Gas exchange responses to constant work rate exercise in chronic cardiac failure

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Abstract

Objective—To examine the time course of changes in minute oxygen consumption and other gas exchange variables and heart rate during constant work rate exercise in patients with chronic cardiac failure.

Design—Treadmill exercise with on line measurement of gas exchange and a target duration of 10 minutes.

Subjects—Seven men in New York Heart Association class II, six in class III, and seven controls.

Main outcome measures—Gas exchange variables and heart rate were averaged for the final two minutes of exercise. Time constants were calculated for the increase in all variables.

Results—Consumption of oxygen at the end of exercise (VO₂) was similar in class II patients (mean (95% confidence interval) 14.9 (13.6 to 16.1) ml kg⁻¹ min⁻¹), class III patients (13.2 (11.2 to 15.1) ml kg⁻¹ min⁻¹), and controls (13.3 (12.5 to 14.2) ml kg⁻¹ min⁻¹). The patients reached this VO₂ more slowly with longer exponential time constants of 0.82 (0.59 to 1.04) min in class II and 1.19 (0.86 to 1.51) min in class III, than the 0.49 (0.35 to 0.64) min in the controls. Time constants of other gas exchange variables and heart rate were also longer in patients. By analysis of covariance, peak VO₂ accounted for the between group difference in the time constant for VO₂, suggesting that circulatory factors may be an important cause of the delayed kinetics.

Conclusions—A delayed rise in VO₂ in response to exercise may be responsible for subnormal values of VO₂ early in exercise in patients with chronic cardiac failure.

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Patients with chronic cardiac failure characteristically have a reduced maximal exercise capacity and hence a diminished oxygen consumption (VO₂) at peak exercise. Also, some investigators have shown that VO₂ during incremental work rate protocols is lower in patients with chronic cardiac failure than in controls, when compared at similar absolute work intensities. This occurs during both treadmill⁷ and cycle ergometer⁸ incremental exercise tests, with the duration of exercise stages ranging from one to six minutes. The discrepancy in VO₂ is more pronounced in severely ill patients.³

These findings are interesting as VO₂ at steady state is closely dependent on the work rate being performed.⁵ The implication is either that there exists a basic abnormality in muscle metabolism in chronic cardiac failure, which leads to abnormal muscle bioenergetics,⁶ or alternatively that the attainment of steady state conditions is simply delayed compared with normal.¹ Meakins and Long found a slow rate of rise in VO₂ in one patient with rheumatic heart disease when walking at the same speed as a control subject, although they did not resolve the question as to whether or not the patient's VO₂ eventually reached that of the control.⁷

We hypothesised that the eventual VO₂ is not different when patients with chronic cardiac failure and normal controls perform equal absolute work rates, but rather that in patients there is a slowing of the rate at which this VO₂ is approached. To test this hypothesis, we used a constant work rate exercise test of 10 minutes in duration, and examined the time course of the resultant changes in gas exchange variables and heart rate.

Patients and methods

PATIENTS

Thirteen men (mean age 64 (range 48–76) with compensated chronic cardiac failure took part in the study. Seven patients were in New York Heart Association (NYHA) class II and six in class III. The mean time from diagnosis was 15.1 (range 3–27) months. All had been clinically stable for a minimum of two months before the study. Mean body weight was 67 (range 49–91) kg. The mean left ventricular ejection fraction as determined by radionuclide angiography was 0.28 (range 0.09–0.43), and the cardiothoracic ratio was >0.50 in all cases. The aetiology of chronic cardiac failure was ischaemic heart disease in 12 and alcoholic cardiomyopathy in one. Three patients were in atrial fibrillation, and 10 in sinus rhythm. All patients were being treated with diuretics (median dose 80 (range 40–120) mg frusemide) and four with digoxin. Also three patients were taking flosequinan (Manoplax, Boots UK) and four were taking captopril. None had significant pulmonary disease (from history or spirometry, defined as forced expiratory volume in one second (FEV₁) <75% or FEV₁/FVC (forced vital capacity) <75% predicted), intermittent claudication, or musculoskeletal disease that

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could lead to a premature end of exercise. Clinical evidence of fluid overload (peripheral oedema, raised venous pressure, basal râles) was absent at the time of the study.

**CONTROLS**

Seven men (mean age 63·4 (range 56–67) with no cardiac or pulmonary disease acted as controls. They were healthy on the basis of normal history, examination, and exercise testing. All were sedentary and were taking no medication. Body weight was 81 (range 68–101) kg.

The Ethics Committee of The Queen's University of Belfast granted approval for the study. All the men gave informed consent.

**PROTOCOL**

Before the study all men were familiarised with the apparatus; all patients had performed at least three and controls at least two previous incremental treadmill exercise tests. Data from our laboratory suggest that, at least in the case of maximal exercise tests, one familiarisation test is adequate. Patients and controls then returned to the laboratory on two separate occasions. On the first occasion a symptom limited maximal graded treadmill test was performed for determination of peak VO\(_2\). On the second day patients and controls went to the laboratory one to two hours after a light breakfast or light lunch. A teflon cannula was inserted into an antecubital vein and they rested supine for 10 minutes. After two minutes standing on the treadmill, a constant work rate treadmill test with on line measurement of expired gas analysis was carried out. Treadmill grade was 5% and speed was 2·7 km h\(^{-1}\) in all tests. The patients and controls were asked not to grip the treadmill bars. The test ended after 10 minutes, or earlier if the patients could not tolerate the full 10 minutes. A blood sample for lactate was taken immediately before the end of exercise. The electrocardiogram was monitored continuously in CM5 configuration, and blood pressure was recorded at three minute intervals with a mercury sphygmomanometer. Patients and controls were asked to indicate their perceived level of exertion at the end of exercise by means of a Borg score.

**MEASUREMENT OF GAS EXCHANGE**

Minute ventilation (VE) was measured with a vane turbine placed on the inspiratory side of a non-rebreathing respiratory valve circuit (dead space 88 ml) in conjunction with a ventilometer (PK Morgan, UK). Interruptions of a light beam by the vane were counted to measure inspired volume that was then converted to expired volume with the Haldane correction and standard formulae. Expired gas was led through lightweight tubing into a five litre mixing chamber. This was sampled continuously and expired \(\text{O}_2\) and \(\text{CO}_2\) concentrations determined by paramagnetic and infrared analysis respectively. The outputs from the ventilometer and gas analysers were fed through an analog to digital converter to an Ericsson microcomputer for on line calculation of \(\text{VO}_2\), \(\text{CO}_2\) production (\(\text{VCO}_2\)), and VE. Data points were averages of 15 second periods. Calibration of the ventilometer was carried out weekly with multiple strokes of a standard one litre syringe, and the gas analysers were calibrated before each test with gases of known concentration. Validation of the system has been previously described.\(^{10}\)

**BLOOD SAMPLING AND ASSAY**

Lactate samples were precipitated immediately in 8% perchloric acid and the supernatant assayed by an enzymatic colourimetric method (Sigma, St Louis, MO, USA). The coefficient of variation for the assay was 0·8%.

**STATISTICAL METHODS**

Differences among groups were assessed by analysis of variance and between group comparisons were made with \(t\) tests with Bonferroni's correction. The relation between variables was described by simple correlation. The degree to which differences between the groups could be explained by confounding variables was determined, where appropriate, by performing an analysis of covariance. A P value <0·05 was taken as the level of significance.

**CURVE FITTING FOR THE EXERCISE RESPONSES**

After initial visual inspection, the 15 s averaged gas exchange data were fitted to a single compartment exponential "wash-in" model of the type:

\[
f(t) = A - Be^{-rt},
\]

where \(f(t)\) represents the value of \(\text{VO}_2\), \(\text{VCO}_2\), VE, or heart rate after \(t\) minutes of exercise, \(A\) represents the asymptotic value to which the function rises, \(B\) represents the total increment from baseline to the asymptote, and \(r\) is the time constant. The longer the time constant, the slower are the kinetics of the physiological variable under consideration. The SigmaPlot graphics package (Jandel Scientific, Corte Madera, CA, USA) was used to select values for \(A, B, r\) and \(t\) by multiple iterations so as to achieve a minimum residual sum of squares.

**Results**

**INCREMENTAL EXERCISE**

Peak \(\text{VO}_2\) during symptom limited maximal exercise was much greater in controls (mean (95% confidence interval (95% CI)) 29·7 (25·7 to 33·6) ml kg\(^{-1}\) min\(^{-1}\)) than in patients with either NYHA class II (21·6 (19·4 to 23·8)) or class III symptoms (15·1 (13·6 to 16·7) ml kg\(^{-1}\) min\(^{-1}\)).

**CONSTANT WORK RATE EXERCISE**

Although no adverse events occurred during constant work rate exercise, three patients with class III symptoms were unable to complete the full 10 minutes of the test because of leg fatigue. All patients and controls completed at least six minutes. Table 1 shows the treadmill time, Borg score of perceived exertion, the lactate value and its increment from
rest at the end of exercise, the average $\dot{V}O_2$, $\dot{V}CO_2$, $R$ (respiratory exchange ratio, $\dot{V}CO_2/\dot{V}O_2$), and heart rate. Differences were more pronounced in the class III group. By contrast $\dot{V}O_2$ was not significantly different between patients and controls. In patients taken as a single group, peak $\dot{V}O_2$ was inversely correlated with lactate at the end of exercise ($r = 0.65$, $P < 0.02$) and $R$ ($r = 0.72$, $P < 0.01$).

Figure 1 shows the average responses of $\dot{V}O_2$, $\dot{V}CO_2$, $VE$, and heart rate for the two patient groups and the controls. Repeated measures analysis of variance was performed for the first six minutes of exercise, the period for which complete data for all subjects were available. The $\dot{V}O_2$ displayed a time dependent difference among the groups ($P < 0.02$), such that there was a trend towards lower values in the class III patients during the first two minutes, with a tendency to slightly higher values in both patient groups after three minutes of exercise. By contrast $\dot{V}CO_2$ was greater overall ($P < 0.005$) in both patient groups over the first six minutes, and there was a time dependent effect ($P < 0.0001$), so that the size of the between group difference became greater with time. The response of $VE$ was similar to $\dot{V}CO_2$, with an overall increase in both patient groups ($P < 0.05$). The responses diverged with time ($P < 0.0001$). Heart rate was greater overall in patients ($P < 0.02$), as was the rise over the first six minutes of exercise ($P < 0.0001$).

Table 2 shows the $r$ values for the exponential curves fitted to the plots of $\dot{V}O_2$, $\dot{V}CO_2$,
VE, and heart rate for the first six minutes of exercise. The model satisfactorily described the plots in all cases, as judged by visual inspection. Figure 2 shows representative plots of the \( V_{O2} \), \( V_{CO2} \), VE, and heart rate responses and the corresponding fitted functions for a patient in class III and a control. In patients, especially those in class III, \( \tau \) was longer for all variables, indicating a slower course towards the respective steady state values.

In patients as a whole, \( \tau \) for \( V_{O2} \) was significantly correlated with peak \( V_{O2} \) (fig 3; \( r = 0.73 \), \( P < 0.005 \)), but only showed a weakly positive, non-significant correlation with lactate at the end of exercise (\( r = 0.54 \), \( P = 0.06 \)). Analysis of covariance was carried out on the values of \( \tau \) for \( V_{O2} \) with peak \( V_{O2} \) and lactate as covariates, and confirmed peak \( V_{O2} \) as the main determinant of \( \tau \). Thus, the mean difference in \( \tau \) between patients and controls (0-50 (0-22-0-77) min) was eliminated (0-00 (0-00-0-02) min) when peak \( V_{O2} \) but not when lactate ((0-33 (0-03-0-64)) was taken into account.

### Discussion
The results of this study show that the \( V_{O2} \) eventually reached by patients with chronic cardiac failure in response to a given absolute work rate is in fact not low when compared with that in normal controls. The time constant for the rise in \( V_{O2} \) is longer in the patients, so that during early exercise the \( V_{O2} \) in the patient group tends to be lower than in the controls. This finding is likely to explain the paradox that \( V_{O2} \) at similar absolute work rates may sometimes be significantly lower during incremental exercise tests in patients with chronic cardiac failure than in controls.1-4

![Figure 2](http://heart.bmj.com/)

**Figure 2** Representative plots for rise in (A) \( V_{O2} \), (B) \( V_{CO2} \), (C) VE, and (D) heart rate for the first six minutes of constant work rate exercise. The fitted exponential curves and their equations are also shown. The same men are used in all illustrations.

**Table 2** Values for the time constants (\( \tau \) (min)) describing the rise in \( V_{O2} \), \( V_{CO2} \), VE, and heart rate during the first six minutes of constant work rate exercise.

<table>
<thead>
<tr>
<th>Patients</th>
<th>NYHA class II</th>
<th>NYHA class III</th>
<th>Controls</th>
<th>P value (ANOVA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>( V_{O2} )</td>
<td>0.82 (0.59 to 1.04)*</td>
<td>1.19 (0.86 to 1.51)***</td>
<td>0.49 (0.35 to 0.64)</td>
<td>&lt;0.0005</td>
</tr>
<tr>
<td>( V_{CO2} )</td>
<td>1.46 (0.97 to 1.96)</td>
<td>2.06 (1.58 to 2.55)***</td>
<td>1.00 (0.59 to 1.41)</td>
<td>&lt;0.005</td>
</tr>
<tr>
<td>VE</td>
<td>1.59 (0.89 to 2.30)</td>
<td>2.80 (1.39 to 4.22)**</td>
<td>0.95 (0.60 to 1.30)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Heart rate</td>
<td>1.01 (0.24 to 1.77)</td>
<td>2.19 (1.14 to 3.25)**</td>
<td>0.40 (0.13 to 0.68)</td>
<td>&lt;0.005</td>
</tr>
</tbody>
</table>

Values are mean (95% CI). *P < 0.05; **P < 0.01; ***P < 0.005, Bonferroni correction, compared with controls.
Differences in actual work, and hence \( \dot{V}O_2 \)
may occur between people performing the same apparent external work, especially on
the treadmill.\(^{11}\) Factors such as stride length,\(^{12}\) 
or whether the bars are gripped may serve to
vary the work done. In particular, patients
with chronic cardiac failure take shorter
strides than controls, which may lead to some
increase in \( \dot{V}O_2 \).\(^{13}\) We attempted to minimise
such variables by familiarisation, and by not
allowing the bars to be gripped. Actual differ-
ences in external work rate are unlikely to be
responsible for much of the difference in \( \dot{V}O_2 
\) between patients and controls, as the finding
that \( \dot{V}O_2 \) in patients may be lower has also
been noted in bicycle studies.\(^{14,15}\) These allow
more accurate measurement of external work with
lower variability of \( \dot{V}O_2 \) than treadmill
studies.\(^{15}\) An advantage of treadmill over cycle
ergometer exercise, however, is that the work
done is related to the mass of the subject.
This is important when comparing people of
different weights, as in the current study, as work
rate per kg is then standardised, and \( \dot{V}O_2 \) is
expressed as a function of body weight.

Whereas we believe that all of our patients
were adequately familiarised with the exercise
testing procedures before the study, some of
the controls had performed one less familiarisa-
tion test than the patients. On this basis, it
could be argued that hyperventilation due to
anxiety on the part of the controls contributed to
their rapid gas exchange responses. This
seems unlikely, however, as it is clear from
inspection of figs 1 and 2 that neither R nor the
ratios \( VE/\dot{V}O_2 \) or \( VE/\dot{V}CO_2 \) increased
everously at any time during exercise. This
implies that ventilation was appropriate to
metabolic requirements.

Some investigators have found no differ-
ce in \( \dot{V}O_2 \) between patients with chronic
cardiac failure and normal controls perform-
ing work of equal intensity.\(^{16,17}\) As pointed out
by Cowley et al, many of the patients in these
studies had mild heart failure, and this may
have masked any tendency to reduced levels
of \( \dot{V}O_2 \).\(^3\) In one study, \( \dot{V}O_2 \) tended to be lower in
patients with NYHA class III than class II,\(^2\)
and other investigators noted a non-signifi-
cant trend towards a lower \( \dot{V}O_2 \) during sub-
maximal exercise in patients with lower peak
\( \dot{V}O_2 \).\(^{14}\) This is supported by our finding that
patients with class III have longer values of \( \tau \)
than in patients with class II chronic cardiac
failure, and that a strong inverse relation
exists between peak \( \dot{V}O_2 \) and the \( \tau \) for \( \dot{V}O_2 \).

The kinetics of \( O_2 \) uptake have previously
been found to be slowed in patients with
chronic lung disease\(^{18}\) and pulmonary vascular
disease,\(^{19}\) and also in controls given \( \beta \)
adrenoceptor antagonists.\(^{20}\) In addition healthy peo-
ple exhibit faster \( \dot{V}O_2 \) kinetics after training
than before training.\(^{21,22}\) A slow circulatory
response to exercise is probably present in all
of the above conditions. Our findings of
slowed kinetics, not only of \( \dot{V}O_2 \) but also of
\( \dot{V}CO_2 \), \( VE \), and in particular heart rate, in a
disease characterised by an inadequate circu-
laratory response to exercise,\(^{23}\) lend support to
this suggestion. Moreover, in our study,
analysis of covariance showed that the slowed
kinetics for \( \dot{V}O_2 \) could be accounted for by dif-
ferences in peak \( \dot{V}O_2 \), a measure of functional
capacity highly dependent on cardiac output.\(^{24}\) This provides further evidence that an
impaired circulatory response may be the fac-
tor responsible. Other factors linked to
the deficiency circuit, such as reduced ventila-
tory efficiency,\(^{25}\) may also contribute to
delayed kinetics. Analogous delays in the
return of \( \dot{V}O_2 \) towards resting levels after exer-
cise in chronic cardiac failure have recently
been reported.\(^{27}\)

The \( \dot{V}CO_2 \), \( VE \), R, heart rate, venous lactate,
and Borg score were all greater at the end of
exercise in the patients with chronic cardiac
failure than in the controls. This suggests that
the exercise intensity constituted a greater
physiological strain to the patient group. The
inverse relation in patients between peak \( \dot{V}O_2 \)
and both R and lactate at the end of exercise
is in line with this conclusion.

Lactic acid has long been known to be
excessive in patients with cardiac failure when
exercising at similar absolute work rates to
controls,\(^7,28,29\) and might be expected to be
associated with a reduced \( \dot{V}O_2 \). The energy
supplied by anaerobic processes is, however,
relatively small, and in view of the rather low
lactate values at the end of exercise in our
study, it is likely that anaerobic metabolism
contributed much less than the 20% of total
energy expenditure quoted for maximal exer-
cise.\(^{30}\) Indeed, it is perhaps not surprising that
the \( \dot{V}O_2 \) eventually reached by the patients and
controls should be similar, as the increased
lactate production in the patients may repre-
sent a \( \dot{V}O_2 \) not much greater than the error of
\( \dot{V}O_2 \) measurement (about 2%).\(^31\) Furthermore,
and more importantly in the patients, some of
the deficit in \( \dot{V}O_2 \) arising during early exercise
from anaerobic processes may be paid back
later on. This, along with other as yet uncer-
tain factors,\(^32\) may tend to cause excess \( \dot{V}O_2 \)
during the latter stages of moderate to heavy
exercise and may explain the tendency for \( \dot{V}O_2 \)
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to be greater later on in the exercise test in our patient groups. It has been proposed that lactate is causally associated with a drift upwards in $V_O_2$ after three minutes of steady state exercise in both controls and patients with heart failure.\(^{33-35}\) We were unable to confirm a major association between the lactate at the end of exercise and the prolonged $V_O_2$ kinetics in our patients. Such conclusions must be treated with caution as we measured peripheral venous lactate concentrations that may be influenced by many factors including cardiac output, lactate washout from working muscle, and lactate metabolism by inactive muscle.\(^{36}\) It is also likely that $V_O_2$ and $V_E$ are affected by kinetics of lactate increase\(^{37}\) as well as by the direct influence of circulatory factors.

In conclusion, we have shown slowed kinetics for oxygen uptake and for other gas exchange variables and heart rate in response to exercise in patients with chronic cardiac failure. It is likely that circulatory factors are implicated, particularly the cardiac output response. These slowed kinetics are likely to explain the finding that $V_O_2$ is sometimes less in patients with chronic cardiac failure than in controls at equivalent work rates during incremental exercise.

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