NUMERICAL ANALYSIS OF HEAT TRANSFER IN MULTILAYERED SKIN TISSUE EXPOSED TO 5G MOBILE COMMUNICATION FREQUENCIES

Jagbir Kaur1,*, S.A. Khan2

ABSTRACT

Rapid growth in wireless communications has triggered the advent of 5G mobile communication systems. The use of millimeter waves (30-300 GHz) in 5G system has generated global concern about its biological safety. In present paper, we have numerically analyzed the heat transfer in a 3D multilayered skin tissue exposed to 5G frequencies. The numerical scheme comprises coupling of solution of Maxwell's equation of wave propagation within tissue to Pennes' bioheat equation. Temperature variations are analyzed at 28 GHz, 38 GHz, and 60 GHz. Additionally, electric field and specific absorption rate distribution are also studied. Highest values of electric field and specific absorption rate are estimated in epidermis layer of skin tissue. For all considered frequencies, highest transient temperature (37.36°C) is predicted in subcutaneous fat layer of the skin. However, the steady state temperature is nearly same as core body temperature (37°C). The results show that 5G mobile phones do not cause any thermal damage to the skin tissue and can be considered safe.

Keywords: Millimeter Waves, Bioheat Transfer, Multilayered Skin Tissue, 5G Mobile Phones

INTRODUCTION

Mobile phones have been an integral part of our lifestyle nowadays. The continuous increasing demand for high mobile data traffic and the proliferation of applications requiring high data rates have triggered the need for 5G mobile communication systems. As the data traffic is increasing tremendously, the sub 3 GHz is becoming heavily crowded. On the other hand, millimeter wave (mmW) frequencies (30-300 GHz) are considered promising for 5G communication networks due to the massive amount of raw bandwidth and potential multigigabit-per-second (Gb/s) data rates [1]. This “next generation” mobile communication system will be capable of responding to an avalanche of traffic, an upsurge in the number of connected devices, and the large variety of use cases. Despite numerous benefits, 5G mobile communication systems have generated a concern among the researchers if they are biologically safe. The electromagnetic (EM) energy carried by mmWs is transferred to the biological tissue, which results in ion acceleration and collisions among molecules. Consequently, the local temperature of the tissue rises [2]. A small change of 1-5°C in tissue may cause malfunctions of organs [3].

Since mmWs have very short wavelength, their penetration in the biological medium is superficial [4, 5]. The shallow penetration of 5G frequencies suggests that they can only infiltrate the skin layer of human body. Thus the EM energy associated with the mmWs will be absorbed within the skin layer; and the temperature changes due to absorption of the radiation will occur in skin. Thermal changes in humans due to RF and microwaves have been reported in many previous studies [6-9]. Kojima et al. studied the ocular damage due to thermal changes in eyes exposed to mmWs [10]. Stewart and co-workers investigated skin heating due to prolonged exposure to 94 GHz [11].
Sasaki et al. used Monte Carlo simulations to investigate the temperature changes in skin tissue due to GHz exposure [12]. However, more investigations are needed for comprehensive knowledge of thermal effects of mmWs.

The present study provides the detailed analysis of thermal response of a three-dimensional multilayered skin tissue to 5G mobile communication frequencies. The frequencies selected for the analysis viz. 28 GHz, 38 GHz, and 60 GHz, are promising frequencies for 5G mobile communications [13, 14]. The skin tissue is exposed to the mmWs from the microstrip antenna of the mobile phone. The output power transmitted by the half wave dipole microstrip antenna is compliant with the power density limit (10 W/m²) set by International Commission on Non-Ionizing Radiation Protection (ICNIRP) [15]. The obtained temperature variations suggest whether 5G mobile phones are safe for the users.

METHODS

A 3D skin model comprising four different layers viz. epidermis, dermis, subcutaneous fat, and inner tissue (muscle) is exposed to 5G mobile communication frequencies (28 GHz, 38 GHz, and 60 GHz) (Fig.1). The microstrip antenna is placed 2 cm away from epidermis layer. The EM energy is transmitted by the lumped port of the antenna and this energy is absorbed by the multilayered skin tissue. Physical and dielectric properties of the different layers of skin tissue are listed in Table 1 and Table 2 respectively. A finite element method (FEM) based software; COMSOL Multiphysics has been employed to numerically analyze the electric field, SAR and temperature variations within the tissue. The analysis is carried in two steps: (a) electromagnetic analysis and (b) bioheat analysis. In electromagnetic analysis electric field and SAR are determined, and the temperature variations are studied in bioheat analysis.

Figure 1. Schematic geometry of multilayered skin tissue with microstrip patch antenna.

Table 1. Physical and thermal properties of skin layers [16]

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Specific Heat, $C_p$ (J/kg. K)</th>
<th>Thermal conductivity, $K$ (W/m.K)</th>
<th>Mass density, $\rho$ (kg/m³)</th>
<th>Perfusion rate, $\omega$ (1/s)</th>
<th>Thickness, $x$ (m)²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epidermis</td>
<td>3600</td>
<td>0.235</td>
<td>1190</td>
<td>0</td>
<td>80*10E-6</td>
</tr>
<tr>
<td>Dermis</td>
<td>3300</td>
<td>0.445</td>
<td>1116</td>
<td>1.5</td>
<td>0.002</td>
</tr>
<tr>
<td>Subcutaneous Fat</td>
<td>2700</td>
<td>0.185</td>
<td>971</td>
<td>1.5</td>
<td>0.01</td>
</tr>
<tr>
<td>Inner tissue (muscle)</td>
<td>4000</td>
<td>0.5</td>
<td>1,000</td>
<td>1.5</td>
<td>0.03</td>
</tr>
</tbody>
</table>
Electromagnetic Analysis

Electric field distribution and SAR are calculated within the exposed tissue using electromagnetic equations. Following assumptions are made for simplification of the problem:
1. Electromagnetic wave propagation is modeled in three dimensions.
2. Interaction between skin tissue and the electromagnetic waves takes place in open region.
3. The free space surrounding the calculated domain is truncated by scattering boundary condition.
4. The dielectric properties of skin tissue are assumed constant for each layer.

As the skin tissue is exposed to mmWs, EM waves propagate inside the skin tissue. Maxwell’s equation governs the propagation of EM waves inside a biological tissue, and electric field (E) inside the tissue can be determined by equation [18, 19]:

$$\nabla \times \frac{1}{\mu_r} (\nabla \times E) - k_0^2 (\varepsilon_r - \frac{j\sigma}{\omega\varepsilon_0})E = 0$$  \hspace{1cm} (1)

where $\mu_r$ is relative magnetic permeability, $\varepsilon_r$ is relative permittivity, $\varepsilon_0 = 8.8542 \times 10^{-12}$ F/m, $k_0$ is the free space wave number, $\sigma$ is conductivity of tissue and $\omega$ is angular frequency.

Boundary conditions

The microstrip patch antenna radiates electromagnetic energy which strikes the skin tissue with a specific power. At the bottom of patch antenna; the stimulator employs a lumped port between two patches in order to generate an electromagnetic field. Lumped port boundary condition is defined as [20]:

$$Z_{in} = \frac{V_1}{I_1} = \frac{E_1}{l}$$  \hspace{1cm} (2)

where $Z_{in}$ is the input impedance, $V_1$ is the voltage along the edges, $I_1$ is the electric current magnitude, $E_1$ is the electric field along the source edge, and $l$ is the edge length.

A perfect electric conductor boundary condition is applied to the side walls of the patch, so that the lumped port acts as a cavity [21]:

$$n \times E = 0$$  \hspace{1cm} (3)

In order to truncate the space outside the calculated domain, scattering boundary condition is applied on outer edges of modeled region (Fig. 2), as

$$n \times (\nabla \times E) - jkn \times (E \times n) = -n \times (E_0 \times jk(n - k)) \exp(-jk\cdot r)$$  \hspace{1cm} (4)

where $k$ is the wave number, $n$ is normal vector, $j = \sqrt{-1}$, and $E_0$ is the electric field intensity of incident plane wave.

The existence of electric field within skin model conveys that EM energy is deposited in the tissues. The energy absorbed by the tissues is expressed in terms of SAR. The local SAR can be calculated using the relation [22, 23, 24]

$$SAR = \frac{\sigma|E|^2}{\rho}$$  \hspace{1cm} (5)

where $\sigma$ is the conductivity and $\rho$ is mass density of the tissue, and $E$ is the electric field within the tissue exposed to 5G frequencies.
Table 2. Dielectric properties of skin layers at different frequencies [25]

<table>
<thead>
<tr>
<th>Layer</th>
<th>28 GHz</th>
<th>38 GHz</th>
<th>60 GHz</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Permittivity $\varepsilon_r$</td>
<td>Conductivity $\sigma$ (S/m)</td>
<td>Permittivity $\varepsilon_r$</td>
</tr>
<tr>
<td>Epidermis</td>
<td>3.39</td>
<td>0.3</td>
<td>3.37</td>
</tr>
<tr>
<td>Dermis</td>
<td>4.28</td>
<td>18.2</td>
<td>4.24</td>
</tr>
<tr>
<td>Subcutaneous fat</td>
<td>3.76</td>
<td>7.1</td>
<td>3.73</td>
</tr>
<tr>
<td>Inner tissue (muscle)</td>
<td>24.4</td>
<td>33.6</td>
<td>19.1</td>
</tr>
</tbody>
</table>

Bioheat Analysis

The heat transferred to the skin tissue is calculated using Pennes’ bioheat equation, the pioneer mathematical model for analyzing heat conduction in the biological tissues. This mathematical model has been widely used in many investigations to predict the temperature changes in biological medium [16, 17, 26, 27, 28]. It is expressed as [29]:

$$\rho C_p \frac{\partial T}{\partial t} = \nabla \cdot (K \nabla T) + \rho_b C_{p,b} \omega_b (T_b - T) + Q_m + Q_{ext}$$  

(6)

where $\rho$, $C_p$, and $K$ are the density, specific heat at constant pressure, and the thermal conductivity of tissue, respectively. $\rho_b$ and $C_{p,b}$ are the density and specific heat (at constant pressure) of the blood, respectively. $\omega_b$ is the blood perfusion rate; $T$ and $T_b$ are tissue and blood temperatures, respectively. $Q_m$ is metabolic heat generation in the tissue and $Q_{ext}$ is the external heat source term (electromagnetic heat-source density). The term $\rho_b C_{b,b} \omega_b (T_b - T)$ describes the heat caused by convection and $\nabla \cdot (K \nabla T)$ represents the heat conduction inside the skin tissue. $Q_{ext}$ represents the resistive heat generated by the electromagnetic source and is expressed as [30]:

$$Q_{ext} = \frac{1}{2} \sigma |E|^2$$  

(7)

where $\sigma$ is the conductivity of the tissue and $E$ is the electric field within the tissue exposed to mmWs. $Q_m$ is neglected and assumed zero in the analysis, since we aim to evaluate the effect of electromagnetic energy associated with mmWs only.

Figure 2. Boundary conditions for numerical analysis.
Following assumptions are made for bioheat transfer analysis:

1. Each layer of tissue is a biomaterial with constant thermal properties.
2. No chemical reaction takes place in the considered domain.
3. The blood perfusion rate remains constant for each layer.
4. There is no change of phase of matter within the tissue.

In the analysis, bioheat equation (Eq. (6)) is coupled to the solution of Maxwell’s equation. Substituting values of $E$, $Q_{ext}$ is calculated using Eq. (7) and temperature rise is determined using Eq. (6).

**Boundary conditions**

The temperature of human body is assumed uniform and its initial value is $37^\circ C$. Therefore, initial condition for bioheat transfer is:

$$T = 37^\circ C \text{ at } t=0 \text{ and } \frac{dT}{dt} = 0 \text{ at } t=0$$  \hspace{1cm} (8)

No heat is exchanged between the calculated domain and the outer region. Hence, thermal insulation boundary condition is employed on the outer boundaries of the model (Fig. 2) [19]:

$$n \cdot (K \nabla T) = 0$$ \hspace{1cm} (9)

**Procedure for Simulation and Grid Independence**

In numerical simulation, the electric field at every point within calculated domain is determined in electromagnetic analysis. The value of electric field is used to calculate the heat generated by the electromagnetic source (Eq. (7)). The temperature at every point within domain is determined using electromagnetic field at that point and the steps are repeated until the steady state of temperature is reached. The numerical domain is discretized using tetrahedral elements with a total number of 140272 elements. A grid independence test is performed to approximate the number of elements above which the solution is independent of mesh elements (Fig. 3b). The grid independence is attained for the number of elements more than 73130. Higher numbers of elements are not tested due to lack of computational memory and performance.
RESULTS AND DISCUSSION

Validation

Considering the ethical restrictions, it is not possible to directly measure the electric field and temperature distribution inside the human body. Therefore, to validate the accuracy of the numerical simulation used in present study, a modified case of simulation results is verified against the numerical results with same geometric model obtained by Nishizawa and Hashimoto [24]. Fig. 4a shows 2D schematic geometry of human abdomen used in validation case. The model consists of three layers viz. skin, fat and muscle. The plane wave E (TM) of frequency 1300 MHz and power density 1 mW/cm², propagating along positive X-direction is incident on left side of model. The local SAR is plotted along cross sectional line through the abdomen; the graph (Fig. 4b) shows good agreement in SAR values between present solution and that of Nishizawa and Hashimoto. Small deviations occurring between two results may be due to different numerical scheme used and input dielectric properties of the tissues. This encouraging comparison gives confidence in accuracy of the simulation used in present study.

Figure 4. (a) 2-dimensional geometry for validation of model from Nishizawa and Hashimoto [24] (b) Comparison of local SAR distribution calculated by present numerical simulation and that by Nishizawa and Hashimoto

Electric Field and SAR Distribution

Penetration of electromagnetic radiation within the layers of skin tissue generates electric field inside the skin. Fig.5 shows the steady state electric field distribution in the multilayered skin tissue exposed to 5G mobile communication frequencies. The electric field distribution pattern is different in each exposure case; At 28 GHz and 38 GHz electric field spreads to larger areas than at 60 GHz and electric field hotspots are located in the vicinity of lumped port of microstrip antenna. Maximum steady state surface electric field at 28 GHz, 38 GHz and 60 GHz is 312.51 V/m, 478.41 V/m and 461 V/m respectively; which conveys that at 38 GHz maximum electromagnetic energy is transferred to the skin tissue. The electromagnetic energy absorbed by a biological tissue is expressed in terms of SAR (W/kg). The local SAR is calculated using Eq. (3). Since SAR at any point is directly proportional to electric field at that point, local SAR distribution pattern can be assumed similar to electric field distribution pattern.

Figure 5. Steady state electric field distribution in skin tissue at (a) 28 GHz, (b) 38GHz, and (c) 60 GHz.
Fig. 6 portrays the variations in electric field and local SAR at different frequencies along cross sectional line (arc length) passing through the different layers of skin tissue. The electric field and SAR curves show similar pattern at a particular frequency. Greatest electric field and local SAR (1050 V/m and 725 W/kg respectively) at 38 GHz and lowest (214.3 V/m and 42.2 W/kg respectively) at 60 GHz suggest that EM energy absorbed is maximum at 38 GHz and least at 60 GHz. In all the exposure cases, the peak values of electric field and SAR are estimated in epidermis layer of skin. Beyond epidermis, the magnitude of electric field and SAR falls rapidly; further than dermis, electric field and SAR are almost zero. The penetration depth of electric field is approximately 2 mm within the skin tissue. This result is in agreement with the fact that the energy absorbed due to 5G radiation exposure is superficial.

Figure 6. Electric field and local SAR distribution versus arc length at (a) 28 GHz, (b) 38 GHz, and (c) 60 GHz; 1: epidermis, 2: dermis, 3: subcutaneous fat, 4: inner tissue.
Temperature Distribution

The EM energy absorbed within the tissue produces oscillations in the molecules, which consequently raise the local temperature of the tissue. The temperature distribution within multilayered skin tissue is estimated using Pennes’ bioheat equation. Fig. 7 shows the steady state temperature distribution in skin tissue exposed to 5G frequencies from a microstrip patch antenna. The initial skin temperature considered for analysis is 34°C and the steady state temperature is slightly higher than core body temperature (37°C) for all exposure frequencies. However, the highest steady state temperature (37.006°C) is estimated at 60 GHz. This finding suggests that exposure to mmWs do not alter the skin temperature significantly, hence the possibility of thermal hazards due exposure to 5G frequencies is negligible.

![Steady state temperature distribution](image)

**Figure 7.** Steady state temperature distribution in skin tissue at (a) 28 GHz, (b) 38GHz, and (c) 60 GHz.

Fig. 8 illustrates the temperature variations at various instants of mmW exposure along cross sectional (arc length) line passing through different layers of skin tissue. The skin tissue responds immediately as it is exposed to the radiation. The temperature within the different layers rises very quickly; temperature achieves its peak value (37.36°C) at 2 s of the exposure. The highest temperature is estimated in subcutaneous fat layer. Low thermal conductivity of subcutaneous fat layer (Table 1) can be held responsible for occurrence of temperature peak in this layer. However the temperature falls quickly, and at 10 s of exposure the temperature falls below 37°C within subcutaneous fat. This behavior of temperature curve indicates the onset of blood perfusion, which sweeps the heat generated locally in the tissue. Thereafter, trivial changes in the temperature take place and it attains its steady state. This may be noted that the temperature behavior is almost identical at all the exposure frequencies, except small variations in the temperature curves at 2 s. This finding signifies that thermal response of the skin tissue does not vary significantly with 5G frequencies.
Figure 8. Temperature distribution at different instants of exposure period versus arc length at (a) 28GHz, (b) 38 GHz, and (c) 60 GHz; 1: epidermis, 2: dermis, 3: subcutaneous fat, 4: inner tissue.
Fig. 9 portrays the temperature changes in different layers w.r.t. the exposure duration at 60 GHz. The highest temperature (37.361 °C) is estimated in subcutaneous fat layer, which may be due to low thermal conductivity of this layer. The peak temperatures in epidermis and dermis layer are 37.205°C and 37.160°C respectively. The peaks in temperature curves occur at 2 s of exposure, thereafter, temperature drops very quickly below 37°C, which is core body temperature. Steady state in temperature (37.006°C) is attained after 140 s of exposure.

Figure 9. Temperature distribution versus exposure duration within different skin layers at 60 GHz.

Role of Dielectric Properties and Their Frequency Dependence

Dielectric properties play an important role in determining the energy deposition and hence, temperature elevation in the biological matter exposed to electromagnetic fields. Eq. (1), Eq. (5) and Eq. (7) show that electric field, SAR and the temperature variations in exposed tissue depend on the permittivity and conductivity of the tissue.

The complex valued, relative permittivity of a tissue is expressed as [31]:

\[ \varepsilon^* = \varepsilon^r - \frac{i\sigma}{\omega \varepsilon_0} = \varepsilon^r' - i\varepsilon^r'' \]  \hspace{1cm} (10)

where \( \varepsilon^r = \varepsilon_r \) and \( \varepsilon^r'' = \frac{\sigma}{\omega \varepsilon_0} \).

The complex conductivity and complex permittivity are related by [31]:

\[ \sigma^* = i\omega \varepsilon^* \]  \hspace{1cm} (11)

\( \varepsilon^* \) and \( \sigma^* \) vary with frequency and such a variation is called dispersion. Biological materials show many dispersions over a wide range of frequencies [32]. \( \alpha \) dispersion exhibited at low frequencies, which is associated with ionic diffusion process at cellular membranes. \( \beta \) dispersion, in hundreds of kilohertz region, is due to polarization of cellular membranes, proteins and other organic molecules. In gigahertz region, \( \gamma \) dispersion is due to polarization of water molecules in the tissue. The water content in biological tissues exists in two forms: bulk and hydrated. Hydration rate is very high in tissues and can reach 80% [32]. The volume fraction of bulk water 18% in epidermis and 28% in dermis [32]. Feldmen et al. showed that in GHz region, frequency dependence of dielectric properties is function of bulk water content [33]. Epidermis has lower water content, thus it has very small variations with
frequency. However, dermis layer has slightly higher frequency dependence, due to its bigger water content. Studies revealed that subcutaneous fat has water content of 18-28.7% [34, 35, 36]. Thus, frequency dependence of fat layer lies between that of dermis and epidermis.

Fig. 8 shows that the temperature curves for skin tissue are almost identical for all the exposed frequencies. This result is in agreement with the fact that in GHz range, the dielectric properties of skin layers vary slightly with frequency. Hence, for each tissue layer, temperature predicted at different frequencies is almost same.

CONCLUSIONS

The explosive growth in mobile communication systems has called for 5G era. The use of mmW frequencies in 5G communication systems has generated concerns about their biological safety. Thermal analysis of exposed tissues is particularly important to evaluate the health risks associated with mmW exposure. Due to “very small” wavelength, the penetration of mmWs is very small. Thus, the potential thermal hazards due to mmW exposure can occur in skin. The present study evaluates the electric field, local SAR and temperature variations within a multilayered skin tissue exposed to 5G frequencies in mmW range. We have estimated the electric field and local SAR at mmW frequencies for exposure power density limit (10 W/m$^2$) set by ICNIRP. The electric field and SAR distribution exhibit similar pattern. Maximum electric field is estimated in skin when the exposure frequency is 38 GHz; while lowest electric field is predicted at 28 GHz. Also, SAR which measures the EM energy absorbed by skin tissue is maximum at 38 GHz and is minimum at 60 GHz.

In addition to electric field and SAR analysis, it is necessary to determine the temperature alterations occurred in the exposed tissue for complete investigation of thermal effects of 5G frequencies. We have used Pennes’ bioheat equation to calculate the temperature variations in exposed multilayer skin tissue. It is found that the SAR and temperature distribution do not follow a similar pattern. Highest value of SAR is predicted in epidermis layer, whereas highest temperature peak is estimated in the subcutaneous fat layer. For all the exposure frequencies, highest transient temperature is predicted at 2s of exposure. Highest transient temperature in epidermis layer, dermis layer and subcutaneous fat is 37.20°C, 37.16°C and 37.36°C respectively. Highest temperature in subcutaneous layer is due to lower thermal conductivity of this layer. The temperature falls very quickly due to onset of blood perfusion and achieves its steady state (approximately 37°C), which is core body temperature. In addition to blood perfusion and thermal conductivity, the dielectric properties of skin layers play an important role in estimating the temperature variations. In GHz region, the dielectric properties do not vary significantly with exposure frequency. As the result, temperature variations are almost similar at different frequencies of exposure.

This study reveals that the maximum temperature change in skin exposed to 5G frequencies is 0.36°C, and the temperature spike is very short lived. The temperature rise is too small to cause any thermal damage to the tissue. The results indicate that the mmWs cannot cause thermal injuries in skin and may be considered safe for 5G mobile communications.

NOMENCLATURE

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>Specific heat capacity, J/(kg .K)</td>
<td></td>
</tr>
<tr>
<td>E</td>
<td>Electric field intensity, V/m</td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>Electric current, A</td>
<td></td>
</tr>
<tr>
<td>K</td>
<td>Thermal conductivity, W/(m.K)</td>
<td></td>
</tr>
<tr>
<td>k</td>
<td>Wave number, 1/m</td>
<td></td>
</tr>
<tr>
<td>l</td>
<td>Edge length, m</td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>Normal vector</td>
<td></td>
</tr>
<tr>
<td>Q</td>
<td>Heat generation, W/m$^3$</td>
<td></td>
</tr>
<tr>
<td>SAR</td>
<td>Specific Absorption Rate, W/kg</td>
<td></td>
</tr>
<tr>
<td>T</td>
<td>Temperature, °C</td>
<td></td>
</tr>
</tbody>
</table>
Voltage, V
x Thickness of layer, m
Z Impedance, Ω

Greek symbols

ε Permittivity, F/m
μ Magnetic permeability, H/m
ρ Mass density, kg/m³
σ Conductivity of tissue, S/m
ωb Blood perfusion rate, 1/s
ω Angular frequency, Hz

Subscripts

0 Free space, initial
b Blood
ext External
in Input
m Metabolic
p Constant pressure
r Relative

Conflict of Interest: The authors declare that they have no conflict of interest.

REFERENCES


