Exercise-Associated Muscle Cramps: Causes, Treatment, and Prevention

Kevin C. Miller, PhD, ATC, CSCS,*† Marcus S. Stone, PhD, ATC,‡ Kellie C. Huxel, PhD, ATC,§ and Jeffrey E. Edwards, PhD||

Context: Exercise-associated muscle cramps (EAMC) are a common condition experienced by recreational and competitive athletes. Despite their commonality and prevalence, their cause remains unknown. Theories for the cause of EAMC are primarily based on anecdotal and observational studies rather than sound experimental evidence. Without a clear cause, treatments and prevention strategies for EAMC are often unsuccessful.

Evidence Acquisition: A search of Medline (EBSCO), SPORTDiscus, and Silverplatter (CINHAL) was undertaken for journal articles written in English between the years 1955 and 2008. Additional references were collected by a careful analysis of the citations of others’ research and textbooks.

Results: Dehydration/electrolyte and neuromuscular causes are the most widely discussed theories for the cause of EAMC; however, strong experimental evidence for either theory is lacking.

Conclusions: EAMC are likely due to several factors coalescing to cause EAMC. The variety of treatments and prevention strategies for EAMC are evidence of the uncertainty in their cause. Acute EAMC treatment should focus on moderate static stretching of the affected muscle followed by a proper medical history to determine any predisposing conditions that may have triggered the onset of EAMC. Based on physical findings, prevention programs should be implemented to include fluid and electrolyte balance strategies and/or neuromuscular training.

Keywords: cramping; dehydration; electrolytes; fatigue; stretching

Skeletal muscle cramps that occur during or shortly following exercise in healthy individuals with no underlying metabolic, neurological, or endocrine pathology have been termed exercise-associated muscle cramps (EAMC). Although controversial, an important differentiation in determining the cause of EAMC may be the number and location of muscles affected. EAMC typically occur in single, multijoint muscles (eg, triceps surae, quadriceps, hamstrings) when contracting in a shortened state, whereas generalized EAMC occur in multiple, usually bilateral muscles. Clinically, EAMC may be recognized by acute pain, stiffness, visible bulging or knotting of the muscle, and possible soreness that can last for several days. Although EAMC can be isolated, athletes often complain of EAMC symptoms up to 8 hours after exercise. This postexercise period of increased susceptibility to EAMC has been termed the cramp prone state. Although some EAMC do not appear to affect athletic performance, other times, EAMC can be completely debilitating. The clinical presentation of EAMC is easily recognized, but its cause continues to be unresolved. A better understanding of the underlying mechanisms causing EAMC may allow better prevention and treatments, thus reducing the incidence rate. Norris et al reported that 95% of physical education students (115 of 121) had experienced spontaneous cramps in their lifetimes with 26% (31 of 121) experiencing cramps after exercise. Kantarowski et al reported that 67% of triathletes (1631 of 2438) complained of EAMC under a variety of training conditions. More recently, a subset of American football players who experienced exertional heat illness reported concomitant skeletal muscle cramping. Thus, EAMC are common in both the recreational and the competitive athlete.

Although EAMC are common in athletes, the cause is unknown and controversial. Traditionally thought to be caused by factors associated with exercise in hot and humid environments (eg, dehydration and/or electrolyte imbalances), evidence suggests a neuromuscular cause. The inference...
from several studies is that EAMC have a singular, unknown cause. Authors have also suggested that there may be different kinds of EAMC and thus different causes (eg, isolated and generalized). Without a clear cause, the treatments and prevention strategies for EAMC vary considerably and have limited perceived effectiveness by health professionals. One approach to determining cause and the effectiveness of treatments is to examine the published studies and determine their level of evidence.

THEORIES FOR THE CAUSE OF EAMC

The dehydration–electrolyte imbalance theory is the most common among health care professionals. Proponents state that the body does not store enough water for exercise and athletes do not ingest enough water to replace the amounts they lose during exercise; EAMC are the result of fluid and electrolyte depletion, which results in the sensitization of select nerve terminals. The resulting contracture of the interstitial space increases the mechanical pressure on select motor nerve endings and finally results in EAMC. Exercise in hot and humid conditions exacerbates the amount of fluid and electrolytes lost, thereby facilitating cramps.

Support for the dehydration–electrolyte imbalance theory comes mainly from research classified as level 4 and 5 evidence. Miners develop cramps because of their sweat losses while working in hot and humid conditions. More recently, researchers observed that the majority of cramping (95%, 87 of 92) occurred in hot months—specifically, when football players exercised in environmental conditions in which the risk of developing heat illness was “high” or “extreme.” Other evidence for this theory comes from case studies and other observational work in which large sweat losses occurred in exercising athletes. Some health professionals postulate that sweat glands are unable to reabsorb sodium at “high” sweat temperatures and that the body does not store enough water for exercise.

The neuromuscular theory of EAMC proposes that muscle overload and neuromuscular fatigue cause an imbalance between excitatory impulses from muscle spindles and inhibitory impulses from Golgi tendon organs (GTOs). These localized EAMC tend to occur when the muscle is contracting in an already-shortened position. The reduced tension in the muscle tendon likely reduces the inhibitory feedback from GTO afferents, thereby predisposing the muscle to cramp from the imbalance between inhibitory and excitatory drives to the alpha motor neuron. This enhanced excitability at the spinal level results in an increase in alpha motor neuron discharge to the muscle fibers, producing a localized muscle cramp.

Study designs that examine the plausibility of the neuromuscular system’s role in EAMC are stronger than those for dehydration (levels 3 to 5): animal, exercising humans, and stretching for EAMC. These varying models and treatment observations are more consistent with the neuromuscular theory than with the dehydration–electrolyte imbalance theory.

In felines, muscle spindle and GTO activity were measured following neuromuscular fatigue induced by supramaximal stimulation (100 Hz). Fifty percent of type Ia (25 of 49) and 55% of Ila (18 of 33) muscle spindle afferents increased their resting discharge following fatiguing electrical stimulation. Similarly, feline GTO discharge rate was lowered and delayed with fatigue induced with a similar protocol. Thus, neuromuscular fatigue appeared to decrease the inhibition from the GTO and increase the excitatory stimuli from muscle fibers.

The treatment for EAMC also fails to support the dehydration–electrolyte imbalance theory. If EAMC were due to dehydration, the simple cure would be fluid replacement. However, when carbohydrate-electrolyte fluids were ingested at a rate that matched sweat loss, EAMC still occurred in 69% of athletes (9 of 13). Moreover, athletes who develop EAMC often ingest similar amounts of fluid during exercise as do their noncramping counterparts. Oral fluid ingestion may be ineffective, and intravenous fluid may provide a faster delivery for athletes suffering from acute EAMC. It is interesting that stretching the affected muscle almost immediately relieves EAMC and yet has no effect on the fluid conditions of the body.

Overall, the dehydration–electrolyte imbalance theory has limitations: First, inferences of cause and effect cannot be made from observational data (eg, field studies); causation may be inferred only from meta-analyses and randomized, experimental research designs (evidence levels 1 and 2, respectively). Second, although EAMC may appear in the presence of significant electrolyte and fluid losses during exercise, numerous other variables associated with exercise may be factors (eg, accumulation of metabolites, intensity of exercise, and acclimatization). Because athletes who experience EAMC often have significant fluid deficits, restoring body fluids is an appropriate precautionary measure against the development of more serious forms of heat illness (eg, exertional hyponatremia, heat stroke).

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spindles. These effects may result in a heightened excitatory state at the spinal level.

In humans, EAMC occurs more frequently at the end of competitions and physical work\textsuperscript{32-35} and when the muscle contracts while it is already shortened.\textsuperscript{36} Stretching, the primary treatment for acute EAMC,\textsuperscript{36} is thought to relieve EAMC via autogenic inhibition. Stretching increases the tension in the muscle’s tendon, resulting in GTO activation and an increase in inhibition of the alpha motor neuron, which may restore the physiological relationship between excitatory and inhibitory impulses to the alpha motor neuron.\textsuperscript{36}

The neuromuscular theory also has limitations. The report of altered muscle spindle and GTO activity relies on difficult methodologies that have produced inconsistent results. The majority of GTO Ib afferents (5 of 8, 63%) have only a slight decline or no change in firing in response to stretching of a fatigued muscle.\textsuperscript{37} Neuromuscular fatigue often induces muscle afferent fatigue with supramaximal electrical stimulation (eg, 100 Hz).\textsuperscript{38,42}\textsuperscript{1} Normal human muscle recruitment patterns indicate stimulation frequencies much lower (eg, < 30 Hz) than those used to induce fatigue in animal studies (eg, 100 Hz).\textsuperscript{43} Low electrical stimulation frequencies closer to normal recruitment patterns (eg, 16 to 32 Hz) have successfully induced cramps in humans.\textsuperscript{39,51,53} Thus, the frequencies used to support the neuromuscular theory\textsuperscript{39,42} do not match normal neuromuscular signaling in humans. Finally, it is unclear how fatigued a muscle needs to become for an EAMC to occur or whether the neuromuscular fatigue is occurring peripherally (ie, in the muscle) and/or centrally (in the spinal cord or brain). Moreover, it is unlikely that neuromuscular fatigue induced with electrical stimulation is the same as fatigue induced with volitional muscle contractions, given that larger diameter motor neurons/units are stimulated first with electrical stimulation and last with volitional contractions.\textsuperscript{30,21}

Muscle fatigue is a continuum rather than an absolute condition. It is likely that the degree of fatigue required to elicit cramping is unique to each athlete.

Because EAMC occur in a variety of situations, environmental conditions, and populations, it is unlikely that a single factor (eg, dehydration, electrolyte imbalance, or neuromuscular factors) is responsible for causing them directly. It is more likely that EAMC are due to a combination of factors that simultaneously occur under specific physiological circumstances in each athlete.

**TREATMENT OF EAMC**

The paucity of experimental data regarding the cause of EAMC has led to a plethora of treatments for EAMC, confirming the lack of understanding and consensus for EAMC etiology. Many of these treatment options are anecdotal and unsupported by experimental research: ingesting mustard, pickle juice, sports drinks, cryotherapy, thermotherapy, massage, decreasing exercise intensity, body position, intravenous infusion, and TENS (transcutaneous electric nerve stimulation) therapy.

The dehydration–electrolyte theory suggests that ingesting fluids containing electrolytes is beneficial to treat and alleviate EAMC. However, owing to the minimal amount of electrolytes in many sports drinks, it may be difficult to sufficiently replace the volume of electrolytes lost during exercise even if the athlete has modest sweat losses and sweat sodium content. Assuming that a relationship between dehydration–electrolyte imbalance and EAMC exists, the National Athletic Trainers’ Association recommends that athletes prone to muscle cramping add 0.3 to 0.7 g/L of salt to their drinks to stave off muscle cramps.\textsuperscript{3} Others have recommended adding higher amounts of sodium (about 3.0 to 6.0 g/L) to sports drinks based on the frequency of EAMC.\textsuperscript{3} Note that fluids and electrolytes are not absorbed immediately after ingestion; that is, even hypotonic fluids require at least 13 minutes to be absorbed into the circulatory system.\textsuperscript{61} Theoretically, intravenous infusion of fluids removes this delay, and it has been used to aid athletes who develop acute EAMC.\textsuperscript{22}

However, experimental evidence regarding the effectiveness of intravenous fluid infusion on EAMC is still lacking.

Stretching, quinine, and beta-blockers have stronger levels of evidence (level 2 or 3) to support their use, based on drug trials with human participants\textsuperscript{30,41} and other research.\textsuperscript{36,51} If the athlete has no underlying illness, then the most common treatment for EAMC is stretching,\textsuperscript{36} which has proven to be effective for EAMC and other types of muscle cramps\textsuperscript{37,38,51,52} but may be ineffective for “heat cramps.”\textsuperscript{4} Therefore, moderate stretching of the affected muscle to alleviate the cramp is recommended.

Once a cramp is alleviated, health care providers should determine what factors may be involved (eg, diabetes mellitus, thyroid disease).\textsuperscript{6}

**PREVENTION OF EAMC**

Despite the lack of direct evidence, maintaining hydration and adequate electrolyte levels is a good prevention strategy for individuals susceptible to EAMC.\textsuperscript{36} Fluid volumes of 1.8 L per hour have been well tolerated by tennis athletes who are susceptible to EAMC.\textsuperscript{6} Health professionals should monitor each athlete’s fluid losses and recommend replacement during and after exercise (eg, obligatory fluid losses). Both the National Athletic Trainers’ Association and the American College of Sports Medicine recommend a volume of fluid that allows for less than a 2% body weight reduction.\textsuperscript{15,39} Monitoring an athlete’s body weight is an easy method of ensuring adequate fluid replacement and individualizes each athlete’s fluid needs.\textsuperscript{13}

An athlete who ingests a liter of water or hypotonic sports drink at least 1 hour before competition can be confident that the majority of the fluid, electrolytes, and nutrients have been absorbed and are available in the body. Fluids should be available and easily accessible throughout practices and competitions. A balanced diet is important given that much of fluid and electrolyte replacement occurs during meals.\textsuperscript{12}
Clinical Recommendation

**SORT: Strength of Recommendation Taxonomy**

**A:** consistent, good-quality patient-oriented evidence  
**B:** inconsistent or limited-quality patient-oriented evidence  
**C:** consensus, disease-oriented evidence, usual practice, expert opinion, or case series

| Clinical Recommendation                                                                 | SORT Evidence Rating |
|----------------------------------------------------------------------------------------|----------------------|
| Clinicians should take a multifaceted approach to treating and preventing EAMC as it is unlikely they are caused by a single factor. | C                    |
| To treat EAMC, clinicians should stretch the cramping muscle. Once the EAMC is alleviated, the clinician should focus on the athlete’s hydration and electrolyte needs. 14,15,39,36 | B                    |
| To prevent EAMC, clinicians should take a detailed medical history of each athlete, determine their level of conditioning, and monitor their fluid and electrolyte levels. 15,57 | C                    |

For more information about the SORT evidence rating system, see www.aafp.org/afpsort.xml and Ebell MH, Siwek J, Weiss BD, et al. Strength of Recommendation Taxonomy (SORT): a patient-centered approach to grading evidence in the medical literature. *Am Fam Physician.* 2004;69:549-557.

A common perception is that level of conditioning is a factor in the development of EAMC.14,26,41 There is a strong theoretical basis for performing exercises that target the neuromuscular system to prevent EAMC.25,31,56 Prevention exercises that target muscle spindle and GTO receptors should be implemented to delay neuromuscular fatigue onset and, hence, EAMC. Plyometric exercises may be beneficial to elicit neural adaptations in muscle spindle and GTO receptor firing, enhancing efficiency and sensitivity of reflexive and descending pathways used for neuromuscular control.14,59,62 Endurance training may also serve as an effective means of preventing EAMC by expanding plasma volume and the extracellular fluid compartment15 and delaying neuromuscular fatigue.25

**SUMMARY**

EAMC are common and affect several different populations. Despite the prevalence of EAMC, few experimental data exist on their cause, treatment, and prevention. Although several theories have been postulated for their cause, much of the evidence is nonscientific or observational; thus, causation cannot be inferred. Numerous untested, anecdotal prevention strategies exist for the prevention of EAMC (eg, pickle juice). The level of evidence for these prevention strategies is low (level 4 or 5). It is likely that the cause of EAMC is multifactorial. Stretching appears to be effective regardless of the cause of EAMC.

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