Thermographic Investigation of Osseous Stress Pathology

Daniel TJ. Arthur, Masood Mehmood Khan and Luke C. Barclay

Abstract—The debilitating pathology of stress fracture accounts for 10% of all athletic injuries[2], with prevalence as high as 20% in modern military basic training cohorts [3]. Increasing concerns surrounding adverse effects of radiology [5], combined with the 12.5% contribution of diagnostic imaging to Australian Medicare benefits paid in 2009-10 [6], have prompted the search for alternative/adjunct electronic decision support systems[7]. Within conducive physioanatomic milieu, thermal infrared imaging (TIRI) may feasibly be used to remotely detect and topographically map diagnostically useful signs of suprathreshold thermodynamic pathophysiology. This paper details a three month clinical pilot study into TIRI-based detection of osseous stress pathology in the lower legs of Australian Army basic trainees. A dataset of over 500 TIRI’s was amassed. The apparent ‘normal’ thermal profile of the anterior aspect of the asymptomatic lower leg is topographically defined and validated against current thermophysiological theory [8] via cadaveric dissection.

I. INTRODUCTION

THERMAL infrared imaging (TIRI) employs a focal plane array (FPA) to remotely detect and topographically map thermal emittance. Suprathreshold thermodynamic pathophysiology occurring in conducive pathoanatomic milieu may feasibly manifest TIR ‘signs’. Osseous stress pathophysiology features focal thermodynamic mechanisms including hyperemia, and both osteocytic and hyperalgesic nitric oxide production [9]. In physioanatomic context of the anteromedial aspects of the tibiae, as pertinent to epidemiological prevalence [3], these mechanisms occur within a thin superficial volume of thermodynamically determinate tissues [10], unoccluded by thermogenic anatomy or dissipative heat-sinking vasculature [11]. The aim of this study was to elucidate the potential of TIRI in the diagnosis and/or management of osseous stress pathology in the lower limbs, via; topographic definition of the ‘normal’ asymptomatic thermal profile of the anterior tibial ROI; compilation of a multi-modality symptomatic dataset for validation of TIR signs and development of a pathoanatomically representative bioheat transfer model; and compilation of longitudinal intrasubject TIR datasets to elucidate the potential for TIR detection of subclinical pathology and general pathological staging. In light of the afore-alluded-to feasibility study, and development of objective clinical protocols via functional isolation of the inherent systems’ architecture, the authors were granted approval for protocols 592-10 and HR62/2010 by the Australian Defence Force and Curtin University human research ethics committees, respectively.

II. MATERIALS AND METHODS

Upon arrival at the Australian Army Recruit Training Centre, three platoons of basic trainees consisting of 56, 48, and 43 recruits were given a comprehensive plain-language study briefing, and allowed 24 hours to anonymously deposit consent forms featuring medical questions pertinent to
contraindication of TIRI into a sensitively located ballot box. Consenting participants were first thermally imaged during their initial inoculation parade, occurring prior to their first physical training session. Thermal infrared images were acquired prior to vaccine administration to avoid vasoactive artifacts. In absence of data pertinent to optimal conditions for pre-TIRI equilibration, participants remained seated, clad in loose fitting gym shorts and T-shirts, without shoes or socks, for a nominal 20-30 minute period, in a carpeted room with ambient conditions as listed in Table 1.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Optimal Thermoneutral Value*</th>
<th>Experimental Value</th>
<th>Meter / Reference for Experimental Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ambient Air Temperature</td>
<td>30°C</td>
<td>28°C ± 1°C</td>
<td>Vaisala HUMICAP® HM34</td>
</tr>
<tr>
<td>Relative Humidity</td>
<td>40%</td>
<td>46% ± 3%</td>
<td>Vaisala HUMICAP® HM34</td>
</tr>
<tr>
<td>Emissivity of surrounding surfaces</td>
<td>0.93</td>
<td>0.92-0.96</td>
<td>[1]</td>
</tr>
<tr>
<td>Temperature of surrounding surfaces</td>
<td>30°C</td>
<td>Not Controlled</td>
<td>N/A</td>
</tr>
<tr>
<td>Ambient Air Velocity</td>
<td>0.05 m/s</td>
<td>Minimized</td>
<td>N/A</td>
</tr>
<tr>
<td>Basal Metabolic Rate</td>
<td>0.8</td>
<td>Minimized</td>
<td>N/A</td>
</tr>
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</table>

*Optimal thermoneutral values as calculated via the IESD-Fiala human thermoregulation model used in conjunction with ANSYS CFX indoor environment program [4].

During equilibration each participant’s legs were visually photographed to screen potentially contraindicative superficial lesions. Following equilibration, patients lay stationary supine upon an unarticulated/flat LINAK OpenBus™ hospital bed, for no less than 3 minutes prior to image acquisition. Several 2cm thick dark grey matt-finish gym mats were kept in redundancy, equilibrating on rotation to 20°C in an anteroom and used as a backing surface between the bed and each participant to prevent residual thermal artifacts. An ad-hoc camera mounting system was constructed to steadily maintain the lens to skin-surface distance (‘D’ Fig. 1), with the optical axis normal to the patient bed. In absence of data concerning optimal ambient conditions for image capture, five wall-mounted Daikin FVXS71H ducted inverter units were used to maintain conditions as close to thermoneutral (Table 1) as practicable. Pending release of the IEC TC/SC/WG5 specifications for medical grade thermal infrared cameras [12], a FLIR ThermoVision™ A40V thermal infrared camera featuring a 7.5 to 13.0 µm uncooled bolometric FPA and thermal sensitivity of 0.08°C was used in the configuration shown in Fig.1. The longest dimension of the region of interest ($L_{roi}$) was anticipated to be ≤ 515mm in accordance with 95th percentile ‘popliteal height’ as defined in ISO/TR 7250-2:2010 [13]. Given the A40V’s 320x240 FPA with $\Theta_{FOV}$ of 24°x18° [14], the lens to skin-surface distance (‘D’ Fig. 1) was set to 1.30 m, providing a 553 mm x 412 mm coplanar field of view, and spatial resolution of 0.58mm² at the skin’s surface. In absence of anatomy-specific data, ThermaCAM™ Researcher’s generic ‘skin’ emissivity value of 0.98 was adopted [15]. A single static TIRI was acquired from each participant. Participants were thermally imaged via the same protocol on three further occasions during their training continuum; in conjunction with scheduled medical check-up parades in weeks four, seven, and nine. Any participants presenting consistently with osseous stress injury during the course of the study were withdrawn from physical training and subject to either MRI or nuclear bone scan in accordance with Australian Army SOPs. Thermal infrared images were acquired from symptomatic subjects within the two hour period immediately prior to MRI or bone scan procedures. Subjects diagnosed with stress fracture were sent home for 6 weeks of convalescent leave.

### III. RESULTS

Among the 474 asymptomatic thermal images obtained, were 116 complete 4-image longitudinal intrasubject TIRI sets taken over the full ten weeks of basic training. Early analysis

![Fig. 2. TIR profile of apathological anterior lower leg; i) greybody temperature map for global emissivity of 0.98; ii) gray-scale negative with coloured crosshairs at geometric centers of emissively distinct regions corresponding to Thermophysiology of underlying anatomy, as explained below.](image-url)

of these asymptomatic images reveals a ‘normal’ thermal topography for the asymptomatic anterior lower leg (Fig. 2i); the
characteristic features of which have been validated via consideration of the thermodynamics of the underlying anatomy (Fig. 3 & 4). As exemplified by Fig. 2, the healthy TIR profile features 5 topographically distinct regions. The blue crosshair in Fig. 2ii demarcates the point within the funnel-shaped proximal region from which emittance is typically at a minimum.

Table II

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>Mode</th>
<th>Standard Deviation</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Inter-subject</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Intra-subject</td>
</tr>
<tr>
<td>Maximum Emittance / Temperature*</td>
<td>31.0 °C</td>
<td>31.2 °C</td>
<td>0.6 °C</td>
</tr>
<tr>
<td>Minimum Emittance / Temperature*</td>
<td>28.0 °C</td>
<td>28.2 °C</td>
<td>0.4 °C</td>
</tr>
</tbody>
</table>

*Typical loci of maxima and minima indicated by the red and blue crosshairs in Fig.2ii respectively

As shown in Fig. 3i, this region overlies no thermogenic anatomy, and as shown in Fig. 5, the avascular interstitial osteocutaneous tissue volume is approximately 3.5 mm thick. The second row of Table 2 characterizes the average quantitative thermal characteristics of this region, as calculated from a dataset of 474 TIRI’s acquired over a period of 3 months. The red crosshair in Fig 2ii demarcates the geometric centre of the point within the almond-shaped lateral region, from which emittance is typically at a maximum. As shown in blue in Fig.3, this lateral region overlies the compartment of the thermogenic tibialis anterior muscle. The white crosshair in Fig. 2ii, and the red regions in Fig. 3 correspond to the thermogenic superficial posterior muscles, with Fig. 3i.

As shown in Fig. 3i, this region overlies no thermogenic anatomy, and as shown in Fig. 5, the avascular interstitial osteocutaneous tissue volume is approximately 3.5 mm thick. The second row of Table 2 characterizes the average quantitative thermal characteristics of this region, as calculated from a dataset of 474 TIRI’s acquired over a period of 3 months. The red crosshair in Fig 2ii demarcates the geometric centre of the point within the almond-shaped lateral region, from which emittance is typically at a maximum. As shown in blue in Fig.3, this lateral region overlies the compartment of the thermogenic tibialis anterior muscle. The white crosshair in Fig. 2ii, and the red regions in Fig. 3 correspond to the thermogenic superficial posterior muscles, with Fig. 3i.
validation of perceived pathological thermal aberrations, and to guide development of a pathoanatomic bioheat transfer environment, facilitating intermodal registration and for MRI’s, and bone scan images was amassed within a controlled environment. The authors’ knowledge, this is the first classification of lower leg emissive topography in the literature. Coarse thermographic tissue discrimination has been demonstrated, highlighting the potential of TIRI to subjectively reveal morphology.

IV. CONCLUSION

A dataset of over 500 normal and symptomatic TIRI’s, MRI’s, and bone scan images was amassed within a controlled environment, facilitating intermodal registration and for validation of perceived pathological thermal aberrations, and to guide development of a pathoanatomic bioheat transfer model. The normal thermal topography of the asymptomatic anterior aspect of the lower leg has been defined, and validated against underlying anatomy. To the authors’ knowledge, this is the first classification of lower leg emissive topography in the literature. Coarse thermographic tissue discrimination has been demonstrated, highlighting the potential of TIRI to subjectively reveal morphology.

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REFERENCES