Device and Medication Preferences of Canadian Physicians for Emergent Endotracheal Intubation in Critically Ill Patients

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ABSTRACT

Objectives: Various medications and devices are available for facilitation of emergent endotracheal intubations (EETIs). The objective of this study was to survey which medications and devices are being utilized for intubation by Canadian physicians.

Methods: A clinical scenario-based survey was developed to determine which medications physicians would administer to facilitate EETI, their first choice of intubation device, and backup strategy should their first choice fail. The survey was distributed to Canadian emergency medicine (EM) and intensive care unit (ICU) physicians using web-based and postal methods. Physicians were asked questions based on three scenarios (trauma; pneumonia; heart failure) and responded using a 5-point scale ranging from “always” to “never” to capture usual practice.

Results: The survey response rate was 50.2% (882/1,758). Most physicians indicated a Macintosh blade with direct laryngoscopy would “always/often” be their first choice of intubation device in the three scenarios (mean 85% [79%-89%]) followed by video laryngoscopy (mean 37% [30%-49%]). The most common backup device chosen was an extraglottic device (mean 59% [56%-60%]). The medications most physicians would “always/often” administer were fentanyl (mean 45% [42%-51%]) and etomidate (mean 38% [25%-50%]). EM physicians were more likely than ICU physicians to paralyze patients for EETI (adjusted odds ratio 3.40; 95% CI 2.90-4.00).

Conclusions: Most EM and ICU physicians utilize direct laryngoscopy with a Macintosh blade as a primary device for EETI and an extraglottic device as a backup strategy. This survey highlights variation in Canadian practice patterns for some aspects of intubation in critically ill patients.

RÉSUMÉ

Objectifs: Différents dispositifs et différents médicaments s’offrent aux médecins pour faciliter l’intubation endotrachéale (IET) en extrême urgence. L’étude décrite ici visait à déterminer quels dispositifs et quels médicaments utilisent les médecins pour l’intubation au Canada.

Méthodes: Une enquête reposant sur des scénarios cliniques a été élaborée afin de déterminer quels médicaments utiliserent-ils les médecins pour faciliter l’IET en extrême urgence, quel serait leur premier choix de dispositif d’intubation et quelle serait leur solution de rechange en cas d’échec. Le questionnaire a été envoyé soit par voie électronique, soit par la poste aux urgentologues et aux intensivistes travaillant au Canada. Trois scénarios (trauma, pneumonie, insuffisance cardiaque) ont été soumis aux médecins, et ceux-ci devaient répondre aux questions à l’aide d’une échelle à 5 points variant de « Toujours » à « Jamais » pour indiquer leur pratique habituelle.

Résultats: Le taux de réponse à l’enquête a atteint 50,2 % (882/1758). La plupart des médecins ont indiqué qu’une lame Macintosh sous laryngoscopie directe serait « Toujours » ou « Souvent » leur premier choix de dispositif d’intubation, et ce, dans les 3 scénarios (moyenne : 85 % [79-94 %]), puis en vidéo-laryngoscopie (moyenne : 37 % [30-49 %]). L’instrument de rechange indiqué le plus souvent était un dispositif extraglotique (moyenne : 59 % [56-60 %]). Quant aux médicaments, les...
Emergent endotracheal intubation practices

A variety of devices and medications are available to facilitate intubation. Direct laryngoscopy (DL) has been considered the gold standard in airway management, with the Macintosh blade being the most common laryngoscope blade utilized for emergent endotracheal intubation (EETI). Indirect laryngoscopy is a relatively new technology and includes video laryngoscopy (VL) and flexible intubation scopes. VL offers improved visualization of the laryngeal inlet compared to DL and there is emerging evidence that in certain circumstances it may bring additional value for EETI in critically ill patients. Although it has been suggested by some that VL is the new standard for EETI, others have expressed concerns over adoption of these newer technologies.

In addition to device choice, consideration of patient physiology and the pharmacology of medications for intubation are essential to planning an EETI strategy. Available sedative and paralytic medications have unique benefits/risks, and there is controversy over the optimal medication to administer. Etomidate is commonly used as an induction agent as part of rapid sequence intubation (RSI) in emergency medicine (EM) due to its reported minimal effects on patient hemodynamics; however, there are concerns regarding associated adrenal suppression. Other agents such as ketamine, fentanyl, midazolam, and propofol are frequently used alone or in combination to facilitate EETI in patients with hemodynamic instability. Rocuronium and succinylcholine are also commonly administered as muscle relaxants for RSI, and debate continues over which is a superior agent for RSI.

Data on the use of devices and medications for EETI is incomplete. More information on these intubation practices is required to better understand how EETIs are performed and to allow for planning of future research studies. The aim of this study was to describe the EETI practices of Canadian EM and ICU physicians treating critically ill patients. Specifically, our objectives were to determine the types of intubation devices, sedatives, and paralytics being used to facilitate EETI.

METHODS

This study was approved by the Nova Scotia Health Authority Research Ethics Board in Halifax, Nova Scotia. The research team developed a clinical scenario-based survey (available in Supplementary Materials) which was revised and validated for face and content validity through an iterative process among study authors and the Canadian Critical Care Trials Group (CCCTG). The survey involved scenarios of patients with congestive heart failure (CHF), pneumonia, or trauma (Appendix 1) and was designed to identify physician preferences for intubation devices and medications. The electronic survey was constructed using the SelectSurvey instrument (www.selectsurvey.net) and administered in both English and French.

A combined web-based and postal strategy was used to administer the survey to all non-trainee physician members of the Canadian Association of Emergency Physicians (CAEP), the Canadian Critical Care Society (CCCS), and the CCCTG. We combined the membership lists from all three societies and removed any duplicate names, physical addresses, or email addresses. The survey was electronically distributed three times over a 3-month period during 2012/2013. Non-responders to the third electronic survey reminder were mailed a hard copy of the survey along with a pre-stamped envelope to complete and return by post. Controls were in place to ensure physicians did not complete and submit more than one survey. A blinded administrative assistant aided in the coordination of survey distribution and ensured all respondents remained anonymous.

Data was collected on physician specialty, affiliation (academic or community hospital), years in practice, fellowship, and if they currently performed EETI in critically ill patients. Specifically, our objectives were to determine the types of intubation devices, sedatives, and paralytics being used to facilitate EETI.
their practice. Physicians were presented questions regarding techniques and medications they would use in each scenario and asked to respond using a 5-point Likert scale ranging from “always” to “never” based on “what they would do if they were managing the scenario in their usual place of work” to allow for possible variation among physician practice, resources, and support. Survey respondents were grouped as EM or ICU physicians. The EM group included the EM specialties (Fellow of the Royal College of Physicians of Canada [FRCPC]), EM (Canadian College of Family Physicians [CCFP]—EM certificate), EM (CCFP or other), and family medicine. The ICU group included intensivists from the specialties of anesthesia, internal medicine, and surgery; in addition, any physician from another specialty who had completed a CCM fellowship was included in the ICU group.

A Microsoft Excel® (Microsoft Corp., Redmond, WA) database was created and populated directly with responses from the electronic survey. One investigator (MB) entered responses from the paper-based survey into the database. The accuracy of data entry from paper-based surveys was confirmed by randomly checking 10% of the responses. In the analysis, we included any survey question that was fully answered; single and multipart questions that were incompletely answered were excluded. We used descriptive statistics (proportions, means, ranges) and graphically represented the stated practices of physicians as diverging stacked bar charts using a compressed 5-point Likert scale (always/often, sometimes, rarely/never). Multivariable logistic regression was used to model the association between predictor variables of physician characteristics (primary specialty [reference: internal medicine], years of practice [reference: <1 year], CCM fellowship [reference: no CCM fellowship]) and dichotomous outcome variables (backup intubation strategy, use of paralytics). Associations identified through the multivariable analyses were expressed as adjusted odds ratios (aORs) and 95% confidence intervals (CIs). A p-value of <0.05 was considered to be significant for all statistical tests. All data analysis was performed using R (version 3.1.0, Spring Dance) in the RStudio GUI (version 0.98.932) and IBM SPSS Statistics Version 21.21

RESULTS

The survey was sent to 1,758 physicians with a response rate of 50.2% (882/1,758). Some respondents did not complete all of the survey questions; thus, the denominator for each question varies based on the number of physicians who addressed it. A quality check of 10% of paper-based survey responses entered in the database revealed 99.8% accuracy. Characteristics of survey respondents are shown in Table 1. Of physicians who provided information about their specialty, 73% (463/634) were grouped as EM physicians and 27% (171/634) as ICU physicians. Most respondents (79%, 521/661) practiced at an academic hospital, and the majority (61%, 403/662) had more than 10 years of experience; 26% (171/657) of respondents had a CCM fellowship.

The clinical scenarios included a 67-year-old male with CHF, a 59-year-old female with pneumonia, and a 29-year-old male trauma patient in a cervical spine immobilization collar with abrasions on his head, chest, and abdomen due to a motor vehicle crash (Appendix 1). Physician preferences for primary intubation strategy are shown in Figure 1. Overall, the device most physicians would “always/often” use as their first choice in the three

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<th>Table 1. Characteristics of Emergency Medicine and Intensive Care Unit Physicians</th>
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<td>Currently performing EETI (n = 657)</td>
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EM = emergency medicine; ICU = intensive care medicine; CCM = critical care medicine; EETI = emergent endotracheal intubation.

*EM physicians included the specialties EM (CCFP-EM) (n = 276), EM (FRCPC) (n = 128), EM (CCFP or other) (n = 42), and family medicine (n = 26); percentages are based on number of EM physicians who answered each question.

†ICU physicians included the specialties internal medicine (n = 96), anesthesia (n = 57), and surgery (n = 108); percentages are based on number of ICU physicians who answered each question.
Figure 1. Primary strategy of emergency medicine (EM) and intensive care unit (ICU) physicians for emergent endotracheal intubation in three clinical scenarios. aOther indirect devices include intubating laryngeal mask airway, optical stylet, airtrac, and lightwand.
The medication most EM physicians would use was ketamine in the pneumonia scenario (61%, 127/208) and etomidate in the CHF (54%, 246/457) and trauma (63%, 284/451) scenarios. In contrast, most ICU physicians would “always/often” administer fentanyl in all three scenarios (CHF 77%, 128/167; pneumonia 76%, 127/168; trauma 75%, 125/166). A greater proportion of ICU physicians indicated they would “always/often” use midazolam (CHF 64%, 107/166; pneumonia 65%, 110/166; trauma 59%, 98/166) compared to EM physicians (CHF 18%, 84/452; pneumonia 17%, 77/447; trauma 16%, 71/453).

Figure 4 shows physician preferences for use of paralytics to facilitate EETI. Succinylcholine was the drug most EM physicians would “always/often” use for paralysis (CHF 72%, 329/459; pneumonia 77%, 356/461; trauma 78%, 358/459). A considerably smaller proportion of ICU physicians would “always/often” use succinylcholine (CHF 18%, 30/166; pneumonia 19%, 31/166; trauma 29%, 49/167). Rocuronium was marginally preferred over succinylcholine for paralysis by ICU physicians in all three scenarios (CHF 27%, 45/165; pneumonia 30%, 50/164; trauma 33%, 54/165). EM physicians chose to paralyze patients more often than ICU physicians (aOR 3.40; 95% CI 2.90-4.00).

**DISCUSSION**

The results of this clinical scenario-based survey demonstrated that a wide variety of practice patterns exist in Canada for EETI in critically ill patients. Most Canadian EM and ICU physicians would use DL with a Macintosh blade as their first choice of device if a similar patient presented in their practice, followed by VL. If their first choice was unsuccessful, most physicians would use an extraglottic device as a backup strategy. The sedatives most physicians would use to facilitate EETI were fentanyl and etomidate; however, there was variation between EM and ICU physicians in their preferences of which sedatives to use in each of the three clinical scenarios. Overall, EM physicians were considerably more likely than ICU physicians to use paralytics for EETI. Choice of paralytic agent varied, with EM physicians preferring succinylcholine for paralysis while ICU physicians preferred rocuronium. These findings provide evidence of variation between the intubation practices of Canadian EM and ICU physicians and reinforce the need for further investigation into the optimal approaches to EETI in critically ill patients.

Unlike intubations in the operating room where the primary objective is induction of anesthesia, EETIs are...
often performed in unstable patients with the goal of securing the airway as a life-saving intervention in a patient with respiratory failure or shock.\textsuperscript{1,2} The procedure is associated with life-threatening adverse events due to multiple factors related to the patient (comorbidities, limited cardiopulmonary...

**Figure 2.** Backup strategy of emergency medicine (EM) and intensive care unit (ICU) physicians for emergent endotracheal intubation if their primary strategy was unsuccessful. *Extraglottic devices include LMA, King LT, etc.*
Figure 3. Sedatives and anesthetic choices of emergency medicine (EM) and intensive care unit (ICU) physicians to facilitate emergent endotracheal intubation.
reserve, etc.), staff (training level of intubator, skill level and experience of nurses), and environment (limited range of airway equipment, crowded bedside).1,22-24 Device and drug choices to facilitate EETI can significantly impact the outcomes of the procedure.2,13,25 While there has been controversy in EM and ICU circles over the pharmacology and approach to EETI, little data is available on actual clinical practice.23,25-31 When comparing intubation devices, it is important to consider the heterogeneity in design of newer technologies being used as alternatives to DL, the availability of intubation equipment, and the training level of the intubator.11 Our finding that most Canadian physicians perform EETI using DL with a Macintosh blade is similar to the results of recent observational studies. In a prospective single-center study of EETI practices in Australia, Phillips and colleagues found EETI was performed most commonly using DL with a Macintosh blade (53.5% of intubations) followed by use of C-MAC VL (45.9% of intubations).26 An earlier prospective single-center study of Australian EETI practices also found intubation was performed most commonly using DL with a Macintosh blade (53.5% of intubations) followed by use of C-MAC VL (45.9% of intubations).27 In a multicenter prospective surveillance study of 17,583 adult emergency intubations in the United States, Canada, and Australia between 2002 and 2012, Brown and colleagues examined 13 centers and found DL was used in 84% of first attempts while use of VL increased from under 1% in the first three years to 27% in the last three years of the study.25 Another prospective multicenter study of EETI practices in Japan found the vast majority of intubations (90.5%) were performed using DL, followed by use of VL in 4.1% of cases.28 In addition, there have been two Canadian surveys of the practices of anesthesiologists in difficult airway scenarios. Both surveys found DL to be the preferred intubation technique,29,30 and that most anesthetists would use VL in an unanticipated difficult intubation situation if DL was unsuccessful.30

Figure 4. Paralytic preferences of emergency medicine (EM) and intensive care unit (ICU) physicians to facilitate emergent endotracheal intubation.
Several studies comparing rates of first pass success and complications between DL and VL in emergency intubations suggest VL may offer advantages for novice intubators and nonanesthesiologists, or in difficult airway situations; however, these improved success rates in comparative studies have not demonstrated a significant difference in complication rates with the exception of one study which found reduced rates of esophageal intubation and oxygen desaturation events with VL, and the reported first pass success rates are not superior to historical benchmarks from large sets of registry data. A systematic review and meta-analysis of DL and VL use in ICU patients included 9 trials (2,133 patients) and found that VL reduced the risk of difficult orotracheal intubation (OR 0.29; 95% CI 0.20-0.44), Cormack 3/4 grades (OR 0.26; 95% CI 0.17-0.41), and esophageal intubation (OR 0.14; 95% CI 0.02-0.81), and increased first-attempt success (OR 2.07; 95% CI 1.35-3.16) compared to DL. In contrast, a systematic review and meta-analysis of DL and VL use in the emergency room or prehospital setting included four trials (1,305 patients) and found no benefit to use of VL over DL with respect to intubation success, time to intubation, and the glottic view achieved by the device. Recently, two randomized controlled trials (RCTs) have compared use of DL and VL in the emergency department (ED). In one trial, 198 patients undergoing EETI were randomly assigned to either DL or VL using a C-MAC device for the initial intubation attempt; the study authors did not detect a difference between VL or DL using the C-MAC device in first-pass success, attempt duration, aspiration pneumonia, or hospital length of stay. A second RCT randomized 140 patients requiring EETI during cardiopulmonary resuscitation to either DL or VL by an experienced intubator; there were no differences in first-pass success, EETI success rate, or time to complete EETI between the DL and VL groups, but VL was better for completing EETI without chest compression interruptions. Despite a call from some authors for the exclusive use of VL for EETI, our study demonstrates that DL is still the primary choice of physicians practicing EETI in Canada.

Similar to our findings, previous studies have reported variability in the sedative medications used for emergency intubation. One prospective observational study from Australia reported thiopentone was used most often (72.9%) for EETI in the ED, followed by use of ketamine (8.5%). Another study of Australian ED and ICU resuscitation practices found most adult intubations were facilitated with fentanyl (67%), followed by propofol (61.6%). A survey of Italian ED intubation practices found the sedatives used most often were midazolam (59.7%) and propofol (46.3%). In their study of 13 EDs (11 in the United States, 1 in Canada, 1 in Australia), Brown and colleagues found the medication used most often for adult intubation was etomidate (91%), followed by midazolam (3.2%). Since the majority of ED intubations in this multicenter study were performed in the United States (96%, 16,910/17,583), these findings suggest that etomidate is widely used for EETI in the United States. Although our study found most Canadian physicians would use fentanyl to facilitate EETI overall, we also observed that most EM physicians would administer etomidate in the CHF (54%) and trauma (63%) scenarios.

Our finding that most Canadian physicians prefer succinylcholine for paralysis in EETI is in accordance with other studies. An Australian observational study of EETI found succinylcholine was used in 86.8% of ED intubations, while a second study reported that succinylcholine was used in 85.7% of ED and ICU intubations. A study of modified RSI practice in the United States surveyed all anesthesia residency training programs and found most residents and physicians preferred succinylcholine (58%) followed by rocuronium (39%) during RSI procedures. Finally, Brown and colleagues found 75% of the adult emergency intubations in their prospective multicenter study were performed using succinylcholine as a paralytic. Collectively, these results suggest succinylcholine is the medication used most widely by physicians for paralysis in critically ill patients. However, our finding that rocuronium was marginally preferred over succinylcholine by ICU physicians in Canada provides evidence of variation in EETI practice between specialties. While we are unable to determine from this study why EM physicians were more likely than ICU physicians to use paralytics, we believe this highlights a practice variation that warrants further investigation.

The use of devices and medications are certainly not the only factors that can affect intubation success. A number of algorithms and guidelines have been proposed to reduce variation in the practice of EETI and decrease the incidence of adverse events. These include the Eastern Association for the Surgery of Trauma Practice Management Guidelines for emergency tracheal intubation, the Montpellier-ICU intubation algorithm, together with the Eastern Association for the Surgery of Trauma Practice Management Guidelines for emergency tracheal intubation, the Montpellier-ICU intubation algorithm.
and recommendations for difficult airway management from the Canadian Airway Focus Group. While there is some evidence that an intubation management protocol can reduce EETI-related adverse events when compared to a conventional strategy, it is unknown how widely such protocols have been implemented in clinical practice. Simulation-based training has been reported to improve intubation success rates in some studies, but not others. Additional methods found to improve intubation success include the use of preprocedural checklists and preoxygenation.

As with all surveys, this study has limitations. The 49.8% survey non-response rate is an important limitation of this study, since the practices of these non-responding physicians may impact the validity of our findings. Most physicians who responded to the survey practiced in academic settings; this may bias the results and limit their generalizability to Canadian physicians, especially those practicing in non-academic settings. Another possible source of bias is the method used to identify study participants. We used the mailing lists of CAEP, the CCCS, and the CCCTG to identify practicing EM and ICU physicians. These organizations do not represent all EM and ICU physicians in Canada; however, these mailing lists were the most comprehensive national listings of EM and ICU physicians that were available to the study team. It is also important to note that we did not specifically ask physicians about their training, access, or familiarity with VL. Furthermore, since physicians self-reported on their resuscitation practices, some discrepancy may exist between their self-reported preferences and the actual frequencies of device/drug use that would be measured if their practice was observed. We attempted to minimize this by instructing physicians to answer survey questions based on what they would do if they were managing a patient in their usual place of work.

Despite these limitations, we believe this national survey of resuscitation practices among EM and ICU physicians provides valuable insight into the devices and drugs currently being used to perform EETI in Canada, including preferences for a backup intubation strategy. The main strengths of this study are that the clinical scenarios and survey questions were developed and refined by experts in EETI and members of the CCCTG using a rigorous methodology based on guidelines for surveying clinicians from the Academy of Critical Care: Development, Evaluation, and Methodology (ACCADEMY) Group, and that the survey was administered in both electronic and paper-based versions to all members (French and English) of three national societies. This study of Canadian physicians demonstrates that while there is consensus in some aspects of EETI practice such as device preference, there is considerable variability in other aspects such as which medications to use. The findings of this survey provide information that will aid in the planning of future investigations.

CONCLUSION

The results of this survey highlight the practice patterns in Canada for intubation in critically ill patients. Despite the availability of novel video devices, most EM and ICU physicians chose DL with a Macintosh blade for EETI. Medications used to facilitate intubation, including the use of paralytics, was variable. Additional work is required to determine the optimal approaches to performing EETI in the critically ill population.

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SUPPLEMENTARY MATERIAL

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REFERENCES


