STUDY SUMMARY

AeroChamber* Brand of Valved Holding Chambers

Updated: March 2014
Foreword

The **AeroChamber**® Brand of Valved Holding Chamber (VHC) has been continually updated and improved upon since it was first introduced in 1983. The product enhancements are designed to increase ease of use for patients while enhancing the consistency of aerosol medication delivery.

The following variants of **AeroChamber Plus**® Valved Holding Chamber are available:

- **AeroChamber Plus**® VHC
- **AeroChamber Plus**® Flow-Vu® Anti-Static VHC

Both of the variants provide comparable fine particle delivery to the MDI alone. *In vitro* performance is not affected by the addition of the product enhancements (anti-static chamber, Flow-Vu® Indicator). The enhancements were made to increase ease of use for patients and their caregivers. The Flow-Vu® Indicator is a feedback tool that helps provide assurance that inhalation is performed correctly and allows caregivers to: ensure a proper seal, coordinate actuation with inhalation and count patient breaths. The anti-static chamber provides consistent drug delivery and can be used right out of package with no pre-treatment required.

The aerosol drug delivery performance of **AeroChamber**® Brand of VHC is supported by hundreds of peer-reviewed scientific studies. This study summary includes some of the more recent and relevant studies.

The following sections are included within the Study Summary:

1. Guidelines recommending Valved Holding Chambers
   - National and International Guidelines recommending the use of VHCs with MDIs

2. Importance of Valved Holding Chambers
   - VHCs help improve medication delivery, reduce oropharyngeal deposition and assist patients overcome difficulties in the co-ordination of actuation of the MDI with inhalation

3. Most Recommended - **AeroChamber**® Brand of Chambers
   - Confidence in Aerosol Drug Delivery
   - List of recommendations from MDI companies

4. Equivalency data
   - *In vitro* data showing comparable fine particle delivery of **AeroChamber Plus**® VHC variants compared to MDI alone

5. Importance of Feedback Features
   - Researchers highlight the importance of a feedback mechanism on aerosol delivery devices

6. Importance of Facemask Seal
   - Facemask seal is one of the most important factors in aerosol drug delivery to infants and children

7. Performance with different Metered Dose Inhaler formulations
   - *Divided by drug formulation, the studies are listed in chronological order with the most recent studies appearing first

8. Large versus Small Volume Valved Holding Chambers
   - Comparable performance between large and small volume VHCs

9. Metered Dose Inhalers and Valved Holding Chambers Versus Nebulizers
   - Advantages of MDI/Valved Holding Chamber versus Nebulizer
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GUIDELINES RECOMMENDING VALVED HOLDING CHAMBERS

The Metered Dose Inhaler (MDI) is the most commonly prescribed aerosol medication system. Valved Holding Chambers are designed to improve medication delivery, reduce oropharyngeal deposition of medication and help patients overcome difficulties in the co-ordination between actuation with an MDI and inhalation. Incorrect inhaler technique is prevalent and is a major issue associated with poor Asthma control.

American Association of Respiratory Care (AARC) (www.aarc.org) – Neonatal and Pediatric Patients 2007
- A spacer/holding chamber should be used with an MDI
- A spacer/holding chamber with facemask is appropriate for patients (usually < 3 years) unable to use a mouthpiece

American College of Chest Physicians (ACCP) / American College of Allergy, Asthma & Immunology (www.chestnet.org) 2005
- For patients who have trouble coordinating inhalation with device actuation, the use of a spacer (with a valve) may obviate this difficulty
- The use of spacers is mandatory for infants and young children

- Children and adults with mild and moderate exacerbations of Asthma should be treated by bronchodilators given from a pMDI + Spacer/Holding Chamber with doses titrated according to clinical response
- In children aged 0-5, pMDI + spacer are the preferred method of delivery of B2 agonists or inhaled steroids
- The spacer should be compatible with the pMDI being used

Canadian Pediatric Asthma Consensus Guidelines (www.cmaj.ca/cgi/content/full/173/6_suppl/S12) 2005
- The use of a holding chamber with pMDI is strongly recommended for children

Canadian Thoracic Society – 2010 Consensus Summary for children six years of age and over, and adults
- The addition of a holding chamber with mouthpiece is helpful in overcoming poor hand-mouth coordination and reducing side effects, with increased drug delivery and lung deposition
- Holding chambers with facemask attachments are useful for the elderly, who can use four to six tidal breaths for each actuation of the medicine

Global Initiative on Asthma (GINA) (www.ginasthma.com) 2011
- pMDI + dedicated Spacer/Holding Chamber with facemask is the preferred delivery system for children 4 years of age and younger
- pMDI + dedicated Spacer/Holding Chamber with mouthpiece is the preferred delivery system for children between 4 and 6 years of age

Global Initiative for Chronic Obstructive Lung Disease (GOLD) (www.goldcopd.com) 2014
- For the MDI, the addition of a large or small volume spacer often overcomes coordination problems, and improves lower airway deposition and clinical benefit

International Primary Care Respiratory Group (IPCRG) (www.theipcrg.org) 2006
- The preferred device for administering inhaled asthma medication for infants and young children is a pressurized MDI with a spacer and face mask
- As the child’s ability to co-operate improves (often around the age of 4-6 years), a spacer with a mouthpiece can be used rather than a face mask

- All patients taking inhaled steroids should use a Spacer/Holding Chamber
- Patients under 5 years should use a Spacer/Holding Chamber with Facemask for inhaled steroids
National Institute for Clinical Excellence, UK (NICE) (www.nice.org.uk) 2010

- Spacer/Holding Chamber recommended with a facemask where necessary for both corticosteroids and bronchodilators (Children under 5)
- A press and breathe pMDI used with an appropriate Spacer/Holding Chamber is first choice for corticosteroids (Children aged 5-15 years)
- COPD – pMDI alone is rarely suitable for use with the elderly
- The spacer should be compatible with the patient’s metered-dose inhaler

BACKGROUND: In practice it is logical that inhalers are prescribed only after patients have received training and demonstrated their ability to use the device. However, many patients are unable to use their pressurised metered-dose inhaler devices (pMDIs) correctly. We assessed the relationship between asthma control and patients’ ability to use their prescribed pMDIs.

METHODS: Evaluation of 3,981 (46% male) primary care asthma patient reviews, which included inhaler technique and asthma control, by specialist nurses in primary care in 2009. The paper focuses on people currently prescribed pMDI devices.

RESULTS: Accurate data on reliever and preventer inhaler prescriptions were available for 3,686 and 2,887 patients, respectively. In patients prescribed reliever inhalers, 2,375 (64%) and 525 (14%) were on pMDI alone or pMDI plus spacer, respectively. For those prescribed preventers, 1,976 (68%) and 171 (6%) were using a pMDI without and with a spacer, respectively. Asthma was controlled in 50% of patients reviewed. The majority (60% of 3,686) were using reliever inhalers, 2,375 (64%) and 525 (14%) were on pMDI alone or pMDI plus spacer, respectively. Incorrect pMDI use was associated with poor asthma control (p<0.0001) and more short burst systemic steroid prescriptions in the last year (p=0.038). Of patients using beclometasone (the most frequently prescribed preventer drug in our sample), significantly more of those using a breath-actuated pMDI device (p<0.0001) and a spacer (p<0.0001) were controlled compared with those on pMDIs alone. CONCLUSIONS: Patients who are able to use pMDIs correctly have better asthma control as defined by the GINA strategy document. Beclometasone via a spacer or breath-actuated device resulted in better asthma control than via a pMDI alone. Patients prescribed pMDIs should be carefully instructed in technique and have their ability to use these devices tested; those unable to use the device should be prescribed a spacer or an alternative device such as one that is breath-actuated.


Whilst the inhaled route is the first line administration method in the management of asthma, it is well documented that patients can have problems adopting the correct inhaler technique and thus receiving adequate medication. This applies equally to metered dose inhalers and dry powder inhalers and leads to poor disease control and increased healthcare costs. Reviews have highlighted these problems and the recent European Consensus Statement developed a call to action to seek solutions. This review takes forward the challenge of inhaler competence by highlighting the issues and suggesting potential solutions to these problems. The opportunity for technological innovation and educational interventions to reduce errors is highlighted, as well as the specific challenges faced by children. This review is intended as a policy document, as most issues faced by
patients have not changed for half a century, and this situation should not be allowed to continue any longer. Future direction with respect to research, policy needs and practice, together with education requirements in inhaler technique are described.


Many children with asthma do not use their inhalers correctly and consequently gain little or no therapeutic benefit from the treatment. The focus of inhalation therapy should be on those inhalers which are easiest to use correctly by various groups of children and the amount of tuition and training required to obtain a correct technique. It is recommended that clinicians focus on a limited number of inhalers. Most children can be taught effective inhalation therapy by using a pMDI, a pMDI with a spacer, or a DPI. Most preschool children can be taught effective use of a pMDI and spacer with a valve system and a face mask. Therefore, this is the preferred mode of delivery in these age groups. When the child is capable of using the spacer without a face mask this administration technique should be adopted. In older children pMDIs are more difficult to use correctly than a pMDI with a spacer, a DPI, or a breath-actuated pMDI. Because DPIs and breath-actuated pMDIs are more convenient to use these devices are normally considered the preferred inhalation devices in these age groups except for administration of beclometasone dipropionate, which for safety reasons should be delivered by a spacer.


Aerosol inhalation is considered the optimal route for administering the majority of drugs for the treatment of obstructive airways diseases. A number of Pressurised Metered-Dose and Dry Powder Inhalers are available for this purpose. However, inhalation of therapeutic aerosols is not without difficulty; it requires precise instructions on the inhalation manoeuvre, which is different from spontaneous normal breathing. Also, the characteristics of the inhaler device have to be suitable for the user. Available data indicate a frequent lack of knowledge demonstrated by health professionals and patients on the inhalation manoeuvre and handling of inhalers, resulting in a reduction of therapeutic benefit. This paper reviews the literature concerning the fundamental aspects of inhaler devices, inhalation manoeuvre and device selection, in an attempt to increase the knowledge of, and to optimise the clinical use of, therapeutic inhalers.


AIM: Spacer devices are inhalation aids of varying dimension and complexity, specifically designed to overcome problems with the use of pressurised metered dose inhalers (pMDIs). The aim of this review is to examine the current understanding about these inhalation devices and discuss their advantages and disadvantages. METHODS: The pertinent literature concerning the characteristics and effects of spacers on delivery and lung deposition of inhaled medications, as well as their clinical efficacy in patients with reversible airway obstruction, is examined. RESULTS: Spacers minimise problems of poor inhalation technique with pMDI, reduce oropharyngeal deposition and increase lung deposition. Spacers improve the clinical effect of inhaled medications, especially in patients unable to use a pMDI properly. Compared to both pMDIs and dry-powder inhalers, spacers may increase the response to beta-adrenergic bronchodilators, even in patients with correct inhalation technique. A pMDI plus spacer has proven to be viable lower cost alternative to the use of a nebuliser for delivering large bronchodilator doses in patients with severe acute asthma or chronic obstructive pulmonary disease. The use of large-volume spacers is recommended for delivering high doses of inhaled corticosteroids, and may permit a lower maintenance dose to be used. CONCLUSION: pMDIs may be routinely fitted with a spacer, especially in situations where correct pMDI use is unlikely.


Objective: Spacer devices (SD) in conjunction with metered dose inhalers (MDI) have been shown to be as effective as saline nebulizers for the delivery of beta-agonists. A preliminary study suggests that SDs are not consistently used. The purpose of this study was to investigate patterns of SD ownership and use to identify potential targets for future educational efforts to increase ownership and use of SD. Methods: Cross-sectional convenience sample survey of patients presenting to an academic Emergency Department (ED) with a history of asthma/COPD (chronic obstructive pulmonary disease). Informed consent was obtained. Survey data included demographics, association with a primary care physician (PCP), SD ownership, patterns of use, opinions of efficacy about SD and disease severity assessed by duration of asthma/COPD, prior ED visits, hospitalizations, and history of prior intubation. Patterns of use are described and univariate and multivariate analyses were used to identify factors associated with SD ownership. Results: Of the 313 patients, 55.9% were female, the mean age was

Although the principles of asthma management are well established in Europe, the available data indicate that asthma in patients is not well controlled. Many patients derive incomplete benefit from their inhaled medication because they do not use inhaler devices correctly and this may compromise asthma control. The Aerosol Drug Management Improvement Team (ADMIT), incorporating clinicians from the UK, Germany, France, Italy, Spain and The Netherlands, reviewed published evidence to examine ways to improve the treatment of reversible airways disease in Europe. Data indicate that there is a clear need for specific training of patients in correct inhalation technique for the various devices currently available, and this should be repeated frequently to maintain correct inhalation technique. Devices which provide reassurance to patients and their physicians that inhalation is performed correctly should help to improve patient compliance and asthma control. Educational efforts should also focus on primary prescribers of inhaler devices. ADMIT recommends dissemination of information on the correct inhalation technique for each model of device by the use of an accessible dedicated literature base or website which would enable to match the appropriate inhaler to the individual patient. There is also a need for standardisation of prescribing practices throughout Europe. Regular checking of inhalation technique by prescribers is crucial as correct inhalation is one of the keystones of successful asthma management.


The two most important differences between inhaled and oral therapy are (1) the lungs have evolved to exclude foreign material while the gut has evolved to take in large amounts of foreign material, and (2) even if patients adhere to a treatment regimen (regimen compliance or adherence), they may fail to derive any benefit from using an inhaler due to failure of drug delivery (poor device compliance). In other words: True compliance = regimen compliance x device compliance. Aerosol scientists, building on the observations of those working in the field of industrial hygiene, have developed devices that largely address the challenge of bypassing the lung's defenses, in that current devices generate aerosols that contain a significant proportion of particles in the range of 1-5 microm. These have a relatively high probability of entering the lungs and depositing through impaction and/or sedimentation. The development of delivery systems for systemically acting drugs has led to further refinement. The second issue, that of patient behavior, has, until very recently, received very little attention from those developing devices. Regimen compliance involves taking the medication at the suggested times. Device compliance (using the device optimally) is dependent on competence and contrivance. A patient taking a tablet before rather than after a meal is likely to receive some therapeutic benefit even if the effect is suboptimal. A patient whose device compliance is poor because either they are not competent to use the device or contrive to use it in an ineffective manner may derive little or no benefit even if they are scrupulously adhering to their treatment regimen. Lack of precision in the use of the terms “compliance” and “adherence” has contributed to the failure to build in features that may help address issues relating to patient behavior. The resurgence of interest in developing devices that can be used to deliver potent systemically acting drugs has, out of necessity, led to the development of systems that help minimize the impact of poor competence or contrivance on drug delivery. There are suggestions, that need to be confirmed, that regimen compliance (adherence) can be influenced by providing feedback. In the absence of formal studies, comparison of the high-tech and low-tech approaches to improving device compliance incorporated into novel devices might provide valuable insights into what aspects of feedback are important in the clinical setting.


Inhalational drug delivery is the primary mode of asthma therapy in children and is the main focus on this article. Pressurized metered dose inhalers (pMDIs) are now the method of choice in infants and children under 5 years old, when used in combination with an appropriate valved holding chamber or spacer. Spacers are particularly important for steroid inhalation to maximize lung deposition and minimize unwanted oropharyngeal deposition. Optimal inhalation technique with a pMDI-spacer in infants is to inhale the drug by breathing tidally through the spacer. Drug delivery to the lungs using pMDIs can vary greatly, depending on the formulation used and the age of the child. Dry powder inhalers (DPIs) are driven by the peak
inspiratory flow of the patient and are usually not appropriate for children under 5 or 6 years of age. Nebulizers continue to play a role in the treatment of acute asthma where high doses of bronchodilator are required, though multiple doses via pMDI spacer may suffice. Important drug delivery issues specific to children include compliance, use of mask versus mouthpiece, lower tidal volumes and inspiratory flows, determination of appropriate dosages, and minimization of adverse local and systemic effects. CONCLUSION: In recent times, pMDI-spacers have become the most commonly used approach to aerosol therapy in children. They can be used for all ages, and can be used both for long-term preventative therapy and for short-term treatment of acute exacerbations. One of the main advantages of pMDI-spacer use is that normal tidal breathing can be used during aerosol administration, which makes them ideal for infants and younger children. However, there are many newer and more innovative devices available that offer therapeutic advantages in terms of increased efficacy, convenience, and compliance, particularly for older children and adolescents. The ability of a child to utilize an inhaler device and to perform the required inhalation technique consistently must be evaluated when making the choice of an appropriate delivery system for children of different ages.


The increasing prevalence of childhood asthma has become a concern among health practitioners. Effective management emphasizes long-term management and inhaled therapy has become the mainstay home management for children. However, proper utilization of medication is pertinent in improving control. Proper asthma education is mandatory in improving skills and confidence amongst parents. To assess the skills of using the metered-dose inhaler (MDI) with a spacer among asthmatic children before and after educational intervention and to analyse any difficulties which may occur amongst the participants in executing the assessment steps. A cross-sectional clinic based study involving 85 parents and children with asthma. A standardized metered-dose inhaler-spacer checklist of eight steps of medication usage and five steps of cleaning the spacer were used as the assessment tools for pre and post intervention. The performance on using the inhaler-spacer and spacer cleaning knowledge pre and two months post intervention was evaluated. One point was given for each correct step and zero points for incorrect answers/steps. The mean score for skills of inhaler technique improved significantly after educational intervention (3.51 to 6.01, p < 0.0001) as did the mean score for parental knowledge of spacer cleaning technique (1.35 to 3.16, p 0.001). Analysis showed only a limited improvement even after an educational session in three steps of inhalation technique: step 5 (23.5%/69.4%), step 6 (28.2%/68.2%) and step 7 (25.9%/61.2%). Parents with asthmatic children had poor skills in utilizing their children's medication. A short-term educational intervention was able to improve overall knowledge and skill but certain skills need more emphasizing and training.


Although most attention has been focused on the drugs used to control asthma, it is increasingly recognised that effective delivery of these drugs to the lungs is just as important. The most effective drugs, β2-agonists and corticosteroids, are given by inhalation so there has been a search for more efficient inhaler devices that are easier for patients to use. A symposium at the European Respiratory Society Annual Meeting in 2005 discussed some of the important issues in inhaler therapy in adults and children. This article summarises the major points of discussion that arose out of this symposium. New more effective inhaler devices are now becoming available and are likely to have an important impact on asthma management.


Aerosol therapy, the preferred route of administration for glucocorticosteroids and short-acting β2-adrenergic agonists in the treatment of paediatric asthma, may be given via nebulisers, metered-dose inhalers and dry powder inhalers. For glucocorticosteroids, therapy with aerosolized medication results in higher concentrations of drug at the target organ with minimal systemic side effects compared with oral treatments. The dose of drug that reaches the airways in children with asthma is dependent on both the delivery device and patient-related factors. Factors that affect aerosol drug delivery are reviewed briefly. Advantages and disadvantages of each device and device-specific factors that influence patient preferences are examined. Although age-based device recommendations have been made, the optimal choice for drug delivery is the one that the patient or caregiver prefers to use, can use correctly and is most likely to use consistently.
DO PEDIATRIC HEALTHCARE PROVIDERS KNOW HOW TO USE METERED DOSE INHALER PLUS SPACER DEVICES?  

We tested whether health practitioners correctly used MDI-spacer devices. Of 122 subjects, 89% had instructed a patient on using a spacer. Whilst performance with the AeroChamber* was the best, only 3% correctly demonstrated all the steps for that device.

HOW CAN WE IMPROVE ASTHMA MANAGEMENT?  

Asthma remains a poorly controlled disease both in Europe and the USA despite the availability of effective asthma treatments. Patient noncompliance, incorrect use of inhaler devices, insufficient treatment of peripheral airway inflammation as well as limitations of the asthma management guidelines themselves may all contribute to this poor control. Asthma control may be improved by improving the consultation process during the visit at the doctor. The ideal consultation would involve critical listening to the patient, accurate assessment of asthma symptoms as indicators for asthma control and prescribing the appropriate medication and dose for the individual patient according to the degree of severity of asthma. In addition, correctness of inhalation technique as performed by the patient should be regularly checked; patients should be educated and trained how to manage their personal condition and should be offered convenient follow-up options. Choosing the right inhaler for the patient may improve patient compliance. Inhaler choice should be based on an evidence-based rationale rather than on an empirical basis. The preference of the patient should also be taken into consideration, as it is the patient who has to use the inhaler daily over a long period of time. The ideal inhaler should demonstrate sufficient drug delivery to the lower airways as well as good drug distribution to both the central and peripheral airways. It should ensure consistency of the emitted dose, be easy to teach and use, be small in size and convenient to handle. It should also be multi-dose, require a low inspiratory airflow for activation, provide feedback to the patient on correct inhalation technique, be re-usable/refillable, have an appealing design, and have a reliable dose counter.

MISUSE OF CORTICOSTEROID METERED-DOSE INHALER IS ASSOCIATED WITH DECREASED ASTHMA STABILITY.

This study assessed whether the improper use of pressurized metered-dose inhalers (pMDIs) is associated with decreased asthma control in asthmatics treated by inhaled corticosteroids (ICS). General practitioners (GPs) included consecutive asthmatic outpatients treated by pMDI-administered ICS and on-demand, short-acting beta2-agonists. They measured an asthma instability score (AIS) based on daytime and nocturnal symptoms, exercise-induced dyspnoea, beta2-agonist usage, patient also was assessed. GPs (n=915) included 4,078 adult asthmatics; 3,955 questionnaires were evaluable. pMDI was misused by 71% of patients, of which 47% was due to poor coordination. Asthma was less stable in pMDI misusers than in good users (AIS: 3.93 versus 2.86, p<0.001). Among misusers, asthma was less stable in poor coordinators (AIS: 4.38 versus 3.56 in good coordinators, p<0.001). To conclude, misuse of pressurized metered-dose inhalers, which is mainly due to poor coordination, is frequent and associated with poorer asthma control in inhaled corticosteroid-treated asthmatics. This study highlights the importance of evaluating inhalation technique and providing appropriate education in all patients, especially before increasing inhaled corticosteroid dosage or adding other agents. The use of devices which alleviate coordination problems should be reinforced in pressurized metered-dose inhaler misusers.

POOR INHALATION TECHNIQUE, EVEN AFTER INHALATION INSTRUCTIONS, IN CHILDREN WITH ASTHMA.  

The aim of this study was to evaluate the effect of instructions to children with asthma (given by general practitioners or by pharmacy assistants) on how to inhale from metered dose inhalers with spacers (MDIs) or dry powder inhalers (DPI). We scored inhalation technique of asthmatic children according to criteria defined by the Netherlands Asthma Foundation, and related the performance to the inhalation instructions given. For each inhaler, a number of steps were considered essential for reliable drug delivery. Patients newly referred for asthma were asked to demonstrate their inhalation technique and to fill out a questionnaire on the inhalation instruction received prior to referral. Children participating in a clinical trial, who had received repeated comprehensive inhalation instructions, served as a control group. Sixty-six newly referred patients (1-14 years of age, median age 5 years; 37 boys) and 29 control patients (5-10 years of age, median age 7 years; 21 boys) completed the study. Sixty patients (91%) had received inhalation instruction prior to referral. Only 29% of these patients, using a dry powder inhaler, performed all essential steps correctly, compared to 67% of children using a metered dose inhaler/spacer combination (P < 0.01). Children who had received comprehensive inhalation instructions with repeated checks of proper inhalation technique at the pharmacy or in the clinical trial setting were more likely to perform all essential steps correctly (79% and 93%, respectively) than children who had received a single instruction by a general practitioner (39%, P < 0.01). Many asthmatic children use their inhalers devices too poorly to result in reliable drug delivery, even after inhalation instruction. Comprehensive inhalation instruction and repeated check-ups are needed to assure reliable inhalation technique.
Many children with asthma use their inhaler device incorrectly even after comprehensive inhalation instruction. The aim of this study was to identify factors associated with correct inhalation technique. Two hundred children with asthma demonstrated their inhalation technique. Patient characteristics and the components of inhalation instructions they had received were compared for children demonstrating a correct or incorrect inhalation technique. In addition, the inhalation technique of 47 newly referred patients was followed-up prospectively after repeated comprehensive instruction sessions. Seventy-eight percent of all patients demonstrated a correct inhalation technique. Patients who had received repeated instruction sessions and patients who had previously been asked to demonstrate the use of their inhaler during an instruction session were more likely than other children to demonstrate a correct inhalation technique (p < 0.001 and p = 0.03, respectively). Multiple logistic regression analysis showed that repetition of instructions was significantly associated with a correct inhalation technique (odds ratio (OR) 8.2, 95% CI 3.2-21.5; p < 0.0001) irrespective of type of inhaler used. Demonstration of the inhaler use by the patient was significantly associated with a correct inhalation technique for patients using a metered dose inhaler plus spacer device (OR 3.5, 95% CI 1.0-12.6; p = 0.05) but not for patients using a dry powder inhaler (OR 1.6, 95% CI 0.4-6.4; p = 0.54). The number of newly referred patients demonstrating a correct inhalation technique improved from 57.4% to 97.9% after three comprehensive instruction sessions. CONCLUSION: Inhalation instruction should be given repeatedly to achieve and maintain correct inhalation technique in asthmatic children.

The objective was to evaluate the correctness of the inhalation technique in a nationwide sample of patients and medical personnel, in order to define targeted educational goals. A total of 1,640 volunteers (746 patients, 466 nurses and 428 physicians) were evaluated. Only 9% of patients, 15% of nurses and 28% of physicians showed a correct inhalation technique. Physicians performed significantly better (mean score 77 +/- 23) than nurses (71 +/- 22) and patients (62 +/- 26). Scores in general practitioners and pediatricians were significantly lower than those of chest physicians and allergists. In conclusion, proper use of metered dose inhalers (MDI) in patients and medical personnel is still faulty. Despite the physician's awareness of the importance of a correct inhalation technique in the use of MDI, this study shows severe deficiencies, showing the need for substantial changes in educational efforts, and particularly addressed to general practitioners.
MOST RECOMMENDED - AEROCHAMBER* BRAND OF VHC

CHAMBERS ARE NOT INTERCHANGEABLE:

“Spacers should not be regarded as interchangeable: patients who use a spacer with their inhaler should use the spacer device named in the Summary of Product Characteristics (where specified by name). Patients whose asthma is well controlled and who are using a spacer should always use the same type of spacer and not switch between spacers. Different spacers may deliver different amounts of inhaled corticosteroid, which may have implications for both safety and efficacy.” Drug Safety Update – Inhaled products that contain corticosteroids, July, 2008. Medicines and Healthcare Products Regulatory Agency (MHRA).

“Commercially produced spacers with well-characterized drug output characteristics are preferable, although spacer devices or face masks differ in their drug delivery and therefore may not be interchangeable” GINA – Global Strategy for Asthma Management and Prevention. Updated 2012.

“Spacers come in different designs and, since the dose received may vary considerably from one device to another, a spacer device that has documented efficacy in young children is recommended.” GINA – Global Strategy for the Diagnosis and Management of Asthma in Children 5 Years and Younger, 2010.


“VHCs are not interchangeable…Parents, clinicians and pharmacists should be educated not to interchange VHCs once a child is stable on a particular ICS dose and VHC combination. Moreover, the initial prescription for a VHC should include language (e.g. “Do not substitute” or “Medically necessary”) to prevent the pharmacist from substituting a different VHC. We suggest that each VHC should be evaluated for relative lung bioavailability prior to their routine use with a particular ICS” Blake K et al. Bioavailability of inhaled fluticasone propionate via chambers/masks in young children. Eur Respir J 2012;39:1.

AEROCHAMBER* BRAND OF VHC IS RECOMMENDED FOR USE WITH THE FOLLOWING METERED DOSE INHALERS:

Airomir – Patient Instruction Leaflet, UK
Some people find it difficult to press their inhaler and breathe in at the same time. A spacer device helps to overcome this problem. The spacer that fits AIROMIR Inhaler is called the AeroChamber Plus* spacer device. If you use the AeroChamber Plus* spacer device please follow the instructions provided with it. Your doctor, nurse or pharmacist will be able to advise you about the AeroChamber Plus* device.

Alvesco® 160/80 Summary of Product Characteristics, EU
To address specific patient needs, such as finding it difficult to press the inhaler and breathe in at the same time, Alvesco® can be used with the AeroChamber Plus* spacer device.

Alvesco® 100/200 Product Monograph, Canada
In patients who find co-ordination of a pressurized metered dose inhaler difficult, a spacer device (AeroChamber Plus*) may be used with Alvesco®.

Alvesco® 40, 80 and 160 Patient Instruction Leaflet, UK
If you find it difficult to use the inhaler, your doctor may recommend the use of a spacer. The spacer that fits the Alvesco inhaler is called AeroChamber Plus*. If you use the AeroChamber Plus* device, please follow the instructions provided with it. Your doctor or pharmacist will be able to advise you about the device.

Atrovent® Inhaler CFC-Free 20 Summary of Product Characteristics, EU
The inhaler can be used with the AeroChamber Plus* spacer device. This may be useful for patients, e.g. children, who find it difficult to synchronise breathing in and inhaler actuation.

Atrovent® Inhaler CFC-Free 20 Patient Instruction Leaflet, UK
If you find breathing in and pressing the inhaler at the same time (step 3) difficult you should talk to your doctor or pharmacist, as you could use a spacer device (AeroChamber Plus*) with your inhaler. A spacer is a device designed to make step 3 easier. A spacer is generally a plastic container with a mouthpiece at one end and a hole for inserting the mouthpiece of the inhaler at the other end. The puff of medicine from your inhaler is sprayed into the spacer and the puff of medicine stays there, inside the spacer,
until you breathe in through your mouth from the spacer with the spacer mouthpiece in your mouth and with your lips closed around it. This means that you do not have to worry about breathing in and pressing the inhaler at the same time.

*Flutiform® pressurized inhalation suspension, Summary of Product Characteristics, EU*
Use of a spacer device with Flutiform inhaler is recommended in patients who find it difficult to synchronise aerosol actuation with inspiration of breath. The AeroChamber Plus® is the only spacer device recommended for use with Flutiform inhaler.

*Flutiform® pressurized inhalation suspension, Patient Instruction Leaflet, EU*
If you have difficulty using your inhaler your doctor or asthma nurse may give you a device called an AeroChamber Plus® spacing device, to help you to breathe your medicine into your lungs properly.

*Fostair 100/6 Inhalation Aerosol – Summary of Product Characteristics – EU*
Patients who find it difficult to synchronise aerosol actuation with inspiration of breath, may use the AeroChamber Plus® spacer device.

*Fostair 100/6 Inhalation Aerosol – Patient Instruction Leaflet – EU*
If you find it difficult to operate the inhaler while starting to breathe in you may use the AeroChamber Plus® spacer device. Ask your doctor, pharmacist or nurse about this device.

*Respimat® Soft Mist™ Inhaler – Kamin et al., 2011*
To ensure standardized dosing, the use of the Respimat® inhaler with spacer (AeroChamber Plus®) is recommended for all children below 5 years of age.

*Sabumalin® Inhaler 100 µg/dose, Summary of Product Characteristics, EU*
Salbutamol may be used with a Vortex® or AeroChamber® Plus spacer device by children and patients who find it difficult to synchronise aerosol actuation with inspiration.

*Sabumalin® Inhaler 100 µg/dose, Package Leaflet, EU*
Some people find it difficult to release a puff of medicine just after they start to breathe in. In this case, as well as for children, the Vortex® or AeroChamber Plus® spacer device can be used. Please refer to the product information of the spacer device for its correct handling.

*Seretide™ Evohaler™ 50 / 125 / 250 – Summary of Product Characteristics – EU*
Use of a spacer device with Seretide inhaler is recommended in patients who have, or are likely to have difficulties to coordinate actuation with inspiration. Either the Volumatic or AeroChamber Plus spacer device can be used (depending on National Guidance).

*Seretide™ Evohaler™ 50 / 125 / 250 - Patient Instruction Leaflet – EU*
If you or your child find it difficult to use Seretide Evohaler, either the Volumatic™ or AeroChamber Plus® spacer device may be used. Before starting to use a spacer device for the first time or if you need to change your make of spacer device, talk to your doctor, nurse or pharmacist.

*Qvar® 50/100 Product Monograph, Canada*
Where a spacer is considered necessary the AeroChamber® is a suitable device for use with QVAR™ MDI as the extrafine particle fraction is maintained.

*Qvar® 50/100 Summary of Product Characteristics, UK*
Where a spacer is considered necessary for specific patient needs, Qvar aerosol can be used with AeroChamber Plus® holding chamber, as the extrafine particle fraction is maintained.

*Qvar® 50/100 Patient Instruction Leaflet, UK*
The spacer that fits Qvar aerosol is called the AeroChamber Plus® spacer device.

*Zenhale™ Product Monograph, Merck Canada Inc.*
Use of the AeroChamber Plus® Anti-Static valved holding chamber is recommended with ZENHALE, in patients who find it difficult to synchronize aerosol actuation with inspiration of breath.
CLINICAL DATA WITH AEROCHAMBER® BRAND OF VHC INCLUDED IN THE FOLLOWING PRODUCT MONOGRAPHS:

- Advair® Product Monograph, Canada, May 2011
  50 / 125 / 250 mcg
- Flovent® HFA Inhalation Aerosol Product Monograph, USA, Jan 2012
  44 / 110 / 220 mcg
- Flovent® HFA Inhalation Aerosol Product Monograph, Canada, May 2011
  50 / 125 / 250 mcg
- Ventolin® HFA Inhalation Aerosol Product Monograph, USA, 2009
  90mcg

CONFIDENCE IN AEROSOL DELIVERY

Sold in over 100 countries around the world

Most studied brand of VHC
  Supported by over 500 in vitro and in vivo studies

30th Anniversary – AeroChamber® Brand of VHC (2013)
  Since its introduction, the AeroChamber® VHC has been continually updated and improved upon. Improvements were made to increase ease of use for patients while enhancing the consistency of aerosol medication delivery.

Winner of Top Ten Innovations in Technology Award (2013)
  AeroChamber Plus® Flow-Vu® Chamber has won an award for being one of the top 10 innovations in technology. The award was given by the nonprofit group - Allergy & Asthma Network Mothers of Asthmatics. A panel of editors, board members, families, and volunteers reviewed innovations and technologies that have changed the world, narrowing it down to the top 10 within the last 25 years. The award was given to products and services that have improved the quality of life for people with asthma and allergy conditions.

Free of Bisphenol A, Phthalates, Latex, Lead and PVC
  In 2010, we adopted a company position to heed the precautionary principle and move toward BPA free components for our products. There continues to be new studies regarding the potential health risks associated with BPA. Since the medications taken with our products are critical to the health of those that need them, it is our responsibility to lead in the development of consumer friendly products that are manufactured from materials that do not contain BPA.

Manufactured to the Highest Quality Standards
  AeroChamber Plus® Flow-Vu® Chambers are manufactured in a Class 8 Cleanroom manufacturing facility. TMI is registered to ISO 13485:2003, which is specific for medical device manufacturers.
EQUIVALENCY DATA – AEROCHAMBER* BRAND OF VHCS

**EQUIVALENT ASTHMA CONTROL WHEN INFANTS WITH ASTHMA USE TWO DIFFERENT VERSIONS OF THE AEROCHAMBER PLUS SPACER.** Chrystyn H, Ammari WG, Chetcuti P, Toor S. Am J Respir Crit Care Med 2013: 187;A4574.

The AeroChamber Plus* Spacer (Trudell Medical International, Canada) [AC] has recently been adapted to include a visual indicator that confirms the inhalation phase during use (AeroChamber Plus* Flow-Vu*; Trudell Medical International, Canada) [FV]. Movement of the indicator also confirms the required seal between the mask and the user’s face. The asthma control of infants with asthma, aged < 5 years, during routine use of these spacers has been compared and patient preference has been obtained. Ethical approval was obtained and all infants and their parents gave signed informed consent. All infants entered a 2 week run-in period using the AC. At visit 2 they were randomised to the AC or FV for the 12 week duration of this study. They returned after 6 and 12 weeks (visits 3 and 4). All infants were trained to use a gentle tidal breathing routine with their spacer. At each visit their asthma quality of life (ACQ; Juniper et al, Eur Respir J, 1996) was obtained by questioning each parent and their inhalation flow (IFR) was measured using the IN-Check Dial (Clement Clark International, UK). At visit 4 each FV parent was asked to rate their preference for the FV using a 5 point Likert scale (5=much better to 1=much worse). The FV was demonstrated to all AC parents and they were then asked the same preference rating. At visit 1 the mean (SD) age of the AC (n=38) and FV (n=38) infants was 3.3 (1.1) and 2.8 (0.9) years, respectively. A summary of the ACQ and IFR is presented in Table 1.

<table>
<thead>
<tr>
<th>Visit</th>
<th>ACQ</th>
<th>IFR (L/min)</th>
<th>IFR (L/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>AC</td>
<td>AC</td>
<td>FV</td>
</tr>
<tr>
<td>1</td>
<td>1.83 (0.88)</td>
<td>1.85 (0.91)</td>
<td>40.1 (16.4)</td>
</tr>
<tr>
<td>2</td>
<td>1.56 (0.91)</td>
<td>1.81 (0.87)</td>
<td>40.6 (14.8)</td>
</tr>
<tr>
<td>3</td>
<td>1.77 (1.16)</td>
<td>1.54 (1.07)</td>
<td>41.1 (15.7)</td>
</tr>
<tr>
<td>4</td>
<td>1.29 (0.90)</td>
<td>1.62 (0.92)</td>
<td>40.1 (16.0)</td>
</tr>
</tbody>
</table>

Two-way analysis of variance revealed no difference in the ACQ or the IFR between the visits and the two groups. Patient preference is shown in Table 2.

<table>
<thead>
<tr>
<th>Likert Scale Rating</th>
<th>AC</th>
<th>FV</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>24</td>
<td>21</td>
</tr>
<tr>
<td>4</td>
<td>10</td>
<td>11</td>
</tr>
<tr>
<td>3</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

There was no difference in asthma control and IFR between the two groups suggesting that the Flow-Vu* did not affect drug delivery, therefore demonstrating clinical equivalency. Overall the IFR values were the required slow flows suggesting that the infants and their parents had retained the clinic training about the use of the spacer. This reflects the lack of any difference in asthma control between the two groups. Despite correct use of the AC in this group all parents preferred FV because it provides them with the reassurance that the correct inhalation procedure is performed and that their child has received their inhaled medication.


Purpose: We sought to determine parent preferences and patient outcomes to chronic pediatric asthmatics using either valved holding chamber (VHC): Aerochamber Plus*¨ (AC) or AC with Flow Vu*¨ indicator (ACF) regarding preference and outcomes of Emergency Department (ED) visits and hospitalizations. Methods: This is a pilot prospective randomized controlled trial of chronic pediatric asthmatics on inhaled corticosteroids that were assigned and educated on AC or ACF. Parents were instructed by nurses and interviewed by research assistants blinded on device assignment. Parents were contacted at day 3-5, 30 and 90 and were questioned on preference using a modification of the Patient Satisfaction and Preference Questionnaire (PASPQ) using the domains of overall performance and satisfaction, the Asthma Control Test (ACT) at 30, 60, and 90 days and an overall open ended question of preference ranked from 0-100. ED visits and hospitalizations were also reviewed during the study period. The ACF was compared to the AC at day 0 and day 90 for ACT score, preference score and PASPQ score using student's t test (two tailed, unpaired). The day 0 to day 90 scores (for PASPQ and ACT) were compared within the ACF and AC groups using student's t test (two tailed, paired). ACT scores were adjusted to a % that just took into account the patient's age.
Demographics of the AC and ACF groups were compared using students t test. Results: There were 48 patients who were assigned to ACF and 38 to AC. The mean age overall was 6.8 years, 61% of the study population was male, and there were no differences between age, sex race, ED visits and hospitalizations in the 2 groups. Both the ACF and AC group had PASPQ scores that improved significantly over the course of the study (Figure 1). There was no difference between spacer type PASPQ scores at either time point. Both the ACF and AC group had preference scores that were significantly higher over time. The ACF spacer had a higher preference score than the AC spacer. Both the ACF and AC group had asthma control scores that improved significantly over time (Figure 2). At day 0, the ACF group had worse control than the AC group. This difference was gone by day 90. Conclusion: VHGs are important adjuncts in pediatric asthma therapy that are well accepted. Both types of VHGs were associated with good asthma control over time. A visual indicator for inhalation was associated with improved patient satisfaction, which may have implications for compliance.

DEVELOPING A “UNIVERSAL” VALVED HOLDING CHAMBER (VHC) PLATFORM WITH ADDED PATIENT BENEFITS

INTRODUCTION: Crompton et al. recently observed that inhalers which provide reassurance to patients and their health care providers that inhalation is performed correctly should help improve patient compliance and control (1). As a result of such concerns, more attention is being being paid the incorporation of attributes into inhalers that assist the user in achieving both optimum compliance and consistent medication delivery (2,3). A progression of such changes with the AeroChamber Plus* (AC-Plus) VHC with mouthpiece (Trudell Medical International, London, Canada), has resulted in three variants being available depending on market needs and the local regulatory environment. These are: 1. the non-conducting AC-plus, 2. the non-conducting AC-plus with Flow-Vu* Inspiratory Flow Indicator (IFI) 3. the anti-static (charge dissipative) AC-plus with IFI. The present study was undertaken to provide users with indicative in vitro performance data based on 8 different hydrofluoroalkane-based formulations that are currently available and which may be prescribed for use with these VHGs.

MATERIALS AND METHODS: All measurements (n=5 VHGs/group) were made using an Andersen 8-stage cascade impactor equipped with USP/Ph.Eur. induction port and operated at 28.3 L/min ± 5%. The non-conducting variants were pre-treated in accordance with manufacturer instructions, thus ensuring that electrostatic charge was not a confounding factor. The anti-static VHGs were evaluated out-of-package, also to instructions. Measurements were made with no delay between pMDI actuation and the onset of sampling, and repeated with new devices, introducing a 2-s delay by means of a proprietary apparatus (4). The latter condition simulated use by an uncoordinated patient (5) in accordance with the Canadian standard for spacers and VHGs (6). The formulations evaluated are listed in Table 1. Recovery and assay for collected API was in each case undertaken by validated HPLC-UV spectrophotometric or fluorescence-based techniques.

<table>
<thead>
<tr>
<th>Trade Name, manufacturer</th>
<th>Active Pharmaceutical Ingredient (API)</th>
<th>Label claim mass/actuation (μg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Advair®, (GSK)</td>
<td>fluticasone propionate (FP)</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td>salmeterol xinafoate (SX)</td>
<td>25</td>
</tr>
<tr>
<td>Alvesco®, Nycomed</td>
<td>ciclesonide</td>
<td>100</td>
</tr>
<tr>
<td>Atrovent®, Boehringer Ingelheim</td>
<td>ipratropium bromide</td>
<td>20</td>
</tr>
<tr>
<td>Cenil®, Trinity Chiesi</td>
<td>beclomethasone dipropionate</td>
<td>100</td>
</tr>
<tr>
<td>Flovent HFA®, GSK</td>
<td>fluticasone propionate</td>
<td>125</td>
</tr>
<tr>
<td>Qvar®, TEVA</td>
<td>beclomethasone dipropionate</td>
<td>100</td>
</tr>
<tr>
<td>Symbicort®, AstraZeneca</td>
<td>budesonide (BUD)</td>
<td>80</td>
</tr>
<tr>
<td></td>
<td>formoterol fumarate (FF)</td>
<td>4.5</td>
</tr>
<tr>
<td>Ventolin HFA®, GSK</td>
<td>salbutamol (base equivalent)</td>
<td>100</td>
</tr>
</tbody>
</table>

RESULTS: Values of total emitted mass/actuation (TEM) and fine particle mass < 4.7 μm aerodynamic diameter/actuation (FPD<4.7μm), are summarised in Tables 2 and 3 respectively. FPD<4.7μm was selected as the metric most appropriate to define the portion of TEM likely to deposit beyond the oropharynx. Equivalent values determined in parallel experiments for the pMDI alone (no delay) are provided as benchmark data. The addition of any of the VHC variants resulted in a large reduction in TEM compared with the pMDI alone, as the result of the removal of most of the coarse fraction. The three AC plus VHC variants provided substantially equivalent values of TEM and FPM<4.7μm at each delay condition. This outcome was anticipated as the IFI is an external visual feedback aid and does not interfere with the aerosol pathway within the VHC. FPM<4.7μm value(s) ex VHC with 2-s delay were within ±25% of the equivalent benchmark data from the pMDI alone.

Total emitted mass / actuation (mean ± SD) for pMDI with and without AC PLUS VHC variants

<table>
<thead>
<tr>
<th></th>
<th>pMDI</th>
<th>Non conducting</th>
<th>Non conducting</th>
<th>Anti-Static AeroChamber</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>


### FPM<4.7µm (mean ± SD) for pMDI with and without AC PLUS VHC variants

<table>
<thead>
<tr>
<th></th>
<th>pMDI alone</th>
<th>Non conducting AeroChamber Plus* VHC</th>
<th>Non conducting AeroChamber Plus* VHC with Flow-Vu* Indicator</th>
<th>Anti-Static AeroChamber Plus* VHC with Flow-Vu* Indicator</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Delay (s)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Advair (FP)</strong></td>
<td>17.4 ± 2.1</td>
<td>26.6 ± 1.4</td>
<td>20.4 ± 1.8</td>
<td>23.0 ± 1.9</td>
</tr>
<tr>
<td><strong>Advair (SX)</strong></td>
<td>8.9 ± 1.3</td>
<td>13.9 ± 0.9</td>
<td>11.1 ± 1.0</td>
<td>12.2 ± 0.8</td>
</tr>
<tr>
<td><strong>Alvesco</strong></td>
<td>42.4 ± 3.8</td>
<td>68.1 ± 2.8</td>
<td>56.5 ± 2.6</td>
<td>64.5 ± 3.1</td>
</tr>
<tr>
<td><strong>Atrovent</strong></td>
<td>6.7 ± 0.4</td>
<td>10.0 ± 0.2</td>
<td>8.1 ± 0.8</td>
<td>9.8 ± 0.8</td>
</tr>
<tr>
<td><strong>Clenil</strong></td>
<td>30.9 ± 1.6</td>
<td>45.0 ± 2.3</td>
<td>33.2 ± 1.0</td>
<td>46.1 ± 1.7</td>
</tr>
<tr>
<td><strong>Flovent HFA</strong></td>
<td>46.2 ± 2.1</td>
<td>64.6 ± 3.0</td>
<td>47.3 ± 1.5</td>
<td>59.9 ± 2.5</td>
</tr>
<tr>
<td><strong>Qvar</strong></td>
<td>41.9 ± 2.2</td>
<td>67.2 ± 5.8</td>
<td>44.4 ± 6.9</td>
<td>71.6 ± 3.5</td>
</tr>
<tr>
<td><strong>Symbicort (BUD)</strong></td>
<td>43.2 ± 1.8</td>
<td>48.6 ± 2.2</td>
<td>42.0 ± 2.4</td>
<td>48.2 ± 2.9</td>
</tr>
<tr>
<td><strong>Symbicort (FF)</strong></td>
<td>2.7 ± 0.2</td>
<td>3.1 ± 0.2</td>
<td>2.6 ± 0.2</td>
<td>3.0 ± 0.1</td>
</tr>
<tr>
<td><strong>Ventolin HFA</strong></td>
<td>34.8 ± 1.4</td>
<td>48.6 ± 4.6</td>
<td>36.3 ± 1.8</td>
<td>42.3 ± 3.4</td>
</tr>
</tbody>
</table>

**CONCLUSIONS:** Two recent improvements made to the AC plus VHC to mitigate electrostatic charge and provide visual feedback to users have not affected in vitro performance compared with the original device. These devices also provide comparable delivery of fine particles to the pMDI alone in the likelihood that the user delays inhalation after actuating their inhaler.
IMPORTANCE OF FEEDBACK FEATURES

A number of researchers highlight the importance of a feedback mechanism on aerosol delivery devices.

- Devices which provide reassurance to patients and their physicians that inhalation is performed correctly should help to improve patient compliance and asthma control. (Crompton GK et al. The need to improve inhalation technique in Europe: A report from the Aerosol Drug Management Improvement Team. Respiratory Medicine 2006;100:1479-1494)

- Devices should be easy to use and incorporate multiple feedback mechanisms which reassure the patient that medication has been delivered. (Barnes P et al. Asthma Management: important issues. European Respiratory Review 2005;14,(97)147-151)

- Providing devices that are quick and easy to use effectively is a basic requirement for future devices, while some form of feedback does appear to have an impact on both regimen compliance and device delivery. (Everard ML. Playing the Game: Designing Inhalers for Pediatric Use. Respiratory Drug Delivery Europe 2007: 71-78)

- Practical technology for patients to self-assess technique would be of value (e.g. a feedback mechanism built into the inhaler device to confirm correct inhalation). (Papi A et al. Inhaler devices for asthma: a call for action in a neglected field. Eur Respir J 2011;37:982-985)

- It is important that feedback be provided to the patient or caregiver to confirm they are operating each device correctly. Steps critical to successful drug delivery should be a particular focus, such as ensuring a facemask seal to the face during use of a pMDI-valved holding chamber. (Mitchell JP et al. Developing Patient-Friendly Devices for Inhalation Therapy. Respiratory Drug Delivery Europe 2011:463-467)

The Flow-Vu* Inspiratory Flow Indicator is a valuable feedback tool

- The Flow-Vu* Indicator prevents the potential for large losses of medication due to facemask-to-face leakage, by guiding the caregiver or patient to seat the facemask on the face correctly – and at the same time, it indicates the optimum timing for inhaler actuation. (Michell JP et al. Letter to the Editors: Improving the odds that patients and caregivers will use inhalers correctly: a manufacturer's response. Prim Care Resp Respir J 2011; 20(2):219-220)

- A visual indicator for inhalation was associated with improved patient satisfaction, which may have implications for compliance. (Baig M et al. Flow-Vu Indicator in Valved Holding Chamber for Pediatric Asthma. American Academy of Pediatrics Conference 2013)


Purpose: We sought to determine parent preferences and patient outcomes to chronic pediatric asthmatics using either valved holding chamber (VHC): Aerochamber Plus* (AC) or AC with Flow Vu* indicator (ACF) regarding preference and outcomes of Emergency Department (ED) visits and hospitalizations. Methods: This is a pilot prospective randomized controlled trial of chronic pediatric asthmatics on inhaled corticosteroids that were assigned and educated on AC or ACF. Parents were instructed by nurses and interviewed by research assistants blinded on device assignment. Parents were contacted at day 3-5, 30 and 90 and were questioned on preference using a modification of the Patient Satisfaction and Preference Questionnaire (PASPQ) using the domains of overall performance and satisfaction, the Asthma Control Test (ACT) at 30, 60, and 90 days and an overall open ended question of preference ranked from 0-100. ED visits and hospitalizations were also reviewed during the study period. The ACF was compared to the AC at day 0 and day 90 for ACT score, preference score and PASPQ score using student's t test (two tailed, unpaired). The day 0 to day 90 scores (for PASPQ and ACT) were compared within the ACF and AC groups using student's t test (two tailed, paired). ACT scores were adjusted to a % that just took into account the patient's age. Demographics of the AC and ACF groups were compared using students t test. Results: There were 48 patients who were assigned to ACF and 38 to AC. The mean age overall was 6.8 years, 61% of the study population was male, and there were no differences between age, sex race, ED visits and hospitalizations in the 2 groups. Both the ACF and AC group had PASPQ scores that improved significantly over the course of the study (Figure 1). There was no difference between spacer type PASPQ scores at either time point. Both the ACF and AC group had preference scores that were significantly higher over time. The ACF spacer had a higher preference score than the AC spacer. Both the ACF and AC group had asthma control scores that improved significantly over time (Figure 2). At day 0, the ACF group had worse control than the AC group. This difference was gone by day 90. Conclusion: VHCs are important adjuncts in pediatric asthma therapy that are well accepted. Both types
of VHC were associated with good asthma control over time. A visual indicator for inhalation was associated with improved patient satisfaction, which may have implications for compliance.


The AeroChamber Plus* Spacer (Trudell Medical International, Canada) [AC] has recently been adapted to include a visual indicator that confirms the inhalation phase during use (AeroChamber Plus* Flow-Vu*, Trudell Medical International, Canada) [FV]. Movement of the indicator also confirms the required seal between the mask and the user’s face. The asthma control of infants with asthma, aged < 5 years, during routine use of these spacers has been compared and patient preference has been obtained. Ethical approval was obtained and all infants and their parents gave signed informed consent. All infants entered a 2 week run-in period using the AC. At visit 2 they were randomised to the AC or FV for the 12 week duration of this study. They returned after 6 and 12 weeks (visits 3 and 4). All infants were trained to use a gentle tidal breathing routine with their spacer. At each visit their asthma quality of life (ACQ; Juniper et al, Eur Respir J, 1996) was obtained by questioning each parent and their inhalation flow (IFR) was measured using the IN-Check Dial (Clement Clark International, UK). At visit 4 each FV parent was asked to rate their preference for the FV using a 5 point Likert scale (5=much better to 1=much worse). The FV was demonstrated to all AC parents and they were then asked the same preference rating. At visit 1 the mean (SD) age of the AC (n=38) and FV (n=38) infants was 3.3 (1.1) and 2.8 (0.9) years, respectively. A summary of the ACQ and IFR is presented in Table 1.

Table 1. Mean (SD) ACQ and IFR data

<table>
<thead>
<tr>
<th>Visit</th>
<th>ACQ</th>
<th>IFR (L/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>AC</td>
<td>FV</td>
</tr>
<tr>
<td>1</td>
<td>1.83 (0.88)</td>
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<td>2</td>
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<td>1.29 (0.90)</td>
<td>1.62 (0.92)</td>
</tr>
</tbody>
</table>

Two-way analysis of variance revealed no difference in the ACQ or the IFR between the visits and the two groups. Patient preference is shown in Table 2.

Table 2. Parent preference for FV (5= much better, 1= much worse)

<table>
<thead>
<tr>
<th>Likert Scale Rating</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
</tr>
<tr>
<td>FV</td>
</tr>
<tr>
<td>AC</td>
</tr>
</tbody>
</table>

There was no difference in asthma control and IFR between the two groups suggesting that the Flow-Vu* did not affect drug delivery, therefore demonstrating clinical equivalency. Overall the IFR values were the required slow flows suggesting that the infants and their parents had retained the clinic training about the use of the spacer. This reflects the lack of any difference in asthma control between the two groups. Despite correct use of the AC in this group all parents preferred FV because it provides them with the reassurance that the correct inhalation procedure is performed and that their child has received their inhaled medication.


Purpose: Poor inhaler compliance is recognized as needing to be addressed. The Flow-Vu* Inspiratory Flow Indicator (IFI) is a feedback aid for those using the AeroChamber Plus* Flow-Vu* Anti-Static VHC (Trudell Medical Inc., London, Ontario). Regulators require that the modification does not affect delivery of the therapeutically beneficial fine particle dose < 4.7 µm diameter from the inhaler. Methods: Measurements (n=5 VHCs/group) of fine particle mass for salbutamol (100 µg/actuation) were made using an Andersen 8-stage impactor equipped with Ph.Eur. induction port and operated at 28.3 L/min. Data were obtained for the pMDI alone and for the pMDI +VHC (2-second delay), simulating poor coordination. The movement of the IFI monitored airflow through the VHC and a proper seal of the mouthpiece in the apparatus. The VHCs were tested out-of-package in accordance with instructions. Recovery and assay for salbutamol was undertaken by HPLC-UV spectrophotometry. Results: Fine particle mass/actuation (FPM2s) for pMDI alone (mean±SD) was 34.8 ± 1.4 µg, compared with 33.2 ± 3.3 µg/actuation for the pMDI +VHC group. The IFI moved from the inhalation valve closed to open position immediately upon initiation of sampling. Conclusions: The IFI provided feedback on the delivery of this widely prescribed ‘rescue’ medication and
did not interfere with the new VHC, delivering substantially comparable FPM$_{2.5}$ to that from the pMDI alone. It should therefore aid patient compliance.


The AeroChamber Plus* with Flow-Vu* (FV) is a new version of the AeroChamber Plus* Spacer (AC). The only difference is that FV has a caregiver feedback mechanism that provides a visual indicator whose movement confirms correct inhalation technique and a secure seal between the facemask and the face. The latter is a critical factor in aerosol drug delivery to infants and children (Amirav et al., 2008). After a run-in period of 2 weeks children <5 years with uncontrolled or partially controlled asthma (GINA Guidelines) were randomized to receive their medication using either AC (n=9) or FV (n=10). Ethical approval was obtained and all children and their parents gave signed consent. Each parent completed the first 6 questions of the asthma control questionnaire (Juniper et al., 1999) on behalf of their child and the Paediatric Asthma Caregiver’s Quality of Life Questionnaire (Juniper et al., 1996) at 0, 6 and 12 weeks. The use of either AC or FV, with proper training, resulted in good asthma control. At the end of the study the FV group parents were asked to rank their preference compared to the AC whilst the FV was demonstrated to the AC group and the parents were asked about the 2 spacers. There was a clear preference among both groups for the FV device.

<table>
<thead>
<tr>
<th>FV Preference</th>
<th>Much Better</th>
<th>Better</th>
<th>No Difference</th>
<th>Worse</th>
<th>Much Worse</th>
</tr>
</thead>
<tbody>
<tr>
<td>FV</td>
<td>5</td>
<td>3</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>AC</td>
<td>6</td>
<td>2</td>
<td>2</td>
<td>0</td>
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</table>

The FV and AC provide similar asthma control but the Flow-Vu* feature was preferred by parents. Devices which provide reassurance to patients and their physicians that inhalation is performed correctly should help to improve patient compliance and asthma control (Crompton et al., 2006).


Delivery of inhaled medication to infants/small children by VHC-facemask can be difficult to verify. An external visual aid (Flow-Vu*) is available with the AeroChamber Plus* (Trudell Medical International) VHCs as an inspiratory flow indicator (IFI) to aid compliance with instructions for use. We report an *in vitro* study in which delivery of salbutamol (Ventolin®; 100µg/actuation, GSK plc) was measured using infant and child models (ADAM-II), in which the soft facial tissues are modeled where the facemask makes contact. The facemask was applied with an appropriate force of 1.6 kg, and tidal breathing was simulated (tidal volume (V$_t$) 50 ml, 30 bpm, 25% duty cycle - VHC-infant facemask; V$_t$ = 155 ml; 25 bpm, 33% duty cycle - VHC-child facemask (n=5 devices/group)). Total emitted mass (TEM) of salbutamol was collected by filter located behind the lips after 1, 2, 3, 4 and 5 inhalations.

| TEM (mean SD; µg) for AeroChamber Plus* VHCs with Flow-Vu* IFI |
|-----------------|----------------|---------------|----------------|----------------|----------------|
| Number of Inhalations | 1      | 2              | 3              | 4              | 5              |
| Infant          | 5.8 ± 2.2 | 13.0 ± 4.0     | 13.8 ± 3.8     | 14.5 ± 3.2     | 15.6 ± 3.6     |
| Child           | 15.9 ± 3.7| 17.6 ± 4.6     | 20.1 ± 3.5     | 20.8 ± 2.9     | 22.6 ± 4.2     |

At least two inhalations were required to achieve consistent medication delivery from the VHC-infant facemask. The first inhalation was sufficient to achieve similar consistency with the VHC-child facemask. However, these tests were undertaken with a well-fitting facemask and no leakage. Manufacturer instructions indicate 5-inhalations be taken as a precaution. The IFI validates an effective seal between facemask-face as well as confirms the number of inhalations, assisting in compliance with instructions.


Delivery of inhaled medication to infants by valved holding chamber (VHC) with facemask may require more than one inhalation to empty the VHC because tidal volumes are typically smaller than chamber capacity. This study investigated the correlation between movement of an integrated inspiratory flow indicator (IFI) as a caregiver feedback aid for a VHC-facemask, number of inhalations and mass of medication, simulating use by a 6-9 month infant (tidal volume (V$_t$) = 50-ml; duty cycle = 25%; 30
cycles/min). Anti-static AeroChamber Plus* VHCs incorporating the IFI feature, with infant facemask (n=5/group, 3 replicates/device; Trudell Medical International, London, Canada) were coupled to a breathing simulator (ASL5000 test lung, IngMar Medical, Pittsburgh, PA, USA). The VHCs were prepared as per manufacturer instructions and the facemask of the device on test was fitted to the ADAM-II flexible infant face model with a clinically appropriate force of 1.6 kg. Aerosol capture took place using an electret filter positioned behind the lips of the face model. Delivery of medication was evaluated from two different pressurized metered dose inhaler formulations likely to be used with paediatric patients (Flovent HFA® 44; 44 μg fluticasone propionate (FP) delivered ex-actuator and Ventolin HFA®; 90 μg salbutamol base equivalent (SAL) delivered ex-actuator, both from GSK plc. One actuation was delivered to the VHC at the onset of inhalation, and the filter removed after 1 complete breathing cycle, observing the movement of the IFI to confirm inhalation valve opening. This procedure was subsequently repeated by removing the filter after 2, 3, 4, 5 and 6 breathing cycles. Assay for FP or SAL was undertaken by HPLC-UV spectrophotometry. During these measurements, the IFI of each device was observed to move in synchrony with valve opening on all occasions, confirming that the facemask sealed onto the face model without leakage of ambient air into the mask during the inspiratory phase of each breathing cycle. Emitted mass after the first breathing cycle (EM1) was 2.1 ± 0.7 μg (FP) and 5.8 ± 2.2 μg (SAL); substantially lower than the corresponding values after 6 cycles (EM6), being 9.0 ± 2.1 μg (FP), and 15.9 ± 3.1 μg (SAL) [paired t-test for each formulation; p < 0.001]. After 2 breathing cycles, values of EM2 (6.9 ± 2.0 (FP) and 13.0 ± 4.0 μg (SAL)), though significantly greater than their corresponding EM1 values [p ≤ 0.002], were still noticeably lower than the corresponding EM6 value for FP (p = 0.028), and barely statistically insignificant for SAL (p = 0.063). After 3 inhalations, EM3 increased further to 7.6 ± 2.0 μg (FP) and 13.8 ± 3.8 μg (SAL), and thereafter were close to the corresponding EM6 values, indicating emptying of the VHC had taken place. We conclude that at least two successive inhalations are required to achieve optimum medication delivery for the ‘infant’ condition under optimum conditions with a well fitted facemask with no leakage. The IFI is an important feature which validates that the facemask is properly sealed to the infant’s face and also confirms the number of inhalations that take place, thereby optimizing the therapeutic dose. Clinical studies are recommended to evaluate the benefit of this aid for the delivery of inhaled medication by VHC to this age group.
IMPORTANT OF FACEMASK SEAL


BACKGROUND: In order to improve patient compliance, the use of charge dissipative materials in VHC construction is becoming the standard of care. A facemask is required as the interface between patient and VHC for young children who cannot breathe through a mouthpiece. Recent studies have emphasized that a well-fitting facemask is critical for optimal drug delivery. We report a laboratory based comparison of aerosol drug delivery between two 'antistatic' VHCs under simulated breathing conditions, using a anatomically correct infant face-upper airway model (ADAM-III, Trudell Medical International (TMI)). METHODS: Delivery of fluticasone propionate (FP; 44 μg/actuation GSK) as evaluated via anti-static AeroChamber Plus® VHC with Flow-Vu® IFI/infant mask (AC-Plus, MMC) and OptiChamber Diamond† VHC/LiteTouch† small-mask (OD, Philips) (n=5 devices/group). Tidal-breathing (tidal-volume (Vt)= 155-mL, duty-cycle=33%, rate= 25-breaths/min) was simulated with an Ingmar ASL 500 test lung. Each facemask was applied to the face with the same clinically-appropriate force (1.6 kg). FP was recovered from the pMDI mouthpiece, VHC, facemask, face and airway of the model as well as the filter at the carinal exit of the model airway (equivalent to lung dose). Delivered mass of FP (DMFP) was quantified by HPLC. RESULTS: DMFP (mean±SD) was significantly greater from AC-plus (11.6±1.4 μg) than OD (7.2±1.4 μg) (unpaired t-test, p=0.002). This difference was largely due to the FP lost on the facemask of the OD facemask (8.8±0.9 μg) compared to that of the AC-plus (4.3±0.3 μg). CONCLUSION: While other factors such as facemask dead volume and device design are important factors in device performances, decreased aerosol delivery from the OD is explicable in terms of leakage between facemask and face, or choice of anti static materials, supported by higher deposition in its facemask. Clinicians should be aware that each VHC-pMDI combination is unique.


Background & Objectives: Leakage between facemask-and-face may result in medication loss by valved holding chamber (VHC)-facemask (1). Our study evaluated how an inspiratory flow indicator (IFI) can be used to avoid leakage. Methods: An infant face with realistic soft-tissue modeling (ADAM-III, Trudell Medical International (TMI), London, Canada (2)) was used to evaluate delivery of fluticasone propionate (FP; 50 μg/actuation, GSK (Canada)) via anti-static AeroChamber Plus® VHC with Flow-Vu® IFI/infant mask (AC-Plus, TMI) or OptiChamber® Diamond® VHC/LiteTouch® small-mask (OD, Philips-Respironics, Parsipanny, NJ, USA) (n=5 devices/group), simulating tidal-breathing (tidal-volume (Vt)=155-mL, duty-cycle=33%, rate= 25-breaths/min). Each facemask was applied to the face with the same clinically-appropriate force (1.6 kg). The IFI of the AC-plus was observed to be moving. FP was recovered from the nasopharynx and base (lung dose) of the model, and delivered mass (DMFP) quantified by HPLC-spectrophotometry as % label claim (LC). Findings: DMFP (mean±S.D.) was significantly greater from AC-plus (25.8±5.3%LC) than OD (17.0±3.7%LC) (unpaired t-test, p=0.019). FP on the facemask of the AC-plus (6.2±1.9% LC), was slightly smaller than that determined with the OD facemask (9.9±2.6%). Discussion/Conclusion: Vt was set larger than normal in order to detect facemask-to-face leakage more precisely. Leakage was eliminated with the AC-plus, by observing IFI movement. However, inasmuch as the OD does not have an IFI, it was not possible to do more than ensure that its facemask was applied with the same force to the model. Decreased aerosol delivery from the OD is explicable in terms of leakage between facemask and face, supported by higher deposition in its facemask.


Inhaled drugs are frequently given to infants and young children with a pressurized metered-dose inhaler (pMDI) attached to a valved-holding chamber (VHC) with face mask. In young children and infants who cannot breathe through a mouthpiece, the face mask serves as the interface between the patient and the VHC. Although the mask interface is one of the most important factors determining the dose of medication delivered from the VHC to the nose and mouth in these patients, its optimal characteristics are not well known. Recent studies clearly identify several face mask factors that determine the success or failure of drug delivery with these devices. This review summarizes the most important features of an optimal mask design such as: face seal/leak, volume of dead space, contour, flexibility, transparency, weight and cost. By optimizing these characteristics it should be possible to improve mask design. This will maximize the magnitude and reduce the variability of the dose presented to the respiratory tract while making the mask more comfortable and patient/caregiver-friendly.

BACKGROUND: Valved Holding Chambers (VHCs) with facemask as patient interface are an important adjunct in delivery of medication particularly to infants. Leakage between facemask and face can result in severe loss of medication because there is no pressure source to drive the aerosol towards the patient once the inhaler has been actuated. Laboratory performance testing therefore needs to be undertaken with the facemask in place. METHODS: We evaluated the effect of two different but similar-sized VHCs (AeroChamber Plus* with small mask - Monaghan Medical Corp.) and Vortex® with Babywhirl™ mask - PARI Respiratory Equipment Inc., Midlothian, PA) on delivery of albuterol (2-actuations of Ventolin® HFA, GlaxoSmithKline) to an oral-breathing 9-12 month anatomical infant face model with simulated in-vivo facial surfaces where the mask was applied with a clinically relevant force of 1.6 kg. The VHCs were prepared in accordance with manufacturer instructions and evaluated at a flow rate of 4.9 L/min, sampling the aerosol by low-flow impactor to determine total emitted mass (TEM) and fine particle fraction (FPF) < 4.7 μm aerodynamic diameter. These conditions were considered to be representative of a 6-12 month old infant having a 50th percentile body weight range from 7.5 to 9.9 kg. Recovered albuterol was determined by HPLC-UV spectrophotometry. RESULTS: TEM/kg (n = 5 devices, mean ± SD) from the AeroChamber Plus* VHCs was in the range 1.6 to 2.2 μg/kg, comparable with data provided in the Patient Information Leaflet for this formulation with infant users, and FPF was 73%. No leakage was detected between facemask and face (100% flow to the face via the VHC). In contrast, a perfect seal could not be achieved between the Vortex® VHCs and associated wider facemask with the model face (97% of the flow bypassed the VHC under best conditions achievable), so that TEM/kg from these devices ranged from 0.07 to 0.09 μg/kg, with FPF of 100%. CONCLUSION: The Aerosol-Delivery-to-Anatomical-Model (ADAM-II) face technique allows laboratory testing of VHC-facemask to be evaluated as a complete system. The data obtained confirm that no leakage between facemask and face must be achieved for reliable delivery of medication from these devices.


Valved holding chambers (VHCs) are widely prescribed for use with pressurized metered dose inhalers (pMDIs) for the treatment of respiratory disease by aerosol therapy. The facemask is the preferred patient interface for use by infants and small children, as well as by geriatric patients, due primarily to poor coordination skills. However, care is required in the design of the facemask-VHC system to optimize the delivery of medication. In particular, it is essential to achieve an effective mask-to-face seal and to minimize the volume of dead space. It is also important to ensure that the fit of the facemask is comfortable to the patient when applied with sufficient force to create a seal. We review each of these design principles and their application in the evolution of a range of VHCs from the same family of devices during the past fifteen years. We also examine the various methods available for evaluating VHC-facemasks as a system, recommending where future work might be directed.

FORCE-DEPENDENT STATIC DEAD SPACE OF FACE MASKS USED WITH HOLDING CHAMBERS. Shah SA, Berlinski AB, Rubin BK. Respir Care 2006;51(2):140-144.

BACKGROUND: Pressurized metered-dose inhalers with valved holding chambers and masks are commonly used for aerosol delivery in children. Drug delivery can decrease when the dead-space volume (DSV) of the valved holding chamber is increased, but there are no published data evaluating force-dependent DSV among different masks. METHODS: Seven masks were studied. Masks were sealed at the valved holding chamber end and filled with water to measure mask volume. To measure mask DSV we used a mannequin of 2-year-old-size face and we applied the mask with forces of 1.5, 3.5, and 7 pounds. Mask seal was determined by direct observation. Intra-brand analysis was done via analysis of variance. RESULTS: At 3.5 pounds of force, the DSV ranged from 29mL to 100mL, with 3 masks having DSV of < 50mL. The remaining masks all had DSV > 60 mL. At 3.5 pounds of force, DSV percent of mask volume ranged from 33.7% (AeroChamber*, p <0.01 compared with other masks) to 100% (Pocket Chamber). DSV decreased with increasing force with most of the masks, and the slope of this line was inversely proportional to mask flexibility. Mask fit was 100% at 1.5 pounds of force only with the AeroChamber* and Optichamber. Mask fit was poorest with the Vortex, Pocket Chamber, and BreatheRite masks. CONCLUSION: Rigid masks with large DSV might not be suitable for use in children, especially if discomfort from the stiff mask makes its use less acceptable to the child.
OBJECTIVE: Masks are an essential interface between valved holding chambers (VHCs), or spacers, and a small child’s face for providing aerosol therapy. Clinical experience suggests that many young children do not cooperate with the VHC treatment or tolerate a mask of any kind. This might impair the mask-face seal and reduce the dose delivered to the child. The objective of this study was to evaluate the ability of parents to provide a good mask-face seal in infants and toddlers using 3 masks provided with commonly used pediatric VHCs and compare this with the seal obtained with the Hans Rudolph pediatric anesthesia mask. METHODS: A preliminary in vitro filter study was conducted to validate the assumption that reduced ventilation as a result of increased facemask leak reduces the drug aerosol dose delivered to the mouth. Facemask leak then was studied in vivo for NebuChamber, AeroChamber*, BabyHaler, and Hans Rudolph masks by measuring ventilation with an in-line pneumotachograph while the facemask was held in place by experienced parents who were asked to demonstrate how they deliver medication to their children without any additional instruction. Thirty children (mean age: 3.2 +/- 1.4 years) performed 4 repeat studies with each mask. The first 10 patients performed the tests once again within 1 month. On the second occasion, the parents were coached continuously and encouraged to hold the mask tightly against the child’s face. RESULTS: The AeroChamber* and Hans Rudolph masks provided the best seal as reflected in the magnitude of the ventilation measured through them. The NebuChamber provided the poorest seal, with 45% less ventilation than the AeroChamber* and Hans Rudolph masks. There was considerable intraindividual variability for all masks (24% to 48%); however, the variability with the NebuChamber mask was 2-fold greater than the other masks. All ventilatory volumes during the coached session were significantly greater than during the uncoached session. Variability during the coached session was significantly less (except for the BabyHaler, which remained unchanged). CONCLUSIONS: VHCs with masks designed for use with small children may provide a poor seal with the face, leading to reduced or more variable dose delivery. The facemask seal is critical for efficient aerosol delivery to infants and young children, and this should be stressed to parents.
AECROCHAMBER PLUS* VHC - PERFORMANCE WITH DIFFERENT MDI FORMULATIONS

Alvesco® (Ciclesonide) Nycomed™


Objective: To evaluate the efficacy and safety of three doses of ciclesonide (with or without spacer) in children with persistent asthma. Patients and methods: This was a multicentre, double-blind, placebo-controlled, 12-week study of ciclesonide 40, 80 or 160 μg (once daily pm). Children (6–11 years) were randomised 1:1 to treatment via a metered dose inhaler (MDI) or MDI plus spacer (AeroChamber Plus*). The primary variable was change from baseline in mean morning peak expiratory flow (PEF). Secondary variables included: time to first lack of efficacy (LOE), asthma control, forced expiratory volume in 1 s (FEV1), asthma symptom score and quality of life (QoL). Safety assessments included: adverse events (AEs), urinary cortisol excretion and body height. Results: In total, 1073 children received treatment. At endpoint, mean morning PEF significantly improved with all doses of ciclesonide vs. placebo. There was no difference over placebo in time to first LOE, but ciclesonide was superior to placebo on asthma control, symptom score, FEV1 and QoL. There were no differences between the spacer or non-spacer subgroups. The incidences of AEs were comparable between treatment groups (approximately 35%) and there were no between-group differences in body height or urinary cortisol. Conclusions: Ciclesonide 40–160 μg once daily is effective and well tolerated in children with persistent asthma; its efficacy and safety are unaffected by the use of a spacer.


Background: Ciclesonide is an inhaled corticosteroid administered by a metered dose inhaler (MDI) to treat bronchial asthma. After inhalation, the inactive ciclesonide is converted by esterases in the airways to active metabolite desisobutyryl-ciclesonide (des-CIC). Aim: To compare the pharmacokinetic (PK) parameters of des-CIC in children after administration of therapeutic dose of ciclesonide with and without spacer (AeroChamber Plus*). Methods: Open-label, 3 period, cross over, repeated dose, PK study in 37 children with mild to moderate stable asthma (age: 6–11 y; body weight: 20–53 kg). During each 7-day treatment period, ciclesonide was inhaled once in the morning; A) 160 μg MDI with spacer, B) 80 μg MDI with spacer, and C) 160 μg MDI without spacer. Serum PK parameters of ciclesonide and des-CIC were determined on Day 7 of each period. The primary PK parameters were the AUC(0-∞) and Cmax for des-CIC. Results: Inhaling ciclesonide with spacer led to a dose proportional systemic exposure (AUC(0-∞)) of des-CIC (0.316 μg*h/L for 80 μg and 0.663 μg*h/L for 160 μg). The dose-normalized systemic exposure for des-CIC (based on AUC(0-∞)) was 27% higher after inhalation of ciclesonide 80 μg or 160 μg with spacer than without spacer; the corresponding Cmax values for des-CIC were, respectively, 63% and 55% higher with spacer. No clinically relevant abnormalities or adverse drug reactions were observed. Conclusions: Inhalation of therapeutic ciclesonide dose with spacer led to a slight increase in the systemic exposure of des-CIC, which does not warrant dose adjustment.


Inhaled corticosteroids (ICS) are recommended as first-line treatment for adults and children with persistent asthma. The Global Initiative for Asthma recommends that patients taking medium- or high-dose ICS delivered by metered-dose inhalers (MDIs) should use a spacer device. This randomized, open-label, 12-week, non-inferiority study compared the efficacy and safety of ciclesonide 160 μg once daily delivered via hydrofluoroalkane-MDI alone (CIC160) or with a spacer (either an AeroChamber Plus* [CIC160P] or an AeroChamber MAX* [CIC160M]) in patients with persistent asthma. The primary efficacy variable was change in forced expiratory volume in 1 s (FEV1) from baseline to study end. Significant improvements in FEV1 were observed from baseline to study end in each treatment group; least squares mean change from baseline ranged between 0.32 and 0.34L in the per-protocol (PP) analysis and similar results were observed for the intention-to-treat (ITT) analysis (p < 0.0001 for all). Non-inferiority of CIC160P and CIC160M to CIC160 was observed for both PP and ITT populations (p < 0.001 [one-sided]). In all groups, daily asthma symptom scores were reduced to 0 and significant reductions were observed in rescue medication use at study end (p < 0.0001 versus baseline for all). Ciclesonide was well tolerated in all groups and no cases of oral candidiasis were reported. Morning serum cortisol levels significantly increased in all groups from baseline to study end (p ≤ 0.0389), with no significant between-treatment differences. In patients with persistent asthma, ciclesonide was shown to have similar efficacy and tolerability when administered via MDI alone or with a spacer.
Ciclesonide is an onsite-activated inhaled corticosteroid (ICS) for the treatment of asthma. This study compared the efficacy, safety and effect on quality of life (QoL) of ciclesonide 160 microg (ex-actuator; nominal dose 200 microg) vs. budesonide 400 microg (nominal dose) in children with asthma. Six hundred and twenty-one children (aged 6-11 yr) with asthma were randomized to receive ciclesonide 160 microg (ex-actuator) once daily (via hydrofluoroalkane metered-dose inhaler and AeroChamber Plus® spacer) or budesonide 400 microg once daily (via Turbohaler(R)) both given in the evening for 12 wk. The primary efficacy end-point was change in forced expiratory volume in 1 s (FEV(1)). Additional measurements included change in daily peak expiratory flow (PEF), change in asthma symptom score sum, change in use of rescue medication, paediatric and caregiver asthma QoL questionnaire [PAQLQ(S) and PACQLQ, respectively] scores, change in body height assessed by stadiometry, change in 24-h urinary cortisol adjusted for creatinine and adverse events. Both ciclesonide and budesonide increased FEV(1), morning PEF and PAQLQ(S) and PACQLQ scores, and improved asthma symptom score sums and the need for rescue medication after 12 wk vs. baseline. The non-inferiority of ciclesonide vs. budesonide was demonstrated for the change in FEV(1) (95% confidence interval: -75, 10 ml, p = 0.0009, one-sided non-inferiority, per-protocol). In addition, ciclesonide and budesonide showed similar efficacy in improving asthma symptoms, morning PEF, use of rescue medication and QoL. Ciclesonide was superior to budesonide with regard to increases in body height (p = 0.003, two-sided). The effect on the hypothalamic-pituitary-adrenal axis was significantly different in favor of ciclesonide treatment (p < 0.001, one-sided). Both ciclesonide and budesonide were well tolerated. Ciclesonide 160 microg once daily and budesonide 400 microg once daily were effective in children with asthma. In addition, in children treated with ciclesonide there was significantly less reduction in body height and suppression of 24-h urinary cortisol excretion compared with children treated with budesonide after 12 wk.


Background: Ciclesonide is an inhaled corticosteroid that provides safe and effective control of patient asthma. Ciclesonide is administered as an aerosol solution in a metered-dose inhaler, using hydrofluoroalkane-134a as a propellant. It is activated in the lung to form its only active metabolite, desisobutyryl-ciclesonide (des-CIC). A spacer may be used in combination with the hydrofluoroalkane metered-dose inhaler (HFA-MDI) to maintain inhaled corticosteroid delivery to the lung in patients with poor inhalation technique. Objective: To determine if the pharmacokinetics of des-CIC and ciclesonide are altered when a spacer is used for ciclesonide inhalation. Methods: A randomized, open-label, 2-period crossover, single-center pharmacokinetic study was conducted in 30 patients with asthma (forced expiratory volume in 1 second ≥ 70% predicted). A single dose of ciclesonide (320 µg ex-actuator; equivalent to 400 µg ex-valve) was administered via the HFA-MDI with and without an AeroChamber Plus® spacer (Trudell Medical International, London, ON, Canada). Serum concentrations of ciclesonide and des-CIC were measured before inhalation and at various intervals until 14 hours after treatment using high-performance liquid chromatography with tandem mass spectrometric detection. Results: The pharmacokinetic properties of the active metabolite, des-CIC, were equivalent after inhalation of ciclesonide with and without the AeroChamber Plus® spacer. Point estimates and 90% confidence intervals (CIs) for the ratio of des-CIC pharmacokinetic properties in the presence or absence of a spacer were within the conventional bioequivalence range of 0.80-1.25 (area under the serum concentration time curve from time zero to infinity 0.96 [0.85, 1.07]; peak serum concentration 1.05 [0.94, 1.18]; elimination half-life 1.04 [0.92, 1.18]). Furthermore, there was no relevant difference in the point estimate and 90% CI of the difference of the time to reach peak serum concentration of des-CIC with or without a spacer. Conclusion: The AeroChamber Plus® spacer did not influence the pharmacokinetics of the pharmacologically active des-CIC. Thus, systemic exposure to the active metabolite is similar when ciclesonide is inhaled with or without a spacer.

Beclomethasone Diproionate


AIM: The purpose of this study was to determine whether local anti-inflammatory therapy with inhaled beclomethasone dipropionate is effective in the outpatient management of acute viral croup. METHODS: Children six months to five years of age, presenting to the Emergency Department (ED) with a croup score of at least 2 participated in the study. All children were assigned in a randomised double-blind fashion to receive either nebulized L-epinephrine (LE), a single intramuscular injection of dexamethasone (D) 0.6 mg/kg, or inhaled beclomethasone dipropionate (BD) 200 mg, via aerochamber. Croup score (CS), heart rate (HR), blood pressure, respiratory rate (RR) and oxygen saturation were recorded at study entry and at 15, 30, 60, 90

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and 120 minutes after treatment. RESULTS: Sixty-four patients were enrolled into the study. Significant improvement of the croup score was noticed at the end of observation time in all groups. The LE group showed significant improvements of CS, HR and RR in comparison to the other two groups. Inhaled BD was as effective as intramuscular D in the treatment of mild to moderate croup in the ED. CONCLUSION: The use of inhaled beclomethasone in the outpatient management of croup was associated with a significant reduction in the severity of illness within 24 h after treatment.

Clenil® HFA (Beclomethasone Dipropionate) Chiesi / Vectura


Background: To which extent volume spacers may influence systemic activity of inhaled beclomethasone dipropionate (BDP) has not been evaluated. Aim: To assess whether the AeroChamber Plus(™) spacer is equivalent to the Volumatic(™) spacer for administration of inhaled hydrofluorokane 134a propelled BDP in terms of lower leg growth rate (LLGR). Patients and methods: Prepubertal children with mild asthma (n=26, ages 6-14 years) were included in a 3-time periods of 2 weeks duration randomised double-blind cross-over study with a single-blind placebo run-in and 2 wash-out periods. LLGR was measured with the knemometer. Interventions were inhaled BDP hydrofluorokane 134a pMDI 100 μg and 200 μg b.i.d. with the AeroChamber Plus and 200 μg b.i.d. with the Volumatic spacer. Results: BDP 200 μg b.i.d. from the AeroChamber Plus was non-inferior to BDP 200 b.i.d. from the Volumatic spacer as the lower margin of confidence interval of the difference between treatments (-0.18 to 0.13 mm/week) was greater than the pre-specified lower limit for non-inferiority (-0.20 mm/week). UFC/creatinine data showed no statistically significant variations. Conclusion: The systemic activity of BDP via the Volumatic(™) and AeroChamber Plus(™) spacers is similar. The AeroChamber Plus spacer may be used in children without risk of increasing systemic activity of BDP.

SHORT-TERM LOWER LEG GROWTH IN 5- TO 11-YEAR-OLD ASTHMATIC CHILDREN USING BECLOMETHASONE DIPROPIONATE INHALERS WITH CHLOROFLUOROCARBON OR HYDROFLUOROKANNE PROPPELLANTS: A 9-WEEK, OPEN-LABEL, RANDOMIZED, CROSSOVER, NONINFERIORITY STUDY. Wolthers OD, Walters EG. Clinical Therapeutics 2011;33(8):1069-1076.

Background: Beclomethasone dipropionate–hydrofluorokane (BDP-HFA) is a non–chlorofluorocarbon (CFC)-propelled metered dose inhaler. Data is needed to support the registration of BDP-HFA in pediatric populations for countries in the European Union. Objective: The aim of the study was to assess short-term lower leg growth in children with asthma during treatment with BDP-HFA 100 μg BID compared with BDP-CFC 200 μg BID. Methods: Children with asthma were included in this open-label, randomized, crossover study with 2-week run-in, active treatment, and washout periods. Lower leg length was measured every second week. As a secondary outcome parameter, 24-hour urine was collected for assessment of free cortisol. Interventions were inhaled BDP-HFA 100 μg BID with AeroChamber Plus spacer and BDP-CFC 200 μg BID with Volumatic spacer. Results: In 63 patients with asthma aged 5 to 11 years, BDP-HFA 100 μg BID was not inferior to BDP-CFC 200 μg BID, as the lower margin of CI (−0.03 to 0.10 mm/wk) of the estimated difference (0.03 mm/wk) was greater than the prespecified lower limit for noninferiority of −0.12 mm/wk. Mean (SD) lower leg growth rate during run-in, BDP-HFA 100 μg BID, and BDP-CFC 200 μg BID was 0.36 (0.17), 0.27 (0.21), and 0.23 (0.18) mm/wk, respectively (BDP-HFA estimate of difference, −0.09 [95% CI, −0.16 to −0.03 mm/wk; P < 0.01]; BDP-CFC estimate of difference, −0.13 [95% CI, −0.19 to −0.06 mm/wk; P < 0.001]). No statistically significant differences were seen in urinary free cortisol assessments. Eight and 6 mild to moderate adverse events in 10 children were reported during treatment with BDP-HFA and BDP-CFC, respectively. One event in each group was judged to be probably related to the study medication; no others were judged to be related. Conclusions: No statistically significant differences were found in lower leg growth between BDP-HFA 100 μg BID with AeroChamber Plus spacer and BDP-CFC 200 μg BID with Volumatic spacer during 2-week treatment. Evidence of differences in systemic activity between the treatments was not found.


Introduction: Factors affecting dose delivery from pMDIs fitted with add-on devices include formulation, device design (e.g., materials, size, incorporation of a non-return valve), cleaning procedures and use-mode. Spacer-mode involves a conventional press-and-breathe manoeuvre whilst inhaling through the mouthpiece of the pMDI-device assembly. The spacer creates a longer path-length, allowing more time for propellant evaporation and slowing the cloud to facilitate lung access. Use in holding-chamber mode requires the device to have a non-return valve and discharging the dose into the chamber where it is held for a period before being inhaled, eliminating the need for press-and breathe co-ordination. This mode also permits the patient to carry out repeated inhalations from the same dose. We were interested in comparing the effects of different add-on devices and
their mode of use because the large size of some holding chamber devices may deter user acceptability. This study compares the dose delivered when beclomethasone dipropionate (BDP) HFA solution type pMDIs were used in conjunction with both small and large volume devices in the two modes. Conclusions: These in vitro results would imply that, when used by patients in association with AeroChamber-Plus, the drug delivery performance for Modulite-BDP pMDIs could be similar to that obtained with Volumatic up to holding times of at least 5s for all three product strengths and up to 10s for the 50µg and 100µg dose strengths.

**Combivent® (Salbutamol & Ipratropium Bromide) Boehringer Ingelheim™ Pharmaceuticals Inc.**


Purpose: To compare the performance of a new small volume VHC (AeroChamber Plus® - 149 ml - Monaghan Medical Corp, Plattsburgh, NY) with a larger VHC (OptiChamber® - 218 ml, Respironics, Cedar Grove, NJ) for the delivery of a combination b-agonist/anticholinergic formulation for the treatment of COPD (Combivent®: 103 µg/dose albuterol sulfate (SAL) + 18 µg/dose ipratropium bromide (IPR) ex actuator mouthpiece (Boehringer-Ingelheim Pharmaceuticals Inc). Methods: Fine particle dose (FPD, particles < 4.7 μm aerodynamic diameter) and total emitted dose (ED) were determined for each VHC (n = 5 devices for each group) by Andersen 8-stage impactor, operating at 28.3 ± 0.5 L/min. The quantities of SAL and IPR components collected in the impator were determined by HPLC-UV spectrophotometry. Results: FPD and ED (% of label claim dose) were as follows. AeroChamber Plus® VHC: SAL - 56.2 ± 3.6%, 59.7 ± 4.1%; IPR - 47.0 ± 4.5%, 51.0 ± 5.0%, OptiChamber® VHC: SAL - 41.1 ± 3.3%, 43.5 ± 3.1%; IPR - 35.0 ± 6.0%, 38.0 ± 7.0%. The AeroChamber Plus® VHCs delivered significantly more of both components as either FPD or ED [un-paired t-test, p £ 0.001]. Conclusions: Increased chamber volume does not necessarily correlate with improved FPD or ED in this fromulation. Other considerations, such as internal geometry and inhalation valve design contribute to performance by controlling internal aerosol losses. Clinical Implications: The differences observed with these specific devices may have significant clinical implications that require further study.

**Flovent® (Fluticasone Propionate) GSK™ Inc.**


BACKGROUND: In order to improve patient compliance, the use of charge dissipative materials in VHC construction is becoming the standard of care. A facemask is required as the interface between patient and VHC for young children who cannot breathe through a mouthpiece. Recent studies have emphasized that a well-fitting facemask is critical for optimal drug delivery. We report a laboratory based comparison of aerosol drug delivery between two ‘antistatic’ VHCs under simulated breathing conditions, using an anatomically infant face-upper airway model (ADAM-II, Trudell Medical International (TMI)). METHODS: Delivery of fluticasone propionate (FP; 44 μg/actuation GSK) as evaluated via anti-static AeroChamber Plus® VHC with Flow-Vu® IFI/infant mask (AC-Plus, MMC) and OptiChamber Diamond® VHC/LiteTouch® small-mask (OD, Philips) (n=5 devices/group). Tidal-breathing (tidal-volume (Vt)= 155-mL, duty-cycle=33%, rate= 25-breaths/min) was simulated with an Ingmar ASL 500 test lung. Each facemask was applied to the face with the same clinically-appropriate force (1.6 kg). FP was recovered from the pMDI mouthpiece, VHC, facemask, face and airway of the model as well as the filter at the carinal exit of the model airway (equivalent to lung dose). Delivered mass of FP (DMFP) was quantified by HPLC. RESULTS: DMFP (mean±SD) was significantly greater from AC-plus (11.6±1.4μg) than OD (7.2±1.4μg) (unpaired t-test, p=0.002). This difference was largely due to the FP lost on the facemask of the OD facemask (8.8±0.9μg) compared to that of the AC-plus (4.3±0.3 μg). CONCLUSION: While other factors such as facemask dead volume and device design are important factors in device performances, decreased aerosol delivery from the OD is explicable in terms of leakage between facemask and face, or choice of anti static materials, supported by higher deposition in its facemask. Clinicians should be aware that each VHC-pMDI combination is unique.


RATIONALE: Adoption of materials to mitigate medication losses due to electrostatic charge accumulation on VHCs has been associated with improved reliability of output. We investigated if this change has resulted in similar in vitro performance, simulating poor coordination of inhalation with inhaler actuation, for which VHCs are prescribed. METHODS: Each VHC (n=5
devices/group) tested out-of-package. 5-actuations of fluticasone propionate (125 µg FP/actuation; Flovent®-HFA, GSK plc) was delivered at 30-s intervals to the VHC on test, sampling via a cascade impactor (CI) after 2, 5 and 10-s delay intervals, using a proprietary “delay” apparatus. FP was recovered from the CI and assayed by HPLC-spectrophotometry. Fine particle mass <4.7 µm aerodynamic diameter (FPM<4.7µm) provided a measure of the therapeutically beneficial medication capable of reaching the lungs. RESULTS: Values of FPM4.7 µm (µg FP; mean ± SD) are summarized in the Table. FPM4.7 µm for the pMDI alone (no delay) was 46.2 ± 2.1 µg.

<table>
<thead>
<tr>
<th>VHC/manufacturer</th>
<th>Delay Interval (s)</th>
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<tbody>
<tr>
<td></td>
<td>2</td>
</tr>
<tr>
<td>AeroChamber Plus*/Trudell Medical International</td>
<td>42.2 ±3.1</td>
</tr>
<tr>
<td>Antistatic Pocket Chamber®/nSpire Health Inc.</td>
<td>24.1 ±4.0</td>
</tr>
<tr>
<td>OptiChamber® Diamond®/Philips-Respironics Inc.</td>
<td>35.0 ±3.2</td>
</tr>
<tr>
<td>Vortex®/PARI Respiratory Equipment Inc.</td>
<td>39.9 ±2.9</td>
</tr>
</tbody>
</table>

CONCLUSIONS: Although antistatic materials enable VHCs to be used without pre-washing, there are still large differences in output. Regardless of delay length, the AeroChamber Plus* VHC most closely matched the benchmark FPM 4.7µm for the pMDI without VHC.


**RATIONALE:** Electrostatic charge mitigation by the use of charge dissipative materials with VHCs is common, since initial pre-washing can be avoided. We compared ‘antistatic’ VHCs; Optichamber® Diamond® (OD), Phillips Healthcare with AeroChamber Plus* Flow-Vu* (AC Flow-Vu) Trudell Medical International) (n=4 devices/group), to determine suitability for patients delaying inhalation post-actuation. METHODS: An abbreviated Andersen impactor that determined fine particle mass < 4.7 µm at 28.3 L/min (FPM<4.7µm) was used with an apparatus simulating 2, 5 and 10 s delay intervals following pMDI actuation (Flovent®, GSK plc, 125 µg/actuation fluticasone propionate (FP)). This approach conforms to guidance from European authorities that testing of VHCs should simulate delayed inhalation. Assay for FP was undertaken by HPLC-UV spectrophotometry. Measurements without delay were undertaken to assess mass recovery for FP, validating the procedure. All values are mean±SD. RESULTS: Mass recoveries (131.5±2.9 and 130.7±3.8 µg/actuation for the OD and ACPlus VHCs respectively) were close to label claim, validating system suitability. The variation of FPM<4.7µm with delay interval is shown in the Table.

<table>
<thead>
<tr>
<th>Delay (s)</th>
<th>AC Flo-Vu</th>
<th>OD</th>
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<tbody>
<tr>
<td>2</td>
<td>42.2±3.1</td>
<td>35.0±3.2</td>
</tr>
<tr>
<td>5</td>
<td>39.7±1.3</td>
<td>29.2±1.7</td>
</tr>
<tr>
<td>10</td>
<td>35.7±2.0</td>
<td>23.0±2.8</td>
</tr>
</tbody>
</table>

The ratio FPM<4.7µm-ACPlus/FPM<4.7µm-OD) increased from 1.2 (2-s) to 1.4 (5-s) and to 1.6 (10-s), demonstrating faster depletion of the therapeutically beneficial medication from the OD. CONCLUSION: Not all VHCs manufactured from anti-static materials provide optimum performance for patients who have poor coordination.


Background & Objectives: Leakage between facemask-and-face may result in medication loss by valved holding chamber (VHC)-facemask (1). Our study evaluated how an inspiratory flow indicator (IFI) can be used to avoid leakage. Methods: An infant face with realistic soft-tissue modeling (ADAM-III, Trudell Medical International (TMI), London, Canada (2)) was used to evaluate delivery of fluticasone propionate (FP; 50 µg/actuation, GSK (Canada)) via anti-static AeroChamber Plus* VHC with Flow-Vu* IFI/infant mask (AC-Plus, TMI) or Optichamber® Diamond® VHC/LiteTouch® small-mask (OD, Philips-Respironics, Parsippany, NJ, USA) (n=5 devices/group), simulating tidal-breathing (tidal-volume (Vt)=155-mL, duty-cycle =33%, rate= 25-breaths/min. Each facemask was applied to the face with the same clinically-appropriate force (1.6 kg). The IFI of the AC-plus was observed to be moving. FP was recovered from the nasopharynx and base (lung dose) of the model, and delivered mass (DMFP) quantified by HPLC-spectrophotometry as % label claim (LC). Findings: DMFP (mean±S.D.) was significantly greater from AC-plus (25.8±5.3%LC) than OD (17.0±3.7%LC) (unpaired t-test, p=0.019). FP on the facemask of the AC-plus

Delivery of inhaled medication to infants by valved holding chamber (VHC) with facemask may require more than one inhalation to empty the VHC because tidal volumes are typically smaller than chamber capacity. This study investigated the correlation between movement of an integrated inspiratory flow indicator (IFI) as a caregiver feedback aid for a VHC-facemask, number of inhalations and mass of medication, simulating use by a 6-9 month infant (tidal volume ($V_t$) = 50-ml; duty cycle = 25%; 30 cycles/min). Anti-static AeroChamber Plus® VHCs incorporating the IFI feature, with infant facemask ($n=5$/group, 3 replicates/device; Trudell Medical International, London, Canada) were coupled to a breathing simulator (ASL5000 test lung, IngMar Medical, Pittsburgh, PA, USA). The VHCs were prepared as per manufacturer instructions and the facemask of the device on test was fitted to the ADAM-II flexible infant face model with a clinically appropriate force of 1.6 kg. Aerosol capture took place using an electret filter positioned behind the lips of the face model. Delivery of medication was evaluated from two different pressurized metered dose inhaler formulations likely to be used with paediatric patients (Flovent HFA® 44; 44 μg fluticasone propionate (FP) delivered ex-actuator and Ventolin HFA®; 90 μg salbutamol base equivalent (SAL) delivered ex-actuator, both from GSK plc. One actuation was delivered to the VHC at the onset of inhalation, and the filter removed after 1 complete breathing cycle, observing the movement of the IFI to confirm inhalation valve opening. This procedure was subsequently repeated by removing the filter after 2, 3, 4, 5 and 6 breathing cycles. Assay for FP or SAL was undertaken by HPLC-UV spectrophotometry. During these measurements, the IFI of each device was observed to move in synchrony with valve opening on all occasions, confirming that the facemask sealed onto the face model without leakage of ambient air into the mask during the inspiratory phase of each breathing cycle. Emitted mass after the first breathing cycle (EM1) was 2.1 ± 0.7 μg (FP) and 5.8 ± 2.2 μg (SAL); substantially lower than the corresponding values after 6 cycles (EM6), being 9.0 ± 2.1 μg (FP), and 15.9 ± 3.1 μg (SAL) [paired test for each formulation; p < 0.001]. After 2 breathing cycles, values of EM2 (6.9 ± 2.0 (FP) and 13.0 ± 4.0 μg (SAL)), though significantly greater that their corresponding EM1 values [p ≤ 0.002], were still noticeably lower than the corresponding EM6 value for FP (p = 0.028), and barely statistically insignificant for SAL (p = 0.063). After 3 inhalations, EM3 increased further to 7.6 ± 2.0 μg (FP) and 13.8 ± 3.8 μg (SAL), and thereafter were close to the corresponding EM6 values, indicating emptying of the VHC had taken place. We conclude that at least two successive inhalations are required to achieve optimum medication delivery for the ‘infant’ condition under optimum conditions with a well fitted facemask with no leakage. The IFI is an important feature which validates that the facemask is properly sealed to the infant’s face and also confirms the number of inhalations that take place, thereby optimizing the therapeutic dose. Clinical studies are recommended to evaluate the benefit of this aid for the delivery of inhaled medication by VHC to this age group.


OBJECTIVE: To evaluate the efficacy and tolerability of fluticasone propionate (FP) hydrofluoroalkane (HFA) in children age 1 to < 4 years with asthma. STUDY DESIGN: Children were assigned (2:1) to receive FP HFA 88 μg (n = 239) or placebo HFA (n = 120) twice daily through a metered-dose inhaler with a valved holding chamber and attached facemask (AeroChamber Plus® VHC) for 12 weeks. The primary efficacy measure was mean percent change from baseline to endpoint in 24-hour daily (composite of daytime and nighttime) asthma symptom scores. RESULTS: The FP-treated children had significantly greater ($P \leq .05$) reductions in 24-hour daily asthma symptom scores (~53.9% vs ~44.1%) and nighttime symptom scores over the entire treatment period compared with the placebo group. Daytime asthma symptom scores and albuterol use were slightly more decreased with FP than with placebo; however, the differences were not statistically significant. Increases in the percentage of symptom-free days were comparable. The percentage of patients who experienced at least 1 adverse event was similar in the 2 groups. Baseline median urinary cortisol excretion values were comparable between the groups, and there was little change from baseline at endpoint. FP plasma concentrations demonstrated that systemic exposure was low. CONCLUSIONS: FP HFA 88 μg twice daily was effective and well tolerated in pre–school-age children with asthma.

Valved holding chambers with masks are often used with metered-dose inhalers in children with asthma to deliver drug to the lungs. Differences in holding chamber design can influence the amount of drug delivered. Lung deposition of fluticasone propionate (FP) using hydrofluoroalkane (HFA) propellant was examined using the AeroChamber Plus® and Babyhaler valved holding chambers. Children 1 to <4 year-olds were randomized in an open-label, 2-way crossover design (no washout between treatments) to receive 88 µg (44 µg/actuation) twice daily (every 12 hours) for 7.5 days (15 doses) using the AeroChamber Plus® VHC and Babyhaler with face-masks (FAS10002). The first and last 4 doses were directly observed by study staff. To limit the amount of blood collected from any one patient, children were randomized to one of three groups for blood sampling: Group 1: pre-dose, and 0.5-1, 1.5-2, 2.5-3, 3.5-4 hours post-dose: Group 2: 2.5-3, 3.5-4, 4.5-5, 6.5-7, 7.5-8 hrs post-dose; Group 3: 7.5-8, 8.5-9, 9.5-10, 11.5-12, post-dose, 12.5-13 hrs (0.5-1 hrs hour post dose #16). FP systemic exposure as described by area under the curve (AUC) was determined by population pharmacokinetics. Seventeen and 18 children completed AeroChamber® and Babyhaler treatments, respectively: one child completed only the Babyhaler treatment. Population mean (95% confidence interval) for FP exposure following dosing with the AeroChamber Plus® VHC was 97pg*h/ml (85, 113) and with the Babyhaler was 52pg*h/ml (34, 64). Lung deposition of FP through the AeroChamber Plus® VHC was higher when compared to the Babyhaler. However, systemic exposure for both devices was well below the threshold observed for decreases in cortisol production. Thus, both devices provide safe delivery of FP HFA to young children.

**FLUTICASONE PROPIONATE HFA IMPROVES ASTHMA CONTROL IN PRE-SCHOOL AGE CHILDREN WITH ASTHMA.**

**SUGERANW RM, TEPER AM, GIRARDI G, SCOTT CA, CLEMENTS DS, WU W, CRIM C. J ALLERGY CLIN IMMUNOL FEB 2005;115(2):S4-S5.**

**RATIONALE:** To evaluate the efficacy of fluticasone propionate HFA 88mcg BID (FP) vs placebo HFA (PLA) via MDI with the AeroChamber Plus® spacer with attached facemask for 12 weeks in pre-school age children with asthma. **METHODS:** One to <4 year-olds with ≥ 2 episodes of increased asthma symptoms requiring medical attention and pharmacotherapy ≤ 12 months prior to screening and a baseline 24-hr daily asthma symptom score (DASS; scale 0 = none to 3 = severe) of ≥ 1.1 were enrolled in this randomized (120 PLA: 239 FP), double-blind, parallel-group, placebo-controlled trial. Efficacy measures included: mean percent change from baseline to endpoint (last 28 days of treatment) in DASS (primary), mean change from baseline in nighttime asthma symptom scores over the entire treatment period (NASS), change from baseline to endpoint in daily rescue albuterol use (DRAB), and time to treatment failure (TF; i.e., time to first asthma exacerbation). **RESULTS:** Baseline mean DASS and NASS were comparable between groups (DASS=1.7 PLA, 1.8 FP; NASS = 1.2 PLA, 1.4 FP). At endpoint, FP-treated patients experienced a greater reduction (improvement) from baseline in DASS (54% FP, 44% PLA; p=0.036) and NASS (-0.56 FP, -0.44 PLA; p=0.049). Baseline DRAB use was comparable across groups (4 inhalations/day [IPD] PLA;5 IPD FP). DRAB decreased by 2 and 3 IPD for the PLA and FP groups, respectively, at endpoint. More PLA patients (12%) discontinued due to TF compared with FP-treated patients (5%) (p=0.034). **CONCLUSION:** Treatment with FP HFA 88 mcg BID for 12 weeks significantly improves asthma control in 1 to < 4 year-olds with asthma.

**SAFETY PROFILE OF FLUTICASONE PROPIONATE HFA IN PRE-SCHOOL AGE CHILDREN WITH ASTHMA.**


**RATIONALE:** To evaluate the safety of fluticasone propionate HFA 88mcg BID (FP) vs placebo HFA (PLA) via MDI with the AeroChamber Plus® spacer with attached facemask for 12 weeks in pre-school age children with asthma. **METHODS:** One to <4 year-olds with symptomatic asthma, receiving maintenance asthma medications (excluding systemic [ICS] or inhaled corticosteroids [ICS]) plus a short-acting beta-agonist (SABA) or SABA alone, were enrolled in this randomized (120 PLA: 239 FP), double-blind, parallel-group, placebo-controlled trial. Children receiving SCS within 10 weeks prior to randomization and/or ICS within 2 (low dose) or 8 (moderate-high dose) weeks prior to Screening were excluded. Safety assessments included: adverse events (AEs), clinical labs, oropharyngeal/nasal exams, asthma exacerbations, and 12-hour, overnight urinary cortisol excretion (U-Cortisol). **RESULTS:** No deaths or treatment-related serious AEs were reported. The percentages and types of AEs were comparable between groups. Events most commonly reported were fever (PLA=24%, FP=28%), nasopharyngitis (PLA=14%, FP=16%) and URTI (PLA=11%, FP=13%), events common in this age-group. Clinical lab results were comparable between groups. Few (PLA=0, FP=2) patients had a negative to positive shift in the oropharyngeal/nasal exam. More PLA-treated patients experienced an asthma exacerbation (11%) compared with FP-treated patients (4%). Baseline median U-Cortisol values were similar between groups (PLA=2.3mcg; FP=2.8mcg); and, there was little change from baseline after 12 weeks (PLA = -0.1mcg; FP = -0.4mcg). **CONCLUSION:** 12-week treatment with FP HFA 88mcg BID was well tolerated in 1 to <4 year-olds with asthma. The safety profile was similar to PLA and there was no evidence of adrenal suppression.

**IN VITRO DEPOSITION OF FLUTICASONE AEROSOL FROM A METERED-DOSE INHALER WITH AND WITHOUT TWO COMMON VALVED HOLDING CHAMBERS.**


**BACKGROUND:** Previous in vitro aerosol deposition experiments indicate that the corticosteroid respirable dose from a metered-dose inhaler (MDI) can vary by threefold depending on the specific valved holding chamber (VHC) MDI combination. **OBJECTIVE:** We compared in vitro aerosol deposition from a fluticasone propionate MDI (Flovent, GlaxoSmithKline, Research
performance of large and small volume valved holding chambers (VHCs) as a function of flow rate.


It is useful from the standpoint of the health care provider, if the performance of add-on devices for use with pressurized metered dose inhalers is characterized within the range of flow rates likely to be achieved by users. VHCs representing smaller (adult AeroChamber Plus®, 149 ml; n = 5) and larger (Volumatic™, 750 ml; n = 5) devices were compared with HFA-formulated fluticasone propionate (125 µg/dose ex metering chamber) at three flow rates, 28.3, 45 and 60 L/min. Measurements were made by Andersen 8-stage impactor. Fine particle fractions (< 4.7 µm, < 4.6 µm and < 4.0 µm aerodynamic diameter at 28.3, 45 and 60 L/min respectively) from both VHCs were close to 90%, significantly greater than that from the pMDI alone. At 28.3 L/min, fine particle dose (FPD) from the smaller VHC (50.5 ± 3.8 µg) was comparable with that from the larger VHC (45.9 ± 7.8 µg) [p = 0.27]. At the higher flow rates, FPD from the smaller VHC (65.5 ± 2.6 µg (45 L/min) and 65.2 ± 6.2 µg (60 L/min) exceeded equivalent values from the larger VHC (53.8 ± 3.7 µg (45 L/min) and 55.3 ± 4.9 µg (60 L/min)) [p < 0.023].


Rationale: To compare the delivery of fluticasone propionate in terms of fine particle (< 4.7 µm aerodynamic diameter (FPD)) and total emitted dose (ED) from 4 VHCs. Methods: FPD and ED were determined for AeroChamber Plus® (Monaghan Medical Corp., Plattsburgh, NY, 149 ml), OptiChamber® (Respironics, Cedar Grove, NJ, 218 ml), Pocket Chamber™ (Ferraris Medical, Inc., Holland, NY, 90 ml) and ACE® (DHD Healthcare, Warminster, PA, 150 ml) VHCs (n=5 devices/group). Particle size measurements were made by Andersen cascade impactor (Graebe Andersen, USA) at 28.3 ± 0.5 L/min. The VHCs were washed with a mild ionic detergent followed by air-drying before testing to minimize the influence of electrostatic effects. The mass of FP was assayed by HPLC-UV spectrophotometry. Results: Both FPD and ED unit doses, normalized to label claim, from the AeroChamber Plus® VHCs (60.2 ± 3.8 µg (FPD); 69.1 ± 3.5 µg (ED)) significantly exceeded equivalent values from the OptiChamber® VHCs (38.4 ± 1.7 µg (FPD); 43.7 ± 1.2 µg (ED)), Pocket Chamber™ (37.2 ± 0.8 µg (FPD); 41.1 ± 0.8 µg (ED)) and ACE® (20.3 ± 4.1 µg (FPD); 22.0 ± 4.5 µg (ED)) (1-way ANOVA, p < 0.001). Conclusion: Chamber volume alone does not necessarily correlate with improved FPD or ED and other considerations, such as internal geometry and inhalation valve design are also important.


We report an in vitro investigation in which the performance of an improved small volume (149 ml) VHC (adult AeroChamber Plus*, Trudell Medical International, London, Canada; n = 5 devices) was compared with a 218 ml VHC (OptiChamber®; Respironics, Cedar Grove, NJ, USA) for a corticosteroid (HFA-fluticasone propionate (HFA-FP) 125 µg/dose, GlaxoSmithKline plc, UK). Measurements of emitted dose (ED), together with fine particle fraction (FPF) and fine particle dose (FPD), were made by Andersen 8-stage impactor equipped with a USP induction port at 28.3 ± 0.5 L/min. We observed the following (mean ± S.D.):

<table>
<thead>
<tr>
<th>VHC</th>
<th>FPD* (mg)</th>
<th>FPF* (%)</th>
<th>ED (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AeroChamber Plus®</td>
<td>43.1 ± 3.8</td>
<td>86.3 ± 3.1</td>
<td>49.9 ± 4.4</td>
</tr>
<tr>
<td>OptiChamber®</td>
<td>19.3 ± 2.7</td>
<td>80.2 ± 1.6</td>
<td>24.1 ± 3.5</td>
</tr>
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</table>

*particles < 4.7 µm aerodynamic diameter

In spite of its smaller volume, the AeroChamber Plus® VHCs delivered significantly more HFA-FP than the OptiChamber® VHCs, both in terms of FPD and ED (un-paired t-test, p < 0.001). The size selectivity of the smaller VHC was also significantly better (p = 0.005). The improved small volume VHC provides highly efficient delivery of particles of HFA-FP in sizes likely to...
penetrate to receptors in the lung, whilst minimizing the release of coarser particles that are likely to deposit in the oropharyngeal region.


The delivery of aerosolized medication from valved holding chambers (VHCs) used with pressurized metered dose inhalers (pMDIs) is dependent on the inspiratory flow rate of the patient. A study was undertaken to evaluate the delivery of a corticosteroid (Flovent®: 110 µg/dose CFC-fluticasone propionate (FP)) through two VHCs (AeroChamber Plus®; Monaghan Medical Corp., Plattsburgh, NY and OptiChamber®; Respironics, Cedar Grove, NJ: n = 10 devices/group) at 5.0 ± 0.5 and 12.0 ± 0.5 l/min, representative of low flow rate patients. A single dose of medication was delivered to a filter square (Filtrate®, 3M Corp., St. Paul, MN) located at the mouthpiece/mask adapter of each device to provide data indicative of optimum performance without ‘dead volume’. The filter was removed 30 s after pMDI actuation and agitated in methanol to release collected medication quantitatively. The mass of FP collected was assayed by HPLC-UV spectrophotometry. The AeroChamber Plus® VHCs delivered total emitted doses of 45 ± 2 and 67 ± 4 µg FP at 5 and 12 l/min respectively, compared with 23 ± 6 and 49 ± 2 µg from the OptiChamber® VHCs at the equivalent conditions. The difference in emitted dose between VHC types was statistically significant at both flow rates (un-paired t-test, p < 0.001). In vitro data based on measurements made at a flow rate of 28.3 l/min or higher should not be taken as predictive of performance with patients that can only achieve low inspiratory flow rates.


From the standpoint of the health care provider, it is helpful if the therapeutically beneficial fine particle dose of a pMDI-delivered formulation such a corticosteroid is comparable with that delivered by the pMDI alone, irrespective of the choice of propellant in the formulation. The performance of an improved valved holding chamber (adult AeroChamber Plus® VHC; n = 5) is reported with HFA-formulated fluticasone propionate (HFA - FP; 125 µg/dose ex metering chamber). As expected, the VHC greatly increased the proportion of the emitted dose (ED) delivered in particles finer than 4.7 µm aerodynamic diameter (FPF) from 39.0 ± 3.0% to 74.7 ± 3.7% (HFA-FP) [paired t-test p < 0.001]. The fine particle dose (FPD) with the VHC (43.5 ± 5.3 µg) was comparable with that delivered by the pMDI alone (47.4 ± 1.1 µg) [p = 0.21]. The portion of the dose contained in coarser particles that might otherwise deposit in the upper respiratory tract and be associated with unwanted side-effects, decreased substantially from 74.5 ± 7.5 µg (pMDI alone) to 14.7 ± 2.2 µg with the addition of the VHC [p<0.001].

Flutiform™ (Fluticasone / Formoterol ) SkyePharma AG

FLUTICASONE PROPIONATE/FORMOTEROL FUMARATE COMBINATION THERAPY IS EQUALLY EFFECTIVE AND WELL-TOLERATED WHEN ADMINISTERED WITH OR WITHOUT A SPACER DEVICE TO PATIENTS WITH ASTHMA. Price D, Papi A, Kaiser K, Grothe B, Lomax M. Presented at the European Respiratory Conference in Amsterdam, September, 2011.

Background: Phase 3 studies involving a new asthma therapy combining fluticasone (FLUT) and formoterol (FORM) in a single aerosol inhaler (FLUT/FORM: flutiform®) either used a spacer or did not. This is the first comparison of the efficacy and tolerability of treatment with or without the use of a spacer device. Methods: Adults and adolescents with mild, moderate or moderate-severe asthma were treated with FLUT/FORM 100/10µg or 250/10µg b.i.d. delivered either with (N=195) or without (N=532) a spacer in 6 randomised, double-blind and open-label, parallel group studies. The endpoint was non-inferiority between spacer and non-spacer groups (concluded if the lower bound of the 95% CI was ≥-0.2L) in terms of changes in morning pre-dose FEV1 and morning pre-dose to 2h post-dose FEV1 over 12 weeks. The incidence of adverse events (AEs) was analysed over 8 weeks. Results: FLUT/FORM was consistently as effective when delivered with or without a spacer at all dose strengths and asthma severities. From baseline to end of study, the LS mean treatment difference in morning pre-dose FEV1 was 0.067L greater without spacer (95% CI: -0.149, 0.015) and in morning pre-dose to 2h post-dose FEV1 was 0.015L greater with spacer (95% CI: -0.051, 0.081). AEs were reported with similar frequency both with and without a spacer (nasopharyngitis: 14 (2.1%) vs 34 (3.2%); asthma: 12 (1.8%) vs 17 (1.6%) patients; cough: 4 (0.6%) vs 10 (0.9%); dysphonia: 5 (0.7%) vs 7 (0.7%).Conclusions: Pooled analysis showed that fluticasone/formoterol may be given with or without a spacer device with both approaches providing similar efficacy and tolerability.


Introduction: Inhaled combination products are the mainstay of therapy in asthma and COPD. Three dosage strengths of Flutiform were developed to offer to the patient the novel combination of the well-established corticosteroid (ICS) fluticasone
and the fast onset, long-acting beta-agonist (LABA) formoterol in a convenient pMDI presentation. Spacers or valved holding chambers (VHC) can be used coupled to pMDIs to address the common issues patients face with; (1) the co-ordination of inhalation and actuation, and (2) the mouth/throat deposition and/or swallowing of steroid drugs. The use of VHCs is intended to remove the larger particles from the emitted aerosol whilst maintaining the delivery of fine particles or lung dose to the patient. Regulatory guidance documents in Europe and the US require the product to be characterized in vitro with any spacer or VHC device recommended in the patient instructions (1,2). More recently, regulators specified that the same principals should be applied to in vitro spacer tests as were applied to comparative pMDI testing (3). Methods: Four different spacers and VHC were selected for in vitro characterization based on design and material considerations and market prevalence. Aerodynamic particle size distribution (APSD) was determined by Andersen Cascade Impactor (ACI. 28.3 L/min configuration) of medium strength Flutiform 125/5 delivered via spacer / VHC, with flutiform delivery without spacer as control. Further characterization of the product performance with the selected spacer was performed by evaluating dose delivery and APSD of Flutiform strengths (50/5, 125/5, 250/10 ug/actuation) at various flow rates (28.3, 60 and 90 L/min) and different operating regimes (0 second time lag between actuation and inhalation and 2 and 5 second time lag to simulate poor coordination). Results and Discussion: In the first part of the study, APSD of flutiform delivery via four different spacers was evaluated for selecting the device providing the best match for further evaluation. As exemplified by Figure 1 and 2, a comparable in vitro APSD of the delivered drugs was found when the product was delivered via its proprietary press-and-breathe actuator with or without the use of the TMI AeroChamber Plus* VHC device, apart from a major proportion of large particles being kept in the spacer which other-wise would be deposited in the USP induction port (‘throat’) of the apparatus. Most importantly, the amount of drug deposited on stage 3 onwards of the ACI (28.3 L/min configuration) was very similar. Further studies were performed with all flutiform strengths to characterize the drug delivery from TMI AeroChamber Plus* VHC at various flow rates and operating regimes. The results showed no relevant impact of the time lag on the fluticasone deposited on grouped stages of the ACI, as exemplified by Figure 3 and 4 for low strength flutiform 50/5. It should be noted that the fine particle fraction (FFP) (stage 3 to filter, < 4.7 µm) is around 40% of the delivered dose across all conditions. Comparable results were found for the delivery of formoterol and for the other flutiform strengths. Conclusions: The in vitro APSD profiles confirm the design intent for improved delivery to patients by showing that use of the TMI AeroChamber Plus* delivers the same estimated lung dose as the standard press and breathe actuator whilst effectively removing the larger particles from the emitted aerosol which would be expected to impact in the patient’s mouth/throat and/or be swallowed. The FPF (expressed as % of delivered dose) was shown to be consistent at around 40% for both drugs under all conditions tested. The TMI AeroChamber Plus was confirmed as a suitable spacer for recommendation in the patient instructions for flutiform.

**Fostair™ (Beclometasone Dipropionate / Formoterol Fumarate) Chiesi Farmaceutici S.p.A**


**WHAT IS ALREADY KNOWN ABOUT THIS SUBJECT** • Use of a spacer minimises oropharyngeal deposition and optimises drug targeting to the airways in subjects with coordination difficulties. However, the increase in pulmonary deposition often observed with spacer devices, could potentially lead to an increase in overall systemic exposure. • EMA guidelines recommend that the development of a pMDI should always include testing of at least one specific spacer for use with a particular pMDI. • The aim of this study was to examine the effect of AeroChamber Plus* VHC on the lung bioavailability and total systemic exposure of a HFA pMDI fixed combination of extrafine beclometasone dipropionate/formoterol (100/6 µg) (Foster® ). WHAT THIS STUDY ADDS • The use of AeroChamber Plus* VHC optimises the lung delivery of beclometasone and formoterol in subjects that find it difficult to synchronise aerosol actuation with the inspiration of breath. • The total systemic exposure of beclometasone 17-monopropionate and formoterol was not significantly increased by the use of the AeroChamber Plus* spacer. • Use of the AeroChamber Plus(TM) spacer device with the extrafine beclometasone dipropionate/formoterol (100/6 µg) fixed combination pMDI can be a valuable option for certain patients groups, such as subjects with difficulties in achieving an adequate inhalation technique. Aim To assess the effect of AeroChamber Plus* on lung deposition and systemic exposure to extrafine beclometasone dipropionate (BDP)/formoterol (100/6 µg) pMDI (Foster® ). The lung deposition of the components of the combination given with the pMDI was also evaluated using the charcoal block technique. Methods Twelve healthy male volunteers received 4 inhalations of extrafine BDP/formoterol (100/6 µg) using (1) pMDI alone, (2) pMDI and AeroChamber Plus* and (3) pMDI and charcoal ingestion. Results Compared with pMDI alone, use of AeroChamber Plus* increased the peak plasma concentrations (C_max) of BDP (2822.3 ± 1449.9 vs 5454.9 ± 3197.1 pg/ml), its active metabolite beclometasone 17-monopropionate (17-BMP) (771.6 ± 288.7 vs 1138.9 ± 495.6 pg/ml) and formoterol (38.4 ± 17.8 vs 54.7 ± 20.0 pg/ml). For 17-BMP and formoterol, the AUC0-30 min, indicative of lung deposition, was increased in the AeroChamber Plus* group by 41% and 45%, respectively. This increase was mainly observed in subjects with inadequate inhalation technique. However, use of AeroChamber Plus* did not increase the total systemic exposure to 17-BMP and formoterol. Results after ingestion of charcoal confirmed that AUC0-30min can be taken as index of lung bioavailability and that more than 30% of the inhaled dose of extrafine BDP/Formoterol 100/6 µg was delivered to the lung using the pMDI alone. Conclusions: The use of AeroChamber Plus* optimises the delivery of BDP and formoterol to the lung in subjects with inadequate inhalation technique. The total systemic exposure was not increased, supporting the safety of extrafine BDP/formoterol pMDI with AeroChamber Plus*.
The effect of AeroChamber Plus* spacer on the systemic exposure to formoterol, BDP and B17MP (beclometasone-17-monopropionate, the main BDP metabolite) was evaluated in healthy male subjects after inhalation of a single 4-puff dose of the extrafine BDP/formoterol 200/6 µg pMDI fixed combination (total inhaled dose 800/24 µg) following a randomized, cross-over design. The use of the spacer increased peak plasma concentration (Cmax) of B17MP (+12%) and formoterol (+39%). The area under the plasma concentration versus time over the first 30 minutes post-dose (AUC0-0.5h), indicative of lung absorption, increased by 25% and 32% for B17MP and formoterol, respectively, in the spacer group. However, the use of AeroChamber Plus* did not increase the total systemic exposure to B17MP and formoterol, as it decreased the amount of drug swallowed and absorbed from the gastrointestinal tract. This result was in accordance with the higher Fine Particle Dose observed in vitro for both components when using the spacer, and with the results of a previous study evaluating the effect of AeroChamber Plus* on FOSTER® (extrafine BDP/formoterol 100/6 µg). Similar systemic exposure was confirmed by comparable pharmacodynamic effects observed after administration with and without spacer. No differences were found between AeroChamber Plus* spacer and the standard actuator in serum and urinary cortisol, both treatments showing a statistically significant decrease compared to placebo, with mean values which remained within the physiological range. Serum potassium, vital signs and QTcB parameters were also not affected by the use of the spacer device.

Proventil™ (Albuterol Sulfate) Key Pharmaceuticals Inc.

RESPONSE TO ALBUTEROL SULFATE KEY PHARMACEUTICALS INC. DELIVERED THROUGH AN ANTI-STATIC CHAMBER DURING NOCTURNAL BRONCHOSPASM. Prabhakaran S, Shuster J, Chesrown S, Hendeles L. Respir Care 2012;57(8):1291-1296.

Background: Decreasing electrostatic charge on valved-holding chambers increases the amount of drug delivered. However, there are no data demonstrating that this increases bronchodilatation. The objective of this study was to investigate the influence of reducing electrostatic charge on the bronchodilator response to albuterol inhaler during nocturnal bronchospasm.

Methods: This randomized, double-blind, double-dummy, crossover study included subjects 18-40 years with nocturnal bronchospasm (20% overnight decrease in peak flow on 3 of 7 nights during run-in), FEV1 60-80% predicted during the day, and ≥ 12% increase after albuterol. Subjects slept in the Clinical Research Center up to 3 nights for each treatment. FEV1 and heart rate were measured upon awakening spontaneously or at 4 am, and 15 minutes after each dose of 1, 2 and 4 cumulative puffs of albuterol MDI. The drug was administered after an anti-static valved holding chamber (AeroChamber Plus Z-Stat®) or a conventional valved holding chamber containing a static charge (AeroChamber Plus®). Results: Of 88 consented subjects, 11 were randomized and 7 completed the study. Most exclusions were due to lack of objective evidence of nocturnal bronchospasm. Upon awakening, FEV1 was 44±9% predicted before the anti-static chamber and 48±7% predicted before the static chamber. The mean (±SD) % increase in FEV1 after 1, 2 and 4 cumulative puffs using anti-static vs static chambers, respectively, were 52%±26% vs 30%±19%, 73%±28% vs 48%±26% and 90±34 vs 64±35%. The point estimates for the difference (95% CI) between devices (anti-static-static) were 21% (4-38) [p=0.026], 23% (6-41) [p=0.018] and 25% (7-42) [p=0.013] for 1, 2, and 4 cumulative puffs, respectively. There was no significant difference in heart rate between treatments.

Conclusion: Delivery of albuterol through an anti-static chamber provides a clinically relevant improvement in bronchodilator response during acute, reversible bronchospasm, such as nocturnal bronchospasm.


Purpose: To compare fine particle (FPD) and total emitted dose (ED) from 149 and 218 ml valved holding chambers (VHCs) with an HFA-formulated bronchodilator. Methods: FPD and ED were determined for 5 OptiChamber VHCs (volume = 218 ml; Respironics, Cedar Grove N.J.) and 5 AeroChamber Plus* VHCs with mouthpiece (149 ml; Monaghan Medical Corp., Plattsburgh, N.Y.) with Proventil™-HFA (108 µg albuterol sulfate ex actuator; Key Pharmaceuticals Inc.) by means of an Andersen 8-stage impactor, sampling at 28.3 ± 0.5 l/min. Results: (ED) and ((FPD), particles finer than 4.7 µm aerodynamic diameter) were as follows: OptiChamber VHC: ED = 52.1 ± 3.2 µg, FPD = 50.4 ± 2.5 µg, AeroChamber Plus*: ED = 68.9 ± 5.2 µg, FPD = 65.8 ± 5.8 µg. The differences in both ED and FPD were statistically significant (unpaired t-test for each variable, p<0.001). Fine particle fraction (FPF) from either type of VHC (OptiChamber®: 96.9 ± 2.3%, AeroChamber Plus* (95.5 ± 1.8%) were insignificantly different (p = 0.30). Conclusion: Increased chamber volume does not necessarily equate with improved FPD or ED. Other considerations, such as internal geometry and inhalation valve design contribute to performance by controlling the internal aerosol losses.
Objectives: The aim of this in vitro study was to determine the delivered dose of budesonide 200µg via a chlorofluorocarbon-free pressurized metered dose inhaler (pMDI) when administered through different spacers in tidal breathing patterns of young children. Methods: Tidal breathing was simulated for toddlers and children. Spacers tested were Babyhaler®, AeroChamber Plus* VHC small and medium; the pMDI was Dudiair® 200µg. Output was measured after one actuation and five inhalations in primed and unprimed spacers. Cumulated output was evaluated after each of five simulated inhalations. Aerosol characteristics – i.e. particle size distribution of the output – were determined in primed spacers with a cascade impactor using high-performance liquid chromatography and UV detection. Results: Total output from primed spacers after five inhalations was determined between 37.9 µg and 40.9 µg with little differences between spacers and breathing patterns. About 58 – 79% of this total output was inhaled with the first breath from the AeroChamber Plus* and about 26% from the Babyhaler®. The fine particles <5µg ranged between 87% and 92% of the delivered dose for all three spacers. Discussion and Conclusion: The nominal dose (200µg) of the Dudiair® 200µg inhaler is reduced to 40µg delivered dose or less by using Babyhaler® and AeroChamber Plus* spacers taking five breaths. With a single breath the delivered dose can be reduced further to a minimum of 10µg using the Babyhaler®. Clinical studies are warranted in the future for decisions on ‘clinical efficacy’, safety, and exact dose adjustment.

QVAR® (Beclomethasone Dipropionate), Graceway™ Pharmaceuticals LLC


Background: The study objective of this pilot study was to determine the lung delivery of HFA-134a beclomethasone dipropionate (HFA-BDP; QVAR™) and CFC-beclomethasone dipropionate (CFC-BDP; Becloforte™) with and without the add-on spacers, AeroChamber*, and Volumatic™. The smaller particles of HFA-BDP were presumed to produce greater lung deposition using spacers, with and without a delay [i.e. metered dose inhaler (MDI) actuation into the spacer and subsequent inhalation 0 and 2 sec later], compared with the larger particles of CFC-BDP. The study included a comparison of breathhold effects (i.e. 1 and 10-sec breathholds) on lung deposition. Methods: The study was an open-label design and utilized healthy subjects (n=12 males). Each arm of the study contained three subjects; thus, outcomes were not powered to assess statistical significance. HFA-BDP and CFC-BDP were radiolabeled with technetium-99m and delivered to subjects. Results: Results showed that the small particle HFA-BDP lung deposition averaged 52% and was not affected by the use of AeroChamber* with or without a spacer delay. The oropharyngeal deposition of HFA-BDP was reduced from approximately 28% to 4% with the AeroChamber*. Lung deposition with the large particle CFC-BDP was 3–7% and generally decreased with AeroChamber* or Volumatic. A 2-sec time delay between actuation and breath plus the spacer reduced lung deposition slightly but reduced oropharyngeal deposition substantially (84% down to 3–20%) using the AeroChamber* or Volumatic with and without a spacer delay. HFA-BDP lung deposition was dependent on the breathhold. Lung deposition with HFA-BDP was reduced by 16% with a 1-sec versus 10-sec breathhold. The difference was measured in the increased exhaled fraction, confirming that smaller particles need time to deposit and are exhaled if there is a reduced breathhold. The large particle CFC-BDP lung deposition was not affected by breathhold. Conclusions: The use of Aerochamber* or Volumatic spacers with HFA-BDP did not alter lung deposition but it did reduce oropharyngeal deposition. However, HFA-BDP displayed reduced oropharyngeal deposition without a spacer.


The aim of the present study was to measure airway, oropharyngeal and gastrointestinal deposition of 99mTc-labelled hydrofluoroalkane-beclomethasone dipropionate after inhalation via a pressurized metered-dose inhaler and spacer (AeroChamber Plus* VHC) in asthmatic children. A group of 24 children (aged 5-17 yrs) with mild asthma inhaled the labeled drug. A total of 12 children took five tidal breaths after each actuation (tidal group). The other 12 children used a slow maximal inhalation followed by a 5-10-s breath-hold (breath-hold group). Simultaneous anterior and posterior planar γ-scintigraphic scans (120-s acquisition) were recorded. For the tidal group, mean ±SD lung deposition (% ex-actuator, attenuation corrected) was 35.4±18.3, 47.5±13.0 and 54.9±11.2 in patients aged 5-7 (n=4), 8-10 (n=4) and 11-17 yrs (n=4), respectively. Oropharyngeal and gastrointestinal deposition was 24.0±10.5, 10.3±4.4 and 10.1±6.2. With the breath-hold technique, lung deposition was 58.1±6.7, 56.6±5.2 and 58.4±9.2. Oropharyngeal and gastrointestinal deposition was 12.9±3.2, 20.1±9.5 and 20.8±8.8. Inhalation of the extrafine formulation with the breath-hold technique showed significantly improved lung deposition
comparing with tidal breathing across all ages. Oropharyngeal and gastrointestinal deposition was markedly decreased, regardless of which inhalation technique was applied, compared with a previous paediatric study using the same formulation delivered via a breath-actuated metered-dose inhaler.


STUDY OBJECTIVE: To compare in vitro aerosol deposition from a beclomethasone dipropionate metered-dose inhaler (MDI) containing hydrofluoroalkane propellant with that of the MDI in combination with two common valved holding chambers (VHCs) to evaluate how these VHCs affect the respirable dose of beclomethasone dipropionate. DESIGN: In vitro aerosol deposition study. SETTING: University research center. DEVICES: Beclomethasone dipropionate hydrofluoroalkane MDI alone, the MDI with OptiChamber† VHC, and the MDI with AeroChamber Plus* VHC. INTERVENTION: The respirable dose (1-5-microm aerosol particles) of beclomethasone dipropionate was determined by sampling 10 80-microg actuations from five runs with each configuration (MDI alone, MDI with OptiChamber†, and MDI with AeroChamber Plus* VHC), using a well-established in vitro cascade impactor method. MEASUREMENTS AND MAIN RESULTS: Beclomethasone dipropionate aerosol was washed from the impactor with 50% methanol and quantified by means of high-performance liquid chromatography. Differences among outcomes were determined by using analysis of variance. Mean beclomethasone dipropionate respirable dose from AeroChamber Plus* VHC (27.2 +/- 10.0 microg/actuation) was not significantly different (p>0.05) from that of the MDI alone (29.0 +/- 7.0 microg/actuation). OptiChamber† respirable dose (12.8 +/- 6.0 microg/actuation) was less than half that produced by either the AeroChamber Plus* VHC or the MDI alone (p=0.013). CONCLUSIONS: The OptiChamber† and AeroChamber Plus* VHCs do not demonstrate equivalent in vitro performance when used with a beclomethasone dipropionate MDI that contains hydrofluoroalkane propellant. The respirable dose of beclomethasone dipropionate aerosol from the hydrofluoroalkane MDI was decreased by only 6% when the MDI was mated to an AeroChamber Plus* VHC and by 56% when used with an OptiChamber† VHC.


Purpose: To demonstrate the reduction in coarse component (particles >4.7 µm aerodynamic diameter) of pressurized metered dose inhaler (pMDI) delivered HFA-beclomethasone dipropionate (BDP) (QVAR100: 100 µg/dose, 3M Pharmaceuticals Inc.), using a small volume VHC (AeroChamber Plus*, (Monaghan Medical Corp.)).METHODS: 5 AeroChamber Plus* VHCs were washed with a mild ionic detergent followed by air-drying prior to testing in order to minimize the influence of electrostatic effects. Particle size measurements were made by Andersen cascade impactor (Graseby Andersen, USA) at 28.3 ± 0.5 l/min and the mass of BDP assayed by HPLC-UV spectrophotometry. 5 doses of medication were delivered 30 s apart.

Results: The AeroChamber Plus* VHCs delivered total unit doses of 62.0 ± 3.4 µg (94 % < 4.7 µm), whereas the pMDI alone provided 74.3 ± 1.6 µg (61% <4.7µm). The amount of BDP >4.7 µm was 28.9 ± 3.0 µg and 3.1 ± 1.1 µg for the pMDI alone and AeroChamber Plus* respectively. Conclusion: Based on these data a holding chamber still appears to be necessary to minimize deposition in the upper airway, even with the introduction of this CFC-free solution formulation.

ratio-Salbutamol® HFA (Albuterol Sulfate) ratiopharm Canada


PURPOSE: VHCs are prescribed for patients that have difficulty coordinating pressurized metered-dose inhaler (pMDI) use, frequently resulting in delayed inhalation following inhaler actuation. Our study introduced a realistic 2-second delay, comparing delivery of a beta-2 agonist via VHCs of similar size (n=5/group), one manufactured from cardboard (LiteAire”, Thayer Medical, Tucson, AZ – 160-ml) the other from rigid polymer (AeroChamber Plus* VHC, Monaghan Medical Corp., Syracuse, NY – 150-ml). METHODS: The AeroChamber Plus* VHCs were pretreated by washing in water containing a mild ionic detergent, rinsed and drip-dried, as recommended prior to use. The LiteAire” VHCs were assembled and used in accordance with manufacturer's instructions. Each VHC was tested using an Andersen 8-stage impactor with USP Induction Port operated at 28.3±0.5 l/min, representative of flow rates seen with adult patients. A shutter that interfaced between the VHC mouthpiece and induction port entrance was used to simulate a 2-s delay interval between pMDI actuation and the onset of sampling. The shutter moved to allow flow from the VHC to the impactor only after the defined delay. 5-actuations of albuterol (Ratiopharm, Mississauga, Canada, 100 µg/dose albuterol base equivalent ex metering valve) were delivered from a pre-primed and shaken pMDI canister at 30-s intervals. The induction port and stages of the impactor were subsequently assayed for albuterol by HPLC-UV spectrophotometry. Benchmark measurements were also made with the pMDI alone. RESULTS: Fine particle
RESPIMAT® Soft Mist™ Inhaler, Boehringer Ingelheim


Purpose: In a non-invasive handling study the Respimat® inhaler was investigated, followed by flow profile recording of children (N=99) below 5 years of age to find out whether inhalation therapy will be possible. Methods: Handling of the active Respimat® inhaler alone or with a spacer and facemask (AeroChamber Plus*) was assessed using a standardised handling questionnaire. Next the inhalation flow profiles were recorded and results summarized by descriptive statistics. Results: The most important flow profile parameter was volume inhaled after release (VA) within 10s using the spacer (acceptable holding time of spray in the spacer, ~5 breaths) and 1.5s using Respimat® alone (spray duration). The criterion for successful inhalation was a minimum volume inhaled of 0.15L which equals the spacer volume and is an acceptable value in accordance with the young age. This criterion was met in all age groups and handling configurations. With Respimat® a single breath resulted in median VA: 0.63L (3-<4yr) and 0.47L (4-<5yr). Using the spacer, median values of VA were: 0.34L (0-<2yr), 0.71L (2-<3yr), 1.12L (3-<4yr), 0.94L (4-<5yr) for ~5 breaths. The median peak flow with Respimat® alone reached 1L/s and with spacer it was below 0.7L/s in all age groups. Conclusions: These data suggest Respimat® is suitable for inhalation therapy in pre-school children. In order to ensure standardized dosing, the use of Respimat® inhaler with spacer in children below 5 years of age is recommended.


Rationale / Introduction: Drug delivery by inhalation is the most effective non-invasive therapy available today. While patients older than 5 years are mostly able to perform correct inhalation, children below that age require special assistance. Respimat® Soft Mist™ Inhaler is a novel, easy to use hand-held multidose propellant-free inhalation device that generates a fine, slow-moving cloud with high fine particle dose. Since spray generation is independent of the inspiratory flow it may offer opportunities to treat young children. The objective of this study was to establish the age at which children below 5 years can use Respimat® inhaler and which degree of help by parents or by using a spacer may be appropriate. Methods: Open two-centre observational handling study. 99 pediatric patients (any respiratory disease) in five age groups were analyzed. Children <2 years started directly the inhalation with Respimat® and spacer (AeroChamber Plus* with facemask). Children 2 years and older started with the use of Respimat® without spacer, with or without help by their parents. If correct handling was not achieved, inhalation was repeated with Respimat® and spacer. Successful handling was defined as: (A) enclosure of the inhaler mouthpiece without covering the airvents, (B) coordination of dose release and inhalation, and for use with spacer (C) correct placement of the spacer followed by inhalation. The primary endpoint was the proportion of correct handling maneuvers. The inhalation profile in the successful handling configuration was verified with a pneumotachograph. Patient satisfaction and preferences were investigated in a questionnaire answered by the parents as one of the secondary endpoints. Results: All children below 3 years of age achieved correct handling of Respimat® with spacer (only one child refused cooperation). 40% (12/30) of the 3 to <4 year old children achieved correct handling of Respimat® without spacer. 85% (23/27) of the 4 to <5 year old children achieved correct handling without spacer. The percentage of correct handling maneuvers is lower without assistance by parents. In general the correct handling maneuvers were confirmed by the inhalation profile assessments. Analysis of satisfaction and preferences showed that most parents were fully satisfied with the handling of Respimat® and more than 90% stated that they were able to handle Respimat®. Conclusion: Children below 3 years of age should use Respimat® with spacer. The majority of 4 to <5 year old patients can handle Respimat® without spacer. Respimat® can be considered as suitable for inhalation therapy of young patients.

Salbutamol (Albuterol Sulfate)

Background: The objective of this work is to numerically evaluate several commercial Valved Holding Chamber (VHC) geometries, in terms of airflow behavior and wall deposition. Also, the Fine Particle Dose (FPD) and Mass Mean Aerodynamic Diameter (MMAD) of the drug dose reaching the lungs were evaluated. Downwards the VHC Mouthpiece, the USP Throat geometry was added. Major detail was included in the VHC components representation. Methods: Through the use of Computational Fluid Dynamics (CFD), the airflow velocity and turbulence fields were calculated for four geometries (i.e. A2A®, Aerochamber Plus® Flow-Vu®, NebuChamber® and Volumatic®). Using a constant flow of 30 L/min and several realistic spray inputs, the deposition was analyzed for three distinct particle size distributions. Results: The Volumatic® presents the higher recirculation in comparison to the other small volume VHC devices. Each VHC Valve and Mouthpiece design leads to different flows entering the Throat, where the NebuChamber® exhibits the higher air velocities (i.e. 34 m/s). The higher is the MMAD of the distribution injected, the greater will be the deposition in the upper walls. Therefore, the lower MMAD distribution results in higher dose available for the patient. Volumatic® showed the higher Body deposition, as well as, the NebuChamber® USP Throat. Although the Aerochamber* Valve presents the higher deposition, it provides the greater amount of drug for the patient lungs. On the other hand, the Volumatic® geometry while yielding the lower MMAD for the patient is far from providing the higher FPD. Conclusions: Based on the results, the VHC components design lead to very distinct airflow patterns. The sudden changes in particle trajectory result in higher deposition at those locations. The Aerochamber* delivers the higher FDM to the patient lungs, while the Volumatic® delivers the smaller MMAD distribution to the lungs.


RATIONALE: Clinical guidelines for asthma and COPD suggest health care providers titrate the patient to the least dose that is efficacious. In mild stable asthma or COPD, the dosing regimen will likely be pMDI+VHC. However, in an exacerbation, nebulizer treatment may be more appropriate. If a dosimetric BAN is used, it is possible to relate the drug mass delivered in a given time to the equivalent number of pMDI actuations. We report such data here for salbutamol, which can be delivered by either pMDI+VHC or nebulizer routes. METHODS: Fine particle mass < 4.7 µm salbutamol ex-AeroChamber Plus* VHC; Trudell Medical International (TMI), London, Canada (FPM<4.7µm; n=5 devices) was determined by Andersen 8-stage cascade impactor following the pharmacopeial method, but simulating a 2-s delay between pMDI actuation and the onset of sampling to mimic the poorly coordinate patient for whom these devices are prescribed. In parallel studies, the fine particle delivery rate (FPM<4.7µm/min) of salbutamol solution (2.5 mg/3mL) from AeroEclipse®II BANs (n=5) with 1.5, 2.0, 2.5 and 3.0 mL fill volumes operated at 50 psig was determined with the mouthpiece of the nebulizer connected via a collection filter to a breathing simulator (ASL5000, Ingmar Medical, Pittsburgh, PA), used to generate adult breathing (tidal volume = 600-mL; duty cycle = 33%; rate = 10-cycles/min). Assay for salbutamol in both studies was by HPLC-UV spectrophotometry. RESULTS: Preliminary studies had confirmed linearity of FPM<4.7µm ex-VHC between 2 and 10 actuations. FPM<4.7µm/min for the BAN was independent of volume fill and linear with time until sputter. The table illustrates the relationships between ex VHC and treatment time ex BAN to achieve the same FPM<4.7µm from pMDI+VHC. Mean values are reported as coefficients of variation were <10%.

<table>
<thead>
<tr>
<th>Number of actuations</th>
<th>FPM&lt;4.7µm (µg)</th>
<th>Treatment time (min:sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>70</td>
<td>0:53</td>
</tr>
<tr>
<td>4</td>
<td>140</td>
<td>1:45</td>
</tr>
<tr>
<td>6</td>
<td>210</td>
<td>2:38</td>
</tr>
<tr>
<td>8</td>
<td>280</td>
<td>3:30</td>
</tr>
<tr>
<td>10</td>
<td>350</td>
<td>4:20</td>
</tr>
</tbody>
</table>

*values calculated based on measured FPM<4.7 µm of 33.2 ± 3.3 µg/actuation for 5-actuations

CONCLUSIONS: The ability to transition to and from pMDI + VHC to BAN offers the clinician new possibilities in titrating the adult tidal-breathing patient through exacerbations of broncho-constrictive diseases such as asthma or COPD, and easing the transition from hospital to the home environment.

EFFICACY OF TWO DIFFERENT SPACE HOLDING CHAMBERS IN THAI CHILDREN WITH ASTHMA: A PILOT STUDY. Kamalaporn H, Thongkum K, Preutthiphan A. Chest 2011;140:371A.
PURPOSE: To compare the clinical efficacy of a newly-designed spacer, AeroHaler® (Aerocare, Thailand) to an established space holding chamber, Aerochamber® (Trudell Medical, Canada) in Thai children with asthma. METHODS: A double-blinded, experimental study was conducted at the Pediatric Chest Clinic, Ramathibodi Hospital. We recruited known cases of asthma, aged 6-15 years with history of significant bronchodilator response by ATS criteria. Patients were double-blinded randomized into 2 groups with blocked allocation. The pulmonary function tests were performed at baseline. 4 puffs of salbutamol MDI was used to test bronchial reversibility. The patients received bronchodilator through either Aerochamber® or AeroHaler® on the first day of study and through another on the next day. The pulmonary function tests were repeated at 15 and 30 minutes after bronchodilator given. Significant bronchodilator response is identified as >12% increment of force expiratory volume in 1 second (FEV1) from baseline. The mean baseline FEV1 increment which reflects the efficacy of each space holding chamber in delivering bronchodilator was compared by paired T test. RESULTS: Twenty children with median age of 9.1 years (range 7.0-12.0) were enrolled. Thirteen children were male. The mean baseline FEV1 in patients using Aerochamber® was 75.1±17.9% predicted and 77.6±16.8 % predicted in patients using AeroHaler® (p=0.07). Using Aerochamber®, 10 children (50%) demonstrated bronchodilator response but only 4 of them (20%) showed bronchodilator response while using AeroHaler®. The FEV1 increment in patients using Aerochamber® was greater than that of AeroHaler® at both time points, 15 and 30 minutes, which maximal value found at 30 minutes. The mean increments of FEV1 using Aerochamber® and AeroHaler® were 9.97±9.36 % and 5.57±8.51 % respectively (p=0.02). CONCLUSIONS: As assessment of FEV1 increment to demonstrate bronchodilator response in Thai children with asthma, Aerochamber® is superior to AeroHaler®.

IMPORTANCE OF PATIENT-FRIENDLY FEATURES TO ADDRESS LACK OF INHALER COMPLIANCE: A LABORATORY EVALUATION OF AN INSPIRATORY FLOW INDICATOR AS A FEEDBACK AID FOR A VALVED HOLDING CHAMBER.


Purpose: Poor inhaler compliance is recognized as needing to be addressed. The Flow-Vu* Inspiratory Flow Indicator (IFI) is a feedback aid for those using the AeroChamber Plus* Flow-Vu* Anti-Static VHC (Trudell Medical Inc., London, Ontario). Regulators require that the modification does not affect delivery of the therapeutically beneficial fine particle dose < 4.7 µm diameter from the inhaler. Methods: Measurements (n=5 VHCs/group) of fine particle mass for salbutamol (100 µg/actuation) were made using an Andersen 8-stage impactor equipped with Ph.Eur. induction port and operated at 28.3 L/min. Data were obtained for the pMDI alone and for the pMDI +VHC (2-second delay), simulating poor coordination. The movement of the IFI monitored airflow through the VHC and a proper seal of the mouthpiece in the apparatus. The VHCs were tested out-of package in accordance with instructions. Recovery and assay for salbutamol was undertaken by HPLC-UV spectrophotometry. Results: Fine particle mass/actuation (FPM2s) for pMDI alone (mean±SD) was 34.8 ± 1.4 µg, compared with 33.2 ± 3.3 µg/actuation for the pMDI +VHC group. The IFI moved from the inhalation valve closed to open position immediately upon initiation of sampling. Conclusions: The IFI provided feedback on the delivery of this widely prescribed ‘rescue’ medication and did not interfere with the new VHC, delivering substantially comparable FPM2s to that from the pMDI alone. It should therefore aid patient compliance.


We have shown that the amount of urinary salbutamol excreted in the first 30 minutes (USAL0.5) represents the relative lung deposition and the 24 hour salbutamol plus its metabolite excretion (USAL24) indicates the total systemic delivery following an inhalation (Hindle and Chrystyn. Brit J Clin Pharmacol 1992; 34: 311-5). We have used these in-vivo methods together with in-vitro characterisation of the emitted dose using an Andersen Cascade Impactor (ACI) to compare the Volumatic (VOL) and AeroChamber Plus* (AERO). Spacers were attached to a salbutamol CFC free metered dose inhaler (MDI). 13 subjects, mean (SD) 31.2(7.6) years and 64.9 (10.9) Kg completed the in-vivo study.

The in-vitro and in-vivo results were:

<table>
<thead>
<tr>
<th>Mean (SD) from two 100µg doses (µg except MMAD µm)</th>
<th>MDI</th>
<th>MDI+VOL</th>
<th>MDI+AERO</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACI Spacer</td>
<td>74.9(6.1)</td>
<td>90.6(6.7)</td>
<td></td>
</tr>
<tr>
<td>TED</td>
<td>176.6(7.6)</td>
<td>94.9(4.6)</td>
<td>85.3(4.5)</td>
</tr>
<tr>
<td>Throat</td>
<td>93.6(7.4)</td>
<td>11.3(1.9)</td>
<td>11.7(1.2)</td>
</tr>
<tr>
<td>FDP</td>
<td>41.5(3.4)</td>
<td>41.8(2.3)</td>
<td>36.8(1.5)</td>
</tr>
<tr>
<td>MMAD</td>
<td>2.69(0.03)</td>
<td>2.76(0.07)</td>
<td>2.91(0.10)</td>
</tr>
<tr>
<td>Urinary salbutamol</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>USAL0.5</td>
<td>5.71(1.9)</td>
<td>16.36(8.2)</td>
<td>14.4(7.6)</td>
</tr>
<tr>
<td>USAL24</td>
<td>100.2(16.7)</td>
<td>97.3(12.7)</td>
<td>84.6(58.8)</td>
</tr>
</tbody>
</table>
TED - total emitted dose; Throat - AC I throat+S0+S1; FPD - fine particle dose, ACI S2-filter; MMAD - mass median aerodynamic diameter. Statistical analysis of the USAL0.5 data revealed no difference between the two spacers (mean difference [95% confidence interval] of 1.9[-4.5,8.3] μg). USAL 0.5 VOL and AERO were each greater (p<0.001) than MDI alone (mean difference [95%CI] of 10.6[4.2,17.1] and 8.7[2.3,15.1] μg, respectively). USAL24 amounts were all similar. The in-vitro characteristics suggest that slightly more salbutamol will be delivered to the lungs from a Volumatic than an AeroChamber Plus VHC. The in-vivo data confirms this but the difference, as predicted by the in-vitro data, is only small. The results are consistent with the smaller size of the AeroChamber Plus.


Previous studies have documented equivalent clinical efficacy of directly inhaled CFC and HFA albuterol MDIs but not whether use of a holding chamber alters this relationship. We compared albuterol delivery to the lungs by an HFA MDI with that of a CFC MDI when used in combination with an Aerocamber Plus* valved-holding chamber (VHC) using a methacholine challenge based bioassay. Seventeen subjects completed this double-blind, randomized, balanced cross-over study. Treatments were 1 or 2 actuations of albuterol CFC MDI (90 mcg/puff) or HFA MDI (100 mcg/puff). One of 4 treatments was administered during each study period with the AeroChamber Plus* VHC. A methacholine challenge (modified Juniper method) was initiated 15 minutes after albuterol administration. Results: (geometric mean PC20FEV1)

<table>
<thead>
<tr>
<th>1 Puff CFC</th>
<th>2 Puffs CFC</th>
<th>1 Puff HFA</th>
<th>2 Puffs HFA</th>
</tr>
</thead>
<tbody>
<tr>
<td>16.96</td>
<td>18.81</td>
<td>15.06</td>
<td>20.79</td>
</tr>
</tbody>
</table>

The dose-response relationship was significant (p=0.034) and parallelism and preparation contrasts were not significant (p=0.93, 0.27, respectively). The relative potency estimated using Finney 2-by-2 bioassay statistics was 0.97 (90% confidence interval [CI] 0.41-2.14). The 90% bias-corrected and accelerated percentile bootstrap CI for this estimate was 0.58-1.75. Removing an outlier from the data, the estimated relative potency was 1.04 (90% CI 0.65-1.73). Conclusion: HFA-and CFC-MDIs deliver equivalent quantities of albuterol to the lung when used with the AeroChamber Plus VHC.

Seretide® (Salmeterol & Fluticasone Propionate) GSK™ Inc.

We report the results from a laboratory study, in which the effect of transitioning a small volume VHC (145-mL) from non-conducting to electrostatic charge-dissipative polymer construction was demonstrated by determining the delivery of a two-component pMDI oral inhaled product comprising a long-acting bronchodilator (LABA) with inhaled corticosteroid (ICS). Measurements of fine particle mass between 1.1 and 4.7 μm aerodynamic diameter made by cascade impactor with 1, 2, and 5 delay between actuation of the inhaler and the onset of sampling have demonstrated substantially equivalent performance to that of the pMDI alone and with an earlier version of the VHC manufactured from non-conducting materials that required pre-washing after removal from its packaging in order to mitigate loss of medication due to electrostatic charge. The ability to use the new VHC without pre-washing is intended to make the device easier to use, with the ultimate goal of improving patient compliance with the prescribing clinician’s intent for therapy.


Thirty-one pediatric subjects (4-11 years) with asthma participated in an open-label, repeat dose, crossover study to compare serial concentrations of serum cortisol, fluticasone propionate (FP) and salmeterol (SAL) after three weeks of administration of Advair Diskus®, Advair® HFA, or Advair® HFA with the valved holding chamber (spacer), AeroChamber Plus®. Following a baseline assessment of serum cortisol and SAL pharmacodynamic parameters, 28 subjects completed the study with each subject receiving two of the three treatments using a randomized incomplete block design. FP systemic exposure was low after all treatments resulting in geometric mean (95% CI) Cmax of 54pg/mL (33,89) for Advair Diskus, 16pg/mL (8, 29) for Advair HFA and 35pg/mL (20, 60) for Advair HFA with spacer.


The treatment of both the bronchoconstriction and inflammatory aspects of asthma simultaneously by a single pressurized metered dose inhaler (pMDI) represents a significant advance in convenience to the patient. However, a valved holding chamber (VHC) may still be needed to reduce the coarse component of the dose that is likely to deposit in the oropharyngeal region, and a small sized device may offer significant advantages to the patient from the standpoint of compliance with therapy. VHCs representing small (adult AeroChamber Plus® with mouthpiece, 149-mL) and large (Volumatic™, 750-mL) devices have been compared in an in vitro evaluation with Seretide®/Advair® (hydro-fluoro alkane [HFA]-formulated fluticasone propionate [FP = 125 µg/dose] and salmeterol xinafoate [SX = 25 µg/dose]) by Andersen Mark-II eight-stage impactor operated at 28.3L/min following compendial methodology. Fine particle fraction, based on the size range from 1.1 to 4.7 μm aerodynamic diameter, from either large or small VHCs with either component (69-79%) was similar [p ≥ 0.08], and significantly greater than that from the pMDI alone (approximately 40%) [p < 0.001]. Fine particle dose emitted by the VHCs for SX (8.2 ± 0.8 μg for the AeroChamber Plus® and 7.7 ± 0.5 μg for the Volumatic™) were comparable, and also similar to the fine particle dose delivered without the pMDI when used without a VHC (7.6 ± 0.6 μg). Fine particle doses for the FP component delivered by the two VHCs (46.4 ± 3.4 μg for the AeroChamber Plus® and 46.3 ± 2.7 μg for the Volumatic™) were equivalent, but were slightly greater than the corresponding fine particle dose from the pMDI alone (39.1 ± 2.6 μg). However, this difference (approximately 20%) is close to the limit of resolution based on intermeasurement variability and is unlikely to have clinical significance, given the interpatient variability seen with inhaled drug therapy. It is therefore concluded that either of these VHCs has equivalent in vitro performance with this combination formulation in terms of the portion of the dose emitted from the pMDI that is likely to reach the receptors in the lungs.

**Serevent® (Salmeterol Xinafoate) GSK™ Inc.**


Salmeterol xinafoate is a widely prescribed long-acting beta-adrenergic agonist. Valved holding chambers (VHCs) improve drug delivery from pressurized metered-dose inhalers (pMDI), particularly with patients having poor coordination. The present study compared a large volume VHC (Volumatic™, GlaxoSmithKline - 750-mL, n=5 devices) with a small volume VHC (AeroChamber Plus®, Monaghan Medical Corp. - 149-mL, n=5 devices) with salmeterol xinafoate (Serevent®): total dose 21 μg ex actuator, GlaxoSmithKline). Measurements were also made with the pMDI without VHC. Total emitted dose (TD), fine particle dose (FPD - particles < 4.7 μm aerodynamic diameter) and fine particle fraction (FPF) were determined by Andersen 8-stage impactor with USP Induction Port at 28.3 ± 0.5 L/min. Assays for salmeterol xinafoate were undertaken by HPLC-fluorescence spectrometry at excitation and emission wavelengths of 226 nm and 296 nm respectively. As expected, both types of VHC greatly reduced the coarse component of the dose from the pMDI (10.5 ± 1.2 μg - pMDI alone; 1.0 ± 0.6 μg - AeroChamber Plus® VHC; 0.9 ± 0.7 μg - Volumatic™ VHC). Both FPD and TD from the AeroChamber Plus® (12.7 ± 1.3 μg and

STUDY OBJECTIVE: To determine whether a spacer device designed as a valved holding chamber with a flow signal increases the efficacy of the long-acting beta(2)-agonist, salmeterol, in patients who use incorrect technique with metered-dose inhaler (MDI) alone. DESIGN: Double-blinded, randomized, placebo-controlled study. SETTING: University hospital outpatient rooms. PATIENTS: Twenty adult outpatients with stable persistent asthma, receiving a daily anti-inflammatory drug.

INTERVENTIONS: Patients were randomized to either salmeterol MDI (incorrect use: 1 s after actuating MDI, inhale rapidly) and placebo plus spacer (correct use: inhale slowly as MDI is actuated, continue to inhale slowly and deeply) or placebo MDI (incorrect use) and salmeterol plus spacer (correct use). The following week, patients received the opposite treatment. The dose was two puffs from each device on each treatment day; each puff was separated by 1 min. MEASUREMENTS AND RESULTS: After baseline peak expiratory flow (PEF), salmeterol was administered and serial PEF determined (0.5, 1, 2, 3, 4, 6, 8, 10, and 12 h). Administration of salmeterol MDI plus spacer resulted in significantly greater increases in PEF from baseline vs MDI at 4 h (44 L/min vs 10 L/min; p < 0.01) and 6 h (49 L/min vs 24 L/min; p < 0.05). Both methods of administration were equally well tolerated. CONCLUSION: We conclude that patients who have poor timing and rapid inhalation with salmeterol MDI alone will have greater increases in PEF at 4 h and 6 h and no additional side effects if the dose is administered with a valved holding chamber that is used correctly. Further study is needed regarding other errors in MDI technique with salmeterol.

Symbicort™ (Budesonide / Formoterol) AstraZeneca™


Objective: Spacers and valved holding chambers (VHCs) were developed to facilitate using pressurized metered dose inhaler (pMDI) by patients who could not coordinate the actions required for successful pMDI use. There is little in vivo evidence about how VHC may affect the bronchodilation from combination drugs in pMDI. This study was to determine the effect, if any, of VHC on the bronchodilating actions of the pMDI budesonide/formoterol combination. Methods: Sixteen adult asthmatic subjects with 15% or greater reversibility of forced expiratory volume in one second (FEV-1), 15 minutes after inhaling 180-360 µg of albuterol, participated. The study had a randomized crossover design with each subject using budesonide-formoterol pMDI as described in the product information one time and on a second occasion using the pMDI with a VHC. Spirometry and impedance oscillometry were measured at baseline and repeatedly over a 12-hour period. This study was approved by IntegReview Institutional Review Board, Austin, TX, USA. The clinical trial number for this study was NCT 009-15538 (http://www.clinicaltrials.gov). Results: The area under the curve of FEV-1, the FEV-1, and the fraction FEV-1/FVC was similar over the 12-hour time frame with both methods. Resistance was not different at any time point. In both procedures, the onset of bronchodilation occurred rapidly within 3 to 5 minutes. Conclusions: In well-trained asthmatic subjects, both tested methods caused equivalent bronchodilation. This suggests a VHC itself has no deleterious effect on the bronchodilator activity in the combined drug. These results may not apply to patients who have coordination problems with the pMDI and further study is indicated.

USE OF A VALVED HOLDING CHAMBER DOES NOT ADVERSELY AFFECT BRONCHODILATION FROM Budesonide/Formoterol Pressurized Meter Dose Inhaler (pMDI). Mansfield LE, Cueto E, Maynes R, Romero E. Journal of Allergy and Clinical Immunology 2010;125(2):S1.

RATIONALE: pMDI are often used clinically with valved holding chambers to facilitate patient coordination and use. There is no available information about the effect of using the valved holding chamber on the bronchodilator activity of combination pMDI of formoterol and budesonide. This study addresses this issue. METHODS: 16 adult asthmatics with demonstrated 15% or greater increase in FEV-1 after 2-4 inhalations of albuterol 90µg were studied in a randomized crossover design. Pulmonary functions were measured for 12 hours after using Symbicort™ pMDI, 160/4.5 in usual fashion or with a valved holding chamber (Aerochamber Plus’). Subjects inhaled from valved holding chamber immediately after actuation of pMDI. 12 hour area under the curve was determined for FEV-1 in liters, and FEV-1/FVC and individual time points compared. RESULTS: AUC FEV-1 pMDI 1937.1 Liters, pMDI plus chamber 1920.7 liters, AUC FEV-1/FVC pMDI 58417%, pMDI plus chamber 57830%. There were no significant differences at any time point. CONCLUSIONS: Use of a valved holding chamber does not adversely affect the bronchodilating activity of formoterol in the budesonide/formoterol combined pMDI when used by adult asthmatics.

A combination of budesonide and formoterol in a single pressurized metered-dose inhaler (pMDI) is available in the United States and elsewhere. This study was designed to evaluate the delivered dose and fine particle dose (FPD; mass of particles <4.7-micrometer diameter) using the pMDI with two valved holding chambers (VHCs), using sampling methods reflecting different patient techniques. FPD was measured using an Andersen Cascade Impactor and delivered dose was measured using a disposable filter. Two VHCs, AeroChamber Plus* and AeroChamber MAX* (Trudell Medical International, London, Ontario, Canada), were evaluated using three sampling methods: (1) immediate collection; (2) collection after up to a 5-second delay; (3) using simulated adult, child, and infant tidal breathing patterns (delivered dose). Decreases in delivered dose were observed using a VHC compared with the pMDI alone. FPD with both VHCs was similar to that with the pMDI alone with minimal delay between actuation and collection. With delays, the antistatic AeroChamber MAX* was more resistant than AeroChamber Plus* to dose losses. Delivered doses from adult and child profiles were comparable with those after a 1-second delay. The infant profile produced lower delivered doses, probably because more breath cycles are required to empty the VHC. Budesonide/formoterol pMDI can be used effectively with AeroChamber Plus* and the antistatic AeroChamber MAX*. With minimal delay between actuation and collection, FPD with both VHCs was similar to that with the pMDI alone, giving physicians a choice of administration regimen and taking into account the needs and skills of the patient.


Background: Symbicort (budesonide/formoterol) Rapihaler® is a novel pressurised HFA metered dose inhaler. This study aimed to evaluate the in vitro delivery of budesonide and formoterol via Symbicort Rapihaler from 2 spacer devices: the NebuChamber® and the AeroChamber Plus* valved holding chamber. Methods: The particle size distribution was analysed using Next Generation Impactor (NGI) at 30 L/min. The effect of different dose regimens on fine particle dose (FPD) was investigated. The following regimens using Symbicort 80/4.5 μg were assessed: single actuation (act) with a 2-s delay between act and collection (1 act, 2-s delay) and 2 acts with a 2-s delay between acts and collection (2 acts, 2-s delay). Six acts were collected/test. Results: The graph shows FPD (as a % of nominal dose) for budesonide. Similar results were observed for formoterol. Conclusion: This in vitro study shows that the fine particle dose from Symbicort Rapihaler pMDI is similar when using either NebuChamber or AeroChamber Plus*.

Ventolin® (Salbutamol) GSK™ Inc.


The AeroChamber Plus® VHC (Trudell Medical International, London, Canada) is a widely prescribed add-on device for patients having poor coordination with their pressurized metered-dose inhaler (pMDI). Changes have been made to its design to make it more patient-friendly, so that compliance with prescribed inhaled medication can be improved. However, there may be concern that these developments may affect medication delivery performance, where the intent is to match as closely as possible the therapeutically important metrics, emitted extra-fine (EPM<1.1μm) and fine particle mass (FPM<4.7μm) ex VHC, with those from the pMDI alone. We report a laboratory study in which both measures were compared, simulating a 2-s delay between pMDI operation and the onset of sampling via each VHC. Published data for the pMDI alone (no delay), the original VHC (non conducting (NC) and non conducting with inspiratory flow indicator (NC-IFI)) were compared with new data for the anti-static, take-apart with IFI (AS). Both non-conducting device groups were pre-washed in ionic detergent and drip-dried in accordance with manufacturer instructions to mitigate surface electrostatic charge, whereas the AS group was evaluated out-of-package without pretreatment. We evaluated these devices with Ventolin®, representing a formulation known to have a high degree of electrostatic charge. We found both measures of performance were consistent between VHCs (EPM<1.1μm = 4.7±0.8 μg (NC); 3.1±0.5 μg (NCIFI); 3.9±2.1 μg (AS); FPM<4.7μm = 36.3±1.8 μg (NC); 30.9±2.0 μg (NC-IFI); 33.4±4.2 μg (AS)). The more important FPM<4.7μm was within ±15% of the benchmark value for the pMDI alone (FPM<4.7μm = 34.8±1.4 μg).

BACKGROUND: Airway obstruction and bronchial hyperactivity often times lead to emergency department visits in infants. Inhaled short-acting beta2-agonist bronchodilators have traditionally been dispensed to young children via nebulizers in the emergency department. Delivery of bronchodilators via metered-dose inhalers (MDIs) in conjunction with holding chambers (spacers) has been shown to be effective. STUDY OBJECTIVE: Safety and efficacy evaluations of albuterol sulfate hydrofluoroalkane (HFA) inhalation aerosol in children younger than 2 years with acute wheezing caused by obstructive airway disease. METHODS: A randomized, double-blind, parallel group, multicenter study of albuterol HFA 180 microg (n = 43) or 360 microg (n = 44) via an MDI with a valved holding chamber and face mask in an urgent-care setting. Assessments included adverse events, signs of adrenergic stimulation, electrocardiograms, and blood glucose and potassium levels. Efficacy parameters included additional albuterol use and Modified Tal Asthma Symptoms Score (MTASS) reduction in MTASS representing improvement. RESULTS: Overall, adverse events occurred in 4 (9%) and 3 (7%) subjects in the 180-microg and 360-microg groups, respectively. Drug-related tachycardia (360 microg) and ventricular extrasystoles (180 microg) were reported in 1 patient each. Three additional instances of single ventricular ectopy were identified from Holter monitoring. No hypokalemia or drug-related QT or QTc prolongation was seen; glucose values and adrenergic stimulation did not significantly differ between treatment groups. In the 180-microg and 360-microg groups, mean change from baseline in MTASS during the treatment period was -2.8 (-49.8%) and -2.9 (-48.4%), and rescue albuterol use occurred in 4 (9%) and 3 (7%) subjects, respectively. CONCLUSIONS: Cumulative dosing with albuterol HFA 180 microg or 360 microg via MDI-spacer and face mask in children younger than 2 years did not result in any significant safety issues and improved MTASS by at least 48%.


Body: Delivery of inhaled medication to infants/small children by VHC-facemask can be difficult to verify. An external visual aid (Flow-Vu*) is available with the AeroChamber Plus* (Trudell Medical International) VHCs as an inspiratory flow indicator (IFI) to aid compliance with instructions for use. We report an in vitro study in which delivery of salbutamol (Ventolin®; 100µg/actuation, GSK plc) was measured using infant and child models (ADAM-II), in which the soft facial tissues are modeled where the facemask makes contact. The facemask was applied with an appropriate force of 1.6 kg, and tidal breathing was simulated (tidal volume (Vt) 50 ml, 30 bpm, 25% duty cycle - VHC-infant facemask; Vt = 155 ml; 25 bpm, 33% duty cycle - VHC-child facemask (n=5 devices/group)). Total emitted mass (TEM) of salbutamol was collected by filter located behind the lips after 1, 2, 3, 4 and 5 inhalations.

<table>
<thead>
<tr>
<th>Number of Inhalations</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infant</td>
<td>5.8 ± 2.2</td>
<td>13.0 ± 4.0</td>
<td>13.8 ± 3.8</td>
<td>14.5 ± 3.2</td>
<td>15.6 ± 3.6</td>
</tr>
<tr>
<td>Child</td>
<td>15.9 ± 3.7</td>
<td>17.6 ± 4.6</td>
<td>20.1 ± 3.5</td>
<td>20.8 ± 2.9</td>
<td>22.6 ± 4.2</td>
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At least two inhalations were required to achieve consistent medication delivery from the VHC-infant facemask. The first inhalation was sufficient to achieve similar consistency with the VHC-child facemask. However, these tests were undertaken with a well-fitting facemask and no leakage. Manufacturer instructions indicate 5-inhalations be taken as a precaution. The IFI validates an effective seal between facemask-face as well as confirms the number of inhalations, assisting in compliance with instructions.


Introduction: Inhaled short-acting bronchodilators are recommended for the quick relief of bronchospasm symptoms in children including those less than five years of age. However, limited safety data is available in this young population. Methods: Safety data were analyzed from a randomized, double-blind, parallel group, placebo-controlled multicenter, study evaluating albuterol HFA 90µg or 180µg versus placebo three times a day for 4 weeks using a valved holding chamber, Aerochamber Plus* and facemask in children birth ≤24 months old with a history of bronchospasm. Results: The overall incidence of adverse events (AE) during treatment was: albuterol 90µg (59%), albuterol 180µg (76%) and placebo (71%). The most frequently reported AEs were pyrexia in 7 (24%), 2 (7%), and 3 (11%) subjects in the albuterol 180µg, albuterol 90µg, and placebo groups, respectively. Upper respiratory tract infection (URTI) occurred in 5 (17%) and 3 (11%) subjects in the albuterol 180µg and placebo groups, respectively. Sinus tachycardia occurred in 5 (17%), 2 (7%) and 2 (7%) subjects receiving albuterol 180µg, albuterol 90µg and placebo, respectively. One subject in each of the albuterol treatment groups experienced drug related agitation and/or restlessness or mild sinus arrhythmia. No drug-related QT prolongation or abnormal serum potassium and glucose levels were reported in the albuterol treatment groups. Conclusion: This study provides additional albuterol HFA safety information for the treatment of children aged birth ≤24 months with a history of bronchospasm.
A NEW VALVED HOLDING CHAMBER (VHC) FOR YOUTH THAT IS MANUFACTURED FROM ELECTROSTATIC CHARGE-DISSIPATIVE MATERIALS PROVIDES EFFICIENT DELIVERY OF MEDICATION IF USED OUT OF THE PACKAGE


Children prescribed a VHC are often reluctant to take inhaled medication for asthma in front of their peers. A new line of equivalent anti-static VHCs (AC Girl* and AC Boyz* holding chambers), based on the AeroChamber Plus* anti-static VHC with Flow-Vu* Inspiratory Flow Indicator (IFI) (Trudell Medical International, London, Canada), has been developed with attractive youth-relevant markings to improve compliance. We report a study in which delivery of Ventolin®-HFA via these VHCs (n=3), was compared with VHCs manufactured from nonconduction polymer (OptiChamber® Advantage, Respironics Inc., Cedar Grove, NJ, USA (n=5 VHCs)) as benchmark devices. Both VHC groups were evaluated immediately after removal from their packaging to simulate use in an urgent care situation, but the OptiChamber® Advantage VHCs were also studied after pre-washing in mild detergent, rinsing and drip-drying in accordance with instructions to minimize electrostatic charge. A proprietary apparatus that interfaced between the VHC mouthpiece and induction port leading to an 8-stage Andersen cascade impactor was used to simulate a 2-s delay between pMDI actuation and the onset of sampling at 28.3 L/min, representing a typical uncoordinated user. Reference data were also obtained with no delay. Fine particle mass (FPM (μg/actuation; mean ± SD)), based on particles < 4.7 μm aerodynamic diameter, was 52.7 ± 5.5 μg and 37.7 ± 3.4 μg for the AC Girl*/AC Boyz* Chambers without and with delay respectively. Equivalent values for the OptiChamber® Advantage VHCs were 29.5 ± 4.3 μg (no delay) and 8.9 ± 2.1 μg (delay) when tested out-of-package and 38.5 ± 3.7 μg (no delay) and 19.7 ± 3.0 μg (delay) after pre-washing. The AC Girl*/AC Boyz* Chambers significantly outperformed the OptiChamber® Advantage VHC group (un-paired t-test at each delay interval, p < 0.001), even when pre-washed. This study indicates the advantage of charge dissipative materials to avoid electrostatic charge-related losses.


Delivery of inhaled medication to infants by valved holding chamber (VHC) with facemask may require more than one inhalation to empty the VHC because tidal volumes are typically smaller than chamber capacity. This study investigated the correlation between movement of an integrated inspiratory flow indicator (IFI) as a caregiver feedback aid for a VHC-facemask, number of inhalations and mass of medication, simulating use by a 6-9 month infant (tidal volume (Vt) = 50-ml; duty cycle = 25%; 30 cycles/min). Anti-static AeroChamber Plus* VHCs incorporating the IFI feature, with infant facemask (n=5/group, 3 replicates/device; Trudell Medical International, London, Canada) were coupled to a breathing simulator (ASL5000 test lung, IngMar Medical, Pittsburgh, PA, USA). The VHCs were prepared as per manufacturer instructions and the facemask of the device on test was fitted to the ADAM-II flexible infant face model with a clinically appropriate force of 1.6 kg. Aerosol capture took place using an electret filter positioned behind the lips of the face model. Delivery of medication was evaluated from two different pressurized metered dose inhaler formulations likely to be used with paediatric patients (Flovent® HFA 44; 44 μg fluticasone propionate (FP) delivered ex-actuator and Ventolin® HFA; 90 μg salbutamol base equivalent (SAL) delivered ex-actuator, both from GSK plc. One actuation was delivered to the VHC at the onset of inhalation, and the filter removed after 1 complete breathing cycle, observing the movement of the IFI to confirm inhalation valve opening. This procedure was subsequently repeated by removing the filter after 2, 3, 4, 5 and 6 breathing cycles. Assay for FP or SAL was undertaken by HPLC-UV spectrophotometry. During these measurements, the IFI of each device was observed to move in synchrony with valve opening on all occasions, confirming that the facemask sealed onto the face model without leakage of ambient air into the mask during the inspiratory phase of each breathing cycle. Emitted mass after the first breathing cycle (EM1) was 2.1 ± 0.7 μg (FP) and 5.8 ± 2.2 μg (SAL); substantially lower than the corresponding values after 6 cycles (EM6), being 9.0 ± 2.1 μg (FP), and 15.9 ± 3.1 μg (SAL) [paired t test for each formulation; p < 0.001]. After 2 breathing cycles, values of EM2 (6.9 ± 2.0 (FP) and 13.0 ± 4.0 μg (SAL)), though significantly greater that their corresponding EM1 values [p ≤ 0.002], were still noticeably lower than the corresponding EM6 value for FP (p = 0.028), and barely statistically insignificant for SAL (p = 0.063). After 3 inhalations, EM3 increased further to 7.6 ± 2.0 μg (FP) and 13.8 ± 3.8 μg (SAL), and thereafter were close to the corresponding EM6 values, indicating emptying of the VHC had taken place. We conclude that at least two successive inhalations are required to achieve optimum medication delivery for the ‘infant’ condition under optimum conditions with a well fitted facemask with no leakage. The IFI is an important feature which validates that the facemask is properly sealed to the infant’s face and also confirms the number of inhalations that take place, thereby optimizing the therapeutic dose. Clinical studies are recommended to evaluate the benefit of this aid for the delivery of inhaled medication by VHC to this age group.


Differences between the size and shape of spacers may affect the emitted dose and provide different effects when interchanged during routine use. Using a urinary pharmacokinetic method we have measured the relative lung and systemic
bioavailability from urinary salbutamol excretion 30 min (USAL0.5) and 24 h (USAL24), respectively, after the inhalation of two 100-mg doses from a Ventolin Evohaler when used alone (MDI) and when attached to the Volumatic (VOL) or the AeroChamber Plus* (AERO) spacers. The in-vitro properties of the emitted dose were determined. The mean (s.d.) USAL0.5 values following MDI, VOL and AERO (n = 13 volunteers) were 5.7 (1.9), 16.4 (8.2) and 14.8 (7.4) mg, respectively. VOL and AERO were significantly greater (P < 0.001 and < 0.01, respectively) than MDI. Comparison of VOL and AERO was similar with a mean ratio (90% confidence interval) of 108.2 (84.5, 138.6)%.

USAL24 values between the three inhalation methods were similar. The values for the mean (s.d.) fine particle dose of two 100-mg doses emitted from MDI, VOL and AERO were 83.0 (6.8), 83.6 (4.6) and 73.6 (2.9) mg and the mass median aerodynamic diameters were 2.7 (0.03), 2.8 (0.07) and 2.9 (0.10) mm, respectively. The results showed that during routine use the Volumatic and the AeroChamber Plus* spacers should provide similar lung and systemic delivery when attached to a Ventolin Evohaler.

Zenhal™ (Mometasone Furoate / Formoterol Fumarate) Merck Sharp


Rationale: The bronchodilatory effect of mometasone furoate/formoterol fumarate (MF/F) administered by metered-dose inhaler (MDI) with or without spacer has not been evaluated previously in children. Methods: This was a randomized, multicenter, placebo (PBO)-controlled, single-dose, 4-period crossover study. Children with persistent asthma aged 5–11 y participated in this study. Subjects used inhaled corticosteroids with/without long-acting β2-agonists for ≥ 12 wk before enrollment and had FEV1 ≥ 70% predicted at screening. Subjects received MF/F-MDI 100/10 μg with/without spacer, F-dry powder inhaler (DPI) 10 μg, and PBO-MDI with/without spacer in separate treatment periods. The primary end point was FEV1 AUC0-12h for the comparison of MF/F with spacer vs PBO. Secondary measurements included MF/F without spacer vs PBO as well as MF/F with spacer vs MF/F without spacer and F-DPI vs PBO. Analysis was performed with an ANCOVA model for a crossover study. Results: A total of 87 subjects completed treatment, and 79 subjects were in the per-protocol analysis set. MF/F with spacer demonstrated a larger change in mean FEV1 AUC0-12h vs PBO (115 vs − 9 mL), with a treatment difference of 124 mL (95% CI, 94 to 154; P < .001). Similarly, MF/F without spacer vs PBO resulted in a 102 mL difference in mean adjusted FEV1 AUC0-12h (95% CI, 73 to 131, P < .001), whereas the difference between MF/F with spacer vs MF/F without spacer was 22 mL (95% CI, − 8 to 52, P = .144). The difference between F-DPI vs PBO was 106 mL (95% CI, 77 to 135, P < .001). No unexpected adverse events were observed. Conclusions: In this trial, MF/F-MDI 100/10 μg demonstrated significant bronchodilation in children aged 5–11 y regardless of the use of a spacer. Similar bronchodilatory profiles were observed for F delivered by DPI and by MDI in combination with MF.
Large versus small volume valved holding chambers


Background: To which extent volume spacers may influence systemic activity of inhaled beclomethasone dipropionate (BDP) has not been evaluated. Aim: To assess whether the AeroChamber Plus (™) spacer is equivalent to the Volumatic (™) spacer for administration of inhaled hydrofluoralkane 134a propelled BDP in terms of lower leg growth rate (LLGR). Patients and methods: Prepubertal children with mild asthma (n=26, ages 6-14 years) were included in a 3-time periods of 2 weeks duration randomised double-blind cross-over study with a single-blind placebo run-in and 2 wash-out periods. LLGR was measured with the knemometer. Interventions were inhaled BDP hydrofluoralkane 134a pMDI 100 µg and 200 µg b.i.d. with the AeroChamber Plus and 200 µg b.i.d. with the Volumatic spacer. Results: BDP 200 µg b.i.d. from the AeroChamber Plus was non-inferior to BDP 200 b.i.d. from the Volumatic spacer as the lower margin of confidence interval of the difference between treatments (-0.18 to 0.13 mm/week) was greater than the pre-specified lower limit for non-inferiority (-0.20 mm/week). UFC/creatinine data showed no statistically significant variations. Conclusion: The systemic activity of BDP via the Volumatic (™) and AeroChamber Plus (™) spacers is similar. The AeroChamber Plus spacer may be used in children without risk of increasing systemic activity of BDP.


Background: The study objective of this pilot study was to determine the lung delivery of HFA-134abeclomethasone dipropionate (HFA-BDP; QVAR) and CFC-beclomethasone dipropionate (CFC-BDP; Becloderm) with and without the add-on spacers, AeroChamber®, and Volumatic™. The smaller particles of HFA-BDP were presumed to produce greater lung deposition using spacers, with and without a delay [i.e., metered dose inhaler (MDI) actuation into the spacer and subsequent inhalation 0 and 2 sec later], compared with the larger particles of CFC-BDP. The study included a comparison of breathhold effects (i.e., 1 and 10-sec breathholds) on lung deposition. Methods: The study was an open-label design and utilized healthy subjects (n=12 males). Each arm of the study contained three subjects; thus, outcomes were not powered to assess statistical significance. HFA-BDP and CFCBDP were radiolabeled with technetium-99 m and delivered to subjects. Results: Results showed that the small particle HFA-BDP lung deposition averaged 52% and was not affected by the use of AeroChamber® with or without a spacer delay. The oropharyngeal deposition of HFA-BDP was reduced from approximately 28% to 4% with the AeroChamber®. Lung deposition with the large particle CFC-BDP was 3–7% and generally decreased with AeroChamber® or Volumatic. A 2-sec time delay between actuation and breath plus the spacer reduced lung deposition slightly but reduced oropharyngeal deposition substantially (84% down to 3–20%) using the AeroChamber® or Volumatic with and without a spacer delay. HFA-BDP lung deposition was dependent on the breathhold. Lung deposition with HFA-BDP was reduced by 16% with a 1-sec versus 10-sec breathhold. The difference was measured in the increased exhaled fraction, confirming that smaller particles need time to deposit and are exhaled if there is a reduced breathhold. The large particle CFC-BDP lung deposition was not affected by breathhold. Conclusions: The use of AeroChamber® or Volumatic spacers with HFA-BDP did not alter lung deposition but it did reduce oropharyngeal deposition. However, HFA-BDP displayed reduced oropharyngeal deposition without a spacer.


Differences between the size and shape of spacers may affect the emitted dose and provide different effects when interchanged during routine use. Using a urinary pharmacokinetic method we have measured the relative lung and systemic bioavailability from urinary salbutamol excretion 30 min (USAL0.5) and 24 h (USAL24), respectively, after the inhalation of two 100-mg doses from a Ventolin Evohaler when used alone (MDI) and when attached to the Volumatic (VOL) or the AeroChamber Plus® (AERO) spacers. The in-vitro properties of the emitted dose were determined. The mean (s.d.) USAL0.5 values following MDI, VOL and AERO (n = 13 volunteers) were 5.7 (1.9), 16.4 (8.2) and 14.8 (7.4) mg, respectively. VOL and AERO were significantly greater (P < 0.001 and < 0.01, respectively) than MDI. Comparison of VOL and AERO was similar with a mean ratio (90% confidence interval) of 108.2 (84.5, 138.6)%. USAL24 values between the three inhalation methods were similar. The values for the mean (s.d.) fine particle dose of two 100-mg doses emitted from MDI, VOL and AERO were 83.0 (6.8), 83.6 (4.6) and 73.6 (2.9) mg and the mass median aerodynamic diameters were 2.7 (0.03), 2.8 (0.07) and 2.9 (0.10) mm, respectively. The results showed that during routine use the Volumatic and the AeroChamber Plus® spacers should provide similar lung and systemic delivery when attached to a Ventolin Evohaler.
Introduction: Factors affecting dose delivery from pMDIs fitted with add-on devices include formulation, device design (e.g., materials, size, incorporation of a non-return valve), cleaning procedures and use-mode. Spacer-mode involves a conventional press-and-breathe manoeuvre whilst inhaling through the mouthpiece of the pMDI-device assembly. The spacer creates a longer path-length, allowing more time for propellant evaporation and slowing the cloud to facilitate lung access. Use in holding-chamber mode requires the device to have a non-return valve and discharging the dose into the chamber where it is held for a period before being inhaled, eliminating the need for press-and-breathe co-ordination. This mode also permits the patient to carry out repeated inhalations from the same dose. We were interested in comparing the effects of different add-on devices and their mode of use because the large size of some holding chamber devices may deter user acceptability. This study compares the dose delivered when beclometasone dipropionate (BDP) HFA solution type pMDIs were used in conjunction with both small and large volume devices in the two modes. Conclusions: These in vitro results would imply that, when used by patients in association with AeroChamber-Plus, the drug delivery performance for Modulite-BDP pMDIs could be similar to that obtained with Volumatic up to holding times of at least 5s for all three product strengths and up to 10s for the 50 µg and 100 µg dose strengths.


RATIONALE: Variability in the clinical use of inhaler devices is high, particularly in children. Optimisation of inhalation therapy should ensure more consistent dose delivery to the airways of young children. We assessed the effect of spacer volume, inhalation technique and training of the parent/child on drug delivery to children using pressurised inhalers. METHODS: Albuterol was delivered via large (Volumatic; VOL) and small (AeroChamber Plus* VHC; AC+) spacers to 21 children (2-14yrs). Children ≥5yrs either took 5 tidal breaths, or one slow maximal inhalation with 10 sec breath-hold. Children <5yrs used tidal breathing only. Training sessions were scheduled ≥12wks apart. Drug delivery was assessed using a low resistance filter attached to the spacer mouthpiece. RESULTS: Mean (SD) drug delivery (% nominal dose) to children of all ages using AC+ [51.5 (14.7)%] was significantly higher (p=0.04) than using VOL [39.3 (10.1)%]. Mean (SD) drug delivery using the single maximal inhalation technique [45.4 (13.7)%] was significantly higher (p=0.01) than that using tidal breathing [32.3 (13.9)]. The improvement in delivery using the single maximal inhalation was most marked in the 5-7yr age group. Training the parent/child to use the spacer correctly gave a small (3.9%) but significant increase (p=0.04) in drug delivery. CONCLUSIONS: AC+ (small volume) delivered more drug than VOL (large volume). This is possibly due to the more efficient construction and design of the AeroChamber Plus* as delivery is normally improved when using large volume spacers. The single maximal inhalation technique increased drug delivery to patients compared to tidal breathing. However, it is easier for children <5yrs to use the tidal breathing technique. Training of the parent/patient resulted in a smaller than expected (albeit significant) increase in drug delivery.


The treatment of both the bronchoconstriction and inflammatory aspects of asthma simultaneously by a single pressurized metered dose inhaler (pMDI) represents a significant advance in convenience to the patient. However, a valved holding chamber (VHC) may still be needed to reduce the coarse component of the dose that is likely to deposit in the oropharyngeal region, and a small sized device may offer significant advantages to the patient from the standpoint of compliance with therapy. VHCs representing small (adult AeroChamber Plus* with mouthpiece, 149-mL) and large (Volumatic™, 750-mL) devices have been compared in an in vitro evaluation with Seretide®/Advair™ (hydro-fluoro alkane [HFA]-formulated fluticasone propionate [FP = 125 µg/dose] and salmeterol xinafoate [SX = 25 µg/dose]) by Andersen Mark-II eight-stage impactor operated at 28.3L/min following compendial methodology. Fine particle fraction, based on the size range from 1.1 to 4.7 um aerodynamic diameter, from either large or small VHCs with either component (69-79%) was similar [p ≥ 0.08], and significantly greater than that from the pMDI alone (approximately 40%) [p < 0.001]. Fine particle dose emitted by the VHCs for SX (8.2 ± 0.8 µg for the AeroChamber Plus* and 7.7 ± 0.5 µg for the Volumatic™) were comparable, and also similar to the fine particle dose delivered by the pMDI when used without a VHC (7.6 ± 0.6 µg). Fine particle doses for the FP component delivered by the two VHCs (46.4 ± 3.4 µg for the AeroChamber Plus* and 46.3 ± 2.7 ug for the Volumatic™) were equivalent, but were slightly greater than the corresponding fine particle dose from the pMDI alone (39.1 ± 2.6 µg). However, this difference (approximately 20%) is close to the limit of resolution based on intermeasurement variability and is unlikely to have clinical significance, given the interpatient variability seen with inhaled drug therapy. It is therefore concluded that either of these VHCs has equivalent in vitro
performance with this combination formulation in terms of the portion of the dose emitted from the pMDI that is likely to reach the receptors in the lungs.

**RANDOMISED CONTROLLED STUDY OF CLINICAL EFFICACY OF SPACER THERAPY IN ASTHMA WITH REGARD TO ELECTROSTATIC CHARGE.** Dompeling E, Oudesluys-Murphy AM, Janssens HM, Hop W, Brinkman JG, Sukhai RN, de Jongste JC. Arch. Dis. Child 2001;84;178-82.

BACKGROUND: Inhalation therapy using a pressured metered dose inhaler (pMDI) and a spacer is frequently used in the treatment of airway disease in children. Several laboratory studies found a clear negative influence of electrostatic charge (ESC) on plastic spacers on the delivery of aerosol. AIMS: To investigate whether ESC on plastic spacers could diminish bronchodilating responses to salbutamol. METHODS: Ninety asthmatic children (aged 4-8 years) were randomised into three groups: metal Nebuchamber, plastic Volumatic, and plastic AeroChamber*. The bronchodilating response was measured by the change in peak expiratory flow rate (PEF) after 100 microgram and 400 microgram salbutamol. Within the Volumatic and AeroChamber* groups, a crossover comparison was made between electrostatic and non-electrostatic spacers. RESULTS: We found no significant effect of ESC on the bronchodilating response to salbutamol with any of the doses in the AeroChamber* and Volumatic groups. For the plastic spacers, the mean difference of the change in PEF after 100 microgram salbutamol between non-electrostatic and electrostatic spacers was only +1.7% (95% CI -1.3% to 4.7%). After 400 microgram salbutamol this was +1.9% (95% CI -1.4% to 5.1%). A comparable efficacy was found for the Nebuchamber, the AeroChamber*, and Volumatic with respect to the change in PEF after 100 and 400 microgram salbutamol. CONCLUSION: This study showed no negative influence of ESC on plastic spacers with regard to clinical efficacy of a beta(2) agonist (salbutamol) in children with asthma. The metal Nebuchamber, plastic AeroChamber*, and plastic Volumatic were equally effective.


It is useful from the standpoint of the health care provider, if the performance of add-on devices for use with pressurized metered dose inhalers is characterized within the range of flow rates likely to be achieved by users. VHCs representing smaller (adult AeroChamber Plus*, 149-ml; n=5) and larger (Volumatic™, 750-ml; n=5) devices were compared with HFA-formulated fluticasone propionate (125 µg/dose ex metering chamber) at three flow rates, 28.3, 45 and 60 L/min. Measurements were made by Andersen 8-stage impactor. Fine particle fractions (< 4.7 µm, < 4.6 µm and < 4.0 µm aerodynamic diameter at 28.3, 45 and 60 L/min respectively) from both VHCs were close to 90%, significantly greater than that from the pMDI alone. At 28.3 L/min, fine particle dose (FPD) from the smaller VHC (50.5 ± 3.8 µg) was comparable with that from the larger VHC (45.9 ± 7.8 µg) [p = 0.27]. At the higher flow rates, FPD from the smaller VHC (65.5 ± 2.6 µg (45 L/min) and 65.2 ± 6.2 µg (60 L/min)) exceeded equivalent values from the larger VHC (53.8 ± 3.7 µg (45 L/min) and 55.3 ± 4.9 µg (60 L/min)) [p < 0.023].


Salmeterol xinafoate is a widely prescribed long-acting beta-adrenergic agonist. Valved holding chambers (VHCs) improve drug delivery from pressurized metered-dose inhalers (pMDI), particularly with patients having poor coordination. The present study compared a large volume VHC (Volumatic™, GlaxoSmithKline - 750-ml, n=5 devices) with a small volume VHC (AeroChamber Plus*, Monaghan Medical Corp. - 149-ml, n=5 devices) with salmeterol xinafoate (Serevent®: total dose 21 µg ex actuator, GlaxoSmithKline). Measurements were also made with the pMDI without VHC. Total emitted dose (TD), fine particle dose (FPD - particles < 4.7 µm aerodynamic diameter) and fine particle fraction (FPF) were determined by Andersen 8-stage impactor with USP Induction Port at 28.3 ± 0.5 L/min. Assays for salmeterol xinafoate were undertaken by HPLC-fluorescence spectrometry at excitation and emission wavelengths of 226 nm and 296 nm respectively. As expected, both types of VHC greatly reduced the coarse component of the dose from the pMDI (10.5 ± 1.2 µg - pMDI alone; 1.0 ± 0.6 µg - AeroChamber Plus* VHC; 0.9 ± 0.7 µg - Volumatic™ VHC). Both FPD and TD from the AeroChamber Plus* (12.7 ± 1.3 µg and 13.6 ± 0.9 µg respectively) and from the Volumatic™ (12.3 ± 1.7 µg and 13.2 ± 2.1 µg respectively) VHCs were comparable (un-paired t-test, p > 0.70). FPD from the pMDI alone was 10.6 ± 1.0 µg, slightly lower but still comparable with the FPD from either type of VHC. The small volume VHC appears to be as effective as the larger chamber for the delivery of this formulation. These data are consistent with the recommendation to use a VHC with this formulation for patients with poor coordination (Demirkan et al. (Chest 2000; 117, 1314-1318)).


If we assume that the only function of a spacer is to facilitate the execution of a spray, its use is limited to small children who do not collaborate (below age 6-7 years). However, spacers seem to improve the effectiveness of drugs and reduce both directly
and indirectly the side effects. To assess if these characteristics have a role in clinical practice, the response to 100 micrograms of salbutamol administered directly by Autohaler was compared to that obtained with the same dose administered with three different spacers, AeroChamber* VHC, Babyhaler, Volumatic. A series of 88 asthmatic subjects with a FEF 25-75 less than 70% of the predicted value was considered. Overall patients provided 118 responses to the bronchodilator: 17 using the Aerotec (Autohaler), 38 the AeroChamber* VHC, 33 the Babyhaler, 30 the Volumatic. The response was evaluated considering the parameters obtained by spirometry just before, 5 and 20 minutes after the inhalation of salbutamol. Heart rate was also measured at the same time points. Heart rate, but not spirometric parameter were increased by the use of the Autohaler, proving that the drug had been inhaled. All the spacers determined a significant increase in the parameters considered. No significant difference was detected among spacers, although the smallest (AeroChamber* VHC, Babyhaler) showed a trend to a better response, in particular before age 7 years. The complete ineffectiveness of direct inhalation and the excellent response to inhalation with spacers show the indispensability of the latter, independent of age. Although no substantial difference among spacers was detected, the trend to obtain a better response with smaller spacers inclines us to use them in particular between 4 and 7 years of age. The negative correlation between the increase in spirometric parameters and the age of the patient would allow to have doses aimed to age or to body weight.
METERED DOSE INHALERS AND VALVED HOLDING CHAMBERS VERSUS NEBULIZERS

BACKGROUND: In acute asthma inhaled beta(2)-agonists are often administered by nebuliser to relieve bronchospasm, but some have argued that metered-dose inhalers with a holding chamber (spacer) can be equally effective. Nebulisers require a power source and need regular maintenance, and are more expensive in the community setting. OBJECTIVES: To assess the effects of holding chambers (spacers) compared to nebulisers for the delivery of beta(2)-agonists for acute asthma. SEARCH METHODS: We searched the Cochrane Airways Group Trial Register and reference lists of articles. We contacted the authors of studies to identify additional trials. Date of last search: February 2013. SELECTION CRITERIA: Randomised trials in adults and children (from two years of age) with asthma, where spacer beta(2)-agonist delivery was compared with wet nebulisation. DATA COLLECTION AND ANALYSIS: Two review authors independently applied study inclusion criteria (one review author for the first version of the review), extracted the data and assessed risks of bias. Missing data were obtained from the authors or estimated. Results are reported with 95% confidence intervals (CIs). MAIN RESULTS: This review includes a total of 1897 children and 729 adults in 39 trials. Thirty-three trials were conducted in the emergency room and equivalent community settings, and six trials were on inpatients with acute asthma (207 children and 28 adults). The method of delivery of beta(2)-agonist did not show a significant difference in hospital admission rates. In adults, the risk ratio (RR) of admission for spacer versus nebuliser was 0.94 (95% CI 0.61 to 1.43). The risk ratio for children was 0.71 (95% CI 0.47 to 1.08, moderate quality evidence). In children, length of stay in the emergency department was significantly shorter when the spacer was used. The mean duration in the emergency department for children given nebulised treatment was 103 minutes, and for children given treatment via spacers 33 minutes less (95% CI -33 to -24 minutes, moderate quality evidence). Length of stay in the emergency department for adults was similar for the two delivery methods. Peak flow and forced expiratory volume were also similar for the two delivery methods. Pulse rate was lower for spacer in children, mean difference -5% baseline (95% CI -8% to -2%, moderate quality evidence), as was the risk of developing tremor (RR 0.64; 95% CI 0.44 to 0.95, moderate quality evidence). AUTHORS’ CONCLUSIONS: Nebuliser delivery produced outcomes that were not significantly better than metered-dose inhalers delivered by spacer in adults or children, in trials where treatments were repeated and titrated to the response of the participant. Spacers may have some advantages compared to nebulisers for children with acute asthma.


OBJECTIVE: To compare the incremental cost and effects (averted admission) of using a metered-dose inhaler (MDI) against wet nebulization to deliver bronchodilators for the treatment of mild to moderately severe asthma in pediatric emergency departments (EDs). METHODS: We measured the incremental cost-effectiveness from the perspective of the hospital, by creating a model using outcome characteristics from a Cochrane systematic review comparing the efficacy of using MDIs versus nebulizers for the delivery of albuterol to children presenting to the ED with asthma. Cost data were obtained from hospitals and regional authorities. We determined the incremental cost-effectiveness ratio and performed probabilistic sensitivity analyses using Monte Carlo simulations. RESULTS: Using MDIs in the ED instead of wet nebulization may result in net savings of Can $154.95 per patient. Our model revealed that using MDIs in the ED is a dominant strategy, one that is more effective and less costly than wet nebulization. Probabilistic sensitivity analyses revealed that 98% of the 10,000 iterations resulted in a negative incremental cost-effectiveness ratio. Sensitivity analyses around the costs revealed that MDI would remain a dominant strategy (90% of 10,000 iterations) even if the net cost of delivering bronchodilators by MDI was Can $70 more expensive than that of nebulized bronchodilators. CONCLUSIONS: Use of MDIs with spacers in place of wet nebulizers to deliver albuterol to treat children with mild-to-moderate asthma exacerbations in the ED could yield significant cost savings for hospitals and, by extension, to both the health care system and families of children with asthma.


Background: Airway obstruction and bronchial hyperactivity oftentimes lead to emergency department visits in infants. Inhaled short-acting β2-agonist bronchodilators have traditionally been dispensed to young children via nebulizers in the emergency department. Delivery of bronchodilators via metered-dose inhalers (MDIs) in conjunction with holding chambers (spacers) has been shown to be effective. Study Objective: Safety and efficacy evaluations of albuterol sulfate hydrofluoroalkane (HFA) inhalation aerosol in children younger than 2 years with acute wheezing caused by obstructive airway disease. Methods: A
randomized, double-blind, parallel center, multicenter study of albuterol HFA 180 μg (n = 43) or 360 μg (n = 44) via an MDI with a valved holding chamber and face mask in an urgent-care setting. Assessments included adverse events, signs of adrenergic stimulation, electrocardiograms, and blood glucose and potassium levels. Efficacy parameters included additional albuterol use and Modified Tai Asthma Symptoms Score (MTASS reduction in MTASS representing improvement). Results: Overall, adverse events occurred in 4 (9%) and 3 (7%) subjects in the 180-μg and 360-μg groups, respectively. Drug-related tachycardia (360 μg) and ventricular extrasystoles (180 μg) were reported in 1 patient each. Three additional instances of single ventricular ectopy were identified from Holter monitoring. No hypokalemia or drug-related QT or QTc prolongation was seen; glucose values and adrenergic stimulation did not significantly differ between treatment groups. In the 180-μg and 360-μg groups, mean change from baseline in MTASS during the treatment period was $-2.8 (-49.8\%)$ and $-2.9 (-48.4\%)$, and rescue albuterol use occurred in 4 (9%) and 3 (7%) subjects, respectively. Conclusions: Cumulative dosing with albuterol HFA 180 μg or 360 μg via MDI-spacer and face mask in children younger than 2 years did not result in any significant safety issues and improved MTASS by at least 48%.


To determine whether parents who deliver albuterol treatments in a pediatric emergency department with a metered dose inhaler with a spacer device (MDIS) report better adherence to MDIS use at home compared to parents whose children undergo standard nebulizer therapy. Children aged 1-5 years were randomized by day to usual treatment with nebulized albuterol (40 children) or to treatment by the parent with albuterol with an MDIS (46 children). All caregivers received standard discharge instructions, a spacer and an MDI. Two weeks following the visit, a trained research assistant blinded to the child's group status, administered a brief telephone questionnaire to each caretaker. At follow-up, children in the MDIS group were 7.5 times more likely to be using the MDIS for their albuterol treatments (95\% CI 1.6-35.6). Involving parents in treatment of asthma exacerbations in the emergency department using an MDIS may improve adherence to MDIS use at home.

**NEBULISERS OR SPACERS FOR THE ADMINISTRATION OF BRONCHODILATORS TO THOSE WITH ASTHMA ATTENDING EMERGENCY DEPARTMENTS?** Mason N, Roberts N, Yard N, Partridge MR. Respiratory Medicine 2008;102:993-998.

Background: Systematic reviews and national guidelines conclude that the nebulised route of administration of bronchodilators has no advantage over the use of a spacer in moderately severe exacerbations of asthma. Whether this recommendation is implemented and whether it might affect use of staff time is unknown. Objectives: To determine the current method of administration of bronchodilators to those with non-life-threatening asthma attending emergency departments (ED) in London, UK and to monitor the implementation of a new policy to administer bronchodilators by spacers in one ED with a special reference to the time taken by nurses to administer the therapy by two different routes. Methods: Thirty-five EDs in Greater London were surveyed regarding their current practice. A time and motion study was then undertaken in one department observing nurses administering bronchodilators in the 3 weeks before and 3 weeks after a departmental policy change to favour the use of spacer devices rather than nebulisers. Results: The majority of EDs (94.3\%) in Greater London were using the nebulised route of administering bronchodilators to the majority of their adult patients. Spacers were more commonly used for the treatment of children (60.3\% of departments using spacers and nebulisers or spacers alone). Over half of the hospitals surveyed (51.4\%) were unaware that the British Guidelines on Asthma Management suggested that outcomes were the same and that there were potential advantages in the use of a spacer for both adults and children. Time and motion studies showed that the use of a spacer took no more nursing time than administration of the bronchodilator via a nebuliser, in fact treatment and set-up time were considerably lower for spacers. Conclusion: Spacer administration of bronchodilators to those with asthma attending EDs utilises less treatment time than use of a nebuliser. A survey of EDs in Greater London has shown that despite guideline conclusions there appears to be little evidence of reduction in use of nebulisers; a fear that use of alternatives might take nurses longer is not supported by this study.


**BACKGROUND:** Metered-dose inhalers with valved holding chambers (MDI-VHCs) have been shown to be equivalent to small volume nebulizers (SVNs) for the delivery of bronchodilators in children. At Seattle Children’s Hospital and Regional Medical Center we sought to implement the conversion from SVN to MDI-delivered albuterol in nonintubated patients receiving intermittent treatments. METHODS: There were 4 distinct interventions used to plan and implement this conversion program: (1) literature review, (2) product selection, (3) policy and operational changes, and (4) staff training. Bronchodilator administration guidelines and clinical pathways for asthma and bronchiolitis were revised to recommend MDI-VHC use in lieu of SVNs. Computerized physician order sets were amended to indicate MDI-VHC as the preferred method of delivering inhaled albuterol in children with asthma and bronchiolitis. Data from administrative case mix files and computerized medication delivery systems were used to assess the impact of our program. RESULTS: MDI-VHC utilization increased from 25% to 77% among all non-intensive-care patients receiving albuterol, and from 10% to 79% among patients with asthma (p < 0.001).
Duration of stay among patients with asthma was unchanged after conversion to MDI-VHC (p 0.53). CONCLUSIONS: Our program was very successful at promoting the use of MDI-VHC for the administration of albuterol in our pediatric hospital. Duration of stay among patients with asthma did not change during or since the implementation of this program.


OBJECTIVE: To compare the efficacy of beta-agonists given by metered-dose inhaler with a valved holding chamber (MDI+VHC) or nebuliser in children under 5 years of age with acute exacerbations of wheezing or asthma in the emergency department setting. STUDY DESIGN: Published (1966 to 2003) randomized, prospective, controlled trials were retrieved through several different databases. The primary outcome measure was hospital admission. RESULTS: Six trials (n=491) met criteria for inclusion. Patients who received beta-agonists by MDI+VHC showed a significant decrease in the admission rate compared with those by nebulizer (OR, 0.42; 95% CI, 0.24-0.72; P=0.002); this decrease was even more significant among children with moderate to severe exacerbations (OR, 0.27; 95% CI, 0.13-0.54; P=0.0003). Finally, measure of severity (eg, clinical score) significantly improved in the group who received beta-agonists by MDI+VHC in comparison to those who received nebulizer treatment (standardized mean difference, -0.44; 95% CI, -0.68 to -0.20; P=0.003). CONCLUSIONS: The use of an MDI+VHC was more effective in terms of decreasing hospitalization and improving clinical score than the use of a nebulizer in the delivery of beta agonists to children under 5 years of age with moderate to severe acute exacerbations of wheezing or asthma.


The aim of this pilot study was to compare the HaloLite Paediatric Nebulizer (HPN) with a pressurized metered dose inhaler and valved holding chamber (pMDI VHC, Aerochamber*) in terms of drug delivery, adherence to treatment, compliance with device, true adherence, and acceptability. Fourteen children aged 11-36 months with asthma on regular treatment with inhaled corticosteroids were enrolled into an open, randomized, crossover trial. They received budesonide for 2 weeks with each delivery system. Both devices incorporated a datalogger which recorded information on how the device was used. The HPN was preprogrammed to deliver 25 microg of budesonide to the patient. A single actuation of budesonide 200 microg was used with the pMDI VHC. The median delivered dose of budesonide was 36 microg (range, 31-45 microg; CV, 15%) for the HPN and 53 microg (range, 17-85 microg; CV, 47%) for the pMDI VHC. The median adherence was 68% (range, 11-96%) with the HPN and 71% (range, 11-100%) with the pMDI VHC. The median device compliance was 30% and 51% and the median true adherence was 23% and 36%, respectively. The shape, size, and weight of the HaloLite Paediatric Nebulizer were generally less acceptable than the shape, size, and weight of the pMDI VHC with datalogger. These results indicate that reproducible quantities of drug can be delivered to very young children using AAD technology such as that incorporated into the HPN. Drug delivery with the pMDI VHC was more variable, but parents preferred this device.


A substantive amendment to this systematic review was last made on 18 February 2003. Cochrane reviews are regularly checked and updated if necessary. Background: In acute asthma inhaled beta-2-agonists are often administered to relieve bronchospasm by wet nebulisation, but some have argued that metered-dose inhalers with a holding chamber (spacer) can be equally effective. In the community setting nebulisers are more expensive, require a power source and need regular maintenance. Objectives: To assess the effects of holding chambers compared to nebulisers for the delivery of beta-2-agonists for acute asthma. Search strategy: We last searched the Cochrane Airways Group trials register in February 2004 and the Cochrane Central Register of Controlled Trials (The Cochrane Library, Issue 1, 2004). Selection criteria: Randomised trials in adults and children (from two years of age) with asthma, where holding chamber beta-2-agonist delivery was compared with wet nebulisation. Data collection and analysis: Two reviewers independently applied study inclusion criteria (one reviewer for the first version of the review), extracted the data and assessed trial quality. Missing data were obtained from the authors or estimated. Results are reported with 95% confidence intervals (CI). Main results: This review has been updated in 2003 and has now analysed 1076 children and 444 adults included in 22 trials from emergency room and community settings. In addition, five trials on in-patients with acute asthma (184 children and 28 adults) have been added to the review. Method of delivery of beta-2-agonist did not appear to affect hospital admission rates. In adults, the relative risk of admission for holding chamber versus nebuliser was 0.88 (95% CI 0.56 to 1.38). The relative risk for children was 0.65 (95% CI 0.4 to 1.06). In children, length of stay in the emergency department was significantly shorter when the holding chamber was used, with a weighted mean difference of -0.47 hours, (95% CI -0.58 to -0.37 hours). Length of stay in the emergency department for adults was similar for the two delivery methods. Peak flow and forced expiratory volume were also similar for the two delivery methods. Pulse rate was lower for holding chamber in children, weighted mean difference -7.6% baseline (95% CI -9.9 to -5.3% baseline). An
OBJECTIVE: To determine if administration of albuterol by a metered-dose inhaler with a spacer device (AeroChamber*) is as efficacious as administration of albuterol by nebulizer to treat wheezing in children aged 2 years and younger. DESIGN: Double-blind, randomized, placebo-controlled clinical trial. SETTING: Pediatric emergency department. PATIENTS: From a convenience sample of wheezing children aged 2 to 24 months, 85 patients were enrolled in the nebulizer group and 83 in the spacer group. INTERVENTIONS: The nebulizer group received a placebo metered-dose inhaler with a spacer followed by nebulized albuterol. The spacer group received albuterol by a metered dose inhaler with a spacer followed by nebulized isotonic sodium chloride solution. Treatments were given every 20 minutes by a single investigator blinded to group assignment. MAIN OUTCOME MEASURES: The primary outcome was admission rate. Pulmonary Index score and oxygen saturation were measured initially and 10 minutes after each treatment. RESULTS: The nebulizer group had a significantly higher mean (SD) initial Pulmonary Index score compared with the spacer group (7.6 [2.5] vs 6.6 [2.0]; P =.002). With the initial Pulmonary Index score controlled, children in the spacer group were admitted less (5% vs 20%; P =.05). Analyses also revealed an interaction between group and initial Pulmonary Index score; lower admission rates in the spacer group were found primarily in children having a more severe asthma exacerbation. CONCLUSION: Our data suggest that metered-dose inhalers with spacers may be as efficacious as nebulizers for the emergency department treatment of wheezing in children aged 2 years or younger.


STUDY OBJECTIVES: To determine the efficacy of albuterol by metered-dose inhaler (MDI) and spacer (AeroChamber*) compared to a nebulizer. DESIGN: A prospective, open-label study. SETTING: Large urban emergency department (ED). PATIENTS: All consecutive adult asthma patients over a 2.5-year period. INTERVENTIONS: ED personnel used a standardized treatment algorithm, which included albuterol administered by nebulization, for patients presenting to the ED during the first 12 months of the study. The treatment algorithm then was switched to one that utilized albuterol administered by MDI/spacer as the primary mode of delivery for the following 18 months. As part of the conversion to MDI/spacer, ED staff counseled patients on self management and supplied patients with a peak flow meter, an MDI/spacer, and an inhaled steroid for home use. MEASUREMENTS: Pulmonary function, clinical outcome, laboratory data, and financial data were assembled and analyzed from 2,342 ED visits and 1,420 patients. RESULTS: While there was no significant difference in hospital admission rates between patients in the MDI/spacer group and the nebulizer group (13.2% and 14.6%, respectively), there was a statistically greater improvement in peak flow rates in the MDI/spacer group (126.8 vs 111.9 L/min, respectively; p = 0.002). The MDI/spacer group also spent significantly less time in the ED (163.6 and 175 min, respectively; p = 0.007), had a lower total albuterol dose (1,125 microg and 6,700 microg, respectively; p < 0.001), and showed a greater improvement in arterial oxygen saturation (p = 0.043). Relapse rates at 14 and 21 days were significantly lower (p < 0.01 and p < 0.05, respectively) among patients treated with the MDI/spacer and were associated with asthma education and the provision of a peak flow meter, a spacer, and an inhaled corticosteroid for patients' home use. CONCLUSIONS: Albuterol administered by MDI/spacer is an efficacious and cost effective alternative to nebulization in adults with acute asthma who present at a large urban ED.


OBJECTIVE: To compare the costs and effectiveness of albuterol by metered dose inhaler (MDI) and spacer versus nebulizer in young children with moderate and severe acute asthma. DESIGN: Randomized, double-blind, placebo controlled trial in an emergency department at a children's hospital. The participants were children 1 to 4 years of age with moderate to severe acute asthma. Patients assigned to the spacer group received albuterol (600 microg) by MDI by spacer (AeroChamber*) followed by placebo by nebulizer (n = 30). The nebulizer group received placebo MDI by spacer followed by 2.5 mg albuterol by nebulizer (n = 30). Treatments were repeated at 20-minute intervals until the patient was judged to need no further doses of bronchodilator, or a total of 6 treatments. RESULTS: Clinical score, heart rate, respiratory rate, auscultatory findings, and oxygen saturation were recorded at baseline, after each treatment, and 60 minutes after the last treatment. Baseline characteristics and asthma severity were similar for the treatment groups. The spacer was as effective as the nebulizer for clinical score, respiratory rate, and oxygen saturation but produced a greater reduction in wheezing (p =0.03). Heart rate increased to a greater degree in the nebulizer group (11.0/min vs 0.17/min for spacer, p <0.01). Fewer children in the spacer group required admission (33% vs 60% in the nebulizer group, p =0.04, adjusted for sex). No differences were seen in rates of
CHILDREN WITH MILD ACUTE ASTHMA. COMPARISON OF ALBUTEROL DELIVERED BY A METERED DOSE INHALER WITH SPACER VERSUS A NEBULIZER IN BRONCHODILATOR RESUSCITATION IN THE EMERGENCY DEPARTMENT PART 1 OF 2: DEVICE SELECTION. CONSENSUS STATEMENT: AEROSOLS AND DELIVERY DEVICES. Respir Care 2000;45(6):589-596.

Because the pMDI and DPI delivery system are the most convenient and produce the lowest cost/dose, they should be the first choice of clinicians. A valved holding chamber should be used with the pMDI whenever the patient cannot demonstrate acceptable hand breath coordination and whenever pharyngeal deposition is of clinical concern (e.g. inhaled steroids). In general, the valved holding chamber often with mask is almost always required in pediatric and geriatric populations. The nebulizer may be used if the drug is only available as a solution or if the pMDI/DPI cannot be used effectively.


This paper reviews the impact of device selection on bronchodilator resuscitation in the emergency department. The pMDI/holding chamber is equivalent to nebulizer therapy for treatment of infants, children, and adults with moderate to severe asthma. There may be some advantage in reduced treatment time and reduced adverse systemic effects of children with pMDI/HC. For treatment of patients with moderate airway obstruction (secondary to acute asthma and COPD), the selection of aerosol device appears to be of less importance in affecting clinical response than for patients with severe airway obstruction. In treating the most severe asthmatic (adult, child, or infant), the pMDI/HC has been demonstrated to be as effective as the nebulizer (or other available devices) in relief of airway obstruction, and appears to offer some advantage in fewer adverse effects. If the pMDI/HC works in the ED, with the sickest of patients, it should be equally effective in other settings as well. The evidence is abundant and clear: The debate on pMDI/HC versus nebulizer appears to no longer be a relevant issue.


OBJECTIVE: In children with mild acute asthma, to compare treatment with a single dose of albuterol delivered by a metered dose inhaler (MDI) with a spacer in either a weight-adjusted high dose or a standard low-dose regimen with delivery by a nebulizer. STUDY DESIGN: In this randomized double-blind trial set in an emergency department, 90 children between 5 and 17 years of age with a baseline forced expiratory volume in 1 second (FEV1) between 50% and 79% of predicted value were treated with a single dose of albuterol, either 6 to 10 puffs (n = 30) or 2 puffs (n = 30) with an MDI with spacer or 0.15 mg/kg with a nebulizer (n = 30). RESULTS: No significant differences were seen between treatment groups in the degree of improvement in percent predicted FEV1 (p =0.12), clinical score, respiratory rate, or O2 saturation. However, the nebulizer group had a significantly greater change in heart rate (p =0.0001). Our study had 93% power to detect a mean difference in percent predicted FEV1 of 8 between the treatment groups. CONCLUSION: In children with mild acute asthma, treatment with 2 puffs of albuterol by an MDI with spacer is just as clinically beneficial as treatment with higher doses delivered by an MDI or by a nebulizer.


Treatment of chronic lung disease of prematurity requires effective aerosol delivery of different therapeutic agents. Aerosols can be generated by a metered dose inhaler (MDI) or a jet nebulizer. An MDI combined with a spacer device is easier to use and avoids undesirable effects noted in conjunction with jet nebulization. We compared the clinical effectiveness of 200 micrograms (2 puffs) salbutamol delivered from an MDI in conjunction with a valved spacer device (AeroChamber®), and 600 micrograms given via jet nebulizer (PariBaby) on 2 consecutive days, the order being randomized. Thirteen spontaneously breathing very pre-term infants [mean (SD) gestational age 27.2 (1.8) weeks; birth weight 0.90 (0.34) kg] were studied at a corrected age of 37 (2.3) weeks. Mean (SD) study weight was 1.83 (0.38) kg. Dynamic lung compliance and resistance were determined from measurements of flows, volumes, and transpulmonary pressures, using a pneumotachometer and a small esophageal microtransducer catheter before and 20 min after salbutamol application. Baseline values before salbutamol administration were similar on both occasions: the mean (SD) compliance was 7.7 (3.0) mL.kPa-1.kg-1 pre-MDI plus-spacer and 8.4 (3.1) pre-jet nebulizer; the resistance was 10.4 (4.0) kPa.L-1.s pre-MDI plus-spacer and 9.7 (3.4) pre-jet nebulizer. Following salbutamol, compliance did not change significantly with either MDI plus spacer or jet nebulizer. Resistance fell significantly with MDI plus spacer (mean -2.2; 99.9% CI -0.35, -4.35) and jet nebulizer (-2.4; 99% CI -0.39, -4.42). We conclude
that even in small pre-term infants 200 micrograms salbutamol via MDI plus spacer improves dynamic resistance as effectively as 600 micrograms via jet nebulizer and may therefore be a preferable mode of aerosol administration.


Objective: To determine whether the administration of β-agonists by metered-dose inhaler (MDI) with a spacer device is as effective as the administration of β-agonists by nebulizer for the treatment of acute asthma exacerbations in children. Design: Randomized trial with two arms. Setting: Urban pediatric emergency department (ED) in Bronx, NY. Patients: Convenience sample of 152 children 2 years and older with a history of at least two episodes of wheezing presenting to the ED with an acute asthma exacerbation. Interventions: Patients were randomly assigned to receive standard doses of an β-agonists (albuterol) by an MDI with spacer (AeroChamber®) or by a nebulizer. Dosing intervals and the use of other medications were determined by the treating physician. Measurements/ Main Results: Baseline characteristics and asthma history were recorded. Asthma severity score, peak expiratory flow rate in children 5 years or older, and oxygen saturation were determined at presentation and before admission or discharge. The groups did not differ in age, sex, ethnicity, age of onset of asthma, or asthma severity score, and peak expiratory flow rate, oxygen saturation, number of treatments given, admission rate. Patients given MDIs with spacers required shorter treatment times in the ED (66 minutes vs. 103 minutes, p<0.001). Fewer patients in the spacer group had episodes of vomiting in the ED (9% vs. 20%, p<0.04), and patients in the nebulizer group had a significantly greater mean percent increase in heart rate from baseline to final disposition (15% vs. 5%, p<0.001). Conclusions: These data suggest that MDIs with spacers may be an effective alternative to nebulizers for the treatment of children with acute asthma exacerbations in the ED.