Serevent nationwide surveillance study: comparison of salbutamol with salmeterol in asthmatic patients who require regular bronchodilator treatment

Win Castle, Rick Fuller, John Hall, James Palmer

Abstract

Objective—To compare safety of salmeterol and salbutamol in treating asthma.

Design—Double blind, randomised clinical trial in parallel groups over 16 weeks.

Setting—General practices throughout the United Kingdom.

Subjects—25180 patients with asthma considered to require regular treatment with bronchodilators who were recruited by their general practitioner (n=3516).

Interventions—Salmeterol (Serevent) (50 μg twice daily) or salbutamol (200 μg four times a day) randomised in the ratio of two patients taking salmeterol to one taking salbutamol. All other drugs including prophylaxis against asthma were continued throughout the study.

Main outcome measures—All serious events and reasons for withdrawals (medical and non-medical) whether or not they were considered to be related to the drugs.

Results—Fewer medical withdrawals due to asthma occurred in patients taking salmeterol than in those taking salbutamol (2.91% vs 3.79%; χ²=13.8, p=0.0002). Mortality and admissions to hospital were as expected. There was a small but non-significant excess mortality in the group taking salmeterol and a significant excess of asthma events including deaths in patients with severe asthma on entry. Use of more than two canisters of bronchodilator a month was particularly associated with the occurrence of an adverse asthma event.

Conclusions—Treatment over 16 weeks with either salmeterol or salbutamol was not associated with an incidence of deaths related to asthma in excess of that predicted. Overall control of asthma was better in patients allocated salmeterol. Serious adverse events occurred in patients most at risk on entry and were probably due to the disease rather than treatment.

Introduction

Inhaled β agonists are accepted as the most effective bronchodilators in current clinical use, although recent evidence in treating asthma has been questioned. The questions do not relate to their use in relieving acute bronchospasm but rather to their use as maintenance treatment. One study suggested that the control of asthma may be worse during treatment with regular fenoterol compared with intermittent β agonists. More recently the safety of β agonists has come under scrutiny. Firstly, a rebound increase in bronchial reactivity has been observed in research that studied regular use of β agonists, although this has not been a consistent finding. Secondly, a reduction in lung function was reported during a two year study of salbutamol and ipratropium bromide, although again this has not been found in other studies.

In addition, epidemiological studies have shown an increase in deaths due to asthma, which some authors have related to the use of β agonists, and a more detailed case control study has related increased use of β agonists to an increased risk of death from asthma. For each additional canister administered each month the odds ratio increased by 2.6. Causal relations have been suggested but the high use of β agonists probably merely reflects severity of asthma and that these patients with more severe asthma are at greater risk of death. Such a relation between severe asthma and the risk of death has indeed been shown in other epidemiological studies.

The development of a long acting β agonist for which regular use is recommended has raised some concerns. Our study compared the safety of salmeterol (50 μg Serevent twice daily) with that of salbutamol (200 μg four times a day) in a large number of patients with asthma (> 25 000). Because of the large numbers we could compare our results with events related to asthma throughout the United Kingdom.

Subjects and methods

PROTOCOL

The trial was a randomised, double blind study in parallel groups over 16 weeks. Randomisation from blocks of six was four patients allocated to salmeterol (50 μg, two puffs morning and bedtime and two puffs of placebo noon and early evening to retain the blinding) to two patients allocated to salbutamol (200 μg, two puffs four times a day). The general practitioners were all given prenumbered treatment packs which they gave to the patients as they were allocated the next consecutive study number. All other drugs including prophylaxis against asthma were continued throughout the study. The supervising general practitioner was asked to prescribe whatever treatment he or she considered appropriate to relieve symptoms. The trial was conducted by 3516 general practitioners and monitored by staff employed by the medical department of Allen and Hanburys.

PATIENTS

At the first visit of the study patients with asthma were recruited by their general practitioner and randomised. After they had shown effective use of an
inhaler they then started the treatment. There was no run in period. Patients had to be over the age of 12 years and have a clinical requirement for regular bronchodilator treatment. Patients taking β blockers were excluded as were patients with serious uncontrolled pulmonary or systemic disease or who were pregnant. The general practitioner completed a demographic questionnaire with details of each patient's present drugs, medical history, etc. The general practitioner also clinically assessed each patient as having mild, moderate, or severe asthma.

The patient returned after four, eight, and 16 weeks of the study treatment, and the investigator was instructed to check for the occurrence of serious adverse events whether or not they were considered to be related to the drug and to record these as well as reason for withdrawals (medical or non-medical) on a serious adverse event or withdrawal form. The general practitioners were reimbursed £50 for each completed patient studied.

**DATA MANAGEMENT METHODS**

A serious adverse event was defined in the protocol as death from any cause, any event requiring admission to hospital or prolonging a stay in hospital, a life-threatening event, a severely disabling or incapacitating event, a congenital abnormality, cancer, or drug overdose. On completing a serious adverse event or withdrawal form the investigator was required to add his or her assessment of causal relation to the study drug. The options were definitely related, probably related, probably unrelated, or definitely unrelated.

Any patient could be withdrawn from the study at any time at the discretion of the investigator or the request of the patient, but the reason for doing so was required for each case. Whenever a patient was withdrawn from the study the investigator had to confirm that the patient remained alive at the end of the study.

Each serious adverse event or withdrawal form was forwarded promptly to the medical department, where its receipt was documented but the code not broken. The investigator was contacted by telephone or letter or visited if follow up information was required. Documentation from death certificates and post-mortem examinations was obtained. The records were photocopied and sent to the international drug surveillance division of Glaxo Group Research, where the randomisation code was broken and the drug identity was recorded on the form. Access to the code was restricted to the people within the division who required the information to generate interim reports to regulatory authorities.

Individual reports of adverse events which were considered serious under the requirements of the regulatory authorities and also assessed by the investigator as definitely or probably related to the study drug were forwarded immediately to the Medicines Control Agency in the United Kingdom and, after prior agreements, periodically to the Food and Drug Administration in the United States. Interim confidential reports were also regularly prepared for the two organisations, as well as for other regulatory authorities as requested.

For the purpose of providing overview tables, the type of report for each patient was classified as death, serious events, or withdrawals. Each death was categorised as respiratory and related to asthma; chronic obstructive airways disease; respiratory but not related to asthma, including lung cancer; cardiovascular or relating to other body systems. Two respiratory physicians employed by Glaxo reviewed all possible respiratory deaths to give their view about the most appropriate medical category without knowing to which drug the patient had been randomised.

For patients with adverse events falling into more than one category, deaths were given precedence over events meeting the regulatory definitions of serious, which in turn took precedence over withdrawals for minor medical reasons. If follow up information was received about a particular patient which materially altered the nature of the case or the outcome—for example, if a patient with a serious adverse event was later found to have died within the study period—this result on the database was updated to reflect the new information.

Events related to asthma were taken to include all asthmatic events irrespective of causality. Chest infections such as pneumonia, when no symptom of bronchospasm was reported, were categorised as respiratory but not related to asthma. Respiratory adverse events were defined according to ICD (ninth revision).

When patients were described as having an event in which different symptoms or diagnoses fell into different body systems, asthma as a cause was given precedence. Otherwise the event was counted against the classification of the body system of the most important symptom in the opinion of Glaxo physicians.

An independent review of the blinded data for deaths related or possibly related to asthma in treatment groups was undertaken by three consultants selected by the National Asthma Campaign. An independent statistician received a copy of the interim tables forwarded to the regulatory authorities and gave independent statistical advice throughout. The standard (demographic) forms completed at entry and at four, eight, and 16 weeks were contracted out for data management and analysis by VAMP Research Ltd. Comparisons between the treatment groups were made by χ² test. The results are expressed as absolute incidence and relative risk between treatments. No formal power calculation on any specific event was made as the primary objective was to compare the incidence of all safety outcomes. The numbers in table I refer to the number of patients in whom full data were available. The larger numbers in table III refer to the total numbers randomised.

**Results**

There were 25 180 patients randomised to treatment, 16 787 to salmeterol and 8393 to salbutamol. All serious adverse event or withdrawal forms were entered and analysed. Randomised treatment groups according to entry criteria (table I) were well balanced.

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**TABLE I—Criteria on entry of patients with asthma randomised to salmeterol or salbutamol. Values are numbers (percentages)**

<table>
<thead>
<tr>
<th>Entry criteria</th>
<th>Salmeterol (n=14 113)</th>
<th>Salbutamol (n=7082)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 18</td>
<td>899 (6.4)</td>
<td>421 (5.9)</td>
</tr>
<tr>
<td>18-30</td>
<td>2752 (19)</td>
<td>1359 (19)</td>
</tr>
<tr>
<td>31-40</td>
<td>1970 (14)</td>
<td>1095 (16)</td>
</tr>
<tr>
<td>41-50</td>
<td>2064 (15)</td>
<td>1060 (15)</td>
</tr>
<tr>
<td>51-60</td>
<td>2435 (17)</td>
<td>1200 (17)</td>
</tr>
<tr>
<td>&gt; 60</td>
<td>3993 (28)</td>
<td>1947 (28)</td>
</tr>
<tr>
<td>General practitioner's classification of severity of asthma:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mild</td>
<td>2386 (17)</td>
<td>1235 (17)</td>
</tr>
<tr>
<td>Moderate</td>
<td>9261 (66)</td>
<td>4592 (65)</td>
</tr>
<tr>
<td>Severe</td>
<td>2466 (17)</td>
<td>1255 (18)</td>
</tr>
<tr>
<td>Event asthma taken*:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oral corticosteroids</td>
<td>679 (4.8)</td>
<td>332 (4.7)</td>
</tr>
<tr>
<td>Inhaled corticosteroids</td>
<td>6969 (69)</td>
<td>4895 (69)</td>
</tr>
<tr>
<td>Methyldopa</td>
<td>2157 (15)</td>
<td>1055 (15)</td>
</tr>
<tr>
<td>β Agonists (oral)</td>
<td>688 (4.9)</td>
<td>377 (5.3)</td>
</tr>
<tr>
<td>β Agonists (inhaled)</td>
<td>10 820 (91)</td>
<td>6415 (91)</td>
</tr>
<tr>
<td>β Agonists (nebulised)</td>
<td>203 (1.7)</td>
<td>103 (1.7)</td>
</tr>
<tr>
<td>Sodium cromoglicate/nedocromil/ketotifen</td>
<td>618 (4.4)</td>
<td>269 (3.8)</td>
</tr>
<tr>
<td>Anticholinergic drugs</td>
<td>780 (5.5)</td>
<td>370 (5.2)</td>
</tr>
<tr>
<td>Combination drugs</td>
<td>1036 (7.3)</td>
<td>475 (6.7)</td>
</tr>
</tbody>
</table>

*Many patients were taking more than one drug at entry.
Table III summarises all deaths, serious events, and withdrawals, whose incidences were similar between the two groups. The only significant difference was the numbers of medical withdrawals due to asthma, which were fewer with salbutamol than with salbutamol (2.91% vs. 3.79%; \( \chi^2 = 13.6; p = 0.0002 \)).

For salbutamol there were 12 deaths from asthma out of 16,787 patients treated (0.07%) and for salbutamol two out of 8,933 (0.02%). When summarization of binomial probabilities was performed and the 2:1 randomisation accounted for the 95% confidence interval for the relative risk of 3:0 between the two groups was 0.7 to 20. The difference between the two groups (0.07% vs 0.02%) was not significant (\( p = 0.105 \)). Obviously the relative risk of 2:0 and 1:0 for other obstructive airways disease and other non-obstructive airways disease respectively are similarly not significantly different, the probability being 0.506 and 1.0 of obtaining a distribution as extreme or more extreme in either direction by chance.

The 14 deaths from asthma were in patients with severe asthma either in the view of the general practitioner or in the view of the independent consultants appointed by the National Asthma Campaign. Five were taking oral steroids and a further patient was taking nebulated \( \beta \) agonists. Another patient had previous attacks described as catastrophic, but in this case inhaled steroids were stopped because of fear of growth suppression. All but two of the patients who died were taking two or more inhalers of \( \beta \) agonist a month at the time of entry to the study. For 10 of the patients who died the independent consultants considered that their asthma could possibly have been more appropriately treated by earlier or higher doses of glucocorticosteroids. Five of the patients who died had been admitted to hospital.

The overall incidence of serious events from all causes (table III) (4.0% for salmeterol and 4.1% for salbutamol) was unremarkable, the incidence of those suspected as being related to the drugs (1.19% vs 1.15%) was low, and importantly there was no evidence of any previously unrecognised side effect of salmeterol or salbutamol. Not surprisingly in a study of this type there was a fairly high rate of patients withdrawing for non-medical reasons, which was balanced between the two groups (table III).

Table IV shows non-fatal events related to asthma by randomised treatment group and also by the estimated rates of these non-fatal events according to classification of severity of asthma before treatment. For each indicator of severity (general practitioner’s assessment, oral steroids, and increase of \( \beta \) agonist inhalers used in the previous four weeks) there was a consistent significant increase in rates of events related to asthma.

*Estimated rate of non-fatal events related to asthma considering severity of asthma before treatment.
with severity within each randomised treatment group (p < 0.001). The difference in rates of events between salmeterol and salbutamol was assessed by using a Mantel-Haenszel test, stratified by each of the three indicators of severity of asthma at entry in turn. For each, there was a significantly lower rate in patients treated with salmeterol than in those treated with salbutamol as follows: general practitioner’s classification $X^2=9.15$ (p = 0.002), oral steroid use $X^2=9.86$ (p = 0.002), and inhaler use $X^2=8.13$ (p = 0.004).

Discussion

This trial is the largest randomised, double blind clinical trial ever conducted in the United Kingdom. The aim was to recruit about 24,000 patients: 16,000 to salmeterol and 8000 to salbutamol. Recruitment started in late 1990 and finished during early December 1991. Waller et al recently criticised surveillance studies sponsored by pharmaceutical companies on the grounds that they did not include comparator groups, that recruitment was slow (so slow in some that the study had to be abandoned), that the total numbers of patients were low compared with the median of 1480 patients presented in original product licence applications, and finally that companies were slow to provide study results.10 We believe this trial had none of these failings. About 16 times the median number in product licence applications were recruited in 14 months. Glaxo reported monthly to the Medicines Control Agency, and the final safety report was sent to the agency within four weeks after the last recruited patient had completed the 16 week study. Inevitably the amount of detail in such a vast study and its overall quality is of a lower standard than that usually obtained from smaller well controlled preregistration studies conducted to good clinical practice.

Comparison of the reported events shows that the only significant differences were the number of medical withdrawals due to asthma, fewer occurring with salmeterol (2-91% v 3-79%). The relative benefits of treatment can be inferred to some degree from the rate of withdrawal. None of the other events in this trial approached differences which were significant, the study being sized appropriately and of adequate duration to detect a clinically meaningful difference.

There was a numerical excess of deaths in the salmeterol group: 0-07% compared with 0-02%, a relative risk of 3. The numbers were very small, however, and the confidence interval of this risk was wide; the difference was therefore not significant. Is the difference then a chance finding? Obviously we cannot answer this, although a number of points need to be considered. The overall number of deaths (14) is in line with that which would have been expected of a sample of patients with asthma of this size in the United Kingdom—that is, 15.11,12 This comparison would be valid only if the distribution of severe asthma in the study population was at least as great as that in the general population. The patients recruited to the study did not predominantly have mild asthma; 17% were classified by their general practitioners as having severe asthma at entry, 69% were taking inhaler steroids, 4-7% were taking oral steroids, 9-5% had used three or more canisters of $\beta$ agonist inhalers during the four weeks before the study, and, of course, all had a clinical requirement for regular bronchodilator treatment. As a comparison, in 1990 52% of adults with asthma were issued with at least one prescription for an inhaler steroid, 3% were given oral steroids for a full 12 months as maintenance treatment (5-1% at some time during the year), and 3-4% were issued with three or more $\beta$ agonist inhalers each month (AHA Medical UK, unpublished data). Therefore the mortality was not biased because the study recruited patients with mild asthma. In addition, there was no detectable pattern in the deaths related to asthma which occurred in patients deemed by the independent assessors to have severe asthma. Also other events related to asthma showed no trend towards being worse with salmeterol than salbutamol. In fact, the only significant difference seen was that in the proportion of patients treated for asthma.

Previous much smaller studies such as those described by Sears et al and van Schayck et al described an apparent deterioration in asthma, albeit over a longer treatment period. Our data are not consistent with this as the overall rates of death and admission to hospital were not excessive, and patients with mild and moderate asthma were not put at equal risk by the use of the $\beta$ agonist as were those with severe asthma. The problem of deaths from asthma and severe events is therefore one of undertreatment of asthma rather than regular use of $\beta$ agonists. As the independent reviewers suggested that the asthma might have been more appropriately treated by earlier or higher doses of gluocorticosteroids in at least 10 of the patients in the study who died, it is appropriate to strengthen the recommendations of how best to treat severe asthma. Glaxo has therefore altered all its data sheets for drugs used to treat asthma, both $\beta$ agonists and inhaled corticosteroids, to improve the information to doctors on the treatment of the disease, especially when it is severe or unstable. This study clearly indicates such patients are at risk.

We suggest that patients with severe and/or unstable asthma are at potential risk of death by asthma. Use of high doses (two or more canisters a month) of $\beta$ agonists is not appropriate as the main or only treatment and is related to severity of disease. Such patients require stabilisation of their asthma with appropriate doses of inhaled or oral gluocorticosteroids (>1 mg per day of beclomethasone dipropionate or equivalent) as their main treatment.

We thank Sir David Cox FRCS, who received a copy of the interim tables forwarded to the regulatory authorities and gave independent statistical advice throughout, and also the National Asthma Council and the three consultants it selected: Dr Maryn Partridge, Martin Hetzel, and Brian Harrison.


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