Non thermal effects of radiofrequency electromagnetic field exposure on neural cells

Rosaria Grasso1,4*, Rosalia Pellitteri2, Santi Armando Caravella3, Francesco Musumeci1,4, Giuseppina Raciti3, Agata Scordino1,4, Giovanni Sposito3, Antonio Triglia1, and Agata Campisi1

1University of Catania, Department of Physics and Astronomy “Ettore Majorana”, 95123 Catania, Italy
2Italian National Research Council, Institute for Biomedical Research and Innovation, 95126 Catania, Italy
3University of Catania Department of Drug Sciences, Section of Biochemistry, 95123 Catania, Italy
4National Institute for Nuclear Physics, Laboratori Nazionali del Sud, 95123 Catania, Italy
5Temix Engineering s.r.l., 95039 Trecastagni, Italy

Abstract. The non-thermal mechanisms, underlying the damage induced on human cells by radiofrequency electromagnetic fields (RF-EMFs), are still unclear and only few studies reported about the effect of RF-EMFs on self-renewal of neural progenitor cells. In this research, we investigated the influence of low-intensity RF-EMFs on Olfactory Ensheathing Cell (OEC) cultures, typical glia cells showing characteristics of stem cells. Cell cultures were exposed, in far-field condition, at 900 MHz continuous and amplitude modulated EMFs for 10, 15 and 20 min at 37°C. The expression of OEC marker (S-100), stem cell marker (Nestin), cytoskeletal proteins (GFAP and Vimentin), apoptotic pathway activation by Caspase-3 cleavage and cell viability, were evaluated. Surprisingly 20 min of exposure to continuous or amplitude modulated 900 MHz EMF induced a different and significant decrease in cell viability, some dynamic changes in the expression of the analysed markers and in the activation of the apoptotic pathway.

1 Introduction

Starting from the Industrial Revolution, the influence of the human being on his environment has gradually grown, introducing significant and perhaps irreversible changes. Just let consider the alterations of the climate that are ascribed to human activities such as the replacement of forests with cultivated areas, the intensive breeding and the increase in the concentration of greenhouse gases and aerosols in the atmosphere. However, the most drastic alteration of the natural conditions occurred with the introduction, in our vital environment, of a huge amount (compared to the pre-industrial period) of electromagnetic fields characterized by a wide frequency range and by intensities some orders of magnitude greater than the natural background. An evident phenomenon in advanced societies is the fast increase in the use of mobile phones and the spread of wireless methodologies in the management of information through computers. These technologies are based on the use of so-called radio-frequency electromagnetic fields (RF-EMFs) having frequencies ranging from hundreds KHz to some GHz. These fields, practically absent on the Earth's surface before the Industrial Revolution, today have permeated totally our living space.

Until a few decades ago it was believed that the non-ionizing electromagnetic radiation was able to induce effects on the biological matter only through overheating of the tissues, inducing the so-called thermal effects. This induced to define the concept of Specific Absorption Rate (SAR), that is the energy absorbed by tissue when exposed to RF-EMF. The SAR value is related to the properties of tissue (electrical conductivity, density), sample volume and the root mean square of the impinging electric field. The SAR definition includes the effect of electromagnetic field frequency only indirectly, through the changes of tissue properties, but does not evaluate the possible effect of the electromagnetic field modulation. On this basis, the security protocols of various organizations, such as the WHO and ICNIRP, responsible for assessing the risks associated with the use of modern technologies, have been developed [1]. However, more recently, experimental evidence of the ability of RF-EMF to generate non-thermal biological effects at the cellular level is increasing [2-5] even if the heterogeneity of the RF-EMF, used for various technological applications, makes it difficult to find a cause-effect correlation.

In the paper, we present some results about our investigation on the non-thermal effect induced by a continuous RF-EMF at 900 MHz (CW 900 MHz) on a particular glial cell type, the Olfactory Ensheathing Cells. The interest versus this cell line is increasing for several reason: an information transfer has been observed in the olfacto-hippocampal network [6]; it appears to be a promising tool for cellular therapy in spinal cord injury [7] and axonal growth [8]; it can stimulate axonal regeneration and functional restoration in the lesion of...
2 Materials and methods

2.1 Materials

Normal Goat Serum, 200 mM L-glutamine, collagenase Penicillin-Streptomycin solution, heat inactivated Foetal Bovine Serum Dulbecco’s Modified Eagles Medium, 0.05% trypsin-0.02% EDTA solution, were purchased by Invitrogen (ThermoScientific, Milan, Italy). Monoclonal mouse anti Vimentin clone V9 (cod. 2012-03) and polyclonal rabbit anti GFAP (cod. 2015-02) were from DAKO. 1-(4,5-Dimethylthiazol-2-yl)-3,5-diphenylformazan (MTT), were from Sigma-Aldrich (Milan, Italy). Monoclonal mouse anti Vimentin clone V9 (cod. 2012-03) and polyclonal rabbit anti GFAP (cod. 2015-02) were from DAKO. 1-(4,5-Dimethylthiazol-2-yl)-3,5-diphenylformazan (MTT), were from Sigma-Aldrich (Milan, Italy).

2.2 Animals and OEC cultures

The mouse pups were provided by Envigo RMS s.r.l. Italy (stock: C57BL6j). Animals were kept in a controlled environment (23 ± 1°C, 50 ± 5 % humidity) with a 12 h light/dark cycle with food and water available ad libitum. The animal experiments were carried out according to the Italian Guidelines for Animal Care (D.L. 116/92 and 26/2014), which are in compliance with the European Communities Council Directives (2010/63/EU) and were approved by the Ethical Committee at the University of Catania (Catania, Italy).

Primary OECs were isolated from mouse P2 olfactory bulbs and purified as previously described [10].

2.3 RF-EMF exposure

The RF- EMF was emitted by a double horn antenna (ETS-Emco- 3115) plugged to an Agilent-8648D signal generator. The amplitude modulation of sinusoidal waveform was obtained by an external oscillator (Philips-PM5127). Two different irradiation conditions at the same amplitude around 6V/m were used: continuous radiofrequency at 900MHz wave (CW 900MHz); cells maintained in water thermal baths, exposed to CW 900 MHz for 10, 15 and 20 minutes (CW 900 MHz); cells maintained in water thermal baths, exposed to AM 900 MHz for 10, 15 and 20 minutes (AM 900 MHz). The MTT assay showed no significant change in cell viability respect to sham versus control cells at 10, 15 and 20 minutes exposure time. The continuous and, in particular, amplitude modulated 900 MHz were able to induce a significant decrease in cell viability respect to respective shams only when the exposure was for 20 minutes.

For what concern immunocytochemistry analysis, the exposure to RF-EMFs for 10 minutes did not induce effect on cell cultures. Therefore, we report about the quantitative analysis of GFAP (fig. 1) and Vimentin (fig. 2) expression levels in OECs un-exposed and exposed to EMFs for 15 and 20 minutes. The control cell showed a low number of GFAP positive cells (fig. 1). The absence of CO2 (Sham) for 20 min induced a light, even if significant, increase in the GFAP positivity cell. The exposure to CW 900 MHz, both for 15 min and 20 min, induced a highly significant increase in GFAP expression level. Instead, the exposure to AM

3 Results and discussion

This study represents the first step of a more complex one, involving a long and deep investigation. We assessed the effect of electromagnetic field exposure on the main cytoskeletal components which are more relevant in terms of OEC physiology. We analysed cytoskeleton protein expression, as it represents the first signal of cellular alteration. It is well known that the cytoskeleton plays a key role in the cell architecture and it is responsible for intracellular trafficking of endogenous and exogenous molecules and stimuli.

We investigated cell viability using MTT assay and, through immunocytochemical procedures, the expression of Glial Fibrillary Acidic Protein (GFAP), Vimentin and Nestin, that are proteins implicated in the formation of Intermediate Filaments, as well as S100 protein, OEC marker, and the activation of apoptotic pathway by analysing caspase-3 cleavage.

For the experiments, cells were divided in four groups: control cells maintained into the incubator at 37°C in ambient of humidified air and CO2 (95%-5%), to verify the normal cellular status (Ctrl); cells maintained in water thermal baths at 37°C for 10, 15 and 20 minutes (Sham); cells maintained in water thermal baths, exposed to CW 900 MHz for 10, 15 and 20 minutes (CW 900 MHz); cells maintained in water thermal baths, exposed to AM 900 MHz for 10, 15 and 20 minutes (AM 900 MHz).

Here we will present, schematically, only the results concerning cell viability, GFAP and Vimentin.

The MTT assay showed no significant change in cell viability in sham versus control cells at 10, 15 and 20 minutes exposure time. The continuous and, in particular, amplitude modulated 900 MHz were able to induce a significant decrease in cell viability respect to respective shams only when the exposure was for 20 minutes.

For what concern immunocytochemistry analysis, the exposure to RF-EMFs for 10 minutes did not induce effect on cell cultures. Therefore, we report about the quantitative analysis of GFAP (fig. 1) and Vimentin (fig. 2) expression levels in OECs un-exposed and exposed to EMFs for 15 and 20 minutes.

The control cell showed a low number of GFAP positive cells (fig. 1). The absence of CO2 (Sham) for 20 min induced a light, even if significant, increase in the GFAP positivity cell. The exposure to CW 900 MHz, both for 15 min and 20 min, induced a highly significant increase in GFAP expression level. Instead, the exposure to AM

Nervous System [9]. To better understand the effect induced by modulation, we exposed cell cultures also to an amplitude modulated 900 MHz (AM 900MHz). Moreover we evaluated the effect of exposure time (10, 15 and 20 minutes).
900 MHz, both for 15 min and 20 min, induced a decrease in GFAP expression level respect to cell cultures exposed to CW 900 MHz, and, at 20 min, also respect to sham.

Fig. 1. % of GFAP positive OECs on control, sham, CW 900 MHz, AM 900 MHz after exposure time of (○) 15 min, (□) 20 min. Values of four separated experiments are reported as mean ± SD.

Both control and sham showed a low positivity for Vimentin (fig. 2). The exposure to continuous 900 MHz induces an high significant increase in Vimentin positive OECs. A recovery effect seems to be activated after 15 min exposure time. The AM 900 MHz induced a highly significant increase of the Vimentin positive cells number at 15 min exposure time and its drastic decrease at 20 min not only respect to the cell cultures exposed to CW 900 MHz but also respect to control and sham.

Fig. 2. % of Vimentin positive OECs in control, sham, CW 900 MHz, AM 900 MHz after exposure time of (○) 15 min, (□) 20 min. Values of four separated experiments are reported as mean ± SD.

In conclusion, these results, together to others not showed here, highlighted that continuous or amplitude modulated 900 MHz exposure was able to differently alter the cytoskeletal proteins at various levels and that these effects were function of exposure time. In particular, the results on GPAF expression levels indicated that the exposure to CW 900 MHz enhances the OECs ability to differentiate versus astroglial cell type, while AM 900 MHz exposure for 20 min induces a decrease respect to the Sham. Moreover, the experimental results on Vimentin expression levels pointed out that the exposure to CW 900 MHz and AM 900 MHz can induce different effect on the cytoskeleton remodelling. It is worth to note that the latter one, highlighted through Vimentin, activates signalling pathways having got a role both as cell viability modulator and cell structure constituents.

References
1. High Frequency review. ISBN 978-3-934994-10-2 (2009)
2. S. Cucurachi, W.L. Tamis, M.G. Vijver, W.J. Peijnenburg, J.F. Bolte, G.R. de Snoo. Environ. Int. 51, 116 (2013)
3. A. Campisi, M. Gulino, R. Acquaviva, P. Bellia, G. Raciti, R. Grasso, F. Musumeci, A. Vanella, A. Triglia, Neurosci. Lett. 473, 52 (2010).
4. C. Chen, Q. Ma, C. Liu, P. Deng, G., Zhu, L. Zhang, M. He, Y. Lu, W. Duan, L. Pei, et al., Sci. Rep. 4, 5103 (2014).
5. M. Eghlidospour, A. Ghanbari, S.M.J. Mortazavi, H. Azari, Anat. Cell. Biol. 50, 115 (2017).
6. B. Gourévitch, L.M. Kay, C. Martin, J. Neurophysiol. 103, 2633 (2010).
7. R.M. Gómez, M.Y. Sánchez, M. Portela-Lomba, K. Ghotme, G.E. Barreto, J. Sierra, J., M.T. Moreno-Flores, Glia. 66, 1267 (2018).
8. F. Chehrehasa, L.C. Windus, J.A. Ekberg, S.E. Scott, D. Amaya, A. Mackay-Sim et al., Mol. Cell. Neurosci. 45, 277 (2010).
9. S. C. Barnett, J.S. Riddell, J. Anat. 204, 57 (2004)
10. R. Pellitteri, M. Spatuzza, A. Russo, S. Stanzani, Neurosci. Lett. 417, 24 (2007).
11. A. Campisi, M. Spatuzza, A. Russo, G. Raciti, A. Vanella, S. Stanzani, R. Pellitteri, Neurosci. Res. 72, 289 (2012).
12. R. Pellitteri, R. Bonfanti, M. Spatuzza, M.T. Cambria, M. Ferrara, G. Raciti, A. Campisi, Mol. Neurobiol. 54, 6785 (2017).