Pharmacological basis for antihypertensive effects of intravenous labetalol

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Labetalol 1.5 mg/kg administered intravenously to normal subjects in the supine position produced an immediate mean fall in systolic (16%) and diastolic (25%) blood pressure with a concomitant increase in heart rate (12%). After graded exercise, intravenous labetalol inhibited increases in heart rate and blood pressure.

Isoprenaline log dose response curves of increase in heart rate and reduction in diastolic pressure after intravenous labetalol shifted to the right in a parallel manner compared with pre-labetalol response curves suggestive of competitive antagonism at beta-adrenoceptor sites. Similarly, phenylephrine dose response curves of increase in systolic pressure before and after intravenous labetalol were suggestive of competitive antagonism at alpha-adrenoceptor sites. The ratio of relative potency alpha: beta adrenoceptor antagonism after intravenous labetalol was approximately 1 : 7, whereas in the same subjects after oral labetalol the ratio was approximately 1 : 3 as previously reported.

Using the inhibition of isoprenaline tachycardia to estimate the potency of the beta-adrenoceptor antagonism of labetalol relative to that of propranolol the potency ratio was 1 : 6. However, using inhibition of Valsalva tachycardia as the index, the estimated ratio was approximately 1 : 3. Estimates of relative potency using inhibition of tilt tachycardia were complicated by the additional effects upon blood pressure after labetalol not seen after propranolol.

Labetalol produced adrenoceptor blockade at both alpha and beta sites in man sufficient to explain its therapeutic antihypertensive effect.

Labetalol is a compound possessing combined alpha and beta adrenoceptor antagonist properties (Farmer et al., 1972). Consequently it has been assessed in clinical practice as an antihypertensive agent. Prichard et al. (1975) and Koch (1976) have shown that labetalol administered intravenously to hypertensive patients exerts an antihypertensive effect with little effect upon either heart rate or cardiac output. Similar results were obtained by Edwards and Raftery (1976) where labetalol was administered orally on a chronic basis in hypertensive patients. Rosei et al. (1975) and Pearson and Havard (1976) administered labetalol intravenously in doses of 1-0 to 2-0 mg/kg to reduce blood pressure in patients with severe and malignant hypertension. A dose of 1.5 mg/kg was particularly effective in reducing blood pressure. This report concerns a study of intravenous labetalol which was used at a dose of 1-5 mg/kg in order to determine the pharmacological properties of the drug which might be contributory to its therapeutic antihypertensive effect. We also report detailed data of a study previously reported in summarised form (Boakes et al., 1971).

Method

Six informed healthy male volunteers who had previously taken part in a similar study involving oral labetalol (Richards et al., 1976) consented to take part. Their ages ranged from 27 to 39 years and weights from 60 to 84 kg. Blood pressure was measured with a London School of Hygiene and Tropical Medicine sphygmomanometer (Rose et al., 1964) and heart rate was taken from a continuously running electrocardiograph. Each subject rested in the supine position for 15 minutes, then heart

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rate and blood pressure were measured at minute intervals for a period of 5 minutes. The subjects then transferred to a bicycle ergometer (Elema) and readings were taken for a further 5 minutes. The subjects exercised at increasing work loads of 50, 100, and 150 Watts for 2 minutes at each level; blood pressures were recorded at minute intervals throughout. The subjects returned to the supine position and rested for a period of 30 minutes during which a left forearm vein was cannulated using a butterfly needle, and heparinised saline was infused throughout in order to maintain patency of the intravenous needle. During the last 5 minutes of this rest period, observations of heart rate and blood pressure were recorded at minute intervals.

Each subject was then infused with phenylephrine and isoprenaline in random order, 3 receiving phenylephrine first; the remainder isoprenaline first. Phenylephrine was infused at increasing doses in order to plot a cumulative log dose response curve for increases in systolic pressure. The starting dose of phenylephrine was 100 μg/minute for 4 minutes followed by 200 μg/minute for 4 minutes, the final dose being 400 μg/minute. Measurements of heart rate and blood pressure were made throughout this period.

When each subject had rested for 30 minutes after the initial infusion, the alternate drug was infused. Isoprenaline was started at 2 μg/minute for 4 minutes followed by 4 μg/minute, with the final dose being 8 μg/minute each for 4-minute periods. As before, heart rate and blood pressure were measured throughout. After a delay of 30 minutes each subject received intravenous labetalol 1-5 mg/kg body weight (average dose 113 mg) injected over 1 minute. Heart rate and blood pressure were recorded throughout the subsequent 15 minutes at minute intervals. Phenylephrine and isoprenaline were then reinfused in random order. Following labetalol on these occasions, the doses of both phenylephrine and isoprenaline were adjusted to produce changes similar to those recorded in the predrug period. After a rest period of 30 minutes after the final infusion, each subject was again exercised at a similar work level as in the predrug period.

More detailed studies were carried out in 3 other subjects. The laboratory procedures were similar to those described above. Each subject received an infusion of isoprenaline followed by intravenous labetalol in doses of 10, 40, and 160 mg, and propranolol in a dose of 10 mg each administered on different occasions. Each dose of drug was given over a period of 5 minutes and was followed 20 minutes later with the repeat infusion of isoprenaline. On separate occasions phenylephrine was infused but here only doses of 40 and 160 mg labetalol were administered. Before the administration of labetalol, each subject had been tilted at 80° for a period of 2 minutes. This was followed by the Valsalva manoeuvre where a column of mercury was maintained at 40 mmHg for 20 seconds. Measurements of blood pressure and heart rate were recorded throughout. These procedures were repeated 6 minutes after administration of each intravenous dose of labetalol and propranolol in each subject.

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**Fig. 1** Effect of intravenous labetalol (1.5 mg/kg) on mean blood pressure and heart rate in 6 normotensive men.
Antihypertensive effects of intravenous labetalol

Results

EFFECTS OF LABETALOL UPON HEART RATE AND BLOOD PRESSURE, SUPINE AND SITTING

Before labetalol administration in 6 subjects in the supine position the mean (±SEM) resting heart rate was 68 (4-3) beats/minute; systolic pressure 108 (6-5) mmHg and diastolic pressure 72 (3-9) mmHg. Within one minute of labetalol (1-5 mg/kg) administration there was a reduction in both systolic and diastolic blood pressure together with a concomitant increase in heart rate. The maximum reduction in systolic pressure was 16 per cent (17 mmHg) and diastolic pressure 25 per cent (18 mmHg) occurring within 3 minutes of the completion of the labetalol injection. The increase in heart rate was maximal 3 minutes after the completion of the drug administration and at maximum was 12 per cent (8 beats/min) above the preinjection value (Fig. 1). Of the 6 subjects, 4 spontaneously mentioned that they experienced a feeling of warmth and/or flushing immediately after the administration of labetalol.

Before labetalol administration and immediately before exercise mean heart rates were 68 (3-3) supine, and 79 (5-2) beats/minute in the sitting position. Blood pressures were 104 (4-2)/71 (2-6) mmHg and 97 (4-6)/67 (4-0) mmHg, respectively. The difference between supine and sitting systolic pressure was significant (P<0-05). One hundred minutes after labetalol the mean heart rates were 76 (4-6) beats/minute supine and 82 (2-9) beats/minute sitting and comparable blood pressures were 108 (2-6)/71 (1-8) mmHg and 89 (4-5)/59 (4-3) mmHg. Again the difference between supine and sitting systolic pressures was significant (P<0-05). In addition systolic and diastolic pressures (sitting) after labetalol were significantly different (P<0-05) from the values before administration of the drug.

EFFECTS OF LABETALOL UPON EXERCISE-INDUCED CHANGES IN HEART RATE AND BLOOD PRESSURE

The mean values for heart rate blood pressure and mean arterial pressure after graded exercise before and after labetalol (Table 1) showed that the drug inhibited the exercise-induced increases in heart rate and systolic blood pressure at each level of exercise (P<0-001) (Fig. 2). In addition, labetalol reduced diastolic pressure at each exercise level so that at the highest exercise level (150 Watts) the mean arterial pressure was 73 (2-4) mmHg compared with 100 (3-4) mmHg before labetalol was administered (P<0-001).

EFFECTS UPON INCREASING DOSES OF INTRAVENOUS ISOUPRENAline

The mean predrug increase in heart rate was 58 (7-4) beats/minute and the reduction in diastolic pressure 36 (4-3) mmHg after the 8 µg/minute dose of the cumulative isoprenaline infusion (Table 2). In 3 subjects, isoprenaline was infused 15 minutes after labetalol (1-5 mg/kg) administration and in the other 3 subjects after 60 minutes. Comparison of the antagonist effect of labetalol in the two groups indicated that there were no significant differences relating to order of administration so that all individual values were used to calculate the mean post-drug values (Table 2). The log dose isoprenaline response curves in respect of increases in heart rate before and after labetalol (Fig. 3) show that there was a parallel shift to the right after labetalol. These curves were plotted using the mean response at each dose level, and from regression lines an estimate of dose ratio was made. After labetalol the dose of isoprenaline had to be increased by a mean of 26-2 times in order to induce similar increases in heart rate. This is defined as the isoprenaline dose ratio.

The log dose isoprenaline response curve for

Table 1  Effects of graded exercise on heart rate and blood pressure before and after intravenous labetalol (n=6)

<table>
<thead>
<tr>
<th>Before drug</th>
<th>After labetalol</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Heart rate (beats/min) ±SE</td>
</tr>
<tr>
<td>Pre-exercise</td>
<td>79 (5-2)</td>
</tr>
<tr>
<td>Exercise 50 Watts</td>
<td>93 (7-0)</td>
</tr>
<tr>
<td>Exercise 100 Watts</td>
<td>112 (5-0)</td>
</tr>
<tr>
<td>Exercise 150 Watts</td>
<td>132 (9-6)</td>
</tr>
</tbody>
</table>

*P<0-05 differences from predrug values; **P<0-001 differences from predrug values.
Fig. 2  Effects of intravenous labetalol (1.5 mg/kg) on exercise induced changes in heart rate and blood pressure (n=6).

reduction in diastolic pressure before and after labetalol (Fig. 4) showed a parallel shift to the right after the drug, and the dose of isoprenaline was increased by a mean of 23.8 times the dose administered before the drug in order to induce similar reductions in diastolic pressure.

More detailed studies in 3 subjects showed that after graded doses of labetalol there were increasing shifts to the right of the log dose isoprenaline response curves (Fig. 5). After labetalol 10 mg the isoprenaline dose ratio was 3.8, after 40 mg 6.0, and after 160 mg 37.4.

The administration of propranolol 10 mg also resulted in a parallel shift of the isoprenaline dose response curve and in this case the isoprenaline dose ratio was 13.2.

EFFECTS OF INCREASING DOSES OF INTRAVENOUS PHENYLEPHRINE
As with isoprenaline, phenylephrine was started 15 minutes after labetalol (1.5 mg/kg) in 3 subjects and 60 minutes after in the remaining 3 subjects.

As before, we examined the data to see if the order of administration and time difference significantly

Table 2  Mean isoprenaline- and phenylephrine-induced changes before and after 1.5 mg/kg labetalol intravenously (n=6)

<table>
<thead>
<tr>
<th>Dose/min</th>
<th>Before labetalol</th>
<th>After labetalol</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Heart rate (beats/min)</td>
<td>Blood pressure (mmHg)</td>
</tr>
<tr>
<td>Pre-isoprenaline</td>
<td>70 (7-6)</td>
<td>103/69 (8.2)/37-7</td>
</tr>
<tr>
<td>Isoprenaline</td>
<td>2 µg</td>
<td>84 (8-4)</td>
</tr>
<tr>
<td></td>
<td>4 µg</td>
<td>110 (8-2)</td>
</tr>
<tr>
<td></td>
<td>8 µg</td>
<td>128 (7-3)</td>
</tr>
<tr>
<td></td>
<td>32 µg</td>
<td>75 (7-7)</td>
</tr>
<tr>
<td></td>
<td>64 µg</td>
<td>95 (2-6)</td>
</tr>
<tr>
<td></td>
<td>128 µg</td>
<td>112 (6-9)</td>
</tr>
<tr>
<td>Pre-phenylephrine</td>
<td>63 (6-8)</td>
<td>106/74 (7-2)/4-0</td>
</tr>
<tr>
<td>Phenylephrine</td>
<td>100 µg</td>
<td>55 (4-1)</td>
</tr>
<tr>
<td></td>
<td>200 µg</td>
<td>47 (2-2)</td>
</tr>
<tr>
<td></td>
<td>400 µg</td>
<td>40 (0-5)</td>
</tr>
<tr>
<td></td>
<td>800 µg</td>
<td>39 (0-8)</td>
</tr>
<tr>
<td></td>
<td>1600 µg</td>
<td>39 (0-8)</td>
</tr>
</tbody>
</table>
Antihypertensive effects of intravenous labetalol

Before 0o 2 4 8 12 32 64 128 ug Isoprenaline sulphate iv

Fig. 3 Mean isoprenaline-induced increases in heart rate before (○) and after (●) intravenous labetalol (1·5 mg/kg) (n = 6).

After -/U 0-0 20 40 60 80 100 120 ug Isoprenaline sulphate iv

Influenced the responses. This was not so and, therefore, all values were pooled. Before labetalol the mean increase in systolic pressure after the cumulative dose of phenylephrine was 69 (6·8) mmHg (Table 2), whereas to induce similar maximum increases in systolic pressure after labetalol (1·5 mg/kg) the dose of phenylephrine had to be quadrupled. The log dose phenylephrine response curves of increasing systolic pressure before and after labetalol (Fig. 6) showed a parallel shift to the right after the drug. Again from the regression lines it was calculated that after labetalol the dose of phenylephrine to produce similar increases in systolic pressure had to be increased by a mean of 3·8 and this was defined as the phenylephrine dose ratio.

In the 3 other subjects there were also parallel shifts of the post-labetalol log dose phenylephrine response curves (Fig. 7). After labetalol 40 mg, the phenylephrine dose ratio was 2·4 and after 160 mg it increased to 3·8.

In order to estimate the relation between the
alpha adrenoceptor blocking component with that of its beta adrenoceptor blocking component, the ratio of alpha:beta antagonist effects was calculated from:  
\[
\frac{\text{isoprenaline dose ratio}}{\text{phenylephrine dose ratio}}
\]

In the series of six subjects the values were 26.2:3.8; thus the mean ratio of alpha:beta antagonist effect was 1:6.9. Thus the alpha blocking component is approximately one-seventh less potent than the beta component under the circumstances. In the series of three subjects the ratios after 40 mg labetalol were 1:2.5 and after 160 mg 1:9.8.

EFFECTS OF LABETALOL AND PROPRANOLOL UPON TILT AND VALSALVA TACHYCARDIA

Before drug administration, tilting was associated with little change in mean arterial pressure after either one or two minutes (Table 3) but resulted in a mean increase in heart rate of 14 to 21 beats/minute after 1 minute and 18 to 20 beats/minute after 2 minutes of tilt. The effects upon blood pressure after labetalol were dose related; the high dose inducing mean reductions in systolic pressure of 8.8 and 10.2 mmHg at 1 and 2 minutes, respectively, compared with predrug values. Propranolol produced only small reductions in systolic pressure of 2.4 and 2.3 mmHg, respectively. The administration of both labetalol and propranolol was associated with an inhibition of tilt tachycardia. Propranolol 10 mg produced approximately a 50 per cent reduction compared with the predrug values whereas labetalol produced smaller reductions at all doses. In contrast, labetalol inhibited the tachycardia induced by the Valsalva manoeuvre to a greater extent and this was dose related. The highest dose produced 90 per cent inhibition compared with predrug values while propranolol produced a 57 per cent reduction.

Discussion

Rosei et al. (1975), Prichard et al. (1975), Koch (1976), and Pearson and Havard (1976) have shown that intravenously administered labetalol in doses of 0.5–2 mg/kg reduced blood pressure in hypertensive patients. The optimum doses to obtain abrupt reductions in blood pressure in the treatment of hypertensive crises appears to lie between 1.0 and 2 mg/kg. In our own study we showed that intravenous labetalol 1.5 mg/kg produced an immediate reduction in blood pressure in normotensive subjects. The accompanying small increase in heart rate was presumably related to parasympathetic withdrawal, consequent upon the reduction in blood pressure. This relatively small increase in heart rate after labetalol contrasts with the much larger increases in heart rate usually seen after diazoxide administration (Prichard et al., 1975) and this is presumably the result of the additional beta adrenoceptor inhibitory effect of labetalol. After graded exercise 100 minutes after intravenous labetalol administration the drug significantly inhibited the increase in heart rate and blood pressure.
Table 3  Effects upon mean arterial pressure and mean heart rate induced by tilting and the Valsalva manoeuvre before and after propranolol and labetalol (n=3)

<table>
<thead>
<tr>
<th></th>
<th>Mean arterial pressure (mmHg) (± SE)</th>
<th>Change in heart rate after tilting (beats/min) (± SE)</th>
<th>Change in heart rate after Valsalva (beats/ min) (± SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Before drug</td>
<td>After drug</td>
<td>Before drug</td>
</tr>
<tr>
<td></td>
<td>Tilt 1 min</td>
<td>2 min</td>
<td>Tilt 1 min</td>
</tr>
<tr>
<td>Propranolol</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>80 mg</td>
<td>80 ± 1</td>
<td>81 ± 4</td>
<td>65 ± 14</td>
</tr>
<tr>
<td>160 mg</td>
<td>83 ± 5</td>
<td>83 ± 1</td>
<td>64 ± 15</td>
</tr>
<tr>
<td>10 mg</td>
<td>(3-2) (3-4)</td>
<td>(4-0) (3-7)</td>
<td>(3-4) (3-7)</td>
</tr>
<tr>
<td>40 mg</td>
<td>(3-1) (3-7)</td>
<td>(3-4) (3-6)</td>
<td>(3-2) (2-9)</td>
</tr>
<tr>
<td>160 mg</td>
<td>(3-0) (3-7)</td>
<td>(3-4) (2-9)</td>
<td>(3-6) (2-9)</td>
</tr>
</tbody>
</table>

normally accompanying exercise. There appeared to be a greater effect upon blood pressure than upon heart rate; these observations, coupled with the other immediate cardiovascular effects observed, are consistent with those resulting from combined alpha and beta adrenoceptor antagonism.

In our studies we infused isoprenaline in order to produce exogenous cardiac (β₁) and peripheral (β₂) adrenoceptor stimulation. The substantial parallel shifts of the isoprenaline log dose-response curves indicate that labetalol is a non-selective competitive beta adrenoceptor blocking drug. Using phenylephrine as an exogenous alpha adrenoceptor stimulant, we showed that after labetalol there were parallel shifts of the log dose-response curves indicative of competitive alpha adrenoceptor antagonism.

In order to construct log-dose response curves of phenylephrine-induced pressor responses in the subjects in our study, it was necessary to use doses up to 400 μg/minute in order to obtain values that could be plotted on the straight part of the sigmoid dose response curve. Lower doses than those used provide points on the lower plateau of the sigmoid curve and a plateau of this type is an unreliable index of agonist activity. In order to induce similar increments in blood pressure after labetalol it was necessary to increase the dose up to 1600 μg/minute which leads to a somewhat greater pressor response than in the prelabetalol phase. This response was not greater than the systolic pressure increase induced by exercise in these subjects on other occasions and thus phenylephrine-induced increases were within levels seen during normal physiological stresses.

Since six subjects in this study had previously been studied under very similar conditions before and after oral labetalol (Richards et al., 1976), we have compared these data with the results in the present study. After 400 mg oral labetalol the isoprenaline dose ratio was 9-63, whereas after intravenous labetalol (average dose 113 mg), this increased to 26-2, suggesting that by this route the drug is approximately three times more potent. However, after oral and intravenous labetalol the phenylephrine dose ratios were 3-36 and 3-80, respectively, suggesting little difference in potency. Estimates of relative potency alpha:beta based upon these data were approximately 1:3 after oral labetalol and approximately 1:7 after intravenous labetalol.

As the dose of labetalol was increased the isoprenaline dose response curves were also shifted progressively to the right. This observation is consistent with previous studies of beta adrenoceptor antagonist drugs where isoprenaline has been used as the alpha agonist (Conway et al., 1976; Aellig, 1976).

The post labetalol phenylephrine dose response curves were also shifted progressively to the right as the dose of labetalol was increased but the shifts in these cases were much less pronounced. Thus, the higher the dose of labetalol the greater became the separation between the alpha:beta antagonist effects. From these it might be assumed that the manifestation of high doses of labetalol in man would be predominantly those known to be associated with beta adrenoceptor antagonists. However, the immediate effect of 1-5 mg/kg labetalol intravenously was a pronounced reduction in blood pressure and a concomitant small increase in heart rate. This is quite unlike the response seen after the acute administration of a beta adrenoceptor blocking drug such as propranolol (Prichard et al., 1975). A dose of 1-5 mg/kg is associated with a degree of alpha blockade that is clinically significant.

Other cardiovascular effects consistent with an alpha blocking effect are the immediate reduction in
blood pressure, the dose-related reduction in mean arterial pressure during tilting and the relatively modest inhibition of tilt tachycardia which appeared to be influenced by the alteration of blood pressure.

Comparing all these data it is possible to assess the relative potency of propranolol and labetalol. The inhibition of tilt tachycardia by labetalol was modified by the drug's effect upon mean arterial pressure, leading to the possibly erroneous conclusion that propranolol was at least 16 times more active. On the other hand, the inhibition of Valsalva tachycardia, probably a fairly good indicator of myocardial beta adrenoceptor blockade, suggests that propranolol was approximately three times more active. This approximation was more consistent with the estimate of potency ratio obtained from studies of isoprenaline-induced tachycardia where the ratio was approximately 6:1.

We conclude that labetalol administered intravenously possesses combined alpha and beta adrenoceptor antagonist properties, confirming our previously reported studies with oral labetalol, and that these properties are sufficient to explain the therapeutic effect of intravenous labetalol in hypertension.

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