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Effects of degradable magnesium on paracrine signaling between human umbilical cord perivascular cells and peripheral blood mononuclear cells†

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Human mesenchymal stem cells (MSC) interact with numerous immune cells that can promote regenerative processes and inhibit inflammatory responses. We hypothesised that the cross-talk between human umbilical cord perivascular cells (HUCPV; an alternative source of MSC) and peripheral blood mononuclear cells (PBMC) could be influenced by degradable transwell magnesium (Mg). To study the correlations between paracrine signaling and specific cellular behaviour during the host response to Mg, we used a transwell coculture system for up to 7 days. The proliferation and viability of both cell types were not significantly influenced by Mg. When HUCPV were cultured with degradable Mg, a moderate inflammation (e.g., lower secretions of pro-inflammatory interleukin 1 beta and IL2, and tumour necrosis factor alpha, interferon gamma, anti-inflammatory interleukins 4, 5, 10, 13, and 1 receptor antagonists and granulocyte colony stimulating factor), and an increased pro-healing M2 macrophage phenotype were observed. Moreover, when PBMC were cultured with degradable Mg, the expression of migration/wound healing related cytokines (interleukin 8, granulocyte-macrophage colony-stimulating factor, monocyte chemoattractant protein 1 and macrophage inflammatory protein 1 α/β) was upregulated, accompanied by an increase in the migration ability of HUCPV (cell scratch assay). In addition, an increased pro-osteogenic potential was demonstrated via an increase of osteoblastic markers (e.g., alkaline phosphatase activity, specific gene expression and cytokine release). These results collectively imply that Mg possesses osteo-immunomodulatory properties. They also help to design Mg-based bone substitute biomaterials capable of exhibiting desired immune reactions and good clinical performance.

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1. Introduction

Magnesium (Mg) and its alloys have been investigated as degradable implants for cardiovascular,^{1,2} orthopaedic^{3–5} and bone regenerative applications^{6,7} because of their desirable bioactive properties, such as stimulated osteoblast adherence⁸ and formation of cartilage.⁹ Usually, immune responses to a biomaterial comprise non-specific inflammation.¹⁰ Peripheral blood mononuclear cells (PBMC), comprising lymphocytes (T cells, B cells, and natural killer cells; 70–90%), monocytes (10–20%) and dendritic cells (1–2%), have been widely used as immune cells to evaluate the *in vitro* inflammatory reaction to biomaterials.^{11,12} In addition, PBMC are able to secrete a

broad range of cytokines in non-specific inflammatory responses, including pro-inflammatory interleukins 1 beta (IL1 β), IL2, tumour necrosis factor alpha (TNF α), and interferon gamma (IFN γ),¹³ as well as anti-inflammatory cytokines, such as IL10.¹⁴ As an *in vitro* model to follow the initial immune reaction after biomaterial implantation, the secretion of cytokines by PBMC could be very informative. The interactions between the immune system and biomaterials rely on the tissue surrounding the implant, which will induce the tissue-specific innate immune response and the subsequent adaptive immune response.¹⁵ After biomaterial implantation, when the blood-biomaterial come into contact, immediately a layer of proteins from the surrounding microenvironment could be adsorbed onto the material surface, forming a surface matrix and attracting specific cells from the innate immune system (natural killer cells, dendritic cells, mast cells, and granulocytes (neutrophils, basophils and eosinophils), and macrophages) to the damaged sites in short time. These immune cells secrete cytokines and chemokines that are

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capable of recruiting tissue repair cells, such as mesenchymal stem cells (MSC) *etc.*, into the injured sites. Normally, the occurring immune response is accompanied by tissue repair reactions in the following 2–10 days.¹⁶ Although a variety of cell types are involved in tissue repair, macrophages, a small group of PBMC, play a key role in the initial immune response by secreting cytokines and chemokines that directly affect tissue repair.¹⁷ As a source of MSC, the human umbilical cord perivascular cells (HUCPV) were selected. HUCPV exhibit a low immunogenicity^{18,19} and potent immunomodulatory effects,²⁰ and a natural primary potential to differentiate into an osteogenic phenotype.^{21–23} The cell-to-cell communication between immune cells and MSC could regulate the immune reaction *via* paracrine signaling, in which a cell could secrete the factors to influence the behaviour and functions of nearby cells.²⁴ Therefore, in the current study, the molecule releases were measured after 1, 4, and 7 days to follow the initial immune reaction to the degradable Mg material.

Promoting tissue regeneration by modulating the immune system using biomaterials is one of the strategies followed in regenerative medicine and in smart biomaterial development. Thus, as immune responses cannot be avoided and are necessary in the early stages of bone implantation, a moderate inflammation and an increase of the pro-healing M2 macrophage phenotype around the implants may be more efficient to promote fracture healing and lasting stability of bone implants.^{25–28} In recent years, numerous reports have proposed that Mg and its alloys play a role in the modulation of immune systems.^{29,30} For instance, *in vitro* studies have demonstrated the direct role of Mg in decreasing the pro-inflammatory cytokine production,³¹ as well as mediating the anti-inflammatory effects.³² The anti-inflammatory effects of Mg salts³³ or nanoparticles³⁴ were confirmed *in vivo*. To date, research studies have confirmed that a faster inflammation resolution stimulated by Mg and Mg-based materials could be desirable to convert the macrophages from the M0 to the pro-healing M2 phenotype *in vitro*.^{35,36} In addition, it has become clear that mesenchymal stem cells (MSC) also possess immunomodulatory properties that can modulate several immune cells,³⁷ especially macrophages.^{38,39} For instance, coculture of MSC was able to induce macrophages to adopt an enhanced regulatory (M2) phenotype *via* increased levels of IL4, IL10, and IL13, and reduced concentrations of TNF α , IFN γ and IL2.⁴⁰ Increasing numbers of clinical tests show that the success of MSC-based immunomodulation relies on the assessment of molecule secretion and their interaction with immune cells.⁴¹ Also, MSC can influence the various pathways of immune responses in a paracrine manner.^{42,43} Furthermore, a recent study about Mg ions (magnesium sulphate) has indicated that Mg could modulate some immunoregulatory properties of MSC.⁴⁴ However little is reported about the role of degradable Mg in the immunomodulatory properties of MSC in a biomaterial scenario. In particular, the effects of degradable Mg on paracrine signaling between MSC and immune cells require further examination.

Indeed, the communication of MSC-immune cells is not a one-way street. Immune cells have the potential to attract stem

cells to the damaged tissue and several mediators have been implicated in the migration of MSC. Among these factors, monocyte chemoattractant protein 1 (MCP-1 also known as C-C motif chemokine ligand 2 or CCL2), macrophage inflammatory protein (MIP-1 α /CCL3), IL-8 and TNF α ^{45,46} can be listed. In addition, macrophage polarisation and release of cytokines and other substances influence tissue healing and remodeling.⁴⁷ These macrophages and their polarised morphological variants always stay at the material-tissue interface *in vivo*. This cell-cell fusion also contributes to the degradation of biomaterials *via* their phagocytosis and released mediators.^{48,49} The released mediators also play crucial roles in MSC differentiation. For instance, pro-inflammatory cytokines (TNF α and IL1 β) inhibited the osteogenic differentiation of stem cells.⁵⁰ Soluble factors, *e.g.*, granulocyte-macrophage colony-stimulating factor (GM-CSF), enhanced osteogenic differentiation in Saos-2 cells⁵¹ or stem cells.⁵² The secretions of basic fibroblast growth factor (bFGF), vascular endothelial growth factor (VEGF), and IL12 could also be markers of osteoblastic differentiation.²³ In addition, numerous reports over recent years have demonstrated that Mg ions can stimulate osteogenic differentiation of mesenchymal stem cells (MSC) and bone regeneration *in vitro*^{53,54} and Mg-based alloys can promote bone formation *in vivo*.^{6,55} Thus, the influence of degradable Mg on immune cell-mediated MSC behaviour (*e.g.*, migration/wound healing and osteogenic differentiation), notably the paracrine signaling, needs further examination.

To answer the above questions (influences of degradable Mg on paracrine signaling between immune cells and MSC) in an interactive microenvironment, a two-way communication, transwell (TW) system, was used.⁵⁶ By employing the TW system, HUCPV and PBMC (and especially their derived macrophages; later referred to as PBMC/macrophages in this manuscript) were grown separately while sharing the same microenvironment and soluble factors. Thus, the contribution of each cell type can be investigated. Initially, the current study focused on the role of Mg degradation in paracrine secretions in relation to inflammation, as well as the M2 macrophage phenotype. Besides, the secreted cytokines, which are responsible for MSC behaviour (recruitment and osteogenic differentiation), were determined. Several involved aspects, wound healing assay and pro-osteogenic activity of HUCPV, were also examined.

2. Experimental

2.1. Cell isolation and culture

HUCPV were derived from the perivascular site (Wharton's jelly) of umbilical cords. Umbilical cord samples were obtained from Asklepios Klinik Altona (Hamburg, Germany) immediately after caesarean sections of consenting donors. The HUCPV isolations were approved by the Ethics Committee of the Hamburg Medical Association (PV4058) and were performed as previously described.⁵⁷ The purity of the isolated HUCPV was validated *via* immunophenotyping by flow cytome-



try (S3e; Bio-Rad Laboratories, Munich, Germany) presented in the ESI (Fig. S1a†). The immunophenotype of HUCPV was characterised using CD105 (SH2) and CD90 (Thy-1) as mesenchymal cell positive and specific markers,²² and CD54⁵⁸ and CD31⁵⁹ to exclude the contamination of fibroblast and endothelial cells, respectively. Additionally, the HUCPV morphology (Fig. S1b†) was evaluated *via* live/dead staining with calcein AM (living cells, green) and ethidium homodimer-1 (dead cells, red; 1 and 2 μM , respectively; Thermo Fisher Scientific, Schwerte, Germany) using an inverted fluorescence microscope (Eclipse Ti, Nikon GmbH, Düsseldorf, Germany). The isolated HUCPV were expanded in an α -minimum essential medium (α -MEM; Invitrogen, Fisher Scientific GmbH, Schwerte, Germany) containing 15% (v/v) foetal bovine serum for human mesenchymal stem cells (FBS; Biological Industries, Beit-Haemek, Israel) and 1% (v/v) penicillin/streptomycin (P/S; Life Sciences, Karlsruhe, Germany) under physiological conditions (5% CO_2 , 20% O_2 , 95% relative humidity, and 37 °C). The medium was changed every 2–3 days and the cells were cultured until approximately 80% confluence was reached. HUCPV at 5–8 passages were used for the biological tests.

Human leukocyte-enriched blood samples were provided by the University Hospital Hamburg-Eppendorf (UKE; Hamburg, Germany), and then PBMC were isolated using the Ficoll Paque 400 (GE Healthcare, Chicago, Illinois, USA) from the buffy coat as described by the former protocol.⁶⁰ Meanwhile, the human plasma (above the buffy coat) was collected, centrifuged at 300g for 10 min, passed through a 0.45 μm sterile filter (Thermo Fisher Scientific, Schwerte, Germany) and frozen at -20 °C for further use. The isolated PBMC subpopulations used in this study are shown in Fig. S2 (ESI†).

Here we used pooled allogeneic human plasma instead of FBS consistent with other research studies.^{61,62} Because of the species difference of biological response modifiers (BRM) between humans and bovines, using autologous human plasma is a more appropriate way to test the effects of biomaterials on human PBMC/macrophages and HUCPV.⁶³ Additionally, compared to human plasma, FBS might contain higher background of non-specific cell stimulatory factors.^{64,65} Importantly, to avoid a very high level of variability between human serum or plasma lots,⁶⁶ we used pooled human plasma throughout our project. Therefore, the α -MEM medium was supplemented with 10% human plasma (v/v) and 1% P/S and used as the cell culture medium in our system.

2.2. Material preparation

Biomaterials were produced at Helmholtz-Zentrum Geesthacht, Geesthacht, Germany. High-purity Mg (99.93%) was prepared by permanent mould gravity casting, followed by a T4 treatment of the produced ingots, and extrusion into rods of 10 mm diameter. Finally, after being machined to obtain a diameter of 9 mm, Mg discs were cut with 1.5 mm thickness and each disc weight was about 0.2 g. After cleaning *via* sonication for 20 min in 100% *n*-hexane, 100% acetone and 100% ethanol as well as sterilisation in 70% ethanol (Merck KGaA, Darmstadt, Germany), each disc was preincubated for

24 hours in 500 μL of cell culture medium (the same as above) prior to use.

2.3. TW systems

In order to elucidate how the interaction between PBMC/macrophages and HUCPV is influenced by Mg biomaterial *in vitro*, a transwell (5 μm pore size; Alvetex™ Strata, Glasgow, United Kingdom) was used (Fig. 1a). An agarose coating prevents cell adhesion and survival on a tissue culture plate. In an agarose-coated 12-well plate (Sigma-Aldrich Chemie GmbH, Taufkirchen, Germany), HUCPV were seeded at a density of 2×10^5 cells (25 μL) on preincubated Mg discs for 30 min (lower compartment or well bottom), and then 1.8 mL of fresh cell culture medium was added. Afterwards, PBMC were seeded at a density of 5×10^5 cells in 0.2 mL of culture medium in the upper compartment. Single culture of HUCPV and PBMC and coculture without Mg disc were selected as controls. To detect the possible artefact or background caused by the cell culture medium and Mg disc, the pure cell culture medium and supernatant from Mg discs without cells were regarded as blank and no-cell controls, respectively (all experimental conditions and their abbreviations are presented in Table 1). In the context of

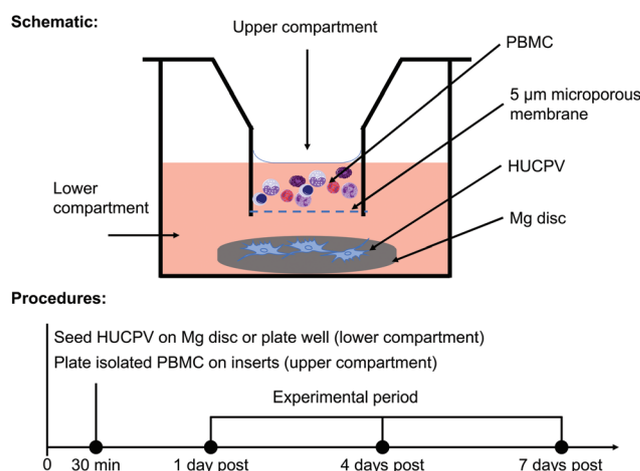


Fig. 1 *In vitro* TW coculture system.

Table 1 Settings and abbreviations of *in vitro* coculture system conditions

	No.	Abbreviation	Experimental condition
TW system	1	Blank	Cell culture medium
	2	Mg	Only Mg disc
	3	H	HUCPV alone
	4	H + Mg	HUCPV on Mg disc
	5	P	PBMC alone
	6	P + Mg	PBMC (upper compartment) with Mg disc (lower compartment)
	7	H + P	Coculture of HUCPV and PBMC
	8	H + P + Mg	Coculture of HUCPV on Mg disc (lower compartment) with PBMC (upper compartment)



transplantation, biomaterials and the surrounding cells, such as cells of innate immune response (*e.g.* monocytes and macrophages) and MSC, interplay with each other during the first two weeks.¹⁶ Considering the validation of the present TW system (data not shown), all systems were cultured up to 7 days. The medium was renewed on day 4. The supernatants and cells were collected on day 1, 4 and 7 and stored at $-80\text{ }^{\circ}\text{C}$ until use.

2.4. pH and Mg concentration

The pH values and Mg contents under two different experimental conditions were quantified at each time point using an ArgusX pH Meter (Sentron Europe BV, Roden, Netherlands) and by atomic absorption spectroscopy (AAS; Agilent 240/280 Series AA (Agilent Technologies, Waldbronn, Germany)), respectively.

2.5. Cell viability and proliferation

2.5.1. Live/dead staining. The cell viability was assessed *via* live/dead staining. The live/dead staining solution (live/dead viability/cytotoxicity kit for mammalian cells, Thermo Fisher Scientific, Schwerte, Germany) is a mixture of two highly fluorescent dyes that differentially label living and dead cells: the live cell dye labels intact, viable cells green, whereas, dead cells with incomplete membrane are presented as red fluorescent cells. To evaluate the cell viability under each experimental condition, cells and cells with Mg disc were washed with phosphate-buffered saline (PBS; 0.137 M NaCl, 0.0027 M KCl, 0.01 M $\text{Na}_2\text{HPO}_4\text{-}2\text{H}_2\text{O}$, and 0.00176 M KH_2PO_4 , pH 7.4; all chemicals were purchased from Sigma-Aldrich Chemie GmbH, Taufkirchen, Germany) and serum-free medium, respectively, and then stained with 1.6 mM calcein-AM and 2.0 mM ethidium homodimer-1 for 20 min in an incubator. Finally the staining solution was renewed with PBS (only cells) or serum-free medium (cells with Mg disc) and photographs were immediately taken with an inverted microscope.

2.5.2. Deoxyribonucleic acid (DNA) content. Proliferation was also estimated by the DNA content. Furthermore, the DNA contents were used as the normalisation factor for further biological measurements to counteract any cell proliferation variation. The cells were digested with 200 μL of lysis buffer (25 mM NaOH and 0.2 mM EDTA, pH 6) for 5 min under physiological conditions. Afterwards, the lysate was shaken at $98\text{ }^{\circ}\text{C}$ and 1000 rpm for 1 h and then cooled down to $15\text{ }^{\circ}\text{C}$ at 700 rpm using a thermomixer (Eppendorf thermomixer comfort, Hamburg, Germany). Then 200 μL of neutralisation buffer (40 mM Tris/HCl, pH 5.5) was added to all the samples. The samples were diluted 1 : 5 (v/v) in DNA dilution buffer (2.5 M NaCl in 19 mM sodium citrate, pH 7). The diluted sample of volume 100 μL was transferred into the wells of a 96-well-plate (flat bottom, black microtiter plate) and measured in duplicate. Then 50 μL of DNA working buffer (2 M NaCl in 15 mM sodium citrate, pH 7) and 50 μL of bisbenzimidazole solution (2 $\mu\text{g mL}^{-1}$) were added and the plates were incubated for 15 min in the dark. The fluorescence emission (excitation wavelength, 355 nm and emission wavelength, 460 nm) was

measured with a VICTOR3 multilabel microtiter plate reader (PerkinElmer, Massachusetts, USA). The DNA content was calculated using the fluorescence values of a corresponding standard curve. All chemicals were purchased from Sigma-Aldrich Chemie GmbH (Taufkirchen, Germany).

2.6. Production of cytokines

To identify the potential secretory production involved in the crosstalk between HUCPV and PBMC/macrophages, the protein concentrations of IL1 β , IL2, IL4, IL5, IL8, IL10, IL12, IL13, IL1RA, TNF α , IFN γ , granulocyte colony stimulating factor (G-CSF), GM-CSF, MCP-1, MIP-1 α/β , bFGF, and VEGF were measured in the supernatants using a Bio-Plex multiplex immunoassay (M500KCAF0Y; Bio-Rad Laboratories, Munich, Germany) according to the manufacturer's instructions. 50 μL of bead solution was added to the wells of a 96-well plate, then mixed (Eppendorf thermomixer comfort, Hamburg, Germany) with 50 μL of diluted standards, blanks and samples and finally incubated at room temperature (RT) with shaking at 850 rpm for 30 min. Afterwards, 25 μL of detection antibody was added and incubated for 30 min at RT with shaking at 850 rpm. Then 50 μL of streptavidin-phycoerythrin (PE) was added and incubated for 10 min at RT with shaking at 850 rpm. The assays were ended and resuspended in 125 μL of assay buffer and shaken at 850 rpm for 30 s. In each shaking step, the plate was covered with a new sheet of sealing tape and protected from light. All plates were read using a Bio-Plex 200 system (Bio-Rad Laboratories, Munich, Germany) and analysed using Bio-Plex Manager (Bio-Rad Laboratories, Munich, Germany; version 6.1). The variation induced by cell numbers under each condition was normalised by the DNA content.

2.7. Flow cytometry

To address the variation in the monocyte/macrophage ratios in single culture or coculture with and without Mg disc, the M2 phenotype was investigated by flow cytometry. The harvested cells were washed twice with ice cold PBS. After 1000 rpm centrifugation for 5 min, the cells were resuspended at a density of 1×10^6 cells in 1 mL of PBS supplemented with 1% bovine serum albumin (BSA; Sigma-Aldrich Chemie GmbH, Taufkirchen, Germany). The antibody anti-human CD 163-PE (scavenger receptor cysteine-rich type 1 protein M130, a high affinity receptor for the hemoglobin-haptoglobin complex; 560933, BD Biosciences, Heidelberg, Germany) was diluted 10 times in PBS. The cells were incubated with the diluted antibody for 1 h on ice with gentle shaking in the dark. Then the cells were washed and fixed with 3.7% formaldehyde (Sigma-Aldrich Chemie GmbH, Taufkirchen, Germany) and permeabilised with 0.7% Tween 20 (Sigma-Aldrich Chemie GmbH, Taufkirchen, Germany). The cells were washed with PBS and stained with the intracellular antibody anti-human CD 68-fluorescein isothiocyanate (FITC; scavenger receptor class D, member 1; 130096964, Miltenyi Biotec GmbH, Bergisch Gladbach, Germany) for 30 min at RT with shaking in the dark. The cells were washed again and resuspended with 1% BSA in PBS. Anti-human CD 68⁺ was selected as the pan



marker for macrophages⁶⁷ and anti-human CD 163⁺⁶⁸ as the surface marker for M2. The expressions of CD 68 and CD 163 were investigated using a flow cytometer. The flow cytometry panel was designed as follows. Positive groups were cells stained with anti CD 68 and anti CD 163 to determine the sub-population of type 2 macrophages in CD 68 positive cells. Single staining of CD 68 and CD 163 was performed for further result compensation. Propidium iodide (PI) drop solution (Bio-Rad Laboratories, Munich, Germany) was used to exclude the dead cells after each analysis. Isotype controls were the corresponding mouse IgG2b (130099119, Miltenyi Biotec GmbH, Bergisch Gladbach, Germany) and IgG1 (559320, Thermo Fisher Scientific, Schwerte, Germany). Cells resuspended in 1% BSA in PBS were used as the blank control to determine the forward scatter (FSC) and side scatter (SSC) parameters during the assays. The proportion of M2-macrophages within the monocyte population ("M2/CD 68⁺") was calculated as follows: (CD 68⁺ CD 163⁺)/CD 68⁺ cells (%).

2.8. Wound healing assay

To clarify whether the migratory ability of HUCPV was affected by the Mg levels in the supernatants and/or stimulatory cytokines from PBMC/macrophages, wound healing or scratch assay of HUCPV was carried out under different conditions. HUCPV were seeded at a density of 1×10^5 cells per well in 12-well plates containing α -MEM supplemented with 15% FBS until the cell monolayer was formed. Then the cells were starved in an α -MEM medium with low serum (1% FBS). After 24 h, the cells were treated with a 0.01 mg mL⁻¹ mitomycin C (Sigma-Aldrich Chemie GmbH, Taufkirchen, Germany) solution to prevent further cell proliferation.⁶⁹⁻⁷¹ The scratch was created with a 1 mL pipette tip. The cells were treated for another 24 h with cell culture medium (α -MEM containing 10% plasma) as the control. The TW (day 1) and CM supernatants collected under Mg, P, P + Mg conditions were also used. The scratch gap was documented using an inverted microscope (Eclipse Ti, Nikon GmbH, Düsseldorf, Germany) at t_0 and t_{24} h under bright light conditions. The relative cell migration distance was calculated from the difference of the widths of the scratch gaps between 0 and 24 h using NIS-Elements imaging software (Eclipse Ti, Nikon GmbH, Düsseldorf, Germany).

2.9. Pro-osteogenic potential of HUCPV

2.9.1. Alkaline phosphatase (ALP) activity. To evaluate the HUCPV osteogenic potential, the alkaline phosphatase (ALP) activity of the supernatant on day 7 was investigated (QuantiChrom Alkaline Phosphatase Assay Kit; BioAssay Systems, Hayward, CA, USA) according to the manufacturer's instructions. 50 μ L of the supernatant was mixed with 150 μ L of ALP assay buffer (pH 10.5) containing 5 mM Mg acetate and 10 mM *para*-nitrophenylphosphate (pNPP) liquid substrate. The activity of ALP enzymes was studied at time $t = 0$ and again after 4 min at $t = 4$ at a wavelength of 405 nm using a microplate reader (Tecan Sunrise; TECAN Deutschland GmbH, Crailsheim, Germany). Each result was normalised by the DNA content of HUCPV according to the conditions.

2.9.2. Quantitative reverse transcription polymerase chain reaction (RT-qPCR). The extraction of total cellular ribonucleic acid (RNA) was performed with a QIA shredder and a Qiagen RNeasy mini kit (Qiagen, Hilden, Germany). The RNA concentration was determined using a Nanodrop 2000c (Thermo Fisher Scientific, Schwerte, Germany). Complementary DNA (cDNA) was produced with 2 μ g of total RNA in the reaction mixture using a Sensiscript RT kit (Qiagen, Hilden, Germany). The RT-qPCR was carried out with SsoFast EvaGreen Supermix (Bio-Rad Laboratories GmbH, Munich, Germany) using a CFX96 Touch real-time PCR detection system (Bio-Rad Laboratories GmbH, Munich, Germany). Beta 2 microglobulin (B2M), glyceraldehyde-3-phosphate dehydrogenase (GAPDH) and ribosomal protein L10 (RPL10) genes were used as reference genes. Expressions of specific osteoblastic genes, collagen type I alpha 1 (COL1A1), osteocalcin (OC) and osteopontin (OPN) were studied. Primers (Table 2) were designed using Primer 3 (version 4.0.0) or found in the RTPrimerDB database and supplied by Eurofins NDSC Food Testing Germany GmbH (Hamburg, Germany). The relative fold of gene expression compared to the expression of the reference gene was calculated by the 2(-Delta CT) method ($\Delta\Delta$ CT).

2.10. Statistical analysis

Overall, at least two wells per condition with technical duplicate of three different donors of HUCPV and PBMC were performed ($n = 6$). The results in the section "pH and Mg concen-

Table 2 Primer sequences

Full name	Abbreviation	GenBank no.	Sequences	Annealing (°C)
Collagen type I alpha 1	COL1A1	NM_000088	Forward: 5'-AAGACATCCCACCAATCACC-3' Reverse: 5'-GCAGTTCTTGGTCTCTCGTCAC-3'	60
Osteocalcin	OC	NM_199173	Forward: 5'-ATGAGAGCCCTCACACTCT-3' Reverse: 5'-TGGACACAAAGGCTGCAC-3'	
Osteopontin	OPN	NM_000582	Forward: 5'-CTCCATTGACTCGAAGCACTC-3' Reverse: 5'-CAGGTCTGCGAACTTCTTAGAT-3'	
B2 microglobulin	B2M	NM_004048.2	Forward: 5'-TGCTGTCTCCATGTTTGTATGATCT-3' Reverse: 5'-TCTCTGCTCCCCACCTCTAAGT-3'	
Glyceraldehyde3-phosphate dehydrogenase	GAPDH	NM_002046.7	Forwards: 5'-GTCGAGTCAACGGATTG-3' Reverse: 5'-TGGGTGGAATCATATTGGAA-3'	
Ribosomal protein L10	RPL10	NM_006013.4	Forwards: 5'-AGTGGATGAGTTTCCGCTTT-3' Reverse: 5'-ATATGGAAGCCATCTTTGCC-3'	



tration" were obtained from three individual experiments with two replicates ($n = 3$). The results were presented as arithmetic mean \pm standard deviation (SD). The significances between the two groups were investigated by the *t*-test. The statistical significances between multiple comparisons were analysed by one-way ANOVA or one-way ANOVA on ranks using the SigmaPlot (Systat software GmbH, Erkrath, Germany; version 13.0). The following *post hoc* multiple comparisons were based on Tukey's when equal variance was assumed or Dunn's when equal variance was not assumed ($\alpha = 0.05$, $*P \leq 0.05$, $**P \leq 0.01$ and $***P \leq 0.001$). For RT-qPCR, to detect the gene expression, the Volcano Plot was used in which the regulation threshold and *P* value were set as 1.5 and 0.05, respectively (calculated using CFX Manager Software version 3.1; Bio Rad, Munich, Germany).

3. Results

3.1. pH and Mg concentration

To characterise the experimental conditions, the pH values and Mg contents of the supernatants at each time point were analysed (Table 3). According to the pH values under all experimental conditions, Mg and/or cells were exposed to pH ranging from 8.05 to 8.32. The Mg contents under conditions without Mg disc were in a similar range (0.60 to 0.78 mM) to the value (0.71 mM) under blank conditions. Considering the treatment with the Mg disc, the Mg contents under cell-seeded conditions (*i.e.* H + Mg, P + Mg and H + P + Mg) fell in the ranges from 4.71 to 6.87, 15.62 to 18.03, and 8.19 to 14.86 mM at various time points, respectively.

3.2. Cell viability and proliferation

Cell live/dead staining was performed to estimate the cell viability and adherence/proliferation behaviour. Bright green represented the living cells, whereas red fluorescence indicated the dead cells (Fig. 2a and c). The nucleic acid content can be a quantifiable indicator of the cell number.⁷² The bisbenzimidazole can specifically bind to the nucleic acid. Therefore, the fluorescence emission of their complexes correlates with the cell number and reflect the cell proliferation.

Adherent HUCPV could be found under all experimental conditions up to 7 days of culture (Fig. 2a, green). After 1 day

of culture, the Mg disc increased the cell number (Fig. 2a) and DNA contents (Fig. 2b) of HUCPV compared to the HUCPV monolayer. In the subsequent culture period, PBMC stimulated the HUCPV growth under H + Mg *vs.* H + P + Mg (day 4) and H + Mg *vs.* H + P (day 7) conditions, rather than Mg discs. Particularly, a reduction was observed on day 7 when HUCPV were directly exposed to the Mg surfaces. Besides, HUCPV and PBMC mixed cultures, HUCPV seeded on the Mg disc with and without PBMC or kept as monocultures, showed no significantly increased number of dead cells and proliferative cells compared with controls. Again, similar observations were made with the corresponding DNA contents.

Besides, green-labelled adherent PBMC were identified, as shown in Fig. 2c, in the presence of the Mg disc, the dead cells were not apparently increased, and meanwhile, the cell number of PBMC increased on day 4 and 7 in the presence of Mg, H or both (Fig. 2d). Specifically, PBMC exposed to Mg could proliferate much faster than those cultured with HUCPV on day 1 and 4.

3.3. Roles of degradable Mg in the immunomodulatory properties of HUCPV

3.3.1. The synergetic effects of Mg and HUCPV on attenuating inflammation via a moderate inflammatory cytokine production. To determine whether degradable magnesium can influence the immunomodulatory properties of HUCPV and its cellular communication with immune cells (PBMC), the inflammatory cytokine expression in the supernatants was investigated. Several key secretory factors involved were selected: pro-inflammatory (IL1 β , IL2, TNF α and IFN γ) and anti-inflammatory (IL4, IL5, IL10, IL13, IL1RA and G-CSF). After comparing the cytokine levels between conditions with Mg and without Mg (+Mg *vs.* -Mg), the fold change of cytokine production was quantified and is summarised in Fig. 3.

A decrease in the levels of pro-inflammatory proteins (IL2, TNF α , IL1 β and IFN γ) on day 4 and an increase in the production of anti-inflammatory (IL13 and G-CSF) cytokines on day 7 were observed in HUCPV in contact with Mg. Thus, Mg could induce HUCPV to an anti-inflammatory state, especially on day 4.

Initially, PBMC were activated when cultured with Mg (day 1), expressing high levels of all the above-mentioned cytokines. However, when PBMC were cocultured with HUCPV, a downre-

Table 3 The pH value and Mg concentration of the supernatants in the TW system. The blank group on day 0 was characterised (pH: 7.60 ± 0.009 ; Mg content: 0.71 ± 0.009 mM). H: HUCPV; P: PBMC

Conditions	pH			Mg contents (mM)		
	Day 1	Day 4	Day 7	Day 1	Day 4	Day 7
Mg	8.27 ± 0.004	8.20 ± 0.013	8.30 ± 0.054	7.66 ± 0.074	11.75 ± 0.079	8.36 ± 0.044
H	8.09 ± 0.012	8.05 ± 0.029	8.09 ± 0.013	0.68 ± 0.007	0.63 ± 0.005	0.60 ± 0.002
H + Mg	8.15 ± 0.041	8.27 ± 0.030	8.24 ± 0.009	5.74 ± 0.001	18.03 ± 0.070	13.81 ± 0.206
P	8.05 ± 0.009	8.09 ± 0.023	8.14 ± 0.015	0.54 ± 0.002	0.64 ± 0.004	0.78 ± 0.004
P + Mg	8.23 ± 0.022	8.31 ± 0.066	8.23 ± 0.036	4.71 ± 0.011	15.62 ± 0.064	8.19 ± 0.057
H + P	8.05 ± 0.020	8.09 ± 0.014	8.25 ± 0.132	0.64 ± 0.001	0.70 ± 0.003	0.69 ± 0.003
H + P + Mg	8.27 ± 0.051	8.32 ± 0.039	8.26 ± 0.058	6.87 ± 0.043	17.91 ± 0.083	14.86 ± 0.107



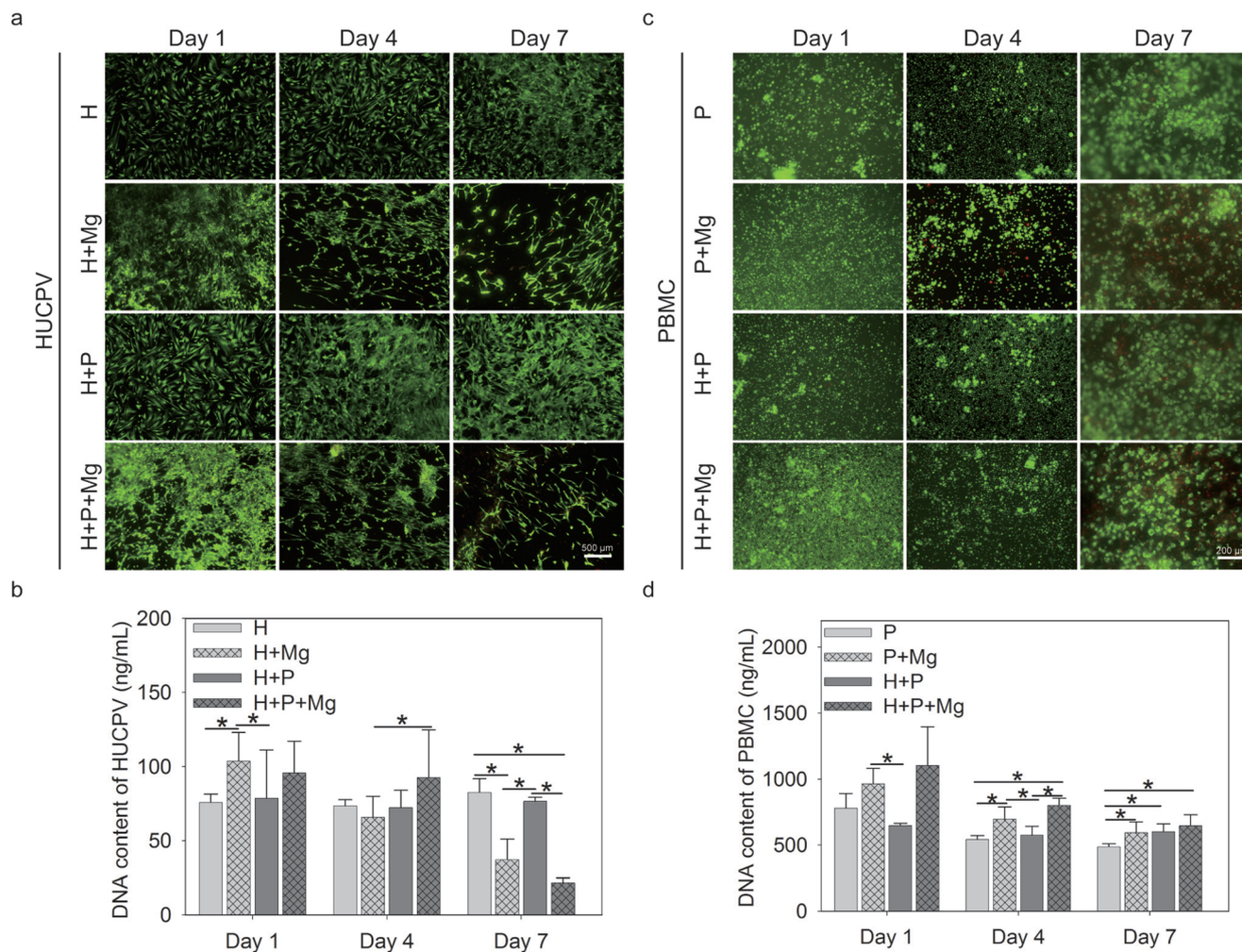


Fig. 2 Cell viability and proliferation of HUCPV and PBMC in the TW system. (a) Live/dead staining of HUCPV and (c) PBMC. Green colour and red spot represent alive and dead cells, respectively. (b) DNA content of HUCPV and (d) PBMC. Scale bars represent 500 μm (a) and 200 μm (c). Stars indicate significant differences between the two conditions (ANOVA; $\alpha = 0.05$, $*p \leq 0.05$). H: HUCPV; P: PBMC.

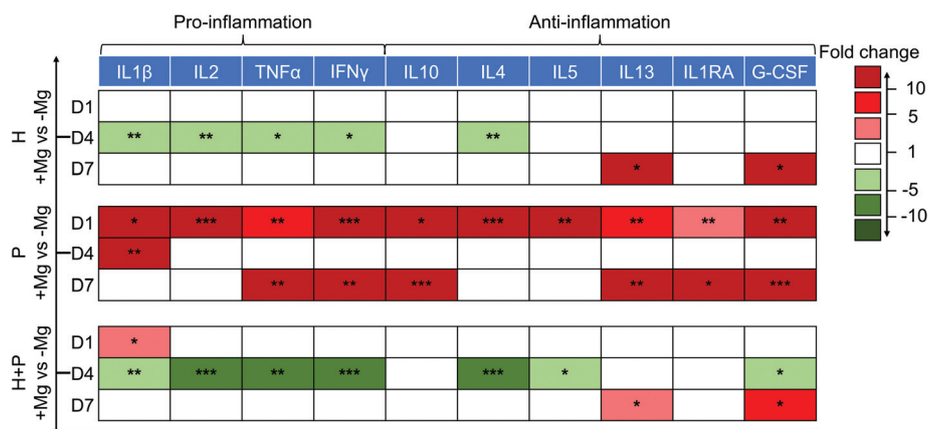


Fig. 3 Cytokine levels in the supernatant of HUCPV, PBMC and HUCPV-PBMC coculture in the presence and absence of Mg disc. The colour scale represents fold change compared between the conditions with Mg disc and controls without Mg disc (red: up-regulation; white: no change; and green: down-regulation). Asterisks indicate significant differences between the two conditions (t-test; $\alpha = 0.05$, $*p \leq 0.05$, $**p \leq 0.01$ and $***p \leq 0.001$). H: HUCPV; P: PBMC.



gulation of pro-inflammatory IL2, TNF α , and IFN γ (fold change -5 to -10) on day 4, and an increase in the anti-inflammatory IL13 and G-CSF levels (fold change 5 to 10) on day 7 were observed (Fig. 3). Besides, a similar increase of IL4/5 caused by Mg was also observed in PBMC. However, a clear decrease of IL4/5 caused by Mg was observed in cocultures. Obviously, a moderate inflammation could be observed, when HUCPV mediates Mg-induced cytokine production in PBMC.

Taken together, these results indicated that when exposed to Mg, HUCPV could create an anti-inflammatory microenvironment. Furthermore, HUCPV play a role in the suppression of Mg-induced inflammatory cytokine production in PBMC. The immune-modulation of HUCPV on PBMC/macrophage behaviour is mainly dominated by the immunomodulatory characteristics of HUCPV.^{73,74}

3.3.2. The synergetic effects of Mg and HUCPV on enhancing the M2 macrophage phenotype in PBMC. In PBMC, the cells, differentiating from monocytes into M2 macrophages, were identified by the phenotype changes after 1, 4 and 7 days *via* flow cytometry. PBMC were labelled with anti-human CD 68⁺ (pan marker for macrophages; Fig. 4a) and anti-human CD 163⁺ (surface marker for the M2 phenotype; Fig. 4b). The flow cytometry (Fig. 4c) revealed that, on day 1, an elevated percentage of M2 in macrophages was exhibited, when PBMC was exposed to HUCPV (P *vs.* H + P). In addition, compared to the

H + P condition, M2/CD 68⁺ cell population increased under the H + P + Mg condition from day 4 onwards. Particularly on day 7, Mg in synergy with HUCPV, promoted the M2 macrophage phenotype. Interestingly, Mg could induce higher number of M2 macrophages than HUCPV alone (P + Mg *vs.* H + P). Thus, it can be postulated that this polarisation may be derived from the effect of Mg on the immunomodulatory properties of HUCPV.

3.4. Roles of degradable Mg in inflammatory secretion-mediated fracture healing

Within the context of tissue repair, after recruitment of stem cells by immune cells at the site of injury, the cell-cell interactions between immune cells and stem cells are vital. Thus the contributions of immune cells (PBMC) to HUCPV behaviour under the stimulation of Mg were studied, including wound healing assay and the evaluation of HUCPV osteogenic potential.

3.4.1. The synergetic effects of Mg and PBMC on stimulating the migratory ability of HUCPV. To elucidate the influence of Mg (\pm PBMC) on MSC migration into the injury site *in vitro*, a multiplex assay of migration/wound healing related cytokines (IL8, GM-CSF, MCP-1 and MIP-1 α/β) was performed (Fig. 5a). On both day 1 and day 7, PBMC exhibited higher production of GM-CSF, IL8, MCP-1 and MIP-1 α/β after Mg exposure.

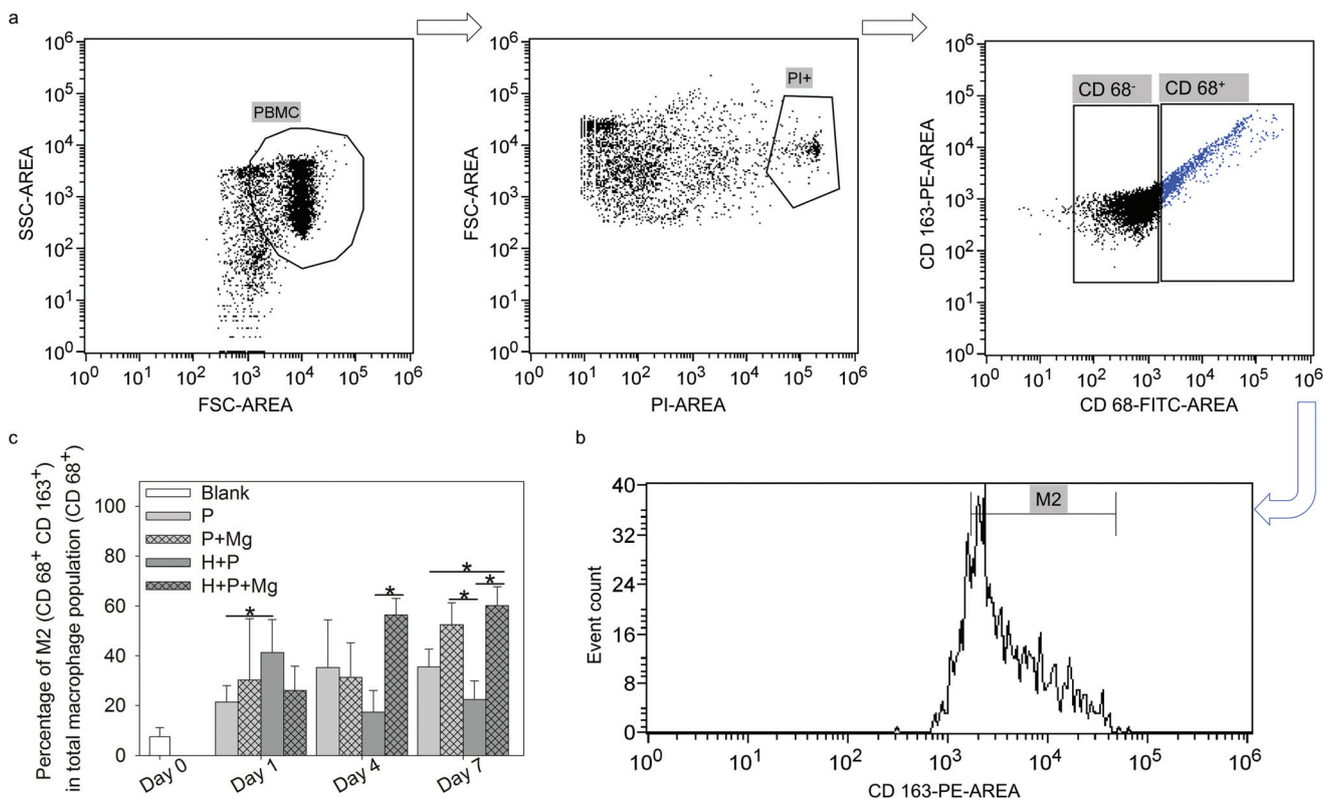


Fig. 4 Flow cytometry analysis of the M2 subpopulation at each time point in the TW system. (a) The selection of macrophage (CD 68⁺) cells. In brief, the living cells were firstly gated out, and then the CD 68⁺ cells were gated out. (b) The selection of M2 macrophages in the CD 68⁺ subgroup, in which, CD 163 was used as the specific marker to identify M2 macrophages from total macrophages. (c) Percentage of M2 in the total macrophage population. Stars indicate significant differences between the two conditions (ANOVA; $\alpha = 0.05$, $*p \leq 0.05$). H: HUCPV; P: PBMC.



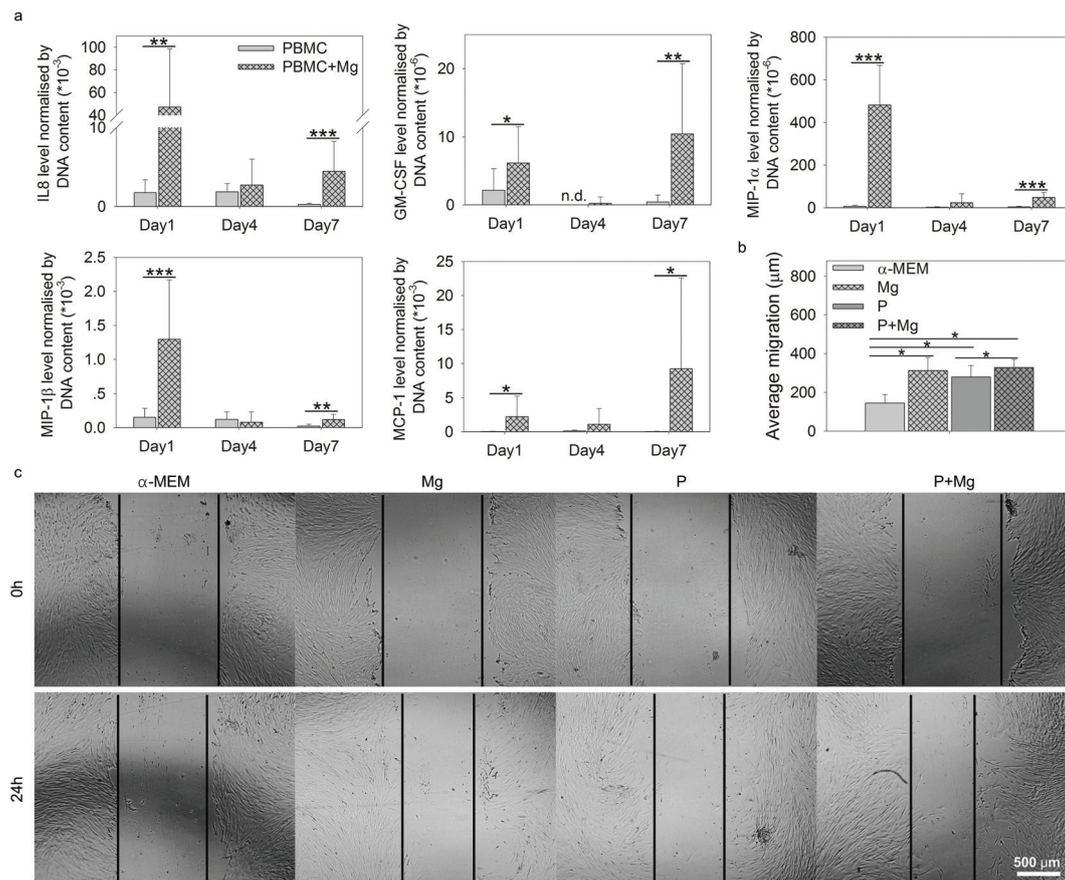


Fig. 5 The migratory ability of HUCPV. (a) Selected cytokine (IL8, GM-CSF, MIP-1 α/β , and MCP-1) production under P and P + Mg conditions at each time point. (b) Average migration distance of HUCPV, when cultured in the supernatant from the TW system. The migration distance was calculated from the area without cell difference between 0 and 24 h. (c) Microscopic examination of a scratch assay. The scratch at 0 and 24 h were quantified by the gap width. Scale bar represents 500 μm . Stars indicate significant differences between the two conditions (*t*-test or ANOVA, $\alpha = 0.05$; **p* ≤ 0.05 ; ***p* ≤ 0.01 ; and ****p* ≤ 0.001). P: PBMC.

The higher levels of these selected 5 cytokines in PBMC (+Mg) indicated the possible promotion of MSC migration into injured sites,^{45,75,76} to stimulate the wound healing process. Thus, attempting to further examine the migration potency of HUCPV affected by Mg, an *in vitro* wound healing assay was carried out with the supernatants collected on day 1 under the TW conditions, comprising Mg, P, and P + Mg.

The migration distance of HUCPV was calculated from the scratch area difference between 0 and 24 h (Fig. 5b). In comparison with the α -MEM control group, Mg or PBMC (P) or both (P + Mg) could significantly increase the migratory ability of HUCPV. Similarly, a stimulated migration capacity of HUCPV was observed under the P + Mg condition, compared with HUCPV under the P condition. A large amount of HUCPV existed at and crossed the edge of scratch in TW (Fig. 5c). These results revealed that HUCPV have more capacity to migrate when cultured with Mg and/or PBMC for 24 h.

3.4.2. Enhanced pro-osteogenic potential of HUCPV by Mg but not PBMC. The pro-osteogenic potential of HUCPV was assessed using the selected cytokine production (e.g., bFGF,

VEGF, and IL12),²³ and ALP activity in the supernatants, as well as the expression of osteogenic genes.

As shown in Fig. 6a, Mg elevated the production of bFGF (H vs. H + Mg on day 1), VEGF (H vs. H + Mg or H + P vs. H + P + Mg on day 7), and IL12 (H vs. H + Mg on day 7). Interestingly, on day 1 or day 4, the involvement of PBMC could significantly reduce the production of these three cytokines (H vs. H + P or H + Mg vs. H + P + Mg).

The ALP activity normalised by DNA is presented in Fig. 6b. The ALP activity on day 7 can be attributed to HUCPV as PBMC/macrophages do not produce a significant amount of ALP.⁷⁷

In TW, compared to the HUCPV monoculture or cocultures, the levels of ALP were significantly increased when the cells were exposed to the Mg disc. In addition, the fold change of osteogenesis-related gene expression of COL1A1, OC and OPN in HUCPV is indicated by the colour scale in Fig. 6b. The coculture with PBMC could enhance the gene expression of OC in HUCPV. Similarly, an increase of COL1A1, OC and OPN was found when Mg was used. Even a synergistic effect of P and Mg could be seen for OC (the regulation fold reached 20).



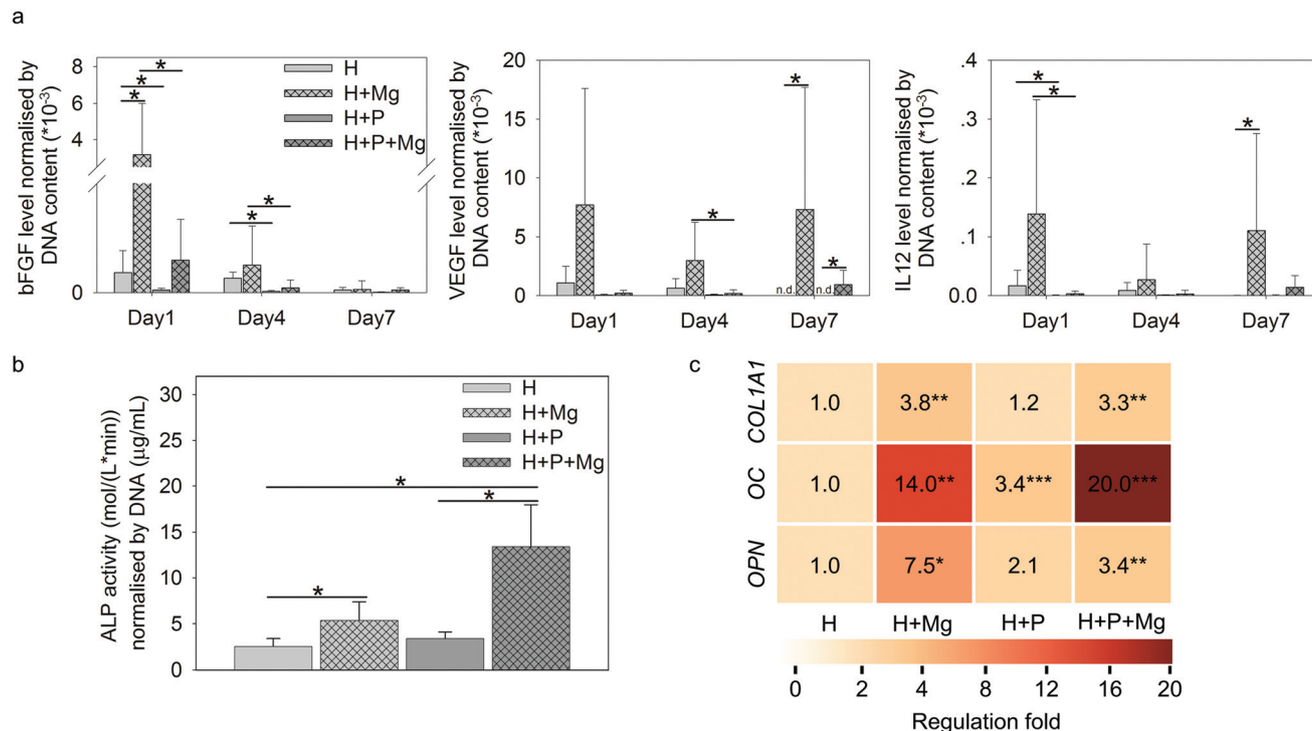


Fig. 6 Pro-osteogenic activity of HUCPV. (a) Selected cytokine (BFGF, VEGF, and IL12) production under H/H + P with and without Mg at each time point. (b) ALP activity in TW on day 7. (c) Osteogenic gene expressions in TW on day 7. Cytokine production and ALP activity were normalised by the DNA content. Fold of gene expression was compared to controls in TW. The significance is represented by asterisks and obtained from *post hoc* multiple comparisons between each condition or comparing to controls (H) (t-test or ANOVA, $\alpha = 0.05$; * $p \leq 0.05$, ** $p \leq 0.01$ and *** $p \leq 0.001$). H: HUCPV; P: PBMC.

4. Discussion

A biomaterial triggers host immune reactions after implantation prior to MSC recruitment.^{16,78} MSC can interact with immune cells by modulating immune responses and promoting bone regeneration (osteo-immunomodulation), and indeed MSC and immune cells can share the secreted cytokines.^{41,42} In addition, the immune responses can affect the mechanical properties and performance of biomaterials.⁷⁹ However, a comprehensive knowledge of the effects of degradable Mg on paracrine signaling between immune cells (PBMC) and MSC is still lacking. Thus, the roles of degradable Mg in the immunomodulatory properties of MSC, as well as immune-mediated MSC behaviour, like MSC migration and osteogenic differentiation, were studied in the current research.

Tissue repair cells, immune cells, and Mg ions should be the three protagonists involved in the cascades of degradable Mg-modulated fracture healing. HUCPV are defined as mesenchymal progenitors, multipotent and allogeneic stem cells.²² In comparison with bone marrow mesenchymal stem cells (BMMSC), HUCPV, as an alternative foetal MSC, possess the differentiation capacity and the immunomodulatory properties as well.^{20–23} Macrophages are one of the key immune cells (PBMC) participating in fracture healing thanks to their polarisation plasticity.³⁸ Mg and Mg-based alloys have been proved to be potential degradable materials for regenerative

applications.^{80–82} Here pure Mg was selected to avoid an increase in the biological complexity due to an alloying element. As a more representative human *in vivo*-like system of biomaterial-modulated wound healing, TW is highly relevant for a microenvironment and allows an in-depth tracking of the effects of degradable Mg on cell-cell interactions. In the TW system, PBMC were typically seeded on the top of the chamber and HUCPV in the lower compartment, and thus the different cells were always interacting with each other, allowing monocytes to differentiate into macrophages while HUCPV can differentiate into osteoblast-like cells. Meanwhile, the Mg contents varied from 0.5 mM to 18 mM as a consequence of Mg degradation. Thus, in the context of Mg degradation, the focus of this study could provide an insight into paracrine secretions between HUCPV and PBMC during the fracture healing process.

In most mammalian cells, the total concentration of Mg is in the range of 14–20 mM, and that in an intracellular medium is between 0.5 and 0.7 mM.^{83,84} Moreover, many cells could maintain the Mg contents at the physiological levels, which is irrelevant to the extracellular Mg concentration.⁸⁵ In the present TW system, the Mg concentration in the cell culture medium (α -MEM) was 0.7 mM. As reported, Mg²⁺ could play various roles in the cell growth: stimulation of the cell proliferation⁸⁶ or inhibitory effect on the cell number.⁸⁷ For instance, in U2OS cells, a slight inhibition of cell prolifer-



ation was observed when the Mg concentration was higher than 5 mM.⁸⁸ Other studies suggested that Mg ions can significantly inhibit the cell growth in the range from 12.5 to 50 mM.^{89–93} Similarly, as calculated in Fig. 2b, an observation was concluded that there is a decreased DNA content of HUCPV in the presence of Mg (around 14 mM), while an increased DNA content of HUCPV when exposed to Mg and P + Mg (6 mM or about 18 mM), separately. Meanwhile, Mg at various concentrations ranging from 0.54 to 17.91 mM promoted PBMC proliferation (Fig. 2d). Thus, the effects of Mg ions on cellular proliferation could be biphasic⁹⁴ or rely on the whole microenvironment between materials and cells.

In the host immune response after Mg implantation, the secretory cytokines were the vital players in the regulation of inflammation. Because of the pro- and anti-inflammatory properties of MSC and PBMC/macrophages,^{40,95} this correlation was evaluated *via* the cytokine production of the inflammatory panel (TNF α , IFN γ , IL1 β , IL2, IL4, IL5, IL10, IL13, IL1RA, and G-CSF) in the current TW coculture. In the presence of Mg, HUCPV secreted a lower number of pro-inflammatory TNF α , IFN γ , IL1 β , and IL2 on day 4, when the Mg contents in the supernatants reached 18 mM. Moreover, on day 1 and day 7 (the Mg concentration varied in the range from 5 mM to 8 mM), Mg-stimulated PBMC exhibited an intensive inflammation state, releasing higher levels of pro- and anti-inflammatory cytokines, and this stimulation probably could be recognised by the inflammatory response and cell activation.⁹⁶ Meanwhile, Mg, at a concentration of 16 mM, had no effects on the inflammatory cytokine production in PBMC (day 4). Therefore, the effects of degradable Mg on the production of inflammatory cytokines in HUCPV monoculture or PBMC was biphasic. Nevertheless, the coculture of HUCPV could switch such Mg-induced higher levels of pro-inflammatory cytokines to a down-expressed state on day 1 (Mg: 7 mM) and day 4 (Mg: 18 mM). Similarly, on day 7, at a concentration of 15 mM, the Mg-induced higher secretions of anti-inflammatory cytokines in PBMC was also attenuated by HUCPV (colour scale: dark red to light red). These comparisons of cocultures indicated the immuno-suppressive role of HUCPV interacting with PBMC under the stimulation of Mg, regardless of the Mg concentration. Thus, the interface of Mg biomaterial–tissue could retain a more moderate status of inflammation. This is consistent with the study of the immuno-suppressive capacity of MSC that MSC can regulate the transition from a T helper type 1 (Th1)-driven response to an anti-inflammatory Th2 response, characterised by lower concentrations of TNF α , IFN γ and IL2, as well as higher production of IL4, IL5, IL10 and IL13.⁴⁰

Macrophages are polarised by various stimuli to generate heterogeneous populations with different characteristics and functions. The main subtypes of macrophages are M1 and M2 phenotype macrophages, which are vital factors in the resolution of inflammation and tissue remodelling.^{97,98} M1 phenotype macrophages, which are associated with pro-inflammatory activities,⁹⁹ are often induced by toll-like receptor (TLR) ligands or Th1 cytokines (TNF α and IFN γ).¹⁰⁰ In addition, the M2 phenotype macrophages are polarised by Th2-derived cyto-

kines, including IL4, IL10, IL13, transforming growth factor beta (TGF β) or prostaglandin E2 (PGE2).¹⁰⁰ M2 macrophages are called “tissue repair macrophages” because they promote tissue repair through immune tolerance and tissue remodelling, removal of debris, and immune regulation.¹⁰¹ Besides, MSC present interactive but complicated immunomodulatory properties for immune cells *via* their cytokines. In the scale of MSC immunomodulation in cells from innate and adaptive immune responses,^{102,103} despite various cell groups, *i.e.*, monocytes, lymphocytes and dendritic cells, many research studies consolidated that immunomodulation by MSC is crucial for macrophage polarisation.^{100,101} For instance, MSC was indicated to trigger the monocyte population towards the pro-healing M2 phenotype,^{104,105} which aids in the resolution of inflammation and tissue remodelling.^{97,98,106} In addition, the correlation between Mg²⁺ and MSC and its effects on bone tissue have been investigated.^{60,107,108} The anti-inflammatory properties of Mg were reported as well.^{109,110} In the past decades, multiple studies have confirmed that MSC could induce anti-inflammatory M2 polarisation,^{38,111,112} which could depend on paracrine signaling through the secretion of soluble factors, such as TGF β , PGE2 and indoleamine 2,3-dioxygenase (IDO).^{113–115} The current results of TW suggested that the subpopulation of M2 was increased by Mg alone or Mg cocultured with HUCPV (\pm Mg). It implied that the M2 macrophage phenotype was not only influenced by Mg, but also modulated by MSC (HUCPV). This effect of MSC is at least partially mediated by soluble factors, such as the widely reported PGE2.^{116,117} Prostaglandin E synthase (PTGES) is a key enzyme in the production of PGE2.¹¹⁸ Our preliminary work has shown a significantly upregulated expression of PTGES2 in HUCPV with magnesium on day 7 (ESI, Fig. S3 \dagger). This result implies that the PGE2 production could be increased by Mg even though the HUCPV proliferation was decreased by Mg. The role of Mg in influencing PGE2 expression in immunomodulation was also certified by a study that Mg²⁺ (5 mM) could regulate immune responses by decreasing the expression of IL1 β and IL6, and by increasing the expression of PGE2 and IL10 in Murine C3H/10T1/2 MSC stimulated with LPS or TNF α .⁴⁴ Furthermore, the secretory ability of MSC is not correlated with their proliferative ability.¹¹⁹ Therefore, the decreased DNA contents of HUCPV on day 7 (Fig. 2b) should not be directly linked to their M2 phenotype polarisation potential.

During the fracture healing process, MSC initially migrate to the bone injury site to participate, before differentiating into osteoblasts-like cells.^{120–123} This recruitment of MSC can be influenced by MCP-1, MIP-1 α , MIP-1 β and IL8.^{45,124} As shown in Fig. 5a, cytokines inducing MSC migration, such as IL8, GM-CSF, MCP-1, and MIP-1 α/β in PBMC, were dramatically upregulated by Mg, especially on either day 1 or day 7. In addition, the supernatants from Mg or PBMC or both could stimulate HUCPV migration into the closure area in TW (Fig. 5b and c). Therefore, the enhanced migratory ability of HUCPV was not only due to Mg itself (Mg contents), but also due to the involvement of Mg-induced paracrine signaling of



IL8, GM-CSF, MCP-1, and MIP-1 α/β . Furthermore, the increased recruitment of HUCPV could also contribute to the understanding of *in vivo* increased bone formation observed after magnesium implantation^{6,125–127} and even the increased osteoblast apposition on the implant surface.^{8,128} Moreover, the higher levels of IL12, bFGF, and VEGF are recognized as markers of foetal-derived osteoblastogenesis.²³ Mg-Stimulated higher secretions of IL12, bFGF, and VEGF, as well as various osteogenic parameters such as ALP activity, collagen synthesis and expression of OC and OPN, indicated that Mg can enhance the osteogenic differentiation of MSC.^{129,130} As reported, monocytes can enhance the osteogenic gene expression of MSC¹³¹ and further promote osteoblast formation as well.^{132,133} Furthermore, it has also been well-documented that macrophages may engage in the degradation of biomaterials and can contribute to the osteogenic behaviour of MSC which aid in bone regeneration.¹³⁴ The desired immune microenvironment can effectively stimulate osteogenesis, and therefore, the roles of degradable Mg *via* paracrine secretions in immune-mediated osteogenesis would need to be further studied for longer periods (as 7 days reflect the early osteogenesis).

5. Conclusions

Our findings emphasised degradable Mg influenced the HUCPV–PBMC interaction. Mg could affect the cell behaviour in cooperation with other cell types. Mg collaborated with HUCPV to attenuate inflammation *via* a moderate cytokine release, accompanied by the pro-healing M2 macrophage phenotype. In turn, Mg in synergy with immune cells (PBMC) could stimulate HUCPV migration, while the pro-osteogenic potential of HUCPV could be influenced by Mg but not PBMC. Thus, a suppressive inflammation and faster bone tissue repair could be expected due to the osteo-immunomodulatory properties of degradable Mg.

Conflicts of interest

There is no conflict of interest to declare.

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Written informed consent from the donor was obtained for the use of these samples in research and all experiments were conducted in accordance with the Declaration of Helsinki. HUCPV were isolated and all experiments were performed with the approval from the local ethical committee Ethik-Kommission der Ärztekammer Hamburg (Hamburg, Germany, PV4058).

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