Haemodynamic and neurohumoral response to exercise in patients with congestive heart failure treated with captopril

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SUMMARY The contribution of the renin-angiotensin system to the cardiovascular response to exercise was studied in 12 patients with congestive heart failure. The haemodynamic effects of captopril were measured at rest and during supine bicycle exercise. After captopril administration, resting systemic vascular resistance fell by 26-6% and mean blood pressure by 16-7% and cardiac index increased by 19-7%. During exercise, captopril decreased systemic vascular resistance by 25-6% and mean blood pressure by 8-2% and increased cardiac index by 24-4%. Pulmonary wedge pressure fell by 25% at rest but was not altered by captopril during exercise. Pretreatment plasma renin activity increased from 13-4(16-0) ng/ml/hr (10-3(12-3) mmol/l/hr) at rest to 20-0(27-8) ng/ml/hr (15-4(21-4) mmol/l/hr) during exercise. Pretreatment plasma noradrenaline concentration increased from 659(433) pg/ml (39(25-6) nmol/l) at rest to 2622(1486) pg/ml during exercise (155(88) nmol/l). Captopril favourably alters systemic vascular resistance and cardiac index during exercise in patients with congestive heart failure. This may reflect inhibition of the increased activity of the renin-angiotensin system during exercise in these patients and a subsequent reduction in systemic vasoconstriction.

The renin-angiotensin system contributes to systemic vasoconstriction in patients with congestive heart failure. After the administration of saralasin (an angiotensin II antagonist) and of teprotide, captopril, and enalapril (angiotensin-converting enzyme inhibitors) resting cardiac function improves as a consequence of systemic vasodilatation. A beneficial haemodynamic response during exercise also occurs after captopril administration. This includes an increase in cardiac output and stroke volume and a fall in systemic vascular resistance. The clinical efficacy of captopril has been shown in placebo controlled trials.

In resting patients the vasodilatation that accompanies converting enzyme inhibition correlates with the pretreatment resting plasma renin activity. While plasma renin activity also increases during exercise in normal subjects, it is not established that plasma renin activity increases in patients with heart failure during exercise. Activation of the renin-angiotensin system during exercise would further contribute to vasoconstriction and as a result limit the increase in cardiac output. Captopril may, therefore, inhibit angiotensin mediated vasoconstriction during exercise and thus improve cardiac function.

Some investigators have reported a fall in plasma noradrenaline concentration after converting enzyme inhibition, though this has not been confirmed by others. During exercise, plasma noradrenaline increases in normal subjects and in patients with congestive heart failure. A decrease in sympathetic nervous system activity may contribute to captopril mediated vasodilatation.

This study was therefore designed to assess the haemodynamic and neurohumoral response to captopril during exercise in patients with congestive heart failure. Plasma renin activity and noradrenaline concentrations were assessed at rest and during exercise.
to determine whether haemodynamic changes seen during exercise after converting enzyme inhibition can be explained by inhibition of the renin-angiotensin or sympathetic nervous systems.

Patients and methods

STUDY POPULATION

Twelve patients with symptoms and signs of severe congestive heart failure participated in the study. The group included 11 men and one woman (mean age 59 (11) years). The cause of congestive heart failure was coronary artery disease in six patients as documented by previous myocardial infarction or coronary angiography or both and hypertension in two. Four patients had a primary cardiomyopathy of unknown aetiology. Patients having sustained a myocardial infarction within three months or those with primary valvar pathology were excluded. The mean duration of heart failure was 5-2 (3-5) years (range one month to seven years). Eight of the patients were categorised as being in New York Heart Association class III and four as being in class IV. All had been receiving digitalis and diuretic drugs. The mean radionuclide ejection fraction was 23-7 (11-6)%.

The protocol of the study had the approval of the Institutional Review Board for Human Research, and each patient gave informed consent.

Patients were admitted to hospital and maintained on a diet containing 86 mmol sodium daily for at least three days before the study. All vasodilators were withdrawn at least 48 hours before the investigation. Digitalis, diuretics, and all medications other than clinically necessary antiarrhythmic agents were withheld on the day of the haemodynamic study.

CENTRAL HAEMODYNAMIC ASSESSMENT

On the evening before the study, patients underwent right heart catheterisation with a Swan-Ganz catheter (Edwards Laboratories) and radial artery cannulation. Pressures were measured using a Hewlett-Packard model 1280 strain gauge transducer and recorded on a direct-writing Hewlett-Packard multigraph. Mean pressures were obtained by electronic integration. Zero reference was chosen at a level 5 cm vertically beneath the sternal angle. Heart rate was determined from the simultaneous electrocardiographic signal. Cardiac output was determined by the Fick method as detailed below. Thus measurements were obtained for systolic, diastolic, and mean blood pressure, mean pulmonary artery pressure, pulmonary capillary wedge pressure, and right atrial pressure. Hemodynamic indices and systemic resistance were calculated from pressure and output values according to standard formulas.

RESPIRATORY GAS ANALYSIS AND DETERMINATION OF CARDIAC OUTPUT

A Beckman metabolic cart comprising a volume transducer, a Beckman OM-11 oxygen analyser, and a Beckman UB-2 carbon dioxide analyser was used to determine minute volume oxygen consumption ($V_{O_2}$), and minute volume carbon dioxide production ($V_{CO_2}$). Oxygen and carbon dioxide concentrations in expired air were delivered via a mouthpiece with a low resistance three way valve to a mixing chamber, volume transducer, and gas analyser. The gas analyser was calibrated with 4% carbon dioxide and 16% oxygen calibration gas. A model 1810 Monroe programmable calculator was used to calculate $V_{O_2}$ and $V_{CO_2}$. Blood was collected for arterial and mixed venous oxygen content. Blood for oxygen content was analysed on a Lex-O2-Con oxygen analyser (Lexington Instruments). Cardiac output was calculated as the ratio of the $V_{O_2}$ to the arteriovenous oxygen content difference.

PLASMA RENIN ACTIVITY AND PLASMA NORADRENALINE CONCENTRATION MEASUREMENTS

Samples of mixed venous blood for measurement of plasma renin activity and noradrenaline concentration were collected while the patients were resting supine and at peak exercise, before and after captopril administration. The collection of blood coincided with the haemodynamic measurements. Blood specimens for plasma renin activity and noradrenaline concentration were immediately placed on ice and centrifuged at 4°C. Plasma renin activity was determined by a radioimmunoassay for angiotensin I generation and plasma noradrenaline concentration by a radioenzymatic assay.

STUDY PROTOCOL

Patients were studied after fasting overnight without premedication. Haemodynamic measurements were made while patients were lying supine. Systemic haemodynamic data were collected until three successive determinations 10 minutes apart showed homeostasis. Simultaneous respiratory gas analysis also assured a stable resting state. Patients then began exercise on a Quinton supine bicycle ergometer at a workload of 33 W (200 kpm), pedalling at 40 to 50 rev/min. The duration of this stage was three minutes. The workload was increased by 8 W (50 kpm) every three minutes. Haemodynamic measurements, collection of arterial and mixed venous blood for oxygen content, and respiratory gas analysis were performed during the last minute of each stage. Exercise was terminated when the patient was unable to continue because of dyspnoea or fatigue. Haemodynamic measurements and respiratory gas analysis were
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Table  Effect of captopril on systemic haemodynamics and oxygen consumption at rest and during exercise. Values are mean (SD)

<table>
<thead>
<tr>
<th></th>
<th>Rest</th>
<th>Captopril</th>
<th>p value</th>
<th>Exercise</th>
<th>Captopril</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>VO₂ (ml/min)</td>
<td>239 (45)</td>
<td>268 (130)</td>
<td>NS</td>
<td>689 (94)</td>
<td>678 (94)</td>
<td>NS</td>
</tr>
<tr>
<td>Systemic vascular resistance (dyn s cm⁻¹)</td>
<td>2138 (802)</td>
<td>1569 (614)</td>
<td>&lt;0.01</td>
<td>1719 (766)</td>
<td>1275 (487)</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Blood pressure (mm Hg)</td>
<td>99 (20)</td>
<td>82 (17)</td>
<td>&lt;0.01</td>
<td>113 (24)</td>
<td>104 (21)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Cardiac index (l/min per m²)</td>
<td>2.1 (0.8)</td>
<td>2.6 (0.9)</td>
<td>&lt;0.05</td>
<td>2.9 (15)</td>
<td>39 (16)</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Stroke volume index (ml/m²)</td>
<td>88 (18)</td>
<td>85 (18)</td>
<td>NS</td>
<td>122 (20)</td>
<td>114 (17)</td>
<td>NS</td>
</tr>
<tr>
<td>Heart rate (beats/min)</td>
<td>30 (11)</td>
<td>21 (9)</td>
<td>&lt;0.05</td>
<td>39 (10)</td>
<td>36 (10)</td>
<td>NS</td>
</tr>
<tr>
<td>Pulmonary capillary wedge pressure (mm Hg)</td>
<td>28 (11)</td>
<td>21 (9)</td>
<td>&lt;0.05</td>
<td>122 (20)</td>
<td>114 (17)</td>
<td>NS</td>
</tr>
</tbody>
</table>

*Significant differences between values obtained during the control period and those obtained after captopril administration.

obtained during the minute before exercise termination. A plateau of VO₂ and a disproportionate rise in VCO₂ signalled the onset of anaerobic metabolism and maximal exercise capacity.

Patients were allowed to rest for at least four hours. After baseline conditions were re-established and homeostasis again assured, captopril 25 mg was given orally. Resting haemodynamic measurements were repeated 30 and 45 minutes later. Incremental supine bicycle exercise was then repeated to maximal tolerance as described above, with haemodynamic measurements, blood collection, and respiratory gas analysis being obtained during the last minute of each stage. Each patient again exercised to fatigue as verified by a plateau in VO₂ and a disproportionate increase in VCO₂.

STATISTICAL ANALYSIS

Values are expressed as mean (SD). Data at rest and during exercise and before and after captopril were analysed by the paired t test or the Wilcoxon rank sum test. Measurements obtained at rest and during maximal exercise before treatment were compared with those obtained at rest and during the comparable exercise stage after treatment. Statistical significance is defined as p<0.05.

Results

RESTING HAEMODYNAMIC RESPONSE TO CATOPRIL (TABLE, FIG. 1)

There was no significant difference in any of the resting haemodynamic variables obtained before exercise on the morning of the investigation and before administration of captopril several hours after the completion of exercise. Thus a resting baseline was re-established before captopril administration. Captopril decreased the mean blood pressure by 16.7% (p<0.01) and the systemic vascular resistance by 26.6% (p<0.01), while the cardiac index increased by 19.7% (p<0.05). Since there was no significant change in heart rate, the increment in cardiac index was secondary to a 16.7% increase in stroke volume index (p<0.05). Captopril reduced the pulmonary capillary wedge pressure by 24.6% (p<0.05).

![Fig. 1 Effect of captopril on (a) blood pressure, (b) cardiac index, (c) systemic vascular resistance, and (d) pulmonary capillary wedge pressure at rest and during exercise. Values are mean (SD). * p<0.05 precaptopril vs postcaptopril; ** p<0.01 precaptopril vs postcaptopril; † p<0.01 rest vs exercise; ‡ p<0.01 precaptopril vs postcaptopril; ‡ p<0.01 rest vs exercise.](http://heart.bmj.com/)

![Fig. 2 Effect of captopril on (a) plasma renin activity and (b) plasma noradrenaline concentration at rest and during exercise. Values are mean (SD). ** p<0.01 precaptopril vs postcaptopril; † p<0.01 rest vs exercise; $ p<0.001 rest vs exercise.](http://heart.bmj.com/)
EFFECT OF CAPTOPRIL ON EXERCISE

HAEMODYNAMICS (TABLE, FIG. 1)

During maximal exercise, captopril reduced the mean blood pressure by 8-2% (p<0.01) and systemic vascular resistance by 25.6% (p<0.05). The cardiac index increased by 24.4% (p<0.05) and the stroke volume index by 32.9% (p<0.05). The heart rate during exercise did not differ after captopril compared with pretreatment values. Captopril did not decrease the pulmonary capillary wedge pressure during exercise. There were no significant changes in exercise duration from a pretreatment time of 5-1(1-9) minutes or in maximal oxygen consumption from a pretreatment value of 689(94) ml/min after captopril. The respiratory quotient, or ratio of VO₂/VO₂, was 1.0(0.1) at maximal exercise before treatment and 1.1(0.1) after captopril (NS).

PLASMA RENIN ACTIVITY AND NORADRENALINE CONCENTRATION (FIG. 2)

The resting plasma renin activity was 13-4(16-0) ng/ml/hr (10-3(12-3) mmol/l/hr). During exercise, before captopril, the plasma renin activity increased to 20-0(27-8) ng/ml/hr (15-4(21-4) mmol/l/hr) (p<0.01). After captopril, the resting plasma renin activity increased further to 32-8(39-9) ng/ml/hr (25-2(30-7) mmol/l/hr) (p<0.01) and during exercise the value was 44-5(46-1) ng/ml/hr (34-2(35-5) mmol/l/hr) (p<0.01 vs control).

The pretreatment resting plasma noradrenaline concentration was 659(433) pg/ml (39(26) nmol/l). During supine bicycle exercise, before captopril, values increased to 2622(1486) pg/ml (155(88) nmol/l) (p<0.001). After captopril, neither the resting nor exercise noradrenaline concentration changed significantly.

Discussion

The improvement in resting cardiac function seen after the administration of captopril in this study is similar to that previously reported.4*8 Cardiac output and stroke volume increased as a consequence of the systemic vasodilatory effects of angiotensin converting enzyme inhibition. The fall in pulmonary capillary wedge pressure may be secondary to venodilatation or improved ventricular compliance. Mild and transient increases in limb venous volume have been found with captopril in patients with congestive heart failure and may account, in part, for the reduction in left ventricular filling pressure.6

Indices of cardiac function improved during exercise after captopril administration. Cardiac output was greater and systemic vascular resistance lower than pretreatment values. Since captopril did not alter the heart rate the increase in cardiac output during exercise is attributed solely to an increase in stroke volume. Captopril has no known positive inotropic properties and therefore the increase in stroke volume must be a consequence of systemic vasodilatation.

In this study resting plasma renin activity and plasma noradrenaline concentration were higher than values usually obtained in normal subjects in our laboratory.21 Both plasma renin activity and plasma noradrenaline concentration increased during exercise before captopril. The increase in plasma noradrenaline concentration during exercise in patients with heart failure has been reported by others.16 17 To our knowledge, an increase in plasma renin activity during exercise has not been previously reported in patients with heart failure. It is, however, known to occur in normal subjects.14 15 Plasma renin activity increased after captopril at rest and during exercise. This increase reflects a loss of negative feedback inhibition consistent with a decrease in angiotensin II concentrations. The vasoactive effects of captopril, however, may also be mediated via the kallikrein-kinin and prostaglandin systems; therefore, modulating effects of these hormonal systems cannot be excluded.22 23 After captopril, plasma noradrenaline concentration did not fall either at rest or during exercise. Thus a decrease in sympathetic nervous system activity cannot explain the observed vasodilatation.

During exercise, when autoregulatory phenomenon as well as activation of the renin-angiotensin system and sympathetic nervous system occur, it is not known whether the increased blood flow associated with captopril administration is distributed to exercising muscle or non-exercising tissue. Exogenous administration of angiotensin II produces varying degrees of vasoconstriction depending on the specific regional circulation.24 25 Pharmacological doses of angiotensin II result in pronounced vasoconstriction in the limbs of normal subjects.26 27 In experimental studies using sodium depleted dogs, converting enzyme inhibition significantly improves adrenal, cerebral, coronary, and renal blood flow but causes only a modest increase in skin blood flow and no improvement in muscle blood flow.28 In patients with heart failure studied at rest, captopril does not have a uniform effect on each regional circulation.8 29 30 The increase in renal blood flow is disproportionate to the negligible changes that occur in limb, splanchic, and coronary blood flow.

Since the renin-angiotensin system is stimulated during effort in patients with heart failure, it is possible that the higher concentrations of angiotensin II will have a greater influence on limb vascular resistance during exercise than at rest. Kugler and colleagues, however, have suggested that blood flow to exercising muscle was not increased by angiotensin
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blockade since femoral vein oxygen content measured during bicycle exercise did not change after captopril. In that study, however, limb blood flow was not actually measured.

If blood flow to exercising muscle is influenced by the renin-angiotensin system exercise capacity after the administration of captopril might be expected to improve. Other investigators have assessed the exercise response to captopril and have not found an improvement in exercise capacity after short term administration. An acute increase in exercise capacity and maximal oxygen consumption has been reported in patients with heart failure after enalapril treatment. Studies of limb blood flow during exercise are required during angiotensin II inhibition for further elucidation.

References

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