratan sulphate**

Endogenous peroxidase was blocked in bead samples with 0.3% (v/v) H$_2$O$_2$ for 10 min then blocking undertaken in DAKO non-protein blocking agent. Aggrecan immunolocalisation were pre-digested with chondroitinase ABC (0.1 U/ml) in 50 mM Tris HCl pH 7.2 + 2% (w/v) BSA for 1 h, type I and II collagen immunolocalisations were pre-digested with proteinase K for 6 min and bovine testicular hyaluronidase (1000 U/ml) for 1 h at 37°C in phosphate buffer pH 5.0. The bead sections were incubated with anti-aggrecan (1/10,000 dilution), anti-type I collagen (1/500 dilution), anti-type II collagen (1/200 dilution) and MAb 5-D-4, anti-KS (1/1000 dilution) in TBS + 2% (w/v) BSA overnight at 4°C then biotinylated anti-mouse and anti-rabbit IgG antibodies and horse-radish peroxidase conjugated streptavidin were added using Nova RED substrate for visualisation. Negative control sections were run omitting primary Ab or using an irrelevant primary antibody. Both yielded negative results. The stained specimens were examined using a Leica photomicroscope linked to a DFC 480 digital camera using bright-field illumination.

**Results**

Loose bodies were observed in 12 of 18 total knee joint replacements, and typically 2.5 mm in size, smooth and glistening. Histological examination demonstrated a high cell density and abundant deposition of proteoglycan in the loose bodies (Figure 1). The loose body contained a central necrotic core where the cells were arranged in clumps. Closer inspection within the loose body sections revealed a tran-
Discussion

The present study arose from a series of failed attempts to isolate articular chondrocytes from the articular remnants of discarded surgical material from total knee arthroplasty patients. This was not an unexpected finding given that these specimens had little residual cartilage, displayed severe eburnation, loss of > 50% of the menisci and marginal osteophytic features of advanced knee-joint degeneration. The chondrocytes that were isolated typically were 50-60% viable, had poor replicative capability, and were obtained in insufficient numbers to warrant further investigation. Incidental observations on a number of these knee replacement tissues drew our attention to small glistening loose bodies present in a significant number of cases (12/18). On closer inspection histologically we were surprised to see that the loose bodies were highly cellular and contained abundant proteoglycan, the same could not be said of the residual articular cartilage of these joints. A necrotic core was a common finding particularly in the loose bodies ≥ 4 mm in size however this was to be expected given the advanced degenerative features of these knee specimens. We subsequently developed a protocol to isolate the loose body chondrocytes at ≥ 95% viability.

The therapeutic use of autologous chondrocytes in isolation13 or in combination with mesenchymal stem cells16 in the development of cartilage repair strategies are of considerable interest in repair medicine.17 The present study describes a simple convenient procedure for the isolation of these cells from loose bod-

Figure 2. Cellular morphologies of selected regions of the loose body depicted in Figure 1. H&E stained loose body tissue sections depicting cellular morphology: (a) the surface zone (SZ) cells are small and of a flattened morphology and are located adjacent to the transitional zone (TZ) where better defined rounded cells are located (b), this leads into the cartilaginous zone (CZ) where the cells are larger again (c) and reminiscent of cells in the cartilage rudiments in human foetal joint development. The cells of the core region (d) are somewhat misshapen and many are dead and arranged in clumps. Toluidine blue-fast green and H&E stained loose body sections of surface zones 1 and 2 (see Figure 1) of the loose body (a-e). Two surface zone areas can be differentiated on the basis of cellular distributions (SZ1, SZ2) and GAG staining (e). The cells in the cartilaginous zone (CZ) are larger, round-ed and surrounded by a matrix containing toluidine blue stained proteoglycan. The surface zone cells have differing morphologies to the deeper chondrocytic cells and do not lay down as high levels of proteoglycan as the CZ cells (f, g, h, i).
ies in the knee joint and could be of further application. We initially examined the cells isolated by flow cytometry, front and side scatter data demonstrated a single cell population with only a small proportion of dead or non-viable cells. Loose body cell numbers were initially expanded in monolayer culture which eliminated these dead cells and demonstrated a homogeneous cell population of cells with a typical cobblestone morphology and of high replicative capability and viability. Further culture of these cells encapsulated in alginate microspheres allowed us to demonstrate the synthesis of type II collagen, aggrecan, and KS but not of type I collagen by the loose body cells. Chondrocytes do not divide in this culture system but produce cartilage specific ECM components. The lack of type I collagen synthesis was further evidence of the chondrocytic pedigree of the isolated cell population. Furthermore morphometric image analysis of the immunolocalised bead sections over a 2-8 week culture period demonstrated a steady increase in chondrocyte ECM products plateauing at 4 weeks of culture. Our laboratory has formerly used this culture system with intervertebral disc cells, meniscal cells and articular chondrocytes. 

The microcarrier cell culture system was first introduced in 1967 by van Wezel. These spheres were typically 125-250 μm in size and made from DEAE-dextran, bioglass, polystyrene, acrylamide, collagen or alginate and were available commercially as dextran beads (Cytodex, GE Healthcare), alginate (GEM, Global Cell Solutions), collagen (Cultispher, Percell) and polystyrene (Solohill Engineering). The surface chemistries of these microspheres were conducive to cell attachment and they were robust enough for use in spinner cultures. Cancer cells were one cell type which was cultured on and within these spheres in an effort to simulate a spheroid cell mass and promote cellular cross-talk to maintain cell viability and provide a culture system suitable for the assessment of anti-cancer agents in vitro. In 1992 alginate microspheres were developed as an encapsulated system for the culture of cells of a chondrogenic background. The high negative charge of the mannuronic and guluronic acid alginate copolymer was envisaged to reproduce the high fixed charge density 3D microenvironment chondrocytes experience in cartilaginous matrices. Chondrocytes rarely undergo division when surrounded by a mature 3D ECM however with maturity terminal differentiation can lead to hypertrophy. Neither of these features were observed by the loose body chondrocytes in the alginate bead cultures examined in the present study (Figures 3 and 4) however these features were evident in histological sections of the loose bodies and typical of the columnar arrangement of growth plate chondrocytes (Figures 1 and 2) suggesting that the loose bodies may also be useful as models of the growth plate. The deer antler has been suggested as a growth plate model, mainly stemming from its impressive rate of growth however deer antler has a tissue organisation quite dissimilar to that of the long bone growth plates whereas, as seen in the present study, loose bodies have similar cellular arrangements to these found in growth plate cartilage. Loose bodies can be stored in standard tissue culture media containing foetal calf serum and antibiotics to maintain cell viability for at least 3 weeks making them a potentially useful research tool and they could be considered as explants without cut edges. In the present study, the distribution of chondrocytes in alginate beads and absence of cell division were well illustrated in the negative control bead sections (Figure 4 h). Long term culture (8 months) of chondrocytes in alginate beads have been reported to lead to morphological changes in the outer bead cells with the appearance of cells of a flattened morphology similar to annular fibrochondrocytes in the intervertebral disc, whereas cells located more centrally within the bead have well

Figure 3. Assessment of the polydispersity of the loose body cell preparation isolated in this study and morphometric image analysis of staining intensities of individual beads stained with antibodies to aggrecan, type I, II collagen in alginate bead culture. a) Flow cytometric analysis of the loose body cell preparation isolated by front and side scatter characteristics; the gated area labelled with an asterisk designates dead or non-viable cells. b) Immunolocalised Collagen I, II and aggrecan in bead sections from 2, 3 and 4 weeks of culture (4 beads per time point) were subjected to morphometric image analysis to quantify the staining using Adobe Photoshop software. Integrated density of the stained pixels were measured, data shown is x 106. Immunolocalisation of aggrecan (c,f) and type II collagen (d,g) in sections of beads from 2 and 3 weeks of alginate bead culture.
defined rounded chondrocytic morphologies.28 A re-evaluation of the microsphere culture system has occurred in the last few years where cancer cells are encapsulated in an environment permissive to cell-cell cross-talk. Various matrix components can also be introduced into the bead in an effort to develop a more appropriate cell culture micro-environment similar to that found in vivo.22,23 This culture system has been used to culture macrophages and fibroblasts in breast cancer,25 invasive hepatocellular cells,32 and epithelial-stromal cells in prostate cancer.36 Porous chitosan-alginate microspheres have also been developed to examine prostate cancer37 and glioma.38 Hepatocarcinoma spheroids have also been prepared using gelatin microspheres,39 Pullulan40 and controlled release rhBMP2 in 3D printed porous hydroxyapatite,41 injectable nanofibrous microspheres,41 and arginine-chitosan BMP-2 nanoparticle cell delivery vehicles for bone repair have also been developed.12,33
These promising initial findings with the loose body chondrocytes warrants further studies to determine the gene expression profiles of the loose body chondrocytes compared to articular and growth plate chondrocytes. Human chondrocytes are difficult to source and it is only relatively recently that knee chondrocytes have become available commercially with most suppliers previously using hip cartilage as a tissue source. Pascual-Garrido et al.42 found the viability of loose bodies from paediatric patients were 88% vs 92% for healthy articular cartilage. Others have also suggested that the loose body cells represent a valuable resource for autologous cell transplantation.43 Thus, the loose bodies examined in the present study should be considered a valuable cell resource rather than as a surgical discard.

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