Clinical practice guidelines for suctioning the airway of the intubated and nonintubated patient

Dina Brooks BScPT MSc PhD1*, Cathy M Anderson BSc BHScPT MSc2,3*, Margaret A Carter BSc RRCP RRT4*, Laurie A Downes BScPT MSc5*, Sean P Keenan MD MSc FRCPC2*, Carol J Kelsey BScPT MSc1*, Janet B Lacy BScPT MSc1*

1Department of Physical Therapy, University of Toronto, Toronto; 2London Health Sciences Centre, London; 3Department of Physical Therapy, University of Western Ontario, London; 4College of Respiratory Therapists of Ontario, Toronto; 5College of Physiotherapists of Ontario, Toronto, Ontario

**GUIDELINES**

OBJECTIVE: To provide physicians, physiotherapists, nurses and respiratory therapists with guidelines for the application of airway suctioning.

DESIGN: This clinical practice guideline was developed using the model by Browman and colleagues. A working group of representatives from four professional colleges (nurses, physicians and surgeons, physiotherapists and respiratory therapists) and research experts was formed to conduct a systematic review, develop evidence-based recommendations and generate clinical practice guidelines. MEDLINE (1966 to 1998), CINAHL (1982 to 1997) and EMBASE (1974 to 1996) as well as the reference lists of identified articles were searched. Inclusion of articles was determined by at least two group members, and studies were classified according to type. Randomized, controlled; randomized; and non-randomized crossover and comparative cohort trials were grouped by type of intervention and population for use in the development of recommendations. Other observational and animal studies dealing with adverse effects of suctioning were included in the review but were not used in the development of recommendations. Input on the evidence-based recommendations was sought and incorporated from members of all four professions and from experts on content and methodology.

SETTING: Any setting (hospital or home) where suctioning is performed.

POPULATION: Intubated and nonintubated adults, infants and children.

RESULTS AND CONCLUSIONS: An attempt was made to develop recommendations in each of the subcategories of suctioning techniques addressed by at least one study. In some subcategories, definite recommendations were made (13 in adults, and three in children and infants); in other subcategories, insufficient evidence precluded recommendations. The recommendations addressed the following aspects of suctioning: preoxygenation, hyperinflation, insufflation, hyperoxygenation, hyperventilation, saline instillation, adaptor use, medication use, open and closed systems, and various types of catheters.

Key Words: Adult; Clinical practice guidelines; Intubation; Pediatric; Suction

*Members of the Interdisciplinary Working Group
Correspondence and reprints: Dr D Brooks, Department of Physical Therapy, University of Toronto, 256 McCaul Street, Toronto, Ontario M5T 1X5. Telephone 416-978-1739, fax 416-978-4363, e-mail dina.brooks@utoronto.ca

Pour le résumé, voir page suivante
Directives de pratiques cliniques pour l’aspiration des voies respiratoires du patient intubé ou non intubé

OBJECTIF : Offrir aux médecins, physiothérapeutes, infirmières et inhalothérapeutes des directives techniques pour l’aspiration des voies respiratoires.


CONTEXTE : Tout contexte (hôpital ou domicile) où l’on effectue l’aspiration des voies respiratoires.

POPULATION : Adultes, nourrissons et enfants, intubés et non intubés.

RÉSULTATS ET CONCLUSIONS : On a tenté de développer des recommandations pour chacune des sous-catégories de techniques d’aspiration mentionnée dans au moins une étude. Dans certaines sous-catégories, des recommandations précises ont été faites (13 pour les adultes et 3 pour les enfants et les nourrissons). Dans d’autres sous-catégories, le manque de preuves a empêché la préparation de recommandations. Ces recommandations visaient les aspects suivants de l’aspiration : pré-oxygénation, hyperinflation, insufflation, hyperoxygénation, hyperventilation, instillation de sérum physiologique, utilisation d’un adaptateur, utilisation de médicaments, systèmes ouverts et fermés et divers types de cathétères.

U nder the Regulated Health Professions Act, enacted in Ontario in 1994, high risk procedures are designated as ‘controlled acts.’ Only certain identified professionals in the legislation are authorized to perform these controlled acts on clients. In Ontario, nurses, physicians, physiotherapists and respiratory therapists are authorized to perform the controlled act of suctioning. Each professional college regulates its members concerning the performance of the controlled act.

To perform suctioning, a catheter or device is inserted via the nasopharynx or oropharynx, or through a nasopharyngeal or an oropharyngeal airway, or an endotracheal or tracheostomy tube. Although suctioning is a common technique employed with critically ill patients, rehabilitation patients and patients at home, controversy exists over the beneficial and harmful effects of the procedure.

A survey of practitioners in Ontario revealed a large variation in practice in the application of oropharyngeal and tracheal suctioning and procedures such as hyperoxygenation, hyperinflation and saline instillation (1). These variations existed despite reports of locally developed guidelines and policies regarding suctioning practices (1), several published review articles (2-13) and one clinical practice guideline (CPG) on this topic (14). Neither the review articles nor the published CPG used a rigorous critical appraisal procedure or a clear explicit process for consensus or investigation and assessment of the evidence.

The goal of the interdisciplinary working group was to answer the following questions, using a model published by the Ontario Cancer Treatment Practice Guidelines Initiative CPG group (15).

What are the clinical indications and precautions for suctioning, and what is the evidence to support the procedures of tracheal and pharyngeal suctioning by health professionals for patients of any age to prevent upper airway obstruction and to clear secretions?

What are the methods, techniques and monitoring tools that optimize benefits and reduce complications when suctioning is performed on patients of all ages who are intubated, nonintubated or tracheotomized?

MATERIALS AND METHODS

Electronic database searches included MEDLINE (1966 to 1998), EMBASE (1974 to 1996) and CINAHL (1982 to 1997). Key words used for the searches were ‘trachea’, ‘pharynx’, ‘tracheostomy’, ‘suction’, ‘artificial airway’, ‘inflation’ and ‘endotracheal’. All searches were limited to English. Printouts from these searches, including title, authors, journal and abstract (where available) were reviewed by at least two group members. Initially, broad, subject-related criteria were used to determine which articles would be retrieved for further review. Retrieved articles consisted of reviews, guidelines, studies of any design and any article for which the reference list was thought to be useful. The Cochrane Database was searched for relevant reviews; none were found.

At least two members of the working group reviewed each article retrieved and agreed on whether it should be included or excluded from further analysis. Excluded articles consisted of those that used interventions not relevant to the questions, that reported on a diagnostic test or that were reviews, guidelines, protocols or other descriptive articles. Included articles were randomized, controlled trials; randomized and nonrandomized crossover trials; comparative cohort studies; case series; and animal studies. The epidemiologist’s decision to exclude any article was ratified by at
least one clinician. The reference lists of both included and excluded studies were reviewed for identification of other relevant articles. Only abstracts with sufficient information to allow judgment of methodology and interpretation of data were included. No attempts were made to locate unpublished data or to contact authors for information that was absent from the published article.

A data collection form was designed by the group. Data extraction was first completed by the clinical epidemiologist, and included information on the study type, allocation methods, subject inclusion/exclusion criteria, when the study was conducted, adequacy of follow-up, comparison of baseline characteristics, specific interventions and outcome measures. At least two group members then reviewed each article and the previously extracted information on the collection form. Additional data extraction involved determining whether the outcome measures of physiological status were appropriate, valid and reliable; setting priorities when there were multiple outcomes reported; and abstracting results. Detailed summaries of each article were composed. The working group used these summaries to formulate recommendations. Modified versions of the data tables have been included in the ‘Discussion’ section of this paper.

To develop evidence-based recommendations, we focused primarily on randomized, controlled trials (RCTs) and comparative cohort studies. Many of the studies used a crossover design. If the authors did not specify randomization of the order of interventions, the study was assumed to be nonrandomized. Because many studies were not randomized and most could not be adequately double-blinded, no formal quality assessment tool was used. No attempt was made to use a formalized system to rate the levels of evidence; however, the number and types of studies were summarized in each category. Due to the heterogeneity in patient populations, differences in the intervention techniques and outcomes measured, and other methodological issues, no meta-analyses were performed.

Case series and animal studies were not considered when making evidence-based recommendations; however, they were reviewed to gain insight into the potentially harmful effects of suctioning. For ethical reasons, it is not possible to use RCTs and comparative cohort studies to examine the possible hazardous effects of suctioning. Once the evidence-based recommendations were formulated, we ensured that none of the recommended approaches had been shown to cause harm in either case series or animal studies.

We generated a full, detailed ‘technical report’ and a short summary of the recommendations (less than five pages). We then sought input on the short summary from 165 clinicians (84 physical therapists, 33 respiratory therapists and 48 nurses). In addition, the full technical report was sent to four clinical experts representing each of the professions and three methodological experts (with recognized expertise in guideline development, systematic overview and/or meta-analysis). Standardized forms were used to seek feedback. The working group considered all feedback, and modifications were made for clarification of the document.

### RESULTS

From the combined searches, 317 articles were retrieved. Of these, 162 were included and 155 were excluded. The number of articles excluded or included and the reasons for this are presented in Table 1.

Only one study dealt specifically with the first question, i.e., the indication for suctioning. The remaining 161 studies were related to the second question (i.e., “What are the methods, techniques and monitoring tools that optimize benefits and reduce complications when suctioning patients of all ages who are intubated, nonintubated or tracheotomized?”), and were either randomized, controlled, crossover or comparative cohort in design. Studies were separated into two sections: infants and children (n=23), and adults (n=139). There were 13 recommendations for adults and three for infants and children. For the infants and children section, articles were categorized according to subpopulation. For the adult population, the studies were categorized on the basis of the technique used. While many studies lacked important outcomes, recommendations were only made for the outcomes reported.

Input on the evidence-based recommendations was received from 103 clinicians (57 physical therapists, 19 respiratory therapists and 27 nurses), and from all seven clinical and methodological experts. The working group discussed all input in detail and incorporated the changes where it was deemed appropriate by the entire working group. Although the feedback did not result in fundamental changes to the recommendations, the document was modified to improve its clarity and readability.

### DISCUSSION

Summaries and recommendations in infants and children: The studies in pediatrics were categorized according to subpopulation: nonintubated neonates with and without meconium aspiration syndrome (MAS), intubated preterm and full-term neonates, and intubated infants and children. Table 2 includes information about the studies that was used to formulate the following recommendations.

#### TABLE 1

| Number of included and excluded articles, the types of included ones and the reasons for exclusion |
|-------------------------------------------------|-------------------------------------------------|
| 317 studies retrieved | 162 articles included | 155 articles excluded |
| 59 randomized, controlled or crossover trials | 43 interventions or outcomes unrelated to suctioning |
| 28 nonrandomized crossovers or comparative cohorts | 16 related to diagnostic tests |
| 49 observational studies of precautions | 50 reviews or descriptive articles |
| 26 animal or test-lung models | 31 observational studies not dealing with precautions |
| | Five animal or test models unrelated to suctioning |
| | Three no data |
| | Seven other |

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### TABLE 2
Summary of studies in children and infants

<table>
<thead>
<tr>
<th>Author, year (reference)</th>
<th>Population</th>
<th>Treatment allocation</th>
<th>First outcome (primary study outcome)</th>
<th>Second outcome (could also be primary)</th>
<th>P</th>
</tr>
</thead>
</table>
| Cohen-Addad et al, 1989 (16) | Deliveries with meconium staining, not intubated (n=60) | Type of catheter
  - RCT
  - Intrapartum suctioning, syringe (bulb)
  - Intrapartum suctioning, DeLee catheter | Incidence of MAS
  - Overall: 7/60 (11.7%)
  - 3/31 De Lee (9.7%)
  - 4/29 Bulb (13.8%) | Meconium recovered
  - means
  - Bulb: 0.24 mL
  - De Lee: 0.16 mL | NS |
| Locus et al, 2000 (17) | Moderate or thick meconium (n=107) | Type of suction device
  - Cohort
  - 1. Bulb suction
  - 2. DeLee suction | Meconium suctioned from below vocal cord (0.22 and 0.24 mL) | MAS or death | 0.22 |
| Daga et al, 1998 (18) | Meconium in utero, not intubated (n=49) | Oropharyngeal versus tracheal
  - RCT
  - 1. OP
  - 2. OP + tracheal suctioning | Incidence of death
  - 0/23 in 1
  - 1/26 in 2 | NS |
| Brogden et al., 1989 (19) | Newborn with 'normal' vaginal delivery, meconium stained (n=726) | Indication-tracheal
  - Cohort
  - 1. Tracheal suction
  - 2. OP | Particulate/pea soup amniotic fluid
  - 40% in 1 and 35% in 2 | NS |
| Estol et al, 1992 (20) | Clear amniotic fluid, not intubated (n=40) | Indication-pharyngeal
  - Cohort
  - 1. Suction
  - 2. No suction | Expiratory and inspiratory dynamic compliance, NS | NS |
| Barnes et al., 1991 (21) | Respiratory distress, intubated and mechanically ventilated (n=32) | Hyperinflation
  - RC
  - 1. CPT, bagging, suction, bagging, suction
  - 2. CPT, suction, bagging, suction, bagging
  - 3. CPT, bagging, suction, bagging, suction, bagging, hyperinflation | TcPO2 values, NS | NS |
| Cunningham et al, 1984 (22) | Premature babies (RDS and non RDS), intubated and ventilated (n=8) | Hyperoxygenation and hyperventilation
  - RC
  - 1. O2 10% increase, 5 breaths/15 s
  - 2. O2 20% increase, 5 breaths/15 s
  - 3. O2 15% increase, 10 breaths/30 s
  - 4. 20% increase, 10 breaths/30 s | TcPO2
  - RDS babies had a decrease in TcPO2 in all groups
  - Non RDS babies in conditions 1 and 3 had slight fluctuation in O2. With conditions 2 and 4, TcPO2 rose with first inflation
  - Condition 1 resulted in more hypoxic events than 2, 3 and 4 | P<0.02 |
| Walsh et al, 1987 (23) | Preterm with RDS, intubated and ventilated (n=21) | Preoxygenation
  - 1. Standard hygiene
  - 2. O2 increased until TcPO2 above 90 | P<0.006 |
| Cabal et al., 1984 (24) | Premature, intubated and ventilated (n=10) | Preoxygenation
  - Cohort
  - 1. Preoxygenation
  - 2. No preoxygenation | PaO2 postinsuffation values
  - 50 mmHg in 1
  - 63 mmHg in 2 | P<0.02 |
| Skov et al., 1992 (25) | Intubated and ventilated newborns, gestational age of 25 to 40 weeks | Preoxygenation
  - Cohort
  - 1. First time suction with preoxygenation, second time without
  - 2. Had preoxygenation before second suction | Hemoglobin oxygenation index and SaO2
  - Preoxygenation reduced the decrease in both | P<0.05 |
| Wilson et al, 1991 (26) | Premature with RDS, intubated and ventilated (n=37) | Suction frequency
  - RCT
  - 1. Suction q6h
  - 2. Suction q12h | Median time on ventilation
  - 28 h in 1
  - 39 h in 2 | NS |
| Fancorri and Duc, 1987 (27) | Prematures intubated, paralyzed, ventilated (n=28) | Muscle relaxant
  - Cohort
  - 1. With muscle paralysis
  - 2. Without muscle paralysis | ICP increase during suctioning
  - when not paralyzed (13.7 to 15.8 in 1 and 12.5 to 28.5 in 2) | P<0.01 |

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### TABLE 2 (continued)
Summary of studies in children and infants

<table>
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<th>First outcome (primary study outcome)</th>
<th>Second outcome (could also be primary)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saarenmaa et al, 1996 (28)</td>
<td>Ventilated, intubated newborns, gestational age of 29-36 weeks (n=10)</td>
<td>Sedation RC 1. Saline 2. Alfentanil 10 g/kg IV 3. Alfentanil 20 g/kg</td>
<td>Pain and behavioural scale less in group 3, but muscle rigidity frequent</td>
<td>P=0.031</td>
</tr>
<tr>
<td>Pokela, 1994 (29)</td>
<td>Preterm or term RDS, less than one week old (n=84)</td>
<td>Sedation RC 1. Sedation (meperidine, 1 mg/kg) 2. No sedation (saline)</td>
<td>Longer duration of hypoxemia in 2 (44 s versus 29 s)</td>
<td>P=0.002 No difference in BP or HR NS</td>
</tr>
<tr>
<td>Ninan et al, 1986 (30)</td>
<td>Respiratory failure, premterm, RDS, ventilation (n=7)</td>
<td>Sedation Cohort 1. Sedation (phenobarbital or chloral hydrate) 2. No sedation Instillation 1. No sedation 2. Sedation + saline instillation</td>
<td>Significant blunting of ICP in 1</td>
<td>P 0.01 Mean arterial and cerebral perfusion pressures blunted in 1 P 0.01 P 0.05</td>
</tr>
<tr>
<td>Shorten et al, 1991 (31)</td>
<td>Newborn, gestational age 25 to 40 weeks, RDS, intubated and ventilated (n=27)</td>
<td>Sedation RC 1. Suction 2. Suction + saline instillation</td>
<td>No effect on HR NS</td>
<td>Difference in mean arterial blood pressure, NS NS</td>
</tr>
<tr>
<td>Beeram and Dhanireddy, 1992 (32)</td>
<td>RDS or MAS, intubated and ventilated (n=18)</td>
<td>Instillation RC 1. Suction no saline 2. Suction + saline (1.0 mL)</td>
<td>Dynamic compliance: no change in RDS or MAS Airway resistance: no change in RDS but decrease in MAS with saline</td>
<td>NS In RDS, significant drop in SaO2 with saline; in MAS, significant drop in SaO2 in both groups P 0.05</td>
</tr>
<tr>
<td>Gunderson et al, 1986 (33)</td>
<td>Prematures with RDS, intubated, ventilated (n=11)</td>
<td>Adaptor Cohort 1. ETAdaptor 2. ETAdaptor with end hole</td>
<td>Change in TcPO2 2.9% in PVETS 21.4% in NVETS</td>
<td>P=0.001 Change in HR 3.4% in PVETS 11.3% in NVETS P=0.047</td>
</tr>
<tr>
<td>Bodai et al, 1989 (34)</td>
<td>Respiratory failure, intubated, ventilated (n=15)</td>
<td>Adaptor RC 1. PVETS 2. NVETS</td>
<td>SaO2 desaturation with NeoSafe 9.6%, with conventional suction 22%</td>
<td></td>
</tr>
<tr>
<td>Zmora and Merritt, 1980 (35)</td>
<td>Newborns, RDS, pneumonia, asphyxia, GI surgery, intubated, ventilated (n=13)</td>
<td>Adaptor Cohort 1. ETAdaptor with side holes 2. ETAdaptor with end hole</td>
<td>TcPO2 decreased less with side-hole adaptor</td>
<td>P=0.01 Occurrence of bradycardia less in six, no different in 11 and considerably worse in 1 P=0.01</td>
</tr>
<tr>
<td>Cabal et al, 1979 (36)</td>
<td>Severe RDS, intubated, ventilated (n=8, 256 procedures)</td>
<td>Adaptor Cohort 1. Disconnected from ventilator, preoxygenated and hyperventilated manually 2. Suctioning adapter, no hyperventilation</td>
<td>Peak drop in HR 127 in 1, 137 in 2. HR deceleration longer in 1 (7.5 s) than 2 (6.8 s)</td>
<td>P=0.001 Peak drop in SaO2 to 86 in 1 and 90 in 2 P=0.001</td>
</tr>
<tr>
<td>Kerem et al, 1990 (37)</td>
<td>Infants in the ICU, intubated and ventilated, one day to 10 years (n=25)</td>
<td>Hyperinflation/Hyperoxygenation RC 1. Suction 2. Preoxygenation by ventilator 3. Hyperinflation presuction 4. Hyperinflation post suction</td>
<td>Drop in PaO2 of 24 mmHg in 1</td>
<td>P=0.001 Drop in SaO2 of 4.4% in 1 P=0.001</td>
</tr>
<tr>
<td>Feaster et al, 1985 (38)</td>
<td>Children requiring chronic respiratory support, eight months to three years, (BPD and myopathy) (n=7)</td>
<td>Hyperinflation/Hyperoxygenation RC 1. Hyperinflation alone 2. Hyperinflation + hyperoxygenation 3. Hyperinflation + hyperventilation 4. Hyperinflation + hyperventilation + hyperoxygenation</td>
<td>Change in SpO2 presuction to 30 s after suction Only statistically significant</td>
<td>NS (clinically)</td>
</tr>
</tbody>
</table>

**BP Blood pressure; BPD Bronchopulmonary dysplasia; CPT Chest physical therapy; ETT Endotracheal tube; FIO2 Fractional inspired concentration of oxygen; GI Gastrointestinal; HR Heart rate; ICP Intracranial pressure; ICU Intensive care unit; IV Intravenous; MAS Meconium aspiration syndrome; NS Not significant; NVETS Nonventilated endotracheal suction; OP Oropharyngeal suctioning; PaO2 Arterial pressure of oxygen; PVETS Partially ventilated endotracheal suction with adapter; q6h Every 6 h; q12h Every 12 h; RC Randomized crossover; RCT Randomized, controlled trial; RDS Respiratory distress syndrome; SaO2 Arterial oxygen saturation; SpO2 Oxygen saturation by pulse oximetry; Stats Statistics; TcPO2 Transcutaneous partial pressure of oxygen**

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**Nonintubated neonates with evidence of MAS:** Meconium is the bowel content of a fetus, usually passed within 48 h of delivery. If meconium is passed in utero, it is possible for the fetus to aspirate the meconium before or during delivery (16,17). This aspirated meconium may cause airway obstruction and air trapping, and may support bacterial growth. The usual practice is to monitor amniotic fluid for the presence of meconium, and to suction the infant at the perineum during delivery (16,17).

Three types of catheters and suctioning systems have been used to suction the upper airway of neonates at birth. With the bulb catheter, the suctioning pressure is provided by intermittent manual compression of the bulb. With the DeLee catheter, the operator provides the suctioning pressure by generating a negative pressure orally. This catheter places operators at risk of suctioning the infant’s airway contents into their mouth. A third system — wall suction — provides suctioning pressure by a central vacuum system.

After the infant with evidence of meconium-stained amniotic fluid has been delivered, the two approaches to clinical care are suctioning of the infant’s oropharynx, and suctioning of the infant’s trachea using direct visualization and a meconium adaptor to apply suction directly to the endotracheal tube (ETT) (18,19).

There were two studies (one RCT [16], one comparative cohort study [17]) comparing DeLee with bulb suctioning in neonates with evidence of MAS. In these two studies, there was no difference between DeLee and bulb suctioning for frequency of MAS and death, or the amount of meconium recovered; however, both studies lacked the statistical power to detect such a difference. No studies used wall suctioning in this patient population.

Two studies compared oropharyngeal suctioning with tracheal suctioning in neonates with evidence of MAS; one was an RCT (18) and the other a comparative cohort study (19). Neither study provided evidence of a benefit for tracheal suctioning over oropharyngeal suctioning alone. In the comparative cohort study (19), which was subject to allocation bias, some harmful effects of tracheal suctioning were noted — ie, the presence of stridor.

**Recommendations:** The DeLee and bulb methods are equally effective in the aspiration of meconium when suctioning nonintubated neonates with evidence of MAS. Because there is a risk to the operator associated with DeLee suction, it is not recommended. In vaginally delivered infants with MAS, no evidence exists to suggest a difference in the effectiveness of oropharyngeal compared with tracheal suctioning.

**Nonintubated neonates without evidence of MAS:** One published comparative cohort study was found on the indication for oropharyngeal suctioning in full-term, vaginally delivered infants without evidence of MAS (20). The study described the effect of oropharyngeal suctioning at birth on the mechanical properties of the respiratory system. There were no differences between the groups with respect to any of the outcomes (dynamic compliance on inspiration and expiration, and total resistance of the respiratory system). Given the paucity of literature, we are unable to make recommendations regarding suctioning of vaginally delivered infants (who have no evidence of MAS immediately after birth).

**Intubated neonates, preterm and full-term — effect of hyperoxygenation, preoxygenation and hyperventilation:** Hyperoxygenation is defined as the administration of oxygen at an inspired fractional concentration (FiO2) that is greater than that which the individual is receiving or usually requires before, during or after suctioning. In an effort to reduce hypoxemia during suctioning, increased levels of oxygenation may be delivered before, during and/or after suctioning. Preoxygenation refers to hyperoxygenation before initiation of suctioning. Hyperventilation is an increase in ventilation but does not imply an increase in oxygen concentration.

The effects of different degrees of hyperventilation and preoxygenation in preterm neonates were examined in two studies — one randomized (21) and one nonrandomized crossover study (22) — neither of which used a control group of no preoxygenation and hyperventilation. Due to differences in outcomes and interventions, and problems with the analysis of the data, we are unable to make a recommendation on the timing of hyperventilation and hyperoxygenation (ie, before, during or after suctioning), or on different combinations of hyperoxygenation and hyperventilation in this patient population.

Three studies examined the effectiveness of preoxygenation alone: one randomized (23) and two nonrandomized crossover studies (24,25). One study (25) included some full-term infants. All three studies showed an improvement in oxygenation with preoxygenation; however, none of these studies addressed the potentially negative effects of hyperoxia in preterm infants (eg, retinopathy of prematurity).

**Recommendation:** Individualized preoxygenation should be applied in intubated and ventilated preterm and full-term infants to improve oxygenation.

**Intubated neonates, preterm and full-term — frequency of suctioning:** One study (RCT) examined the effect of suctioning frequency (either every 6 h or every 12 h) (26). Although the study reported a relative reduction in death of 7% in those suctioned every 6 h, the difference was not statistically significant and might have resulted from differences in other confounding variables. The high variability in death rate in the preterm population precludes a recommendation based solely on this study.

**Intubated neonates, preterm and full-term — drugs for muscle paralysis:** The relation between cerebral perfusion and hypoxic-ischemic lesions such as periventricular leukomalacia and intraventricular hemorrhage is not well understood. Some researchers have speculated that maintaining a consistent cerebral perfusion pressure (CPP) may reduce the potential for hypoxic-ischemic lesions (27). One nonrandomized crossover study (27) examined the effect of muscle paralysis...
using pancuronium intravenous bromide on variations in CPP in preterm infants. This study showed that muscle paralysis in ventilated preterm infants minimizes increases in intracranial pressure (ICP) during suctioning; however, the long term benefit of a transient minimization in ICP is unknown. Before the use of muscle paralyzing drugs can be recommended, further study of this intervention is warranted.

**Intubated neonates, preterm and full term – drugs for pain relief during suctioning:** Term and preterm infants experience pain during suctioning, and local and systemic analgesics are widely used. Because crying is not easily assessed in intubated infants, physiological parameters (vital signs, arterial oxygen saturation [\(\text{SaO}_2\]), stress hormones, cerebral blood flow) are used as indicators of pain.

Three studies – one randomized crossover (28), one RCT (29) and one nonrandomized crossover trial (30) – examined pain relief during suctioning using a variety of drugs (alfentanil, meperidine, phenobarbital). Although these studies do not provide clear evidence of benefit or harm of analgesic use, clinicians should consider the need for pain relief for infants on an individual basis.

**Intubated neonates, preterm and full term – instillation:** Instillation refers to the instillation of aliquots of saline directly into the ETT or tracheostomy tube to facilitate secretion removal. Two randomized crossover studies examined the effect of instillation (31,32). Neither study made comparisons between groups; they only reported differences over time. No recommendation is offered on instillation of saline in ventilated preterm and full-term infants.

**Intubated neonates, preterm and full term – adaptor:** Adaptors with capped apertures have been a part of the patient ventilator circuit for many years. When opened, the aperture permits suctioning of intubated, ventilated patients without disconnection from the ventilator circuit. A meconium adaptor is used to connect the ETT to the suction tubing, and a porthole on the adaptor allows control over the application of suction.

Two randomized crossover (33,34) and two nonrandomized crossover (35,36) trials examined the effects of adaptors designed to maintain pressure and/or ventilation during suctioning. Two studies (33,34) used a modified Wye adaptor with an end hole for suctioning, while the other two (35,36) used a modified ETT adaptor that allows suctioning through a side hole. All studies showed a beneficial effect for the use of an adaptor on oxygenation.

**Recommendation:** Adaptors should be used in mechanically ventilated preterm and full-term infants with respiratory distress syndrome for whom hypoxemia is a concern.

**Intubated infants and children – hyperventilation and hyperoxygenation:** There were two randomized crossover studies on the effect of hyperventilation and hyperoxygenation in infants and children during suctioning (37,38). One of the studies (37) examined intubated children, aged one day to 10 years (nontyanotic heart disease, central nervous system diseases, respiratory failure and sepsis), while the other (38) examined children aged eight months to three years requiring chronic respiratory support. These studies compared several protocols using combinations of hyperventilation and hyperoxygenation. Neither study performed comparisons between groups; however, in pre-post suction analysis, the studies indicated that no oxygen desaturation occurred with suctioning with hyperoxygenation and hyperventilation, or a combination of these techniques. There is insufficient evidence to make recommendations concerning hyperventilation and hyperoxygenation in infants and children.

**Summaries and recommendations in adults:** In the adult population, we examined the evidence related to the following techniques and tools for suctioning: preoxygenation, hyperoxygenation, hyperinflation, insufflation, methods of delivery, use of adaptors, open versus closed systems, catheter designs and types, instillation and use of medication. Where study populations differed significantly, evidence was summarized based on patient population. Tables 3 and 4 contain information about the studies that was used to formulate the following recommendations.

**Preoxygenation:** In addition to evacuating secretions, endotracheal suctioning removes oxygen-containing air from the tracheobronchial tree. Consequently, a decrease in the arterial pressure of oxygen (\(\text{PaO}_2\)) is associated with suctioning. Various methods of preoxygenating patients, particularly those who are intubated, have been used to ameliorate this reduction.

Two randomized crossover (39,40) and two comparative cohort (41,42) studies on preoxygenation were retrieved. Both of the randomized crossover studies involved patients with respiratory failure. One of the comparative cohort studies examined the effects of preoxygenation in postoperative cardiac surgery (42), while the other focused on patients undergoing lung resections (41). Two of the studies used ventilators to deliver the preoxygenation (39,40), one used a manual resuscitation bag (MRB) with 15 L/min oxygen flow (42), and another study used either an MRB or a ventilator (41). All four studies showed a beneficial effect of preoxygenation on \(\text{PaO}_2\) or \(\text{SaO}_2\).

**Recommendation:** Mechanically ventilated adult patients should receive additional oxygenation before suctioning.

**Hyperoxygenation and hyperinflation:** Five randomized crossover trials examined the combination of hyperoxygenation and hyperinflation (43-47). Because the outcomes of interest were different, recommendations for medical/surgical patients were separated from those for head-injured patients. Hyperoxygenation and hyperinflations were delivered by either MRB or ventilator. Studies varied in the attention paid to the volumes and amount of oxygen actually being delivered to the patient. Only some studies accounted for ventilator washout time.

There were four studies in mechanically ventilated medical and surgical patients, which showed that hyperoxygenation helps in maintaining oxygen levels (43-46); however, the added value of hyperinflation was unclear. In one study involving patients with chronic obstructive pulmonary dis-
TABLE 3
Summary of studies in adults related to preoxygenation, hyperoxygenation, hyperinflation, oxygen insufflation and the use of adaptors

<table>
<thead>
<tr>
<th>Author, year (reference)</th>
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<th>Treatment allocation</th>
<th>Primary study outcome</th>
<th>Secondary outcome (could also be primary)</th>
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<tbody>
<tr>
<td>Craig et al, 1984 (39)</td>
<td>Intubated, ventilated for acute respiratory failure (n=17)</td>
<td>Preoxygenation</td>
<td>Decrease in $\text{SaO}_2$ greater with no preoxygenation</td>
<td>$P&lt;0.04$ (clinically NS)</td>
</tr>
<tr>
<td>Brown et al, 1983 (40)</td>
<td>Respiratory failure, ventilated, copious secretions (COPD) (phase 1, n=12; phase 2, n=10; phase 3, n=9)</td>
<td>Preoxygenation/use of adaptor</td>
<td>Phase 1</td>
<td>$P&lt;0.05$ Mean desaturation time was shortened by either four post-breaths or using adaptor</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Phase 2</td>
<td>$P&lt;0.05$ No differences in desaturation time or recovery</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Phase 3</td>
<td></td>
</tr>
<tr>
<td>Downes et al, 1981 (41)</td>
<td>Adolescents and adults intraoperative lung resections, anesthetized (n=10, intact pleurae)</td>
<td>Preoxygenation Cohort</td>
<td>Mean changes in $\text{SaO}_2$ between hyperventilated and nonhyperventilated were all different</td>
<td>$P&lt;0.01$ Mean change in $\text{SaO}_2$ not significantly different between 1 and 2</td>
</tr>
<tr>
<td>Belknap et al, 1980 (42)</td>
<td>Postop CABG, ventilated (n=13)</td>
<td>Preoxygenation or bag versus ventilator Cohort</td>
<td>Mean $\text{PaO}_2$, 157 mmHg when hyperinflated with ventilator; 188 mmHg when hyperoxygenated with bag</td>
<td>$P&lt;0.05$ No differences found in $\text{PCO}_2$</td>
</tr>
<tr>
<td>Lookinland and Appel, 1991 (43)</td>
<td>Non head-injured trauma patients, no underlying lung disease (n=24)</td>
<td>Hyperoxygenation and hyperinflation</td>
<td>Mean arterial pressure, pulmonary artery pressure, mean capillary wedge pressure and cardiac output: no differences among the four groups</td>
<td>NS Mean $\text{PaO}_2$ in 2 and 4 greater than in 1 and 3 at end of succioning</td>
</tr>
<tr>
<td>Rogge et al, 1989 (44)</td>
<td>Patients with x-ray evidence of COPD, ventilated (n=11)</td>
<td>Hyperoxygenation and hyperinflation</td>
<td>BP, $\text{SpO}_2$, HR, cardiac rhythm: no difference in any variable during either protocol</td>
<td>NS</td>
</tr>
<tr>
<td>Skelley et al, 1980 (45)</td>
<td>Postoperative cardiac surgery, ventilated (n=11)</td>
<td>Hyperoxygenation and hyperinflation</td>
<td>Immediately after and at 90 s and 180 s, significant difference in $\text{PaO}_2$ between 1 and 3, and between 2 and 3</td>
<td>$P&lt;0.01$, but clinically NS</td>
</tr>
<tr>
<td>Langrehr et al, 1981 (46)</td>
<td>Postoperative cardiac surgery, ventilated (n=10)</td>
<td>Hyperoxygenation and hyperinflation/insufflation</td>
<td>No difference in $\text{PaO}_2$ between 3 and 4 at any point</td>
<td>$P&lt;0.05$ Immediately after succioning, significant differences between 2 and 3, 2 and 4, and 1 and 2</td>
</tr>
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### TABLE 3 (continued)
Summary of studies in adults related to preoxygenation, hyperoxygenation, hyperinflation, oxygen insufflation and the use of adaptors

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<tr>
<td>Rudy et al, 1991 (47)</td>
<td>Severe head injury, intubated and ventilated (n=30)</td>
<td><strong>Hyperoxygenation and hyperinflation</strong>&lt;br&gt;RC&lt;br&gt;1. Two ET suctioning sequences, hyperoxygenation and hyperoxygenation with hyperinflation (135% TV)&lt;br&gt;2. Three ET suctioning sequences, hyperoxygenation (four breaths of 100% O₂) and one with hyperoxygenation and hyperinflation (135% TV)</td>
<td>Mean ICP&lt;br&gt;No difference with addition of hyperinflation in group 1&lt;br&gt;Significant differences across time&lt;br&gt;No significant difference between 2 and 3 suction sequences</td>
<td>MAP, HR and CPP all behaved similarly to mean ICP&lt;br&gt;P&lt;0.001 NS</td>
</tr>
<tr>
<td>Kelly et al, 1987 (48)</td>
<td>Before cardiac surgery, fully anesthetized (n=38)</td>
<td><strong>Oxygen insufflation</strong>&lt;br&gt;RC&lt;br&gt;1. 20 s suctioning with O₂ using double lumen catheter&lt;br&gt;2. 20 s of continuous suctioning without O₂ insufflation</td>
<td>With insufflation, PaO₂ increased by 5 torr&lt;br&gt;Without insufflation, PaO₂ dropped 16 torr</td>
<td>PaO₂&lt;br&gt;P&lt;0.01 NS</td>
</tr>
<tr>
<td>Dam et al, 1994 (49)</td>
<td>Postoperative CABG surgery, ventilated (n=20)</td>
<td><strong>Oxygen insufflation</strong>&lt;br&gt;RC&lt;br&gt;1. Suction with a double lumen insufflation catheter 100% O₂&lt;br&gt;2. Suction with a single lumen catheter, ventilator 100% O₂</td>
<td>Arterial pressure differences favouring insufflation</td>
<td>PaO₂&lt;br&gt;P&lt;0.02 NS</td>
</tr>
<tr>
<td>Smith et al, 1987 (50)</td>
<td>Trauma, medical, or neurosurgical patients, ventilated (n=18)</td>
<td><strong>Oxygen insufflation</strong>&lt;br&gt;RC&lt;br&gt;1. Double lumen (Jinotti) catheter, 100% O₂, and no preoxygenation&lt;br&gt;2. Preoxygenation, 3 min ventilator at 100% O₂, then suction</td>
<td>No difference in SpO₂ between groups</td>
<td>NS</td>
</tr>
<tr>
<td>Taft et al, 1991 (51)</td>
<td>Patients in the ICU, intubated and ventilated (n=23)</td>
<td><strong>Oxygen insufflation</strong>&lt;br&gt;RC&lt;br&gt;1. Conventional method: bag, five breaths 100% O₂ before and after suction two times&lt;br&gt;2. Insufflation method: 15 s of O₂ at 15 L/min with single lumen catheter</td>
<td>PaO₂ and SaO₂: significant interaction between preoxygenation method and time on mean PaO₂ and SaO₂ statistics for main effects of method not reported</td>
<td>PaO₂ and SaO₂&lt;br&gt;P&lt;0.001 No significant difference in HR with either method or time&lt;br&gt;NS</td>
</tr>
<tr>
<td>Stone et al, 1989 (52)</td>
<td>Postcardiac surgery, intubated and ventilated (n=8)</td>
<td><strong>Hyperinflation (degree)</strong>&lt;br&gt;RC&lt;br&gt;Hyperinflated at five different weight-based inspiratory volumes (TV 12, 14, 16, 18 mL/kg)</td>
<td>Mean arterial pressure: no differences between groups; mean increase of 15 mmHg over three hyperinflation sequences (not volume dependent)</td>
<td>No significant difference in PaO₂ with different volumes&lt;br&gt;P&lt;0.001 NS</td>
</tr>
<tr>
<td>Stone, 1990 (53)</td>
<td>Postcardiac surgery, intubated and ventilated (n=26)</td>
<td><strong>Hyperinflation (degree)</strong>&lt;br&gt;RC&lt;br&gt;Hyperinflated at five different weight-based inspiratory volumes (TV 12, 14, 16, 18 mL/kg)</td>
<td>No clinically significant difference in PaO₂ with volume&lt;br&gt;(clinically)</td>
<td>No differences in HR based on lung volume&lt;br&gt;NS</td>
</tr>
<tr>
<td>Preusser et al, 1988 (54)</td>
<td>Stable postcardiac surgery (n=10)</td>
<td><strong>Bag versus ventilator</strong>&lt;br&gt;RC&lt;br&gt;Three lung inflation breaths, 100% O₂ by:&lt;br&gt;1. A preprimed manual bag at two different volumes/kg&lt;br&gt;2. A preprimed vent at two different volumes/kg (same as in 1)</td>
<td>MAP showed no difference between groups 1 and 2</td>
<td>No significant difference in PaO₂ between groups 1 and 2&lt;br&gt;NS</td>
</tr>
<tr>
<td>Stone et al, 1991 (55,56)</td>
<td>Postcardiac surgery, intubated and ventilated (n=34)</td>
<td><strong>Hyperinflation (degree)</strong>&lt;br&gt;RC&lt;br&gt;Hyperinflated at five inspiratory volumes (TV 12, 14, 16, 18 mL/kg)</td>
<td>MAP increased with sequences, not volume dependent</td>
<td>Cardiac output and PAP increased with sequences, not volume dependent&lt;br&gt;NS</td>
</tr>
<tr>
<td>Kerr et al, 1997 (57)</td>
<td>Severe head injured, ventilated, intubated (phase 1: n=29; phase 2: n=3 to 37)</td>
<td><strong>Hyperventilation</strong>&lt;br&gt;RC&lt;br&gt;Phase 1&lt;br&gt;1. Hyperventilation (100%), four breaths&lt;br&gt;2. Hyperventilation (100%), eight breaths&lt;br&gt;Phase 2&lt;br&gt;1. Hyperventilation (100%), four breaths&lt;br&gt;2. Hyperventilation (100%), 30 breaths</td>
<td>ICP and cerebral perfusion pressure: no difference in phase 1; increasing rate and number of breaths resulted in significant difference up to 10 min after in phase 2</td>
<td>P&lt;0.001 NS</td>
</tr>
</tbody>
</table>
TABLE 3 (continued)

Summary of studies in adults related to preoxygenation, hyperoxygenation, hyperinflation, oxygen insufflation and the use of adaptors

<table>
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<th>Secondary outcome (could also be primary)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Campbell, 1991 (58)</td>
<td>Severe head injured patients, ventilated (n=10)</td>
<td>Hyperinflation RC</td>
<td>No effect of volume on mean ICP</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Hyperinflation at three different volumes (TV 14, 16, 18 mL/kg)</td>
<td>Mean ICP increased over time and through the hyperinflation and suctioning sequences</td>
<td></td>
</tr>
<tr>
<td>Grap et al, 1996 (59)</td>
<td>Medical, surgical, trauma ICU patients, moderate or severe lung injury, intubated, ventilated (n=29)</td>
<td>Bag versus ventilator RC</td>
<td>Significant difference in PaO2 between groups</td>
<td>P&lt;0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1. Hyperoxygenation with bag using 15 L/min O2 flow before and after suctioning</td>
<td>Mean PaO2 was higher during ventilated breaths immediately and 30 s post suctioning</td>
<td>No significant difference in mean between the groups NS</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Three mechanical breaths of 100% O2 before and after suctioning</td>
<td>PaO2 increase greater in group 2</td>
<td>P&lt;0.05</td>
</tr>
<tr>
<td>Pierce and Piazza, 1987 (60)</td>
<td>Cardiac surgery patients (n=30)</td>
<td>Bag versus ventilator RC</td>
<td>PaO2 rose after suctioning compared with control for both methods</td>
<td>P=0.0007 SaO2 behaved similarly but may not be clinically meaningful NS</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1. Postoxygenation with bag at 100% O2</td>
<td>PaO2 increase greater with ventilator</td>
<td>P=0.001</td>
</tr>
<tr>
<td>Conforti, 1982 (61)</td>
<td>Postop coronary revascularization and/or valve replacement, ventilated without PEEP (n=33)</td>
<td>Bag versus ventilator RC</td>
<td>Baseline to post suctioning change in SaO2 significantly greater with ventilatory sighing than with bagging</td>
<td>P&lt;0.028</td>
</tr>
<tr>
<td>Anderson, 1989 (62)</td>
<td>Critically ill (mixed medical/trauma), intubated, ventilated (n=28)</td>
<td>Bag versus ventilator Cohort</td>
<td>Mean decrease in PaO2 through ventilator 24.6%; mean decrease off ventilator 67.4%</td>
<td>P&lt;0.001 Suctioning time through adapter 2.3 min; off ventilator time 1.8 min NS</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1. Hyperinflation by bag (15 L/min O2)</td>
<td>P=0.001</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Hyperinflation by sigh on ventilator (100% O2)</td>
<td>P=0.001</td>
<td></td>
</tr>
<tr>
<td>Belling et al, 1978 (63)</td>
<td>Postoperative cardiac surgery, ventilated (n=20)</td>
<td>Use of adaptor</td>
<td>SaO2 fell less during use of adaptor</td>
<td>P&lt;0.01 Difference in HR response to the two protocols, NS NS</td>
</tr>
<tr>
<td>Jung and Newman, 1982 (64)</td>
<td>Acute lung disease in ICU, intubated (n=18)</td>
<td>Use of adaptor RC</td>
<td>Mean decrease in PaO2 through adaptor 24.6%; mean decrease off ventilator 67.4%</td>
<td>P&lt;0.001 Suctioning time through adapter 2.3 min; off ventilator time 1.8 min NS</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1. Suctioned via standard ETT (disconnected from ventilator)</td>
<td>SaO2 fell less during use of adaptor</td>
<td>P&lt;0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. Suctioned through modified suction adaptor</td>
<td>Difference in HR response to the two protocols, NS</td>
<td>NS</td>
</tr>
<tr>
<td>Douglas and Larson, 1985 (65)</td>
<td>Medical/surgical ICU patients, no history of chronic lung disease, ventilated (n=12)</td>
<td>PEEP valve RC</td>
<td>No significant effect of PEEP on PaO2</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1. PEEP adaptor on the bag</td>
<td>No significant difference in mean SaO2</td>
<td>P=0.005</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. No adaptor on bag</td>
<td>PaO2 increased from 84 to 122 mmHg with PEEP valve and returned to baseline by 5 min</td>
<td></td>
</tr>
<tr>
<td>Schumann and Parsons, 1985 (66)</td>
<td>ARDS patients, ventilated (n=15)</td>
<td>PEEP valve RC</td>
<td>Without PEEP, PaO2 increased from 92 to 106 mmHg and returned to baseline by 5 min</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1. PEEP valve</td>
<td>P=0.005</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. No PEEP valve</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

ARDs Acute respiratory distress syndrome; BP Blood pressure; bpm Beats/min; CABG Coronary artery bypass graft; COPD Chronic obstructive pulmonary disease; CPP Cerebral perfusion pressure; ET Endotracheal; ETT Endotracheal tube; FIO2 Fractional inspired concentration of oxygen; HR Heart rate; ICP Intracranial pressure; ICU Intensive care unit; MAP Mean arterial pressure; NS Not significant; PaO2 Arterial pressure of oxygen; PEEP Positive end expiratory pressure; Postop Postoperative; RC Randomized crossover; RCT Randomized controlled trial; SaO2 Arterial oxygen saturation; SpO2 Oxygen saturation by pulse oximetry; TV Tidal volume

Recommendations: For mechanically ventilated trauma, cardiac and chronic obstructive pulmonary disease patients, hyperoxygenation should be used during suctioning to maintain oxygen levels. For severely head-injured patients, hyperoxygenation should be used during suctioning to maintain oxygen levels. For severely head-injured patients, hyperoxygenation should be used during suctioning to maintain oxygen levels.

ease (44), lower levels of hyperoxygenation were safe and adequate; this suggests that hyperoxygenation should be individualized. For patients with severe head injury where ICP was a concern (47), there was no difference between hyperoxygenation alone and the combination of hyperoxygenation and hyperinflation on ICP, arterial blood pressure, heart rate or saturation.
### TABLE 4
Summary of studies in adults related to open and closed systems, types and designs of catheters, instillation of saline, drugs and jet ventilation

<table>
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<tr>
<th>Author, year (reference)</th>
<th>Population</th>
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<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adams et al, 1997 (68)</td>
<td>Postoperative liver transplant, ventilated (n=20)</td>
<td>Open versus closed RCT 1. Open 2. Closed</td>
<td>Microbiological assessment of secretions (colony count in aspirates); no significant difference at extubation</td>
<td>No cases of pneumonia</td>
<td>P=0.05</td>
</tr>
<tr>
<td>Ritz et al, 1986 (70)</td>
<td>Ventilated, with positive sputum culture</td>
<td>Open versus closed Prospective crossover 1. Open 2. Closed</td>
<td>No significant difference between sputum culture from catheter tips and sputum sample</td>
<td>No significant difference between sputum culture from catheter tips and sputum sample</td>
<td>NS</td>
</tr>
<tr>
<td>Cobley et al, 1991 (71)</td>
<td>Ventilated, colonized with Gram-negative bacteria but without significant infection (n=11)</td>
<td>Open versus closed Prospective crossover 1. Open 2. Closed</td>
<td>On average, 18.6 fewer colonies with closed from the air sampled</td>
<td>Increased colony counts (settle) with open system at 50 and 100 cm</td>
<td>P&lt;0.001</td>
</tr>
<tr>
<td>Wu et al, 1993 (72)</td>
<td>Anesthetized patients (n=8)</td>
<td>Open versus closed Randomized crossover 1. Open 2. Closed</td>
<td>Significantly greater drop in PaO2 with open system</td>
<td>No significant difference in PaO2 with open system</td>
<td>P&lt;0.05</td>
</tr>
<tr>
<td>Hardie and Kirchhoff, 1990 (73)</td>
<td>Ventilated, respiratory failure (n=28)</td>
<td>Open versus closed Randomized crossover 1. Open 2. Closed</td>
<td>Stable O2 saturation during and following suction for both</td>
<td>No significant difference in O2 saturation during and following suction for both</td>
<td>NS</td>
</tr>
<tr>
<td>Carlson et al, 1987 (74)</td>
<td>Intubated, ventilated (n=20)</td>
<td>Open versus closed Cohort 1. Open 2. Closed</td>
<td>No clinically important difference detected in arterial blood gases</td>
<td>No clinically important difference detected in arterial blood gases</td>
<td>NS (clinically)</td>
</tr>
<tr>
<td>Johnson et al, 1997 (75)</td>
<td>Intubated, ventilated (n=35; 276 procedures)</td>
<td>Open versus closed Prospective cohort Quasirandomized 1. Open 2. Closed</td>
<td>O2 saturation 30 s postsuctioning 1.4% increase with closed</td>
<td>PaO2 96 with open versus 99 with closed</td>
<td>P&lt;0.0001 NS clinically</td>
</tr>
<tr>
<td>Clark et al, 1990 (76)</td>
<td>Intubated but not all ventilated (n=189)</td>
<td>Open versus closed Prospective cohort 1. Open 2. Closed</td>
<td>4% fall in SvO2 with open; no change with closed</td>
<td>No change in SvO2 with open; no change with closed</td>
<td>P=0.0001 NS</td>
</tr>
<tr>
<td>Witmer et al, 1991 (77)</td>
<td>Ventilated patients in the ICU (n=25)</td>
<td>Open versus closed Randomized crossover 1. Open 2. Closed</td>
<td>No significant difference in mass of sputum</td>
<td>No significant difference in mass of sputum</td>
<td>P=0.88</td>
</tr>
<tr>
<td>Harris and Hyman, 1984 (78)</td>
<td>Newly tracheotomized patients (n=209)</td>
<td>Sterile versus clean Retrospective chart review 1. Sterile (five hospitals) 2. Clean (five hospitals)</td>
<td>Infection rate higher with sterile procedure (versus clean and mixed) based on tool developed by investigators</td>
<td>Significant difference in group 4, cannulation of left bronchi as detected by chest x-ray</td>
<td>Not stated</td>
</tr>
<tr>
<td>Freedman and Goodman, 1982 (79)</td>
<td>Intubated, ventilated (n=126)</td>
<td>Type of catheter Cohort 1. Straight 2. Coudé directed to left 3. Straight with traction of trachea to right 4. Coudé directed to left</td>
<td>X-ray of catheter position Straight: entered left 3/35 regardless of procedure Coudé: reached goal in 50% of cases</td>
<td>X-ray of catheter position</td>
<td>Not stated</td>
</tr>
<tr>
<td>Kirimi et al, 1970 (80)</td>
<td>Postoperative, intubated, ventilated (n=12)</td>
<td>Type of catheter Prospective crossover 1. Straight catheter 2. Coudé catheter</td>
<td>X-ray of catheter position</td>
<td>X-ray of catheter position</td>
<td>Not stated</td>
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<tr>
<td>Kubota et al, 1980 (81)</td>
<td>Postanesthetic with major surgery, intubated (n=104)</td>
<td>Type of catheter Prospective crossover 1. Portex straight and Coudé 2. Rubber catheter straight and Coudé</td>
<td>X-ray of catheter position  Portex Straight + neutral 10/52 (19%), rotation 17/52 (31%) Coudé + neutral 21/52 (40%), rotation 13/52 (25%) Rubber catheter Straight + neutral 8/52 (15%), rotation 15/52 (29%) Coudé + neutral 26/52 (50%), rotation 12/52 (23%)</td>
<td>No statistics</td>
<td></td>
</tr>
<tr>
<td>Jung and Gottlieb, 1976 (82)</td>
<td>Nonintubated with four tracheotomized (n=20)</td>
<td>Type of catheter Prospective crossover 1. Single eyelet 2. Two eyelet 3. Coudé 4. “Beaded”</td>
<td>Mucosal trauma with still and cine photography Negligible difference between groups 4 and 1 Gross traumatic effects – petechiae Sucking effect more pronounced with group 2 compared with 4 Little mucosal trauma with either catheter</td>
<td>No statistics</td>
<td></td>
</tr>
<tr>
<td>Valles et al, 1995 (83)</td>
<td>Intubated (n=190)</td>
<td>Catheter design RCT 1. Subglottic aspiration 2. Control</td>
<td>Decreased relative risk of pneumonia – relative risk of 1.76 compared with control</td>
<td>No difference in hospital mortality</td>
<td>NS</td>
</tr>
<tr>
<td>Mahul et al, 1992 (84)</td>
<td>Patients intubated for more than three days (n=145)</td>
<td>Catheter design ETT and use of a syringe RCT 1. Control 2. Subglottic drainage</td>
<td>Nosocomial pneumonia Control: 29% Group: 2: 13%</td>
<td>Time to onset of pneumonia: Control: 8.3 days Group: 2: 16.2 days</td>
<td>P&lt;0.05</td>
</tr>
<tr>
<td>Schmidt et al, 1995 (85)</td>
<td>Intubated, ventilated (n=12 for study A; n=28 for study B)</td>
<td>Catheter type/saline Randomized crossover Study A PaO&lt;sub&gt;2&lt;/sub&gt; – during 1. PaO&lt;sub&gt;2&lt;/sub&gt;: 6.6 mmHg fall 2. PaO&lt;sub&gt;2&lt;/sub&gt;: 21.1 mmHg fall but increased after suction Study B PaO&lt;sub&gt;2&lt;/sub&gt; – immediately after 1. 1.0 mmHg fall 2. 25.9 mmHg fall</td>
<td>No value</td>
<td>Secretion volume Study A 1. 11.0 mL 2. 0.84 mL Study B 1. 10.6 mL 2. 2.5 mL</td>
<td>P&lt;0.001</td>
</tr>
<tr>
<td>Isea et al, 1993 (86)</td>
<td>Ventilated patients who required frequent suctioning (n=20)</td>
<td>Catheter type/saline 1. Conventional catheter and 10 mL of saline 2. Irrigated catheter with 40 mL of saline</td>
<td>Sputum volume (dry weight) Conventional: 0.36 mL Irrigation catheter: 1.04 mL</td>
<td>No difference in O&lt;sub&gt;2&lt;/sub&gt; saturation between groups</td>
<td>NS</td>
</tr>
<tr>
<td>Doorley et al, 1990 (87)</td>
<td>Ventilated (n=24)</td>
<td>Type of catheter Cohort 1. Single lumen (Ballard) 2. Double lumen (Concord Sleri-catheter)</td>
<td>Effective secretion removal (subjective report from suctioned) – group 1 more effective</td>
<td>Fall in SpO&lt;sub&gt;2&lt;/sub&gt;85% (worse drop in group 2)</td>
<td>P&lt;0.05</td>
</tr>
<tr>
<td>Bostick and Wendelgass, 1987 (88)</td>
<td>Postoperative open heart surgery, ventilated (n=45)</td>
<td>Saline Randomized crossover 1. No saline 2. 5 mL saline 3. 10 mL saline</td>
<td>No significant difference in PaO&lt;sub&gt;2&lt;/sub&gt; between groups when controlled for pre-O&lt;sub&gt;2&lt;/sub&gt; levels</td>
<td>No difference in weight of sputum between groups</td>
<td>NS</td>
</tr>
<tr>
<td>Gray et al, 1990 (89)</td>
<td>Critically ill, ventilated (n=15)</td>
<td>Saline Randomized crossover 1. 5 mL saline 2. No saline</td>
<td>No difference in gas exchange between groups Statistically different amount of secretion removed</td>
<td>No difference in HR or BP between groups</td>
<td>NS</td>
</tr>
<tr>
<td>Ackerman and Gugerty, 1990 (90)</td>
<td>Ventilated (n=26)</td>
<td>Saline Randomized crossover 1. No saline 2. 5 mL saline</td>
<td>Fall in SaO&lt;sub&gt;2&lt;/sub&gt; in group 2</td>
<td>Significant difference in sputum weight between groups (greater in saline)</td>
<td>P&lt;0.05</td>
</tr>
</tbody>
</table>

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**Insufflation:** During endotracheal suctioning, insufflation catheters (usually double lumen catheters) permit oxygen-enriched air to be blown into the tracheobronchial tree.

Five randomized crossover trials were identified that dealt with the technique of insufflation as a method of oxygenation. Three of the studies involved cardiac surgery patients (two postoperative [46,49], one intraoperative [48]). The others involved a mixture of trauma, medical, surgical and neurological patients (50,51). Double lumen insufflation catheters were used in the studies. With the exception of one study (46), all studies employed commercially available insufflation catheters. In general, insufflation resulted in maintenance of oxygen levels throughout the procedure, at flow rates of 10 to 15 L/min.

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### TABLE 4 (continued)
**Summary of studies in adults related to open and closed systems, types and designs of catheters, instillation of saline, drugs and jet ventilation**

<table>
<thead>
<tr>
<th>Author, year (reference)</th>
<th>Population</th>
<th>Treatment allocation</th>
<th>Primary outcome</th>
<th>P</th>
<th>Secondary outcome (could also be primary)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ackerman, 1993 (91)</td>
<td>Critically ill males, ventilated (n=40)</td>
<td>Saline Randomized crossover In-line suction for both groups 1. 5 mL saline 2. No saline</td>
<td>Statistical analysis of SaO2 not interpretable</td>
<td></td>
<td>No difference in SaO2 between groups</td>
<td></td>
</tr>
<tr>
<td>Hudak and Bond-Domb, 1996 (92)</td>
<td>Postoperative major head and neck surgery with tracheotomy, nonventilated (n=20)</td>
<td>Saline Randomized crossover 1. No saline 2. 5 mL saline</td>
<td>Significant difference in sputum weight with saline</td>
<td>P&lt;0.02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Werba et al, 1993 (93)</td>
<td>Head injury or spontaneous subarachnoid hemorrhage, ventilated (n=18)</td>
<td>Drugs to reduce ICP RCT 1. Vecuronium (0.12 mg/kg) 2. Atropine (0.4 mg/kg)</td>
<td>No change in ICP for both drugs Significant fall in ICP without either drug</td>
<td>P&lt;0.05</td>
<td>Diaphragmatic movement seen in two patients in both drug groups</td>
<td>Not stated</td>
</tr>
<tr>
<td>White et al, 1984 (94)</td>
<td>Ventilated with brain injury with ICP 20 mmHg with suctioning (n=15)</td>
<td>Drugs to reduce ICP Randomized crossover 1. 2 mL normal saline (IV) 2. 1 mg/kg fentanyl (IV) 3. 3 mg/kg thiopental (IV) 4. 1.5 mg/kg lidocaine (IV) 5. 1.5 mg/kg lidocaine (intratracheal) 6. 1 mg/kg succinylcholine (IV)</td>
<td>Intratracheal lidocaine reduced peak ICP with suctioning but initially increased ICP Succinylcholine prevented increase in ICP with suctioning compared with saline No difference between IV lidocaine and control</td>
<td>P&lt;0.05*</td>
<td>Rise in MAP was similar in all groups</td>
<td>No value</td>
</tr>
<tr>
<td>Donelan and Bedford, 1980 (95)</td>
<td>Ventilated head injury, comatose (n=10)</td>
<td>Drugs to reduce ICP Randomized crossover 1. Lidoacaine 2% IV (1.5 mg/kg) 2. Saline placebo</td>
<td>Reduced ICP with lidocaine given presuctioning, but both had increased ICP post suctioning</td>
<td>P&lt;0.05</td>
<td>BP increased in both groups; no difference between 1 and 2</td>
<td>NS</td>
</tr>
<tr>
<td>Brown and Peeples, 1992 (96)</td>
<td>Ventilated with head injury (n=11)</td>
<td>Drugs to reduce ICP Randomized crossover 1. Hyperventilation 2. IV lidocaine (2%, 1.5 mg/kg, 50 mg/min) 3. Aerosolized lidocaine (4%, 5 mg/kg over 15 min)</td>
<td>No difference in ICP between groups</td>
<td>NS</td>
<td>No difference in CPP between groups</td>
<td>NS</td>
</tr>
<tr>
<td>Hanowell et al, 1993 (97)</td>
<td>Head injury (n=7)</td>
<td>Drugs to reduce ICP Randomized crossover 1. IV saline 2. Alfentanil 15 g/kg 3. Alfentanil 30 g/mg</td>
<td>No difference in CPP between groups Greater increase in MAP with combined alfentanil groups versus control</td>
<td>NS</td>
<td>Greater increase in MAP with combined alfentanil versus saline</td>
<td>NS</td>
</tr>
<tr>
<td>Winston et al, 1987 (98)</td>
<td>Pulmonary parenchymal disease and sepsis, ventilated (n=6)</td>
<td>Drugs for bradycardia Randomized crossover 1. Nebulized atropine sulphate (0.05 mg/kg of ideal body weight) 2. Parenteral atropine sulphate (1 mg by IM or slow IV) 3. Saline</td>
<td>Bradycardia prevented with use of either drug; tachycardia occurred in group 2 but rarely in group 1</td>
<td>P&lt;0.001</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Guntupalli et al, 1984 (99)</td>
<td>Ventilated with jet ventilator and jet tube (n=15)</td>
<td>Jet ventilation Prospective crossover 1. Suction with continuous ventilation 2. Stop ventilation during procedure</td>
<td>PaO2 decreased from 417 to 327 mmHg with stopping jet ventilation</td>
<td>P&lt;0.001</td>
<td>PaCO2 increased from 36 to 38 mmHg</td>
<td>P&lt;0.05</td>
</tr>
</tbody>
</table>

BP Blood pressure; bpm Beats/min; COPD Chronic obstructive pulmonary disease; CPP Cerebral perfusion pressure; ETT Endotracheal tube; FiO2 Fractional inspired concentration of oxygen; HR Heart rate; ICP Intracranial pressure; ICU Intensive care unit; IM Intramuscular; IV Intravenous; MAP Mean arterial pressure; NS Not significant; PaO2 Arterial pressure of oxygen; PEEP Positive end-expiratory pressure; RCT Randomized controlled trial; SaO2 Arterial oxygen saturation; SpO2 Oxygen saturation by pulse oximetry; TDPO2 Transcutaneous partial pressure of oxygen; TV Tidal volume
One study (54) showed no difference between the two methods, and four studies (59-62) favoured ventilator-delivered breaths to support oxygenation during suctioning. There was no difference noted between the two methods of oxygenation for arterial blood pressure in the one study that reported this outcome (54). In three of the studies (60-62), it was assumed (not confirmed by calibration) that the MRB actually delivered hyperinflations and an oxygen concentration of 100%. Only two studies (54,59) measured the two factors, and only one (54) controlled for these measures in the design of the trial. The reason that use of a ventilator was found to be superior to use of an MRB may be that the MRB is simply an inefficient method of breath delivery.

**Recommendation:** A ventilator is more effective for the delivery of oxygen than an MRB and should be used when oxygen delivery is a concern.

**Use of an adaptor:** Swivel adaptors with capped apertures have been an integral part of the patient ventilator circuit for many years. When opened, the aperture permits suctioning of intubated, ventilated patients without complete disconnection from the ventilator circuit. This may be advantageous in reducing the deleterious effects of endotracheal suctioning; however, suctioning through the adaptor with a conventional, disposable catheter may be technically more difficult.

Three randomized crossover trials dealt with the use of an adaptor in suctioning (40,63,64). One of the studies involved postoperative cardiac surgery patients (63), another acute lung disease or acute on chronic lung disease patients (64), and one acutely ill patients with respiratory failure (40). In two reports, the type of swivel adaptor used was not identified (40,63), while one study (64) noted the use of a suction adaptor by Novametrix Medical Systems Inc, USA.

Two studies (63,64) found that the use of an adaptor attenuated the change in oxygenation observed when patients are disconnected from the ventilator while being suctioned (without hyperoxygenation). The third study (40) found that suctioning through an adaptor was no different from disconnection from the ventilator with pre- and posthyperoxygenation.

**Recommendation:** To preserve oxygenation in mechanically ventilated medical and surgical patients, suctioning through an adaptor may be as effective as disconnecting the ventilator circuit and hyperoxygenating before and after suctioning. However, if the clinician does not have access to hyperoxygenation and/or hyperinflation, use of an adaptor is preferable to disconnection from the ventilator.

**Positive end expiratory pressure adaptor:** The application of positive end expiratory pressure (PEEP) to airways is a commonly used technique, particularly in acutely ventilated patients. PEEP is designed to increase functional residual capacity and enhance oxygen transport across the alveolar-capillary membrane, and is part of the management of condi-

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**Recommendation:** The use of oxygen insufflation (at 10 to 15 L/min flow) to maintain oxygenation is recommended for relatively stable patients who are mechanically ventilated.

**Hyperinflation alone:** As previously mentioned, while secretions are being removed from the tracheobronchial tree during the suctioning procedure, air is also being removed. Hypoxemia and atelectasis are among the possible harmful consequences. One manoeuvre designed to correct or reduce these effects is hyperinflation – ie, the provision of a breath or sequence of breaths at volumes greater than the usual tidal volume. Hyperinflations may be delivered with an MRB or a ventilator.

Five randomized crossover studies of hyperinflation alone involved postoperative cardiac surgery patients (52-56). The results of one study were published twice (55,56) and so were combined. Two randomized crossover studies involved head-injured populations (57,58). An MRB or a ventilator was used to deliver the hyperinflations in all five studies. In each case, the volumes delivered were measured and confirmed regularly by testing and calibration. Because the outcomes of interest were different, recommendations about hyperinflation for cardiac surgery patients were separated from those for head-injured patients.

The studies did not show a consistent beneficial effect of hyperinflation on oxygen levels in patients after cardiac surgery. An increase in blood pressure was observed to accompany hyperinflation. In patients with head injury, studies showed that an increase in volume (but not an increase in rate of inflation) may result in an increase in ICP. When suctioning these patients, caution should be exercised because ICP increases over time with additional suctioning sequences.

**Recommendation:** The use of hyperinflation delivered by any method is not recommended as a means to improve oxygenation in preoxygenated patients who undergo coronary artery bypass graft surgery. If hyperinflation is used in other populations (eg, unstable cardiac patients), caution should be employed because its use may be associated with increases in mean arterial blood pressure. Minute ventilation should be increased by rate but not volume to minimize increases in ICP in head-injured patients during suctioning.

**MRB versus ventilator delivery:** Several methods of delivering hyperoxygenation and/or hyperinflation to the patient during the suctioning procedure have been suggested. The MRB and ventilator are two such commonly used methods, particularly in intubated patients.

There were four randomized crossover trials (54,59-61) and one comparative cohort study (62). Three of the study populations involved postoperative cardiac surgery patients (54,60,61). The other studies (59,62) had a mixed group of medical, surgical and trauma patients. All of the trials examined patients who were oxygenated using an MRB connected to an oxygen supply (commonly set at a flow rate of 15 L/min) and a ventilator delivering various volumes and fractions of inspired oxygen (commonly 100%).

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tions such as adult respiratory distress syndrome (ARDS) and pulmonary edema. Disconnection from the ventilator for procedures such as endotracheal suctioning disrupts the positive airway pressure being exerted. Conventional manual bagging to provide hyperinflation and/or hyperoxygenation during the technique of suctioning does not sustain the end expiratory pressures maintained by the ventilator. A modified PEEP adaptor used in conjunction with a manual MRB system has been designed to permit the maintenance of PEEP while patients are manually bagged.

One randomized crossover study (65) and one non-randomized crossover study (66) involving use of a PEEP adaptor were identified. The first study involved a mixture of medical and surgical patients. The second focused on patients with ARDS. One study (65) found no difference in the use of a PEEP adaptor, while the other (66) reported a positive effect on PaO2. There is insufficient evidence to make a recommendation regarding the use of a PEEP adaptor in medical/surgical or ARDS patients.

**Open versus closed suctioning systems:** Suctioning patients on a ventilator generally requires either disconnection from the ventilator and use of a conventional disposable catheter (open), or use of a specialized closed-system suction catheter without removal from the ventilator (closed). In the closed system, the catheter is connected to a specially designed T piece attached to the airway. Possible benefits of the closed system include prevention of desaturation and decreased incidence of cross-contamination.

Eleven studies comparing closed with open suctioning were retrieved (67-77); three RCTs (67-69), three randomized crossover studies (72,73,77), one quasirandomized, controlled trial (74) and four comparative cohort studies (70,71,75,76). Studies included patients with respiratory failure, or patients following cardiac or vascular surgeries; some patients were studied during surgery.

Five studies examined the effect of open versus closed suctioning system on infection (67-71). Although some studies found a greater rate of colonization in patients with the closed suctioning system (67,70), there was no difference in the rates of nosocomial pneumonia (67-69) and mortality (67). The clinical relevance of an increase in the surrogate outcome of colonization rate is unclear. All studies lacked the power to detect a difference in the incidence of pneumonia. One study examined the effect of closed versus open system on environmental spread of infection and found lower colony counts (air) with a closed system (71). The clinical relevance of this finding is not clear.

Some studies found a statistically significant difference in levels of oxygenation (72,75,76), whereas others found no difference for open versus closed systems (73). When a statistically significant difference was found, the difference was not clinically important (72,75,76). One small study (74) found that the closed suctioning system attenuated desaturation in ventilated patients with PEEP with greater than 10 cm H2O.

Only one study (77) examined the effect of closed versus open systems on secretion removal. No difference was found in the amount of secretions removed.

There is insufficient evidence on which to base a recommendation regarding the effectiveness of closed or open systems on oxygenation, secretion removal or frequency of colonization for pneumonia.

**Clean versus sterile technique:** The controversy continues on whether suctioning should be performed with clean or sterile technique. Clean technique involves the use of clean gloves, whereas sterile technique involves the use of sterile gloves. Only one study (78) examined clean versus sterile technique. In a retrospective comparative cohort study (chart review), infection rates were reported to be higher in the sterile than in the clean group (78). These results are questionable because of methodological problems such as lack of standardization of techniques, lack of control of confounders and use of a nonvalidated outcome tool. There is insufficient evidence on which to base a recommendation concerning clean versus sterile technique.

**Straight versus Coudé catheter:** Directed cannulation of either mainstem bronchus is the purpose of using a Coudé (curved) tipped catheter, particularly when it is used in intubated or tracheotomized patients. The left mainstem bronchus is more difficult to enter due to its smaller diameter and more acute angle. Failure to adequately suction the left lower lobe may result in an increased incidence of left lower lobe secretions and atelectasis (79).

Four comparative cohort studies (79-82) examined the use of straight versus Coudé catheters. Two of the studies examined the use of these catheters with anesthetized, intubated patients at the end of surgery (80,81). One studied patients with tracheostomies who required fibreoptic bronchoscopy for diagnostic purposes (82), and another studied patients with either an ETT or a tracheostomy (79). The additional effect of head positioning was evaluated in two of these studies (80,81).

Directed cannulation of the left mainstem bronchus was more successful in patients with a tracheostomy and with the use of a Coudé catheter (79) than with a straight catheter. The use of a curved catheter with the head in midline was also effective in cannulating the left bronchus in intubated patients after surgery (81). The investigators in one study (80) found that the straight catheter entered the left main bronchus three of 35 times regardless of the intended bronchus. This reflects the difficulty of intubating the left mainstem bronchus selectively. The Coudé catheter was somewhat more successful in cannulating the left mainstem but reached its goal less than 50% of the time. In the one study that examined the effect of the catheters on mucosal injury (82), the authors used a subjective outcome measure that made the data of limited use. There is insufficient evidence to base a recommendation on the best catheter design for the prevention of mucosal injury.

**Recommendation:** A curved catheter rather than a straight catheter (in tracheotomized or intubated patients) is more successful in entering the left main bronchus; however, there is no evidence as to whether suctioning the left versus the right bronchus provides an advantage to patient outcome.
Brooks et al

**Catheter designs:** Double lumen ETTs have been developed to assist with drainage of subglottic secretions and, thus, to prevent ventilator-associated pneumonia in intubated patients. Similarly, double lumen suction catheters with the additional lumen for instillation of saline have been studied.

Two RCTs examined the effectiveness of a double lumen ETT (83,84) to assist with drainage of subglottic secretions. One study (83) used a design that allowed continuous aspiration of subglottic secretions (ie, below the glottis and above the cuff of the ETT) and found a decreased relative risk of pneumonia without a difference in hospital mortality. A second study (84) used a similar type of ETT but with hourly subglottic drainage via syringe and found a greater time to onset and lower incidence of nosocomial pneumonia in the group with the subglottic drainage. Further study is needed to assess the cost effectiveness of this technology and its use in high risk groups.

**Recommendation:** There is some evidence to support the use of modified ETT with capabilities for continuous or intermittent subglottic suctioning for the prevention of subglottic aspiration.

Three studies examined the effectiveness of double lumen catheters – two randomized crossover trials (85,86) and one cohort study (87). Although one study (85) found a difference in oxygenation and suctioned volume that favoured the double lumen catheter, interpretation of the results was difficult due to the apparent lack of data for parts of the analysis. Another study (86) also reported that the dry weight of the secretions collected was greater with the double lumen catheter. In contrast, one study (87) found that the single lumen catheter was better in terms of prevention of fall in SaO2, need to increase FiO2 and effective secretion removal. There is insufficient evidence to make a recommendation regarding the use of single versus double lumen catheters for the purpose of instillation.

**Instillation of saline:** Instillation of saline may loosen secretions, increase the amount of mucus plugs and secretions removed, elicit a strong cough and aid in removal of thick secretions.

Five studies (one RCT [88] and four randomized crossover studies [89-92]) assessed the effectiveness of the instillation of saline during suctioning on the recovery of secretions and on oxygenation. Four of the studies were conducted on ventilated patients (88-91) and one involved nonventilated patients with tracheostomies (92). All subjects had evidence of good gas exchange and no underlying hypoxia.

The studies compared instillation of 5 mL and/or 10 mL of saline with no saline (88-92). Most studies found no difference in oxygenation, gas exchange, heart rate or blood pressure, but an increased weight of returned secretions in the saline groups; however, the authors did not control for the added weight due to saline instillation. There is insufficient evidence on which to base recommendations regarding the use of the instillation of saline for improving oxygenation or secretion removal. There are no known detrimental effects associated with instillation. Further research is needed to examine the effectiveness of instillation in patients with marginal gas exchange; this research should control for the weight and amount of saline in the suctioned secretions.

**Drugs to prevent increased ICP or CPP during suctioning:** Increases in ICP as a result of suctioning may reduce cerebral perfusion and worsen ischemic insult in patients with brain injury. While sustained rises in ICP and decreases in CPP are potentially harmful, the impact of transient changes is unknown. There is no evidence to suggest that short term, acute increases in ICP or decreases in CPP during suctioning have a significant impact on neurological recovery. Various drugs are used to attenuate the rise in ICP.

One RCT (93) and four randomized crossover trials (94-97) were designed to assess the effectiveness of a variety of pharmacological agents on ICP and CPP in patients with acute head injury. The studies included patients who had severe brain injury with decreased levels of consciousness requiring ventilatory support. The interventions assessed were neuromuscular blocking agents (93,94), topical and parenteral anesthetic agents (94-96), barbiturates (94) and narcotics (94,97). One study assessed the effect of hyperventilation as a co-intervention (96).

The goal of neuromuscular blocking agents is muscle paralysis and, thus, the prevention of the strong muscle contractions associated with coughing during suctioning. Two studies (93,94) examined the effectiveness of neuromuscular blocking agents and found that vecuronium (0.12 mg/kg), atracurium (0.4 mg/kg) and succinylcholine (1 mg/kg) during suctioning prevented the rise in ICP.

Topical and parenteral anesthetic agents are used to depress the cough response and thus prevent the associated increase in ICP. Three studies (94-96) examined the effectiveness of topical and parenteral anesthetic agents (lidocaine) and found that intravenous lidocaine (1.5 mg/kg) had no effect on ICP rise during suctioning. Intratracheal administered lidocaine (1.5 mg/kg) was more effective in suppressing the cough reflex but triggered an initial rise in ICP due to coughing caused by instillation of the fluid.

It is hypothesized that the deep sedatory effect of barbiturates prevents a rise in ICP. Only one study (94) compared thiopental (3 mg/kg) with saline control. This drug did not prevent a rise in ICP associated with suctioning. There is insufficient evidence to recommend the use of thiopental (3 mg/kg) in head-injured patients.

Finally, a rise in ICP may be prevented through the sedatory effect of narcotics. Two studies examined the effectiveness of alfentanil and fentanyl (94,97). One study (97) found that alfentanil (15 μg/kg and 30 μg/kg) resulted in a decrease in CPP; in the other, fentanyl (1 μg/kg) did not result in a difference in peak ICP (94). There is insufficient evidence to recommend the use of narcotics in head-injured patients. Because alfentanil may have a detrimental effect on CPP, clinicians should use caution when administering this agent.
**Recommendations:** Intravenous vecuronium (0.12 mg/kg), atracurium (0.4 mg/kg) or succinylcholine (1 mg/kg) may be considered in head-injured patients to prevent an increase in ICP with suctioning. Intravenous lidocaine (1.5 mg/kg) does not appear to prevent a rise in ICP caused by coughing in head-injured patients. There may be a role for intratracheally administered lidocaine, but further investigation is needed.

**Drugs to minimize or decrease the likelihood of bradycardia during suctioning:** Suctioning can be complicated by bradycardia in some patients. Stimulation of vagal afferent fibres and hypoxemia has been hypothesized to be the cause of this response. A fall of greater than 20% in resting heart rate is considered significant bradycardia. Atropine sulphate is frequently used for prevention and treatment of bradycardia, and may be used in patients who exhibit this tendency during suctioning. Only one randomized crossover trial (98) assessed the ability of drugs to prevent bradycardia in ventilated patients with pulmonary parenchymal disease and sepsis that showed a 20% decrease in resting heart rate with suctioning. These investigators compared the effect of nebulized (0.05 mg/kg ideal body weight) and parenteral (1 mg by intramuscular injection or slow intravenous) atropine sulphate with a control of nebulized saline (0.05 mL/kg ideal body weight). Although bradycardia was prevented in all patients with either form of atropine sulphate (P<0.001), tachycardia occurred in all patients receiving the parenteral form. Bradycardia with suctioning is not common but may result in important clinical problems.

**Recommendation:** Nebulized atropine (0.05 mg/kg ideal body weight) may prevent bradycardia in patients exhibiting a decreased heart rate with suctioning.

**Jet ventilation and suctioning:** Jet ventilation or high frequency jet ventilation uses smaller tidal volumes and more rapid rates than conventional ventilation. The rates are generally defined as greater than 50 breaths/min and tidal volumes less than or equal to dead space volume.

Only one small nonrandomized study examined suctioning during jet ventilation (99). PaO2 decreased when jet ventilation was discontinued during suctioning (P<0.001), while the partial pressure of arterial carbon dioxide increased slightly. Maintaining jet ventilation during suctioning reportedly did not affect gas exchange to the same extent. There is insufficient evidence to make a recommendation on suctioning during jet ventilation.

**CONCLUSIONS**

We found no literature on several suctioning procedures. For example, there were no studies on the optimum route for suctioning when the patient is not intubated (nasal versus oral); use of airway (nasal or oral); catheter size; frequency of suctioning, suction pressure, duration of suction, landmark for applying suction (ie, when resistance is encountered or af-

**Practice guidelines for suctioning**

...ter the catheter has been withdrawn slightly); intermittent versus constant suction; rotation of catheter; and the use of lubrication (ie, water or gel). We did not develop evidence-based recommendations in these areas.

When examining the effectiveness of suctioning procedures, ideal outcome measures should reflect long term consequences, such as the effects of the suctioning technique on survival or the incidence of certain complications (eg, bacteremia or pneumothorax). There are inherent limitations to the use of these outcomes. For example, due to the low rate of most complications, a large sample size is needed to assess survival and the frequency of other complications. Another outcome is the effect of suctioning approaches on quality of life; however, it is difficult to measure quality of life in acutely ill patients. For these reasons, the majority of studies employed physiological outcomes that were impairment-based (such as arterial blood gases). These physiological measures may be considered surrogate measures for long term outcomes. Although a drop in oxygen saturation postsuctioning may result in higher mortality or frequency of complications, we are not aware of any studies that have examined the correlation between changes in short term physiological measures and long term outcomes. We are aware that all studies did not report on all outcomes that were measured. This was taken into account in the careful formulation of the recommendations. Because none of the studies included cost information, no information about costs is included in this guideline. The literature reviewed gave no indication of patient preferences; thus, this aspect of the guideline is also absent.

This CPG will guide interdisciplinary clinical practice when suctioning adults and children. As with any guideline, it is not designed to supplant clinical judgment in specific clinical circumstances. While there is a paucity of good quality evidence regarding many aspects of suctioning, we feel that such a synthesis of the research serves a number of purposes. First, it has focused attention on areas where there is, in fact, sufficient evidence to guide practice. Second, this effort has clearly identified many gaps in the literature, and should be used to guide future research on this topic. An update of this CPG is planned in five years. Implementation strategies have been developed to enhance the use of this guideline in practice and aid in translating the evidence into clinical practice.

**ACKNOWLEDGEMENTS:** Members of the Working Group express their appreciation to the following individuals for their contribution to this project: Kathy Badali, Catherine Barbeau, Corinne Berinstein, George Browman, Ian Graham, Toni Newman, Sherra Solway, Anna Tsang, and all the clinicians and experts who provided feedback on the evidence-based recommendations. The College of Nurses of Ontario and the College of Respiratory Therapists of Ontario participated with the College of Physiotherapists of Ontario to develop the CPGs. Staff representatives from the College of Physicians and Surgeons of Ontario consulted with respect to the guideline development process. The College of Physiotherapists of Ontario provided most of the financial support for this project. No funding was obtained from product manufacturers.
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