Lipoprotein metabolism influenced by training-induced changes in human skeletal muscle.

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The influence of training-induced adaptations in skeletal muscle tissue on lipoprotein metabolism was investigated in six healthy men. The knee extensors were studied at rest and during exercise after 8 wk of dynamic exercise training of the knee extensors of one leg, while the other leg served as a control. The trained and nontrained thighs were investigated on different occasions. In the trained knee extensors, muscle (m) lipoprotein lipase activity (LPLA) was 70 +/- 29% higher compared with the nontrained (P less than 0.05), and correlated positively with the capillary density (r = 0.84). At rest there was a markedly higher arteriovenous (A-V) VLDL triacylglycerol (TG) difference over the trained thigh, averaging 55 mumol/liter (range 30-123), than over the nontrained, averaging 30 mumol/liter (4-72). In addition to the higher LPLA and VLDL-TG uptake in the trained thigh, a higher production of HDL cholesterol (C) and HDL2-C was also observed (P less than 0.05). Positive correlations between m-LPLA and A-V differences of VLDL-TG (r = 0.90; P less than 0.05) were observed only in the trained thigh. During exercise with the trained thigh the venous concentration of HDL2-C was invariably higher than the arterial, and after 110 min of exercise a production of 88 mumol/min (54-199) of HDL2-C was revealed. Even though a consistent degradation of VLDL-TG was not found during […]

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Abstract

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In the trained knee extensors, muscle (m) lipoprotein lipase activity (LPLA) was 70±29% higher compared with the nontrained (P < 0.05), and correlated positively with the capillary density (r = 0.84).

At rest there was a markedly higher arteriovenous (A-V) VLDL triacylglycerol (TG) difference over the trained thigh, averaging 55 μmol/liter (range 30–123), than over the nontrained, averaging 30 μmol/liter (4–72).

In addition to the higher LPLA and VLDL-TG uptake in the trained thigh, a higher production of HDL cholesterol (C) and HDL2-C was also observed (P < 0.05). Positive correlations between m-LPLA and A-V differences of VLDL-TG (r = 0.90; P < 0.05) were observed only in the trained thigh.

During exercise with the trained thigh the venous concentration of HDL2-C was invariably higher than the arterial, and after 110 min of exercise a production of 88 μmol/min (54–199) of HDL2-C was revealed. Even though a consistent degradation of VLDL-TG was not found during exercise, the total production of HDL-C across the trained and nontrained thigh, estimated from A-V differences times venous blood flow for the whole exercise period, correlated closely with the total estimated degradation of VLDL-TG (r = 0.91).

At the end of 2 h of exercise m-LPLA did not differ from the preexercise value in either the nontrained or the trained muscle.

We conclude that changes in the lipoprotein profile associated with endurance training to a large extent are explainable by training-induced adaptations in skeletal muscle tissue.

Introduction

Several studies have shown that the pattern of serum lipoproteins is related to the level of habitual physical activity. Results of longitudinal training studies strongly suggest that lipoprotein changes can be induced in initially sedentary people by physical training in the absence of changes in body weight (1, 2). Furthermore, in a recent investigation of well-trained men with high serum levels of HDL a further increase in HDL was noted during 6 mo of enhanced training (3). The HDL fraction consists of two major components, HDL2 and HDL3. Very physically active people have higher HDL2 levels than sedentary subjects (4). After conditioning for 10 wk a significant rise in HDL2-cholesterol (C) concentration was observed in initially sedentary men without weight changes, whereas HDL3-C tended to decrease (5). The mechanisms by which exercise and training influence the serum lipoprotein pattern have not been fully clarified. It is known, however, that HDL2 is derived from HDL3 during the hydrolysis of triacylglycerol (TG)-rich lipoprotein by the activity of lipoprotein lipase (LPLA) (6). Furthermore, higher LPLA is found in physically active than in sedentary men (7). Therefore, training-induced increase in muscle (m) LPLA might provide the functional basis for an increased formation of HDL-C in trained subjects.

Thus, the present study was designed to test the hypothesis that training-induced changes in skeletal muscle may contribute to the altered serum lipoprotein pattern. To approach this problem in man we chose to compare, in the same individual, the lipoprotein metabolism in the femoral quadriceps muscle of one leg after an 8-wk period of endurance dynamic knee extension exercise training with that in the contralateral, nontrained quadriceps muscle.

Methods

Experimental model

Subjects were seated on a specially designed ergometer that permitted exercise to be confined to the quadriceps femoris muscle group. (Fig. 1). This experimental model approaches an ideal model in man for studies of muscle alterations induced by physical training. The model allows direct measurements of blood flow and arteriovenous (A-V) differences over the contracting muscle, and muscle biopsies are easy to obtain. The knee extensor exercise is performed dynamically with 60 contractions/min. The contractions cause the lower part of the leg to move from 90 to 170° flexion. A previously published paper details the design of the ergometer and documents the evidence that exercise is confined to the knee extensors (8).

Subjects

The study was performed on six healthy, normal men with an average age of 23 yr (range 21–26) and average weight of 75 kg (range 69–81). Their maximal total body oxygen uptake (V02) during two-legged bicycle exercise averaged 3.65 liters/min (range 3.35–3.84). The subjects

1. Abbreviations used in this paper: A-V, arteriovenous; C, cholesterol; LPLA, lipoprotein lipase activity; m-LPLA, muscle LPLA; TG, triacylglycerol; V-A, venous-arterious; V02, oxygen uptake; w.w., wet weight.
were all fully informed about the nature of the study and any risks involved before they volunteered to participate. A seventh subject participated in the study, but only muscle biopsies were taken in this subject.

The subjects were asked not to do any training other than the knee extensor exercises. They were initially familiarized with the model by practicing with both legs separately about five times. To determine the maximal work and peak VO₂ for the knee extensors, a maximal test was performed with each leg separately. Total body VO₂ increased linearly with increasing work loads until an increase in work load caused a steeper increase in total body VO₂. The sudden steeper rise in whole body VO₂ is due to recruitment of extraneous muscles used to stabilize the body during knee extension. Total body VO₂ at this point minus total body VO₂ at rest was regarded as the peak VO₂ for the muscle studied.

Training period
Training consisted of knee extensor exercise with one leg during a total period of 8 wk. During the first 3 wk the subjects exercised 3 d/wk, increasing the duration of each session from 0.5 to 1.5 h during the 3-wk period. From week 4 they exercised 4 d/wk for 2 h at each session and this was maintained during the final 5 wk. The work load was increased progressively in relation to the changes in maximal performance assessed every 2 wk, being kept at 65% of peak VO₂ for the knee extensors. In four subjects the right leg was trained and in three others the left leg.

Experimental procedure
In alternate subjects the trained or the nontrained leg was tested first. Dietary intake was controlled during days 1 and 2 (D-1 and D-2; see below). The controlled diet provided an average 37±5 energy % of fat, 47±4 energy % of carbohydrates, and 16±3 energy % of proteins (mean±SE). The mean ratio of polysaturated to saturated fatty acids was 0.54±0.16 and the total caloric intake was 11.7±2.3 MJ per day. No alcohol was allowed.

D-1. No physical exercise was performed.

D-2. The subject arrived in the morning after an overnight fast of 10–12 h. After 20 min rest in the supine position blood was drawn from an arm vein while venous stasis was applied. A muscle biopsy was then taken from the lateral portion of one of the vastus femoris muscles 15 cm above the patella (9) for determination of LPLA and for histochemical analysis. The subject then performed knee extension exercise with the biopsied leg for 2 h at ~65% of the maximal work capacity of the nontrained leg, which had been determined one week earlier. Pulmonary VO₂ was measured during the work.

D-3. The subject arrived in the morning after an overnight fast of 10–12 h. Catheters were inserted into the femoral artery and vein of the test leg 2 and 4 cm distal to the inguinal ligament, respectively, and advanced centrally. A thermistor catheter (Eds laboratory TD-probe; American Edwards Laboratories) was inserted through the venous catheter and advanced 12–15 cm anterogradely for measurement of blood flow. The subject then rested for 45 min in the supine position. After another 15 min of rest in the sitting position blood samples were drawn from the femoral artery and vein simultaneously. Thereafter, a muscle biopsy was taken from the lateral vastus muscle for determination of LPLA and histochemical analyses.

The subjects then repeated the knee extension exercise for 2 h at the same work level as on D-2. Arterial and venous blood samples were drawn simultaneously, and venous blood flow and pulmonary VO₂ were measured during and at the end of the exercise period. The heart rate was monitored continuously. An inflatable cuff was placed just below the knee of the working leg and inflated to 280 mmHg during each period of measurement (2–2.5 min) to exclude admixture of blood from the lower leg. Another biopsy was taken from the lateral vastus muscle of the quadriceps femoris at the end of exercise for determination of LPLA.

4 d later the same protocol sequences were repeated on D-1, D-2, and D-3 with the other leg, with the same absolute work load.

Analytical methods
Blood analysis. Lipoproteins were separated by ultracentrifugation. A detailed description of the analyzing procedures for lipoproteins has been published previously (10). To minimize any transfer of lipids from VLDL and LDL to HDL during the preparative ultracentrifugation the following procedure was applied. An MgCl₂ phosphotungstate technique was used to precipitate the VLDL and LDL. The supernatant was ultracentrifuged (LKB-65; Beckman Instruments, Inc., Palo Alto, CA) at a density of 1.125 to separate HDL₂ and HDL₃.

VLDL in serum were isolated in the top fraction in a separate ultracentrifugation run at a density of 1.006. The VLDL fraction was analyzed for TG and C concentrations (see below) and also for free glycerol. To correct for contamination of free glycerol in the VLDL, the free serum glycerol was subtracted from the value for VLDL-TG. LDL were precipitated from the bottom fraction after the ultracentrifugation step and HDL were isolated in the supernatant after the precipitation.

TG and C concentrations were assayed in serum and in the different isolated lipoprotein subclasses using enzymatic methods (Boehringer Mannheim GmbH, Mannheim, FRG) in a Multistat III (Instrumentation Laboratory, Inc., Lexington, MA). The TG method is based on assay of glycerol after hydrolysis of the TG. The HDL subclasses HDL₃ and HDL₄ (see above) were assayed in quadruplicate and the mean values were used in the calculations. The analytical error (coefficient of variation) determined from 264 samples run in 19 series was 1.3 and 1.4% for TG and C analyses, respectively.

Insulin in serum was assayed by modified radioimmunoassay technique (Phadebas, Pharmacia Diagnostics AB, Uppsala, Sweden). Epinephrine and norepinephrine were measured by a single isotope-derivative assay (11).

All analyses of lipoproteins, insulin, epinephrine, and norepinephrine were done without knowing whether the samples were from the trained or nontrained leg.

Muscle analyses. A piece of the muscle biopsy specimen was immediately freed from blood and visible connective tissue, rapidly frozen in liquid nitrogen, and stored at −80°C for subsequent biochemical analyses. The other part of the specimen was mounted in embedding medium, frozen in isopentane, cooled to its freezing point in liquid nitrogen, and then stored at −80°C for subsequent histochemical analyses. LPLA was measured as described by Lithell and Boberg (12) with slight methodological modifications (13). The tissue was incubated in a hep- arin-containing medium with a serum-activated triolein emulsion with [³H]triolacte as trace substance (14). The release of [³H]oleic acid was used as a measure of LPLA. 1 nmol of the fatty acid released per min was equal to 1 mU of enzyme activity, and the activity is expressed per g wet weight (w.w.).

For histochemical analysis transverse sections (10 μm) were cut with a microtome at −20°. The amylase-periodic acid-Schiff method was used to visualize muscle capillaries (15).
Blood flow was measured by the thermodilution technique. At rest a bolus of ice-cold saline was injected, but during exercise constant infusion of ice-cold saline was used as described by Andersen and Saltin (16). Pulmonary \( VO_2 \) during rest and exercise was measured by collecting expired air in Douglas bags. A Tissot spirometer was used for volume analysis and the \( O_2 \) and \( CO_2 \) contents of the bags, collected during exercise, were determined with a paramagnetic (Taylor Servomex Limited, Crawborough, Sussex, UK) and infrared (LB-2; Beckman Instruments, Inc.) system, respectively. The \( O_2 \) and \( CO_2 \) contents of the bags filled at rest were measured by the Scholander microtechnique. Heart rate was recorded on an Elema minograph (Siemens Corp., Iselin, NY).

Statistics. Results are given in the following as mean values and ranges. For statistical analyses the paired Wilcoxon signed rank test, Friedman’s two-way analysis of variance, and linear regression analysis were used.

Results

Muscle biopsies (n = 7)

LPLA. In muscle biopsies taken on D-2 the mean resting LPLA was found to be 34.4 mU/g w.w. (16-78) in the nontrained muscle (Table I). In the trained muscle of six subjects LPLA was markedly higher compared with the nontrained muscle. In the seventh subject the LPLA was high in the nontrained muscle and was not higher in the trained muscle. Thus the mean of LPLA in the trained muscles was 49.3 mU/g w.w. (36-63). The mean difference between nontrained and trained muscles was 70% (\( P < 0.05 \)).

Capillary density. A difference in capillary density was observed between the nontrained and the trained muscle. Thus, the number of capillaries surrounding each fiber averaged 1.82 (1.38-2.22) in the nontrained muscle but 2.25 (1.87-2.53) in the trained muscle (\( P < 0.05 \)). There were 343 (276-401) capillaries/mm\(^2\) in nontrained and 417 (350-495)/mm\(^2\) in trained muscle (\( P < 0.05 \)). Comparing m-LPLA and the capillary density, increased m-LPLA was correlated with an increased capillary density (\( r = 0.85 \); Fig. 2).

Peak \( VO_2 \). The \( VO_2 \) of the knee extensors (peak \( VO_2 \)) was determined by subtracting resting total body \( VO_2 \) from exercising total body \( VO_2 \). Peak \( VO_2 \) in the nontrained muscle averaged 0.870 (0.780-0.950) liters/min, whereas in the trained muscle it averaged 1.06 (0.920-1.14 liters/min; \( P < 0.05 \)).

Resting study

The arterial VLDL-TG concentration in the nontrained muscle was 361 \( \mu \)mol/liter (236-584) and 360 \( \mu \)mol/liter (210-650) in the trained muscle. The A-V VLDL-TG difference over the nontrained muscle amounted to 30 \( \mu \)mol/liter (4-72), but was markedly higher over the trained muscle, averaging 55 \( \mu \)mol/liter (30-123) (Table II).

The arterial concentration of HDL-C averaged 1.00 mmol/liter (0.83-1.20) and 1.04 mmol/liter (0.74-1.34) in the nontrained and trained muscle, respectively. Mean femoral vein concentration of HDL-C was 1.01 mmol/liter (0.83-1.20) in the nontrained and 1.10 mmol/liter (0.80-1.36) in the trained muscle. Thus a nonconsistent venous-arterious (V-A) HDL-C difference was found over the nontrained muscle (10 \( \mu \)mol/liter) (−30+50), whereas a consistent positive V-A difference of HDL-C was obtained over the trained muscle averaging 53 \( \mu \)mol/liter (20-110) (\( P < 0.05 \), Table II). The arterial concentration of HDL\(_2\)-C in the nontrained muscle averaged 324 \( \mu \)mol/liter (231-440) and no consistent A-V HDL\(_2\) differences were noted. In the trained muscle the arterial HDL\(_2\)-C concentration was 362 \( \mu \)mol/liter (230-630) and was not different from the arterial concentration in the nontrained muscle. However, in the trained muscle the concentration of HDL\(_2\)-C was markedly higher in the femoral vein compared with the artery, averaging 400 \( \mu \)mol/liter (291-500). Thus, over the trained muscle the V-A difference in HDL\(_2\)-C was 38 \( \mu \)mol/liter (10-73), which was significantly higher than that over the nontrained muscle (\( P < 0.05 \)) (Table II, Fig. 3).

The arterial LDL-C concentration was 2.79 mmol/liter (1.67-3.33) and 2.81 mmol/liter (1.66-3.48) in the nontrained and trained muscle, respectively. The V-A difference in this variable was 113 \( \mu \)mol/liter (90-160) over the nontrained muscle, but lower over the trained muscle, averaging 92 \( \mu \)mol/liter (20-150) (Table II).

The mean resting femoral venous blood flow was 0.620 liters/min in both legs. The uptake of VLDL-TG calculated from arterial minus venous VLDL-TG concentration times plasma flow averaged 8 \( \mu \)mol/min in the resting nontrained muscle and 19 \( \mu \)mol/min in the resting trained muscle. The mean formation of HDL\(_3\)-C in plasma, calculated as arterial minus venous HDL\(_3\)-C content times plasma flow was 15 \( \mu \)mol/min in the trained muscle at rest, but was not significantly different from zero in the nontrained muscle.

At rest no significant correlations were found over the nontrained leg between m-LPL activity and A-V differences of VLDL-TG (\( r = -0.31 \)), between m-LPL activity and A-V differences of HDL-C and HDL\(_2\)-C (\( r = 0.13 \) and \( r = -0.13 \), respectively), or between A-V differences of VLDL-TG and HDL-C (\( r = 0.13 \)). However, over the trained leg a positive correlation was found between m-LPL activity and A-V differences of VLDL-TG (\( r = 0.71 \)), between m-LPL activity and

Table I. m-LPLA, Mean±SE of Seven Subjects

<table>
<thead>
<tr>
<th></th>
<th>Nontrained</th>
<th>Trained</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mU/g w.w.</td>
<td>mU/g w.w.</td>
</tr>
<tr>
<td>D-2</td>
<td>34.4±8.2</td>
<td>49.3±3.6*</td>
</tr>
<tr>
<td>D-3</td>
<td>36.7±7.4</td>
<td>46.6±6.0</td>
</tr>
<tr>
<td>Before</td>
<td>36.7±8.1</td>
<td>49.4±6.3</td>
</tr>
<tr>
<td>After</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Differences between nontrained and trained muscle (\( P < 0.05 \)).

The values were obtained in the morning at rest on D-2 and again in the morning on D-3 at rest and after 2 h of exercise in nontrained and trained muscle (vastus lateralis muscle).
Table II. Resting Values for Concentrations of Lipoproteins in Femoral Arterial and Venous Serum in the Nontrained and Trained Muscle, mean±SE of Six Subjects

<table>
<thead>
<tr>
<th></th>
<th>Nontrained muscle</th>
<th>Trained muscle</th>
<th>(A-V)</th>
<th>(A-V)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
<td>V</td>
<td>A</td>
<td>V</td>
</tr>
<tr>
<td>HDL-C (mmol/liter)</td>
<td>1.00±0.08</td>
<td>1.01±0.05</td>
<td>-0.01±0.01</td>
<td>1.04±0.07</td>
</tr>
<tr>
<td>HDL2-C (μmol/liter)</td>
<td>324±28</td>
<td>322±26</td>
<td>2±7</td>
<td>362±57</td>
</tr>
<tr>
<td>LDL-C (mmol/liter)</td>
<td>2.79±0.24</td>
<td>2.91±0.24</td>
<td>-0.12±0.01</td>
<td>2.81±0.26</td>
</tr>
<tr>
<td>VLDL-C (μmol/liter)</td>
<td>214±40</td>
<td>247±60</td>
<td>-34±12</td>
<td>186±39</td>
</tr>
<tr>
<td>VLDL-TG (μmol/liter)</td>
<td>361±66</td>
<td>330±72</td>
<td>30±12</td>
<td>360±50</td>
</tr>
</tbody>
</table>

* Differences between nontrained and trained (P < 0.05).

A-V differences of HDL-C (r = 0.80, P < 0.05) and HDL2-C (r = 0.83, P < 0.05), and between A-V differences of VLDL-TG and HDL-C (r = 0.90, P < 0.05).

Exercise study

Total body VO2 averaged 795 ml/min (623–1,075) 10 min after the start of exercise with the nontrained muscle and increased toward the end of the 2-h work period to 890 ml/min (648–1,212) (P < 0.05). During exercise with the trained muscle total body VO2 was 807 ml/min (583–961) at 10 min and remained at that level for the rest of the exercise period.

The heart rate averaged 110 beats/min (88–141) after 10 min exercise, with both the nontrained and the trained muscle. During the remaining period of exercise with the nontrained muscle it increased continuously, averaging 128 beats/min (100–160) at 110 min, whereas during exercise with the trained muscle it remained constant.

The femoral venous blood flow remained constant during exercise. In the exercising nontrained leg the blood flow was 5.51 liters/min, but in the exercising trained leg it was lower, averaging 4.65 liters/min (P < 0.05).

In the nontrained leg the arterial concentration of VLDL-TG was 417 μmol/liter (220–1,168) after 60 min of exercise and increased to 495 μmol/liter (266–1,116) at 110 min. In the exercising trained leg the arterial concentration of VLDL-TG increased to 392 μmol/liter (180–655) at 60 min and to 439 μmol/liter (210–761) at 110 min (P < 0.05). Small but inconsistent A-V differences of VLDL-TG were found over the exercising nontrained and trained muscle.

During exercise the arterial concentration of HDL-C increased nonsignificantly from resting values in both the nontrained (1.06 mmol/liter [0.89–1.19] at 60 min and 1.05 mmol/liter [0.89–1.19] at 110 min) and trained thigh (1.08 mmol/liter [0.83–1.28] at 60 min and 1.07 mmol/liter [0.83–1.25] at 110 min).

Positive V-A HDL-C concentration differences were observed over both the exercising nontrained (12 μmol/liter [−60–40] at 60 min and 18 μmol/liter [−40–70] at 110 min) and trained thigh (10 μmol/liter [0–30] at 60 min and 30 μmol/liter [10–80] at 110 min) except in one subject in the nontrained situation, where negative V-A HDL-C concentrations were found. Difference in production between the nontrained and trained thigh were not significant.

At rest the arterial concentration of HDL2-C was similar on the day when the nontrained muscle was studied (324 μmol/liter) and on the day the trained muscle was studied (362 μmol/liter). A small, nonsignificant increase was observed during the exercise period in both situations.

No consistent V-A differences in HDL2-C were observed over the exercising nontrained muscle, whereas in the exercising trained muscle there was a mean V-A difference at 60 min of 30 μmol/liter (0–60) and 34 μmol/liter at 110 min (20–80) (Fig. 3).

Before exercise the LPLA was 36.7 mU/g w.w. in the nontrained muscle and remained unchanged at the end of the exercise period (36.7 mU/g w.w.). In the trained muscle the preexercise LPLA value averaged 49.3 mU/g w.w. and no change was observed at the end of exercise (49.6 mU/g w.w.) (Table I).

Hormones. The mean arterial insulin concentrations at rest were similar on the days of experiments with the nontrained (7.8 mU/liter) and the trained muscle (7.5 mU/liter) (Table III). During exercise the arterial insulin level decreased continuously to 3.7 mU/liter in the nontrained and 4.4 mU/liter in the trained muscle at 110 min (Table III).

The arterial concentrations of epinephrine were similar on the days of experiments with the nontrained and trained muscle at rest (0.12 and 0.11 ng/ml, respectively; Table III). Changes from resting values were not observed during exercise with either muscle category (Table III). The resting arterial norepinephrine concentrations were also similar on the days of experiment with the nontrained (0.38 ng/ml) and the trained muscle (0.33 ng/ml) (Table III). 10 min after exercise with the nontrained muscle, arterial norepinephrine had risen to 0.46 ng/ml (P < 0.05) and an insignificant increase followed during
the remaining exercise period. During exercise with the trained muscle an increase to 0.37 ng/ml was found at 10 min (P < 0.05) and a further increase to 0.45 ng/ml at 60 min (P < 0.05), and the value then remained at that level (Table III).

**Table III. Mean Values±SE of Norepinephrine (NE), Epinephrine (E), and Insulin in Femoral Artery Blood during 2 h of Exercise with the Trained (T) and Nontrained (NT) Thigh (n = 6)**

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>Arterial NE (ng/ml)</th>
<th>Arterial E (ng/ml)</th>
<th>Arterial insulin (μU/liter)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rest</td>
<td>T: 0.33 NT: 0.38</td>
<td>T: 0.11 NT: 0.12</td>
<td>T: 7.8 NT: 7.5</td>
</tr>
<tr>
<td></td>
<td>(0.09) (0.02)</td>
<td>(0.05) (0.03)</td>
<td>(0.9) (1.2)</td>
</tr>
<tr>
<td>10</td>
<td>0.37* (0.04)</td>
<td>0.12 (0.03)</td>
<td></td>
</tr>
<tr>
<td>60</td>
<td>0.45* (0.05)</td>
<td>0.13 (0.03)</td>
<td>5.5* (0.7)</td>
</tr>
<tr>
<td>110</td>
<td>0.47* (0.05)</td>
<td>0.14 (0.02)</td>
<td>4.4* (0.8)</td>
</tr>
</tbody>
</table>

* Difference compared with resting levels (P < 0.05).

Discussion

In the present study we have investigated the influence of 8 wk of dynamic exercise training of one quadriceps femoris muscle on muscle lipoprotein metabolism at rest and during exercise, using the nontrained contralateral muscle as control. The major findings were a training-induced increase in m-LPLA and a significant formation of HDL₂-C in trained muscle but not in nontrained muscle.

An increase in the activity of skeletal m-LPLA by ∼ 47% was previously observed after a period of physical training in nonobese, healthy men (17). In that study, however, the state of nutrition, which may significantly influence LPLA (18–20), was not controlled. In the present dietary-controlled study the m-LPLA was 70% higher in the trained muscle compared with the nontrained muscle of the same individual, which may at least partly be the result of the increase in capillary density in the trained muscle. The physiological site of LPLA is the luminal surface of the capillary endothelium, where the enzyme is bound to the endothelial cell surface. In the trained muscle the increased number of capillaries around each fiber type compared with the nontrained muscle provides more binding sites for the enzyme. Moreover, LPLA correlated with the capillary density (Fig. 2), which further supports the idea of a causal relationship between capillary density and LPLA. In the present study both were determined in tissue from the same biopsy sample. This was not the case in an earlier study, which probably explains why such a relationship was not found then (17).

It has been suggested that the breakdown of VLDL may contribute to the formation of HDL₂ in the circulation. The primary HDL particle (nascent HDL) is produced in the intestine and liver, but immediately after its delivery to the circulation it is transformed to HDL₃. Upon assimilation of phospholipids, C, and apoproteins from degraded VLDL, the HDL₃ is transformed to particles designated HDL₂ (6). Since the degradation of TG-rich particles is largely dependent upon the LPLA in peripheral tissues (21, 22), an increase in tissue LPLA is expected to result in transference of an increased amount of surface material to HDL in the plasma compartment. The question arises whether the formation of HDL₂-C in the trained leg can be ascribed to transference of surface material from VLDL-TG as a result of LPLA. Such a contention is in agreement with the finding of significantly larger A-V differences for VLDL-TG at rest in trained muscle compared with nontrained, while significant formation of HDL-C and HDL₂-C was only observed in trained muscle (Table II). It also agrees with the finding of a training-induced increase in m-LPLA (Table I). Still, it might seem strange that no consistent formation of HDL-C and HDL₂-C was observed in the nontrained leg in spite of significant degradation of VLDL-TG (Table II). However, it is interesting to note that at rest a correlation between VLDL-TG degradation and m-LPLA as well as between LPLA and HDL-C and HDL₂-C formation was obtained only in trained muscle. Furthermore, a significant correlation between degradation of VLDL-TG and formation of HDL-C was only found in trained and not in nontrained muscle at rest. Since arterial concentrations of insulin, catecholamines, and VLDL-TG as well as blood flow were similar when the trained and nontrained muscles were studied at rest, the above-mentioned correlations suggest that in addition to the training-induced increase in m-LPLA as measured in vitro, training may also have elicited qualitative changes in LPLA. Such qualitative changes could explain the apparent difference in expression of LPLA in vivo in the trained and nontrained muscle. Obviously, the above described differences in vivo expression of LPLA could also be ascribed to a diminished inhibition of LPLA in the trained compared with the nontrained muscle. Support for such a contention may be found in a study by Olivecrona and Bengtsson (23), who suggest that when lipoproteins encounter LPL at the endothelium, lipid hydrolysis is initially very rapid and the lipolytic products are rapidly and efficiently used by the tissue cells, probably after removal from the site of hydrolysis by lateral diffusion in the cell membrane (24). However, when the capacity for product use in the tissue is exceeded, hydrolysis slows down primarily as a result of product inhibition of LPLA (22). The capacity for using fatty acids as an energy source obviously was improved in trained muscle, as the activity of the enzyme β-OH-acyl CoA-dehydrogenase, which is needed for β-oxidation of fatty acids was 40% higher in trained than in nontrained muscle (25). Moreover, the diffusion distances decreased in trained muscle as judged by the increase in capillary density. Thus, in trained muscle the existence of a greater potential for removal and use of fatty acids from the site of TG hydrolysis makes it likely that in vivo LPLA is less inhibited.
than in nontrained muscle, in turn accounting for the observed increased femoral venous concentration of HDL-C (and HDL-C) in trained compared with nontrained muscle.

During exercise a significant formation of HDL-C and HDL-C was only detectable in the trained leg, but significant A-V differences for VLDL-TG were measurable neither in trained nor in nontrained muscle. Probably due to the much higher blood flow during exercise than rest, which in turn decreases capillary transit time drastically, VLDL-TG differences were not consistently detectable. However, in an effort to obtain a rough estimate of total leg degradation of VLDL-TG during exercise, the area under the arterial and venous concentration curves was integrated from 0 to 110 min of exercise. The difference between these areas multiplied by the plasma flow thus represents an estimate of the total leg degradation of VLDL-TG. Similar calculations were performed for total HDL-C formation. The estimates for VLDL-TG degradation and for formation of total HDL-C using values from both trained and nontrained legs correlated closely ($r = 0.91$; Fig. 4). This correlation strengthens the probability that HDL-C formation was due to hydrolysis of VLDL-TG during exercise as well.

During exercise diminished product inhibition of LPL could be a factor explaining why HDL-C formation was found only across the trained muscle. Apart from the training-induced increased capacity for fatty acid oxidation and the possibility that increased capillarization improved diffusion conditions, the training-induced increased capillary density has other possible effects. Thus, the larger capillary density in conjunction with the lower blood flow in the trained muscle during exercise implies that the mean capillary transit time was longer in trained than in nontrained muscle. This may be of importance for the transport of gas and substrates within the muscle, but it also appears likely that a longer mean transit time may be crucial for the contact between lipoprotein particles and LPL, and thus for the formation of HDL. Such factors may help to explain why A-V differences in HDL-C and HDL-C were found only in the trained muscle during exercise.

Previous studies have shown that a single bout of whole body exercise results in increased m-LPLA and the response seems to be related more to the duration than to the intensity of exercise (26, 27). It may also be related to the status of intramuscular substrate levels (28, 29). In a previous study it was found that during days of heavy exercise afternoon values of LPLA were significantly higher than morning values (30), and this exercise-induced increase appeared to persist for at least 12 h. In the present study no changes in LPLA were found on the morning after (18 h) 2 h of submaximal exercise was performed (Table I), either in trained or in nontrained muscle. Moreover, m-LPLA was not altered immediately after 2 h of one-legged knee extensions (Table I). In a previous study m-LPLA was found to be positively correlated to the excretion of epinephrine in the urine after heavy work (30). However, we have recently shown that one-legged knee extensions cause no increase in m-LPLA during exercise, but that an increase in LPLA is found 4 h after exercise in the exercised leg but not in the contralateral control leg (31). Furthermore, the increase was transient as it was no longer detectable 8 h after exercise. Thus, muscle contractions per se cause a delayed, local, and transient increase in m-LPLA.

The relationship between physical training and HDL-C is clear from several longitudinal studies of both sedentary and well trained subjects (e.g., 1–3), and a relationship between HDL-C concentration and maximal $V_O_2$ has been documented. However, it was recently reported that an increase in HDL-C occurred during increased training in already well trained men without a further increase in maximal $V_O_2$ (3). This indicates that local factors play a major role in lipoprotein metabolism. In the present study training did not have any effects on whole body maximal $V_O_2$, although peak $V_O_2$ increased in the trained muscle. Nevertheless, 8 wk of intensive dynamic exercise training with one muscle group, weighing on the average 2.5 kg, resulted in a significantly higher (8%) HDL-C. This increase is small in absolute terms but is of the same magnitude as reported after moderately intense training of sedentary men (1) and larger than the difference between subjects who have suffered myocardial infarction and those who have not (32). Our findings of changes in systemic HDL-C offer further support for the view that changes in lipoproteins induced by endurance training are caused to a large extent by local adaptations in skeletal muscle.

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