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Exercise intensity and duration effects on in vivo immunity

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ABSTRACT

Purpose: To examine the effects of intensity and duration of exercise stress on induction of in-vivo immunity in humans using experimental contact hypersensitivity (CHS) with the novel antigen Diphenylcyclopropenone (DPCP). Methods: Sixty-four healthy males completed either 30 minutes running at 60% \dot{V} O_{2peak} (30MI); 30 minutes running at 80% \dot{V} O_{2peak} (30HI); 120 minutes running at 60% \dot{v} O_{2peak} (120MI) or seated rest (CON). Twentyminutes later subjects received a sensitizing dose of DPCP and four-weeks later the strength of immune reactivity was quantified by measuring the cutaneous responses to a low, doseseries challenge with DPCP on the upper inner-arm. Circulating epinephrine, norepinephrine and cortisol were measured pre, post and 1h post-exercise or CON. Next, to better understand whether the decrease in CHS response on 120MI was due to local inflammatory or T-cell mediated processes, in a cross-over design, eleven healthy males performed 120MI and CON and cutaneous responses to a dose-series of the irritant, croton oil (CO) were assessed on the upper inner-arm. Results: Immune induction by DPCP was impaired by 120MI (skin-foldthickness -67% vs CON; P<0.05). However, immune induction was unaffected by 30MI and 30HI despite elevated circulating catecholamines (30HI vs pre: P<0.01) and greater circulating cortisol post 30HI (vs CON: P<0.01). There was no effect of 120MI on skin irritant responses to CO. Conclusions: Prolonged, moderate-intensity exercise, but not shortlasting high or short-lasting moderate-intensity exercise, decreases the induction of *in-vivo* immunity. No effect of prolonged, moderate-intensity exercise on the skin's response to irritant challenge points towards a suppression of cell-mediated immunity in the observed decrease in CHS. DPCP provides an attractive tool to assess the effect of exercise on in-vivo immunity.

Key words: stress; running; immune; contact hypersensitivity; diphencyprone; irritant

INTRODUCTION

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against pathogenic and environmental assaults (8). Measures of in vivo immunity at the skin 3 4 include delayed type hypersensitivity (DTH) responses to intradermal injection of antigens, or 5 the less invasive contact hypersensitivity (CHS) responses to epicutaneous application of 6 antigens. These in vivo measures are considered more informative than the commonly used in 7 vitro measures where immune cells, typically from peripheral blood, are extracted from their normal environment and analyzed in artificial cultures (2). Isolated measures of immune 8 function may react differently to a whole-body immune challenge because they lack the highly 9 10 integrated neural and hormonal components within the specific tissue environment in which immune responses usually take place (1). Studies using *in vivo* cutaneous immune measures 11 12 have shown impaired responses in individuals exposed to psychological stress (3), physical stress (17), during acute infectious illness e.g. Epstein-Barr virus (5) and in diabetes and 13 psoriasis (4). Furthermore, *in vivo* cutaneous immune measures have been shown to predict 14 mortality in critically ill HIV-infected patients (12) and in patients with surgical infections (31). There is a need to better understand *in vivo* cutaneous immune measures for investigators 16 examining the influence of exercise stress on immunity in humans. 17 18 Physical exercise provides a well-controlled model to study the effects of stress on immune responses. Given the obvious ethical constraints of studying experimental infection in humans, 20 animal models have provided valuable insight into the effects of exercise on clinically relevant 21 responses to viral infection. The work in animals indicates that prolonged and high intensity 22 23 exercise is associated with higher mortality rates whereas short, moderate intensity exercise lowers mortality rates, compared with controls (21). The research evidence on immune 24 responses after short, moderate intensity exercise in humans is not definitive and tends to indicate immune enhancement only in individuals with sub-optimal immune status (14, 30).

The skin constitutes the body's largest immunological organ, providing the first line of defense

Work in humans indicating that a single bout of short duration, high intensity exercise and prolonged duration, moderate intensity exercise decreases immunity, is largely based upon results of studies examining in vitro immune measures (26, 32). Little is known about the impact of a single bout of exercise on cutaneous measures of in vivo immunity in humans. One such study showed that after an acute bout of prolonged, continuous exercise (lasting ~6.5 h), DTH reactions to common recall antigens in the Mérieux CMI Multitest® were reduced but this test is no longer commercially available (6). Moreover, the use of common recall antigens does not permit the assessment of the effects of stress on the induction of new immune memory and findings may be confounded by the lack of control over immunological memory: both the sensitizing dose and time elapsed since sensitization influence immunological memory. To the best of our knowledge, no study has investigated the impact of the intensity and duration of continuous exercise stress on in vivo immunity in humans. Challenging the skin using novel antigens such as keyhole limpet hemocyanin (KLH) (35) or diphenylcyclopropenone (DPCP) (17) permits the investigation of the influence of stressors on in vivo immunity and allows rigorous control of both the dose and timing of sensitization. Using topical DPCP, we have recently shown that 2 h of moderate intensity exercise decreases both the induction of immunity (-53%) in those with no prior exposure to DPCP and elicitation of immunity (-19%) in those who received repeated monthly DPCP exposures to boost responses to a reproducible plateau (17). Possible mechanisms include the activation of the hypothalamic-pituitary-adrenal axis and sympatheticoadrenal-medullary axis, which is widely acknowledged to occur following prolonged stress (typically lasting hours) and in-turn increases glucocorticoids and catecholamines, previously shown to decrease the induction of CHS in mice (10, 33). It has yet to be determined whether the inhibitory effects of prolonged exercise on immune responses to DPCP are due to systemic effects on the dendritic cell/T cell axis between the skin and lymph nodes or whether they involve local effects on cutaneous inflammatory processes mediated principally via innate immune mechanisms. The levels of

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local cutaneous cytokines known to facilitate (e.g. IL-1β) and inhibit (e.g. IL-10) dendritic cell (DC) migration are considered to play a central role in the early DC-dependent events of CHS induction, namely, antigen processing and DC trafficking (38). One experimental approach to this problem is to investigate the effect of prolonged, moderate intensity exercise on cutaneous responses to a topically applied irritant such as croton oil (CO). Unlike DPCP, which ultimately stimulates an antigen-specific, T-cell-mediated immune response, CO is an irritant, which stimulates a non-T-cell mediated, inflammatory response after a single exposure (27). CO has no sensitizing properties but is capable of producing similar cutaneous erythema responses to those seen after CHS challenge (40). Here we present the findings from two studies, starting with the effects of intensity and duration of exercise stress on in vivo immune induction by DPCP. We hypothesized that a prolonged, moderate intensity exercise bout (120 minutes at 60% $\dot{V}O_{2peak}$) and a short, high intensity exercise bout (30 minutes at 80% $\dot{V}O_{2peak}$) would decrease the CHS responses to DPCP compared with a short, moderate intensity exercise bout (30 minutes at 60% \dot{V} O_{2peak}) and seated rest. Then, to examine whether exercise-related effects on local cutaneous inflammatory processes play a role in the inhibitory effect of prolonged, moderate intensity exercise on the CHS response we investigated irritant responses to a CO patch test.

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METHODS

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trials (17).

Subjects. All subjects were healthy, non-smoking, recreationally active males with no previous history of exposure to DPCP. Subjects were excluded if they were taking any medication or dietary supplements, had a history of atopy or any other immune-related or inflammatory dermatological conditions. Subjects were required to abstain from caffeine, alcohol, and exercise for 24 h before and 48 h after the experimental trials. All subjects gave written informed consent to participate after being fully briefed and informed of the study's procedures. The study received Local University Ethics Committee approval and was conducted in accordance with the Declaration of Helsinki principles. The effect of exercise intensity and duration on induction of DPCP immune memory. Subjects were matched for age and aerobic fitness (gas exchange threshold (GET) and \dot{V} O_{2peak}) before being randomly assigned to one of four experimental groups: 1) 120 minutes of seated rest (CON); 2) 30 minutes of moderate intensity (60% \dot{V} O_{2peak}) exercise (30MI); 3) 30 minutes of high intensity (80% \dot{V} O_{2peak}) exercise (30HI); or 4) 120 minutes moderate intensity (60% \dot{V} O_{2peak}) exercise (120MI) (Fig. 1). These exercise intensities and durations were chosen to allow comparison with the relevant literature (17), to assess the in vivo immune response to exercise recommended to healthy adults for fitness and health (e.g. the ACSM recommends 30 minutes, moderate-intensity exercise on most days), to best separate intensity and duration effects on in vivo immunity; and finally, with feasibility in mind (e.g. our subjects could complete 30 minutes at 80% \dot{V} O_{2peak}). There were no significant differences between groups for characteristics (Table 1). The study was performed between February, 2011 and April, 2012 and no data was taken from our previous investigation that also included 120MI and CON

*** Fig. 1 near here *** 96 97 *** Table 1 near here *** 98 99 **Preliminary measures and familiarization.** Anthropometric measures were recorded on 100 101 arrival at the laboratory. Body composition assessment was completed by whole body Dual Energy X-ray Absorptiometry (DEXA: Hologic QDR Series-4500, USA). Following this, V 102 103 O_{2peak} was estimated by means of a ramped exercise test on a treadmill (h/p/cosmos Mercury 4.0, Nussdorf-Traunstein, Germany). Following 3- minutes of walking at 5 km·h⁻¹ with an 104 incline of 1 %, speed increased at a rate of 1 km·h⁻¹·min⁻¹ to a maximum of 18 km·h⁻¹, after 105 which the incline increased at a rate of 1 %·min⁻¹ until volitional exhaustion. Pulmonary gas 106 exchange was measured breath-by-breath for the duration of the test (Cortex Metalyser 3B, 107 Biophysik, Leipzig, Germany). The \dot{V} O_{2peak} was taken as the highest 30-s average value before 108 the subject's volitional exhaustion and the speed equivalent to 60 % or 80 % of the \dot{V} O_{2peak} 109 110 was calculated. The GET was also determined from the ramped exercise test using the V-slope 111 method. 112 At least 24 h after the preliminary test, each subject's calculated exercise intensity was verified 113 by running for 50 % of their allocated exercise duration and all subjects were familiarized with 114 laboratory equipment. 115 116 117 **Experimental procedures.** Dietary intake was controlled during the 24 h before the main 118 experimental trial by providing subjects with their estimated daily energy requirement using

DEXA determined fat free mass as described (mean \pm SD: 11.2 ± 1.1 MJ day⁻¹) (9), multiplied

by a physical activity factor (37), and water proportional to 35 mL·kg⁻¹·d⁻¹ body mass.

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Within 3 weeks of the preliminary testing, on the day of the exercise trial, all subjects were transported to the laboratory at 0730 h and provided with a standard breakfast (0.03 MJ·kg⁻¹). Subjects were permitted to perform light activity before commencing the intervention. Nude body mass (NBM) was recorded before and after exercise on a digital platform scale to determine water allowance (Model 705; Seca, Hamburg, Germany). Exercising subjects received 5ml·kg⁻¹NBM of water immediately before and after the exercise, 2 ml·kg⁻¹NBM at 15 minutes intervals throughout, and any additional exercise fluid loss was replaced following exercise. Subjects assigned to the 120MI began running on a treadmill at 1100 h and those assigned to 30HI and 30MI began at 1230 h, so that all subjects completed the exercise at the same time of day (1300 h; Fig. 1). Immediately after the trial, exercising subjects showered and returned to the laboratory within 15 minutes of completion. The CON, non-stress condition, consisted of 2 h seated, passive rest in the same laboratory, in the same ambient conditions of 20 °C, at the same time of day, with a fluid intake proportional to 35 mL·kg⁻¹·d⁻¹ body mass.

Induction of contact sensitivity. Subjects were sensitized to DPCP at 1320 h, exactly 20 minutes after exercise cessation or equivalent seated rest, as described previously (17). This sensitization time was chosen to allow cutaneous blood flow to return to baseline (19). The sensitizing exposure to the novel antigen DPCP involved application of an occluded patch, constituting a 12 mm aluminum Finn chamber (Epitest Oy, Tuusula, Finland) on scanpor hypoallergenic tape containing an 11 mm filter paper disc. The paper disc was soaked in 22.8 μ l of 0.125 % DPCP in acetone (patch = 30 μ g·cm⁻² DPCP) and allowed to dry for 5 minutes before being applied to the skin on the lower back, for exactly 48h.

Elicitation. The magnitude of *in vivo* immune responsiveness was quantified by measuring the responses elicited by secondary exposure to the same antigen (Fig. 1). Twenty eight days after the initial sensitization to DPCP, all subjects received a challenge with a low concentration,

dose-series of DPCP on individual patches, each comprising an 8mm aluminium Finn chamber on scanpor hypoallergenic tape containing a 7 mm filter paper disc. Patches were applied to the inner aspect of the upper arm in the following concentrations: 10 μl of DPCP: 0.0048 %, 1.24 μg·cm⁻²; 0.0076 %, 1.98 μg·cm⁻²; 0.012 %, 3.17 μg·cm⁻²; 0.0195 %, 5.08 μg·cm⁻²; 0.0313 %, 8.12 μg·cm⁻² and 10 μl 100 % acetone control patch for background subtraction. Patches were applied in randomly allocated order at the local site in order to minimize any anatomical variability in responses. Elicitation patches were removed after 6 h and the strength of immune reactivity was assessed as cutaneous responses at 48 h post-application.

Blood collection and analysis. Blood samples (venepuncture from an antecubital vein) were collected into one K_3EDTA coated vacutainer, and one lithium heparin coated vacutainer (Becton Dickinson, Oxford, UK) pre, immediately post and 1 h post exercise. The samples were spun at 1500 g for 10 minutes in a refrigerated centrifuge. Plasma was aliquoted into Eppendorf tubes, and immediately frozen at -80°C for later analysis.

Plasma epinephrine and norepinephrine concentrations in K₃EDTA plasma were determined using a commercially available CatCombi ELISA (IBL International, Hamburg, Germany). Aliquots of lithium heparin plasma were used to determine cortisol concentration by ELISA, performed according to the manufacturer's instructions (DRG Instruments, Marburg, Germany). The intra-assay coefficient of variation for plasma epinephrine, norepinephrine and cortisol was 4.1 %, 4.1 % and 4.4 %, respectively.

The effect of prolonged, moderate intensity exercise on the cutaneous response to the irritant, croton oil. To investigate the possibility that the inhibitory effect of 120MI on CHS induction was mediated via local effects on cutaneous inflammatory processes, 11 healthy

males (age 24 ± 5 years; height 179 ± 8 cm; body mass 79.0 ± 9.9 kg; \dot{V} O_{2peak} 53 ± 6 ml.kg⁻¹.min⁻¹) completed a follow-up study to investigate the cutaneous responses to the non-specific irritant, CO.

In a randomized, counterbalanced, repeated measures design, subjects performed 120MI-CO or CON-CO separated by 7 - 14 d. Subjects received a CO challenge at 1320 h, exactly 20 minutes after exercise cessation or seated rest. This involved the topical application of a dose-series of CO on individual patches comprising 8mm aluminium Finn chambers mounted on hypoallergenic scanpor tape and 7 mm filter paper discs. Patches were applied in duplicate to the inner aspect of the upper arm in the following concentrations: $10~\mu l$ of CO in ethanol: 0.3~%, 0.55~%, 1.0~% and 3~% and $10~\mu l$ 100~% ethanol control patch (23). To account for local anatomical variability, the location of each concentration was randomized. Patches remained in place for exactly 24 h and the assessment of cutaneous responses was performed 2 h after removal of the CO patches, as described (23).

Assessment of cutaneous responses. Skin edema (inflammatory swelling) is considered the key measure of CHS elicitation responses (17). This was assessed as mean skin-fold thickness from triplicate measurements at each elicitation site using modified spring-loaded skin callipers (Harpenden Skin-fold Calliper, British Indicators, England), as described (17). Skin-fold thickness was recorded to the nearest 0.1 mm by the same investigator by placing the jaws of the calliper at the outer diameter of the response site and measuring skin thickness only (no subcutaneous fat).

Dermal thickness was determined at each patch site using a high-frequency ultrasound scanner (Episcan, Longport Inc., Reading, UK). The ultrasound probe was placed over the centre of each patch site together with ultrasound gel. The mean of three measurements was taken from each 12 mm scan image by an independent investigator, who was blinded to the trial assignment. Due to a delay in the availability of this equipment, dermal thickness was assessed in a subpopulation of 50 subjects who completed the DPCP patch test (CON = 13, 30MI = 14, 30HI = 12, 120MI = 11) and all subjects who completed the CO patch test.

Skin erythema is an objective measure of skin redness, which is considered the key measure of irritant responses (29). This was determined from triplicate measurement at each patch site using an erythema meter (ColorMeter DSM11, Cortex Technology, Hadsund, Denmark) as previously described (17).

Mean background values were determined from triplicate measurements at the vehicle only patch site for thickness and redness. In order to determine the increase in thickness and redness in response to DPCP or CO, the value from the vehicle-only site was subtracted from each patch site value. The values for increase in skin-fold thickness, dermal thickness and erythema over all the doses were summed, which gave an approximation of the area under the dose-response curve, representative of the overall reactivity of each subject to DPCP or CO, respectively.

Statistical analysis. Data in the results are presented as mean \pm SD, unless otherwise stated and statistical significance was accepted at P < 0.05. Data were checked for normality and sphericity. Greenhouse-Geisser adjustments to the degrees of freedom were applied where necessary (skin-fold dose-series response to DPCP, epinephrine, norepinephrine and cortisol).

All statistical analysis was conducted using SPSS software. The mean difference with 95 % confidence intervals is presented for the main outcome measures.

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Sample size was estimated using data from a previous study examining the effect of prior exercise stress on CHS responses to DPCP (17). The alpha (Type I error rate) was set at 0.05, and power at 0.95 (1 - Type II error rate) (G*Power software, version 3.1.2). For the CO element, a minimum important difference using biological variation data of the summed CO erythema response was used to estimate an effect size (0.91). A one-way ANOVA was used to assess differences between the groups in physical characteristics. The effect of exercise intensity and duration was analyzed using a one-way ANOVA to determine differences in the summed increase in responses to DPCP between the CON, 30MI, 30HI and 120MI trials. A two-way, mixed model ANOVA (DPCP data) or a repeated measures ANOVA (CO data) was used to analyze the skin-fold and dermal thickness responses across the full dose-series challenge (trial × dose). A two-way mixed model ANOVA (trial × time) was used to compare the circulating stress hormone data. Significant differences were identified using post hoc Tukeys HSD or Bonferroni corrected t-tests, where appropriate. To further investigate the differences between CON and 120MI, independent t-tests (DPCP data) or paired t-tests (CO data) were used to assess summed increases. Logarithmic transformation was performed on the DPCP data to allow for the calculation of the x-intercept when y = 0, utilizing linear regression on the linear portion of the dose response curve. A threshold dose for a response to DPCP was then calculated by back transformation (anti-log). Simple linear regression and a calculation of the standard error of the estimate (SEE) were performed to assess the validity of skin-fold measurement, using skin-fold callipers, as a practical method to determine dermal thickening compared with the objective criterion, high-frequency ultrasound. This was performed on the

sum of the 5 elicitation sites for a sub-population with complete data sets at the 48 h time point

245 (n=50).

RESULTS

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The effect of exercise intensity and duration on induction of DPCP immune memory. 248 **Assessment of CHS responses.** The skin-fold response, summed from five challenge doses, 249 250 was significantly different between groups (F(3,60) = 3.6, P < 0.05). Tukeys post hoc analysis 251 revealed that skin-fold thickness was reduced 67% by 120MI compared with CON (P < 0.05; 252 Fig. 2a). The mean difference between 120MI and CON was 3.17 mm (95% confidence 253 intervals 0.31 to 6.03 mm). There was no significant difference between the short duration 254 30MI or 30HI exercise groups compared with CON. The full, dose-series response to DPCP for each group was also determined for the increase in skin-fold thickness (Fig. 2b). The skin-255 256 fold thickness responses from the five individual doses revealed a significant trial × dose interaction (F(7.3,145.1) = 3.0, P < 0.01). Post hoc analysis revealed that skin-fold thickness 257 was significantly lower in 120MI compared with CON at the 1.98 μ g·cm⁻² dose (P < 0.05). 258 5.08 μ g·cm⁻² and 8.12 doses (P < 0.01) and approached significance at the 3.17 μ g·cm⁻² dose 259 (P = 0.058). To further investigate the differences between CON and 120MI, the threshold 260 261 dose for a positive response to DPCP was calculated using the linear part of the dose response curve, as 0.48 and 2.09 µg·cm⁻² for the CON and 120MI groups, respectively. This suggests 262 that to elicit a positive response, 120MI required a 4.4 times greater DPCP dose in 263 comparison with CON. Skin-fold thickness assessed using skin-fold callipers was strongly 264 related with high-frequency ultrasound readings of dermal thickness (r = 0.93, $r^2 = 0.86$, SEE 265 = 1.3 mm; P < 0.01). 266 267 *** Fig. 2 near here*** 268

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Circulating stress hormones. At baseline, pre-exercise, there were no significant differences between groups for circulating epinephrine, norepinephrine or cortisol concentration. A significant trial \times time interaction was observed for circulating epinephrine (F(4.6,88.5) = 7.0,

P < 0.01; Fig. 3a), norepinephrine (F(3.4,67.1) = 24.0, P < 0.01; Fig. 3b) and cortisol concentration (F(4.6,90.6) = 7.0, P < 0.01; Fig. 3c). The raised circulating epinephrine and norepinephrine concentration observed immediately post on both 120MI and 30HI (P < 0.01) had returned to pre-exercise levels by 1 h post exercise. Circulating epinephrine concentration was greater at post on 120MI compared with CON (P < 0.01) and circulating norepinephrine concentration was greater at post on 30HI compared with CON (P < 0.01). Circulating cortisol concentration was greater at post and 1 h post on 120MI and at post on 30HI compared with CON (P < 0.01). The typical diurnal response in circulating cortisol concentration is shown, whereby levels were lower at post (1300) and 1 h post (1400) compared with pre-exercise (1100) on both 30MI and CON (P < 0.01).

*** Fig. 3 near here ***

memory and cutaneous responses to the irritant, croton oil. The aim here was to examine whether the inhibitory effect of 120MI on CHS is due to local effects on cutaneous inflammatory processes mediated principally via innate immune mechanisms. To this end, Fig. 4 shows the summed responses to all challenge doses for induction of DPCP immune memory (5 doses) and irritant responses to CO (4 doses). Results are presented as dermal thickness, considered a key measure of CHS responses (17), and erythema, considered a key measure of irritant responses (29). Here we show that 120MI significantly decreased DPCP responses measured as dermal thickness (t(22) = 3.5, P < 0.01; Fig. 4b) and erythema (t(30) = 2.1, P < 0.05; Fig. 4a). The mean difference for dermal thickness was 3.17 mm (95% confidence intervals 1.27 to 5.07) and for erythema was 18.61

The effect of prolonged, moderate intensity exercise on the induction of DPCP immune

AU (95% confidence intervals 0.41 to 36.82). No effect of 120MI-CO on irritant responses

measured as erythema (t(10) = 0.2, P = 0.826; Fig. 4c) or dermal thickness (t(10) = 1.2, P = 0.826) 0.253; Fig. 4d) points to an inhibitory effect of 120MI on cell-mediated processes rather than local inflammatory processes in the decrease in CHS. It is noteworthy that the erythematous response to the top challenge dose of CO was comparable to the erythematous response to the top dose of DPCP (mean \pm SD: 11.75 \pm 5.28 AU and 11.25 \pm 4.84 AU, respectively). As would be expected, dermal thickening response to the dose-series of the irritant, CO was small compared with DPCP (Fig. 4 d). For visual comparison, the increase in erythema responses to the full, dose-series of CO is also presented (Fig. 5). There was no significant trial × dose interaction observed between 120MI-CO and CON-CO for erythema responses (F(3,30) = 1.4,P = 0.267). ***Fig. 4 near here *** ***Fig. 5 near here ***

DISCUSSION

The advantages of, and the need for further research utilizing, *in vivo* immune measures in humans have recently been highlighted (1, 39). The primary aim of this work was to determine the unknown effects of the intensity and duration of continuous exercise stress on the induction of *in vivo* immunity in humans. In line with our hypothesis, prolonged, moderate intensity exercise (120MI) decreased the induction of *in vivo* immunity; however, short lasting moderate intensity (30MI) or high intensity (30HI) exercise did not influence this response despite elevated circulating catecholamines on 30HI and greater circulating cortisol on 30HI compared with CON. We then demonstrated that prolonged exercise had no effect on cutaneous responses to the irritant, CO. These findings support the notion that the observed decrease in *in vivo* immune induction to DPCP represents an effect on T-cell mediated immune responses rather than exercise-effects on local expression of inflammatory effector processes.

This is the first study to compare the effects of intensity and duration of continuous exercise stress on *in vivo* immunity assessed by use of an experimental CHS model in humans. In keeping with our previous findings, we observed that 120MI had a significant inhibitory effect on the induction of new immunity via the skin (17). Our finding that 30MI had no effect on *in vivo* immune induction is at odds with one hypothesis underpinning the J-shaped model (25), whereby a moderate dose of exercise is proposed to be immune-enhancing, but in accordance with recent research showing no effect of a moderate dose of exercise on the response to vaccination in young, healthy adults (20). While other studies have shown that a moderate dose of exercise can enhance antibody responses to vaccination, thereby supporting one hypothesis underpinning the J-shaped model, this typically occurs in individuals with sub-optimal immune status or when a half dose of vaccine is administered (14, 30). We also acknowledge that

exercise might differentially affect CHS, a cutaneous T-cell mediated response, and the antibody response to vaccination, a systemic B-cell mediated response.

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We hypothesized that 30HI would decrease *in vivo* immune induction to DPCP, based upon evidence from *in vitro* work showing that short lasting high intensity exercise decreases indicators of both lymphocyte and neutrophil function (26, 32). However, our results do not support this despite elevated circulating catecholamines on 30HI and greater circulating cortisol on 30HI compared with CON. These findings provide little support for an involvement of circulating stress hormones in the mechanisms associated with altered in vivo immune responses to DPCP at the skin. For example, circulating norepinephrine was highest after 30HI when there was no immunosuppression suggesting that circulating norepinephrine has little immunosuppressive effect on the CHS system. Although circulating cortisol tended to be higher on 120MI compared with 30HI this did not reach statistical significance. In addition, circulating cortisol exceeded the purported binding capacity (~552 nmol·L⁻¹)(22) at postexercise before DPCP application in a similar proportion of subjects on 30HI (11 of 16) and 120MI (12 of 16) yet 30HI did not decrease immune induction by DPCP. There is clear evidence from murine models that high doses of these stress hormones can have significant immune-modulating effects. Intradermal injections of high dose corticosterone or catecholamines, both locally or distant from the sensitization site, inhibit the antigen-presenting capability of cutaneous DCs, reduce the number of T cells in draining lymph nodes and ultimately suppress DTH and CHS responses (11, 15, 33). Results from human studies are less consistent with some authors reporting a lack of association between stress hormones and in vivo immune responses (3, 13, 28). One frequently proposed explanation is that human studies typically rely on individual snapshot assessments of circulating stress hormones, thus missing important information regarding the kinetics of these responses. In this regard, we

acknowledge a limitation of the current study is that we applied the DPCP sensitization patch 20 minutes after exercise at a time when circulating cortisol likely reached a peak but circulating catecholamines would likely have returned to pre-exercise levels. At the outset, we considered the strengths and weaknesses of DPCP sensitization at the cessation of exercise to coincide with the peak in circulating catecholamines. After careful consideration, we chose to delay sensitization until 20 minutes after exercise to avoid possible confounding due to raised skin blood flow and sweating. One might also argue that another limitation is that we only took blood samples to characterize circulating cortisol at immediately post and 1 h post exercise yet the DPCP sensitizing patch remained in place for 48 h. Work in young adults showed the inhibitory effect of stress on the development of immune memory is particularly evident when stress is experienced at, or close to, the time of sensitization: this supports our choice of sample timing to characterize the circulating cortisol response in close proximity to the exercise stress (35).

The findings from the current study show that 120MI had no impact on cutaneous inflammatory responses to CO. This suggests that the inhibitory effect of 120MI on CHS induction with DPCP is likely associated with cell-mediated events rather than exercise effects on local inflammatory processes. Further research is required to better understand the mechanisms associated with the inhibitory effect of 120MI on *in vivo* responses to DPCP. Research should target the interactions between DC's and T cells in terms of antigen processing and presentation and activation of T cells and the subsequent balance between effector and regulatory T cells considered central to the successful induction of CHS (38). Also, the duration of the inhibitory effect of prolonged, heavy exercise on CHS induction in humans remains unknown and could be determined in a study that manipulates the timing of DPCP sensitization after 120MI. Given the reported sex differences in immune responses to

exercise (16), we recognize the limitation of using only males in this study and encourage the investigation of *in vivo* immune responses to exercise using this CHS model in females.

Experimental-CHS provides an attractive measure of *in vivo* immunity, not only because the skin is immediately accessible but because it overcomes many of the limitations of commonly used *in vitro* measures which are lacking in terms of clinical significance and practicality. We recognize that there are limitations with using DPCP in the CHS model described. Given that DPCP is benign, determining the clinical significance of the response, with specific regard to infection (skin and other) is an important avenue for future research. Preferably, the strength of the cutaneous recall response to DPCP could be generalized beyond skin immunity to indicate the immune system's general ability to respond to an infectious challenge. The available evidence in this regard is supportive as cutaneous immune measures are impaired in individuals with acute infectious illness (5), diabetes and psoriasis (4), and predict mortality in critically ill HIV-infected patients (12). An alternative viewpoint is that the benign characteristic of DPCP actually overcomes the ethical constraints associated with using live pathogens, such as rhinovirus to assess *in vivo* immunity. We also recognize the limitation that experimental-CHS requires purposefully inducing CHS; nevertheless, the selected doses we use are low and the mild elicitation responses are temporary.

Experimental-CHS with DPCP is practical, safe, and can be administered without the need for expensive equipment, invasive injections or blood sampling, making it a suitable immunological tool for both laboratory and field investigations. Moreover, the use of a novel antigen such as DPCP provides investigators with rigorous control over the timing and dose of sensitizing exposure, enabling the effects of various stressors on the primary immune response to be studied. The measurement of DTH responses to KLH is an alternative per-cutaneous *in*

vivo method, also reported to represent a primary immune response (36). However, since KLH is derived from a shellfish this may explain why some individuals exhibit significant responses to KLH prior to immunization (34). Experimental-CHS with DPCP is not restricted to examining the effects of stress on the induction phase. Recently we have shown that this approach can be used to assess the effect of exercise stress on the elicitation phase in subjects who, following repeated monthly DPCP skin challenges, achieved a reproducible plateau in responses (17). Furthermore, the standardized CHS model we describe overcomes some of the limitations of vaccine models of *in vivo* immunity including variable immunogenicity (e.g. hepatitis B (18)), annual changes in vaccine (e.g. influenza (7)) and difficulty when comparing the circulating antibody results from different studies using in-house ELISAs. Nevertheless, a standard protocol for measuring CHS elicitation responses in humans has yet to be established. The use of erythema to quantify CHS elicitation has been questioned, particularly at sites of stronger responses, where yellow vesicles can interfere with the erythema (redness) readings (17, 24). Erythema is typically the preferred measure of irritant responses which, as we show (Fig. 4 d), induce less edema than CHS responses (29). Notwithstanding the degree of subjectivity, a particular strength of the current findings is that skin-fold thickness was strongly related with dermal thickness measured by a high-frequency ultrasound scanner and read by a blinded investigator (r = 0.93). Hence, we agree with the recommendation of others that, skin-fold callipers present a simple and cost-effective measure of CHS edema (24). In conclusion, using experimental CHS with DPCP, these results demonstrate that prolonged, moderate intensity exercise, but not short-lasting high or short-lasting moderate intensity exercise, decreases the induction of in vivo immunity in healthy humans. No effect of prolonged, moderate intensity exercise on the skin's response to the irritant, CO points towards

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a suppression of cell-mediated immunity in the observed decrease in CHS response. The

topical application of DPCP provides an attractive tool to assess the effect of exercise stress on in vivo immunity in humans. Acknowledgements Bethany Diment's PhD was supported by a 125th anniversary research scholarship from Bangor University. This study received no external funding. None of the authors had a conflict of interest. The results of the present study do not constitute endorsement by the American College of Sports Medicine.

| 455 | | References |
|--------------------------|-----|---|
| 456 | | |
| 457 458 459 | 1. | Akbar AN, Reed JR, Lacy KE, Jackson SE, Vukmanovic-Stejic M, Rustin MH. Investigation of the cutaneous response to recall antigen in humans in vivo. <i>Clin Exp Immunol.</i> 2013; 173:163-172. |
| 460 461 462 | 2. | Albers R, Bourdet-Sicard R, Braun D et al. Monitoring immune modulation by nutrition in the general population: identifying and substantiating effects on human health. <i>Br J Nutr.</i> 2013; 110 Suppl. 2:S1-30. |
| 463 464 | 3. | Altemus M, Dhabhar FS, Yang R. Immune function in PTSD. <i>Ann N Y Acad Sci.</i> 2006; 1071:167-183. |
| 465 466 | 4. | Bangsgaard N, Engkilde K, Menne T et al. Impaired hapten sensitization in patients with autoimmune disease. <i>Clin Exp Immunol</i> . 2011; 165:310-317. |
| 467 468 469 | 5. | Bennett BK, Hickie IB, Vollmer-Conna US et al. The relationship between fatigue, psychological and immunological variables in acute infectious illness. <i>Aust N Z J Psychiatry</i> . 1998; 32:180-186. |
| 470 471 472 | 6. | Bruunsgaard H, Hartkopp A, Mohr T et al. In vivo cell-mediated immunity and vaccination response following prolonged, intense exercise. <i>Med Sci Sports Exerc</i> . 1997; 29:1176-1181. |
| 473 474 | 7. | Burns VE. Using vaccinations to assess in vivo immune function in psychoneuroimmunology. <i>Methods Mol Biol.</i> 2012; 934:371-381. |
| 475 476 | 8. | Clark RA, Ghosh K, Tonnesen MG. Tissue engineering for cutaneous wounds. <i>J Invest Dermatol</i> . 2007; 127:1018-1029. |
| 477 478 479 | 9. | Cunningham JJ. Body composition as a determinant of energy expenditure: a synthetic review and a proposed general prediction equation. <i>Am J Clin Nutr.</i> 1991; 54:963-969. |
| 480 481 | 10. | Dhabhar FS. Psychological stress and immunoprotection versus immunopathology in the skin. <i>Clin Dermatol.</i> 2013; 31:18-30. |
| 482 483 | 11. | Dhabhar FS, McEwen BS. Enhancing versus suppressive effects of stress hormones on skin immune function. <i>Proc Natl Acad Sci U S A</i> . 1999; 96:1059-1064. |
| 484 485 486 487 | 12. | Dolan MJ, Clerici M, Blatt SP et al. In vitro T cell function, delayed-type hypersensitivity skin testing, and CD4+ T cell subset phenotyping independently predict survival time in patients infected with human immunodeficiency virus. <i>J Infect Dis.</i> 1995; 172:79-87. |
| 488 489 490 | 13. | Edwards KM, Burns VE, Reynolds T, Carroll D, Drayson M, Ring C. Acute stress exposure prior to influenza vaccination enhances antibody response in women. <i>Brain Behav Immun.</i> 2006; 20:159-168. |

- 491 14. Edwards KM, Pung MA, Tomfohr LM et al. Acute exercise enhancement of 492 pneumococcal vaccination response: a randomised controlled trial of weaker and 493 stronger immune response. *Vaccine*. 2012; 30:6389-6395.
- 494 15. Flint MS, Valosen JM, Johnson EA, Miller DB, Tinkle SS. Restraint stress applied prior 495 to chemical sensitization modulates the development of allergic contact dermatitis 496 differently than restraint prior to challenge. *J Neuroimmunol.* 2001; 113:72-80.
- 497 16. Gillum TL, Kuennen MR, Schneider S, Moseley P. A review of sex differences in 498 immune function after aerobic exercise. *Exerc Immunol Rev.* 2011; 17:104-121.
- Harper Smith AD, Coakley SL, Ward MD, Macfarlane AW, Friedmann PS, Walsh NP.
 Exercise-induced stress inhibits both the induction and elicitation phases of in vivo T-cell-mediated immune responses in humans. *Brain Behav Immun*. 2011;
 25:1136-1142.
- 503 18. Hernandez-Bernal F, Aguilar-Betancourt A, Aljovin V et al. Comparison of four recombinant hepatitis B vaccines applied on an accelerated schedule in healthy adults. *Hum Vaccin*. 2011; 7:1026-1036.
- 506 19. Kenny GP, Webb P, Ducharme MB, Reardon FD, Jay O. Calorimetric measurement of 507 postexercise net heat loss and residual body heat storage. *Med Sci Sports Exerc*. 508 2008; 40:1629-1636.
- 509 20. Long JE, Ring C, Drayson M et al. Vaccination response following aerobic exercise: can 510 a brisk walk enhance antibody response to pneumococcal and influenza 511 vaccinations? *Brain Behav Immun.* 2012; 26:680-687.
- 512 21. Martin SA, Pence BD, Woods JA. Exercise and respiratory tract viral infections. *Exerc Sport Sci Rev.* 2009; 37:157-164.
- 514 22. Mccarthy DA, Dale MM. The Leukocytosis of Exercise A Review and Model. *Sports* 515 *Med.* 1988; 6:333-363.
- 516 23. Memon AA, Friedmann PS. 'Angry back syndrome': a non-reproducible phenomenon. *Br J Dermatol.* 1996; 135:924-930.
- 518 24. Narbutt J, Lesiak A, Skibinska M et al. Suppression of contact hypersensitivity after repeated exposures of humans to low doses of solar simulated radiation.

 520 *Photochem Photobiol Sci.* 2005; 4:517-522.
- 521 25. Nieman DC. Exercise, infection, and immunity. *Int J Sports Med.* 1994; 15 Suppl 3:S131-S141.
- 523 26. Nieman DC, Miller AR, Henson DA et al. Effect of high- versus moderate-intensity 524 exercise on lymphocyte subpopulations and proliferative response. *Int J Sports* 525 *Med.* 1994; 15:199-206.
- 526 27. Nosbaum A, Vocanson M, Rozieres A, Hennino A, Nicolas JF. Allergic and irritant contact dermatitis. *Eur J Dermatol.* 2009; 19:325-332.

- 528 28. Oliver SJ, Macdonald JH, Harper Smith AD et al. High altitude impairs in vivo immunity in humans. *High Alt Med Biol.* 2013; 14:144-149.
- 29. Parslew R, Friedmann PS. The irritancy of anthralin is inhibited by repeat applications of a subirritant concentration. *Br J Dermatol*. 1999; 141:469-474.
- 532 30. Pascoe AR, Fiatarone Singh MA, Edwards KM. The effects of exercise on vaccination responses: A review of chronic and acute exercise interventions in humans. *Brain Behav Immun.* 2014; 39:33-41.
- 535 31. Poenaru D, Christou NV. Clinical outcome of seriously ill surgical patients with intraabdominal infection depends on both physiologic (APACHE II score) and immunologic (DTH score) alterations. *Ann Surg.* 1991; 213:130-136.
- Robson PJ, Blannin AK, Walsh NP, Castell LM, Gleeson M. Effects of exercise
 intensity, duration and recovery on in vitro neutrophil function in male athletes.
 Int J Sports Med. 1999; 20:128-135.
- 541 33. Seiffert K, Hosoi J, Torii H et al. Catecholamines inhibit the antigen-presenting capability of epidermal Langerhans cells. *J Immunol.* 2002; 168:6128-6135.
- 543 34. Smith A, Vollmer-Conna U, Bennett B, Wakefield D, Hickie I, Lloyd A. The relationship 544 between distress and the development of a primary immune response to a novel 545 antigen. *Brain Behav Immun*. 2004; 18:65-75.
- 546 35. Smith AJ, Vollmer-Conna U, Bennett B, Hickie IB, Lloyd AR. Influences of distress and alcohol consumption on the development of a delayed-type hypersensitivity skin test response. *Psychosom Med.* 2004; 66:614-619.
- 549 36. Smith TP, Kennedy SL, Fleshner M. Influence of age and physical activity on the
 primary in vivo antibody and T cell-mediated responses in men. *J Appl Physiol*.
 2004; 97:491-498.
- Todorovic VE, Micklewright A. The Parenteral and Enteral Nutrition Group of the
 British Dietetics Association: A Pocket Guide to Clinical Nutrition. 1st ed.
 Rochester: British Dietetics Association; 2004. 1-12 p.
- 555 38. Toebak MJ, Gibbs S, Bruynzeel DP, Scheper RJ, Rustemeyer T. Dendritic cells: biology of the skin. *Contact Dermatitis*. 2009; 60:2-20.
- 557 39. Walsh NP, Gleeson M, Shephard RJ et al. Position statement. Part one: Immune function and exercise. *Exerc Immunol Rev.* 2011; 17:6-63.
- 559 40. Willis CM, Stephens CJ, Wilkinson JD. Differential patterns of epidermal leukocyte 560 infiltration in patch test reactions to structurally unrelated chemical irritants. *J* 561 *Invest Dermatol.* 1993; 101:364-370.

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Figure Legends

FIGURE 1. Schematic for the effect of exercise intensity and duration on induction of DPCP immune memory. Samples; venepuncture blood.

FIGURE 2. Effect of exercise stress prior to induction of contact hypersensitivity with DPCP on responses to elicitation challenge 28 d later. Shown here as (a) summed increase in skinfold thickness (callipers: mean \pm SD) and (b) responses to the full dose-series challenge with DPCPn-fold thickness (callipers: mean \pm SEM) # P < 0.05 and ## P < 0.01 vs CON.

FIGURE 3. Circulating epinephrine (a), norepinephrine (b) and cortisol (c) response to exercise or seated rest. \downarrow = induction of contact sensitivity by DPCP application. ** $P < 0.01 \ vs$ pre-exercise; ## $P < 0.01 \ vs$ CON. Data are mean \pm SEM.

FIGURE 4. Effect of prolonged exercise stress (120MI) or seated rest (CON) prior to induction of contact sensitivity with DPCP or irritant challenge with CO. Shown here are the summed responses to: DPCP elicitation challenge 28 d later, measured as (a) erythema and (b) dermal thickness (ultrasound); and CO challenge applied 20 minutes after exercise or equivalent seated rest, measured as (c) erythema and (d) dermal thickness (ultrasound). # P < 0.05 and ## P < 0.01 vs CON. Data are mean \pm SD.

FIGURE 5. Effect of prolonged exercise stress (120MI-CO) or seated rest (CON-CO) on erythema responses to irritant challenge with CO. Shown here are the responses to the full

dose-series challenge with CO applied 20 minutes after exercise or equivalent seated rest. Data are mean \pm SEM.

TABLE 1. Subject information. Values are mean \pm SD.

| | CON | 30MI | 30HI | 120MI |
|---|-----------------|-----------------|-----------------|-----------------|
| N | 16 | 16 | 16 | 16 |
| Age (years) | 23 ± 4 | 20 ± 2 | 22 ± 4 | 22 ± 4 |
| Height (cm) | 180 ± 7 | 180 ± 5 | 179 ± 7 | 180 ± 7 |
| Body mass (kg) | 77.3 ± 11.3 | 74.5 ± 10.1 | 76.3 ± 12.8 | 78.8 ± 12.1 |
| Body fat (%) | 15.2 ± 3.7 | 15.1 ± 4.5 | 15.0 ± 4.7 | 15.9 ± 4.3 |
| $VO_{2peak}\ (ml{\cdot}kg^{\text{-}1}{\cdot}min^{\text{-}1})$ | 57 ± 7 | 58 ± 5 | 58 ± 6 | 56 ± 5 |
| GET (L·min⁻¹) Weekly | 3.04 ± 0.31 | 3.09 ± 0.59 | 3.08 ± 0.60 | 3.11 ± 0.51 |
| exercise (h) | 6 ± 4 | 6 ± 2 | 5 ± 2 | 6 ± 3 |

GET, gas exchange threshold

















