کاهش یپری سلول‌های بنیادی مزانشیمی با استفاده از نیز سولفات در معرض میدان الکترومغناطیس با فرکانس بسیار پایین

روش استفاده: سلول‌های بنیادی مزانشیمی مشتق شده از بافت چربی موش صحرایی به واسطه سولفات روی به عنوان یک عنصر کمیاب ضروری و میدان الکترومغناطیس با فرکانس بسیار پایین تأثیر می‌گذارند. در این مطالعه، سلول‌های بنیادی مزانشیمی مشتق شده از بافت چربی موش و در معرض میدان الکترومغناطیس به واسطه القاء بیان ژن تلومراز در زمان واقعی با استفاده از روش RT-PCR شناسایی گردید که سولفات روی ممکن است باعث کاهش پیری و افزایش بیان ژن تلومراز و کاهش پیری سلول‌های بنیادی مزانشیمی با استفاده از میدان الکترومغناطیس با فرکانس بسیار پایین شود. به نظر می‌رسد که سولفات روی ممکن است عامل مؤثری در تأخیر افتاده‌پیری سلول‌های بنیادی مزانشیمی با استفاده از میدان الکترومغناطیس با فرکانس بسیار پایین باشد.

واژه‌های کلیدی: پیری، سلول‌های بنیادی مزانشیمی، سولفات، میدان الکترومغناطیس
Introduction

Everyone is exposed to a complex mix electromagnetic field (EMF) that permeates our environment. The effects of extremely low frequency electromagnetic fields (ELF-EMFs) on biological systems including anti-oxidative enzymes, cell proliferation and differentiation of stem cells have been investigated. Today, the hazardous or beneficial biological effects of EMF on human and animals are the subject of many studies. In recent years, some studies have shown an association between exposure to ELF-EMF and changes in the expression of genes involved in the metabolic processes, response to stress and cell proliferation, aging and death. It has been reported that EMF of 75 Hz, 2 mT increases cell proliferation rate of human mesenchymal stem cells (hMSCs). Moreover, it has been shown that EMF of 50 Hz, 1 mT can enhance cell proliferation rate and inhibit programmed cell death in human neuroblastsoma and rat pituitary cells. On the contrary, it was found that EMF of 50 Hz, 20 mT can inhibit growth and metabolism of hMSCs.

Reportedly, telomeres and telomerase provide a way to delay aging. Telomeres are the ends of linear chromosomes that protect the chromosomes from end-to-end fusion and maintain genomic integrity. Telomerase is a ribonucleoprotein enzyme that uses its RNA as a template for synthesizing telomeric DNA on chromosomal ends. Telomerase reverse transcriptase (TERT) is the catalytic component of telomerase enzyme considered to play an essential role in the activation of telomerase. TERT expression is tightly regulated and it is only found highly expressed in germ cells, stem cells and ~90% of cancer cell lines containing a functional telomerase. There is a correlation between telomere length and aging. On the other hand, aging is characterized by presenting cells with critically short telomeres. Furthermore, some studies have suggested that aging is the result of oxidative damage accumulation caused by free radicals generated as by-products during normal metabolism. Moreover, the increase in free radicals are known to induce apoptosis and aging and ELF-EMF has been shown to increase free radical levels. With these explanations, much attention has been given to the elements preventing cellular aging in the last decade. It has been shown that L-carnitine as an antioxidant can be used as a good candidate for extending the replicative life-spans of aged MSCs by increasing TERT gene expression and telomere length. Further, it has been documented that radio electric asymmetric conveyer (REAC) directly influences the expression of TERT and telomere length during aging of hMSCs by transcriptionally enhancing the expression of TERT and counteracting telomere shortening.

It is believed that Zinc ion (Zn\(^{2+}\)) plays a critical role in various biological processes like cell growth and cycling and has an important role in many processes that are related to the scavenging of reactive oxygen species, brain aging and onset of age-related neurodegenerative diseases. There are some reports that Zn\(^{2+}\) deficiency causes apoptosis in various cell and tissue types A number of studies have shown the correlation between Zn\(^{2+}\) and apoptosis and suggested that Zn\(^{2+}\) may function as an antioxidant in cells. One anti-apoptotic mechanism of Zn\(^{2+}\) is its ability to minimize oxidative damage to cellular organelles.

The use of human beings as the test model is impossible and the long-term observation of a test group could be affected by other factors such as dietetic habit and inherited disease. Certainly, stem cells could be the best choice for experimental studies. Any small influence on stem cells may cause unexpected results to their related adult cells. Furthermore; studies on the influence of ELF-EMF on stem cells are rare. To date, the effects of ELF-EMF in the presence of zinc sulfate (ZnSO\(_4\)) are yet to be reported on the TERT gene expression and senescent adipose-derived mesenchymal stem cells (ADSCs). This study was carried out to evaluate the effect of ZnSO\(_4\) on TERT gene expression and aging of ELF-EMF-exposed rat primary adipose MSCs.

Materials and Methods

All chemicals, unless specified, were purchased from Sigma-Aldrich (St. Louis, USA). All cell culture plasticware was from SPL Life Sciences (Pocheon, Korea). The type of this study was experimental study.

**Isolation and culture of ADSCs.** In this study, consent was given by the ethical committee of the University of Tabriz (Tabriz, Iran). Male rats (6 to 8 week-old) were euthanized using ketamine (87 mg kg\(^{-1}\); Alfasan, Woerden, The Netherlands) and xylazine (13 mg kg\(^{-1}\); Alfasan) and epididymal adipose tissues were collected under sterile conditions. The ADSCs were isolated from adipose tissue as previously described. Briefly, adipose tissues were transported to the laboratory in phosphate-buffered saline (PBS) solution supplemented with 2% (v/v) fetal bovine serum (FBS) within 2 hr post-operation, carefully dissected and minced using a sterile scissor. Subsequently, the fat was washed extensively with Dulbecco’s modified Eagle’s Medium (DMEM) supplemented with 5% penicillin/streptomycin. Tissue was then enzymatically dissociated for 30 min at 37 °C using 0.075% (w/v) collagenase type I (Invitrogen Ltd., Paisley, UK). The solution was neutralized by addition of DMEM containing 10% (v/v) FBS and centrifuged at 800 g for 5 min. The cell pellet was filtered through a 75-µm nylon mesh to remove cellular debris and re-suspended in DMEM containing 10% (v/v) FBS and 1% (v/v) penicillin/streptomycin solution. Following incubation, the plates were washed extensively with PBS to remove non-adherent cells, red blood cells and remaining debris. Cultures were maintained in sub-confluent levels in a
37 °C incubator with 5% CO₂ and passaged with trypsin ethylenediaminetetra acetic acid (EDTA; Invitrogen) when required.²⁶,²⁷

Detection of ADSCs markers by Immunocytochemistry. A total of 4 × 10⁴ cells from passage four were seeded in a 24-well culture plate. After one day of culture, cells were washed three times with PBS and fixed in 4% paraformaldehyde for 30 to 60 min at room temperature. After fixation, the parafomaldehyde was removed and cells were then washed two times with PBS and once with PBS and 1% bovine serum albumin (BSA). Cells were incubated overnight at 4 °C with a 1:100 dilution of monoclonal antibodies (mAbs) against CD73, CD90, CD45 and CD56 (BD Biosciences, San José, USA) in PBS and 1% BSA. Cells were then washed three times with PBS and 1% BSA and incubated with a 1:500 dilution of biotin-conjugated mouse monoclonal IgG1 antibody against rat in PBS and 1% BSA for one hr. After three washings with PBS, a 1:500 dilution of Streptavidin Alexa Fluor™ 488 conjugate (Molecular Probes, Eugene, USA) was added for 1 hr. Cells were washed three times with PBS and nuclei were stained with 7.50 µM propidiod iode (PI) for 15 min. After washing three times with PBS, cells were covered with Vectashield mounting medium (Vector Laboratories Inc., Burlingame, USA) and visualized under the fluorescence microscope. The pluripotent capacity of the isolated MSCs was confirmed with adipogenic and osteogenic differentiation.

Adipogenic differentiation and Oil Red-O staining. Cells at passage four were seeded at a density of 20 × 10³ cells per cm². Sub-confluent cells were incubated in adipogenic induction medium containing 0.50 mM 1-methyl-3-isobutylxanthine, 1 µM dexamethasone, 10 µg mL⁻¹ insulin and 200 µM indomethacin; the medium was changed every 3 to 4 days. At the end of the day 21, formalin-fixed cells were washed in 50% isopropanol and stained with Oil Red-O for 15 min and lipid droplets were observed by a light microscope.²⁸

Osteogenic differentiation and alizarin red staining. The ADSCs were plated in the same manner as described for the adipogenesis. After ADSCs reached about 90% confluency, osteogenesis was induced by osteogenic induction medium containing 10% FBS, 10 nM dexamethasone, 100 U mL⁻¹ penicillin, 100 µg mL⁻¹ streptomycin, 10 mM b-glycerophosphate, and 0.05 mM L-ascorbic acid-2-phosphate for 21 days. To confirm the successful osteogenic differentiation, after 21 days of culture, calcium depositions were stained with alizarin red staining. Briefly, cells were washed twice with excess PBS and fixed in a solution of 2% (v/v) formaldehyde. After 15 min, alizarin red (40 mM, pH 4.10) was added to each well. The plates were incubated at room temperature for 20 min and then, they were washed two to three times with PBS for 5 min to reduce non-specific staining.²⁹

RNA extraction and reverse transcription (RT)-PCR analysis of bone and adipose tissue-specific genes expressions. Total RNA from the osteogenic and adipogenic differentiated cells was isolated using trizol reagent (Invitrogen).²⁶ Extracted cellular RNA was dissolved in diethyl phosphorocyanidate-treated water. After DNase treatment by DNase I amplification grade kit (Invitrogen), 2 µg RNA were used for the first strand cDNA synthesis in a total volume of 20 µL according to the manufacturer’s guidelines. The thermal program for PCR was an initial denaturation for 15 min at 94 °C followed by 28 cycles of 1 min at 94 °C, 1 min at 55 °C, 1 min at 59 °C, 58 °C and 56 °C (an optimised annealing temperature for alkaline phosphatase (ALP), osteocalcin (OCN), PPAR-alpha, PPAR-gamma and GAPDH primers, respectively), and 1 min at 72 °C with a final 10 min extension at 72 °C. Reaction mixtures for PCR included 20 ng cDNA, 10 µL TEMPase Hot Start 2x Master Mix I Blue (Ampliqon A/S, Odense M, Denmark), 0.50 µM of each antisense and sense primer as follows: ALP (NM_013059.1): 5'- CCTTGGAAAAT GCCCTGAA-3' (forward), 5'- CTGGAGAGAGCCCAAAGG -3' (reverse), osteocalcin (OCN) (NM_001278484.2): 5'- GTCCCCACACCACTGC-3' (forward), 5'- CGAAGGGCTGA AGCTGCGG-3' (reverse), PPAR-alpha (NM_013196.1): 5'- CCCGGTCTCCGTGGAAC TGAC-3' (forward), 5'- GGGACTCATCCTGACTGTGGGGGAC-3' (reverse). The PPAR-gamma (NM_013124.3): 5'- GGTGA AACTCTGGGAGATCC-3' (forward), 5'- TGAGGAGTTTGA GACTCTTC-3' (reverse), GAPDH (NM_017008.4): 5'- ATGACTCTACCCAGGCAAG-3' (forward), 5'- CTTGAGATGT GTGATGATT-3' (reverse) and 9. Amplified PCR products were analyzed by ethidium bromide staining after 1.5% agarose gel electrophoresis.³⁰,³¹ The PCR primers were designed with Oligo primer design software (version 7.0; National Biosciences, Plymouth, USA).

Determination of suitable concentration of ZnSO₄ and cell proliferation by methylthiazol tetrazolium (MTT) assay. The tetrazolium-based colorimetric assay MTT test measures the mitochondrial activity in the cell culture, which reflects the number of viable cells. Briefly, at passages three to aux cells were trypsinized, counted with hemocytometer and seeded at a density of 4 × 10³ cells per well to a 96-well culture plate. Cells were incubated for 24 hr at 37 °C in a humidified environment with 5% CO₂ to grow the cells in a monolayer. The ZnSO₄ was added to the wells at final concentrations of 1.40, 0.14 and 0.014 µg mL⁻¹. Control wells were prepared by addition of corresponding medium. The stock MTT dye solution (2 mg mL⁻¹) was added to each well after the plates were incubated at 37 °C in a 5% CO₂ incubator for 21 days. Following incubation for 4 hr, the supernatant was removed and dimethyl sulfoxide (100 µL) was added. The optical density of each well was measured in an ELISA reader at a wavelength of 570 nm.³²,³³

The ELF-EMF system and exposure protocol. The EMF device was designed and manufactured by University
of Tabriz in Iran. The EMF was generated by a parallel set of Helmholtz solenoid coils with 500 turns of 0.70 mm coated copper wire. Each solenoid diameter was 27 cm. The coil was placed into the cell incubator; the field was set to 50 Hz electromagnetic field frequency and generated a magnetic flux density of 20 mT as previously described.4 The EMF-exposed plates were placed in the center half way between the plains of coils to receive a uniform field for 30 min per day for 21 days. Sham-exposed control samples were kept under the same conditions but in another incubator, without using EMF. The CO2 concentration, temperature and humidity of the sham-exposed control samples were similar to EMF-exposed samples. The value of the alternating magnetic field was measured continuously using an EFA-2 field analyzer with a 3 cm diameter probe.

**Real-time PCR for analyzing TERT gene expression.** The ADSCs (from passage four) were cultured at a concentration of 30 × 10^4 cells per well in 6-well plates for 21 days consist of four groups including group I (control without ZnSO4 treatment and EMF exposure), group II (50 Hz, 20 mT EMF exposure without ZnSO4 treatment), group III (0.14 μg mL^-1 ZnSO4 treatment without EMF exposure) and group IV (50 Hz, 20 mT EMF exposure with 0.14 μg mL^-1 ZnSO4 treatment). Exposure time duration for EMF-exposed groups was 30 min per day for 21 days. After termination of EMF exposure at the end of 21th day, total RNA was isolated from each group of cells using TRIzol reagent (Invitrogen Life Technologies, Burlington, Canada). The cDNA synthesis was carried out using RevertAid™ first strand cDNA synthesis kit (K1622; Fermentas Life Science, Frankfurt, Germany). The resulting cDNA was stored frozen at −20 °C until assayed by real-time PCR. All PCR reactions were performed using Corbett Rotor-Gene (model 6000 HRM; Corbett Research, Sydney, Australia) in a total volume of 20 μL containing 10 μL Power SYBR Green master mix (2x) (TaKaRa Bio, Sydney, Australia) in a total volume of 20 μL containing 10 μL Power SYBR Green master mix (2x) (TaKaRa Bio, Sydney, Australia) in a total volume of 20 μL containing 10 μL Power SYBR Green master mix (2x) (TaKaRa Bio, Sydney, Australia) in a total volume of 20 μL containing 10 μL Power SYBR Green master mix (2x) (TaKaRa Bio, Sydney, Australia). The β-actin (NM_031144) and TERT transcripts in samples were measured in relation to β-actin. 

**Senescence-associated β-galactosidase (SA-β-gal) activity assay.** To determine the percentage of senescent cells in the presence of ELF-EMF and ZnSO4, SA-β-gal staining was used. The assay is based on cytochemical staining for β-galactosidase activity at pH 6. The ADSCs from passages 3, 5, 7 and 9 were plated (2 × 10^3 cells per cm²) in 24-well plates for 21 days consist of four groups including group I (control without ZnSO4 and ELF-EMF exposure); group II (ELF-EMF-exposure without ZnSO4); group III (ZnSO4 treatment without ELF-EMF exposure) and group IV (ELF-EMF exposure with ZnSO4). After termination of EMF exposure at the end of 21th day, cells were washed with PBS and fixed for 5 min with the fixation solution containing formaldehyde and glutaraldehyde. The cells were then, washed twice with PBS, incubated with freshly staining solution containing 40 mM citric acid/Na phosphate buffer, 5 mM potassium ferricyanide, 5 mM potassium ferrocyanide, 150 mM sodium chloride, 2 mM magnesium chloride and 1 mg mL^-1 of 5-bromo-4-chloro-3-indolyl-D-β-galactoside (X-gal) in distilled water overnight (12 to 16 hr) at 37 °C without CO2. After incubation, the cells were washed twice with PBS and once with methanol and the dishes were allowed to air dry. A minimum of 100 cells was counted by light microscopy in 10 random fields to determine the percentage of SA-β-gal-positive cells which were appeared as bluestained cells.31,37

**Statistical Analysis.** The results were analyzed using the software program Graph Pad Prism (version 6.01; Graph Pad Prism, Chicago, USA). We used one-way and two-way ANOVA followed by Dunnett’s post hoc test to determine the significant difference among groups. Statistical significance was determined at p < 0.05. All experimental procedures were repeated for three times.

**Results**

**Phenotypical characterization of ADSCs.** The ADSCs had the capacity to adhere to plastic flasks in culture and morphologically, they appeared as spindle-shaped cells both as scattered individuals and in small colonies. After three passages, red blood cells were seldom seen by microscopy. Immunocytochemical images revealed that cultured cells were consistently positive for CD73 and CD90 and negative for CD45 and CD56 (Fig. 1 A to D). The ADSCs were able to differentiate towards the adipogenic and osteogenic lineages.

**Adipogenic differentiation of ADSCs.** Positive adipogenic differentiation was determined by Oil Red-O staining and molecular analysis. Treated ADSCs with adipogenic differentiation media were stained positive with Oil Red-O staining (Fig. 1E). The expression of peroxisome proliferator-activated receptor-gamma (PPAR-γ) and PPAR-α as adipocyte-specific genes were detected by RT-PCR analyses. These results confirmed that our isolated cells were MSCs.
Osteogenic differentiation of ADSCs. The osteogenic differentiation was evident in alizarin red staining. After staining, redness of the nodules indicated the presence of mineralized compartments as a result of the osteogenic treatment (Fig. 1F). The RT-PCR analysis confirmed the expression of bone-specific genes including ALP and OCN in the treated cells.

Determination of ZnSO$_4$ effect on cell proliferation by MTT assay. As shown in Figure 2, ZnSO$_4$ had no significant effect on ADSCs proliferation at concentrations of 1.40 and 0.014 µg mL$^{-1}$, but at concentration of 0.14 µg mL$^{-1}$ significant proliferative effect was seen ($p < 0.05$). We used 0.14 µg mL$^{-1}$ ZnSO$_4$ in a complete culture medium to treat the cells.

Discussion

Due to increased use of ELF-EMFs for domestic and industrial appliances in the last few years, different *in vivo* and *in vitro* studies have been conducted on the influence of these fields on various biological functions such as tumor growth, neurodegenerative diseases, free radical production, gene expression, growth, aging and differentiation potential of MSCs. Since the MSCs are attractive candidates for cell-based tissue regeneration, finding EMFs effects on MSCs is important for researchers. Aging of MSCs is one of the conditions that may be caused by EMFs. Aging is an inherently complex process that is manifested within an organism at molecular
and cellular levels. While the fundamental mechanisms are still poorly understood, telomere shortening and reduction of TERT expression are one of the cellular markers of aging and/or cell death. Molecular control of stem cell aging is highly regulated by two different, telomerase-independent and telomerase-dependent pathways. Epigenetic events play a role in the telomerase-independent senescence pathway. Therefore, finding ways to reduce the replicative senescence is definitely attractive in cell transplantation approaches. The results of EMFs studies are different because of the difference in the magnetic field intensity and frequency and duration of exposure. Reportedly, REAC acts on a gene and protein expression program of both telomerase-independent and telomerase-dependent patterns to optimize stem cell ability to cope with senescence progression. It seems that gene expression regulation may be directly involved in differentiation potential and aging in stem cells. It has been demonstrated that TERT over-expression can prevent the human MSCs senescence and the cells showed significantly higher and unlimited proliferation capacities. It was found that 50 Hz, 0.50 and 1 mT magnetic fields can promote survival and proliferation of the human ADSCs. There is no substantial evidence for the effects of ELF-EMF on TERT expression. In this regard, an understanding of the TERT expression underlying the regulation of telomerase activity might allow the modulation of telomerase expression and thus, cell lifespan. At present, finding ways to reduce the reduction of TERT expression and cellular senescence caused by ELF-EMF is definitely attractive. In the present study, 0.14 μg mL⁻¹ was used as the final concentration of ZnSO₄. To determine a suitable range for ZnSO₄ under this study experimental conditions, MTT assay was used to evaluate cell viability. In this study, it was discovered that ZnSO₄ has no impact on the viability of cells below 0.14 μg mL⁻¹ and above 0.14 μg mL⁻¹. The previously described exposure condition was used in this study. Real-time PCR was used to explore the mechanism of ELF-EMF in the presence of ZnSO₄ on ADSCs aging at the molecular level and the mRNA expression of TERT was detected. The results of this study showed that in cultured cells under EMF in the presence of 0.14 μg mL⁻¹ ZnSO₄ (group IV), TERT mRNA expression remarkably increased (4.60 fold) when compared with EMF-exposed group (group II). Previous reports provided the first evidence that metals, especially Zn⁺² can modulate telomerase in cancer cells by inducing an enhancement of its activity. On the contrary, it has been reported that zinc phthalocyanine (ZnPc) has the ability to inhibit telomerase activity. In this study, β-galactosidase staining method was used as a common method to study cells senescence in vitro, in order to examine the percentages of the senescent ADSCs in our studied culture condition. According to our observations, β-galactosidase positive cells were first observed at passage three and their number increased as the passage number increases. The results of the present study showed that senescence related β-galactosidase positive cells at passages 3, 5, 7 and 9, under EMF in the presence of 0.14 μg mL⁻¹ ZnSO₄ (group IV) decreased significantly when compared with the EMF-exposed group (group II). The effects of ZnSO₄ on TERT gene expression and aging of ADSCs under ELF-EMF are yet to been investigated, but the role of Zn⁺² in growth regulation was evaluated. However, the findings of this study are in agreement with the hypothesis that 0.14 μg mL⁻¹ ZnSO₄ as an antioxidant can prevent the aging of ADSCs under 50 Hz, 20 ELF-EMF by increasing TERT gene expression and ADSCs aging reduction. Although the underlying mechanism remains unclear, but few studies have been published confirming the effectiveness of zinc on telomeres. Reportedly, octacationic zinc phthalocyanine (ZnPc) can induce intramolecular G-quadruplex telomeric structure transition from the antiparallel to parallel form. It has been shown...
that 80 µm L⁻¹ ZnSO₄ helps to maintain and shorten the telomere length of hepatocytes L-02 and hepatoma cells SMMC-7721.² It was found that the molecular mechanism might be related to inhibition of telomerase activity in SMMC-7721 cells. It might also be the result of p53 protein activation, a zinc-binding transcription factor that responds to multiple forms of stress and controls proliferation, survival, DNA repair and cells differentiation. Therefore, supplementation of zinc could affect the transcription of DNA and telomere length by activation of p53 in hepatoma cells.⁴⁶ According to the aforementioned reports, in the present study, it was suggested that ADSCS aging prevention in the presence of ZnSO₄ may be due to changes in TERT gene expression, telomerase activity, telomere structure and telomere length.

In conclusion, the findings of this study showed that the TERT gene expression significantly increased while the percentage of senescent cells decreased under ELF-EMF of 50 Hz, 20 mT in ZnSO₄-treated ADSCs when compared to untreated cells. The results suggested that 50 Hz, 20 mT magnetic field could affect the senescence and cellular aging due to TERT gene expression; although, the detailed mechanisms are still unclear. More studies are recommended on the effects of magnetic fields on telomere length and telomerase activity of stem cells.

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