Research Article

Therapeutic Benefit for Late, but Not Early, Passage Mesenchymal Stem Cells on Pain Behaviour in an Animal Model of Osteoarthritis

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Background. Mesenchymal stem cells (MSCs) have a therapeutic potential for the treatment of osteoarthritic (OA) joint pathology and pain. The aims of this study were to determine the influence of a passage number on the effects of MSCs on pain behaviour and cartilage and bone features in a rodent model of OA.

Methods. Rats underwent either medial meniscal transection (MNX) or sham surgery under anaesthesia. Rats received intra-articular injection of either $1.5 \times 10^6$ late passage MSCs labelled with $10 \mu g/ml$ SiMAG, $1.5 \times 10^6$ late passage mesenchymal stem cells, the steroid Kenalog ($200 \mu g/20 \mu L$), $1.5 \times 10^6$ early passage MSCs, or serum-free media (SFM). Sham-operated rats received intra-articular injection of SFM. Pain behaviour was quantified until day 42 postmodel induction. Magnetic resonance imaging (MRI) was used to localise the labelled cells within the knee joint.

Results. Late passage MSCs and Kenalog attenuated established pain behaviour in MNX rats, but did not alter MNX-induced joint pathology at the end of the study period. Early passage MSCs exacerbated MNX-induced pain behaviour for up to one week postinjection and did not alter joint pathology.

Conclusion. Our data demonstrate for the first time the role of a passage number in influencing the therapeutic effects of MSCs in a model of OA pain.

1. Introduction

Osteoarthritis (OA) is the most common joint disease in adults, and current prevalence is 12% in the population > 60 years, which will escalate over the next 20 years [1, 2]. Although there is controversy in the field, it is acknowledged that a broad spectrum of proinflammatory pathways and catabolic factors contributes to the initiation of OA, which impacts upon both the joint cartilage, synovium, and bone [1]. Pain is one of the first symptoms of knee OA; it can progress to be continuous, reducing movement and quality of life [1].

The mechanisms underlying OA pain involve structural changes and alterations in peripheral transduction and central processing of painful sensory inputs. Current treatments for OA pain have limited efficacy [3], and total joint replacement (TJR) surgery is a common outcome [1]. TJR surgery often reverses central sensitization, indicating that nociceptive output from the joint is fundamental in driving central pain mechanisms [4]. However, surgery is not suitable for patients < 55 years [5], and it remains critical that numbers of people with OA pain reliant on joint replacement as a treatment are reduced.

An alternative approach is the development of more effective cell-based therapies that limit the joint pathology and reduce synovial inflammation, which is significantly associated with OA pain [6].

Mesenchymal stem cells (MSCs) have a potential as a therapy for OA [7–10]. MSCs readily differentiate into bone,
cartilage, and adipose cells and release soluble factors (such as growth factors and chemokines) which harbour a regenerative environment through a variety of mechanisms [9, 11]. Animal models mimicking pathological and pain components of OA are widely used [12–14]; intra-articular injection of rat bone marrow-derived MSCs reduced pain behaviour in the absence of an effect on joint pathology in the monosodium iodoacetate model of OA in the rat [15] and had significant chondroprotective and anti-inflammatory effects in a rat surgical model of OA, but pain was not assessed [16]. Similarly, local delivery of adult MSCs was associated with regeneration of meniscal tissue and reduced joint destruction in a caprine model of OA [17]. Increased passage number from 4 to 9 of human adipose-derived adult stem cells increases the chondrogenic potential of cells [18]; whether this translates into improved benefit in vivo has yet to be addressed.

Maximising the therapeutic potential of cell-based therapies for the treatment of OA pain requires further understanding of the conditions required to maximise the potential therapeutic effect of MSCs, and knowledge of their sites and mechanisms of action, which requires monitoring of implanted cells within the joint. The aim of the present study was to compare effects of early versus late passage MSCs on pain behaviour, structural changes to the knee joint, and circulating levels of tumor necrosis factor alpha (TNFα) and interleukin 10 (IL-10) in a surgical model of OA in the rat. Tracking and imaging of the MSCs within the joint were achieved using magnetic resonance imaging (MRI) of superparamagnetic iron oxide nanoparticles (SPION) internalised by MSCs in a subset of the groups within the study.

2. Materials and Methods

2.1. Cell Isolation, Expansion, and Characterisation. Early passage bone marrow murine MSCs (P3) were isolated from Balb/c mice, and late passage cells (P9) were isolated from C57Bl/6 mice as previously described [19]. Both sets of cells were fully characterised for membrane receptor expression of CD31, CD44, CD11b, CD45, CD105, and Ly-6A (Sca-1) on C57Bl/6 mice as previously described [19]. The surgical MNX model of OA in rats has been demonstrated to induce symptoms comparable to that seen in human OA for pain behaviour, weight-bearing asymmetry, and disease pathology, through synovitis, pathology of the subchondral bone, and chondropathology, as well as the development of osteophytes [13, 22–24]. Rats were anaesthetised with isoflurane (3% induction, 2–2.5% maintenance; 1 L/min O2), and local anaesthetic EMLA cream was applied to the left hind limb. A full thickness cut through the medial meniscus of the left knee was performed. Sham-operated rats had their meniscus exposed, but not transected. Recovery from anaesthesia was monitored, and weight gain and general behaviour were monitored throughout the postinjury period.

2.2. Magnetic Nanoparticle (MNPs) Labelling. MSCs were labelled with SiMag (1000 nm; particle size) (Chemical, Germany). These are commercially available MNPs consisting of a maghemite iron oxide core (Fe3O4) and an unmodified silica surface with terminal negatively charged silanol groups. Cell were labelled using a passive incubation method as described by Markides et al. [20]. In brief, MNPs were suspended in serum-free CIM (cell isolation media) and added directly to cells in culture. Following a 24-hour incubation period, cells were washed three times with phosphate buffered saline (PBS) to remove noninternalised MNPs.

2.3. CellTracker™ CM-DiI Fluorescent Dye Labelling. A 1 mg/ml stock solution of the red fluorescent CellTracker CM-DiI (Molecular Probes, UK) was prepared in dimethyl sulfoxide (DMSO). MSCs were trypsinized, washed with PBS, and incubated with CM-DiI (2.5 μl of stock per 1 ml of PBS) for 5 minutes at 37°C, and then for an additional 15 minutes at 4°C, in darkness.

Unincorporated dye was then removed by centrifugation at 300g for 5 minutes and 2 washes in PBS. Cells were resuspended in serum-free IMDM (Iscove’s Modified Dulbecco’s Medium) and maintained at 4°C until injection.

2.4. MSC-Conditioned Medium Studies. Cells used for conditioned medium studies were previously isolated from C57Bl/6 and Balb/c mice and expanded in vitro. For testing, conditioned medium was prepared using murine bone marrow-derived MSCs at P3 and P10. Conditioned medium was prepared using serum-free IMDM (SF-IMDM) (GIBCO, Life Technologies) with no supplements added as described previously [21]. Briefly, cells were expanded to confluence using cell expansion medium (CEM, comprising IMDM with 9% FBS, 9% Horse Serum, and 1% Pen/Strep). Flasks were then rinsed three times with DPBS and once with SF-IMDM before adding 12 ml SF-IMDM. Following this, flasks were incubated at 37°C, 5% CO2, with SF-IMDM without cells used for controls. After 48-hour incubation, the medium was removed and centrifuged for 5 minutes at 1200g to remove cell debris. 11 ml supernatant was then passed through 3kDa centrifugal filters (amicon ultra 15 centrifugal filter tubes, Merck Millipore, Hertfordshire, UK) at 4000 rpm for 30 minutes at 4°C, and residual supernatant was removed from filters and immediately frozen to −80°C until testing.

2.5. Rat Model of Osteoarthritis Pain. Male Sprague Dawley rats (weighing 160–190 g) were purchased from Charles River UK. Studies were carried out in accordance with UK Home Office Animals (Scientific Procedures) Act (1986) and the guidelines of the International Association for the Study of Pain. Further, all works were conducted under Home Office project licence number 40-3647. Studies were undertaken in a blinded fashion. Rats underwent meniscal transection (MNX), or sham surgery, as previously described [22]. The surgical MNX model of OA in rats has been demonstrated to induce symptoms comparable to that seen in human OA for pain behaviour, weight-bearing asymmetry, and disease pathology, through synovitis, pathology of the subchondral bone, and chondropathology, as well as the development of osteophytes [13, 22–24]. Rats were anaesthetised with isoflurane (3% induction, 2–2.5% maintenance; 1 L/min O2), and local anaesthetic EMLA cream was applied to the left hind limb. A full thickness cut through the medial meniscus of the left knee was performed. Sham-operated rats had their meniscus exposed, but not transected. Recovery from anaesthesia was monitored, and weight gain and general behaviour were monitored throughout the postinjury period.

2.6. Intervention Studies and Pain Behaviour. Baseline measurements were taken prior to surgery (day 0) and from day 3 onwards. Behavioural assessment of changes in weight distribution and sensitivity to mechanical stimuli applied to the hindpaw were performed for up to 42 days postsurgery.
(see Supplementary Information). Two separate intervention studies were undertaken at 14 days postsurgery (MNX and sham). Prior to treatment, rats were stratified according to weight bearing and paw withdrawal thresholds (PWTs) (days 3–14) to ensure balanced groups. Under brief isoflurane anaesthesia (3% L/min O₂), rats received one intra-articular injection.

2.6.1. Study 1: Late Passage MSCs (P.10). MNX rats received intra-articular injection of either 1.5 × 10⁶ mesenchymal stem cells labelled with 10 μg/ml SiMAG (MSC-MNP; n = 11 rats), 1.5 × 10⁶ mesenchymal stem cells (MSC-VEH; n = 12 rats), or serum-free media (SFM; n = 12 rats). Sham-operated rats received intra-articular injection of serum-free media (SFM; n = 8 rats).

2.6.2. Study 2: Early Passage MSCs (P.3) versus Steroid Treatment. MNX rats received intra-articular injection of 200 μg/20 μL Kenalog (n = 8), 1.5 × 10⁶ MSC (n = 10 rats), or serum-free media (n = 10 rats). Sham-operated rats received serum-free media (SFM; n = 8 rats).

Following intra-articular injection, rats recovered from anaesthesia and were returned to the home cage. Weight bearing and PWTs were assessed on days 21, 28, 31, 35, and 38 post sham/MNX surgery. Experiments were terminated on day 42.

2.7. Magnetic Resonance Imaging (MRI). The in vivo MRI visibility threshold was determined previously by intra-articular injection of either 1 × 10⁶ or 2 × 10⁶ MSCs labelled with 0, 1, 5, and 10 μg/ml SiMAG into the joint of nonarthritic cadaveric 18-week-old Wistar rats. Rats were MR imaged using a Brucker 2.3 T animal scanner with the following sequence parameters; T2-weighted GEFI sequences, TR = 700 ms, TE = 5.5 ms, Flip angle = 30° and FoV = 7.9 × 7.9 cm, and matrix size = 256 × 192 to determine the location of the MNPs. MR images and signal loss profiles were compared to the untreated control groups and also between the treatment groups. Study 1 rats were sacrificed on day 42 and immediately MR imaged using the same system described above. Signal loss profiles were obtained and compared across all the groups. For details on data analysis, see Supplementary Information.

2.8. Histology. At sacrifice, tibiobular joints were removed and postfixed in neutral buffered formalin (4% formaldehyde) decalcified in ethylenediaminetetraacetic acid (EDTA) [24]. Histomorphometry was performed by an observer blinded to treatment. Coronal tissue sections (Osteoarthritis Research Society International (OARSI) guideline for histological assessment for OA in the rat) were cut at 5 μm [25].

Haematoxylin and eosin (H&E) stained sections were scored for joint morphology [26]. To validate the MRI results from study 1, mid-sagittal serial sections (4 μm) were obtained and stained with H&E and the fluorescent dye DAPI (1:200 dilution prepared in PBS) in order to visualise implanted CM-DiI labelled MSCs (see Supplementary Information).

2.9. ELISA. At sacrifice, blood was taken via cardiac puncture, aliquots spun for 20 minutes at 1000g, and serum supernatant was collected. Serum samples were diluted 2-fold, TNFα using 75 μL with calibrator diluent (R&D Systems, RD5-17), IL-10 using 50 μL with assay diluent (R&D Systems, RD1-21), and βNGF using assay diluent made up using 10% heat inactivated FBS (Life Technologies Ltd., 10500-064). Serum levels of TNFα and IL-10 were determined using commercially available Enzyme-linked immunosorbant assay (ELISA) kits (R&D Systems, Minneapolis MN) as per manufacturer’s instructions. Each serum sample was repeat tested n = 2, and absorption read at 450 nm with correction at 540 nm applied. Proprietary kits for measurement of βNGF in rat serum were not available, so components were sourced individually with the basic application using Duoset ELISA Development Kit (R&D Systems Europe, Ltd., DY556) and recommended components as per kit instructions (all R&D Systems Europe Ltd.) except for DPBS (GIBCO, Life Technologies, 14190-169).

2.10. Statistics. Data were analysed using GraphPad Prism 5.0. All data were tested for normality and for nonparametric testing: Kruskall-Wallis one way ANOVA with Dunn post hoc testing or 2-way ANOVA with Tukey’s post hoc testing was applied where appropriate, with probability values considered significant at *p < 0.05, **p < 0.01, and ***p < 0.001.

3. Results and Discussion Results

3.1. MSC Selection and Characterisation. Murine MSCs were freshly isolated and expanded in vitro to passage 3 (early passage) or passage 10 (late passage) with passaging taking place at 80–90% confluence. Stem cell properties were confirmed with flow cytometry for a panel of recognised MSC markers (CD105+, CD44+, CD73+, CD90+, CD11b+, CD45−, and CD34−) (Supplementary Information Figure 1) and selected for use. Cells were successfully differentiated towards adipogenic, osteogenic, and chondrogenic lineages after 21 days in culture with relevant differentiation media (Supplementary Information Figure 2). CFU-F assay was used to assess the proliferative capacity of the cells being expanded in culture, and the results showed that late passage cells retained a high proliferation rate in culture (early passage 49 ± 18% versus late passage 51.5 ± 12%). Early and late passage cells were cultured for conditioned medium collection, and levels of key cytokines (IL-10, TNFα, and βNGF) were measured (Supplementary Information Figure 3). Levels of IL-10 in the conditioned medium were minimal and did not vary between early and late passage cells. Levels of βNGF were significantly higher in conditioned medium from late passage MSC, compared to early passage MSC (Student’s t-test, p < 0.05, n = 6). TNFα was not detected in the conditioned medium under either condition (Supplementary Information Figure 3).

3.2. MRI of MSCs Labelled with SiMAG In Vivo. The presence of iron oxide MNPs was detected as a decrease in signal intensity when MR imaged using T2-weighted MRI sequences. This signal loss is portrayed visually as black areas on grey scale MR images, referred to as hypointense regions. The optimal cell number and SiMAG ratio to ensure good
MRI visibility over a prolonged period of time were determined to be 1.5 × 10^6 cells labelled with 10 μg/ml of SiMAG (Supplementary Information Figure 4). This labelling combination allows for improved visibility thresholds over 1 × 10^6 whilst minimising excessive blooming as seen with 2 × 10^6 cells and was taken forward for subsequent in vivo MRI tracking studies.

3.3. Effects of MSC Treatment versus Steroid on Pain Behaviour in a Model of OA. Both sham surgery and MNX surgery resulted in early changes in weight bearing on the operated hindlimb. By day 14 postsurgery, there was a clear difference between the extent of weight-bearing asymmetry between the sham and MNX groups, indicative of pain behaviour in the MNX group. At days 28–38, weight-bearing asymmetry remained significantly increased in the MNX group, but had returned to baseline in the sham group (Figure 1(a)). Area under the curve analysis of the last three timepoints tested (days 31–38) revealed a significantly greater weight-bearing asymmetry in the MNX group compared to the sham control group. Consistent with the studies in our group, hindpaw withdrawal thresholds (PWTs) were lowered in both sham and MNX rats following surgery (Figure 1(c)). There were no significant differences in the PWTs between the two groups of rats (Figure 1(d)).

The effect of intra-articular injection of 1.5 × 10^6 of late passage MSCs on established pain behaviour in the MNX model was determined. MSC treatment did not alter weight-bearing asymmetry in the week (days 14–21) immediately following treatment (data not shown). By contrast, there was a significant reduction in weight-bearing asymmetry at later timepoints (days 31, 35, and 38) in the MNX group treated with late passage MSCs (Figure 2(a)). PWTs were not altered by the intra-articular injection of late passage MSCs (Figure 2(b)). The effects of intra-articular injection of early passage MSCs versus a steroid treatment

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**Figure 1:** MNX surgery for induction of OA pain, or sham surgery, was performed on day 0. (a) MNX rats receiving SFM exhibited significant differences in weight-bearing asymmetry, compared to the sham-SFM rats. (b) Area under the curve (AUC) data calculated for days 31, 35, and 38 postsurgery revealed significant weight-bearing asymmetry in MNX rats at the later timepoints of the model. (c) Paw withdrawal thresholds were not significantly different in MNX rats, compared to sham controls at individual timepoints, or following AUC analysis. (d) Statistical comparison of the groups at each timepoint: two-way ANOVA with Bonferroni’s post hoc. Comparison of AUC used a Mann–Whitney’s nonparametric unpaired t-test. *p < 0.05, **p < 0.01 MNX versus sham. Data are mean ± SEM, n = 8–10 per group.
3.4. Joint Pathology, Inflammation, and Pain-Related Cytokines. Analysis of joint histology at the end of the study revealed that MNX surgery resulted in a significant chondrocyte score (Figure 4(a)), inflammation score of the synovium (Figure 4(b)), and an increase in the number of osteophytes (Figure 4(c)). At this final timepoint, none of the treatments significantly altered joint chondropathy or inflammation (Figures 4(a), 4(b), and 4(c)). Serum levels of three cytokines were measured in the different treatment groups at the end of the study (day 42 postsurgery). There were no differences in IL-10 expression between the treatment groups (Figure 4(d)). There was a significant increase in serum TNFα in the MNX-MSC early passage treatment group, compared to the sham-SFM controls and MNX-MSC late passage treatment group (Figure 4(e)). There were no significant differences in serum βNGF expression between the groups (Figure 4(f)).

3.5. MRI Tracking. A subset of rats received intra-articular injection of SimAG-labelled MSCs (MSC-MNP) for terminal MRI imaging at 29 days post cell implantation. MRI revealed regions of increased hypointensity (black areas) localised to the synovial cavity of rats treated with MSC-MNP (Figure 5). In contrast, no hypointense regions were evident in the groups treated with MSCs (MSC-VEH) or serum-free media (SFM). To further validate these data, signal loss profiles were plotted for the different treatment groups and revealed a significant signal loss in the MSC-MNP group, compared to unlabelled MSCs (MSC-VEH) and SFM which had a relatively high signal intensity across the joint (Figure 5). In a subset of rats, it was confirmed that there were no differences between the effects of SimAG-labelled MSCs and unlabelled MSCs on pain behaviour (data not shown). H&E staining was used to identify key structural features of the knee joint whilst identifying the location of the fluorescently labelled MSCs. CM-Dil-labelled MSCs were identified in the synovium of all MSC-treated groups (MSC-MNP and MSC-VEH) but not in the SFM-treated group (Figure 5).

4. Discussion

This study presents new evidence that passage number of MSCs markedly influences the effects of these cells on pain behaviour following their injection into the knee joint in a surgical model of OA pain. We report that late passage MSCs significantly reduced weight-bearing difference, a surrogate index of pain on loading, whereas early passage MSCs exacerbated weight-bearing difference for a period of 7 days postinjection in the MNX model. Despite the beneficial effects of the late passage MSCs on pain behaviour in the MNX model, there was no evidence for an alteration in the progression of joint pathology or inflammation at the end of the study. Nevertheless, a peripheral site of action of the MSCs was supported by the demonstration that SimAG-labelled MSCs were detected within the synovial cavity at 29 days postinjection.

As joint degeneration progresses, a variety of surgical procedures can rebuild the degenerated cartilage lesions, but do not necessarily reduce the generalised joint inflammatory processes. Chondrocytes as a cell-based therapy (autologous chondrocyte implantation (ACI)) were successfully developed and used widely over the past 10 years, but
towards an inhibition of weight-bearing asymmetry by Kenalog, this was only signi
timepoints (days 31, 35, and 38). mMSC treatment did not alter weight-bearing asymmetry in MNX rats. Although there was a trend
Data are expressed as a % of the mean MNX-SFM AUC for timepoints 31, 35, and 38 days postsurgery. Statistical analysis used a

![Figure 3](image-url)

Figure 3: (a) Timecourse of the effects of intra-articular injection of the steroid Kenalog (200 μg/20 μL) versus 1.5 × 10⁶ of early passage
mMSC on weight-bearing asymmetry. Rats received the active treatments or 50 μL SFM (vehicle) on day 14 postsurgery. Kenalog had a
rapid inhibitory effect on MNX-induced weight-bearing asymmetry at 17 and 21 days post model induction, compared to the MNX-SFM
group. mMSC treatment significantly increased weight-bearing asymmetry at early timepoints (days 17 and 21) post model induction. At
later timepoints, weight-bearing asymmetry was comparable between the MNX-MSC group and the MNX-SFM group. Data was analysed
using a 2-way ANOVA with Tukey’s post hoc test. *p < 0.01 MNX-SFM versus sham-SFM, ##p < 0.01 MNX-MSC versus MNX-SFM;
@p < 0.05, @@@p < 0.0001 MNX steroid versus MNX-MSC. (b) Area under the curve (AUC) analysis of the effects of intra-articular
injection of steroid Kenalog versus 1.5 × 10⁶ of early passage mMSC on MNX-induced weight-bearing asymmetry for the last three
timepoints (days 31, 35, and 38). mMSC treatment did not alter weight-bearing asymmetry in MNX rats. Although there was a trend
towards an inhibition of weight-bearing asymmetry by Kenalog, this was only significantly compared to the MNX-mMSC group.
Data are expressed as a % of the mean MNX-SFM AUC for timepoints 31, 35, and 38 days postsurgery. Statistical analysis used a
Kruskal-Wallis test with Dunn’s post hoc, *p < 0.05. Data are mean ± SEM, n = 9-10 per group.

this treatment relies on damaging healthy cartilage to pro-
vide the cell sources. The therapeutic potential of alternate
sources of cells, such as MSCs derived from the bone
marrow, which have anti-inflammatory and immunosup-
pressive properties, has been investigated. Our finding that
late passage MSCs attenuated established weight-bearing
asymmetry in the MNX model is consistent with the report
that intra-articular injection of MSCs reversed pain behav-
ior compared to pretreatment values, but did not alter
structural damage or synovial inflammation in the chemical
monosodium iodoacetate model of OA pain [27]. The
clinical validity of animal models of OA continues to be
debated, both in terms of the aetiology of the joint damage
and the temporal progression of the structural changes seen
in these models, compared to disease progression in
patients. The MNX model of OA is believed to replicate
some of the key biomechanical events that lead to clinical
joint pathology, as well as displaying many of the features
associated with joint pathology in OA (see refs in [28]).
Our evidence that intra-articular injection of early passage
MSCs exacerbated pain behaviour in the model of OA pro-
vides important new knowledge of the conditions under
which the therapeutic potential of MSCs for OA pain can
be harnessed.

Our data were built upon the previous studies that
focused on the potential for MSCs to mediate joint repair.
Indeed, therapeutic benefit of intra-articular injection of
autologous MSCs has been reported in a surgical model of
OA in the goat [17], and more recent studies report beneficial
effects of MSCs from bone marrow and adipose in models of
OA and extend the initial evidence by demonstrating repara-
tive effects of the cells on the cartilage [29, 30]. The pro-
gression of MNX-induced joint pathology was not halted by
intra-articular injection of late passage MSCs, suggesting that
at least in this model the effects of MSC treatment on pain
behaviour are not associated with increased joint repair.
Despite this lack of effect on joint pathology, our study did
provide evidence for changes in systemic inflammation.
There was a trend towards an increase in serum TNFα in
the MNX model of OA pain, compared to the sham control
group at day 42 postsurgery. Interestingly, MNX rats treated
with the early passage MSCs had significantly increased
serum TNFα, compared to the sham group, consistent with
the exacerbation of pain behaviour by this treatment. By con-
trast, serum TNFα was significantly lower in MNX rats
receiving MSCs compared to the early passage MSCs.
These data are consistent with the ability of late pas-
sage, but not early passage, MSCs to reduce pain behaviour.
Pharmacological studies using comparable methods demon-
strated a significant increase in plasma TNFα in the MIA
model of OA pain compared to control rats, with these
changes reversed by an analgesic treatment [31]. Although
both IL-10 and NGF were detected in the serum, there were
no significant differences between the treatment groups, sug-
gest that changes in TNFα do not reflect a generalised
change in inflammation.
Figure 4: (a–c) MNX surgery was associated with significant chondropathy (a), joint inflammation (b), and increased presence of osteophytes (c), compared to sham controls at 42 days postsurgery. None of the treatments in MNX rats significantly altered the extent of chondropathy, inflammation, or osteophyte number. Statistical analysis used Kruskal-Wallis test with Dunn’s post hoc, **p < 0.01, ***p < 0.001. Data are mean ± SEM, 2–4 sections per rats were analysed, and total numbers of sections are sham-SFM: 37; MNX-SFM: 55; MNX-MSC early passage: 44; MNX Kenalog: 42; MNX-MSC late passage: 20. (d–f) Serum levels of cytokines in MNX- and sham-operated rats at 42 days postsurgery. There were no differences in IL-10 expression between the treatment groups (d). There was a significant increase in serum TNFα in the MNX-MSC early passage treatment group, compared to the sham-SFM controls and MNX-MSC late passage treatment group (e). There were no significant differences in serum βNGF expression between the groups. Statistical analysis used Kruskal-Wallis test with Dunn’s post hoc, *p < 0.05, **p < 0.001. Data are overall mean ± SEM, whilst mean values from duplicate samples per rat were analysed; total numbers of mean values are sham-SFM: 15; MNX-SFM: 22; MNX-MSC early passage: 10; MNX Kenalog: 9; and MNX-MSC late passage: 23.
Understanding the differences in properties of the early versus late passage MSCs that lead to the behavioural outcomes will help refine MSC treatment strategies. Prolonged in vitro culture of bone marrow-derived MSCs leads to a loss of MSC phenotype, multipotency, decreased wound homing properties [32], and self-renewal by around passage 15–20, associated with the onset of cellular senescence [33–36]. As a result, the use of early passage cells is recommended; however, early passage cell populations have increased likelihood of heterogeneity whilst late passage cells retain characteristic markers for MSC phenotype in a selectively more homogeneous population [36, 37]. The immunomodulatory properties of MSCs in a long-term culture have been reported. Late passage MSCs of umbilical origin (passage 15) show significant upregulation of anti-inflammatory mediator HMOX-1, a modulator of IL-10 and NO activity [38, 39]; downregulation of proinflammatory IL-1α, IL-1β, and IFN-γ; reduced proliferation of PHA-stimulated peripheral blood mononuclear cells, implicated in expression of proinflammatory cytokines; and no change in expression of

Figure 5: (a): Sagittal MRI scans; Location of SiMAG-labelled cells are depicted as areas of hypointense signal loss and highlighted by the red ring over the synovial cavity. (b) Corresponding histological sections. Fluorescent images correspond to location marked X on H&E images (inset). Implanted DiI-labelled MSCs are shown in red whilst all native materials are show by DAPI in blue. (c) Corresponding MRI signal loss profile. Groups include (i) SiMAG-labelled MSCs (MSC-MNP), (ii) MSCs only (MSC-VEH), and (iii) serum-free media. T: tibia; F: femur; S: synovial lining; M: meniscus. Scale bars = 100 μm.
TGF-β, a cytokine critical to the immunosuppressive capabilities of transfused MSC [39]. It may therefore follow in our study that IFN-γ production is impaired and macrophage balance shifts from M1 to M2, as seen previously with in vivo MSC transplantation [40]. Production of IL-6 is increased in the late passage human bone marrow-derived MSC compared to early passage cells, whilst CXCL8 levels fall [41]. IL-6 has both anti- and proinflammatory activities and may function here to activate IL-10 and bind to toll-like receptors to inhibit proinflammatory cytokine production, for example, TNF-α and IL-1, whilst also providing protection against bacterial proliferation [42, 43]. More analyses of the cytokine profile are needed to define exact pathways in the variation in potential immunomodulation between early and late passage MSCs in our model of OA pain.

There is evidence of other phenotypic shifts in MSCs with prolonged culture which may provide insight. For example, prolonged culture of MSCs reduces STRO-1 and BMP7 expression, changes alkaline phosphatase activity, and reduces osteogenic capability [36, 44, 45]. Earlier studies, examining passage-dependent differences to chondrogenic potential of MSCs, have produced varied results, with some studies reporting a maintenance of the chondrogenic potential of the cells up to passage 20 [46], or increase in COL2A1 and AGC1 expression from P4 to P9 in human adipose-derived adult stem [18]. Another study reported a reduction of chondrogenic capabilities of MSCs at late passage [33, 35, 47]. With an increasing passage number, MSC isolated from synovium shows migratory behaviour similar to chondrocytes, whilst at low passage (p < 4), a reduced ability to undergo chondrogenesis was observed [48]. Our results revealed that late passage MSCs attenuated established pain behaviour (weight-bearing asymmetry) in MNX rats, but did not alter MNX-induced joint pathology. This suggests that higher passage cells may have an increased potential for therapeutic application. What is clear is that selection of populations of cells for therapy will depend on the level of precurture with significant changes in phenotype and potential therapeutic activity as a result of prolonged culture. The trade-off between efficacy and cell number generated by scale-up of cell numbers will be an important consideration in a therapeutic design. Future studies of earlier experimental timepoints will shed light on the effects of treatments on the alleviation of pain and moderation of inflammatory response following MSC treatment.

The key to enabling the translation of MSC treatment for OA to the clinic is the tracking of the cells following injection into the knee joint to initially demonstrate effective delivery of cells and to monitor cell retention and biodistribution thereafter. Previously, we have shown that SiMAG, a commercially available superparamagnetic iron oxide nanoparticles (SPIONs), can be used to track the migration of MSCs over 7 days and localise them to the knee joint in a mouse model of rheumatoid arthritis (RA) [20]. Importantly, the same study demonstrated no adverse effects in terms of in vitro MSC properties, cell viability, and proliferation as a result of SiMAG labelling. Furthermore, the delivery of SiMAG-labelled MSCs was also well tolerated by mice thereby encouraging the use of SiMAG as an imaging and tracking agent in this current study. Although a number of noninvasive imaging modalities can be applied to track cells post implantation, SPION- and MRI-based tracking techniques benefit from relatively long-term and longitudinal monitoring of implanted cell populations. In a recent study by van Buul et al. [27], implanted MSC populations were monitored by bioluminescence and MRI with MSCs transfected with firefly luciferase and labelled with a particular SPION known as Endorem. A gradual and complete loss in bioluminescence signal was observed over a 3-week period. By contrast, the SPION-based component of this system has the potential to allow SPION-labelled cells to be monitored for up to 12 weeks as demonstrated by 2 independent studies by Jing et al. [49] and Chen et al. [50]. In these studies, Endorem-labelled MSCs and chondrocytes were monitored in vivo for up to 12 weeks within the knee joint [49, 50]. Nevertheless, unlike bioluminescence, the viability of implanted cells cannot be determined by MRI-based modalities. In our study, SiMAG-labelled cell was successfully observed up to 29 days post implantation.

5. Conclusions

We have demonstrated differences in pain responses in a rat surgical model of OA following cell therapy using populations of MSCs at early and late passages following isolation from the bone marrow. Late passage MSCs significantly reduced weight-bearing difference, a surrogate index of pain on loading (days 31, 35, and 38), whereas early passage MSCs exacerbated weight-bearing difference for a period of 7 days postinjection in the MNX model. Neither treatment altered progression of joint pathology nor inflammation quantified at the end of the study. Our data provide further evidence for the need for characterisation of this cell type prior to clinical use due to its multifunctional nature and the changing phenotype from repair to immunomodulation with time in culture.

Ethical Approval

All studies were approved by local ethics committee. Experiments were carried out in accordance with UK Home Office Animals (Scientific Procedures) Act (1986) and the guidelines of the International Association for the Study of Pain.

Conflicts of Interest

The authors declare that there is no conflict of interest regarding the publication of this article.

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Supplementary Materials

Supplementary Figure 1: Cell marker profile for early and late passage MSC, characterised through CD105⁺, Sca-1⁻, CD31⁻, CD11b⁻, CD45⁻, CD44⁺, CD34⁻. CD105 is known to vary in expression in murine MSC. In this study, cells at late passage expressed lower levels of CD105 whilst all other markers were consistently expressed. Supplementary Figure 2: Tri-lineage differentiation for early (A) and late (B) passage cells along adipogenic (1), osteogenic (2) and chondrogenic (3) lineages. Positive differentiation was detected for both early and late passage MSC. Scale bars = 200 μm. Supplementary Figure 3: Measurement of cytokines conditioned media from early and late passage cultured MSCs. There were no differences in IL10 levels between the two conditions. TNFα was not detected in conditioned medium. Expression of βNGF was significantly higher in late passage MSC compared to early passage MSC (Student’s t-test, p < 0.05) (n = 6 for all samples). Supplementary Figure 4: In vivo dose response: Signal loss profiles and corresponding sagittal MR images following the implantation of (A) 1 × 10⁶ and (B) 2 × 10⁶ MSCs labelled with (i) 10 μg/ml, (ii) 5 μg/ml, and (iii) 1 μg/ml SiMAG into knee joints of cadaveric rats and compared to the (iv) untreated control. Location of SiMAG labelled cells is depicted as areas of hypointense signal loss and highlighted by the red ring over the synovial cavity. (Supplementary Materials)

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