Wet-Side Breath-Enhanced Jet Nebulization: Controlling Drug Delivery During Mechanical Ventilation

Ann D Cuccia, Sunya Ashraf, Michael McPeck, Joshua Samuel, and Gerald C Smaldone

BACKGROUND: The present study tested a novel nebulizer and circuit that use breath enhancement and breath actuation to minimize ventilator influences. The unique circuit design incorporates “wet-side” jet nebulization (the nebulizer connected to the humidifier outlet port) to prevent unpredictable aerosol losses with active humidification. The system was studied using several ventilator brands over a wide range of settings, with and without humidification. METHODS: During treatment, a 2-position valve directed all ventilator flow to the nebulizer, providing breath enhancement during inspiration. Aerosol was generated by air 50 psi 3.5 L/m triggered during inspiration by a pressure-sensitive circuit. Particles were captured on an inhaled mass filter. Testing was performed by using active humidification or bypassable valved heat and moisture exchanger (HME) over a range of breathing patterns, ventilator modes, and bias flows (0.5-5.0 L/m). The nebulizer was charged with 6 mL of radiolabeled saline solution. Mass balance was performed by using a gamma camera. Tidal volume was monitored by ventilator volume (exhaled V_T) and test lung volume. The Mann-Whitney test was used. RESULTS: A total of 6 mL was nebulized within 1 h. Inhaled mass (% neb charge): mean ± SD (all data) 31.1% ± 6.45; no. = 83. Small significant differences were seen with humidification for all modes (humidified 36.1% ± 5.60, no. = 26; bypassable valved HME 28.8% ± 5.51, no. = 57 [P < .001]), continuous mandatory ventilation modes [P < .001], and pressure support airway pressure release ventilation modes [P < .001]. Mass median aerodynamic diameter ranged from 1.04 to 1.34 μm. The V_T was unaffected (exhaled V_T -5.0 ± 12.9 mL; P = .75) and test lung (test lung volume 25 ± 14.5 mL; P = .13). Bias flow and PEEP had no effect. CONCLUSIONS: Breath enhancement with breath actuation provided a predictable dose at any ventilator setting or type of humidification. Preservation of drug delivery during active humidification is a new finding, compared with previous studies. The use of wall gases and stand alone breath actuation standardizes conditions that drive the nebulizer independent of ventilator design. Wet-side nebulizer placement at the humidifier outlet allows delivery without introducing aerosol into the humidification chamber. Key words: aerosols; nebulizers and vaporizers; administration; inhalation; ventilators; mechanical; humidifiers; drug delivery; bias flow. [Respir Care 2020;65(8):1077–1089. © 2020 Daedalus Enterprises]
in the ventilator circuit,6,7 humidification,2,5-8 duty cycle,6,7,9 bias flow,1,5 nebulizer technology,1,2,6,7,10,11 nebulizer driving pressure and flow,5,12 internal ventilator valves,3,4 device reliability,14 fill volume,4,15 and treatment time.6,7 Output data from commercially available devices

The present article describes a new jet nebulizer-ventilator circuit combination (i-AIRE, InspiRx, Somerset, New Jersey) designed to mitigate the problems described above and deliver a predictable dose to the patient who is intubated and on mechanical ventilation. The device incorporates the concept of breath enhancement into the ventilator circuit. The nebulizer is powered by wall gases at 50 psi at a relatively low flow of 3.5 L/min. The new nebulizer–circuit design directs all inspiratory ventilator flow directly through the top of the nebulizer during active nebulization. A Venturi effect within the nebulizer enhances aerosol generation only during inspiration (Fig. 2). Breath enhancement, therefore, should increase nebulizer output, reducing both treatment time and the influence of expiratory phenomena on aerosol losses, for example, duty cycle.

Combined with a pressure-controlled independent breath-actuated circuit, providing wall gases to the nebulizer only during inspiration (breath actuation), dose delivery to patients who are intubated may be improved. Active humidification can result in unpredictable losses of generated aerosol in ventilator tubing and, depending on nebulizer location, in the humidifier.1 Clinical studies have avoided humidification entirely to prevent these effects.5,22-24 Modern use of dry-side humidification may lead to humidifier contamination. To prevent humidifier losses and potential contamination, we placed our nebulizer on the “wet” side of the humidifier (humidifier outlet). To test this nebulizer–system design, the effects on aerosol delivery by

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What this paper contributes to our knowledge

Drug delivery to the patient who is intubated and on mechanical ventilation is uncontrolled, with published studies reporting values that range from 2.7% to 41% of the nebulizer dose. This variability may affect the results of clinical studies for drugs that cannot be assessed at the bedside, for example, antibiotics. Important variables include nebulizer type, duty cycle, bias flow, nebulizer position, and methods of humidification.

Methods

Nebulizer

Jet nebulizers have a nebulizing chamber or reservoir that contains liquid medication; a jet of pressurized gas is directed through a narrow orifice, which results in a reduction in pressure and medication being drawn up through the capillary tube from the nebulizer reservoir into the gas stream. Baffles placed in the aerosol stream act to produce smaller particles and remove larger particles, returning them to the reservoir. Breath-enhanced jet nebulizers that are currently used in spontaneously breathing patients are designed to generate more aerosol during inspiration25 by entraining the patient’s inspiratory flow through the top of the nebulizer into the nebulizing chamber, increasing nebulizer output during inspiration relative to expiration.26 The i-AIRE system jet nebulizer (Fig. 3) functions in a similar manner.

The i-AIRE system is designed to accept all inspiratory flow going from the ventilator to the patient. This flow-enhanced aerosol effect operates only during inspiration (breath-enhanced nebulization). To generate aerosol, the nebulizer requires wall air or oxygen using a standard flow

Quick Look

Current knowledge

Drug delivery to the patient who is intubated and on mechanical ventilation is uncontrolled, with published studies reporting values that range from 2.7% to 41% of the nebulizer dose. This variability may affect the results of clinical studies for drugs that cannot be assessed at the bedside, for example, antibiotics. Important variables include nebulizer type, duty cycle, bias flow, nebulizer position, and methods of humidification.

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What this paper contributes to our knowledge

The i-AIRE system with breath-enhancement and stand-alone breath-actuation provided predictable drug delivery during invasive ventilation, independent of the ventilator type, across a wide range of commonly used settings, with and without humidification. Wet-side nebulizer placement at the humidifier outlet avoided humidifier effects on aerosol delivery and contamination of the humidifier.
meter at 3.5 L/min, 50 psi, is controlled by a pressure-actuated solenoid, activated only during inspiration (breath-actuated nebulization). The 3.5 L/m flow is delivered to the nebulizer gas inlet nipple and is controlled by the solenoid, which actively opens and delivers the 3.5 L/m flow during a defined percentage of inspiration, and actively closes and ceases flow to the nebulizer during the expiratory phase. At the same time, during inspiration, all inspiratory gas flow from the ventilator is directed through the nebulizer (breath enhancement) (Fig. 2); this added flow through the top of the nebulizer lowers the local pressure in the nebulizer, causing a further decrease in pressure, which augments the rate at which liquid is drawn through the capillary tube and increasing the rate of aerosol formation and nebulizer output.

The i-AIRE jet nebulizer was filled (nebulizer charge) with 6 mL of normal saline solution mixed with technetium-99m and connected either to the wet side (humidifier outlet) of the humidifier or to the inspiratory outlet of the ventilator via a unique 2-way breath-actuated–breath-enhanced valved circuit (Figs. 4 and 5). The 6-mL nebulizer charge was chosen to assess delivery of volumes consistent with antibiotic solutions. The run time during early experiments varied between 30 and 60 min, depending on the duty cycle and the presence or absence of humidity (estimated, from the time that the 2-position valve was turned to engage breath enhancement to the point of cessation of aerosol production determined by visual inspection). Based on these observations, run time was standardized to 60 min for all the experiments defined from the time that the 2-position valve was turned to engage breath enhancement and ended at 60 min (measured by a stopwatch).

Solenoid

The pressure-activated solenoid (ARO Solenoid M182SD-012, Ingersoll-Rand, Bryan, Ohio) actively opened and actively closed via dual electromagnetic controls triggered by an electronic circuit. This was accomplished by using an Arduino controller board to sense pressure and control the solenoid. First, for several breaths, the circuit assessed the average inspiratory time (T1). Then, once the circuit sensed the increase in airway pressure during inspiration, the

Fig. 1. Inhaled mass data from current nebulizer technologies reported in the literature. Inhaled mass as percentage of nebulizer charge versus duty cycle (inspiratory time/total cycle time [T1/Ttot]).

Fig. 2. Breath enhancement. Venturi effect created by added flow through the top of nebulizer causes a further decrease in pressure across the nebulizer chimney (P0) inversely proportional to local air flow (V). The drop in pressure augments the rate at which liquid is drawn through the capillary tube (Q), which results in an increase in rate of aerosol formation and nebulizer output during inspiration.
solenoid opened in ~20 ms and remained open until ~75% of inspiration was completed, and the valve actively closed.

The Circuit

Our model is outlined in Figures 4 and 5. Breath enhancement required that, during aerosol therapy, all ventilator inspiratory flow be directed to the nebulizer. During active humidification (Fig. 4), the system was positioned on the wet side (humidifier outlet) of the humidifier (MR850, Fisher Paykel, Auckland, New Zealand; or ConchaTherm Neptune, Teleflex Medical, Morrisville, North Carolina). During a treatment, the 2-position valve was turned to the nebulizer and the breath-actuation circuit engaged. Inspiratory gas passed through the humidifier in the usual manner and after exiting the humidifier, was diverted through the top of the nebulizer. After treatment was completed (Fig. 4, insert), the breath-acting circuit was turned off and the 2-position valve turned, which bypassed the nebulizer and directed flow to the conventional circuit.

During passive humidification (Fig. 5), the i-AIRE system was positioned at the inspiratory outlet of the ventilator. During aerosol delivery, the bypassable valved heat and moisture exchanger (HME) was turned to the aerosol position, and the air flow was diverted in the same manner by using the 2-position valve. After treatment was completed (Fig. 5, insert), flow was redirected to the conventional circuit by turning the 2-position valve, and the HME turned to the HME position. Ventilatory parameters were measured during nebulization with an ASL 5000 test lung (Ingmar Medical, Pittsburgh, Pennsylvania). Positioning the nebulizer on the wet side (humidifier outlet) allows aerosol to be delivered through the circuit without coming in contact with the humidifier chamber. To avoid potential effects of nebulized cold aerosol and wall gas on humidification, the temperature probe of the humidifier was relocated from the standard location on the proximal end of the conventional inspiratory line to the point where the 2-position valve was placed on the humidifier (Fig. 4).

Three test ventilators were used in this study: Avea (Vyaire Medical, Mettawa, Illinois), Dräger V500 (Dräger Medical, Lubeck, Germany), and Servo-i (Maquet, Getinge, Solna, Sweden). Each ventilator was connected via a standard 22-mm ventilator circuit, heated wire circuits were used for experiments that used active humidification (Evaqua, Fisher Paykel; or Hudson RCI ISO-GARD, Teleflex Medical). Heated humidification experiments were tested by using either the MR850 humidifier (Fisher Paykel) or ConchaTherm Neptune humidifier (Teleflex Medical) set at 37°C. For the HME configuration, the circuit contained a HME (Curaplex ClearPath HME, Tri-anim, Dublin, Ohio).

During active humidification, before aerosolization, the heated wire circuit was heated and humidified for ~20 min to reach a stable 37°C. To complete the circuit, a closed suction system (Halyard Health, Alpharetta, Georgia) and a 7.5-mm endotracheal tube (Rusch, Teleflex, Medical) were connected to an ASL 5000 test lung. An inhaled mass (IM) filter (Pari, Starnberg, Germany) placed in the circuit distal to the endotracheal tube measured the aerosol particles that would be inhaled by a patient (IM) under similar conditions. A similar filter was placed in the expiratory line for mass balance measurements.

Ventilator settings are listed in Tables 1–6. Ventilator modes were chosen to encompass relevant clinical settings, including controlled mechanical ventilation, pressure support, and patient-triggered breaths. Most experiments were conducted at a bias flow of 2.0 L/min. The ASL 5000 test lung served 2 roles, (1) it monitored ventilatory parameters during aerosol delivery, and (2) it served to trigger the ventilator during assisted and pressure-supported modes of ventilation. Each mode was tested by using all 3 ventilator brands at varying airway pressures, tidal volumes (VT), frequencies, T₁ (duty cycle), and a PEEP of 5 cm H₂O at 21% oxygen with 2.0 L/m bias flow during HME or active humidification. In a separate series of experiments, the effect of PEEP was tested by using 4 ventilator settings, and bias flow was tested with HME by using 2 ventilator settings.

Particle Distribution

In another series of experiments, particles were sampled via a cascade impactor (2 L/min flow, Marple 8-stage impactor,
Thermo Fisher Scientific, Waltham, Massachusetts) with the device located distal to the endotracheal tube; experiments were conducted with both HME and active humidification. For these measurements, the i-AIRE nebulizer was charged with 6 mL of normal saline solution mixed with a known amount of radioactivity. Aerosols were sampled over a 9-min period for a single ventilator setting (Avea, volume control–continuous mandatory ventilation, frequency 18 breaths/min, VT 500 mL, flow 43 L/min, TI 0.7 s, PEEP 5 cm H2O, bias flow 2.0 L/min). Radioactivity on the cascade stages was measured by using a collimated rate meter (Ludlum Measurements, Sweetwater, Texas), and the distribution was plotted on log probability paper. Activity at the median defined the mass median aerodynamic diameter.

**Bias Flow**

A bias flow of 2.0 L/min (a common default value) was used for most experiments. The effect of varying bias flow was measured at bias flows of 0.5, 3.5, and 5.0 L/min at 2 ventilator settings: volume control–continuous mandatory ventilation, frequency 15 breaths/min, Vt 460 mL, flow 55 L/min, T I 0.50 s, and I:E 1:6.5; and volume control-continuous mandatory ventilation, frequency 20 breaths/min, Vt 650 mL, flow 40 L/min, T I 0.97 s, and I:E 1:2.1. These experiments were carried out by using HME with an Avea ventilator, which allows adjustment of bias flow over this range.

**PEEP**

A PEEP setting of 5 cm H2O was used for the majority of experiments. To test the potential effect of varying PEEP conditions, 10 and 15 cm H2O was used with T I of 0.5, 0.7, 1.0 s; duty cycle of 0.20, 0.33 using the Avea and Dräger ventilators.

**Analysis**

Aerosol delivery was defined as the IM% or radioactivity on the IM filter reported as a percentage of nebulizer charge. To relate IM% to the range of ventilator test conditions, IM% was plotted as a function of T I. T I was read directly on the ventilator and varied widely with different ventilator settings. Data were also plotted against...
duty cycle (Ti/Ttot), the fraction of the breath that was inspiration; this parameter was calculated from ventilator-derived parameters and allowed comparison with data reported from other studies that measured aerosol delivery during mechanical ventilation.

To describe the distribution of radioactivity throughout the ventilator circuit a complete mass balance was carried out for 2 experiments. This process included measurement of IM, nebulizer residual, deposition in ventilator tubing, expiratory filter, and endotracheal tube for the same settings during HME and active humidification. Radioactivity deposited in all parts of the circuit (nebulizer, tubing, and filters) was quantified by using a gamma camera (Maxi Camera 400 [General Electric, Horsholm, Denmark]; model 604/150/D [Power Computing, Austin, Texas]; Nuclear MAC, version 4.2.2 [Scientific Imaging, Campbell, California]).

Statistics

Data were analyzed by using GraphPad Prism version 8.1 for Mac OS X (GraphPad Software, San Diego, California) and reported as mean ± SD. Group data were compared by using the Mann-Whitney rank sum test.

Results

IM

IM% (% neb charge): mean ± SD [all data] 31.1% ± 6.45, no. = 83; humidified 36.1% ± 5.60, no. = 26; HME 28.8% ± 5.51, no. = 57. Variation in IM percentage with different settings and modes of ventilation is shown on Figure 6. On the left, data are plotted against T1, which illustrates IM% with controlled ventilator settings for different ventilators. On the right, are data that could not be plotted against T1 because the mode contained spontaneous-pressure supported breaths triggered by the ASL 5000 test lung and, therefore, there was no single value for T1. The latter data are grouped by ventilatory mode.

In general, aerosol delivery was preserved for all settings and conditions with limited variability. A trend of increasing aerosol delivery was seen, with increasing duty cycle. Although there is considerable overlap between HME and humidified aerosol delivery, there is a small systematic shift that favors humidification, which was significant between humidification type (HME and active...
humidification) for all data \((P < .001)\), CMV modes \((P < .001)\), and pressure support–airway pressure release ventilation modes \((P < .001)\). Bias flow had no effect as illustrated by the points at \(T_I\) of 0.5 and 1.0 s. (Fig. 6). PEEP had no effect, 10 and 15 cm H\(_2\)O were tested as illustrated by points at \(T_I\) 0.5, 0.7, and 1.0 s (Fig. 6).

Table 1. Ventilator Settings: VC-CMV Mode

<table>
<thead>
<tr>
<th>Ventilator</th>
<th>Ventilator frequency, breaths/min</th>
<th>Set (V_T), mL</th>
<th>Flow, L/min</th>
<th>(T_I), s</th>
<th>I:E</th>
<th>Duty Cycle ((T_I/T_{tot}))</th>
<th>Waveform</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avea 12</td>
<td>700</td>
<td>42</td>
<td>1.00</td>
<td>1:3.8</td>
<td>0.20</td>
<td>Const</td>
<td></td>
</tr>
<tr>
<td>Avea 15</td>
<td>460</td>
<td>55</td>
<td>0.54</td>
<td>1:6.5</td>
<td>0.14</td>
<td>Const</td>
<td></td>
</tr>
<tr>
<td>Avea 18</td>
<td>500</td>
<td>60</td>
<td>0.70</td>
<td>1:3.8</td>
<td>0.21</td>
<td>Const</td>
<td></td>
</tr>
<tr>
<td>Avea 20</td>
<td>650</td>
<td>40</td>
<td>0.97</td>
<td>1:2.1</td>
<td>0.32</td>
<td>Const</td>
<td></td>
</tr>
<tr>
<td>Dräger 12</td>
<td>700</td>
<td>45</td>
<td>1.00</td>
<td>1:4.0</td>
<td>0.20</td>
<td>Decel</td>
<td></td>
</tr>
<tr>
<td>Dräger 15</td>
<td>460</td>
<td>59</td>
<td>0.53</td>
<td>1:6.5</td>
<td>0.13</td>
<td>Const</td>
<td></td>
</tr>
<tr>
<td>Dräger 18</td>
<td>500</td>
<td>Autoflow</td>
<td>1.00</td>
<td>1:3.7</td>
<td>0.21</td>
<td>Decel</td>
<td></td>
</tr>
<tr>
<td>Dräger 20</td>
<td>650</td>
<td>44</td>
<td>0.98</td>
<td>1:2.1</td>
<td>0.33</td>
<td>Const</td>
<td></td>
</tr>
<tr>
<td>Servo 15</td>
<td>460</td>
<td>59</td>
<td>0.55</td>
<td>1:6.3</td>
<td>0.14</td>
<td>Decel</td>
<td></td>
</tr>
<tr>
<td>Servo 18</td>
<td>500</td>
<td>43</td>
<td>0.70</td>
<td>1:3.8</td>
<td>0.21</td>
<td>Decel</td>
<td></td>
</tr>
<tr>
<td>Servo 20</td>
<td>650</td>
<td>40</td>
<td>1.00</td>
<td>1:2.1</td>
<td>0.33</td>
<td>Decel</td>
<td></td>
</tr>
</tbody>
</table>

Three ventilator brands that use heated wire humidification or bypassable–valved HME, PEEP 5.0 cm H\(_2\)O, bias flow 2.0 L/min.

VC-CMV = volume control–continuous mandatory ventilation

\(T_I\) = inspiratory time

I:E = inspiratory to expiratory ratio

\(T_{tot} = T_I + T_e\)

Const = constant flow waveform

Decel = decelerating waveform

Table 2. Ventilator Settings: PC-CMV Mode

<table>
<thead>
<tr>
<th>Ventilator</th>
<th>Ventilator frequency breaths/min</th>
<th>Set (V_T), mL</th>
<th>PIP (PC), cm H(_2)O</th>
<th>(T_I), s</th>
<th>I:E</th>
<th>Duty Cycle ((T_I/T_{tot}))</th>
<th>Waveform</th>
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</thead>
<tbody>
<tr>
<td>Avea 18</td>
<td>470</td>
<td>15 (10)</td>
<td>0.85</td>
<td>1:5.3</td>
<td>0.26</td>
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<td></td>
</tr>
<tr>
<td>Servo 20</td>
<td>500</td>
<td>15 (10)</td>
<td>0.90</td>
<td>1:2.3</td>
<td>0.30</td>
<td>Decel</td>
<td></td>
</tr>
<tr>
<td>Dräger 18</td>
<td>415</td>
<td>15 (10)</td>
<td>0.70</td>
<td>1:3.2</td>
<td>0.24</td>
<td>Decel</td>
<td></td>
</tr>
</tbody>
</table>

Three ventilator brands that use heated wire humidification or bypassable–valved HME, PEEP 5.0 cm H\(_2\)O, bias flow 2.0 L/min.

PC-CMV = pressure control–continuous mandatory ventilation

PIP = peak inspiratory pressure

PC = pressure control level above PEEP

\(T_I\) = inspiratory time

I:E = inspiratory to expiratory ratio

\(T_{tot} = T_I + T_e\)

Decel = decelerating waveform

Table 3. Ventilator Settings: VC-IMV + PS Mode

<table>
<thead>
<tr>
<th>Ventilator</th>
<th>Frequency (ASL/vent), breaths/min</th>
<th>Set (V_T/ASL) (PS)VT, mL</th>
<th>PS, cm H(_2)O</th>
<th>Flow, L/min</th>
<th>Set (T_I/s)</th>
<th>I:E</th>
<th>Waveform</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avea</td>
<td>12/5</td>
<td>700/620</td>
<td>10</td>
<td>43</td>
<td>1.0</td>
<td>1:5.0</td>
<td>Const</td>
</tr>
<tr>
<td>Dräger</td>
<td>12/4</td>
<td>500/390</td>
<td>8</td>
<td>Autoflow</td>
<td>0.70</td>
<td>1:6.0</td>
<td>Decel</td>
</tr>
<tr>
<td>Servo</td>
<td>18/10</td>
<td>500/460</td>
<td>10</td>
<td>52</td>
<td>0.65</td>
<td>1:5.7</td>
<td>Decel</td>
</tr>
</tbody>
</table>

Three ventilator brands that use heated wire humidification or bypassable–valved HME, PEEP 5.0 cm H\(_2\)O, bias flow 2.0 L/min.

VC-IMV + PS = volume control–intermittent mandatory ventilation + pressure support

ASL = rate triggered by ASL test lung

Vent = rate set on ventilator

ASL (PS)VT = tidal volume generated by ASL triggered breaths

I:E = inspiratory to expiratory ratio

\(T_I\) = inspiratory time

Const = constant flow waveform

Decel = decelerating waveform

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### Table 4. Ventilator Settings: PRVC-IMV + PS Mode

<table>
<thead>
<tr>
<th>Ventilator</th>
<th>Frequency (ASL/vent), breaths/min</th>
<th>Set $V_T$/ASL (PS) $V_T$, mL</th>
<th>PS, cm H$_2$O</th>
<th>PIP, cm H$_2$O</th>
<th>Mand $T_i$, s</th>
<th>I:E</th>
<th>Waveform</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aave</td>
<td>15/5</td>
<td>350/310</td>
<td>5</td>
<td>15</td>
<td>0.85</td>
<td>1: 3.7</td>
<td>Decel</td>
</tr>
<tr>
<td>Aave</td>
<td>18/10</td>
<td>500/350</td>
<td>7</td>
<td>12</td>
<td>0.70</td>
<td>1: 3.9</td>
<td>Decel</td>
</tr>
<tr>
<td>Servo</td>
<td>18/10</td>
<td>550/312</td>
<td>5</td>
<td>15</td>
<td>0.9</td>
<td>1: 5.7</td>
<td>Decel</td>
</tr>
</tbody>
</table>

Two ventilator brands that use heated wire humidification or bypassable–valved HME, PEEP 5.0 cm H$_2$O, bias flow 2.0 L/m.

PRVC-IMV + PS = pressure-regulated volume control with synchronized intermittent mandatory ventilation + pressure support

ASL = rate triggered by ASL test lung
PS = pressure support
PIP = peak inspiration pressure
Mand $T_i$ = $T_i$ for set ventilator rate
I:E = inspiratory to expiratory ratio
Const = constant flow waveform
Decel = decelerating waveform

### Table 5. Ventilator Settings: CPAP + PS Mode

<table>
<thead>
<tr>
<th>Ventilator</th>
<th>Frequency (ASL/vent), breaths/min</th>
<th>ASL (PS)$V_T$, mL</th>
<th>PS, cm H$_2$O</th>
<th>$T_i$, s</th>
<th>Waveform</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aave</td>
<td>12/0</td>
<td>280</td>
<td>10</td>
<td>1.08</td>
<td>Decel</td>
</tr>
<tr>
<td>Dräger</td>
<td>12/0</td>
<td>598</td>
<td>10</td>
<td>~1.1</td>
<td>Decel</td>
</tr>
<tr>
<td>Dräger</td>
<td>12/0</td>
<td>475</td>
<td>8</td>
<td>~1.1</td>
<td>Decel</td>
</tr>
<tr>
<td>Servo</td>
<td>12/0</td>
<td>480</td>
<td>8</td>
<td>~1.1</td>
<td>Decel</td>
</tr>
<tr>
<td>Servo</td>
<td>12/0</td>
<td>480</td>
<td>5</td>
<td>~1.1</td>
<td>Decel</td>
</tr>
</tbody>
</table>

Three ventilator brands that use heated wire humidification or bypassable–valved HME, PEEP 5.0 cm H$_2$O, bias flow 2.0 L/m.

CPAP + PS = continuous positive airway pressure + pressure support
ASL = rate triggered by ASL test lung
PS = pressure support
$T_i$ = inspiratory time
Const = constant flow waveform
Decel = decelerating waveform

### Table 6. Ventilator Settings: APRV and APRV (Bi-Vent)

<table>
<thead>
<tr>
<th>Ventilator</th>
<th>Frequency (ASL/vent), breaths/min</th>
<th>$V_T$/ASL (PS) $V_T$, mL</th>
<th>$P_{high}$/$P_{low}$, cm H$_2$O</th>
<th>$T_{high}$, s</th>
<th>$T_{low}$, s</th>
<th>Waveform</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aave</td>
<td>12/0</td>
<td>488/500</td>
<td>25/0</td>
<td>4.4</td>
<td>0.60</td>
<td>ND</td>
</tr>
<tr>
<td>Dräger</td>
<td>12/0</td>
<td>591/537</td>
<td>25/0</td>
<td>4.4</td>
<td>0.60</td>
<td>ND</td>
</tr>
<tr>
<td>Servo</td>
<td>12/0</td>
<td>480/500</td>
<td>25/0</td>
<td>4.4</td>
<td>0.60</td>
<td>ND</td>
</tr>
</tbody>
</table>

Three ventilator brands that use heated wire humidification or bypassable–valved HME, PEEP 5.0 cm H$_2$O, bias flow 2.0 L/m.

* APRV (Bi-Vent).

APRV = airway pressure release ventilation (or BiVent)
ASL = rate triggered by ASL test lung
PS = pressure support
$P_{high}$/$P_{low}$ = tidal volume generated by ASL triggered breaths
$T_{high}$ = inspiratory time
$T_{low}$ = expiratory time or release time
$T_i$ = inspiratory time

**NOTE:** The above tables provide ventilator settings for different modes and brands, along with explanations for abbreviations used. The settings include ventilator frequency, tidal volume, pressure support, inspiratory to expiratory ratio, and waveform types. The tables also note the use of heated wire humidification or bypassable–valved humidification modalities and specify PEEP and bias flow settings along with ventilator brands.
Residual Volume

Residual volume percentage (% neb charge remaining in the nebulizer after treatment completed) was the following: mean SD [all data] 29.6% ± 11.2, no. = 81; humidified 16.3% ± 7.04, no. = 26; and HME 35.9% ± 6.00, no. = 56, with a significant difference between HME and active humidification (P < .001).

Nebulizer Flow: Effect on Ventilator Settings

Data for values of $V_T$ measured for 4 different ventilator settings before and during nebulization are listed in Table 7. All extraneous filters and attachments to the circuit were removed to minimize circuit leaks. $V_T$ was reported in 3 ways: (1) defined by the set volume ($V_T$), (2) monitored by the ventilator (Exhaled $V_T$), and (3) measured directly by the ASL 5000 test lung volume. Added nebulizer flow during breath-actuated nebulization resulted in small, insignificant changes in $V_T$ (mL) measured by both the ventilator (Exhaled $V_T$ =5.0 ± 12.9 mL; $P = .75$) and the ASL test lung volume (25 ± 14.5 mL; $P = .13$).

Particle Distribution

The particle distributions of captured aerosol at the distal tip of the endotracheal tube are shown in Figure 7. The mass median aerodynamic diameter was 1.04 μm in the HME mode and 1.34 μm during active humidification. Approximately 2% of the particles were > 5 μm.

Mass Balance

The distribution of radioactivity throughout the ventilator circuit for active humidification and HME are listed in Table 8. For the same ventilator setting (frequency 18 breaths/min, $V_T$ 500 mL, inspiratory flow 43 L/m, constant wave form), nebulizer residual was reduced during humidification by > 50% (39.4–14.6%). With active humidification, circuit deposition was increased throughout the circuit but, because of the reduced nebulizer residual, IM% was increased (35.0 vs 29.4%). The latter findings were seen for all measurements of nebulizer output reported in Figure 6. Endotracheal tube losses were small, consistent with the terminal particle distribution. Losses on the exhalation filter were relatively low, an effect of breath actuation.
Table 7. Effect of Added Nebulizer Flow on Delivered $V_T$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Set $V_T$, mL</th>
<th>Exhaled $V_T$, mL*</th>
<th>ASL $V_T$, mL†</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V_T = 460$ mL/$T_I = 0.70$ s/DC = 0.20</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No NEB</td>
<td>460</td>
<td>410</td>
<td>426</td>
</tr>
<tr>
<td>With NEB</td>
<td>460</td>
<td>400</td>
<td>447</td>
</tr>
<tr>
<td>$\Delta$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$V_T = 500$ mL/$T_I = 0.70$ s/DC = 0.20</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No NEB</td>
<td>500</td>
<td>460</td>
<td>455</td>
</tr>
<tr>
<td>With NEB</td>
<td>500</td>
<td>460</td>
<td>486</td>
</tr>
<tr>
<td>$\Delta$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$V_T = 690$ mL/$T_I = 0.80$ s/DC = 0.24</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No NEB</td>
<td>690</td>
<td>630</td>
<td>629</td>
</tr>
<tr>
<td>With NEB</td>
<td>690</td>
<td>610</td>
<td>636</td>
</tr>
<tr>
<td>$\Delta$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$V_T = 650$ mL/$T_I = 1.01$ s/DC = 0.33</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No NEB</td>
<td>650</td>
<td>610</td>
<td>593</td>
</tr>
<tr>
<td>With NEB</td>
<td>650</td>
<td>620</td>
<td>634</td>
</tr>
<tr>
<td>$\Delta$</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Set tidal volume (SET $V_T$), monitored by ventilator ($E\;V_T$), measured directly by ASL test lung (ASL $V_T$).

* Average $\Delta\;E\;V_T = -5.0$ mL.

† Average $\Delta\;ASL\;V_T = 25.0$ mL.

$V_T = $ tidal volume

ASL $V_T = $ tidal volume generated by ASL triggered breaths

$T_I = $ inspiratory time

DC = duty cycle

NEB = nebulizer

No NEB = $V_T$ measured when nebulizer bypassed

With NEB = $V_T$ measured during nebulization

Fig. 7. Aerodynamic particle distribution measured via Marple cascade impactor placed in line between the endotracheal tube and inhaled mass filter. Volume control–continuous mandatory ventilation (VC-CMV), rate 18, tidal volume ($V_T$) 500 mL, flow 43 L/min, inspiratory time ($T_I$) 0.7 s, PEEP 5 cm H$_2$O, bias flow 2.0 L/m. The log particle size for each cascade stage was plotted against probability. Mass median aerodynamic diameter (MMAD): bypassable–valved HME = 1.04 µm; active humidification = 1.34 µm. HME = heat and moisture exchanger.
WET-SIDE BREATH-ENHANCED JET NEBULIZATION

Table 8. Mass Balance

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Active Humidification (%)</th>
<th>HME (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nebulizer residual</td>
<td>14.6</td>
<td>39.4</td>
</tr>
<tr>
<td>Inhaled mass filter</td>
<td>35.0</td>
<td>29.4</td>
</tr>
<tr>
<td>Expiratory filter</td>
<td>13.6</td>
<td>9.2</td>
</tr>
<tr>
<td>Ventilator circuit</td>
<td>27.0</td>
<td>13.2</td>
</tr>
<tr>
<td>ETT</td>
<td>2.8</td>
<td>2.7</td>
</tr>
<tr>
<td>Total recovery</td>
<td>93.0</td>
<td>95.2</td>
</tr>
</tbody>
</table>

Tidal volume 500 mL, 18 breaths/min, flow 43 L/min, inspiratory time 0.7 s; PEEP 5 cm H2O, bias flow 2.0 L/min; HME = heat and moisture exchanger

Discussion

This study demonstrated that the i-AIRE nebulizer with the combination of breath enhancement and breath actuation provided predictable drug delivery over a wide range of ventilator settings, regardless of the type of ventilator or mode of humidification. The i-AIRE nebulizer was far more efficient than previously reported jet-driven devices. For example, Miller et al5 reported similar values of IM% for a single ventilator setting when using breath actuation (provided directly by the ventilator) and high nebulizer jet flow (8 L/min), but they reported a treatment time of 1 h for only 2 mL of nebulizer charge. The i-AIRE nebulized 6 mL in the same time period, which suggests at least a 300% increase in output for the i-AIRE system effected by breath enhancement (eg, compared with AeroTech II5 (Biodex Medical Systems, Shirley, New York).

To our knowledge, there are no approved medications for inhalation specifically on a ventilator circuit, and all commonly used drugs are used off label. To develop approved aerosolized antibiotic therapy, our group focused on controlling nebulizer delivery in patients who were intubated because of the general understanding that nebulizer function is affected by many variables. This point is illustrated in Figure 1, which contains output data for many reported studies that contained enough information to calculate effectiveness of a delivery cycle (Ti in general, was not available).1,2,5,8,16–21 As shown in the figure IM% varied widely (2.7-41%), based on nebulizer technology, breathing pattern, position in the circuit, and the presence of humidification.

Without some form of superimposed control, it is easy to see why a dose-dependent drug with a delayed clinical response would be hard to study and to prove efficacy. Previous studies performed at a single site indicated that the ventilator type, use of humidification, and ventilator modes must be rigorously controlled for clinical antibiotic success.5,22,23,27 This type of control was not possible in multi-center studies, and failure to control dose delivery may have contributed to the recent reported failures of multi-center antibiotic protocols.28-30 The i-AIRE system reduces the dosing variability (23-41% vs 2.7-41%) seen with current technologies and functions reliably for most described forms of invasive ventilation across a wide range of commonly used settings, with preservation of dose delivery during active and passive humidification.

Ehrmann et al13 evaluated inspiratory synchronization during ventilator-integrated, breath-actuated nebulization. They assessed nebulizer flow at the beginning and end of inspiration and found that nebulization continued up to 1 s after the end of inspiration, which resulted in as much as 80% of nebulization occurring during the expiratory phase.13 Ehrmann et al13 stated that gas compression and decompression proximal to the nebulizer are probably major determinants of persistent nebulization during expiration. Analysis of their data suggests that pressure in the nebulizer line due to the high resistance in the nebulizer orifice prevents closure of the ventilator solenoid.13 Failure of solenoid closure in the ventilator allows nebulization to continue during expiration. To prevent this phenomenon, our system used an independent solenoid that actively opened and closed, which prevented increased expiratory losses.

Jet nebulizers require a defined flow of gas and pressure to generate aerosol. Commercial ventilators do not universally provide breath actuation or standardized flow and pressure.12 The i-AIRE system relies on wall gas and pressure, and a standardized nebulizer flow of 3.5 L/m, avoiding unpredictable nebulizer-ventilator combinations. Jet nebulizers powered from hospital wall gas may affect ventilator displays because nebulizer gases enter the expiratory line. However, the brief introduction of nebulizer gas during breath-actuated inspiration was not readily detected (Table 7). The average added VT to the circuit of 25 mL is 5% of a 450 mL VT; this added volume is much less than that resulting from a conventional jet nebulizer, which runs at 6–8 L/m at any VT.

Added humidity has been reported to be an unpredictable factor that affects aerosol delivery.2,5,7,16,31-33 In early studies, our group was one of the first to report losses in aerosol delivery during active humidification, of ~50%.3,7 In the present study, when using the i-AIRE nebulizer and specialized wet-sided nebulization, we observed similar dose delivery between both active humidification and HME. Analysis of these data indicates that dosing of the patient would not be significantly affected with i-AIRE, with or without active humidification. The mass balance analysis and particle distribution data show that wet-side humidification significantly increased nebulizer output. As shown in the mass balance data, with humidification there is a decrease in residual volume.

Most jet nebulizers operate by using dry gas. The residual volume represents liquid that contains a drug that coats the walls of the nebulizer with some liquid drying on the
walls and failing to be recycled into the nebulizer bowl. The i-AIRE system, ventilated with humidified air, demonstrates reduced wall losses, which makes more liquid available for nebulization. The increase in nebulizer efficiency is balanced by increased losses in the circuit (likely due to particle growth). These effects result in similar IM% and mass median aerodynamic diameter, with and without active humidification.

The effect of bias flow on aerosol delivery has been previously studied. Miller et al. measured an effect with the nebulizer located in the inspiratory limb. However, they tested bias flows of 10–15 L/m. Ari et al. assessed the effect of 2.0 and 5.0 L/m in 2 circuit positions that compared a jet nebulizer and a vibrating mesh nebulizer. This group found that bias flow had a statistically significant effect when placed 15 cm from the Y-piece in the inspiratory limb but not when placed close to the ventilator. In the current study, 2.0 L/min was used for all routine experiments because this is a common default value on adult ventilators that both allow for adjustment of bias flow and with a fixed bias flow. The effect of bias flow (0.5, 3.5, and 5 L/m) was tested by using an Aeva ventilator, which provided for adjustment of bias flow. No effect was detected on IM%.

The nebulizer-circuit combination is a prototype design and subject to untested clinical variables. We did not test all commercially available ventilators and clinically relevant situations. This study established a potential design that, if carried forward, may allow therapeutic trials of important drugs whose clinical effects cannot be assessed in real time at the bedside.

Conclusions

Because aerosol delivery device-mechanical ventilator combinations are not standardized for drug delivery, there can be substantial variation among practitioners in the practice of aerosol administration. The i-AIRE system minimizes interprovider variability, with a nebulizer-circuit combination that provides predictable drug delivery, with and without active humidification, and avoiding potential humidifier contamination.

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REFERENCES


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