Water intake after dehydration makes muscles more susceptible to cramp but electrolytes reverse that effect

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**Recommended Citation**

Lau, W. Y., Kato, H., & Nosaka, K. (2019). Water intake after dehydration makes muscles more susceptible to cramp but electrolytes reverse that effect. DOI: [https://doi.org/10.1136/bmjsem-2018-000478](https://doi.org/10.1136/bmjsem-2018-000478)

[Lau, W. Y., Kato, H., & Nosaka, K. (2019). Water intake after dehydration makes muscles more susceptible to cramp but electrolytes reverse that effect. BMJ Open Sport and Exercise Medicine, 5(1). Available here.](https://ro.ecu.edu.au/ecuworkspost2013/6047)
ABSTRACT

Objective  No previous study has compared water and oral rehydration solution (ORS) intake after dehydration induced by exercise in the heat for the effect on muscle cramps. The present study tested the hypothesis that water ingestion after dehydration would increase muscle cramp susceptibility, but this would be prevented by ORS ingestion.

Methods  Ten men performed two bouts of downhill running (DHR; −5%) in the heat (35°C–36°C) until their body mass was reduced by 2%. Ten minutes after DHR, either spring water or electrolyte water similar to ORS (OS-1®) was ingested in a counter-balanced order on two different days separated by a week. Muscle cramp susceptibility was assessed by a threshold frequency (TF) of electrical train stimulation to induce cramp before, immediately after (0), and 30 and 60 min after the ingestion. Blood samples were taken before, immediately and 80 min after DHR to measure serum electrolyte concentrations.

Results  Muscle cramp susceptibility assessed by TF did not change from baseline to immediately after DHR for both conditions (water: 24.6 ± 2.1 Hz, OS-1®: 24.7 ± 1.4 Hz). TF decreased after water intake by 4.3 Hz (30 min) and 5.1 Hz (60 min post-ingestion), but increased after OS-1® intake by 3.7 and 5.4 Hz, respectively. Serum sodium and chloride concentrations decreased after water intake but maintained after OS-1® intake.

Conclusion  These results suggest that water intake after dehydration makes muscles more susceptible to electrical simulation-induced muscle cramp, probably due to dilution of electrolytes, and when OS-1® is consumed, the susceptibility to muscle cramp decreases.

INTRODUCTION

Exercise-associated muscle cramp (EAMC) is a painful, involuntary muscle contraction induced during or after exercise.1–4 EAMC can affect sports performance,5–6 thus prevention of EAMC is necessary. To do so, the aetiology of EAMC should be considered, but the mechanisms underpinning EAMC are not necessarily clear, although neural theory and dehydration-electrolytes theory have been mainly proposed.2,3 7 It has been documented, in recent studies, that EAMC stems from an imbalance between excitatory drive from muscle spindles and inhibitory drive from Golgi tendon organs to the alpha motor neurons, rather than dehydration or electrolytes deficits.2,7

In association with the dehydration-electrolytes theory, several studies have reported that dehydration does not affect muscle cramp.8–10 Miller et al9 did not find changes in threshold frequency (TF) of electrically induced muscle cramp after 3% dehydration. It should be noted that they reported an increase in serum sodium concentration from baseline (138.6±0.2 mmol/L) to post-dehydration (145.1±0.5 mmol/L). Similarly, Braulick et al10 reported an increase in serum sodium concentration in 3%–5% dehydrated condition (149.5±1.8 mmol/L) in comparison to euhydrated condition (141.9±3.1 mmol/L) and found no difference in TF between the conditions. The increases in serum sodium concentration suggest that body water loss was more than sodium loss in these studies. Hyponatremia is a condition in which sodium concentration is less than 135 mmol/L, and one of its typical symptoms is muscle cramp.11 Thus, it is possible that water intake after dehydration dilutes sodium and other electrolytes in the blood and extracellular fluid in the body and could decrease sodium concentration, inducing muscle cramp.
Our pilot study showed that no significant changes in TF were evident immediately after downhill running (DHR) in the heat to induce 1% or 2% loss of body mass by sweat, but when water (600 mL for 1% and 1200 mL for 2% dehydration condition) was ingested immediately after DHR, a significant decrease in TF was observed at 30 and 60 min later. Interestingly, when no water was ingested immediately after DHR, no such decrease in TF was found at 30 min later, but ingesting water at this time point, a decrease in TF was evident at 30 min later. These suggest that water intake in dehydrated condition may increase muscle cramp susceptibility, probably due to dilution of electrolytes in the body. Jung et al reported that half a tablespoon of sodium chloride added to a litre of carbohydrate–electrolyte beverage (Gatorade) taken before and during exercise in a hot environment, delayed onset of EAMC and prolonged exercise time, when compared with hypohydration condition in which no fluid was taken. However, no previous study has investigated the effect of oral rehydration solution (ORS) intake after dehydration on EAMC.

Therefore, the present study investigated the hypothesis that ingestion of spring water after 2% dehydration induced by DHR in the heat would decrease TF, but no changes in TF would be observed after ingestion of ORS. Previous studies have used TF of the abductor hallucis muscle,13–15 or the plantar flexors,16 and used a decrease in TF as a marker of increased muscle cramp susceptibility. In the present study, a TF measure of calf muscles was established and used to assess muscle cramp susceptibility after DHR in the heat, and OS-1® (Otsuka Pharmaceutical Factory, Japan) was used as ORS.

METHODS
Experimental design
This study used a cross-over study design to compare between water and OS-1® conditions for changes in TF of electrical stimulation to induce calf muscle cramp before and after DHR in the heat that resulted in 2% loss of body mass by increased sweating, using 10 healthy young men. The two conditions (water, OS-1®) were separated by a week in a counter-balanced order (five participants: water condition first, five participants: OS-1® condition first).

OS-1® contains sodium (2970 mg/L), potassium (794 mg/L), magnesium (25 mg/L), chloride (1801 mg/L), glucose (18 300 mg/L) and others (eg, phosphorus). For the water condition, a commercially available spring water was used, which contains a small amount of sodium (2 mg/L), potassium (0.5 mg/L), magnesium (18 mg/L), chloride (1.2 mg/L), and calcium (39 mg/L).

In addition to TF, serum sodium, potassium, magnesium and chloride concentrations were measured before, immediately after and 80 min after DHR (figure 1). Haematocrit (Hct), haemoglobin (Hb) and serum osmolality were assessed as dehydration parameters, and heart rate (HR), rate of physical exertion (RPE), thermal sensation, tympanic temperature, blood pressure and body mass were measured during DHR.

Participants
This study was approved by the Institutional Human Research Ethics Committee and complied with the Declaration of Helsinki. All participants signed an informed consent form and completed a medical questionnaire before participating in the study. It was confirmed that all of them had good health and fitness and did not have any pathological conditions such as peripheral neuropathy and liver cirrhosis that are known to cause muscle cramps. They did not have current or previous (in 2 years at least) lower limb injuries. Their mean±SD (range) age, height and body mass were 25.0±2.7 (22–31) years, 173.7±6.4 (165–184) cm and 74.0±12.0 (57.2–89.3) kg, respectively. Using the data from our pilot study showing that TF to induce muscle cramp for normal condition (no dehydration) was 21.3±1.5 Hz, and TF decreased by 15% (3 Hz) after water intake, it was estimated that nine participants were necessary. Considering a potential dropout and estimation error, 10 participants were recruited for this study.
All participants were instructed to refrain from any strenuous exercise for 1 week before the study. They were asked to consume 600 mL of spring water at 2 hours before coming to the laboratory, and refrain from any food intake for at least 2 hours before each exercise session. The food intake before the first session was recorded by each participant, and he was asked to have the same food before the second session. The participants were requested not to change their lifestyle and diet, not to take any anti-inflammatory drugs or nutritional supplements and not to perform any strenuous exercise during the week between the sessions.

**Threshold frequency**

The participants were familiarised with the electrical stimulation to induce muscle cramp in a familiarisation session at 1 week before the first experimental session. Here, we checked whether muscle cramp was induced by the stimulation. Electrical train stimulation was delivered to calf muscles of the kicking (dominant) leg by using a portable electrical stimulator (Compex 2, Compex Medical, Switzerland). To deliver the stimulation, one electrode (cathode) was placed over the tibialis posterior nerve in the popliteal fossa, and the other electrode (anode) was placed at the ankle between the end of posterior soleus muscle and tibialis tendon. The locations of the electrodes were marked by a semi-permanent marker to ensure the consistent electrode placement between measures in the same day and between sessions separated by a week.

Each participant was lying prone on a massage bed, and the instep was placed on the bed, which kept the ankle joint in a plantar-flexed position. Each stimulation consisted of 0.5 s duration of rise time and 2.0 s bursts of stimuli of 300 μs duration. The stimulation started at a frequency of 10 Hz, and two stimulations were given at this frequency during which the stimulation intensity was increased to a preset level (18–60 mA). The stimulation intensity for each participant was determined in the familiarisation session, and the intensity was set for the baseline TF to be between 22 and 26 Hz. The intensity was the same for the rest of the measurements and between the sessions. Thereafter, the stimulation frequency was automatically increased by 2 Hz every 30 s until muscle cramp was induced. The muscle cramp was identified by a visibly taut muscle, pain and calf muscle contraction and also reported by the participant. As soon as muscle cramp was confirmed, the investigator provided passive dorsiflexion of the foot to release the cramp. The TF at which cramp was induced was recorded for further analysis.

The test-retest reliability of the TF to induce muscle cramp was assessed between two baseline measurements within the same day separated by 15 min and between days separated by 24 hours. The coefficient of variation was 3% for the two measures in the same day as well as the two different days, and SE of the measurement was 0.8 Hz for the same day and 0.6 Hz for the different days.

**Downhill running**

The participants performed two bouts of DHR (slope: 5%) in a climate chamber set at 35°C–36°C and 27%–34% relative humidity until their body mass was decreased by 2%. To increase sweating, all participants wore a sauna suit for the upper body. The running velocity was between 7.5 and 9.7 km/h depending on the participants, and the velocity was changed on the condition of each participant such that the velocity was reduced when the participant struggled to maintain the velocity. The body mass was measured by a scale (Mettler Tolto ID1, Columbus, OH, USA), when the participants stopped after the first 20 min of DHR, took off all clothes and wiped sweat. This was repeated every 10 min thereafter until the body mass was decreased by 2% of the baseline body mass of each participant.

HR, RPE, and thermal sensation were measured before, then every 5 min during DHR. A HR monitor (Model S610i; Polar Electro Oy, Finland) was used to record HR during DHR, and the 6-point to 20-point Borg Scale was used for RPE. Thermal sensation was assessed by a thermal sensation scale (eight-point scale ranging from unbearably cold [0] to unbearable hot [8]). Blood pressure and tympanic temperature were measured by an automatic sphygmomanometer and a temperature probe, respectively, before, after the first 20 min and every 10 min during DHR, and at the end of DHR.

**Blood sampling and analyses**

Approximately 8 mL of blood was drawn by a standard venepuncture technique from the antecubital vein before, immediately after and 80 min after DHR. Using a portion of the blood sample (1.5 mL), Hct and Hb were measured by a capillary method and a HemoCue (Hb 201 System, Sweden), respectively. The rest of the blood was centrifuged for 10 min at 3000 rpm to obtain serum to be analysed for electrolyte concentrations of sodium, chloride, potassium and magnesium, and osmolality.

**Statistical analysis**

Two-way repeated measures analysis of variance (ANOVA) was used to compare the changes in the TF between conditions (water vs OS-1®) before, immediately after, and 50 and 80 min after DHR. Changes in other measures (body mass, HR, RPE, thermal sensation, blood pressure, tympanic temperature) during DHR, and changes in the blood dehydration parameters (Hct, Hb, serum osmolality), serum electrolyte concentrations (sodium, potassium, magnesium, chloride) before, immediately after and 80 min after DHR were also compared between conditions using two-way repeated measures ANOVA. When the ANOVA showed a significant time effect and/or interaction effect, a Tukey’s post-hoc test was performed for multiple comparisons. Statistical significance was set at p<0.05, and all data were presented as mean±SD.

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Table 1 Changes (mean±SD) in Hct and Hb, and serum concentration of sodium, potassium, magnesium and chloride before (pre), and immediately (0) and 80 min following DHR for the spring water (water) and electrolyte water (OS-1®) intake at 10 min after DHR

|          |          |          |        |          |
|----------|----------|----------|--------|----------|
|          | Pre      | 0        | 80     | ANOVA    |
| Hct (%)  | Water    | 47.3±2.7 | 48.2±2.1| 47.6±1.8| F=2.37   |
|          | OS-1     | 46.3±1.6 | 47.8±2.0*| 45.6±1.1| p=0.12   |
| Hb (g/dl)| Water    | 15.3±1.1 | 15.9±0.9*| 15.4±0.7| F=2.91   |
|          | OS-1     | 15.5±0.8 | 16.1±0.7*| 15.1±1.2| p=0.08   |
| Osmolarity (mOsm/kg H₂O) | Water | 290.9±5.4 | 296.2±3.5*| 285.0±3.4*| F=7.73 |
|          | OS-1     | 292.3±5.4 | 296.2±3.5*| 294.0±2.8| P=0.004† |
| Sodium (mmol/L) | Water | 141.7±1.5 | 142.3±1.9| 138.4±1.7*| F=33.85 |
|          | OS-1     | 141.8±0.9 | 142.6±1.4| 141.8±1.0| P=0.001† |
| Potassium (mmol/L) | Water | 4.5±0.4 | 4.4±0.3| 4.0±0.4*| F=1.51 |
|          | OS-1     | 4.5±0.4 | 4.4±0.4| 4.2±0.4| p=0.24 |
| Magnesium (mmol/L) | Water | 0.88±0.02 | 0.87±0.02| 0.84±0.02*| F=1.35 |
|          | OS-1     | 0.88±0.04 | 0.86±0.04| 0.85±0.05*| P=0.28 |
| Chloride (mmol/L) | Water | 101.0±1.1 | 102.6±2.3*| 99.5±2.1*| F=19.51 |
|          | OS-1     | 101.9±1.3 | 103.7±1.5*| 103.3±1.5*| P=0.001† |

*significant (p<0.05) difference from the prevalue. †significant (p<0.05) difference between conditions.

RESULTS

Downhill running

To reduce the 2% of body mass, three participants ran for 40 min, two for 50 min and five for 60 min for both conditions. The distance covered in the time was 5.4–8.4 km, and the average velocity was 7.9±1.2 km/h. The amount of body mass loss at immediately after exercise, thus the amount of the fluid intake was 1140–1600 (1324±17) ml for the water and 1100–1500 (1285±17) ml for the OS-1® condition.

The average HR, RPE and thermal sensation during DHR were 159.1±3.0 bpm, 5.9±0.9 and 13.0±2.0, respectively, without a significant difference between conditions. Systolic blood pressure increased and diastolic blood pressure decreased by approximately 10 mm Hg from the baseline during DHR. The tympanic temperature increased by 2°C from the baseline (36.6°C±0.2°C) during DHR for both conditions.

As shown in table 1, no significant differences were found for the baseline values of Hct, Hb and serum osmolality between the water and OS-1® conditions. Immediately after DHR, Hct, Hb and serum osmolality increased significantly for both conditions similarly, and returned to the baseline values at 80 min post-DHR.

Changes in TF

Figure 2 shows the changes in TF following DHR. No significant difference was found for the baseline TF between the water (25.1±1.5 Hz) and OS-1® (25.2±1.5 Hz) conditions. TF did not change significantly at immediately after DHR for both conditions (24.6±2.1 and 24.7±1.4 Hz, respectively). Changes in TF were significantly different between the conditions such that at 30 min after water intake (50 min post-DHR), TF decreased by 4.3 Hz from the baseline, but TF increased by 3.7 Hz after OS-1® ingestion. At 80 min after DHR, TF was still lower than the baseline for the water condition (20.0±1.6 Hz) but higher for the OS-1® condition (30.6±4.8 Hz).
Changes in serum electrolytes concentrations

As shown in table 1, a significant interaction effect was found for the changes in serum sodium and chloride concentrations, and post-hoc analyses showed significantly lower values at 80 min post-DHR (60 min after fluid ingestion) for the water than OS-1® condition. Serum chloride concentration increased significantly from baseline to immediately post-DHR for both conditions, without a significant difference between them. Serum potassium concentration decreased for the water condition only, and serum magnesium concentration decreased from baseline to 80 min post-DHR for both conditions similarly.

DISCUSSION

The results of the present study showed that (1) dehydration itself did not change the TF as previous studies reported,8-10 (2) spring water intake after dehydration decreased TF, which is an indicative of increased muscle cramp susceptibility, (3) ingestion of fluid containing electrolytes (OS-1®) increased TF, showing muscles became less susceptible to cramp and (4) serum sodium and chloride concentrations were decreased after spring water intake, but maintained after OS-1® intake. It appears that dilution of electrolytes was a cause of the increased muscle cramp susceptibility.

In the previous studies that investigated the effects of dehydration on muscle cramp, serum sodium concentration was increased9 10 and they found no changes in TF of electrical stimulation to induce muscle cramp at immediately after 3%-5% dehydration. In the present study, no significant changes in serum electrolyte concentrations were observed after 2% dehydration induced by DHR in the heat. This may be due to the lower dehydration level (2%) than that of the previous studies (3%-5%). In fact, the increases in Hct and Hb at immediately after exercise were smaller in the present study (table 1), when compared with those in the previous studies.9 10 It is possible that sodium, chloride and other electrolytes were lost with sweat, but because the decrease in plasma volume was proportional to that of the electrolytes, no changes in the electrolytes concentrations were observed. To the best of our knowledge, this was the first study to examine changes in TF for more than 30 min after dehydration and water intake, and show that TF decreased at 30 and 60 min after spring water intake (figure 2). The magnitude of the decrease in TF in the present study (4-5 Hz) was comparable to the previous study15 in which an increase in muscle cramp susceptibility was found after volitionally induced muscle cramp (~4 Hz). It is not known how long this effect would last, so it is important to extend the TF measures to 2 hours or longer after water intake in the future studies.

It is interesting that TF increased at 30 and 60 min after ingesting OS-1® (figure 2). This was not expected, since we hypothesised that TF would not change for the OS-1® condition. The magnitude of the TF increase (~4 to 5 Hz) in the present study was similar to that found after a neuromuscular electrical stimulation intervention for 6 weeks to reduce calf muscle cramp.16 To the best of our knowledge, this was the first study to show that ORS intake may reduce muscle cramp. It should be noted that serum sodium and chloride concentrations did not increase after OS-1® intake. This suggests that the increase in TF was not directly associated with serum sodium and chloride concentrations, although they showed a significant difference between the water and OS-1® conditions (table 1). It is important to note that OS-1® contains glucose (18 300 mg/L); thus, it might have been better to add the same amount of glucose to spring water to examine the effects of glucose on muscle cramp. If the electrolytes in OS-1 were the main reason for the increase in TF, judging from the differences in serum electrolytes concentrations between the OS-1® and water conditions (table 2), it appears that sodium and chloride are more important than potassium and magnesium. Further studies are necessary to investigate the mechanisms of the increased TF after OS-1® intake.

The main reason for the choice of DHR in the present study was that metabolic demand is smaller during DHR than level or uphill running,19 but it still induces a lot of sweating when it is performed in the heat. It has been documented that increased excitatory afferent activity due to increased muscle spindle activity and decreased inhibitory afferent activity due to decreased Golgi tendon activity lead to increased alpha motor neuron activity, increasing muscle membrane activity, inducing muscle cramp.14 DHR has been known to induce muscle damage due to eccentric (lengthening) muscle contractions.20 Thus, it was thought that muscle spindle and Golgi tendon organs might be more affected by eccentric (lengthening) muscle contractions performed in DHR. It is interesting to investigate further the effects of eccentric muscle contractions on EAMC in comparison to concentric (shortening) muscle contractions.

The present study showed a possibility that intake of OS-1® before and during exercise would reduce EAMC. To investigate this, it is necessary to establish a reliable marker of muscle cramp susceptibility during exercise. It is also important to understand the relationship between TF and actual muscle cramp. Even if the cause of EAMC is more associated with neural especially spinal mechanisms, it is likely that dilution of electrolytes is also involved in the mechanisms directly or indirectly.1 It is

What this study adds

- Water intake after dehydration with exercise makes muscles more susceptible to muscle cramp.
- Oral rehydration solution (ORS) intake after dehydration with exercise makes muscles less likely to cramp.
- It seems possible that dilution of electrolytes especially sodium and chloride in the body may increase muscle cramp susceptibility.
- Ingestion of ORS instead of water during and after exercise may reduce exercise-associated muscle cramp.

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also interesting to know whether the findings of our study will apply to other muscle cramps such as those affecting older people and pregnant women. We conclude that the involvement of electrolytes in EAMC should not be ignored.

Acknowledgements The authors appreciate the participants of the studies.

Contributors KN designed the study, the data collection was performed mainly by WYL and HK, and the data analyses were done by WY and KN. KN drafted the manuscript and the final manuscript was checked by all authors.

Funding The authors appreciate the support from the Otsuka Pharmaceutical Factory Incorporation (Japan), which provided us a grant and OS-1® to perform the study. However, this manuscript was not oversighted by the company.

Competing interests None declared.

Patient consent for publication Obtained.

Provenance and peer review Not commissioned; externally peer reviewed.

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