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1 ABSTRACT

Background & Aims. Dietary flavonoids, such as those present in black tea, are associated with reduced risk of cardiovascular disease (CVD), possibly through improving nitric oxide (NO) mediated vascular function. The aim of this study was to examine the effect of acute black tea ingestion on cutaneous microvascular function.

Methods. Twenty healthy participants (58±5 yr, 9 men) attended two experimental 6 trials (tea, placebo), 7-days apart in a randomised, controlled, double-blind, cross-over 7 design. Participants ingested a single dose of 200ml black tea or placebo, followed by 8 assessment of forearm cutaneous perfusion using laser-Doppler flowmetry (LDF) 9 using three distinct heating protocols, enabling us to distinguish between axon- and 10 endothelium-dependent vasodilation: 1. rapid 42 °C, 2. rapid 39 °C and 3. gradual 42 °C. 11 12 On the contralateral arm, full-field laser perfusion imaging (FLPI) was used to assess forearm perfusion during gradual 42°C. Data were presented as cutaneous vascular 13 conductance (CVC; flux/mean arterial pressure, MAP) and CVC expressed as a 14 percentage of maximal CVC (%CVC_{max}). 15

Results. Rapid local heating to 39°C or 42°C demonstrated no effect of tea for flux,
CVC or %CVC_{max} (all P>0.05). Gradual local heating to 42°C, however, produced a
higher skin blood flow following black tea ingestion for absolute CVC (P=0.04) when
measured by LDF, and higher absolute flux (P<0.001) and CVC (P<0.001) measured</p>
with FLPI. No effect of tea was found for %CVC_{max} when assessed by either LDF or
FLPI.

Conclusions. Acute tea ingestion enhanced cutaneous vascular responses to gradual local heating to 42°C in healthy, middle-aged participants, possibly through a mechanism related to activation of endothelium-derived chemical mediators, such as

- NO. These improvements may contribute to the cardiovascular health benefits ofregular tea ingestion.
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- Key words: tea; flavonoids; cardiovascular health; vascular function;
 microcirculation.

30 **INTRODUCTION**

Cardiovascular disease (CVD) remains the leading cause of global mortality, 31 representing ~30% of all deaths.¹ The role of dietary factors on CVD risk has been 32 frequently explored in recent years, with a high dietary flavonoid intake being 33 associated with a reduction in CVD risk.² Tea, produced from the plant Camillia sinesis, 34 is the major dietary source of flavonoids in many countries globally³ and can be found 35 as catechins and flavonols in green tea and theaflavins, thearubigins and flavonols in 36 37 black tea.⁴ Accordingly, several studies have revealed a strong, inverse relation between regular intake of tea and cardiovascular risk.5,6 38

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A frequently cited explanation for the cardioprotective effects of black and green tea 40 ingestion relates to the reduction in blood pressure following chronic consumption.⁵⁻⁷ 41 Further research found that acute and regular tea ingestion improves nitric oxide-42 mediated, endothelium-dependent dilation of conduit arteries.^{6, 8-11} Both conduit and 43 resistance vessels have demonstrated improved endothelial function following tea 44 ingestion in both healthy individuals^{6, 8} and in those with CVD.¹⁰ Thus, the general 45 consensus is that regular tea ingestion improves blood pressure by virtue of a 46 generalised improvement of endothelial function and lowering of peripheral vascular 47 resistance.^{6, 8, 10-12} 48

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50 Despite encouraging data supporting a beneficial effect of tea ingestion in larger 51 (conduit) vessels, no previous study has explored the effect of black tea on small 52 vessels (skin microcirculation). Therefore, our aim was to examine cutaneous vascular 53 responses to local skin heating. Given the complexity of the cutaneous vascular 54 system and contribution of distinct mechanisms for skin dilation when gradually or rapidly heating the skin, we adopted a comprehensive approach of using rapid *and* gradual local skin heating protocols simultaneously. We hypothesised that black tea ingestion would be associated with increased cutaneous microcirculation responses for both rapid and gradual heating protocols.

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60 MATERIALS AND METHODS

61 **Participants**

Twenty middle-aged male (n=9) and post-menopausal female (n=11) participants 62 63 were recruited through local advertisement. All participants were healthy and nonsmokers (58±5yrs, height 1.70±0.1m, weight 75.9±16.1kg, BMI 26±4 kg/m², baseline 64 mean arterial pressure 104±8mmHg). Individuals with a medical history of 65 hypercholesterolaemia (total cholesterol >6.5mmol/l),¹³ cardiovascular disease and 66 hypertension (systolic blood pressure ≥140mmHg, diastolic blood pressure 67 ≥90mmHg)¹⁴ were excluded. Participants were not taking any vasoactive medications 68 or supplements. After being fully informed of the methods, written informed consent 69 was obtained from all participants. The study conformed to the Declaration of Helsinki 70 and was approved by the Research Ethics Committee of Liverpool John Moores 71 University. 72

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74 Experimental Design

All participants performed two experimental trials (tea and control), 7-days apart in a randomised, controlled, double-blind, cross-over design (figure 1). The cross-over design was chosen to eliminate between-participant variability, taking into account a 6-day washout period between the two interventions to avoid any carry-over effects, which is in accordance with previous similar designed cross-over tea vascular function

studies.^{6, 7} Computer-generated randomisation was used to reduce potential selection 80 bias. Upon arrival to the laboratory, and 2h prior to microvascular assessment, 81 participants ingested a tea drink (containing 300 mg flavonoids, 75 mg caffeine and 82 2.8 g sucrose) or a taste and appearance matched placebo drink (0 mg flavonoids, 75 83 mg caffeine, 2.7 g sucrose, tea flavour and caramel colour), prepared by dissolving 84 two sachets in 200 ml hot water. Participants subsequently rested for 2h prior to 85 commencement of testing to match peak plasma concentrations of flavonoids and 86 other metabolites such as phenolic acids, with testing of skin microcirculation. During 87 88 each testing day, baseline and thermally stimulated forearm cutaneous blood flow was examined simultaneously using rapid (to 39 and 42°C) and gradual (to 42°C) local 89 heating protocols. Since these protocols reflect different dilator mechanisms and a 90 distinct role of the NO-pathway, they provide complementary insight into the impact of 91 black tea on cutaneous microvasculature. Rapid local heating was performed at two 92 different sites (i.e. two different local heating protocols) on the dominant forearm and 93 examined using laser Doppler flowmetry (LDF). Gradual local heating to 42°C was 94 performed on the dominant forearm using LDF and on the contralateral (non-dominant) 95 arm using laser speckle imaging to provide whole forearm cutaneous microcirculation 96 function. 97

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99 Experimental Measures

All participants fasted for at least six hours and refrained from alcohol, food products high in polyphenols (dark chocolate, red wine), caffeine and exercise for 24h prior to testing.¹⁵ Participants were asked to refrain from drinking all types of tea for a period of one week prior to each trial. Sips of water were permitted prior to testing to ensure

that participants were well hydrated. All trials were conducted in a guiet, temperature 104 controlled environment (23.4±0.6°C)^{15, 16} and at the same time of day to reduce any 105 circadian influences on vascular function.¹⁵ Following a 20-minute stabilisation period, 106 the LDF equipment was calibrated using two generic points, 0 and 250 PU, a zeroing 107 disk and motility standard, according to manufacturer's guidelines (Perimed AB, 108 Järfälla, Stockholm, Sweden). Two hours following tea ingestion, participants 109 assumed a comfortable, supine position on a bed, with the head slightly elevated and 110 the hand of each testing arm relaxed, supinated and supported by a vacuum cushion 111 to minimise microcirculatory fluctuations resulting from motion artefact.^{15, 16} If 112 necessary, forearm measurement sites were shaved 24h prior to testing to avoid any 113 inflammatory response that may affect cutaneous blood flow; we inspected the 114 forearms prior to each trial to ensure that no skin damage was present that may 115 adversely influence cutaneous blood flow responses. Participants were instrumented 116 for LDF measurements on the dominant forearm; three heating discs (Perimed 355, 117 Perimed AB, Järfälla, Stockholm, Sweden) were placed ~5cm apart on the dominant 118 forearm, with a 7-laser array probe (PF 413, Perimed AB, Järfälla, Stockholm, Sweden) 119 placed into each heater and firmly attached to the skin using adhesive stickers and 120 medical tape. Following sterilisation of the non-dominant arm measurement site, 121 participants were instrumented for laser speckle imaging using the technique of full-122 123 field laser perfusion imaging (FLPI); a water-filled clear heating probe (Moor VHP3, Moor Instruments, Axminster, UK) was placed on the skin and attached using an 124 adhesive sticker to obtain a good seal. Measurement sites were randomly chosen, 125 avoiding visible veins, hair follicles and dermatological lesions.¹⁶ Upon completion of 126 the first experimental trial, the location of the LDF and FLPI assessment sites was 127 marked on the skin, with digital photographs and measurements taken to the nearest 128

millimetre using anatomical and skin-surface landmarks for reference, to ensure accurate re-selection of probe sites for the second trial. Stature (seca 217 stadiometer, seca UK, Birmingham, UK) and body mass (seca 767 calibrated electronic scales, Germany) were recorded using standardised protocols. Body mass index was calculated (BMI) as the body weight (kg) divided by the height squared (m²).

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Both LDF and FLPI provide non-invasive, continuous measures of cutaneous blood 135 flow.¹⁷ By using a combination of these techniques, it is possible to simultaneously 136 evaluate superficial (<300 micron) and deeper (1-1.5mm) skin blood flow via FLPI and 137 LDF, respectively. Rather than assessing overall microvascular function, using local 138 thermal hyperaemia provides a more comprehensive assessment of microvascular 139 140 reactivity to acute tea ingestion and the complex neural and chemically-mediated pathways underlying microvascular function. Distinct rapid and gradual local heating 141 protocols all provide a different type of vasodilation that likely relates to different 142 vasodilator pathways. 143

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Rapid local heating. First, we adopted the classic local heating protocol rapid 42°C 145 (0.5°C per 5s, 30-min at 42°C, 20-min at 44°C), which induces a rapid, transient axon-146 reflex, followed by a more gradual, but sustained, heating response. The plateau 147 phase represents 80-90% of the maximal response, and is partly (60-70%) NO-148 mediated.^{18, 19} Secondly, we examined a more recently introduced protocol; rapid 39°C 149 (0.5°C per 5s, 30-min at 39°C, 20-min at 44°C), that also induces an axon-reflex and 150 gradual plateau during the heating response. By stopping the heating protocol at 39°C, 151 the plateau phase is largely NO-mediated and only represents 50% of the maximal 152

response.²⁰ The *rapid 42* °C and *rapid 39* °C protocols were examined simultaneously
on the dominant arm using LDF.

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Gradual local heating. We examined an adapted version of the gradual local heating protocol that increases to $42 \,^{\circ}C$ (0.5°C per 2min30s, 30-min at 42°C, 20-min at 44°C), and induces a slow heating response that is largely NO-mediated and reflects 80-90% of the maximal response.²¹ This protocol was examined on the dominant arm using LDF, covering 6mm² of skin at a penetration depth of ~1-1.5mm, and on the contralateral arm using FLPI, which covers an area of skin up to 30,000mm² at a depth of ~0.3mm.²²

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Laser Doppler Flowmetry (LDF). Laser Doppler flowmetry is a non-invasive technique 164 that is routinely used to study microvascular function,^{16, 18, 22} and is sensitive in 165 detecting changes in skin perfusion over a period of time and in response to a stimulus, 166 such as local thermal hyperaemia.¹⁶ LDF is concerned with the reflection of a laser 167 168 beam that undergoes a change in wavelength, or Doppler shift, when it detects moving red blood cells, the magnitude and frequency of which is related to the concentration 169 and velocity of blood cells and is recorded as a signal of red blood cell flux (RBCF).^{16,} 170 ²² Following a 20-minute acclimation period, cutaneous blood flow was measured as 171 RBCF at the chosen probe sites using a laser Doppler flowmeter (Periflux system 5000, 172 Perimed AB, Järfälla, Stockholm, Sweden). The local heating discs were connected 173 to a heating unit (Peritemp 4005 heater, Perimed AB, Järfälla, Stockholm, Sweden) 174 which was manually controlled to perform the temperature stages of the local heating 175 protocols. Baseline skin RBCF was recorded with the local heating disc temperature 176

set at 33°C for 10-minutes for each measurement site. Subsequently, local skin
temperature was heated using the three distinct protocols.

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Full-field Laser Perfusion Imaging (FLPI). The FLPI technique, also known as laser 180 speckle contrast imaging, exploits the fact that the random speckle pattern that is 181 generated when tissue is illuminated by laser light, changes when blood cells move 182 within the region of interest.²³ High levels of movement (fast flow) produce a more 183 blurred pattern, associated with a reduction in contrast in that region. Low contrast 184 corresponds with high flow and high contrast corresponds with low flow. The strengths 185 of this technique are that video frame rate blood flow images (up to 25 per second) 186 187 enable the tracking of fast transient blood flow changes and provides high spatial and 188 temporal resolution. This device works with a near infra-red laser diode (785nm) and is able to scan skin surfaces from 5mm x 7mm to 15cm x 20cm, to a depth of 189 approximately 150-300 micron and is safe for human use. Following a 20-minute 190 acclimation period, FLPI recordings of the non-dominant forearm were performed 191 using a blood flow imaging system (moorFLPI-1, Moor Instruments, Axminster, UK) 192 with a laser wavelength of 785nm and sampling frequency of 25Hz. The distance 193 between the laser head and skin surface was fixed at 15cm.²⁴ A skin heater module 194 (moorVMS-HEAT, Moor Instruments, Axminster, UK) was used to manually set the 195 baseline temperature at 33°C for 10-minutes and to perform the incremental local 196 heating protocol; gradual 42°C_{FLPI} (0.5°C per 2min30s, 30-min at 42°C, 20-min at 197 44°C).²¹ 198

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200 Haemodynamics. Heart rate (HR) and blood pressure were recorded at the beginning and at the end of the 20-minute acclimation period using an automated 201 sphygmomanometer (Dinamap V100, GE Healthcare, UK) positioned on the ankle, 202 corresponding to the same laterality as their dominant arm. Thereafter, mean arterial 203 pressure (MAP, mV/mmHg) and HR were recorded at 5-minute intervals throughout 204 the local heating protocols. MAP was used to calculate cutaneous vascular 205 conductance (CVC=RBCF/MAP), thereby accounting for changes in skin blood flow 206 resulting from variations in blood pressure.^{16, 18, 25} 207

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209 Data Analysis

Data analysis was performed blind. Cutaneous RBCF (PU) was expressed as 210 cutaneous vascular conductance (CVC), as described previously.¹⁶ Artefact in the 211 data, due to unwanted subject movement, was identified and removed prior to 212 analysis. Baseline laser Doppler RBCF was averaged over a stable 10-minute 213 baseline period. For the rapid 42°C and rapid 39°C protocols, following initiation of 214 heating, initial peak and nadir CVC values were calculated over a stable 60-second 215 216 period,¹⁸ with the initial peak identified as the highest value and the nadir as the lowest value during the first 5-10 minutes of local heating.²⁶ A clear nadir was not detected in 217 all measurement traces, which is typical of this type of thermal provocation test. In 218 those traces (~5%), data was included from a 60-second period, 1-minute after the 219 initial peak. This value was always lower than the initial peak. CVC was calculated 220 over a stable 60-second period for the final minute of each temperature increment (34-221 41°C) of the gradual 42°C local heating protocol. For each of the three protocols, rapid 222 42°C, rapid 39°C and gradual 42°C, plateau phases during heating (42°C, 39°C and 223 maximal 44°C) were averaged over the last 5-minutes of each phase. Data at baseline 224

and at the various plateau phases were also normalised to the maximal CVC achieved at 44°C (%CVC_{max}=[CVC/CVC_{max}] x 100).²⁵ All data were collected in LabChart 7.0 (ADInstruments, Dunedin, New Zealand).

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229 Statistical Methods

Data were expressed as mean±SD and statistical significance was set at P<0.05. For 230 all protocols, linear mixed models (main effects of condition and time) were used to 231 examine the impact of acute tea ingestion on blood pressure and forearm skin 232 microcirculation. The repeated covariance type was Unstructured and Condition, Time 233 and Condition*Time was specified as Fixed Effects (intercept was included) and as 234 235 Estimated Marginal Means. We interpreted the Test of Fixed Effects Condition*Time 236 interaction. Significant main effects of Time or Condition or a Time*Condition interaction were followed up with a simple main effects analysis and the least 237 significant difference (LSD) approach to multiple comparisons.²⁷ Data were stored and 238 transformed within Microsoft Excel (Microsoft Office 2010, Microsoft Corporation), and 239 statistical analyses were performed using SPSS 22.0 (SPSS, Chicago, IL, USA). 240

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243 **RESULTS**

One participant was removed from the *gradual* $42 \,^{\circ}C_{LDF}$ analysis for both experimental trials (due to probe failure) and five participants were removed from the *gradual* $42 \,^{\circ}C_{FLPI}$ analysis for both trials (linked to excessive movement artefacts), giving a population of n=19 and n=15, respectively. No participants were removed from the *rapid* 39 $^{\circ}C$ and *rapid* 42 $^{\circ}C$ analysis (both n=20). Baseline MAP was not different between conditions (108±11, 108±11, P=0.73) and showed no change across time (P=0.52). There were no differences in baseline cutaneous perfusion between trials for measurement sites that underwent *rapid* 39°C or *rapid* 42°C local heating for absolute flux, CVC or %CVC_{max} (Table 1). Also the site that underwent *gradual* 42°C local heating using LDF showed no difference in baseline cutaneous blood flow between trials for absolute flux, CVC or %CVC_{max} (Table 2). However, using FLPI, a significantly higher baseline perfusion was found after tea ingestion for cutaneous flux and CVC, but not for %CVC_{max} (Table 2).

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258 Rapid local heating: impact of tea

Rapid 39°C. Local heating induced a typical pattern of an initial peak, nadir and plateau in cutaneous blood flow. Therefore, a main effect of time was demonstrated for absolute flux, CVC and %CVC_{max} (Table 1). However, we found no effect of the intervention or a timeXintervention-interaction for absolute flux, CVC or %CVC_{max} (Table 1).

Rapid 42 °C. Local heating induced a typical pattern of an initial peak, nadir and plateau
in cutaneous blood flow. Consequently, a main effect of time was demonstrated for
absolute flux, CVC and %CVC_{max} (Table 1), whilst no main effect of intervention or
timeXintervention-interaction was found for absolute flux, CVC or %CVC_{max} (Table 1).

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269 Gradual local heating: impact of tea

Gradual 42 °*C*_{*LDF*}. Local heating induced a gradual, slow heating response with no detectable initial axon reflex-induced peak or nadir with a main effect of time (Table 2). A higher skin blood flow throughout the heating protocol was observed during the trial preceded by black tea for absolute CVC (P=0.04), with a trend towards significance when data were presented as absolute flux (P=0.06, Table 2). No effect of tea was found when CVC was normalised for maximum perfusion (%CVC_{max}, P=0.82, Table 276 2). No timeXintervention-interaction was found for absolute flux (P=0.93), CVC 277 (P=0.95) or %CVC_{max} (P=0.98, Table 2).

Gradual 42 °C_{*FLPI*}. Local heating induced a gradual, slow heating response with no detectable initial axon reflex-induced peak or nadir (Table 2). Tea ingestion was associated with a significantly higher absolute flux (P=0.00) and CVC (P=0.00), but not when CVC was normalised to maximum CVC (%CVC_{max}, P=0.35, Table 2). No timeXintervention-interaction was present for absolute flux (P=0.50), CVC (P=0.66) or %CVC_{max} (P=1.00, Table 2).

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285 Our statistical analysis revealed no presence of a carry-over effect.

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287 **DISCUSSION**

The aim of this study was to test the hypothesis that a single dose of black tea 288 ingestion improves cutaneous microcirculation following both rapid and gradual local 289 skin heating. We found that gradual local heating of the skin to 42°C induced a greater 290 vasodilatory response following tea ingestion compared to placebo when expressed 291 as absolute flux and CVC. The ability of tea to improve local gradual heating responses 292 in the skin was reinforced by the observation that both LDF and FLPI, two distinct but 293 accepted techniques to assess skin perfusion, detected this effect. Conversely, rapid 294 local heating did not demonstrate a significant increase in cutaneous microcirculation 295 with tea ingestion, either for the rapid 39°C or rapid 42°C protocols. Taken together, 296 our study provides some further evidence that regular tea ingestion may mediate its 297 potential cardiovascular benefits via improvements in (cutaneous) microvascular 298 function. 299

Our study is the first to explore the acute effects of tea ingestion on the cutaneous 301 microcirculation whilst adopting a rigorous protocol involving blind analysis of rapid 302 303 and gradual heating protocols as well as two distinct, accepted techniques. This observation fits with the general observation of tea being able to enhance endothelial 304 function in conduit vessels when assessed by flow-mediated dilation (FMD).^{6, 9} Taken 305 together, these findings suggest that acute tea ingestion improves vascular function 306 across the vascular tree, including skin microvessels, possibly via upregulation of 307 308 vasodilator mechanisms.

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In contrast to gradual local heating, rapid heating of the skin did not alter cutaneous 310 vascular function following tea ingestion when compared to placebo. Our findings were 311 similar for both rapid heating protocols (rapid 39°C and rapid 42°C). Interestingly, a 312 recent observational study²⁸ found improved microvascular function following regular 313 consumption of green tea (14 days) using rapid heating (whilst no measure of gradual 314 heating was included). Important differences were present between studies, especially 315 since this previous study did not include a placebo control, did not fully adhere to 316 guidelines for vascular assessment (e.g. control of menstrual cycle),¹⁵ and was limited 317 by a lack of control of dietary habits.⁶ Furthermore, whilst our study investigated the 318 319 acute (2h) effects of tea, they examined a protocol of 14 days of green tea. Despite the rapid 39°C and gradual 42°C protocols both being linked to the release of NO, 320 distinct responses are clearly evident between the gradual and rapid heating protocols 321 in our study. Different vasodilator pathways directly influence the cutaneous 322 microcirculation, including neurogenic reflexes and local chemical mediators.^{18, 21, 25} 323 The rate at which the skin is heated, alters the contribution of these vasodilator 324

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pathways, with rapid (0.5°C per 5s) local heating inducing a transient axon-reflex 325 mediated vasodilation that is produced via activation of heat sensitive 326 sensory/nociceptive nerves releasing calcitonin gene-related peptide (CGRP) and 327 substance P and adrenergic nerves releasing norepinephrine and neuropeptide Y.^{18,} 328 ²⁹ This initial neurogenic response is followed by a more gradual, sustained 329 vasodilation. In both phases, vasodilation occurs through complex pathways that lead 330 to the production of NO and smooth muscle relaxation via hyperpolarization from 331 endothelial derived hyperpolarization factors (EDHFs),²⁹ with a greater (but not 332 exclusive) contribution of NO during the plateau phase.^{18, 20} Furthermore, the relative 333 contribution of NO to the vasodilation during the plateau phase of the rapid heating 334 protocols depends upon the target heating temperature, as the heating response to 335 39°C seems to depend more on NO than the response to 42°C.^{18, 20} These studies, 336 therefore, demonstrate that the underlying mechanism for cutaneous vasodilation 337 differ based on the rate and maximum level of heating. The different vasodilator 338 pathways for these heating protocols may contribute to the distinct findings in our study. 339 From a methodological perspective, the differences between rapid and gradual local 340 heating highlight the importance of using multiple heating protocols simultaneously 341 when exploring the impact of an intervention on skin perfusion. 342

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The higher vasodilatory responses that we observed following gradual heating of the skin were demonstrated for arbitrary flux and CVC values, for both LDF and FLPI techniques. However, the difference in responses between the tea and placebo trials was not significant when data were expressed as %CVC_{max}. The skin is commonly heated to 44°C to reach maximal vasodilation and expressing CVC as a percentage of maximal perfusion is often considered the preferred method of data expression,¹⁶ with improved reproducibility compared to flux or CVC.²⁵ Despite a main effect of tea on flux and CVC, post-hoc analyses revealed no differences between trials at 44°C (LFD: flux=0.17 and CVC=0.19; FLPI: flux=0.09 and CVC=0.08). However, the magnitude of differences in flux and CVC between tea and placebo are larger than one may expect based on day-to-day variation.²⁵ This provides some indication that the tea intervention may have altered cutaneous perfusion at 44°C local heating.

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Clinical Relevance. Tea consumption is known to have cardiovascular benefits, 357 358 including a reduction in blood pressure after short- to long-term intervention, possibly mediated (in part) by improved endothelial function of conduit vessels.^{8, 9, 12} In our 359 study, cutaneous microcirculation responses to gradual heating improved following tea 360 ingestion. We speculate that these findings may have implications for individuals with 361 microvascular complications and skin endothelial dysfunction, such as type 2 diabetes 362 mellitus. Interestingly, consumption of tea has been associated with a reduced risk for 363 type 2 diabetes mellitus.³⁰ Our findings thus support the hypothesis that regular tea 364 consumption may have potential benefit in such patient groups. Future studies are 365 warranted to explore this hypothesis. 366

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Limitations. Due to our modest sample size, we are unable to generalise our findings towards the wider populace. Furthermore, although we included a middle-aged population who likely are at an increased risk of CVD, we cannot simply extrapolate our findings to clinical groups. Moreover, our population may have impaired endothelial function as blunted cutaneous NO-mediated vasodilation has been demonstrated in older individuals,²¹ suggesting that young healthy volunteers may exhibit different results than our older population. Therefore, future work is required to

explore the potential impact of acute as well as chronic tea ingestion on cutaneous 375 vascular function in both individuals with compromised endothelial function and in 376 young, healthy individuals. A further limitation is that we did not obtain plasma 377 measures of flavonoids or NO compounds and, therefore, our study does not provide 378 any biochemical or biomolecular insight into the mechanisms underlying the 379 improvement in cutaneous microvascular function. However, it is important to 380 emphasise that this was not the purpose of our study, particularly given that we are 381 the first to explore the effects of acute tea ingestion on the cutaneous microcirculation. 382

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In conclusion, our findings suggest that acute tea ingestion enhances cutaneous 384 vascular function in a healthy, middle-aged population, when measured following 385 gradual local heating to 42°C. Therefore, these data suggest that acute tea ingestion 386 has a beneficial impact on vascular function at the microcirculatory level, which is likely 387 achieved through a mechanism related to activation of endothelium-derived 388 vasodilators. These improvements in cutaneous microvascular function may 389 contribute to the potential cardiovascular health benefits of regular tea ingestion. 390 Future studies are required to explore the acute and chronic effects of tea on 391 individuals with increased CVD risk and in clinical populations with a priori endothelial 392 dysfunction. 393

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395 STATEMENT OF AUTHORSHIP

K. A. W., D. A. L., D. H. J. T., N. D. H., R. D. and Y. d. G. designed research; K. A.
W. conducted research; K. A. W., D. A. L. and Y. d. G. analysed data; K. A. W., D. A.
L., D. H. J. T., R. D. and N. D. H. wrote the paper; R. D. and D. H. J. T. had primary
responsibility for final content. All authors read and approved the final manuscript.

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484 for the *rapid 39°C* and *rapid 42°C* protocols for placebo and tea interventions.

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Rapid 39°C	Placebo	Теа	time	LMM tea	time*tea
Absolute flux(PU)					
Baseline	22 ± 11	21 ± 8			
Axon-reflex	108 ± 38	103 ± 50			
Nadir	57 ± 25	52 ± 26	<0.001*	0.14	0.76
Plateau 39°C	136 ± 53	123 ± 70		-	
Plateau 44 °C	288 ± 61	263 ± 61			
Absolute CVC (PU/mmHg)					
Baseline	0.21 ± 0.12	0.21 ± 0.10			
Axon-reflex	1.03 ± 0.39	0.99 ± 0.47			
Nadir	0.54 ± 0.25	0.50 ± 0.26	<0.001*	0.27	0.91
Plateau 39°C	1.29 ± 0.52	1.17 ± 0.65			
Plateau 44 °C	2.70 ± 0.67	2.52 ± 0.59			
Maximal CVC (%CVC _{max})					
Baseline	8 ± 4	8 ± 3			
Axon-reflex	39 ± 15	39 ± 15			
Nadir	20 ± 10	20 ± 10	<0.001*	0.76	0.99
Plateau 39°C	48 ± 15	46 ± 21			
Rapid 42°C					
Absolute flux(PU)					
Baseline	22 ± 9	25 ± 16			
Axon-reflex	199 ± 60	208 ± 60			
Nadir	165 ± 64	177 ± 74	<0.001*	0.51	0.99
Plateau 42°C	252 ± 72	253 ± 67			
Plateau 44 °C	300 ± 79	302 ± 63			
Absolute CVC (PU/mmHg)					
Baseline	0.21 ± 0.10	0.25 ± 0.16			
Axon-reflex	1.90 ± 0.61	2.00 ± 0.61			
Nadir	1.57 ± 0.64	1.71 ± 0.74	<0.001*	0.29	1.00
Plateau 42°C	2.39 ± 0.74	2.43 ± 0.66			
Plateau 44 °C	2.81 ± 0.81	2.91 ± 0.74			
Maximal CVC (%CVC _{max})					
Baseline	8 ± 3	8 ± 4			
Axon-reflex	67 ± 11	68 ± 11	0.004	a a-	
Nadir	55 ± 16	57 ± 19	<0.001*	0.65	0.95
Plateau 42 °C	85 ± 8	83 ± 12			

486 Data are mean ± SD. *Main effect of time *P*<0.001 vs baseline.

- 487 **Table 2.** Cutaneous blood flow responses to local heating for the gradual_{LDF} (42 °C)
- 488 and gradual_{FLPI} (42 °C) protocols for placebo and tea interventions.

489

Intervention (mean ± SD)						
			LMM			
Gradual _{LDF} (42°C)	Placebo	Теа	time	tea ti	me*tea	
Absolute flux(PU)						
Baseline	26 ± 11	24 ± 9				
Plateau 42°C	268 ± 79	278 ± 61	<0.001*	0.06	0.93	
Plateau 44 °C	302 ± 84	319 ± 45				
Absolute CVC (PU/mmHg)						
Baseline	0.25 ± 0.11	0.23 ± 0.09				
Plateau 42°C	2.51 ± 0.76	2.61 ± 0.64	<0.001*	0.04^	0.95	
Plateau 44 °C	2.80 ± 0.82	2.93 ± 0.51				
Maximal CVC (%CVC _{max})						
Baseline	9 ± 5	8 ± 3				
Plateau 42°C	90 ± 7	89 ± 14	<0.001*	0.82	0.98	
Gradual _{FLPI} (42°C)						
Absolute flux(PU)						
Baseline	30 ± 9	36 ± 8				
Plateau 42°C	197 ± 51	222 ± 50	<0.001*	<0.001	^ 0.50	
Plateau 44 °C	216 ± 65	253 ± 68				
Absolute CVC (PU/mmHq)						
Baseline	0.29 ± 0.09	0.36 ± 0.07				
Plateau 42 °C	1.85 ± 0.55	2.10 ± 0.57	<0.001*	<0.001	^ 0.66	
Plateau 44 °C	2.01 ± 0.64	2.34 ± 0.72				
Maximal CVC (%CVC _{max})						
Baseline	17 ± 11	17 ± 8				
Plateau 42 °C	94 ± 10	91 ± 6	<0.001*	0.35	1.00	

490 Data are mean ± SD. *Main effect of time *P*<0.001 vs baseline. ^Main effect of intervention; placebo

491 vs. tea *P*<0.05.

492 **FIGURE LEGENDS**

493 Figure 1. CONSORT diagram showing the flow of participants through each stage of494 the randomised trial.

495

Figure 2. Study overview and schematic depicting the stages of the local heating protocols. Light grey shading denotes local heating, mid grey shading represents the plateau and dark grey shading represents the maximal plateau.

500

Figure 3. Cutaneous vascular conductance (CVC) responses across time points
(baseline at 33 °C, axon peak, axon nadir, plateau at 39/42 °C and maximal
plateau at 44 °C) following rapid local heating for A. *rapid 39* °C and B. *rapid 42* °C in 20 healthy volunteers when heating was preceded by
ingestion of placebo (open squares) or tea (solid triangles). Data are
presented as means, with error bars representing SE.

507

Figure 4. Cutaneous vascular conductance (CVC) responses across time points
(from baseline at 33 °C to maximal plateau at 44 °C) following gradual local
heating using A. laser-Doppler flowmetry (LDF) and B. full-field laser
perfusion imaging (FLPI) in 20 healthy volunteers when heating was
preceded by ingestion of placebo (open squares) or tea (solid triangles).
Data are presented as means, with error bars representing SE. *Main effect
of condition *P*<0.05 placebo vs tea.