## 1 Acetaminophen (paracetamol) induces hypothermia during

2 acute cold stress.

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- Abbreviations: APAP, acetaminophen,  $T_c$ , core temperature,  $T_{sk}$ , skin temperature, COX,
- 29 cyclooxygenase

#### **KEY POINTS**

- 31 Accidental hypothermia was the primary or secondary diagnosis in over 100,000 hospital
- admissions from 2005 to 2015 in the United Kingdom. In this study we sought to determine
- whether acetaminophen, a non-prescription drug used to manage mild pain and fever, reduced
- core temperature stability during a 2-hour passive cold or thermoneutral exposure.
- 35 Acetaminophen had no effect on core temperature in thermoneutral conditions compared with a
- placebo, but reduced core temperature by up to 0.57°C after 2-hours cold exposure. The present
- 37 results improve our knowledge about the side-effects of acetaminophen and provides important
- information of relevance for hypothermia pathology.

#### **ABSTRACT**

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- Background: Acetaminophen is an over-the-counter drug used to treat pain and fever, but it has
- also been shown to reduce core temperature  $(T_c)$  in the absence of fever. However, this side-
- effect is not well examined in humans, and it is unknown if the hypothermic response to
- acetaminophen is exacerbated with cold exposure.
- 45 Objective: To address this question, we mapped the thermoregulatory responses to
- acetaminophen and placebo administration during exposure to acute cold (10°C) and thermal
- 47 neutrality (25°C).
- 48 Methods: Nine healthy Caucasian males (age: 20 to 24 years) participated in the experiment. In a
- double-blind, randomised, repeated measures design, participants were passively exposed to a
- thermo-neutral or cold environment for 120-minutes, with administration of 20 mg/kg lean body
- mass acetaminophen or a placebo 5-minutes prior to exposure.  $T_c$ , skin temperature  $(T_{sk})$ , heart
- rate, and thermal sensation were measured every 10-minutes, and mean arterial pressure was
- recorded every 30-minutes. Data were analysed using linear mixed effects models. Differences in
- thermal sensation were analysed using a cumulative link mixed model.
- Results: Acetaminophen had no effect on  $T_c$  in a thermo-neutral environment, but significantly
- reduced  $T_c$  during cold exposure, compared with a placebo.  $T_c$  was lower in the acetaminophen
- 57 compared with the placebo condition at each 10-minute interval from 80 to 120-minutes into the
- trial (all p < 0.05). On average,  $T_c$  decreased by  $0.42 \pm 0.13$ °C from baseline after 120 minutes of
- cold exposure (range 0.16 to 0.57°C), whereas there was no change in the placebo group (0.01  $\pm$
- 60 0.1°C).  $T_{\rm sk}$ , heart rate, thermal sensation, and mean arterial pressure were not different between
- 61 conditions (p > 0.05).
- 62 Conclusion: This preliminary trial suggests that acetaminophen-induced hypothermia is
- exacerbated during cold stress. Larger scale trials seem warranted to determine if acetaminophen
- 64 administration is associated with an increased risk of accidental hypothermia, particularly in
- vulnerable populations such as frail elderly individuals.

- 67 Key Words: acetaminophen, paracetamol, thermoregulation, cold exposure, thermogenesis,
- 68 hypothermia

#### 1.1 INTRODUCTION

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71 Accidental hypothermia is characterised by an unintended core temperature (T<sub>c</sub>) reduction to 72 35°C or lower. Such a fall in  $T_c$  can induce ventricular fibrillation and ultimately cardiac arrest if  $T_c$  declines to < 28°C [1, 2]. In the United States, hypothermia was the cause or contributing 73 cause of death in over 5500 cases between 2006 and 2010 [3], but this is likely underestimated 74 75 since  $T_c$  needs to be measured at or near the time of death. Nonetheless, data from United Kingdom hospital episode statistics indicate that hypothermia was the primary or secondary 76 diagnosis in over 100,000 hospital admissions from 2005 to 2015 [4]. Although death from 77 hypothermia is rare, it remains a significant health risk in elderly and very young individuals, 78 79 particularly during winter months and unaccustomed cold spells [1]. Interestingly, there is a growing body of evidence demonstrating that acetaminophen could reduce  $T_c$  stability during 80 cold exposure (discussed below), placing users at an increased risk of accidental hypothermia. 81 Acetaminophen is an over-the-counter drug marketed as paracetamol in Europe and Tylenol in 82 the United States. It is best known for its ability to decrease pain perception and reduce  $T_c$  during 83 a fever; each of these actions are in part mediated through an inhibition of cyclooxygenase 84 (COX) enzyme activity [5]. However, there is evidence of a 'hypothermic' action of 85 acetaminophen, which refers specifically to an acetaminophen-induced reduction in  $T_c$ 86 independent of febrile status. In mice, high doses (150 to 300 mg/kg body mass) administered 87 intravenously reduced  $T_c$  by 2 to 4°C [6–8]. In humans, there have been 246 reports in Vigibase<sup>©</sup> 88 (the World Health Organisation international database of adverse drug reactions) specific to 89 90 acetaminophen-induced accidental hypothermia [9]. In addition, several case studies report profound hypothermia following therapeutic doses [10] and high doses of acetaminophen when 91 ingested orally [11, 12]. Finally, oral acetaminophen administration (20 mg/kg lean body mass) 92 reduced  $T_c$  in young adults by ~0.2°C (range, 0.10 to 0.39°C) during exposure to mild cold [13]. 93 Although the  $T_c$  reductions were small, this hypothermic side-effect of acetaminophen occurred 94 in all thirteen participants. Despite this data, additional criteria, such as the environmental 95 temperature, are needed to accurately predict when acetaminophen poses the greatest risk for 96 hypothermia development. Since the COX pathway could be involved in non-febrile 97 thermogenesis [14, 15], inhibition of this enzyme by acetaminophen might cause  $T_c$  to fall during 98 cold exposure, while exerting negligible effects on  $T_c$  while exposed to a warm environment. 99

If acetaminophen-induced hypothermia is a risk during cold exposure, this can have major implications for public health recommendations. Each year in the United States, ~6% of adults are prescribed acetaminophen at doses of more than 4 g/day [16], while it is also available overthe-counter without prescription. Acetaminophen is recommended as the first line analgesic for the elderly because it has minimal drug interactions and is well tolerated when taken at recommended doses [17]. It is also recommended for use in neonatal intensive care units following minor procedures and circumcision [18, 19]. Each of these age groups have a high incidence of accidental hypothermia due to a decreased ability to produce heat and make perceptually driven behavioural changes [20, 21]. Due to its hypothermic effects, use of acetaminophen in these populations could decrease their  $T_c$  to the point in which they are clinically hypothermic. However, the question remains as to whether acetaminophen exerts its hypothermic effect by increasing heat loss, or decreasing heat production. If the COX pathway is required for full heat production, inhibition of its activity by acetaminophen would cause  $T_c$  to fall during cold exposure while exerting no hypothermic action during a thermo-neutral exposure (in which no heat production is required). The aim of this trial was to examine the thermoregulatory response to acetaminophen administration (20 mg/kg of lean body mass) during a 120-minute exposure to a thermo-neutral and cold environment in healthy adult humans. Due to a potential role of COX in non-febrile thermogenesis [14, 15], it was hypothesised that acetaminophen would reduce  $T_c$  in cold conditions, but have no effect on  $T_c$  in thermo-neutral conditions relative to a placebo.

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## 1.2 METHODS

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## 1.2.1 Ethical Approval 121 Experimental procedures were approved by the University Research Ethics committee (approval 122 123 code 2014ISPAR011). All experimental procedures conformed to the standards set by the World Association Declaration of Helsinki 'Ethical Principles for Research Involving Human Subjects'. 124 1.2.2 Sample Size Calculation 125 Power analyses were conducted with GPower software version 3.1 (Heinrich University, 126 Düsseldorf, Germany) to determine the sample necessary to achieve two-tailed statistical 127 significance ( $\alpha = 0.05$ ), with a power of 0.90 and a partial eta-squared ( $\eta^2$ ) of 0.42. Using $T_c$ data 128 from a previous experiment where acetaminophen was tested as a hypothermic agent [22], it was 129 determined that nine participants were required to reach the statistical power. If acetaminophen 130 131 exerts the hypothesised hypothermic response, a larger project within vulnerable populations may be warranted to determine if acetaminophen contributes to accidental hypothermia 132 admissions. 133 1.2.3 Participants 134 Nine Caucasian males [age: $22 \pm 1$ years, height: $179 \pm 5$ cm, body mass: $80.7 \pm 11.9$ kg, body 135 fat $(20 \pm 5\%)$ ] volunteered to take part in this study. Participants were provided with written 136 information regarding the experimental procedures, with supporting oral explanations from the 137 principal investigator. All participants subsequently provided written informed consent. The 138 participants were non-smokers, non-febrile (resting $T_c < 38^{\circ}$ C), and free from musculoskeletal 139 140 injury. 1.2.4 Inclusion & Exclusion Criteria 141 Prior to each laboratory visit, participants completed an alcohol use disorder identification test 142 [AUDIT; [23]], a breathalyser test (AlcoSense, One, Berkshire, UK), and an acetaminophen risk 143 assessment questionnaire. To avoid any risk of liver damage inflicted by acetaminophen, 144

participants were not able to participate in the research if they scored above ten on the AUDIT

questionnaire or alcohol was present in their bloodstream (i.e. > 0% blood alcohol content). In addition, the acetaminophen dose was relative to lean body mass, as it is a closer indicator of liver volume than total body mass [24]. No participants presented with any pre-existing medical conditions that may have put them at an increased risk of acetaminophen toxicity. Due to potential thermoregulatory adaptations [25, 26], individuals were not permitted to take part in any experimental procedures if they were heat/cold acclimated or acclimatised. Thus, those who had travelled to a hot/cold climate or participated in a laboratory based heat/cold acclimation protocol less than three weeks before the experiment were not permitted to take part. All participants presented to the laboratory with a stable resting  $T_c$  of 36.5-37.5°C.

## 1.2.5 Experimental Design

A schematic of the experimental design is displayed in Figure 1. To determine if acetaminophen reduces  $T_c$  stability during cold stress compared to a placebo, the participants visited the laboratory on five occasions, each separated by at least seven days. On visit 1, participants arrived fasted (overnight) and their body fat was assessed via air displacement plethysmography (Bod Pod, 2000A, Birmingham, UK). The body fat reading from this test was used to determine the participant's dose of acetaminophen received in the experimental trials. Visits 2-5 (experimental trials) were randomised (SPSS Inc., Chicago, USA), double blinded (drug only), and followed a repeated measures design. On these visits, participants were exposed to either cold [10°C, 40% relative humidity (r.h)] or thermo-neutral (25°C, 40% r.h.) environmental temperatures for 120 minutes, having been administered acetaminophen (20 mg/kg of lean body mass) or a placebo (dextrose). Acetaminophen (Paracetamol, Aspar Pharmaceuticals, London, UK) was administered via the oral route. The placebo was matched in terms of appearance i.e. the same number of capsules were provided to the participants. The average dose of acetaminophen administered in the present work was 1,287  $\pm$  173 mg (range, 1,082 to 1,486 mg).

\*\*\*please insert Figure 1 near here\*\*\*

## 1.2.6 Experimental Protocol

All participants arrived at the laboratory at 10:00. Upon arrival, participants were instrumented 174 for the measurement of  $T_c$ , skin temperature  $(T_{sk})$ , and heart rate (see "Instrumentation and 175 Equations" for details). Thirty minutes after arrival participants consumed a standardised 176 breakfast [cornflakes (50 g), milk (250 ml) and 1 litre of tap water] and ingested acetaminophen 177 or a placebo one hour after the meal was consumed. Participants remained rested in an upright, 178 seated position between meal consumption and acetaminophen or placebo ingestion to ensure 179 resting physiological status was attained. Participants were wheeled into the environmental 180 181 chamber immediately following drug administration, and remained in the seated position for the duration of the protocol. Clothing was shorts and calf length socks, representing a Clo value of 182  $\sim$ 0.1. Resting measurements of  $T_c$ ,  $T_{sk}$ , heart rate and thermal sensation were collected five 183 minutes prior to acetaminophen and placebo ingestion, and subsequently every 10 minutes for 184 185 120 minutes' post-ingestion. Blood pressure was measured prior to chamber entry and every 30 minutes (pre-ingestion, 30, 60, 90, 120 minutes post-ingestion) until the end of the trial. Data in 186 Tables 1 and 2 provide the mean and range for each variable ( $T_c$ ,  $T_{sk}$ , heart rate, and MAP) at 30-187 minute intervals. 188

## 1.2.7 Instrumentation and Equations

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- $T_c$  was measured via insertion of a rectal thermistor (Henleys Medical Supplies, 400H/4491H, Hertfordshire, UK) 10 cm beyond the anal sphincter. The thermistor was connected via cable to a portable data logger (Libra Medical, ET402, Birmingham, UK), in which  $T_c$  was continuously displayed throughout each experimental protocol. This was only visible to the researchers, not the participants.
- Copper based thermocouples (Grant, EUS-U-VS5-0, Dorset, UK) connected to a wireless data logger (Grant, Squirrel Series, Dorset, UK) recorded  $T_{\rm sk}$  at four sites: calf, thigh, chest, and triceps [27]. Thermocouples were securely attached to the belly of each muscle by hypafix surgical adhesive tape (BSN medical, D-22771, Hamburg, Germany). The weighted  $T_{\rm sk}$  of four sites was subsequently calculated using the equation below [27]:

Mean  $T_{sk} = 0.3 \times (T_{arm} + T_{chest}) + 0.2 \times (T_{calf} + T_{thigh})$ 

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- Thermal sensation was obtained using a 0 to 8 scale ranging from unbearably cold (0) to
- unbearably hot (8). Heart rate was measured during all tests using short-range telemetry (Polar,
- FS1, Warwick, UK), and was expressed as beats per minute (b/min).
- Blood pressure was measured using a portable blood pressure monitor (Omron M5-1, Omron,
- Milton Keynes, UK). Measurements were taken at baseline (pre), and every 30 minutes of the
- 208 120-minute exposure period. Mean arterial pressure (MAP) was later calculated as  $[(2 \times DBP) +$
- 209 SBP]/3 mmHg.

## 1.2.8 Statistical Analysis

- All statistical analyses were performed using the 'nlme', 'ordinal', 'ez', 'sjPlot' and 'stats'
- packages in R version 3.3.2 (R Core Development Team 2014). Normality assumptions were
- checked using quantile-quantile plots [28] and were plausible in all instances. Central tendency
- and dispersion are reported as means  $\pm$  standard deviation (SD). The Akaike information criteria
- 215 (AIC) was used to determine model fit [29]. The correlation structure with the lowest AIC was
- chosen based on this procedure. A linear mixed model with fixed ('drug', 'time') and random
- 217 ('subject i.d') effects was fitted with an autoregressive correlation structure (to account for
- autocorrelation) to examine the effect of acetaminophen on  $T_c$ ,  $T_{sk}$ , and heart rate in thermo-
- 219 neutral and cold conditions [Time (13 levels): pre, 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 110,
- 220 120 minutes × Drug (2 levels): placebo, acetaminophen]. The same model with different levels
- of time [Time (5 levels): pre, 30, 60, 90, 120 minutes) × Drug (2 levels): placebo,
- acetaminophen] was fitted to determine the effect of acetaminophen on MAP in thermo-neutral
- and cold conditions. A cumulative link model was used to compare thermal sensation scores
- between placebo and acetaminophen in the thermo-neutral and cold conditions. The two-tailed
- alpha level of significance testing was set as  $p \le 0.05$ . 95% confidence intervals (CI) are
- presented to denote the imprecision of the point estimate.

## **1.3 RESULTS**

- 228 *1.3.1 Thermo-neutral*
- There was no main effect for drug or interaction effect (drug  $\times$  time) for  $T_c$ ,  $T_{sk}$ , heart rate, TSS,
- or MAP. A main effect for time was present in each of these variables apart from MAP, showing
- that  $T_c$ ,  $T_{sk}$ , heart rate and TSS changed (p < 0.05) over time with no differences observed
- between acetaminophen and placebo. Descriptive (mean  $\pm$  SD) data for each 30-minute interval
- is shown in Table 1.
- 234 *1.3.2 Cold*
- The  $T_c$  response during cold exposure differed between the acetaminophen and placebo
- conditions. An interaction effect ( $F_{1.12} = 2.25$ , p = 0.01), main effect for drug ( $F_{1.2} = 2.25$ , p < 0.01)
- 237 0.01), and main effect for time ( $F_{1,12} = 8.33$ , p < 0.01) was found between placebo (37.06 ±
- 238  $0.20^{\circ}\text{C}$ ; 95% CI = 36.99 to 37.12°C) and acetaminophen (36.90 ± 0.32°C; 95% CI = 36.79 to
- 239 37.01°C). Specifically,  $T_c$  was 0.18, 0.19, 0.22, 0.27, 0.29 and 0.35°C lower in the
- acetaminophen trial at time points 70 to 120 minutes compared with the placebo. The peak  $T_c$
- reduction in the nine participants (120 minute compared with baseline) was 0.16 to 0.57°C
- (mean =  $0.40 \pm 0.15$ °C). Mean and individual  $T_c$  responses over the 120-minute exposure period
- are displayed in Figures 2 and 3 respectively.
- There were no main effects for drug or interaction effect between drug and time for  $T_{\rm sk}$ , heart
- rate, TSS, or MAP. A main effect for time was present in each of these variables excluding
- MAP. All descriptive data for each 30 minute interval is shown in Table 2. For  $T_c$ , Table 3
- 247 displays the model's fixed effects coefficients and random effect variances.
- 248 \*\*\*please insert Table 1 & 2 near here\*\*\*
- \*\*\*please insert Figure 2 near here\*\*\*
- 250 \*\*\*please insert Figure 3 near here\*\*\*
- 251 \*\*\*please insert Table 3 near here\*\*\*

# 1.4 DISCUSSION

254	It was hypothesised that acetaminophen would reduce $T_c$ in cold conditions, but have no effect
255	on $T_c$ in thermo-neutral conditions relative to a placebo. The experimental hypothesis was
256	accepted. The major finding of the present study was that, compared with a placebo,
257	acetaminophen administration reduced T <sub>c</sub> (0.16 to 0.57°C decrease after 120 minutes exposure)
258	during an acute cold stress (10°C), while it appeared to have no effect on thermoregulation at a
259	thermo-neutral ambient temperature (25°C). During cold exposure, acetaminophen caused $T_{\rm c}$ to
260	fall by $\sim 0.40$ °C compared with the baseline value at 120 minutes, while it did not decline in the
261	placebo trial. The variability in the response may be due to between subject differences in the
262	rate of acetaminophen absorption, but unfortunately this was not analysed in this trial. The
263	hypothermic response to acetaminophen ingestion observed in the current study corroborates our
264	prior work in humans, in which acetaminophen reduced $T_c$ by $\sim 0.19$ °C in humans exposed to
265	mild cooling [13]. Furthermore, this is the first study to demonstrate that the ambient temperature
266	can dictate the degree of hypothermia induced by acetaminophen. During cold exposure, this
267	trial shows that healthy young adults could not defend their $T_c$ following acetaminophen
268	administration (Figure 2). Given that elderly individuals already struggle to defend their $T_{\rm c}$
269	without prior drug ingestion [20], it is reasonable to suspect that acetaminophen would cause $T_{\rm c}$
270	to decline at a faster rate, increasing the risk of accidental hypothermia.
271	The notion that ambient and skin temperature dictates the magnitude of acetaminophen's
272	hypothermic action is in line with previous research. In a recent experiment, acetaminophen (20
273	mg/kg) had no effect on sweat output and $T_c$ during 1-hour exercise in hot conditions (34°C, 52%)
274	r.h.) at a fixed rate of heat production (8 W/kg) [30]. In that study, the mean skin temperature
275	increased by 1°C during the trial (up to ~35°C), a condition in which no heat producing
276	mechanisms will be active [31, 32]. Because the mean skin temperature during cold stress was
277	~24°C at the end of the trial (Figure 2), cutaneous vasoconstriction and active thermogenesis
278	were required for $T_c$ to remain stable [33, 34]. The presence of thermogenesis and
279	vasoconstriction indicates that acetaminophen may reduce $T_{\rm c}$ through inhibition of at least one of
280	these mechanisms, but the precise mechanism needs to be confirmed in future work. Previous
281	data demonstrated that acetaminophen reduced $T_c$ by 0.10 to 0.39°C (mean $\pm$ SD, 0.19 $\pm$ 0.09°C)
282	at rest when the mean skin temperature was ~27°C [13]. Similar reductions in skin temperature

induce shivering thermogenesis [33], which, if inhibited by acetaminophen, may explain the 283 small reduction in  $T_c$  seen previously [13]. 284 Studies in mice have shown  $T_c$  fell by 0.40, 0.80, and 2°C following 1-hour acetaminophen 285 infusion of 100, 200, and 300 mg/kg body mass respectively [14]. Thus, acetaminophen-induced 286 hypothermia is not only dependent on ambient temperature, but also on the dose administered. It 287 288 is important to note here that mice are often housed in environments of 18 to 20°C, which is 8 to 10°C beneath their normal thermo-neutral zone [35]. These housing conditions are consistent in 289 290 experiments concerning acetaminophen-induced hypothermia in rodents [6, 8, 14], such that these animals constantly produce heat to maintain their  $T_c$ . Inhibition of this heat production 291 292 through acetaminophen may explain its hypothermic action, a notion that should be confirmed through the administration of high dose acetaminophen in mice housed within and below their 293 294 thermo-neutral zone (i.e. 30°C and 20°C, respectively). It is possible that the acetaminophen-induced reduction in  $T_c$  observed in the present study was 295 due to inhibition of cyclooxygenase (COX). There are two COX isoforms (COX-1 and -2), and 296 their function is to convert arachidonic acid to prostaglandin (PG) H<sub>2</sub> [36], which cell-specific 297 isomerases and synthases then convert to prostanoids [(PGE<sub>2</sub>, PGF<sub>2</sub>, PGD<sub>2</sub>, PGI<sub>2</sub>, and 298 thromboxane A<sub>2</sub> (TXA<sub>2</sub>)]. The strongest evidence that acetaminophen-induced hypothermia is 299 mediated through COX inhibition was provided by Ayoub and colleagues [14], who 300 demonstrated that acetaminophen reduced  $T_c$  by 4°C in wild-type mice, but by only 1.5°C in a 301 COX-1<sup>-/-</sup> strain. In addition, they showed a strong relationship between brain PGE<sub>2</sub> 302 concentrations and  $T_c$ , where the maximum reduction in  $T_c$  was met with a 96% reduction in 303 brain PGE<sub>2</sub>. Data supporting a role for a COX-1 splice variant (COX-1b) in the hypothermic 304 effect of acetaminophen is equivocal. While infusion of putative COX-1b inhibitors aminopyrene 305 and antipyrene exert a similar hypothermic effect to acetaminophen [14, 37], genetic studies 306 307 suggest that the human COX-1b gene produces a non-functional protein because it retains intron-1 [38]. Even when this was corrected via site-directed mutagenesis, acetaminophen did not 308 309 inhibit COX-1b activity [39]. Taken together, these studies suggest that COX-1 mediated PGE<sub>2</sub> production may be required for normal  $T_c$  maintenance in mammals housed in sub-neutral 310 311 ambient temperatures, while COX-1b is unlikely to be involved. If this were true, similar hypothermic responses would be expected with non-selective COX inhibitors Ibuprofen and 312

Aspirin, or SC-560, a COX-1 specific inhibitor. Whether these drugs initiate a loss of  $T_c$  control 313 during cold exposure has not yet been determined. 314 Given acetaminophen reduced  $T_c$  stability in healthy adult males (Figure 2), its hypothermic 315 effect is likely to be larger in populations already considered vulnerable in sub-neutral ambient 316 temperatures (i.e. the very young and the elderly). Accidental hypothermia is a rising global 317 318 health concern. In the USA, the Centre for Disease Control and Prevention report that hypothermia was the cause of nearly 17,000 deaths from 1999 to 2011 [40]. In the UK, hospital 319 320 episode statistics show that there were over 108,000 admissions to NHS hospitals from 2005 to 2014, where hypothermia was the primary or secondary cause [4]. This database also shows that 321 322 the very young (0-4 years; 43,868 admissions) and the elderly ( $\geq$  65 years; 48,477 admissions) make up 85% of the total admissions. This is concerning for two reasons. Firstly, acetaminophen 323 324 is the most frequently administered analgesic among frail and pre-frail elderly individuals [41], with no age-related delay in drug absorption [42]. Secondly, acetaminophen is commonly used 325 326 for neonatal pain management [43]. In the perioperative setting,  $T_c$  monitoring after acetaminophen administration in these vulnerable groups is recommended. A 2011 study showed 327 that intravenous acetaminophen (~20 mg/kg) did not cause hypothermia in 93 neonates [44]. 328 However, the ambient temperature was not reported (presumably 23-25°C), and only the skin 329 330 temperature was measured. This is problematic since our work showed a clear reduction in  $T_c$ without a change in skin temperature between acetaminophen and placebo [13]. Moreover, 331 neonates are exposed to cold stress when wet with amniotic fluid, during transportation, or 332 during surgery. Based on our data, we propose that acetaminophen may increase the risk of 333 neonatal hypothermia only when coupled with one of these cold stressors, and not in a thermo-334 335 neutral environment. 1.4.1 Limitations 336 This study has limitations that should be considered in future work. Firstly, we did not measure 337 metabolic heat production or cutaneous blood flow, key parameters that control  $T_c$  during cold 338 stress. Although a reduction in  $T_c$  from resting value is the primary variable of interest from a 339 medical standpoint, it is still unknown what aspect of the thermoregulatory system 340 acetaminophen targets to exert this effect. Measuring metabolic heat production and changes in 341 cutaneous blood flow in future studies of a similar design may help to elucidate the mechanism 342

that regulates acetaminophen's hypothermic action. Secondly, no pharmacokinetic parameters are reported in this experiment. Disparity in the plasma concentration of acetaminophen throughout each trial may have explained the between subject variability in the hypothermic response elicited by acetaminophen i.e. a low plasma concentration may result in a reduced hypothermic response. We administered a dose relative to lean body mass to reduce the variability in acetaminophen absorption, and our previous experiment showed that a 20 mg/kg lean body mass dose was appropriate for therapeutic plasma concentrations to be reached within the 120-minute exposure period [13].

#### 1.4.2 Conclusions

In conclusion, this preliminary trial demonstrated that acute acetaminophen ingestion (20 mg/kg lean body mass) reduced  $T_{\rm c}$  maintenance during acute cold exposure in healthy young adults. We are the first to show that the hypothermic action of acetaminophen is strongly influenced by the ambient temperature in which it is administered. Future research should determine if this effect is amplified in new-borns and in elderly individuals, placing them at risk of accidental hypothermia. It should also be determined if hypothermic effects are limited to acetaminophen, or are present in COX inhibitors such as Ibuprofen and Aspirin (non-selective COX inhibitors), or COXIBs (COX-2 selective inhibitors). If all COX inhibitors induce hypothermia during cold exposure, the prescription of these medications to individuals vulnerable to hypothermia should be carefully considered during cold spells and in the perioperative period.

**Additional information** 363 364 **Competing interests** 365 The authors declare they have no competing interests. **Author contributions** 366 JF, LT, and ARM contributed to the conception and design of the study. JF, LT, DH, AG and JH 367 contributed to data interpretation and manuscript revision. JF collected the data. All authors 368 agree to be accountable for all aspects of the work in ensuring that questions related to the 369 accuracy or integrity of any part of the work are appropriately investigated and resolved. All 370 authors approved the final version of the manuscript and all authors qualifying for authorship are 371 listed. 372 Acknowledgements 373 The authors thank Miss Katie Thomasson and Mr Jack Field for their assistance with data 374 collection. The authors also thank those who participated in this experiment. 375 376 **Funding** None declared 377 378

## Figure Captions:

- Fig 1. A flowchart of the study design. APAP = Acetaminophen. Visits 2-5 completed in a
- randomised order for each participant. Visits separated by 1-week and drug administration
- double blinded.

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- Fig 2. Mean and SD of the  $T_c$  (A, C) and  $T_{sk}$  (B, D) response during the 120-minute exposure to
- 384 25°C (left panel i.e. A, B) and 10°C (right panel i.e. C, D). The triangles and squares represent
- the placebo and acetaminophen trials, respectively. \* Main effect for condition. # Main effect for
- time. † Interaction effect. Significance set at p < 0.05.
- Fig 3. Change in  $T_c$  during cold exposure in each participant following administration of a
- placebo (A) or acetaminophen (B).

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**Table 1.** Descriptive data for each of the five response variables in the thermo-neutral condition (25°C). Descriptive data are the mean values (± standard deviation) during the 120-minute exposure period. The range is provided in parentheses.

		Time-point (minutes)				
		Pre	30	60	90	120
	D1 1	$37.00 \pm 0.13$	$36.93 \pm 0.15$	$36.95 \pm 0.15$	$36.94 \pm 0.14$	$36.94 \pm 0.16$
T (0C)	Placebo	(36.80 - 37.15)	(36.72 - 37.13)	(36.73 - 37.15)	(36.75 - 37.15)	(36.74 - 37.21)
<i>T</i> <sub>c</sub> (°C)		$37.04 \pm 0.20$	$36.95 \pm 0.22$	$36.93 \pm 0.21$	$36.91 \pm 0.23$	$36.89 \pm 0.19$
	Acetaminophen	(36.78 - 37.25)	(36.78 - 37.14)	(36.77 - 37.05)	(36.68 - 37.10)	(36.62 - 37.10)
	Distrik	$30.6 \pm 0.9$	$30.9 \pm 0.7$	$30.8 \pm 0.7$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$30.7 \pm 0.7$
<b>T</b> (0C)	Placebo	(28.7 - 31.8)	(29.9 - 31.9)	(29.8 - 31.7)	(29.5 - 31.8)	(29.3 - 31.7)
$T_{\rm sk}$ (°C)	A 4	$30.3 \pm 0.6$	$30.8 \pm 0.5$	$30.7 \pm 0.4$	$30.7 \pm 0.5$	$30.6 \pm 0.6$
	Acetaminophen	(29.0 - 31.1)	(29.9 - 31.4)	(29.9 - 31.2)	(29.9 - 31.5)	(29.6 - 31.6)
	DI 1	$65 \pm 8$	59 ± 8	58 ± 10	$58 \pm 9$	$60 \pm 10$
HD (1 / ' )	Placebo	(53 - 79)	(50 - 76)	(46 - 79)	(48 - 74)	
HR (b/min)		$68 \pm 8$	$62 \pm 10$	$65 \pm 10$		$59 \pm 10$
	Acetaminophen	(53 - 81)	(49 - 80)	(50 - 84)	(49 - 68)	(42 - 71)
		$4.0 \pm 0.1$	$4.1 \pm 0.3$	30       60       90       120 $36.93 \pm 0.15$ $36.95 \pm 0.15$ $36.94 \pm 0.14$ $36.91 \pm 0.23$ $36.89 \pm 20.23$	$4.4 \pm 0.6$	
TS (0 to 8	Placebo	(4.0 - 4.5)				
scale)		$4.0 \pm 0.2$		,	,	$4.6 \pm 0.5$
,	Acetaminophen	(3.5 - 4.5)	(4.0 - 5.0)	(4.0 - 5.0)	(4.0 - 5.0)	(4.0 - 5.0)
		91 ± 7	91 ± 9	91 ± 10	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
MAP	Placebo	(83 - 103)				
(mmHg)	Acetaminophen	$88 \pm 6$	,	` /		
` 0/		(80 - 97)				(85 - 104)

Core temperature  $(T_c)$ , Skin temperature  $(T_{sk})$ , Heart rate (HR), Thermal sensation (TS), Mean arterial pressure (MAP)

**Table 2.** Descriptive data for each of the five response variables in the cold condition (10 $^{\circ}$ C). Descriptive data are the mean values ( $\pm$  standard deviation) during the 120-minute exposure period. The range is provided in parentheses.

	Time-point (minutes)					
		Pre	30	60	90	120
T <sub>c</sub> (°C)  T <sub>sk</sub> (°C)  HR (b/min)  TS (0 to 8 scale)	DI I	$36.98 \pm 0.20$	$37.09 \pm 0.19$	$37.03 \pm 0.22$	$36.97 \pm 0.23$	$36.96 \pm 0.25$
T (0C)	Piacebo	(36.70 - 37.13)	(36.79 - 37.38)	(36.72 - 37.34)	(36.71 - 37.29)	(36.64 - 37.19)
$I_{\mathfrak{c}}({}^{\mathfrak{c}}\mathfrak{C})$	Acataminanhan	$36.97 \pm 0.21$	$37.05 \pm 0.26$	$36.94 \pm 0.31$	$36.76 \pm 0.30$ *	$36.58 \pm 0.23*$
HR (b/min)  TS (0 to 8 scale)  MAP (mmHg)	Acetaminophen	(36.61 - 37.36)	(36.59 - 37.49)	(36.52 - 37.45)	(36.33 - 37.29)	(36.11 - 36.87)
		Pre 30 60 90 12 36.98 ± 0.20 37.09 ± 0.19 37.03 ± 0.22 36.97 ± 0.23 36.96  (36.70 - 37.13) (36.79 - 37.38) (36.72 - 37.34) (36.71 - 37.29) (36.64  (36.97 ± 0.21 37.05 ± 0.26 36.94 ± 0.31 36.76 ± 0.30* 36.58 ± 0.30* 36.59 - 37.49) (36.52 - 37.45) (36.33 - 37.29) (36.11  Placebo 30.5 ± 0.5 25.8 ± 1.0 24.9 ± 1.0 24.4 ± 1.0 24.2 ± 0.4 (29.6 - 31.3) (24.7 - 27.6) (23.8 - 26.9) (23.2 - 26.5) (22.8 30.7 ± 0.7 26.1 ± 1.0 25.1 ± 1.0 24.5 ± 1.2 24.3 ± 0.2 (29.6 - 31.8) (24.7 - 28.2) (23.7 - 26.6) (23.0 - 26.3) (22.4 ± 0.4 (29.6 - 31.8) (24.7 - 28.2) (23.7 - 26.6) (23.0 - 26.3) (22.4 ± 0.4 (29.6 - 31.8) (24.7 - 28.2) (23.7 - 26.6) (23.0 - 26.3) (22.4 ± 0.4 (20.6 ± 1) (50.79) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.64) (41.70) (39.73) (42.70) (10.70) (39.73) (42.70) (10.70) (39.73) (42.70) (10.70) (39.73) (42.70) (10.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (42.70) (39.73) (39.70) (39.73) (39.70) (39.73) (39.70) (39.73) (39.70) (39.73) (39.70) (39.73) (39.70) (39.7				
	Dlasaka	$30.5 \pm 0.5$	$25.8 \pm 1.0$	$24.9 \pm 1.0$	$24.4 \pm 1.0$	$24.2 \pm 1.0$
T (0C)	Piacebo	(29.6 - 31.3)	(24.7 - 27.6)	(23.8 - 26.9)	(23.2 - 26.5)	(22.8 - 26.6)
$I_{\rm sk}$ (°C)	A aataminanhan	$30.7 \pm 0.7$	$26.1 \pm 1.0$	$25.1 \pm 1.0$	$24.5 \pm 1.2$	$24.3 \pm 1.3$
	Acetaminophen	(29.6 - 31.8)	(24.7 - 28.2)	(23.7 - 26.6)	(23.0 - 26.3)	(22.4 - 26.5)
	Dlaasha	$68 \pm 7$	$62 \pm 9$	$61 \pm 4$	$57 \pm 8$	$60 \pm 9$
HR	Flacebo	(54 - 79)	(48 - 74)	(55 - 67)	(48 - 68)	(51 - 75)
(b/min)	Acataminanhan	$66 \pm 11$	$59 \pm 9$	$58 \pm 10$	$54 \pm 7$	$57 \pm 9$
,	Acetaminophen	(50 - 79)	(41 - 70)	(39 - 73)	(42 - 64)	(41 - 70)
	Placebo					$1.8 \pm 0.4$
`		,	,		,	(1.0 - 2.0)
scale)	Acetaminophen					$1.7 \pm 0.5$
	riccummophen	(3.5 - 4.0)	(2.0 - 3.0)	(1.5 - 3.0)	(1.0 - 3.0)	(1.0 - 2.5)
		02 ± 10	07 + 0	00 + 8	07 ± 7	$105 \pm 8$
MAD	Placebo					
		` /	,	,	,	(92 - 117)
(mmHg)	Acetaminophen					$99 \pm 6$
	-	(/8 - 102)	(74 - 102)	(91 - 111)	(88 - 104)	(77 - 104)

Core temperature ( $T_c$ ), Skin temperature ( $T_{sk}$ ), Heart rate (HR), Thermal sensation (TS), Mean arterial pressure (MAP). \* denotes significant difference between the APAP and placebo condition (p < 0.05).

**Table 3.** Beta coefficients (*B*), 95 % confidence intervals (CI), alpha values (*p*), and the Phi coefficient are reported for the fixed components (drug & time) during exposure to cold stress (10°C). The standard deviation of the intercept and residual are reported for the random effect (subject ID).

	Co	Core Temperature (°C)			
	В	CI	p		
Fixed Parts			-		
Intercept	36.95	36.71 to 37.13	<.001		
Drug×Time Interaction					
DRUG:TIME10	0.03	-0.11 to 0.17	.694		
DRUG:TIME20	-0.03	-0.17 to 0.11	.672		
DRUG:TIME30	-0.06	-0.20 to 0.09	.442		
DRUG:TIME40	-0.10	-0.24 to 0.04	.179		
DRUG:TIME50	-0.12	-0.26 to 0.02	.109		
DRUG:TIME60	-0.13	-0.28 to 0.01	.076		
DRUG:TIME70	-0.18	-0.32 to -0.03	.021		
DRUG:TIME80	-0.21	-0.36 to -0.07	.006		
DRUG:TIME90	-0.24	-0.38 to -0.10	.002		
DRUG:TIME100	-0.29	-0.43 to -0.15	<.001		
DRUG:TIME110	-0.31	-0.45 to -0.17	<.001		
DRUG:TIME120	-0.36	-0.50 to -0.22	<.001		
Phi Coefficient					
0.938					
Random Parts (Subject ID)	Random Parts (Subject ID)				
		Standard Deviation			
Intercept	0.13				
Residual	0.16				