Adrenal suppression with inhaled budesonide and fluticasone propionate given by large volume spacer to asthmatic children

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Abstract

Background – The aim of this study was to compare the systemic bioactivity of inhaled budesonide (B) and fluticasone propionate (F), each given by large volume spacer, on a microgram equivalent basis in asthmatic children.

Methods – Ten stable asthmatic children of mean age 11 years and forced expiratory volume in one second (FEV₁) 81.6% predicted, who were receiving treatment with ≤400 µg/day of inhaled corticosteroid, were studied in a placebo controlled single blind (investigator blind) randomised crossover design comparing single doses of inhaled budesonide and fluticasone propionate 400 µg, 800 µg, and 1250 µg. Doses were given at 20:00 hours with mouth rinsing and an overnight 12 hour urine sample was collected for estimation of free cortisol and creatinine excretion.

Results – The results of overnight 12 hour urinary cortisol output (nmol/12 hours) showed suppression with all doses of fluticasone propionate (as geometric means): F400 µg (11.99), F800 µg (6.49), F1250 µg (7.00) compared with placebo (24.43), whereas budesonide caused no suppression at any dose. A comparison of the drugs showed that there were differences at 800 µg and 1250 µg levels for urinary cortisol: B800 µg versus F800 µg (2.65-fold, 95% CI 1.26 to 5.58), B1250 µg versus F1250 µg (2.94-fold, 95% CI 1.67 to 5.15). The results for the cortisol/creatinine ratio were similar to that of urinary cortisol, with fluticasone causing suppression at all doses and with differences between the drugs at 800 µg and 1250 µg.

Conclusions – Single doses of inhaled fluticasone produce greater systemic bioactivity than budesonide when given by large volume spacer on a microgram equivalent basis in asthmatic children. The systemic bioactivity of fluticasone, like budesonide, is due mainly to lung bioavailability.

Keywords: asthmatic children, adrenal suppression, fluticasone propionate, budesonide, inhaled corticosteroid.

The systemic activity of inhaled corticosteroids is determined by several factors including glucocorticoid receptor potency, receptor binding affinity, plasma elimination half life, lipophilicity, and systemic tissue retention. The systemic effects of inhaled corticosteroids may arise by absorption from either the gastrointestinal tract or the lung.¹ Using a large volume spacer results in approximately 20% lung deposition and less than 10% oropharyngeal deposition.² As first pass hepatic metabolism is estimated at 99% for fluticasone propionate and 89% for budesonide,³ it is clear that, when using a large volume spacer, the main determinant of overall systemic bioavailability will be the lung bioavailability which avoids first pass hepatic metabolism for both fluticasone and budesonide.¹

Although systemic side effects from inhaled corticosteroids are not considered a problem at doses up to 400 µg/day, it is not uncommon for asthmatic children to require higher doses to maintain control of their disease. The aim of this study was to compare the systemic bioactivity of single doses of budesonide and fluticasone propionate given by large volume spacer to asthmatic children. As far as we are aware, no dose-ranging studies have previously addressed this issue in children.

Methods

Patients
Ten asthmatic children (outpatients, two girls) of mean (SD) age 11 (2.11) years (range 8–14) completed the study. The mean forced expiratory volume in one second (FEV₁) was 81.6 (6.26)% predicted (range 72–91% predicted) and mid forced expiratory flow (FEF25–75) was 74.8 (16.5)% predicted. All were using inhaled corticosteroids in a dose of ≤400 µg/day: beclomethasone dipropionate 200 µg/day (two children), 300 µg/day (one child), 400 µg/day (three children); budesonide 400 µg/day (two children), and fluticasone propionate 200 µg/day (two children). None had received oral corticosteroids in the preceding three months.

Both parents and children gave written informed consent. Approval for the study was obtained from the Tayside medical ethics committee for measurement of overnight
excretion and cortisol/creatinine ratios of 800 *p < 0.05; Table
Fluticasone Budesonide
15.38 (9.47 to 24.75)
17.21 (10.69 to 27.54)
20.53 (12.76 to 33.04)
24.43 (17.26 to 34.51)
Fluticasone 7.00 (4.94 to 9.16)**
Budesonide 9.89 (7.02 to 12.75)**
Budesonide 3.36 (2.74 to 4.00)**
Fluticasone 3.40 (2.74 to 4.00)**
Budesonide 2.26 (1.50 to 3.00)**
Fluticasone 2.21 (1.47 to 3.32)**
Budesonide 5.24 (3.18 to 8.61)
Fluticasone 6.49 (4.58 to 9.16)**
Budesonide 17.21 (10.69 to 27.54)
Fluticasone 11.99 (8.47 to 16.94)*
Budesonide 15.38 (9.47 to 24.75)

[p < 0.05; **p < 0.01; ***p < 0.001.

urinary cortisol but not plasma cortisol. All the children were continent of urine day and night.

PROTOCOL
A single blind (investigator blind) randomised (Latin square) crossover design was used. Patients were randomised to receive single inhaled doses of either placebo, budesonide (Pulmicort MDI, 50 μg and 200 μg per actuation, Astra Pharmaceuticals, Hertfordshire, UK), or fluticasone propionate (Fluicotide MDI, 25 μg, 50 μg, 125 μg, and 250 μg per actuation, Allen and Hanburys, Uxbridge, Middlesex, UK) taken in conjunction with their respective large volume spacer (Nebulhaler or Volumatic). Budesonide and flutica-
sone were given in identical single doses of 400 μg, 800 μg, and 1250 μg. A second placebo day was included one week after the last randomised treatment in order to check the reproducibility of urinary cortisol excretion between the two study days. Each study dose was taken at 20.00 hours at night with mouth rinsing, 15 minutes after the patient's usual bronchodilator treatment, with at least three days washout between each study treatment. The dose of inhaled steroid was kept constant throughout the study except for omitting it on the evening of the study day. Before taking the study drug patients emptied their bladder and collected all their overnight urine for 12 hours until 08.00 hours the following morning.

MEASUREMENTS
Overnight 12 hour urinary free cortisol excretion and cortisol/creatinine ratio were measured. The urinary cortisol was measured using a commercial radioimmunoassay kit (Incstar, Wokingham, Berkshire, UK). The coefficients of variability for intra-assay and interassay analytical imprecision were 8.50% and 6.65%, respectively. Urinary creatinine was measured on a Cobas-Bio autoanalyzer (Roche Products Ltd, Welwyn Garden City, Hertfordshire, UK). The intra-assay and interassay coefficients of variability were 4.55% and 0.63%, respectively.

STATISTICAL ANALYSIS
All data were analysed using multifactorial analysis of variance (MANOVA) and Duncan's multiple range testing with subjects, treatments, period, and doses as factors. All data were log transformed as they were not normally distributed. For comparison of the proportion of cortisol values below 10 nmol/12 hours the χ² test was used.

RESULTS
Data for 12 hour urinary free cortisol and cortisol/creatinine ratio are given in table 1. The overnight 12 hour urinary cortisol output (nmol/12 hours) showed suppression for all doses of fluticasone propionate compared with placebo (as fold difference from placebo with 95% CI): F400 μg (2.04-fold, 95% CI 1.13 to 3.36), F800 μg (3.77-fold, 95% CI 1.89 to 7.50), F1250 μg (3.49-fold, 95% CI 1.98 to 6.15), whereas budesonide caused no suppression at any dose (fig 1). Differences were seen between the drugs at dose levels of 800 μg and 1250 μg for urinary cortisol: B800 μg versus F800 μg (2.65-fold, 95% CI 1.26 to 5.58), B1250 μg versus F1250 μg (2.94-fold, 95% CI 1.67 to 5.15). The results for the cortisol/creatinine ratio were identical to those of urinary cortisol with suppression by flutica-
sone at all doses and with differences between the drugs at 800 μg and 1250 μg. The data for all three doses are also presented as individual values to show dispersion and outliers (fig 2).

Figure 1 Geometric mean (SE) 12 hour urinary free cortisol levels with single doses of budesonide (BUD), fluticasone (FP) and placebo. Crosses denote a significant difference (p < 0.05) between budesonide and fluticasone at a given dose level. Asterisks denote a significant difference between the steroid dose and placebo (**p < 0.05, ***p < 0.01, ****p < 0.001).

Figure 2 Individual values of urinary free cortisol for fluticasone (FP) and budesonide (BUD). Six of 30 budesonide and 18 of 30 fluticasone measurements were below 10 nmol/12 hours (p < 0.005).
Adrenal suppression with budesonide and fluticasone propionate

and show that six of 30 budesonide measurements and 18 of 30 fluticasone measurements were below 10 nmol/12 hours (p < 0.005).

There was no significant difference in urinary cortisol excretion between the two placebo days: placebo day 1 (18.58 nmol/12 hours) versus placebo day 2 (18.71 nmol/12 hours). The intrasubject coefficient of variation for the placebo days was 5.37%.

Discussion

The results of this study show that, when given by large volume spacer to asthmatic children, single inhaled doses of fluticasone propionate of 400–1250 μg produce suppression of urinary cortisol compared with placebo, whereas corresponding doses of budesonide do not.

The greater systemic bioactivity of fluticasone propionate revealed by this study can be explained by its basic pharmacology. The estimates for topical glucocorticosteroid activity as assessed by the MacKenzie skin vasoconstrictor assay vary. However, it is generally accepted that fluticasone propionate is at least twice as potent as budesonide and beclomethasone dipropionate. Fluticasone also has a greater glucocorticoid receptor affinity than budesonide, higher lipophilicity, and a longer glucocorticoid/receptor complex half life. In adults the plasma half life of fluticasone is 14.4 hours compared with 2.3 hours for budesonide. Interestingly, in children there is 40% greater clearance of budesonide with a shorter elimination half life at 1.5 hours compared with adults. This increased clearance in children probably explains why no suppression of overnight urinary cortisol compared with placebo was found at any of the doses of budesonide studied when we have previously found suppressed overnight urinary cortisol with single doses of 1000 μg in adults. The absence of significant dose related suppression of urinary free cortisol with budesonide has been demonstrated in a previous paediatric study with doses ranging from 200 μg to 800 μg/day. From first principles, greater systemic bioactivity effects would be predicted for a steroid which is more potent, has greater affinity and, when given by the inhaled route, has no first pass metabolism in the lung.

In adult studies budesonide delivered by Nebuhaler causes increased adrenal suppression compared with metered dose inhaler alone. This presumably reflects the effect of enhanced lung bioavailability with the spacer which outweighs the effect of reduced oral bioavailability, at least for budesonide where first pass metabolism is 89%. The present finding of adrenal suppression caused by single doses of inhaled fluticasone compared with budesonide, using a large volume spacer and mouth rinsing, further highlights the point that lung bioavailability is also the determinant of its systemic bioactivity in asthmatic children.

In conclusion, our study provides clear evidence of adrenal suppression with single doses of inhaled fluticasone propionate given by spacer to asthmatic children. It is likely that, with repeated dosing, the differences between fluticasone and budesonide would be greater due to the longer plasma half life and receptor binding affinity as well as enhanced systemic tissue retention for fluticasone than for budesonide. Further chronic dose ranging studies are now indicated to resolve this issue.

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