Exercise metabolism in healthy volunteers taking celiprolol, atenolol, and placebo

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Abstract
Objective—Previous studies have shown that β, selective agents have fewer adverse effects on exercise metabolism than non-selective β blockers, and this has been attributed to their reduced blockade of β, receptors. This study aimed at determining whether a β blocker with partial agonist activity at β, and β, receptors (celiprolol) was better than a conventional β, receptor-blocker (atenolol) in prolonging exercise capabilities.

Methods—After four days of treatment with celiprolol 200 mg, atenolol 50 mg, or placebo, 22 healthy volunteers exercised on a treadmill for two hours at 50% of their maximal oxygen uptake. Resting heart rate and blood pressure were recorded before and after exercise. During exercise, fat oxidation, plasma free fatty acids, glycerol, glucose, and ammonia were measured together with heart rate and perceived exertion.

Results—Mean exercising heart rate was significantly lower in those taking either of the β blockers than in those taking placebo, and significantly lower for those taking atenolol rather than celiprolol. Fat oxidation was significantly lower for those taking celiprolol (38.8 (SD 12.2)% P<0.01) and atenolol (36.6 (15.9)% P<0.01) compared with placebo (45.6 (14.1)%). For the first 15 minutes of exercise, fat oxidation was significantly lower for those taking atenolol (24.6 (12.8)% P<0.01) than celiprolol (29.6 (14.3)%). The rise in plasma free fatty acids and glycerol during exercise was also significantly attenuated by both β blockers in comparison with the rise in those taking placebo (P<0.01).

Conclusions—Both celiprolol and atenolol reduced fat oxidation compared with placebo. For the first 15 minutes of exercise fat oxidation was preserved by celiprolol, but not atenolol. This preservation of fat oxidation during the early part of exercise may confer some small benefit to patients who take β blockers and intend to exercise regularly. However, we did not detect significant differences between atenolol and celiprolol in overall mean fat oxidation or perceived exertion in this study.

Keywords: atenolol; celiprolol; exercise; β blockade; fat oxidation

Prolonged aerobic exercise and β blockade both have important roles in the prevention and management of cardiovascular disease. However, fatigue is a commonly reported side effect of β blockade, and may be due to reduced cardiac output, reduced liver and muscle glycoegenolysis, and reduced lipolysis, all of which may reduce the capacity for exercise to some degree.

As the therapeutic effects of β blockers are due to their β, receptor blocking properties, some reduction in adipose lipolysis (β, and β, mediated) must remain an unavoidable side effect of β, receptor blockade.

Celiprolol is a selective β, receptor blocker with some partial agonist activity at both β, and β, receptors. The purpose of this study was to examine exercise metabolism, and particularly fat oxidation, during submaximal exercise preceded by four days’ oral administration of a β, selective drug (atenolol), a β, selective drug with β, agonist properties (celiprolol), or placebo.

Method

SUBJECTS
Twenty four healthy subjects (12 male, 12 female, body mass index ≤32) underwent a routine medical examination and a blood test. No abnormalities were found. Two subjects (one male, one female) were later excluded from the trial when it was discovered that they were taking previously undisclosed drugs. The remaining 22 subjects then carried out a fitness test (constant speed and increasing gradient protocol, two minute stages until exhaustion) to measure maximal oxygen uptake (V02 max).

Subjects were non-specifically trained with a V02 max of less than 60 ml/kg/min. On another day, a habituation walk was completed on a motorised treadmill at a speed and gradient producing an exercise intensity of 50% of their V02 max. No blood samples were taken during this stage of the trial, but heart rate, perceived exertion (category ratio scale during steady state exercise and Borg scale during incremental exercise) subjective feeling, expired air were measured every 15 minutes. Subjects also completed a mood state questionnaire before and after exercise. The study was approved by the South Birmingham Health Authority research ethics committee. All subjects gave written, informed consent to participate in the study.

STUDY DESIGN
The study was a double blind, randomised crossover design, with all subjects completing all conditions: celiprolol 200 mg daily, atenolol
Impact and atenolol monitored was Trial days final dose minimum of patterns walking utilisation. mg at treadmill standardised samples (7 ml) expired exercise, 100 terminated for exercise, 21 calorimetry. Oxalate.

ENERGY EXPENDITURE DURING EXERCISE TRIALS The mean energy expenditure during the two hours' exercise trial was 3645 (993) kJ for those taking placebo, 3579 (1077) kJ for atenolol, and 3523 (913) kJ for celiprolol. There were no significant differences between treatments.

HEART RATE AND BLOOD PRESSURE Table 1 shows the mean resting heart rates and systolic and diastolic blood pressures (mmHg). Resting heart rate before exercise was higher for those taking celiprolol (68.7 (9.6) beats/min, P<0.05) and lower for atenolol (54.0 (7.9) beats/min, P<0.01) than for placebo (65.2 (7.7) beats/min). Resting systolic pressure was also higher for celiprolol (126.1 (7.1) mmHg, P<0.01) and lower for atenolol (116.0 (9.0) mmHg, P<0.01) than for placebo (121.8 (9.4) mmHg).

Figure 1 shows the heart rate at 15 minute intervals during exercise with each treatment. The mean heart rate was significantly lower for celiprolol 200 mg (114 (7) beats/min, P<0.01) and for atenolol 50 mg (99 (7) beats/min, P<0.01) than for placebo (132 (7) beats/min), and significantly lower for atenolol than celiprolol (P<0.01). The same relation continued during the incremental exercise period, with mean heart rate reaching 178 (9) beats/min at 90% Vo2 max for placebo, 153 (11) beats/min for celiprolol, and 141 (9) beats/min for atenolol. Heart rate remained significantly lower for both celiprolol and atenolol (P<0.01 at all time points) than for placebo, and significantly lower for atenolol than celiprolol.

During rest after exercise, both celiprolol and atenolol produced a reduction in systolic and diastolic pressures, though the reduction was significantly greater with atenolol than with celiprolol.
Table 1  Resting heart rate and systolic and diastolic blood pressures, before and after exercise for 22 subjects taking atenolol 50 mg, placebo, or celiprolol 200 mg. (Results are shown as means (SD))

<table>
<thead>
<tr>
<th>Treatment</th>
<th>15 Minutes before exercise</th>
<th>15 Minutes after exercise</th>
</tr>
</thead>
<tbody>
<tr>
<td>Placebo</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heart rate (beats/min)</td>
<td>65.2 (7.7)</td>
<td>85.1 (10.5)</td>
</tr>
<tr>
<td>Systolic BP (mm Hg)</td>
<td>121.8 (9.4)</td>
<td>120.8 (7.8)</td>
</tr>
<tr>
<td>Diastolic BP (mm Hg)</td>
<td>66.9 (7.6)</td>
<td>65.8 (7.3)</td>
</tr>
<tr>
<td>Celiprolol 200 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heart rate (beats/min)</td>
<td>68.7 (9.6)*</td>
<td>83.5 (9.0) NS</td>
</tr>
<tr>
<td>Systolic BP (mm Hg)</td>
<td>126.1 (7.1)*</td>
<td>118.4 (8.1)*</td>
</tr>
<tr>
<td>Diastolic BP (mm Hg)</td>
<td>66.1 (5.5) NS</td>
<td>62.7 (7.6)†</td>
</tr>
<tr>
<td>Atenolol 50 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heart rate (beats/min)</td>
<td>54.0 (7.9)†‡</td>
<td>70.2 (9.6)†‡</td>
</tr>
<tr>
<td>Systolic BP (mm Hg)</td>
<td>116.0 (9.0)†‡</td>
<td>111.1 (6.8)†‡</td>
</tr>
<tr>
<td>Diastolic BP (mm Hg)</td>
<td>68.0 (7.1)†‡</td>
<td>58.5 (7.3)†‡</td>
</tr>
</tbody>
</table>

*Significantly different from placebo; †p < 0.05; ‡p < 0.01.
Significantly lower than celiprolol †p < 0.01.

INDIRECT CALORIMETRY
The contribution of fat as a fuel towards energy expenditure during the two hours' exercise trials is expressed as a percentage of the total energy expenditure during each trial (fig 2). Fat oxidation at rest was not measured. During exercise, mean fat oxidation was significantly lower for celiprolol 200 mg (38.8 (12.2)%, P<0.01) and atenolol 50 mg (36.6 (15.9)%, P<0.01) than for placebo (45.6 (14.1)%). The difference in overall mean fat oxidation with celiprolol and atenolol was not significant.

After 15 minutes' exercise, fat oxidation was significantly lower for atenolol 50 mg (24.6 (12.8)%, P<0.01) than for placebo (32.2 (12.6)%, but the difference for celiprolol 200 mg was not significant (29.6 (14.3)%, P = NS). After 120 minutes, fat oxidation was lower for both atenolol (46.8 (14.0)%, P<0.01) and celiprolol (47.4 (7.5)%, P<0.01) than for placebo (57.7 (12.1)%). During the subsequent incremental exercise stages the RER was significantly higher for both celiprolol and atenolol than for placebo during each two minute stage. However, as this was no longer steady state exercise, calculation of fat oxidation rates from the RER was no longer valid. There were no significant differences between celiprolol and atenolol at any of the higher exercise intensities.

PLASMA VOLUME CHANGES
During exercise, plasma volume did not change significantly from resting values and there were no significant differences between treatments.

PLASMA FREE FATTY ACID (FFA) AND GLYCEROL CONCENTRATION
Figure 3 shows the plasma FFA concentration (µmol/l). There were no significant differences between treatments at rest. As exercise progressed, there was a significant increase in plasma FFA concentration with all treatments (P<0.01), confirming known responses to prolonged exercise. During exercise, there were significant differences between treatments (P<0.01). After 120 minutes of exercise, plasma FFA concentration was significantly lower for celiprolol 200 mg (493.2 (196.9) µmol/l, P<0.01) and atenolol 50 mg (388.3 (238.9) µmol/l, P<0.01) than for placebo (700.1 (245.6) µmol/l).

During the incremental exercise stages, the FFA concentration remained lower for both celiprolol and atenolol than for placebo (P<0.01), and lower for atenolol than for celiprolol (P<0.01) at 122 minutes but not at 128 minutes (P = NS).

Plasma glycerol (µmol/l) (fig 3) increased during exercise regardless of treatment (P<0.01), confirming known responses to prolonged exercise. At rest there were no significant differences between treatments. After 120 minutes of exercise, plasma glycerol was significantly lower for both celiprolol (207.0 (80.8) µmol/l, P<0.05) and atenolol (195.3 (94.2) µmol/l, P<0.01) than for placebo (244.9 (85.8) µmol/l). During the incremental exercise period, and during rest after exercise, there were no significant differences between celiprolol, atenolol, and placebo (P = NS). Venous blood samples proved difficult to draw during the incremental exercise period, particularly from subjects receiving active drug treatment, and therefore many samples were missed. The data are therefore from nine subjects only.

PLASMA AMMONIA CONCENTRATION
Plasma ammonia (µmol/l) (fig 3) increased during exercise regardless of treatment (P<0.01). After 60 minutes, plasma ammonia was higher for celiprolol (76.1 (39.3) µmol/l, P<0.05) and atenolol (83.5 (49.7) µmol/l, P<0.01) than for placebo (52.8 (28.7) µmol/l). After 120 minutes, ammonia was still higher for atenolol (97.0 (59.0) µmol/l, P<0.01) and celiprolol (87.0 (63.8) µmol/l, P<0.01) than for placebo (63.1 (24.5) µmol/l). During the incremental exercise stages, plasma ammonia was higher for atenolol at 122 minutes and 128 minutes (P<0.01) than for placebo, but the difference between celiprolol and placebo was not significant.

PLASMA GLUCOSE CONCENTRATION
Plasma glucose (mmol/l) fell from rest (5.2 (1.0)) as exercise started regardless of treatment (P<0.01), but euglycaemia was maintained throughout steady state (4.9 (0.48) at T120) and incremental exercise periods. There were no significant differences between treatments.

PLASMA LACTATE CONCENTRATION
With all treatments, plasma lactate (µmol/l) progressively and significantly fell from rest (1932.0 (560.0)) during steady state exercise at 50% V̇O₂ max (858.0 (298.0) at T120, P<0.01), and there were no significant differences between treatments. During the incremental exercise stages, plasma lactate increased significantly with all treatments (P<0.01), but
Impact of celiprolol and atenolol on exercise metabolism

**Figure 1** Heart rate during two hours’ exercise at 50% VO$_2$ max, followed by two minutes at 60, 70, 80, and 90% VO$_2$ max while taking placebo, celiprolol 200 mg, atenolol 50 mg. Significantly lower than placebo *P<0.01, significantly lower than celiprolol †P<0.01.

**Figure 2** Fat oxidation during two hours’ exercise at 50% VO$_2$ max, followed by two minutes at 60, 70, 80, and 90% VO$_2$ max while taking placebo, celiprolol 200 mg, or atenolol 50 mg. Significantly lower than placebo *P<0.01, significantly lower than celiprolol †P<0.05. RER = respiratory exchange ratio.

Again there were no significant differences between treatments.

**PLASMA POTASSIUM CONCENTRATION**
With all treatments, plasma potassium increased significantly from rest (4.2 (0.67) mmol/l) during steady state exercise at 50% VO$_2$ max (P<0.01 for all treatments). After 120 minutes’ exercise mean plasma potassium levels on each treatment were: placebo 5.13 (0.48) mmol/l, celiprolol 5.14 (0.40) mmol/l, atenolol 5.22 (0.47) mmol/l, and there were no significant differences between treatments.

**RATE OF PERCEIVED EXERTION (RPE) FOR LEG EFFORT**
Figure 4 shows the RPE for leg effort during the two hours’ exercise, which increased as exercise progressed (P<0.01). After 120 minutes, perceived leg effort was significantly higher for celiprolol (3.6 (2.1), P<0.01) and atenolol (3.4 (2.1), P<0.01) than for placebo (2.7 (1.6)), and significantly higher for celiprolol than atenolol (P<0.05).

**RPE FOR RESPIRATORY EFFORT**
The RPE for respiratory effort increased as exercise progressed (P<0.01), and after 120 minutes was higher for atenolol (2.5 (1.4), P<0.05) and celiprolol (2.7 (1.7), P<0.01) than for placebo (2.1 (1.5)).

**RPE FOR LEG PAIN**
The RPE for leg pain increased as exercise progressed (P<0.01), and after 120 minutes' exercise leg pain was higher for atenolol (2.5 (2.2), P<0.01) and celiprolol (2.8 (2.3), P<0.01) than for placebo (1.7 (1.6)), and higher for celiprolol than atenolol (P<0.05).

**FEELING SCALE**
Reported feeling was from -5 (very bad) to +5 (very good) during two hours’ steady state exercise at 50% VO$_2$ max. Subjects generally reported feeling worse as exercise progressed (P<0.01), and there was a trend for subjects taking drug treatments to report that they felt worse than reported by those taking placebo (at 120 minutes: placebo +1.6 (1.2), celiprolol +0.5 (1.3), atenolol +1.0 (1.3)), but differences between treatments were not statistically significant.

**RPE DURING INCREMENTAL EXERCISE**
The RPE increased with exercise intensity (P<0.01). At 60% VO$_2$ max (122 minutes) RPE was higher for celiprolol (12.3 (2.3), P<0.01) than for placebo (11.4 (1.3)), but the difference between placebo and atenolol was not significant (11.9 (1.5)). At 90% VO$_2$ max (128 minutes) RPE was higher for atenolol (16.5 (2.7), P<0.01) and celiprolol (16.3 (2.5), P<0.01) than for placebo (15.6 (2.1)).

**Discussion**
This study compared celiprolol and atenolol at doses with comparable clinical efficacy. In the healthy normotensive subjects in this trial, each drug had contrasting effects on the resting heart rate before exercise (when endogenous adrenergic stimulation would be minimal); celiprolol produced a slight increase and atenolol produced a decrease. Similar differences were evident in resting systolic blood pressure before exercise. The increase in resting heart rate and systolic blood pressure for patients taking celiprolol can be explained by the drug’s partial agonist properties at β1 and β2 receptors. During recovery after exercise, when endogenous adrenergic stimulation would be higher (30 minutes after exercise, plasma adrenaline would still be twice resting values before exercise), systolic and diastolic blood pressures were reduced by both atenolol and celiprolol.

The partial agonist effect of celiprolol at β1 and β2 receptors may also be responsible for the observed trend toward higher resting levels of plasma FFA for those taking celiprolol (58.0 (4.18) µmol/l) rather than atenolol (32.8 (21.3) µmol/l), which continued throughout exercise (fig 3). Although this trend might also be expected to be observed with plasma glycerol levels, the fact that glycerol is cleared more slowly (hepatic clearance) from plasma than FFA (rapid uptake by exercising muscle) might account for the similarity in glycerol levels for both celiprolol and atenolol throughout exercise.
Both celiprolol and atenolol significantly reduced fat oxidation compared with placebo. For the first 15 minutes of exercise, fat oxidation with celiprolol and placebo was not significantly different, but was significantly lower with atenolol than with placebo. There was a trend for atenolol to reduce fat oxidation more than celiprolol over the first hour, but the difference was not significant.

The very low level of resting mean plasma FFA for placebo (53 (48) μmol/l) might have been a response to the carbohydrate meal before exercise. Plasma FFA and plasma glycerol (fig 3) concentrations were not significantly different between celiprolol and atenolol for the same early period of exercise. In fact, for the first 30 minutes of exercise, plasma FFA and plasma glycerol values were almost identical for both drug treatments. Therefore it is unlikely that the early differences noted in the rates of fat oxidation between celiprolol and atenolol are due to different effects on adipose lipolysis and plasma FFA concentration. Possibly, changes in intramuscular lipolysis may be responsible.

Cleroux and Leenen have suggested that intramuscular lipolysis is under β2 receptor control.11 We believe that the higher rate of fat oxidation with celiprolol during the early part of exercise is due to the β2 receptor stimulation, which increases the rate of intramuscular lipolysis during the onset of exercise when endogenous adrenergic stimulation is low. However, possibly, the β2 agonist effect in this study was minimal, and a higher dose of 400 mg might have increased β2 receptor stimulation, producing greater changes in fat metabolism. A future study might address this point. The maintenance of fat oxidation delays the depletion of glycogen stores, thereby reducing any metabolic basis for premature fatigue. The plasma ammonia data appear to support this hypothesis, as there is a trend for plasma ammonia to be higher for atenolol during steady state exercise, and during the incremental exercise period.

The origin of the increased plasma ammonia during exercise of moderate intensity is still debated, but it is well known that plasma ammonia rises with increased exercise intensity and duration, and with depleted glycogen stores,30 31 and may be a useful marker of metabolic stress. There are two possible explanations for the observed increases in plasma ammonia during prolonged exercise. One is that branch chain amino acid oxidation increases,32 and the other (perhaps more likely) is that when substrate supply to exercising muscle fat in this case) is restricted, degradation of ADP to AMP and IMP occurs, producing ammonia by the action of AMP deaminase.31 32 β Blockade,33 and particularly non-selective β blockade,37 significantly increases plasma ammonia during exercise of moderate intensity, suggesting that metabolic stress is increased in those exercising and taking β blockers.

In previous studies, reduced fat oxidation correlated well with increased perceived exertion.14 35 However, in this study perceived

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**Figure 3** Plasma free fatty acid (FFA), glycerol, and ammonia (NH₃) during two hours' exercise at 50% Vᵒ₂ max, followed by two minutes at 60, 70, 80, and 90% Vᵒ₂ max, while taking placebo, celiprolol 200 mg, atenolol 50 mg. Significantly lower than placebo: *P<0.05, †P<0.01 significantly lower than celiprolol ‡P<0.01.

**Figure 4** Rate of perceived exertion (RPE) for leg effort during two hours' exercise at 50% Vᵒ₂ max while taking placebo, celiprolol 200 mg, or atenolol 50 mg. Significantly higher than placebo *P<0.01, significantly higher than atenolol †P<0.05.
exertion for leg effort and leg pain during steady state exercise was higher for celiprolol than for atenolol despite fat oxidation being better maintained by celiprolol. The reason for this disparity is unclear.

In conclusion, both atenolol and celiprolol attenuated the normal rise in fat oxidation during prolonged exercise. Any reduction or delay in fat oxidation places increased demand on glycogen stores, and could lead to a reduction in exercise capacity. Celiprolol appeared to maintain fat oxidation at least during the first 15 minutes of exercise. Further investigation of $\beta$ blockers with $\beta_1$ agonist properties is warranted.

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