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3

4 **Hydration marker diagnostic accuracy to identify mild intracellular and extracellular dehydration**

5

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22 **Running title:** Intra and extracellular dehydration markers

23

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25

26 **Abstract**

27 Identifying mild dehydration ($\leq 2\%$ of body mass) is important to prevent the negative effects of
28 more severe dehydration on human health and performance. It is unknown whether a single
29 hydration marker can identify both mild intracellular and extracellular dehydration with adequate
30 diagnostic accuracy (≥ 0.7 receiver operating characteristic-area under the curve (ROC-AUC)). Thus, in
31 15 young healthy men, we determined the diagnostic accuracy of 15 hydration markers after three
32 randomized 48-h trials; euhydration (EU, water $36 \text{ ml}\cdot\text{kg}\cdot\text{d}^{-1}$), intracellular dehydration caused by
33 exercise and 48 h of fluid restriction (ID, water $2 \text{ ml}\cdot\text{kg}\cdot\text{d}^{-1}$), and extracellular dehydration caused by
34 a 4 h diuretic-induced diuresis, begun at 44 h (ED, Furosemide $0.65 \text{ mg}\cdot\text{kg}^{-1}$). Body mass was
35 maintained on EU and dehydration was mild on ID and ED ($1.9 (0.5)\%$ and $2.0 (0.3)\%$ of body mass,
36 respectively). Urine color, urine specific gravity, plasma osmolality, saliva flow rate, saliva osmolality,
37 heart rate variability and dry mouth identified ID (ROC-AUC; range 0.70-0.99) and postural heart rate
38 change identified ED (ROC-AUC 0.82). Thirst 0-9 scale (ROC-AUC 0.97 and 0.78 for ID and ED) and
39 urine osmolality (ROC-AUC 0.99 and 0.81 for ID and ED) identified both dehydration types. However,
40 only thirst 0-9 scale had a common dehydration threshold (≥ 4 ; sensitivity and specificity of 100%,
41 87% and 71%, 87% for ID and ED). In conclusion, using a common dehydration threshold ≥ 4 , the
42 thirst 0-9 scale identified mild intracellular and extracellular dehydration with adequate diagnostic
43 accuracy. In young healthy adults' thirst 0-9 scale is a valid and practical dehydration-screening tool.

44

45 **Keywords:** hypohydration, thirst, urine, plasma, saliva, tear, ROC curve.

46 **Introduction**

47 No consensus currently exists on the best method to assess dehydration and prescribe fluid intake
48 (Armstrong, 2007; Cheuvront & Kenefick, 2014; Cotter et al., 2014). This is in part because
49 dehydration is a complex condition that manifests as different types. When fluid intake is
50 inadequate, and the concentration of body fluids lost is hypoosmotic relative to plasma (e.g. exercise
51 sweat loss), the body fluid redistribution that occurs results in a relatively larger loss of intracellular
52 than extracellular fluid (Sawka, 1992). Consequently, this type of dehydration is referred to as
53 intracellular dehydration and characterized by an increased plasma osmolality (hyperosmolality). In
54 contrast, extracellular dehydration, is caused by iso-osmotic fluid loss and is characterized by volume
55 depletion (hypovolemia) and the absence of hyperosmolality. Extracellular dehydration often occurs
56 when people are ill, take medications (e.g. diuretics), are immersed in water, or exposed to cold
57 and/or hypoxia (Cheuvront & Kenefick, 2014; Cotter et al., 2014). Whether hydration markers
58 identify intracellular or extracellular dehydration is likely to depend on the relationship between the
59 marker and the distinct physiological characteristics of each dehydration type.

60

61 Potential candidate markers to identify both types of dehydration are urine, saliva, ratings of thirst
62 and cardiovascular parameters, including resting and postural changes in heart rate and blood
63 pressure, and heart rate variability (HRV) (Cheuvront et al., 2012; Cotter et al., 2014; Fitzsimons,
64 1976; Oliver et al., 2008). These markers may respond directly to osmotic and volume stimuli, or
65 indirectly to the subsequent alterations in autonomic tone (Charkoudian et al. 2005, Oliver et al.
66 2008, Sands & Layton 2009). While most of these hydration markers have shown promise to identify
67 moderate and severe intracellular dehydration (>3% body mass; Armstrong et al. 1994, 2014, Walsh
68 et al. 2004, Cheuvront et al. 2012), limited research has investigated the validity and diagnostic
69 accuracy of these hydration markers to identify more mild extracellular or intracellular dehydration
70 ($\leq 2\%$ of body mass). Mild dehydration is important to identify, as it is beyond this threshold that

71 human performance has been consistently shown to decline (Cheuvront & Kenefick, 2014; Goulet,
72 2012; Savoie et al., 2015).

73

74 The aim of this study was therefore to determine hydration marker diagnostic accuracy to identify
75 mild intracellular and extracellular dehydration. Based on previous research examining hydration
76 markers after moderate and severe dehydration (Cheuvront et al., 2012; Fortes et al., 2011; Oliver et
77 al., 2008; Shirreffs et al., 2004), we hypothesized that urine, thirst, dry mouth, saliva and HRV
78 markers would identify both types of mild dehydration with adequate diagnostic accuracy (ROC-AUC
79 ≥ 0.7 ; Hooper et al. 2016). Based on this research we also hypothesized that plasma osmolality and
80 tear osmolarity would identify mild intracellular dehydration, but not mild extracellular dehydration;
81 and postural heart rate and blood pressure change would identify extracellular dehydration, but not
82 intracellular dehydration.

83

84 **Materials and Methods**

85 ***Participants***

86 Fifteen healthy males volunteered to complete the study (age 22.8 (5.4) years, height 180.4 (5.0) cm,
87 mass 78.9 (8.6) kg, BMI 24.2 (1.8) $\text{kg}\cdot\text{m}^{-2}$, VO_2max 52.3 (6.9) $\text{ml}\cdot\text{kg}^{-1}\cdot\text{min}^{-1}$). Participants were
88 excluded if they were, smokers, had abnormal blood chemistry or renal function, suffered from
89 diabetes, asthma, bronchitis, epilepsy, hypertension, dental or oral disease or were receiving any
90 medication or treatment. Informed written consent was obtained from each participant. The study
91 was approved by the Institutional Ethics Committee and adhered to the Declaration of Helsinki.

92

93 ***Preliminary measures***

94 As body mass loss during the 48-h trials was the reference standard in this study, we standardized
95 energy intake and physical activity 24 h before and during trials. Energy intake was calculated as the
96 product of resting metabolic rate and an estimated physical activity factor. Resting metabolic rate

97 was estimated from anthropometry (Harris & Benedict, 1918) and adjusted by a general daily
98 physical activity and diet induced thermogenesis factor coefficient of 1.6, which was determined
99 from the activities completed on trials (Todorovic & Micklewright 2004). Participants were also
100 habituated with the hydration assessment techniques and completed a graded cycle exercise test to
101 determine their peak power output, which was used to prescribe the workload for the experimental
102 trial cycling exercise (Excalibur Sport, Lode, Netherlands).

103

104 ***Study protocol***

105 The study followed a crossover design. Separated by seven days, participants completed three trials
106 in a random order including a euhydrated control trial (EU), an intracellular dehydration (ID) trial,
107 and an extracellular dehydration (ED) trial. Each trial consisted of a baseline hydration assessment,
108 an exercise bout, one of the three 48-h interventions, and a second hydration assessment (Figure 1).
109 Hydration assessments and exercise was performed in an air-conditioned laboratory, temperature
110 and humidity, 19.4 (1.0) °C and 42 (6)%, respectively.

111

112 The day before each experimental trial participants abstained from alcohol, caffeine or strenuous
113 physical activity and consumed a standardized individually prescribed diet (energy and sodium
114 intake 3034 (245) kcal and 2.2 (0.1) g; 62%, 25%, 13% carbohydrate, fats and protein, respectively).
115 Daily energy intake was the same for the duration of the trials except on day one participants
116 consumed additional food (391 (193) kcal) to replace energy expended during the cycling exercise.
117 This was calculated from indirect calorimetry during the habituation visit cycling exercise test (Cortex
118 MetaLyzer 3B, Germany).

119

120 On day one of each trial participants woke at 07:00 h and drank water equal to 6 ml·kg⁻¹ of body
121 mass (471 (52) ml). On arrival to the laboratory at 08:00 h participants received a further bolus of
122 water equal to 6 ml·kg⁻¹ of body mass and a standardized breakfast (690 kcal, sodium 0.8 (0.1) g;

123 62%, 23% and 15% carbohydrate, fat and protein, respectively). To monitor and standardize physical
124 activity on the trial's participants were fitted with pedometers and provided with step count targets
125 (Digi-Walker SW200, Yamax, Japan). At 12:00 h participants returned to the laboratory for the
126 baseline hydration assessment. Immediately after, dehydration was induced via cycling exercise at
127 70% peak power output until exhaustion. After the cycling exercise, the participants began one of
128 three 48-h trials. The calculated sweat loss from the cycling exercise was replaced with water on EU
129 and ED but not on ID. Drinking water was restricted on ID to 2 ml·kg⁻¹ of body mass per day (total
130 314 (35) ml). In contrast, on EU and EH participants drank water equal to 36 ml·kg⁻¹ of body mass per
131 day (total for 48 h 5728 (600) ml). This fluid intake strategy was adapted from those previously used
132 in our laboratory to maintain euhydration (Oliver et al., 2007; 2008; Walsh et al., 2004). On day
133 three, participants reported to the laboratory at 07:30 h. At 08:00 h, and after a standardized
134 breakfast, on EH participants consumed the diuretic Furosemide as a liquid equal to 0.65 mg·kg⁻¹ (51
135 (6) mg Frusol, Rosemount Pharma, UK). All urine voided between 08:00 h and 12:00 h was collected
136 to measure total urine volume. At 12:00 h on all trial's participants began the hydration assessment
137 2.

138

139 **Hydration assessments**

140 Hydration markers were obtained in the same order on each trial and at each hydration assessment.
141 First, participants completed subjective ratings of thirst and dry mouth on 100 mm visual analogue
142 scale (VAS), and the 0-9 thirst sensation scale (0 = "not-at-all" to 9 = "severe"; Engell et al. 1987).
143 Participants were instructed to respond to the scale based on how they felt at that moment. Second,
144 a urine sample was collected in a container and immediately analyzed for urine color by an 8-point
145 chart (Armstrong et al., 1994), urine specific gravity (USG) was measured in duplicate using a
146 handheld refractometer (Atago, Japan) and urine osmolality was measured in triplicate by a freezing
147 point depression osmometer (Model 3300, Advanced Instruments, USA). Third, nude body mass was
148 determined to the nearest 50 g using a digital platform scale (Model 705 Seca, Germany). Fourth,

149 participants were fitted with a heart rate monitor (Polar RS800, Finland), after 2 min of seated rest,
150 beat-to-beat heart rate was recorded for 10 min for the determination of HRV (Marek, 1996). All R–R
151 series were extracted with a processing program (Polar Precision Performance, Polar Electro,
152 Finland) and analyzed in the time and frequency-domain after automatic removal of occasional
153 ectopic beats (Kubios, BSAMIG, Finland). Fifth, the participants sat quietly for 5 min before a tear
154 fluid sample was analyzed for tear osmolality from the right eye as previously described (Fortes et
155 al. 2011, TearLab™ Osmolarity System, USA). Sixth, after 5 min supine rest, blood pressure and heart
156 rate were recorded (Tango, SunTech Medical Ltd, USA). These measures were then repeated after
157 exactly 1 min of standing for the determination of postural change measures of blood pressure and
158 heart rate calculated as the difference between lying and standing measures. Seventh, a seated 5
159 min unstimulated saliva sample was collected for the determination of saliva flow rate and
160 osmolality as previously described (Oliver et al., 2008). Finally, after 10 min seated rest, a venous
161 blood sample was collected by venipuncture without venostasis into a vacutainer tube containing
162 lithium heparin (Becton Dickinson, UK). This blood was immediately used to determine, in triplicate,
163 hematocrit (packed cell volume) by microcentrifugation (Hawksley and Sons Ltd., Sussex, UK) and
164 hemoglobin by automated analyzer (B-Hemoglobin, Hemocue, Sweden). Plasma volume change was
165 then estimated from the change in hemoglobin and hematocrit values between hydration
166 assessment 1 and 2 (Dill & Costill, 1974; Strauss et al., 1951). The remaining blood was centrifuged at
167 1500 g for 10 min at 5 °C and plasma was analyzed for osmolality in triplicate. If any of the intra-
168 sample osmolalities differed by more than 1% a further sample was measured and the mean of the
169 four samples was used.

170

171 **Statistical analysis**

172 Hydration marker diagnostic accuracy to identify mild ID and ED was determined from hydration
173 assessment 2 data by ROC-AUC with 95% CIs (MedCalc Software bvba, Belgium) as recommended
174 (Zweig & Campbell, 1993). Body mass change was used as the mild dehydration reference standard

175 as it is a precise measure of body fluid change in controlled laboratory studies (Cheuvront et al.,
176 2010; Oliver et al., 2008). Body mass loss was calculated on all trials to ensure euhydration was
177 maintained on EU and mild dehydration was achieved on ID and ED. A 1% threshold was used as this
178 has previously been reported as the typical day-to-day variability of body mass in active men
179 (Cheuvront et al., 2010). Hydration markers were also given a qualitative ROC-AUC descriptor that
180 relates to the quantitative diagnostic accuracy statistic as poor (0.6), adequate (0.7), moderate (0.8),
181 high (0.9), near perfect (0.95) and perfect (1.0) (Obuchowski et al., 2004). For hydration markers to
182 be considered to have adequate diagnostic accuracy it has also previously been specified that ROC-
183 AUC should be ≥ 0.7 (Hooper et al., 2016). A value of 0.5 indicates that a hydration marker has no
184 better ability than chance to discriminate between euhydration and dehydration whereas 1.0
185 indicates that the marker has perfect discrimination (Zweig & Campbell, 1993). A sample size of 15
186 was selected, to allow for drop-out, and based on a balanced design (i.e. equal numbers of
187 participants with and without dehydration) that indicated a sample size of 14 was sufficient to
188 enable a marker with a diagnostic accuracy of ≥ 0.7 to be statistically discriminated from 0.5, i.e. no
189 better than chance. For hydration markers with adequate diagnostic accuracy (≥ 0.7) a secondary
190 analysis was performed where the Youden Index was used to generate an objective mild
191 dehydration threshold (Schisterman et al., 2005). Hydration markers at the hydration assessments
192 were also compared between trials by one-way analysis of variance (ANOVA) with planned multiple
193 comparisons by Tukeys (GraphPad Prism version 6.0, USA). Unless stated all values are mean (SD)
194 and statistical significance was accepted at $P < 0.05$.

195

196 **RESULTS**

197 **Hydration assessment 1 and trial physical activity**

198 Standardization of pre-trial fluid and energy intake was successful as indicated by consistent
199 euhydrated hydration status at hydration assessment 1 (CON, ID and ED: plasma osmolality 287 (4),
200 289 (5), 287 (3) mOsm·kg⁻¹, $P=0.10$; urine specific gravity 1.009 (0.004), 1.009 (0.004), 1.007 (0.003)

201 $\text{g}\cdot\text{ml}^{-1}$, $P=0.34$; body mass 78.4 (8.4), 78.3 (8.3), 78.4 (8.7) kg, $P=0.89$; coefficient of variation for
202 plasma osmolality, urine specific gravity and body mass were 1.0%, 0.3% and 0.6%, respectively).
203 Also similar on all trials was the cycling exercise time and sweat loss (CON, ID and ED: time to
204 exhaustion 1200 (377), 1339 (415), 1323 (431) s, $P=0.15$; sweat loss 470 (200), 540 (150), 590 (200)
205 ml, $P=0.10$) and trial physical activity (CON, ID and ED: 15299 (4172), 17182 (5106), 17982 (4625)
206 steps $\cdot\text{trial}^{-1}$, $P=0.08$).

207

208 **Hydration assessment 2**

209 Body mass, plasma osmolality and volume were stable during EU confirming euhydration and
210 supporting that the decreased body mass on ID and ED represents mild dehydration and not an
211 energy deficit (Table 1, $P<0.001$). Intracellular dehydration was confirmed on ID by increased plasma
212 osmolality (Table 1). Extracellular dehydration was confirmed on ED by decreased plasma volume
213 without a change in plasma osmolality (Table 1). Further, after the diuretic on ED urine production
214 was increased compared to EU and ID as expected (1677 (338) vs. 772 (311) and 138 (54) ml,
215 $P<0.001$). Increased urine production on ED ceased before hydration assessment 2 as indicated by a
216 similar urine volume on all trials at hydration assessment 2 (Mean (SD) CON, ID and ED: 143 (110), 97
217 (57), 189 (120) ml, $P=0.13$). Compared to EU, the HRV index LF/HF ratio was increased after ID but
218 not ED (Table 1). Further cardiovascular and renal differences between ID and ED, and the
219 descriptive statistics for other hydration markers studied for diagnostic accuracy are outlined in
220 Table 2.

221

222 **Hydration marker diagnostic accuracy**

223 Thirst 0-9 and urine osmolality had adequate diagnostic accuracy to identify both mild intracellular
224 and extracellular dehydration (Table 3). The diagnostic accuracy of these markers was near perfect
225 to identify mild intracellular dehydration and moderate for mild extracellular dehydration. For thirst
226 0-9, the Youden index derived the same threshold for both mild intracellular and extracellular

227 dehydration (≥ 4). The sensitivity and specificity of this threshold was 100% and 87% for ID and 71%
228 and 87% for ED (Table 3). For urine osmolality, the Youden index derived two different thresholds
229 depending on the type of dehydration (Table 4).

230

231 Several other hydration markers identified mild intracellular dehydration with adequate diagnostic
232 accuracy (ROC-AUC ≥ 0.7 , Table 3). The discriminatory accuracy was perfect for urine markers (color
233 and specific gravity), near perfect for plasma osmolality, high for thirst (VAS) and dry mouth (VAS)
234 and adequate for heart rate variability, saliva flow rate and osmolality. The mild intracellular
235 dehydration thresholds for these hydration markers and their sensitivity and specificity to identify
236 mild intracellular dehydration are shown in Table 4. In addition to thirst 0-9 scale and urine
237 osmolality, postural change in heart rate was the only other hydration marker to identify mild
238 extracellular dehydration with adequate diagnostic accuracy (ROC-AUC ≥ 0.7).

239

240 **DISCUSSION**

241 This study extends current hydration marker understanding by using diagnostic accuracy statistics to
242 evaluate several markers' validity to identify mild intracellular and extracellular dehydration. A
243 particular strength of this study is the standardization of energy intake and physical activity during
244 the experimental trials, which alongside the maintenance of body mass within typical day-to-day
245 variation (Cheuvront et al., 2010) on the euhydrated control trial, provides confidence that individual
246 participant body mass losses on ID and ED represent mild fluid rather than energy deficits. The
247 primary finding of this study is that thirst 0-9 and urine osmolality were the only hydration markers
248 with adequate diagnostic accuracy to identify both mild intracellular and extracellular dehydration,
249 caused by exercise and 48 h of fluid restriction and a 4 h diuretic-induced diuresis, respectively.
250 However, thirst 0-9 was the only marker with a common dehydration threshold to identify mild
251 intracellular and extracellular dehydration (≥ 4 for ID and ED, Table 4).

252

253 Notably, the present study is the first to determine the validity of thirst ratings using diagnostic
254 accuracy statistics (Table 3). As hypothesized, thirst had adequate diagnostic accuracy to identify
255 both types of mild dehydration, which may be expected as it is the major homeostatic effector
256 mechanism for restoring euhydration. Further, that thirst identified both intracellular and
257 extracellular dehydration, is in agreement with known physiological regulators whereby thirst is
258 sensitive to changes in both osmotic and volume stimuli (Fitzsimons, 1976). Osmolality is the
259 principal thirst regulator (Cheuvront & Kenefick, 2014) and this may explain the better diagnostic
260 accuracy of thirst to identify intracellular dehydration than extracellular dehydration in this study
261 (Table 3). Indeed, plasma osmolality was increased by 3.5% after intracellular dehydration, which
262 exceeds the reported 2% osmotic threshold of thirst (Table 1, Zerbe & Robertson 1983). The blood
263 volume reduction is the most likely stimuli for the increase in thirst after mild extracellular
264 dehydration as other thirst regulators plasma osmolality, dry mouth and saliva flow rate were similar
265 after the ED and EU control trials.

266

267 In agreement with our hypothesis, plasma osmolality, saliva flow rate and osmolality, dry mouth,
268 urine markers and HRV showed adequate diagnostic accuracy to identify mild intracellular
269 dehydration, whilst postural change in heart rate showed adequate diagnostic accuracy to identify
270 mild extracellular dehydration (Table 3). The diagnostic accuracy of these markers compares
271 favorably to that previously reported after more severe dehydration (ROC-AUC range, 0.89-0.98;
272 Bartok et al. 2004, Cheuvront et al. 2010, 2012, Armstrong et al. 2014). Identifying milder
273 dehydration with similar diagnostic accuracy is practically advantageous. Contrary to our hypothesis,
274 tear osmolarity did not identify intracellular dehydration and saliva osmolality, HRV and postural
275 blood pressure change did not identify extracellular dehydration with adequate diagnostic accuracy.
276 The reason for the poorer than anticipated diagnostic accuracy in these markers compared to
277 previous studies (equivalent to $\geq 3\%$ of body mass; Oliver et al. 2008, Fortes et al. 2011, Ely et al.
278 2014) may relate to the smaller fluid-deficit and osmotic, volume and autonomic nervous system

279 (ANS) alterations. In addition, our HRV results highlight that ANS alterations, when compared with
280 euhydration, may be greater after intracellular than extracellular dehydration of the same
281 magnitude (Table 1; $P=0.04$ CON vs ID; $P=0.14$ CON vs ED). Given the postulated role of ANS system
282 in saliva control (Oliver et al. 2008) this may explain why saliva parameters' diagnostic accuracy was
283 adequate to identify ID but not ED.

284

285 As thirst 0-9 and urine osmolality were the only markers to identify mild intracellular and
286 extracellular dehydration with adequate diagnostic accuracy, they might be considered the most
287 suitable to identify persons that require simple oral rehydration to prevent the negative
288 consequences of more severe dehydration to performance. Practically, thirst 0-9 has some
289 additional advantages to urine osmolality. This includes a common threshold to identify mild
290 dehydration regardless of the dehydration type. Further, thirst can be assessed instantly, and is easy
291 to assess repeatedly, which could be particularly useful to help guide daily fluid intake, and
292 rehydration from exercise, with persons aiming to achieve thirst ratings below or equal to 4. Urine
293 osmolality in contrast has a lengthy collection and analysis process that requires the collection of a
294 urine sample, which is not always possible, and specialist laboratory analysis. We therefore
295 recommend that the thirst 0-9 scale is used as the initial screening tool to identify mild dehydration,
296 and where determining the type of dehydration is important, plasma osmolality and postural change
297 in heart rate are used to confirm if the dehydration is intracellular or extracellular, respectively.

298

299 Our hydration marker findings should be considered carefully within the context they were
300 obtained, i.e. dehydration methods used, environmental conditions and population studied. Urine
301 volume at the second hydration assessment was similar and suggests overall fluid balance was stable
302 at the time when hydration marker diagnostic accuracy was determined. However, the time to mild
303 dehydration was much longer on ID than ED (48 h ID and 4 h ED), and consequently, fluid
304 redistribution between body fluid compartments may have been more complete after ID than ED

305 (Sawka, 1992). As extracellular dehydration is typically acute, e.g. when people are ill, take
306 medications (e.g. diuretics), are immersed in water, or exposed to cold and/or hypoxia, it is a
307 practical strength of this study that we determined hydration marker diagnostic accuracy after acute
308 rather than chronic extracellular dehydration. In contrast, intracellular dehydration may occur
309 chronically, as in this study, or acutely, e.g. sweating from passive heating and/or exercise sweat. As
310 these different dehydration methods may influence fluid regulation and redistribution (Sawka,
311 1992), and hydration marker diagnostic accuracy, future studies are warranted comparing the
312 diagnostic accuracy of hydration markers to identify different dehydration methods, particularly that
313 occur across different time courses. As in the present study, these future studies would benefit from
314 measuring fluid compartments to confirm fluid redistribution by isotope or dye tracer techniques
315 (e.g. bromide, Evans blue). Given the potential of thirst as a practical hydration marker, studies are
316 needed to compare the diagnostic accuracy of thirst to identify acute and chronic mild intracellular
317 dehydration. These studies are important as causes of acute intracellular dehydration including
318 exercise, and exposure to hot and dry environments may alter thirst independently of dehydration
319 due to direct effects of high ventilation, heat and drying of the oral cavity. Future studies should also
320 determine the diagnostic accuracy of thirst in other populations e.g. females, children and the
321 elderly. In the elderly, the diagnostic accuracy of thirst may be poorer than in young healthy adults
322 as ageing and disease impair kidney and saliva gland function; in addition, the elderly are more likely
323 to take medications that induce dry mouth which may alter thirst independently of dehydration
324 (Kenney & Chiu, 2001; Scully, 2003). Further, elderly persons with dementia and young children may
325 not interpret the thirst scale as young healthy adults.

326

327 In conclusion, thirst 0-9 scale was the only hydration marker, with a common dehydration threshold,
328 to identify both mild intracellular and extracellular dehydration with adequate diagnostic accuracy in
329 young healthy males, residing in a thermoneutral environment. The practical utility of thirst is

330 reinforced because it is a free and simple to use hydration marker that could also guide fluid intake
331 to maintain euhydration.

332

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339

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Table 1. Characterization of experimental hydration status after mild intracellular and extracellular dehydration

	Euhydration (EU)	Intracellular dehydration (ID)	Extracellular dehydration (ED)
Body mass change (%)	0.0 (0.6)	-1.9 (0.5) **	-2.0 (0.3) **
Body mass change range (%)	+0.9 to -0.7	-1.2 to -2.9	-1.5 to -2.5
Body mass change (kg)	0.0 (0.5)	-1.5 (0.5) **	-1.6 (0.3) **
Blood volume change (%)	0.8 (4.7)	0.0 (4.3)	-3.5 (2.8) ‡
Plasma volume change (%)	1.7 (6.2)	-0.3 (5.7)	-6.6 (4.0) ‡‡
Plasma osmolality (mOsm·kg ⁻¹)	287 (4)	297 (7) ††	286 (5)
HRV (LF/HF ratio)	1.8 (1.1)	3.4 (2.2) *	2.9 (2.1)

Note: HRV, Heart rate variability; LF/HF ratio, low-to-high frequency heart rate variability power ratio. Values represent mean (SD). Post hoc test differences indicated by * $P < 0.05$ vs. EU, ** $P < 0.01$ vs. EU, †† $P < 0.01$ vs. EU and ED, ‡ $P < 0.05$ vs. EU and ID, ‡‡ $P < 0.01$ vs. EU and ID.

Table 2. Hydration markers after mild intracellular and extracellular dehydration

	Euhydration (EU)	Intracellular dehydration (ID)	Extracellular dehydration (ED)
Thirst (0-9)	3 (1)	6 (1) ++	4 (1) **
Thirst (VAS)	33 (19)	69 (17) †	43 (17)
Dry mouth (VAS)	27 (17)	60 (21) ++	36 (12)
Urine osmolality (mOsm·kg ⁻¹)	267 (138)	1054 (127) ++	402 (110) ‡
Urine specific gravity (g·ml ⁻¹)	1.008 (0.004)	1.028 (0.005) ++	1.010 (0.004)
Urine colour (1-8)	2 (1)	6 (1) ++	2 (1)
Saliva flow rate (μL·min ⁻¹)	365 (241)	196 (165) †	425 (321)
Saliva osmolality (mOsm·kg ⁻¹)	56 (12)	64 (13) †	55 (12)
Tear osmolality (mOsm·l ⁻¹)	296 (12)	300 (11)	292 (12)
Postural change in HR (b·min ⁻¹)	14 (8)	19 (10)	26 (12) ‡
Postural change in SBP (mmHg)	8 (12)	4 (14)	0 (9)
Supine HR (b·min ⁻¹)	56 (10)	56 (12)	57 (15)
Supine SBP (mmHg)	112 (8)	111 (10)	108 (10)

Note: HR, heart rate; SBP, systolic blood pressure. Values represent mean (SD). Post hoc test differences indicated by * $P < 0.05$ vs. EU, ** $P < 0.01$ vs. EU, † $P < 0.05$ vs. EU and ED, ++ $P < 0.01$ vs. EU and ED, ‡ $P < 0.05$ vs. EU and ID.

Table 3. Diagnostic accuracy of hydration markers to identify mild intracellular and extracellular dehydration

Hydration marker	Intracellular dehydration (ID)			Extracellular dehydration (ED)		
	ROC-AUC	95% CI	SE	ROC-AUC	95% CI	SE
1. Urine osmolality (mOsm·kg ⁻¹)	0.99*	0.88-0.99	0.01	0.81*	0.63-0.93	0.09
2. Thirst (0-9)	0.97*	0.84-0.99	0.02	0.78*	0.59-0.90	0.08
3. Urine specific gravity (g·ml ⁻¹)	0.99*	0.88-0.99	0.01	0.68	0.48-0.83	0.10
4. Thirst (VAS)	0.92*	0.76-0.98	0.04	0.66	0.47-0.83	0.10
5. Dry mouth (VAS)	0.88*	0.69-0.97	0.06	0.66	0.47-0.83	0.10
6. Urine colour (1-8)	0.99*	0.88-0.99	0.01	0.52	0.33-0.70	0.11
7. Plasma osmolality (mOsm·kg ⁻¹)	0.96*	0.82-0.99	0.03	0.53	0.34-0.71	0.11
8. Postural change in HR (b·min ⁻¹)	0.66	0.47-0.82	0.10	0.82*	0.64-0.93	0.08
9. HRV (LF/HF ratio)	0.72*	0.52-0.87	0.09	0.64	0.45-0.81	0.11
10. Saliva osmolality (mOsm·kg ⁻¹)	0.70*	0.51-0.85	0.09	0.55	0.36-0.73	0.11
11. Saliva flow rate (μl·min ⁻¹)	0.70*	0.51-0.85	0.09	0.55	0.36-0.73	0.11
12. Tear osmolality (mOsm·l ⁻¹)	0.61	0.41-0.78	0.11	0.61	0.42-0.82	0.11
13. Postural change in SBP (mmHg)	0.56	0.37-0.74	0.11	0.65	0.46-0.82	0.10
14. Supine SBP (mmHg)	0.56	0.37-0.74	0.11	0.64	0.44-0.80	0.11
15. Supine HR (b·min ⁻¹)	0.53	0.34-0.72	0.11	0.52	0.33-0.70	0.11

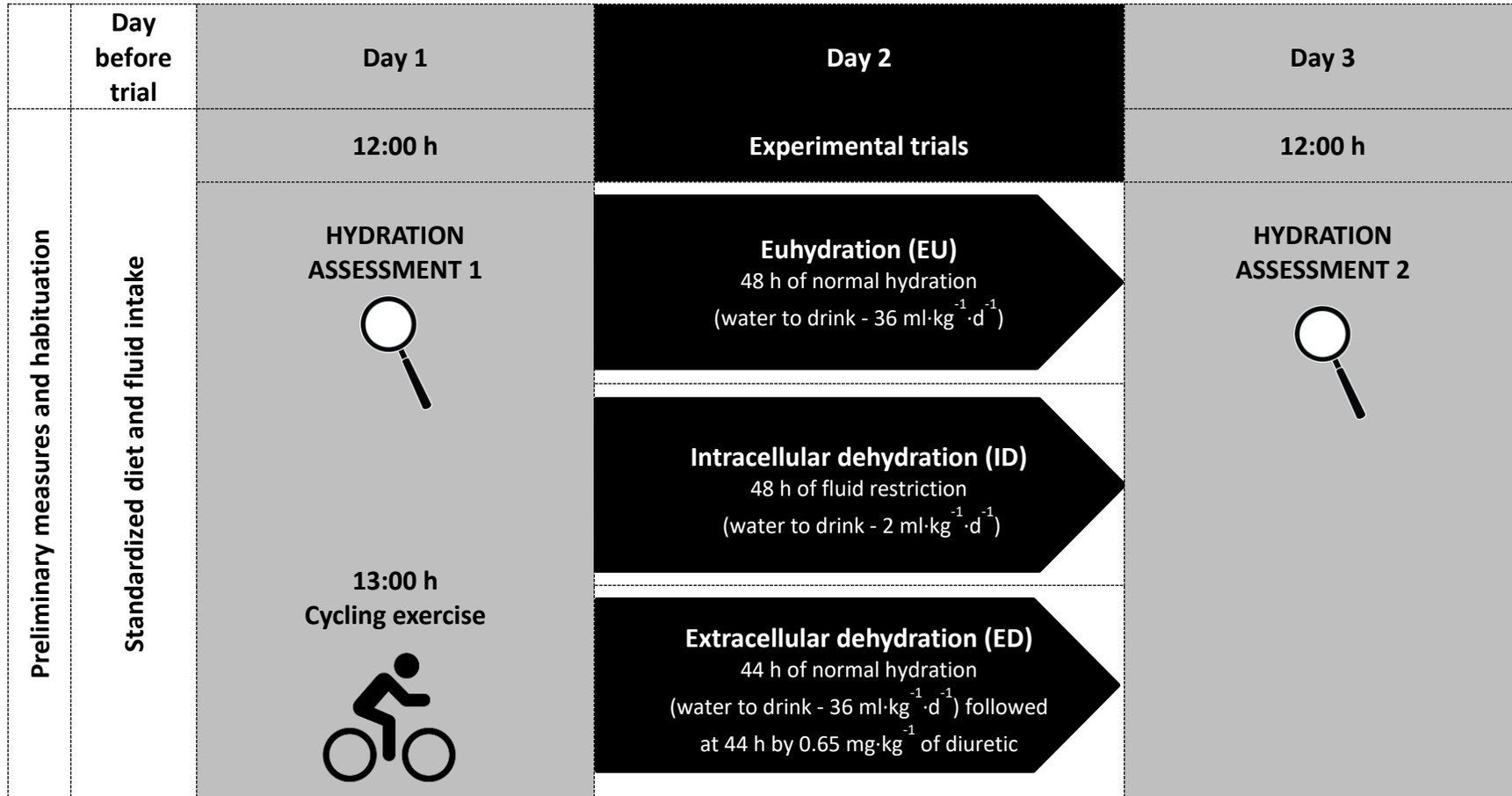
Note: HRV, Heart rate variability; LF/HF ratio, low-to-high frequency heart rate variability power ratio; ROC, receiver operating characteristic; ROC AUC, area under the ROC curve; CI, binomial exact confidence interval for AUC; SE, standard error (Hanley & McNeil, 1982); * indicates that the hydration biomarker identifies dehydration type better than chance. Note: hydration markers are ranked by combined diagnostic accuracy.

Table 4. Sensitivity and specificity of Youden derived mild dehydration thresholds for hydration markers

Hydration marker	Intracellular dehydration (ID)			Extracellular dehydration (ED)		
	Mild Dehydration Threshold ^b	Sensitivity (%)	Specificity (%)	Mild Dehydration Threshold ^b	Sensitivity (%)	Specificity (%)
Urine Osmolality (mOsm·kg ⁻¹)	>595	99	99	>341	80	87
Thirst (0-9)	≥4	99	87	≥4	71	87
Urine specific gravity (g·ml ⁻¹)	>1.016	99	99	No	-	-
Thirst (VAS)	>47	93	80	No	-	-
Dry mouth (VAS)	>40	79	80	No	-	-
Urine colour (1-8)	≥4	99	99	No	-	-
Plasma osmolality (mOsm·kg ⁻¹)	≥291	93	87	No	-	-
Postural change in HR (b·min ⁻¹)	No	-	-	>14	93	60
Saliva osmolality (mOsm·kg ⁻¹)	≥57	73	67	No	-	-
Saliva flow rate (μl·min ⁻¹)	≤137	67	67	No	-	-
HRV (LF/HF ratio)	>2.8	57	93	No	-	-
Tear osmolality (mOsm·l ⁻¹)	No	-	-	No	-	-
Postural change in SBP (mmHg)	No	-	-	No	-	-
Supine HR (b·min ⁻¹)	No	-	-	No	-	-
Supine SBP (mmHg)	No	-	-	No	-	-

Note: HR, heart rate; SBP, systolic blood pressure; HRV, Heart rate variability; LF/HF ratio, low-to-high frequency heart rate variability power ratio. ^bYouden derived mild dehydration threshold, where ROC-AUC ≥0.70.

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Figure 1. Schematic representation of experimental trial. The cycling exercise intensity was 70% of peak power output until exhaustion. Hydration assessments and exercise was performed in an air-conditioned laboratory, temperature and humidity, 19.4 (1.0) °C and 42 (6)%, respectively.