The cardiovascular effects of beta adrenergic agonist drugs administered by nebulisation

A. Flatt, J. Crane, G. Purdie, T. Kwong, R. Beasley and C. Burgess

Departments of Medicine and Community Health, Wellington School of Medicine, Wellington, New Zealand.

Summary: The cardiovascular effects of equal doses (5 mg) of nebulised fenoterol, salbutamol and terbutaline were compared in 12 healthy individuals in a double-blind, placebo-controlled study. Measurements of heart rate, blood pressure, systolic time intervals, QTc interval and T-wave amplitude were made at baseline and at 15, 30, 45, 60 and 90 minutes after nebulisation. Fenoterol caused significantly greater chronotropic electrocardiographic and inotropic effects than either salbutamol or terbutaline. The peak effects after terbutaline occurred later than those after fenoterol or salbutamol.

Introduction

The inappropriate use of nebulised beta-2-adrenergic agonist drugs has been proposed as a possible factor contributing to asthma mortality in New Zealand.1 This hypothesis was supported by the finding of a subsequent study which investigated the management of patients dying from asthma in New Zealand.2,3 In those patients who had a home nebuliser and its use during the final attack was known, over one-third had used it inappropriately.2 Indeed, over one-quarter of all the patients who died from asthma during the two year period of this study had a home nebuliser available for self-administration of beta-adrenergic agonist drugs. This is in marked contrast with the situation in England, where similar asthma mortality studies have demonstrated that death from asthma outside hospital is not associated with home nebuliser use.4,5

Analysis of the specific beta-2 adrenergic agonist self-administered by the patients who died, demonstrated that there was a significantly greater proportion of fenoterol than salbutamol used when compared with national sales of these drugs, suggesting that these agents may differ in their extra-pulmonary effects or toxicity. We have previously shown that fenoterol (administered by metered dose inhaler) has significantly greater electrocardiographic, inotropic and chronotropic effects than equal doses of either salbutamol or isoprenaline.6 In this present study we have assessed the cardiovascular effects of equal doses of fenoterol, salbutamol and terbutaline as these are the beta-adrenergic agents most commonly administered by nebulisation in New Zealand.

Subjects and methods

Twelve healthy subjects (eight male), mean age 30 years (range 21–41 years) were studied on four occasions, at least 2 days apart, after a 4-hour fast. None of the subjects smoked and all had abstained from alcohol and caffeine-containing substances for 12 hours prior to each study. After resting supine for 10 minutes, control recordings of the systolic time intervals, high speed electrocardiogram (ECG) and blood pressure were made. The subjects then inhaled 5 mg (1 ml) of either salbutamol, fenoterol, terbutaline or an equal volume of saline via a Pari inhalerboy nebuliser. The treatments were administered according to a cross-over, randomized, double-blind, Latin square design. Further recordings were made 15, 30, 45, 60 and 90 minutes following completion of nebulisation.

The systolic time intervals were measured from simultaneous high speed (100 mm/s) photographic recordings of the ECG, phonocardiogram and carotid pulse trace. From these recordings, total electromechanical systole (QS), left ventricular ejection time (LVET) and pre-ejection period (PEP = QS₂ – LVET) were made as previously described.7 QS was corrected for heart rate using an equation developed in our laboratory (QS₁ = QS₂ + 1.2 × HR; QS₁ is the QS index corrected for heart rate). From the ECG the QT interval and T-wave amplitude were measured. The end of the QT interval was measured by extrapolating the slope of the T-wave to the baseline (P-P interval). This interval (QTc) was corrected for the heart rate.
using the Bazett formula. The percentage change in T-wave amplitude from the control measurements \((\Delta T\%)\) was noted on each day. Blood pressure was measured automatically using a Vita Stat recorder.

All subjects gave written informed consent and the study was approved by the Wellington Hospital Research Ethical Committee.

**Statistical analysis**

A three-way analysis of variance with subject, treatment and time effects was performed on the change from control data. The significance level was adjusted for multiple comparisons by using Bonferroni’s inequality \((P < 0.05 \div \text{number of comparisons})\). Hence, when a significant treatment-time interaction was present, a probability value of \(P < 0.0017\) was considered significant when comparisons between treatments were made at each time.

**Results**

There were no significant differences in the control measurements between the treatment days (Table I).

When compared to placebo (see Figure 1 and Table II), all the active agents increased heart rate, decreased electromechanical systole \((QS_2I)\), pre-

<table>
<thead>
<tr>
<th>Table I</th>
<th>Mean (s.e.m.) control values in the twelve subjects prior to treatment.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Placebo</td>
</tr>
<tr>
<td>Heart rate (beats/min)</td>
<td>63.9 (2.2)</td>
</tr>
<tr>
<td>QS_2I (ms)</td>
<td>489.5 (7.5)</td>
</tr>
<tr>
<td>PEP (ms)</td>
<td>111.4 (5.3)</td>
</tr>
<tr>
<td>QTc (ms)</td>
<td>382.2 (9.0)</td>
</tr>
<tr>
<td>Systolic blood pressure (mmHg)</td>
<td>115.7 (2.1)</td>
</tr>
<tr>
<td>Diastolic blood pressure (mmHg)</td>
<td>77.5 (2.8)</td>
</tr>
</tbody>
</table>

Data analysis with s.e.m. values.

\(QS_2I\) - Total electromechanical systole corrected for heart rate.

PEP - Pre-ejection period.

Figure 1  Effect of the treatments on heart rate (HR), total electromechanical systole \((QS_2I)\), QTc interval and systolic blood pressure \((SBP)\). □, Placebo; ■, fenoterol; ○, salbutamol; ●, terbutaline.
Table II  Effect of placebo, fenoterol, salbutamol and terbutaline on pre-ejection period (PEP), T wave height and diastolic blood pressure (mean ± s.e.m.).

<table>
<thead>
<tr>
<th>Time (mins)</th>
<th>P</th>
<th>ΔPEP (ms)</th>
<th>T</th>
<th>P</th>
<th>ΔT%</th>
<th>T</th>
<th>P</th>
<th>ΔDBP (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>3.5</td>
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<tr>
<td></td>
<td>± 2.5</td>
<td>±4.8</td>
<td>±3.8</td>
<td>±4.5</td>
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<td>30</td>
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<td>-5.2†</td>
<td>-32.7†</td>
<td>-28.2†</td>
<td>-6.1</td>
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<tr>
<td></td>
<td>± 2.1</td>
<td>±3.9</td>
<td>±3.1</td>
<td>±3.6</td>
<td>±3.7</td>
<td>±5.4</td>
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<tr>
<td>45</td>
<td>1.8</td>
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<td>-21.2†</td>
<td>-6.3</td>
<td>-35.4†</td>
<td>-29.7†</td>
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<tr>
<td></td>
<td>± 2.8</td>
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<td>60</td>
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<td>-22.1†</td>
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<td>-2.6</td>
</tr>
<tr>
<td></td>
<td>± 2.5</td>
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<td>±3.4</td>
<td>±3.8</td>
<td>±3.2</td>
<td>±5.3</td>
</tr>
</tbody>
</table>

P = placebo, F = fenoterol, S = salbutamol, T = terbutaline. †P < 0.0001

ejection period and T wave amplitude at all times (P < 0.0001). Fenoterol and terbutaline increased the QTc interval at all times (P < 0.0001), whereas salbutamol increased this interval from 15 to 60 minutes (P < 0.005 to P < 0.0001). Fenoterol increased systolic blood pressure from 15 to 60 minutes (P < 0.0001), whereas salbutamol increased this parameter at 15 minutes only (P < 0.0001). None of the drugs differed in their effect on diastolic blood pressure from placebo.

Comparison of active agents

(i) Systolic time intervals  Fenoterol caused significantly greater decreases in Q50i than either salbutamol (15–90 minutes) or terbutaline (15 and 30 minutes); it also significantly decreased pre-ejection period at 45–90 minutes when compared to salbutamol or terbutaline (15 minutes). Terbutaline significantly decreased Q50i and pre-ejection period when compared to salbutamol (60 and 90 minutes). Salbutamol decreased pre-ejection period at 15 minutes when compared to terbutaline.

(ii) ECG  Fenoterol significantly increased heart rate when compared to salbutamol (30, 45 and 90 minutes) and terbutaline (15 and 30 minutes) and increased the QTc interval when compared to salbutamol (30–90 minutes) or terbutaline (30 minutes). Terbutaline increased the QTc interval and decreased T wave amplitude at 90 minutes when compared to salbutamol.

(iii) Blood pressure  Fenoterol caused a significantly greater rise in systolic pressure than either salbutamol (30 and 45 minutes) or terbutaline (15 to 45 minutes). Terbutaline caused a significantly greater decrease in diastolic blood pressure than salbutamol at 60 minutes.

Discussion

In this study we have shown that the administration of equal doses of nebulised beta-2 adrenergic agonists results in different pharmacodynamic effects. In particular, nebulised fenoterol produced significantly greater inotropic, chronotropic and electrophysiological effects than equivalent doses of either salbutamol or terbutaline. The predominant difference between salbutamol and terbutaline related to a more gradual onset and a longer duration of action of the latter agent resulting in greater effects 90 minutes after nebulisation.

The marked increase in systolic blood pressure, heart rate and reduction in Q50i following administration of fenoterol indicates direct stimulation of cardiac beta-1 and beta-2 receptors.112 Fenoterol also caused a significant increase in the QT interval and decrease in T-wave height. These electrocardiographic changes are of interest, for they may be associated with increased ventricular excitability. The increased risk of ventricular arrhythmias associated with QTc interval prolongation may be more common in the situation of an increase in heart rate, as we observed with fenoterol. When anti-arrhythmic agents, such as amiodarone, cause prolongation of the QTc interval, this is invariably associated with bradycardia.14 The changes in T-wave height may be consequent to the prolongation of the QT interval, or may be associated with hypokalaemia which has been demonstrated with beta-2 agonists.1516

In contrast to fenoterol, the cardiovascular effects of salbutamol and terbutaline are less marked. This confirms our previous observation, that repeated inhalation of fenoterol by metered dose inhaler results in greater inotropic, chronotropic and electrophysiological effects than equal doses of salbutamol.1 These differences are primarily due to greater beta-1 adrenergic receptor stimula-
tion following fenoterol, and are consistent with previous in vitro and in vivo studies. Although the cardiovascular effects of salbutamol and terbutaline are similar in magnitude, there were significant differences in their duration of action. The effects due to terbutaline were of slower onset and of greater duration than those following salbutamol. As a result, at the end of the 90 minute course the cardiovascular effects of terbutaline were greater than those after salbutamol. We have observed similar differences in the time courses of the hypokalaemic effects of these agents when administered to some of these subjects. This difference is likely to be due to differences in bioavailability which in turn is likely to be due to differences in the preparation of these agents.

These results are of importance as they point to major differences in the clinical extra-pulmonary effects of these agents. Fenoterol is more likely to cause increased myocardial oxygen consumption, and perhaps in the setting of hypoxia may be more likely to predispose an asthmatic patient to ventricular arrhythmias than either salbutamol or terbutaline. Indeed Tandon demonstrated that equal doses of fenoterol were associated with a greater frequency of ventricular ectopic beats than salbutamol and we have recently shown that the relative risk of asthma death is increased in asthmatics prescribed fenoterol either by metered dose inhaler or nebuliser when compared to other bronchodilator therapy, thus stressing the clinical relevance of this present study. Our results raise the possibility that in asthmatic patients with home nebulisers the disproportionate use of nebulised fenoterol among those who die may be due to the direct toxic effect of this agent, rather than different prescribing habits as previously postulated.

Acknowledgement

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References

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