Adaptation of mitochondrial ATP production in human skeletal muscle to endurance training and detraining

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WIBOM, R., E. HULTMAN, M. JOHANSSON, K. MATHEREI, D. CONSTANTIN-TEODOSIU, AND P. G. SCHANTZ. Adaptation of mitochondrial ATP production in human skeletal muscle to endurance training and detraining. J. Appl. Physiol. 73(5): 2004-2010, 1992.—The adaptation of mitochondrial ATP production rate (MAPR) to training and detraining was evaluated in nine healthy men. Muscle samples (~60 mg) were obtained before and after 6 wk of endurance training and after 8 wk of detraining. MAPR was measured in isolated mitochondria by a bioluminometric method. In addition, the activities of mitochondrial and glycolytic enzymes were determined in skeletal muscle. In response to training, MAPR increased by 70%, with a substrate combination of pyruvate + palmitoyl-L-carnitine + α-ketoglutarate + malate, by 50% with only pyruvate + malate, and by 92% with palmitoyl-L-carnitine + malate. With detraining, MAPR decreased by 12–28% from the posttraining rate (although not significantly for all substrates). No differences were found when MAPR was related to the protein content in the mitochondrial fraction. The largest increase in mitochondrial enzyme activities induced by training was observed for cytochrome-c oxidase (78%), whereas succinate cytochrome c reductase showed only an 18% increase. The activity of citrate synthase increased by 40% and of glutamate dehydrogenase by 45%. Corresponding changes in maximal O₂ uptake were a 9.6% increase by training and a 6.0% reversion after detraining. In conclusion, both MAPR and mitochondrial enzyme activities are shown to increase with endurance training and to decrease with detraining.

β-oxidation; carbohydrate metabolism; fat metabolism; luminescence; maximal oxygen uptake; oxidative enzymes; oxidative phosphorylation

SKELETAL MUSCLE is a remarkably plastic tissue that responds rapidly to increases and decreases in the degree of physical activity. Changes in metabolic capacity have been related to mitochondrial density (22), enzyme activities (20, 27), and rate of tissue O₂ uptake (13, 20). The studies performed are in agreement that contractile activity and inactivity lead to increases and decreases, respectively, in all these forms of mitochondrial expression. It has long been known that the maximal O₂ uptake increases after training. This was shown to be associated with increases in mitochondrial enzymes in human muscle in the late 1960s and early 1970s (for references see Ref. 27). Thus the activities of cytochrome-c oxidase and succinate dehydrogenase were found to be increased by training. However, no direct studies of mitochondrial respiration in human muscle before and after training have been done.

The reason for this is that the classic method for measuring mitochondrial respiration (and ATP formation rate) requires the use of muscle of ~0.5–1 g tissue obtained by surgical procedures (7, 14). This amount of tissue cannot easily be obtained repeatedly in healthy subjects. However, a sensitive bioluminometric method has recently been developed that permits mitochondrial ATP production rates (MAPR) to be determined by using different substrates (36, 37). The amount of tissue needed is 40–60 mg, a sample size that can be obtained repeatedly by the percutaneous needle biopsy technique described by Bergström (3). With this method it was recently shown that the MAPR was ~80% higher in well-trained than in sedentary subjects (36).

Our aim in this study was to evaluate the response of MAPR to endurance training and subsequent detraining. At the same time we investigated several commonly used mitochondrial marker enzymes, thus enabling comparative analyses of these variables to be made.

MATERIALS AND METHODS

Subjects. Nine healthy untrained men [age 20 ± 1 (SD) yr, height 1.81 ± 0.07 m, and weight 74 ± 5 kg] volunteered for the study. The subjects were performing their military service, and all were carrying out duties of a sedentary character, such as office work or car driving. They had not done any regular physical training for the 3 mo immediately preceding this study. They all were fully informed about the purpose of the study and the risks and discomfort associated with the experiments before they volunteered to participate. The study was approved by the Ethics Committee of the Karolinska Institute.

Experimental protocol. The experiments were carried out during a 9 wk period. In the first 6 wk the subjects underwent an endurance training program on a bicycle ergometer. This was immediately followed by a detraining period of 3 wk without any physical activities other than those involved in everyday living. Submaximal and maximal work tests were performed, and muscle biopsy samples were taken before and after the training, as well as after the detraining.

Training protocol. The subjects trained, on the average, 4 × 36 min/wk for 6 wk using bicycle ergometers (Monark, Varberg, Sweden or Cardionics, Stockholm, Swe-
den). The absolute work load and the length of the training sessions were increased from 175 ± 24 W and 30 min/day in week 1 to 200 ± 30 W and 40 min/day in week 6. The work load was adjusted so as to induce an O₂ uptake corresponding to 70% of the maximal O₂ uptake during the whole training period. This was achieved by monitoring each subject’s heart rate (Sporttester PE 3000, Polar Electro, Kempele, Finland) during training and comparing it with the individual heart rate per O₂ uptake relationship observed in laboratory experiments. The blood lactate levels (33) after 15 min of training were measured during weeks 2 and 6 and corresponded to 4.1 ± 1.2 and 4.6 ± 1.9 mmol/l (n = 7), respectively.

Submaximal and maximal cycle ergometer tests. The subjects were made familiar with the testing procedures on a separate occasion before the experiments were started. Before and after the training period, as well as after the detraining period, one maximal and two submaximal (100 W, 150 W) work loads were performed on a cycle ergometer (Monark) at 60 rpm. O₂ uptake, heart rate, and lactate concentration in blood were determined for each work load.

O₂ uptake was determined by the Douglas bag technique (1a), and the O₂ and CO₂ contents in the expired air were analyzed in a mass spectrometer (Centronic 200). Heart rate was measured with a pulse watch (Sporttester PE 3000). The blood lactate concentration was analyzed by a slight modification of Barker-Summerson’s method (33). The linear relationship between O₂ uptake and heart rate was determined for each subject at pretraining and was used to calculate the absolute work load during training.

Muscle biopsy sampling. The muscle biopsy samples were taken percutaneously under local anesthesia from the middle portion of the vastus lateralis, using a Bergström-Stille biopsy needle (6 mm diam) (3). The muscle samples, weighing ~100 mg, were dissected free from a fat and connective tissue on a glass plate cooled on ice. The samples were divided into four portions: two were frozen in liquid nitrogen and stored at -80°C for up to 3 mo for enzyme assays.

**Determination of MAPR.** The MAPR was determined by the bioluminescence technique, as originally described for rat skeletal muscle (37) and later applied to human skeletal muscle (36). Mitochondria were isolated from the middle portion of the vastus lateralis, using a Bergström-Stille biopsy needle (6 mm diam) (3). The muscle samples, weighing ~100 mg, were dissected free from visible fat and connective tissue on a glass plate cooled on ice. The samples were divided into four portions: two were frozen fresh for determination of MAPR and succinate-cytochrome c reductase (SCR) + NADH-cytochrome c reductase (NCR), respectively; see below), and two were frozen in liquid nitrogen and stored at ~80°C for up to 3 mo for enzyme assays.

**Calculations of MAPR.** The MAPR was determined as millimoles ATP per minute per liter mitochondrial suspension. The intramitochondrial activity of glutamate dehydrogenase (GDH) was determined in the same suspension. Total GDH activity in the crude muscle homogenate was also determined (see below). The relative mitochondrial yield was calculated from the intramitochondrial GDH activity in the suspension and the GDH activity in the whole muscle, and with the use of this ratio the MAPR was referred to the muscle mass (mmol · min⁻¹ · kg⁻¹) (36).

**Protein.** MAPR was also related to the content of alkali soluble protein in the mitochondrial suspension, as determined by the method of Lowry et al. (24).

**Enzyme assays.** SCR (EC 1.3.99.1) and NCR (EC 1.6.99.3). Fresh muscle (~15 mg wet wt) was homogenized in a Potter-Elvehjem homogenizer (1:50 wt/vol) in a solution consisting of (in mmol/l) 100 KCl, 50 tris(hydroxymethyl)aminomethane, 5 MgCl₂, 1.8 ATP, and 1 EDTA, pH 7.2. SCR was spectrophotometrically determined at 25°C by the method of Cooperstein et al. (11). NCR was essentially determined as SCR, but the reaction was started with NADH (0.5 mmol/l) instead of succinate. The extinction coefficient of cytochrome c was taken as 21.1 l · mmol⁻¹ · cm⁻¹ (550 nm).

**GDH (EC 1.4.1.4).** Pieces of frozen muscle (~5 mg wet wt) were thawed and homogenized in a Potter-Elvehjem homogenizer. In the mitochondrial suspension the GDH activity was determined with and without the disruption of intact mitochondria, and the intramitochondrial GDH activity was calculated. The extramitochondrial fraction arises from mitochondria disrupted during the preparation of mitochondria. The procedures were described in detail in Ref. 36. The GDH activity was analyzed spectrophotometrically at 35°C (31).

For the remaining enzymes a frozen muscle sample (~20 mg wet wt) was thawed and homogenized (1:25 wt/vol) in a Potter Elvehjem homogenizer containing ice-cold potassium phosphate buffer (0.3 mol/l) and bovine serum albumin (0.05%, wt/vol), pH 7.7. The homogenate was stored at ~80°C for subsequent analyses of the following enzymes.

**6-Phosphofructokinase (PFK, EC 2.7.1.11), β-hydroxyacyl-CoA dehydrogenase (HAD, EC 1.1.1.35), and citrate synthase (CS, EC 4.1.3.7).** PFK, HAD, and CS were determined spectrophotometrically at 25°C, as described by Opie and Newsholme (26), Bass et al. (2), and Alp et al. (1), respectively.

**Malate dehydrogenase (MDH, EC 1.1.1.37).** MDH was determined spectrophotometrically at 25°C by the method of Bischer et al. (5). The cytoplasmic and mitochondrial isoenzymes of MDH (cMDH, mMDH, respectively) were assayed as described in detail by Schantz (28).

**Cytochrome c oxidase (COX, EC 1.9.3.1).** COX was determined polarographically at 30°C by the method of Tottmar et al. (31).

**Statistical analyses.** All statistical calculations were based on one-way analysis of variance (ANOVA). Differences were determined with Scheffe’s test, and P < 0.05
was considered significant. Results are presented as means ± SD.

RESULTS

Posttraining and detraining values are described here in terms relating to pretraining levels. Only significant differences are given here. Relative changes with detraining were calculated from the posttraining value.

Physiological responses to training and detraining. The responses in heart rate and blood lactate concentration to sub maximal and maximal work loads, as well as the maximal O2 uptake, are given in Table 1. The pretraining maximal O2 uptake was 44 ± 4 ml·kg⁻¹·min⁻¹. The 6 wk of endurance training and subsequent 3 wk of detraining resulted in an increase (9.6 ± 2.3%) and decrease (6.0 ± 6.7%), respectively, in the maximal O2 uptake, expressed as liters per minute. Training resulted in a significantly lower heart rate and blood lactate concentration in response to the higher of the two sub maximal work loads, changes that were not reversed by the detraining (Table 1).

Mitochondrial yield. The protein yield of the mitochondrial suspension (determined as g protein isolated/kg muscle homogenized) was 3.90 ± 0.64 and 6.88 ± 1.20 g/kg (n = 9) before and after training, respectively, and corresponded to an average increase of 80 ± 39%. After detraining the protein yield was 5.60 ± 1.25 g/kg (n = 8), which was 46 ± 35% above the pretraining level.

The efficiency of the isolation procedure of mitochondria was determined as the percent intramitochondrial protein was used as the reference base.

MAPR in muscle. With training the MAPR, expressed as liters per minute. Training resulted in a significantly different from the increase in P+M, 50 ± 36% (P < 0.001). No other differences in increases were found between the substrates. Malate is added to the reaction mixtures to maintain a high intramitochondrial concentration of this substance, which enables the tricarboxylic acid cycle to function. After detraining, the MAPR of the substrates PPKM, T+A, and S+R decreased significantly (17 ± 16% to 28 ± 16%) compared with after training. After detraining, the MAPRs were still higher than the pretraining level for the following substrates: PPKM, 40 ± 35%; T+A, 37 ± 32%; α-KG, 41 ± 46%; and PC+M, 70 ± 54%.

The MAPRs were also related to the protein content in the mitochondrial suspension (Table 3). No significant differences were found between the untrained, trained, and detrained conditions when the mitochondrial protein was used as the reference base.

Enzyme activities in muscle. The enzyme activities before and after training and after detraining are presented in Table 4. Enzyme activities of cMDH and PFK were not changed by training and detraining, except for a 10 ± 8% increase in PFK after detraining. As expected, the training-induced increases in the activities of mMDH, CS, and GDH were 28 ± 28, 43 ± 36, and 47 ± 25%, respectively. After detraining, the activities were still 39 ± 40 and 30 ± 32% above the pretraining levels for CS and GDH but not for mMDH. No significant changes were seen in the activity of HAD.
The lack of homogeneity in the muscle tissue analyzed. mmol ATP
ANOVA (P < 0.05) between * before and after training, t after training
mitochondria having different localizations within the
MAPR and enzyme activities can be due to the muscle
found to be 10% in two samples from the same leg (36). A highly active group in the previous study (36).
noticed for enzyme activities (16). lism during submaximal work in response to endurance
changes in enzyme activities reflect changes in MAPR,
changes in enzyme activities and after detraining. These changes were highly active subjects (mainly elite ice-hockey players)
large variations. To some extent, this can be explained by
parameters investigated showed relatively small increases. To some extent, this can be explained by the
An increased contribution of fat to the energy metabo-
and after detraining, and $ before training and after detraining.
membrane-bound enzyme complexes varied from 21 ±
21% for SCR and 48 ± 47% for NCR up to 78 ± 36% for
COX. No significant changes occurred with detraining.
However, after detraining, only the COX activity was sig-
nificantly higher (50 ± 39%) than the pretraining level.
Comparison between changes in enzyme activities and
MAPR. Several of the enzymes included in this study are
used as markers for different mitochondrial processes. It
is therefore of interest to estimate to what extent
changes in enzyme activities reflect changes in MAPR.

<table>
<thead>
<tr>
<th>Enzymes</th>
<th>Before Training</th>
<th>After Training</th>
<th>After Detraining</th>
</tr>
</thead>
<tbody>
<tr>
<td>PFK</td>
<td>36.7±3.2</td>
<td>34.2±7.6</td>
<td>40.1±5.6†‡</td>
</tr>
<tr>
<td>cMDH</td>
<td>161±25</td>
<td>149±37</td>
<td>103±34</td>
</tr>
<tr>
<td>mMDH</td>
<td>80±14</td>
<td>95±29</td>
<td>80±15</td>
</tr>
<tr>
<td>HAD</td>
<td>16±3.0</td>
<td>20±3.3</td>
<td>18±3.2</td>
</tr>
<tr>
<td>CS</td>
<td>13±2.7</td>
<td>18±3.3*</td>
<td>16.4±1.8†</td>
</tr>
<tr>
<td>GDH</td>
<td>1.33±0.17</td>
<td>1.94±0.24*</td>
<td>1.69±0.22†</td>
</tr>
<tr>
<td>NCR</td>
<td>5.1±1.6</td>
<td>7.8±1.5*</td>
<td>7.3±1.0</td>
</tr>
<tr>
<td>SCR</td>
<td>2.46±0.34</td>
<td>2.91±0.24*</td>
<td>2.67±0.38</td>
</tr>
<tr>
<td>COX</td>
<td>1.84±0.27</td>
<td>3.15±0.46*</td>
<td>2.71±0.60‡</td>
</tr>
</tbody>
</table>

Values are means ± SD in mmol·min⁻¹·kg muscle⁻¹; n = 9 men
(except for NCR, SCR, and COX, where n = 8 men). PFK, phospho-
fructokinase; cMDH, mMDH, cytoplasmic and mitochondrial isoen-
zymes of malate dehydrogenase, respectively; HAD, β-hydroxyn-a-
CoA dehydrogenase; CS, citrate synthase; GDH, glutamate dehydro-
gease; NCR, SCR, NADH and succinate cytochrome c reductase,
respectively; COX, cytochrome-c oxidase. Significant differences with
ANOVA (P < 0.05) between * before and after training, † after training
and after detraining, and $ before training and after detraining.

Discussion
In the present study a 6-wk endurance training pro-
gram was followed by 3 wk of detraining. Classic effects
of physical training were seen, e.g., increased maximal O2
uptake and decreased heart rate and blood lactate levels
at submaximal work (1a). An increase of 9% and a
decrease of 6% in maximal O2 uptake was found in response
to the training and detraining, respectively. Further-
more, the well-known pattern of changes in oxidative en-
zymes induced by physical activity and inactivity was es-
sentially reproduced (18, 29). All these findings are in
accord with previous training and detraining studies, and
the intended effects of the training were thus attained.

In the present investigation mitochondria were iso-
lated from muscle samples obtained by a percutaneous
needle biopsy technique. The weight of the samples was
~60 mg. After the 6-wk training period the mitochondrial
fraction had increased by ~80%, measured as pro-
tein content in the suspension. After the detraining pe-
riod, the content was 46% higher than the pretraining content. The efficiency of the isolation procedure (mea-
sured as the intramitochondrial GDH activity in the
mitochondrial suspension compared with that in the muscle
sample) was constant: on the average it was 26–27% on
the three measurement occasions. The changes in mito-
chondrial protein (although the change with detraining
was not significant) were matched by similar changes in
MAPR with the different substrate combinations, indi-
cating that the MAPR, expressed per gram mitochondrial
protein, was unchanged (see Table 3).

These findings are in conformity with studies in ani-
mals. Holloszy (20) and Davies et al. (13) reported 60–
70% increases in mitochondrial protein with physical
training of rats. Such changes were directly related to
increases in mitochondrial respiration, and thus the res-
piration, expressed per unit of mitochondrial protein,
was unchanged.

The present results bear out the observation in a pre-
vious cross-sectional study where a group of physically
highly active subjects (mainly elite ice-hockey players)
showed a 67% higher MAPR with PPKM than a group of
sedentary subjects (36). The 70% increase in the corre-
sponding MAPR is remarkable in light of the short train-
ing period (6 wk). The subjects reached a rate of 12.4
mmol ATP·min⁻¹·kg⁻¹ after training, which is compara-
ble to the level of 11.0 mmol ATP·min⁻¹·kg⁻¹ for the
highly active group in the previous study (36).

An increased contribution of fat to the energy metabo-
ism during submaximal work in response to endurance
training has been shown (9, 17, 19). The mechanism for
this is not clear, but it has been viewed as a result of the
total increase in mitochondrial density as such (15, 21).
An increase in the mitochondrial density would lead to an enhanced muscle capacity to phosphorylate ADP, which would reduce the ADP content in muscle during submaximal exercise. The effect of the lower level of ADP and secondarily AMP would also give a lower stimulation of glycolysis at the trained state. The contribution of fat to the energy metabolism during exercise would thereby automatically increase, even without a change in the mitochondrial composition (15). However, specific changes in various mitochondrial functions induced by training cannot be excluded.

Of interest are the present results showing that the largest increase in MAPR occurred with the substrate PC+M (92%). The substrate P+M gave a lower increase (50%). Similar results have been obtained in a study of rats where the oxidation rate of PC+M in homogenates of whole muscle was increased by 127% in the endurance-trained group compared with the untrained group, whereas the corresponding value for P+M was 67% (13). In the same report it was calculated that the contribution of lipid oxidation during exercise was three times higher in a group of trained than in a group of untrained rats. The difference in increase of MAPR between the two substrates is surprising as the catalytic product in the mitochondria for both substrates is acetyl-CoA. The difference would consequently depend on the rate of acetyl-CoA formation from the two substrates, being less increased from pyruvate than from palmitoyl-L-carnitine. In a training study of rats, Molé et al. (25) found 100% increases of the activity of the enzymes palmitoyl-CoA synthetase, carnitine palmitoyl-transferase, and palmitoyl-CoA dehydrogenase. This would give a similar increase of the capacity to form acetyl-CoA from palmitoyl-L-carnitine. The rate of pyruvate utilization is dependent on the pyruvate uptake by the mitochondria and the activity of pyruvate dehydrogenase (PDH). The PDH activity is low in resting muscle, and only a few studies on the effect of training have been performed. Thus a training study of rats showed a 70% increase of the active form of PDH in resting muscle but no increase of total PDH (4). Similar changes were observed in human muscle by Ward et al. (35).

In a recent study by Constantin-Teodosiu et al. (10) it was shown that exercise with submaximal work loads increased the PDH activity (the active form) by four to five times, with a similar increase of acetyl-CoA formation from pyruvate. The effect of exercise on the availability of acetyl-CoA from pyruvate should be taken into account when pyruvate oxidation in exercising humans is compared with values observed in mitochondrial suspensions obtained from resting muscle. The relationship between carbohydrate and fat utilization in exercising muscle can consequently not be established from in vitro studies of oxidative capacities. However, the large increase in the capacity of isolated mitochondria to produce ATP from fat could have an impact on fat utilization during exercise after training.

Analyses of marker enzyme activities for estimating the training-induced increases in the MAPR have frequently been used. On the basis of average values it appears from this study that GDH, COX, and CS are suitable marker enzymes, whereas SCR and HAD are poor mitochondrial markers. The smaller training-induced increase in HAD than in other mitochondrial enzymes has been observed previously (6, 29, 30), and the striking difference between changes in HAD activity and the rate of ATP production from palmitoyl-L-carnitine further indicate that HAD activity also does not accurately reflect the mitochondrial capacity for fat utilization.

The activities of mMDH and NCR were not compared with the MAPR because mMDH is a component in the malate-aspartate shuttle and therefore also depends on
cytosolic factors, and the method for NCR determination used in the present study was not specific for the mitochondrial enzyme activity (32).

To our knowledge no other studies have been presented concerning the effects of detraining on O₂ consumption or the ATP production rate by isolated mitochondria. In the present study a decline was observed in the MAPR of all substrates (12-28%) over a 3-wk period of detraining, although it was not significant in some cases. Costill et al. (12) measured the respiratory capacity, with pyruvate as the substrate, in crude homogenates of biopsy samples from the posterior deltoid muscle of swimmers during detraining. Even during the 1st wk of detraining they noted a 50% decline, but no other changes occurred during the subsequent 3 wk. The decline was larger than in the present study, which may be related to the fact that Costill et al. studied a nonpostural muscle, whereas we studied a postural muscle.

The decreases in MAPR with detraining were paralleled by decreases in mitochondrial enzyme activities, as reported previously (8, 18, 29). The rates of retrogression varied, however, between the different studies. For example, Henriksson and Reitman (18) found that the COX activity had returned to the pretraining level 2 wk after the cessation of training, whereas both the present results and those of Klausen et al. (23) indicate a slower rate of retrogression.

In conclusion, this study further extends our knowledge of the adaptability of the mitochondrial function in human skeletal muscle. For the first time it is shown that the MAPR in human muscles increases with physical training and decreases with detraining. This is matched in all essentials by changes in mitochondrial enzyme activities. The largest increase in the MAPR after training was observed when PC+M was used as the substrate.

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