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1	Intensive	exercise	does not	preferen	tially m	obilise	skin-ho	ming T	cells a	nd NK	cells.
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2 **Running head:** CLA+ lymphocytes and exercise

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#### 22 Abstract

23

lymphocyte antigen (CLA) - a homing molecule for endothelial cell leukocyte adhesion molecule 1 (ELAM-24 25 1), which enables transmigration to the skin – are selectively mobilised in response to acute exercise. *Methods*: Nine healthy males (mean  $\pm$  SD age: 22.1  $\pm$  3.4 years) completed two exercise sessions: high-26 intensity continuous cycling ('continuous exercise' at 80%  $\dot{V}O_{2_{MAX}}$  for 20 min) and low-volume high-27 intensity interval exercise ('HIIE' at 90%  $\dot{V}O_{2_{MAX}} 10 \times 1$  min repetitions with 1 min recovery intervals). 28 29 Blood was collected before, immediately- and 30 min post-exercise for cryo-preservation of peripheral blood mononuclear cells. CLA+ and CLA- cells were quantified within NK subpopulations (CD56<sup>bright</sup> 'regulatory' 30 and CD56<sup>dim</sup> 'cytotoxic' cells) as well as the following CD8+ T cell subpopulations: naive ('NA'; 31 CD45RA+CCR7+), central memory ('CM'; CD45RA-CCR7+), effector-memory ('EM'; CD45RA-CCR7-) 32 33 and CD45RA-expressing effector-memory cells ('EMRA'; CD45RA+CCR7-). Results: CLA+ NK cells and CD8+ memory T cells increased in response to both exercise bouts, but, overall, their numerical contribution 34 to the exercise lymphocytosis was inferior to CLA- cells, which increased to a much greater extent during 35 exercise. Tellingly, the most exercise-responsive cells – effector memory CD8+ cells and CD56<sup>dim</sup> cells – 36 were CLA-. Conclusions: A small subset of CLA+ lymphocytes are mobilised into blood during acute 37 intensive exercise, but CLA+ cells are not major contributors to exercise lymphocytosis, thus providing 38 preliminary evidence that the skin is not a major origin, or homing-destination, of exercise-sensitive 39 40 lymphocytes.

Purpose: This study investigated whether natural killer (NK) cells and CD8+ T cells expressing cutaneous

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42 Key words: T cells, inflammation, skin, homing

#### 44 INTRODUCTION

Memory CD8+ T cells and natural killer (NK) cells are the largest contributors to the acute and 45 transient lymphocytosis that occurs in peripheral blood during intensive exercise (6, 20, 36). Largely driven 46 47 by their relative high density cell surface expression of  $\beta$ 2-adrenergic receptors that induce detachment from 48 the vascular endothelium upon stimulation by catecholamines, the mobilisation of NK cell and CD8+ T cell subsets is considered to be an evolved mechanism that facilitates effector cell recruitment to sites of potential 49 50 or ongoing injury (10). Built on findings from animal models and using an analogy of immune cells as 51 'soldiers', it is proposed that naïve and central memory CD8+ T cells – which are slightly increased during 52 exercise – traffic from the boulevards (i.e., 'bloodstream') back to the 'barracks' (i.e., the spleen, lymph nodes). On the other hand, tissue-specific memory CD8+ T cells – which mobilise to a larger extent during 53 exercise - home to specific peripheral tissues (e.g., lungs, gut) to conduct immune-surveillance against 54 potential pathogenic challenge. Finally, it is hypothesised that effector CD8+ T cells – which are mobilised to 55 the greatest extent during exercise – are redeployed to 'battlefield' sites of wound healing (e.g. in the skin) 56 following acute exercise (9, 10). However, whether these highly cytotoxic effector NK and CD8+ T cells, 57 58 which are selectively increased in an acute and transient manner by acute exercise, have the propensity to 59 migrate to peripheral cutaneous sites remains unknown.

60 In support of this model, it has been demonstrated in murine models that exercise redeploys large 61 numbers of T cells to the Peyer's patches, lungs and bone marrow, reinforcing the idea that T cells are 62 mobilised to sites of potential antigen encounter (i.e., lungs, gut), as well as to serve other important functions 63 such as the provision of additional stimuli for haematopoiesis in the bone marrow (20). However, the 64 aforementioned study was unable to measure homing to cutaneous sites or areas of ongoing inflammation. It 65 has been shown that T cells and NK cells do show increased infiltration into sites of experimental 66 inflammation (i.e., subcutaneous implantation of a surgical sponge treated with the lymphocyte-specific chemokine lymphotactin) in mice 24 to 48 hours after acute psychological stress (40). However, it remains 67 unclear whether this increased cell infiltration is driven by stress-induced effector cell redistribution, or other 68 69 mechanisms occurring in the days after the acute stressor.

70 An approach that is commonly used to investigate lymphocyte homing propensity in humans is the 71 assessment of cell surface adhesion molecule expression on cells; this approach can be used to reveal the 72 probable trafficking patterns of cells mobilised into the bloodstream during exercise. For example, studies 73 have shown that there is a selective influx of CD8+ memory cells into the bloodstream that exhibit lower 74 levels of lymphoid homing markers such as CD62L and CCR7 (6), thus providing evidence that exercise 75 mobilises CD8+ T cells with a homing capacity for peripheral tissues. Further research has shown that these 76 cells mobilised by exercise express adhesion molecules such as CD11a (15, 22), CD11b (17, 18), VLA-4 77 (very late antigen-4) and LPAM-1 (lymphocyte Peyer's patch adhesion molecule-1) (15), which enable 78 migration to peripheral sites including the bone marrow (23), Peyer's patches (43) and lungs (38). However, 79 the aforementioned adhesion molecules cannot be used exclusively for identifying skin-homing potential and it remains uncertain whether exercise-responsive CD8+ T cells and NK cells have a skin-homing phenotype. 80

81 Cutaneous lymphocyte antigen (CLA) expression on lymphocytes can be used to determine whether 82 lymphocytes preferentially mobilised during intensive exercise exhibit a homing phenotype for cutaneous sites. CLA is a specialised form of P-selectin glycoprotein ligand-1 (PSGL-1; CD162), a surface glycoprotein 83 84 expressed constitutively on all human peripheral-blood T cells. After post-translational modification, PSGL-1 bears a Sialyl-Lewis<sup>X</sup> (sLe<sup>X</sup>) moiety, termed CLA, which avidly binds CD62E (E-selectin), an adhesion 85 molecule which initiates the transmigration cascade to the skin, and which is also upregulated during 86 cutaneous inflammation (4, 27, 28, 32). Thus, most T cells in both normal and diseased cutaneous sites are 87 CLA+ (7, 29, 30). In support, leukocyte infiltration to inflammatory sites can be largely inhibited by a CLA 88 89 modifier (11). Taken together, analyses of CLA+ cell mobilisation can be used as proxy marker to reveal 90 whether cells mobilised by exercise have the phenotypic capacity to migrate to 'battlefield' sites in the skin 91 (10).

The primary objective of this study was to investigate whether CD8+ T cells preferentially mobilised during intensive exercise stress exhibit a homing phenotype for cutaneous sites. To fulfil this objective, we investigated the number of CLA-positive cells among CD8+ T cell subsets, to establish the contribution of CLA-positive cells to the stepwise CD8+ lymphocytosis pattern previously observed in response to exercise (6). We also extended the analyses of CLA+ and CLA- cells to NK cell populations – the largest responders

97	to exercise stress (6). The second objective of this study was to compare the magnitude of CD8+ T cell and
98	NK cell mobilisation in response to continuous high-intensity exercise (continuous exercise) and high-
99	intensity intermittent exercise (HIIE). HIIE is often referred to as High Intensity Interval Training (HIIT)
100	when repeated frequently over several weeks or months. This form of exercise has received considerable
101	attention as a short-duration and low-volume means of achieving similar health benefits to continuous
102	exercise. Given the increasing health benefits that have been observed with HIIE, and although we have
103	established in a prior report that CD8+ T cell and NK cell subset mobilisation is intensity dependent (6), the
104	effects of HIIE on changes to these cell subsets remains unknown.

#### 105 METHODS

#### 106 *Participants*

107 Ten healthy males were recruited to take part in this study as previously described (41, 42). Peripheral 108 blood mononuclear cells (PBMCs) were available from nine of the ten participants, and were isolated from blood samples collected before, immediately after, and 30 min after two different forms of exercise, described 109 below. All nine participants (age:  $22.1 \pm 3.4$  years; height:  $180.5 \pm 6.1$  cm; weight:  $78.1 \pm 11.0$  kg; body mass 110 index: 24.0  $\pm$  3.1 kg.m<sup>2</sup>;  $\dot{VO}_{2 \text{ MAX}}$ : 43.8  $\pm$  4.1 ml.kg.min<sup>-1</sup>) included in this study were non-smokers, and 111 refrained from taking vitamin supplements and anti-inflammatory medication for fourteen days, and did not 112 exercise and consume alcohol or caffeine for two days prior to experimental trials. All participants provided 113 114 written informed consent and the study was approved by the Science, Technology, Engineering and 115 Mathematics Ethical Review Committee at University of Birmingham (reference: ERN 12-0830).

116

### 117 Preliminary measurements

Height and weight were assessed using standard methods and cardiorespiratory fitness ( $\dot{V}O_{2MAX}$ ) was 118 119 measured during an incremental exercise test on an electromagnetically braked cycle ergometer (Lode Excalibur Sport, Groningen, Netherlands). Following a three-minute warm up at 30 watts, workload was 120 increased by 30 watts every minute, until volitional exhaustion. A facemask was fitted throughout preliminary 121 exercise tests in each main trial, and expired air measured breath-by-breath averaged every 20 seconds for 122 oxygen uptake and carbon dioxide production (Oxycon Pro, Jaeger, Wuerzberg, Germany). Heart rate (HR) 123 was monitored every minute using a Polar Vantage heart rate monitor (Polar Vantage, Kempele, Finland); 124 HR<sub>peak</sub> represents the maximum heart rate achieved during each trial. The following criteria were used to 125 indicate that  $\dot{V}O_{2MAX}$  had been reached: a fall in cadence below 60 rpm, a respiratory exchange ratio ( $\dot{V}CO_2/$ 126  $\dot{V}O_2$ ) >1.10-1.15, a plateau in oxygen consumption and a heart rate >220 beats min<sup>-1</sup> minus age. 127

128

### 129 *Experimental trials*

Experimental trials were undertaken at least seven days after preliminary measurements, in the morning, and following an overnight fast. Each trial was separated by at least three days in a randomised

132 design. Prior to each trial, but after the baseline blood sample, participants undertook a warm up (5 minutes) at a workload eliciting 40%  $\dot{V}O_{2 \text{ MAX}}$ . Exercise trials were either vigorous steady state cycling at 80%  $\dot{V}O_{2 \text{ MAX}}$ 133 for 20 minutes ('continuous exercise') or high intensity interval exercise ('HIIE') trial. HIIE consisted of ten 1 134 minute cycling phases at a workload to elicit 90%  $\dot{VO}_{2MAX}$ , with 1 minute of low intensity cycling at 40% 135  $\dot{VO}_{2MAX}$  between each phase. Workload was expressed as watts, and relative to body mass (i.e., watts/kg). 136 Values presented are the average workload over the entire exercise protocol (i.e., in the vigorous trial: the 137 duration of cycling at 80%  $\dot{VO}_{2_{MAX}}$ ; and in the HIIE protocol: each 1-minute sprint phase of cycling at 90% 138 139  $\dot{VO}_{2MAX}$ ). Values were obtained directly from the electromagnetically braked cycle ergometer (Lode 140 Excalibur Sport, Groningen, Netherlands). Energy expenditure (Kcal) was estimated by indirect calorimetry from calculations of fat and carbohydrate oxidation (g/min) and data were averaged over the 1-minute phases 141 142 of the protocols and summed to provide total energy expenditure from the entire duration of each trial, expressed relative to body mass (Kcal/kg). Ratings of perceived exertion (RPE) were recorded using the Borg 143 Scale every 1 minute during continuous exercise, and after each 90%  $\dot{V}O_{2MAX}$  interval during HIIE; the 144 highest RPE score was selected as the final RPE result for each exercise trial (RPE<sub>peak</sub>). 145

146

147 Blood sampling

An intravenous cannula (*Becton & Dickson, Oxford, UK*) was inserted into an antecubital vein and
blood samples were drawn into potassium ethylene diaminetetraacetic acid (EDTA) vacutainer tubes (*Becton & Dickson, Oxford, UK*). The cannula was kept patent with saline (0.9% NaCl). The leukocyte differential
was assessed using an automated haematology analyser (*Coulter Analyser, Beckman-Coulter, High Wycombe, UK*).
UK).

153

#### 154 Blood Cell Isolation

Approximately 15 ml of blood from each time point (pre-exercise, post-exercise and 30 min postexercise) was diluted 1:1 with Roswell Park Memorial Institute Media (RPMI), and then layered on top of Ficoll paque PLUS (GE Healthcare) (2 blood : 1 Ficoll), before centrifuging at  $500 \times g$  for 30 minutes at 21°C. PBMCs were aspirated and washed three times in RPMI by centrifuging at  $400 \times g$  for 5 minutes. The

cell pellet was re-suspended in 1-ml of freezing mixture (70% RPMI, 20% fetal calf serum (FCS) and 10%
dimethyl sulfoxide (DMSO)) and frozen at -1°C / min using a freezing container (Nalgene "Mr Frosty"
Thermoscientific). Cells were stored at -80°C and analysed within six months.

162

### 163 *Flow cytometry*

164 Samples were thawed rapidly at 37°C and washed twice in PBS containing 2% FCS and 2mM EDTA by centrifuging at  $400 \times g$  for 5 minutes. Cells were counted using a haemocytometer and approximately 165 166 300,000 PBMCs were added to tubes for incubation with fluorescently conjugated antibodies to identify 167 specific lymphocyte populations using eight-colour flow cytometry (FACS-CANTO, Becton-Dickinson, San 168 Jose, USA). The following monoclonal antibodies (mAbs) were used: anti-CD45RA-FITC clone # HI100, anti-CD197 (CCR7)-PE clone # 150503, anti-CD56-PE-Cy7 clone # B159, anti-CD8-APC clone # RPA-T8, 169 170 anti-CD3-APC-Cy7 clone # SK7 (BD Pharmingen, San Diego, USA), anti-CLA-V450 clone # HECA-452 171 (BioLegend, San Diego, USA) and anti-CD16-V500 clone # 3G8 (BD Horizon, San Diego, USA). In addition, 7-aminoactinomycin D (7-AAD; PerCP channel; BD Pharmingen, San Diego, USA) was used to exclude 172 necrotic and apoptotic cells. For validation purposes, fluorescence-minus-one (FMO) tubes were used on 173 separate samples from three healthy donors to establish negative and positive gating strategies for CLA 174 175 expression.

Flow cytometry data were analysed using FlowJo version 7.6.5 (FlowJo LLC, Oregon, USA). Briefly, 176 lymphocytes were gated on the forward versus sideways scatter, and 7AAD- cells were divided into CD3+ or 177 CD3- cells. Cytotoxic T cells were identified as being CD3+CD8+ and further differentiated into naïve 178 ('NA'; CCR7+CD45RA+), central memory ('CM'; CCR7+CD45RA-), effector memory ('EM'; 179 CCR7-CD45RA-) or CD45RA+ effector memory cells ('EMRA'; CCR7-CD45RA+). CD3- cells were 180 further differentiated into CD56<sup>dim</sup> 'cytotoxic' NK cells or CD56<sup>high</sup> 'regulatory' NK cells, as proposed by 181 Cooper and colleagues (2001), using CD16 and CD56 dotplot gating (8). Within each subpopulation identified 182 above, cells were subdivided based on their expression of CLA (CLA+ or CLA-) in SSC (side scatter) versus 183 184 V450-CLA mode. Absolute cell counts for each subpopulation were computed by multiplying the percentage of cells within the CD3+ or CD3- gates by the total lymphocyte count (obtained via the Coulter principle). 185

This value and subsequent absolute subpopulation counts were multiplied by percentage values for gateddaughter subpopulations.

188

### 189 Statistical analyses

190 All results are presented as mean ± standard deviation unless otherwise stated. Statistical calculations were performed on IBM SPSS for Windows Version 21. Kolmogorov-Smirnov tests confirmed that all data 191 were normally distributed at all time points and for all variables. Repeated Measures Analysis of Variance 192 (ANOVA) tests were used to contrast changes over time (baseline, exercise and 30 min post-exercise), and 193 between the two exercise conditions (HIIE and continuous exercise) for each cell type. Within-trial main 194 effects of time for each cell type were calculated in separate individual ANOVAs. Post-hoc pairwise 195 comparisons were made with Bonferroni adjustments for multiple comparisons. Between trial differences at a 196 197 given time point were assessed by paired *t*-tests. Statistical significance was accepted at the p < .05 level.

#### 198 RESULTS

All participants completed both the continuous exercise and HIIE tasks. Total energy expenditure was significantly higher after continuous exercise compared to HIIE (HIIE:  $2.69 \pm 0.31$  Kcal/kg; continuous exercise:  $3.54 \pm 0.48$  Kcal/kg;  $F_{(1,17)} = 20.0$ , p < .05). Similarly, RPE<sub>peak</sub> was significantly higher after continuous exercise compared to HIIE (HIIE:  $16 \pm 2$  (Borg scale); continuous exercise:  $18 \pm 1$  (Borg scale);  $F_{(1,17)} = 4.6$ , p < .05). Workload was higher after HIIE compared to continuous exercise (HIIE:  $2.74 \pm 0.54$ Watts/kg; continuous exercise:  $2.18 \pm 0.33$  Watts/kg;  $F_{(1,17)} = 7.1$ , p < .05). No differences were found between trials for HR<sub>peak</sub> (HIIE:  $178 \pm 14$  beats/min; continuous exercise:  $187 \pm 7$  beats/min; p > .05).

206

### 207 *Effects of HIIE and continuous exercise on the number of total leukocytes and leukocyte subpopulations*

Total leukocyte counts increased (p < .05) from baseline after both the HIIE ( $5.31 \pm 1.15 [x10^9/L]$  to 8.92 ± 2.77) and continuous exercise ( $5.57 \pm 0.94$  to  $10.73 \pm 2.14$ ) conditions. Similarly, total lymphocyte counts also increased (p < .05) from baseline after both the HIIE ( $1.83 \pm 0.47$  to  $3.64 \pm 1.46$ ) and continuous exercise ( $1.91 \pm 0.41$  to  $4.83 \pm 1.33$ ) conditions, with a larger mean change observed during continuous exercise, compared to HIIE, but these differences were not significant (p > .05). 30 min after each exercise condition, lymphocyte numbers between trials were similar (HIIE:  $1.67 \pm 0.37$ ; continuous exercise:  $1.84 \pm$ 0.35) and had returned to pre-exercise levels (p > .05, compared to baseline).

215

## 216 Effects of HIIE and continuous exercise on the number of total CD3+ T cells and CD3+CD8+ T cells

Table 1 displays total CD3+ T cells and CD3+CD8+ T cell numbers during both exercise tasks. Total CD3+ T cells increased significantly during both conditions (p < .01), with larger mean increases during continuous exercise, compared to HIIE, but these between trial differences were not significant (p > .05). 30 min after each condition, CD3+ T cell numbers between trials were similar and had returned to pre-exercise levels. CD3+CD8+ T cells increased significantly during both conditions (p < .05), with larger increases during continuous exercise compared to HIIE; again, these differences between trials were not significant (p > .05). 30 min after each condition, CD3+CD8+ numbers between trials were similar and had returned to pre-223 .05). 30 min after each condition, CD3+CD8+ numbers between trials were similar and had returned to pre-

224	exercise levels. Replicating previous findings (6, 39), CD3+CD8+ T cell subsets were mobilised in a stepwise
225	manner, with CD3+CD8+ EMRA T cells mobilising more than EM, CM and NA cells. Within these subsets,
226	there were no significant differences in the magnitude of mobilisation between continuous exercise or HIIE
227	trials.

228

*Effects of HIIE and continuous exercise on the numbers of CLA+ and CLA- CD3+ T cells and CD3+CD8+ T cells*

Approximately 14% of total CD3+ T cells were CLA+ at baseline, and despite an increase in the number of CLA+ CD3+ T cells during both exercise conditions (p < .05), the proportion of CLA+ CD3+ T cells decreased slightly during exercise due to a larger influx of CLA- CD3+ T cells (p = .05) (see Table 1). At 30 minutes post-exercise the number and proportion of CLA+ CD3+ T cells returned to pre-exercise values.

Approximately 9% of CD3+CD8+ T cells were CLA+ at baseline, and, in a similar manner to total CD3+ T cells, the number of CLA+ CD3+CD8+ T cells increased during both exercise conditions (p < .05), before returning to baseline levels at 30 minutes post-exercise (see Table 1). However, there was no change (p> .05) in the proportion of CLA+ CD3+CD8+ T cells during exercise. No differences were observed in the magnitude of mobilisation between CLA+ and CLA- cells in any of these broad T cell subsets between HIIE and continuous exercise.

242

243 Effects of HIIE and continuous exercise on the numbers of CLA+ and CLA- CD3+CD8+ T cell subsets

In accordance with prior literature (7, 31), very few of the CD3+CD8+ NA cell population expressed CLA (median = 1.35%) and the few CLA+ naïve cells did not change during exercise (data not shown). The largest proportion of CLA+ CD3+CD8+ T cells were of the CM (~20%) and EM (~23%) phenotypes, with only a small fraction of EMRA cells expressing CLA (<7%). As illustrated in Figure 1, the percentage change in the number of CLA+ cells increases in a stepwise mobilisation pattern (i.e., EMRA > EM > CM) in

response to both exercise conditions, and this pattern is also evident in CLA– cells. However, as illustrated in Figure 2, the proportion of CLA+ cells per memory CD3+CD8+ T cell subset actually shows a marginal and significant (p<0.05) decline during exercise, before returning to baseline levels at 30 min post-exercise. Thus, CLA+ cells do not appear to be major contributors to the large influx of effector cells observed in the peripheral bloodstream during exercise; this is illustrated by a greater mobilisation of CLA–, compared to CLA+, EMRA CD3+CD8+ T cells in Figure 1.

255

*Effects of HIIE and continuous exercise on the numbers of NK cells, and CLA+ and CLA- cells within the NK cell subsets*

258 Table 2 displays changes in the numbers of CD3-CD56+ NK cells in response to continuous exercise 259 and HIIE. As expected, NK cells were extremely sensitive to exercise stress, exhibiting significant differences 260 between the two exercise modalities (p < .05), with HIIE resulting in a ~400% increase and continuous exercise resulting in a ~600% change during exercise. With regards to the major CD56+ NK cell subsets, 261 CD56<sup>dim</sup> 'cytotoxic' cells were the most sensitive to exercise stress, with HIIE resulting in a 550% increase 262 and continuous exercise resulting in a 725% increase (p < .05). As expected, CD56<sup>bright</sup> 'regulatory' NK cells 263 264 were less sensitive to exercise stress and no differences were found between exercise conditions (p > .05); 265 HIIE resulted in a 100% increase and continuous exercise resulted in a 200% increase.

At baseline, approximately ~22% of CD3-CD56+ cells were CLA+, with a higher proportion of 266 CD56<sup>bright</sup> 'regulatory' NK cells positive for CLA (~67%) than CD56<sup>dim</sup> 'cytotoxic' NK cells (~17.5%). 267 During exercise, the total number of CLA+ NK cells increased (HIIE: ~230%; continuous exercise: ~350%), 268 and, as expected, this was driven by a larger increase in CLA+ CD56<sup>dim</sup> cells compared to the CLA+ 269 CD56<sup>bright</sup> cells. As a consequence of a greater influx of CLA- NK cells (HIIE: ~500%; continuous exercise: 270 ~650% compared to baseline), primarily comprised of CLA-CD56<sup>dim</sup> cells during exercise (Figure 3), the 271 proportion of CLA+ cells in the total NK cell pool was actually reduced during exercise (Figure 4). Thus, 272 CLA+ NK cells do not appear to be major contributors to the large influx of effector cells observed in the 273 274 peripheral bloodstream during exercise.

#### 275 DISCUSSION

This study assessed the mobilisation propensity of CLA+ CD8+ T cells into the peripheral 276 277 bloodstream in response to intensive exercise. A preferential mobilisation of this cell phenotype would fulfil a 278 component of the stress redistribution theory (10), which hypothesises that effector memory cells 279 preferentially mobilised by exercise have a phenotype that enables post-exercise migration to cutaneous sites of wound healing or active inflammation (e.g., injured skin). We examined the number of CLA+ and CLA-280 281 cells because CLA avidly binds CD62E; CD62E is thought to be the primary initiator of routine CD8+ T cell 282 transmigration to the skin, and is also central to T cell tethering to cytokine-activated endothelium at inflamed 283 cutaneous sites (14). We found that exercise resulted in an increase in the number of memory CLA+ CD3+CD8+ T cells. However, the numerical contribution of CLA+ memory cells to exercise-induced 284 285 lymphocytosis was inferior to CLA- cells, and, as a consequence, the proportion of CLA+ cells amongst the 286 total memory CD8+ T cell pool showed a decline during exercise. Furthermore, the most exercise-sensitive subset (EMRA CD8+ T cells) were largely CLA-, suggesting that the most exercise-responsive T cell subset 287 288 mobilised by exercise do not have a phenotype that would enable rapid transmigration to sites of active 289 cutaneous inflammation.

290 Until now, the effect of exercise on the mobilisation of CLA+ cells has not yet been investigated. A 291 study investigating the effects of acute psychological stress on circulating CLA+ CD8+ cells found a marginal decline in the number of CLA+ CD8+ cells during the stressor (2). These aforementioned results differed to 292 293 the findings of our study, as we observed an approximate doubling in the number of CLA+ CD3+CD8+ cells 294 during exercise. It was proposed by Atanackovic et al. (2006) that CLA+ CD8+ T cells had already initiated 295 transmigration to sites of inflammation during acute stress. However, evidence exists to suggest that elevated 296 levels of epinephrine selectively decreases adhesion of CD8+ T cells (12) and NK cells (3) to endothelial 297 cells, and may thus be a contributing mechanism for the maintenance of CLA+ cells in the peripheral 298 bloodstream, as observed in our exercise trials. We extended our CLA+ analyses to four distinct CD8+ T cell subsets conventionally identified using the cell-surface markers CCR7 and CD45RA (17). In agreement with 299 prior literature showing that the majority of CLA+CD3+ cells are CD45RO+ (7), we found that few naïve 300 (CCR7+CD45RA+) CD8+ T cells expressed CLA (31). This did not change in response to exercise. In further 301

302 agreement with Clark et al. (2006), we found a majority of CLA+ cells were of the central memory (CCR7+ 303 CD45RA-) phenotype and fewer were effector memory (CCR7-CD45RA-) CD8+ T cells; these CLA+ 304 effector memory cells showed the largest increase during exercise, though the numerical increase was small. 305 On one hand, the findings of this study are supportive of the stress redistribution model in that 306 exercise evoked an increase in the total number of peripheral blood CLA+ CD8+ T cells, which are 307 phenotypically consistent with skin-homing T cells. On the other hand, the absolute number of CLA+ cells 308 mobilised was small, and the relative magnitude of mobilisation was much less than CLA- CD8+ cells. 309 Furthermore, few EMRA CD8+ T cells – conventionally the most exercise-sensitive CD8+ subset – expressed 310 CLA, and did not increase in number as much as CLA- EMRA CD8+ T cells. Thus, the CD8+ T cells with the greatest propensity for exercise-induced mobilisation and extravasation are CLA- and do not exhibit a 311 phenotype characteristic for rapid homing to cutaneous sites or sites of endothelial inflammation. Given that 312 313 the large majority of CLA+ cells reside in the skin (7), it is possible to conclude that the skin is not a major 314 contributor/source of cells to exercise-induced lymphocytosis. The findings of this study are not surprising: CLA+ cells are not as susceptible to immunosenescence (25), and are thus strikingly different to the most 315 316 exercise-sensitive cells, which have shorter telomeres, high expression of CD57 (35, 36); and exhibit exaggerated exercise-induced mobilisation in participants seropositive for cytomegalovirus (39). In addition, it 317 318 has been demonstrated in rodents that exercise redeploys large numbers of T cells to the lungs (20). Lungderived T cells are all CLA-, but positive for other homing molecules including CCR5 and CXCR3 (5). 319 Similarly, exercise also redeploys T cells to the Peyer's patches and bone marrow (19, 20) and this is likely 320 governed by other site-specific homing molecules like LPAM-1 (37) or VLA-4 (23, 26) rather than CLA. This 321 allocation of certain cells to defined parts of the body might represent a homeostatic immune-surveillance 322 323 response (10), or, instead, it has been hypothesised that senescent T cells are mobilised into the blood to 324 facilitate their subsequent apoptosis in peripheral tissues (21, 34), which may contribute to progenitor cell 325 mobilisation after exercise (24). With regards to cutaneous surveillance against tumours and stressed tissue 326 cells, it is unlikely this is tasked by  $\alpha\beta$  CD8+ T cells alone (13), and is likely supported by  $\gamma\delta$  T cells and NK 327 cells, which are highly responsive to acute stress (1, 6), and have the migratory capacity to enter cutaneous sites (13). We found that CLA+ CD56+ NK cells were substantially mobilised by exercise, but, much like 328 329 CD8+ T cells, were outnumbered by the mobilisation of CLA- NK cells.

330 This is the first study to investigate and compare the effects of HIIE (also referred to as HIIT when 331 repeated over several weeks or months) and continuous exercise on T cell and NK cell subset mobilisation 332 responses. HIE typically involves shorter and more intense bouts of exercise than more traditional forms 333 (e.g., 30 minutes of moderate intensity running or cycling) and is thought to be more attractive and better 334 tolerated by participants. HIIE has received considerable recent attention as an effective means of achieving 335 certain physiological adaptations (e.g., improved insulin sensitivity and cardiorespiratory fitness, but probably 336 not weight loss) in healthy and diseased populations that are similar or superior to traditional endurance-based 337 exercise (16).

Although we found a trend whereby a greater mobilisation of lymphocytes was observed during continuous exercise compared to HIIE, these differences were not significant for the majority of lymphocyte subpopulations investigated. We found that CD56 NK cells – conventionally the most exercise responsive cells – were mobilised to a greater extent during continuous exercise compared to HIIE, a response that was driven by larger increases to CD56<sup>dim</sup> NK cells. It may be that similar intensity-dependent effects would have been found for CD8+ T cells, and other subsets, if a larger group of participants had been tested.

A limitation of this study was that post-exercise blood samples were collected 30 minutes after 344 exercise cessation, rather than after 60 minutes, when NK and CD8+ T cell lymphocytopenia is more 345 346 pronounced (6, 39). This was a consequence of practical and logistical constraints imposed by the broader study that was being undertaken (41). At 30 min post-exercise, we found that all cell phenotypes were present 347 in peripheral blood in similar numbers to pre-exercise levels. It would be of interest to investigate the 348 contribution of CLA+ T cells to lymphocytopenia at later time points. An additional practical limitation was 349 the cryopreservation of PBMCs, which may disproportionately affect the viability of some cell phenotypes -350 351 however this is unlikely to affect the within-subject model used in our study. A further consideration for future studies surrounds the ongoing debate over the optimal phenotyping of skin-homing CD8+ T cells. In 352 353 addition to CLA, CCR4 and CCR8 may be useful in differentiating between CD8+ T cells involved in normal 354 cutaneous immune-surveillance to those involved in acute or chronic inflammation (7, 33). Finally, our 355 investigations were limited to healthy individuals with no apparent cutaneous inflammation. Investigation of exercise-induced lymphocyte skin-homing in experimental models of infection or in clinical populations 356

including psoriasis is warranted, and more invasive techniques such as biopsy may provide further insightsinto exercise-induced homing to cutaneous or inflammatory sites.

359

## 360 CONCLUSION

361 A greater number of CLA- T cells and NK cells were mobilised into peripheral blood than CLA+ counterpart T cells and NK cells during exercise. Furthermore, the majority of EMRA T cells and CD56<sup>dim</sup> 362 cells - i.e., conventionally the most exercise-responsive cells - did not express CLA. Together, these findings 363 demonstrate that CLA+ cells are not major contributors to exercise lymphocytosis, thus providing preliminary 364 evidence that the skin is not a major origin, or homing-destination, of exercise-sensitive lymphocytes. We 365 conclude that the most exercise-sensitive lymphocytes likely migrate from, and to, non-cutaneous sites post-366 367 exercise. 368 369 ACKNOWLEDGEMENTS: The results of this present study do not constitute endorsement by ACSM. The 370 flow cytometric analyses for this study were funded by the University of Birmingham Clinical Immunology 371 Service.

372

373 **CONFLICTS OF INTEREST:** None declared

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# 485 FIGURE CAPTIONS

486

487	<b>Figure 1.</b> Mean ± SEM changes in the proportion of CD8+ T cell subsets (CM = central memory; EM =					
488	effector memory; EMRA = CD45RA+ effector memory) that were either positive or negative for cell surface					
489	CLA, at baseline and immediately after Continuous exercise. It was found that CLA+ CD45RA+ effector					
490	memory CD8+ T cells were mobilised to a greater magnitude than CLA- CD45RA+ effector memory CD8+					
491	T cells; similar patterns were found for HIIE (data not shown).					
492	* Indicates significant difference between CLA+ and CLA– ( $p < .05$ ).					
493						
494	<b>Figure 2.</b> Mean ± SEM changes in the proportion of CD8+ T cell subsets (NA = naïve; CM = central					
495	memory; EM = effector memory; EMRA = CD45RA+ effector memory) that were positive for CLA, from					
496	pre- to immediately post-continuous exercise. A reduction in the proportion of CLA+ CM, EM and EMRA					
497	cells was observed immediately after continuous exercise compared to baseline; similar patterns were					
498	observed during HIIE (data not shown).					
499	* Indicates significant differences in the proportion of CLA+ cells between pre- and post-continuous exercise					
500	(p < .05).					
501	<b>Figure 3.</b> Mean ± SEM changes in the proportion of NK cell subsets that were either CLA+ or CLA-, at					
502	baseline and immediately after continuous exercise. A trend was observed whereby CLA- CD56 <sup>dim</sup> cells					
503	were mobilised to a greater extent than CLA+ CD56 <sup>dim</sup> NK cells ( $p = .058$ ); similar patterns were found for					
504	HIIE (data not shown).					
505	<b>Figure 4</b> Mean + SEM changes in the proportion of NK call subsets that were CLA+ from pre- to					
505	Figure 4. Mean - SEM changes in the proportion of NK cen subsets that were CLA+, non pre- to					
506	immediately post-continuous exercise. A reduction in the proportion of CLA+ CD56 <sup>bright</sup> and CD56 <sup>dim</sup> NK					
507	cells was observed from pre- to post-exercise; similar patterns were observed during HIIE (data not shown).					
508	* Indicates significant differences in the proportion of CLA+ cells between pre- and post-continuous exercise					

509 (*p* < .05).

510

**Table 1.** Changes in the numbers of T-lymphocytes and T-lymphocyte subpopulations in response to HIIE and continuous exercise (mean  $\pm$  SD).

Cells per µL	Condition	Baseline	Exercise	+30min	Main effects of time	Time × task interaction	
CD3+	HIIE	$1494.4 \pm 406.2$	$2508.8 \pm 1076.5 *$	$1330.2\pm293.2$	$F_{(2,16)} = 16.18; p = .002$	$E_{n} = 2.06; n = NS$	
CD3+	Continuous Ex	$1454.1\pm321.9$	$3049.6 \pm 794.4 {***}$	$1416.3\pm363.1$	$F_{(2,16)} = 42.97; p < .001$	$\Gamma_{(2,32)} = 2.00, p = 103$	
CD2+CLA+	HIIE	$189.2\pm89.7$	$316.2 \pm 237.0$	$186.8 \pm 95.7$	$F_{(2,16)} = 6.80; p = .030$	$F_{(2,22)} = 37 \cdot n = NS$	
CD3+ CLA+	Continuous Ex	$183.0\pm91.4$	$350.6 \pm 166.6 **$	$196.7\pm123.2$	$F_{(2,16)} = 26.06; p < .001$	$\Gamma_{(2,32)} = .57, p = 143$	
CD2+CLA	HIIE	$1305.2 \pm 376.2$	$2192.6 \pm 894.2 **$	$1143.4 \pm 260.2$	$F_{(2,16)} = 17.72; p = .002$	E = 2.22; n = NE	
CD3+ CLA-	Continuous Ex	$1271.0\pm293.3$	$2699.0 \pm 276.0 {***}$	$1219.6\pm291.7$	$F_{(2,16)} = 41.40; p < .001$	$\Gamma_{(2,32)} = 2.55, p = 185$	
CD2+ CD2+	HIIE	$509.5\pm181.9$	$1000.9\pm654.7$	$463.1\pm174.7$	$F_{(2,16)} = 9.46; p = .014$	E = 20, MC	
CD3+ CD8+	Continuous Ex	$491.2\pm162.3$	$1185.0\pm 398.8^{***}$	$490.4\pm195.4$	$F_{(2,16)} = 48.99; p < .001$	$F_{(2,32)} = .89; p = INS$	
	HIIE	41.3 ± 21.6	77.9 ± 65.6	43.2 ± 29.3	$F_{(2,16)} = 6.22; p = .035$		
CD3+ CD8+ CLA+	Continuous Ex	$40.7\pm26.2$	86.9 ± 53.8**	$43.8\pm32.7$	$F_{(2,16)} = 23.95; p = .001$	$F_{(2,32)} = .28; p = NS$	
	HIIE	$468.2 \pm 163.4$	923.0 ± 591.5	$420.0 \pm 148.7$	$F_{(2,16)} = 9.76; p = .013$		
CD3+ CD8+ CLA-	Continuous Ex	$450.4\pm141.2$	$1098.1 \pm 355.0 ***$	$446.5\pm166.9$	$F_{(2,16)} = 48.96; p < .001$	$F_{(2,32)} = .95; p = NS$	
N	HIIE	$239.9\pm93.1$	$341.6 \pm 144.1*$	$192.8\pm62.2$	$F_{(2,16)} = 14.14; p < .001$	E 1.51 NG	
Naive CD8+	Continuous Ex	$215.9\pm76.1$	$377.9 \pm 69.0 ***$	$198.5\pm56.4$	$F_{(2,16)} = 55.18; p < .001$	$F_{(2,32)} = 1.51; p = NS$	
	HIIE	3.4 ± 1.4	5.2 ± 2.6	3.0 ± 1.4	$F_{(2,16)} = 10.20; p = .001$		
Naive CD8+ CLA+	Continuous Ex	$3.5 \pm 1.8$	$6.3 \pm 3.4*$	$3.5 \pm 2.2$	$F_{(2,16)} = 17.67; p = .002$	$F_{(2,32)} = .97; p = NS$	
	HIIE	$236.5\pm92.2$	336.4 ± 142.3*	$189.8\pm62.0$	$F_{(2,16)} = 14.13; p < .001$	E 1.40 MG	
Naive CD8+ CLA-	Continuous Ex	$212.4\pm75.8$	$370.5 \pm 70.3 ***$	$195.0\pm55.4$	$F_{(2,16)} = 54.09; p < .001$	$\mathbf{F}_{(2,32)} = 1.49; p = NS$	
CM	HIIE	$118.8\pm37.2$	$221.8\pm97.1*$	$124.7\pm55.9$	$F_{(2,16)} = 21.50; p = .001$	$E = 2.95 \cdot n = NC$	
CM	Continuous Ex	$121.9\pm64.2$	$278.6 \pm 96.2 ***$	$132.3\pm85.4$	$F_{(2,16)} = 101.54; p < .001$	$\Gamma_{(2,32)} = 5.85, p = 185$	
CM CD9+ CLA+	HIIE	$20.2 \pm 8.3$	$35.6\pm25.0$	21.7 ± 12.7	$F_{(2,16)} = 7.01; p = .026$	E 20 NG	
CM CD8+ CLA+	Continuous Ex	$20.0\pm11.8$	$38.9 \pm 20.7 **$	$21.3\pm14.7$	$F_{(2,16)} = 25.16; p < .001$	$F_{(2,32)} = .29; p = INS$	
CN CD9+ CLA	HIIE	$98.6\pm30.4$	186.2 ± 72.9**	$103.1 \pm 44.2$	$F_{(2,16)} = 27.81; p < .001$	F 5 (0 017	
CM CD8+ CLA-	Continuous Ex	$101.9\pm53.5$	$239.7 \pm 77.4 ***$	$110.6\pm71.8$	$F_{(2,16)} = 123.76; p < .001$	$F_{(2,32)} = 5.69; p = .017$	
EM	HIIE	$64.4\pm29.9$	$144.9 \pm 70.8 **$	$69.5\pm37.2$	$F_{(2,16)} = 26.74; p = .001$	$E_{-2,20,} = NS$	
EM	Continuous Ex	$66.6\pm39.3$	$184.8 \pm 71.0$ ***	$75.6\pm52.0$	$F_{(2,16)} = 80.73; p < .001$	$F_{(2,32)} = 3.50; p = 1NS$	
	HIIE	$12.3 \pm 10.5$	$24.7\pm27.5$	13.7 ± 13.4	$F_{(2,16)} = 4.73; p = NS$	E 12 NG	
EM CD8+ CLA+	Continuous Ex	$12.0\pm10.8$	$26.9\pm24.5*$	$13.6\pm14.0$	$F_{(2,16)} = 11.22; p = .010$	$F_{(2,32)} = .13; p = NS$	
	HIIE	52.1 ± 20.5	120.2 ± 44.7***	55.8 ± 24.6	$F_{(2,16)} = 40.39; p < .001$	$F_{(2,32)} = 5.08; p = .026$	
EM CD8+ CLA-	Continuous Ex	$54.6\pm29.7$	157.8 ± 51.3***	$62.2\pm40.4$	$F_{(2,16)} = 90.05; p < .001$		
EMDA	HIIE	$88.3\pm87.0$	$292.6\pm390.3$	$76.1\pm69.7$	$F_{(2,16)} = 4.07; p = NS$	E = 12; n = NS	
EMKA	Continuous Ex	$86.6\pm57.7$	$344.6 \pm 296.6*$	$83.8\pm52.1$	$F_{(2,16)} = 9.16; p = .016$	$F_{(2,32)} = .13; p = NS$	
	HIIE	$5.4 \pm 4.0$	$12.4 \pm 12.6$	4.9 ± 3.7	$F_{(2,16)} = 5.35; p = .047$	$E = 20 \cdot n - NC$	
EMIKA UD8+ ULA+	Continuous Ex	$5.4 \pm 4.4$	$14.6 \pm 10.9 **$	$5.4 \pm 4.6$	$F_{(2,16)} = 17.13; p = .003$	$\Gamma_{(2,32)} = .29; p = NS$	
	HIIE	$80.9\pm84.0$	$280.2\pm378.7$	71.2 ± 66.8	$F_{(2,16)} = 4.02; p = NS$	$E = 10 \dots - M^{O}$	
EMIKA UD8+ ULA-	Continuous Ex	$81.4\pm54.4$	$329.9 \pm 291.0*$	$78.4\pm48.6$	$F_{(2,16)} = 8.75; p = .018$	$\mathbf{F}_{(2,32)} = .12; p = \mathbf{NS}$	

\*p < .05 in comparison to baseline

\*\* p < .01 in comparison to baseline

\*\*\* p < .001 in comparison to baseline

NS p > .05

Cells per µL	Condition	Baseline	Exercise	+30min	Main effects of time	Time × task interaction	
	HIIE	$112.4 \pm 44.8$	591.9 ± 361.0**	$115.2 \pm 59.5$	$F_{(2,16)} = 18.27; p = .003$	F = -4.47; n = 05	
CD30+	Continuous Ex	$164.3\pm144.2$	$1130.6 \pm 707.5 **$	$157.6\pm159.9$	$F_{(2,16)} = 22.64; p = .001$	$\Gamma_{(2,32)} = 4.47, p = .05$	
CD56+CLA+	HIIE	$28.2 \pm 10.4$	92.7 ± 47.8**	$29.2 \pm 12.4$	$F_{(2,16)} = 23.96; p = .001$	$F = -3.37 \cdot n - NS$	
CD30+CEA+	Continuous Ex	$31.7\pm9.5$	$142.1 \pm 69.2 **$	$31.9\pm10.9$	$F_{(2,16)} = 26.10; p = .001$	$\Gamma_{(2,32)} = 5.57, p = 143$	
CD56+CLA-	HIIE	84.1 ± 44.6	499.3 ± 352.6*	86.0 ± 56.5	$F_{(2,16)} = 14.13; p = .005$	$E = -2.75 \cdot n - NS$	
CD30+CLA-	Continuous Ex	$132.6\pm136.6$	$988.4 \pm 698.6 **$	$125.7\pm153.2$	$F_{(2,16)} = 18.14; p = .003$	$\Gamma_{(2,32)} = 5.75, p = 105$	
CD56 <sup>bright</sup>	HIIE	$11.0 \pm 7.4$	$23.9 \pm 19.4 *$	$15.2\pm13.0$	$F_{(2,16)} = 9.98; p = .011$	$F_{(2,32)} = 2.17; p = NS$	
CD30	Continuous Ex	$12.8\pm7.0$	$36.0 \pm 25.7*$	$15.6\pm7.8$	$F_{(2,16)} = 11.05; p = .009$		
CD56 <sup>bright</sup> CI A+	HIIE	7.6 ± 5.5	$15.0 \pm 12.7*$	$10.2 \pm 9.1$	$F_{(2,16)} = 9.05; p = .015$	$F_{(2,32)} = 1.32; p = NS$	
CD30 CLA+	Continuous Ex	$8.4 \pm 4.7$	$20.3 \pm 15.8*$	$10.2 \pm 5.7$	$F_{(2,16)} = 8.88 \ p = .015$		
CD56 <sup>bright</sup> CLA-	HIIE	3.4 ± 2.0	8.9 ± 7.2*	$5.0 \pm 4.0$	$F_{(2,16)} = 10.14; p = .011$	E = -1.70; n - NS	
CD30 - CLA-	Continuous Ex	$4.4 \pm 2.7$	$15.8\pm14.4$	$5.4\pm2.5$	$F_{(2,16)} = 6.35; p = .035$	$F_{(2,32)} = 1.79; p = 1.8$	
CD56 <sup>dim</sup>	HIIE	$101.4 \pm 44.5$	$568.0 \pm 343.6 **$	$99.9\pm55.6$	$F_{(2,16)} = 18.41; p = .003$	F = -4.40; n = .05	
CD30	Continuous Ex	$151.5\pm47.2$	$1094.5 \pm 229.9 **$	$141.9\pm53.1$	$F_{(2,16)} = 22.72; p = .001$	$\Gamma_{(2,32)} - 4.49; p = .05$	
CD56 <sup>dim</sup> CL A+	HIIE	$20.6\pm10.0$	77.7 ± 43.8**	$19.0 \pm 7.6$	$F_{(2,16)} = 22.68; p = .001$	$F_{(2,32)} = 3.21; p = NS$	
CD30 CLA+	Continuous Ex	$23.3\pm7.8$	$121.8 \pm 60.8 **$	$21.7\pm8.9$	$F_{(2,16)} = 25.51; p = .001$		
CD56 <sup>dim</sup> CL A-	HIIE	$80.7\pm44.0$	$490.3 \pm 345.8*$	81.0 ± 55.0	$F_{(2,16)} = 14.16; p = .005$	$E = -2.75 \cdot n - NS$	
CD30 CLA-	Continuous Ex	$128.2\pm135.3$	$972.7 \pm 689.1 **$	$120.3\pm152.3$	$F_{(2,16)} = 18.19 \ p = .003$	$\Gamma_{(2,32)} = 5.75, p = 1NS$	

Table 2. Changes in NK cell numbers in response to HIIE and continuous exercise (mean  $\pm$  SD).

\*p < .05 in comparison to baseline

\*\* p < .01 in comparison to baseline

NS p > .05



Figure 1









Figure 4

