A Comparative Analysis of the In Vitro Effects of Pulsed Electromagnetic Field Treatment on Osteogenic Differentiation of Two Different Mesenchymal Cell Lineages

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Abstract

Human mesenchymal stem cells (MSCs) are a promising candidate cell type for regenerative medicine and tissue engineering applications. Exposure of MSCs to physical stimuli favors early and rapid activation of the tissue repair process. In this study we investigated the in vitro effects of pulsed electromagnetic field (PEMF) treatment on the proliferation and osteogenic differentiation of bone marrow MSCs (BM-MSCs) and adipose-tissue MSCs (ASCs), to assess if both types of MSCs could be indifferently used in combination with PEMF exposure for bone tissue healing. We compared the cell viability, cell matrix distribution, and calcified matrix production in unstimulated and PEMF-stimulated (magnetic field: 2 mT, amplitude: 5 mV) mesenchymal cell lineages. After PEMF exposure, in comparison with ASCs, BM-MSCs showed an increase in cell proliferation (p < 0.05) and an enhanced deposition of extracellular matrix components such as decorin, fibronectin, osteocalcin, osteonectin, osteopontin, and type-I and -III collagens (p < 0.05). Calcium deposition was 1.5-fold greater in BM-MSC-derived osteoblasts (p < 0.05). The immunofluorescence related to the deposition of bone matrix proteins and calcium showed their colocalization to the cell-rich areas for both types of MSC-derived osteoblast. Alkaline phosphatase activity increased nearly 2-fold (p < 0.001) and its protein content was 1.2-fold higher in osteoblasts derived from BM-MSCs. The quantitative reverse-transcription polymerase chain reaction (qRT-PCR) analysis revealed up-regulated transcription specific for bone sialoprotein, osteopontin, osteonectin, and Runx2, but at a higher level for cells differentiated from BM-MSCs. All together these results suggest that PEMF promotion of bone extracellular matrix deposition is more efficient in osteoblasts differentiated from BM-MSCs.

Key words: human adipose-derived stem cells; human mesenchymal stem cells; osteogenic differentiation; pulsed electromagnetic field

Introduction

The use of stem cells in regenerative medicine is an appealing area of research that has received a great deal of attention in recent years. Ideally, stem cells for regenerative medicinal applications should meet the following set of criteria: they should be (i) found in abundant quantities (millions to billions of cells); (ii) readily collected and harvested by a minimally invasive procedure; (iii) differentiated along multiple cell lineage pathways in a reproducible manner; (iv) safely and effectively transplanted to either an autologous or allogeneic host.¹,² The use of totipotent and multipotent stem cells in regenerative medicine has opened the way for autologous tissue transplantation. However, mesenchymal stem cell (MSC) concentration in bone marrow (BM) transplants is lower with respect to cultured cells. It also has to
be considered that cell amplification by culture is not free from the dangers of bacterial contamination, xenogenic risk, or cellular transformation, influencing MSC differentiation capacities. It is still necessary to perform more clinical trials to select the best treatment modality to repair bone tissue.1,4

Although BM-derived MSCs (BM-MSCs) are being widely used in cell-based therapy and tissue engineering, adipose-derived MSCs (ASCs) might serve as an alternative source of MSCs. In the setting of an autologous implant, the use of ASCs as a cell source for bone therapeutic applications presents many potential advantages in comparison with BM-MSCs. Subcutaneous adipose depots are accessible, abundant, and replenishable, thereby providing a potential adult stem cell reservoir for each individual.5–10 Aside from the use of MSCs, different types of approach have been applied in order to improve osteoinduction and consequently to reduce the differentiation period. Many biomaterials have been tested,11–13 and following a “biomimetic” approach, different bioreactors have been used to improve osteogenesis.14–16 Biophysical stimulation (i.e., pulsed electromagnetic fields [PEMF] or low-intensity pulsed ultrasound) has been used in clinical settings to accelerate and finalize the healing process of a fresh fracture, or a fracture at risk of nonunion, and to enhance the spontaneous repair capability of the bone tissue (i.e., to reactivate the healing process in pathological conditions such as delayed union or pseudoarthrosis).17 In particular, PEMFs have been widely used in orthopedics for at least three decades,18,19 but as far as we know a comparative analysis to evaluate their potential effects on in vitro culture of both BM-MSCs and ASCs has not been performed.

PEMF therapy is approved for bone disorders in animals and in humans, including nonunited bone fracture healing, pseudoarthrosis, and osteoporosis.20–22 PEMF treatment reportedly aids the healing of osteotomies23 and has been applied clinically to promote bone healing for many years.24 Clinical effectiveness was initially thought to be due to the accelerated formation of bone matrix by the weak electric current generated by the magnetic field25 in animal experiments.26,27 Based on studies that indicated that electromagnetic fields may heal bone fractures and slow down bone matrix loss. Current studies on stem cells have revealed that the regeneration of human body tissues and supplementation of mature functioning cells are due to the proliferation and differentiation of stem cells.28–31 The mechanism of action underlying how PEMF promotes the formation of bone in an in vitro environment remains elusive. The electromagnetic stimulation raises the net Ca2+ flux in human osteoblast-like cells,32 and, according to Pavalko’s diffusion-controlled/solid state signaling model,33 the increase in the cytosolic Ca2+ concentration is the starting point for signaling pathways targeting specific bone matrix genes. Considering the model of Pavalko, the analysis of the transcripts specific for decorin, osteopontin, and type-I collagen revealed, concordantly, that the application of the electromagnetic wave caused an increase in gene transcription. In recent published studies, it was reported that PEMF exposure could enhance early cell proliferation in BM-MSC–mediated osteogenesis and accelerate osteogenesis.34-35

In view of a tissue engineering approach for bone repair, the aim of this study was to assess if the application of a physical stimulus could have a biological effect on human BM-MSCs and ASCs. Therefore we investigated the influence of PEMF on the proliferation and differentiation of both types of MSCs in terms of osteoblast morphology, proliferation, and deposition of a mineralized extracellular matrix (ECM). Considering the possible clinical application of PEMF in support of skeletal therapy, the final scope of this study was to perform a comparative analysis on the efficacy of electromagnetic treatment in both types of stem cell lineage for in vivo bone development. To the best of our knowledge, this is the first comparative analysis on this topic.

Materials and Methods

Electromagnetic bioreactor apparatus

The electromagnetic bioreactor consisted of a supporting structure custom-designed in a tube of polymethylmethacrylate; the windowed tube had a well-plate and two solenoids (Helmoltz coils, the planes of which were parallel).35 In this experimental setup, the magnetic field and the induced electric field were perpendicular and parallel to the scaffold surfaces, respectively. The cell surfaces were 5 cm away from the solenoid plane, and the solenoids were powered by a Biostim SPT pulse generator (Igea, Carpi, Italy), which generated PEMFs. Given the position of the solenoids and the characteristics of the pulse generator, the electromagnetic stimulation had the following parameters: magnetic field intensity equal to 2 ± 0.2 mT, induced electric tension amplitude equal to 5 ± 1 mV, signal frequency of 75 ± 2 Hz, and pulse duration of ~1.3 msec.36 The magnetic field was measured with a Hall Effect transverse gaussmeter probe (Sypris Solutions, Louisville, KY) and gaussmeter (Laboratorio Elettrofisico, Milan, Italy), the induced electric tension was measured with a standard coil probe, and the temporal pattern of the electromagnetic signal was evaluated by a digital oscilloscope (LeCroy, Chestnut Ridge, NY). In clinical settings the PEMF parameters were similar, but a period of 30–40 min represented the exposure time for tissues or organs. As shown in this in vitro study, the PEMF biological effect was directly evaluated on both stem cells and the exposure time was determined experimentally. In our experimental settings, the electromagnetic bioreactor was placed into a standard cell culture incubator in a 37°C, 5% CO2 environment.

Isolation, expansion, and culture of BM-MSCs and ASCs

The design of this study was approved by the Institutional Review Board of the Fondazione IRCCS Policlinico San Matteo and the University of Pavia (2011).

BM-MSCs. BM aspirates were harvested from healthy pediatric hematopoietic stem cell donors after obtaining written informed consent. Thirty milliliters of BM from each donor was assigned to BM-MSC generation; heparin was added as an anticoagulant. Mononuclear cells were isolated from BM aspirates (30 mL) by Ficoll density gradient centrifugation (density, 1.077 g/mL; Lymphoprep, Nycomed Pharma, Oslo, Norway) and plated in noncoated 75- to 175-cm2 polystyrene culture flasks (Corning Costar, Celsio, Milan, Italy) at a density of 16 × 10⁶ cells/cm². Cells were cultured in Mesencult medium (Stem Cell Technologies, Vancouver, Canada) supplemented with 2 mM L-glutamine, 50 μg/mL gentamycin, and 10% fetal calf serum. Cultures were maintained at 37°C in a humidified atmosphere containing 5% CO2. After 48 h, nonadherent cells were discarded and culture medium was replaced twice a week. After reaching 80%
confluence as a minimum, the cells were harvested and replated for expansion at a density of 4000 cells/cm² until the fifth passage. The colony-forming unit fibroblast assay (CFU-F) was performed as described previously.³⁷ CFU-F formation was examined after 12 days of incubation in a humidified atmosphere (37°C, 5% CO₂); the clonogenic efficiency was calculated as the number of colonies per 10⁶ BM mononuclear cells seeded. According to the International Society for Cellular Therapy on the nomenclature of mesenchymal progenitors, the cells cultured for this study were defined as multipotent stromal cells. To phenotypically characterize BM-MSCs and to define their purity, FACS analysis was performed as previously described.³⁷ After reaching 80% confluence at a minimum, the cells were harvested and replated for expansion at a density of 2.5 x 10⁵ cells/cm². The cells were cultured at 37°C, 5% CO₂, and three fifths of the medium was renewed every 3 days.

Adipose-tissue MSCs. Subcutaneous adipose tissue was obtained from healthy pediatric donors during orthopedic surgery. Informed consent was obtained before the surgical intervention. In brief, the tissue was finely minced and then incubated in digestion buffer (0.01% collagenase type-II in Dulbecco’s modified Eagle’s medium [DMEM] F12-HAM medium supplemented with 10% fetal calf serum, 100 U penicillin/streptomycin, amphotericin) for 1 h at 37°C with vigorously shaking. At the end of the incubation, five volumes of DMEM F12-HAM were added to neutralize the collagenase, and the suspension was centrifuged at 800 g for 10 min. The resulting pellet, containing ASCs, was suspended in DMEM F12-HAM supplemented with 10% fetal bovine serum, and 100 U penicillin/streptomycin and amphotericin (control medium, CM). The ASCs were initially cultured in CM up to 95% confluence in a humidified atmosphere, 95% air with 5% CO₂ at 37°C. The adherent cells were trypsinized and 1 x 10⁵ ASCs per 100-mm² tissue culture plate were seeded in flasks. These passages were repeated three times. To phenotypically characterize ASCs and to define their purity, FACS analysis was performed as described.³⁸

Cell culture conditions: unstimulated and PEMF stimulated

A suspension (5 x 10⁵) of both types of stem cell in 200 μL of proliferative medium (PM) was seeded in 24-well tissue culture plates and allowed to attach overnight. Then, the medium was replaced with osteogenic medium (OM) and changed twice per week (RT) and incubated with phalloidin (Alexa-Fluor-488 phalloidin, Invitrogen) for 20 min. Then the cells were further incubated overnight at 4°C with the primary and antivinculin clone (hVIN-1, Sigma-Aldrich). Finally, after staining with Alexa-Fluor-633–conjugated secondary antibodies (Invitrogen) for 1 h at RT, samples were mounted and nuclei were counterstained with Hoechst (Sigma Aldrich).

Adhesion and morphological analysis. For adhesion and morphological analysis, paraformaldehyde fixed cells were permeabilized with 0.1% Triton X-100 for 1 h at room temperature (RT) and incubated with phalloidin (Alexa-Fluor-488 phalloidin, Invitrogen) for 20 min. The “electromagnetic culture” was stimulated with PEMF for 5, 10, and 30 min and 1, 4, and 8 h per day for 7 days in PM, whereas the “static culture” or non–PEMF-stimulated culture was incubated for the same amount of time in PM in standard well-plates culture placed in a CO₂ incubator far from the electromagnetic bioreactor.

The osteogenic differentiation of both types of stem cell lineages was performed for 21 days with or without PEMF exposure in OM.

3-(4,5-dimethylthiazole-2-yl)-2,5-diphenyl tetrazolium bromide test

To evaluate the mitochondrial activity of both cultured cell conditions, a test with 3-(4,5-dimethylthiazole-2-yl)-2,5-diphenyl tetrazolium bromide (MTT; Sigma-Aldrich, St. Louis, MO), was performed on day 7 in PM as previously reported⁴⁰ on the unstimulated and PEMF-stimulated cultures at different exposure times. The viability assay was also performed on days 7, 14, and 21 (end of the culture period) on the unstimulated and PEMF-stimulated cultures of both cell types. Aliquots of 200 μL were sampled, and the related absorbance values were measured at 570 nm by a microplate reader (BioRad Laboratories, Hercules, CA). A standard cell viability curve was used and the results were expressed as percentage referred to both type of unstimulated cells, respectively.

Apoptosis

Apoptosis is defined as programmed, physiological cell death and plays an important role in tissue homeostasis. The Annexin V technique detects apoptosis by targeting the loss of phospholipid asymmetry in the plasma membrane. The loss of plasma membrane asymmetry is an early event in apoptosis, independent of cell type, resulting in the exposure of phosphatidylserine (PS) residues at the outer plasma membrane leaflet.⁴¹

The Annexin V-FITC Apoptosis Detection Kit (Bender Medsystems, Vienna, Austria) was used according to the manufacturer’s instructions.

DNA content

At the end of incubation, the cells were lysed by a freeze-thaw method in sterile deionized distilled water. The released DNA content was evaluated with a fluorometric DNA quantification kit (PicoGreen; Molecular Probes, Eugene, OR). A DNA standard curve⁴⁰ obtained from a known amount of osteoblasts was used to express the results as cell numbers per well.

Confocal laser scanning microscopy analysis

For immunological studies, paraformaldehyde solution in 0.1 M phosphate buffer (pH 7.4) for 8 h at room temperature and washed with phosphate-buffered saline (PBS) three times for 15 min. Then, the unstimulated and PEMF-stimulated cells were treated as described in the following sections.

Immunological studies. For immunological studies, paraformaldehyde fixed cells were blocked by incubating with PAT (PBS containing 1% [w/v] bovine serum albumin and 0.02% [v/v] Tween 20) for 2 h at room temperature and washed. Anti–type-I and -III collagens, antidecorin, anti-osteopontin, anti-osteocalcin, anti-osteonectin, and anti–alkaline phosphatase (anti-ALP) rabbit polyclonal antisera were used as the primary antibodies diluted to 1:500 in...
PAT. The same dilution was used with anti-fibronectin (anti-FN) rabbit polyclonal IgG. The incubation with the primary antibodies was performed overnight at 4°C, whereas the negative controls were incubated with PAT alone. The samples and the negative controls were washed and incubated with Alexa-Fluor-488 goat antirabbit IgG (H&L; Invitrogen) at a dilution of 1:750 in PAT for 1 h at room temperature. At the end of the incubation, the samples were washed in PBS, counterstained with a Hoechst solution (2 μg/mL) to target the cellular nuclei, and then washed.

The images were taken by the TCS SPII confocal microscope (Leica Microsystems, Bensheim, Germany) equipped with a digital image capture system at 40× magnification.

Scanning electron microscopy analysis

Both types of cell were also seeded on plastic cell culture coverslip disks (Thermanox Plastic, Nalge Nunc International, New York, NY) and cultured as previously indicated. After 3 weeks of culture, cells were fixed with 2.5% (v/v) glutaraldehyde solution in 0.1 M sodium cacodylate buffer (pH 7.2) for 1 h at 4°C, washed with sodium cacodylate buffer, and then dehydrated at room temperature in an ethanol gradient series up to 100%. The samples were kept in 100% ethanol for 15 min, and then critical-point dried with CO2. The specimens were spatter-coated with gold and observed at 50× and 1000× magnification respectively with a Leica Cambridge Stereoscan 440 microscope (Leica Microsystems, Bensheim, Germany) at 8 kV.42

Gene expression analyses

Total RNA from the stimulated samples (treat) and the nonstimulated samples (control) was extracted with the Trizol reagent (Invitrogen) and retrotranscribed into cDNA with the iScript cDNA Synthesis Kit (BioRad Laboratories) as previously reported.42

Quantitative reverse-transcription polymerase chain reaction (qRT-PCR) analysis was performed in a 48-well optical reaction plate using a MiniOpticon Real-Time PCR System (BioRad Laboratories). Oligonucleotide primers were designed with gene sequences published in GenBank and are indicated in Table 1. Reactions were performed in 20 mL with 2 mL of cDNA, 10 mL Brilliant SYBER Green qPCR Master Mix (Stratagene, La Jolla, CA), 0.4 mL of each primer, and 7.2 mL H2O. PCR conditions were as follows: 3 min at 95°C, 30 cycles of 5 sec at 95°C, and 23 sec at 60°C. Gene expression was normalized to the GAPDH housekeeping gene expression. Each sample was analyzed in triplicate and correlated against a standard curve. The reaction mixture, without cDNA, was used as a negative control in each run.

### Purified proteins

Decorin, type-I collagen, and fibronectin were purified as described previously42; osteocalcin was acquired from Biomedical Technologies, Inc. (Stoughton, MA), osteopontin and osteonectin were obtained from Assay Designs, Inc. (Ann Arbor, MI); and type-III collagen and ALP were purchased from Sigma-Aldrich, Inc.

### Polyclonal antisera

Dr. Larry W. Fisher (http://csdb.nidcr.nih.gov/csdb/antisera.htm, National Institutes of Health, Bethesda, MD) provided us with the rabbit polyclonal anti-type-I and -III collagen, antidecorin, anti-osteopontin, anti-osteocalcin, anti-osteonectin, and anti-ALP. Polyclonal antibody against human fibronectin was produced as previously described.40

### Extraction of ECM proteins from the cultured scaffolds and enzyme-linked immunosorbent assay

At the end of the culture period, in order to evaluate the amount of ECM produced, the samples were washed extensively with sterile PBS to remove culture medium, and then incubated for 24 h at 37°C with 1 mL of sterile sample buffer (20 mM Tris-HCl, 4 M GuHCl, 10 mM EDTA, 0.066% [w/v] sodium dodecyl sulfate [SDS], pH 8.0). At the end of the incubation period, the total protein concentration in both culture systems was evaluated with the BCA Protein Assay Kit (Pierce Biotechnology, Inc., Rockford, IL). The total protein concentration was 90.1 ± 7.1 μg/mL for the unstimulated BM-MSCs and 118.5 ± 8.2 μg/mL for the PEMF-stimulated BM-MSCs, 85.7 ± 2.5 μg/mL for the unstimulated ASCs and 105.0 ± 3.3 μg/mL for the PEMF-stimulated ASCs.

Calibration curves to measure type-I and -III collagens, decorin, osteopontin, osteocalcin, osteonectin, fibronectin, and APT were performed as previously described.39 In order to measure the ECM amount of each protein an enzyme-linked immunosorbent assay (ELISA) was performed as previously reported.42

We have taken into consideration that an underestimation of the absolute protein deposition is possible because the sample buffer, used for matrix extraction, contains SDS, which may interfere with protein adsorption during the ELISA. The amount of ECM constituents from both samples was expressed as pg/(cells per well).

### Quantification of calcium

To evaluate the calcium deposition, fluorescent calcein detection and calcium–cresolphthalein complexone methods were performed as described previously.42

### Calcein.

At the end of cell incubation, each sample was rinsed with sterile PBS and stained with a calcein solution.

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### Table 1. Primers Used for Quantitative Reverse-Transcription Polymerase Chain Reaction

| Gene | Upstream primer | Downstream primer |
|------|-----------------|-------------------|
| BOSP | 5'-GGG CAG TAG TGA CTC ATC CG-3' | 5'-TCA GCC TCA GAG TCT TCA TCT TC-3' |
| GAPDH | 5'-AGC CTC AAG ATC ATC AGC AAT GCC-3' | 5'-TGTT GGT CAT GAG TCC TTC CAC GAT-3' |
| OP | 5'-GTC ATT TGC TTT TGC CTC CT-3' | 5'-GCC ACA GCA TCT GGG TAT TT-3' |
| RUNX2 | 5'-ACA GTA GAT GGA CCT CGG GA-3' | 5'-ATA CTG GGA TGA GGA ATG CG-3' |
| OSN | 5'-CTT CAG ACT GCC CGG AGA-3' | 5'-GAA AGA AGA TCC AGG CCC TC-3' |
(5 mM in PBS; Invitrogen, Carlsbad, CA) for 30 min at 22°C. The samples were counterstained with a Hoechst solution (2 μg/mL) to target the cellular nuclei, and then washed with PBS. The images were taken by the TCS SPII confocal microscope (Leica Microsystems) equipped with a digital image capture system at 40× magnification.

Calcium–cresolphthalein complexone method. In order to evaluate calcium deposition, the calcium–cresolphthalein complexone method was performed as previously described on cells cultured with or without PEMF. Briefly, the calcium content of each sample was assayed to quantify the amount of mineralized matrix present, using a Calcium Fast kit (Mercury SPA, Naples, Italy) according to the manufacturer’s instructions as previously described. Samples were run in triplicate and compared with the calibration curve of standards.

ALP activity

ALP activity was determined using a colorimetric end point assay. The assay measures the conversion of the colorless substrate p-nitrophenol phosphate (PNPP) by the enzyme ALP to the yellow product p-nitrophenol; the rate of color change corresponds with the amount of enzyme present in solution. The test was performed as previously described on cells cultured with or without PEMF stimulation. Samples were run in triplicate and compared with the calibration curve of p-nitrophenol standards. The enzyme activity was expressed as micromoles of p-nitrophenol produced per minute per milligram of enzyme.

Statistics

Each experiment was repeated three times. Results are expressed as the mean ± standard deviation. In order to compare the results between the two types of sample unstimulated or PEMF-stimulated, the one-way analysis of variance (ANOVA) with post hoc Bonferroni test was applied, with a significance level of 0.05.

Results

The isolated, expanded, and characterized human stem cells harvested from BM aspirates or obtained surgically from subcutaneous adipose tissue of pediatric donors were seeded onto 24-well culture plates and then cultivated in proliferative culture conditions in the absence (control) or presence of an electromagnetic bioreactor for 7 days (Fig. 1). To determine the best exposure time for PEMF stimulation, an MTT test was performed at day 7 on both stem cell cultures stimulated for 5, 10, and 30 min and 1, 4, and 8 h/day with a PEMF in proliferative medium. The controls were represented by unstimulated BM-MSCs and ASCs and were set as 100% cell viability. The data were represented as a percentage of the control. Bars indicate mean values ± standard error of the mean of results from three experiments.

FIG. 1. Effect of pulsed electromagnetic field (PEMF) exposure on bone marrow mesenchymal stem cell (BM-MSC) and adipose tissue mesenchymal stem cells (ASC) viability. A cell viability assay was performed at day 7 of culture stimulating both types of stem cells for 5, 10, and 30 min and 1, 4, and 8 h/day with a PEMF in proliferative medium. The controls were represented by unstimulated BM-MSCs and ASCs and were set as 100% cell viability. The data were represented as a percentage of the control. Bars indicate mean values ± standard error of the mean of results from three experiments.

Genic culture conditions are observed to express genes and proteins associated with the osteoblast phenotype. As indicated in Figure 1, the two MSCs lineages were exposed to PEMF for 10 min per day and compared to unstimulated cells. This culture system allowed the study of both stem cell types as they proliferated and differentiated to osteoblasts producing a calcified ECM in an inactive or electromagnetically active environment. We compared the cell viability, cell matrix distribution, and calcified matrix production of the unstimulated and PEMF-stimulated culture systems for both types of mesenchymal cell lineages.

Cell viability and morphology

On days 7 and 14 and at the end of the culture (on day 21), the average cell viability for the unstimulated or PEMF stimulated was in the 88%–95% range with no statistically significant differences between cell types, unstimulated or PEMF-stimulated (p > 0.05). The percentage of viability was as follows: on days 7, BM-MSCs + PEMF, 88% ± 6.1%; ASCs + PEMF, 90% ± 8.2%; on day 14, BM-MSCs + PEMF, 94% ± 9%; ASCs + PEMF, 93% ± 7.5%; and on day 21, BM-MSCs + PEMF, 95% ± 9%; ASCs + PEMF, 94.4% ± 7.8%. Additional experiments were then performed.

After 21 days in comparison with unstimulated cells, greater proliferation of both cell types was observed with PEMF exposure: the measurement of the DNA content in the static culture showed an increase in cell number per well from 3.7 × 10⁴ ± 8.2 × 10² up to 5.1 × 10⁴ ± 8.4 × 10² (p < 0.05) in the dynamic culture for BM-MSC-derived cells. A similar increase was observed for ASC-derived cells: from 4.2 × 10⁴ ± 8.2 × 10² up to 5.4 × 10⁴ ± 8.4 × 10² (p < 0.05) cell number per well. Because the DNA may remain entrapped in the calcified matrix, an underestimation of culture cellularity is possible.
To determine whether PEMF treatment provoked cell apoptosis, Annexin V and propidium iodide (PI) staining was performed. On the same days as the MTT test, confocal laser scanning microscopy (CLSM) analysis was performed on both unstimulated and PEMF-stimulated cells. It is known that in the early stages of apoptosis, the plasma membrane excludes viability dyes such as PI, therefore cells that display only Annexin V staining (PI negative) are in the early stages of apoptosis. During late-stage apoptosis, loss of cell membrane integrity allows Annexin V binding to cytosolic PS, as well as cell uptake of PI. In view of this, CLSM analysis of unstimulated and PEMF-stimulated BM-MSCs and ASCs were negative after Annexin V and PI staining (data not shown), indicating that the PEMF stimulation did not induce apoptosis in either type of cell.

To qualitatively evaluate cell viability and morphology, cells were also observed by CLSM (Fig. 2B) and by scanning electron microscopy (SEM; Fig. 2A) at the end of the culture period. CLSM observations revealed an equivalent cell morphology in both types of culture condition (Fig. 2B), suggesting that the PEMF stimulation did not alter cell viability or proliferation. In particular, stimulated cells (Fig. 2B-ii, iv) showed better cytoskeleton organization with F-actin-containing fibers arranged as straight, cable-like cords transversing the cytoplasm.

Figure 2A is a representative SEM image of 21 days in cell culture showing adherence and morphology of both types of cell, unstimulated and PEMF-stimulated. In particular, independently of the stem cell source, osteoblasts grown in PEMF (Fig. 2A-ii, iv) homogeneously covered the wells in comparison with unstimulated cells (Fig. 2A-i, iii). At higher magnification, no significant morphological differences for either cell type or culture condition were observed: cells were flat and compact.

**Characterization of gene expression in bone**

To characterize the gene expression in both types of cell and culture system condition, RT-PCR and qRT-PCR analyses were performed after 7 and 21 days of culture. On day 7, the qualitative RT-PCR and qRT-PCR did not show a significant difference in the gene expression levels for ECM proteins in either cell or culture system condition (data not shown). The main difference in gene expression levels between cell types and culture conditions was observed after 21 days of culture.

At the end of the cell culture period, the qRT-PCR performed on both cell types and culture system condition showed some differences. It revealed a bright band on the agarose gel for all indicated genes and particularly for the transcripts specific for bone sialoprotein, osteopontin, osteonectin (data not shown).

To further examine these data, a qRT-PCR for gene expression profiles of bone-specific proteins and transcriptional factor Runx2 was performed after 21 days using the ΔΔCt method. The results showed a significantly enhanced fold difference for bone sialoprotein (BOSP), osteopontin (OP), osteonectin (OSN), and Runt-related transcription factor 2 (Runx2) \( (p<0.001; \text{Fig. 3A}) \) in PEMF-stimulated cells of BM origin in comparison with the unstimulated cells. No evident fold difference was detected for gene expression profiles of the same proteins between the unstimulated and PEMF-stimulated cells deriving from human adipose tissue \( (p>0.05; \text{Fig. 3B}) \).

**Characterization of calcified ECM deposition**

At the end of the cell culture, immunolocalization of osteopontin and osteocalcin showed a more intense green fluorescence in osteoblasts differentiated from BM-MSC and exposed to PEMF (Fig. 4A-ii and B-ii), revealing the stimulatory effect of the physical stimulation in terms of higher cell proliferation and more intense fluorescent staining of the ECM. Unstimulated cells were few and surrounded by a thin and discontinuous ECM (Fig. 4A-i, B-i). No difference in fluorescence intensity was observed between unstimulated (Fig. 4A-iii, B-iii) and PEMF-stimulated cells (Fig. 4A-iv, B-iv) both originating from ASCs.

Regarding the origin of the cell source for the differentiated cells, the immunolocalization of other proteins such as type-I

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**FIG. 2.** Effect of PEMF exposure on osteoblast morphology differentiated from BM-MSCs and ASCs after 21 days of incubation, as determined by scanning electron microscopy (SEM) (A) and confocal laser scanning microscopy (CLSM) (B) analysis. SEM and CLSM observations confirmed equivalent cell morphology in both culture conditions, suggesting that PEMF stimulation did not alter cell viability or proliferation. SEM images (A) are at 200× magnification (insets, 1000×); scale bars represent 100 μm (insets, 10 μm). CLSM images (B) are at 40× magnification; scale bar represents 50 μm.
collagen, type-III collagen, osteonectin, or FN showed similar patterns (data not shown).

To evaluate the amount of ECM constituents produced by both types of cell in unstimulated or PEMF-stimulated system conditions, an ECM extraction was performed on days 7 and 21. Unfortunately, on day 7 even if the total protein content was determined, the levels of the specific bone proteins were too low to be detected in either sample types. At the end of the culture period, the detection of bone proteins showed some differences. In comparison with the unstimulated culture system, the deposition of bone proteins in the PEMF-treated cells differentiated from BM-MSCs was considerably enhanced ($p < 0.05$) (Table 2). In contrast, this protein deposition was not as marked for osteoblasts differentiated from PEMF-treated ASC compared with unstimulated ASC ($p > 0.05$; Table 2). These data are in accordance with the immunofluorescence analysis performed on unstimulated and PEMF-stimulated culture systems for both cell types (Fig. 4).

Protein deposition was particularly enhanced for ALP, which was 1.7-fold greater compared with unstimulated cells derived from BM-MSCs (Table 2). Again, this difference was not significant between unstimulated and PEMF-stimulated cells differentiated from ASCs. Figure 5 shows the ALP activity measured in both cell types at the end of the culture period: the level of ALP activity was consistently higher in cells differentiated from BM-MSCs and treated with PEMF than in unstimulated cells ($p < 0.001$; Fig. 5B). In contrast, the ALP level was not different in cells of ASC origin, PEMF-stimulated or not (Fig. 5B). These data for cells differentiated from BM-MSCs or ASCs are in accordance with the immunofluorescence analysis for osteocalcin (Fig. 4B) and type-I collagen (Fig. 4A) for both cell types (Fig. 5A).

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**FIG. 3.** Effect of PEMF exposure on bone gene expression of the indicated bone-specific markers as determined by quantitative reverse transcriptase–polymerase chain reaction. (A) Representative image of osteoblasts differentiated from BM-MSCs. (B) Cells differentiated from human adipose tissue. The graph shows the fold induction of gene expression expressed in arbitrary units setting the expressions of the indicated genes in cells grown in absence of PEMF as equal to 1. *$p < 0.001$ was considered statistically significant; # $p > 0.05$. Data are representative of one of three experiments performed.

**FIG. 4.** Effect of PEMF exposure on BM-MSC and ASC osteogenic differentiation as determined by immunolocalization of type-I collagen (A) and osteocalcin (B) at day 21 of incubation at 40× magnification; scale bar represents 50 μm. The dense layer of bone extracellular matrix was clearly evidenced by an intense fluorescence signal in the PEMF stimulated cells derived from BM-MSCs (ii) in comparison with unstimulated BM-MSCs (i) or unstimulated ASCs (iii)/PEMF stimulated (iv) but derived by ASCs.
The relative amount of calcium for both types of cells and culture system condition was evaluated by the calcine method (Fig. 6A) and quantified by the calcium-cresolphthalein complexone method (Fig. 6B) to evaluate matrix calcification. Figure 6 shows a qualitative assessment of cell and calcium deposition by both cell types in unstimulated and PEMF-stimulated culture conditions with some significant differences: the mineralization of ECM produced by osteoblasts gave a more intense green fluorescence signal colocalized within the cells on PEMF-stimulated than on unstimulated BM-MSCs or PEMF-stimulated cells originating from ASCs. The qualitative evaluation of calcium deposition was confirmed: calcification of the deposited ECM was considerably greater in PEMF-stimulated cells of human BM origin (Fig. 6A-ii) than in unstimulated controls (Fig. 6A-i; p < 0.05). In contrast, no difference was observed between unstimulated and PEMF-stimulated cells of human adipose tissue origin (Fig. 6A-iii, iv).

**Discussion**

Research oriented at isolation of stem cells, their culture for purposes such as developing cell or tissue therapies, differentiation studies, the comprehension of the factors necessary to direct cell specialization to specific pathways, and other developmental studies is an absolute priority in tissue engineering approach. The increase in differentiation observed for both types of MSCs independent of their tissue origin could be of interest in other clinical settings or in a tissue engineering approach.

In a previous study by our group we showed that cell stimulation using an electromagnetic bioreactor (magnetic field intensity, 2 mT; frequency, 75 Hz) significantly increased SAOS-2 human osteoblast proliferation and bone ECM deposition.36 To the best of our knowledge, the PEMF effects either in stem cells isolated from adipose tissue and/or comparative studies with BM-MSCs have not been investigated. To this end, the main aim of this study was to show the greater efficacy of PEMF exposure on the osteogenic differentiation of BM-MSCs in comparison with ASCs, by evaluating cell proliferation, morphology, and calcified matrix deposition. The increase in differentiation observed for both types of MSCs independent of their tissue origin could be of interest in other clinical settings or in a tissue engineering approach.

In a previous study by our group we showed that cell stimulation using an electromagnetic bioreactor (magnetic field intensity, 2 mT; frequency, 75 Hz) significantly increased SAOS-2 human osteoblast proliferation and bone ECM deposition.36 In this study, we investigated the effects of electromagnetic stimulation on osteogenesis of stem cells isolated from BM and adipose tissue. When we performed
experiments for the same length of time as that utilized for SAOS-2 cell stimulation, the BM-MSCs and ASCs did not survive. In fact, the best results in terms of cell proliferation were achieved by stimulating with the electromagnetic bioreactor both stem cells for 10 min a day with the same physical parameters. Even though, the SAOS-2 cell line is useful to screen different types of culture condition, primary cells (osteoblasts or MSCs) are more relevant and should be used to assess in vitro the best physical parameters for electromagnetic stimulation to increase or improve biological mineralization.36

A temporal and functional pattern of gene expression characterizes the osteoblast maturation process, which can be divided into proliferation, differentiation, and mineralization stages,34 in turn the effects of a pulsed electromagnetic field depend on the maturation stages of the osteoblasts.35,45 PEMF-stimulated osteoblasts of BM origin exhibited a more evident increase in cell proliferation than those of adipose tissue origin. These results are in line with previous data demonstrating that application of PEMF to BM-MSCs cultured in OM results in early onset of cell proliferation and, consequently, higher cell densities.35 PEMF not only affects osteoblast cellular proliferation and differentiation of bone cells in vitro by enhancing DNA synthesis,34,45,46 but also increases expression of bone marker genes during differentiation and mineralization and enhances calcified matrix production.34 In order to study the effects of PEMF on ECM deposition for both types of culture condition, the fundamental bone matrix constituents such as type-I and -III collagen, decorin, osteopontin, osteocalcin, osteonectin, and fibronectin were investigated. Both types of PEMF-stimulated cells were able to significantly increase ECM deposition with respect to unstimulated cells. In a comparative quantitative protein analysis between BM-MSCs and ASCs we observed some differences. In both cell types exposed to PEMF, the deposition of type-I and -III collagen and decorin was approximately 1.4-, 1.2-, and 1.1-fold greater, respectively, in wells seeded with BM-MSCs as determined by CLSM (A) (40 × magnification; scale bar represents 50 μm) and by quantification of calcium content (B). Results are expressed on a per-scaffold basis and are presented as an average ± standard deviation (*p < 0.05).

All together these results suggest that the main effect of cell exposure to PEMF is a greater efficiency in promoting bone ECM deposition by osteoblasts differentiating from BM-MSCs.

Interestingly, the qRT-PCR analysis showed an increase in bone sialoprotein, osteopontin, osteonectin, and Runx2 gene expression levels from both types of PEMF-stimulated osteoblasts, at a higher level for BM-MSC differentiated cells in comparison with adipose tissue maturated stem cells. Since osteonectin and osteopontin are critical in mediating the signal cascade for the full expression of the mature osteoblast phenotype and mineralization of the ECM,55,56 the higher OSN expression in both cell types could be related to the cell’s ability to better differentiate toward mature osteoblasts and to deposit mineralized bone matrix as a result of PEMF exposure. This concept is also true for BOSP, which is a significant component of the bone ECM and is suggested to constitute approximately 8% of all noncollagenous proteins found in bone and cementum. Furthermore, we decided to evaluate the gene expression level of Runx2, also known as core binding factor 1 (cbfa1; subsequently renamed Runx2 as runt-related transcription factor 2).57 Runx2 is a transcriptional factor involved in osteoblastic and skeletal morphogenesis. Runx2 has an essential role in the maturation of osteoblasts by binding to its target promoters and enhancers of various other bone-specific target genes, including collagen type-I, osteopontin, and bone sialoprotein through its runt homology domain.58,59

The Runx2 gene expression in both types of unstimulated cell perfectly correlates with the cell’s
ability to differentiate towards osteoblasts. However, Runx2 expression was more enhanced by PEMF exposure in BM-MSCs than ASCs. Since Runx2 is considered the transcriptional target of Wnt/b-catenin pathway, its up-regulation in both types of stem cells during PEMF exposure may due to the activation of the Wnt/b-catenin signaling pathway. However, the electromagnetic application seems to show a stimulatory effect in osteoblasts differentiated from ASCs, but slightly lower when compared with osteoblasts originating from BM-MSCs. Following PEMF exposure, the Wnt/b-catenin signaling pathway is possibly activated in BM-MSCs but other signaling pathways may be stimulated in ASCs. Recently, it was shown that the exposure of ASCs cells to a 45-Hz electromagnetic field induced osteogenic marker expression via bone morphogenetic protein, transforming growth factor β, and Wnt signaling pathways based on microarray analyses.

In summary, the increased transcription of these genes could indicate that PEMF exposure drives a more rapid commitment of stem cells toward osteoblastic differentiation, in particular for BM-MSCs, but to a lower extent for ASCs.

The increase in transcript levels of the BOSP, OCN, and OP genes was supported by the mineralization data. Quantitative analysis of the calcium mineral content showed that both types of PEMF-stimulated cells were able to deposit a significantly higher amount of newly mineralized matrix compared with unstimulated cells. In a quantitative comparative analysis, an approximate 1.5-fold difference in matrix calcification was detected for BM-MSC-derived osteoblasts with respect to ASCs. Furthermore, confocal microscopic analysis evidenced calcification and bone matrix proteins in the cell-rich area for both types of cells, with a stronger fluorescence signal for cells differentiated from BM.

In this study, the increase in calcium deposition was consistent with the rise in ALP expression in both types of PEMF-exposed MSCs. In a comparison analysis with ASC-derived osteoblasts, the protein content was 1.3-fold greater for cells differentiated from BM-MSCs. ALP activity also increase by almost 2-fold. ALP is recognized as an important component in hard tissue formation and is highly expressed in mineralized tissue cells. Osteoblast cells grown with ascorbic acid sequentially express osteoblastic marker proteins such as ALP and then form a mineralized ECM as a consequence of osteoblastic differentiation. The importance of ALP in the mineralization of ECM has been previously reported. Its mechanism of action is not completely known, but it appears to promote both the local concentration of inorganic phosphate, a mineralization promoter, and to decrease the concentration of extracellular pyrophosphate, an inhibitor of mineral formation. Previous studies have reported that ALP is expressed in large amounts in osteoblasts in vivo. The elevated expression of ALP (which occurs at the end of the cell proliferative state), osteopontin and bone sialoprotein may suggest that the MSC-derived osteoblasts are more differentiated than ASC osteoblasts and have already started to promote bone ECM deposition.

Although ASCs and BM-MSCs share many biological characteristics, there are some differences in their immunophenotype, differentiation potential, transcriptome, proteome, and immunomodulatory activity. A possible explanation of why BM-MSCs showed a more rapid osteogenic differentiation than ASCs when treated with PEMF could be related to their differences such as specific features of BM-MSCs and ASCs, or it could be due to the inherent heterogeneity of both BM-MSC and ASC populations or to a different activation of a signaling pathway.

In summary, these results suggest that PEMF enhances the commitment of BM-MSCs to osteoblasts more efficiently in comparison with ASCs. Both types of mesenchymal stem cells are osteogenic (Table 2), but in an tissue engineering approach, which combines biomaterials, growth factors, and cells, the ASC exposure to PEMF seem to be less effective. Furthermore, PEMF in vitro studies should be considered as groundwork for in vivo bone development that may support skeletal therapy. Taken all together, these findings may provide insights on the development of PEMF as an effective technology for regenerative medicine.

Acknowledgments

This work was supported by the Compagnia S. Paolo (Genova) Grant titled “Uso di cellule staminali mesenchimali sotto-poste a campo elettromagnetico: un approccio innovativo per l’osteointegrazione di impianti di titanio” (2011) to L.V. and by an INAIL grant entitled “Effetti dei campi elettromagnetici sulla salute umana: modelli sperimentali in vitro” (2011) to G.C. We are grateful to P. Vaghi (Centro Grandi Strumenti, University of Pavia) and D. Picenoni (Politecnico di Milano, Milano, Italy) for technical assistance in the CLSM and SEM studies, respectively. A special thanks to Laurene Kelly (University of Pavia) for correcting the English in the manuscript.

Author Disclosure Statement

All the authors state that no competing financial interests exist.

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