

Research Paper

Microwave Irradiation Affects Gene Expression in Plants

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KEY WORDS

tomato, microwave, non-ionizing radiation, bZIP, MSRC, EMF

ABBREVIATIONS

LebZIP1 *Lycopersicon esculentum* basic leucine zipper
MSRC Mode Stirring Reverberation Chamber
TEM cell transverse electromagnetic cell
EMF electromagnetic field

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ABSTRACT

The physiological impact of nonionizing radiation has long been considered negligible. However, here we use a carefully calibrated stimulation system that mimics the characteristics (isotropy and homogeneity) of electromagnetic fields present in the environment to measure changes in a molecular marker (mRNA encoding the stress-related *bZIP* transcription factor), and show that low amplitude, short duration, 900 MHz EMF evokes the accumulation of this mRNA. Accumulation is rapid (peaking 5–15 min after stimulation) and strong (3.5-fold), and is similar to that evoked by mechanical stimulations.

INTRODUCTION

High frequency nonionizing radiation is becoming increasingly common in the environment because of the exponential use of mobile phone technology and wireless communication devices. While many reports point out its lack of effects on living organisms,^{1,2} it can be argued that most studies have used unsuitable stimulation devices, primarily TEM-cells,³ custom-made tools^{4,5} or even commercial cell phones,⁶ and address the problem at a very general level with little concern for underlying molecular-level events.⁷ Most stimulation devices are inadequate, since they emit a signal as a plane wave with a fixed polarization and incidence. Studies have therefore been undertaken to develop appropriate technology for EMF studies on living systems by adapting reverberating chambers, which are widely used in acoustics. This has led to the Mode Stirring Reverberation Chamber (MSRC), a facility specially designed to create isotropic and homogeneous EMF that irradiates the subject from all directions. The reflections of the original signal on the metallic walls of the chamber (that isolates the subject from external radiation) randomize the polarization of the EMF without influencing its amplitude (homogeneity). This mimics Nature, where there are multiple reflections and diffractions of EMF from buildings, mountains and trees. Thus, this equipment is extremely well suited to study the effect of EMF on life.

Two aspects of these biological experiments need special attention: the organism itself and the parameter, preferably a molecular marker,⁸ to be studied. Plants may be appropriate experimental subjects, in the sense that they are highly sensitive to environmental signals⁹⁻¹¹ and plant studies raise less emotional concern than studies on animals or humans. Moreover, radiation from a GSM telephone or from a 105 GHz Gunn oscillator has been shown to have an effect comparable to that of a variety of environmental stimuli (such as manipulation stress, drought, wind and cold shock) in inducing physiological modifications (production of numerous epidermal meristems in the hypocotyls) or modifications of the proteome (pI shift, appearance or disappearance of a spot in 2D electrophoresis) in flax seedlings;¹²⁻¹⁵ modifications of the proteome have also been observed in *Arabidopsis* seedlings subjected to stimuli such as cold shock or radiation from a GSM telephone.¹⁵ Here, we have studied the effect of mobile-phone intensity microwave radiation on another type of plant (the tomato), taking a particular care to the conditions of irradiation and studying the accumulation of a stress-related transcript (mRNA) that responds very rapidly to even small environmental stimulations.¹⁶ We are going to show that such microwave radiation enhances accumulation of the transcript encoding a specific, wound-related transcription factor, *LebZIP1*.¹⁷

MATERIAL AND METHODS

Plant culture and treatment. Tomato plants (*Lycopersicon esculentum* cv VFN-8) were germinated in the greenhouse and transferred to an EMF-permeable culture chamber and grown under controlled conditions (Light/Dark 16 h/8 h 26°C/21°C, light intensity of 175 $\mu\text{mol}\cdot\text{s}^{-1}\cdot\text{m}^{-2}$ at plant level) for three weeks, until the 4th terminal leaf had formed. The culture chamber containing the plants was transferred to the stimulation chamber at least 18h before treatment. Stimulation was made using the MSRC (Fig. 1A). This facility is a large room (8.4 x 6.7 x 3.5 m, about 195 m³) enclosed in double-layered metal walls which act as a Faraday cage to protect the experiment from external (environmental) EMF background. For plant stimulation, a single frequency (900 MHz) was produced by a signal synthesizer (Anritsu model 68147C), verified with a signal analyser (Anritsu model MS2665C) and emitted into the chamber with a log-periodic emission antenna. A rotary stirrer was used to create different patterns of multiple reflections on the chamber walls, thus randomizing the polarization of the electromagnetic waves. The resulting electromagnetic field is statistically isotropic and homogeneous within a defined “working” volume. The stimulation (5 V/m, 10 min) was given to the plant in the middle of the light period. The 4th terminal leaf was collected at various times after the end of the stimulation and immediately frozen in liquid nitrogen. Control plants were collected before stimulation. Because of the limited size of the culture chamber, only one plant could be used for each time point. For some experiments, the culture chamber was shielded in a polymer mesh covered with an aluminium layer that causes a 45 dB signal attenuation at 900 MHz (more than 87%).

RNA isolation and quantitative PCR. RNA was isolated from frozen tissue using Tri-Reagent (Sigma) and total RNA (1 μg) was used to drive cDNA synthesis (Advantage RT for PCR, BD Bioscience) for 1 h at 42°C both according to the manufacturers’ instructions. The cDNA was diluted 5-fold and used as DNA template for quantitative PCR analysis (Two step qPCR Mastermix Plus for SYBR-Green, Eurogentech). The chosen primers (sense: 5'-GGGATGGA-GAAGTTTGGTGGTGG-3' Anti-sense:5'-CTTCGACCAAGGG-ATGGTGTAGC-3') amplify just *LebZIP1* cDNA¹⁷ (Genbank accession number AF176641). The reactions were performed and analysed using the 2^{- $\Delta\Delta\text{Ct}$} method¹⁸ with actin as internal control, and values calculated relative to the nonexposed control plants. Actin and *LebZIP1* fragments amplify with the same efficiency in our experimental conditions (data not shown).

RESULTS AND DISCUSSION

Experiments were conducted inside the MSRC within the working volume (Fig. 1A, grey area) containing the plant culture chamber, where the EMF was statistically isotropic and homogeneous (Fig. 1B, a) in contrast to a polarized, nonhomogeneous EMF (Fig. 1B, b). These characteristics were determined by measuring the values of the standard deviations (σ_x , σ_y and σ_z) of the three spatial components

Figure 2. Field characteristics in the plant culture chamber. The tunable antenna was used to generate EMF of 5 Vm⁻¹ from 800–1000 MHz (A), or 893–911 MHz (B). (A) Standard deviation of amplitude for the total field (σ_{xyz} , bold line) and for each of the Cartesian axes (σ_x , σ_y , σ_z); dashed line: 3dB limit specified by the IEC 61000-4-21 standard. (B) Effective values measured within the nonshielded (○) and shielded (△) culture chamber. Dashed line: 5 V/m input signal.

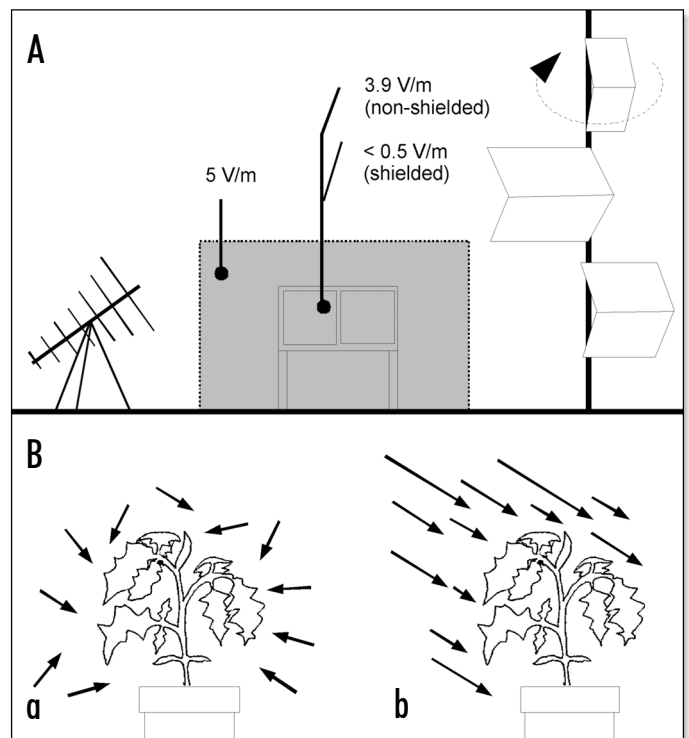
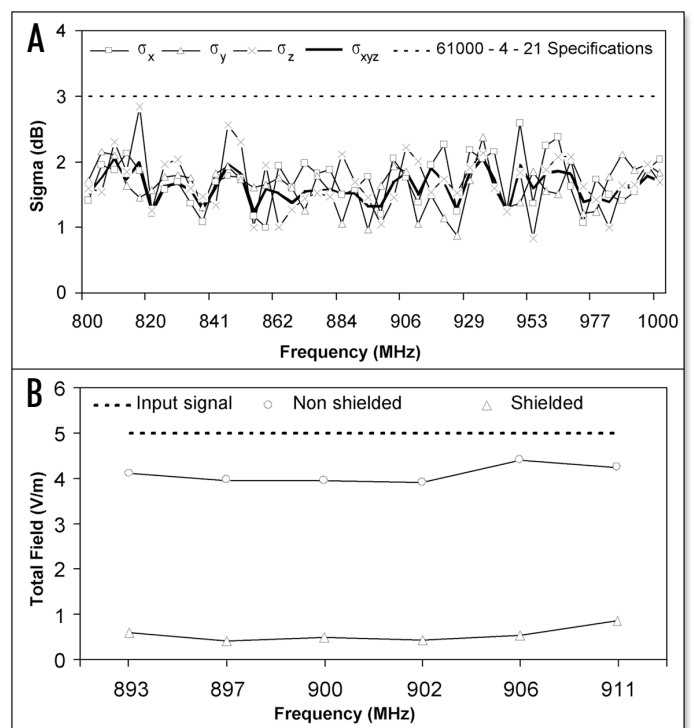


Figure 1. (A) The mode stirring reverberation chamber. This is a large room with metal walls (dark lines) to exclude external EMF, an antenna (lower left) to emit tunable EMF, a rotary stirrer to make the EMF homogeneous (right side) and a plant culture chamber placed within the working volume (gray area). (B) Schematic representation of EMF types. a, non-polarized (isotropic) and homogeneous field, where the field components align in all possible directions and the field has the same amplitude at all points. b, polarized, nonhomogeneous field, where the field components align in a single direction while the amplitude varies (heterogeneity).



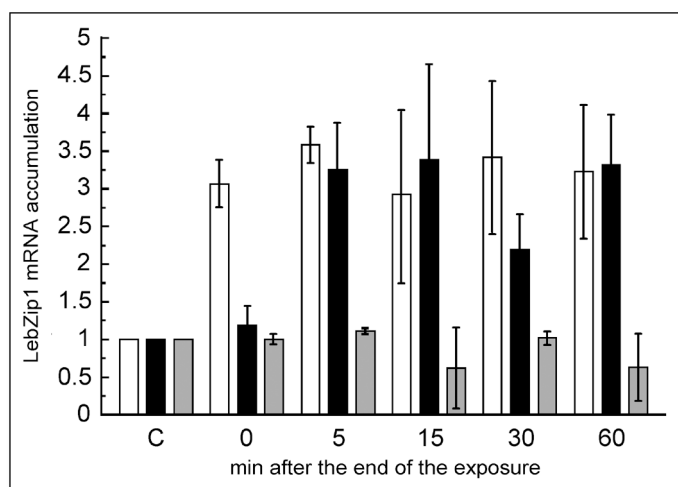


Figure 3. Accumulation of *LebzIP1* transcript after EMF-stimulation in the nonshielded culture chamber. Plant show either an immediate response (white bars) or a 5 min delayed response (black bars). Plants stimulated in the shielded culture chamber (gray bars). Each value is expressed relative to the nonexposed control (C) and normalized to the actin mRNA and is the average of at least 3 independent repetitions \pm the standard error.

of the electric field in 8 locations within the culture chamber, and the σ_{xyz} calculated from these 24 components.

To verify the isotropy and homogeneity of the EMF, we excited the chamber with single frequency signals varying from 800 to 1000 MHz and made 18 measurements of the generated field per stirrer rotation for each tested frequency and the results are shown in Figure 2A. The standard deviation for each of the three spatial components (σ_x , σ_y and σ_z) and of the combined (σ_{xyz}) never exceeded the 3 dB limit listed in the 61000-4-21 specification,¹⁹ proving that the field quality is not affected by interactions with the culture chamber walls. However, they do cause a signal attenuation of about 20%, since the input amplitude was 5 V/m (Fig. 2B dashed line), whereas the measured amplitude inside the chamber was 3.9 V/m (Fig. 2B, “non-shielded”). Furthermore, the shielding material decreased the field amplitude by 87% inside the chamber 0.5 V/m (45 dB attenuation) at 900 MHz (Fig. 2B, “shielded”). In the unshielded culture chamber, the plants were therefore exposed to an EMF similar to that occurring in the natural environment in terms of amplitude, isotropy and homogeneity, indicating that the MSRC is an outstanding facility to generate such conditions.

Exposure of the plant to EMF induced a rapid (maximum after 5–15 min) and strong (3.5-fold) accumulation of the stress-related *LebzIP1* mRNA in the 4th terminal leaf (Fig. 3). In some experiments, the accumulation began immediately after the end of the stimulation (Fig. 3, white bars), while in others there was a short delay (Fig. 3, black bars). In all cases, the response was maximal at 5–15 min after the end of stimulation, in some cases (black bars) declined somewhat at 30 min, but generally remained at high levels until 60 min. Since, space limitations within the MSRC precluded the use of more than 1 plant (1 leaf) per time point, and since each experiment used a different batch of plants, we are amazed at the relative consistency of the results. When plants were placed in the shielded culture chamber, no significant accumulation of *LebzIP1* transcript was seen (Fig. 3, gray). These results indicate that: (1) the cellular responses are directly linked to exposure of plants to the EMF; and (2) that the

remaining EMF (0.5 V/m) present in the shielded culture chamber is insufficient to evoke *LebzIP1* mRNA accumulation. Although the treated tissue did not display any apparent damage, the rapidity and amplitude of the response are comparable to those observed after strong stimulation such as flaming.¹⁷

These results are quite surprising and strongly question the mechanism of interaction between the plant and the EMF. The energy associated with the EMF radiation is extremely low²⁰ and insufficient to evoke plant defence mechanisms involving the genesis of free radicals or molecule ionization. Variations of cytosolic Ca^{2+} concentration have been implicated after EMF stimulation¹³ and might constitute the initial signal that evokes the observed molecular responses. Thermal effects²¹ are unlikely to arise considering the frequency (900 MHz) and the very low power (0.1 W) dissipated in the large volume of the MSRC. The response is therefore triggered by a mechanism which is unlikely to be based on a simple energy transfer from the wave to the plant. The amplitude (3.9 Vm⁻¹) of the signal that evoked this rapid stress-related response is within the range used for mobile phone communication or to the EMF background present in an urban environment.

The major points arising from this study are that high frequency low amplitude EMF cause enhanced expression of at least one plant-wound gene. This response is reproducible and exceedingly rapid, in all instances peaking within 15 min following the end of exposure. This rapid response and its suppression by an EMF-proof shield allows us to make a formal link between the EMF stimulation and the accumulation of the *bZIP* mRNA.

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