DOES EXERCISE ALTER ANAEROBIC THRESHOLD IN CORONARY ARTERY DISEASE DURING BETA BLOCKADE?

S. N. KOYAL, PhD, FACSM, R. J. STUART, MD, Ruth LUNDSTROM, MS, RPT, V. THOMAS, BA and M. H. ELLESTAD, MD, FACC

Division of Cardiology, Memorial Medical Center of Long Beach (University of California – Irvine),
2801 Atlantic Avenue, Long Beach, California 90801-1428

ABSTRACT

The effect of propanolol on cardiac patients undergoing exercise training is reported to increase exercise tolerance and maximum oxygen uptake (VO2 max) but its effect on anaerobic threshold (AT) is unknown. It was the purpose of this study to determine the role of exercise training with propanolol on AT in patients with coronary artery disease (CAD). Eight men and one woman with significant (CAD) were selected for this study. Each patient completed a maximum treadmill stress test (MTST) following the Bruce protocol on propanolol 40-160 mg/day as a control study. Cardiorespiratory variables were measured at rest and at each stage of the treadmill test. These patients underwent an exercise training programme for 12-16 weeks on the same dose of propanolol. Training sessions were for a minimum of 30-40 minutes, 3 times a week, with training heart rate of 75%-85% of the pretraining peak heart rate. Training heart rate ranged from 98 to 128 beats/min. They were retested with a MTST after the training programme, on the same dose of propanolol. AT was calculated noninvasively by measuring respiratory variables every 30 seconds in relation to work increment. AT was identified by measuring the time course of VE, VCO2, VE/V02, etc. in relation to incremental work. The mean values of VO2, O2P and % VO2 max at AT before and after training on propanolol were as follows: VO2 = 1.43 L/min ± .25 and 1.86 L/min ± .44, O2P = 14.39 ± 2.40 and 18.73 ± 4.00 ml/beat, % of VO2 max = 68.20 ± 6.31 and 73.59 ± 5.84. The mean changes of VO2, O2P, and % of VO2 max were +.43 L/min ± .20 (P < .003), + 4.38 ± 2.55 (P < .003) and ± 5.07% ± 4.84 (P < .001). After exercise training on propanolol, the mean peak exercise tolerance time and absolute VO2 max increased by 2.8 min (from 9.0 to 11.8 min) (P < .001) and 22.7% (P < .007), respectively. We conclude that the increase in anaerobic threshold in patients with coronary artery disease may be due to improvement in VO2 max, increased stroke volume, and peripheral O2 extraction.

Key words: Coronary artery disease, Exercise training, Propanolol, Anaerobic threshold, Maximum oxygen uptake, Oxygen pulse, Minute ventilation, Ventilatory equivalents.

INTRODUCTION

The use of beta adrenergic blocking agents like propanolol is well established in the treatment of symptomatic patients with coronary artery disease. Propanolol reduces blood pressure and heart rate during physical or emotional stress. Controversy however, has developed regarding the effect of propanolol on cardiovascular training.

The adequacy of circulatory and metabolic responses to exercise in man can be described by the determination of maximum oxygen uptake (VO2 max) and other cardiosrespiratory variables; e.g. cardiac output, blood pressure, O2 pulse, CO2 production and ventilatory equivalents for O2 and CO2.

In the normal subject, during a steady-state or progressive exercise test, there is an interaction between the cardiovascular and pulmonary system to meet the metabolic demands of working muscles. Energy sources for moderate levels of exercise, where oxygen demand is matched by oxygen supply, are usually derived from aerobic metabolism. With incremental exercise, oxygen delivery eventually becomes inadequate, and anaerobic metabolism occurs. At this point, anaerobic threshold (AT) is reached and accompanied by an increase in VCO2 and lactate production (Wasserman et al, 1973). With progressive work increments in exercise the observation of heart rate, blood pressure, electrocardiogram (ECG) and respiratory changes identify the time of onset of the anaerobic threshold. Respiratory parameters like minute ventilation (VE) gas exchange ratio, and ventilatory equivalents for oxygen and carbon dioxide also provide useful information for the detection of anaerobic threshold. A number of authors, Wasserman et al, 1973; Wasserman and Whipp, 1975; Davis et al, 1979; Matsumura et al, 1983, have defined anaerobic threshold as the level of work or oxygen consumption just below the point at which metabolic acidosis and other associated changes in
gas exchange take place. Hossack et al., 1981, in their studies on propanolol on exercise training “prescription” demonstrated a minor but insignificant change in the ratio of maximal oxygen uptake to maximum heart rate. Contrary to Hossack et al., 1981; Obma et al., 1979; Thadani and Parker, 1979; Pratt et al., 1981; Vanhees et al., 1982; and recently Gordon et al., 1983, showed an improved cardiovascular and ventilatory responses after training in patients with coronary heart disease during beta-adrenergic blockade.

It was the purpose of this investigation to determine the effect of exercise training with a long term beta-adrenergic blocking agent on maximal oxygen uptake and the anaerobic threshold in patients with coronary artery disease.

METHODS

Eight ambulatory men and one woman with significant coronary artery disease, documented by angiographic study, were selected at random for this study from among patients attending our clinic. The physical characteristics of the patients were as follows: (a) their ages varied from 42 to 63 years with an average of 52.1 ± 6.7 yr; (b) Height ranged from between 165 to 185 cms with a mean height of 177.6 ± 5.8 cm; and (c) Body weight varied from 64 to 99 kg with an average weight of 82.8 ± 11.4 kg. Patients were receiving orally a nonselective beta adrenergic blocker, — propanolol 40-160 mg/day. It was started prior to the initial test and maintained during the final treadmill test for a 12-16 week period of cardiovascular training, and through the final treadmill stress test. The exercise training consisted of three sessions a week of 20 min/pump/jog or cycle-ergometer exercise at a level of approximately 75%-85% of maximal exercise capacity demonstrated during the initial maximal treadmill test for a minimum of 30 minutes. A pre-exercise warm-up of 15-20 minutes with stretching calisthenics preceded the formal cardiovascular training session and ended with a cool-down period of 10-15 minutes.

Before joining the cardiovascular training programme all patients underwent a maximum treadmill stress test following the Bruce protocol (Bruce et al., 1963) while on their usual dose of propanolol. Oxygen uptake and other cardiorespiratory variables were continuously monitored by the Beckman Metabolic Cart (Beckman, Fullerton, CA). Cardiorespiratory variables such as oxygen uptake (V02), heart rate (HR), expired pulse (O2P), metabolic equivalent (met), respiratory gas exchange ratio (RER), ventilatory equivalents for O2 and CO2 (VE/V02 and VE/CO2), tidal volume, respiratory ratio, rate of respiratory dead space and tidal volume ratio (VD/VT) were measured and printed out at 30 s intervals at rest and during each stage of the treadmill exercise.

During the test, the patient had a rubber mouthpiece and nose clip, and was connected through a nonbreathing Rudolph valve (dead space = 45 ml). The expired air was sent through the mixing chamber into an assembly for measuring expired volume, breath counting, and measurement of expired air temperature. For all sampling conditions, a pump drew off the sample through O2 and CO2 analysers and a nominal flow rate of 50 ml/min. Although a mouthpiece is objectionable to some patients, the diagnostic value of gas exchange information justified its use. The loss of verbal communication with a subject during the test was overcome with prearranged hand signals. The ambient temperature of the laboratory in which the patients were tested was 22.4 ± 1.5°C.

Measurement of maximal oxygen uptake (V02 max) and other cardiorespiratory variables were repeated on each individual patient using an identical protocol after 12-16 weeks of supervised physical training.

### TABLE I

<table>
<thead>
<tr>
<th>Age (yr)</th>
<th>Before</th>
<th>After</th>
<th>% Diff.</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>52.1 ± 6.75</td>
<td>60.1 ± 6.35</td>
<td>9.7 ± 0.05</td>
<td>.005</td>
<td>.02</td>
</tr>
</tbody>
</table>

### RESULTS

Table I represents the mean values with the standard deviation of physical characteristics and responses to maximal treadmill test before and after training in patients with coronary artery disease (CAD). The data demonstrate a significant increase in V02 max, maximum O2 pulse, absolute treadmill time (duration) and reduction in maximum heart rate before and after training of 12-16 weeks.

### TABLE II

<table>
<thead>
<tr>
<th>Maximal Oxygen Uptake (V02, Max)</th>
<th>Before</th>
<th>After</th>
<th>% Diff.</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>F.T.</td>
<td>2.73</td>
<td>2.85</td>
<td></td>
<td></td>
</tr>
<tr>
<td>J.R.</td>
<td>2.00</td>
<td>2.65</td>
<td></td>
<td></td>
</tr>
<tr>
<td>E.F.</td>
<td>2.49</td>
<td>3.25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>K.L.</td>
<td>2.35</td>
<td>2.88</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R.Y.</td>
<td>1.74</td>
<td>2.80</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B.W.</td>
<td>1.46</td>
<td>1.54</td>
<td></td>
<td></td>
</tr>
<tr>
<td>E.H.</td>
<td>1.87</td>
<td>1.79</td>
<td></td>
<td></td>
</tr>
<tr>
<td>J.H.</td>
<td>2.39</td>
<td>3.34</td>
<td></td>
<td></td>
</tr>
<tr>
<td>D.T.</td>
<td>1.87</td>
<td>2.23</td>
<td></td>
<td></td>
</tr>
<tr>
<td>X</td>
<td>2.07</td>
<td>2.53</td>
<td></td>
<td>.007</td>
</tr>
<tr>
<td>SD</td>
<td>± 0.41</td>
<td>± 0.81</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SEM</td>
<td>± 0.14</td>
<td>± 0.20</td>
<td></td>
<td>.000</td>
</tr>
</tbody>
</table>

Table II Individual data of maximum oxygen uptake (V02 max) of the patients with coronary artery disease before and after endurance training with propanolol.

p = probability, (P) = propanol
TABLE III

VO₂ at Anaerobic Threshold (AT)

<table>
<thead>
<tr>
<th>Name</th>
<th>Before Ex. Training $\bar{\theta}$ (P)</th>
<th>After Ex. Training $\bar{\theta}$ (P)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F.T.</td>
<td>60.1</td>
<td>67.3</td>
</tr>
<tr>
<td>J.R.</td>
<td>57.5</td>
<td>74.0</td>
</tr>
<tr>
<td>E.F.</td>
<td>66.1</td>
<td>69.0</td>
</tr>
<tr>
<td>K.L.</td>
<td>68.0</td>
<td>70.7</td>
</tr>
<tr>
<td>R.Y.</td>
<td>69.0</td>
<td>65.3</td>
</tr>
<tr>
<td>B.W.</td>
<td>78.0</td>
<td>81.0</td>
</tr>
<tr>
<td>E.H.</td>
<td>70.0</td>
<td>76.0</td>
</tr>
<tr>
<td>J.H.</td>
<td>76.0</td>
<td>81.0</td>
</tr>
<tr>
<td>D.T.</td>
<td>72.0</td>
<td>78.0</td>
</tr>
</tbody>
</table>

$\bar{x} = 1.43 \text{L/min}$
SD $\pm 0.22$ = 0.44
SEM $\pm 0.08$ = 0.14

Table III: Oxygen uptake at anaerobic threshold of the individual patients before and after exercise training with propranolol.
$p = probability, (P) = propranolol$

TABLE IV

Percentage of VO₂ Max at A.T.

<table>
<thead>
<tr>
<th>Name</th>
<th>Before Ex. Training $\bar{\theta}$ (P)</th>
<th>After Ex. Training $\bar{\theta}$ (P)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F.T.</td>
<td>60.1</td>
<td>67.3</td>
</tr>
<tr>
<td>J.R.</td>
<td>57.5</td>
<td>74.0</td>
</tr>
<tr>
<td>E.F.</td>
<td>66.1</td>
<td>69.0</td>
</tr>
<tr>
<td>K.L.</td>
<td>68.0</td>
<td>70.7</td>
</tr>
<tr>
<td>R.Y.</td>
<td>69.0</td>
<td>65.3</td>
</tr>
<tr>
<td>B.W.</td>
<td>78.0</td>
<td>81.0</td>
</tr>
<tr>
<td>E.H.</td>
<td>70.0</td>
<td>76.0</td>
</tr>
<tr>
<td>J.H.</td>
<td>76.0</td>
<td>81.0</td>
</tr>
<tr>
<td>D.T.</td>
<td>72.0</td>
<td>78.0</td>
</tr>
</tbody>
</table>

$\bar{x} = 68.2 \pm 6.31$ = 5.84
SEM $\pm 2.23$ = 1.84

Table IV: The percentage of maximum oxygen uptake at anaerobic threshold before and after exercise training with (P) of the individual patients.
$p = probability, (P) = propranolol$

TABLE V

VO₂ Max/Body Wt. (ml/Kg)

<table>
<thead>
<tr>
<th>Name</th>
<th>Before Ex. Training $\bar{\theta}$ (P)</th>
<th>After Ex. Training $\bar{\theta}$ (P)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F.T.</td>
<td>37.0</td>
<td>33.2</td>
</tr>
<tr>
<td>J.R.</td>
<td>22.1</td>
<td>22.0</td>
</tr>
<tr>
<td>E.F.</td>
<td>25.1</td>
<td>26.3</td>
</tr>
<tr>
<td>K.L.</td>
<td>30.3</td>
<td>27.5</td>
</tr>
<tr>
<td>R.Y.</td>
<td>23.3</td>
<td>30.5</td>
</tr>
<tr>
<td>B.W.</td>
<td>15.1</td>
<td>18.4</td>
</tr>
<tr>
<td>E.H.</td>
<td>21.9</td>
<td>25.8</td>
</tr>
<tr>
<td>J.H.</td>
<td>27.1</td>
<td>37.2</td>
</tr>
<tr>
<td>D.T.</td>
<td>20.5</td>
<td>33.0</td>
</tr>
</tbody>
</table>

$\bar{x} = 24.57 \pm 6.34$ = 5.87
SEM $\pm 2.11$ = 1.96

Table V: The maximum oxygen uptake for kg body weight before and after exercise training with propranolol.
$p = probability, (P) = propranolol$

Figure 2 shows the individual and mean increase in VO₂ at anaerobic threshold before and after training with propranolol. Figures 3 and 4 represent the individual and mean data of percentage of VO₂ max and oxygen pulse at anaerobic threshold before and after training with propranolol.
DISCUSSION
The beta adrenergic blocking agent propranolol has been shown to induce effects that might influence the matching of ventilatory and circulatory responses to increased metabolic rate. Propranolol has been shown to reduce heart rate, cardiac output, mean arterial pressure and to increase central venous pressure in normal subjects (Epstein et al, 1965; Furberg and Schmalensee, 1968; Brown et al, 1976). It has also been responsible for lowering muscle lactate and carbon dioxide production during exercise in normals and patients with coronary artery disease as well (Twentyman et al, 1981). In addition, it has been associated with a decreased alactic exercise O2 debts, (Barnard and Foss, 1973) and favours an increased arteriovenous oxygen difference by reducing venous O2 content by increasing peripheral oxygen extraction (Detry et al, 1971).

We found that maximum oxygen uptake in patients on propranolol with coronary artery disease clearly increased after an exercise training programme. Bruce et al, 1979; Hessack et al, 1981; Sklar et al, 1982; Sable et al, 1982 and more recently Petersen et al, 1983 and Wilmore et al, 1983, observed no change, or a slight decrease in VO2 max in normal subjects after acute injection or oral administration of a beta blocker.

On the other hand, Obma et al, 1979; Pratt et al, 1981; Vanhees et al, 1982 and Gordon et al, 1983 found significant increase in VO2 max after training in coronary artery disease during long term beta adrenergic blockade. Our findings (Table II and Fig 1) agree with the data of Gordon et al, 1983 and above groups. Of special interest was the alteration of anaerobic threshold after training. Davis et al, 1979, showed a significant increase in VO2 at an anaerobic threshold (AT) on middle aged men after endurance training of 9 weeks duration with corresponding increase in VO2 max. They demonstrated that anaerobic threshold (AT) was profoundly influenced by endurance training on previously sedentary normal subjects. Our data (Table III, Fig 2) on VO2 at AT in patients with CAD show the similar results.

The percentage of VO2 max at AT in our patients increased from 68.2% to 73.6% after training. Wasserman et al, 1973, found the AT in patients with congestive heart disease (CHD) Class II and Class III was approximtely 30% to 40% of VO2 max. Matsumura et al, 1983; found the mean AT to be between 68% to 71% of VO2 max in patients with CHD Class I, II and III. The reasons for this difference might be that Wasserman et al, 1973, used the cycle ergometer for determination of VO2 max and AT, and other authors used the treadmill. There is a significant difference in VO2 max between different modalities of stress testing. Koyal et al, 1976; Hermansen et al, 1979; Åstrand and Rodahl, 1977, found nearly 20% difference in VO2 max between the cycle ergometer and treadmill. The mean VO2 max/kg body weight (Table V, Fig 4) increased significantly after exercise training.

In these patients the determinations may not be exactly the same VO2 max as in normal trained subjects where the VO2 plateaus at near max exercise isotips of increased work increments near termination of exercise due to fatigue and exhaustion. The VO2 max in patients with CAD may be symptom-limited and therefore, usually do not achieve the same physiologic plateau as in normal highly trained subjects.

Our criteria for noninvasive determination of anaerobic threshold was based upon the nonlinear increase of minute ventilation (VE), carbon dioxide production (VCO2) and ventilatory equivalents for oxygen and carbon dioxide (Wasserman and Mcllroy, 1979; Wasserman et al, 1973; Koyal et al, 1983). In normal subjects, studies with normal subjects, patients with CAD the gas exchange ratio (RQ) is a less reliable noninvasive determinant of anaerobic threshold. It was reported by Twentyman et al, 1981, that propranolol decreased CO2 production at the onset of exercise and early stages of incremental work due to reduced muscle perfusion as a consequence of decreased cardiac output as well as a direct effect on muscle blood supply. Cronin, 1967; Barnard and Foss, 1973, suggested that propranolol also inhibits glycolysis and lipolysis in working muscle and reduces the size of both lactic and alactic O2 debts of exercise. As a consequence of these factors, anaerobic metabolism is increased further as the rate of CO2 removal from the exercising muscle is delayed with lowering of gas exchange ratio during the onset and early stages of incremental exercise. During progressive work at the higher workloads anaerobic metabolism increases and the effects of increasing local metabolite concentrations supercede the effects of the beta blocker on muscle perfusion. The resultant increased blood flow facilitates carbon dioxide removal and, therefore, increases the gas exchange ratio.

Furburg and Schmalensee, 1968 and Reybrouck et al, 1977 demonstrated an increased arteriovenous oxygen difference with beta adrenergic blockade. The mechanism of venous O2 decrease was explained by reduced venous O2 content resulting from increased peripheral O2 extraction during exercise. Patients with CAD using long term beta blockers in addition to an exercise training programme might, we suggest, possibly augment an increased arterial O2 content and may thereby further widen an already increased A-V O2 difference. This would, in turn, induce a higher VO2 max, longer exercise time, and shifting anaerobic threshold to higher workload. Our data on VO2 max in CAD patients compare well with the findings of Obma et al, 1979; Pratt et al, 1981; Vanhees et al, 1982 and Gordon et al, 1983.

The oxygen pulse (O2 P) is one of the important cardiorespiratory variables used for fitness classification. O2 P is defined as the amount of O2 carried to the tissues in the blood by each left ventricular systole. It can be estimated by the ratio of oxygen uptake (VO2) and heart rate (HR). It is also equal to the product of stroke volume and arteriovenous oxygen (A-V O2) difference. We did not measure A-V O2 difference and cardiac output but we did estimate O2 pulse by measuring VO2 and HR. Hanson et al, 1968; Åstrand and Rodahl, 1977; Wasserman et al, 1973 and Ellestad, 1981, postulated that O2 P increased at a work rate below and above AT in middle aged normal subjects from rest. However, Twentyman et al, 1981, believed increased A-V O2 difference is the major contributing factor for increased VO2 max and
exercise tolerance in patients with CAD. Whatever the reason, our patients increased their O₂ at anaerobic threshold after 4 months of a supervised training regime.

In summary, this study reveals an alteration of anaerobic threshold 

which may be responsible for a significant increase in VO₂ max. We suggest, tentatively that alteration of AT may also be due to an increase in stroke volume and perhaps also may be responsible for increased peripheral oxygen extraction.

ACKNOWLEDGEMENTS

The authors wish to express their appreciation to Ophelia Woodfork and Carol Fraser for their excellent secretarial assistance. We are also indebted to Ben Zinser, Public Relations Department, Memorial Medical Center for his assistance in the preparation of this manuscript.

References


