INTRODUCTION

Water is an essential component of the human body with important metabolic, transport, temperature control, structural, and mechanical functions (Lorenzo, Serra-Prat, & Yébenes, 2019). In skeletal muscle, for example, fluid flows are crucial in the regulation of ion concentrations and pH, which affect muscle contraction and hence force development during exercise (Sjogaard & Saltin, 1982). In addition to the importance of muscle fluids for the biochemistry of muscle contraction, recent studies suggest that fluid plays important mechanical roles within muscle, influencing passive force development during stretch (Gindre, Takaza, Moerman, & Simms, 2013; Sleboda & Roberts, 2017), work production during active contraction (Azizi, Deslauriers, Holt, & Eaton, 2017), and dynamic changes in muscle shape that influence muscle speed and force (Azizi, Brainerd, & Roberts, 2008; Eng, Azizi, & Roberts, 2018).

It is well documented that with the onset of muscle activity, fluid moves from blood plasma into muscle due to increased hydrostatic (Kjellmer, 1964) and osmotic (Lundvall, Mellander, Westling, & White, 1972) forces. These findings have been confirmed in sympathectomized cat muscle (Björnberg, 1990), in perfused, isolated cat muscle (Ward, Hamilton, & Watson,
Activity-induced increases in muscle volumes have also been observed in vivo in different kinds of (sport) activities, for example, after short-term, high-intensity exercises (Ploutz-Snyder, Convertino, & Dudley, 1995; Raja, Raymer, Moran, Marsh, & Thompson, 2006; Shi, Zheng, Chen, & Huang, 2007; Sjøgaard & Saltin, 1982). Furthermore, increased muscle thickness was observed after a prolonged isometric contraction of the supraspinatus muscle (Jensen, Jørgensen, & Sjøgaard, 1994).

Additional evidence for the shift of fluids into the intracellular space results from the proton transverse relaxation time (T2) increases observed by magnetic resonance imaging after exercise (Meyer & Prior, 2000). Increases in T2 times with exercise have been frequently reported in the literature (Fisher, Meyer, Adams, Foley, & Potchen, 1990; Fleckenstein et al.; Shellock, Fukunaga, Mink, & Edgerton, 1991). It has been suggested that increases in T2 relaxation times are related to fluid content within a tissue, even though the exact mechanism underlying T2 time increases with muscle activity is not well understood (Archer et al., 1992; Cole, Leblanc, & Hingran, 1993; Fleckenstein et al., 1991; Ploutz-Snyder, Nyren, Cooper, Potchen, & Meyer, 1997). However, T2 changes cannot quantify absolute muscle volume increases due to fluid shifts. Consequently, segmentation of muscle outlines within MRI images is currently considered the gold standard for muscle volume determination (Pons et al., 2018). Furthermore, this method has been proven to be able to detect intervention-related changes in leg muscle volumes (Hudelmaier et al., 2010).

During prolonged running, it was found that total muscle water of the vastus lateralis muscle increased at 10 min and then stayed almost constant at 120 min (Costill, Coté, Fink, & Handel, 1981). However, neither this nor other studies addressing fluid shifts during prolonged running exercise (Senay & Pivarnik, 1985) have analyzed the time course of muscle fluid volume changes at a higher temporal resolution. Furthermore, despite the importance of calf muscles for propulsion and support during running (Hamner, Seth, & Delp, 2010), analyses of volume changes in these muscles during locomotion are rare. This gap of knowledge limits our current understanding of how muscle fluid volumes of calf muscles change in distance running.

Therefore, the purpose of the present study was to describe the time course of muscle volume changes during a prolonged bout of running. Based on results from single fiber stimulation (Rapp et al., 1998) and results from T2-relaxation time changes after exercise, we hypothesized that whole-organ muscle volumes would increase rapidly with the onset of activity and then approach a plateau at a slower rate.

2 MATERIALS AND METHODS

We included 18 recreationally active participants (11 men and 7 women; height: 1.77 ± 0.08 m; body mass: 70.6 ± 7.5 kg; age: 29 ± 3 years) in this study. All participants were free of injuries for at least 12 months before the study. The study protocol was approved by the local university ethics committee. All procedures were carried out in compliance with the Declaration of Helsinki. Before the data collection, we obtained written, informed consent from all participants.

All participants underwent a protocol consisting of 30 min of resting in a supine position to achieve an uninfluenced baseline volume followed by in total 75 min of running at a self-selected speed on a treadmill (Figure 1). The running speeds were selected individually such that they could be kept for the entire 75-min running protocol. Running speeds varied between 2.2 and 3.6 m/s (mean: 2.7 m/s; standard deviation: 0.4 m/s) between participants. The treadmill was positioned directly in front of the MR room (about 10 m) with a 3.0-Tesla magnetic resonance imaging (MRI) scanner (Philips Ingenia 3.0T, Philips Healthcare, Best, Netherlands). A 16-channel knee coil was chosen to achieve the best possible resolution as well as fixation and reproducibility of the positioning. In addition, the feet were fixed to a plate connected to the MR Table to allow fast and accurate repositioning between scans. A T2 spin echo sequence was used for the anatomical delineation of the different muscles (echo time: 10 ms; repetition time: 2000 ms; flip angle 90 degree; field of view: 150 × 197 × 185 mm; slice thickness: 5 mm; number of slices: 10; gap: 15 mm; acquisition voxel: 1.5 × 1.5 × 1.5 mm; reconstruction voxel: 0.61 × 0.61 mm; turbo factor: 16; parallel imaging acceleration factor (SENSE): 1.7). In previous studies, both T1- and T2-weighted sequences have been successfully validated for muscle segmentation (McColl, Fleckenstein, Bowers, Theriault, & Peshock, 1992; Orgiu et al., 2016; Warfield, Mulkern, Winalski, Jolesz, & Kikinis, 2000). T2-weighted sequences allow not only the anatomical visualization of the muscles but also a subjective assessment of a possible delimitable intramuscular edema as a potential correlate for volume increase. Two saturation slabs (cranial and caudal the field of view) were used to decrease artifacts due to inflowing blood. The cranio-caudal field of view was 15 cm, starting proximally at the fibular head. The scanning time was 2:16 ± 0:27 min, additionally it took 0:42 ± 0:09 min between pause of the run and start of the scan as well as further 0:42 ± 0:18 min after completion of the scan until the restart of the run. We performed MRI scans of the shank before and after 2.5, 5, 10, 15, 45, and 75 min of running in a standardized ankle and knee joint configuration (Figure 1). The run was only interrupted for the duration of the scans and resumed with the same speed. The ankle joint was positioned neutrally (90° joint angle) while the knee was positioned in a slightly (about 20°) flexed configuration. In...
this position, we took ten 5-mm thick transversal MRI slice images of the shank.

To quantify muscle volume, a board-certified radiologist with special focus on musculoskeletal imaging (6 years of experience) manually segmented the medial (GM) and lateral gastrocnemius (GL), soleus (SO), peroneus longus (PER), tibialis posterior (TP), tibialis anterior (TA), and extensor digitorum longus (EDL) muscles within each MRI slice and for every time point. Similar to Ploutz-Snyder et al. (Ploutz-Snyder et al., 1995), we quantified muscle volumes through the average cross-sectional areas of each muscle. For this step, we excluded the two most proximal and two most distal slices, since they were contaminated with artifacts in some participants and muscles.

We performed one factor (time) repeated measures analysis of variance (ANOVA) in order to identify the effects of running time on the muscles anatomical cross-sectional area (ACSA). In case of a significant main effect of time, we performed post hoc comparisons between time points while using Tukey’s Honestly Significant Difference procedure to control for alpha-error accumulation.

3 | RESULTS

We observed a significant increase in ACSA of more than 5% compared to the baseline of both gastrocnemius muscles after 2.5 min of running \( (p < .032; \text{Figure 2; Table 1}). \) Subsequently, ACSAs of the gastrocnemii further increased at a slower rate and reached peak values of about 109% and about 107% of resting values after 10 min of running for gastrocnemius lateralis and gastrocnemius medialis, respectively (Figure 2; Table 1). We observed a similar pattern of the initial increase in soleus, tibialis anterior, and peroneus longus muscles. However, the amplitudes of ACSA changes were much smaller in these muscles (< 3%), and we found no significant differences compared to baseline (Figure 2; Table 1). Tibialis posterior and extensor digitorum longus showed no initial increase in ACSA (Figure 2; Table 1).

After having reached peak ACSA values between 5 and 15 min of running, all muscles showed a reduction over the later stages of the run (Figure 2, Table 1). For all muscles except gastrocnemius medialis, ACSAs were lower compared to baseline after 75 min of running. Except for soleus and extensor digitorum, the ACSAs after 75 min of running were significantly \( (p < .05) \) lower compared to the maximum
ACSA observed within the first 15 min of running (Figure 2; Table 1).

**4 | DISCUSSION**

The purpose of the present study was to provide a detailed insight into the time course of calf muscle volumes during prolonged running. We found increases in gastrocnemii ACSA of up to 9% above resting baseline in the early stages of the 75-min run and reductions of up to 4% below baseline at the end of the running protocol (for the tibialis posterior muscle). Therefore, we can only partly accept that muscle volumes would increase rapidly with the onset of activation and then approach a plateau at a slower rate, since we observed ACSA reductions instead of a plateau in the later stages of the running protocol.

The fluid exchange between extravascular and intramuscular compartments is a complex interplay between hydrostatic and osmotic forces, governed by anatomical and biochemical factors (Lorenzo et al., 2019; Senay & Pivarnik, 1985; Sjogaard & Saltin, 1982). In line with previous literature, we found a rapid initial increase of ACSA with the onset of exercise after 2.5 min of running. Several studies have reported that with the onset of muscle activity, the hydrostatic pressure gradient from the vascular towards the interstitial space of muscle rapidly drops, which theoretically would cause an efflux of fluid from muscle towards the vascular space (Folkow, Haglund, Jodal, & Lundgren, 1971; Höjensgård & Stürup, 1953; Nicolaides & Zukowski, 1986; Pollack & Wood, 1949). However, these changes in hydrostatic pressures are offset by alterations in osmotic pressure gradients as a result of muscular activity and the accumulation of exercise-related metabolites (Björnberg, 1990; Lundvall et al., 1972; Meyer & Prior, 2000). The intensity of this opposing osmotic pressure gradient is considered to be proportional to the intensity of muscle action (Jenner, Foley, Cooper, Potchen, & Meyer, 1994). Senay and Pivarnik (Senay & Pivarnik, 1985) conclude from their own (Senay, 1970, 1972) and the work of others (Beaumont, Underkofler, & Beaumont, 1981; Greenleaf et al., 1977) that the osmotic pressure gradient exceeds the opposing hydrostatic pressure gradient approximately when “muscle exercises at 40–60 percent of its maximum oxygen consumption”. We did not obtain the maximum rate of oxygen uptake (\(\dot{VO}_2\) max) of the runners in our study. However, when using the regression equation developed from a very similar sample of recreational runners to predict the oxygen uptake from treadmill running speed, and by considering the average \(\dot{VO}_2\) max of these runners (Williams & Nute, 1983), we can estimate that our runners were running at approximately 58% of their \(\dot{VO}_2\) max. Consequently, fluids might have shifted from extracellular compartments (mostly from blood plasma (Ploutz-Snyder et al., 1995; Senay & Pivarnik, 1985), towards interstitial and intracellular spaces due to increased osmotic drive towards the muscle.

We observed differences in the initial effects of running on ACSA between muscles, where both gastrocnemius muscles pronouncedly increased their ACSA, while peroneus, tibialis anterior, and soleus increased their ACSA to a lesser extent. Tibialis posterior and flexor digitorum longus did not increase or even decreased their ACSA (Figure 2). These differences could result from multiple factors. Since the increase in ACSA is related to the relative work rate of a muscle (Adams, Duvoisin, & Dudley, 1992; Fisher et al., 1990; Jenner et al., 1994), different relative involvement of muscles might explain these differences in part. Furthermore, the accumulation of metabolites within muscle cells is related to fiber type distribution (Prior, Ploutz-Snyder, Cooper, & Meyer, 2001). Since gastrocnemius muscles contain a higher relative volume of fast-twitch muscle fibers as compared to, for example, the soleus muscle (Alway, MacDougall, Sale, Sutton, & McComas, 1988), it is likely that fiber type distribution is one of the underlying factors explaining the more pronounced shift of fluid to the gastrocnemii observed in the initial stages of running (Stallknecht, Vissing, & Galbo, 1998).

### TABLE 1 Average (± one SD) cross-sectional areas of the analyzed muscles at the respective time points

| Time (min) | Gastrocnemius lateralis (cm²) | Gastrocnemius medialis (cm²) | Soleus (cm²) | Peroneus longus (cm²) | Tibialis posterior (cm²) | Tibialis anterior (cm²) | Extensor digitorum longus (cm²) |
|-----------|-----------------------------|----------------------------|-------------|----------------------|------------------------|------------------------|-------------------------------|
| 0         | 5.3 ± 1.7                  | 10.6 ± 2.6                 | 16.2 ± 3.3  | 4.6 ± 1.1            | 2.8 ± 0.7              | 4.7 ± 0.8              | 2.4 ± 0.6                     |
| 2.5       | 5.7 ± 0.75                 | 11.2 ± 2.9                 | 16.1 ± 3.4  | 4.6 ± 1.2            | 2.7 ± 0.7              | 4.7 ± 0.8              | 2.3 ± 0.6                     |
| 5         | 5.7 ± 0.75                 | 11.3 ± 2.8                 | 16.4 ± 3.5  | 4.6 ± 1.2            | 2.8 ± 0.75             | 4.8 ± 0.9              | 2.4 ± 0.6                     |
| 10        | 5.8 ± 2.0                  | 11.4 ± 2.9                 | 16.3 ± 3.4  | 4.6 ± 1.275          | 2.7 ± 0.8              | 4.8 ± 0.95             | 2.4 ± 0.6                     |
| 15        | 5.7 ± 2.0                  | 11.3 ± 2.9                 | 16.4 ± 3.5  | 4.7 ± 1.75           | 2.7 ± 0.75             | 4.8 ± 0.95             | 2.4 ± 0.6                     |
| 45        | 5.7 ± 1.8                  | 11.0 ± 2.8                 | 16.1 ± 3.4  | 4.6 ± 1.1            | 2.7 ± 0.7              | 4.6 ± 0.9              | 2.3 ± 0.6                     |
| 75        | 5.3 ± 1.8                  | 10.6 ± 2.7                 | 15.9 ± 3.5  | 4.4 ± 1.175          | 2.6 ± 0.75             | 4.5 ± 0.8              | 2.3 ± 0.6                     |

Note: 0, 2.5, 5, 10, 15, 45, and 75 indicate a significant (\(p < .05\)) difference to the volume at baseline, 2.5 min, 5 min, 15 min, 45 min and 75 min of running, respectively.
Furthermore, the anatomical arrangement of muscles within the leg may influence fluid movement. The gastrocnemius, for example, is arranged superficial to the deeper soleus and tibialis posterior muscles. Muscles exert transverse forces on each other during contraction (Siebert, Günther, & Blickhan, 2012), and we speculate that deeper muscles may experience higher intramuscular pressures that might oppose fluid influx. These factors remain speculative, however, and will require experimental verification through future studies.

Beyond 15 min of running, the ACSA of all analyzed muscles began to decrease, and by the end of the running trials, all muscles but the soleus had lower ACSA values compared to baseline measurements. We believe that sweat losses of total body water might be an important factor explaining some of the almost linear reduction in muscle ACSA. Sweat losses can affect the osmolality of blood (Duvillard, Braun, Markofski, Beneke, & Leithäuser, 2004). Hence, with accumulated sweat losses, the osmotic pressure gradient might change with prolonged running such that the overall pressure gradient (combined osmotic and hydrostatic) favors movement of water from muscle to the vascular space. With prolonged running, fluids likely shift from intracellular and interstitial muscle spaces to the vascular space in order to conserve blood plasma volume and to protect vital function (Costill, 1977; Costill et al., 1981; Nose, Mack, Shi, & Nadel, 1988).

Next to sweating-related changes in osmotic pressure gradients, it is conceivable that changes in metabolite contents within intra- and extramuscular spaces induce osmotically driven fluid shifts away from active muscle tissues. With prolonged submaximal exercise, it has been observed that lactate is removed from active muscle to the vascular space at a higher rate, potentially leading to changes in osmotic pressure gradients (Gladden, 2000; Stallknecht et al., 1998).

Changes in muscle volume may alter the force generating potential of muscle. Studies on isolated muscles have demonstrated an increase in passive muscle tension with increased muscle fluid volume (Sleboda & Roberts, 2017). Passive muscle force at a given length changes in proportion to volume change, and a measurable change in force can be observed with volume changes as small as 5% in isolated muscles (Sleboda, Wold, & Roberts, 2019). Changes in muscle volume may also influence active contractile force via influences on muscle pressure (Sleboda & Roberts, 2020). We speculate that the changes in muscle volume observed in the present study may have been accompanied by changes in the passive mechanical behavior of running muscles. Passive force development contributes considerably to moment and power generation in human locomotion (Whittington,ilder, Heiderscheit, & Thelen, 2008), and changes in passive muscle tension have the potential to influence the kinematics and energetics of locomotion. Whittington et al. (Whittington et al., 2008) demonstrated that passive force contributions occur at the end ranges of joint motions (i.e., peak hip extension, peak knee extension, and peak ankle dorsiflexion), and so fluid-related changes in passive muscle mechanics have the greatest potential to influence force development during the late swing and early stance phase in running, potentially altering the foot strike pattern of runners (Willwacher, Regniet, Fischer, Oberländer, & Brüggemann, 2014). Combining interventions that affect muscle fluid content in humans in vivo with measurements of passive force development seems to be an essential step in future studies. Potentially, this could address other muscle groups, for example, the hip flexors.

However, the present study faces several limitations. We did not standardize the running speed, for example, to VO2 max. Future studies should more carefully address this issue in order to develop the dose–response relationship between running exercise and muscle fluid changes. Related to this, we also did not determine total body water losses due to sweat by, for example, measuring body mass losses after the running protocol. In order to better understand the reductions in muscle ACSA at the later stages of prolonged running, future research needs to address this.

Furthermore, while there is good evidence from animal muscle preparations suggesting that changes in muscle fluid volume are related to passive stiffness (Sleboda & Roberts, 2017; Sleboda et al., 2019), we did not measure changes in passive muscle (or total joint) stiffness. Since these measurements take time, and we did not want to interrupt the running protocol longer than needed for the MRI measurements, we did not include them in our experimental protocol. However, future studies could address this relationship with dedicated experimental designs. Finally, it might be beneficial to quantify the runners’ biomechanics during prolonged running. Recent research suggests that energy generation patterns within the lower extremity can change throughout a prolonged run (Sanno, Willwacher, Epro, & Brüggemann, 2018). In order to differentiate whether fluid shifts away from muscle tissue are due to dehydration or due to altered muscle activity, experimental designs should consider changes in running mechanics.

In conclusion, we provide evidence that calf muscle fluid volumes change throughout a prolonged running protocol, which might affect the force generation ability of muscle through biochemical and biomechanical pathways.

5 | PERSPECTIVE

Calf muscle volumes undergo initial increase and subsequent decrease during prolonged, moderate intensity running in young, healthy, recreational runners. Recent evidence suggests that muscle fluid volume changes can affect passive force development and therefore the physiological performance of muscles (Sleboda & Roberts, 2017;
Sleboda et al., 2019). Consequently, next to the well-known importance of fluid shifts for cell biochemistry and cardiovascular performance (Maughan, 2010), the biomechanical effects of fluid shifts should be considered during sporting activities. This finding might have important implications for understanding distance running performance as well as injury development. 

ACKNOWLEDGMENT 
We like to thank Richard L. Marsh, Brown University for a fruitful discussion on an early version of the paper. This work was partly funded through a grant from Brooks Running Inc., Seattle, WA, USA.

CONFLICT OF INTEREST 
None declared.

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How to cite this article: Willwacher S, Sleboda DA, Mählich D, Brüggemann G-P, Roberts TJ, Bratke G. The time course of calf muscle fluid volume during prolonged running. Physiol Rep. 2020;8:e14414. https://doi.org/10.14814/phy2.14414