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MMP13 and TIMP1 are functional markers for two different potential modes of action by mesenchymal stem/stromal cells when treating osteoarthritis

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Abstract
Mesenchymal stem cells (MSCs) have been investigated as a potential injectable therapy for the treatment of knee osteoarthritis, with some evidence of success in preliminary human trials. However, optimization and scale-up of this therapeutic approach depends on the identification of functional markers that are linked to their mechanism of action. One possible mechanism is through their chondrogenic differentiation and direct role in neo-cartilage synthesis. Alternatively, they could remain undifferentiated and act through the release of trophic factors that stimulate endogenous repair processes within the joint. Here, we show that extensive in vitro aging of bone marrow-derived human MSCs leads to loss of chondrogenesis but no reduction in trophic repair, thereby separating out the two modes of action. By integrating transcriptomic and proteomic data using Ingenuity Pathway Analysis, we found that reduced chondrogenesis with passage is linked to downregulation of the FOXM1 signaling pathway while maintenance of trophic repair is linked to CXCL12. In an attempt at developing functional markers of MSC potency, we identified loss of mRNA expression for MMP13 as correlating with loss of chondrogenic potential of MSCs and continued secretion of high levels of TIMP1 protein as correlating with the maintenance of trophic repair capacity. Since an allogeneic injectable osteoarthritis therapy would require extensive cell expansion in vitro, we conclude that early passage MMP13+, TIMP1-secreting high MSCs should be used for autologous OA therapies designed to act through engraftment and chondrogenesis, while later passage MMP13−, TIMP1-secreting high MSCs could be exploited for allogeneic OA therapies designed to act through trophic repair.

KEYWORDS
chondrogenesis, mesenchymal stem cells, MMP13, TIMP1, trophic repair
1 | INTRODUCTION

The concept of multipotent mesenchymal stem cells (MSCs) was established by Caplan in the early 1990s\(^1\) following the seminal work of Friedenstein in the 1960s and 1970s, demonstrating the presence of osteogenic precursor cells in bone marrow.\(^2\)-\(^4\) Building on this early work, there have been many studies demonstrating the capacity of MSCs to differentiate in vitro with clear evidence for multipotent skeletal lineage differentiation,\(^5\) including chondrogenesis,\(^6\)-\(^13\) osteogenesis,\(^14\)-\(^17\) and adipogenesis\(^9\),\(^18\),\(^19\) as well as more limited evidence for pluripotent differentiation including endodermal and ectodermal pathways.\(^1\),\(^20\) In one of these studies, we generated clonal populations of bone marrow MSCs and showed that individual cells retained the capacity for chondrogenesis, with varying degrees of potency.\(^9\)

More recently, evidence has grown that MSCs may support tissue repair through mechanisms that do not directly relate to their multipotential differentiation capacity.\(^21\) Caplan has described MSCs as having “trophic” capacity by which, following implantation, they induce neighboring cells to secrete active molecules, for example, in the treatment of stroke, myocardial infarction, or in meniscal cartilage repair.\(^22\) Trophic repair is most likely mediated through the production by MSCs of large amounts of growth factors and other mediators.\(^20\),\(^22\)-\(^25\) We have previously developed a therapeutic strategy for meniscal cartilage repair based on the trophic properties of MSCs\(^26\) that has shown some evidence of efficacy in preclinical and clinical trials.\(^27\) A second mechanism contributing to trophic repair is the ability of MSCs to suppress immune responses by a range of mechanisms including downregulation of T cell proliferation.\(^9\),\(^24\),\(^28\)-\(^31\) This important property of MSCs has been used clinically to support the engraftment of donated hematopoietic cells and to prevent graft vs host disease.\(^24\),\(^32\)

There have been several studies describing the loss of differentiation capacity with increasing passage of MSCs in vitro,\(^33\)-\(^36\) with other studies suggesting that in vivo aging also leads to a loss of differentiation capacity after ex vivo isolation of the aged cells.\(^37\)-\(^39\) These observations combined with parallel observations of shortening in telomere length with in vitro aging of MSCs\(^40\) have been interpreted as indicating a rapid senescence process that compromises cell function and limits their clinical utility. Conversely, no comparative aging-related data have been reported for trophic repair or immunoregulation.

The aim of this study was to identify markers of MSC chondrogenic potency (measured by cartilage tissue engineering) and trophic potency (measured by meniscal cartilage integration as well as by suppression of T-cell proliferation), using extensive in vitro aging of the cells as a method of regulating their specific functional activity.

2 | MATERIALS AND METHODS

2.1 | Isolation and expansion of human marrow-derived MSCs for in vitro studies

The model used as the basis for all the experiments reported here was long-term culture of human bone marrow-derived MSCs. Bone marrow plugs were collected from the femoral heads of patients undergoing total hip replacement. All patients gave their informed consent and the study was carried out according to local ethical guidelines (North Bristol NHS Trust Research Ethics Committee). Patient details can be seen in Table S1. Cells were suspended in stem cells expansion medium consisting of low glucose Dulbecco's Modified Eagles Medium (Sigma-Aldrich) supplemented with 10 vol%/vol% fetal bovine serum (FBS, Sigma-Aldrich), 1 vol%/vol% Glutamax (Gibco), and 1 vol%/vol% Penicillin/Streptomycin (Sigma-Aldrich). The serum batch was selected to promote the growth and differentiation of MSCs.\(^41\) The medium was also supplemented with 10 ng/mL FGF-2 (Peprotech). This growth factor has been previously shown to enhance the MSC proliferation rate in vitro,\(^12\),\(^42\) to retain MSCs as undifferentiated cells during proliferation\(^6\),\(^43\) and to enhance chondrogenic differentiation when the FGF-2 expanded MSCs are subsequently exposed to differentiation conditions.\(^12\),\(^42\) The cell suspension was separated from any bone in the sample by repeated washing with media. The cells were centrifuged at 500g for 5 minutes and the supernatant/fat removed. The resulting cell pellet was resuspended in medium, and then plated at a seeding density of between 1.5 \(\times 10^5\) and 2.0 \(\times 10^5\) nucleated cells per \(\text{cm}^2\). These flasks were incubated at 37°C in a humidified atmosphere of 5% CO\(_2\) and 95% air. Four days were allowed before the first medium change and then the medium was changed every other day until adherent cells reached 90% confluence and were ready for passaging.

2.2 | Cell passaging and calculation of population doublings and doubling time

At the end of each passage, the MSCs were harvested using 0.25% trypsin-EDTA (Sigma-Aldrich), pooled, counted, and then divided into different centrifuge tubes for reseeding and further growth, for immediate use in measurement of % integration of meniscal cartilage, for storage in liquid nitrogen for subsequent use in differentiation protocols as well as for genomic and proteomic analysis. The cells for each

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**Significance statement**

This study has shown that mesenchymal stem cell (MSC) chondrogenesis is a transient property of these cells which is lost as they age in vitro, whereas trophic repair potency is maintained until the MSCs cease to grow. These findings are significant as they highlight the importance of defining the intended mode of action when preparing MSCs for injection into osteoarthritic joints. These studies demonstrate that the development of injectable MSC therapies for osteoarthritis must take into account the transient nature of chondrogenic potency relative to their sustained trophic potency with increasing passage and specific strategies should be adopted to exploit one or other of these mechanisms of action.
patient were passaged continuously without freezing, until growth arrest, defined as no detectable increase in cell number between passages (see Table S1). At each passage, the total number of harvested MSCs was determined. The first cell harvest after seeding of fresh bone marrow was taken as passage 0. The number of cells reseeded at the start of passage 1 was used as the baseline for calculation of the first population doubling (PD) value at the end of passage 1. Downstream analyses of the MSCs were undertaken from passage 1 onward.

The number of PDs was calculated using the following formula:

$$PDs = \frac{\log(\text{Number of Harvested MSCs}) - \log(\text{Number of seeded MSCs})}{\log(2)}$$

The PD for each passage was calculated and added to the PD of the previous passages to generate data for Cumulative PD at each passage.

Population doubling time (PDT) was calculated for each passage using the formula:

$$PDT = \frac{t \times \log(2)}{\log(\text{cells harvested}/\text{cells seeded})}$$

where $t$ = the time between cell seeding and cell harvesting.

### 2.3 Detection of cell-surface phenotypic markers

MSCs (100 000 cells from each patient at each of passages 1, 5, 10, and 15) were suspended in a 1:500 dilution of Zombie (Biolegend), a live/dead cell dye, and incubated for 20 minutes in the dark. Nonspecific antigens were then blocked by incubating the cells at room temperature for 1 hour in 1 wt%/vol% bovine serum albumin (Sigma-Aldrich), 5 vol%/vol% FBS. The cells were washed by centrifugation in three volumes of Dulbecco’s phosphate-buffered saline (PBS; Sigma-Aldrich), and the cell pellet was suspended in 100 μL of a primary antibody solution containing 20 to 100 μg/mL of antibody in blocking solution. All the primary antibodies were fluorescent-labeled mouse anti-human IgGs: anti-CD105-fluorescein isothiocyanate (FITC), anti-CD90-phycoerythrin (PE), and anti-CD45-PE were from R&D Systems; anti-CD34-FITC was from eBioscience, Life Technologies; IgG1-FITC and IgG1-PE isotype controls were from R&D Systems; anti-CD105-fluorescein isothiocyanate (FITC), anti-CD90-phycoerythrin (PE), and anti-CD45-PE were from R&D Systems; anti-CD34-FITC was from eBioscience, Life Technologies; IgG1-FITC and IgG1-PE isotype controls were from R&D Systems.

### 2.4 Chondrogenesis

#### 2.4.1 Cartilage tissue engineering

The chondrogenic capacity of MSCs from each passage was assessed by performing three-dimensional cartilage tissue engineering, as previously described. Briefly, 300 000 cells were loaded drop-wise onto 5 mm diameter × 2 mm thick polyglycolic acid scaffold disks (Biomedical Structures) which had been pre-coated with 100 μg/mL fibronectin (Sigma-Aldrich). Constructs were then cultured in chondrogenic differentiation medium consisting of Dulbecco’s Modified Eagle Medium (DMEM) containing 4500 mg/L (high) glucose (Sigma-Aldrich), supplemented with 10 ng/mL transforming growth factor-β3 (R&D Systems), 100 nM dexamethasone, 80 μM ascorbic acid 2-phosphate, 1 mM sodium pyruvate, 1 vol%/vol% Penicillin/Streptomycin (all from Sigma-Aldrich), 1% insulin-transferrin-selenium-G (ITS), and 2 mM Glutamax (both from Gibco). After 7 days, the medium was further supplemented with 10 μg/mL human insulin (Sigma-Aldrich) until the end of culture. The constructs were incubated at 37°C for a total of 35 days on a rotating platform and medium was changed every 3 days.

#### 2.4.2 Biochemical analysis

Cartilage constructs were freeze-dried and weighed at the end of the 35-day tissue engineering period. The extracellular matrix was fully solubilized by overnight digestion with 2 mg/mL bovine pancreatic trypsin (Sigma-Aldrich) which was then boiled for 15 minutes to inhibit the action of the enzyme. In order to obtain the dry weight of extracellular matrix in the construct, remaining undigested scaffold material was freeze-dried, weighed, and subtracted from the original dry weight. The amounts of proteoglycan in the digests was measured as sulfated glycosaminoglycan (GAG) using a dimethylmethylene blue (Sigma-Aldrich) colorimetric assay.

### 2.5 Osteogenesis and adipogenesis

Whole MSC populations or MSC clones were grown in monolayer until 50% to 70% confluent prior to osteogenic differentiation or 100% confluent prior to adipogenic differentiation. In both cases, control cells were then cultured in minimum essential medium (α-MEM; Sigma-Aldrich) basal medium containing 10% FBS, 1 vol%/vol% Penicillin/Streptomycin, and 2 mM Glutamax. Cells stimulated to undergo differentiation were cultured in basal medium containing either osteogenic supplement or adipogenic supplement (both from R&D Systems) for 21 days. Following osteogenic differentiation cells were fixed in 70% ethanol and stained with 40 mM alizarin red S (Sigma-Aldrich), pH 4.1, for 5 minutes. Following adipogenic differentiation cells were fixed in 4% paraformaldehyde and stained with 0.3% oil red O (Sigma-Aldrich) for 30 minutes. The extent of differentiation was scored under blind conditions and classified as – (no staining), +, ++, or +++ according to increasing area and number of mineralized deposits following osteogenic differentiation (see Figure S1) and increasing number of lipid droplets following adipogenic differentiation (see Figure S3).
2.6 | Meniscal repair

2.6.1 | Preparation of sheep meniscal cartilage

Sheep legs were purchased from Edge & Son Butchers, Wirral, Merseyside and the meniscal cartilage removed under sterile conditions. Meniscal cartilage cylinders (5.0 mm in diameter and 3.0 mm thick) were harvested from the avascular (white zone) of the ovine menisci using a dermal biopsy punch. They were rinsed and incubated with PBS containing 10 vol%/vol% penicillin/streptomycin and 1 vol%/vol %, 250 μg/mL Amphotericin B (Sigma-Aldrich) for 20 minutes. Viability of the fibrocartilage disks was maintained by culture in basic medium consisting of low-glucose DMEM with 10 mM Hepses buffer (Sigma-Aldrich), 1 vol%/vol% penicillin/streptomycin, nonessential amino acids (Sigma-Aldrich), 1 vol%/vol% Glutamax, and 10% amphotericin B at 37°C in a 5% CO₂ environment. The explants were used in the integration experiments within 3 days of culture.

2.6.2 | Cell seeding

Collagen scaffolds (Ultrafoam Collagen Sponge; Bard) were cut into 6 mm diameter disks and seeded with human MSCs at a concentration of 1 × 10⁶ cells/cm². The suspension was loaded drop wise onto the scaffold placed in ultra-low attachment wells of a 24-well plate (Corning). After 4 hours, 1.5 mL of expansion medium containing 10 ng/mL FGF-2 was added and changed daily. Seeded scaffolds were incubated for 48 hours at 37°C in an orbital shaker at 50 rpm.

2.6.3 | Assembling and culture of constructs

Sandwich constructs of two ovine meniscal cartilage disks interposed with a seeded scaffold were assembled as previously described²⁷ using skin clips and cultured in vitro in ultra-low attachment 6-well plates in expansion medium with 10 ng/mL FGF-2 for 7 days followed by culture in an integration medium consisting of high glucose DMEM containing 10 vol %/vol% FBS, 1 vol%/vol% Glutamax, 1 vol%/vol% penicillin/streptomycin, insulin, and 80 μM ascorbic acid 2-phosphate for 33 days. The medium was replenished twice every week. The constructs were incubated at 37°C on a rotating platform throughout the culture period. At the end of culture, the constructs were prepared for histological analysis by fixation in 10 vol%/vol% neutral buffered formalin (Sigma-Aldrich).

2.6.4 | Histomorphometric analysis

Histomorphometry was carried out using a method that we developed and characterized in previous studies.²⁶,²⁷,⁴⁴ Fixed constructs were dehydrated and paraffin embedded. Samples were cut into 4 μm sections and stained with H&E for the study of morphological details. All histological sections were scanned using a Leica Aperio slide scanner and histomorphometric analysis was performed under blind conditions, using ImageScope software (Leica). Two perpendicular sections, one at the edge and another one at the center of each construct, were used. For each section, the entire length of the implant/meniscus interface was measured, as well as the length of any areas of integration at the interface. The repair index was then determined as:

\[ \% \text{Integration} = \frac{\text{Length of integrated interface}}{\text{Total interface length}} \times 100 \]

2.7 | Immunosuppression

Human peripheral blood mononuclear cells (PBMCs) were isolated from donor blood samples from healthy volunteers. All the volunteers gave their informed consent and the study was carried out according to local ethical guidelines (University of Liverpool Research Ethics sub-committee for Physical Interventions). PBMCs were isolated by centrifugation of the blood on 1.077 g/mL Ficoll-Paque PLUS (GE Healthcare Life Sciences) and cultured in RPMI-1640 containing L-glutamine (Sigma-Aldrich) and supplemented with 10% Human AB Serum (Sigma-Aldrich) and 1 vol%/vol% Penicillin/Streptomycin. PBMCs were stained with CellTrace Violet (ThermoFisher) to monitor T-cell proliferation. Labeled PBMCs were then stimulated with 3.75 μg/mL anti-human CD3 (HIT3a) and 2 μg/mL anti-human CD28 (CD28.2) (both from Fisher Scientific Affymetrix eBioscience) and cocultured with MSCs from four individual donors at passages 1, 5, 10, and 15 for 72 hours. The T-cell proliferation profile for each population was analyzed by flow cytometry following exclusion of non-viable cells stained with 7-amino-actinomycin D (7-AAD; BD Biosciences).

2.8 | Quantitative PCR

Real-time quantitative polymerase chain reaction (RT-qPCR) for MMP13 mRNA and for the housekeeping gene Tata Binding Protein (RBP) was performed using the CellsDirect One-Step qRT-PCR Kit (ThermoFisher), with which reverse transcription and PCR amplifications were performed in the same reaction tube. Primers specific for MMP13 (Hs00942584_m1) and TBP (Hs00427621_m1) were purchased from ThermoFisher TaqMan. The reaction was started by synthesizing cDNA at 50°C for 15 minutes, followed by 2 minutes at 95°C to denature RNA-cDNA hybrids and deactivate reverse transcriptase. The thermal cycling program consisted of 50°C for 15 minutes, 95°C for 2 minutes, and 40 two-step cycles of 95°C for 10 seconds and 60°C for 30 seconds. MMP13 expression relative to TBP was determined at each of four time points for each MSC sample and the results normalized to the time (passage 2), which was taken as a Fold-expression of 1.0.

2.9 | TIMP-1 ELISA

TIMP-1 protein was measured in the secretome of MSCs using the Quantikine enzyme-linked immunosorbent assay (ELISA) Kit for
human TIMP-1 (R&D Systems). MSCs were seeded into 6-well plates at 2.25 million cells/well (three replicates) and cultured in 2 mL of DMEM for 24 hours. The medium was then replaced with 1 mL of phenol-free culture medium for a further 24 hours, after which the secretome was collected and assayed at appropriate dilution using the ELISA kit.

2.10 | Acquisition of genomic and proteomic data

2.10.1 | Transcriptomics

Transcriptomics was performed by the Centre for Genomic Research on mRNA extracted from the mRNA of all four patients at P1, P5, P10, and P15. When MSCs were harvested at the end of each passage, 1 × 10^6 cells were isolated and resuspended in RNAprotect Cell Reagent (Qiagen). The cells were stored at −80°C until the complete set of samples from all donors and time points had been collected. RNA was then extracted from selected time points using the RNeasy Plus Mini Kit (Qiagen), according to the manufacturer’s instructions. The concentration of RNA in the extract was determined using a NanoDrop 2000 spectrophotometer (Thermo). Extracted RNA was stored at −80°C prior to analysis.

Ribosomal RNA depletion was performed using the Ribo-Zero H/M/R Kit (New England Biolabs) and RNASeq libraries were then prepared using the NEB Next Ultra Directional RNA Library Prep Kit (New England Biolabs). Paired-end sequencing of the RNASeq libraries was performed by the Illumina HiSeq4000 platform using V4 chemistry.

2.10.2 | Proteomics

Proteomics was performed on secretome prepared from the MSCs of all four patients at P1, P5, P10, and P15. When MSCs were harvested at the end of each passage, 1 × 10^6 cells were isolated and resuspended in 4 mL of serum-free, phenol red-free DMEM supplemented with 4500 mg/L (high) glucose (Sigma-Aldrich), 1 vol%/vol% % Glutamax, and 1 vol%/vol% Penicillin/Streptomycin for 24 hours at 37°C. The conditioned medium recovered at the end of the incubation was harvested from each flask and stored at −80°C prior to analysis.

Proteomic analysis of the secretome samples was undertaken in the "Centre for Proteome Research" facility at the University of Liverpool. Protein solutions were concentrated by adding consecutive 1 mL aliquots of each sample to 10 μL of Strataclean beads (Agilent). After each aliquot had been added, the sample was vortexed for 1 minute, centrifuged at 2000g for 2 minutes and the protein-depleted supernatant removed. After the final aliquot had been added, the beads were washed two times with 1 mL of 25 mM ammonium bicarbonate (ambic) prior to digestion. For on-bead digestion, the beads were resuspended in 80 μL of 25 mM ambic and 5 μL of 1 wt%/vol% Rapigest (Waters) was added. The samples were heated at 80°C for 10 minutes and then reduced by the addition of 5 μL of 9.2 mg/mL dithiothreitol (Roche) in 25 mM ambic and heating at 60°C for 10 minutes. Following cooling, 5 μL of 33 mg/mL iodoacetamide (Sigma-Aldrich) in 25 mM ambic was added and the samples incubated at room temperature for 30 minutes in the dark. Porcine trypsin (sequencing grade, Promega) (1 μg) was added and the sample was incubated at 37°C overnight on a rotary mixer. The digests were acidified by the addition of 1 μL of trifluoroacetic acid (TFA) and incubated at 37°C for 45 minutes. Samples were then centrifuged at 17 200g for 30 minutes and supernatants transferred to 0.5 mL low-binding tubes. They were centrifuged for a further 30 minutes and 10 μL transferred to total recovery vials for liquid chromatography–mass spectrometry (LC-MS) analysis.

Data-dependent LC-MS analyses were conducted on a QExactive HF quadrupole-Orbitrap mass spectrometer coupled to a Dionex Ultimate 3000 RSLC nano-liquid chromatograph (Hemel Hempstead, UK). Sample digest (1-2 μL) was loaded onto a trapping column (Acclaim PepMap 100 C18, 75 μm × 2 cm, 3 μm packing material, 100 Å) using a loading buffer of 0.1 vol%/vol% TFA, 2 vol%/vol% acetonitrile in water for 7 minutes at a flow rate of 12 μL min⁻¹. The trapping column was then set in-line with an analytical column (EASY-Spray PepMap RSLC C18, 75 μm × 50 cm, 2 μm packing material, 100 Å) and the peptides eluted using a linear gradient of 96.2% A (0.1 vol %/vol % formic acid): 3.8% B (0.1 vol%/vol% formic acid in water : acetonitrile [80:20] [vol/vol]) to 50% A:50% B over 90 minutes at a flow rate of 300 nL min⁻¹, followed by washing at 1% A: 99% B for 5 minutes and re-equilibration of the column to starting conditions. The column was maintained at 40°C, and the effluent introduced directly into the integrated nano-electrospray ionization source operating in positive ion mode. The mass spectrometer was operated in data dependent acquisition mode with survey scans between m/z 350-2000 acquired at a mass resolution of 60 000 (full width at half maximum) at m/z 200. The maximum injection time was 100 ms, and the automatic gain control was set to 3e6. The 16 most intense precursor ions with charges states of between 2+ and 5+ were selected for MS/MS with an isolation window of 2 m/z units. The maximum injection time was 45 ms, and the automatic gain control was set to 1e5. Fragmentation of the peptides was by higher-energy collisional dissociation using a normalized collision energy of 30%. Dynamic exclusion of m/z values to prevent repeated fragmentation of the same peptide was used with an exclusion time of 20 seconds. Raw data files from the mass spectrometry were imported into Progenesis QI for Proteomics v.2.0 software (Waters Ltd, Newcastle upon Tyne, UK) for alignment and peak detection. An aggregate file was generated which contained all the peaks from all runs in an experiment so that there were no missing values. The data were filtered and charges +1 and ≥ +8 were removed. The msms fragmentation data were searched against the UniProt human reviewed DB using Mascot v. 2.4.1 software (Matrix Science, London, UK). The precursor ion mass tolerance was set to 10 ppm and the product ion tolerance to 0.01 Da. Oxidation of methionine was selected as a dynamic modification and carbamidomethyl cysteine as a fixed modification. One missed cleavage was permitted. The Mascot search returned 2594 proteins at 2.17% false discovery rate (FDR: psm’s above homology). A 1% FDR was set
and 2366 proteins were exported as an .xml file (FDR type: distinct psms) and imported into Progenesis and peptides assigned to proteins. Protein quantification was based on averaging the individual abundances for each protein per donor at each passage and comparing proteins that were differentially expressed between the four passages.

2.11 Quantification and statistical analysis

2.11.1 Bioinformatics

Data processing, integration and analyses were undertaken by the Computational Biology Facility at the University of Liverpool. RNAseq data were acquired as described above. The raw Fastq files were trimmed for the presence of Illumina adapter sequences using Cutadapt version 1.2.1, option -O 3. Reads were further trimmed using Sickle version 1.200 with a minimum window quality score of 20. Reads shorter than 10 base pairs after trimming were removed. Sequence quality metrics were assessed using FastQC version 0.11.4. No samples were removed. Sequence data were aligned to the human genome version GRCh38 from NCBI using Bowtie2 version 1.1.2 with recommended parameters. Gene level count data were generated from the Bowtie2 alignments using htseq-count version 0.9.0. The R library DESeq2 was used to produce rlog transformed count data. These were filtered to remove genes with less than 1 average count. Statistical analyses were performed in R version 3.4.4 and graphical representations were done using the R package ggplot2.

Normalized proteomics and transcriptomics data were integrated and preliminary exploratory analyses revealed a relevant heterogeneity between patients. In order to discriminate between changes related to patient heterogeneity and changes related to passage, differentially expressed variables over passage were calculated with a two-way analysis of variance (ANOVA) to account for patient variability as confounder. This followed multiple testing corrections using the Benjamin-Hochberg method. Significant variables (338 of which 38 were proteins and 300 were genes) uniquely over passage were selected for further analysis in the Ingenuity Pathway Database (IPA, Qiagen, Analysis 2015).

Base 2 log fold changes of the 338 significant variables were calculated with respect to passage 1 and used for a IPA (Qiagen), where analysis was performed according to the manual to predict possible upstream regulators involved in the changes observed, possible pathways affected and a number of functions and terms significantly enriched (database access on 9 December 2018). IPA was also used to download knowledge terms involved in selected key processes, that is, (a) Cell migration in MSCs, (b) cell movement in MSCs, and (c) wound healing. All these variables were mapped to our experimental

FIGURE 1 Growth and phenotypic characteristics of MSCs from four patients. A. The number of population doublings reached after each passage using MSCs from each patient was recorded until growth ceased (for PN242 data were collected until passage 30, when the cells continued to grow). B. The population doubling time is shown up to passage 17 for all four patients. C. Typical cell morphology is shown for PN251 MSCs at passage 4 and passage 16 (growth arrest was at passage 17 for this patient) and for PN242 at passage 19 (no growth arrest observed for this patient). Scale bars = 200 μm. D. The percentage of cells expressing MSC markers CD90 and CD105 and hematopoietic stem cell markers CD34 and CD45 were determined by fluorescence-activated cell sorting. MSCs, mesenchymal stem cells.
data. Then we assessed the number of variables changing over passage within these terms and contextualize the biological findings.

Principal component analysis (PCA) was performed in mean centered and scaled data via singular value decomposition using the function `prcomp` within the statistical software R.

### 2.11.2 Statistics

Differentiation variables (Tissue engineering, GAG, and osteogenesis and adipogenesis scores) were compared over passage by calculating the non-parametric Spearman correlation embedded in the function `cor.test` within the stats package in R. Osteogenesis and adipogenesis capabilities were measured with semiquantitative data based on image analysis. These were translated into integers 0 to 1 to undertake the calculation.

### 3 RESULTS

#### 3.1 Patient characteristics and MSC growth potential

Bone marrow was collected from patients undergoing knee or hip arthroplasty. All patients gave informed consent and the study was performed in full accordance with local ethics guidelines (Southmead Research Ethics Committee Ref 078/01). Table S1 shows the patient characteristics. All four were male with a mean age of 49 years (range: 38-70 years) at the time of operation. MSCs were isolated from each bone marrow by plastic adhesion and grown under standard culture conditions until they failed to proliferate any further. MSCs from PN241 and PN242 continued to proliferate for a larger number of passages than MSCs from PN251 and PN264, with PN242 cells showing no sign of growth arrest even at passage 30 (Figure 1A and Table S1).

Figure 1B shows the MSC PDT at each passage for each patient up to passage 17. The PDT of MSCs from all four patients was similar at early passages and became longer at later passages, with MSCs from one patient (PN264) showing particularly slow growth at higher passage numbers. Early passage MSCs had a typical small size and stellate appearance, but became larger and less spindly as the cells became more senescent (Figure 1C, PN251). However, PN242 cells had a small size and spindly, stellate appearance even at very high passage number, when they were continuing to proliferate well (Figure 1C, PN242). These results are consistent with previous studies showing that cell enlargement and loss of stellate shape are associated with increased senescence.

Cell surface marker expression was determined by fluorescence-activated cell sorting (FACS) using MSCs from all four patients at each of passages P1, P5, P10, and P15. The expression of MSC markers...
CD105 and CD90 was maintained at >90% at late passage in MSCs from all four patients. Hematopoietic stem cell marker CD45 remained at <10% at all passages; however, there was a small increase in the expression of hematopoietic stem cell marker CD34 with increasing passage number in two of the patients (Figure 1D).

3.2 The tri-lineage differentiation potential of MSCs falls with increasing passage

MSCs from selected passages of each patient were tested for their chondrogenic potential in a three-dimensional cartilage tissue engineering assay, with the amount of engineered cartilage measured as the dry weight of tissue and quality of the cartilage measured as the GAG content expressed as a percentage of dry weight. There was clear evidence of a loss of chondrogenic potential with in vitro aging of the cells. The typical macroscopic appearance of tissue engineered cartilage over a range of passages for MSCs from each of the patients can be seen in Figure 2, clearly demonstrating a reduction in the average size of cartilage constructs when produced using late passage MSCs. This macroscopic observation was supported by quantitative analysis. There was a significant negative correlation between the mean dry weight of cartilage formed and the cumulative population doublings of MSCs recorded at the time of tissue engineering for three out of the four patients and a significant negative correlation between the mean cartilage GAG content and the cumulative population doublings of MSCs recorded at the time of tissue engineering for all four patients (Figure 3A-D). Furthermore, both the dry weight and the GAG content of all the individual tissue engineered cartilage constructs were significantly negatively correlated with the passage number of the MSCs used for tissue engineering (Figure 3E,F).

MSCs were also tested for their osteogenic potential in monolayer culture by semiquantitative analysis of the staining for alizarin red. Typical images of the staining patterns and the scoring systems used are shown in Figure S1. There was some evidence of a gradual loss of osteogenic potency of MSCs; however, the results were inconsistent between patients. For patient PN241, there was no apparent change in the mean osteogenic score with cumulative population doublings (Figure S2A). For patient PN242, there was a significant fall in osteogenic potential with cumulative population doublings (Figure S2A). For patient PN242, there was a significant fall in osteogenic potential with cumulative population doublings
(Figure S2B). PN251 osteogenic scores had already declined to 0 by 10 population doublings and remained at 0 for higher doublings, although the change was not statistically significant (Figure S2C). Patient PN264 exhibited no observable osteogenic activity even at the earliest number of population doublings (Figure S2D). Overall, the individual osteogenic scores for the repeated assays were not significantly correlated with the passage number of the MSCs used (Figure S2E); however, the different patterns observed with each of the patients are indicative of a gradual, if variable decline in osteogenesis with in vitro aging of the cells.

The adipogenic potential of MSCs was tested in monolayer culture by semiquantitative analysis of the staining for Oil-Red-O. Typical images of the staining patterns and the scoring systems used are shown in Figure S3. For patient PN251, there was a small but significant fall in adipogenic potential with cumulative population doublings (Figure S4C), whereas there was no significant change for the other three patients (Figure S4A-C). Overall, the individual adipogenic scores for the repeated assays were not significantly correlated with the passage number of the MSCs used (Figure S4E) and therefore only very limited evidence for a fall in adipogenesis with in vitro aging of the cells.

Collectively, these data indicate a clear loss of differentiation potential of MSCs, with early passage cells tending to show full tri-lineage potential and later passage cells tending to retain only adipogenic potential or adipogenic plus osteogenic but not chondrogenic potential. These data suggest that multipotential differentiation is not a fundamental property of CD105+ve, CD90+ve, CD34−ve, CD45−ve MSCs, but rather a feature of freshly isolated MSCs that is lost with cell expansion in vitro.

3.3 | Trophic repair by MSCs is maintained with increasing passage

We used meniscal cartilage integration and suppression of T-cell proliferation as assays relating to different aspects of trophic repair.
The in vitro integration of two pieces of meniscal cartilage by MSCs was based on our previous studies of this system in vitro, in an in vivo sheep model and in patients with torn meniscus. MSCs from all four patients at each of 17 passages were tested for their potency in our meniscal repair potency assay. The variation in % integration of meniscus in these studies was in the same range as previously described, as illustrated in typical histological images (Figure 4A-C). In contrast to our observations of tri-potential differentiation (see above), there was no significant change in the mean potency of meniscal repair with cumulative population doublings for any of the four patients (Figure 4D-G). Furthermore, the % integration measured for the individual meniscal constructs showed no correlation with the passage number of the MSCs used (Figure 4H).

As well as stimulating tissue repair responses, MSCs are also able to suppress immunity through inhibition of T-cell proliferation and other mechanisms. We therefore measured the % inhibition of T-cell proliferation by MSCs from selected passages for all four patients. As with meniscal repair, there was no significant change in the immunoregulatory capacity of MSCs with increasing passage (Figure 4I).

### 3.3.1 Transcriptomics and proteomics data show significant differences over passage that can be linked to loss of multipotency

We reserved mRNA from the undifferentiated MSCs of all four patients at each passage. Based on growth and differentiation characteristics, we selected mRNA from passages P1, P5, P10, and P15 for gene array comparison. For each of these passages, we also collected conditioned medium for proteomic comparison. The genomic and proteomic data were combined and analyzed for patterns of change in

![Figure 5](image-url)

**Figure 5** Transcriptomics and proteomics. A, Overview of analytical steps performed. Transcriptomics and proteomics data were processed according to data standards and integrated. The integrated data were further analyzed by Ingenuity Pathway Analysis to identify key genes and proteins that change over passage and map them to biological functions and predict possible upstream regulators (see Figures 6 and 7). B, PCA score plot of first two principal components of all genes and proteins. There is a segregation of patients (see blue and orange ellipses) that can also be linked to changes in osteogenic capabilities (see Figure S2), while changes related to passage are capture mainly in PC2. C, Venn diagram of significant variables after two-way analysis of variance revealing 338 genes and proteins that vary significantly and independently of the patient group variation. D, PCA of the 338 significant variables over passage. Shown the score plot of the first two principal components (capturing approximately 70% of variance). As expected, there is a marked segregation over passage. Heatmap of scaled data from the 338 significant proteins and genes that are represented in rows and clustered via Ward hierarchical clustering. Columns represent biological samples, each column is a patient sample and they are ordered by passage with red being passage 1, green passage 5, turquoise passage 10, and purple passage 15. PCA, principal component analysis.
related genes and proteins. The methodological approach to transcrip
tomic and proteomic analysis is illustrated in Figure 5A. Data
structure was appraised via PCA. A score plot of the first two principal
components is shown in Figure 5B. The changes captured over pas-
sage are accounted for in PC2 while PC1 captures considerable vari-
tion between patients which can be stratified in two groups (shown
with blue and orange ellipses). These two groups of patients also
show differences in osteogenic capabilities (see above). With the aim
of identifying genes and proteins changing solely over passage, inde-
dependently of this patient variability, we performed a two-way ANOVA
for each variable, as described in the Section 2. Figure 5C shows the
significant variables of these tests. Only 338 genes and proteins are
significant uniquely over passage, independently of patient to patient
variation or any related iteration. These 338 genes and proteins were
used to calculate PCA, of which the first two principal components
are shown in Figure 5D together with a heatmap of the 338 variables
selected. As expected, these genes and proteins show a very clear
separation of all the patient samples by passage with a subset
decreasing their expression/abundance over passage (top half of the
heatmap) while others increase their expression/abundance (bottom
half of the heatmap).

For each of these 338 significant proteins and genes, we cal-
culated the fold change with respect to passage 1 and analyzed these
data with Ingenuity Pathway Analysis (IPA) (QIAGEN, Inc., https://
www.qiagenbioinformatics.com/products/ingenuity-pathway-
analysis). IPA’s core analysis, overlaid with the global molecular net-
work within the software resulted in the identification of a number of
canonical pathways, functions, and upstream regulators found to be
significantly over-represented within this list and therefore linked to
the loss of multipotential differentiation capacity of the cells. The larg-
est and most significant change was in the cell-cycle master regulator
FOXM1 gene pathway, with clear evidence from transcriptomic data
for downregulation of the FOXM1 gene itself (Figure 6A) and with six
out of seven of its downstream effectors also predicted by IPA to be
downregulated (Figure 6B). Other upstream regulators predicted to be
deactivated were the prostaglandin receptor PTGER2, and members
of the Vascular endothelial growth factor (VEGF) family, while
upstream regulators that were also predicted to be positively acti-
vated over passage included the proliferation regulator NUPR and the
cytoskeleton regulator MYOC. Quantitative data from our trans-
criptomic analyses and the associated IPA predictions of changes in
the downstream effectors of these regulators are shown in Figure S5;

FIGURE 6  Ingenuity Pathway Analysis of integrated transcriptomics and proteomics data. A, Ingenuity pathway analysis identified FOXM1
and four other master regulators genes (see Figure S5) that vary with increasing passage number. FOXM1 data from transcriptomics analysis
show inhibition of gene expression with increasing passage number. Each bar is the mean ± SEM for results using MSCs from each of the four
patients. B, Ingenuity pathway analysis of the integrated transcriptomics and proteomics data set predicts inhibition of five out of six of the
identified downstream regulators in the FOXM1 canonical pathway. C, Ingenuity pathway analysis identified multiple genes and proteins that
mapped to the search terms “Cell Movement,” “Cell Migration,” or “Wound Healing.” Most of these identified genes and proteins did not vary
significantly over passage as determined by analysis of variance. The most highly expressed genes and proteins are listed for each of the search
terms. D, CXCL12 data from transcriptomics analysis show continuous gene expression with increasing passage number. Each bar is the
mean ± SEM for results using MSCs from each of the four patients. E, CXCL12 data from proteomics analysis show continuous protein secretion
with increasing passage number. Each bar is the mean ± SEM for results using MSCs from each of the four patients. MSCs, mesenchymal stem
cells.
however, none of the changes and predicted downstream effects were as clear-cut as for FOXM1.

3.4 Regulators of cell migration and wound healing can be linked to the trophic properties of MSCs

As shown in Figure 4, the trophic repair capacity of MSCs remains unchanged with increasing passage. With the aim of further investigating this phenomenon, we hypothesized that (a) the proteins and genes involved in regulating this function would not present a significant change in abundance/expression over passage and (b) functions related to cell movement, cell migration, and wound healing are likely to be mechanistically involved in trophic repair. To investigate this hypothesis, we used the IPA database to identify the proteins and genes involved within the three terms listed above. Then we mapped those lists to our data, extracted overlapping variables, and appraised their significance over passage. The large majority of the proteins and genes mapping to these search-terms were found to be unchanged over passage and expressed at a consistent level across all four passages of our MSC cultures (Figure 6B), supporting the hypothesis that genes and proteins expected to be necessary for trophic repair continue to be expressed in aging cells, when multipotent differentiation capacity has been lost but trophic repair capacity remains high. We considered CXCL12 (also called stromal cell-derived factor 1) to be of particular importance at it is associated with all three of our search terms (Figure 6C) and was expressed consistently highly at both gene and protein level (Figure 6D,E).

3.5 Marker genes and proteins

The IPA analysis outlined above demonstrated downregulation of the FOXM1 canonical pathway with increasing passage/loss of multipotent differentiation and continuous expression of CXCL12 and other cell migration and wound healing genes and proteins with increasing passage. However, we consider it necessary also to identify genes and proteins that may not be part of canonical pathways or gene/protein families, but that can be used as specific markers of...
cellular aging in vitro. Such markers would aid in comparison of studies of cells from one laboratory to another or in determining the functionality of an MSC population being used for therapeutic purposes. We therefore analyzed the gene array and protein data to identify candidate markers.

Within the significant variable genes identified by transcriptomic analysis, there was a significant decrease with increasing passage in the gene for matrix metalloproteinase 13 (MMP13; Collagenase 3; Figure 7A and Table S2) and a significant increase with increasing passage in the gene for Insulin-like growth factor binding protein 5 (IGF-binding protein 5; Table S2). There was no significant change in other MMP genes (Table S2) or IGFBP genes (Table S3), nor in any of the genes of the transforming growth factor family (Table S4). Because the transcriptomic data indicated that the rate of fall in MMP13 gene expression with passage mirrored closely the fall in chondrogenic potential (cf Figure 7A with Figure 3), we went on to validate these results using quantitative PCR to determine more accurately the changes in MMP13 gene expression with increasing passage. The results confirm a continuous decline in MMP13 gene expression with increasing passage (Figure 7B), demonstrating that loss of MMP13 gene expression could be used to help determine the extent to which MSCs have aged through in vitro proliferation.

Within the proteomic data set, we identified those proteins expressed at highest abundance at all passages (Table S5). The most abundant of these proteins was metalloproteinase inhibitor 1 (tissue inhibitor of metalloproteinase 1 [TIMP1]; Table S5 and Figure 7C). We went on to validate these results using an ELISA kit assay to determine more accurately the changes in TIMP-1 protein secretion with increasing passage (Figure 7D), demonstrating that expression of TIMP-1 is a defining characteristic of MSCs, irrespective of the extent to which they have aged through in vitro proliferation. Furthermore, when MSCs were seeded onto collagen scaffolds and cultured for 24 hours, the cells continued to secrete high levels of MSCs from the cell/scaffold constructs into their culture medium; however, if the constructs were freeze-thawed under conditions that reduced their viability, the secreted TIMP levels were substantially reduced (Figure 7E).

These results indicate that MMP13+, TIMP1-secreting high MSCs can be used as the basis of injectable therapies for osteoarthritis that are intended to act primarily through engraftment and chondrogenic differentiation. In contrast, MMP13-, TIMP1-secreting high MSCs can be used for OA therapies that are intended to act primarily through trophic repair.

4 | DISCUSSION

We have identified two functional markers relating to MSC aging in vitro, namely the MMP13 (Collagenase 3) gene, which is down-regulated with increasing passage, and TIMP-1 protein, which is the most abundant protein in the secretome at all passages. It is particularly interesting that these two markers, which were identified independently of each other (one from transcriptomics data and one form proteomics), are biologically related to each other as TIMP-1 is an inhibitor of MMP13 as well as other metalloproteinases. This relationship is important because it could explain how production of a catabolic proteinase (MMP13) correlates to an anabolic capacity of MSCs (chondrogenesis). The very high levels of TIMP-1 produced by undifferentiated MSCs at all passages will most likely inhibit any activated MMP13, rendering it catabolically inert. In an OA cell therapy context, TIMP-1 protein secretion could be used as a specific marker of MSC identity and viability as it is constitutively expressed at very high levels at all passages whether growing on tissue culture plastic or seeded into three-dimensional scaffolds. MMP13 gene expression could be used as a functional marker to distinguish chondrogenic MSCs from those that have good trophic capacity but poor chondrogenic capacity.

These results also have important implications for the strategies that should be used in the development of injectable OA therapies. Since MMP13+, TIMP1-secreting high MSCs were lost as early as passage 5, it seems likely that OA therapies that are intended to act through engraftment and chondrogenic differentiation will be restricted to the use of autologous cells, because viable allogenic products would require far more extensive expansion. While we cannot rule out the possibility that alternative culture condition to those used here could be identified as a method of sustaining the chondrogenic phenotype even after in vitro aging, the present data nevertheless lead us to conclude that allogeneic injectable stem cell therapies for OA should rather focus on the use of MMP13+, TIMP1-secreting high MSCs acting through trophic repair mechanisms.

Our results also have important biological implications. They clearly indicate that the loss of multipotent differentiation capacity of MSCs with extensive passage in vitro cannot be the result of a generalized loss of cell function as the MSCs senesce, since another important property of MSCs, trophic repair, is not significantly reduced even after a very high number of population doublings and reduced chondrogenic differentiation capacity was observed many passages before cessation of proliferation. Furthermore, families of genes and proteins from a range of critical signaling pathways show clear changes that correlate with increasing passage number and loss of differentiation, whereas genes and proteins known to be involved with wound repair and cell movement/migration do not vary significantly with increasing passage number. Downregulation of the Forkhead Box M1 (FOXM1) canonical pathway showed a clear relationship to passage number, indicating a potential role for FOXM1 in the maintenance and then loss of multipotency. This observation is consistent with previous studies of its biological function. It is a proto-oncogene that is a key master regulator in the survival of cancer stem cells. It has also been shown to be highly expressed in multipotent and pluripotent stem cells and to be critical to the maintenance of stem cell potency and to the induction of pluripotency through reprogramming. CXCL12 (also known as SDF-1) was found to be associated with all three of our search terms related to trophic repair and its gene and protein levels were maintained even at very late passage numbers, indicating a potential role for CXCL12 in MSC-mediated trophic repair. This observation is consistent with previous
studies which have demonstrated its critical role in MSC-mediated induction of spinal cord repair\textsuperscript{53} and myocardial repair,\textsuperscript{54} as well as enhancing nerve cell survival in vitro\textsuperscript{55} and mediating trophism between endothelial cells and tumor cells.\textsuperscript{56} We have identified FOXM1 and CXCL12 through combined proteomic and genomic analysis, but while this in silico approach is a powerful tool, any mechanistic involvement of these regulators must await experimental confirmation.

Previous studies have clearly shown that multipotency of MSCs declines with both in vitro and in vivo aging.\textsuperscript{33-39} Our observation here that chondrogenesis declines, osteogenesis tends to decline, but adipogenesis is retained at higher passage number is in agreement with the work of Yang et al.\textsuperscript{34} Muraglia et al\textsuperscript{26} also showed a loss of multipotent differentiation with passage, but in their experiments, osteogenesis was retained with early loss of adipogenesis. However, they were working with clonal MSC cell-lines whereas the work reported here and by Yang et al investigated the whole MSC population.

Haynesworth et al first described the unique cytokine expression pattern of MSCs\textsuperscript{57} and they demonstrated a reduction in cytokine production after stimulating differentiation using dexamethasone. A decade later, Caplan and Dennis coined the term “trophic effect” which they defined as “those chemotactic, mitotic, and differentiation-modulating effects which emanate from cells as bioactive factors that exert their effects primarily on neighboring cells and whose effects never result in differentiation of the producer cell.”\textsuperscript{22} They cited, as a typical example of this effect, the support provided by MSCs in the bone marrow niche for growth and differentiation of hematopoietic cells. They also summarized evidence for MSCs providing trophic support in a range of cell therapy settings including the treatment of stroke, myocardial infarction, and meniscal cartilage regeneration. Numerous other studies have gone on to describe the importance of MSC-induced trophic repair.\textsuperscript{20,22-25} We exploited the trophic effects of MSCs in devising a new method of treating fresh meniscal cartilage tears using a stem cell/collagen-scaffold implant to promote integration of the damaged tissue. Our original in vitro studies demonstrated that the method depends on cell migration from the implant into surrounding tissue and interaction with the endogenous meniscal cells.\textsuperscript{26,46} Importantly, we found that MSCs that have been stimulated with transforming growth factor-\( \beta \) to undergo chondrogenic differentiation are much less potent at promoting meniscal repair than the undifferentiated MSCs.\textsuperscript{26} We went on to describe the use of undifferentiated MSCs seeded on a collagen scaffold to repair meniscal injury in a sheep preclinical model and in a first in human trial.\textsuperscript{27} In the current study, we used our in vitro semiquantitative meniscal cartilage integration assay\textsuperscript{26} as a model for trophic repair and made the very surprising observation that MSCs that have been cultured for up to 30 passages retain the same capacity for trophic repair as very early passage cells. These functional data were supported by our analysis of genes and proteins involved with cell movement and migration and with wound healing, showing that unlike those linked to differentiation, there was no significant change in their expression with increasing passage.

Another aspect of MSC trophism is the immunoregulatory effects of MSCs. Although a range of mechanisms are involved, it is clear that inhibition of T-cell proliferation is a critical component of their suppressive activity.\textsuperscript{9,24,28-31} Human T cells can be strongly stimulated to proliferate using a combination of anti-CD3 and anti-CD28 antibodies.\textsuperscript{58} Their rate of proliferation can be monitored by covalently labeling intracellular molecules with a fluorescent dye that is then diluted out by 50% with each cell division, tracked by FACS.\textsuperscript{59} Adding MSCs into cultures of labeled, stimulated T cells suppresses the lymphocyte proliferation, so prolonging the accumulation of the dye.\textsuperscript{9} In the current study, we have used this method to measure the immunoregulatory effects of MSCs at early and late passage and found no loss of potency with increasing passage. For human MSCs, the mechanism of T-cell suppression has been described as involving indoleamine 2,3-dioxygenase-mediated tryptophan degradation.\textsuperscript{60} Taken together, these results demonstrate that, as with trophic repair, immunoregulation is a fundamental property of MSCs that is not lost with increasing passage.

Previous studies have proposed that the loss of multipotent differentiation capacity of MSCs with increasing passage is related to progression of the cells to a senescent end-state.\textsuperscript{33-36} However, the results described here demonstrate that even after extensive passaging in vitro, there is no apparent loss of trophic repair. The term “Mesenchymal Stem Cells” was coined by Caplan,\textsuperscript{1} who went on to describe MSCs as “An injury Drugstore”\textsuperscript{23} and more recently advocated a change in their name to “Medicinal Signaling Cells.”\textsuperscript{61} Other studies have questioned the definition of MSCs as stem cells because of the lack of rigorous confirmatory biological evidence.\textsuperscript{20,25,31,62,63} With all of these studies calling for more experimental data before reaching a conclusion on the nomenclature. Others have been more forthright in concluding that MSCs are not stem cells and have called for an immediate change in nomenclature, in order to avoid the overhyped marketing of MSCs as “miracle cures.”\textsuperscript{64-66} Prockop emphasized that the essence of a stem cell should not be determined by its status at a single point in time.\textsuperscript{25} In this study, we have investigated the in vitro differentiation and trophic behavior of MSCs across many passages over time and in this way, have reached the conclusion that the property of multipotency is relatively transient whereas the trophic effects of these cells is apparently permanent all the way through to the time of growth arrest.

5 | CONCLUSION

These studies demonstrate that the development of injectable MSC therapies for OA must take into account the transient nature of chondrogenic potency relative to their sustained trophic potency with increasing passage and specific strategies should be adopted to exploit one or other of these mechanisms of action.

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CONFLICT OF INTEREST
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AUTHOR CONTRIBUTIONS
A.S., K.B.: collection and/or assembly of data, data analysis and interpretation, manuscript writing; M.R., C.L.: collection and/or assembly of data, manuscript writing; E.C.-G., F.F.: data analysis and interpretation, manuscript writing; A.W.B., M.R.W.: provision of study material or patients, manuscript writing; A.P.H.: conception and design, financial support, data analysis and interpretation, manuscript writing.

DATA AVAILABILITY STATEMENT
The accession number for the RNA-seq data reported in this article and deposited at the NCBI Gene Expression Omnibus (GEO) database is: GSE137186. The proteome data reported in this article have been and deposited at the NCBI Gene Expression Omnibus (GEO) database.

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