Secretion Management in the Mechanically Ventilated Patient

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Summary

Secretion management in the mechanically ventilated patient includes routine methods for maintaining mucociliary function, as well as techniques for secretion removal. Humidification, mobilization of the patient, and suctioning are all routine procedures for managing secretions in the ventilated patient. Early ambulation of the post-surgical patient and routine turning of the ventilated patient are common secretion-management techniques that have little supporting evidence of efficacy. Humidification is a standard of care and a requisite for secretion management. Both active and passive humidification can be used. The humidifier selected and the level of humidification required depend on the patient’s condition and the expected duration of intubation. In patients with thick, copious secretions, heated humidification is superior to a heat and moisture exchanger. Airway suctioning is the most important secretion removal technique. Open-circuit and closed-circuit suctioning have similar efficacy. Instilling saline prior to suctioning, to thin the secretions or stimulate a cough, is not supported by the literature. Adequate humidification and as-needed suctioning are the foundation of secretion management in the mechanically ventilated patient.

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Mr Branson presented a version of this paper at the 39th RESPIRATORY CARE Journal Conference, “Airway Clearance: Physiology, Pharmacology, Techniques, and Practice,” held April 21–23, 2007, in Cancún, Mexico.

The author reports no conflicts of interest related to the content of this paper.

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Interruption therapy for secretion removal includes techniques either to simulate a cough, to mechanically loosen secretions, or both. Patient positioning for secretion drainage is also widely used. Percussion and postural drainage have been widely employed for mechanically ventilated patients but have not been shown to reduce ventilator-associated pneumonia or atelectasis. Manual hyperinflation and insufflation-exsufflation, which attempt to improve secretion removal by simulating a cough, have been described in mechanically ventilated patients, but neither has been studied sufficiently to support routine use. Continuous lateral rotation with a specialized bed reduces atelectasis in some patients, but has not been shown to improve secretion removal. Intrapulmonary percussive ventilation combines percussion with hyperinflation and a simulated cough, but the evidence for intrapulmonary percussive ventilation in mechanically ventilated patients is insufficient to support routine use. Secretion management in the mechanically ventilated patient consists of appropriate humidification and as-needed airway suctioning. The technique selected should remedy the suspected etiology of the secretion retention (e.g., insufflation-exsufflation for impaired cough). Further research into secretion management in the mechanically ventilated patient is needed. Key words: secretion management, heated humidification, heat and moisture exchanger, percussion, postural drainage, insufflation, exsufflation, airway suctioning, closed-circuit suctioning, saline instillation, manual hyperinflation, silver coated endotracheal tube, Mucus Shaver, Mucus Slurper, intrapulmonary percussive ventilation. [Respir Care 2007;52(10):1328–1342. © 2007 Daedalus Enterprises]

**Introduction**

Secrecion management in the mechanically ventilated patient consists primarily of adequate humidification and airway suctioning. The role of mechanical aids to improve cough and mobilize secretions is poorly defined and lacks supporting evidence in the intensive care unit (ICU). Even commonly performed procedures such as percussion and postural drainage have little support from the literature. This paper will review the evidence for secretion-management techniques in the mechanically ventilated patient and introduce some new techniques that are early in development.

**Secretion Retention in the Ventilated Patient**

Mechanically ventilated patients are at risk for retained secretions from a myriad of causes. Endotracheal intubation disrupts the mucociliary escalator and predisposes the patient to infection, which increases the volume and tenacity of mucus. Relative immobility of the mechanically ventilated patient confined to bed can lead to atelectasis, impaired cough, and retained secretions. Upper-abdominal and thoracic surgery frequently lead to postoperative atelectasis, weak cough, and secretion retention. Muscle weakness associated with prolonged ICU stay may also contribute to secretion retention. Finally, fluid status, particularly fluid restriction in the mechanically ventilated patient, may contribute to thickened secretions. When any or all of these coexist, the problem is compounded.

**Secretion Clearance Techniques**

Management of secretions in the mechanically ventilated patient involves routine standard of care therapies such as humidification, suctioning, and mobilization. When these routine methods fail, intermittent therapy is employed, with a variety of techniques, to simulate a cough, loosen secretions, drain secretions via positioning, or a combination of the three. Common respiratory therapy techniques for mechanical loosening of secretions include manual percussion and high-frequency chest wall compression, with or without postural drainage. These techniques can be combined with hyperinflation, as is the case with intrapulmonary percussive ventilation. A simulated cough maneuver, with either manual hyperinflation or insufflation-exsufflation, may aid in expelling secretions in the patient with secretion retention and neuromuscular weakness. These will be addressed separately.

**Routine Secretion Management in the Ventilated Patient**

**Mobilization**

“Up and out of bed” is a common postoperative order aimed at preventing atelectasis, stimulating cough, and improving circulation. This common sense approach is widely practiced and uniformly supported, but has little scientific evidence. Recently, Browning and colleagues found that in the first 96 postoperative hours the duration and quality of “up time” was poor, averaging only 3 min
on postoperative day 1. They also showed no difference in pulmonary complications based on the duration of up time.9 Interestingly, greater duration of up time was associated with shorter stay. It remains to be proven if this is a cause-and-effect phenomenon or a “true, true, and unrelated” phenomena. It may simply be that patients who are improving tolerate more up time than their sicker counterparts. Another study suggested that early mobilization is equivalent to postoperative coughing and deep breathing,10 which raises the question of whether these treatments are equally effective or equally ineffective.11 Clearly, investigation of early postoperative ambulation is warranted.

### Humidification

Heating and humidifying the inspired gas is an established standard of care during mechanical ventilation,12,13 but the minimum requirements and optimum devices and settings for humidification are less clear. The ability of any device, regardless of operation, to prevent drying of secretions depends on delivered gas temperature and relative humidity.14,15 Absolute humidity is the amount of water vapor present in a gas. Relative humidity is the ratio of the absolute humidity to the maximum absolute humidity. Relative humidity is perhaps most important, as any deficit must be provided by the tracheobronchial tree.

Assessing the adequacy of humidification is difficult. A number of potential surrogates for comparing humidification techniques have been suggested, including secretion volume and consistency, incidence of endotracheal tube (ETT) occlusion, changes in ETT effective diameter and/or resistance, suctioning frequency, and requirement for normal saline instillation.14,16–18 Measurements of secretion volume are inherently flawed. Secretion volume may change with the number of suctioning attempts, patient position, use of aerosolized medications, and saline instillation. Excessive humidification may cause an increase in secretion volume, and insufficient humidification may result in a decrease in secretion volume because mucus becomes encrusted in the airways.16 In my experience, secretion volume measurements are poorly reproducible and often reflect the individual practice of the clinician rather than the condition of the patient, so secretion volume is not reliable for comparing secretion-management techniques.

### Heated Humidification Versus Heat and Moisture Exchanger for Humidification and Secretion Management

A comparison of the ability of heated humidifiers versus heat and moisture exchangers (HMEs) to optimize secretion management requires measurable and reproducible variables. Those most frequently described are ETT occlusion and changes in effective inner diameter of the ETT resulting from encrusted secretions. Narrowing of the ETT and ETT occlusion have been described with both heated humidifiers19–21 and HMEs,22–25.

During use of a heated humidifier and heated-wire circuit, ETT occlusion is associated with an increase in gas temperature from the humidifier chamber to the patient. As gas temperature rises, relative humidity falls. Gas entering the ETT at a humidity deficit absorbs moisture from secretions in the ETT and large airways.19 This problem can be avoided by maintaining a constant temperature from the chamber to the airway, and by using a connecting tube between the heated-wire circuit and the airway, which allows for cooling and a relative humidity of 100%. The problem of caregivers creating a large temperature difference between the chamber and the airway to reduce condensate in the heated-wire circuit, which increased the risk of ETT occlusion, undoubtedly led to introduction of heated humidifiers with no such clinician-set control.26

Occlusion of the ETT during HME use occurs secondary to poor HME performance, changes in ambient conditions, leaks, and patient conditions (eg, body temperature, minute ventilation, and fluid status).22–25,27,28 HME performance plays a large role, and hygroscopic HMEs clearly outperform hydrophobic HMEs.29–31 Even the most efficient HME allows a net loss of heat and moisture from the respiratory tract, so prolonged use is associated with a greater incidence of ETT occlusion. There is also evidence that HMEs are less effective in patients with chronic lung disease, although this is not well understood.32

Hess evaluated studies of HMEs and heated humidifiers with and without a heated-wire circuit, using a meta-analysis to determine the risk of ETT occlusion53 Figure 1 depicts the results of that analysis. The studies in that meta-analysis represent a cumulative total of over 1,000 patients and the analysis indicates that the risk of ETT occlusion is nearly 4 times greater with HME than with heated humidification. That finding argues against the use

<table>
<thead>
<tr>
<th>First Author</th>
<th>HME (N)</th>
<th>HH (N)</th>
<th>RR (95% CI)</th>
<th>RR (95% CI)</th>
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<td>1/81</td>
<td>7.15 (0.96, 53.18)</td>
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<tr>
<td>Martin</td>
<td>6/31</td>
<td>9/42</td>
<td>17.47 (1.02, 290.95)</td>
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<td>Mollot</td>
<td>4/50</td>
<td>2/26</td>
<td>1.73 (0.35, 8.71)</td>
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<tr>
<td>Rostain</td>
<td>9/55</td>
<td>0/61</td>
<td>21.94 (1.25, 353.20)</td>
<td></td>
</tr>
<tr>
<td>Dreyfuss</td>
<td>1/61</td>
<td>0/70</td>
<td>3.64 (0.14, 82.82)</td>
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<tr>
<td>Vitaline</td>
<td>3/6</td>
<td>1/7</td>
<td>2.62 (0.35, 19.88)</td>
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<tr>
<td>Kinton</td>
<td>0/140</td>
<td>3/140</td>
<td>0.14 (0.01, 2.74)</td>
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<tr>
<td>Humi</td>
<td>2/9</td>
<td>0/95</td>
<td>4.73 (0.23, 98.82)</td>
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</tr>
<tr>
<td>TOTAL</td>
<td>40/554</td>
<td>7/480</td>
<td>3.94 (1.92, 89.82)</td>
<td>p &lt; 0.001</td>
</tr>
</tbody>
</table>

Fig. 1. Meta-analysis of risk of endotracheal tube occlusion with a heat and moisture exchanger (HME) versus with a heated humidifier (HH). RR = relative risk. (Adapted from Reference 33.)
of HMEs in patients with retained secretions, and for limiting
HME use to < 5 days.

Another issue with HME use in patients with increased
secretions is the possibility of occlusion of the HME. This
has not been specifically studied, but frequent soiling of
the HME has been reported as a trigger for switching to
heated humidification. In our experience, HMEs are most
frequently occluded by blood or pulmonary edema fluid.
However, in the presence of copious sputum production an
HME can become completely or nearly completely oc-
cluded (Fig. 2).

A series of studies have compared the effects of humid-
dification devices on in vivo and in vitro ETT resistance,
inner diameter, and surface area. In an early study,
Villafane et al measured the effective inner diameter by
measuring flow and pressure at the proximal ETT and
threading a hollow catheter to the distal tip of the ETT.
This allowed measurement of the pressure drop across the
tube. They made these measurements in patients on a daily
basis. Three groups of patients were studied: group 1 used
a heated humidifier, group 2 used a hygroscopic HME,
and group 3 used a hydrophobic HME. ETT resistance
increased with duration of use in all 3 study groups, but the
hydrophobic HME group had the highest resistance.
Figure 3 shows these changes in 4 patients in that study.

Several authors have used acoustic reflectometry to eval-
uate changes in ETT inner diameter. Boqué et al measured
loss of effective inner diameter and found that within
48 hours > 60% of ETTs lost ≥ 10%. All patients used an
HME.

Shah and Kollef found similar losses of intraluminal sur-
facer area when they compared unused to used ETTs. The
most convincing evidence comes from Jaber et al, who used
acoustic reflectometry to compare ETT volume and resis-
tance over 10 days. Half the patients used heated humidifiers
and half used HMEs. At day 5 there was no difference in the
changes in ETT resistance. However, at day 10 the HME
group had a 19 ± 18% increase in resistance, compared to an
8 ± 12% increase in the heated humidification group (Fig. 4).
The authors concluded that ETT resistance increases with
duration of use and that HMEs increase ETT resistance more
than do heated humidifiers.

These findings are clinically important because several
authors have described ETT resistance as a cause of wean-
ing failure. The findings also reinforce the concept
that the selection of a humidification device should be
based on the expected duration of use and the presence of
thickened secretions. In the mechanically ventilated pa-
tient with secretion-management issues (thick and/or co-
pious sputum), heated humidification is preferred. When
mechanical ventilation is expected to last more than 96
hours, a heated humidifier should be used from the outset.

Maintaining the Endotracheal Tube Lumen

Humidification maintains mucociliary function and as-
sures that secretions remain hydrated so they can be ex-
pectatorated. In the mechanically ventilated patient, the ETT cuff abruptly stops the mucociliary escalator. At this point the second most important secretion-management technique, suctioning, becomes required. The suctioning procedure may include pre-oxygenation and/or hyperinflation, depending on the technique. Suctioning methods include open-circuit suction, closed-circuit suction, and minimally invasive (or shallow) suctioning. One contentious issue in suctioning is instillation of normal saline either to loosen secretions or to stimulate a cough as an aid to secretion mobilization. Studies are also now being done on when to suction. These issues are reviewed below.

**Suctioning**

Removal of tracheobronchial and upper-airway secretions to maintain airway patency and reduce the risk of silent aspiration is a standard of care.\(^42,43\) Suction catheters differ greatly in design but have the same general characteristics. Most adult catheters are 48–56 cm in length, to allow the catheter to reach the mainstem bronchi. The distal tip of the catheter has several openings for secretion removal, and the proximal portion contains a thumb port that the practitioner occludes to activate the suction. The distal tip is blunt, to avoid trauma to the mucosa or perforation of the tracheobronchial tree. The side holes in the distal tip of the catheter also serve to minimize the risk of local tissue damage. If the catheter had only one opening at the distal end, the mucosa could be drawn into the catheter tip and torn during withdrawal of the catheter. Suction catheters should be transparent to allow visual inspection of secretions, rigid enough to pass through the ETT, yet pliable enough to traverse the airways without damaging the mucosa.

<table>
<thead>
<tr>
<th>Catheter</th>
<th>Side hole diameter</th>
<th>Size</th>
<th>Whistle Design</th>
<th>Side Hole Design</th>
</tr>
</thead>
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<tr>
<td>Cardinal Tri-Flo</td>
<td>3 mm</td>
<td>14 Fr</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kendall Regu-vac</td>
<td>4 mm, 6.35 mm apart</td>
<td>14 Fr</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kendall Sensi-vac</td>
<td>5 mm, 9.52 mm apart</td>
<td>14 Fr</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Medline Delee</td>
<td>4 mm, 9.52 mm apart</td>
<td>14 Fr</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Medline Whistle Tip</td>
<td>4 mm</td>
<td>14 Fr</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Portex suction tray</td>
<td>4 mm</td>
<td>14 Fr</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Few comparative evaluations of suction catheter designs have been accomplished.\(^44,45\) Shah et al compared six 14 French suction catheters in a bench study that evaluated the characteristics (side-hole placement) that facilitated the removal of simulated mucus.\(^45\) The viscosity of the simulated mucus was altered to represent thin and thick secretions. Shah et al found that the major factors that affect secretion removal are the position and size of the catheter side holes. Offset side holes were associated with better mucus clearance. The catheters they tested are depicted in Figure 5. These findings are important for future catheter designs.

**Open Versus Closed Suctioning.** Two decades ago the standard of care for suctioning was a single-use, disposable, open-circuit suction catheter. The patient was disconnected from the ventilator, hyperventilated, and hyperoxygenated with a self-inflating manual resuscitator. The patient was then disconnected from the manual resuscitator and the suction catheter was passed into the ETT to remove secretions. The manual resuscitator was also used to simulate a cough, by stacking breaths or giving large volumes. This procedure was also known to result in both hemodynamic instability and hypoxemia.

In the last decade, closed-circuit suction catheters have become popular for a number of reasons, including prevention of problems associated with disconnecting the patient from the ventilator, reduced cost, and reduced exposure of caregivers to infectious materials. Comparisons of closed-circuit and open-circuit suctioning techniques suggest that there is no difference in their ability to evacuate secretions.\(^46,47\) Because the patient does not need to be disconnected from the ventilator during closed-circuit suctioning, the positive end-expiratory pressure (PEEP) is
maintained and hypoxemia and ventilator malfunction may be reduced. There is also evidence that the closed-circuit system reduces caregiver and environmental contamination, although the evidence regarding this is weak. Studies of prolonged use of closed-circuit suction catheters suggest that, compared to open-circuit suctioning, the incidence of ventilator-associated pneumonia is either lower or unchanged. Interestingly, many clinicians have suggested that, by preventing disconnection of the circuit from the ETT, closed-circuit suctioning reduces the risk of ventilator-associated pneumonia. Although this is an attractive hypothesis, it has not been studied. When closed-circuit suctioning was introduced, it was my (RDB) opinion that closed-circuit suctioning removed less secretions than open-circuit suctioning, and that eliminating the manual resuscitator not only prevented the clinician from “feeling” the compliance, but removed the ability to create a cough to propel mucus cephalad. I (RDB) was often told that the sound of closed-circuit suctioning is different because the airway is closed. The satisfying sound of secretions being removed with open-circuit suctioning was also lost. However, if PEEP is preserved during closed-circuit suctioning, it means that while we are trying to suction secretions out, the ventilator is blowing them back into the patient. So closed-circuit suctioning, despite its advantages, may be an inferior secretion-removal device.

This hypothesis was recently tested by Lasocki and colleagues, who compared the effects of open and closed suctioning in patients with respiratory failure. They found that closed suctioning prevented suction-related hypoxemia, but that secretion removal was reduced. They suggested that during open suctioning the disconnection from the ventilator and loss of PEEP simulates a cough and propels mucus upward. They also postulated that the disconnection from the ventilator during open-circuit suctioning increases the pressure differential, which enhances suctioning, whereas during closed-circuit suctioning the ventilator gas delivery that maintains the PEEP forces secretions away from the suction catheter. In that study, secretion volume was reduced with closed-circuit suctioning unless the suction pressure was increased to −400 mm Hg, which restored secretion removal volume (Fig. 6). However, this greater negative pressure was not associated with hypoxemia. They suggested the use of a post-suctioning recruitment maneuver to restore alveolar recruitment. Despite these findings, closed-circuit suctioning appears to have more advantages than disadvantages. In patients with retained secretions, increasing the vacuum pressure may be required to improve secretion removal, but gas exchange and ventilator performance should be monitored closely.

**Bronchial Suctioning.** During routine endotracheal suctioning, the suction catheter most likely enters the right main bronchus if the catheter is advanced far enough, because the angle from the trachea into the left main bronchus is more acute than that into the right main bronchus. As such, the left lung is less likely to be suctioned. Attempts at suctioning the left mainstem have been described and have ranged from simple maneuvers to the use of special catheters. One simple way of suctioning the left main bronchus is by turning the head to the right to increase the likelihood of the catheter entering the left mainstem. The same effect may be gained by placing the patient in the left lateral position and attempting to use gravity to guide the catheter to the left lung.

Specialized catheters that have a curved tip enter the left main bronchus in up to 90% of cases. The success of bronchial suctioning can be affected by tube position, patient body and head position, and type of tube (ETT vs tracheostomy tube). We have not found selective endobronchial suctioning to be a necessary routine technique. Frequent changes in patient body position facilitate movement of secretions to the carina, where they can be suctioned. In patients with infectious processes confined to the left lung, selective endobronchial suctioning may prove useful.

**Deep Versus Shallow Suctioning.** Prior to suctioning a neonate, the length of the suction catheter is often measured to prevent traversing the tip of the ETT and thus to prevent trauma to the neonatal tracheobronchial mucosa. Shallow (or minimally invasive) suctioning means passing the suction catheter tip to the end of the ETT, but no farther. Suctioning past the ETT tip is therefore called deep suctioning. In adults this issue is less frequently discussed. A recent meta-analysis of this topic found that the supporting literature is poor and that no definitive conclusions could be made. Ahn and Hwang examined secretions from neonates following deep and shallow suctioning and found evidence of detached ciliated airway cells with deep suctioning, but no more secretions were removed than with shallow suctioning. They suggested...
that deep suctioning has no advantage, causes unnecessary trauma, and should be avoided.

In adults, van de Leur et al found that minimally invasive suctioning resulted in fewer recollections of suctioning by ventilated patients. However, this was not associated with less discomfort with the suctioning procedure. In a large study of adult patients, they also found that minimally invasive suctioning resulted in fewer hemodynamic and gas-exchange adverse effects but had no effect on duration of ventilation and other outcomes. In the latter study the traditional suction catheter was 49 cm long, compared to 29 cm for the minimally invasive catheter.

This issue essentially pits deep suctioning (to remove the most secretions) against shallow suctioning (just keeping the ETT clear of secretions). The limited evidence seems to suggest that minimally invasive suctioning is as effective at secretion removal, but has fewer adverse effects, so the “first, do no harm” principle suggests that minimally invasive suctioning should be preferred, but further study is necessary.

**Saline Instillation.** During the suctioning procedure it is common for some practitioners to instill 5–10 mL of normal saline in an attempt to thin the tracheobronchial secretions. This practice remains a point of contention, and studies have failed to show any advantage from saline instillation. Our studies of humidification techniques revealed that the only correlation between saline instillation and patient care is practitioner preference. That is, our research failed to show that saline instillation is used uniformly or has any benefit in terms of liquefying secretions. Saline instillation frequently does cause the patient to cough violently, which may aid in the secretion-removal process. From a conceptual standpoint, instilling saline to stimulate a cough makes sense, but the current literature does not support the routine use of saline instillation. From a mucus rheology perspective, the properties of mucus are unlikely to change with the addition of water unless some physical means of mixing the two is accomplished. As well, severe coughing episodes and bronchospasm occasionally may result from saline instillation.

There is also some concern that the use of saline may dislodge bacteria-laden biofilm from the ETT and into the small airways. A host of recent studies found that saline instillation failed to produce any of the intended effects, while potentially causing more pulmonary infections. Based on this evidence, saline instillation to thin secretions is, at best, unsupported and, at worst, dangerous.

**When to Suction.** Routine suctioning should be avoided. Instead, patient assessment, including auscultation and visual inspection, should be used to determine the need for suctioning. In recent years, ventilator graphics have been used to detect the need for endotracheal suctioning. Jubran and Tobin, Guglielminotti et al, and Zamanian and Marini all described alterations in the pressure and flow curves that might suggest the need for suctioning. Figure 7 depicts changes in the expiratory flow signal that suggest airway secretions. The most common finding is a sawtooth pattern in the expiratory flow signal, caused by secretions in the large airways. This finding is also seen with condensate in the expiratory limb of the ventilator circuit, so visual inspection of the circuit should be included in the evaluation.

Visaria and Westenskow demonstrated the ability to detect ETT occlusion using an automated evaluation of pressure and flow signals. They were able to distinguish airway obstruction from bronchospasm and changes in chest wall compliance. Similar systems might be developed to determine when suctioning is required.

Endotracheal suctioning is associated with many complications, and should be undertaken only when necessary, keeping the potential complications in mind. Minimizing these complications by minimally invasive suctioning and suctioning only when necessary, based on reliable detection methods, may both be routine in the future.

**Novel Methods for Secretion Removal**

**From the Endotracheal Tube**

**The Mucus Slurper**

Intermittent closed-circuit suctioning is the current standard of care in mechanically ventilated patients. Kolobow and colleagues recently described a system (the “Mucus Slurper”) that provides automated, intermittent suctioning of the ETT lumen (Fig. 8). The Mucus Slurper is a...
modified continuous subglottic suctioning ETT. The end of the ETT is modified by cutting off the portion beyond the cuff, and attaching a plastic ring with eight 1.3-mm holes. The suction lumen is extended to apply suction to these 8 holes.

In Figure 8, the pressure and flow waveforms show the effects of activating suction to the 8 holes at the end of inspiration. The system draws 135 mL over 0.3 seconds. In this preliminary evaluation in an animal model, the system did not affect ventilator performance. There was some concern about auto-triggering, but most ventilators have a “lockout” period following completion of the inspiratory time, which is near the 0.3-second time frame. Figure 9 shows the system for control of suction pressure and timing.

This very preliminary evidence indicates that in an animal model without pulmonary disease, the ETT lumen remains clean. Presumably, as the secretions approach the ETT, they are removed through the 8 narrow lumens. Recently, the Mucus Slurper system was tested for 72 hours in an animal model that compared every-2-min activation with the mucus slurper to every-6-hour suctioning. There were no differences in airway colonization between the groups. However, inspection of the ETT showed less secretion accumulation in the Mucus Slurper group. Kolobow and colleagues measured protein content in the expiratory condensate as a marker of secretion movement up the ETT. The protein concentrations were lower in the Mucus Slurper group. How this system will perform in a patient with copious, tenacious secretions remains to be seen.

The Mucus Shaver

Kolobow and colleagues also described the Mucus Shaver system for removing secretions from the inner wall of the ETT. The Mucus Shaver is a manually operated system that scrapes the inside of the ETT to remove secretions (Fig. 10). The Mucus Shaver is placed inside the ETT, much like a stylet. The balloon is inflated and the shaving heads are forced against the inner wall; the device is then withdrawn over 3–5 seconds to remove secretions. The intention is to return the ETT resistance characteristics to their pre-use values. In an animal model with normal lungs, the Mucus Shaver maintained a clean internal
lumen and reduced biofilm accumulation. The clinical role for this device remains uncertain.

Special Endotracheal Tubes

ETTs are designed for a number of special situations. There are wire-reinforced ETTs, ETTs for laser surgery, specially shaped ETTs to improve the operative field, and ETTs for high-frequency ventilation. Silver-coated and silver-impregnated ETTs are designed to reduce bacterial colonization and maintain a patent lumen.

Silver has long been appreciated for its bacteriostatic properties. Older clinicians will remember the time when all tracheostomy tubes were silver or silver-plated stainless steel. Silver-coated and silver-impregnated urinary catheters and central venous catheters have been used for over a decade. Silver has other medically useful properties, including prevention of biofilm formation, reduction in bacterial burden, and reduction in inflammation. To test the potential bacterial burden reduction in the respiratory tract with silver-coated ETTs, Olson et al performed an experimental study in 11 ventilated dogs. The silver-coated ETTs reduced biofilm formation and there was a significant lumen-narrowing difference between the ETTs. Five of the 6 noncoated ETTs (83%) and none of the 5 coated tubes had a narrowing of > 50%. The coated tubes not only reduced the bacterial burden, with a statistically minor risk of colonization, but also delayed the formation of luminal side colonization from 1.8 ± 0.4 d to 3.2 ± 0.8 d.

A recent prospective randomized phase II pilot study tested silver-coated ETTs in ICU patients. The main objective was to determine whether a silver-coated ETT reduced the incidence and/or delayed the onset of coloniza-

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**Fig. 9.** Aspirated mucus is collected in a 25-mL mucus trap connected to the external suction line of the Mucus Slurper. A water trap prevents accumulation of water in the solenoid valve. The controller is connected via a pressure transducer to the respiratory circuit, and synchronizes the opening of the solenoid valve during the early expiratory phase. When the valve opens, the circuit is connected to a vacuum source (suction level is adjustable with a vacuum controller). The frequency of activation and the duration of suction are adjustable via an electronic controller. An alarm activates when no synchronized aspiration is detected. (From Reference 94, with permission.)
tion, compared to a noncoated ETT, in mechanically ventilated patients. There was a significant reduction in microbiologic burden with the silver-coated ETT. The high bacterial concentration on the inner surface of a standard ETT can play a role in the development of late-onset ventilator-associated pneumonia when biofilm fragmentation occurs. This fragmentation can be facilitated by airway suctioning or installation of saline. The future role of silver-coated or silver-impregnated ETTs remains to be determined.

Intermittent Techniques to Enhance Secretion Removal

The term respiratory therapy often refers to “treatments” provided by the respiratory therapist to aid in lung expansion, prevent atelectasis, and mobilize retained secretions. Within the realm of respiratory therapy for secretion mobilization are techniques that simulate a cough, that loosen mucus through external forces (chest clapping, vibration), and positioning. In many cases these methods combine two or three of these techniques. Each will be considered.

Techniques to Simulate a Cough

Manual Hyperinflation. Manual hyperinflation during suctioning was at one time a common procedure. By stacking small breaths or providing a large breath and holding it, the respiratory therapist could simulate and/or stimulate a cough. This is separate from the technique of manual hyperinflation aimed at lung recruitment.

Manual hyperinflation with a self-inflating bag for secretion removal is a popular practice in the United Kingdom and some former United Kingdom colonies in Asia. Recent studies indicate that the mechanical ventilator can also be used to achieve hyperinflation, with similar results.

As a separate technique, manual hyperinflation is not commonly used for secretion removal in the mechanically ventilated patient in the United States. Manual hyperinflation delivers a large-tidal-volume breath over a prolonged inspiratory time, followed by an inspiratory hold and rapid release of pressure. The goal is to simulate a cough and propel mucus cephalad. Most studies have suggested that as peak expiratory flow is increased, secretion removal is enhanced. The evidence for the efficacy of manual hyperinflation is scant. Several studies have shown improved compliance and oxygenation after manual hyperinflation, but the findings about changes in secretion volume have been inconsistent.

Manual hyperinflation is not without risk. High airway pressure and large lung volumes produce adverse hemodynamic effects and can injure the lung via barotrauma and/or volutrauma.

Insufflation-Exsufflation. Mechanical insufflation-exsufflation (in-exsufflation) is done with the CoughAssist In-Exsufflator (Respironics, Murrysville, Pennsylvania), which is a stand-alone device that insufflates a fairly large volume of gas into the lungs, and then rapidly reverses to a negative pressure that exsufflates the gas from the lungs, which simulates a cough and propels secretions cephalad. The active expiratory portion of the CoughAssist In-Exsufflator’s operation separates it from manual hyperinflation.

To date, the CoughAssist In-Exsufflator has primarily been used in patients with neuromuscular weakness who are on noninvasive ventilation. There are no clinical trials of the CoughAssist In-Exsufflator in mechanically ventilated patients in the ICU. In our experience, we have used the CoughAssist In-Exsufflator in the neurosurgical ICU, in patients with head and spinal cord injuries and

Fig. 10. A: Mucus Shaver. B: Mucus Shaver inflated. C: Mucus Shaver inflated inside an endotracheal tube. Shaded areas = silicone rubber. (From Reference 96, with permission.)
who retained secretions. These have primarily been tra-
cheostomized patients who cannot cough because of mus-
cle weakness or altered mental status. The intermittent use 
of the CoughAssist In-Exsufflator appears to assist in se-
cretion removal. Studies of in-exsufflation in these pa-
tients are warranted.

External Application of Force to Loosen Secretions

These techniques include percussion and postural drain-
age, high-frequency chest wall vibration, rib-cage com-
pression, and chest physiotherapy. Each of these therapies 
uses patient positioning and postural drainage to facilitate 
secretion removal, so these will be considered together.

Percussion and Postural Drainage. Though percussion 
and postural drainage are widely used in patients with 
cystic fibrosis in the out-patient setting, use of percussion 
and postural drainage in mechanically ventilated patients is 
sporadic. Conceptually, the mechanisms are similar. In 
the mechanically ventilated patient with retained secre-
tions, chest clapping is supposed to mechanically loosen 
secretions. It is interesting to note that percussion and 
postural drainage are often used to treat or prevent atelec-
tasis in the absence of secretion retention. There is no 
known mechanism by which percussion and postural drain-
age can improve atelectasis, unless it is by removal of 
mucus plugs.

The literature on percussion and postural drainage in 
mechanically ventilated patients is poor. In fact, more pa-
pers have described complications from percussion and 
postural drainage (pain, anxiety, atelectasis, elevated ox-
gen consumption) than have found any positive ef-
fects. At present, in mechanically ventilated pa-
tients the use of percussion and postural drainage, or 
chest physiotherapy, or other external vibration meth-
ods is unfounded.

High-Frequency Chest Wall Compression. This topic 
was covered by Chatburn in his contribution to this con-
ference. Suffice it to say that at present there is no 
evidence for the use of chest wall compression in the 
mechanically ventilated patient. One case report described 
removal of bronchial casts with high-frequency chest wall compression but it has not been systematically tested in mechanically ventilated patients.

Manual Rib-Cage Compression

Manual rib-cage compression (also known as “squeez-
ing”) refers to manually applying external force to the rib 
cage to increase expiratory flow and thus facilitate secre-
tion removal. The literature is confined to a couple of 
reports from Japan, which found no increase in secretion 
removal with squeezing. As with various other tech-
niques, anecdotal reports of success predominate.

Kinetic Therapy. The literature is replete with descrip-
tions of the use of rotating beds designed to prevent ven-
tilator-associated pneumonia and treat atelectasis. 
However, study of these beds’ effect on secretion clear-
ance has been limited. Davis et al compared secretion 
clearance, lung mechanics, and gas exchange in paralyzed 
patients with acute respiratory distress syndrome random-
ized to receive manual turning, manual turning with per-
cussion and postural drainage, continuous lateral rotation, 
or continuous lateral rotation with percussion and postural 
drainage performed by the pneumatic cushions of the bed.
Each of 20 patients received each treatment, in a random-
ized order, for 6 hours. The combination of continuous 
lateral rotation and percussion and postural drainage pro-
duced more secretions in this small group of paralyzed 
patients, but no other benefits were identified. As with 
most of the intermittent techniques discussed here, contin-
uous lateral rotation seems to make common sense for 
secretion management, but there is no evidence to support 
its efficacy.

Intrapulmonary Percussive Ventilation

Chatburn’s contribution to this conference also dis-
cussed intrapulmonary percussive ventilation, which com-
bines the secretion-loosening effects of mechanical per-
cussion with hyperinflation to facilitate cough and propel 
mucus cephalad. Study of intrapulmonary percussive ven-
tilation in mechanically ventilated patients has been lim-
ited to a few case reports.

Summary

Management of secretions in the mechanically venti-
lated patient requires adequate humidification and appro-
priate suctioning. The level of humidification required has 
been not well defined, but it is clear that in a patient with 
 thick and copious secretions a heated humidifier is pre-
ferred to an HME. A heated humidifier should also be used 
for prolonged ventilation (> 5 d). Suctioning should be 
done when clinical signs (auscultation, visible secretions 
in the tube, graphic displays) suggest secretions in the 
large airways. Closed-circuit suctioning has some advan-
tages over open-circuit suctioning with regard to prevent-
ing de-recruitment, but neither technique is superior for 
removing secretions. Some interesting new suctioning 
methods have been described, and future studies should 
determine if these have merit.

The use of intermittent secretion-removal techniques, 
including percussion and postural drainage, manual hyper-
inflation, insufflation-exsufflation, and intrapulmonary per-
cussive ventilation, have a dearth of supporting evidence and should be used cautiously. Routine use of these techniques is not supported by the literature.

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Discussion

**MacIntyre:** That was terrific, as always. I have a bunch of questions, let me just start. There’s been discussion about the ventilator as a mucous clearance tool, specifically the use of high-frequency ventilation, and Kolobow has been very keen on tracheal-gas-insufflation-type techniques.¹ What’s your take on using HFO [high-frequency oscillation] or tracheal gas insufflation as a mucous clearance?


**Branson:** Well, I think they’re separate. If you remember, when Miroslav Klain first came out using high-frequency jet ventilation,¹ one of the claims was you could ventilate the patient with the ETT cuff not inflated and it would prevent aspiration; and along with that was the idea that if there was mucus in the airway you would propel it forward. I think either of those techniques, high-frequency or TGI [tracheal gas insufflation] to remove secretions, are really dependent on how well you humidify that gas. Because if you blow a high flow of gas in tracheal gas insufflation into the ETT with the thought of propelling mucus forward, like with the reverse thrust catheter that Kolobow developed, and it’s not adequately humidified, you will make secretion problems worse. I don’t know that high-frequency oscillation is any different than IPV [intrapulmonary percussion ventilation] or anything else that we do in terms of secretion clearance via “shaking”—the question is, does that shaking actually eliminate or loosen secretions? I have to tell you, when you do percussion postural drainage with the Hill-Rom bed, it can create a lot of pressure, and we actually had a pa—
tient in a study who had a pulmonary artery catheter who could not tolerate the percussion, because whenever he got the percussion the catheter would whip against his heart and would cause him to go into runs of ventricular fibrillation. So it does actually create quite a bit of percussion into the airway. I don’t know if that answers your first question or not.


MacIntyre: I have just been intrigued with Ted Kolobow’s ideas over the years, and he’s come up with a lot of them. He has these ETTs with gills on them that don’t really seal; and, as you say, he argues that you can ventilate people without cuffs, which he believes facilitates movement of the mucus out of the airway. It is an interesting area of research, but probably not ready for prime time yet.


Chatburn: Rich, can you say anything about the relationship between the loss of effective ETT diameter and any kind of observable clinical effect on patients’ work of breathing?

Branson: There are a couple of papers that look at the changes in the work of breathing associated with these changes in ETT diameter. We had our lesson in physics yesterday, so anything that decreases the diameter is going to increase the work on the patient. There are a couple of papers, most of them from Orlando Kerton1–5 that suggest that there are patients who fail spontaneous breathing trials because their ETTs have become narrowed by retained secretions, and that there is an iatrogenic reason for weaning failure.


Rubin: Rich, that was a brilliant summary. I would like to continue on a rant that you started, vilifying the addition of saline, not only in terms of the extra time. What interests me, and what I think should require severe corporal punishment is: if saline really is effective in loosening up secretions so they can be suctioned, I can’t understand why so many people put down the saline and then bag the secretions and the saline way down deep in the lungs, well beyond the reach of any suction catheter. I’ve never been able to understand why people would follow saline instillation with bagging. Is this done at other people’s institutions?

Branson: All the time. I think the normal way that is done is that somebody puts saline down, the patient starts to cough, they bag them real good, and then try to give them a big breath so that they kind of simulate a cough and then suction it out. I’ve actually talked to Dean Hess about this before. I know that we don’t want to lose PEEP in recruitment in the mechanically ventilated patient, but I think when we went to closed-circuit suctioning, we lost the old way where we bagged the patients and did a simulated cough that, I thought, seemed to help clear secretions. I realize there are other reasons, ventilator-induced lung injury, not to give big volumes, but I always thought that—in some patients even now I will take them off the ventilator and try to give them a simulated cough. But I agree with you; we probably ought to change the policy to be, put the saline in and as soon as the patient starts to cough, suction, and then if you want to do some other technique for hyperventilation or manual inflations, you should do it.

Schechter: Studies of neonates and children on mechanical ventilation failed to provide support for the routine use of airway-clearance therapies for the prevention of atelectasis. However, there is some suggestion that it helps to promote resolution of atelectasis when it develops in these children. Is there any adult literature on that?

make it open. And I don’t know, and not to put Dave [Pierson] on the spot, but Dave actually has a case report that’s about 20-something years old now, where they did chest PT to remove a foreign body.2 So there are things like that. I will ask the pulmonologists—I thought I would for sure get into the literature and find that a therapeutic bronchoscopy was a really good evidence-based technique for removing secretions; and it’s not! There’s not a single study that suggests that a clean-out procedure with bronchoscopy helps, and we do this all the time! Go in to do a bronchoscopy for a bronchoalveolar lavage for culture, and while we’re in there we see that the lung is a mess, and so whoever the ICU person is spends the next 20 minutes cleaning and suctioning mucus; and we all feel better about that when we’re done. But there is no evidence that I can find that that does anything.


Pierson:* At Harborview we did a study in the late 1970s, when John Marini was a fellow,1 in which we prospectively randomized patients with acute lobar atelectasis detected on a chest x-ray to receive vigorous chest PT, which included postural drainage, bronchodilator inhalation, and clapping—with and without immediate bronchoscopy to try to suck out the “mucus plug” and relieve the atelectasis. First of all we found that to do the study we had to take a chest x-ray after randomization and immediately before we did anything, because so often the atelectasis just went away on its own, which obviously would have skewed the results. But in that relatively small and probably underpowered study, we were unable to detect any difference in the rate of resolution of the atelectasis when bronchoscopy was added to chest PT.

We embarked on a subsequent study in which we tried to do chest PT with and without the clapping, in exactly that same setting, and unfortunately that wound up getting abandoned about two thirds of the way through for poor recruitment of patients. And that’s, I think, where the literature stands on this. You are certainly correct that bronchoscopy is very often performed in patients who have perceived secretions, perceived atelectasis, radiographic abnormalities that might be atelectasis, or fever or hypoxia that could be due to atelectasis, but that is not an evidence-based practice.


Rogers: What is the mechanism of action whereby silver inhibits bacterial biofilm in bacterial formation in the tubes?

Branson: I don’t know if I can answer that fully. There are some good studies in the engineering and biomaterials literature1–3 that silver prevents bacterial growth and adhesion to surfaces—to this day we use silver-coated Foley catheters for patients who have need to be chronically catheterized. We used silver-coated central venous catheters because the silver prevents infection—it’s not bactericidal, it’s bacteriostatic. Again, from the historical standpoint the interesting thing was that in the early part of our country’s history, prostate problems were as common as they probably are today. Supposedly Ben Franklin had a silver straight catheter that he used on himself to empty his bladder and preventative infection. So the idea that silver is somehow bacteriostatic has been around for 200 or more years.


Rogers: When you say bacteriostatic, what does that mean compared with bactericidal?

Branson: A bactericidal actively kills bacteria, whereas a bacteriostatic prevents further growth. Like I said, there are some interesting studies in the literature about silver-coated tubes and preventing the adhesion to the surface of the ETT. Bruce, I don’t know if you can understand that or explain that better from an engineering perspective?

Rubin: No, that was all absolutely correct. That is exactly it. Ionized silver apparently prevents the growth of bacteria in biofilms. It’s not very good at killing them, but it doesn’t allow new growth and development in planktonic forms floating off and doing more bad stuff. But that’s exactly what it does. Just as you described.

Branson: All of you have been around as long or longer than I have, and when Dave was talking about that paper in the 1970s, I said, “Well I missed that, because I was in the 8th grade.” But when I first started in respiratory therapy, all the heated humidifiers for ventilators had copper wool in them. So it looked like a Brillo pad, only it was all copper, and the idea was that the copper would prevent the growth of any bacteria inside the humidifier. So these ideas that met-

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* David J Pierson MD FAARC, Division of Pulmonary and Critical Care Medicine, Harborview Medical Center, University of Washington, Seattle, Washington.
als somehow can be, I guess, helpful, aren’t new. There are studies going on now looking at this very issue: does the silver-coated tube prevent this adherence of secretions to the ETT?

**Myers:** Maybe not as often as the saline debate comes up, but about every 3 times the saline debate comes up, comes up the debate about patients who have thick, tenacious secretions using a diluted sodium bicarbonate down there to break up those secretions. Is there anything in the literature that supports that, or not?

**Branson:** No, there’s not. There are treatments described where they administer Mucomyst down a bronchoscope channel directly onto secretions, with the idea of breaking up the mucus. But I have to tell you, if you have ever seen Mucomyst applied directly to the bronchial mucosa, when done it’s all irritated and red and inflamed and looks terrible. There’s a group that used to do what’s called BBS, “bicarbonate, bag, and suction,” where they put sodium bicarbonate down the ETT. It’s the same idea with hypertonic saline solution to try and somehow change the secretion volume, but there’s no evidence that I know of that any of that works.

**Fink:** Just a comment Rich. I appreciate the mechanical concept that was shown by the Rouby study using the open or closed suction catheter, but as our discussions went yesterday, the quantity of secretions isn’t nearly as important in this case as whether or not the use of the different catheter affected the frequency for the requirement to suction. Obviously, if you have secretions in the airway, you want to clear them, but if you clear them and you have no greater time component difference or frequency difference, does size really matter?

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**Branson:** I will be careful. I think what the study shows is that if you have secretions that have moved themselves to within reach of the suction catheter, you’d like to remove them. Whether or not that makes some long-term difference, I don’t have any idea, and I don’t know that we could ever even study that.

**Fink:** The other question I had was the depth of suctioning. There’s a difference between being at the predicted end of the ETT, and the method of extending the catheter till it stops, withdrawing about a centimeter, and then applying suction. Have those techniques been differentiated?

**Branson:** No comparisons. There are 3 or 4 studies in the literature; the Dutch study looked at adult patients, and compared the secretion volume that was eliminated. And then they went in and did outcomes, too. How long did the patient stay on the ventilator? How many days of care? How much time in the ICU? All that other stuff—incidence of ventilator-associated pneumonia—no difference between just placing the suction catheter to the end of the ETT or further down than that; and potentially, obviously there is probably less damage, and I guess you could make the argument that poking the carina with the suction catheter probably is not a good thing and might actually damage the carina and lead to greater infection.

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**Wojtczak:*** Very nice presentation, Rich. I’m a pediatric pulmonologist, and one of the treatments that I’ve started using, in conjunction with my pediatric surgery colleagues, is prophylactically placing selected children, postoperatively on HFCWC [high-frequency chest-wall compression]. Specifically, children who have undergone posterior spinal fusion. In a medicine-based-evidence way, I plan to take a closer look to see how these children fare in the immediate postoperative period.

There is an abstract; it’s a small study done by the group at Oakland Children’s Hospital, that looked at using the HFCWC via the Vest, postoperatively in adolescents after posterior spinal fusion. The authors reported that HFCWC with the Vest is a clinically safe and effective airway-clearance technique in ventilated or post-extubation patients who have undergone spinal fusion, and suggested that further studies should be performed. The other question I had was, Do you know of any studies that have looked at the use of HFCWC for nonventilated children with tracheostomy tubes?


**Branson:** Not that I’m aware of. It’s almost always done in acutely ill patients with ETTs, but I would assume the same thing.

**Wojtczak:** So often we will have a child come in, acutely ill with a tracheostomy in and spend 10–14 days on a ventilator in the pediatric ICU. But that’s not been studied as far as you know?

**Branson:** No, it has not.

**Wojtczak:** Thank you.

**McMahon:*** It seems that, due to the depressed respiratory system and
all of the complications that occur with mechanical ventilation, that adult patients coming off the ventilator, either out of surgery or post-extubation would be great candidates for airway-clearance techniques, devices, probably a combination of positive airway pressure, aerosol, and oscillation. Are there any studies ongoing or prompted based off of this discussion to look at airway-clearance devices for earlier extubation, keeping the patient stable, just to support the patient post-surgery to be able to get them out of the ICU earlier, to extubate earlier—similar to getting them up out of the bed, studies based on how they fare overall?

Branson: I think the answer is no. In critical care what we do is look to people who fail extubation, and we look for ways to prevent them from being re-intubated. And that’s being studied, noninvasive ventilation, mask CPAP [continuous positive airway pressure] to prevent those things from happening. But, clearly, once the patient gets extubated in the ICU, the respiratory care group mobilizes, essentially—to make a pun about this entire conference—to the bedside, and our therapists routinely will include aerosol therapy, PEP [positive expiratory pressure], and, if need be, nasotracheal suctioning in patients who are recently extubated, who have a secretion problem. My patient population—I have all of these surgery and trauma patients, and except for one in a while, like when we get a 76-year-old with a hundred pack years of smoking, in a car accident—I don’t see very many people who come in with secretion problems. When you are perfectly normal until you get shot, stabbed, or hit by a car, you don’t have a lot of respiratory mechanics issues, unless you develop pneumonia. I think it’s different in the surgical ICU, in the medical ICU, and in the neurological ICU, where patients with spinal cord injury and head injury—that, to me, is an area that just really needs investigation with all of these devices.

Pierson: Following on that, I agree that the circumstance you’ve identified is clearly one in which we desperately need more knowledge, and working in a surgery trauma unit like you do, Rich, one of the populations of patients who I think most needs to be studied and most needs help is the nonintubated elderly patient with a couple of rib fractures or other injuries who has either not been intubated deliberately from the beginning to try to avoid all of that, or has just been extubated but now has to contend with the effects of heavy sedation, recent trauma, impaired clearance mechanisms because of their rib fractures, their pain, or other things, with or without a history of previous lung disease or smoking.

Branson: This is a different issue, but in my experience the elderly trauma patients—“elderly” gets older all the time, right?—but to me, somebody who’s more than 70, who’s been in an automobile accident and requires mechanical ventilation, I would prefer that we trach [tracheostomize] all of those people, because I think it helps them get off the ventilator faster. I think it helps them get out of the ICU faster; that’s just my clinical impression. It’s really hard to go to families and say, “Hey, we’re doing a study where there is a 50–50 chance we’re going to trach Dad, and, no, we don’t know that he needs it.” It’s just kind of hard to do, but I think there are retrospective reviews that suggest early tracheostomy in the elderly patient actually improves outcomes.1 And secretion clearance becomes infinitely easier when you have a channel to suction through.

Hess: Rich, you talked about clearing secretions from below the ETT, but what about clearing them from above the ETT?

Branson: I didn’t address the idea that you have to suction above the ETT because you have this collection of oropharyngeal secretions, which we believe are the reservoir for creating ventilator-associated pneumonia. Some of us have experience with the CASS [continuous aspiration of subglottic secretions] ETTs, the continuous subglottic suction where you have a little hole in the ETT just above the cuff, and it continuously suctions out the material there. Early studies1 suggested that in patients who were recently intubated without infection, that reduces the incidence of early pneumonia, and the early pneumonias are probably due to aspiration. But it doesn’t reduce the incidence of late pneumonia, which are usually nosocomial infections—Pseudomonas, MRSA [methicillin-resistant Staphylococcus aureus], Acinetobacter, or whatever, whichever one you have. And that’s because those are probably spread through another mechanism, poor hand-washing by us, contamination of the environment, and things like that.

Restrepo: I have a couple of comments. I don’t know if you’ve noticed this, but once you read the Cochrane review published in 2004 on normal saline instillation and the use of deep endotracheal suction technique,1 it just ends with a comment, that even though there is no study that could actually just be analyzed, or just summarized, they consider it to be unethical to use the deep suction technique in light of anecdotal data. Three years ago we conducted a survey, which of course carries the usual limitations of valid-

ity associated with questionnaires, at Children’s Healthcare of Atlanta at Egleston. We did survey 102 clinicians, 27 RTs [respiratory therapists] in this group. Back then, suctioning and saline instillation were considered to be the most common procedures performed in any ICU—and I believe they still are. It might have changed a little now that we performed SBTs [spontaneous breathing trials] on a daily basis and we keep RTs busier than what it was 3 years ago.

All we found—in the process of just writing the manuscript—was that there is a lot of variability on what the nurse’s perception is in terms of the indications from normal saline instillation on endotracheal suction, and the frequency. What we could not believe was that, even among RTs, there were several unknowns in terms of the amount of negative pressure that needs to be applied, the amount of normal saline recommended according to age groups, and for how long these suction events should last. Some clinicians were confused about differences between deep and shallow suctioning.

Also, as a comment, it is amazing that when you review literature, several publications that have evaluated the effects of normal saline instillation go back to the 1970s. It has been 30 years of documenting the adverse effects of both normal saline instillation and endotracheal suctioning, and we still haven’t been that proactive in terms of just trying to create guidelines that are followed by everyone; as Richard said, there’s a lot of variability. Some clinicians are going to be using normal saline no matter what. What is funny about this survey is that when you compare this response to the ones we got 5 years before, the current trend to respond, “No, we don’t use normal saline,” is quite different with the routine use a few years back. Of course, as you turn around, some clinicians will do the complete opposite to what their answer was: deep suction and normal saline instillation. Just a comment.


Branson: The people around the room who do ICU and ventilator patients, how many of your therapists normally put saline down the ETT? Are you guys sure about that? I think you made some good points. Like I said; there are a couple of therapists who work at my place who have been there 25, 30 years, and they are going to put saline down every single time, and there’s nothing you are going to do to change that practice. It’s going to take more than explaining that when you put saline down the tube in a patient with a head injury and it increases their intracranial pressure when they cough. These isolated incidents showing that there is a difference. I think most people just have the impression that it might help, and it probably doesn’t hurt, and that’s why they continue to do it.

Restrepo: One final comment. So far—I’ve been very interested in this for years—the only real indication of a routine normal saline instillation was when we conducted studies on partial liquid ventilation.

Branson: Totally different idea.