Bench Study of a New Device to Display and Maintain Stable Artificial Airway Cuff Pressure

William R Howard MBA RRT

BACKGROUND: Artificial airway cuff pressure should be maintained within a narrow range. Excessive cuff pressure presents a risk of tracheal damage and stenosis. Insufficient cuff pressure adds the risk of secretion leak past the cuff, tidal-volume leakage, and accidental extubation. The available cuff-inflation devices do not address these problems. OBJECTIVE: In the laboratory I developed and evaluated a new cuff-inflation device that continuously displays the cuff pressure and maintains stable cuff pressure. METHODS: The cuff-inflation device evaluation included: test the manometer accuracy; compare the displayed pressure to the pressure delivered to the pilot balloon; determine the device’s response to cuff-pressure changes with the addition of 5 mL or 10 mL of air after achieving a 30 cm H2O baseline; measure the VT leak in an intubated artificial trachea by comparing the device results to benchmark measurements; and determine the stability of baseline cuff pressure during routine cuff checks. RESULTS: The mean ± SD bias and precision of device’s display, compared to the calibration analyzer, was 1.3 ± 2.6 cm H2O. The pressure delivered by the cuff-inflation device’s gas-sampling line to the pilot balloon was equal to the pressure displayed by the cuff-inflation device. With the cuff-inflation device the cuff pressure was unchanged, compared to baseline, after adding 5 mL or 10 mL of air. With 2 current cuff methods, cuff pressure increased to means exceeding 160 cm H2O and 300 cm H2O, respectively. Compared to the benchmark, the difference in exhaled VT mean ± SD bias and precision were: cuff-inflation device 1.4 ± 4.8 mL, and syringe-inflation method 2.4 ± 6.2 mL. Representing a single cuff pressure check, disconnecting the endotracheal-tube pilot balloon from the cuff-inflation device’s gas-sampling line and then reconnecting it had no effect on baseline cuff pressure at 2 seconds or 60 seconds. CONCLUSIONS: The cuff-inflation device demonstrated possible improvements over available cuff-inflation devices and cuff-pressure-control methods. Key words: artificial airway; cuff pressure; tracheal damage; stenosis; tidal volume; accidental extubation. [Respir Care 2011;56(10):1506–1513. © 2011 Daedalus Enterprises]

Introduction

It is recommended to maintain artificial-airway cuff pressure within 20–30 cm H2O.1-6 Excessive cuff inflation is a recognized risk factor for tracheal injury and stenosis.7

Insufficient cuff pressure can increase the leakage of subglottic secretions past the cuff, contribute to air leak, and increase the risk of accidental extubation.8-13 Techniques such as palpation of the pilot balloon or performing routine checks using the minimal occlusive volume technique have not provided protection against these risk factors.14-20

SEE THE RELATED EDITORIAL ON PAGE 1625

Cuff pressure must be monitored to ensure that the pressure remains within the recommended range.21-29 Unfortunately, the existing techniques and devices for monitoring cuff pressure are inadequate and do not lessen the risk factors associated with artificial airways.30,31
I designed a new device to continuously measure and display cuff pressure, and we evaluated the device’s effectiveness in the laboratory. The cuff-inflation device is designed to comply with the American Thoracic Society’s and Infectious Disease Society’s ventilator-associated pneumonia level-II recommendation for maintaining cuff pressure above 20 cm H2O.32

Methods

Fig. 1 shows the new cuff-inflation device that was evaluated. The applied pressure from the cuff-inflation device is adjustable, and in the tested configuration it has an operational range of 0–80 cm H2O. The device attaches to a standard flow meter, and the flow from it is directed through a gas-sampling line to the artificial airway’s pilot balloon. The cuff-inflation device is intended to apply a selectable and consistent pressure to the cuff, without cuff-pressure fluctuation, and to accurately display the cuff pressure.

The device consists of a disposable pressure manometer and a modified, adjustable pressure-relief valve (Ventlab, Mocksville, North Carolina) that has a 22 × 22-mm inner-diameter connector (Airlife, Cardinal Health, Dublin, Ohio). The standard adjustable PEEP valve was modified from the original 0–20 cm H2O range by adding 3 more springs, which increased the cuff-inflation device’s pressure limit to 80 cm H2O for this study. The cuff-inflation device also uses a gas-sampling connector (Smiths Medical North America, Dublin, Ohio), a 3-m length of a male-luer to male-luer gas sampling line (Smiths Medical North America, Dublin, Ohio), and an oxygen tubing connector (Salter Labs, Arvin, California).

Test Setup

The cuff-inflation device was attached to the Diameter-Index Safety System outlet of a standard 0–15 L/min, back-pressure-compensated, oxygen flow meter (Precision Medical, Northampton, Pennsylvania). One end of the gas-sampling line was attached to the gas-sampling port of the cuff-inflation device. The male luer connection on the opposite end of the gas-sampling line was attached to a series of 3-way stopcocks (Fig. 2A). The assembly was used as a connection hub for the cuff-inflation device, the artificial airway pilot balloon, a cuff-inflation syringe, a standard analog pressure gauge (DHD Healthcare, Canastota, New York), and a pressure calibration analyzer (DPM2Plus, Fluke Biomedical, Everett, Washington). The stopcocks were turned, as determined by the tests described below, so that the pilot balloon, the cuff-inflation device gas-sampling line, the syringe-pressure gauge, and the pressure analyzer were in open communication.

A 15-cm length of standard 22-mm inner-diameter aerosol hose was used as an artificial trachea. Three artificial airways were used: a 9.0-mm inner-diameter (12.2-mm outer-diameter) endotracheal tube (ETT) (Sheridan, Teleflex Medical, Research Triangle Park, North Carolina); a 9.0-mm inner-diameter (12.6-mm outer-diameter) ETT (Microcuff, Kimberly-Clark Health Care, Roswell, Georgia); and a 9.0-mm inner-diameter (12.6-mm outer-diameter) cuffed tracheostomy tube with disposable inner cannula (Sims Portex, Keene, New Hampshire). The artificial trachea was intubated with these water-soluble lubricated airways. The opposite end of the artificial trachea was connected to a ventilator (Puritan Bennett 840, Covidien-Nellcor, Boulder, Colorado) with a standard adult patient circuit.

Test 1

Part 1 of Test 1 compared the pressure in the cuff-inflation device to the pressure at the end of the 3-m gas-
A sampling line that was attached to the test hub. This comparison was to determine if the cuff-inflation device’s displayed pressure was a reliable indicator of the applied cuff pressure.

Part 2 of Test 1 compared the cuff-inflation device’s pressure-gauge measurement to the measurement from the calibration analyzer, to establish the accuracy of the manometer.

For Test 1 only, the cuff-inflation device and the calibration analyzer were connected via a single stopcock (see Fig. 2B). In Test 1, Part 1, the stopcock was adjusted to open Port A of the cuff-inflation device and to close Port B. This provided a direct communication between the calibration analyzer and the cuff-inflation device. The cuff-inflation device was attached to the flow meter, which was set to deliver a gas flow of 2 L/min. The cuff-inflation device’s pressure-relief valve was adjusted sequentially to calibration-analyzer readings of 20, 30, 40, 50, and 60 cm H₂O. The cuff-inflation device pressures were measured with the calibration analyzer and recorded at each level. All of the pressure measurements were taken from the calibration analyzer, to avoid intra-device error, and recorded in a spreadsheet (Analysis ToolPak, Excel SP-2 professional edition, Microsoft, Redmond, Washington).

The stopcocks were then adjusted with Port B of the cuff-inflation device open to the calibration analyzer and closed to Port A. The same cuff-inflation-device adjustments were repeated to measure the pressure delivered by the gas-sampling line. These data were also recorded in the spreadsheet.

In Part 2 of Test 1, and for the remainder of the evaluation, the cuff-inflation device and calibration analyzer were reconnected to the main test hub. The 3 stopcocks were adjusted with Ports A and B open (see Fig. 2C), to establish communication between the cuff-inflation device, calibration analyzer, and pilot balloon. PEEP of 5 cm H₂O was applied. The cuff-inflation device was sequentially adjusted to its display value of 20, 30, 40, 50, and 60 cm H₂O at test-lung compliance settings of 10, 20, 30, 40, and 50 mL/cm H₂O. These displayed cuff-inflation-device values and the simultaneously measured calibration-analyzer values were recorded in the spreadsheet. This test was repeated at 10 cm H₂O and repeated with a second cuff-inflation device, which produced 100 cuff-pressure measurements.

**Test 2**

Test 2 was designed to determine the cuff-inflation device’s response to cuff-pressure changes that occur from the diffusion of N₂O into the cuff during intraoperative administration, and the device’s response to cuff-pressure changes during a patient’s daily activities (eg, movement, suctioning, and coughing).

Part 1 of Test 2 evaluated the cuff-inflation device’s response to the addition of air injected into the ETT cuff, after achieving an initial baseline cuff pressure of 30 cm H₂O. End-expiratory pressure of 5 cm H₂O was applied to the Teleflex Medical ETT. The cuff-inflation device was adjusted sequentially until the calibration analyzer displayed values of 20, 30, 40, 50, and 60 cm H₂O at the test-lung compliance settings of 10, 20, 30, 40, and 50 mL/cm H₂O. Twenty-five mL of air was injected from the syringe into the pilot balloon (see Fig. 2A) over a
60-second period. After a 60-second period of stabilization, the pressure measurements from the calibration analyzer were recorded. This process was repeated at an end-expiratory pressure of 10 cm H2O, and with the second cuff-inflation device. The calibration analyzer measurements were recorded for comparison at each pressure.

Part 2 of Test 2 was designed to compare the performance of 2 other cuff-pressure management methods to that of the cuff-inflation device. Additional volumes of air were injected into the cuff after an initial baseline cuff pressure of 30 cm H2O was established. Baseline cuff pressure of 30 cm H2O was initially achieved with 3 types of cuff-pressure management methods: the cuff-inflation device, syringe inflation, and a commercially available cuff-pressure controller (PressureEasy, Medex/Smith’s Medical, St Paul, Minnesota). The response of adding further amounts of air into the cuff after initial baseline was established was then compared.

The tracheal model was intubated sequentially with the 3 artificial airways. The cuff was inflated until a calibration-analyzer cuff-pressure reading of 30 cm H2O was obtained. This was followed by injecting 5 mL and then 10 mL of air sequentially into the cuff through the test hub assembly.

There was one exception in the set-up with the PressureEasy device. It was connected, per the manufacturer’s recommendations, between the pilot balloon and the test hub. It was evaluated for the effect on cuff pressure after adding 5 mL and 10 mL of air to the cuff after baseline inflation.

Fig. 3. The PressureEasy device was added between the pilot balloon and the test hub. It was evaluated for the effect on cuff pressure after adding 5 mL and 10 mL of air to the cuff after baseline inflation.

Test 3

Test 3 was designed to evaluate the cuff-inflation device’s ability to maintain a tracheal-cuff seal at various levels of compliance and PEEP. The integrity of the tracheal-cuff seal was deemed effective if the exhaled Vt measurement was no less than the benchmark measurement, which was with the patient circuit connected directly to the test lung, with the artificial airway and artificial trachea removed. The cuff-inflation device and syringe-inflation method of sealing the cuff against the artificial trachea were evaluated and compared to the benchmark exhaled Vt.

The cuff-inflation device was evaluated while mechanically ventilating the test lung (see Fig. 2A), with both types of cuffed ETT and the tracheostomy tube. The artificial airways were coated with a water-soluble lubricant (Johnson & Johnson, New Brunswick, New Jersey) prior to insertion into the artificial trachea. The test-lung compliance was sequentially adjusted to 10, 20, 30, 40, and 50 mL/cm H2O. At each compliance the ventilator was set to Vt of 500, 600, 700, and 800 mL, respiratory rate of 15 breaths/min, PEEP of 5 cm H2O and 10 cm H2O, and a peak flow rate to maintain the inspiratory-expiratory ratio at 1:2. The syringe and analog pressure gauge were closed to the hub, and Stopcock 3 was opened to allow the cuff-inflation device to inflate the cuff. The cuff-inflation device was adjusted and set to approximately 5 cm H2O above the peak airway pressure.

With the syringe-inflation method, Stopcock 3 of Port A (see Fig. 2A) was turned off to Stopcocks 1 and 2, which isolated the cuff-inflation device and calibration analyzer from the test hub. Stopcocks 1 and 2 were open to the syringe, pilot balloon, and the analog pressure gauge, which allowed air injection from the syringe into the cuff. Approximately 5 cm H2O above the peak airway pressure was targeted at each setting and measured with the analog gauge. After 30 breaths were delivered, the next 10-breath exhaled Vt average from the ventilator’s measurement display was recorded. The step-by-step process, as described in the first phase of this test, was repeated, and the measurements were recorded in the spreadsheet.

The artificial trachea was then removed and the patient circuit was connected directly to the test lung. The first phase of this test was repeated to represent the benchmark of Vt, and these measurements were recorded.

Test 3 produced 120 exhaled Vt measurements each from the syringe-inflation method and the cuff-inflation device.
These measurements were compared to the VT measurements from the direct circuit connection to the test lung.

Test 4

Test 4 was designed to determine whether the cuff-inflation device would maintain a baseline cuff pressure if used intermittently for routine checks. The baseline cuff pressure was compared to the cuff pressure that resulted after a single disconnection and reconnection of the cuff-inflation device to the pilot balloon.

With 0 PEEP, the cuffs were sequentially inflated to 20, 30, 40, 50, and 60 cm H2O. The test-lung compliance was also adjusted in sequence to 10, 20, 30, 40, and 50 mL/cm H2O. The stopcocks were open to allow communication between the pilot balloon, the calibration analyzer, and the cuff-inflation device. After a 60-second period of stabilization at each setting, the ETT pilot balloon was disconnected. After another 60 seconds it was reattached to the test hub and the cuff pressure was recorded at 2 seconds and 60 seconds. These same tests were repeated with PEEP of 5 cm H2O and 10 cm H2O. A total of 150 measurements were recorded.

Statistical Analysis

The data from Test 1 and Test 3 were analyzed with limits-of-agreement analysis. In Test 1, bias was calculated as the mean difference between the cuff-pressure measurement from the calibration analyzer and that from the cuff-inflation device. Precision was calculated as the range included between ± 2 standard deviations from the mean difference. In Test 2, the cuff-pressure measurements were averaged and compared to actual cuff pressure with the calibration analyzer at 20, 30, 40, 50, and 60 cm H2O, with a 1-sample t test.

In Test 3, bias was calculated as the mean difference in measured exhaled VT between the direct ventilator-circuit connection to the test lung and the measured exhaled VT after inflating the cuffs with the cuff-inflation device or syringe method. Precision was taken as the range included between ± 2 standard deviations from the mean difference.

In Test 4, the cuff pressure measured at 2 seconds and 60 seconds after reconnection was averaged at the 3 PEEP and test-lung compliance settings. This was compared to the respective baseline cuff pressure of 20, 30, 40, 50, and 60 cm H2O, which were measured with the calibration analyzer, with a 1-sample t test.

All descriptive statistics were calculated with the spreadsheet software. Differences were considered statistically significant when P < .05.

Results

Test 1

In Part 1 of Test 1 the pressure measured in the cuff-inflation device was the same as the pressure measured at the test hub that was delivered by the 3-m cuff-inflation device gas-sampling line. The Pearson correlation was 1.0. In the second Part of Test 1 the cuff-inflation device pressure-gauge display and calibration analyzer comparison mean ± SD bias and precision was 1.3 ± 2.6 cm H2O (Fig. 4).

Test 2

In Part 1 of Test 2, after 25 mL of air was injected from the syringe and a 60-second period for equilibration, the mean ± SD bias and precision was 1.3 ± 2.6 cm H2O, or identical results to Test 1. The Pearson correlation was 1.0.

In Part 2 of Test 2 there was no difference in the cuff pressure after adding 5 mL or 10 mL of air into the cuff, compared to the baseline cuff pressure from the cuff-inflation device. With the PressureEasy device the cuff pressure increased to a mean of 162 cm H2O after adding 5 mL of air, and to 301 cm H2O after adding 10 mL of air. With the syringe-inflation method, cuff pressure increased to a
mean of 167 cm H\textsubscript{2}O by adding 5 mL of air, and to 335 cm H\textsubscript{2}O by adding 10 mL of air (Fig. 5).

**Test 3**

The tracheal-cuff seal at different compliance and PEEP levels was analyzed by comparing the measured VT. The comparison of directly connecting the patient circuit to the test lung was made against inflating the cuffs with the cuff-inflation device and the syringe-inflation method. Compared to the benchmark, the difference in exhaled VT mean ± SD bias and precision were: cuff-inflation device 1.4 ± 4.8 mL, syringe-inflation method 2.4 ± 6.2 mL (Fig. 6).

**Test 4**

After disconnecting the ETT pilot balloon from the cuff-inflation device gas-sampling line and then reconnecting it, there was no change at 2 seconds or 60 seconds from the baseline cuff pressure. The Pearson correlation was 1.0.

**Discussion**

The cuff-inflation device display compared favorably to the calibration analyzer benchmark measurements. The displayed information may increase cuff-pressure awareness as an alternative to the pilot-balloon-palpation and minimal-occlusive-volume techniques, in which pressure is not measured with a manometer. However, without further study and clinical trials, it is unknown whether this information will be beneficial.

Cuff pressure was unchanged from the initial baseline when air was added to the cuff with the cuff-inflation device. This experiment was intended to determine the cuff-inflation device’s response to the cuff compression that occurs during daily activities\textsuperscript{34} and the intra-operative cuff diffusion of N\textsubscript{2}O.\textsuperscript{12,23,24} This finding of cuff pressure stability with the cuff-inflation device compares to the syringe-inflation method and the PressureEasy device. With those 2 methods an excessive and unacceptably high pressure was found above their initial baseline inflation (see Fig. 5).

The cuff-inflation device formed an effective seal between the cuff and artificial tracheal wall. This was demonstrated by comparing favorably to the benchmark measurement of exhaled VT. Finally, when compared to traditional methods used for cuff checks, the cuff pressure was stable after the removal and reconnection of the cuff-inflation device to the pilot balloon. This finding compares to the cuff volume loss that occurs with the devices that are currently used for routine baseline cuff-pressure measurement.\textsuperscript{30}

**Limitations**

The continuous cuff pressure displayed by the cuff-inflation device suggests that there is a potential for heightening clinician awareness of and adherence to maintaining recommended cuff pressure, but this study does not demonstrate that potential benefit. This study also did not determine whether the findings and inferences made from it are transferable to the clinical setting, or if improvements in outcomes will occur. For example, we do not know if using the cuff-inflation device clinically will prevent secretion leakage past the cuff or otherwise protect against ventilator-associated pneumonia or tracheal injury. Further study is necessary, with a 510K-approved version of the cuff-inflation device, to determine whether any of the findings from this bench study will be of value or impact clinical outcomes.
Conclusions

Artificial airways require vigilant monitoring to ensure that cuff pressure is maintained within an acceptable range. Unfortunately, the existing techniques and devices for monitoring cuff pressure are inadequate. Maintaining the cuff pressure is recommended, but in the process of routine checking, the existing techniques and devices substantially lower the cuff volume. We designed a new device to apply continuous cuff pressure, and the device demonstrated a stable, continuous, and accurate display of cuff pressure. As a test of the seal between the cuff and the artificial trachea, the measured exhaled $V_T$ with the cuff-inflation device compared favorably to the benchmark setup in which the patient circuit was connected directly to the test lung.

We do not know, without further study with a clinically approved version of the cuff-inflation device, if even one ventilator-associated pneumonia or tracheal injury can be prevented. It needs to be determined if clinical benefit will result from cuff-pressure management with a 510K-approved cuff-inflation device. Further study is warranted if this becomes available, to identify if such a design will have any effect on patient outcomes, compared to the current devices and techniques used for cuff-pressure management.

REFERENCES