Contribution of Intrinsic Skeletal Muscle Changes to $^{31}\text{P}$ NMR Skeletal Muscle Metabolic Abnormalities in Patients With Chronic Heart Failure

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Patients with heart failure frequently exhibit abnormal skeletal muscle metabolic responses to exercise, as assessed with $^{31}\text{P}$ NMR. To investigate whether these metabolic abnormalities are due to intrinsic skeletal muscle changes, we performed gastrocnemius muscle biopsies on 22 patients with heart failure (peak $\text{VO}_2$, $15.4\pm4.7 \text{ ml/kg/min}$; ejection fraction, $20\pm7\%$) and on eight normal subjects. Biopsies were analyzed for fiber type and area, capillarity, citrate synthase, phosphofructokinase, lactate dehydrogenase, and $\beta$-hydroxyacyl CoA dehydrogenase activity. All patients with heart failure also underwent $^{31}\text{P}$ NMR studies of their calf muscle during plantarflexion at three workloads. Muscle pH responses and the relation of the ratio of inorganic phosphate to phosphocreatine ($\text{Pi}/\text{PCr}$) to systemic $\text{VO}_2$ were then evaluated. Compared with normal subjects, patients with heart failure exhibited a shift in fiber distribution with increased percentage of the fast twitch, glycolytic, easily fatigable type IIB fibers (normal subjects, $22.7\pm10.1\%$; heart failure, $33.1\pm11.1\%; p<0.05$), atrophy of type IIA (normal subjects, $5,477\pm1,109\; \text{heart failure, 4,239}\pm1,247\; \mu\text{m}^2$; $p<0.05$) and type IIB fibers (normal subjects, $5,957\pm1,388\; \text{heart failure, 4,144}\pm945\; \mu\text{m}^2$; $p<0.01$), and decreased activity of $\beta$-hydroxyacyl CoA dehydrogenase (normal subjects, $5.17\pm1.44\; \text{heart failure, 3.67}\pm1.68 \text{ mol/kg protein/hr}$; $p<0.05$). No significant linear correlation could be identified between the slope of the $\text{Pi}/\text{PCr}$ to $\text{VO}_2$ relation and muscle histochemistry or enzyme activities. Similarly, no linear relation was found between intracellular pH at peak exercise and any muscle variable. These data suggest that patients with heart failure develop intrinsic skeletal muscle changes but that these intrinsic muscle changes do not contribute significantly to the abnormal skeletal muscle $^{31}\text{P}$ NMR metabolic responses observed in such patients. (Circulation 1989;80:1338–1346)

Exertional fatigue in patients with heart failure has traditionally been attributed to skeletal muscle underperfusion. However, recent investigations suggest that intrinsic skeletal muscle abnormalities may be operative, as well. Abnormal forearm $^{31}\text{P}$ NMR responses to forearm exercise in patients with heart failure have been demonstrated without substantial decreases in limb blood flow.1,2 Massie et al3 have shown that patients exhibit abnormal forearm muscle responses during ischemic exercise. Lipkin et al4 performed quadriceps muscle biopsies on nine patients with heart failure. A variety of different abnormalities were noted in individual patients, including atrophy of type I and II fibers, increased interstitial cellularity, increased lipid deposition, and increased acid phosphatase staining. However, no characteristic histochemical pattern or enzyme abnormality emerged. In preliminary reports, Drexler et al5,6 described reduced mitochondrial cristae volume in patients with heart failure, suggesting a decrease in oxidative enzymes.

In the present study, we sought to further investigate whether intrinsic skeletal muscle changes occur in patients with heart failure. In addition, we
sought to examine the relation of skeletal muscle biopsy results to $^{31}$P NMR responses to exercise.

**Methods**

**Subjects**

Twenty-two men with chronic New York Heart Association class I–IV heart failure were studied. Average age was $57\pm14$ years ($\pm$SD). All patients were receiving treatment with digoxin and diuretics. In addition, 12 patients were also receiving vasodilators. None had peripheral edema, intermittent claudication, diabetes mellitus, angina, or valvular heart disease. All patients were screened for peripheral vascular disease by observation of physical signs (e.g., hair loss, dependent rubor), palpation of peripheral pulses, and determination of rest ankle and brachial pressures. All participants had preserved peripheral pulses and an ankle to brachial pressure ratio of 1 or more.

Etiology of heart failure was coronary artery disease in eight patients based on a documented myocardial infarction, cardiac catheterization, or both. The remaining 14 patients were presumed to have a cardiomyopathy, seven due to excessive alcohol intake. There was no evidence of myocardial infarction on the electrocardiogram, and no history of myocardial infarction in these 14 patients. In addition, four patients had normal coronary arteries documented by coronary angiography, and three had a normal thallium exercise test. Left ventricular ejection fraction averaged $20\pm7\%$. Peak exercise oxygen consumption during treadmill exercise averaged $15.4\pm4.7$ ml/min/kg.

Eight sedentary normal male subjects also underwent a calf skeletal muscle biopsy. The age of the subjects averaged $60\pm9$ years. Peak exercise oxygen consumption averaged $28.3\pm1.4$ ml/kg/min.

The protocol was approved by the Committee on Studies Involving Human Beings at the University of Pennsylvania. Written informed consent was obtained from all subjects.

**Protocol**

All patients and control subjects underwent skeletal muscle biopsy. $^{31}$P NMR measurements were performed within 1 week of biopsy in all patients with heart failure.

**Skeletal Muscle Biopsy**

Percutaneous muscle biopsies were obtained from the gastrocnemius muscle using the biopsy technique described by Bergstrom. Gross blood and connective tissue were removed from the samples. One biopsy (approximately 40 mg) was immediately frozen in liquid nitrogen and stored at $-80^\circ$ C, pending enzymatic analysis. Another sample (approximately 20 mg) was mounted in embedding medium (OCT) with fibers aligned in longitudinal direction. This specimen was then frozen in isopentane, cooled to its freezing point with liquid nitro-
within a section, and only those fibers that did not appear cross sectional or intact were excluded. In 10 biopsy samples, the mean coefficient of variation for four determinations of fiber area was ±3.6%.

**Skeletal Muscle Metabolic Measurements (NMR Protocol)**

Exercise was performed at least 3 hours after eating. The protocol has been described previously. In brief, after arrival in the NMR facility, the patient was seated and his leg was positioned with the limb under study in an 11.5-in. bore, 1.9-T superconducting magnet (Oxford Research Systems, Oxford, England). Because the homogeneous area of the magnetic field was 37 cm from each end, the leg was inserted up to the groin and the other leg widely abducted. Data acquisition was accomplished with the application of radiofrequency pulses (pulse width optimized at 25–35 μsec) applied every 5 seconds. After optimization of field homogeneity, a 3-minute rest scan was recorded. The subject then withdrew his leg from the magnet, as all exercise was performed outside the magnet. A rapidly inflating pneumatic cuff (D.E. Hokanson, Issaquah, Washington) was applied to the study leg. The patient then exercised next to the magnet, as herein described. At the completion of each workload, the pneumatic cuff was inflated to a suprasystolic pressure (250 mm Hg) to occlude blood flow into and out of the leg and, thereby, maintain pH, inorganic phosphate (Pi), and phosphocreatine (PCr) levels at end-exercise levels. After the cuff was inflated, the patient rapidly (<1 minute) repositioned his leg within the magnet. After acquisition of a 3-minute 31P NMR scan, the cuff was deflated. Recovery scans were recorded every minute for 5 minutes. At the end of the study, calf circumference at the site of scanning was measured.

Exercise consisted of two-footed upright plantarflexion (toe ups). Three different workloads were performed in each subject: plantarflexion every 3, 6, and 9 seconds. Contractions were sustained for approximately 1 second. Each workload was performed for 4 minutes with continuous monitoring of heart rate and measurement of cuff blood pressure at the end of each workload. Simultaneous oxygen consumption measurements were performed to provide quantification of each level of exercise. Measurements of mixed expired oxygen, mixed expired carbon dioxide, and expired volume were determined at rest and every 30 seconds throughout exercise using a metabolic measurement cart (SensorMedics, Anaheim, California). After each exercise workload, 31P NMR data was obtained, as previously described. The subject then rested for 10 minutes before performing the next workload.

Previously, we have demonstrated that metabolism can be temporarily "arrested" by inflating a limb cuff to suprasystolic pressures at end exercise. Forearm exercise was performed in the magnet by seven normal subjects followed by cuff inflation to suprasystolic pressure. Pi/PCr ratio and intracellular pH were monitored for 4 minutes after exercise and remained stable.

Harris et al also investigated the effect of pneumatic cuff inflation on PCr resynthesis in the quadriceps muscle of humans after exhaustive exercise. By performing serial muscle biopsies, these investigators demonstrated that arterial occlusion abolished resynthesis of PCr throughout 6 minutes of occlusion. Release of arterial occlusion was followed by restoration of PCr to normal levels.

**Spectral Analysis**

Quantification of metabolic components was obtained from the Fourier transformed NMR spectra. An exponential multiplication equivalent to a line broadening of 15 Hz was used, yielding a width at half height for PCr of less than 1 ppm. Triangulation was used to assess Pi and PCr signal area. Intracellular pH was calculated from the chemical shift difference of Pi from PCr.

**Statistical Analysis**

Data were compared using unpaired Student’s t testing or analysis of variance, as appropriate. The relation between variables was examined by linear regression analysis. A p value less than 0.05 was considered significant. All data are expressed as mean±SD.

**Results**

**Skeletal Muscle Histology**

When compared with control subjects, patients with heart failure exhibited an increased percentage of type IIB fibers (normal subjects, 22.7±10.1; heart failure, 33.1±11.1%; p<0.05) (Table 1 and Figure 1). No significant differences were observed between the two groups in the percentage of type IIA fibers. The percentage of type I fibers tended to be lower in patients with heart failure; however, this difference did not achieve statistical significance. Eleven patients with heart failure exhibited type IIC fibers, averaging 1.8±3.1% (range, 0–11.7%), compared with none of the control subjects.

In patients with heart failure, a reduction in type IIA fiber area (normal subjects, 5,477±1,109; heart failure, 4,239±1,247 μm²; p<0.05) and IIB fiber area (normal subjects, 5,957±1,388; heart failure, 4,144±945; p<0.01) was observed. No significant difference was noted in the area of type I fibers.

To determine the relative contribution of each fiber type to a given cross-sectional area of muscle, the total area for 100 fibers was calculated by multiplying the percentage of each muscle fiber type by its respective area and then adding the three products. The area occupied by each fiber type was then divided by this total area. With this approach, no significant difference was observed between normal and heart failure populations. In the normal subjects, the relative percentage contributions to
TABLE 1. Comparison of Skeletal Muscle Biopsy Results in Normal Control Subjects and in Patients With Heart Failure

<table>
<thead>
<tr>
<th>Fiber distribution (%)</th>
<th>Normal</th>
<th>Heart failure</th>
</tr>
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<tbody>
<tr>
<td>Type I</td>
<td>51.2±16.3</td>
<td>44.6±16.3</td>
</tr>
<tr>
<td>Type IIa</td>
<td>26.1±12.7</td>
<td>20.7±10.3</td>
</tr>
<tr>
<td>Type IIb</td>
<td>22.7±10.1</td>
<td>33.1±11.1*</td>
</tr>
<tr>
<td>Type IIc</td>
<td>0</td>
<td>1.8±3.1†</td>
</tr>
<tr>
<td>Fiber area (μm²)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Type I</td>
<td>5,369±951</td>
<td>5,067±1,154</td>
</tr>
<tr>
<td>Type IIa</td>
<td>5,477±1,109</td>
<td>4,239±1,247*</td>
</tr>
<tr>
<td>Type IIb</td>
<td>5,957±1,388</td>
<td>4,144±945†</td>
</tr>
</tbody>
</table>

| Capillarity            |        |              |
| Capillarity/fiber      | 1.67±0.25 | 1.85±0.35    |
| Capillarity/mm²        | 260±46  | 435±66†      |

| Enzymatic activity     | (mol/kg protein/hr) |                  |
| CS                     | 4.65±1.49          | 4.49±1.79        |
| BOAC                   | 5.17±1.44          | 3.67±1.68*       |
| LDH                    | 19.4±9.8           | 15.9±9.8         |
| PFK                    | 10.2±2.7           | 13.2±6.5         |
| Protein (mg/g)         | 216±42             | 198±46           |

CS, citrate synthase; BOAC, β-hydroxyacyl CoA dehydrogenase; LDH, lactate dehydrogenase; PFK, phosphofructokinase.

*p<0.05, †p<0.01, normal subjects versus patients with heart failure.

Muscle area for type I, IIa, and IIb fibers were 50±15%, 26±12%, and 25±11% versus, in the patients with heart failure, 50±16%, 19±9%, and 31±10% (p=NS for all).

Microscopic sections stained for myofibrillar ATPase at pH of 4.6 in a normal subject and two representative patients with heart failure are shown in Figure 2. The black fibers are type I; the grey fibers, type IIb; and light fibers, IIa. The shift toward an increased percentage of type IIb fibers is demonstrated in both patients with heart failure. Type II fibers are generally larger than or equal in size to type I fibers. Thus, the marked type II fiber atrophy is readily apparent in the second patient with heart failure.

Capillarity data revealed no difference between the two groups in the number of capillaries surrounding each fiber. However, an increase in capillaries/mm² was noted in patients with heart failure (normal subjects, 260±46; heart failure, 435±66 capillaries/mm²; p<0.001).

Skeletal Muscle Enzyme Activity

The protein (normal subjects, 216±42; heart failure, 198±46 mg/g; p=NS) and myoglobin content (normal subjects, 4.04±0.89; heart failure, 4.41±1.54 mg/g; p=NS) of skeletal muscle were not significantly different between the normal subjects and patients with heart failure (Table 1 and Figure 1). Statistical results were comparable when enzyme activity was expressed as moles per kilograms protein per hour as compared with moles per kilogram myoglobin per hour. Therefore, all enzyme activity is so reported.

When compared with the normal subjects, patients with heart failure demonstrated reduced activity of the mitochondrial enzyme BOAC (normal subjects, 5.17±1.44; heart failure, 3.67±1.68 mol/kg protein/hr; p<0.05), suggesting a reduced ability for oxidation of fats. The enzymatic activity of the mitochondrial enzyme CS, and glycolytic enzymes PFK and LDH were not significantly different in the two groups (Figure 1).

Relation Between Skeletal Muscle Biopsy Results, Etiology of Heart Failure, and Exercise Capacity

Patients were divided into groups based on the etiology of heart failure. Eight patients had coronary artery disease, seven had idiopathic dilated
cardiomyopathy, and seven had alcoholic cardiomyopathy. These populations were well matched with respect to age, ejection fraction, and exercise intolerance (Table 2). When these groups were compared, no difference was observed among groups in regard to fiber type distribution, fiber area, and enzyme concentration (Figure 3).

Peak exercise \( VO_2 \) range was 9.4–25.8 ml/kg/min in the patients with heart failure. A significant inverse relation was noted between the percentage of type IIb fibers and the peak exercise \( VO_2 \) (Figure 4). In contrast, a significant positive correlation was observed between percentage of type I fibers and peak \( VO_2 \) (Figure 4). No significant correlations were observed between peak \( VO_2 \) and enzyme activities or fiber area.

Only four patients exhibited skeletal muscle biopsies that were normal in regard to every parameter measured. These four subjects were young (age, 36±6.0 years) and were New York Heart Association class I or II (peak \( VO_2 \), 20.7±5.4 ml/kg/min).

Table 2. Characteristics of Normal Subjects and Patients With Heart Failure

<table>
<thead>
<tr>
<th></th>
<th>( n )</th>
<th>Age (yr)</th>
<th>EF (%)</th>
<th>( VO_2 ) (ml/kg/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>8</td>
<td>60±9</td>
<td>. . .</td>
<td>28.3±1.4</td>
</tr>
<tr>
<td>CAD</td>
<td>8</td>
<td>59±14</td>
<td>17±5</td>
<td>14.5±4.2*</td>
</tr>
<tr>
<td>IDC</td>
<td>7</td>
<td>56±17</td>
<td>22±10</td>
<td>17.1±6.0*</td>
</tr>
<tr>
<td>ETOH</td>
<td>7</td>
<td>57±13</td>
<td>19±5</td>
<td>15.0±4.1*</td>
</tr>
</tbody>
</table>

CAD, coronary artery disease; IDC, idiopathic cardiomyopathy; ETOH, alcoholic cardiomyopathy.

Relation Between Skeletal Muscle Biopsy Results and \( ^{31}P \) NMR Responses to Exercise

To investigate the contribution of skeletal muscle intrinsic changes to muscle metabolic responses to exercise, all patients underwent \( ^{31}P \) NMR studies to determine their calf metabolic responses to exercise. Upright plantar flexion resulted in a progressive rise in the ratio of P_i to PCr and decline in intracellular pH (Table 3).

To assess patients' metabolic response to exercise, \( VO_2 \) was correlated with the P_i/PCr ratio, as described previously. The P_i/PCr ratio provides an estimate of ADP concentration. ADP level is closely linked to mitochondrial respiration. Thus, the P_i/PCr ratio provides an index of oxidative metabolism. As described in the transfer function of Chance et al., the relation during low-level exercise between power output (oxygen consumption) and the P_i/PCr ratio is linear. Therefore, calculation of the slope of this relation affords a simple way of comparing oxidative metabolism between subjects. The average slope of the work–P_i/PCr relation of all patients was 0.0064±0.0046, significantly more than the mean slope previously noted by us in normal subjects (normal subjects, 0.0014±0.0005 min/ml; \( p<0.05 \)).

Linear regression analysis was performed using the work–P_i/PCr relation versus each biopsy variable. No statistically significant relation was demonstrated. Similarly, linear regression analysis using the pH noted at peak exercise versus each biopsy...
variable was performed. Again, no statistically significant relation was noted.

Discussion

Patients with heart failure are frequently limited by muscular fatigue. Recent $^{31}$P NMR studies suggest that this fatigue may be due, in part, to intrinsic skeletal muscle changes.1–3 In the present study, we sought to establish whether patients with heart failure exhibit such intrinsic skeletal muscle changes and to determine if these changes might contribute to $^{31}$P NMR metabolic abnormalities.

Skeletal Muscle Biopsies

Results of this study demonstrate that patients with heart failure exhibit several skeletal muscle abnormalities. These include a shift in fiber type distribution with a significant increase in the proportion of type IIb fibers, type II fiber atrophy, the presence of type IIC fibers, and reduced activity of the enzyme BOAC. Although an increase in capillaries per millimeter squared was also noted, this was probably due to fiber atrophy because no difference in the number of capillaries surrounding each fiber was observed.

Type IIb fibers represent fibers that are fast twitch, have a low aerobic potential, and are easily fatigued. An increase in the percentage of such fibers might be expected to reduce muscle performance during strenuous exercise; slowly contracting motor units are primarily recruited during low-level exercise. The inverse relation noted between percentage of type IIb fiber and peak exercise $V_o_2$ is consistent with this possibility. However, the area of type IIb fibers was reduced, so that the contribution of type IIb fibers per unit muscle area was not significantly different from the normal subjects, making it difficult to predict the overall impact of the changes in type IIb fibers.

The significance of the increased percentage of type IIC fibers is also difficult to predict. Type IIC fibers are extremely rare in normal subjects except in the neonatal period and during intense physical training.19 Type IIC fibers may represent a population of transitional fibers shifting from type I or IIA to IIb. Alternatively, some investigators have spec-
ulated that they represent fibers in the earliest stages of cell death.

BOAC is a mitochondrial-based enzyme involved in β-oxidation of fatty acids. During rest and light exercise, skeletal muscle uses fat and carbohydrates in approximately equal proportions. With prolonged heavier exercise, fatty acid oxidation contributes significantly more to the energy supply than do carbohydrates. A reduction in enzymes involved in β-oxidation, therefore, potentially could force higher than normal use of carbohydrates during prolonged exercise. This could, in turn, augment lactate accumulation by a mass-action effect and reduce endurance because carbohydrate stores are much smaller than fat stores.

The reduction in BOAC activity also could reflect an overall decrease in mitochondrial-based enzymes. However, we found no reduction in CS activity, a mitochondrial-based enzyme involved in the citric acid cycle.

Our finding of type II fiber atrophy is consistent with prior observations by Dunnigan et al in patients with idiopathic cardiomyopathy. These investigators examined skeletal muscle histochemistry and electron microscopy in 22 young patients (age, 4–39) with idiopathic cardiomyopathy, presenting either as ventricular arrhythmias or cardiac failure. Type II fiber atrophy was found almost exclusively in patients with symptoms of heart failure, occurring in seven of the 11 patients in this group. In contrast, patients with and without symptoms of heart failure exhibited lipid deposition (15 of 22 patients) and endomysial fibrosis (14 of 22 patients). Dunnigan et al speculated that the skeletal muscle changes in their patients may reflect a generalized myopathic process affecting both the heart and skeletal muscle. This seems unlikely, at least for the type II fiber–atrophy changes, because this abnormality was found primarily in patients with symptoms of heart failure. If such atrophy was due to a generalized myopathic process, it should also have been observed in patients presenting with ventricular arrhythmias.

Lipkin et al performed quadriceps muscle biopsies on nine patients with heart failure. Marked generalized fiber atrophy was found in two patients, marked type II fiber atrophy in one patient, increased acid phosphatase staining in six patients, and increased lipid deposition in four patients. The explanation for the more consistent abnormalities noted in our patients compared with the findings of Lipkin et al is uncertain; differences in patient selection and population size may play a role.

Several investigators have also reported preliminary muscle biopsy findings in patients with heart failure. Drexler et al described reduced mitochondrial cristae volume in 21 patients with heart failure, suggesting a reduction in mitochondrial enzyme activity. Sullivan et al reported marked reduction in CS and succinate dehydrogenase activity in 11 patients with heart failure but normal glycolytic enzyme activity. Yancy et al reported reduced skeletal muscle capillarization and succinate dehydrogenase activity in six patients. Although these investigators appear to be observing a different pattern of changes than in the present study, final comparison of our results with these preliminary observations should await completion of the studies.

### Mechanism of Skeletal Muscle Changes

The present study does not indicate the mechanism responsible for the observed skeletal muscle abnormalities. Nevertheless, the most likely mechanism is muscle deconditioning due to inactivity. Previous studies have clearly demonstrated that inactivity can produce fiber atrophy. In studies of totally immobilized limbs, atrophy of both type I and II fibers has usually been noted. However, our population maintained a constant low activity level. Weight analyses of immobilized muscles have demonstrated that immobilization in a stretched position will limit disuse atrophy, whereas, in a shortened position, atrophy is accelerated. Tonic stretch is especially important for the maintenance of type I fibers. For example, when the anterior tibial muscle of the rat is immobilized in a slightly stretched position, a transient hypertrophy of type I fibers with concomitant atrophy of type II fibers occurs. Interestingly, during rat hindlimb suspension, changes in fiber composition of the soleus muscle also occur with transformation of type I to II fibers. Training increases BOAC and CS activity, and inactivity results in decreased enzyme activity. Thus, the particular pattern of atrophy we observed is probably consistent with inactivity.

Malnutrition may also contribute to the abnormalities observed in our patients. Skeletal muscle biopsies of severely malnourished patients have demonstrated extensive necrosis of muscle fibers, neurogeniclike grouping of atrophic type II fibers, and predominant type II fiber atrophy.
Another potential mechanism for the muscle changes observed in our cardiomyopathic outpatients is a generalized myopathic problem involving both the muscle and the heart. This seems unlikely given the absence of any significant difference in histochemical skeletal muscle profile when patients were divided into groups on the basis of the etiology of heart failure. Despite this, we cannot exclude the possibility that some of our patients had a component of chronic alcoholic myopathy. Many of our alcoholic patients continued to abuse alcohol. Ethanol has both direct effects on skeletal muscle, as well as indirect effects from its metabolites, acetaldehyde and acetate, which are oxidized by skeletal muscle. Even a low level of alcohol use can expose the skeletal muscle to an altered pattern of substrate availability and, thus, an altered muscle intermediary metabolism. Nutritional deprivation may lead to an imbalance between protein catabolism and synthesis. These factors may contribute to the type II fiber atrophy frequently observed in chronic alcoholic myopathy. Unlike previous reports on patients with chronic alcoholic myopathy, our patients did not exhibit reduced glycolytic enzyme activity. Other possible explanations for intrinsic skeletal muscle changes include an effect of low muscle flow, alterations in muscle innervation, or hormonal effects. Steroid myopathy, for example, produces type IIb fiber atrophy. Conceivably, stress-related elevations of cortisol by heart failure could influence skeletal muscle.

Relation Between Muscle Biopsies and 
$^{31}$P NMR Observations

$^{31}$P NMR provides a noninvasive method of assessing intracellular metabolic behavior during exercise. In particular, this technique allows assessment of intracellular $p$H, an index of glycolytic activity, and of mitochondrial respiratory control. The relation between P/PCr ratio and work rate provides a noninvasive index of mitochondrial respiratory control that is sensitive to both muscle mitochondrial content and muscle oxygen delivery. During calf exercise, the patients in the present study exhibited higher slopes of the P/PCr ratio to $V_{O_2}$ relation and greater decreases in muscle pH than observed previously in normal subjects; $^{31}$P NMR studies were not performed on the normal subjects in this study for logistic reasons. This is consistent with previous observations that both forearm and calf metabolic responses to exercise are abnormal in patients with heart failure. To test the hypothesis that the abnormal metabolic responses observed in our patients were due to intrinsic muscle changes, we correlated each muscle variable with the work–P/PCr slope and with the pH level noted at peak exercise. No significant relation was noted for any of the variables. Such a finding suggests that the intrinsic muscle changes do not contribute in a major way to the abnormal metabolic responses.

If the changes observed on muscle biopsy do not produce the abnormal metabolic responses, what then is responsible for these responses? One potential factor is muscle underperfusion. However, the levels of systemic $V_{O_2}$ achieved during exercise were modest and unlikely to exceed the capacity of the circulation to deliver blood to working muscle. A more likely explanation for the altered metabolic responses is a reduction in muscle mass. In a previous study of calf exercise, we reported a significant decrease in calf circumference in patients with heart failure. This suggests a reduction in total calf muscle. Lipkin et al. have also reported evidence consistent with a reduced muscle mass. Such a reduction in muscle mass would subject each fiber to an increased load, in turn producing a greater change in the P/PCr ratio and in muscle $p$H.

It should be emphasized, however, that our failure to find a relation between metabolic responses to exercise and muscle biopsy characteristics does not totally exclude such a relation. Within a given individual, there is considerable variability of muscle biopsy results when repeated biopsies are taken from the same muscle. Therefore, it is possible that the single-biopsy results obtained in our patients do not precisely reflect their muscle characteristics. Such inaccuracies might obscure a weak but significant relation between muscle biopsy results and muscle metabolism.

Clinical Implications

Our findings establish the presence of intrinsic skeletal muscle changes in patients with heart failure. To what extent these muscle changes influence the exercise capacity of patients remains to be determined. If these muscle changes do impair exercise performance, it may be possible to improve the exercise capacity of patients by reversing the process directly responsible for the changes. For example, if these changes are due to inactivity, exercise training may reverse the abnormalities and improve the exercise performance of patients. If these changes are due to malnutrition, protein supplementation may be beneficial. Indeed, in several recent nonrandomized studies, patients with heart failure have demonstrated improved exercise capacity with exercise training. An effect of deconditioning on exercise performance may also partially account for the frequent clinical observation of a delay in improvement in maximal exercise performance with therapeutic interventions. These interventions may improve submaximal exercise capacity, permitting patients to increase their activity levels and consequently reverse muscle changes due to inactivity. Alternatively, circulatory dysfunction may be the primary factor limiting the exercise capacity of patients, with intrinsic muscle changes representing an epiphenomenon that does not add appreciably to the exercise intolerance of patients.
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