Skeletal muscle ATP turnover and single fibre ATP and PCr content during intense exercise at different muscle temperatures in humans

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ABSTRACT

The effect of temperature on skeletal muscle ATP turnover, pulmonary oxygen uptake and single fibre ATP and PCr content was studied during intense cycling exercise. Six healthy male subjects performed 6 min intense (Δ50%LT-VO2peak) cycling, at 60 revs.min-1, under conditions of normal (N) and elevated muscle temperature (ET). Muscle biopsies obtained from the vastus lateralis at rest, 2 and 6 min were analysed for homogenate ATP, PCr, lactate and glycogen, allowing estimation of anaerobic ATP turnover. Freeze dried single fibres from biopsies were characterised according to their MHC composition [type I, IIA or IIAX] and analysed for ATP and PCr content. Pulmonary gas exchange was measured throughout. There was no difference in pulmonary oxygen uptake between the trials. The elevation of muscle temperature resulted in a lower (P<0.05) PCr content, higher (P<0.05) lactate content and greater (P<0.05) anaerobic ATP turnover after 2 min of exercise. There was no effect of temperature on these measures at 6 min. In single fibres it was observed that in ET there was a lower (P<0.05) PCr content in type I fibres after 2 min with no differences between conditions after 6 min. The present study demonstrates that elevation of muscle temperature results in a greater anaerobic ATP turnover and type I fibre PCr degradation during the initial 2 min of intense exercise.

INTRODUCTION

It has been well established that muscle temperature (Tm) is an important determinant of the mechanical and metabolic performance in both amphibian and mammalian skeletal muscle (e.g. (7; 35)). This is significant because muscle temperature can vary considerably depending on the environmental conditions and the metabolic heat liberated from the muscle itself (1; 37). Early work in this area by Edwards *et al* (13) showed, in humans, that during isometric exercise there was a greater anaerobic ATP turnover, when performed at elevated muscle temperatures. However, the majority of contractile activity, particularly during athletic events and daily activities, involves dynamic contractions and during isometric contractions energy turnover has been demonstrated, in frog muscle, to be different than during dynamic exercise (17).

In another investigation in humans it was found that during 2 min cycling exercise Febbraio et al (16) found that there was greater glycogenolysis, glycolysis and ATP hydrolysis when exercise was performed after passive warming of the active musculature. Further human work has demonstrated that during 20 min of moderate intensity cycling increasing skeletal muscle temperature results in an increased glycogenolysis, with no change in high energy phosphate degradation (42). During exercise of this duration, however, the main energy source is oxidative phosphorylation (33; 34). Investigations into the effects of muscle temperature on oxygen uptake have revealed somewhat ambiguous results. Indeed, it has been shown in two investigations, both in humans, that passively elevating muscle temperature has no appreciable effect on the oxygen uptake response during moderate and heavy cycling exercise (10; 30). However, Ferguson *et al* (18) found in humans that at 60 revs.min-1 there was an elevation in oxygen uptake and a decrease in mechanical efficiency, whereas at 120 revs.min-1 the converse was true. On a cautionary note Ferguson and colleagues used blood lactate as a measure of anaerobic ATP turnover (12). This may not reflect actual muscle ATP turnover since lactate can be taken up and metabolised by various tissues, including the heart and inactive muscles (20; 21).

It has been proposed that these differences may be due to a fibre type specific effect of alterations in skeletal muscle temperature (23; 39). It is known that phosphocreatine (PCr) splits immediately upon contraction in amounts energetically sufficient to account for the amount of external work done (28). Based on this we have previously employed single fibre measures of PCr to show a preferential effect of elevations in skeletal muscle temperature on human muscle fibres with a predominance of myosin heavy chain (MHC IIA). This effect was noted during a 6 s maximal sprint where the average pedal rate was 120-140 revs.min-1 (24). However at lower pedal rates (60-100 revs.min-1) it has been suggested that the effect of muscle temperature is directed at type I fibres (39). There is no data investigating the effect of passive elevation muscle temperature on the response of different fibre types during exercise, performed at relatively slow contraction frequencies.

Therefore, the aim of this study was to investigate the effects of passive elevation of Tm on skeletal muscle metabolism and single fibre metabolismduring intense cycling exercise performed at 60 revs.min-1. It was hypothesised that elevating Tm would lead to a greater anaerobic ATP turnover, concomitant with a greater PCr degradation, specifically in type I fibres after the elevation of Tm.

**METHODS**

# Subjects

Six healthy male subjects (age 25 ± 3 yrs, height 1.85 ± 0.06 m, mass 87 ± 16 kg; means ± S.D.) with no history of muscle or metabolic disorders, volunteered for the study. All subjects were habitually physically active but not specifically trained. The subjects were fully informed of the purposes, risks and discomfort associated with the experiment before providing written, informed consent. This study conformed to current local guidelines and the Declaration of Helsinki and was approved by the University of Strathclyde Ethics Committee.

**Pre-Experimental Procedures**

Prior to the experimental trials subjects’ O2peak and lactate threshold (LT) were determined on an electromagnetically braked cycle ergometer (Excalibur Sport, Lode, BV, Groningen) at 60 revs.min-1using a continuous incremental test to volitional fatigue, modified from the treadmill protocol described previously (Jones, 1998). Power output was increased every three minutes by between 20 and 35 W. A fingertip blood sample was taken at the end of each 3 min stage and analysed immediately for lactate concentration (Lactate Pro, Akray KDK, Koyota, Japan). HR and RPE were also recorded at the end of each stage with pulmonary gas exchange continuously measured breath-by-breath using an on-line gas analysis system (Oxycon Pro, Jaeger, Hoechberg, Germany). O2peak was taken asthe highest recording of O2 over a 30s period. Lactate threshold was calculated by simultaneously solving two linear regression equations relating blood lactate to workload following identification of the visually observed breakpoint in blood lactate by two experienced reviewers (3), and is expressed as power output at LT (W). O2 at LT was calculated by solving the linear regression equation relating workload to O2 using power output at LT as the x-variable. An external power output half way between LT and O2peak (Δ50 %) was then calculated and used in the subsequent experimental trials. At least 7 days later a familiarisation trial was carried out to fully habituate the subjects to the experimental exercise trials.

**Experimental protocol**

Subjects attended the laboratory on 4 separate occasions, each separated by a minimum of seven days, to perform a 6 min and a 2 min cycle exercise bout under normal (N) and elevated muscle temperature (ET) conditions. Prior to these trials participants were instructed to refrain from strenuous physical exercise, caffeine and alcohol for the preceding 48 hour period. During this period they were also instructed to record their dietary intake and replicate this prior to subsequent trials. The order of conditions was counterbalanced, with the 6 min trial always preceding the 2 min trial in each condition.

On each occasion subjects arrived at the laboratory following an overnight fast. On arrival they inserted a rectal thermistor probe to allow continuous monitoring of rectal temperature. In N, subjects then rested for 30 min at normal room temperature (20-22 °C) during which time the muscle biopsy site was prepared and a flexible muscle temperature probe inserted into the vastus lateralis. A resting biopsy was not taken in the 2 min trial to minimise the number of biopsies taken from each subject. Pilot work demonstrated that with the nutritional replication employed a second resting biopsy was not necessary. In ET, after insertion of the rectal probe muscle temperature was elevated through a combination of hot water and subsequently electrically heated blankets as described previously (23). The same biopsy and temperature probe preparation to the legs was performed by opening the electric blanket for a minimal duration to avoid excessive heat loss. The heating protocol lasted approximately 1 hour.

In the 6 min trials, under N or ET, a resting biopsy was taken at this point. Subjects then moved to the cycle ergometer where they sat at rest for 3 min (in ET the blankets remained wrapped around the legs). They then began cycling at the predetermined power output (Δ 50%) at 60 revs.min-1, for either 2 or 6 min. All subjects completed the full duration of every trial. Immediately after exercise, in all trials, a post exercise biopsy was taken with the subject remaining seated on the bike. Pulmonary gas exchange was recorded continuously at rest and throughout the 6 min trials (the mouthpiece was removed ~15 s before the end of exercise to allow muscle biopsies to be taken).

**Measurements**

*Muscle Temperature*

Muscle temperature (Tm) was measured with a flexible muscle temperature probe (Ellab, Copenhagen, Denmark) which was inserted to a depth of 3 cm in the medial portion of the vastus lateralis muscle. The probe was inserted through a flexible venflon cannula (18 G, Becton Dickinson, Dublin, Ireland) and advanced ~0.5 cm beyond the end of the cannula into the muscle.

*Muscle Biopsies*

Muscle samples were obtained from the medial part of the vastus lateralis using the needle biopsy technique (8). Briefly, after injection of local anaesthetic (1% lidocaine) an incision (~1 cm) was made through the skin and fascia. When two biopsies were taken in the same trial two separate incisions were made approximately 2-3 cm apart. The biopsy needle was introduced, sample collected and frozen immediately in liquid nitrogen and stored at -80 °C for subsequent analyses. At all time points, including after the cessation of exercise, samples were frozen in less than 10s, with no difference in time to freezing between treatments or timepoints. Contralateral legs were used for samplingin each of the N and ET trials, the order ofwhich was counterbalanced.

*Homogenate Muscle analyses*

The samples were freeze dried and dissected free of blood and connective tissue, powdered and prepared for metabolite analysis. Approximately 25mg of powdered tissue was extracted in 2ml of 0.5 M perchloric acid and 1 mM EDTA, neutralized to pH 7.0 with 2.2 M KHCO3, and stored at -80°C. PCr and ATP content was determined in duplicate (CV of 1.3 and 1.4 % for ATP and PCr, respectively) spectrophotometrically (26). Lactate content was determined in duplicate (CV of 1.5%) flourometrically (32). To determine muscle glycogen approximately 4mg tissue was extracted in 300μl 1 M HCl and determined in duplicate (CV of 1.3%) flourometrically (32).

*Single Fibre Analyses*

In four subjects, selected because all biopsies for those subjects were deemed a suitable size to allow a sufficient number of single fibres to be dissected, approximately 40-50 fragments (3-4mm) from each biopsy, were manually dissected under a low powered light microscope (Leica Microsytems, Wetzlar, Germany). Each fibre was then cut in half, with the first fragment used for myosin heavy chain (MHC) determination and the second used for single fibre ATP and PCr measurements.

MHC content was determined by SDS-PAGE using a method derived from Fauteck and Kandarian (14). For approximately each millimetre of fibre fragment, 10 µl of SDS buffer was added. Electrophoresis (Bio-Rad Mini-Protean 3) was then carried out on 6% (crosslinking 2.7%) polyacrylamide resolving gels with 4% (crosslinking 2.7%) stacking gels, which were electrophoresed at 8°C for 18 h at a constant 100 V. Protein bands were visualized by silver staining, using a method modified from Oakley et al (1980), and quantified by densitometry (Bio-Rad GS8000 calibrated densitometer). Fibres were classified as either type I (100% MHC I), IIA (100% MHC IIA) or IIAX (all IIAX hybrid fibres).

The second fibre fragment was weighed on a quartz fibre fish-pole balance (Lowry & Passonneau, 1972) calibrated by the spectrophotometrical determination of weighed p-nitrophenol crystals. Fibre fragments, weighing 1-5 µg, were then analyzed luminometrically for ATP and PCr content using the luciferase method described by Wibom et al (44). Briefly, each fibre was extracted in 200μl of trichloroacetic acid (2.5%) followed by neutralization in 20μl 2.2M KHCO3. 50 µl of the extract was then added to a sucrose buffer containing D-luciferin and the assay carried out on a luminometer (1251, Bio Orbit Oy, Turku, Finland). The coefficient of variation (CV) for duplicate determination of ATP and PCr content in the same fibre extracts (n=26) was 7.2 and 4.5 % respectively.

*Calculations*

Anaerobic ATP turnover was calculated as: ΔPCr + 1.5Δ muscle lactate + 2 ΔATP (40). The small quantity of ATP produced, or utilized, relating to the accumulation of glycolytic metabolites (e.g. pyruvate and glyceraldehyde-3-phosphate) are neglected as they represent <2 % of ATP turnover in all cases (41). The estimation of anaerobic ATP turnover is an underestimation since lactate efflux, nor the amount metabolised by other tissues, is not measured (2). Lactate release has, however, been shown to be the same when Tm is passively elevated by about 3°C during knee extensor exercise (19) so this will not significantly affect comparisons between conditions.

**Statistical analysis**

Data were analysed by either two-way (temperature and time) analysis of variance (ANOVA) with repeated measures or paired *t*-tests. Where a significant effect was detected, differences were located with *post-hoc t*-tests with Bonferroni correction. Significance was accepted at P<0.05. Data are presented as means ± S.D.

**RESULTS**

*O2peak and LT*

O2peak was 3.4 ± 0.5 l.min-1 and was achieved at an external power output of 281 ± 54 W. LT occurred at 164 ± 49 W (58 ± 11 % of O2peak), with the external power output required to elicit Δ50 % was calculated to be 219 ± 50 W. At exhaustion the average heart rate was 181 ± 12 bpm (92 ± 5 % of predicted max heart rate) and average RER was 1.19 ± 0.04.

*Dietary Data*

Energy intake for the 24h period prior to experimental trials was 2432 ± 204kcal, with 26 ± 3% fat, 59 ± 4% carbohydrate and 15 ± 1% fat.

*Rectal and muscle temperature*

The heating protocol resulted in a higher Tm prior to the onset of exercise (34.7 ± 1.1 °C in N vs. 37.4 ± 0.2 °C in ET; P<0.05). Rectal temperature was not different (P>0.05) between trials (37.1 ± 0.2 and 37.3 ± 0.2 °C in N and ET, respectively).

*Homogenate muscle metabolites and ATP turnover*

Elevation of Tm had no effect (P>0.05) on the resting content of any metabolites (Table 1). ATP content did not change during exercise under either condition.

After 2 min of exercise PCr content was lower (P<0.05) in ET compared to N, decreasing by 32.0 ± 8.3 mmol.kg-1 (dm) in N and by 37.8 ± 10.6 mmol.kg-1 (dm) in ET. PCr content continued to decrease (P<0.05) and was the same after 6 min in both conditions. Lactate content was higher (P<0.05) at 2 min in ET compared to N, increasing by 13.4 ± 3.9 mmol.kg-1 (dm) and 8.6 ± 3.6 mmol.kg-1 (dm), respectively. Lactate content continued to increase (P<0.05) with no further difference at 6 mins. Muscle glycogen content was decreased (P<0.05) as a result of exercise. There were no differences in glycogen content between N and ET at any time point. After 6 min of exercise there were no differences in the concentrations of metabolites between temperature conditions.

From these changes in metabolites it was calculated that during the first 2 min of exercise there was a 28% greater (P<0.05) total anaerobic ATP turnover during ET (Figure 1). This increase was the result of a greater (P<0.05) glycolytic ATP turnover and a greater (P=0.05) ATP turnover from PCr hydrolysis. Between 2-6 min as well as the overall period of exercise (0-6 min) elevation of Tm had no further effect on anaerobic ATP turnover, although there was a tendency (P=0.06) for a greater rate of glycolytic ATP turnover between 0-6 mins in ET, compared to N (Figure 1).

*Single fibre ATP and PCr content*

Elevation of Tm had no effect (P>0.05) on resting single fibre ATP content (Table 2). There was also no difference in resting ATP content between fibre groups. After 2 min of exercise, in both N and ET, ATP content was lower (P<0.01) compared to rest in all fibre types. Thereafter ATP content was only reduced in type IIA fibres in both N and ET. When muscle temperature was elevated ATP content was 3.0 ± 0.6 mmol.kg-1 (dm) lower (P<0.01), compared to N, in type I fibres after 6 min of exercise.

Elevation of Tm had no effect (P>0.05) on resting single fibre PCr content, with this being higher in type IIA and IIAX compared to type I fibres (Table 3). After 2 min of exercise PCr content was lower (P<0.01) in all fibre types in both N and ET. In N PCr content reduced (P<0.01) further from 2-6 min in type I and IIA fibres, with PCr content only reduced in IIA fibres at this time point.

The main finding of the single fibre analysis was that after 2 min of exercise there was a 6.7 ± 1.5 mmol.kg-1 (dm) lower (P<0.01) PCr content in type I fibres in ET compared to N. This reflects a 25% greater PCr degradation in type I fibres during ET, compared to N. There were no differences between conditions after 6 min of exercise.

*Pulmonary O2*

When pulmonary oxygen uptake was compared between N and ET no difference was observed at any time point. (Figure 2).

**DISCUSSION**

The present investigation has demonstrated that passively elevating the temperature of exercising muscle before the performance of sustained heavy exercise results in an increased anaerobic ATP turnover, with no differences in muscle glycogen, during the initial 2 min of exercise. The increase in anaerobic ATP turnover was associated with a greater PCr utilisation in fibres with a predominance of MHC I.

Previous work investigating the effects of elevating muscle temperature on anaerobic ATP turnover has demonstrated a greater anaerobic metabolism during intense isometric exercise (13), maximal sprint exercise (23) and intense cycling exercise (16). The present investigation observed similar results, although muscle glycogen levels in the current study were unaffected by temperature, during the initial 2 min of exercise, however when exercise continued there were no differences in anaerobic ATP turnover between temperature conditions. This is a similar finding to that of Ferguson *et al* (19) who did not observe an effect of passively increasing muscle temperature on energy turnover during 10 min of knee extensor exercise. It is worth pointing out that in that study muscle biopsies were only obtained at the end of exercise, potentially missing any temperature induced changes present in the initial few minutes of exercise.

The reason for the greater anaerobic ATP turnover observed at higher Tm is likely to be due to a greater cross-bridge cycling between actin and myosin. Indeed, Karatzaferi *et al*. (29) have clearly demonstrated the temperature dependency of cross-bridge cycling in relation to the force produced during the power stroke of the cycle. This itself is likely to be the result of an effect of the increase in temperature on mATPase activity (27; 43) as well as several other enzymes involved in ATP turnover. For example, creatine kinase is known to have an optimum temperature of around 42 °C (45) and its activity is, therefore, likely to be enhanced after heating. The finding of a similar glycogen breakdown when comparing N and ET was a surprising result. An increase in lactate accumulation with heating has been demonstrated in previous studies (13; 16), although interestingly other work has found an increase in glycogen breakdown without a concomitant increase in muscle lactate (42). The authors of the latter study suggested that this could be due to an increase in muscle lactate efflux in heated trials, although this was not directly measured. This assertion would not explain our findings and work from our group has found that lactate efflux from muscle isn’t altered with heating during exercise (19).

As the exercise duration proceeds beyond the initial period contribution from aerobic sources becomes more significant, as can be seen from the reduced anaerobic ATP turnover from 2-6 mins of exercise. Previous work has demonstrated that during sustained cycling at a similar intensity and pedal cadence (60 revs.min-1) passively elevating Tm resulted in a greater pulmonary O2 (18). The present study, however, found that elevated Tm does not increase pulmonary O2, an observation also made by Burnley *et al* (9) and Koga *et al* (30), albeit at slightly faster pedal rates, as well as during single leg dynamic knee-extensor exercise (19). The reason for these contradictory results is unclear as the mode and intensity of exercise were similar between these investigations. It had previously been suggested that elevations in Tm could increase O2 via a Q10 effect on mitochondrial respiration and/or through the back-leak of protons through the inner mitochondrial membrane. The current study would support the observation that mitochondrial function and physical performance only appear to be compromised at muscle temperature exceeding 40°C (15).

One possible explanation for the lack of Tm effect on pulmonary O2, as well as anaerobic ATP turnover as exercise progresses may be related to the exercise induced rise in Tm (37). It is possible that a temperature effect is observed at the onset of exercise but not at the end as Tm in N will approach the levels reached in ET, due to metabolic heat production. Whilst, due to technical difficulties, no measure of Tm was made post-exercise in the present study, previous work has shown that when Tm is raised by ~3°C prior to exercise it is still approximately 1.5 °C higher at the end of heavy knee-extensor (19) and cycling exercise (30). It may therefore be the case that this smaller Tm difference may not be of sufficient magnitude to alter energy turnover in the latter part of exercise, as was the case in the first two minutes.

Whilst the majority of previous research has focused on investigating the effect of alterations in muscle temperature in homogenate muscle samples it is important to take into account the different muscle fibres that have diverse contractile and metabolic properties. It has been estimated that at the workload of the current study the power required will be less than 50% of maximal power (38), which could likely be generated exclusively from type I fibres. However, it is clear from the single fibre data of the present study, that type II fibres are recruited during this exercise. It is therefore prudent to discuss the effect of temperature on different fibre types. Measurements of single fibre metabolism have now been used in a variety of studies and are providing more detailed information on the metabolic response to exercise (5; 6; 25). These methods are based on the knoweldege that the PCr content of single muscle fibres gives a good indication of the activity of muscle fibres (11; 28).

Based on the early findings of Ranatunga *et al* (36), in animal muscle, it was hypothesised that type I fibres were more sensitive to perturbations in temperature. This assertion was supported, in humans, by the findings of Sargeant and Rademaker (39) and Ferguson *et al* (18). However, both these studies relied on correlations between data to show a cause and effect, making the reliability of the thesis unclear. Furthermore, we have recently shown that during short-term (6 sec) maximal sprint exercise (cadence of ~160 revs.min-1) fibres with a predominance of MHC IIA have a greater PCr and ATP utilisation when muscle temperature is passively elevated (24). From this it appears likely that there may be a velocity dependent effect of muscle temperature on different fibre types. Such a view is supported by the current findings. During cycling at 60 revs.min-1, the current study has shown that, single fibre PCr utilisation was greater in type I fibres in the initial two minutes during ET. This coincides with the greater anaerobic ATP turnover observed in the analysis of muscle homogenates in the initial part of the exercise (Figure 1). One caveat to this analysis was that due to lack of muscle tissue collected we were only able to carry out the single fibre analysis in four subjects and so further work in a greater number of subjects is necessary to support these assertions. Furthermore it is worth pointing out that the results of the single fibres do not entirely match with the homogenate data. However, making comparisons between these two sets of data is problematic as the numbers of each fibre type used in the analysis doesn’t correlate with the average MHC composition of our subjects (e.g. 38.2 ± 12.5% MHC I, 58.7 ± 11.4% MHC IIA and 3.1 ± 2.7% MHC IIX).

The greater PCr utilisation in type I fibres may be explained further by considering the individual efficiency-velocity relationships of different fibre types. In the present study the pedal rate during cycling was maintained at 60 revs.min-1. This cadence is thought to be close to the optimum velocity for maximum efficiency of type I fibres (38). An increase in temperature would induce a shift to the right of the efficiency-velocity relationship for the recruited fibre population, reflecting the change in the force-velocity and power-velocity relationships of muscle that occurs when temperature is elevated (36). For type I fibres such a shift would mean the fibres would be working on the ascending curve of the efficiency-velocity relationship. This would therefore lead to a decrease in efficiency and an increased energy turnover for a given mechanical power output. On the other hand, at this cadence no such affect is seen in type II fibres. These fibres are likely, at 60 revs.min-1, to be operating at the lower part of the ascending limb of the efficiency-velocity curve where a rightward shift will have minimal effect.

From a practical view point it could be considered that as anaerobic energy turnover, and type I fibre metabolism, is elevated in the initial part of exercise in ET that warming the muscles before an athletic event may not be advised. There may be particularly deleterious effects for endurance athletes who have a high percentage of type I fibres (22). In support of this previous work has found that temperature induced changes in mechanical efficiency correlates with the proportion of type I fibres (4). In the current study if we estimated mechanical efficiency (Table 4), as we have previously (23), we did not observe any correlations between the magnitude of change in anaerobic ATP turnover or mechanical efficiency. The reasons for this are not clear but may be due to the participants in the study of Bell and Ferguson (2009) being female, although further work would be needed to confirm this. Another point of consideration for athletes would also be the contraction frequency at which exercise is performed as the response to changes in muscle temperature has been shown to be velocity dependent, with efficiency being higher at greater cadences (18). This may also explain why endurance athletes tend to adopt such high cadences which may have a beneficial effect on mechanical efficiency.The current study is not without limitations. It is well established now that a continuum of different fibre types with a variety of contractile properties exist (31). Whilst we have separated out fibres into three discreet groups, including hybrid fibres, there is still a relatively wide range of MHC type content within each group. Ideally, one would be able to acquire enough fibres to further subdivide each group into pure and hybrid fibre subtypes, allowing a more detailed analysis. This was not possible in the current study as the amount of tissue was limited and the dissection of the single fibres is a laborious and skilled task. A further limitation to the current investigation is the relatively low subject numbers, due to the laborious nature of the methods and the requirements for large amounts of quality muscle tissue. While the current single fibre analysis has provided valuable information on the metabolic response in single fibres to exercise at elevated muscle temperature, a greater number of subjects would have reduced the chances of a type II error.

In conclusion, the present study has demonstrated that passively increasing the temperature of the working muscle leads to an increase in anaerobic ATP turnover during the first two minutes of heavy exercise. This occurs alongside a greater PCr degradation in muscle fibres expressing a predominance of MHC type I. There was, however, no effect of elevating Tm on anaerobic ATP turnover as exercise progressed or on pulmonary O2 at any point throughout the exercise period.

FIGURE LEGENDS

Fig 1. Glycolytic, PCr and Total Anaerobic ATP turnover between 0-2, 2-6 and 0-6 min during heavy exercise under conditions of normal (N) and elevated (ET) muscle temperature. Values are means ± S.D. (N=6). Significant difference (P<0.05) between conditions denoted by †. Values are expressed as mmol.kg-1 (dm)s-1

Fig 2. Pulmonary O2 during 6 min of heavy exercise under conditions of normal (N, filled circles) and elevated (ET, closed circles) muscle temperature.

TABLE LEGENDS

Table 1. Muscle metabolites before and after 2 and 6 min of heavy exercise under conditions of normal (N) and elevated (ET) muscle temperature. Values are means ± S.D. (N=6). Significant difference (P<0.05) from preceding time point denoted by \*. Significant difference (P<0.05) between conditions denoted by †. Values are expressed as mmol.kg-1 (dm) except for glycogen which is in mmol glycosyl units.kg-1 (dm).

Table 2. Single fibre ATP content in MHC characterised muscle fibres under control (N) and elevated (ET) muscle temperatures.Values are expressed as mean ± S.D with the number of fibres analyzed represented by (n). Significant difference (P<0.01) from preceding time point denoted by \*. Significant difference (P<0.01) between conditions denoted by †. Values are mmol.kg.-1 (dm).

Table 3. Single fibre PCr content in MHC characterised muscle fibres under control (N) and elevated (ET) muscle temperatures. Values are expressed as mean ± S.D with the number of fibres analyzed represented by (n). Significant difference (P<0.01) from preceding time point denoted by \*. Significant difference (P<0.01) between conditions denoted by †. Values are mmol.kg.-1 (dm).

Table 4. Estimates of mechanical efficiency during exercise under control (n) and elevated (ET) muscle temperatures. Values are expressed as mean ± S.D. Significant difference (P<0.01) between conditions denoted by \*. TABLE 1

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  |  | Rest | 2 min | 6 min |
| ATP | N | 22.7 ± 1.8 | 21.7 ± 1.9 | 19.8 ± 1.8 |
| ET | 23.2 ± 0.8 | 22.8 ± 2.0 | 21.5 ± 1.7 |
| PCr | N | 72.9 ± 7.0 | 40.9 ± 8.3\* | 29.4 ± 8.4\* |
| ET | 72.8 ± 9.7 | 35.0 ± 8.8\*† | 26.9 ± 7.4\* |
| Lactate | N | 5.4 ± 0.7 | 14.0 ± 3.8\* | 26.2 ± 4.4\* |
| ET | 6.0 ± 0.6 | 18.4 ± 4.4\*† | 32.6 ± 7.2\* |
| Glycogen | N | 399.7 ± 139.0 | 332.8 ± 79.1\* | 242.7 ± 78.8\* |
| ET | 417.5 ± 125.7 | 322.3 ± 97.9\* | 262.7 ± 53.3 |

TABLE 2

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | N |  |  | ET |  |  |
|  | 0 min  (n) | 2 min  (n) | 6 min  (n) | 0 min  (n) | 2 min  (n) | 6 min  (n) |
| I | 25.1 ± 4.1 (19) | 23.4 ± 4.1\* (42) | 22.8 ± 4.1 (30) | 26.2 ± 4.8 (17) | 21.8 ± 3.1\* (32) | 19.8 ± 4.1† (30) |
| IIA | 26.5 ± 3.7 (29) | 23.7 ± 4.8\* (36) | 20.6 ± 4.4\* (32) | 26.1 ± 3.1 (28) | 23.1 ± 3.1\* (28) | 21.2 ± 4.6\* (30) |
| IIAX | 26.6 ± 3.6 (37) | 22.6 ± 1.9\* (17) | 23.3 ± 4.1 (15) | 27.9 ± 3.0 (23) | 23.0 ± 4.6\* (24) | 20.7 ± 6.1 (16) |

TABLE 3

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | N |  |  | ET |  |  |
|  | 0 min  (n) | 2 min  (n) | 6 min  (n) | 0 min  (n) | 2 min  (n) | 6 min  (n) |
| I | 69.8 ± 9.0 (19) | 24.2 ± 9.7\* (42) | 15.5 ± 10.1\* (30) | 73.0 ± 18.7 (17) | 17.7±10.3\*† (32) | 15.4 ± 7.9 (30) |
| IIA | 89.1 ± 12.2 (29) | 34.2 ± 14.9\* (36) | 21.4 ± 10.0\* (32) | 91.8 ± 14.4 (28) | 32.8 ± 14.3\* (28) | 23.3 ± 17.7\* (30) |
| IIAX | 89.5 ± 15.5 (37) | 35.9 ± 29.0\* (17) | 35.5 ± 16.8 (15) | 94.9 ± 15.8 (23) | 43.8 ± 13.3\* (24) | 42.6 ± 14.1 (16) |

Table 4.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
|  | N |  |  | ET |  |  |
|  | 0-2 min | 2-6 min | 0-6 min | 0-2 min | 2-6 min | 0-6 min |
| Mechanical Efficency | 28.1 ± 3.4 | 24.0 ± 1.6 | 25.7 ± 1.8 | 25.4 ± 3.1\* | 24.8 ± 2.4 | 25.4 ± 2.1 |

Reference List

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