



LJMU Research Online

Low, DA, Green, DJ, Cable, NT, Carter, HH, Atkinson, C, Birk, P, Dawson, EA and Thijssen, DHJ

Localised cutaneous microvascular adaptation to exercise training in humans

<http://researchonline.ljmu.ac.uk/id/eprint/7554/>

Article

Citation (please note it is advisable to refer to the publisher's version if you intend to cite from this work)

Low, DA, Green, DJ, Cable, NT, Carter, HH, Atkinson, C, Birk, P, Dawson, EA and Thijssen, DHJ (2018) Localised cutaneous microvascular adaptation to exercise training in humans. European Journal of Applied Physiology. ISSN 1439-6319

LJMU has developed **LJMU Research Online** for users to access the research output of the University more effectively. Copyright © and Moral Rights for the papers on this site are retained by the individual authors and/or other copyright owners. Users may download and/or print one copy of any article(s) in LJMU Research Online to facilitate their private study or for non-commercial research. You may not engage in further distribution of the material or use it for any profit-making activities or any commercial gain.

The version presented here may differ from the published version or from the version of the record. Please see the repository URL above for details on accessing the published version and note that access may require a subscription.

For more information please contact researchonline@ljmu.ac.uk

Localised cutaneous microvascular adaptation to exercise training in humans

Ceri L. Atkinson¹
Howard H. Carter²
Dick H.J. Thijssen^{2,3}
Gurpreet K. Birk²
N. Timothy Cable^{2,4}
David A. Low²
Floortje Kerstens³
Iris Meeuwis³
Ellen A. Dawson²
Daniel J. Green^{1,2}

¹ School of Human Sciences (Exercise and Sport Science),
The University of Western Australia, Crawley, Western Australia, 6009

² Research Institute for Sport and Exercise Science, Liverpool John Moores University,
Tom Reilly Building, Byrom Street, Liverpool L3 3AF, United Kingdom

³ Radboud Institute for Health Sciences, Department of Physiology, Radboud University Medical
Centre, Nijmegen, the Netherlands

⁴ Department of Sport Science, Aspire Academy, Qatar

Running Title: Exercise training and human skin circulation

Funding: Australian Research Council (DP 130103793), E. Dekker stipend (Netherlands Heart
Foundation, 2009 T064)

Author for Correspondence:

Daniel Green

School of Human Sciences (Exercise and Sport Science),
The University Of Western Australia,
Crawley, Western Australia, 6009
Email: danny.green@uwa.edu.au

37 Abstract

38 **Purpose.** Exercise training induces adaptation in conduit and resistance arteries in humans,
39 partly as a consequence of repeated elevation in blood flow and shear stress. The stimuli
40 associated with intrinsic cutaneous microvascular adaptation to exercise training have been less
41 comprehensively studied.

42 **Methods.** We studied 14 subjects who completed 8-weeks cycle ergometer training, with partial
43 cuff inflation on one forearm to unilaterally attenuate cutaneous blood flow responses during
44 each exercise-training bout. Before and after training, bilateral forearm skin microvascular
45 dilation was determined using cutaneous vascular conductance (CVC: skin flux/blood pressure)
46 responses to gradual localised heater disk stimulation performed at rest (33, 40, 42 and 44°C).

47 **Results.** Cycle exercise induced significant increases in forearm cutaneous flux and
48 temperature, which were attenuated in the cuffed arm (2-way ANOVA interaction-effect;
49 $P<0.01$). We found that forearm CVC at 42°C and 44°C was significantly lower in the uncuffed
50 arm following 8-weeks of cycle training ($P<0.01$), whereas no changes were apparent in the
51 contralateral cuffed arm ($P=0.77$, interaction-effect $P=0.01$).

52 **Conclusions.** Lower limb exercise training in healthy young men leads to lower CVC-responses
53 to a local heating stimulus, an adaptation mediated, at least partly, by a mechanism related to
54 episodic increases in skin blood flow and/or skin temperature.

55

56 **Keywords:** skin microcirculation; blood flow; exercise training

57

58 Abbreviations

- 59 BMI – body mass index
60 CVC – cutaneous vascular conductance
61 HRMax- Heart rate max
62 PU – Perfusion units
63 SNP – sodium nitroprusside
64

Introduction

Exercise training has strong and independent cardioprotective effects (Blair and Morris 2009) that may be partly mediated through the direct impact of exercise on the vasculature (Green et al. 2008; Joyner and Green 2009; Green et al. 2017). In conduit and resistance arteries, dynamic exercise training using large muscle groups leads to generalised effects on the vasculature (Thijssen et al. 2012; Green et al. 2011; Silber et al. 1991; Snell et al. 1987). For example, cycle ergometer exercise training, which specifically avoided hand-gripping or use of the upper limb musculature, induced brachial artery adaptation (Birk et al. 2012; Tinken et al. 2008). Interestingly, when exercise-mediated increases in brachial artery blood flow were attenuated during exercise bouts (via unilateral partial cuff inflation), these adaptations were no longer apparent, implying a role for haemodynamic forces, such as shear stress, as important stimuli to conduit artery adaptation in response to exercise training in humans (Birk et al. 2012; Tinken et al. 2010; Green et al. 2017).

Whilst conduit and larger resistance arteries adapt to exercise training by enhancing function and then remodelling to increase their diameter (Green et al. 2017; Tinken et al. 2008; Laughlin 1995), studies of microvascular adaptation in skeletal muscle have proposed that increases in capillarity in response to training may counteract reductions in red cell transit time, enabling increased oxygen extraction (Krustrup et al. 2004). In this context, it is well established that the mode of adaptation to training differs according to rank order along the arterial tree; larger arteries primarily respond to shear stress and other hemodynamic stimuli via cross sectional enlargement (Rowley et al. 2011; Newcomer et al. 2011), whereas microvessels respond to these

stimuli (and hypoxia) by budding and sprouting (Cocks et al. 2013; Andersen and Henriksson 1977; Brown 2003).

The role of repeated exercise and shear stress in the adaptation of cutaneous microvessels is less well understood, despite the obvious relevance of changes in the structure and function of these vessels to microvascular disease (Simmons et al. 2011; Holowatz et al. 2008). Some evidence suggests that exercise training induces intrinsic adaptation in the cutaneous microvasculature (Black et al. 2008; Simmons et al. 2011), whilst we recently completed a series of studies, involving repeated localised and systemic heating, which suggested that cutaneous microvascular adaptation may depend upon local skin temperature changes in addition to changes in blood flow and shear stress that occur as a consequence of elevation in core temperature (Green et al. 2010; Carter et al. 2014). The contribution of these important physiological stimuli to cutaneous adaptation in response to leg exercise training is currently unknown.

The primary aim of this study was to examine the role played by repeated increases in blood flow in localised adaptations in the skin microcirculation of the upper limbs, following 8-weeks of cycling exercise. To address this aim, we partially inflated a pneumatic cuff around one forearm during each exercise bout, to unilaterally attenuate exercise-induced hyperaemia (Tinken et al. 2009; Tinken et al. 2010). Before and after the intervention, skin blood flow responses were assessed at rest using a local heating protocol. We chose to use a localised disk heating protocol as a tool to assess intrinsic microvascular adaptation in the skin because it is well established that this approach provides an intrinsic microvascular index that is independent of reflex neurovascular control (Minson et al. 2001; Black et al. 2008). We hypothesised that cycle

exercise training, by inducing repeated episodic increases in cutaneous blood flow would increase forearm skin vasodilator capacity in response to a localised heating stimulus.

Materials and Methods

Ethical Approval

All study procedures complied with the *Declaration of Helsinki* and were approved by the Human Research Ethics Committee at Liverpool John Moores University and The University of Western Australia. All subjects provided written, informed consent before participating in the study.

Subjects

Fourteen healthy men (Age: 25 ± 2 years, Height: 180 ± 5 cm, Weight: 76 ± 13 kg and BMI: 23 ± 3 kg.m^{-2}) were recruited to participate in an exercise training experiment. A further 6 male subjects (27 ± 6 years) undertook an acute cycle exercise experiment to characterise forearm skin blood flow changes. No subject had cardiovascular disease, diabetes, insulin resistance or possessed cardiovascular risk factors such as hypercholesterolaemia or hypertension. Subjects who smoked or were on medication of any type were excluded. Participants at pre-study entry were either classified as sedentary or recreationally active (<5 hours per week). All sedentary participants only took part in the prescribed 3 x 30 min sessions per week across the 8 weeks and no new activity was introduced outside of this. All recreationally active participants were asked to keep their activity levels consistent throughout the 8 week intervention so that the additional 3 x 30 mins a week were in addition to pre-entry exercise levels.

Acute experiment (n=6): Effect of leg cycle exercise on skin microvascular responses ± forearm cuff inflation

Participants undertook 30 mins of cycle exercise (Monark 874E, Sweden) at 80% HRmax (age-predicted). Throughout the exercise bout a pneumatic cuff (6cm in width: SC5D Hokanson cuff, Bellevue USA) was placed around one forearm immediately below the cubital crease and inflated to 60mmHg. Previous studies have demonstrated that placement and inflation of a forearm cuff in this manner attenuates brachial artery blood flow (Birk et al. 2012; Thijssen et al. 2009; Tinken et al. 2010; Tinken et al. 2009). The contralateral arm remained uncuffed during the exercise bout. Subjects rested for 10-mins on the ergometer in a quiet, temperature controlled room. Baseline bilateral forearm skin blood flows using laser-Doppler flowmetry (Model 413, Periflux 5001 System, Perimed AB) were then recorded for 2.5 mins, followed by unilateral cuff inflation to 60 mmHg for 2.5 mins. Skin blood flow data were recorded in both arms on the ventral side of the forearm during the exercise bout and averaged in 2.5-min bins along with skin temperature using thermistors (Squirrel 1000 series, Grant Instruments, Cambridge, United Kingdom). Placement of skin thermistors and laser Doppler probes was the same on both arms and always distal to the cuff site. In order to avoid any instances of handgripping, participants were instructed to rest both arms on the handle bars of either side of the bike.

Chronic experiment (n=14): Effect of leg cycle exercise on skin microvascular responses ± forearm cuff inflation

Exercise training was performed over an 8-week period with subjects visiting the laboratory 3 times per week. Each of these 24 sessions was supervised and consisted of 30 mins of cycle

exercise (80% age-predicted HRmax, cadence 60 rpm), performed at the same time of day. As above, a pneumatic blood pressure cuff was placed below the cubital crease on one forearm of each subject and inflated to 60 mmHg throughout each 30 min exercise training bout. The arm selected for cuff placement was randomised, but once selected, it remained consistent across the 8-week training period for any given participant.

Gradual local heating protocol

Assessments were performed before and after the 8-week cycling exercise training program at rest. All studies were conducted in a quiet, temperature controlled environment and each visit for a given subject was performed at the same time of day. Subjects were asked to fast for 8 hours, abstain from alcohol and caffeine for 16 hours, and not to perform any exercise for 24 hours. Probe placement sites were selected on each forearm and the location of these sites was recorded using photographs and distances from anatomical landmarks, so that repeated measures were collected at the same skin site. In a previous study, we examined the day-day reproducibility of forearm skin responses to our local heating protocol. Coefficients of variation of 17.1 and 14.4% were obtained when skin responses were presented as absolute cutaneous vascular conductance (CVC; at 42 and 44°C, respectively) (Dawson et al. 2015).

Upon arrival, subjects were seated and instrumented for ~20 mins before beginning the ~90 min heating protocol. Local heater disks (Perimed 455, Stockholm, Sweden) were placed on the forearms using double-sided adhesive rings. The laser Doppler probes, each with a 7-laser array (Model 413, Periflux 5001 System, Perimed AB), were then fitted into the middle of these localised heating disks to record change in red blood cell flux (in perfusion units, PU)

(Cracowski et al. 2006). Once instrumented, recording commenced and the heater disk temperatures were increased to 33°C and maintained at this temperature for a 10 min baseline period. Upon completion of this baseline period, increments in heater disk temperature were gradual, so as to minimise the influence of the axon reflex on the skin blood flow response (Minson et al. 2001). Heat increased in controlled increments of 0.5°C every 2.5 mins until 42°C was reached. Once the probes reached 42°C, they remained at this temperature for a period of 30 mins. Finally, heater disk temperatures were increased to 44°C for another 20 mins to induce a maximal skin blood flow response. Previous studies have suggested that heating to the levels used in the present study results in similar CVC increases to those observed in response to infusion of SNP (Kellogg et al. 1999; Cracowski et al. 2011), indicating that maximal responses are indeed obtained and that these reflect the structural capacity of the microvasculature. Resting blood pressure was recorded every 15 min using an automated sphygmomanometer placed around the ankle (Dinamap; GE Pro 300V2, Tampa, FL). Blood pressure measures taken at the ankle were later corrected for the hydrostatic column by multiplying the vertical height difference between the right atrium and the ankle (cm) by 0.766 to convert to mmHg (Groothuis et al. 2008), this value was then subtracted from the raw blood pressure value acquired at the ankle. Converted blood pressure values were then used to calculate CVC, which accounts for skin blood flow changes occurring as a result of changes in blood pressure (Cracowski et al. 2006). All laser Doppler measurements were relayed and graphed in real time via a Powerlab onto a laptop running LabChart 7 (AD Instruments, Sydney, Australia).

Data analysis and statistics

Skin blood flow (PU) from the cuffed and uncuffed arms was averaged across 1-minute intervals. Calibration of the laser Doppler probes was undertaken before and after the experiments using two generic points, 0 and 250 PU, in accordance with calibration guidelines using a zeroing disk and motility standard (Periflux System, Perimed AB). We measured and analysed the 2.5 min increment at 40°C, as well 5-min averages of data from the plateau-phases at 42°C and 44°C. Measurements in PU were converted to CVC which was calculated as PU/Dinamap mean arterial pressure (MAP).

Statistical analyses were performed using SPSS 21.0 (SPSS, Chicago, Illinois) software. All data are reported as mean (SD) unless stated otherwise, while statistical significance was assumed at $P \leq 0.05$. Initially, a linear mixed model was conducted (with time and cuff inflation as factors) for both skin temperature and skin blood flow during the acute study. A second linear mixed model was used with local heating (33, 40, 42 and 44°C) and time (Week 0 & 8) as factors in order to assess change in CVC responses to the 8-week intervention in the uncuffed and cuffed arm. Finally, a linear mixed model was performed on delta change CVC scores (Week 8 minus Week 0) with cuff and heating as factors. Post-hoc analysis was performed using the least significant difference (LSD) method for pair-wise multiple comparisons when a significant main effect was observed (Perneger 1998).

Results

Acute experiment (n=6): Effect of leg cycle exercise on skin microvascular responses ± forearm cuff inflation

Results of a linear mixed model revealed that heart rate increased significantly during the leg cycling exercise bout (Figure 1A). The increase in HR observed prior to exercise from the -2.5 to 0 min time points may be attributable to an anticipatory response (Williamson et al. 2006). Skin temperature in the uncuffed forearm increased modestly, but significantly, during exercise (by 1.8°C, Figure 1B). However, no change in skin temperature was observed during leg cycling exercise in the cuffed arm (Figure 1B). No significant differences were evident in skin blood flow between the cuffed and uncuffed arms in the baseline period before the cuff was inflated (13.1 ± 3.5 vs. 15.5 ± 9.1 PU, respectively, $P=0.41$). After the onset of leg cycling exercise, there was a significant increase in skin blood flow in the uncuffed arm, whilst the increase in flow was significantly attenuated in the cuffed arm (linear mixed model; interaction effect between exercise and cuff inflation of $P<0.01$, Figure 1C).

Chronic experiment (n=14): Effect of leg cycle exercise on skin microvascular responses ± forearm cuff inflation

Across the 8-week exercise training period, there was 97% adherence to the training sessions in the 14 subjects. The efficacy of our training intervention is indicated by a decrease in the average resting heart rate after 8 weeks of cycle exercise training (67 ± 11 bpm vs. 58 ± 9 ; t-test: $P=0.01$). In addition, to maintain exercise HR at 80% HRmax, the workload performed in the exercise sessions increased significantly over the 8 weeks of training, from 129 ± 19 W to

154±21W at week 8 ($P<0.01$). Resting MAP did not change after training (89±8 vs 88±6 mmHg, $P=0.57$).

Gradual local heating protocol

Figure 2 shows a representative response to local heating in one subject throughout the temperature increments while Figure 3 presents the responses to our local heating protocol before and after training in the cuffed and uncuffed arms. Figure 3 presents CVC data in absolute values, rather than normalised to the peak (44°C) maximum. This is because we wished to understand whether the peak responses *per se* were elevated, possibly indicative of changes in microvascular capacity. Values for CVC at baseline (33°C), heating (40°C) and during the plateau at 42°C and 44°C, in the cuffed and uncuffed limbs were similar prior to training (P -values: 0.32, 0.74, 0.51 and 0.76, respectively). A linear mixed model revealed a significantly lower CVC following training at 42°C and 44°C (Figure 3A) in the uncuffed arm (interaction effect $P=0.01$, time effect 42°C $P\leq0.01$, time effect 44°C $P\leq0.01$). However, in the contra-lateral cuffed arm, CVC did not change as a result of training (Figure 3B) (interaction effect $P=0.69$). A linear mixed model for changes in CVC (expressed as CVC Week 8 minus CVC Week 0) between uncuffed vs. cuffed revealed a significant effect for cuff inflation ($P<0.01$).

Discussion

The aims of the present study were to describe the impact of cycle exercise training on maximal forearm skin blood flow responses to local heating and to determine whether adaptations were dependent upon repeated exposure to increases in blood flow. Our experimental manipulation,

involving partial cuff inflation on one arm, was successful in unilaterally manipulating forearm skin blood flow and temperature during leg exercise. Our principal finding, in contrast to our hypothesis, is that cycle exercise training significantly decreased forearm CVC-responses to local heating in the uncuffed forearm. Our findings suggest that this adaptation was mediated by repeated increases in skin blood flow and/or temperature, as no changes were observed in the cuffed forearm which was exposed to significantly lower levels of blood flow/temperature during each cycle exercise training session (Figure 1).

Elicitation of skin blood flow responses via local heating is commonly used for assessment of the cutaneous vasculature (Boignard et al. 2005; Minson et al. 2002; Smith et al. 2011; Colberg et al. 2005). However, few studies have focussed on modification of the localised cutaneous microvasculature following exercise training in humans (Boegli et al. 2003; Black et al. 2008). Previous studies by our group have utilised exercise-independent interventions to determine the role of repeated increases in skin blood flow on cutaneous vascular function, and found that episodic increases in flow are obligatory for enhanced responsiveness to the local heating test above (Green et al. 2010; Carter et al. 2014). It is also well documented that repeated increases in shear stress act as a potent stimulus for improved endothelial vascular adaptation and NO availability in larger conduit and resistance arteries in humans (Hambrecht et al. 2003; Tinken et al. 2010). These findings informed the rationale for the present study, in which exercise training was used as a stimulus, but the changes in forearm blood flow and shear stress associated with cycle exercise were clamped unilaterally.

No previous study has attempted to investigate the role of repeated increases in blood flow on

skin vessels following an exercise intervention. We hypothesised that cycle exercise training would increase skin blood flow responsiveness to local heating, an adaptation mediated by repeated increases in skin blood flow and/or temperature, consistent with the studies above (Green et al. 2010; Carter et al. 2014). On first pass, our finding in the current study of lower peak CVC responses following the intervention suggests that exercise training may have decreased the release of vasodilator stimuli in response to local heating. However, the skin circulation is highly complex, governed by a number of local and systemic mechanisms (Charkoudian 2003), and the possibility that exercise training induces decreased microvascular function seems unlikely. Moreover, the responses to sustained skin heating to 44°C provide an index of peak microvascular vasodilation, commonly used as an index of structural, rather than functional, vascular adaptation in humans (Kellogg et al. 1999; Choi et al. 2014; Johnson et al. 1986; Taylor et al. 1984; Green et al. 1994). Of interest, a recent study by Brunt *et al.* (Brunt et al. 2016) involving repeated whole body heating for 8 weeks induced improvements in local skin vascular function, but non-significant *decreases* in maximal local heating responses (5.5% decrease in CVCmax), consistent with our data. Collectively, these findings argue against the notion that our findings infer detrimental functional adaptations in the skin microvessels and that, rather, they may reflect changes in microvascular structure.

The possibility that exercise training induces structural adaptations in the skin microcirculation is supported by established findings in both resistance and conduit arteries, that exercise training induces arterial remodelling and structural arterial enlargement (Haskell et al. 1993; Naylor et al. 2005; Green et al. 1994; Rowley et al. 2011; Tinken et al. 2008). Such findings are supported by extensive animal literature (Langille and O'Donnell 1986). In the current study, the decrease in

absolute values for CVC that we observed in the uncuffed arm is not likely to be the result of a decrease in total blood flow to the forearm, since resting brachial artery blood flow [published in (Birk *et al.* 2012)] did not decrease after 8 weeks cycle exercise training (43.3 ± 20.2 vs 57.0 ± 28.5 mL/min, $P=0.14$). We therefore speculate that the decrease in skin flux that we observed following training in response to local skin heating may reflect enlargement of microvessels and/or an increase in the number of capillaries, in that larger microvascular beds should be associated with prolonged red cell transit time. Whilst we admit that such structural adaptation is a novel and speculative proposal in terms of the human cutaneous vasculature, some supportive evidence for microvascular adaptation after exercise training is provided by studies of skeletal muscle capillary density (Fernandes *et al.* 2012a; Fernandes *et al.* 2012b). Indeed, it has been suggested that increased microcirculatory volume in skeletal muscle following exercise training would counteract higher muscle blood flows, allowing sufficient time for extraction of oxygen (Krustrup *et al.* 2004). In support of this notion, Glieman *et al.* observed a training-induced increase in skeletal muscle capillary to fibre ratio as well as increases in capillary lumen area (Gliemann *et al.* 2015). These interesting findings in skeletal muscle suggest that whilst larger (conduit and feed) arteries adapt both functionally and structurally to increase flow (Green *et al.* 2017), there may be simultaneous changes in the microvessels that favour prolongation in red blood cell transit time to optimise gas exchange. This is consistent with our findings and the suggestion, albeit speculative at this stage, that skin adaptation to exercise or whole body heating may induce structural adaptations that are reflected as diminished maximal flux and CVC responses. Whilst changes in microvascular morphology in skeletal muscle may act to prolong transit time and enhance gas exchange, similar adaptation in the human skin would have the

effect of enhancing heat exchange. Further studies should examine these hypotheses and the suggestion that exercise training modifies cutaneous microvascular morphology in humans.

Our finding that changes in CVC were only present in the limb that was exposed to repeated increases in blood flow, infers that shear stress may be an important stimulus for adaptation in human microvessels. Although shear stress decreases during passage down the arterial tree, it is nonetheless likely that the vessels imaged in the present study are exposed to physiologically relevant increases in levels of flow and shear in response to exercise. *In vitro* animal studies have reported that coronary arterioles between 80-130 μ m dilated by 50% and arterioles between 25-70 μ m dilated by 20-30% in response to shear stress (Kuo et al. 1995). The vessels imaged in this study are likely >30-50 μ m in diameter (Fredriksson et al. 2009; Braverman 1989). Also, Laughlin et al. reported that arterioles of smaller size possess more eNOS protein than those >50 μ m (Laughlin et al. 2003). Regardless of the magnitude of shear the skin microcirculation was exposed to, blood flow through these areas was significantly increased, and this is associated with increased haemodynamic forces, that can act as mediators of adaptation (Brown 2003).

Limitations. Cuff inflation to 60mmHg was used to primarily attenuate blood flow in one limb during the exercise bouts. It is conceivable that cuff inflation may have induced unilateral venous distension which, in turn, elicited a reflex cardiovascular response impacting upon skin perfusion or flow (Cui et al. 2012). For example, previous studies that have inflated a cuff to 250mmHg and subsequently distended the veins via infusion, have reported a systemic reflex sympathetic impact (Cui et al. 2011; Cui et al. 2009; Cui et al. 2012). If such a reflex was induced in the present experiment, it might explain the contralateral uncuffed limb responses we observed.

However, our experimental design was different to these studies because it involved cuff inflation to just 60 mmHg. Nonetheless, we performed a sub-study to determine potential systemic effects of unilateral cuff inflation to 60mmHg for 30 minutes (n=5). We observed no effects on blood pressure, HR or skin blood flows in the contralateral limb. We therefore think it unlikely, in this particular experiment, that the effects of cuff inflation could drive a large sympathetic response that would induce adaptation in contralateral skin blood flow responses, at rest, to local heating pre- versus post-training. Furthermore, any central reflex effect, if apparent, would manifest in both arms, cancelling out the likely impact in terms of our experimental design. If a unilateral vasoconstrictor reflex was evident in just the cuffed limb as a result of venous distension, then the microvascular adaptation to repeated episodic vasoconstriction should have been greater in the cuffed arm, yet the opposite was the case in this study. We observed no changes in skin microvascular responses to local heating at rest in the arm that had undergone cuffing during the training intervention (Figure 2 and 3). We therefore suggest that elicitation of reflex responses cannot explain the differences we observed between limbs. Another limitation is that we did not examine core temperature responses during acute cycle exercise, before and after exercise training. We cannot, therefore, relate our findings to changes in thermoregulatory capacity, but this was not our aim. The purpose of our study was related to intrinsic adaptations in cutaneous responses following exercise training, and the role of blood flow and temperature in such localised adaptations, which may be relevant to microvascular health and disease. A further limitation relates to our subject group selection. Whilst we studied subjects who were either sedentary or recreationally active (all <5 hrs regular exercise per week), we cannot exclude the possibility that more active or fit subjects may have exhibited different outcomes. Finally, the local heating protocol employed in the present study was not identical to

that used by Black *et al.*. Whilst the rate of temperature increase has been shown to have little effect on the absolute peak skin blood flow response (Choi *et al.* 2014), we cannot entirely rule out the possibility that the use of a different local heating protocol may have induced distinct results to our study.

Conclusion

We found that cycle exercise training decreased forearm skin blood flow responses to localised heating. This adaptation appears to be dependent upon repeated increases in blood flow during exercise bouts, as adaptation in the skin microcirculation was not apparent in the contralateral limb in which flow responses and skin temperature were attenuated via cuff inflation. We speculate that an increase in cutaneous red cell transit time may result from structural adaptations that, correspondingly, lower maximal CVC responses. Although we cannot provide a definitive evidence mechanism for such an adaptation at this stage, due to methodological limitations with the direct assessment of cutaneous microvascular structure, future studies utilising advanced imaging technology (Carter *et al.* 2016) may be able to test this proposal.

REFERENCES

- Andersen P, Henriksson J (1977) Capillary supply of the quadriceps femoris muscle of man: adaptive response to exercise. *J Physiol* 270 (3):677-690. doi:10.1113/jphysiol.1977.sp011975
- Birk GK, Dawson EA, Atkinson C, Haynes A, Cable NT, Thijssen DHJ, Green DJ (2012) Brachial artery adaptation to lower limb exercise training: role of shear stress. *J Appl Physiol* 112 (10):1653-1658. doi:10.1152/japplphysiol.01489.2011
- Black MA, Green DJ, Cable NT (2008) Exercise training prevents age-related decline in nitric oxide (NO)-mediated vasodilator function in human microvessels. *J Physiol* 586:3511-3524
- Blair S, N., Morris J, N. (2009) Healthy Hearts—and the universal benefits of being physically active: Physical activity and health. *Ann Epid* 19:253-256
- Boegli Y, Gremion G, Golay S, Kubli S, Liaudet L, Leyvraz P-F, Waeber B, Feihl F (2003) Endurance training enhances vasodilation induced by nitric oxide in human skin. *J Invest Dermatol* 121:1187-1204
- Boignard A, Salvat-Melis M, Carpentier PH, Minson CT, Grange L, Duc C, Sarrot-Reynauld F, Cracowski J-L (2005) Local hyperemia to heating is impaired in secondary Raynaud's phenomenon. *Arthritis Res Ther* 7:R1103. doi:10.1186/ar1785
- Braverman IM (1989) Ultrastructure and Organization of the Cutaneous Microvasculature in Normal and Pathologic States. *J Investig Dermatol* 93 (2s):2S-9S
- Brown MD (2003) Exercise and coronary vascular remodelling in the healthy heart. *Exp Physiol* 88:645-658

- Brunt VE, Eymann TM, Francisco MA, Howard MJ, Minson CT (2016) Passive heat therapy improves cutaneous microvascular function in sedentary humans via improved nitric oxide-dependent dilation. *J Appl Physiol* 121 (3):716
- Carter HH, Gong P, Kirk RW, Es, haghian S, Atkinson CL, Sampson DD, Green DJ, McLaughlin RA (2016) Optical coherence tomography in the assessment of acute changes in cutaneous vascular diameter induced by heat stress. *J Appl Physiol* 121 (4):965-972
- Carter HH, Spence AL, Atkinson CL, Pugh CJA, Cable NT, Thijssen DHJ, Naylor LH, Green DJ (2014) Distinct Effects of Blood Flow and Temperature on Cutaneous Microvascular Adaptation. *Med Sci Sport Exer* 46 (11):2113-2121.
- Charkoudian N (2003) Skin Blood Flow in Adult Human Thermoregulation: How It Works, When It Does Not, and Why. *Mayo Clinic Proceedings* 78 (5):603-612.
doi:<http://dx.doi.org/10.4065/78.5.603>
- Choi PJ, Brunt VE, Fujii N, Minson CT (2014) New approach to measure cutaneous microvascular function: an improved test of NO-mediated vasodilation by thermal hyperemia. *J Appl Physiol* 117 (3):277
- Cocks M, Shaw CS, Shepherd SO, Fisher JP, Ranasinghe AM, Barker TA, Tipton KD, Wagenmakers AJM (2013) Sprint interval and endurance training are equally effective in increasing muscle microvascular density and eNOS content in sedentary males. *J Physiol* 591 (3):641-656. doi:10.1113/jphysiol.2012.239566
- Colberg SR, Parson HK, Nunnold T, Holton DR, Swain DP, Vinik AI (2005) Change in cutaneous perfusion following 10 weeks of aerobic training in Type 2 diabetes. *J Diabetes Complicat* 19 (5):276-283.

Cracowski J-L, Minson CT, Salvat-Melis M, Halliwill JR (2006) Methodological issues in the assessment of skin microvascular endothelial function in humans. Trends Pharmacol Sci 27:503-508

Cracowski JL, Gaillard-Bigot F, Cracowski C, Roustit M, Millet C (2011) Skin microdialysis coupled with Laser Speckle Contrast Imaging to assess microvascular reactivity. Microvascular Research 82 (3):333-338. doi:<http://dx.doi.org/10.1016/j.mvr.2011.09.009>

Cui J, Leuenberger UA, Gao Z, Sinoway LI (2011) Sympathetic and cardiovascular responses to venous distension in an occluded limb. Am J Physiol - Reg I 301 (6):R1831

Cui J, McQuillan P, Moradkhan R, Pagana C, Sinoway LI (2009) Sympathetic responses during saline infusion into the veins of an occluded limb. J Physiol 587 (14):3619-3627. doi:10.1113/jphysiol.2009.173237

Cui J, McQuillan PM, Blaha C, Kunselman AR, Sinoway LI (2012) Limb venous distension evokes sympathetic activation via stimulation of the limb afferents in humans. Am J Physiol - Heart C 303 (4):H457

Dawson EA, Low DA, Meeuwis IHM, Kerstens FG, Atkinson CL, Cable NT, Green DJ, Thijssen DHJ (2015) Reproducibility of Cutaneous Vascular Conductance Responses to Slow Local Heating Assessed Using seven-Laser Array Probes. Microcirculation 22 (4):276-284. doi:10.1111/micc.12196

Fernandes T, Magalhães FC, Roque FR, Phillips MI, Oliveira EM (2012a) Exercise Training Prevents the Microvascular Rarefaction in Hypertension Balancing Angiogenic and Apoptotic Factors. Hypertension 59 (2):513

Fernandes T, Nakamura JS, Magalhães FC, Roque FR, Lavini-Ramos C, Schettert IT, Coelho V, Krieger JE, Oliveira EM (2012b) Exercise training restores the endothelial progenitor

cells number and function in hypertension: implications for angiogenesis. *Journal of Hypertension* 30 (11):2133-2143. doi:10.1097/HJH.0b013e3283588d46

Fredriksson I, Larsson M, Strömberg T (2009) Measurement depth and volume in laser Doppler flowmetry. *Microvascular Research* 78 (1):4-13.

Gliemann L, Buess R, Nyberg M, Hoppeler H, Odriozola A, Thaning P, Hellsten Y, Baum O, Mortensen SP (2015) Capillary growth, ultrastructure remodelling and exercise training in skeletal muscle of essential hypertensive patients. *Acta Physiologica* 214 (2):210-220. doi:10.1111/apha.12501

Green DJ, Cable NT, Fox C, Rankin JM, Taylor RR (1994) Modification of forearm resistance vessels by exercise training in young men. *J Appl Physiol* 77 (4):1829-1833

Green DJ, Cable NT, Joyner MJ, O'Driscoll G (2008) Exercise and cardiovascular risk reduction: Updating the rationale for exercise. *J Appl Physiol* 105:766-768

Green DJ, Carter HH, Fitzsimons MG, Cable NT, Thijssen DHJ, Naylor LH (2010) Obligatory role of hyperaemia and shear stress in microvascular adaptation to repeated heating in humans. *J Physiol (Lond)* 588:1571-1577

Green DJ, Hopman MTE, Padilla J, Laughlin MH, Thijssen DHJ (2017) Vascular Adaptation to Exercise in Humans: Role of Hemodynamic Stimuli. *Physiol Rev* 97 (2):495

Green DJ, Spence A, Halliwill JR, Cable NT, Thijssen DHJ (2011) Exercise and vascular adaptation in asymptomatic humans. *Exp Physiol* 96:57-70

Groothuis JT, Poelkens F, Wouters CW, Kooijman M, Hopman MTE (2008) Leg intravenous pressure during head-up tilt. *J Appl Physiol* 105 (3):811-815. doi:10.1152/japplphysiol.90304.2008

- Hambrecht R, Adams V, Erbs S, Linke a, Krankel N, Shu Y, Baither Y, Geilen S, Thiele H, Gummert JF, Mohr FW, Schuler G (2003) Regular physical activity improves endothelial function in patients with coronary artery disease by increasing phosphorylation of endothelial nitric oxide synthase. *Circulation* 107:3152-3158
- Haskell WL, Sims C, Myll J, Bortz WM, St Goar FG, Alderman EL (1993) Coronary artery size and dilating capacity in ultradistance runners. *Circulation* 87 (4):1076-1082
- Hollowatz LA, Thompson-Torgerson CS, Kenney WL (2008) The human cutaneous circulation as a model of generalized microvascular function. *J Appl Physiol* 105:370-372.
doi:10.1152/japplphysiol.00858.2007
- Johnson JM, O'Leary DS, Taylor WF, Kosiba W (1986) Effects of local warming on forearm reactive hyperaemia. *Clin Physiol Oxf* 6::337-346
- Joyner MJ, Green DJ (2009) Exercise protects the cardiovascular system: effects beyond traditional risk factors. *J Physiol (Lond)* 587:5551-5558
- Kellogg DL, Liu Y, Kosiba IF, O'Donnell D (1999) Role of nitric oxide in the vascular effects of local warming of the skin in humans. *J Appl Physiol* 86:1185–1190
- Krustrup P, Hellsten Y, Bangsbo J (2004) Intense interval training enhances human skeletal muscle oxygen uptake in the initial phase of dynamic exercise at high but not at low intensities. *J Physiol (London)* 559 (1):335-345. doi:10.1113/jphysiol.2004.062232
- Kuo L, Davis MJ, Chilian WM (1995) Longitudinal Gradients for Endothelium-Dependent and - Independent Vascular Responses in the Coronary Microcirculation. *Circulation* 92 (3):518-525. doi:10.1161/01.cir.92.3.518
- Langille BL, O'Donnell F (1986) Reductions in arterial diameter produced by chronic decreases in blood flow are endothelium-dependent. *Nature* 231:405-407

- Laughlin MH (1995) Endothelium-mediated control of coronary vascular tone after chronic exercise training. *Med Sci Sports Exerc* 27 (8):1135-1144
- Laughlin MH, Turk JR, Schrage WG, Woodman CR, Price EM (2003) Influence of coronary artery diameter on eNOS protein content. *Am J Physiol* 284:H1307-1312
- Minson CT, Berry LT, Joyner MJ (2001) Nitric oxide and neurally mediated regulation of skin blood flow during local heating. *J Appl Physiol* 91 (4):1619-1626
- Minson CT, Holowatz LA, Wong BJ, Kenney WL, Wilkins BW (2002) Decreased nitric oxide- and axon reflex-mediated cutaneous vasodilation with age during local heating. *J Appl Physiol* 93 (5):1644-1649
- Naylor LH, Weisbrod CJ, O'Driscoll G, Green DJ (2005) Measuring peripheral resistance and conduit arterial structure in humans using Doppler ultrasound. *J Appl Physiol* 98 (6):2311-2315. doi:01047.2004 [pii]
10.1152/japplphysiol.01047.2004
- Newcomer S, Thijssen DHJ, Green DJ (2011) Effects of exercise on endothelium and endothelium/smooth muscle crosstalk: Role of exercise-induced hemodynamics. *J Appl Physiol* 111:311-320
- Perneger TV (1998) What's wrong with Bonferroni adjustments. *BMJ : British Medical Journal* 316 (7139):1236-1238
- Rowley NJ, Dawson EA, Birk GK, Cable NT, George K, Whyte G, Thijssen DH, Green DJ (2011) Exercise and arterial adaptation in humans: uncoupling localized and systemic effects. *J Appl Physiol* 110 (5):1190-1195.
- Silber D, McLaughlin D, Sinoway L (1991) Leg exercise increases peak forearm blood flow. *J Appl Physiol* 71:1568-1573

- Simmons GH, Wong BJ, Holowatz LA, Kenney WL (2011) Changes in the control of skin blood flow with exercise training: where do cutaneous vascular adaptations fit in? *Exp Physiol* 96 (9):822-828. doi:10.1113/expphysiol.2010.056176
- Smith CJ, Santhanam L, Bruning RS, Stanhewicz A, Berkowitz DE, Holowatz LA (2011) Upregulation of Inducible Nitric Oxide Synthase Contributes to Attenuated Cutaneous Vasodilation in Essential Hypertensive Humans. *Hypertension* 58 (5):935
- Snell PG, Martin WH, Buckey JC, Bloomqvist CG (1987) Maximal vascular leg conductance in trained and untrained men. *J Appl Physiol* 62:606-610
- Taylor WF, Johnson JM, Leary D, Park MK (1984) Effect of high local temperature on reflex cutaneous vasodilation. *J Appl Physiol* 57 (1):191
- Thijssen Dick H J, Cable N T, Green Daniel J (2012) Impact of exercise training on arterial wall thickness in humans. *Clinical Science (London, England : 1979)* 122 (Pt 7):311-322. doi:10.1042/CS20110469
- Thijssen DHJ, Dawson EA, Tinken TM, Cable NT, Green DJ (2009) Retrograde flow and shear rate acutely impair endothelial function in humans. *Hypertension* 53:986 - 992
- Tinken TM, Thijssen DHJ, Black MA, Cable NT, Green DJ (2008) Time course of change in vasodilator function and capacity in response to exercise training in humans. *J Physiol* 586 (20):5003-5012. doi:10.1113/jphysiol.2008.158014
- Tinken TM, Thijssen DHJ, Hopkins ND, Black MA, Dawson EA, Minson CT, Newcomer SC, Laughlin MH, Cable NT, Green DJ (2009) Impact of shear rate modulation on vascular function in humans. *Hypertension* 54:278-285
- Tinken TM, Thijssen DHJ, Hopkins ND, Dawson EA, Cable NT, Green DJ (2010) Shear stress mediates vascular adaptations to exercise training in humans. *Hypertension* 55:312-318

Williamson JW, Fadel PJ, Mitchell JH (2006) New insights into central cardiovascular control during exercise in humans: a central command update. *Experimental Physiology* 91 (1):51-58. doi:10.1113/expphysiol.2005.032037

Acknowledgements

D.J.G was funded by the Australian Research Council (DP 130103793). D.H.J.T was a recipient of the E. Dekker stipend (Netherlands Heart Foundation, 2009 T064).

Figure Legends.

Figure 1. Heart rate (A, in beats per minute), and forearm skin temperature (B, in °C) and skin blood flow (C, in perfusion units (PU)) in the uncuffed (solid squares) and cuffed arm (open squares) before and during cycle exercise (at 2.5-min intervals) in healthy, young volunteers (n=6). Post-hoc significantly different at P<0.05 from baseline (*) or between the cuffed and uncuffed arm (#). Error bars represent SE.

Figure 2. A representative response of skin blood flow change presented as cutaneous vascular conductance (CVC) during the gradual local heating protocol in the uncuffed (A) and cuffed (B) arms at Weeks 0 and 8 of the exercise intervention. PU: perfusion units; MAP: mean arterial pressure.

Figure 3. Forearm skin cutaneous vascular conductance (CVC) at baseline (33°C) and during gradual heating. Submaximal heating (40°C), 42°C (Plateau) and 44 °C (Max) are presented in the uncuffed (panel A) and cuffed arms (panel B) at 0 and following 8 weeks of exercise training (black and white bars, respectively) in healthy, young volunteers (n=14). Post-hoc significantly different at P<0.05 from baseline (*). Error bars represent SE.