ASSESSMENT OF MAXIMUM INSPIRATORY PRESSURE IN NON-COOPERATIVE CRITICAL PATIENTS: COMPARISON BETWEEN TWO METHODS

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ABSTRACT

Background: Although mechanical ventilation is necessary for treating acute respiratory insufficiency, it may be associated with deconditioning and respiratory muscle dysfunction. Maximal inspiratory pressure (MIP) evaluation is used to estimate inspiratory muscle strength in artificially ventilated patients, but there is no definition as to the best way to make this measurement. Objective: To compare two methods for MIP evaluation, using four different protocols, among non-cooperative artificially ventilated patients. Method: Thirty non-cooperative patients undergoing the process of weaning off mechanical ventilation were evaluated. In accordance with block randomization, the simple occlusion method (OM) or the unidirectional valve method (UV) was applied to each patient for time periods of 20 and 40 seconds. Additionally, during the 40s measurements, the MIP value at 30s was recorded. Results: The MIP values were higher at 40s than at 20s, both from OM (48.2 ± 21.7 vs. 36 ± 18.7 cmH2O; p< 0.001) and from UV (56.6 ± 23.3 vs. 43.4 ± 24 cmH2O; p< 0.001) and from UV (56.6 ± 23.3 vs. 43.4 ± 24 cmH2O; p< 0.001). The MaxIP values were higher from UV at 40s (OM40) than from OM at 40s (OM40) (56.6 ± 23.3 vs. 48.2 ± 21.7 cmH2O; p< 0.001). There was a difference between UV at 30 and 40s (51.5 ± 20.8 vs. 56.6 ± 23.3 cmH2O; p< 0.001). Conclusion: Among non-cooperative patients, higher MIP values were obtained from the unidirectional valve method with 40s of occlusion than from the other protocols evaluated.

Key words: respiratory muscles; evaluation; weaning from respirator; physical therapy.

INTRODUCTION

It is well established that mechanical ventilation is an essential therapy for patients with acute respiratory insufficiency. In such a context, ventilation support is necessary when ventilation demand becomes superior to the capacity of respiratory muscles as a consequence of several clinical conditions1. Mechanical ventilation reduces or eliminates overload of respiratory muscles. However, it can also be associated with deconditioning and respiratory muscle dysfunction, along with other factors, such as polyneuropathy of the critical patient, sepsis and dysfunction of multiple organs and systems2,3. When problems that produced increased ventilation demand are solved, weaning must be initiated. However, in many patients respiratory muscle weakness can impede withdrawal from mechanical ventilation4. Assessment of patients before weaning is important to avoid respiratory muscle fatigue and to define the most appropriate strategies of withdrawal from mechanical ventilation. The most common form of assessment of respiratory muscle function in critical patients is the measurement of maximum inspiratory pressure (MaxIP)5,6.

The usefulness of MaxIP measures in mechanically ventilated patients has been demonstrated through use of MaxIP values to predict success of withdrawal from mechanical ventilation. In 1962, Bendixin and Bunker1 associated the measurement of maximum inspiratory pressure with ventilatory reserve in anesthetized dogs. Later, Wescott and Bendixin2 verified that maximum inspiratory pressure values between –20 and –30 cmH2O obtained in patients under neuromuscular blockade were sufficient to consider transferring the patients to the post-anesthetic recovery room. After such papers, the use of this type of measurement for patients in the weaning phase was established in the classical study of Sahn and Lakshminarayan3. The authors observed that patients with MaxIP values lower than – 30 cmH2O...
presented good prognosis for withdrawal from mechanical ventilation, while patients with MaxIP values greater than – 20 cmH₂O failed to wean off. Due to several factors such as study design (prospective or retrospective) weaning method and definition of success and failure, the accurateness of MaxIP measures varies considerably between different studies. In general, studies report sensibility over 80% for cutoff values of – 30 cmH₂O⁹,¹⁰,¹¹, - 20 cmH₂O¹⁰,¹¹,¹²,¹³ and even - 15 cmH₂O¹¹. Because success of withdrawal depends on other factors, specificity is low in all studies. The low specificity indices indicate that patients who fail weaning from mechanical ventilation do not necessarily present reduced MaxIP.

In an attempt to make MaxIP measures reliable, Marini et al.¹⁴ described the unidirectional valve method after the assessment of 20 artificially ventilated patients. After a short period of spontaneous ventilation, a device that allowed exhalation only was attached to the airway opening (unidirectional valve). This way, patients were forced to make progressively increasing effort while approaching residual volume. At residual volume MaxIP corresponded to maximum the capability of the patient. The maximum value was reached in 15 or 20 seconds of occlusion. In spite of this result, MaxPI values measured in critical patients are usually underestimated and have poor reliability¹⁵,¹⁶. The identification of reliable methods of assessment of respiratory muscle strength can contribute to better monitoring of withdrawal from mechanical ventilation as well as to better definition of treatment strategies. The objective of the present study was to compare two methods of assessment of maximum inspiratory pressure implemented trough four different protocols, in non-cooperative patients receiving artificial ventilation.

MATERIALS AND METHODS

Sample
Thirty patients admitted to the Intensive Care Unit of Rio de Janeiro Military Police Central Hospital (Table 1) were assessed in a cross-over randomized trial. Patients included in the study were receiving invasive ventilatory assistance and were in process of weaning from mechanical ventilation. Individuals with hemodynamic instability, PaO₂/FiO₂ ratio inferior to 200, intracranial hypertension (ICP > 20 mmHg), deep sedation (Ramsay ≥5), curarization, abdominal surgeries with risk of evisceration, coronary arterial disease or severe cardiac insufficiency were not included in the study. Sample size was calculated with the software SigmaStat 3.1, according to results of Caruso et al.¹⁷, considering the statistical test ANOVA, power of 80%, significance level of 5%, expected difference of 28% and standard deviation of 30%. The estimated sample size was 23 individuals.

The research project was approved by the Augusto Motta University Centre Ethics Committee (approval report number 04/06). Relatives responsible for all patients included in this study signed the informed consent form, according to the CONEP 196/96 resolution.

Procedures
Patients were positioned in dorsal decubitus with the head of the bed elevated at 45° for the measurement of maximum inspiratory pressure with the Simple Occlusion Method (OM). The cuff was hyperinsufflated to avoid escape during measurement. After tracheal aspiration, patients rested for five minutes while connected to the mechanical ventilator. To perform the measurement, the mechanical ventilator was disconnected and after 10 seconds a connector attached to the artificial airway was manually occluded at the end of a normal expiration (functional residual capacity level). MaxIP was measured with an Instrumentation Industries analogical manometer, with a measurement range of ± 120 cmH₂O at 2 cmH₂O steps. Occlusion was maintained for 20 or 40 seconds, according to randomization. The maximum value of each measurement was used for analysis. Three measurements with 2- minute intervals were performed (during intervals the patient was connected to the mechanical ventilator to rest). In order to measure MaxIP with the Unidirectional Valve Method (UV) the same procedure was adopted. At the moment of measurement, however, a low

Table 1. Data related to age and PaO₂/FiO₂ are presented as mean and standard deviation. The severity score (APACHE II) is presented as median (minimum – maximum). M: Male, F: Female, PaO₂/FiO₂: arterial oxygen partial pressure to inspired oxygen fraction ratio, APACHE II: severity score.

<table>
<thead>
<tr>
<th>Subjects Characteristics</th>
<th>Sex</th>
<th>Age (years)</th>
<th>PaO₂/FiO₂</th>
<th>APACHE II</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td>16 M e 14 F</td>
<td>62.6 ± 17.7</td>
<td>334.4 ± 87.8</td>
<td>16 (10-26)</td>
</tr>
</tbody>
</table>

Admission Cause
resistance unidirectional valve (that allowed exhalation only) was connected to the airway opening.

Procedures were performed according to randomization in blocks of 10, considering two techniques of measurement of maximum inspiratory pressure and measurement times (20 or 40 seconds). After the order of measurement techniques (OM or UV) was determined, the respective measurement times were randomized (20 or 40 seconds). The order of measurement times was used for the two measurement techniques for the same patient. Therefore, all patients were assessed with all techniques and times, but in different order according to randomization. Additionally, during the 40-second measurements, maximum inspiratory pressure values at 30 seconds were recorded, without interruption of the test. The two forms of assessment were separated by an interval of thirty minutes, during which patients remained connected to the mechanical ventilator. Ventilatory parameters were not modified during the assessment protocol, and patients were being monitored for cardiac function and oxygen saturation. All assessments were performed by the same examiner.

Statistical analyses were performed with the software SigmaStat 3.1. The Friedmand test was used to compare MaxIP values between the different techniques and measurement times. The Paired t Test was used for the post hoc analysis to compare results of the UV method at 30 and 40 seconds. Differences were considered significant at p< 0.05. Pearson or Spearman tests were used to test for correlations, depending on the type of variable analyzed and results of normality tests.

**RESULTS**

Data regarding patients included in this study are available in Table 1. No differences were found for MaxPi values between OM at 20s (OM20) and UV at 20s (UV20) (36 ± 18.7 and 43.1 ± 24 cmH2O; p> 0.05). Values at 40s were greater than at 20s for both the OM (48.2 ± 21.7 and 36 ± 18.7 cmH2O; p< 0.001) and UV (56.6 ± 23.3 and 43.4 ± 24 cmH2O; p< 0.001) methods (Figure 1). MaxPI values were greater with the UV at 40s (UV40) than with the OM at 40s (OM40) (56.6 ± 23.3 vs 48.2 ± 21.7 cmH2O; p< 0.001) (Figure 1). Considering the three measurements performed to obtain the final value of each patient, the mean coefficient of variation was calculated for each protocol: OM20= 0.2; OM40= 0.1; UV20= 0.16; UV= 0.13. The pos hoc analysis revealed a significant difference between UV at 30 and UV at 40s (51.5 ± 20.8 and 56.6 ± 23.3 cmH2O; p< 0.001) (Figure 1). A significant correlation between patient age and MaxIP value measured with UV40 was found (r= -0.74; p< 0.001) (Figure 2). However, no correlation was found between protocols and severity scores (APACHE). MaxIP mean was lower in females (42.3 cmH2O) than in males (69.1 cmH2O). Considering the Black & Hyatt18 equation for the calculus of predicted MaxIP for each individual of the sample, mean percents of predicted values were calculated for men (60.4 %) and women (62.5 %), with no significant differences between sexes (p= 0.36). Additionally, no significant differences were found between mean percents of predicted values for patients admitted for clinical or surgical reasons (60.9 % and 61.8 %; p= 0.44).

**DISCUSSION**

Assessment of muscle strength is a common and necessary practice for various physical therapy interventions. Regarding weaning from mechanical ventilation, respiratory muscle function may be affected, and this may cause failure in the process of withdrawal from mechanical ventilation. Due to the high sensitivity to predict success at weaning, the measurement of MaxIP is commonly performed, although the accurateness and reliability of the procedure are questionable. The unidirectional valve method proposed by Marini et al.14 was found to be more reliable and capable of detecting higher MaxPI values compared to the simple occlusion method at the end of expiration. In this study, in opposition to the study by Marini et al.14, no significant differences were observed between the two methods in the assessment performed at 20s. Values registered with the simple occlusion method in the present study were on average 19.39% lower than values registered with the unidirectional valve method, in comparison to a 34% mean difference reported by Marini et al.14 (n= 20). The design of the present study may have attenuated the differences between methods even with randomization and rest intervals, since individuals underwent several measurements in the same day and this may have caused some fatigue. Marini et al.14 also observed that the method of the occluding valve could be used independently of patient active participation. Their study, however, was not specifically designed to test this possibility. In 1992, Truwit & Marini20 finally validated the method for patients with difficulties to cooperate adequately, as long as they had an occlusion pressure higher than 2 cm H2O at 100ms (P0.1). In the present study the objective was to measure MaxPi without any patient cooperation or special instructions. However, as P0.1 was not assessed, patients with reduced ventilatory drive may have been included. In this case, 20s may have been insufficient for the ventilatory drive to increase significantly in the two methods, and that may have attenuated the differences between them. Results of the study by Truwie & Marini20 reinforce this hypothesis, as authors found MaxIP values even greater than the values found in this study with the unidirectional valve in patients with P0.1 < 2 cmH2O (49.9 ± 24.6 cmH2O versus 43.8 ± 25.1 cmH2O).

Inter-rater reliability for MaxIP measures have been questioned in several studies15-17. In this study, all measurements were performed by the same examiner with the objective of minimizing inconsistencies. The coefficients of variation found for the three measurements of each method were acceptable.
Figure 1. Comparison between the methods and protocols for Maximal inspiratory pressure measurement. * values significantly different from the correspondent method at 20s (p<0.001). ** values significantly different from the Simple Occlusion Method at 40s (p<0.001). # values significantly different from Unidirectional Valve Method at 30s (p<0.001) (post-hoc analysis – Paired t-test).

Figure 2. Correlation between age and maximal inspiratory pressure recorded by the Unidirectional Valve Method with an occlusion time of 40s.

![Graph showing correlation between age and MaxIP](image)

The use of corticosteroids and other drugs that may influence function of respiratory muscles as well as nutritional status of patients and time since mechanical ventilation was initiated were not controlled during the study. Probably because of heterogeneity of the sample in relation to these factors no significant difference was found for MaxIP percent value predicted by the Black and Hyatt equations between patients admitted to the hospital for clinical or surgical reasons. Despite the multifactorial character of the determination of MaxIP, a significant correlation between age and MaxIP was found. Differences between sexes were also found (values were 38.8% lower for females). These results are in accordance with the classical study by Black and Hyatt in which authors described an equation to estimate MaxIP according to age and sex. Despite the differences between absolute values of MaxIP, when the equation was used no significant differences were found for the mean percentage of predicted values between sexes.

As mentioned before, criteria to define weakness of inspiratory muscles and capacity to undergo withdrawal of mechanical ventilation (values between -20 and -30 cmH2O) are arbitrary and do not take age and sex into consideration. Perhaps the utilization of these variables is relevant to establish specific criteria for each patient, and to increase, in this way, the predictive value of MaxIP in the assessment for weaning.
In face of results it is concluded that, when compared to the other protocols, the unidirectional valve method with occlusion during 40 seconds registers greater values of Maxlp for non-cooperative patients.

REFERENCES