AN APPLICATION OF PULSED INFRARED THERMAL IMAGING IN THE DIAGNOSIS OF DIABETIC MICROANGIOPATHY

by

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Diabetic microangiopathy is an important cause of morbidity and mortality of diabetes foot ulcers. Its early detection is very important for early intervention to avoiding severe clinical symptoms. In this article, theoretical study on a pulsed infrared thermal imaging technology detecting early diabetic microangiopathy in lower extremity was carried out. The working principle of pulsed infrared thermal imaging technology was described and the 3-D thermal conduction model for atherosclerotic plaque in microvessel of distal lower extremity using pulsed infrared thermal imaging technology was established and calculated. The effect of atherosclerotic plaque geometry size including length and thickness to the measurement parameter was studied, and the influence law has been got, which can provide a theoretical basis for the diagnosis of diabetic microangiopathy using pulsed infrared thermal imaging technology.

Key words: heat conduction, thermal imaging, geometry size, diabetic cutaneous microangiopathy

Introduction

Diabetic microangiopathy is a major vascular complication in diabetic patients. Diabetic retinopathy, diabetic nephropathy, diabetic peripheral neuropathy and diabetic foot ulcers are the main clinical manifestations in diabetic microangiopathy. Fundus fluorescein angiography was used to examine diabetic retinopathy, urine microprotein detection can help diagnose diabetic nephropathy, diagnosis of diabetic peripheral neuropathy is mainly based on clinical symptoms and nerve conduction measurements [1]. While diabetic foot ulcers can only be diagnosed when the clinical symptoms are obvious, which often faces the risk of amputation at the same time [2]. Studies have shown that diabetic foot ulcers was the first cause of non-traumatic amputation, with the amputation rate as high as 26.4% [3]. Now it was believed that diabetic foot ulcers was the result of many factors, including neuropathy, vascular disease, infection and so on in the distal lower extremity [4-5].
them, early distal microangiopathy of lower extremity may be a major factor affecting distal peripheral neuropathy and peripheral macroangiopathy. The main characteristic of early distal microangiopathy of lower extremity is atherosclerotic changes of microangiopathy caused by basement membrane thickening and deposition of transparent substances [6]. Unfortunately, there was still a lack of method to detect atherosclerotic changes in microangiopathy. Only the presence of atherosclerotic plaques in the large vessels of the lower extremities can be detected by doppler ultrasound or CT angiography [7], however, the treatment was irreversible when macrovascular disease occurs. Therefore, early detection of lower extremity microvascular atherosclerotic lesions is helpful for early intervention, reversal of pathological changes of microvascular disease, improvement of prognosis and quality of life.

Based on the thermal effect theory of biological tissue, the researches of medical application of infrared technology and biomedical heat transfer were carried out, and formed the Infrared Medical Thermal Imaging Technology (IMTIT). IMTIT is a product of the combination of medical technology, infrared technology and computer multimedia technology, which can record the temperature field of human body. Although current structural imaging detection (MR, CT, X-ray and ultrasound) can provide pathological information of abnormal body structure, it can not reflect the functional changes of body tissues. When diseases occurred, functional changes are earlier than structural changes. The dynamic observation of body function is more precursory than static observation. The technology using long-pulsed infrared thermal wave for testing breast and skin diseases was mature, and increasing studies of this technology theoretical in these disease emerged in [8-9]. However, whether the long-pulsed infrared thermal wave testing technology can be used to detect diabetic microangiopathy was seldom studied. Only fewer clinical case-control studies occurred in this field [10]. Moreover, there was lack of relevant theoretical research of long-pulsed infrared thermal wave technology for detecting diabetic microangiopathy.

In this paper, the application of pulsed infrared thermal imaging in the detection of atherosclerotic plaque lesions located in capillary in papillary dermis of lower extremity skin will be carried out.

The principle of pulsed infrared thermal imaging technology

Infrared thermal imaging technology is a new nondestructive testing method developed in recent years, which has the advantages of non-contact, high efficiency, real-time, etc. The technology has developed rapidly and has been applied more and more widely in the field of biomedicine. Active infrared thermal imaging technology can often achieve better detection results and various external thermal excitation modes can be used. The diagnostic principle of pulse infrared thermal imaging technology is: active heating of tissue by pulse excitation source, and heat flow propagates within the tissue. Because the thermophysical parameters of lesions and non-lesions are different, the corresponding surface temperatures are different. Therefore, according to the difference of temperature distribution on the tissue surface, we can judge whether the lesion is or not and the degree of the lesion.

3-D thermal conduction model
Heat transfer in biological tissues is a complicated process that involves heat conduction, blood perfusion and metabolic heat generation. In this section, a 3-D thermal conduction model for pulsed infrared thermal imaging in the diagnosis of diabetic microangiopathy is built. The structure size of the geometric model is shown in fig. 1. and tab. 1.

![Figure 1. Structure of the geometric model](image)

**Table 1. Structure size of the geometric model**

<table>
<thead>
<tr>
<th>Tissue layers</th>
<th>Epidermis</th>
<th>Papillary dermis</th>
<th>Plaque</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epidermis</td>
<td>$L_E$ [mm]</td>
<td>0.05, 0.10</td>
<td></td>
</tr>
<tr>
<td>Papillary dermis</td>
<td>$L_{Pd}$ [mm]</td>
<td>0.7</td>
<td></td>
</tr>
<tr>
<td>Blood vessels</td>
<td>$d_{BvPd}$ [mm]</td>
<td>0.26</td>
<td></td>
</tr>
<tr>
<td>ddBvPd [mm]</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plaque</td>
<td>$TH_{pla}$ [mm]</td>
<td>0.04, 0.06, 0.08, 0.10</td>
<td></td>
</tr>
<tr>
<td>$L_{pla}$ [mm]</td>
<td>2, 4, 6, 8</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The process of the heat transfer in biological tissues can be represented as follows [11]:

$$\rho \cdot c \frac{\partial T}{\partial t} = k \cdot \nabla^2 T + \omega_b \cdot c_b (T_a - T) + Q_m + Q(t)$$

(1)

where $\rho$ is the tissue density, $c$ is the specific heat of the tissue, $T$ is the local temperature of the tissue, $k$ is the tissue thermal conductivity, $\nabla^2$ is the second-order three-dimensional differential operator, $\omega_b$ is the blood perfusion rate, $c_b$ is specific heat of the blood, $Q_m$ is the metabolic heat generation rate, and $Q(t)$ is the external excitation heat source of the human skin surface, which can be written as follows:

$$Q(t) = \begin{cases} 5W & 0 < t \leq 2s \\ 0 & t > 2s \end{cases}$$

(2)

For the skin surface, the boundary condition is as follows:

$$-k \frac{\partial T(x, y, z, t)}{\partial x} |_{\text{skin surface}} = Q(t) + h_t \left[ T_a - T_{(\text{skin surface}, y, z, t)} \right] + \alpha \left[ T_a^4 - T^4(H, y, z, t) \right]$$

(3)

The initial temperature is normal human body temperature, and the ambient temperature is 293.15 K. Convective heat transfer coefficient between human skin surface and surrounding air is set as 3 W/m²·K. The bottom surface temperature of papillary dermis is
310.15 K. Other edge is assumed as insulated. The 3-D thermal conduction model is calculated by the COMSOL software.

**Results and discussions**

The influence of epidermis thickness

Set papillary dermis thickness 0.7 mm, plaque thickness 0.08 mm, plaque length 4 mm, and epidermis thickness 0.05 mm and 0.1 mm, respectively. Fig. 2 and Fig. 3 shows the temperature distribution on tissue surface at t=1.5s. As can be seen from the figures, the influence of epidermis thickness on the detection effect is relatively small. According to the calculation, the difference between the surface temperature of plaque corresponding tissue and that of normal tissue is about 2 K. And the temperature difference of epidermis thickness 0.05 mm is slightly higher than that of 1.0 mm.

![Figure 2](image1.png)

(a) Temperature distribution       (b) Temperature distribution across the Line 1

**Figure 2.** The temperature distribution on tissue surface at t=1.5s ($L_E=0.05mm$)

![Figure 3](image2.png)

(a) Temperature distribution       (b) Temperature distribution across the Line 2

**Figure 3.** The temperature distribution on tissue surface at t=1.5s ($L_E=0.10mm$)

The influence of plaque thickness

Set epidermis thickness 0.1 mm, papillary dermis thickness 0.7 mm, plaque length 4 mm, and plaque thickness 0.04, 0.06, 0.08, 0.10 mm, respectively. Fig. 4 and Fig. 5 shows the temperature distribution and difference between the surface temperature of plaque corresponding tissue and that of normal tissue according to different plaque thickness at t=1.5s. From Fig. 4 and Fig. 5, it can be seen that the temperature difference increases first and then decreases with the increase of plaque thickness. As the thermal conductivity of plaque is smaller than that of other normal tissues, the surface temperature difference of corresponding tissues increases with the increase of plaque thickness; on the other hand, when
the plaque increases to a certain extent, the blood flow is greatly hindered and the shear
viscous heat generation decreases, so the surface temperature difference begins to decrease.

![Temperature distribution images](image)

Figure 4. The temperature distribution on tissue surface versus plaque thickness at t=1.5s
\(L_{pla}=4\text{mm}\)

![Temperature difference graph](image)

Figure 5. The temperature difference on tissue surface versus plaque thickness at t=1.5s
\(L_{pla}=4\text{mm}\)

The influence of plaque length

Set epidermis thickness 0.1 mm, papillary dermis thickness 0.7 mm, plaque thickness 0.08 mm, and plaque length 2, 4, 6, 8 mm, respectively. Fig. 6 shows the temperature distribution on tissue surface at t=1.5s. Tab. 2 shows the temperature difference between the surface temperature of plaque corresponding tissue and that of normal tissue according to different plaque length at t=1.5s. From fig. 6 and tab. 2, it can be seen that the
temperature difference decreases firstly, then increases, and then decreases with the increase of plaque length. It indicates that the effect of plaque length on the surface temperature difference of tissues is complex.

Figure 6. The temperature distribution on tissue surface versus plaque thickness at t=1.5s

\(TH_{pla}=0.1\text{mm}\)

Table 2. The temperature difference on tissue surface versus plaque length at t=1.5s

<table>
<thead>
<tr>
<th>(L_{pla}) (mm)</th>
<th>2</th>
<th>4</th>
<th>6</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\Delta T) (K)</td>
<td>1.56</td>
<td>1.4873</td>
<td>1.5983</td>
<td>1.4179</td>
</tr>
</tbody>
</table>

Conclusions

In the present study, the atherosclerotic plaques in microvessels of papillary dermis were used to simulate diabetic microangiopathy, and the plaque was modeled in three-dimension. The pulsed infrared thermal imaging technology was used to detect the plaque models of different geometric sizes in the microvascular. The results showed that the epidermis temperature of the normal regions and abnormal regions in the microvascular is different. This result can be provided as the theoretical support for the clinical application.

Acknowledgments

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Nomenclature

\( \rho \) - density, \([\text{kg/m}^3]\)  
\( k \) - tissue thermal conductivity, \([\text{W/(m·K)}]\)  
\( c \) - specific heat, \([\text{J/(kg·K)}]\)  
\( Q(t) \) - long pulsed heat flux, \([\text{W}]\)  
\( \omega_b \) - blood perfusion rate, \([\text{s}^{-1}]\)  
\( c_b \) - specific heat of the blood, \([\text{J/(kg·K)}]\)  
\( Q_m \) - metabolic heat generation rate, \([\text{W/m}^3]}\)

References


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