Effect of High-Frequency Oscillations on Cough Peak Flows Generated by Mechanical In-Exsufflation in Medically Stable Subjects With Amyotrophic Lateral Sclerosis

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BACKGROUND: Mechanically assisted coughing with mechanical in-exsufflation (MI-E) is recommended for noninvasive management of respiratory secretions in amyotrophic lateral sclerosis (ALS). To improve the effectiveness of the technique, a new device combining MI-E with high-frequency oscillations (HFO) has been developed. This work aimed to assess the effect of HFO on the cough peak flow generated by MI-E in medically stable subjects with ALS.

METHODS: This was a prospective study that included subjects with ALS in a medically stable condition. Cough peak flow generated by MI-E was measured in 4 situations: without HFO, with HFO during insufflation, with HFO during exsufflation, and with HFO in both cycles. The parameters used were: insufflation pressure of \(40 \text{ cm H}_2\text{O}\), exsufflation pressure of \(40 \text{ cm H}_2\text{O}\), insufflation time 2 s, exsufflation time 3 s, amplitude of oscillations 10 cm H\(_2\)O, and frequency of oscillations 15 Hz.

RESULTS: Forty-seven subjects with ALS were included: 66% males, 68.2 ± 9.2 y, 40% with bulbar onset, FVC 1.7 ± 1.1 L, percent-of-predicted FVC = 54.4 ± 26.6%, cough peak flow = 3.8 ± 2.2 L/s, \(P_{\text{max}}\) = −39.4 ± 26.4 cm H\(_2\)O, revised ALS scale = 28.5 ± 9.3, Norris bulbar subscore = 26.1 ± 10.4. No statistical differences were found in cough peak flow generated by MI-E in the 4 situations (without HFO = 4.0 ± 1.2 L/s, with insufflation HFO = 3.9 ± 1.2 L/s, with exsufflation HFO = 4.1 ± 1.2 L/s, with in-exsufflation HFO = 3.9 ± 1.1 L/s).

CONCLUSIONS: The addition of HFO to mechanically assisted coughing with MI-E does not have an effect on the cough peak flow of medically stable subjects with ALS.

Key words: cough-assist, amyotrophic lateral sclerosis, neuromuscular disease, respiratory failure [Respir Care 0;0(0):1–. © 0 Daedalus Enterprises]
chospopy or even a tracheostomy. Coughing aids can be both manually assisted and mechanically assisted. Mechanically assisted coughing techniques use a mechanical device that generates positive pressure and provokes the insufflation of the lungs. This is followed almost immediately by an exsufflation with the application of negative pressure. This sudden and rapid change from positive to negative pressure generates air flows during the exsufflation phase, which are capable of clearing the respiratory secretions. However, despite the fact that various studies have demonstrated the usefulness of mechanically assisted coughing, there are still some important issues that are yet to be resolved: Day-to-day clinical practice has shown that the clearance of thick secretions adhering to the tracheobronchial walls is problematic for mechanically assisted coughing. To address this problem, a new device (Cough-Assist E70, Philips Respironics, Murrysville, Pennsylvania) has recently been designed, which combines mechanical in-exsufflation (MI-E) with high-frequency oscillatory vibrations (HFO). During the cycles of insufflation, exsufflation, or both, this device applies high-frequency oscillatory vibrations, generated by air pulses, to the positive or negative pressures or to both. The frequency settings of these oscillations range from 1 to 20 Hz, and the amplitude settings range from 1 to 10 cm H2O. Theoretically, HFO may facilitate the loosening of these thick secretions to the bronchial wall, enabling them to be cleared. The application of HFO to the airway has been shown to change the viscoelastic properties of the secretions, making them more mobile. Therefore, the addition of HFO to MI-E may be able to facilitate the removal of these tenacious secretions in a noninvasive way during respiratory tract infections in patients with ALS, avoiding the need to resort to invasive methods to expel such secretions.

However, bulbar dysfunction is the main cause of ineffective cough peak flow generated by MI-E in patients with ALS, and if the main bulbar dysfunction derives from upper motor neuron damage, hyperreflexia episodes in the upper airway may be triggered by mechanical stimuli such as HFO; this could interfere with the effectiveness of the cough peak flow generated. Therefore, before any evaluation of the clinical effectiveness of the device, the aim of this study was to assess whether the addition of HFO to MI-E could modify the cough peak flow generated by mechanically assisted coughing in medically stable subjects with ALS, either positively or negatively, and whether the oscillations strengthen the effect of ALS patient bulbar dysfunction on the cough peak flow generated.

**Methods**

We performed a prospective study from April 2013 to April 2015 that included all subjects diagnosed with ALS according to the Revised El Escorial criteria who were managed at the Respiratory Care Unit at the Hospital Clínico Universitario (Valencia, Spain). All subjects had to have been in a medically stable condition in the 3 months before inclusion in the study. Informed consent was obtained, and the protocol was approved by the hospital’s ethics committee. Exclusion criteria were: refusal to participate in the study, the presence of bronchial disease, a contraindication to the use of MI-E, or the presence of severe frontotemporal dementia associated with ALS that could interfere with the procedures. After inclusion in the study, the subjects underwent a clinical evaluation, mainly oriented toward assessing bulbar impairment, and a functional respiratory evaluation, which included spirometry, the measurement of respiratory muscle strength, and an evaluation of cough capacity.

**Clinical Assessment**

Clinical assessment data were collated using the revised ALS scale, and bulbar dysfunction was evaluated in accordance with the Norris scale bulbar subscore.

**Functional Assessment**

Spirometry was performed (MS 2000 spirometer, C. Schatzman, Madrid, Spain) using a mouthpiece and a nose clip. To prevent air leaks from the mouthpiece for those subjects with severe facial weakness, the respiratory function test was performed using an oronasal mask (Matin Vecino, Madrid, Spain). FVC, FEV1, FEV1/FVC, and supine FVC were recorded in accordance with European...
Respiratory Society guidelines and suggested reference values. The maximum inspiratory pressure and the maximum expiratory pressure were measured in the mouth (Electrometer 78.905, Hewlett-Packard, Andover, Massachusetts) while the check valve was held closed. The maximum inspiratory pressure was recorded close to residual volume, and maximum expiratory pressure was recorded close to total lung capacity. The pressures sustained for 1 s were measured. Three measurements with < 5% variability were recorded, and the highest value was used for the data analysis. Sniff nasal inspiratory pressure was measured in an occluded nostril during a maximal sniff through the contralateral nostril (Micro RPM, MicroMedical Ltd, Rochester, Kent, United Kingdom). Sniff nasal inspiratory pressure was measured during 10 maximal sniffs performed at functional residual capacity, and the highest recorded pressure was used.

Cough peak flow was measured using an oronasal mask (Martin Vecino, Madrid, Spain) connected to a pneumotachograph spirometer (MS 2000) when the subjects performed a maximal cough effort after a deep inspiration. The highest cough peak flow measurement obtained from ≥3 cough maneuvers with <5% variability was recorded. The peak velocity time, or time necessary to reach cough peak flow, was also recorded. Maximum insufflation capacity by air stacking was achieved by the subject taking a deep breath, holding it, and then air-stacking consecutively delivered volumes of air from a manual resuscitator (Revivator, Hersill, Madrid, Spain) through the oronasal mask up to the maximum volume that could be held with a closed glottis. The subject then exhaled the maximally held volume of air into the pneumotachograph for volume measurement. Manually assisted cough peak flow was also measured with the pneumotachograph connected to the mask and the manual resuscitator.

Interventions

The Cough-Assist E70 (Philips Respironics, Murrysville, Pennsylvania) was used with an oronasal mask (Martin Vecino) and set at an insufflation pressure of 40 cm H₂O and an exsufflation pressure of ~40 cm H₂O, an insufflation/exsufflation ratio of 2:3, and a pause of 1 s after each cycle. The subjects were asked to try to keep their airways open but to otherwise remain passive and let the Cough-Assist device act unimpeded. When HFO were applied, they were set with a frequency of 15 Hz and an amplitude of 10 cm H₂O. A pneumotachograph was placed between the mask and the MI-E circuit, and the cough peak flow generated by MI-E, the insufflation volume, and the exsufflation volume were measured. Three measurements with < 5% variability were recorded, and the highest value was used for the data analysis. Four sessions of MI-E (without oscillation, with oscillations only during insufflation, with oscillations only during exsufflation, and with oscillations during both insufflation and exsufflation) were carried out in a random order using a simple randomization method (random number generator by a computer). Washout time sessions of 30 min were applied to avoid tiredness and hyperventilation of the subject. The acceptability of each MI-E session was measured using a visual analog scale. Subjects were asked to provide a rating in response to the question “How comfortable is this treatment for you?” by making a mark on an uncalibrated 10-cm horizontal line, the left end of which was labeled as “uncomfortable” and the right end of which was labeled “very comfortable.” Graphs (flow, pressure, and volume) produced by the device software were analyzed (Direct-View. Philips Respironics. Murrysville, Pennsylvania) (Fig. 1), and cough peak flow (unassisted and assisted) was considered ineffective when values were <2.7 L/s.

Statistical Analysis

Assuming a risk of 5% and a power of 90% to detect differences and basing in our previous results on the SD in cough peak flow generated by MI-E in medically stable subjects and the mean changes produced in cough peak flow generated by MI-E when set parameters are changed, a minimum sample size of 26 subjects was calculated as necessary to detect differences.

Data were expressed as mean ± SD for continuous normally distributed variables and as frequency counts and percentages for binary and categorical variables. Data comparisons were performed using the Student t test and Mann-Whitney test for normally and non-normally distributed data, respectively; categorical data were compared using the chi-square test. Comparisons for cough peak flow generated by MI-E, insufflation volume, and exsufflation volume between the 4 different sessions of MI-E were performed with an analysis of variance for repeated measurements. Those factors that could influence the effectiveness of mechanically assisted coughing, such as bulbar dysfunction (ALS onset, Norris scale bulbar subscore, revised ALS scale) and the use of mechanically assisted coughing at home, were considered as covariates in the analysis. The level of statistical significance was set at P < .05.

Results

Forty-seven consecutive subjects with ALS in a medically stable condition were enrolled in the study. Time from ALS diagnosis to commencement of the study was 44.8 ± 50.2 months. Twenty-one subjects (44.7%) used noninvasive ventilation and 22 (46.8%) used mechanically assisted coughing previously at home. Data on demographics, respiratory function, and functional assessment are
shown in Table 1. In the spinal onset group, 71.4% of the subjects presented an effective cough peak flow, and in the bulbar onset group, 31.5% presented an effective cough peak flow ($P = 0.004$).

In the analysis of variance, no statistical differences were found in cough peak flow generated by MI-E, insufflation volume, and exsufflation volume between the 4 sessions tested (no oscillation, with oscillation during insufflation, with oscillation during exsufflation, and with oscillation in both phases) (Table 2). No statistical differences were found in cough peak flow generated by MI-E during the 4 sessions when factors related to the severity of bulbar dysfunction were used as covariates (bulbar onset, $P = .58$; revised ALS scale, $P = .79$; Norris scale bulbar subscore, $P = .99$). Using as covariate the presence of an effective cough peak flow, no statistical differences in cough peak flow generated by MI-E were found between the 4 tested sessions ($P = .95$). No statistical differences were found in the acceptability of each session measured using a visual analog scale (without HFO, 6.6 ± 2.2 cm; HFO during insufflation, 6.4 ± 2.2 cm; HFO during exsufflation, 6.1 ± 2.3 cm; HFO in both phases, 6.1 ± 2.1 cm, $P = .35$). The acceptability of each session was compared with the severity of bulbar dysfunction and previous use of mechanically assisted coughing at home, and no significant differences were found (bulbar onset, $P = .23$; revised ALS scale, $P = .81$; Norris scale bulbar subscore, $P = .50$; previous mechanically assisted coughing at home, $P = .92$).

At the beginning of the exsufflation cycle, a large increase in the flow wave reaching a maximal value was produced; this maximal value represents cough peak flow generated by mechanical in-exsufflation; arrow indicates elastic recoil. Fig. 1. Flow and pressure waves recorded by the device during the 4 sessions. A = without oscillations; B = oscillations during insufflation; C = oscillations during exsufflation; D = oscillations during insufflation and exsufflation. * = cough peak flow generated by mechanical in-exsufflation; arrow indicates elastic recoil.

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Table 1. Demographic and Pulmonary Function Parameters

<table>
<thead>
<tr>
<th></th>
<th>Total Population (N = 47)</th>
<th>Spinal Onset (n = 28)</th>
<th>Bulbar Onset (n = 19)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male/female sex, n</td>
<td>31/16</td>
<td>23/5</td>
<td>8/11</td>
<td>.008</td>
</tr>
<tr>
<td>Age, mean ± SD y</td>
<td>68.2 ± 9.2</td>
<td>67.0 ± 10.9</td>
<td>70.1 ± 6.0</td>
<td>.28</td>
</tr>
<tr>
<td>BMI, mean ± SD kg/m²</td>
<td>26.1 ± 4.4</td>
<td>27.1 ± 5.2</td>
<td>24.9 ± 2.8</td>
<td>.10</td>
</tr>
<tr>
<td>FVC, mean ± SD L</td>
<td>1.7 ± 1.1</td>
<td>2.0 ± 1.0</td>
<td>1.4 ± 1.1</td>
<td>.07</td>
</tr>
<tr>
<td>FVCs, mean ± SD % predicted</td>
<td>54.4 ± 26.6</td>
<td>59.6 ± 27.7</td>
<td>48.1 ± 24.5</td>
<td>.16</td>
</tr>
<tr>
<td>MIC, mean ± SD L</td>
<td>2.0 ± 1.0</td>
<td>2.31 ± 1.0</td>
<td>1.65 ± 0.9</td>
<td>.039</td>
</tr>
<tr>
<td>Cough peak flow, mean ± SD L/s</td>
<td>3.8 ± 2.2</td>
<td>4.5 ± 2.3</td>
<td>2.8 ± 1.7</td>
<td>.01</td>
</tr>
<tr>
<td>Peak value time, mean ± SD s</td>
<td>0.2 ± 0.3</td>
<td>0.1 ± 0.0</td>
<td>0.3 ± 0.5</td>
<td>.16</td>
</tr>
<tr>
<td>Cough peak flow generated by MI-E, mean ± SD L/s</td>
<td>3.8 ± 2.2</td>
<td>4.6 ± 2.2</td>
<td>2.8 ± 1.8</td>
<td>.008</td>
</tr>
<tr>
<td>Pmax, mean ± SD cm H₂O</td>
<td>−39.4 ± 26.4</td>
<td>−44.5 ± 29.8</td>
<td>−32.4 ± 19.5</td>
<td>.14</td>
</tr>
<tr>
<td>SNIP, mean ± SD cm H₂O</td>
<td>−35.4 ± 24.7</td>
<td>−42.5 ± 27.9</td>
<td>−25.5 ± 14.7</td>
<td>.01</td>
</tr>
<tr>
<td>Pmax, mean ± SD cm H₂O</td>
<td>68.1 ± 58.3</td>
<td>82.1 ± 69.5</td>
<td>48.2 ± 28.3</td>
<td>.064</td>
</tr>
<tr>
<td>ALSFRS-R, mean ± SD</td>
<td>28.5 ± 9.3</td>
<td>29.1 ± 9.6</td>
<td>27.8 ± 9.2</td>
<td>.64</td>
</tr>
<tr>
<td>NBS, mean ± SD</td>
<td>26.1 ± 10.4</td>
<td>31.3 ± 7.0</td>
<td>18.7 ± 10.0</td>
<td>&lt;.001</td>
</tr>
</tbody>
</table>

BMI = body mass index  
FVCs  = supine FVC  
MIC  = maximal insufflation capacity  
MI-E = mechanical in-exsufflation  
Pmax  = maximal inspiratory pressure  
SNIP  = sniff inspiratory pressure  
Pmax  = maximal expiratory pressure  
ALSFRS-R  = revised amyotrophic lateral sclerosis functional rating score  
NBS  = Norris scale bulbar subscore

Table 2. Data Generated by Mechanical In-Exsufflation During the 4 Sessions

<table>
<thead>
<tr>
<th></th>
<th>Without HFO</th>
<th>HFO During Insufflation</th>
<th>HFO During Exsufflation</th>
<th>HFO During In-Exsufflation</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cough peak flowMI-E, mean ± SD L/s</td>
<td>4.0 ± 1.2</td>
<td>3.9 ± 1.2</td>
<td>4.1 ± 1.2</td>
<td>3.9 ± 1.1</td>
<td>.72</td>
</tr>
<tr>
<td>V insufflation, mean ± SD L</td>
<td>2.2 ± 1.2</td>
<td>2.2 ± 1.2</td>
<td>2.2 ± 0.9</td>
<td>2.2 ± 1.2</td>
<td>.81</td>
</tr>
<tr>
<td>V exsufflation, mean ± SD L</td>
<td>2.7 ± 1.2</td>
<td>2.5 ± 1.2</td>
<td>2.5 ± 1.3</td>
<td>2.4 ± 1.2</td>
<td>.68</td>
</tr>
</tbody>
</table>

HFO  = high-frequency oscillation  
Cough peak flowMI-E  = generated by mechanical in-exsufflation  
V insufflation  = volume generated during insufflation  
V exsufflation  = volume generated during exsufflation

Discussion

The findings of this study show that the addition of HFO to mechanically assisted coughing with MI-E does not have an effect on the cough peak flow of medically stable subjects with ALS, regardless of the severity of bulbar dysfunction. Moreover, we found that the presence of air leaks can interfere with the application of oscillations during the insufflation cycle.

The usefulness and effectiveness of mechanically assisted coughing have been proven in the noninvasive management of respiratory secretions in patients with ALS, and it is the recommended treatment for those ALS patients with an ineffective unassisted cough. However, mechanically assisted coughing is not always an effective technique. The response of the upper airway when there is severe bulbar impairment completely neutralizes the effectiveness of the procedure in clinical practice, and clinical practice also demonstrates that the increase in secretions and the changes in viscoelastic properties during a respiratory tract infection can also reduce the effectiveness of mechanically assisted coughing, making fibrobronchoscopy necessary for the removal of the secretions.

Although mechanically assisted coughing’s effectiveness derives from the quick shift from positive to negative pressure during cycling from insufflation to exsufflation and the large air flows produced, the manufacturer of the Cough-Assist E70 (Philips Respironics) hypothesizes that adding oscillations can contribute to the removal of tenacious secretions, facilitate the process of moving them
toward the central airways, and improve the amount ultimately expelled.20 A review of the general literature shows that, within the range of the respiratory pathologies in which hypersecretion of mucus occurs, physical therapies using HFO have been used to mobilize and remove airway secretions.21,22 In certain patients, these oscillations may change the viscoelastic properties of the secretions, making it easier to move the mucus up to the proximal airway, where secretion removal is then effected by coughing. In fact, oscillations are not a substitute for an effective cough, but rather their use is meant to enhance the effectiveness of coughing.23 This is supported by our current findings because the analysis of the flow waves generated during the exsufflation cycles revealed that the oscillations were unable to reach the necessary magnitude to create an effective cough flow for the removal of the respiratory secretions. Thus, in neuromuscular patients with weak respiratory muscles, other techniques must be used to clear secretions once they have been mobilized centrally by the oscillations.24,25 Two previous studies26,27 have assessed the effect of high-frequency chest wall oscillations in subjects with ALS. Although the results are not conclusive, both studies show that in those subjects with a decreased cough capacity, respiratory secretions cannot be managed with oscillations alone.

With regard to the use of the device at the bedside, an important finding of the present study is that a delay between the beginning of the insufflation cycle and the onset of the oscillations occurs. When the cycle starts, all of the flow is dedicated to delivering the set pressure because this is considered to be the most important factor, and then, once the set pressure has been reached, oscillations start to be generated. Thus, in the presence of air leaks, the set pressure is reached later, making the delay in the commencement of the oscillations greater. A recent study by Frigerio et al28 found that the presence of air leaks can affect the performance of conventional MI-E. Another feature seen in the flow wave analysis and described previously18 is that at the beginning of the pause after the end of exsufflation cycle, a drop in the flow wave occurs due to the elastic recoil of the lung.

Taking into account that bulbar impairment can lead to the passive collapse of the upper airway during the exsufflation cycle in subjects with ALS who are managed with mechanically assisted coughing,7 neutralizing its potential therapeutic effect, a question that needs to be addressed is whether the addition of HFO might in fact increase the harmful effect of bulbar impairment on the cough peak flow generated by MI-E. Furthermore, if the bulbar dysfunction derives predominantly from damage to the upper motor neurons, then the bulbar symptoms will be characterized by hyperreflexia episodes in the upper airways.11 Thus, from a theoretical point of view, the possibility exists that the HFO could trigger mechanical receptors and induce upper airway reflexes that may impair the performance and the effectiveness of mechanically assisted coughing. Evidence against this hypothesis (and in favor of the device) can be seen in our results because they show that, regardless of the severity of bulbar dysfunction as measured in terms of different parameters, the addition of HFO to mechanically assisted coughing did not produce a change in the cough peak flow generated, and the effectiveness of this technique did not decrease in the ALS subjects studied.

Another finding of the present study is the good level of subject tolerance of the addition of HFO to mechanically assisted coughing. The overall acceptability of the technique was similar with and without oscillations and was not affected by the severity of bulbar dysfunction or the previous use of mechanically assisted coughing at home.

This study has some limitations. The main objective of the present study conducted with medically stable subjects...
with ALS unencumbered with secretions was to evaluate the response of the mechanically assisted coughing-generated cough peak flow when HFO were added to MI-E. This is the first protocol from a long-term study in which this technique will be tested in patients with ALS who have an acute respiratory infection. Thus, the results must be taken with caution with regard to acute respiratory episodes, because subjects with ALS suffer an increase in respiratory muscle weakness, a decrease in thoracopulmonary compliance and an increase in airway resistance during a lower respiratory tract infection.8 Indeed, these features may lead to the effect of MI-E with HFO being different from the one we have observed here. Another limitation concerns our use of a single value for the frequency and amplitude of the oscillations; we chose these values because they are the ones that are most often employed with the different devices using high-frequency oscillations.29 Future studies will be necessary to evaluate the effect of different frequencies and amplitudes of high-frequency oscillations added to MI-E. Finally, the lack of a separate control group represents another limitation; instead of using a group of healthy subjects, our control group consisted of the data from the session of MI-E without HFO.

Conclusions

In subjects with ALS in a medically stable condition, the cough peak flow generated by MI-E, independent of the severity of bulbar dysfunction, does not change despite the use of HFO during insufflation, during exsufflation, or in both cycles. The presence of air leaks can delay the commencement of the oscillations in the insufflation cycle to a considerable extent. Whether the addition of HFO to mechanically assisted coughing with MI-E will positively impact cough peak flow during acute respiratory infections and improve long term outcomes is still to be determined.

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