Effect of beta blockade on exercise response after cardiac transplantation

RODNEY S BEXTON,* JOHN R MILNE, RICHARD CORY-PEARCE,†
TERENCE A H ENGLISH, A JOHN CAMM‡

From Department of Cardiology, St Bartholomew's Hospital, West Smithfield, London; and British Heart Foundation Heart Transplant Research Unit, Papworth Hospital, Papworth Everard, Cambridge

SUMMARY Six cardiac transplant recipients underwent maximal exercise testing before and after the administration of intravenous propranolol to assess the effect of beta blockade on their exercise heart rate response and exercise capacity. Before propranolol the patients were capable of a mean of 6-8 minutes of exercise and heart rate increased from a resting value of 102±25 a minute to 138±34 at peak exercise—a mean increase of 35%. All tests were terminated because of tiredness or muscle weakness. After one hour's rest, intravenous propranolol (0-2 mg/kg over 10 minutes) was administered with a reduction in resting heart rate from 109±28 a minute to 83±16. During the repeat exercise test the patients were capable of a mean of 4-5 minutes of exercise and all tests were terminated by extreme exhaustion and/or unsteadiness requiring immediate cessation of the treadmill. Heart rate increased from a resting value of 83±16 a minute to 96±18 at peak exercise.

The exercise capability of the denervated heart is conspicuously reduced by beta blockade, presumably because of its reliance on circulating catecholamines.

Although reinnervation of the transplanted heart has been well documented after orthotopic transplantation of the canine heart,1,2 the transplanted human heart appears to remain both functionally and anatomically denervated indefinitely.3 4 In the absence of the normal cardioacceleratory sympathetic stimulation of exercise, the exercise heart rate response of the denervated heart is gradual and delayed both in onset and offset.5 6 7 Haemodynamic studies, however, both in animals5 8 9 and in man1 6 7 have indicated that denervation neither reduces the capacity for exercise nor the associated increase in cardiac output. In the denervated heart the increase in cardiac output is initially mediated by an increase in stroke volume3 6 8 through the Frank-Starling mechanism,10 11 and only later, with severe exercise, does an increase in heart rate and contractility contribute, associated with increases in the levels of circulating catecholamines.6 12

Studies on the effect of beta blockade on the exercise capacity in intact man have produced conflicting results13 14 15 16 17 18 though in the majority of studies maximal work capacity was not significantly reduced. Theoretically beta blockade should significantly reduce the maximal exercise capacity of the denervated heart, with its reliance on circulating catecholamines. Though limited studies have been performed in denervated19 20 and isolated19 canine hearts, no study on the effect of beta blockade on the exercise response of human cardiac transplant recipients has been reported.

Patients and methods

Six cardiac transplant recipients underwent investigation seven to 18 (mean 13) months after transplantation. Their ages ranged from 28 to 53 (mean 40) years and five were men and one was a woman (Table 1). All patients were functionally well with no haematological, biochemical, or electrocardiographic evidence of rejection. All patients were taking prednisolone and azathioprine as routine immunosuppressive therapy. No patient was taking cardioactive drugs. Written informed consent was obtained from each patient before beginning the study.

*Recipient of British Heart Foundation Fellowship.
†British Heart Foundation Senior Research Fellow.
‡Welcham Senior Lecturer.

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Table 1  Individual patient data

<table>
<thead>
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<th>Case No.</th>
<th>Age (y)</th>
<th>Sex</th>
<th>Pre-transplant diagnosis</th>
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<td>M</td>
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</table>

CM, cardiomyopathy; CAD, coronary artery disease.

Before starting the exercise tests a standard 12 lead electrocardiogram was recorded to ascertain the most suitable six leads for recording during exercise. After orientation to the equipment, the patients underwent maximal graded exercise testing using a Quinton treadmill* and a standard Bruce protocol. At one minute intervals six leads of electrocardiogram were recorded at a paper speed of 50 mm/s. The blood pressure was also measured at three minute intervals. The patients were exercised until forced to stop by a subjective endpoint such as fatigue, exhaustion, dyspnoea, or muscle weakness. Electrocardiographic recordings were continued at one minute intervals after the cessation of exercise until the heart rate had returned to within 5% of the pre-exercise resting rate.

After approximately a one hour rest, intravenous propranolol (0-2 mg/kg) was administered over 10 minutes with electrocardiogram and blood pressure recordings at one minute intervals. The repeat exercise test was started five minutes after completion of the injection. The patients were again maximally exercised with electrocardiographic and blood pressure recordings as previously.

STATISTICAL ANALYSIS
Results are presented as the mean value ± one standard deviation from the mean, and the Student’s two-tailed t test for paired data was used to determine probable differences, which were considered significant when p<0.05.

Results

Before the initial exercise test the resting heart rates ranged from 62 to 138 (102±25) bpm and the resting blood pressure was 134±27/88±16 mmHg. The patients were capable of from five to eight (6.8±1.2) minutes of exercise and all tests were terminated by tiredness or muscle weakness. Increases in heart rate from resting to peak exercise ranged from 16 to 57 (36±14) bpm with peak exercise heart rates of 88 to 178 (138±34) bpm (Fig. 1). Expressed as a percentage of the resting heart rate the increases varied from 17% to 56%, with a mean of 35%. After three minutes of exercise the heart rate had increased by 16% with a further 19% increase over the remaining minutes of exercise (Fig. 2). Both systolic and diastolic blood pressure rose slightly during exercise, though not significantly, to 148±29/90±17 mmHg. The time required to regain control heart rate varied from 12 to 22 (mean 18) minutes.

The administration of propranolol resulted in a reduction in resting heart rate from 109±28 bpm to 83±16 bpm (p<0.01) and in blood pressure from 125±30/78±12 mmHg to 105±15/75±15 mmHg (not significant).

During the repeat exercise test the patients were capable of from 2.75 to 6 minutes of exercise (4.5±1.1 minutes; p<0.01 compared with control exercise test). On this occasion, however, the endpoint of the test was dramatically sudden with all but one of the patients developing pronounced unsteadiness which required immediate cessation of the treadmill to prevent them falling off. None of these patients would have been capable of further exercise. Recovery, however, took place quickly after the termination of exercise. Increases in heart rate ranged from 5 to 25 bpm (13±9 bpm; p<0.01 compared with control exercise

Table 2  Individual patient results

<table>
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<tr>
<th>Case No.</th>
<th>Resting HR (bpm)</th>
<th>Peak exercise HR (bpm)</th>
<th>Increase in HR (bpm)</th>
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<th>Exercise duration (min)</th>
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<td>35 16</td>
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</table>

HR, heart rate; bpm, beats/min; min, minutes; C, control; Prop, propranolol; SD, standard deviation; p, p value (Student’s two-tailed t test for paired data), propranolol vs control.

test), with peak exercise heart rates of 62 to 109 (96±18) bpm (Fig. 1). The percentage increase in heart rate ranged from 5 to 32% with a mean increase of 16% (p<0-01 compared with control exercise test). The heart rate had increased by 12% after three minutes of exercise with little or no increase after this time (Fig. 2). There was again no significant change in blood pressure during the period of exercise.

Table 2 summarises the individual and the overall results and Fig. 3 illustrates typical electrocardiographic recordings during an exercise test before and after propranolol.

Discussion

The physiology of exercise of the denervated heart has been extensively studied both in dogs5 8 9 19 20 and in humans.3 6 7 12 22 Though the heart rate response of the denervated heart to graduated exercise is delayed,

Fig. 2  Percentage increase in heart rate (expressed as a percentage of the resting heart rate) for each minute of exercise before and after the administration of propranolol.

Fig. 3  Typical electrocardiographic recordings during an exercise test before and after propranolol. The heart rate for each minute of exercise is indicated. bpm, beats per minute.
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gradual, and reduced there is no impairment of either exercise capacity or augmentation of cardiac output and oxygen consumption. Early in exercise the pattern of response of the denervated heart is, however, almost the complete opposite to that of the innervated heart. The innervated heart responds to exercise with an almost immediate increase in heart rate with little or no increase in stroke volume, the Frank-Starling mechanism only being important with severe exertion. In the denervated heart in early, and with mild, exercise there is little change in heart rate. The increase in cardiac output is almost entirely mediated through an increase in stroke volume as a result of increased venous return, augmented preload, and the Frank-Starling mechanism.

Haemodynamic studies have shown a pronounced increase in left ventricular end-diastolic pressure and volume associated with this increase in stroke volume in human transplant recipients. Later, and with more severe exercise, increase in heart rate tends to be more pronounced and, together with an increase in contractility, plays an important role in augmenting cardiac output.

The time course of the exercise response of the transplanted heart to severe exercise suggests a humoral mechanism. Normal subjects have been shown to increase plasma catecholamine levels during exercise with the most pronounced increases occurring during maximal exercise. Similarly, Pope et al. have shown a dramatic increase in plasma noradrenaline levels in transplant recipients with the onset of severe exercise which correlated very closely with the increase in heart rate and velocity of circumferential fibre shortening.

Although circulating catecholamines appear to account for the increase in heart rate and contractility occurring with severe exercise in transplant patients, various theories have been proposed to account for the slight increase in heart rate occurring in early, and with mild, exercise. Studies in denervated dogs have shown that this increase is not related to an increase in intravascular temperature or right atrial transmural pressure, and is not affected by bilateral adrenalectomy, and is not abolished by beta blockade. It appears to be an inherent property of the myocardium whereby an acceleration in heart rate occurs proportional to the work performed. The mechanism of this acceleration is still, however, unknown.

The results of this study have confirmed in human transplant recipients the results of previous animal studies. In two studies, one under laboratory conditions and the other using trained, highly motivated racing greyhounds, Donald et al. showed that neither denervation nor the administration of propranolol to normal dogs caused any significant reduction in the capacity for exercise. The administration of propranolol after denervation, however, resulted in the animals either failing to complete the previously attained level of exercise in 85% of tests, with the dogs collapsing on the moving treadmill, or finishing races in a totally exhausted condition at walking pace.

It is apparent from Fig. 1 and 2 that though propranolol has attenuated the heart rate response of the patients early in exercise, its effect is relatively minor. As the severity of exercise is increased after three minutes, the effect of the beta blockade is more apparent, completely abolishing the more pronounced increase in heart rate that occurs after this point during the control tests. This more pronounced increase in heart rate after approximately three minutes of the control test confirms previous reports and is consistent with the onset of action of endogenously released catecholamines. The ability of the denervated heart to perform anything more than mild exercise appears to be critically dependent on this release of catecholamines into the blood stream to augment cardiac output through its effect on heart rate and contractility.

Although catecholamines also appear to be of importance in the response of the innervated heart to severe exercise, studies on the effect of beta blockade in such subjects have only reported a reduction in exercise tolerance of a significant magnitude in one instance, despite significant reductions in exercise tachycardia in all studies. The innervated heart is apparently capable of appropriate haemodynamic readjustments to maintain maximal physical working capacity after beta blockade.

This investigation has confirmed earlier animal studies showing a distinct reduction in the exercise capacity of the denervated heart after the administration of a beta blocking agent. Though having little effect during mild exercise, circulating catecholamines appear to be instrumental in enabling cardiac transplant recipients to retain a virtually normal capacity for exercise. Beta blockade reduces exercise performance conspicuously in such patients as evidenced in this study by a reduction in exercise endurance of 34% and an impressive endpoint to the exercise test. Beta blockers should be used with caution for the treatment of hypertension or myocardial ischaemia in patients after cardiac transplantation.

References

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Requests for reprints to Dr R S Bexton, Department of Cardiology, St Bartholomew's Hospital, London EC1A 7BE.
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R S Bexton, J R Milne, R Cory-Pearce, T A English and A J Camm

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