Lung deposition of aerosol—a comparison of different spacers

H J Zar, E G Weinberg, H J Binns, F Gallie, M D Mann

Abstract

Aims—To investigate (1) aerosol lung deposition obtained from two small volume conventional spacers (Babyhaler and Aerocam) and a homemade spacer (modified 500 ml plastic cold drink bottle); (2) the effect of using a face mask or mouthpiece; and (3) the relation between age and pulmonary deposition.

Methods—Lung deposition of aerosolised technetium-99m DTPA inhaled via spacer was measured in 40 children aged 3–7 years with stable asthma. Each patient performed a single randomly assigned inhalation using two spacers. Three studies were performed: Babyhaler compared to Aerocam (with facemasks); Babyhaler with facemask compared to Babyhaler with mouthpiece; and Babyhaler with mouthpiece compared to a 500 ml bottle.

Results—Median lung aerosol deposition from a Babyhaler and Aerocam—with masks were similar (25% vs 21%, p = 0.9). Aerosol lung deposition from a Babyhaler with mask compared to a Babyhaler with mouthpiece was equivalent (26% vs 26%, p = 0.5). Lung deposition was higher from a 500 ml bottle compared to a Babyhaler in both young (25% vs 12.5%, p = 0.005) and older children (42% vs 22.5%, p = 0.003). A notable reduction in pulmonary deposition occurred at 50 months of age.

Conclusion—A Babyhaler or Aerocam produces equivalent lung deposition of aerosol. There is no difference in lung deposition when a mask or mouthpiece is used. A modified 500 ml plastic bottle produces greater pulmonary aerosol deposition than a conventional small volume spacer.

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A metered dose inhaler (MDI) with spacer provides the optimal system for delivery of maintenance therapy for childhood asthma. A spacer, used in conjunction with an MDI improves drug delivery to the lungs and significantly reduces local and systemic side effects from drugs, especially inhaled corticosteroids. Recent studies suggest that an MDI with spacer is also the preferred delivery system for children with acute asthma. Equivalent or better bronchodilatation is obtained when β agonists are delivered via MDI with spacer compared to nebulisation in children with acute asthma, even when the degree of airways obstruction is severe.

Young children with asthma are unable to use an MDI effectively without the addition of a spacer. Small volume spacers such as the Babyhaler (GlaxoWellcome) and Aerocam (Trudell Medical, Canada) have been developed for this purpose. These spacers consist of a facemask connected to a cylindrical extension device via a one way valve. The facemask can be replaced by a mouthpiece. Home made spacers, principally a 500 ml plastic cold drink bottle have been adapted for use in developing countries. Although these spacers are widely used there is little data on their comparative efficacy or on the effect of a mouthpiece compared to a facemask for delivery of aerosolised medication.

We have previously developed a system to measure aerosol lung deposition from different spacers in older children. This technology was adapted in order to: (1) compare lung deposition of aerosol from the Babyhaler and Aerocam; (2) investigate the effect of a facemask or mouthpiece on lung deposition of aerosol; (3) compare pulmonary aerosol deposition obtained from the Babyhaler and modified 500 ml plastic bottle; and (4) examine the relation between age and pulmonary deposition.

Methods

PATIENTS

Forty patients with moderate asthma who regularly attend the asthma clinic at Red Cross Children’s Hospital, participated in the study. Inclusion criteria were a requirement for regular inhalation therapy for control of asthma, absence of an acute asthma exacerbation within the preceding six weeks, and age 3–5 years (studies 1–3) or 5–7 years (study 3 only). Patients were excluded if they had clinical signs or symptoms of airways obstruction on the day of the test. In addition, children older than 5 years had a peak expiratory flow rate of greater than 85% of the predicted normal value on the day of the study. Informed consent was obtained from a parent or legal guardian. The study was approved by the Ethics Review Committee, UCT Medical School.

DELIVERY AND MEASUREMENT OF AEROSOL

Technetium-99m DTPA (code N108, Amersham International plc, Amersham, UK) aerosol was used to measure deposition in the lungs. The delivery system consisted of a 2 litre anaesthetic rebreathing bag, which could be sealed. A two way tap (isolation valve) of the type used in medical gas lines was attached to
the proximal end of the bag and a gas flow controller from a Baumanometer to the distal end. The bag was filled with aerosol by connecting it to a micro Cirrus nebuliser (Amersham Health Care code N 1430, Amersham International plc, Amersham, UK) with a 1 m length of corrugated inspiratory tubing and passing oxygen vertically up through the nebuliser, corrugated tubing, and bag at a flow rate of 6 l/min for a minimum of 10 minutes. A plastic tube on the flow valve on the distal end of the bag carried all aerosol passing through the bag to a trapping system. The bag was sealed by closing the Baumanometer flow valve and then the two way tap before being removed from the circuit and attached by this tap to the particular spacer device being tested.

The activity available for inhalation was calculated from static one minute images of the bag recorded before and after the patient breathed from the delivery system and spacer combination. Deposition of aerosol in the lungs was calculated from a dynamic series of posterior images recorded at one second per frame for two minutes while the patient breathed deeply from the spacer and attached bag. All activity in the lungs was calculated from static one minute images of the study were taken as total pulmonary activity.

Each patient performed two separate inhalations in order to compare the efficacy of different spacers. The spacers were tested in two sequential inhalations. Immediately before beginning the second inhalation, the residual activity in the patient’s lungs was measured and later subtracted from the total pulmonary activity measured in the second study. Subtraction of residual activity was necessary to obtain an accurate measurement for aerosol lung deposition from the second spacer.

STATISTICAL ANALYSIS
Statistical analysis was performed using the Statistical Package for the Social Sciences (SPSS). Differences in drug delivery from the spacers in a single study were planned comparisons and assessed for significance using the Wilcoxon matched pairs test.

Results
The median (95% confidence intervals) ages of children in the first, second, and third studies were 52 (47–56), 51 (44–54), and 61 (54–72) months respectively. In the third study, children in the younger (3–5 year) group were 46 (42–51) months while those in the older group were 79 (75–85) months.

Table 1 summarises the results of lung deposition of aerosol. The median (95% confidence intervals) aerosol deposition in the lungs from the Babyhaler and Aerochamber with face-masks was 25% (15.2–28.6) and 21% (18.1–27.3) respectively, which were not significantly different. Median aerosol deposition from the Babyhaler with mouthpiece was 12.5% (6.5–19.5) compared to 25% (19.7–31.5) for the Babyhaler with mouthpiece for both young (25% (19.7–31.5) compared to 12.5% (6.5–19.5), p = 0.005) and older children (42% (36.0–44.0) compared to 22.5% (15.9–27.7), p = 0.003).

The age of the child affected the amount of aerosol deposited in the lungs, with decreased deposition occurring in younger children. At approximately 50 months of age, there was a notable reduction in the amount of pulmonary deposition from the second spacer.

<table>
<thead>
<tr>
<th>Spacing</th>
<th>Median (95% CI)</th>
<th>p value</th>
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<tbody>
<tr>
<td>Study 1</td>
<td></td>
<td></td>
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<tr>
<td>Babyhaler with mask</td>
<td>25 (15.2–28.6)</td>
<td>0.9</td>
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<tr>
<td>Aerochamber with mask</td>
<td>21 (18.1–27.3)</td>
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<tr>
<td>Study 2</td>
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<tr>
<td>Babyhaler with mask</td>
<td>26 (21.4–36.6)</td>
<td>0.5</td>
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<tr>
<td>Babyhaler with mouthpiece</td>
<td>26 (22.0–31.4)</td>
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<td>Study 3</td>
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<tr>
<td>Age 3–5 years</td>
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<tr>
<td>Babyhaler with mouthpiece</td>
<td>12.5 (6.5–19.5)</td>
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<td>500 ml bottle</td>
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<td>Age 5–7 years</td>
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<tr>
<td>Babyhaler with mouthpiece</td>
<td>22.5 (15.9–27.7)</td>
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<td>500 ml bottle</td>
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Values expressed as percentage aerosol deposited.
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conventional spacer, the Babyhaler. This plastic bottle produced greater pulmonary discussion; this probably accounts for the lower studies were younger than those in the second. Children in the 3–5 year age group of the third study were younger than those in the second study; this probably accounts for the lower aerosol deposition obtained from a Babyhaler in the former study.

Figure 1 Pulmonary aerosol deposition from a bottle used as a spacer by age.

deposition obtained when the bottle was used as a spacer (fig 1). Of the children younger than 50 months of age, one had pulmonary deposition of more than 30%, while only a single child older than 50 months had deposition less than 30%. This pattern of deposition was similar for the Babyhaler but at a lower level. For the Babyhaler, only one child younger than 55 months had deposition of more than 19%, while seven of 10 children older than 55 months had deposition of more than 19%. Children in the 3–5 year age group of the third study were younger than those in the second study; this probably accounts for the lower aerosol deposition obtained from a Babyhaler.

Discussion

This study showed that a modified 500 ml plastic bottle produced greater pulmonary aerosol lung deposition than a small volume conventional spacer, the Babyhaler. This effect was seen in children of various ages. Two widely available small volume spacers, the Babyhaler and Aerochamber produced equivalent aerosol lung deposition in young children. Furthermore, aerosol deposition did not differ whether a mask or mouthpiece was used.

Many properties of a spacer may affect pulmonary deposition of aerosol, including the volume and shape of the device, the electrostatic charge of the side walls, volume of dead space, and presence or type of valves. The efficiency of a 500 ml bottle may be partly a result of its physical characteristics. A 500 ml bottle was selected in preference to a larger volume bottle as it more closely approximates the optimum chamber dimensions for a spacer for children (experimentally found to be a cylinder approximately 11 cm in length by 3.5 cm in diameter). Furthermore, a 500 ml bottle has been previously reported to be an effective spacer, producing equivalent aerosol deposition and bronchodilatation to that obtained from a conventional spacer in older children. Our results suggest that the absence of a one way valve (as occurs with a bottle) does not adversely affect aerosol deposition but may in fact increase the amount of aerosol delivered. Valveless spacers have been reported to enhance the delivery of aerosol to the lungs in infants with chronic lung disease. Similarly, absence of a valve may prove beneficial for children as the need to overcome the resistance of the valve on inspiration is obviated.

Although the Aerochamber is less than half the volume of the Babyhaler, equivalent aerosol deposition was obtained from both devices. The choice of spacer size must balance the need for a small volume with the loss of aerosol because of impaction on the side walls which increases as the volume diminishes. Previous studies have reported that larger doses of aerosol are obtainable from the Babyhaler than the Aerochamber. However, these studies measured the drug collected on filter paper interposed after the inspiratory valve of the spacer; thus they reflect the dose of aerosol delivered at the child’s mouth and not the amount deposited in the lungs. The reduction in lung aerosol deposition because of the smaller volume of the Aerochamber may be countered by the large dead space in the Babyhaler, thus explaining the equivalence of these spacers.

Characteristics of the child such as their tidal volume, breathing pattern, and technique of spacer usage may further affect drug deposition. As evidence of this, reduced deposition of aerosol occurred at approximately 50 months of age with all spacers. Reasons for this may be related to the ability of the child to cooperate as well as age dependent differences in pulmonary mechanics. Lower tidal volumes and inspiratory flow rates generated with younger age, decrease pulmonary aerosol delivery resulting in substantially less lung deposition in young children compared to older subjects. An MDI with mouthpiece rather than facemask has been recommended for children 3–5 years old. The current British Thoracic Guidelines for the management of childhood asthma advise use of a spacer with mask for children under 2 years. Our study suggests that the mouthpiece and mask produce equivalent aerosol deposition. Although aerosol leakage from the perimeter of the mask where it fits onto the face may result in lower lung deposition, this did not appear to be a problem. However, we were careful to use a well fitting mask that was firmly applied to the face. Choice of the delivery method should be based on the availability and cost of the system as well as the ability and preference of the individual child. Whether a facemask or mouthpiece is selected, the correct technique for use of the delivery system should be carefully demonstrated to the care giver.

Exposure of children to radiation raises ethical concern. However, the total radiation dose for each child in this study was extremely small. The estimated dose of radiation that children were exposed to is far less than that obtained from a single abdominal x ray and may be similar to the amount of naturally occurring radiation that is acquired daily.

The method of aerosol delivery in our study differs from that of an MDI. We have previously developed and tested this delivery system in older children. The aerosol used was generated via an nebuliser that produces particles with a mass median aerodynamic diameter of 2–4 µm. Lung deposition of aerosol from an MDI might differ from that attained...
using our delivery system. However, the percentage deposition obtained is not important, but rather the relative deposition from different spacers. As the delivery system was kept constant and the particular spacer varied, a valid comparison of the efficacy of different spacers could be made.

In conclusion, a modified 500 ml plastic bottle produced superior aerosol deposition to that obtained from a conventional small volume valved spacer. The Aerochamber and Babyhaler were equivalent in the amount of pulmonary aerosol deposition produced. Use of a mouthpiece rather than a facemask did not increase the amount of aerosol deposited in the respiratory tract in children older than 3 years.

The choice of spacer and use of mask or mouthpiece should be made based on consideration of factors such as patient preference, cost of the device, ability of the care giver or patient to use the spacer correctly, and type of MDI used. Use of a 500 ml modified bottle as a spacer for children requiring inhaled therapy should be encouraged.

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