Osteogenic commitment and differentiation of human mesenchymal stem cells by low-intensity pulsed ultrasound stimulation

Viviana Costa1 | Valeria Carina1 | Simona Fontana2 | Angela De Luca1 | Francesca Monteleone2 | Stefania Pagani3 | Maria Sartori4 | Stefania Setti5 | Cesare Faldini6 | Riccardo Alessandro2 | Milena Fini3,4 | Gianluca Giavaresi1,3

1 Rizzoli Orthopedic Institute, Innovative Technological Platforms for Tissue Engineering, Theranostic and Oncology, Palermo, Italy
2 Biology and Genetics Unit, Department of Biopathology and Medical Biotechnologies, University of Palermo, Italy
3 Rizzoli Orthopedic Institute, Laboratory of Preclinical and Surgical Studies, Bologna, Italy
4 Rizzoli Orthopedic Institute, Laboratory BITTA, Bologna, Italy
5 IGEA SpA, Carpi (Modena), Italy
6 Rizzoli Orthopedic Institute, 2nd Orthopaedic and Traumatologic Clinic, Bologna, Italy

Correspondence
Viviana Costa, PhD, Rizzoli Orthopedic Institute, Via Divisi 83, Palermo (ITALY) 90133.
Email: viviana.costa@lor.it

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INTRODUCTION

Mesenchymal Stem Cells (MSCs) are multipotent adult stem cells commonly used for tissue engineering applications, which have been demonstrated to differentiate into osteoblasts, adipocytes, or chondrocytes depending on the culture condition (Alvarez, Lee, Wang, & Hong, 2015; McNamara et al., 2010). MSCs are characterized by self-renewal capability, the process by which stem cells proliferate and create further stem cells, maintaining their undifferentiated, and multipotent state (Kuhn & Tuan, 2010; Pittenger et al., 1999).

Low-intensity pulsed ultrasound (LIPUS) as an adjuvant therapy in in vitro and in vivo bone engineering has proven to be extremely useful. The present study aimed at investigating the effect of 30 mW/cm² LIPUS stimulation on commercially available human mesenchymal stem cells (hMSCs) cultured in basal or osteogenic medium at different experimental time points (7, 14, 21 days). The hypothesis was that LIPUS would improve the osteogenic differentiation of hMSC and guaranteeing the maintenance of osteogenic committed fraction, as demonstrated by cell vitality and proteomic analysis. LIPUS stimulation (a) regulated the balance between osteoblast commitment and differentiation by specific networks (activations of RhoA/ ROCK signaling and upregulation of Ribosome constituent/Protein metabolic process, Glycolysis/Gluconeogenesis, RNA metabolic process/Splicing and Tubulins); (b) allowed the maintenance of a few percentage of osteoblast precursors (21 days CD73+/CD90+: 6%; OCT-3/4+/NANOG+/SOX2+: 10%); (c) induced the activation of osteogenic specific pathways shown by gene expression (early: ALPL, COL1A1, late: RUNX2, BGLAP, MAPK1/6) and related protein release (COL1a1, OPN, OC), in particular in the presence of osteogenic soluble factors able to mimic bone microenvironment. To summarize, LIPUS might be able to improve the osteogenic commitment of hMSCs in vitro, and, at the same time, enhance their osteogenic differentiation.

KEYWORDS
low-intensity pulsed ultrasound, mesenchymal stem cells, osteogenic commitment, osteoblast differentiation, proteomic analysis

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The way stem cells participate in tissue generation, maintenance, and repair, saving themselves from depletion and protecting the host from over-exuberant stem-cell proliferation, is regulated by a specific anatomic entity defined by Schofield in 1978 and described by Scadden in 2006 as stem-cell niche (Scadden, 2006; Schofield, 1978). Stem-cells receive specific signals from both physical and biochemical clues of this niche environment by direct contact with basal membrane proteins and by binding with soluble mediators like growth factors and cytokines, or insoluble extracellular matrix proteins, acting like paracrine regulators of stem-cell function (Griffin, Butler, Seifalian, & Kalaskar, 2015).

Alternatively, adult stem or progenitor cells, such as MSCs, represent a promising resource for new bone formation via cell-based tissue engineering. These cells can be isolated from various adult tissues and are able to promote tissue repair through trophic effects on other cell types and immune-regulatory activities (Griffin et al., 2015). Until now, tissue engineering approaches were characterized by the use of hMSCs isolated by different tissues, alone or in combination with a biomimetic scaffold and/or a physical approach (such as ultrasound, pulsed electromagnetic fields or others), in order to accelerate bone restoration (Baker, Sohn, & Tuan, 2015).

Therefore, it is fundamental to understand how to control and guide the fate of stem cells in culture by providing external stimuli, that is using specific surface topography and chemistry with the help of biomimetic biomaterials, or delivering chemical, and genetic clues in the form of growth factors and cytokines (Griffin et al., 2015; Papadimitropoulos et al., 2014). The loss of MSC properties in vitro, probably due to the absence of the niche environment, might limit the utility of MSCs in cell-based regenerative medicine and long-term LIPUS stimulation was analyzed in terms of (a) changes in the expression of protein markers, through MS-based quantitative proteomic methods, it was attempted to define the expression profiles of membrane proteins in MSCs undergoing osteoblast differentiation, illustrating the effects of many growth factors on this process (Foster et al., 2005; Kratchmarova, Blagoev, Haack-Sorensen, Kassem, & Mann, 2005; Salaszyk et al., 2005). A comparison of the protein expression profiles in undifferentiated and fully differentiated hMSC might reveal expressional differences that provide important clues into the mechanism underlying the process from stem cells to osteoblasts.

The overall purpose of this study was to verify whether LIPUS stimulation was able to allow the osteoblast lineage commitment of hMSCs and differentiation. To this end, LIPUS with spatially averaged and temporally averaged (SATA) intensity of 30 mW/cm² was applied to hMSCs cultured in basal or osteogenic medium at different experimental time points. The MSC differentiation response to short- and long-term LIPUS stimulation was analyzed in terms of (a) changes in the expression of protein markers, through MS-based quantitative proteomic technologies, in order to identify protein networks; and (b) the modulation of different stem cell markers (CD73, CD90, CD105, OCT-3/4, NANOG, SOX2) and osteogenic pathways; both activated by LIPUS (Kuhn & Tuan, 2010; Liu & Lee, 2014; Pricola, Kuhn, Halem-Smith, Song, & Tuan, 2009; Scherzer et al., 2016; Yoon et al., 2014).

2 | MATERIAL AND METHODS

2.1 | Cell culture

Commercially available Human Mesenchymal Stem Cells (hMSC, Lonza, Walkersville, MD) were cultured in Mesenchymal Stem Cell Growth Medium (MSCGM™ Bullet Kit, Lonza) to expand them without inducing differentiation. The culture medium was changed every 3 days, and cells were split at 80–90% of confluence using StemPro Accutase (Gibco by Life Technologies, Grand Islands, NY). hMSCs were used at an early passage (P5) for all experiments. To obtain osteogenic differentiation, hMSCs were treated with hMSC Mesenchymal Stem Cell Osteogenic Differentiation Medium (OM) (hMSC Osteogenic Differentiation BulletKit™, Lonza).

2.2 | LIPUS treatment

The LIPUS exposure device, manufactured by IGEA SpA (Carpì-Modena, Italy), consists of an array of five transducers...
which are specifically designed for stimulating cell cultures in a 48-well plate (Figure 1). The LIPUS signal consisted of 200 µs burst of 1.5 MHz sine waves repeating at 1 kHz and delivering 30 mW/cm² SATA intensity. Twenty-four hours before LIPUS treatment, hMSC cells were seeded in 48-well plates (diameter = 11.0 mm, surface area = 95 mm², and thickness of the well bottom = 1.2 mm) at different concentrations according to the different time points: 24 hr: 10000 cells/well; 7 days: 8000 cells/well; 14 days: 4000 cells/well; and 21 days: 2000 cells/well. For each experimental time point, hMSC cultures were divided in four groups: (a) Untreated MSCGM group (Control): cells were cultured in MSCGM and not exposed to LIPUS treatment; (b) LIPUS MSCGM group: cells were cultured in MSCGM and treated with LIPUS; (c) Untreated OM group: cells were cultured in OM and not exposed to LIPUS treatment; (d) LIPUS OM group: cells were cultured in OM and treated with LIPUS. Culture plates were then placed on the ultrasound transducer array with a thin layer of standard ultrasound gel and exposed to LIPUS for 20 min/day for five consecutive days/week. The untreated groups were handled in the same way, but the ultrasound generator was switched off. At the end of LIPUS stimulation time, a culture plate for each group was cultured for further 7 days at the same condition in the incubator but without LIPUS exposure (indicated as “off”).

2.3 | hMSC viability (WST-1 test)

WST-1 colorimetric reagent (Roche Diagnostics GmbH, Manheim, Germany) was used to evaluate cell viability. Briefly, WST-1 reagent (10% vol/vol) was added to the cell monolayer in each well. After 4 hr of incubation, the formazan dye produced by viable cells was quantified spectrophotometrically at 450 nm by Bio-Rad Microplate Reader (Bio-Rad Laboratories, Hercules, CA) and results were reported as percentage of viable cells compared to Untreated MSCGM group.

2.4 | DNA concentration (PicoGreen assay)

DNA content was quantified by fluorometry using a Quant-iT PicoGreen dsDNA Assay Kit (Invitrogen™, Life Technologies—EuroClone S.p.A, Pero-Milan, Italy). After sample washing with PBS, 100 µl of lysis solution was added to each well and cell lysis was then completed by three freezes–thaw cycles at −80°C. DNA content was calculated from the lysates by adding 100 µl of fluorescent nucleic acid stain to each.

**FIGURE 1** LIPUS set-up experiment: (a) Top and lateral view of plexiglas transducers support; (b) Characteristic of the ultrasound signal: 200 µs burst of 1.5 MHz sine waves repeated at 1 kHz
sample. Fluorescence was measured using a GloMax multiwell plate reader (GloMax, Promega Corporation Madison, WI).

2.5 hMSC characterization

To demonstrate hMSC phenotype maintenance, CD73, CD90, and CD105 antigen expressions were assessed using a Human MSC Analysis Kit (BD Bioscience 562245), whereas to analyze the stem cell transcriptional factor expression a Human Pluripotent Stem Cell Transcription Factor Analysis Kit (BD Bioscience 560589, Milan, Italy) was used. Briefly, a suspension of $1 \times 10^6$ cells was washed in PBS and re-suspended in flow cytometry buffer (FCB: 2% BSA in PBS) containing 0.5 µg/ml of conjugated antibody directed against CD73, CD90, and CD105, or specific conjugated IgG as isotype control. For the stem cell transcription factors analysis, a manufacturing protocol was followed. Cell fluorescence was evaluated by FlowSight (Amnis Corporation, Merck Millipore) and data were analyzed using a Quantitative Imaging software (Amnis Corporation).

Mineralization of differentiating hMSC cultures was assessed by the OsteoImage™ Mineralization Assay kit (Lonza) based on fluorescent staining of hydroxyapatite deposited by cells in bone-like nodules, which was then measured by a Glomax multiwall plate reader at 492/520 nm excitation/emission.

2.6 Reverse transcriptase–quantitative polymerase chain reaction (RT-qPCR) analysis

Total RNA was extracted using a PureLink™ RNA Micro Kit (Invitrogen™, Pero-Milan, Italy) and reverse-transcribed with a High Capacity cDNA Reverse Transcription Kit (Applied Biosystems™, Life Technologies—EuroClone S.p.A) following the manufacturer’s instructions. Each cDNA sample was tested in duplicate.

qRT-PCR analysis was performed by Light Cycler 2.0 Instrument (Roche Diagnostics) using the SYBR® Green Real-Time PCR Master Mix (Applied Biosystems™), QuantiTect Primers (Qiagen Srl, Milan, Italy) and custom made primers (Invitrogen™) (see Tables 1 and 2). The mean threshold cycle was used for the calculation of relative expression using the $2^{-\Delta\Delta Ct}$ method, against GAPDH as housekeeping gene (Livak & Schmittgen, 2001). Results were expressed as relative fold changes calculated using untreated MSCGM samples as calibrators for each experimental time point.

### TABLE 1 Qiagen gene primers specific for osteogenic differentiation or involved in the differentiating process

| Gene   | Qiagen primers         | Catalog number | Annealing temperature (°C) | PCR products size |
|--------|------------------------|----------------|---------------------------|-------------------|
| RUNX2  | Hs_RUNX2_1_SG QuantiTect primer assay | QT00020517     | 60                        | 101 bp            |
| ALPL   | Hs_ALPL_1_SG QuantiTect primer assay | QT00012957     | 60                        | 110 bp            |
| COL1A1 | Hs.COL1A1_1_SG QuantiTect primer assay | QT00037793     | 60                        | 118 bp            |
| BGLAP  | Hs_BGLAP_1_SG QuantiTect primer assay | QT00232777     | 60                        | 90 bp             |
| SPP1   | Hs_SPP1_1_SG QuantiTect primer assay | QT01008798     | 60                        | 115 bp            |

Their expression was normalized to the GAPDH housekeeping gene (Table 2).

2.7 ELISA assays

Protein release in the culture medium was measured for alkaline phosphatase (ALP kit-SEB472Hu), collagen type I alpha 1 (COL1a1 kit-SEA350Hu), osteopontin (OPN kit-SEA899Hu) together with the cellular content of osteocalcin (OC kit-SEA471Hu) using a CloudClone Corp ELISA KIT assay (Cloud-Clone Corp. Houston, TX), according to the manufacturer’s instructions. The values obtained were normalized to medium protein content evaluated by Bradford assay.

2.8 Proteomic analyses

2.8.1 Protein extraction and digestion

All chemicals used for protein extraction and digestion were of analytical grade, and Milli-Q water was employed in all buffers and solutions. At 14 days (Carina et al., 2017), the cells of all groups were dissolved in 100 µl of 50% tetrafluoroethylene (Sigma–Aldrich) in PBS, vigorously vortexed and sonicated for 7 min in an ice bath and finally incubated with constant shaking for 2 hr at 60°C. Proteins were reduced with 5 mM dithiothreitol (Sigma–Aldrich SRL, Milan, Italy) for 30 min at 60°C and alkylated with 25 mM iodoacetamide (Sigma–Aldrich) for 30 min in the dark at room temperature. Before adding mass spectrometry-grade trypsin (Pierce™ Trypsin Protease, Fisher Scientific Italia, Rodano-Milan, Italy), the samples were diluted fivefold with 100 mM ammonium bicarbonate pH 8.0. Protein samples were digested by adding trypsin at a ratio of 1:50 (w/w) for 18 hr at 37°C with constant shaking, in presence of 2 mM CaCl$_2$. To stop digestion 50 µl of 2.5% trifluoroacetic acid (TFA HPLC Grade, Sigma–Aldrich) were added to the samples. Digested samples were then centrifuged at $14,000 \times g$ for 10 min at 4°C. The resulting supernatant, containing the peptide mixture, was extracted using the 100 µl Bond Elute OMIX C18 pipette tips (Agilent Technologies Italia SpA, Cernusco sul Naviglio—Milan, Italy). Eluted peptide mixtures were vacuum dried and reconstituted in 5% acetonitrile 0.1% formic acid for mass spectrometry analyses. Duplicates of each sample were used for the following proteomic analysis.

2.8.2 Generation of the reference spectral library

Approximately, 2 µg of tryptic peptides of each sample were run for Data-Dependent Acquisition (DDA) analysis. The resulting list of protein/peptides was used for construction of the Sequential Window
Acquisition of all THeoretical (SWATH) reference spectral library. The sample was analyzed via reverse-phase high-pressure liquid chromatography electrospray ionization tandem mass spectrometry (RP-HPLC-ESI-MS/MS) using a TripleTOF® 5600 mass spectrometer (AB SCIEX, Milan, Italy). The mass spectrometer was coupled to a nanoLC Eksigent 425 system (AB SCIEX). RP-HPLC was performed with a trap and elution configuration using an Acclaim™ PepMap™ 100 Nano Trap Column 100 µm × 2 cm, C18, 5 µm, 100 Å (Fisher Scientific Italia) and an Acclaim™ PepMap™ RSLC Nano Column 75 µm × 250 mm, C18, 2 µm, 100 Å (Fisher Scientific Italia). The reverse-phase LC solvents were: solvent A (0.1% formic acid in water) and solvent B (2% water and 0.1% formic acid in acetonitrile). The sample was loaded in the trap column at a flow rate of 5 µl/min for 10 min using a solvent, from loading pump, containing 2% acetonitrile and 0.1% v/v TFA in water and eluted at a flow rate of 300 nI/min using a gradient method according to which solvent B is linearly increased from 2% to 10% within 10 min, from 10% to 30% within 110 min, and then to 60% within 15 min; afterwards, phase B is further increased to 95% within 2 min. Phase B is maintained at 95% for 10 min to rinse the column. Finally, B is lowered to 2% over 2 min and the column re-equilibrated for 21 min (170 min total run time). The eluting peptides were on-line sprayed in the Triple TOF 5600 Plus mass spectrometer, that it is controlled by Analyst TF 1.7 software (AB SCIEX).

Each of the four samples used to generate the SWATH-MS spectral library was subjected to two DDA runs. For these eight experiments, the mass range for MS scan was set to m/z 400–1250 and the MS/MS scan mass range was set to m/z 230–1,500. Using the mass spectrometer, a 0.25 s survey scan (MS) was performed, and the top 50 ions were selected for subsequent MS/MS experiments employing an accumulation time of 0.065 s per MS/MS experiment for a total cycle time of 3.5485 s. Precursor ions were selected in high resolution mode (Δ30,000), tandem mass spectra were recorded in high sensitivity mode (resolution >15,000). The selection criteria for parent ions included an intensity of greater than 500 cps and a charge state ranging from +2 to +5. A 15 s dynamic exclusion was used. The ions were fragmented in the collision cell using rolling collision energy, and a collision energy spread (CES) of 5 V.

Eight DDA MS raw files were combined and subjected to database searches in unison using ProteinPilot™ 4.5 software (AB SCIEX) with the Paragon algorithm by using the following parameters: iodoacetamide cysteine alkylation, digestion by trypsin and no special factors. The search was conducted through identification efforts in a UniProt Swiss-Prot database (http://www.uniprot.org/, downloaded in July 2014, with 137216 protein sequence entries) containing whole Homo sapiens proteins. A false discovery rate analysis was also performed.

### 2.8.3 | SWATH-MS analysis and targeted data extraction

Two replicates of each sample were subjected to the cyclic data independent acquisition (DIA) of mass spectra. Data were acquired by repeatedly cycling through 40 consecutive 15-Da precursor isolation windows (swaths). For these experiments, the mass spectrometer was operated using a 0.1 s survey scan (MS). The subsequent MS/MS experiments were performed across the mass range of 100–1600 m/z on all precursors in a cyclic manner using an accumulation time of 0.03 s per SWATH window for a total cycle time of 1.3490 s. Ions were fragmented for each MS/MS experiment in the collision cell using rolling collision energy, and CES was set to 15. The spectral alignment and targeted data extraction of DIA samples were performed using PeakView v.2.2 (AB SCIEX) with the reference spectral library. All eight DIA files were loaded and exported in .txt format in unison using an extraction window of 15 min and the following parameters: three hundred peptides/protein, seven transitions/peptide, peptide confidence level of 90%, excluded shared and modifies peptides, and an extracting ion current (XIC) width of 75 ppm. This export procedure generated three distinct files containing the quantitative output for (1) the peak area under the intensity curve for individual ions; (2) the summed intensity of individual ions for a given peptide; and (3) the summed intensity of peptides for a given protein. For each protein, seven individual ion intensities were summed as peptide intensity, until three hundred peptides intensities were summed as protein intensity. The protein list with False Discovery Rate (FDR) lower than 5% generated by analyzing SWATH data with PeakView 2.2, was exported to MarkerView 1.2.1 (AB SCIEX) for statistical analysis.

### 2.9 | Statistical analysis

Statistical analysis was performed using the IBM SPSS Statistics 23 software. Data are reported as mean ± standard deviation (SD) with of \( p < 0.05 \) considered significant. The Kolmogorov–Smirnov test was performed to test variables normality. The General Linear Model (GLM) with adjusted Sidak’s multiple comparison test with “group” (Untreated OM, LIPUS MCGM, and LIPUS OM) and “experimental time” (24 hr, 7 days, 14 days, 14 days on +7 days off, 21 days, and 21 days on +7 days off) as fixed effects, was performed to assess the influence of LIPUS treatment on hMSCs osteogenic differentiation.
Precision of the reported coefficients was assessed by creating 1,000 bootstrap samples from the entire data set and repeating the estimation process. Standard errors of the means and significances were obtained using this bootstrap method of the corresponding sampling distributions. In particular, the following comparisons were taken into account:

1. within each "experimental time": LIPUS OM versus Untreated OM and LIPUS MSCGM;
2. within each "group": 14 days versus 7 days; 21 days versus 14 days; 14 days on +7 days off versus 14 days; 21 days on +7 days off versus 21 days; 14 days on +7 days off versus 21 days; 21 days on +7 days off versus 14 days.

Regarding the analysis of protein list, a pairwise t-test and principal component analysis (PCA) were done by using MarkerView 1.2.1 software. Heat map clustering of all quantified proteins was performed by using Heatmapper free available web server (http://www.heatmapper.ca). Mean of replicates was used for performing the following paired comparison: (i) Untreated MSCGM versus Untreated OM; (ii) Untreated MSCGM versus LIPUS MSCGM; (iii) LIPUS MSCGM versus Untreated OM; (iv) LIPUS OM versus Untreated OM; (v) LIPUS MSCGM versus LIPUS OM of all different growth conditions. Fold Change (FC) thresholds at 1.5 with an adjusted p-value <0.05 were used to consider a protein up- or down-regulated. The molecular interaction networks among the significantly modulated proteins was analyzed by STRING v10 (Search Tool for the Retrieval of Interacting Genes/Proteins; http://string-db.org/) using the confidence level >0.7. Of note, the reported networks not include all proteins significantly up- or down-modulated since disconnected nodes are hidden for visualizing molecular interactions.

3 | RESULTS

3.1 | Proteomics analysis

To understand the proteomes patterns of hMSCs maintained in MSCGM and OM or after LIPUS treatments, a SWATH quantitation for all samples was performed. A list of 787 proteins was identified, as reported in Supplementary Table S1 online (sheet "Spectral reference library"). In addition, data were validated by using Peakview 2.2 software, obtaining a quantitative information for 555 proteins (Supplementary Table S1 online, sheet "Protein quantification"). Then, PCA and heat map clustering were carried out to acquire a comprehensive structure of the behavior of all quantified proteins under different growth conditions. PCA projection demonstrates that the maximum variability in the dataset, regardless of exposure to LIPUS, occurs between cells grown in presence of MSCGM or OM with the first component covering 68.1% of the data variance (Figure 2a). This result is also reflected in the Heat Map, where two major clusters separating the protein abundance profile of hMSCs grown in presence of MSCGM (both Untreated and LIPUS) and OM (both Untreated and LIPUS) can be observed (Figure 2b). The lists of proteins obtained for each paired comparison were found differentially expressed (Supplementary Table S1, sheet "Protein quantification").

The comparison "Untreated MSCGM versus Untreated OM" was considered as the "reference comparison" since cells in MSCGM maintained their sternness, while in OM were induced

![FIGURE 2](image-url)
to osteogenic differentiation. Thus, the protein profile of LIPUS MSCGM cells was compared to which of cells grown in Untreated OM, as well as to Untreated MSCGM cells in order to better define the molecular strategies through which LIPUS specifically affects the hMSCs behavior. Venn diagrams reported in Figure 3 highlighted that about 38% of proteins modulated in the Untreated MSCGM versus LIPUS MSCGM comparison (13 up-regulated and five down-regulated out of 50 differentially expressed) showed the same trend in the reference comparison, indicating that compared to the basal condition (Untreated MSCGM) the addition of LIPUS induced effects comparable to osteogenic differentiation, as already reported in several published studies (Azuma et al., 2001; Rutten, Nolte, Korstjens, & Klein-Nulend, 2009). In details, SWATH-MS analysis showed some significantly LIPUS-modulated proteins specifically related to osteogenic differentiation: (a) Cytoskeletal reorganization proteins that directly stimulates osteoblastic differentiation (CCT2, CALD1, and CTTN 9) (Higuchi, Nakamura, Yoshikawa, & Itoh, 2009); (b) Galectin 3, which is involved in osteogenesis and formation of bone nodules (LGALS3) (Aubin, Gupta, Bhargava, & Turksen, 1996); (c) Profilin 1, which is down-regulated in osteogenic differentiation process (PFN1) (Lin et al., 2016); and (d) Osteonectin, a marker of osteogenic differentiation (SPARC) (Delany, Kalajzic, Bradshaw, Sage, & Canalis, 2003; Jundt, Berghäuser, Termine, & Schulz, 1987; Kessler & Delany, 2007) (Supplementary Table S2 and Figure S1).

In the LIPUS MSCGM versus Untreated OM comparison it was interesting to note that about 58% of modulated proteins (45 up-regulated and 30 down-regulated out of 128 differentially expressed) showed the same trend in the reference comparison. Moreover, STRING analysis of the modulated protein dataset, showing only high confidence functional connections (confidence score ≥0.7), highlighted two interesting aspects.

The first one concerned the down regulation in LIPUS MSCGM cells, of proteins associated to Ras homolog gene family, member A (RhoA) protein and its activity as a positive regulator of hMSCs osteogenic differentiations (Figure 4a) (Chen et al., 2011). In particular, it was found that the triple functional domain protein (Trio), a specific RhoA activator, was down-regulated in undifferentiated hMSCs (−14 fold/p = 0.0106 in Untreated MSCGM cells vs. Untreated OM cells; Supplementary Table S1, sheet “Protein quantification”), as well as in presence of LIPUS (−25.4 fold/p = 0.0060 in LIPUS MSCGM cells vs. untreated OM cells; Supplementary Table S1, sheet “Protein quantification”). It was interesting to find that also in hMSCs grown in LIPUS OM there was a consistent down-regulation of TRIO in comparison to cells grown in Untreated OM (−42.6 fold/p = 0.0058 in LIPUS OM cells vs. untreated OM cells; Supplementary Table S1, sheet “Protein quantification”).

The second aspect emerged from STRING analysis was that in comparison to OM stimulation, Untreated MSCGM cells (Figure 4b) and LIPUS MSCGM cells (Figure 4c) were characterized by the up-regulation of proteins forming four overlapping functional networks: Ribosome constituent/Protein metabolic process (green area: 36 proteins in Untreated MSGCMG cells and

FIGURE 3 Venn diagram showing the proteins up- and down-regulated in the indicated comparisons. The numbers in brackets represent the total number of up- or down regulated proteins in the respective comparison.
FIGURE 4  Protein–protein interaction network analysis of proteins down-regulated in LIPUS MSCGM vs untreated OM comparison (a), and up-regulated in Untreated MSCGM versus Untreated OM comparison (b) and in LIPUS MSCGM versus Untreated OM comparison (c). The protein–protein interaction network of the proteins listed in Supporting Information Table S1 was constructed using the STRING v10 database, and shows only high confidence functional connections (confidence score >0.7). Among proteins up-regulated in Untreated MSCGM versus Untreated OM and in LIPUS MSCGM versus Untreated OM four functional overlapping networks are identified: Ribosome constituent/Protein metabolic process (green area), Glycolysis/Gluconeogenesis (blue area), RNA metabolic process/Splicing (pink area), tubulins (yellow area).

FIGURE 5  Results of cell viability (a) and dsDNA content (b) of Untreated OM (○), LIPUS MSCGM (●), and LIPUS OM (■) cells, expressed as percentage of Untreated MSCGM values (100%) (Mean ± SD, n = 3 duplicates). GLM analysis with "group" and "experimental time point" as fixed effects showed for: Cell viability—a main effect of "experimental time," $F = 12.32, p < 0.0005$; dsDNA content—an interaction between "group" and "experimental time," $F = 12.42, p < 0.0005$. Adjusted Sidak multiple comparison test (*$p < 0.05$; **$p < 0.005$; ***$p < 0.0005$).
27 in LIPUS MSGCMG cells). Glycolysis/Gluconeogenesis (blue area: four proteins in both cells), RNA metabolic process/Splicing (pink area: nine proteins in Untreated MSGCMG cells and three in LIPUS MSGCMG cells), tubulins (yellow area: three proteins in Untreated MSGCMG cells and two in LIPUS MSGCMG cells).

3.2 | Cell viability and dsDNA content

Cell viability was higher in all groups in comparison with Untreated MSCGM at 7, 14, and 21 days (Figure 5a), followed by a constant decrease. A progressive increase in dsDNA content within 21 days was
observed in Untreated OM (Figure 5b), even though dsDNA content values were lower than those in Untreated MSCGM (except at 21 days). When LIPUS stimulation was applied, an increase in dsDNA content was observed compared to Untreated MSCGM, where dsDNA remained constant even after LIPUS stimulus was stopped. The simultaneous presence of OM medium and LIPUS stimulation (LIPUS OM) determined significant changes in hMSC dsDNA content compared to Untreated OM (21 days: −73%, p < 0.0005; 14 days "on" + 7 days “off”: 139%, p < 0.0005) and LIPUS MSCGM (7 days: −32%, p < 0.05; 14d: d = −33%, p < 0.005; 21 days: d = −68%, p < 0.0005).

3.3 | hMSC characterization

Flow cytometry results showed that hMSCs were positive for the MSC markers CD73 (40–54%), CD90 (27–48%) and CD105 (31–84%), and for the stem cell transcriptional factors NANOG (10–34%), SOX2 (10–34%) and OCT-3/4 (28–61%) at 21 days. Figure 6a-d shows a percentage of hMSC positive gates for CD73, CD90, and CD105 at 21 days. The double positive hMSC populations co-expressing stem cell markers showed a similar decreasing trend over time and among cultures, whereas those co-expressing stem cell transcriptional factors showed an increasing trend with the highest values in LIPUS-treated cultures (Figure 6e).

OsteoImage™ Mineralization assay showed that hMSCs maintained in OM had more calcium nodule formation than those cultured in MSCGM medium (Figure 7). In all groups, significant increases in mineralization were observed over time. The LIPUS OM group presented the highest hydroxyapatite content compared to Untreated OM (7 days: 39%; 14 days: 20%; 21 days: 7%) and LIPUS MSCGM (7 days: 30%; 14 days: 64%; 21 days: 37%) groups at each experimental time.

3.4 | RT-qPCR analysis

LIPUS stimulation induced an increase in gene expression of RUNX2, ALPL, COL1A1, and BGLAP compared to Untreated MSCGM at seven (COL1A1 and BGLAP) and 21 days (RUNX2, ALPL, and COL1A1) (Figure 8). In Untreated OM and LIPUS OM a lower expression of RUNX2 compared to Untreated MSCGM was found, which increased after the cessation of LIPUS stimulation and/or differentiating medium (Figure 8a). A significantly lower value in the expression of RUNX2 at 21 days was observed between LIPUS OM and LIPUS MSCGM. The combination of LIPUS stimulation with OM (LIPUS OM) led to an increase in ALPL expression compared to Untreated OM (40%, p < 0.05) and LIPUS MSCGM (97%, p < 0.05) (Figure 8b). Changes in the expression of ALPL observed over time showed a maximum increase in expression at 21 days with a significant decrease 7 days after the cessation of any stimuli (p < 0.005). The expression of COL1A1 was greater in LIPUS MSCGM than in LIPUS OM (68%, p < 0.05) and it was lower in LIPUS OM and Untreated OM compared to the Untreated MSCGM group, except for the expression at 21 days (Figure 8c). SPP1 presented the highest expression at 7 days (46-fold increase), decreasing after 14 days of culture with values lower than the control group (Untreated MSCGM) (Figure 8d). Similarly, BGLAP expression levels (Figure 8e) decreased significantly from 7 to 14 days, but in LIPUS OM the decrease did not occur until the cessation of LIPUS and OM stimuli, but remained constant at 7, 14, and 21 days.

LIPUS treatment and OM medium induced an up-regulation of MAPK1 in all groups in comparison with Untreated MSCGM, reaching its highest values 7 days after the cessation of stimuli administered up to day 21 (Figure 9a). In particular, the highest MAPK1 expression was found in Untreated OM at 21 days “on” + 7 days “off” (p < 0.0005) compared to LIPUS OM. Conversely, both stimuli determined a down-regulation of MAPK6 in Untreated OM and LIPUS MSCGM, which resolved after the cessation of stimuli (Figure 9b). The MAPK6 expression in LIPUS OM group showed a different trend with a progressive up-regulation, reaching the highest value at 14 days compared to LIPUS MSCGM (p < 0.05), followed by a down-regulation overtime.

3.5 | ELISA assays

ELISA results are reported in Figure 10. No significant increases in the production of ALP were observed in any of the groups other than Untreated MSCGM. In Untreated OM, COL1a1 synthesis was increased by about 2.4%, compared to Untreated MSCGM, whereas LIPUS stimulation increased COL1a1 synthesis by 0.5% (LIPUS MSCGM). COL1a1 release increased by 2.6% in LIPUS OM compared to Untreated MSCGM; this was very different from LIPUS MSCGM (369%, p < 0.005). In all groups, high and modest increases in OC and OPN release, respectively, were observed. OC synthesis results indicated that OM medium (Untreated OM) or LIPUS treatment (LIPUS MSCGM and LIPUS OM) produced on average a similar OC increase by approximately 30% compared to Untreated MSCGM.
FIGURE 8  Results of RUNX2 (a), ALPL (b), COL1A1 (c), SPP1 (d), and BGLAP (e) gene expression of Untreated OM (■), LIPUS MSCGM (□), and LIPUS OM (▲) cells expressed as relative fold changes (RF) of Untreated MSCGM values (1) (Mean ± SD, n = 3 duplicates). GLM analysis with “group” and “experimental time” as fixed effects revealed for: RUNX2-an interaction between “group” and “experimental time,” F = 3.07, p < 0.05; ALPL-main effects of “group” F = 5.09, p < 0.05 and “experimental time” F = 11.94, p < 0.0005; COL1A1-main effects of “group,” F = 9.18, p < 0.005 and “experimental time” F = 4.92, p < 0.05; SPP1-main effects of “experimental time” F = 19.78, p < 0.0005; BGLAP-an interaction between “group” and “experimental time,” F = 3.22, p < 0.05. Adjusted Sidak multiple comparison test (*p < 0.05; **p < 0.005; ***p < 0.0005)
Conversely, in Untreated OM or LIPUS MSCGM, OPN release was 2% compared to Untreated MSCGM, whereas the combined OM and LIPUS stimuli (LIPUS OM) determined only a 0.5% increase, which was significantly lower by about 70% ($p < 0.05$) compared to the Untreated OM and LIPUS MSCGM groups.

4 | DISCUSSION

Different studies have been performed on the effects of LIPUS on cells attempting to describe the cellular response to biomechanical stimuli (Frairia & Berta, 2011) or transient membrane permeability (Engler, Sen, Sweeney, & Discher, 2006; Huang, Holfeld, Schaden, Orgill, & Ogawa, 2013; McBeath, Pirone, Nelson, Bhadriraju, & Chen, 2004). In the present study, the role of LIPUS stimulation (SATA intensity at 30 mW/cm$^2$) in hMSC osteogenic commitment and differentiation was investigated using an in vitro model, where LIPUS stimulation was evaluated in terms of proteome, gene expression, and proteins release modulation.

PCA and heat map clustering analyses of the obtained proteomic data, suggested that the most significant differences in protein expression profiles of MSCs are due to the specific growth medium (MSCGM or OM) rather than the exposure to LIPUS. Very few studies were focused indeed on osteogenic differentiation of hMSC (Gao, Walmsley, Cooper, & Scheven, 2016; Hu et al., 2014; Yue et al., 2013) mediated by LIPUS treatment. In this sense, our proteomics analysis revealed the existence of differences in osteoblast commitment and differentiation potential between all experimental groups. Thus, deeper proteomic data analysis showed the peculiar ability of LIPUS to positively regulate several proteins and pathways associated with stemness, such as the autophagosomes proteins SQSTM1, SRSF2, TMEM109, or Glycolysis/Gluconeogenesis proteins (CKAP4,COX4I, ALDOA) (Folmes & Terzic, 2016; Folmes et al., 2011), and to negatively regulate signaling networks associated to the differentiation of MSCs into osteoblasts, such as the RhoA/ROCK pathway (Chen et al., 2011; Fang et al., 2008; Yun et al., 2016). In particular, our quantitative proteomic analysis indicated that, regardless the grown medium, LIPUS exposure induced a consistent down-regulation of Trio, a specific RhoA activator. The Rho family of small GTPases controls a diverse array of cellular processes, including cytoskeletal dynamics, cell polarity, membrane transport, and gene expression. The RhoA is a molecular switch that respond to cell surface receptors for various cytokines, growth factors, adhesion molecules, and G-protein-coupled receptors by cycling between an inactive guanosine diphosphate (GDP)-bound and an active guanosine triphosphate (GTP)-bound form. Recently, Chen et al. (2011) demonstrated that during the differentiation process in bone marrow stem cell lines, RhoA protein expression increases on the membrane in the activated form of GTP-binding protein activating cellular actin remodeling and cytoskeleton rearrangement.

Regarding hMSC viability and proliferation, LIPUS stimulation or the presence of OM determined an increase in cell viability over time. Even though dsDNA content levels did not reach those observed in Untreated MSCGM, LIPUS MSCGM cells showed higher dsDNA values than those treated with LIPUS stimulation and OM. In fact, the simultaneous presence of OM and LIPUS stimulation (LIPUS OM) inhibited cell proliferation, thus resulting in significantly lower dsDNA content at each experimental time point, which was likely due to the start of the differentiation process (Wang, Peng, et al., 2014).

The analysis of specific markers of hMSC, confirmed the proteomic data of Untreated MSCGM and LIPUS MSCGM. The phenotypic characterization permitted to identify a functional multi-potent hMSC stemness potency maintenance (CD73+/CD90+/CD105+) in the tested groups until 14 days, which decreased at 21 days concurrently with osteoblast differentiation (Kern & Shibata, 2007; Pittenger et al., 1999; Ringdén & Le Blanc, 2006; Rutten et al., 2009; Saalbach, Haustein, & Anderegg, 2000; Yoshimura et al., 2006, 2007). In addition, LIPUS stimulation markedly promoted the
expression of stem cell transcriptional factors OCT-3/4, SOX2, and NANOG related to stemness maintenance, as recently reported by Kusuyama et al. (Alvarez et al., 2015; Kusuyama et al., 2015). As for the tendency of MSCs to lose their multipotency over time in culture, flow cytometry data recorded at different experimental time points might be consistent with long term culture stress, as reported by Stolzing, Coleman, & Scutt, 2006. In particular, according to Haasters et al. (2009) the loss of CD73 in hMSCs culture might be due to either a down-regulation of CD73 or a beginning of differentiation with the formation of a subpopulation of cells not expressing CD73.

These data suggested that hMSCs treated with LIPUS might maintain the expression of mesenchymal markers and increase the hMSC osteoblast lineage commitment (Kulterer et al., 2007) and differentiation especially after LIPUS and OM stimulation.

Finally, an increase in mineralization in Untreated OM and, interestingly, in LIPUS OM occurred after only 7 days, thus confirming a prompt transition from a proliferative state (hMSCs) toward a differentiating state, mostly after LIPUS treatment (LIPUS OM showed the highest calcium nodule formation capability).

The current results suggested that in vitro LIPUS stimulation induced the activation of the osteogenic specific pathways and relative protein release (Claes & Willie, 2007), favoring bone formation, in particular in the presence of osteogenic soluble factors that mimic bone microenvironment. RUNX2 expression was analyzed because it is (a) a fundamental transcription factor for osteogenic differentiation (Ducy, Zhang, Geoffroy, Ridall, & Karsenty, 1997; Otto, Lübbert, & Stock, 2003); and (b) the target of mechanical signals by which physical stimuli act on the metabolic activities of osteoblasts (Ziros et al., 2002). RUNX2 mRNA expression level was increased in comparison to Untreated MSCGM at experimental times next to 21 days. The analysis of gene targets of RUNX2 and their protein release showed that higher values of COL1A1 mRNA expression were found in LIPUS MSCGM compared to LIPUS OM, whereas COL1a1 protein release increased in LIPUS OM compared to LIPUS MSCGM. This is suggestive of an
The combination of LIPUS stimulation with OM led to a statistically significant increase in ALPL mRNA expression at 21 days of treatment both in comparison to Untreated OM group and LIPUS MSGCM, indicating an additive effect of LIPUS on ALPL gene regulation. The release of ALP did not increase with the OM and/or LIPUS stimulation and the highest ALP release was observed in Untreated MSGCM. Most likely, Untreated MSGCM showed an increase in ALP release because there were more osteoblasts committed cells compared to other groups, as demonstrated by phenotypic characterization and gene expression analysis. In addition, ALP data suggested the presence of intracellular control to maintain the balance between mRNA transcription and transduction, in order to allow the synthesis of a correct quantity of specific osteoblast protein throughout the differentiation pathway.

As the effect of LIPUS on MAPK signaling is concerned, LIPUS stimulation and OM induced an up-regulation of MAPK1 mRNA expression in all groups over time, but both stimuli determined a down-regulation of MAPK6. MAPK6 expression in the LIPUS OM group showed a different trend with a progressive up-regulation, reaching its highest value at 14 days of culture compared to LIPUS MSGCM, followed by down-regulation over time. Therefore, an effect of LIPUS on MAPK1/6 up-regulation was detected at gene expression level only after the ultrasound stimulus was stopped. This is probably due to the fact that at 7 and 14 days MAPK1/6 synthesis could have been already stimulated by the previous exposition to LIPUS (Sato et al., 2015). At later time points, after the end of stimulation (7days off), gene up-regulation could be a direct consequence of ultrasounds-induced mechanotransduction, confirming the role of LIPUS as an activator of multiple mechano-sensitive pathways.

To sum up, the current study suggests that LIPUS might be able to maintain, in vitro for a long time, the niche of osteoblast committed hMSCs. Indeed, all data indicated that compared to Untreated OM, LIPUS exposure down regulates some pathways associated to osteogenic differentiation and allowed cells to maintain a protein expression profile characterizing the MSCs, providing new indication on the specific role that LIPUS could have in supporting the osteoblast commitment and differentiation of hMSCs.

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SUPPORTING INFORMATION

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