Superiority of spacer/mask topical anaesthetic compared with conventional spray and gargle method for fibreoptic bronchoscopy

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OBJECTIVE: To compare the safety and efficacy of a new spacer-oral nasal mask device with those of the standard needle nozzle spray method for the delivery of aerosolized lidocaine to the upper airway for pre-bronchoscopic anaesthesia in a tertiary care hospital.

DESIGN: Single-blind randomized control trial.

SETTING: University affiliated tertiary care hospital, ambulatory care bronchoscopy unit.

SUBJECTS: Thirty consecutive consenting patients referred for fibreoptic bronchoscopy for various indications.

INTERVENTION: Thirty randomized subjects received 150 mg of topical 1% aerosolized lidocaine via standard long needle nozzle applicator (group A) or via a new oral/nasal mask with spacer device (group B). Bronchoscopists, blinded as to the preprocedure topical anaesthetic method used, gave additional topical lidocaine at their discretion.

MEASUREMENTS: The study nurse recorded the total dose of lidocaine (mg), timing of the procedure (s), cough frequency expressed as coughs per minute (c/min), vital signs, time for return of gag reflex and patients’ subjective comments.

RESULTS: Fifteen patients were randomized to each group. The lidocaine dose required for insertion through the vocal cords (mean ± SD) was 282.6±66.3 mg in group A and 203.3±70.6 mg in group B (P<0.005). Total lidocaine dose required for the procedure was 330.6±70.2 mg in group A and 256.6±75 mg in group B (P<0.01). The mean time for passage of the bronchoscope from mouth entry to through the vocal cords was 82.7±54.5 s in group A and 110.5±64.4 s in group B (P>0.1). The mean total time for the procedure was 699.7±377.5 s in group A and 697.2±409.1 s in group B (not significant). The mean cough frequency was 8.2±6.1 c/min in group A and 7.0±5.7 c/min in group B (not significant). There were no statistically significant differences in heart rate, in return of gag reflex time or in complication rate between the two groups.

CONCLUSIONS: A statistically significant reduction in the dose of lidocaine is required to achieve equivalent topical anaesthetic for bronchoscopy with a new mask and spacer device compared with a more conventional method. Since no other variables related to the procedure showed a significant difference, the new method appears to be superior to the previous method. (Pour résumé voir page 177)

Key Words: Anaesthesia, Aerosols, Bronchoscopy

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Supériorité de la technique d’anesthésie locale par masque avec dispositif d’espacement comparée à la technique d’anesthésie classique par pulvérisation et gargaramis pour pratiquer une fibroscopie bronchique

OBJECTIF : Comparer un nouveau système de masque oro-nasal muni d’un dispositif d’espacement avec la méthode classique de vaporisation par cône applicateur, sur le plan de la sûreté et de l’efficacité, pour administrer de la lidocaine en aérosol dans les voies respiratoires supérieures en vue d’une anesthésie prébronchoscopique dans un hôpital de soins tertiaires.

MODÈLE : Essai contrôlé randomisé à simple insu.

CONTEXTE : Hôpital de soins tertiaires affilié à une université, unité des soins ambulatoires pour la bronchoscopie.

SUJETS : Une série de trente patients consentants adressés pour une fibroscopie bronchique pour des indications variées.

INTERVENTION : Trente sujets randomisés ont reçu 150 mg de lidocaine à 1 % en aérosol administrée localement à l’aide d’un appareil nasal classique (groupe A) ou par un nouveau système de masque oro-nasal muni d’un dispositif d’espacement (groupe B). La prémédication a été faite suivant l’une des deux méthodes à l’insu des bronchoscopistes qui pouvaient, à discrétion, administrer localement des doses supplémentaires de lidocaine.

MESURES : L’infirmière de l’essai a consigné la dose totale (mg) de lidocaine administrée, le temps (s) de l’intervention, la fréquence de la toux exprimée en nombre de fois par minute (t/minute), les signes vitaux, l’heure de réapparition du réflexe pharyngé et les commentaires subjectifs du patient.

RÉSULTATS : Quinze patients ont été randomisés en deux groupes. La dose de lidocaine requise pour l’insertion du bronroscope à travers les cordes vocales (moyenne ± écart-type) était de 282,6±66,3 mg dans le groupe A et de 203,3±70,6 mg dans le groupe B (P<0,005). La dose totale de lidocaine requise pour l’intervention était de 330,6±70,2 mg dans le groupe A et de 256,6±75 mg dans le groupe B (P<0,01). Le temps moyen de passage du bronroscope de la bouche jusqu’à la traversée des cordes vocales était de 82,7±54,5 s dans le groupe A et de 110,5±64,4 s dans le groupe B (P=0,1). Le temps total moyen pour l’intervention était de 699,7±377,5 s dans le groupe A et de 697,2±40,1 s dans le groupe B (non significatif). La fréquence moyenne de la toux était de 8,2±6,1 t/minute dans le groupe A et de 7,0±5,7 t/minute dans le groupe B (non significatif). Il n’y avait aucune différence significative dans le rythme cardiaque, la réapparition du réflexe laryngé ou le taux de complications entre les deux groupes.

CONCLUSIONS : Le nouveau système de masque muni d’un dispositif d’espacement permet de réduire considérablement la dose de lidocaine requise pour procéder à une anesthésie prébronchoscopique locale équivalente à celle pratiquée par une méthode plus classique. Puisque l’on a observé aucune différence dans les autres variables associées à l’intervention, la nouvelle méthode d’anesthésie semble donc supérieure à celle utilisée antérieurement.

Fibreoptic bronchoscopy is indicated in numerous clinical scenarios and provides a vital diagnostic tool in current respiratory medicine. Unfortunately, considerable patient discomfort is associated with this procedure. Many patients object to the unpleasant taste of lidocaine, and to coughing and gagging during the gargling and spraying of the oropharynx. This standard spray method often delays the starting of the procedure due to marked patient discomfort and anxiety. This may lead to a prolonged, poorly tolerated procedure with suboptimal sample collection. Furthermore, patients who have suffered through one procedure are reluctant to repeat the procedure when necessary.

Numerous authors have evaluated various techniques for anaesthetizing the upper airway with aerosolized or nebulized lidocaine (1). Administration of lidocaine by nebulizer has been advocated, but this method requires large doses of medication and has not been very efficient or effective (2-4). Clark and Pond (5) have recently described atomization of the drug.

Aerosol therapy with beta-agonists using a metered dose inhaler (MDI) (especially with holding chamber) is 2.5 to six times as efficient as continuous nebulization for bronchodilation in asthmatics (6,7). Other investigators have developed new methods of efficiently ‘targeting’ the respiratory tract (eg, spacer devices) (8).

This study compared the dose of lidocaine, the time required and the subject tolerance for a new spacer/mask device versus a conventional spray applicator device in the delivery of aerosolized lidocaine to the oropharynx before bronchoscopy.

PATIENTS AND METHODS

Subjects: Thirty consecutive subjects were entered into the study. Excluded subjects had contraindications to one or more protocol medications (ie, atropine, midazolam and lidocaine). All subjects were informed of all procedures and signed appropriate consent forms. The Hospital Ethics Committee for Human Research approved the protocol.

Premedication: Subjects received atropine (0.4 to 0.6 mg/s) 30 mins before the procedure. After subjects consented to participate in the study, all patients received midazolam (5 mg intramuscularly or 2 mg intravenously) just before starting the procedure.

Randomization and blinding: A nurse research assistant using a random numbers table assigned subjects to the standard spray method (group A) or to the spacer/mask method (group B). Subjects were then given the topical anaesthetic by the nurse before the physician entered the endoscopy suite. The bronchoscopist was therefore blinded to the anaesthetic technique used.

Procedure: Subjects received 150 mg (15 actuations) of 1% lidocaine (Xylocaine; Astra) (10 mg/mL) directed onto the pharynx via the standard spray device or the new spacer/mask device. The standard method used a standard aerosol MDI with a long nosed metal applicator (Figure 1) to spray the posterior pharynx and the vo-
oral cords. If the patient started to cough during the application the nurse would wait until coughing ceased before giving the next actuation. Subjects who received lidocaine using the new method placed the spacer/mask device over their mouth and nose (Figure 2). The oral nasal mask allowed the patients to use the open mouth technique for breathing in the medication. With each actuation of the device, subjects were directed to take rapid deep breaths through their widely open mouth to total lung capacity to achieve relatively high inspiratory flow rates (approximately 60 L/min). They were monitored during the procedure to ensure that they did not inhale via the nose. With the new technique, the lidocaine MDI was actuated at 10 s intervals, thus delivering the 150 mg of lidocaine over approximately 3 mins.

The bronchoscope was introduced through a hollow bite block between the teeth and passed through the vocal cords. The bronchoscopist used additional 4% lidocaine (40 mg/mL), instilled directly onto the vocal cords through the bronchoscope, as needed, to control cough and laryngospasm. Past the vocal cords the procedure was carried out in the usual manner with instillation of 1% topical lidocaine solution applied through the bronchoscope, 1 mL at a time, as required to minimize cough. The study nurse recorded the total dose of lidocaine required to allow the bronchoscope to be passed through the vocal cords, as well as the total dose for the whole procedure.

**Outcomes:** The primary outcome measure was the total lidocaine dose required (mg) to complete the procedure. Secondary outcome measures included the lidocaine dose required to pass the bronchoscope through the vocal cords (mg), the time from introduction of the bronchoscope into the mouth to the bronchoscope passage through the vocal cords (s), total procedure time (s) and cough frequency (c/min). The number of coughs and various times were recorded by the nurse research assistant using a manual counter and stopwatch.

Other outcomes evaluated during the procedure were maximum change in heart rate and complications (bleeding, pneumothorax, etc). Patients were also asked for any general comments regarding any discomfort from the procedure.

**Data analysis:** The arithmetic means of the total dose

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**TABLE 1**

<table>
<thead>
<tr>
<th>Patient profile</th>
<th>Spray method</th>
<th>Mask/spacer method</th>
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</thead>
<tbody>
<tr>
<td>Male/female</td>
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<td>8/7</td>
</tr>
<tr>
<td>Age (mean years)</td>
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<td>60</td>
</tr>
<tr>
<td>Nonsmokers/smoker</td>
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<td>4/11</td>
</tr>
</tbody>
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**TABLE 2**

<table>
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<th>Indications for bronchoscopy</th>
<th>Spray method</th>
<th>Mask/spacer method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cough</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Hemoptysis</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Abnormal radiograph</td>
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<td>7</td>
</tr>
<tr>
<td>Other</td>
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<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>15</td>
<td>15</td>
</tr>
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</table>

**TABLE 3**

<table>
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<tr>
<th>Procedures performed during bronchoscopy</th>
<th>Spray method</th>
<th>Spacer/mask method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Washings for cytology/microbiology</td>
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<td>5</td>
</tr>
<tr>
<td>Washings and brushings</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>Washings, brushings and biopsy</td>
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<td>4</td>
</tr>
<tr>
<td>Total</td>
<td>15</td>
<td>15</td>
</tr>
</tbody>
</table>

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**Figure 1** Standard method uses an aerosol metered dose inhaler with a long nosed metal applicator

**Figure 2** New method places the spacer/mask device over the mouth and nose
of lidocaine topical anaesthetic required, the mean time to pass through the vocal cords, mean cough frequency and mean total number of coughs were compared between the two groups using unpaired t test analysis.

RESULTS

Subject characteristics: There were 15 subjects randomized to each group. There were no significant differences with regard to sex, age or smoking history (Table 1). No significant differences were detected between groups with respect to indications for bronchoscopy (Table 2) or to the type of procedures performed (Table 3).

Lidocaine dose: The lidocaine dose required for insertion of the bronchoscope through the vocal cords (mean ± SD) was 282.7±66.4 mg in group A and 203.3±70.7 mg in group B (P<0.005). Total lidocaine dose required for the procedure was 330.7±70.2 mg in group A and 256.7±75.7 mg in group B (P<0.01).

Length of time for procedure: The mean duration from the time the bronchoscope passed the lips until insertion through the vocal cords was 82.7±54.5 s in group A and 110.5±64.4 s in group B (P>0.1). The total time for the procedure was 699.7±377.5 s in group A and 697.3±409.1 s in group B (P>0.1) (Table 4).

Other parameters: Cough frequency was 8.2±6.2 c/min in group A and 7.0±5.7 c/min in group B (P>0.1). There were no statistically significant differences in changes in heart rate, in return of gag reflex or in complication rate between the two groups. Subjective comments about taste and tolerability of anaesthetic and bronchoscopy procedures were similar in the two groups.

DISCUSSION

Topical anaesthesia remains a fairly cumbersome and time consuming technique which, using our standard spray procedure, involves either applying lidocaine gel to the nasal airway for per nasal introduction and/or gargling 2 mL of unpleasant tasting 4% lidocaine solution in approximately 8 mL of water for per oral insertion of the bronchoscope. This is followed by blindly spraying a 1% solution onto the pharynx, epiglottis and vocal cords and then giving additional 4% lidocaine solution 1 mL at a time directly onto the cords under direct vision through the bronchoscope.

Topical anaesthesia above and onto the cords usually requires the administration of up to 300 mg of lidocaine, provided in 40 mg aliquots (40 mg/mL). Ongoing topical anaesthesia of the lower respiratory tract generally requires an additional 100 to 300 mg of 1% lidocaine, the larger doses for prolonged procedures such as fluoroscopically guided transbronchial biopsy (total maximum lidocaine dose 400 mg over 20 to 30 mins). Gargling with and spraying unpleasant tasting lidocaine onto the oropharynx and upper airway causes many patients to gag and cough, which delays the start of the procedure and creates significant patient anxiety. This anxiety may exacerbate coughing and gagging, leading to a prolonged poorly tolerated procedure with suboptimal sample collection. Patients who have suffered through one procedure are reluctant to repeat the procedure when needed.

The current project tested a unique spacer/mask delivery system to study patient acceptance, reduction of bronchoscope insertion time, total bronchoscopy time and anaesthetic dose requirements compared with a technique currently used by many bronchoscopists.

The new method attempted to anaesthetize the airway from the tongue to the periphery by taking advantage of dispersion of the droplet aerosol produced by these canisters via the spacer/mask device. The larger droplets in the aerosol spray (which contains particles of many sizes from approximately 30 mm down to 1 mm) are more likely to be deposited in the upper respiratory tract and central airways because of high inspiratory flow rates that favour impaction, while the smaller particles (less than 5 mm) are likely to be carried to more peripheral airways (9).

Thus the spacer/mask method should provide topical anaesthesia to the whole respiratory tract, from the mouth to the peripheral airways, depositing the majority of the dose in more proximal airways at the level of the gag and cough receptors. These receptors are located mainly in the pharynx and upper respiratory tract around the vocal cords and in the central airways of the lower respiratory tract (10). It was postulated that this spacer dispersed aerosol would be more evenly distributed to these receptors than would be the case with the nozzle spray method, thus providing effective topical anaesthesia with a smaller total dose. Minimizing total topical lidocaine dose is desirable because there are reported systemic side effects (11-13).

This study confirms that the spacer/mask device provides equivalent symptomatic benefit to the spray method with a 25% decrease in the total dose of lidocaine. No significant differences were observed between the two methods in terms of time to pass the endoscope through the cords, total time of the proce-
dure, cough frequency or change in heart rate. The dose reduction translates into cost savings. The safety and ease of premedication with lidocaine using the new spacer/mask method allows endoscopy nurses to readily perform this procedure, thus saving physician time.

We did not include data regarding the time taken to administer the 15 actuations of 1% lidocaine either via the standard or new method. Typically it took longer to use the standard method because the nurse would often have to wait for the patient to stop coughing between actuations. The spacer mask method caused very little cough response during the actuations. Because it was not possible to do this part of the study in a blinded fashion, we felt that there was significant potential for bias by the study nurse in terms of deciding when to resume giving the lidocaine using the standard method.

The nurse who gave the premedication anaesthetic also counted coughs using a manual counter. The fact that the nurse was not blinded to the type of premedication administered could theoretically lead to biased counting of coughs. No difference was noted between the two groups with respect to cough frequency, which could be interpreted to mean that there was no true difference between the two groups or that the nurses under-counted or over-counted coughs in one or the other group. Cough frequency was one of many measures recorded to compare patient tolerance of the procedure between the two treatment groups. Other measures such as change in heart rate, time to complete the procedure and subjective comments of patients indicate that patient tolerance was comparable between the two groups.

We did not measure serum lidocaine levels in this study. Previous studies (4,13) have demonstrated that systemic serum levels of lidocaine are well below the recommended therapeutic range of 2 to 6 mg/mL when the lidocaine is delivered by ultrasonic nebulizer plus direct instillation of lidocaine onto the mucosa of the upper and lower airways. Total doses of lidocaine delivered to the airways were in the range of 450 to 540 mg in one study (4) and over 1600 mg in another study (13). These values are significantly higher than the maximum lidocaine doses used in our study (approximately 400 mg). It is unlikely that the dose of drug delivered to the tracheobronchial tree is higher with the spacer device because a significant portion of the dose is captured in the spacer device itself. We propose that the spacer device technique results in delivery of a more uniform distribution of the medication to the airways. Furthermore, neither group demonstrated evidence of lidocaine toxicity, particularly central nervous system excitation or depression, or cardiovascular compromise such as bradycardia or hypotension.

Although the patients’ subjective comments regarding the actual anaesthetic procedure are the ‘softest data’, these comments are potentially the most useful clinical information as to whether the spacer/mask method offers any perceived symptomatic benefit to the patient. Not surprisingly, in this study there is little to distinguish the nature of these comments between the two groups because patients were anaesthetized with one method or the other and thus could not compare them.

**CONCLUSIONS**

This study demonstrated that initial topical anaesthesia delivered as a widely dispersed aerosol from a spacer/mask device resulted in the need for a reduced total dose of lidocaine to achieve equivalent cough control compared with a highly focused droplet spray directed against the pharynx. This suggests that the more dispersed aerosol spray is able to anaesthetize the posterior pharynx and fauces better than the older method. Since the primary outcome variable was the additional local anaesthetic necessary to achieve optimum cough control it is not surprising that no differences could be demonstrated in any of the other outcome measures.

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