Fibrin sealant promotes migration and proliferation of human articular chondrocytes: Possible involvement of thrombin and protease-activated receptors

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Abstract. Fibrin sealant (FS), a biological adhesive material, has been recently recommended as an adjunct in autologous chondrocyte implantation (ACI). While FS has been shown to possess osteoinductive potential, little is known about its effects on chondrogenic cells. In this study, we assessed the bioactivity of FS (Tisseel®) on the migration and proliferation of human articular chondrocytes in vitro. Using a co-culture assay to mimic matrix-induced ACI (MACI), chondrocytes were found to migrate from collagen membranes towards FS within 12 h of culture, with significant migratory activity evident by 24 h. In addition, 5-bromo-2'-deoxyuridine (BrdU) incorporation experiments revealed that thrombin, the active component of the tissue glue, stimulated chondrocyte proliferation, with maximal efficacy observed at 48 h post-stimulation (1-10 U/ml). In an effort to elucidate the molecular mechanisms underlying these thrombin-induced effects, we examined the expression and activation of protease-activated receptors (PARs), established thrombin receptors. Using a combination of RT-PCR and immunohistochemistry, all four PARs were detected in human chondrocytes, with PAR-1 being the major isoform expressed. Moreover, thrombin and a PAR-1, but not other PAR-isotype-specific peptide agonists, were found to induce rapid intracellular Ca2+ responses in human chondrocytes in calcium mobilization assays. Together, these data demonstrate that FS supports both the migration and proliferation of human chondrocytes. We propose that these effects are mediated, at least in part, via thrombin-induced PAR-1 signalling in human chondrocytes.

Introduction

Fibrin sealants (FSs) have long been utilised as an adjunct in a variety of surgical procedures to promote hemostasis and tissue sealing (1-4). In orthopaedics, FSs are used as a tissue adhesive for the fixation of osteochondral fragments and fractures (5), in spinal surgery (6), and securing perichondral grafts (7). More recently, fibrin preparations have been employed as biological vehicles for delivering chondrocytes directly to cartilage defects in order to stimulate repair processes, however with contradicting outcomes reported in the literature. Studies by Homminga et al demonstrated that chondrocytes encapsulated in FS retained their morphology and actively synthesised matrix suggesting that the adhesive served an effective matrix (8). Similarly, Hendrickson et al reported that FS-bound allogenic chondrocyte grafts displayed significantly higher glycosaminoglycan and collagen II content 8 months post-implantation (9). On the other hand, in vivo studies by Brittberg et al reported opposite effects suggesting that FS-derived scaffolds were not suitable for osteochondral healing (10). While ambiguity remains over the application of FSs in articular cartilage repair, recent studies indicate that these biological adhesives possess unique osteoinductive properties (11). Furthermore, there is increasing evidence to suggest that FSs support the growth and migration of chondrocytes (12-15). Based on the collation of these and recent studies, the use of FSs as a component of autologous chondrocyte implantation (ACI) has now been advocated.

Thrombin, a coagulative serine protease, is an active ingredient of FSs. Thrombin is ubiquitously expressed at...
sites of vascular injury where it serves to accelerate the coagulation process via proteolytic cleavage of fibrinogen (16). In addition to its role in wound healing, thrombin has been shown to induce a variety of cellular responses including proliferation (17-21), migration (22-25) and survival (26,27). These diverse biological effects are mediated through specific interaction(s) with cell surface receptors. Among the candidate thrombin receptors, members of the seven transmembrane G protein-coupled protease activated-receptor (PAR) family (PAR-1, PAR-2, PAR-3, PAR-4) are perhaps the best characterised (28). Thrombin is known to activate intracellular signalling of PAR-1, 3 and 4 via cleavage of the extracellular N-terminal domain, which unmask a ‘tethered ligand’ sequence which binds intramolecularly to a receptor domain thereby activating G protein-coupled signal transduction pathways (29,30). By comparison, PAR-2 is activated by trypsin and tryptase-associated proteases, but not thrombin (31-33). PAR-1, -2 and -4 can also be activated without proteolytic cleavage, using five to six amino-acid residue peptides corresponding to the new amino termini of the cleaved receptors (28).

The goal of this study was to assess the bioactive properties of commercial FS (Tisseel®), with particular emphasis on the thrombin component, on autologous human chondrocyte migration and proliferation in vitro. In addition, we examined the expression and localisation of PARs in cultured human chondrocytes. Our findings indicate that FS induces strong chemotactic and mitogenic responses in cultured chondrocytes. Furthermore, we provide evidence to suggest that these effects are mediated, at least in part, via thrombin-induced activation of PAR-1 signalling.

Materials and methods

Materials. Tisseel FS was purchased from Baxter AG (Vienna, Austria). Tissue culture reagents and molecular biology reagents were purchased from Life Technologies (Melbourne, Australia) and Stratagene (La Jolla, CA, USA) respectively. The Biotrak® cell proliferation ELISA system was purchased from Amersham Life Sciences (Buckinghamshire, UK). Synthetic agonist PAR peptides with amidated C termini (PAR-1, SFLRN-NH₂; PAR-2, TFLLRN-NH₂; PAR-3, TFTRGAP-NH₂; PAR-4, GYPGQV-NH₂; purity >85%) were synthesized by the Protein Facility, University of Western Australia, Perth, Australia. All other chemicals were purchased from Sigma-Aldrich (St. Louis, MO, USA) unless stated otherwise.

Cell culture. Cartilage biopsies obtained from healthy human patients were used as a source of chondrocytes. All patients consented and ethics approval was obtained through the University of Western Australia Human Research Ethics Committee. In brief, biopsies were mechanically disaggregated and then digested with 0.3% (w/v) collagenase type II (Worthington Biochemical Corp., NJ, USA) at 37°C for 5-8 h with shaking to release chondrocytes. Following digestion, resulting chondrocyte suspensions were passed through a 100 μm cell strainer (Becton Dickinson and Co., NJ, USA) before being cultured in a T75-culture flask containing DMEM F-12 media (Gibco, NY, USA) supplemented with 10% (v/v) fetal calf serum (FCS) (Gibco), 292 μg/ml l-glutamine, 10,000 U/ml penicillin G, 10,000 U/ml streptomycin sulfate, 25 μg/ml amphotericin B, and 50 μg/ml ascorbic acid. Cells were sub-cultured routinely as required per experimental condition.

In vitro MACI co-culture assay. To simulate the in vivo conditions following matrix-induced autologous chondrocyte implantation (MACI) (12), primary human chondrocytes (5.0x10⁶ cells/ml) were seeded onto a 1.5 cm² type I/III collagen membrane (Matrical®, Germany) in 6-well plates and left to attach for at least 24 h. Following attachment, Tisseel FS (containing 500 U/ml thrombin) was applied to the cell-seeded surface of the membrane scaffold using the supplied duojector. The resulting membrane-cell-FS ‘sandwiches’ were then cultured in complete growth media for 12, 24, 48 h, and 15 days to promote cell migration. Seeded collagen I/III membranes in the absence of FS served as controls. Migration patterns of chondrocytes towards the FS were scored either histologically or by phase-contrast light microscopy (Nikon Diaphot).

Histology. MACI-FS ‘sandwiches’ were carefully removed from culture medium and washed twice in 1X PBS before being fixed in ice-cold paraformaldehyde (4%; 15 min) at room temperature. Following fixation, the ‘sandwiches’ were washed 3x with 1X PBS before undergoing routine paraffin processing and embedding. All samples were embedded in vertical orientation to the cutting plane so that both surfaces of the ‘sandwiches’ were displayed during tissue sectioning. Sections were cut (4-6 μm), placed onto glass slides and then de-waxed (xylene: 2x 3-4 min; 100% ethanol: 2x 3-4 min; 95% and 70% ethanol: 1x 3 min each). All sections were rehydrated in 3x with 1X PBS before undergoing routine paraffin processing and left to attach for at least 24 h. Following attachment, Tisseel FS ‘sandwiches’ were carefully removed from culture medium and washed twice in 1X PBS before being fixed in ice-cold paraformaldehyde (4%; 15 min) at room temperature. Following fixation, the ‘sandwiches’ were washed 3x with 1X PBS before undergoing routine paraffin processing and embedding. All samples were embedded in vertical orientation to the cutting plane so that both surfaces of the ‘sandwiches’ were displayed during tissue sectioning. Sections were cut (4-6 μm), placed onto glass slides and then de-waxed (xylene: 2x 3-4 min; 100% ethanol: 2x 3-4 min; 95% and 70% ethanol: 1x 3 min each). All sections were stained with Gill’s haematoxylin and eosin, mounted with Depex, and examined by light microscopy.

Proliferation assay. Cell proliferation was assayed by 5-bromo-2'-deoxyuridine (BrdU) using a commercial available Biotrak cell proliferation ELISA system. Briefly, human chondrocytes (5x10⁴ cells/well) were cultured in 96-well plates in complete growth medium overnight. Following attachment, cells were washed twice with 1X phosphate-buffered saline (PBS) and deprived of serum for an additional 24 h before the addition of thrombin (0.1, 0.5, 1, 10 U/ml) or vehicle (CaCl₂, 40 mM) in combination with serum-free DMEM-F-12 for an additional 24-48 h. BrdU-labeling solution was then added to each well, and cells were re-incubated for an additional 16 h. Time points (24 and 48 h) were staggered so that addition of BrdU occurred on the same day. Cells were fixed, and incorporated BrdU was detected using immunoperoxidase and tetramethylbenzidine (TMB) according to the manufacturer’s protocol. Absorbance was read at 450 nm.

Reverse transcription (RT)-PCR. Total RNA was extracted from the primary human chondrocytes using RNAzol B according to the manufacturer’s instructions (Tel-test, TX). cDNA was synthesised from 2 μg of total RNA using the RETROscript™ First-strand synthesis kit (Ambion). Primers against human PAR isoforms were designed based on published sequence data (34) and purchased from Genset
Confocal sequences were collected as Bio-Rad PIC files and equipped with a krypton-argon laser or argon ion laser coupled scanning microscopy (CLSM) (MRC-1000, Bio-Rad), Detection of fluorochromes was carried out by confocal laser Probes Inc.). Cell nuclei were visualised by counter-staining immunoglobulin G conjugated to Alexa Fluor 488 (Molecular bodies (dilution of 1:1000): goat anti-rabbit or goat anti-mouse previously outlined in Pavlos controls respectively. Zymed Laboratories Inc.) served as negative and positive for the chondrocytic marker S-100 (rabbit polyclonal 18-0046, visualised using streptavidin peroxidase and diaminobenzidine following the addition of secondary antibodies (either bio-

**Immunocytochemistry.** Immunodetection of PARs was conducted on human chondrocytes cultured in 8-well chamber slides (LAB-Tek II; Nunc) according to methods previously described by Asokananthan et al (35). Briefly, upon reaching approximately 80% confluence, cells were washed twice with PBS before being fixed with 4% (v/v) paraformaldehyde. Endogenous peroxidase activity was quenched by incubating chamber slides in 3% (v/v) H2O2 for 5 min, and non-specific binding was blocked by incubation in 10% FCS in PBS for 1 h. Cells were then incubated with primary antibodies raised against specific PAR isoforms: mouse monoclonal anti-human PAR-1 (ATAP2: sc-13503, Santa Cruz), mouse monoclonal anti-human PAR-2 (SAM11: sc-13504, Santa Cruz), rabbit polyclonal anti-PAR-3 (raised against peptide 37TLPIKT FRGAPPNSFEFP35; and rabbit polyclonal anti-PAR-4 (raised against peptide 3EDDSTPSLAPPYGPQVV35) (35). Following the addition of secondary antibodies (either biotinylated anti-mouse or anti-rabbit IgG), PAR expression was visualised using streptavidin peroxidase and diaminobenzidine (DAB). Incubation with either pre-immune serum or staining for the chondrocytic marker S-100 (rabbit polyclonal 18-0046, Zymed Laboratories Inc.) served as negative and positive controls respectively.

Immunolocalisation studies were performed essentially as previously outlined in Pavlos et al (36) using secondary antibody
dilutions of 1:1000: goat anti-rabbit or goat anti-mouse immunoglobulin G conjugated to Alexa Fluor 488 (Molecular Probes Inc.). Cell nuclei were visualised by counter-staining with Hoechst 33342 (1:10,000) (Molecular Probes Inc.). Detection of fluorochromes was carried out by confocal laser scanning microscopy (CLSM) (MRC-1000, Bio-Rad), equipped with a krypton-argon laser or argon ion laser coupled to an epifluorescence Nikon Diaphot 300 inverted microscope. Confocal sequences were collected as Bio-Rad PIC files and processed using Confocal Assistant 4.02. All images were collected under non-saturating conditions set up by the use of an output look-up table (LUT).

**Intracellular Ca**\textsuperscript{2+} mobilization. Intracellular calcium mobilization was measured fluorimetrically using Fura-2/AM (Molecular Probes, Eugene, OR, USA). Cells were trypsinised and seeded onto 10-mm coverslips in 35-mm culture dishes. Upon reaching confluence, cells were washed twice with freshly prepared physiological rodent saline (PRS, 138 mM NaCl, 2.7 mM KCl, 1.8 mM CaCl\textsubscript{2}, 1.06 mM MgCl\textsubscript{2}, 12.4 mM HEPES, 5.6 mM glucose, and 1 mM probenecid; pH 7.3) and loaded for 45 min at (37°C) with Fura-2/AM (3 μM) and 0.0125% F-127 (w/v) in PRS (1x10\textsuperscript{4} cells/ml). Following loading, cells were washed and incubated in the dark (30 min, room temperature). Coverslips were then removed and carefully placed into a heated bio-chamber (37°C) attached to an inverted epifluorescence microscope (Nikon TE2000, Japan). Fluorescence emission (510 nm) at 340- and 380-nm excitation was measured using a spectrophotometer (Cairn, UK). Thrombin (1 U/ml) and PAR-1, 3 and 4 agonists (400 μM) were added into the chamber following 1 min background recording and corresponding fluorescence emission ratios were recorded for 5 min. The PAR-2 agonist was not assessed because of its known activation by trypsin and tryptase (31-33). All intracellular calcium concentrations were expressed as the ratio of emission following excitation of 340 and 380 nm respectively.

**Statistical analyses.** Unless stated otherwise, all data are expressed as mean ± SEM. Statistical significance between means was determined by ANOVA or the Student’s t-test using GraphPad Prism (GraphPad Software, San Diego, CA, USA). P-values <0.01 were considered significant.

**Results.** Fibrin sealant promotes chondrocyte migration in vivo. Our previous in vitro studies indicate that FS (Tisseel\textsuperscript{6}) stimulates migration of autologous chondrocytes to osteochondral defects suggesting that the sealant possesses chemotactic properties (15). In order to determine whether the observed chondrocytic migration was directly related to chemotactic activity of FS, or a subsidiary effect of the repair process in vivo, we sought to replicate MACI using an in vitro co-culture system. For this purpose, autologous human chondrocytes, grown on a type I/III collagen membrane (Matricel), were ‘sandwiched’ between FS and cultured in vitro for 12-48 h. Following incubation, the ‘cell sandwiches’ were fixed and the migratory activities assessed histologically. As shown in Fig. 1A, chondrocytic migration from the collagen scaffold toward the FS was observed as early as 12 h co-culture, although the majority of cells remained on the superficial surface of the collagen membrane, reminiscent to that of the control. At 24 and 48 h time points, substantial cell migration was evident, with no breaching of the FS surface observed. We also assessed chondrocytic migration in co-cultures incubated for up to 15 days. As shown in Fig. 1B, extensive migration of chondrocytes towards the FS-collagen interface is evident, with some cells clearly invading the fibrin matrix. Collectively,
these data corroborate the notion that FS promotes the migration of autologous chondrocytes.

**Thrombin stimulates proliferation of human chondrocytes.** Having established that FS stimulates chondrocytic migration we next asked the question whether the sealant also possessed mitogenic potential. Given that thrombin, the active constituent of FS, has been previously shown to induce proliferation of a variety of cell types including endothelial cells (37), neutrophils (38), and osteoblasts (20), we examined whether the thrombin component similarly influenced the proliferative capacity of articular chondrocytes. To this end, human chondrocytes were incubated with increasing concentrations of thrombin (0.1-10 U/ml) for 24 and 48 h time points and its effect on cell proliferation was assessed using an ELISA-based BrdU incorporation assay. As shown in Fig. 2, thrombin induced a marked increase in BrdU incorporation with maximal response observed at doses of 10 U/ml (24 h: p<0.005; 48 h: p<0.0005). Significant proliferative activity was also detected in cells cultured in the presence of 1 U/ml thrombin, however only for the 48 h time point (p<0.005). On the other hand, lower concentrations of thrombin (<1 U/ml) failed to elicit any significant proliferative response. Comparable results were obtained using Alamar blue cell proliferation assays (data not shown). In all, these data indicate that the thrombin component of FS supports chondrocyte proliferation *in vitro*.

**PAR-1 is the major PAR isoform expressed in human chondrocytes.** Numerous studies indicate that thrombin elicits its biological responses via its interaction with, and subsequent activation of, PARs (28). Therefore, as an initial step towards understanding the role of PARs in the proliferation of human chondrocytes, we used reverse transcription polymerase chain reaction (RT-PCR) to measure the mRNA expression of PAR-1, PAR-2, PAR-3, and PAR-4. As shown in Fig. 3, PAR-1 (708 bp) and PAR-3 (382 bp) are highly expressed in chondrocytes whereas PAR-2 (582 bp) and PAR-4 (392 bp) exhibit moderate to weak expression respectively. ß-actin (240 bp) served as an internal loading control.
elucidating the molecular mechanism(s) underlying thrombin-induced chondrocyte proliferation, we examined the mRNA expression of PARs in chondrocytes by RT-PCR using isoform-specific primers. As shown in Fig. 3, PAR-1 and -3 mRNAs were highly expressed in cultured human chondrocytes. By comparison, PAR-2 and PAR-4 exhibited moderate to weak expression respectively.

To confirm the PCR data, we next assessed PAR protein expression levels by immunocytochemistry using a number of well-characterised PAR isoform-specific antisera (35) (Fig. 4). Consistent with the observed mRNA expression patterns, both PAR-1 and PAR-3 were strongly expressed in human chondrocytes, with PAR-1 exhibiting a slightly higher level of expression. On the other hand, PAR-2 displayed moderate staining whereas PAR-4 expression was detectable albeit weakly. Immunostaining for S-100 served as a positive marker for chondrocytes. Sections were viewed at ×250 and ×400 magnification.

Figure 4. Immunocytochemical detection of PARs in cultured human chondrocytes. Strong expression of both PAR-1 and PAR-3 is evident in human chondrocytes. PAR-2 displayed moderate chondrocytic expression whilst negligible PAR-4 staining was detectable. Immunostaining for S-100 served as a positive marker for chondrocytes. Sections were viewed at ×250 and ×400 magnification.

Figure 5. Subcellular localisation of PARs in human chondrocytes. Chondrocytes were grown on 8-well chamber slides, fixed with 4% paraformaldehyde, and then immunostained for PAR isoforms (green). All slides were counter-stained with Hoechst 33342 to visualise nuclei (blue) and images were recorded by confocal microscopy. Marked staining of the plasma membrane and cytosol was detected for PAR-1, -3 and -4. PAR-2 also shows plasma membrane staining as well as a sub-population of perinuclear vesicles (inset). Bar = 10 μm.

Figure 6. Intracellular calcium responses in human chondrocytes elicited by thrombin and PAR agonist peptides. Human chondrocytes were loaded with Fura-2/AM and incubated either with thrombin (1 U/ml) or PAR-1, -3, and -4 agonist peptides at 400 μM. Thrombin (A) and PAR-1 agonist peptide SFLLRN-NH₂ (B) induced steep intracellular calcium elevations upon addition, but thrombin elevation did not return to baseline. All responses were measured over a 5-min period and results were expressed as a fluorescence ratio (340/380 nm). Traces are representative of at least 5 independent experiments.

not shown). We also examined the subcellular localization of the PAR isoforms by confocal microscopy (Fig. 5). Whereas PAR-1, -3 and -4 isoforms were largely localized to the plasma membranes, with diffuse/reticular-like staining throughout the cytosol, PAR-2 predominantly associated with the plasma membrane and a population of small juxta-nuclear vesicular structures that were reminiscent of endosomes/lysosomes. Together, these data demonstrate that PARs are both differentially expressed and localized in human chondrocytes with...
PAR-1 being the major isotype expressed (PAR-1>PAR-3>PAR-2>PAR-4).

**Thrombin and PAR-1 agonists induce \([\text{Ca}^{2+}]\), influx in human chondrocytes.** The primary upstream signalling pathway of the PARs includes intracellular calcium mobilization (39). Given that \([\text{Ca}^{2+}]\) mobilization have been well-documented to correlate with cell growth and proliferation (20) we hypothesised that the thrombin-induced proliferation might reflect changes in intracellular calcium signalling, possibly via interactions with PARs. To explore this notion, we monitored for changes in free cytosolic calcium concentration in response to thrombin and specific PAR agonist peptides using the \(\text{Ca}^{2+}\) indicator Fura-2. Thrombin at a concentration of 1 U/ml elicited a large \([\text{Ca}^{2+}]\), elevation within a few seconds (mean amplitude of fluorescence ratio transient: 0.65±0.09; mean time to peak: 22.0±1.4 sec, n=8) in approximately 75% of human chondrocytes examined, before gradually decaying (mean half decay time: 32.4±6.6 sec; n=8) to near-baseline level (Fig. 6A). Similarly, the PAR-1 agonist (SFLLRN-NH$_2$, 400 μM) induced a significant and abrupt increase in intracellular \([\text{Ca}^{2+}]\) in an equivalent proportion of human chondrocytes. However, the PAR-1-induced \(\text{Ca}^{2+}\) transients were significantly smaller in amplitude (mean amplitude of fluorescence ratio transient: 0.11±0.01; p=0.0002, n=6) than that elicited by thrombin, but exhibited a steeper \([\text{Ca}^{2+}]\) elevation phase (mean time to peak: 8.5±1.6 sec; p<0.0001, n=6). Nonetheless, the decay phase was comparable to the thrombin-induced \(\text{Ca}^{2+}\) transients (mean half decay time: 28.5±5.1 sec, p=0.58, n=6) (Fig. 6B). Similar responses were also observed using the TFLLRN-NH$_2$, PAR-1 agonist peptide (data not shown). In contrast to PAR-1 agonist peptides, the PAR-4 (n=6) agonist peptide failed to elicit any visible \(\text{Ca}^{2+}\) response in human chondrocytes (Fig. 6B). Predictably, the PAR-3 agonist (n=5) did not evoke calcium mobilization, consistent with its reported inability to activate PAR-3 and low-affinity to thrombin (40). Together, the analogies in \([\text{Ca}^{2+}]\) responses between thrombin and PAR-1 agonists hint that the thrombin-mediated effects on chondrocytes might act via PAR-1 signalling.

**Discussion**

The diverse application of FSs has been advocated by numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have conducted numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have numerous studies from skin grafts through to attachment of FSs has been advocated by numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41). We and others have numerous studies from skin grafts through to attachment of osteochondral fragments (2-4,41).

In addition to promoting chemotaxis, thrombin has been shown to stimulate the proliferation of a variety of cell types (20,26,27,48-50). Consistently, our dose- and time-course-dependent studies chondrocyte conclusively demonstrate that thrombin directly induces chondrocyte proliferation. Cellular proliferation was observed 24-h post-stimulation with a maximum response at 10 U/ml thrombin, suggesting a threshold response concentration close to this point. Moreover, we demonstrate that this increase in cellular proliferation correlated with the mobilization of intracellular calcium.

It is well-established that changes in cellular calcium elicit several physiological sequelae, one of which is to stimulate cell mitotic activity. Thrombin has previously been shown to induce the proliferation of various cell types via the modulation of intracellular calcium (28). In osteoblastic cells, this calcium influx is mediated largely through the interaction of thrombin and activation of PARs suggesting that a similar mechanism may exist in ontogenically-related chondrocytes (20,27,33,35). Indeed, our expression studies document, for the first time, the expression of PARs in human chondrocytes thus extending chondrocytes to the list of PAR-expressing cells. PAR-1 was identified as the predominant isotype followed by PAR-3, PAR-2 and PAR-4. This finding is in accordance with the expression patterns reported for other mesenchymal-lineage cells (20,26,32,33).

Thrombin is known to activate PARs 1, 3 and 4 in various cell types, whereas PAR-2 is activated by trypsin and tryptase (20,26,31,52). Jenkins et al (53) demonstrated in osteoblasts that intracellular calcium underwent a sustained rise after treatment with PAR-1-activating peptide, whereas thrombin caused a sharp peak followed by a rapid return to baseline. In contrast, our results show sustained \([\text{Ca}^{2+}]\), after thrombin treatment, and a sharp transient increase after PAR-1-activating peptide treatment. It has been shown that ligand cross-reactivity exists in PARs whereby PAR-1-activating peptides are able to activate both PAR-1 and -2 (54). This is contradictory to the data reported here. It is possible that the elevated and sustained \([\text{Ca}^{2+}]\), response observed after thrombin treatment (in comparison to the PAR-1 agonist responses) is due to a number of factors. This discrepancy may be explained by either the presence of concurrent
intermolecular PAR activation between the tethered ligand of PAR-1 and neighboring PARs (i.e., PAR-2) (55); combined thrombin activation of PARs 1, 3, and 4; downstream interplay between activated G-protein-coupled PAR pathways; or the possible presence of an unidentified chondrocyte thrombin-sensitive PAR. Based on previous studies on sustained elevation of \( [\text{Ca}^{2+}]_i \) after PAR-2 treatment (33), the former two are considered the more likely possibilities.

The apparent lack of calcium sensitivity observed upon addition of the PAR-4 agonist peptide also implies that PAR-4 may not be crucial to chondrocyte physiology. This notion is supported by the comparatively weak expression levels of PAR-4 detected in human chondrocytes. Moreover, these findings are in accordance with calcium mobilization studies conducted with PAR-4 agonists in gingival fibroblasts which also exhibit strong PAR-1 and -3 but weak PAR-2 and -4 expression (56). Further studies will be required to unravel the precise roles of specific PAR isoforms in chondrocytes.

In summary, we demonstrate that the commercial FS Tisseel promotes the migration and proliferation of primary human chondrocytes. Moreover, we demonstrate that the thrombin component alone is sufficient to stimulate chondrocyte proliferation and elicit intracellular calcium mobilization.

In addition, we document for the first time, the expression and localisation of PARs in chondrocytes and provide evidence to suggest that PAR-1 is the primary thrombin-acting PAR in chondrocytes and, thus, might account for the observed migratory and proliferative responses. This hypothesis will form the basis of more detailed studies in the future. Nonetheless, the data presented in this study endorse the use of FSs in autologous chondrocyte implantation for cartilage injury.

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