Ultrasound Assessment of Ventilator-induced Diaphragmatic Dysfunction in Paediatrics

ABSTRACT

Introduction: Invasive mechanical ventilation contributes to ventilator-induced diaphragmatic dysfunction, delaying extubation and increasing mortality in adults. Despite the possibility of having a higher impact in paediatrics, this dysfunction is not routinely monitored. Diaphragm ultrasound has been proposed as a safe and non-invasive technique for this purpose. The aim of this study was to describe the evolution of diaphragmatic morphology and functional measurements by ultrasound in ventilated children.

Material and Methods: Prospective exploratory study. Children admitted to Paediatric Intensive Care Unit requiring mechanical ventilation > 48 hours were included. The diaphragmatic thickness, excursion and the thickening fraction were assessed by ultrasound.

Results: Seventeen cases were included, with a median age of 42 months. Ten were male, seven had comorbidities and three in seventeen had malnutrition at admission. The median time under mechanical ventilation was seven days. The median of the initial and minimum diaphragmatic thickness was 2.3 mm and 1.9 mm, respectively, with a median decrease in thickness of 13% under pressure-regulated volume control. Diaphragmatic atrophy was observed in 14/17 cases. Differences in the median thickness variation were found between patients with sepsis and without (0.70 vs 0.25 mm; \( p = 0.019 \)). During pressure support ventilation there was a tendency to increase diaphragmatic thickness and excursion. Extubation failure occurred for diaphragmatic thickening fraction ≤ 35%.

Discussion: Under pressure-regulated volume control there was a tendency for a decrease in diaphragmatic thickness. In the pre-extubation stage under pressure support, there was a tendency for it to increase. These results suggest that, by titrating ventilation using physiologil levels of inspiratory effort, we can reduce the diaphragmatic morphological changes associated with ventilation.

Conclusion: The early recognition of diaphragmatic changes may encourage a targeted approach, namely titration of ventilation, in order to reduce ventilator-induced diaphragmatic dysfunction and its clinical repercussions.

Keywords: Child; Diaphragm/ultrasonography; Respiration, Artificial/adverse effects; Ultrasonography

RESUMO

Introdução: A ventilação mecânica invasiva condiciona disfunção diafragmática, atrasando a extubação e aumentando a mortalidade em adultos. Em pediatria, apesar de eventualmente mais relevante, essa disfunção não é sistemicamente avaliada. A ecografia diafragmática tem sido proposta como uma técnica não invasiva e segura para esse fim. O objetivo deste estudo foi descrever a evolução dos índices ecográficos de morfologia e função diafragmáticas em crianças ventiladas.

Material e Métodos: Estudo exploratório, prospetivo. Foram incluídas crianças admitidas num Serviço de Cuidados Intensivos Pediátricos sob ventilação mecânica invasiva > 48 horas e realizadas medições ecográficas de espessura, excursão e fração de espessamento diaphragmáticos.

Resultados: Foram incluídos 17 casos. Mediana de idades: 42 meses. Eram do género masculino 10/17, tinham comorbilidades 7/17 e manifestavam desnutrição na admissão 3/17 casos. Mediana do tempo sob ventilação invasiva: sete dias. Medianas das espessuras diafragmáticas inicial e mínima: 2,3 e 1,9 mm, respectivamente, tendo-se observado uma diminuição mediana da espessura de 13% sob volume controlado regulado por pressão. Observou-se atrofia diafragmática em 14/17 casos. Verificaram-se diferenças na mediana da variação da espessura entre os grupos com e sem sépsis (0,70 vs 0,25 mm; \( p = 0,019 \)). Durante a ventilação em pressão de suporte, observou-se uma tendência para aumento da espessura e excursão diafragmáticas. Ocorreu falência de extubação para fração de espessamento ≤ 35%.

Discussão: Sob volume controlado regulado por pressão verificou-se tendência para diminuição da espessura diafragmática. Sob pressão de suporte, verificou-se uma tendência para o seu aumento. Estes resultados sugerem que, titulando a ventilação, podemos reduzir as alterações morfológicas diafragmáticas associadas à ventilação.

Conclusão: O reconhecimento precoce de alterações diafragmáticas poderá fomentar uma abordagem dirigida, de forma a limitar a disfunção diafragmática induzida pelo ventilador e suas repercussões.

Palavras-chave: Criança; Diafragma/ultrassonografia; Respiração Artificial/efeitos adversos; Ultrassonografia

INTRODUCTION

Invasive mechanical ventilation (IMV) is widely used in intensive care units.\(^1,2\)

Even though it is crucial to the management of critically ill patients, deleterious effects on respiratory muscles have been shown by multiple studies, even with a short-term use.\(^3,4\) These are mainly reflected in myofibrillar and mitochondrial changes, leading to sarcoderm disruption and intracellular lipid accumulation.\(^3,5,6\) There is a decrease in diaphragm thickness, the main respiratory muscle, leading to atrophy and progressive loss in diaphragm contractile function. This is known as ventilator-induced diaphragmatic dysfunction (VIDD).\(^1\)

Other factors apart from IMV seem to add to the diaphragmatic dysfunction found in critically ill patients, namely...
the development of sepsis, the use of corticosteroids, aminoglycoside antibiotics and neuromuscular blocking agents, as well as the patient's nutritional status.2,3,6,7

The definition of VIDD in critically ill adult patients is a relatively recent concept, even though its frequency and relevance has been widely described in different publications.1,2,6 A high incidence of diaphragmatic dysfunction has been found, with an interference on the outcomes, delaying extubation, extending the length of stay, increasing mortality4 and leading to poorer outcomes in the paediatric population, in whom accessory muscles are more fragile and susceptible to fatigue, involving a more difficult approach to diaphragmatic dysfunction.8

However, despite its relevance, diaphragmatic function is not systematically monitored in intensive care units, suggesting that VIDD could be underdiagnosed.2,3

Invasive and troublesome bedside procedures are involved in current gold standard methods for the evaluation of diaphragm function in adults, including the measurement of trans-diaphragmatic pressure with oesophageal and gastric balloon or magnetic phrenic nerve stimulation.2,3 Therefore, diaphragm ultrasound imaging has emerged as a promising technique for the evaluation of diaphragm morphology and contractility.2,9 It is a safe, non-invasive, painless and radiation-free technique which has proved to be cost-effective and easy-to-use in patients receiving IMV.1,2,4,5 Diaphragm thickness, excursion and thickening fraction (DTf) are the major ultrasound parameters,3 allowing for the identification of dysfunction / paralysis, the evaluation of change in diaphragm thickness in ventilated patients and prediction of extubation outcome.2

Even though ultrasound criteria have already been published for the evaluation of diaphragm function in adults,2,3 studies in the paediatric population are still scarce and the prevalence of VIDD in mechanically ventilated paediatric patients is still unknown, in addition to the contribution of other related factors and the impact of dysfunction on clinical outcome.

This study was primarily aimed at describing the progression of ultrasound diaphragmatic morphological and functional indices – thickness, thickening fraction and excursion — in mechanically ventilated paediatric patients admitted to the intensive care unit. The assessment of the impact of ventilation modes on ultrasound indices was the secondary aim of the study.

**MATERIAL AND METHODS**

This was an exploratory observational study with prospective data based on diaphragm ultrasound imaging in paediatric patients admitted to the Paediatric Intensive Care Unit [Serviço de Cuidados Intensivos Pediátricos (CIPE)] of the Hospital Pediátrico de Coimbra (HP).

The study was approved by the ethics committee of the institution, according to the Helsinki declaration. A written informed consent has been obtained from each patient’s legal representative prior to medical examination.

**Study sample**

All paediatric patients admitted to the CIPE between 1 Jun 2017 and 31 Mar 2018 (ten-month study period) and meeting the following criteria were included in the study: 1) patients aged between one month and 18 years; 2) patients in need for IMV for more than 48 hours.

These were the exclusion criteria: 1) patients in need for IMV for at least 12 hours within three months prior to the study; 2) neuromuscular disorder or brain stem injury; 3) congenital thoracic malformation; 4) subphrenic abscesses; 5) thoracic surgery within the six months prior to the study; 6) chest drain at the site of examination; 7) pneumothorax; 8) post liver transplant.

A total of 329 paediatric patients were admitted to the CIPE during the study period (101 were aged under one month, 85 were not receiving IMV and 109 had received IMV for less than 48 hours, in addition to two patients who had been in need for IMV for over 12 hours within the previous three months and two patients who presented with neuromuscular dysfunction and five who were admitted for liver transplant were excluded from the study, such as two patients who presented with neuromuscular dysfunction and five who were recovering from liver transplant were excluded from the study). Eight patients were not assessed due to technical drawbacks and a total of 17 patients were included in the study.

**Patient characteristics and clinical outcomes**

Data were based on the analysis of the patient’s clinical records, by using FileMaker Pro-6® and B-ICU.Care® database of the CIPE of the HP.

The following variables were obtained and analysed: patient’s age (months), gender, admitting diagnosis, comorbidities, body weight (kg), height/length (cm), therapy (sedoanalgesia, neuromuscular blockade, aminoglycoside antibiotics and corticosteroids) and presence of sepsis. Data regarding duration of IMV (hours/days), as well as regarding ventilation parameters and modes, Paediatric Index of Mortality 3 (PIM3)10 and mortality were also obtained.

Daily doses in mg/kg of each drug used during IMV were obtained. All prescribed corticosteroids were converted to equivalent doses of prednisolone, according to the conversion formulae described by Martindale: The Complete Drug Reference.11

Body mass index (BMI) and BMI-for-age percentile (P) were obtained according to the World Health Organization reference curves. The presence of malnutrition was defined at BMI ≤ the third percentile (P3).

Extubation failure was defined as the need for re-intubation within 48 hours and ventilator-free days were calculated and defined as the number of days from successful weaning from mechanical ventilation (invasive or non-invasive) up to day 28 after study inclusion.

**Ultrasound assessment**

Diaphragm ultrasound imaging was carried out with a portable ultrasound Siemens Acuson X300® probe using
two-dimensional (B-mode) and M-mode imaging.

Three measurements of the ultrasound parameters — thickness, excursion and thickening fraction - were obtained in each evaluation— and the average value of the three measurements was considered as the final value, in order to establish its reproducibility. Only measurements made on the right hemi-diaphragm were obtained.12

Ultrasound parameters were assessed considering the different phases of invasive mechanical ventilation. Pressure regulated volume control (PRVC) mode is usually used in paediatric patients, which is a mode of ventilation that combines volume and pressure control ventilation. Pressure support ventilation (PSV) or spontaneous breathing through an endotracheal tube (ETT) are predominantly used in pre-extubation, aimed at maintaining the patient’s inspiratory effort and subsequent inspiratory muscle training.

Considering that the ultrasound evaluation of diaphragm thickness does not depend on maintaining the patient’s inspiratory effort, its evaluation was carried out in every phase of the ventilation process, while the first measurement was obtained within the first 24 hours of IMV and subsequently on a daily basis over a 24 ± 8 hour period, up to the day of extubation or day 28 of intubation, or in case of transfer to another hospital or death.

On the contrary, the preservation of the patient’s inspiratory effort is required for an accurate ultrasound evaluation of diaphragmatic excursion and thickening fraction. Therefore, these were only evaluated on a daily basis in patients in pre-extubation, during PSV or spontaneous breathing through an endotracheal tube.

In the presence of extubation failure, in need for a new ventilation cycle, a daily evaluation of these ultrasound parameters was carried out in pre-extubation, while each cycle was considered as an independent evaluation moment.

All ultrasound imaging evaluations were carried out by the same operator, a paediatric cardiologist, sub-specialist in paediatric intensive care, with adequate training and qualifications.

a) Diaphragm thickness

All evaluations were carried out in B-mode according to a standardised technique6 by using a 10 – 5 MHz transducer linear ultrasound probe. Patients were lying supine in a 30 – 45° position while the probe was positioned perpendicular to the rib surface between the eighth and the tenth intercostal spaces, between midaxillary and right anterior axillary lines, for the acquisition of a bi-dimensional coronal image of the zone of apposition.

The diaphragm was visualised at a 2-3.5 cm depth, showing a hypo-echoic muscular layer amongst two echo-genic layers, one above and one below, corresponding to the diaphragmatic pleura and the parietal peritoneum, respectively (Fig. 1A).

Diaphragm thickness was defined as the vertical distance in millimetres between the midpoint of the diaphragmatic pleura and the midpoint of the parietal peritoneum, measured on the most perpendicular axis to the longitudinal plane.1,6

Airway pressure and flow real-time charts were analysed in each evaluation and three consecutive measurements were obtained during the expiration within the same cycle.

Change in diaphragm thickness from baseline to nadir was obtained for each patient. The presence of diaphragm atrophy was considered in this study at ≥ 10% decrease in thickness from baseline.8

b) Diaphragmatic excursion

The measurement of diaphragmatic excursion was obtained by use of a 6 – 2 MHz convex probe positioned perpendicular to the subcostal area, between midclavicular and right anterior axillary lines.

The diaphragm was visualised in B-mode showing a hyper-echoic line produced by the diaphragmatic pleura adherent to the muscle, while diaphragmatic excursion was assessed in M-mode, corresponding to the vertical distance in millimetres of the diaphragmatic line between maximum inspiration (peak of the curve) and maximum expiration (base of the curve) (Fig. 1B).

Whenever two or more measurements of this parameter were available, the change in diaphragmatic excursion was obtained, corresponding to the difference between peak and baseline values.

c) Diaphragmatic thickening fraction

DTF, corresponding to the change in diaphragm thickness with breathing movements was mathematically obtained as a percentage, considering the values of diaphragm thickness at maximum inspiration and maximum expiration, by using the formula:

\[ \text{DTF} = \left( \frac{\text{diaphragm thickness at end of inspiration} - \text{end of expiration}}{\text{diaphragm thickness at end of expiration}} \right) \times 100. \]

A 10 – 5 MHz linear probe has been used in M-mode, positioned at the same location as for the measurement of diaphragm thickness (Fig. 1C).

Whenever two or more measurements of this parameter were available, the change in DTF was obtained, corresponding to the difference between peak and baseline DTF values.

Statistical analysis

A descriptive analysis was carried out, while the group of patients was characterised through the calculation of measures of central tendency and dispersion for quantitative variables: mean and standard deviation (SD) for variables with normal distribution or median and interquartile range (IQR) for variables without a normal distribution. Shapiro-Wilk test has been used as normality test.

The determination of absolute frequencies for qualitative variables was also carried out and Mann-Whitney’s U-test was used to compare medians of quantitative variables.
A 0.05 level of significance has been considered in this study.

The statistical analysis was carried out by use of SPSS Statistics® version 25 software.

RESULTS

A 42-month median age has been found in our group of 17 patients (10 male) (IQR 9.5 – 155.5).

Principal diagnoses underlying the admission to the CIPE were divided into groups: respiratory (n = 8), traumatic (n = 5), postoperative/postprocedure (n = 2), cardiovascular (n = 1) and infectious diseases (n = 1). Seven patients presented with an underlying condition or comorbidity, namely cancer (n = 3), prematurity with no diagnostic criteria for bronchopulmonary dysplasia (n = 2), cystic fibrosis bronchiectasis (n = 1) and trisomy 21 associated with hypotonia (n = 1).

BMI percentiles were presented into six different ranges (Table 1). Three patients presented at admission with BMI ≤ P_{3} and 5 with BMI ≤ P_{15}.

Fifteen patients were on midazolam (at a daily mean dose of 2.13 ± 1.51 mg/kg), 16 on morphine (at a daily mean dose of 0.21 ± 0.12 mg/kg) and eight patients on fentanyl (at a daily median dose of 0.05 mg/kg) (IQR 0.05 – 0.06).

Rocuronium was administered to 16 patients (at a daily mean dose of 3.34 ± 1.84 mg/kg).

Nine patients were receiving corticosteroid therapy at a daily mean dose of 2.03 ± 0.92 mg/kg of prednisolone and aminoglycoside antibiotics were administered to two patients, while seven patients presented with sepsis during their stay in the hospital.

A seven-day median duration of IMV has been found (IQR 6 – 10.5). Pressure regulated volume control (PRVC) mode was initially used in all the patients, with a 5.9-day median duration (IQR 4.1 – 10.3), while the ventilation mode was not changed in any patient up to pre-extubation.

As regards the ventilation parameters with PRVC, an average maximum respiratory rate of 32 ± 12 breaths/minute, 7 mL/kg median maximum tidal volume (IQR 6.7 – 7.5), 6 cmH\textsubscript{2}O (IQR 5 – 8) median positive end-expiratory pressure (PEEP) and 25 ± 7 cmH\textsubscript{2}O average peak inspiratory pressure (PIP) have been obtained.

Fifteen patients were extubated and pressure support ventilation (PSV) mode was used in pre-extubation in 14 of these patients, with a 23-hour median duration (IQR 7.5 – 45.5). Only one patient remained in spontaneous breathing.

<table>
<thead>
<tr>
<th>Percentile (P)</th>
<th>n (17)</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤ P_{3}</td>
<td>3</td>
</tr>
<tr>
<td>&gt; P_{3} and ≤ P_{15}</td>
<td>2</td>
</tr>
<tr>
<td>&gt; P_{15} and ≤ P_{50}</td>
<td>6</td>
</tr>
<tr>
<td>&gt; P_{50} and ≤ P_{85}</td>
<td>2</td>
</tr>
<tr>
<td>&gt; P_{85} and ≤ P_{97}</td>
<td>2</td>
</tr>
<tr>
<td>&gt; P_{97}</td>
<td>2</td>
</tr>
</tbody>
</table>

n: absolute number; P: percentile
A 16 ± 3.4 cmH₂O average PIP was obtained during PSV.

A 11-day median length of stay at the CIPE (IQR 9 – 17) and a 0.04 mortality rate according to the PIM₃ (IQR 0.02 – 0.15) have been found, while only one of the patients in our group has died in the hospital. A 22-day median ventilator-free time has been found (IQR 20 – 23).

A total of 149 ultrasound evaluations of the diaphragm thickness, 36 of the diaphragmatic excursion and 36 of the diaphragmatic thickening fraction were carried out throughout the study.

A 2.3 mm median baseline diaphragm thickness has been found (IQR 2 – 3.5). A trend towards a decrease in diaphragm thickness was found throughout the time in which the patients were receiving IMV with PRVC mode (Fig. 2).

A 1.9 mm median nadir thickness has been found with PRVC mode (IQR 1.6 – 3.1) and this has occurred at a median four days of IMV (IQR 2 – 7.5). A 0.3 mm median change in diaphragm thickness has been found (IQR 0.2 – 0.7), corresponding to a 13% decrease.

Diaphragmatic atrophy has been found in 14 patients.

The comparison of changes in diaphragm thickness according to patient’s clinical and pathological and pharmacological characteristics is shown in Table 2, showing significant differences regarding median change in thickness between the group with vs. without sepsis (0.70 vs. 0.25 mm; p = 0.019).

As regards diaphragmatic excursion, a 7.8 ± 3.9 mm mean baseline and 10.3 ± 5.4 mm mean peak values have been found, corresponding to 3.0 ± 1.6 mm mean change in diaphragmatic excursion.

Five out of the 14 patients in pre-extubation with PSV remained with this ventilation mode for more than 48 hours. As the ultrasound assessments were carried out daily throughout that phase, two or more assessments of each parameter were obtained in those patients, allowing for the analysis of clinical progression (Fig. 3). A mild trend towards an increasing thickness (Fig. 3A) and diaphragmatic excursion (Fig. 3B) were found as time with PSV mode increased.

A 41.14 ± 11% average baseline value and 48.8 ± 3.7% average peak value of the diaphragmatic thickening fraction (DTf) have been found, with a 19 ± 6.6% average change in DTf.

Extubation failure has been found at values of DTf ≤ 35%, which has occurred within two out of the 17 pre-extubation periods that were evaluated (Fig. 4).

DISCUSSION

This is one the first studies in paediatric patients receiving IMV aimed at the ultrasound assessment of diaphragm morphological and contractile activity.

Between 55 and 60% of the paediatric patients admitted to the CIPE of the Hospital Pediátrico de Coimbra were in need for invasive mechanical ventilation, which is one of the most frequently used techniques in the approach to critically ill paediatric patients.

The complications associated with IMV, namely VIDD, started immediately after the intubation, with a clear interference on the outcomes, as suggested by different studies with adult patients.1,3,6,13 The systematic use of a sensitive and specific diaphragm evaluation is therefore crucial, allowing for the early identification of VIDD.

As the constraints of the remaining modalities of diaphragm assessment were overcome with diaphragm ultrasound imaging, this was considered for the study.1,2,4

 Ultrasound imaging allows for an accurate diaphragm morphological assessment, according to a systematic revision on its relevance in the approach to diaphragmatic dysfunction in critically ill patients,2 allowing for the measurement of diaphragm thickness and the identification of atrophy, one of the major characteristics of VIDD. In addition, functional evaluation is also allowed, with the measurement of the indices of contractility — diaphragmatic excursion.
Table 2 – Comparison of the change in diaphragm thickness regarding the different variables

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Change in Diaphragm Thickness (mm)</th>
<th>p-value (Mann-Whitney’s U-test)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Median (IQR)</td>
<td></td>
</tr>
<tr>
<td>Comorbidities</td>
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<td>0.475</td>
</tr>
<tr>
<td>Yes (n = 7)</td>
<td>0.20 (0.20 – 0.70)</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>0.40 (0.28 – 0.63)</td>
<td></td>
</tr>
<tr>
<td>Malnutrition</td>
<td></td>
<td>0.432</td>
</tr>
<tr>
<td>Yes (n = 3)</td>
<td>0.20</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>0.30 (0.20 – 0.63)</td>
<td></td>
</tr>
<tr>
<td>Fentanyl</td>
<td></td>
<td>0.481</td>
</tr>
<tr>
<td>Yes (n = 8)</td>
<td>0.40 (0.23 - 0.68)</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>0.30 (0.20 - 0.65)</td>
<td></td>
</tr>
<tr>
<td>Corticosteroids</td>
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</tr>
<tr>
<td>Yes (n = 9)</td>
<td>0.30 (0.25 - 0.70)</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>0.25 (0.20 - 0.58)</td>
<td></td>
</tr>
<tr>
<td>Aminoglycoside antibiotics</td>
<td></td>
<td>0.824</td>
</tr>
<tr>
<td>Yes (n = 2)</td>
<td>0.45</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>0.30 (0.20 - 0.60)</td>
<td></td>
</tr>
<tr>
<td>Sepsis</td>
<td></td>
<td>0.019</td>
</tr>
<tr>
<td>Yes (n = 7)</td>
<td>0.70 (0.30 - 0.70)</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>0.25 (0.20 - 0.35)</td>
<td></td>
</tr>
</tbody>
</table>

The change in diaphragm thickness (in mm) is presented as median and interquartile range (IQR). n: absolute number.

Figure 3 – Trend of ultrasound indices measurements with pressure support ventilation (PSV). (A) Change in diaphragm thickness (mm) over time (in hours), with PSV mode. (B) Change in diaphragmatic excursion (mm) over time (hours), with PSV mode.
and thickness fraction. These measurements could be repeated over time, allowing for the comparison of values during follow-up.

Age-related reference values of diaphragm thickness in healthy children have been established by a cross-sectional prospective study. A 2.3 mm median baseline thickness has been found in our study, below the mean reference value for any age group. Despite the differences in measures of central tendency and dispersion that were used in both studies, this finding could be explained by the fact that some patients presented with a chronic disease/comorbidity (7/17) or malnutrition at admission (3/17), which could represent diaphragm fragility and therefore with reduced baseline thickness.

A 13% median decrease in diaphragm thickness has been found in this study throughout the time in which patients were receiving IMV with PRVC mode, with nadir thickness found at a median four days of IMV, while the presence of diaphragmatic atrophy was found in 14 patients. These findings are in line with those described by Glau et al. who had found a -13.8% median change in diaphragm thickness in one of only two studies carried out to date with paediatric patients, with a -3.4% daily atrophy rate. Rapid onset of diaphragmatic atrophy is shown by these findings, with an exponential decline in diaphragm thickness.

An 8.8% decrease in mean diaphragm thickness has also been described by Lee et al. within the first 24 hours upon intubation, with a 0.68% mean gradual daily decline up to day 7 of IMV. However, ventilation parameters that were used in the study do not correspond to lung protective ventilation strategies, except in patients admitted with acute respiratory distress syndrome, which is in contrast with our study, in which a 7 mL/kg median peak tidal volume has been found. Therefore, the discrepancies in these findings could be explained by our small group of patients and by the differences regarding ventilation modes and parameters that were used.

In our study, patients mainly received IMV with PRVC mode up to pre-extubation, in which the patient’s inspiratory effort is detected by the ventilator, assisting the patient in achieving a set tidal volume. Curare replacement in the patient was achieved by administrating low-dose curare. IMV was optimized to make use of PRVC mode, with measurements taken from the patient’s inspiratory effort. This mode involved a combination of IMV with a set pressure level and patient-triggered pressure support. Curare replacement was performed by the ventilator to maintain a constant tidal volume. At the time of pre-extubation, most patients received PSV mode, with settings optimized to maintain a baseline thickness. This mode involved providing additional pressures to the patient’s inspiratory effort to maintain a constant tidal volume. During pre-extubation, PSV mode was used in most patients (14/15). PSV is frequently used in weaning from IMV, aimed at relieving the workload imposed to the respiratory muscles, preserving its spontaneous contraction and preventing atrophy.

A trend towards a decrease in diaphragm thickness has been found during the use of PSV. Instead, a small trend towards an increasing diaphragm thickness has been found with PSV in pre-extubation. These findings suggest that IMV-related diaphragm morphological changes could be reduced by titrating ventilation, using physiological levels of inspiratory effort.

An association between diaphragmatic activity and atrophy in ventilated patients has been suggested by Goligher et al. in a large-scale observational study including the analysis of this parameter in patients receiving different ventilation modes for up to one week. Low diaphragmatic contractile activity was associated with quick reduction in diaphragm thickness, as shown in that study, while a high contractile activity was associated with an increased thickness as contractile activity is known to decrease with increasingly controlled ventilation modes.

Other factors apart from IMV may contribute to VIDD, including sepsis, multiple organ dysfunction and...
malnutrition.\textsuperscript{3,6} The administration of drugs including corticosteroids, aminoglycoside antibiotics and neuromuscular agents is also involved.\textsuperscript{3,5} A significantly higher change in diaphragm thickness has been found in the group of patients with sepsis in our study. These findings are in line with those described in other studies\textsuperscript{8,13} in which sepsis was considered as one of the factors most strongly associated with diaphragmatic dysfunction.

As the assessment of diaphragm thickness regards only one of its morphological characteristics, which is not necessarily correlated to its functional capacity, the ultrasound indices of diaphragmatic contractility — diaphragmatic excursion thickening fraction (DTf) were also evaluated.

DTf was shown by Umbrello et al.\textsuperscript{9} as the most reliable ultrasound indicator of change in the inspiratory muscle effort in response to modifications in the level of ventilation support, in a study aimed at assessing the performance of diaphragmatic excursion and DTf when compared to the traditional indices of inspiratory muscle effort with IMV, as it is only influenced by active muscle contraction.

Additionally, different studies have been carried out to analyse the accuracy of ultrasound indices of diaphragmatic function — excursion and DTf — in predicting the success or failure of extubation, while DTf was considered as the most accurate index.\textsuperscript{2,4,18,19} In our study, extubation failure was found at values of DTf ≤ 35%, in line with a recent systematic revision\textsuperscript{2} describing a range of DTf between 30% – 36% as predictive of extubation failure. Conflicting values regarding extubation failure were found in the study by Lee et al.\textsuperscript{16} in paediatric patients, at values of DTf <17%.

Up to date, there is no established approach to VIDD. The strategies for the use of PSV and earlier extubation need to be analysed and further studies aimed at determining whether IMV could progress from lung-protective to muscle-protective.

Some limitations to our study are worth mentioning: the small group of patients does not allow for any inferences or robust conclusions; all the measurements of diaphragm thickness were carried out a few hours upon intubation and therefore no real baseline diaphragm thickness value was available; all ultrasound assessments were carried out by the same operator while intra and inter-observer reproducibility was not assessed.

Despite these limitations, this study could contribute to a better understanding on the presence of VIDD in paediatric patients and the usefulness of diaphragm ultrasound imaging in the approach to critically ill paediatric patients. It could also represent a model and forerunner for a larger scale study with wider inclusion criteria, namely duration of IMV >24 hours, allowing for stronger conclusions.

CONCLUSION
Diaphragm thickness and diaphragmatic thickening fraction could provide relevant data on the presence of diaphragmatic atrophy and dysfunction while a 13% median decrease in thickness has been found in paediatric patients receiving PRVC ventilation, while nadir thickness was found at a median four days of IMV, with a higher decrease found in patients who developed sepsis. A mild trend towards an increasing diaphragm thickness has been found in pre-extubation in patients receiving PSV ventilation and extubation failure was found at values of DTf ≤ 35%.

These parameters could have been systematically assessed in critically ill paediatric patients in order to titrate ventilation, minimising the presence of diaphragmatic dysfunction and extubation failure.

HUMAN AND ANIMAL PROTECTION
The authors declare that the followed procedures were according to regulations established by the Ethics and Clinical Research Committee and according to the Helsinki Declaration of the World Medical Association.

DATA CONFIDENTIALITY
The authors declare that they have followed the protocols of their work centre on the publication of patient data.

CONFLICTS OF INTEREST
The authors declare that there were no conflicts of interest in writing this manuscript.

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