Evaluation of the influence of patient age and spacer device volume on aerosol lung deposition

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Abstract

Objective: To evaluate the efficacy of three frequently used spacer devices to deliver aerosol to the lung, and to compare radioaerosol deposition with each device in different age groups.

Methods: Nine healthy, non-smoking volunteers were recruited: three adults and six children, including three toddlers and three school age children. Qualitative and semi-quantitative analysis of radioaerosol deposition in the lung were carried out. Yet, two small-volume devices (Aerochamber® and Inal-Air®) and one large-volume device (Flumax®) were compared. Each patient inhaled 99mtechnetium-phytate. The device was filled during 30 seconds with radioaerosol. Oxygen was used as the driving gas. During 10 seconds, the patients inhaled the radioaerosol. The radiation emitted at the front and back of the chest was measured. The radiation inside the device was also measured.

Results: The quantitative evaluation of lung deposition revealed that the younger the patient, the less aerosol was deposited in the lung with the large-volume spacer device (Flumax®). The difference between small-volume devices (Aerochamber® and Inal-Air®) was not significant.

Conclusion: Small-volume spacers are the most appropriate for children. Large-volume devices should only be used by adolescents and adults.


Introduction

Nowadays inhalation therapy is one of the principal therapeutic weapons in the arsenal of respiratory diseases treatment. Although it was used even before the Christian era, its use has only become widespread over the last 30 to 40 years with the advent of ever more efficient devices in terms of pulmonary deposition. Added to this is a growing number of available medications, highly effective and with low incidence of collateral effects. Inhalation therapy by metered-dose aerosols is quick and easy to administer. In addition to this it offers better pulmonary deposition with a lower cost than conventional nebulizers. One obstacle that is encountered when metered-dose aerosols are employed is the need for good patient coordination, in order that the medication is adequately deposited within the lungs.

Spacing devices are valved devices capable of overcoming these difficulties. The correct choice and adequate utilization of spacing devices is a pre-requisite for the successful treatment of asthma in children. Spacing devices facilitate medication deposition within the lungs and eliminates the need for precise coordination, which is hard to achieve with certain patients.1-3 Over recent years, a variety of different spacing devices have become available on the market, with bivalve versions being most effective in the attempt to maximize the availability of the

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medication to the patient. It is known that their physical characteristics, such as size, format, volume and electrostatic charge directly affect this availability.4

One of the main dilemmas in inhalation therapy is choosing a spacer that unites the ideal characteristics that lead to the greatest efficacy in terms of the pulmonary deposition of the medication inhaled from among those most employed within the health services. Nowadays we have innumerable spacing devices available in Brazil, of varying shapes and sizes. However, we lack studies testing their efficacy. Rubim et al., employed scintigraphic techniques with the objective of evaluating a large volume spacer.5 Studies using radioisotopes for the comparative evaluation of spacing device efficacy in terms of pulmonary deposition have not yet been performed in our country. Starting from these assumptions, the study in question has the objective of comparing, by means of a clinical trial, three of the most often prescribed spacing devices at health services: Flumax®, a large volume, plastic spacer (700 ml); Aerochamber®, a small volume, acrylic space (190 ml); and the Inal-Air®, small volume, aluminum, and, therefore, electrostatically uncharged, spacer (230 ml).

**Methods**

Nine healthy, non-smoking patients with no history of respiratory diseases were studied. Three adults and six children were chosen, three pre-school age (3 and 4 years) and three school-age (6 and 7 years), who voluntarily offered to participate in the study, with the signed authorization given by the candidate themselves or their legal guardian. The study and consent form were approved by the Ethics Committee at our institution.

Aerosol particles containing 99mtechnetium-phytate were impelled into the spacing devices for 30 seconds by an oxygen flow at 10 l/min. Images were made of the spacing devices for 30 seconds by an uncharged, spacer (230 ml).

Employing the methods described above, the aerosol pulmonary deposition index was calculated for each spacing device. Thus, each individual was analyzed three times, each time with a different spacer, making a total sample of 27 events.

Significant variation was observed in terms of pulmonary deposition, with a large standard deviation, from patient to patient and from spacer to spacer (Table 1). In eight of the nine patients, irrespective of age group, there was less pulmonary deposition when the large volume spacer was used (Flumax®), compared with the small volume ones (Aerochamber® and Inal-Air®). The children exhibited significant differences in pulmonary deposition between small and large volume spacing devices, with significantly greater deposition occurring when the Aerochamber® and Inal-Air® small volume spacing devices were used. In these children, pulmonary deposition was statistically greater using the Inal-Air small volume, uncharged spacer when compared with the Flumax® large volume spacer with electrostatic charge, with a p value of 0.0313 (Figure 2).

Nevertheless, pulmonary deposition was similar for the adults, with no significant difference being found between the three spacing devices used for this age group. Comparing the Inal-Air and Aerochamber small volume spacing devices, we found that there was no statistically significant difference in pulmonary deposition (p = 0.2188).

* p = 0.0313.

**Figure 1** - Retention of aerosol particles containing 99mtechnetium-phytate into the spacing devices for 30 seconds after they were filled
Discussion

In world literature studies using radioisotopes to assess the efficacy of spacers remain scarce and with disparate results. Pedersen et al. showed deposition of 4 to 8% of aerosols inhaled via spacing devices and nebulizers. In adult patients, studied by Dolovich et al., deposition by metered-dose aerosols varied from 7 to 14%. It is important to note that in all these studies, including ours, there is underestimation of pulmonary aerosol deposition when radioactively marked material is employed, due to radiation absorption by thoracic structures.

Our study has a number of limitations. Among these is the small number of patients studied. Despite this the statistical power was calculated for around 80%. Pulmonary deposition may be different with chronic respiratory disease patients. Asthmatic children, during an acute crisis, or even during the inter-crisis period, may exhibit a pulmonary deposition pattern that is different from normal patients. We would also point out that the observed differences in pulmonary deposition do not necessarily indicate greater clinical efficacy. In order to determine this, studies with greater patient numbers which assess the clinical effectiveness of each spacer.

Spacer volume may impact on the availability of medication for inhalation, which itself may vary depending on the medication employed. Clinicians should be aware that deposition data for a given spacer, originating from studies of a particular drug may not be applicable to others. It is estimated that the tidal volume of children is around 8 to 10 ml/kg. A four-year-old child weighing 20 kg would therefore have tidal volume of between 160 and 200 ml. It is not difficult to imagine that such a child, using a large volume spacer (500 to 800 ml), would not manage to inhale its entire contents. The child’s respiratory capacity is insufficient and does not permit the inhalation of the whole contents. Furthermore, after a few seconds, progressive decantation of the medication occurs within the spacer reservoir and this portion is no longer available to the patient. We had the opportunity to document this finding when analyzing the images of the spacing devices filled with radioisotopes (Figure 1). Greater deposition is clearly observable within the larger volume spacer, due to the decantation of this aerosol. Another important factor to take into account is the electrostatic charge present on the inner surface of each spacer. Synthetic spacing devices present a negatively charged surface. Aerosols have a positive charge, which fact facilitates their adherence to the wall of the reservoir.

Table 1 - Mean, median, standard deviation and p value of \(^{99m}\)Tc-phytate deposition of Flumax\textsuperscript{®}, Aerochamber\textsuperscript{®} and Inal-Air\textsuperscript{®} spacing devices, according to age group

<table>
<thead>
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<th>Patients</th>
<th>Mean</th>
<th>Median</th>
<th>Standard deviation</th>
<th>p</th>
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<td>11.97</td>
<td>15.33</td>
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<td>Aerochamber\textsuperscript{®}</td>
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<td>Children (n = 6)</td>
<td>28.56</td>
<td>25.26</td>
<td>15.65</td>
<td>0.0238</td>
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<tr>
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<td>5.99</td>
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<td>Inal-Air\textsuperscript{®}</td>
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<td>40.91</td>
<td>42.41</td>
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<td>11.81</td>
<td>5.28</td>
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Significant p value < 0.05.

Figure 2 - Comparison of pulmonary deposition of \(^{99m}\)Tc-phytate radioaerosol in children and adults using Flumax\textsuperscript{®}, Aerochamber\textsuperscript{®} and Inal-Air\textsuperscript{®} spacing devices
comparable with international standards, with the Inal-Air® spacing devices present deposition indices that are deposition adequacy. We found that domestically sourced devices should be restricted to adolescents and adults internationally recognized devices. Large volume spacing devices which offer pulmonary deposition similar to documented the efficacy of low cost, Brazilian spacing confirm data from earlier studies. Additionally, we electrostatic charge. The results obtained in this study with pre-school aged children due to the absence of we found better deposition with the former, particularly Acknowledgements

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