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Kay, Alasdair Gawain orcid.org/0000-0003-1953-331X, Fox, James Martin, Hewitson, James Philip orcid.org/0000-0002-3265-6763 et al. (7 more authors) (2022) CD317-Positive Immune Stromal Cells in Human "Mesenchymal Stem Cell" Populations. Frontiers in immunology. 903796. ISSN: 1664-3224

https://doi.org/10.3389/fimmu.2022.903796

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CD317-Positive Immune Stromal Cells in Human "Mesenchymal Stem **Cell**" Populations

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- Keywords: Mesenchymal stromal cells, MSC subtypes, heterogeneity, immunomodulation, 15
- 16 CD317, BST2, tetherin
- 17 1 **Abstract**
- Heterogeneity of bone marrow mesenchymal stromal cells (MSCs, frequently referred to as 18
- 19 "mesenchymal stem cells") clouds biological understanding and hampers their clinical development.
- 20 In MSC cultures most commonly used in research and therapy, we have identified an MSC subtype
- 21 characterised by CD317 expression (CD317^{pos} (29.77±3.00% of the total MSC population),
- comprising CD317^{dim} (28.10±4.60%) and CD317^{bright} (1.67±0.58%) MSCs) and a constitutive 22
- interferon signature linked to human disease. We demonstrate that CD317^{pos} MSCs induced 23
- 24 cutaneous tissue damage when applied a skin explant model of inflammation, whereas CD317^{neg}
- 25 MSCs had no effect. Only CD317^{neg} MSCs were able to suppress proliferative cycles of activated
- human T cells *in vitro*, whilst CD317^{pos} MSCs increased polarisation towards pro-inflammatory Th1 26
- 27 cells and CD317^{neg} cell lines did not. Using an *in vivo* peritonitis model, we found that CD317^{neg} and
- CD317^{pos} MSCs suppressed leukocyte recruitment but only CD317^{neg} MSCs suppressed macrophage 28
- 29 numbers. Using MSC-loaded scaffolds implanted subcutaneously in immunocompromised mice we
- were able to observe tissue generation and blood vessel formation with CD317^{neg} MSC lines, but not 30
- 31
- CD317^{pos} MSC lines. Our evidence is consistent with the identification of an immune stromal cell,
- 32 which is likely to contribute to specific physiological and pathological functions and influence
- 33 clinical outcome of therapeutic MSCs.

2 Introduction

- 37 Mesenchymal stromal cells (MSCs) exist in bone marrow at a frequency of approximately 0.001-
- 38 0.01%(1) and are typically self-renewing for 10-50 population doublings(2, 3). MSCs can
- 39 differentiate into skeletal lineages (osteogenic, adipogenic, chondrogenic) and regulate immune cell
- 40 function(4) predominantly through the release of cytokines and other immunosuppressive factors(5).
- 41 The International Society for Cell & Gene Therapy (ISCT) guidelines identifies MSCs as cells that
- 42 exhibit tri-lineage differentiation in vitro and plastic adherence, alongside an expression profile of
- 43 selected cell surface epitopes (e.g. typically presence of CD105, CD73 and CD90, and absence of
- 44 CD45, CD34, CD14 or CD11b, CD79alpha or CD19 and HLA-DR)(6). There has been some
- progress in identifying in vivo markers of MSC populations in mouse and human systems, which
- include LEPR, nestin, CD271, CD146 and CD164(7), however, no single marker for MSCs exists in
- 47 general use. Cells labelled as "MSCs" are used internationally in clinical trials but are rarely
- characterised (using ISCT or any other criteria(8)) and delivery variable success(9). The majority of
- 49 trials assessing efficacy of MSCs currently aim to harness immunomodulatory properties (10), though
- widespread clinical translation is greatly hindered by insufficient data demonstrating strong and
- 51 consistent clinical effect, mechanisms of action and diverse application of selection criteria(11). In
- addition, MSCs from different origins have been applied in clinical trials with varied outcomes for
- disorders including osteoarthritis(12-15), osteoporotic fracture repair(16), rheumatoid arthritis(17-
- 54 19), type 1 diabetes mellitus(20), diabetic kidney disease(21), multiple sclerosis(22, 23), liver
- failure(24-26), amyotrophic lateral sclerosis(27-30) and COVID-19(31-33). Notably, although
- serious adverse events are extremely rare, mild, transient or acute adverse events occurring are often
- 57 related to acute inflammation(13-16, 19, 21, 25, 29, 30), fever (pyrexia)(17, 19, 22, 24, 26, 30, 34),
- infection(12, 16, 21, 23, 30), allergic reactions/hypersensitivity(13, 15, 16, 19) and haematoma(13),
- all of which are implicated in immune responses.
- 60 Studies examining heterogeneity in MSCs have identified multiple subpopulations of MSCs with
- varied potency for both differentiation and immunomodulation (35-40). Heterogeneous populations of
- MSC-like cells have been isolated from both adult and neonatal sources (e.g. bone marrow(41, 42),
- peripheral blood(43), adipose tissue(44, 45), synovial membrane and fluid(46, 47), dental pulp(48),
- endometrium(49), periodontal ligament(50), tendon(51), trabecular bone(52), umbilical cord(53, 54),
- umbilical cord blood(55, 56), placenta(57)). There are further indications that MSC-like cells may be
- present in most vascularised tissues in some form(58, 59). This widespread distribution of MSC-like
- present in most vascularised distinct in some form(36, 37). This widespread distribution of MSC-inco
- cells with varied differentiation capacities and fluctuations in the expression levels of characterising
- 68 surface markers has prompted increasing reports of unipotent tissue-specific MSCs, yet bone
- 69 marrow-derived MSCs are generally considered to be a population composed entirely of cells
- 70 possessing tripotent differentiation capacity(6). This raises the hypothesis that heterogeneous cell
- 71 populations may collectively characterise as MSCs using ISCT (and other) criteria but comprise
- subsets of cells specialised to perform different functions. The widespread reporting of
- 73 immunomodulatory capacities of MSCs and the impact of immune responses during tissue formation
- and comorbidity in degenerative disease highlights the likelihood of a nascent, endogenous
- 75 population of cells that operate primarily to convey or control immune function. This population has
- the potential to support tissue regeneration rather than contributing to it.
- We previously demonstrated the heterogeneity of human MSCs through the identification of multiple
- subpopulations using a clonal isolation and immortalisation strategy that enabled in-depth and
- 79 reproducible characterisation(60). These populations included an immune-primed MSC subtype
- 80 identifiable through positive expression of CD317 (bone marrow stromal antigen-2 (BST2) or
- 81 tetherin) and possessing enhanced immunomodulatory capacity. Here, we tested the hypothesis that

- 82 CD317 positive (CD317pos) stromal cells function primarily to direct the immune response and do
- 83 not contribute to tissue generation or repair in both physiological and pathological processes and
- 84 therefore represent an identifiable MSC subtype.

85 3 Materials and Methods

86 3.1 Cell culture

87 3.1.1 Immortalised MSC lines and primary bone marrow derived human MSCs

- MSC lines immortalised with human telomerase reverse transcriptase (hTERT) were maintained in
- 89 culture as previously described(60). Clonal hTERT-MSCs included the CD317^{pos} Y202 and Y102
- 90 lines, and the CD317^{neg} Y201 and Y101 lines. Low-passage (p1-p5) primary MSCs were isolated
- 91 from femoral heads, obtained with informed consent during routine hip replacement or as explant
- 92 cultures from human tibial plateaux after routine knee replacement(60). Primary MSCs were also
- established from bone marrow aspirates purchased from Lonza. Cells were cultured at 37°C in 5%
- 94 CO₂ humidified atmosphere incubaters using DMEM (Gibco) culture medium supplemented with
- 95 10% foetal bovine serum and 1% penicillin-streptomycin. Cells were routinely passaged at 80%
- confluence and re-seeded at approximately 3500 cells/cm². hTERT cell lines have a consistent
- 97 population doubling time of approximately 25 hours. Growth kinetics varied between primary
- 98 donors.

99 3.1.2 Isolation of primary T cells from tonsillectomy tissue

- 100 Primary donor T cells were retrieved from tonsillectomy donations according to ethical approval. For
- primary MSC co-cultures, cryopreserved CD4+ human cord blood T cells were purchased from Stem
- 102 Cell Technologies. T cells were isolated from mixed T and B cell cultures using nylon wool
- separation(61). T cells were seeded at a density of 1.0×10^6 cells/ml in an appropriately sized tissue
- 104 culture flask. MSC co-cultures with isolated T cells were set up within 24 hours or cells were
- 105 cryopreserved in 10% dimethylsulfoxide (DMSO) in RPMI1640 medium and re-established in
- culture a minimum of 24 hours prior to use.

107 3.2 Rohart test for independent confirmation of MSC status

- The Rohart MSC test was used as an independent measure for distinguishing MSCs from non-
- MSCs(62). The classifier has previously been validated against 1,291 samples from 65 studies
- derived on 15 different platforms, with >95% accuracy with 97.7% accuracy(62).

111 3.3 Flow cytometry

- MSCs were labelled using optimised concentrations of the required primary antibody or isotype
- 113 control (Table S1). After washing, cells were stained with a fluorescent secondary antibody (Table
- 114 S1), where conjugated primaries were not used. As appropriate, cells were washed as required prior
- to incubation with 1:1000 diluted sytox blue for 5 minutes. Analysis was conducted immediately
- 116 following staining.
- 117 Intracellular flow cytometry of MSC was performed on 4% paraformaldehyde (PFA) fixed cells in
- the presence of 0.1% saponin (Sigma). All flow cytometry was performed on a Beckman Coulter
- 119 CyAn ADP flow cytometer and analysed with Summit v4.3 software, or using a Cytoflex S or LX
- and analysed with FCS Express 7. Cell sorting was undertaken using a Beckman Coulter MoFlo
- 121 Astrios and analysed with summit v6.2 software or FCS Express 7. Sorted primary donor MSCs were

- separated based on CD317 expression with CD317^{neg} represent by lowest CD317 expression in 26.26
- $\pm 4.84\%$ of cells and CD317^{pos} representing the highest 2.20 $\pm 0.50\%$ CD317-expressing cells to
- ensure no overlap between subpopulations. Intermediate CD317^{dim} cells were not included in primary
- donor cell testing.

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3.4 Processing of mouse femurs

- Femurs were dissected from C57BL/6J female mice at ages 8-12 weeks immediately after sacrificing.
- All work was carried out under ethical approval from the University of York Department of Biology
- 129 Ethics Committee and Animal Welfare Ethical Review Body. Muscle tissue was removed and femurs
- were fixed in 4% PFA for 24 hours at 4°C, followed by washing with PBS. Bones were then
- decalcified using 10% EDTA in PBS at pH 7.5 for 24 hours at 4°C. After decalcification, femurs
- were cryoprotected by submerging in 30% sucrose in PBS for 24 hours at 4°C. Bones were embedded
- in Optimal Cutting Temperature compound and sectioned using an OTF5000 cryostat (Bright
- 134 Instruments Ltd.). Sections were collected on SuperFrost plus microscope slides (Thermofisher) and
- 135 stored at -70°C.

3.5 Immunofluorescent staining of mouse bone tissues

- 137 Slides were allowed to reach room temperature. Sections were blocked for 45 minutes in 10% goat
- serum (Sigma) + 0.1% Tween-20 in PBS (10% donkey serum (Sigma) + 0.1% Tween-20 in PBS
- where goat primary antibody was used). Primary antibodies (LEPR, CD31, CD317) were diluted in
- 140 1% IgG-free Bovine Serum Albumin (Sigma) + 0.05% Tween-20 (Sigma) in PBS and sections
- incubated in the dark at 4°C overnight in a humidified chamber. All secondary antibodies were added
- at 1:300 dilution in PBS for 1 hour at room temperature in the dark then stained for 10 minutes with
- 143 0.2 μg/ml 4′,6-diamidino-2-phenylindole (DAPI) in PBS. Dried slides were mounted with Prolong
- Gold antifade mounting medium (Invitrogen) and #1.5 thickness glass coverslip (Scientific
- Laboratory Supplies). Slides were left to cure at room temperature in the dark for 24 hours prior to
- image capture using LSM880 or LSM780 (Zeiss) confocal microscopes with excitation wavelengths
- of 405 nm, 488 nm, 561 nm and 633 nm.

148 **3.6** Proteomic analysis of MSC plasma membranes

- 149 Plasma membranes were isolated from the hTERT immortalised clonal lines following the protocol
- of Holley *et al*(63) before mass spectrometry and comparative proteomic analyses were performed by
- the Proteomics laboratory within the University of York Bioscience Technology Facility using LC-
- MS/MS(64) and Scaffold 4 proteome software for initial analysis using 3% false discovery rate.
- 153 Further in-depth examination of protein expression was conducted using the Knime analytics
- platform and ProteoWizard MSOpen technology(65).

155 3.7 Transwell cell migration assays

- 156 Migration assays were performed in transwell polycarbonate membrane cell culture inserts with a
- 157 5μm pore (Corning, Sigma-Aldrich) using 1.25x10⁵ hTERT and primary MSCs, and monocyte-like
- 158 THP-1 and T cell-like HUT-78 (ECACC 88041901) cells in 6 well plates with 1.5 ml of serum-free
- DMEM. After 24 hours, 600 µl of supernatant or DMEM was added in duplicate to the wells of the
- transwell plates. Polycarbonate filters were carefully placed above supernatant and 2.5×10^5 of the
- appropriate cells in 100 µl serum-free RPMI-1640 were applied to the top of the filter and incubated
- for 5 hours before removing transwells. Migrated cells were assessed by flow cytometry. The
- percentage cells undergoing migration towards stimuli was calculated. For CCR2 testing, 500 nM

164 CCR2 inhibitor was used (Teijin compound 1) in supernatant. Inhibition of migration was calculated as a percentage of cell total.

3.8 Examination of Gene Ontology (GO) terms in disease states for comparison with hTERT MSC lines

- A bioinformatics comparison of the hTERT MSC lines gene expression data with publicly available
- transcriptomic data from a range of autoimmune and related disorders was undertaken to identify
- disease states that correlated with upregulated GO terms associated with the CD317^{pos} Y102 and
- 171 Y202 clonal MSC lines(60). Cross-platform validation was performed using Python and GeneSpring
- software was used to analyse outcomes. Datasets that were analysed on Affymetrix microarray
- platforms were normalized to 75th percentile. For data analysed on Agilent microarray platforms,
- 174 Robust Multichip Algorithm normalization was used which included background correction,
- normalization and calculation of expression values. The differing normalization methods were due to
- 176 GeneSpring default settings, but both methods reduce the level of environmental factors affecting the
- 177 results. In all datasets baseline transformation was to the median of all samples, for each probe the
- 178 median of the log summarized values from all the samples was calculated and subtracted from each
- of the samples. Differentially expressed genes were identified as greater than 2-fold upregulation in
- disease state compared to healthy controls, and GeneSpring was used to identify significance
- 181 (p<0.05) in GO term occurrence. The 10 most upregulated GO terms were identified and
- comparisons made between autoimmune disease states and hTERT immortalised MSC lines.

183 **3.9 Quantitative polymerase chain reaction (qPCR)**

- 184 RNA was isolated from cells using TRIzol for cell lysis and Machery-Nagel RNA Nucleospin II kit
- for RNA isolation, with RNA converted to cDNA for gene expression analyses using Superscript IV
- reverse transcriptase enzymes (Invitrogen). Specific primers for gene expression analyses were
- designed and optimised (Table S2). Gene expression analyses were performed as previously
- described(60). Gene expression of eight IFN-γ regulated genes, namely Ly6E, HERC5, IFI44L,
- 189 ISG15, Mx1, Mx2, EPST11 and RSAD2 were amplified in qPCR and fold changes were calculated
- relative to the expression of the housekeeping gene RPS27a and relative to the Y201 cell line or
- 191 CD317^{neg} cells. The $\Delta\Delta$ CT fold changes were log2-transformed and averaged to calculate IFN- γ
- scores, as previously described (66, 67).

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193 3.10 Enzyme-linked immunosorbent assays

- To detect secreted proteins, supernatants from 100,000 cells incubated in 2.5 ml of serum free
- DMEM for 24 hours was analysed for secreted proteins by enzyme-linked immunosorbent assays
- 196 (ELISA) using ELISA kits for CXCL10, CXCL11 (BioLegend); CCL2 (eBioscience); and SAA4
- 197 (Stratech) following manufacturers instructions.

3.11 PCR molecular diagnostics for infectious disease

- Samples of hTERT lines Y201 and Y202 were tested externally and independently (Charles River)
- 200 for viral contaminants using the Human Comprehensive cell line examination and report (CLEAR)
- 201 Panel to detect RNA transcripts for 26 viral components, including virions commonly linked with
- autoimmune disorders (HIV, hepatitis, herpes simplex and herpesvirus, Epstein-Barr virus, BK virus,
- 203 human T-Lymphotropic virus, Lymphocytic choriomeningitis virus and Cytomegalovirus)(68, 69). A
- low copy exogenous nucleic acid was added to sample lysis prior to nucleic acid isolation to serve as
- both a control to monitor for nucleic acid recovery and PCR inhibition. An RNA NRC was used to

- 206 monitor reverse transcription for RNA virus assays. Nucleic acid recovery and PCR inhibition was
- 207 monitored by a PCR assay specific for the NRC template.

208 3.12 T cell activation assay

209 3.12.1 MSC immunomodulation for deactivation and suppression of T cell proliferation

- 210 Co-culture of primary human tonsil T cells with hTERT MSC lines was used to assess the potential
- immunomodulatory impact of CD317^{neg} (Y101, Y201) and CD317^{pos} (Y102, Y202) cell lines on T
- 212 cell proliferation and T helper differentiation. Continual proliferative capacity was used as a measure
- of T cell deactivation. hTERT MSC lines or CD317-sorted primary MSCs were seeded at a ratio of
- 214 1:10 with T cells with 1.0x10⁴ MSCs seeded into a 96-well U bottomed plate and cultured for 24
- 215 hours at 37°C, 5% CO₂. Primary human MSC were sorted for CD317 expression and co-cultured
- with commercially sourced cryopreserved CD4+ human cord blood T cells (Stem Cell
- 217 Technologies).

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244

- For assessment of proliferation, T cells were stained for 15 minutes at 37°C using 1 uM VPD450
- Violet proliferation dye (eBioscience, Inc.). Unstained cells were used as a control. T cells were
- activated using anti-CD3ɛ/CD28 Dynabeads (Thermo Fisher) at a bead-to-cell ratio of 1:1 then
- seeded onto the MSC at a density of 1.0x10⁵/well (ratio 10:1) in 200 μl RPMI-1640 with 10% FBS,
- 222 0.05 μg/mL IL-2 (Peprotech, Inc) or seeded alone (no MSCs) as a control. Plates were cultured for 5
- 223 days at 37°C. T cell proliferation was assessed following removal of Dynabeads with the DynaMag-2
- as per manufacturer's recommendations. Plates were cultured for 5 days at 37°C. T cell proliferation
- 225 was assessed with flow cytometry, with reduction in signal intensity visualised for repeated
- proliferation peaks. Proliferation was assessed through VPD450 dilution (diminished staining
- intensity) described through a proliferative index (PI) calculated from the fluorescence intensity at
- each cell division as described previously(70). Proliferative cycles undertaken were calculated on
- 229 50% fluorescence intensity reduction peaks, measuring from fluorescence intensity of the first
- 230 division and the final division detected.

3.12.2 MSC immunomodulation to direct effector T cell polarisation

- For assessment of T helper differentiation, T cells were activated and cultured with hTERT MSC
- 233 monolayers, as described above. The following reagents and antibodies for reactivation, transport
- inhibition and staining were sourced from eBioscience. Following 5 days of culture, T cells were re-
- stimulated using a combination of phorbol 12-myristate 13-acetate (PMA) (50 ng/ml) (Sigma
- 236 Aldrich) and Ionomycin (1 µg/ml) (Invitrogen) and intracellular cytokines retained using transport
- 237 inhibitor cocktail with 10 µg/ml brefeldin A and 2 µM Monensin (Invitrogen). Cells were cultured for
- 4 hours at 37°C then stained for surface marker CD4. Intracellular staining for helper T cells was
- undertaken for anti-human IFN-γ (Th1), IL-4 (Th2) or IL17a (Th17) or CD4 and CD25 then
- fixation/permeabilisation and staining for nuclear protein FOXP3 for regulatory T cells. All cells
- were measured using the CyAn ADP or Cytoflex LX flow cytometer and analysed with FCS Express
- 7. Comparisons were drawn for percentage of T helper differentiation within the CD4+ cell
- 243 population and signal intensity (Median) for each antibody tested.

3.13 In vitro human skin explant model to assess cutaneous tissue damage

- 245 The human skin explant assay is an *in vitro* model previously used for evaluation of tissue damage
- induced by T cell or pro-inflammatory cytokine mediated immmunopathological responses (71, 72).
- We used this assay to investigate the *in situ* activities of CD317^{neg} Y201 and CD317^{pos} Y202 MSCs.
- 248 Skin samples were obtained with informed consent and approval of the local research ethics

- committee (REC14/NE/1136, NRES Committee North East, IRAS project ID 129780). Following 48
- 250 hours stimulation with IFN-γ or TNF-α (both at 5 ng/ml), Y201 and Y202 MSCs were harvested,
- washed and plated at a density of 1×10^5 cells/well in a 96 well round-bottomed plate. The cells were
- incubated for 3-4 hours to allow for adherence to the plastic. Two punch skin biopsies at 4 mm
- 253 diameter taken from healthy volunteers were dissected into 10-12 sections of equal size. Each section
- was co-cultured with stimulated or unstimulated Y201 or Y202 in duplicate in a 200 μl total volume
- of DMEM supplemented with 20% heat–inactivated pooled human AB serum at 37°C and 5% CO₂.
- 256 Skin sections cultured in the culture medium containing 200 ng/ml IFN-γ or culture media alone were
- used as positive and background controls respectively. After 3-day culture, the skin sections were
- 258 fixed in 10% formalin, then paraffin embedded and sectioned at 5 μm onto microscopic slides. The
- skin sections were stained with haematoxylin and eosin (H&E) following routine protocols. The
- severity of histopathological tissue damage was evaluated by two independent evaluators according
- 261 to the Lerner scoring criteria (73) as follows: grade 0, normal skin; grade I, mild vacuolization of
- 262 epidermal basal cells; grade II, diffuse vacuolization of basal cells with scattered dyskeratotic bodies;
- grade III, subepidermal cleft formation; grade IV, complete epidermal separation(73). Grade II or
- above were considered positive while Grade I changes considered as background, which is observed
- in skin sections cultured in medium alone.

3.14 In vivo assessment of immunomodulatory capacity of hTERT MSC lines in a murine peritonitis model

- 268 To determine the immunomodulatory properties of hTERT MSC lines, an in vivo zymosan-induced
- peritonitis model was used in C57BL/6J mice aged 8-10 weeks as described previously(74, 75).
- 270 These experiments were carried out in accordance with the Animals and Scientific Procedures Act
- 271 1986, under UK Home Office Licence (project licence number PPL PFB579996 approved by the
- 272 University of York Animal Welfare and Ethics Review Board). At day 0, mice were administered
- with an intraperitoneal infusion of 1 mg of zymosan A (Merck) in 100 µl of PBS. Immediately
- 274 following the administration of zymosan, test condition mice were administered an intraperitoneal
- infusion of 2.0x10⁶ cells of either Y201 (CD317^{neg}) or Y202 (CD317^{pos}) in 100 μl of PBS; negative
- 276 control mice were given PBS vehicle only.
- 277 After 24 hours, mice were euthanised using CO₂ overdose and cervical dislocation. Intraperitoneal
- 278 injection of 4 ml of ice cold RPMI-1640 was administered as peritoneal lavage. The process was
- 279 repeated with a second 4 ml RPMI-1640 wash and wash solutions pooled to form the peritoneal
- 280 exudate cells (PEC).

266267

- For each animal tested, red blood cells were lysed using Red Cell Lysis buffer (Merck) and a cell
- count performed. Spleens were retrieved from the mice and cell counts were recorded and a measure
- of spleen cellularity calculated. PEC samples were initially stained for Ly6C (APC), Ly6G (FITC),
- 284 F4/80 (PE-Cy7) CD45 (PerCP-Cy5.5) (BioLegend) and Ly6G (FITC), CD11b (BUV395) and
- SiglecF (BV421) (BD). Both PEC and spleen samples were then stained for TCRb (AF488), CD3
- 286 (APC-Cy7), CD4 (PerCP-Cy5.5), CD62L (APC) and CD44 (PE) (BioLegend). Although at an early
- 287 timepoint, spleen samples were additionally examined for T cell polarisation looking at T effector
- 288 cells CD8 (PerCP-Cy5.5), CD4 (APC), IL4 (AF488), IFN-γ (PE) and IL17a (BV421) (BioLegend)
- and T reg cells using CD8 (PerCP-Cy5.5), CD4 (APC), CD25 (PE) and FOXP3 (AF488)
- 290 (BioLegend). For all tests, Zombie Aqua (BioLegend) was used to exclude dead cells (Table S1).

291 3.15 In vivo assay to assess tissue forming capacity of hTERT MSC lines

- All procedures used were approved by the University of Leeds Ethics Committee and under the UK
- 293 Home Office Project License (PPL:70/8549). The tissue-forming capacity of CD317^{neg} and CD317^{pos}
- 294 hTERT cell lines CD317^{neg} Y201 and CD317^{pos} Y202 was assessed in CD1 nude mice (Charles
- River) aged 8-10 weeks in an *in vivo* transplantation assay(76). 2.0 x 10⁶ MSC cell suspension in 1
- 296 ml medium was added to 40 mg hydroxyapatite (HA) synthetic bone particles (Zimmer Biomet) of
- 297 250-1000 μm size and rotated at approximately 25 rpm at 37°C for 100 minutes to allow cells to
- attach. HA particles were bound using fibrin glue comprising 30 µl thrombin (400 I.U./ml in DMEM
- 299 medium) mixed 1:1 with fibrinogen (115 mg/ml in 0.85% saline solution). Implants were delivered
- 300 subcutaneously into immunocompromised nude mice with two constructs placed into each mouse.
- 301 Transplants were harvested at 3 and 8 weeks, fixed in 4% PFA, decalcified for 7 days in 10% EDTA
- then stored overnight in 70% ethanol prior to paraffin embedding, sectioning and staining with H&E,
- 303 Alcian Blue and Syrius Red (Thermo Fisher).

3.16 Statistical analysis

- Data were tested for equal variance and normality using D'Agostino & Pearson omnibus normality
- 306 test. Differences between groups were compared using two-tailed 1-way ANOVA for parametric data
- or Kruskall-Wallis for non-parametric testing. For two factor analysis, data was analysed with a two-
- tailed 2-way ANOVA. Bonferroni post-hoc testing was conducted to compare between groups. All
- 309 statistical analysis was carried out using IBM SPSS Statistics 24.0, or GraphPad Prism version 5.0-
- 9.0 with P<0.05 deemed statistically significant. Results are annotated as *p<0.05, **p<0.01,
- ***p<0.001 and all averaged values are expressed as mean \pm standard error of the mean (SEM).

312 4 Results

304

313

4.1 MSC identity of CD317-expressing stromal cells

- In our previous work we isolated nullipotent, CD317^{pos} MSC lines (Y102 and Y202) alongside
- differentiation-competent, CD317^{neg} MSC lines (Y101 and Y201) from the same heterogeneous
- donor source suggesting that a subpopulation of stromal cells exists in typical MSC preparations but
- may not contribute to 'classic' MSC functions. Here, we examined the stromal phenotype the
- 318 CD317^{pos} and CD317^{neg} MSC lines. An *in silico* assessment using the Rohart Test(62) was applied to
- accurately discriminate MSCs from fibroblasts, other adult stem/progenitor cell types and
- differentiated stromal cells. This test uses 16 key MSC marker genes as a proven panel of identifiers
- that has independently confirmed MSC status with 97.85% accuracy in 635 cell samples(62). All of
- the immortalised CD317^{neg} and CD317^{pos} stromal cell lines maintained gene expression patterns that
- independently confirmed their MSC status (Figure S1A and Table S3).
- Next, we used mass spectrometry to determine cell surface protein expression profiles across the
- different cell lines. We identified a high number of commonly expressed proteins alongside cell line-
- specific variations. Using a false detection rate of 3%, we found 2338 proteins expressed across all
- MSC lines, with 584 (65.2%) of these commonly expressed (Figure S1B), which may reveal a
- 328 common stromal surfaceome signature (Table S4). Percentage similarity at the surfaceomic level
- ranged from 76.0% to 83.5% (Figure S1C). Unique proteins were identified in Y101 (20 proteins,
- 330 2.2%); Y102 (30 proteins, 3.3%); Y201 (36 proteins, 4.0%); and Y202 (21 proteins, 2.3%). These
- analyses also confirmed that CD317 (BST2) was only identified on Y102 and Y202 MSC lines
- 332 (Table S4). Principle component analysis (PCA) was used to aid interpretation of mass spectrometry
- data through dimensionality reduction. Analysis highlighted that MSC lines clustered distinctly
- within the whole population but were on a similar spectrum of observation, with Y102 and Y202

- lines lying further from the mean of the whole population (Figure S1D). Together, these data
- demonstrate that the CD317^{neg} Y101 and Y201 cell lines, and the CD317^{pos} Y102 and Y202 cell lines
- have broadly similar protein expression profiles in common with other MSC preparations and may be
- used as models for different MSC subtypes.

4.2 Identification of CD317^{dim} and CD317^{bright} populations in primary MSCs

- We previously reported a CD317^{pos} MSC subset with average frequency of 1-3% in low passage
- primary MSCs(60). Here, using flow cytometry analysis with Y201 and Y202 populations gating for
- primary cells as either CD317^{neg} or CD317^{pos}, we were able to demonstrate that CD317 positivity can
- be subdivided into CD317^{dim} and CD317^{bright} populations in primary MSC cultures (Figure 1A, S1E).
- Further examination of n=24 primary MSC populations (passages 1-4) recorded proportions at
- $CD317^{\text{neg}}$ (70.57±5.09%) and $CD317^{\text{pos}}$ (29.77±3.00%), comprising $CD317^{\text{dim}}$ (28.10±4.60%) and
- 346 CD317^{bright} (1.67±0.58%) (Figure 1B). We observed a decrease in CD317 expression in these cells
- over time in culture (passages 1-4), however this trend did not reach statistical significance due to the
- variability of initial proportions of CD317^{pos} cells when CD317^{dim} was included as a CD317 positive
- result (means passage $1 = 50.66\pm27.63\%$, passage $2 = 30.35\pm6.03\%$, passage $3 = 26.07\pm11.78\%$,
- passage $4 = 22.18\pm12.26\%$; n=2,12,7,3) (Figure S1F). We made a similar observation when
- examining subsets of CD317^{dim} and CD317^{bright} cells, with CD317^{bright} cells almost absent by passage
- 4 (Figure 1C). CD317 expression in isolated primary MSCs from passage 3 to 4 reduced by $49.01 \pm$
- 353 11.84% (n=5); with a freeze/thaw cycle at passage 3, this reduction was recorded at $63.94 \pm 3.64\%$ in
- 354 the same cells (n=5) (Figure S1G). Therefore, human primary MSC isolates express CD317 on a
- 355 spectrum that varies from cell to cell and from individual to individual; the overall proportion of
- 356 CD317^{pos} MSCs, as a composite of CD317^{dim} and CD317^{bright}, is 28-29% in heterogeneous MSC
- cultures (combining all analyses of primary cell donors, percent CD317^{pos} MSCs is 28.44±3.82%
- 358 (mean \pm SEM), range of 0.01-93.03%; median=19.89%; n=52). Within CD317^{pos} cells, there was no
- difference in percentage CD317 expression based upon donor gender (mean expression female
- 360 40.02±5.27; male 24.77±6.51; Mann Whitney T-test p=0.051, n=52) or correlation between donor
- age and CD317 expression (mean age: 69.75±1.29 years; range 45-88; Pearson correlation p=0.141,
- n=52),) (Figure 1D, 1E). There was, however, a significant negative correlation between CD317
- expression and BMI (mean 28.06±0.78; range 17-44; Spearman correlation p<0.05, n=52) (Figure
- 364 1F). Y201 cells represent CD317^{bright} subpopulations, so for all subsequent tests using primary donor
- cells, CD317^{pos} represents only CD317^{bright} cells and CD317^{dim} cells were excluded from testing.
- We previously demonstrated that the hTERT immortalised MSC lines display typical (ISCT) surface
- marker profiles(60). Here, we also examined surface markers commonly associated with human
- stromal progenitor cells or subsets, including CD146, CD271 and CD164, within CD317^{neg} and
- 369 CD317^{pos} primary MSC populations. Isolated MSCs from human primary donors showed CD317^{pos}
- 370 (CD317^{dim} and CD317^{bright} populations combined) with mean % expression values of CD317^{pos}
- 371 $(52.90\pm5.89\%)$, CD146^{pos} $(19.46\pm3.07\%)$, CD271^{pos} $(4.025\pm0.71\%)$ and CD164^{pos} $(95.03\pm2.11\%)$
- 372 (n=27) (Figure 1G). Examination of the CD317^{pos} population only showed similar proportions of
- each marker to those seen in the whole population: CD146^{pos} (24.21±3.23%), CD271^{pos}
- $(7.78\pm1.35\%)$ and CD164^{pos} $(97.18\pm0.66\%)$ (n=27) (Figure 1H). These findings demonstrate that
- expression of these markers is independent of CD317 positivity and that CD164 identifies virtually
- all CD317^{neg} and CD317^{pos} MSCs.
- 377 Comparative gene expression analysis has previously demonstrated a correlation between murine
- peri-sinusoidal stromal cells and CD317^{pos} MSCs(77). LEPR has been shown to mark peri-sinusoidal
- stromal cells in mouse tissue(78). Here we investigated CD317^{pos}/LEPR^{pos} stromal cells in mouse

- bone marrow to identify the *in vivo* location of this subpopulation. CD317 expression was detected
- throughout the bone marrow with low frequency colocalisation of CD317 with LEPR restricted to
- peri-sinusoidal regions adjacent to CD31-positive endothelial cells (Figure 11).

383 4.3 Immune profile of CD317^{pos} MSCs

- Our previous transcriptomic data indicated that CD317^{pos} Y102 and Y202 MSC lines display a
- constitutive immunostimulatory expression profile(60), which we sought to define here using the
- MSC lines and primary cells sorted based on CD317 expression. We confirmed by qPCR that
- 387 ICAM1 (CD54) mRNA levels were significantly elevated in CD317^{pos} Y102/Y202 compared to
- 388 CD317^{neg} Y101 (Figure 2A). Although ICAM1 mRNA expression levels appeared similar in primary
- 389 MSCs sorted for CD317 positivity (Figure 2A), flow cytometric analysis demonstrated that cell
- 390 surface ICAM1 expression, as shown by mean fluorescence intensity (MFI), was significantly
- increased on CD317^{pos} primary MSCs versus CD317^{neg} MSCs and CD317^{pos} Y102/Y202 versus
- 392 CD317^{neg} Y101/Y201 (Figure 2B). Comparative analysis of CXCL10 and CXCL11 mRNA levels in
- immortalised MSC lines and primary MSCs sorted for CD317 demonstrated significantly increased
- expression in all CD317-positive MSCs compared to CD317-negative counterparts (n=7;
- 395 experiments performed in triplicate) (Figure 2C, 2D).
- 396 CD317, ICAM-1 and CXCL10 are regulated by interferon-gamma (IFN-γ). We analysed expression
- 397 levels of the IFN-γ receptor by flow cytometry and demonstrated that it was expressed at similar
- levels in all four MSC lines, independent of CD317 expression (MFI, Y101=9.11, Y201=8.41,
- 399 Y102=9.60, Y202=9.84; p>0.05) (Figure S2A). This finding suggested that all MSC lines were
- 400 capable of responding to IFN-γ stimulation in a similar manner, but CD317-positive MSCs may be
- 401 primed to transduce IFN-γ stimulation more effectively. Secretion of CXCL10 was measured in
- 402 immortalised MSC lines with (Figure 2E) and without (Figure 2F) IFN-γ exposure. Under basal,
- 403 unstimulated conditions, CD317^{pos} Y102/Y202 MSCs secrete larger amounts of CXCL10 compared
- 404 to CD317^{neg} Y101/Y201. Following IFN-γ priming, CD317^{pos} MSC lines demonstrate a significantly
- increased ability to secrete additional amounts of CXCL10 compared to CD317^{neg} MSC lines.
- 406 However, IFN-γ has a proportionally much larger stimulatory effect on CXCL10 secretion by
- 407 CD317^{neg} Y101/Y201 cells, suggesting that constitutive interferon signalling is a feature of CD317^{pos}
- 408 MSC lines (Figure 2F).
- 409 Examination of a further panel of eight IFN-γ related genes showed remarkably different expression
- between CD317^{pos} and CD317^{neg} MSCs (Figure 2G, 2H). Using a method described by Raterman *et*
- 411 al(67), we generated an IFN- γ signature score for CD317^{pos} and CD317^{neg} MSCs using the average of
- the log base-2 normalised relative fold changes of the eight IFN-y related genes. We demonstrated
- 413 that CD317^{pos} MSC lines and primary MSCs had a significantly increased IFN-γ signature score
- 414 compared to CD317^{neg} MSCs (Figure 2I & 2J).
- We have previously provided a detailed analysis of trancriptomic data from Y101, Y201, Y102 and
- 416 Y202 MSC lines (60). Here, we examined combined CD317^{neg} and CD317^{pos} datasets and any
- 417 association with human disease conditions. Bioinformatics analysis of differentially expressed genes
- 418 (DEGs) using combined transcriptomic data(60) from CD317^{neg} (Y101 & Y201) and CD317^{pos}
- 419 (Y102 & Y202) MSC lines identified 2340 significantly upregulated genes in CD317^{pos} MSC
- samples (FC>2, p<0.05) with clear clustering of the Y01 group (Y101, Y201) and the Y02 group
- 421 (Y102, Y202) (Figure S2B). The 10 most significantly upregulated genes in the CD317^{pos} group were
- immune-related and/or interferon-regulated, including OAS1, OASL, RSAD2 and CD317 (BST2)
- 423 (Figure S2C). IFN signalling and elevated IFN-signatures are associated with different human

- disease states (79). When comparing the upregulated Y102/Y202 gene sets with six publicly available
- 425 transcriptomic databases for autoimmune and related disorders (Table S5, Table S6), we identified a
- significant association between DEGs and GO terms that were enriched in Y102/Y202 MSC lines
- and psoriasis, eczema and, to a lesser extent, rheumatoid arthritis and osteoporosis (Table S7).
- 428 Similar observations were made when comparing enriched signalling pathways across Y102/Y202
- and disease datasets (Table S8).
- Therefore, a resident MSC subtype can be identified as CD317^{pos}ICAM-1^{hi}CXCL10^{hi} with apparent
- 431 constitutive interferon signalling, which is likely to contribute to specific physiological and
- pathological immune functions.

4.4 Roles of CD317^{pos} and CD317^{neg} MSCs in monocyte and T cell function

- Immunomodulation may be affected through paracrine signalling altering cell recruitment and
- retention in response to signalling molecule expression. The CCL2 receptor, CCR2, is a monocyte
- chemoattractant receptor protein involved in macrophage activation in cells expressing high levels of
- 437 CCL2. Significantly higher CCL2 mRNA expression and protein secretion was detected in CD317
- 438 expressing MSCs versus CD317-negatives (Figure 3A & B).
- In the presence of an antagonist for CCR2, migration of monocytic cells (THP-1) towards
- supernatant from CD317-expressing MSC lines was selectively inhibited compared to CD317-
- 441 negative MSC lines (Y101, Y201 vs Y102, Y202; 19.37±9.57, 19.61±8.89 vs 39.01±6.57,
- 442 41.02±4.79) (Figure S3A). We tested whether the supernatant of CD317^{pos} and CD317^{neg} MSCs
- could induce the migration of both monocytic (THP-1) and T cell (HUT-78) lines in transwell assays.
- We demonstrated that both THP-1 and HUT-78 cells migrated towards MSC supernatants suggesting
- that MSCs secrete both monocyte and T cell chemoattractants (Figure S3B).
- 446 MSCs have previously been shown to suppress activated T cell proliferation whilst maintaining
- inactivated T cell viability in co-culture(80). Several mechanisms are proposed that provide evidence
- 448 for IFN-γ mediated immunosuppression(81), potentially achieving MSC deactivation of T cells
- 449 through IFN-γ receptor targeting or IFN-γ-mediated induction of indoleamine 2,3-dioxygenase (IDO)
- 450 from MSCs, whereby tryptophan is catabolised leading to suppression of T cell proliferation and
- subsequent apoptosis of activated T cells, leaving inactivated T cells in a viable state(82, 83). In this
- work, T cell proliferation was assessed for peaks of gradual division (proliferative index)(70) and
- proliferative cycles (population doublings)(84) over 5 days of co-culture with or without CD317^{pos}
- and CD317^{neg} MSC cell lines (Figure S3C). T cells do not proliferate in culture, unless activated with
- anti-CD3/CD28, and undergo cell death in absence of IL-2, which is produced in vivo by activated T
- 456 cells(85). Compared to T cells alone, all MSC lines and CD317^{neg} primary MSCs significantly
- reduced proliferative index scores, whereas CD317^{pos} primary MSCs had no significant effect on T
- 458 cell proliferative index (Figure 3C, 3D). Assessment of T cell proliferative cycles showed significant
- reductions when cultured with CD317^{neg} Y101/Y201 and CD317^{neg} primary MSCs (Figure 3C, 3E)
- compared to T cells alone. However, CD317^{pos} Y102/Y202 MSCs and CD317^{pos} primary MSCs did
- not significantly reduce the number of proliferative cycles, although a decline was observed (Figure
- 462 3C, 3E). These results demonstrate that CD317^{pos} MSCs are capable of inactivating a proportion of
- proliferating T cells, although this effect is not sufficient to reduce the number of proliferative cycles
- 464 that the residual activated cells achieve, pointing to a diminished immunosuppressive function for
- 465 CD317^{pos} MSCs.
- Next, we determined the effect of CD317^{neg} and CD317^{pos} MSCs on the polarisation of naïve T cells
- into effector lineages with immunosuppressive/anti-inflammatory function. CD317^{pos} MSC lines

- induced a significant increase in the development of pro-inflammatory Th1 cells. Both Y102 (20.32 ±
- 469 0.92%, p<0.001) and Y202 (15.11 \pm 1.46%, p<0.05) increased Th1 polarisation, as indicated by IFN-
- 470 γ expression, in comparison to T cells alone (8.79±2.30%), CD317^{neg} Y101 (9.25±0.42%, p < 0.001
- 471 (Y102)) and Y201 (7.31 \pm 0.60%, p <0.001 (Y102), p <0.01 (Y202)) (One way ANOVA with
- Bonferroni post hoc test). An increase was also observed in Th2 cells for all MSC lines (p>0.05,
- n.s.). Both Th17 and Treg cells, as indicated by IL17a and CD25/FOXP3 expression respectively,
- increased slightly with CD317^{pos} MSC lines, but not statistically significantly. By examining total
- 47.4 Increased slightly with CD31.7 Wise lines, but not statistically significantly. By examining total
- proportions of differentiating cells, it was notable that a large proportion of CD4+ T cells cultured
- alone did not commit to any lineage when compared to co-culture with MSC lines. When proportions
- are summated, only 48.49% of T cells cultured alone differentiated into the 4 lineages examined,
- 478 whilst approximately 75% (Y101), 90% (Y201) and 100% (Y102, Y202) differentiation into these
- lineages was observed when T cells were co-cultured with MSC lines (Figure 3F).

480 **4.5** Pro-inflammatory and Immuno-regulatory potential of CD317^{neg} and CD317^{pos} MSCs in vitro and in vivo

- 482 Considering the stark differences in immune profiles of CD317^{neg} and CD317^{pos} MSCs, we tested
- 483 their effects in different inflammatory models. Prior to in vitro and in vivo testing, we confirmed the
- 484 representative CD317^{neg} and CD317^{pos} MSCs (Y201, Y202) were not affected by viral contamination
- as a potential origin or contributor to constitutive IFN-γ expression. All cell samples were tested in
- 486 triplicate and returned negative results for molecular diagnostics of infectious diseases (Human
- 487 Comprehensive CLEAR Panel, Charles River) using PCR for RNA representing a panel of 26
- 488 virions.
- Initially, we investigated the potential pro-inflammatory property of CD317^{neg} Y201 and CD317^{pos}
- 490 Y202 MSCs in a skin explant model, which is an *in vitro* tool to detect the presence of cutaneous
- 491 tissue damage following a pro-inflammatory insult(86, 87). CD317^{neg} Y201 and CD317^{pos} Y202
- 492 MSCs were primed with IFN- γ or TNF- α and co-cultured *in vitro* with skin explants.
- In this assessment, no tissue damage was observed after skin co-incubation with CD317^{neg} Y201 cells
- in all conditions tested (Figure 4A top panel and Figure 4B left panel). In contrast, cutaneous tissue
- 495 damage was detected when skin was co-cultured with unstimulated or TNF-α stimulated CD317^{pos}
- 496 Y202 cells showing clear cleft formation in the basal layer between the dermis and epidermis (Figure
- 497 4A bottom panel and Figure 4B right panel). When comparing the ability to cause tissue damage,
- 498 Y202 cells caused significantly increased damage compared to Y201 cells in unstimulated and TNF-
- 499 α stimulated conditions (p<0.05) whilst no cutaneous tissue damage was observed when skin was co-
- 500 cultured with IFN-γ stimulated Y202 cells.
- Interferon signalling genes are regulated by interferon in host-pathogen interactions. It is
- 502 hypothesised that constitutive interferon signalling occurs to provide a rapid response to pathogen
- infections through pre-established interferon signature (79), such as that observed here in CD317^{pos}
- MSCs. To investigate the potential for constitutive IFN-γ related signalling on innate immune
- responses *in vivo*, we evaluated immune regulation by CD317^{neg} and CD317^{pos} MSCs in a zymosan-
- induced peritonitis model of acute inflammation that promotes the recruitment of monocytes and
- neutrophils to the peritoneal cavity. Following zymosan treatment, peritoneal exudate cells (PEC)
- were collected by lavage and analysis performed on the cell content. A gating strategy was devised
- for flow cytometric analysis of multiple PEC cell types focusing on haematopoietic, myeloid and
- 510 lymphoid cells including monocytes, macrophages and T cells (Figure S4A & S4B). Treatment with
- either Y201 or Y202 MSC lines suppressed the recruitment of inflammation-related cells to the area.

- There was a significant reduction in total cells recruited in both Y201 (3.552±1.543 x 10⁶) and Y202
- 513 (2.076±0.421 x 10⁶) treated conditions compared to zymosan-induced peritonitis without treatment
- 514 (9.686±1.894 x 10⁶) (p<0.05), with no significant difference between MSC-treated animal PEC
- numbers and PBS controls $(4.420\pm1.790 \times 10^5)$ (Figure 4C).
- 516 Examination of the composition of PEC showed that zymosan-induced peritonitis prompted a
- significant increase in haematopoietic cells (p<0.05). No difference in recruitment of eosinophils or
- 518 neutrophils was observed in MSC-treated mice when compared to zymosan alone or PBS controls
- 519 (Figure S4C & S4D). Examination of the production of monocytes and macrophages in PEC samples
- showed no differences in monocyte recruitment, however both zymosan alone and zymosan plus
- Y202 showed significant increases in macrophage proportions compared to PBS controls (p<0.001,
- 522 p<0.05 respectively) whilst Y201 treatment suppressed macrophage numbers (p<0.05) (Figure 4D).
- Within these monocyte and macrophage populations, the proportions of Ly6C positive and negative
- 524 cells matched the proportions seen in zymosan treatment only animals (Figure S4F & S4G). Ly6C
- 525 positive monocytes and macrophages are linked with pro-inflammatory responses by CCR2/CCL2
- mediated homing to sites of tissue injury, whilst Ly6C low or negative monocytes and macrophages
- are reparative, guided by VCAM-1 and other adhesion proteins(88, 89).
- 528 Spleens retrieved from MSC-treated and control mice were homogenised and analysed for naïve and
- 529 polarised T cells, and memory T cells. No differences were found in the mass or cellularity of spleens
- between controls and MSC-treated animals (data not shown). When tested, a significant increase was
- found in activated CD4+ central memory T cells (TcM) in CD317^{neg} Y201 cell treated conditions
- 532 (14.23±0.06%) in comparison to PBS controls (4.53±0.18%) or Y202 treated animals (5.89±4.30)
- 533 (Figure 4E). CD4+ effector T cell polarisation was not altered by introduction of zymosan or MSC
- treatments within the 24 hour time period measured. However, treatment with either CD317^{neg} Y201
- 535 (1.51 \pm 0.57%) or CD317^{pos} Y202 (0.84 \pm 0.25%) MSCs suppressed CD8a/b+ expression
- representative of cytotoxic T cell production in mice in comparison to CD8a/b+ expression in
- untreated animals $(5.42 \pm 1.10\%)$ (Figure 4F).

538 **4.6** In vivo tissue formation is enhanced in CD317^{neg} MSC lines when compared to CD317^{pos} subpopulations

- We hypothesised that the immunomodulatory enhancements observed in CD317-positive MSCs
- would impact on their tissue-forming capacity. To test this hypothesis, CD317^{neg} (Y201) and
- 542 CD317^{pos} (Y202) MSC lines were loaded onto hydroxyapatite (HA) scaffolds and implanted
- subcutaneously in nude mice. Scaffolds were retrieved at 3 and 8 weeks post-implantation and
- examined using histological analysis for *de novo* tissue formation by deposition of extracellular
- matrix (ECM), collagen and neoangiogenesis.
- 546 CD317^{neg} Y201 MSCs showed clearly advanced ECM and collagen deposition in histological stains
- using Sirius Red for collagen formation and Alcian Blue for proteoglycan synthesis (Figure 5A, 5B
- 548 & 5C), suggestive of a more stable capacity for tissue formation. Haematoxylin and eosin staining
- showed evidence of tissue formation from 3 weeks post implantation in CD317^{neg} MSCs alongside
- evidence at 8 week timepoints of capillary tube structures containing blood cells indicative of
- neoangiogenesis (Figure 5D). Although there was some evidence of tissue formation in CD317^{pos}
- Y202-loaded scaffolds, the tissue formed appeared less continuous or cohesive compared to
- 553 CD317^{neg} Y201 samples and by 8 weeks post-implantation there was clear evidence of
- disaggregation and cleft formation at the surface of HA particle clusters following histological

- 555 staining for ECM formation (Alcian Blue and Sirius Red) with no detectable vessel formation (Figure
- 556 5A, 5B, 5C & 5D).

Discussion 4.7

- 558 This study investigated the characteristics and properties of a CD317^{pos} subpopulation within
- 559 heterogeneous MSCs and their ability to contribute to immune responses and tissue repair. We used
- 560 immortalised MSC model lines and primary MSCs isolates to elucidate the biology and potential
- impact on the therapeutic application of these cells. Here, we confirm CD317^{pos} MSCs represent a 561
- 562 subpopulation of cells commonly found in human MSCs preparations with an equal distribution in a
- 563 range of demographic groups and health conditions. We found that rare CD317-expressing cells
- 564 colocalised with LEPR-positivity adjacent to endothelial cells at marrow sinusoids, and in vivo
- 565 location consistent with other bone marrow stromal cell preparations (78, 90). CD317pos MSCs may
- therefore interact with and be regulated by endothelial cells in a perivascular niche, similar to those 566
- 567 described for other stem and progenitor cell types (91, 92), but further investigation is required. Using
- in vitro and in vivo functional assays, we demonstrate that CD317^{pos} MSCs have reduced 568
- 569 immunomodulatory and tissue-forming capacity compared to CD317^{neg} MSCs, suggesting that
- 570 CD317^{pos} cells will not contribute to tissue repair or *de novo* tissue formation. Any contribution of
- 571 CD317^{pos} cells in therapy, when delivered within an undefined heterogeneous MSC culture, is
- 572 therefore likely to be through immunomodulatory influence, and the contribution to the regenerative
- 573 process is dependent upon the therapeutic target and the inflammatory environment present in the
- 574 recipient at the time of transplantation. Given the potential for CD317^{pos} MSCs to respond to the
- inflammatory environment in vivo, these cells may serve a positive function in assisting the repair of 575
- damaged tissues by CD317^{neg} MSCs when transplanted as part of a heterogeneous population.
- 576
- 577 However, our in vivo results demonstrate that CD317^{neg} cells are capable of inducing both anti-
- inflammatory immunomodulation and tissue regeneration in the absence of CD317^{pos} counterparts, 578
- 579 suggesting the support function is not vital to successful repair of damaged tissue by CD317^{neg} MSCs
- 580 alone. Of note, when supplied in sufficient numbers CD317^{pos} MSCs are capable of causing tissue
- 581 damage, as observed in our skin explant model, which may be linked to their distinctive immune
- profile and functional differences to CD317^{neg} MSCs. Qualitative histological analysis of tissue 582
- 583 generation was not subjected to quantification. Further work should be done to fully evaluate the
- extent and quality of tissue repair formed using both CD317^{pos} and CD317^{neg} MSC lines and primary 584
- donor cells to enable more firm conclusions to be drawn. 585
- 586 Inflammation serves a dual role in tissue repair. Cells in the immune response, such as neutrophils,
- 587 function to initiate the repair process. Neutrophils cause tissue breakdown during inflammation but in
- 588 the absence of neutrophils, macrophages rapidly recruited to the site of injury will display reduced
- 589 rate of tissue regeneration owing to the presence of cell debris normally phagocytosed by
- 590 neutrophils(93). Our results from MSC treatment of zymosan-induced peritonitis in mice showed a
- 591 neutrophil population present in PEC suspensions from PBS injected mice, and significantly
- 592 increased neutrophils present in the PEC of both zymosan-only and MSC-treated mice. However,
- 593 examination of subsequent macrophage populations showed that whilst no macrophages were
- detected in the PBS control mice, both zymosan-only and CD317^{pos} MSC plus zymosan conditions 594
- 595 displayed significant increases in macrophage numbers. Significantly fewer cells, including
- 596 macrophages, were recruited in the presence of CD317^{neg} MSCs compared to zymosan only
- 597 induction, therefore CD317^{pos} MSCs fail to inhibit macrophage recruitment.
- The influence of CD317^{pos} MSCs on T cells appears to be highly modulated in comparison to 598
- 599 CD317^{neg} MSCs. MSCs have been widely shown to deactivate T cells *in vitro* and suppress T cell

600 proliferation whilst directing CD4+ effector T cells from Th1 to Th2 profile(80, 94-101). However, 601 in activated T cells in cell to cell contact with CD317^{pos} MSCs, we observed minimal deactivation of 602 T cells and continued T cell proliferation, in conjunction with an active increase in Th1 polarisation, 603 contrary to the widely accepted immunosuppressive properties of MSCs. IFN-y stimulation of MSCs 604 has been shown to induce activation through upregulation of HLA class II, pushing the MSC towards 605 antigen-presenting capability for immune regulation, promoting T cell interactions and potentially 606 influencing CD8+ T cell activation(102). This may go towards explaining the results we observe 607 when CD317^{pos} cells interact with T cells in vitro and T and B cells in vivo. CD317^{pos} MSCs show minimal interaction with T cells in vitro, yet function more effectively in a pro-inflammatory in vivo 608 609 environment. CD317 promotes an immune response through stimulating activation of NFκB(103) 610 which in turn contributes to B cell development (104). MSC immunomodulation is intrinsically tied to 611 interactions with dendritic cells (DCs), with MSCs inhibiting DC maturation, resulting in reduced 612 migration, cytokine secretion, antigen presentation to T helper cells and cross-presentation to 613 cytotoxic T cells(105) through interrupting entry into the cell cycle, inhibiting DC differentiation and 614 function (106). DCs also mediate the MSC immunosuppressive effect through the induction of 615 regulatory T cells(107, 108).

- Deeper analysis of the CD317^{pos} subset of MSCs identified a heightened interferon signature that was
- not related to IFN-γ receptor expression levels, suggestive of constitutive IFN signalling. Pre-
- established, low level constitutive IFN signalling contributes to rapid pathogen responses in the
- innate immune system and conveys a protective effect to de novo IFN exposure in these cells(109).
- 620 CD317^{pos} MSCs, if maintained at appropriate levels, may therefore contribute to enhanced innate
- immunomodulation. Of interest, CD317^{pos} MSCs may also serve as a useful tool in the investigation
- of host tropism in viral infection, a particularly prevalent issue with the advent of COVID-19.
- Indeed, the presence of BST2/CD317 on the cell surface has been shown to convey a protective
- effect by tethering coronavirus virions to the cell surface or intracellular membranes and decreasing
- budding of progeny virus(110). These cells may therefore provide an enhanced response to viral
- 626 infection that facilitates tissue regeneration as well as immunomodulation. However, whilst
- 627 constitutive IFN signalling may convey a protective effect to cells experiencing de novo IFN in the in
- 628 vivo environment, there also exists the potential for a link between unregulated constitutive IFN
- signalling and tissue damage in human disease conditions including autoimmunity. It is therefore
- highly significant that we show the baseline gene expression levels of CD317^{pos} MSCs aligns them
- with cells present in autoimmune and related conditions.
- In this report we characterise a subset of human MSCs that favour immunomodulatory interactions
- over tissue regeneration, yet identify as MSCs through both independent tests (e.g. Rohart) and ISCT
- 634 guidelines(111). These cells display a distinct immune profile and operate in contrast to the
- expectations of MSC's immunosuppressive function. Further *in vivo* investigation is necessary to
- elucidate the probability of pro-inflammatory outcomes when using CD317^{pos} MSCs as a therapeutic.
- We have demonstrated that the proportion of CD317^{pos} MSCs varies considerably between donor
- MSC preparations, which could reflect individual inflammatory state and/or infection history. We
- propose that the success of therapeutic applications for tissue regeneration may be associated with the
- numbers of CD317^{pos} MSCs present in the administered cell dose. There is also the possibility that
- 641 CD317^{pos} MSCs can bring therapeutic benefits in the inflamed environment. The expression of
- 642 CD317 on MSCs serves as a positive marker for cells that display all the characteristics of an
- immune stromal cell and targeted therapies should aim to harness the knowledge of this cell type as
- novel approaches to the treatment of degenerative, and inflammatory conditions.

5 Conflict of Interest

There are no conflicts of interest with respect to this work.

6 Author Contributions

- AGK designed, performed and analysed T cell experiments. AGK and JPH designed, performed and
- analysed peritonitis experiments. AS designed, performed and analysed MSC localisation
- experiments. JMF, SR and SJ designed, performed and analysed ELISA, Interferon signature, Rohart
- 651 testing, cell migration experiments and bioinformatics. XY and EK performed subcutaneous HA
- scaffold implantation in vivo whilst AGK performed the associated cell culture and analysis of
- explants. PG designed experiments and was responsible for conceptualisation, funding acquisition,
- supervision and writing (review and editing). XW designed, performed and analysed the *in vitro* skin
- explant model. AK, JMF and PG wrote the paper.

656 7 Funding

647

- This work was funded by the Biotechnology and Biological Sciences Research Council (BBSRC)
- United Kingdom, Doctoral Training Partnership grant (BB/M011151/1) and the Tissue Engineering
- and Regenerative Therapies Centre Versus Arthritis (21156). LK and XY are partially funded by
- 660 funding by the 'EPSRC CDT in Tissue Engineering and Regenerative Medicine' at the University of
- Leeds (Grant number EP/L014823/1). The York Centre of Excellence in Mass Spectrometry was
- created thanks to a major capital investment through Science City York, supported by Yorkshire
- Forward with funds from the Northern Way Initiative, and subsequent support from EPSRC
- 664 (EP/K039660/1; EP/M028127/1).that aided the efforts of the authors.

665 8 Acknowledgments

- The authors thank the staff and patients of Clifton Park Hospital for samples. We are grateful to the
- University of York Technology Facility for support with flow cytometry, cell sorting, confocal
- microscopy bioinformatics and proteomic analysis. We also thank the York Centre of Excellence in
- Mass Spectrometry which was created thanks to a major capital investment through Science City
- York, supported by Yorkshire Forward with funds from the Northern Way Initiative, and subsequent
- support from EPSRC (EP/K039660/1; EP/M028127/1). We thank Emily Taylor for assistance in
- isolating primary T cells.

673 **9 Data Availability Statement**

- The raw data supporting the conclusions of this article will be made available by the authors, without
- undue reservation.

676

10 Contribution to the Field

- Mesenchymal stromal cells (MSCs) are the most widely studied cell type in clinical trials for
- musculoskeletal diseases, but outcomes at clinical trials are variable due to the heterogeneity in
- stromal cell populations. We have identified consistent subpopulations of MSCs occurring within
- 680 human bone marrow-derived MSCs with varied tissue regenerative and immunomodulatory
- properties. Here we define a new immune stromal cell with previously unidentified immune and non-
- regenerative characteristics based on in vitro and in vivo evidence. CD317-positive cells are present
- at variable levels in most MSC preparations currently used in research and therapy. The presence of
- 684 CD317-positive MSCs may impact upon outcomes of clinical trials and will influence interpretation

- of clinical and research data describing outcomes using heterogeneous cell populations. We believe
- these cells contribute to variability in trial outcomes and may therefore impede clinical translation of
- novel therapies in regenerative medicine.

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992 Figure Legends

- 993 **Figure 1** Analysis of CD317-expressing MSC populations within primary cell isolates. **(A)** The
- CD317 expressing populations can be divided into CD317^{bright} and CD317^{dim} with CD317^{bright} MSCs.
- 995 **(B)** Average proportions of CD317^{neg} and CD317^{pos}, comprising CD317^{dim} and CD317^{bright}, in
- primary MSCs lines. (C) Expression of CD317 over early passages 1 to 4 in Primary MSCs with
- 997 CD317^{neg} increasing, CD317^{dim} and CD317^{bright} decreasing during in vitro culture (n=2-12). Variation
- of CD317 expression with gender (**D**), age (**E**) and BMI (**F**) in primary donors (n=52). (**G**) Isolated
- MSCs from human primary donors showed CD317^{pos} (CD317^{dim} and CD317^{bright} combined) with
- mean values of CD317^{pos}, CD146^{pos}, CD271^{pos} and CD164^{pos} (n=27). (H) Examination of the
- 1001 CD317^{pos} population only, showed similar proportions of each marker to those seen in the whole
- population only, showed similar proportions of each market to those seen in the whole
- population (n=27). (I) CD317 expression was detected throughout the bone marrow of mice with low
- frequency colocalization of CD317 and LEPR+ in peri-sinusoidal regions (arrows).
- Figure 2 Examination of the immune profile of CD317pos MSCs. (A) Comparative mRNA
- expression of ICAM-1 in MSC lines and primary cells sorted by CD317 expression (RNA was
- extracted from 3 different donors or 5 cell line passages; qPCR performed in triplicate, mean shown
- ± SEM). (B) Mean fluorescence intensity of ICAM-1 expression on the cell surface of MSC lines and
- primary MSCs differentially gated by CD317 staining (MSCs from 5 different donors or 4 different
- passages of MSC lines were stained for flow cytometry, mean shown \pm SEM). (C)/(D) Comparative
- 1010 (mean ± SEM) mRNA expression of CXCL10 (red) and CXCL11 (blue) in MSC lines/ primary
- MSCs sorted for CD317 expression (RNA was extracted from 7 different donors/7 different cell
- passages; experiments were performed in triplicate). (E/F) CXCL10 secretion by MSC lines prior to

- 1013 IFN- γ priming and after priming with baseline (unprimed) secretion subtracted (mean \pm SEM, n=2).
- 1014 (**G/H**) Comparative mRNA expression of 8 IFN-γ signature genes in MSC lines/primary MSCs
- sorted by CD317 expression (RNA was extracted from 5 different donors/5 different cell passages;
- experiments were performed in triplicate, mean shown \pm SEM). (I)/(J) IFN- γ score for MSC
- lines/primary MSCs sorted by CD317 expression (n=5)*/** = significance at P<0.05/0.01 using an
- appropriate statistical test.
- 1019
- Figure 3 Influence of CD317^{neg} MSCs and of CD317^{pos} MSCs on immune cell function (A)
- 1021 Comparative mRNA expression of CCL2 in primary MSCs sorted by CD317 expression (RNA was
- extracted from 7 different donors; experiments performed in triplicate, mean shown \pm SEM). (B)
- 1023 CCL2 secretion in primary MSCs sorted by CD317 expression and MSC lines (from 4 different
- donors/4 different cell line passages; experiments performed in triplicate, mean shown \pm SEM). (C)
- In vitro co-culture of hTERT immortalised lines Y201 and Y202 and primary CD317^{neg} and
- 1026 CD317^{pos} cells with activated T cells. CD317^{neg} cells reduce proportion of proliferating T cells and
- number of cell cycles achieved (**D**) hTERT cell lines significantly reduce proportion of proliferating
- 1028 cells as demonstrated through proliferative index (E) CD317^{neg} cell lines reduce proliferative cycles
- achieved by activated T cells in comparison to CD317^{pos} or T cell alone controls. (**F**) assessment of
- the influence of MSC on T cell polarisation in co-culture demonstrates CD317^{pos} cells influence
- activated T cells to preferentially polarise towards IFN-γ expressing (Th1) subset with indications of
- increased IL17a+ and CD25+FOXP3+ expressing cells.
- Figure 4 In vitro and in vivo immunomodulation by CD317^{neg} Y201 or CD317^{pos} Y202 MSCs. (A)
- Representative images of skin explants independently assessed for damage to tissues, examining
- keratinocytes, basal cells, keratotic bodies, the appearance of sub-epidermal clefts at the junction
- with the dermis and in highly damaged tissue the appearance of complete epidermal separation
- following treatment with MSCs primed with IFN- γ or TNF- α and co-cultured in vitro. (B) Y201 co-
- culture did not prompt damage to the tissue in any conditions whilst Y202 cell line demonstrated
- marked tissue damage in untreated cells and TNF-α treated cell lines. Both Y201 and Y202 cell lines
- retained the ability to inhibit tissue damage when primed with IFN-γ. (C) MSCs subsequently
- applied to an *in vivo* peritonitis model of inflammation showed immunomodulation through reduced
- immune cell recruitment, (**D**) reduced macrophage development following Y201 treatment, (**E**)
- increased central memory T cell development following Y201 treatment and (F) reduced CD8+
- 1044 cytotoxic T cell development following Y202 treatment. n=3, *p<0.05, **p<0.01, ***p<0.001
- Figure 5 In vivo tissue generation in HA scaffolds loaded with CD317^{neg} Y201 or CD317^{pos} Y202
- 1046 MSCs. (A, B) Histological staining of recovered implants using Sirius Red for collagen formation
- and (C) Alcian Blue for proteoglycan synthesis at 3 and 8 weeks post-implantation in HA scaffolds
- loaded with either CD317^{neg} Y201 MSCs and CD317^{pos} Y202 MSCs. (**D**) Haematoxylin and eosin
- staining comparting tissue and blood vessel formation at 3 and 8 weeks post-implantation in HA
- scaffolds loaded with CD317^{neg} Y201 MSCs and CD317^{pos} Y202 MSCs. Scale bars = 250µm (Part A
- Scale bars = 500µm). Asterisks = HA particles, arrows = blood vessels.